## Detecting vegetation stress in mixed forest ecosystems through the joint use of tree-water monitoring and land surface modeling

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February 16, 2024

#### Abstract

Recent European heatwaves have significantly impacted forest ecosystems, leading to increased plant water stress. Advances in land surface models aim to improve the representation of vegetation drought responses by incorporating plant hydraulics into the plant functional type (PFT) classification system. However, reliance on PFTs may inadequately capture the diverse plant hydraulic traits (PHTs), potentially biasing transpiration and vegetation water stress representations. The detection of vegetation drought stress is further complicated by the mixing of different tree species and forest patches. This study uses the Community Land Model version 5.0 to simulate an experimental mixed-forest catchment with configurations representing standalone, patched mixed, and fully mixed forests. Biome-generic, PFT-specific, or species-specific PHTs are employed. Results emphasize the crucial role of accurately representing mixed forests in reproducing observed vegetation water stress and transpiration fluxes for both broadleaf and needleleaf tree species. The dominant vegetation fraction is a key determinant, influencing aggregated vegetation response patterns. Segregation level in PHT parameterizations shapes differences between observed and simulated transpiration fluxes. Simulated root water potential emerges as a potential metric for detecting vegetation stress periods. However, the model's plant hydraulic system has limitations in reproducing the long-term effects of extreme weather events on needleleaf tree species. These findings highlight the complexity of modeling mixed forests and underscore the need for improved representation of plant diversity in land surface models to enhance the understanding of vegetation water stress under changing climate conditions.

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## 1 Detecting vegetation stress in mixed forest ecosystems through the joint use of tree-2 water monitoring and land surface modeling

3

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#### 9 Key Points:

- We show that the model's dominant fraction of a mixed ecosystem masks the water status of smaller fractions within a grid cell.
- We demonstrate that refining the plant hydraulic traits based on species presence improves the representation of mixed forests in CLM5.
- We evidence the limitations of CLM5 in reproducing the needleleaf water stress using tree water deficit measurements.
- 16

#### 17 Abstract

Recent European heatwaves have significantly impacted forest ecosystems, leading to increased 18 plant water stress. Advances in land surface models aim to improve the representation of 19 vegetation drought responses by incorporating plant hydraulics into the plant functional type 20 (PFT) classification system. However, reliance on PFTs may inadequately capture the diverse 21 22 plant hydraulic traits (PHTs), potentially biasing transpiration and vegetation water stress representations. The detection of vegetation drought stress is further complicated by the mixing 23 of different tree species and forest patches. This study uses the Community Land Model version 24 5.0 to simulate an experimental mixed-forest catchment with configurations representing 25 standalone, patched mixed, and fully-mixed forests. Biome-generic, PFT-specific, or species-26 specific PHTs are employed. Results emphasize the crucial role of accurately representing mixed 27 forests in reproducing observed vegetation water stress and transpiration fluxes for both 28 29 broadleaf and needleleaf tree species. The dominant vegetation fraction is a key determinant, influencing aggregated vegetation response patterns. Segregation level in PHT parameterizations 30 shapes differences between observed and simulated transpiration fluxes. Simulated root water 31 potential emerges as a potential metric for detecting vegetation stress periods. However, the 32 model's plant hydraulic system has limitations in reproducing the long-term effects of extreme 33 weather events on needleleaf tree species. These findings highlight the complexity of modeling 34 mixed forests and underscore the need for improved representation of plant diversity in land 35 surface models to enhance the understanding of vegetation water stress under changing climate 36

37 conditions.

#### 38 Plain Language Summary

39 Numerical simulation models for large-scale ecosystems often oversimplify mixed forests,

40 neglecting the diversity of species and structural complexity. This oversight impacts the accuracy

41 of simulated plant water use, especially during droughts and heatwaves. This study focused on

42 the specific traits of key tree species at a Luxembourg site, aiming to enhance the model's ability

43 to represent the vegetation response to extreme conditions. By incorporating detailed plant traits,

the model improved in replicating observed tree water use and identifying periods of water

45 deficit. The findings highlight the importance of considering the functional diversity of mixed

46 forest ecosystems for accurate simulations. Moreover, the study introduces a simple metric using

47 the model's structure to pinpoint periods when different species experience severe water deficit.

48 The proposed metric provides a practical tool for identifying critical periods of water stress for

49 various species within mixed forests. This approach enhances our understanding of mixed forest

50 dynamics under extreme conditions, emphasizing the need for nuanced representations in large-

51 scale models.

#### 52 **1 Introduction**

53 The increasing frequency and intensity of droughts and heatwaves in Europe pose challenges to

vegetation (He et al., 2020), affecting various physiological processes (Arend et al., 2021; Hajek

et al., 2022; Obladen et al., 2021; Senf et al., 2020). Dendrochronological studies indicate that

56 coniferous trees (e.g., Spruce or European Larch) are generally more susceptible to drought than

57 broadleaf trees (e.g., Beech or deciduous Oaks) (Vitasse et al., 2019). However, recent severe

events, like the 2018 drought in central Europe, have endangered even the typically resilient Oak

<sup>59</sup> and Beech trees, with projections suggesting worsening conditions in the future (Schuldt et al.,

60 2020; Shekhar et al., 2024). These extreme events disrupt the plant water transport system

61 (Dietrich et al., 2019; Kannenberg, Novick, & Phillips, 2019; Tomasella et al., 2019), leading to

62 constraints on carbon assimilation, delayed phenological development, and reduced plant

63 growth. The negative effects of drought have lasting implications for a forest's long-term

resilience to environmental changes. (Kannenberg et al., 2020; Van Passel et al., 2022). The

combination of the duration of these impacts and the extent of disruption in physiological
 processes is termed as legacy effects. Recognizing and understanding these enduring

processes is termed as legacy effects. Recognizing and understanding these enduring
 consequences are crucial for assessing the overall health and adaptability of forest ecosystems in

the face of evolving environmental conditions. (Kannenberg, Novick, Alexander, et al., 2019;

69 Yu et al., 2022).

The responses of forest to changing environmental conditions is heavily influenced by 70 tree competition (e.g., Fernández-de-Uña et al., 2015; Ford et al., 2017), where tree demography 71 (e.g., size, age) and diversity (e.g., species) are the key features determining the individual tree 72 interactions (Felton et al., 2010; Jactel et al., 2017; Pretzsch et al., 2013). Mixed forests, 73 featuring multi-generational tree populations in the same area (del Río et al., 2018), exhibit 74 structural complexity that enhances tree survival during prolonged droughts or heatwaves (Ma et 75 76 al., 2023). The dynamic equilibrium in these ecosystems involves a balance between niche separation and resource competition among different tree species, with finite resources such as 77 water, light, space, and nutrients influencing the vegetation response (Bravo-Oviedo et al., 2014). 78 Resource competition among tree species in mixed forests is a dynamic process influenced by 79 latitude, forest successional stage, and environmental conditions (McDowell et al., 2020). The 80 intra-specific competition (Bosela et al., 2015) is amplified when environmental extremes, such 81 as heatwaves and droughts occur, resulting in more frequent and massive die-off in the forests 82 (Hammond et al., 2022). Furthermore, when exploring the effects of species mixing, site-83 dependent conditions like geology, species spatial distribution or forest successional stage must 84 be taken into account (Gong et al., 2020; Han et al., 2022; Mina et al., 2018) to isolate the 85

<sup>86</sup> response related to species interactions from the effect of environmental factors.

The forest response to severe weather conditions, like heatwaves, has been monitored 87 using remote sensing products that analyze long-term changes in land surface temperature, 88 89 vegetation health index, or normalized difference vegetation index (e.g., Bento et al., 2020; Prăvălie et al., 2022). Although these products enable the simultaneous monitoring of large 90 areas, their spatial resolution is inadequate to investigate physiological processes at the tree or 91 organ level, such as stem dehydration. (e.g., stem dehydration). In this regard, in-situ 92 measurements carried out with stem psychrometers, sap flow sensors, and tree dendrometers 93 (e.g., Brinkmann et al., 2016; Guan et al., 2022; Mohanty et al., 2017; Zweifel et al., 2005) 94 95 provide an inside view of multiple physiological aspects of the trees. Stem psychrometers measure water potential (Dixon & Tyree, 1984), revealing how plants respond to specific 96 environmental stressors like droughts (Steppe, 2018). Sap flow sensors gauge sap velocity that 97 98 can be upscaled to tree and stand transpiration ( $E_{\rm T}$ ) estimates (Poyatos et al., 2021). Tree dendrometers measure changes in tree diameter, allowing for the calculation of growth and the 99 tree water deficit (TWD) (Zweifel et al., 2014). Therefore, the continuous monitoring of 100 individual trees generates a detailed data set describing how the diverse forest tree population 101 respond to harsh meteorological conditions that cannot be retrieved with remote sensing 102 103 products.

Latest versions of land surface models (LSMs) have implemented a plant hydraulic 104 system (PHS) to reproduce the water transport of the vegetation (e.g., Bonan et al., 2014; Eller et 105 al., 2020; L. Li et al., 2021a), reduce the bias of soil-moisture based formulations of plant water 106 stress (Sloan et al., 2021) and improve the simulated transpiration rates (e.g., Kennedy et al., 107 2019; L. Li et al., 2021a). The PHS uses the plant vulnerability curve (PVC) to characterize the 108 hydraulic stress in the vegetation by describing plant hydraulic conductivity as a function of the 109 water potential in a given plant organ (i.e., percent loss of hydraulic conductivity, 110 PLC) (Venturas et al., 2017). The implementation of the PHS in LSMs allows to mimic the 111 interspecific variability of plant transpiration response across forest stands and sites (e.g., 112 (Jiménez-Rodríguez et al., 2024; Kennedy et al., 2019; Wu et al., 2020). Furthermore, the 113 explicit simulation of the water potential in different plant segments (i.e., roots, stem, and leaves) 114 provides the opportunity to compare the simulated plant water stress with novel measurements 115 such as the TWD by exploiting experimental evidence on the link between tree diameters and 116 water status (Dietrich et al., 2018; Schäfer et al., 2019). 117 The implementation of the PHS approach in LSMs relies on the homogenization of plant 118 hydraulic traits (PHTs) for each plant functional type (PFT). The PFT classification uses a three-119

level system based on latitudinal location, phenology, and plant type (Colin Prentice et al., 120 1993) assuming homogeneous conditions for some biophysical variables of the forest (e.g., 121 aerodynamic conductance). However, the use of the PFT classification system is currently under 122 scrutiny due to its limitations in representing the functional diversity of forest ecosystems 123 (Cranko Page et al., 2024). Therefore, the PHT parameterization of individual PFTs is a daunting 124 task because the PHTs vary between individual trees and even more so between species of the 125 same PFT (Domec et al., 2012; Hochberg et al., 2018; Rosner et al., 2019). This highly diverse 126 nature of PHTs has become a major obstacle for PFT classification, highlighting the need to 127 adopt alternative approaches (e.g., habitat filtering) where PHTs of different vegetation types are 128 restricted according to trait-environment relationships (Chatanga & Sieben, 2020; Fry et al., 129 2019; Verheijen et al., 2015; Violle et al., 2014). In this regard, the use of the PFT classification 130 to evaluate the PHS may be less problematic for mono-specific stands and more challenging for 131 more complex ecosystems such as mixed forests that require a more detailed representation of 132 their PHTs (Bohrer & Missik, 2022). For instance, in boreal mixed forests composed of Spruce 133 and Birch species or in the Mesophytic deciduous forests dominated by Pedunculate Oak and 134 Hornbeam (Barbati et al., 2014), there are examples in which the PHTs (e.g.,  $\Psi_{p50}$ ) of the main 135 tree species differ considerably (Laughlin et al., 2023). This issue is further complicated in 136 highly segregated landscapes where multiple PFTs, located within the same model grid cell, 137 interact through the shared resources (e.g., soil water). While previous studies have tackled the 138 challenge of representing sub-grid heterogeneity to understand the impacts of topographical 139 gradients (Tesfa & Leung, 2017), hydrological processes (Torres-Rojas et al., 2022), or 140 vegetation types (Ke et al., 2013) on the estimation of energy and water fluxes, no study has yet 141 evaluated the impact of sub-grid heterogeneity on the simulated plant water stress in LSMs. In 142 particular, the effect of homogenizing PHTs within single PFTs and/or across the landscape on 143 144 the simulated plant water stress is still poorly understood.

This manuscript aims to assess the impact of standardizing the PHTs within the PFT
classification system for diverse sub-grid representations of the landscape. The study investigates
the transpiration and plant water status simulated by the advanced Land Surface Model, CLM5.0.
It utilizes species- and region-specific PHTs, examining standalone, mixed patched, and fullymixed forest stands. The underlying hypothesis suggests that a more detailed characterization of

150 PHTs in mixed forest ecosystems will better represent observed water fluxes and stress

- 151 compared to the generic PFT generic parameterization. The study conducted a series of point-
- scale numerical experiments employing generic, PFT-specific, and species-specific plant
- 153 hydraulic parameterizations. It considered both shared and separated access of forest stands to
- soil water resources. The research was conducted in the Weierbach catchment (Luxembourg),
  which has been extensively monitored in the last two decades (Hissler et al., 2021). The mixed
- forest cover of the Weierbach catchment allows for characterizing PHTs of two PFTs (i.e.,
- 157 broadleaf deciduous temperate trees (BDT) and needleleaf evergreen temperate trees (NET) and
- 158 four tree species (i.e., Oak, Beech, Spruce, and Douglas fir). These species, representative of
- BDT and NET PFTs in Europe, have been monitored using sap flow and dendrometer sensors,
- 160 providing information on the fluxes and plant water status under various weather conditions,
- 161 including extreme weather events.

## 162 2 Materials and Methods

Forests in Europe cover 35% of the land area (FOREST EUROPE, 2020), and most of 163 these stands are dominated by broadleaf deciduous (BDF) and needleleaf evergreen (NEF) 164 forests (Poulter et al., 2015). The BDFs are represented mainly by Beech (Fagus sylvatica L.), 165 Birch (Betula sp.), Hornbeam (Carpinus betulus L.), Maple (Acer sp.), and Oak (Quercus sp.) 166 trees (de Rigo et al., 2016; Nascimbene et al., 2013). In Europe, the NEFs are mainly represented 167 by Pine (Pinus sp.) and Spruce (Picea sp.) trees (Buras & Menzel, 2019; de Rigo et al., 2016) 168 with commercial plantations of the non-native Douglas fir (Pseudotsuga menziesii (Mirbel) 169 Franco) (Thomas et al., 2022). The Weierbach catchment in Luxembourg is covered by BDF 170 and NEF forests, spatially distributed as mixed and standalone patches, respectively. The 171 availability of continuous measurements of sap flow and tree diameters between 2019 and 2021 172 makes this case study suitable for evaluating the impact of the recent climate extremes on 173 Europe's most representative forest types. 174

### 175 2.1. Study Site

The Weierbach catchment is located in the western part of Luxembourg (N: 49° 49' 38", 176 E: 5° 47' 44"), covering an area of 0.45 km<sup>2</sup>, and an elevation range from 450 m a.s.l. to 500 m 177 a.s.l. The catchment contributes to the Alzette river basin with a mean annual stream discharge of 178 478 mm yr<sup>-1</sup> (Hissler et al., 2021). This catchment features a "V" shape topography and is 179 180 located on top of a schist's bedrock (Pfister et al., 2017), with a plateau feature on the upper catchment characterized by deep periglacial deposits and the hillslope with shallow soils 181 (Martínez-Carreras et al., 2016). Soil characteristics in the plateau allow a larger storage capacity 182 than in the shallow soils on the hillslope. The first 30 cm of soil have a silty clay texture, 183 changing towards a loam texture at deeper layers (Glaser et al., 2016). The catchment has a semi-184 marine climate (Hissler et al., 2021) with a mean annual precipitation of 953 mm yr<sup>-1</sup> and a mean 185 annual temperature of 8.9 °C. The forested catchment is covered by a mixed deciduous forest of 186 Beech (Fagus sylvatica L.) and Oak trees (mostly hybrids of Quercus petraea (Matt.) Liebl. and 187 Quercus robur L.) covering 77.6% of the catchment, with the remaining 22.4% covered by small 188 patches of coniferous stands of Spruce (Picea abies (L.) H. Karst.) and Douglas fir (Pseudotsuga 189 menziesii (Mirbel) Franco) (Figure 1). 190





194 Figure 1. Map of the forested land cover at the Weierbach catchment obtained from the aerial

195 LIDAR survey (ACT, 2020) describing the canopy height variation. The map also shows the

spatial distribution of the environmental sensors (i.e., soil moisture, sap flow, and dendrometers)

installed in the catchment. The schema on the left side of the figure represents the

implementation of the catchment in the CLM5.0 modeling framework.

199 2.2 Transpiration estimates

Sap velocities (cm hr<sup>-1</sup>) were measured in 13 Oak, 14 Beech, six Spruce, and three 200 Douglas fir trees during the 2019 and 2020 growing season. Technical details of sensor 201 202 deployment and calculation of sap velocities of the trees monitored at Weierbach can be found in 203 Schoppach et al. (2021) and Fabiani et al., (2022). Sap flux per tree per cross sectional area (Q,  $cm^3 hr^{-1}$ ) was quantified by multiplying sap flow velocities by the sapwood area ( $cm^2$ ) of each 204 tree equipped with a sap flow sensor. The daily sap flux was calculated per tree  $(m^3 d^{-1})$  and 205 averaged per species ( $\overline{Q}$ ). The transpiration estimates ( $E_T$ ) were calculated in mm d<sup>-1</sup> based on 206 Equation 1, where  $\Lambda_{g.sp}$  is the basal area of the measured trees (m<sup>2</sup>),  $\Lambda_{G.sp}$  is the total basal area 207  $(m^2)$  of the species,  $\Lambda_r$  is the reference surface area (i.e., whole catchment or individual forest 208 209 stand), and 1000 is the conversion factor from m to mm. The diameter class distribution and basal area was assessed with two surveys carried out with temporal plots in both forest types, the 210

- 211 needleleaf evergreen and broadleaf deciduous stands (see Table S1).
- 212

$$E_T = \sum_{sp=1}^{n} 1000 \cdot \left(\frac{\overline{Q}}{\Lambda_{g.sp}}\right) \cdot \frac{\Lambda_{G.sp}}{\Lambda_r}$$
 Equation 1

#### 213 2.3 Tree Water Deficit

The tree water deficit (TWD) was calculated based on tree diameter increment 214 215 measurements on individual trees, carried out using high-resolution band dendrometers (model: DBL60 ICT) between 2019 and 2022. A total of 22 sensors were installed across the Weierbach 216 catchment in nine Beech trees, nine Oak trees, three Spruce trees, and one Douglas fir tree. We 217 218 selected the most representative trees from large (50–75 cm) and small (25–50 cm) diameter classes for Beech and Oak. Spruce and Douglas fir stands are evenly aged because they are a 219 forest plantation. For this reason, we equipped trees from one diameter class only, ranging from 220 20 cm to 35 cm. The data was processed with the treenetproc R package (Haeni et al., 2020; 221 222 Knüsel et al., 2021; Treenetproc, 2019/2023; Wickham et al., 2019), which allows the automated detection and removal of outliers. For each tree, we calculated the TWD, which is the difference 223 between the theoretical radius at full hydration and the current radius (Zweifel et al., 2016). The 224 225 tree water deficit of each forest type cover was determined using a weighted average of the species based on the area covered by each of them (see Table 2). 226

#### 227 2.4 Model Setup

The Community Land Model version 5.0 (CLM5, Lawrence et al., (2019)) was implemented for the Weierbach catchment using a point-scale setup. Multi-year spin-up runs were performed until the soil moisture and temperature reached a dynamic equilibrium. Each spin-up covered a 11-year time window using the atmospheric forcing and land surface parameterization from 2011 to 2021. A production model run of the 11-year time window was carried-out after the spin-up period for each experiment; see section 2.5 and section 2.6 for additional details.

### 235 2.4.1 Atmospheric Forcing

Hourly atmospheric forcing was retrieved from the Roodt meteorological station (N: 49° 236 48' 21.91", E: 5° 49'52.49"), which is located 3.5 km south-east of the catchment. This data set 237 includes precipitation (P) in mm hr<sup>-1</sup>, wind speed (u) in m s<sup>-1</sup>, air temperature (T) in  $^{\circ}$ C, relative 238 humidity (RH) in %, and incoming shortwave radiation ( $R_s$ ) in W m<sup>-2</sup>. To fill occasional gaps in 239 T and P, we used the Aarsdorf (N: 49° 51' 36.49", E: 5° 50'32.9") and Useldange stations (N: 240 49° 46' 2.74", E: 5° 58'2.17"). Gaps in atmospheric pressure (p) and downwelling longwave 241 radiation  $(R_{\rm I})$  were filled using the ERA5 reanalysis product (Muñoz Sabater, 2019, 2021), 242 downloaded from the Copernicus Climate Change Service (C3S) Climate Data Store. The 243 atmospheric CO<sub>2</sub> concentration was kept constant at 411.6 ppm. This value is the average of 244

daily measurements at Mauna Loa (NOAA-ESRL, 2022) between 2018 and 2022.

246 2.4.2 Soil and Bedrock Parameterization

The lower boundary condition describes the geopedological characteristics of the catchment, where soil texture and depth to bedrock are documented by Glaser et al. (2016). The clay content decreases with depth, from 45.2% (at 12 cm deep) to 22.27% below 32 cm deep. Meanwhile, the sand percentages increase considerably with depth, from 10% at the surface up to 31.41% below 32 cm (Table 1). The unsaturated zone at this site reaches down to 3.0 m where the soil Cv soil horizon is located. However, the default depth to bedrock used by CLM5 based

- on Pelletier et al. (2016) is 95.4 cm at the Weierbach catchment making necessary to increase the
- unsaturated zone. Consequently, we modified the depth to bedrock by adding 150 cm and setting
- the clay and sand percentages of the newly added soil layers to 10.0% and 90.0%, respectively,
- following the approach proposed by Jiménez-Rodríguez et al. (2022a).
- Table 1. Vertical distribution of the soil texture characterization used as the lower boundary condition for the numerical experiments carried out at the Weierbach catchment.

Depth [cm]	Description	Clay	Sand	Source
		Fraction	Fraction	
		[%]	[%]	
0 - 12	Soil Horizon Ah	45.20	10.04	(Glaser et al., 2016)
12 - 32	Soil Horizon Ah-B	37.70	16.04	(Glaser et al., 2016)
32 - 92	Soil Horizon B1/B2	22.27	31.41	(Glaser et al., 2016)
92 - 245	Weathered Bedrock	10.00	90.00	(Jiménez-Rodríguez et al. 2022a)

#### 259 2.4.3 Land Surface Parameterization

The broadleaf deciduous (BDF) and needleleaf evergreen (NEF) forests were associated 260 with the temperate plant functional type (PFT) broadleaf deciduous tree (BDT) and needleleaf 261 evergreen tree (NET), respectively. The monthly leaf area index (LAI,  $m^2m^{-2}$ ) of the BDT was 262 determined based on MODIS (ORNL DAAC, 2018). The retrieved LAI is based on the mean 263 value for the broadleaf deciduous classification of the 1.0 km area around the catchment 264 boundaries, done using the MODISTools package (Hufkens, 2022; Tuck et al., 2014). The LAI 265 of the BDT was set to  $0 \text{ m}^2\text{m}^{-2}$  between November and March to represent the dormant season. 266 Default LAI values of CLM5 were used for the NET forest stand because MODIS does not 267 correctly depict the signal due to the small sizes of needleleaf evergreen patches. To assess the 268 proportion of Beech and Oak canopy to the total LAI, we employed the allometric equations 269 from Pretzsch (2019) along with the data from the forest survey carried out in 2022 (Fabiani et 270 al., 2022). We found that Oak and Beech contribute to 19.6% and 80.4%, respectively, of the 271

total LAI of BDT (Text S1 and Figure S1).

273 2.4.4 Plant Hydraulic Parameterization

The plant hydraulic system (PHS) implemented in CLM5 aims to mimic the plant water 274 transport in the model using Equation 2 to calculate the hydraulic conductance  $k (mm_{H2O} mm_{H2O})$ 275 276  $1^{1}$ s<sup>-1</sup>) of each plant segment (i.e., roots, stem, and sunny and shaded leaves). Equation 2 represents the plant vulnerability curve (PVC) and uses the xylem pressure inducing 50% loss of hydraulic 277 conductance ( $\Psi_{p50}$ , MPa), the non-dimensional sigmoidal shape parameter of the curve ( $c_k$ ), the 278 water potential of the lower segment ( $\Psi$ ), and the maximum plant hydraulic conductance ( $k_{\text{max}}$ , 279  $mm_{H2O} mm_{H2O} - s^{-1}s^{-1}$ ). Each of the plant hydraulic parameters (i.e.,  $\Psi_{p50}$ ,  $c_k$ ,  $k_{max}$ ) can be adjusted 280 per plant segment and PFT and are considered static over time. In this study, we considered that 281 the plant segments (i.e., roots, stems, leaves) within each PFT (and tree species classification) 282 have the same plant hydraulic parameterization (i.e.,  $k_{\text{max}}$ ,  $\Psi_{p50}$ , and  $c_k$ ). 283

$k = k_{max} 2^{-\left(\frac{\Psi}{\Psi_{p50}}\right)^{c_k}}$	Equation 2
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285 CLM5 has a generic model parameterization (GMP) describing the main PFTs (Table 2).

286 In this study, we created two additional parameterizations for the Weierbach forest stands. The

PFT adjusted parameterization (PAP) describes the BDT and NET forest stands lumping the
 species present in each PFT fraction. The species-specific parameterization (SSP) characterizes

- the PHS of the individual tree species present in each forest fraction and do not lump the species
- per PFT. To determine the PHT parameters (i.e.,  $\Psi_{p50}$  and  $c_k$ ) of each additional parameterization
- 291 (i.e., PAP and SSP), we sampled the  $\Psi_{p12}$ ,  $\Psi_{p50}$ , and  $\Psi_{p88}$  values from the Xylem Functional
- 292 Traits (XFT) database (Choat et al., 2012) for each tree species investigated. To improve the
- 293 representation of Beech we did an additional search for PHT data, which resulted in an improved
- data set with 23 records (see Table S2). We followed the procedure described by Jiménez-
- Rodríguez et al. (2024) to determine the  $c_k$  used in the plant vulnerability curve of CLM5 for the
- <sup>296</sup> mean  $\Psi_{p50}$  values of PAP and SSP (Table 1). Figures S2 and S3 show the differences in the
- 297 PVCs of the different forest stand classification (i.e., BDT, NET, Oak, Beech, Spruce, Douglas fir) per peremeterization (i.e., CMP, PAP, SSP)
- fir) per parameterization (i.e., GMP, PAP, SSP).

Table 2. Plant hydraulic traits used for each model parameterization and forest stand. The

300 percentage of land cover is estimated with respect to the catchment area. The area percentage

301 value between brackets in Oak and Beech expresses the proportion of the BDT area covered by

302 each species.

Parameterization	Forest Stand	Area [%]	Ψ <sub>p50</sub> [MPa]	Ck
Generic Model Parameterization	BDT	77.6	-2.65	3.95
[GMP]	NET	22.4	-5.20	3.95
PFT-Adjusted Parameterization	BDT	77.6	-3.24	2.93
[PAP]	NET	22.4	-3.28	5.82
	Oak	15.2 [19.6]	-3.49	3.69
Species-Specific Parameterization	Beech	62.4 [80.4]	-3.15	3.12
[SSP]	Spruce	10.2	-3.5	3.69
	Douglas fir	12.2	-3.25	3.12

#### 303 2.5 Numerical Experiments

Several numerical experiments were carried out by implementing standalone and mixed 304 forests under different PHT parameterizations (Figure 2). In these experiments, the standalone 305 forest parameterization is represented by an individual forest stand composed of only one PFT or 306 species that covers 100% of the area and does not compete for water resources with other PFTs 307 or tree species. In the mixed patched configuration, the forest is characterized by two or more 308 forest types sharing the same lower boundary conditions (i.e., soil water resources). The fully-309 mixed configuration corresponds only to the broadleaf deciduous forest where Beech and Oak 310 trees are randomly distributed. Different PHT parametrizations according to the forest stand 311 classification (i.e., BDT, NET, Oak, Beech, Spruce, and Douglas fir) are used within each set of 312 experiments. The basal area proportions of Oak and Beech were used to partition the observed 313 314 leaf area index (LAI) at the site into species-specific LAI values in the model (see section 2.4.3).

Experiment 1 evaluates the difference in the forest response between standalone forest patches using GMP, PAP, and SSP PHT parameterization. This set of numerical experiments assesses the impact of using a different level of detail in the PHT parameterization; from biome-

generic PFT classification to species-specific hydraulic trait characterization. In this experiment 318 319 the observed LAI of the site is assumed to represent only the proportion assigned to the one simulated PFT or species. Experiment 2 repeats the same assessment as Experiment 1, assuming 320 a mixed patched forest where all patches share a common soil water store. In this second set of 321 experiments the GMP and PAP parameterizations use the basal area proportion of NET and BDT 322 in the catchment, which corresponds to 22.4% and 77.6%, respectively (Table 2). In the mixed 323 patched forest configuration using the SSP parametrization, the area is subdivided according to 324 the following percentages: Spruce (10.2%), Douglas fir (12.2%), Oak (15.2%), and Beech 325 (62.4%). For Spruce and Douglas fir, these percentages were obtained from aerial photographs 326 because these stands are established as mono-species plantations. The percentage of Oak and 327 Beech was determined according to the proportion of the basal area contribution of each species 328 (see Text S1), because these species grow in a mixed stand and their cover fractions cannot be 329 easily distinguished in aerial photographs. Finally, Experiment 3 evaluates the effect of a fully-330 mixed forest stand that is represented only by the BDT in the catchment. This experiment 331 considers the GMP and PAP PHT parameterizations of the BDT and the SSP of Oak and Beech. 332 This forest stand comprises Beech (80.4%) and Oak (19.6%) trees, allowing us to compare the 333

representation of a fully-mixed condition.

342

For each set of numerical experiments, we examined the impact of the different model parameterizations on the simulated transpiration and tree water stress as a consequence of plant

337 hydraulic traits and competition for available water resources. This evaluation is carried out by

338 separating normal and extremely hot years. We hypothesize that a higher level of PHTs

- refinement for the selected PFTs and tree species leads to a better agreement between simulated
- 340 and measured signals of vegetation states (i.e., water stress) and fluxes (i.e., transpiration) than 341 the generic PFT parameterization.



- Figure 2. Schema of the three sets of numerical experiments carried out at the Weierbach
- 344 catchment. A solid line frame represents the total ground area and a shared water resource, while

the dotted line frames represent the reference ground area for a given forest stand. Columns

346 represent different experiments, rows different PHT parameterizations. GMP: generic model

347 parameterization; PAP: PFT-adjusted parameterization; SSP: species-specific parameterization;

BDF: broadleaf deciduous forest; NEF: needle-leaf evergreen forest. The tree crown images were

- retrieved from pch.vector (2023).
- 350 2.6 Data analysis

351 The data analysis was carried out for the extended summer period (i.e., May to September) due to the deciduous nature of the larger proportion of forest in the catchment (i.e., 352 353 BDT). The analyses involving the measured tree transpiration ( $E_{\text{T-mea}}$ ) are based on the years 2019 and 2020, while the evaluation of tree water deficit (TWD) is based on the years 2020 and 354 2021, because this is the only period in which all the species were simultaneously monitored. 355 The classification of years into normal and hot is based on the total number of days classified as 356 a heatwave for Luxembourg according to the European Drought Observatory (EDO, 2023). This 357 classification uses the definition given by Lavaysse et al., (2018) who defined a heatwave as the 358 359 period of at least three consecutive days with daily minimum and maximum air temperatures above the 90<sup>th</sup> percentile of the baseline period. We used the same baseline period implemented 360 by the EDO to determine the threshold between normal and hot years in Luxembourg (1980-361 2010). This threshold corresponds to the average plus one standard deviation (5.8 + 3.3 days), 362 leaving a value of 10 days per year. Consequently, the years 2019 and 2020 were classified as 363 364 hot years and 2021 as a normal year.

365 2.6.1 Principal Component Analysis

A principal component analysis (PCA) was carried out to evaluate the dependency of  $E_{T}$ . 366 mea and TWD on environmental factors. The environmental factors evaluated were air 367 temperature (T), relative humidity (RH), incoming short-wave radiation ( $R_s$ ), wind speed (u), and 368 soil water content ( $\theta$ ). The meteorological variables used for the PCAs were extracted from the 369 model atmospheric forcing (see section 2.4.1). The  $\theta$  was measured at seven monitoring clusters 370 established in the catchment (Hissler et al., 2021), obtaining the daily values by averaging soil 371 moisture across depths (10 cm, 20 cm, 40 cm, and 60 cm) and monitoring clusters (see Figure 372 S4). 373

374 2.6.2 Index of Agreement

375 Measured ( $E_{\text{T-mea}}$ ) and modeled ( $E_{\text{T-mod}}$ ) transpiration estimates were aggregated at the daily scale because sap flow does not translate directly to transpiration rates due to tree 376 capacitance. These estimates were compared against each other using the index of agreement 377 378  $(I_{oa})$  proposed by Duveiller et al., (2016). The  $I_{oa}$  (Equation 3) is the product of an  $\alpha$  coefficient and the Pearson correlation coefficient (r). The  $\alpha$  coefficient quantifies the bias existing between 379 measured and modeled daily transpiration rates, where a value of 1.0 represents a perfect 380 agreement between both data sets and a 0 value means no agreement between them. This 381 coefficient is determined using the standard deviation of measured and modeled transpiration ( $\sigma_X$ 382 and  $\sigma_{Y}$ , respectively) and their mean values ( $\overline{X}$  and  $\overline{Y}$ ). For the mixed forest conditions (i.e., 383 Experiments 2 and 3) we discretize the  $I_{oa}$  between the different forest fractions ( $I_{oa-F}$ ) to be 384 compared with the I<sub>oa</sub> at grid cell level (I<sub>oa-G</sub>). 385

$$I_{oa} = \alpha r \text{ where } \alpha = f(x) = \begin{cases} 0, & \text{ if } r \le 0\\ \frac{2}{\frac{\sigma_X}{\sigma_Y} + \frac{\sigma_Y}{\sigma_X} + \frac{(\overline{X} - \overline{Y})^2}{\sigma_X \sigma_Y}}, \text{ otherwise} \end{cases}$$
Equation 3

387 2.6.3 Plant Water Stress Analysis

The model results include the estimations of water potential in roots ( $\Psi_{root}$ ), stem ( $\Psi_{stem}$ ) 388 and leaf ( $\Psi_{\text{leaf}}$ ). However, we selected only  $\Psi_{\text{root}}$  and  $\Psi_{\text{leaf}}$  to compare with the TWD considering 389 the lack of differences between  $\Psi_{stem}$  and  $\Psi_{leaf}$  in the model results and the vulnerability 390 segmentation hypothesis (Tyree & Ewers, 1991) which states that the distal portions of the plant 391 (i.e., roots and leaf) are the most vulnerable sections of the plant. The tree water deficit (TWD) 392 was correlated with the selected water potentials (i.e.,  $(\Psi_{root} \text{ and } \Psi_{leaf})$  during the hot year and the 393 normal year (i.e., 2020 and 2021, respectively). The  $\Psi_{root}$  corresponds to the integral of water 394 potentials of all root segments along the soil profile. The  $\Psi_{\text{leaf}}$  is the weighted average of shaded 395 and sunlit leaves according to their LAI values. The simulated matric potential data (i.e.,  $\Psi_{root}$ ) 396 and  $\Psi_{\text{leaf}}$ ) and canopy stress factor ( $\beta$ ) were averaged only for daytime conditions (i.e., from 397 06:00 to 18:00). The model uses Equation 4 to estimate the leaf water stress factor ( $\beta_x$ ) as the 398 ratio of the actual stomatal conductance  $(g_{x,s})$  over the unstressed stomatal conductance  $(g_{x,max})$ 399 of each component of the canopy (i.e., sunlit and shaded leaf). Then, Equation 5 determines  $\beta$  as 400 401 the weighted average of the canopy component and its corresponding LAI. The  $\beta$  ranges from 0 when the canopy is fully stressed to 1 when the canopy is unstressed. More details on the 402 mathematical formulation of  $\beta$  in CLM5 are provided by Kennedy et al. (2019). 403

 $\beta_x = \frac{g_{x.s}}{g_{x.max}}$  Equation 4

$\beta_{\beta} = \beta_{sunlit} \cdot LAI_{sunlit} + \beta_{shaded} \cdot LAI_{shaded}$	Equation 5
$p = \frac{1}{LAI_{sunlit} + LAI_{shaded}}$	Equation 5

406 2.6.4 Identification of stress periods based on the plant hydraulic system

We performed an additional assessment to test the feasibility of using the plant hydraulic 407 system (PHS) of CLM5 to detect the historical stress periods linked to hot years according to the 408 EDO classification at Weierbach. Here we referred to the mixed patched forest configuration 409 used in Experiment 2 and adopted the plant specific hydraulic parameterization (see Section 2.5). 410 We used the simulations of the 11-year time window (2011-2021), which contains four years 411 classified as hot (i.e., 2015, 2018, 2019, and 2020) and the remaining seven years as normal 412 years (see Section 2.6). Different physiological metrics extracted from the PHS of CLM5 namely 413  $\beta$ ,  $\Psi_{root}$  and  $\Psi_{leaf}$ , were assessed as suitable indicators for detecting stressed forest ecosystems. 414

#### 415 **3 Results**

416 3.1 Forest response to environmental conditions

In the Weierbach catchment, two key tree physiological variables, forest transpiration 417 418  $(E_{\rm T})$  and tree water deficit (TWD), were measured. Figure 3.A and Figure 3.B illustrate their primary differences on an annual basis, facilitating a comparison of transpiration responses and 419 tree water status at different levels such as catchment, plant functional type (i.e., NET, BDT), 420 421 and species (i.e., Spruce, Douglas fir, Oak, Beech). Figure 3.C and Figure 3.D present the outcomes of the principal component analysis (PCA), a crucial tool for identifying the main 422 environmental variables influencing the variations in both  $E_{\rm T}$  and TWD. The PCA analysis is 423 pivotal in determining the most suitable indicators of tree physiological stress effects, which can 424 then be utilized in the analysis of results obtained from numerical experiments. 425

426 The  $E_{\rm T}$  estimates of all the forest stands were scaled up to the catchment area, revealing significant differences in contribution among various forest stands, such as Beech versus Oak, 427 with minor interannual changes (Figure 3.A). The interquantile range of catchment-scale  $E_{\rm T}$  (i.e., 428 Wei. in Figure 3A) ranges between 0.5 mm d<sup>-1</sup> and 3.0 mm d<sup>-1</sup> for 2019 and 1.0 mm d<sup>-1</sup> and 4.0 429 mm  $d^{-1}$  for 2020, reaching a maximum of 5.8 mm  $d^{-1}$  on certain days in June 2020 (see Figure 430 S5). Segregation of  $E_{\rm T}$  per plant functional type (PFT) indicates that broadleaf deciduous trees 431 (DBT) contribute more significantly than needleleaf evergreen trees (NET). Among individual 432 species, Beech stands out as the largest contributor to catchment  $E_{\rm T}$ , reaching up to 3.0 mm d<sup>-1</sup>, 433 with peak rates during 2020. Oak transpiration is at most 1.0 mm d<sup>-1</sup>, with its highest  $E_{\rm T}$ 434 occurring in 2019 (see Figure S5). In contrast, needleleaf trees (Douglas Fir and Spruce) exhibit 435 similar transpiration rates between the two years, not exceeding 0.5 mm d<sup>-1</sup> due to the smaller 436 area they occupy in the catchment. 437

The PCA highlights that  $E_{\rm T}$  of all forest stands is primarily influenced by solar radiation 438  $(R_s)$  and relative humidity (RH) (Figure 3.C).  $R_s$  lies parallel to the first dimension of the PCA, 439 which explains more than 70% of the variability of the  $E_{\rm T}$  and environmental variables (Figure 440 3.C).  $R_s$  has correlation coefficients larger than 0.8 with the  $E_T$  from all the forest stands (see 441 Figure S6). Conversely, the RH is negatively correlated with  $E_{\rm T}$ , showing values below -0.65. It 442 is noteworthy that  $\theta$  has on  $E_{\rm T}$  for all the forest stands, with negative correlation coefficients 443 444 close to -0.3, which are not statistically significant (p > 0.05). This indicates that the  $E_T$  response 445 at the selected site, for all tree stands, is primarily an atmospheric-driven process.

In the hot year (2020), all species and PFTs experienced higher TWD compared to the 446 normal year (2021) (Figure 3B). Notably, the distribution of TWD does not overlap between the 447 448 two years for all forest stands, with only some outliers in the normal year reaching the median value of the hot year. At the PFT level, the bimodal distribution of TWD in Douglas fir during 449 the hot year (2020) indicates a significant shrinkage of the monitored tree during the summer, 450 followed by a sudden swelling process in autumn. Oak, Beech, and Douglas fir show a distinct 451 452 breaking point where the TWD measurements do not overlap between years (except for outliers), with values of approximately 300 µm for Beech, 400 µm for Oak, and 100 µm for Douglas fir. In 453 454 contrast, Spruce does not exhibit such a breaking point. The BDT displays the most significant 455 differences between years, with the TWD values in the hot year being twice as high as those 456 recorded during the normal year. At the species level, Oak and Beech trees show the most substantial differences, influencing the response of the entire BDT forest stand (Figure 3B). 457 458 Among needleleaf trees, Douglas displays the largest variability in TWD between years, while

the differences for Spruce are less pronounced. The PCA of TWD indicates that variations are

460 primarily explained and negatively correlated with  $\theta$  (Figure 3D). Other environmental variables

have a lower influence on the TWD response of the different forest stands (see Figure S7). The

462 proximity of all forest stands in the PCA highlights a consistent pattern where TWD is dependent

463 on changes in  $\theta$ , making this measurement a robust indicator of the soil water stress experienced

464 by forest ecosystems. However, it is less indicative of atmospheric water stress, as variables like

465  $R_{\rm s}$ , u (wind speed), and T (air temperature) are orthogonal to the TWD of different species.



467 **<u>Figure 3</u>**. Summary of the observed forest response to environmental drivers. The upper plots

468 (i.e., A and B) represent the interannual variability of transpiration ( $E_T$ ) and tree water deficit

(TWD), respectively. The lower plots (i.e., C and D) are the correlation circles of the principal

470 component analysis (PCA). Each PCA shows the two main dimensions of the analysis that

471 explain at least the 50% of all the data variability. The arrows labelled with species or PFT names

472 represent either  $E_{\rm T}$  (Panel C) or TWD (Panel D) for each species or PFT.

473 3.2 Standalone Forest Experiment

474 When examining each forest stand as an independent entity (Experiment 1), the  $E_{\rm T}$  shows larger values across PHT parameterizations (i.e., GMP, PAP, SSP) when the canopy stress factor 475  $(\beta)$  ranges between 0.6 and 0.8. The results highlight the absence of periods with a stressed 476 canopy ( $\beta$ <0.5). Among the needleleaf forest stands, NET-GMP exhibits the lowest index of 477 agreement ( $I_{oa}$ ) between simulated  $E_T$  and observed sap flux ( $I_{oa}$ : 0.48), indicating a consistent 478 479 overestimation of  $E_{\rm T}$  throughout the growing season (see Figure S8). The performance of the NET standalone parameterization significantly improves when using the PAP parameterization, 480 481 with  $I_{oa}$  increasing to 0.75. This improvement reduces the  $E_T$  bias throughout 2019, yet the overestimation persists during the second half of the growing season of 2020 (see Figure S8). 482 483 Conversely, employing a SSP PHT parameterization for needleleaf species leads to a marginal and minor improvement on the  $I_{0a}$  compared to the GMP parameterization for Douglas fir ( $I_{0a}$ : 484 0.563) and Spruce (I<sub>0a</sub>: 0.692), respectively. The PHT parameterization of Douglas fir maintains 485 486 a low bias in 2019 but struggles to reproduce  $E_{\rm T}$  in 2020, resulting in a large bias (see Figure S8). 487 Transitioning from GMP to PAP in the BDT PHT parameterization has a negligible effect on I<sub>oa</sub>, increasing from 0.757 (GMP) to 0.774 (PAP). However, both GMP and PAP parameterizations 488 489 systematically underestimate  $E_{\rm T}$  for both years (2019 and 2020). With the implementation of the SSP parameterization, the performance of the individual BDT forest stands deteriorates in both 490 years (see Figure S8) compared to the GMP and PAP parameterizations. The SSP for Oak mostly 491 underestimates  $E_{\rm T}$ , with a mean absolute error of 0.9 mm d<sup>-1</sup>, while SSP-Beech underestimates 492  $E_{\rm T}$  with a mean absolute error of up to 1.8 mm d<sup>-1</sup>, resulting in low I<sub>oa</sub> values of 0.58 and 0.55, 493 respectively (see Figure S8). 494



497 **Figure 4**. Modeled ( $E_{\text{T-mod}}$ ) and measured transpiration rates ( $E_{\text{T-mea}}$ ) of the standalone forest 498 stands using the generic model parameterization (GMP), the PFT-adjusted parameterization 499 (PAP), and the species-specific parameterization (SSP). The canopy stress factor ( $\beta$ ) is used to 500 visually characterize stress at the leaf level.

The simulated  $\Psi_{root}$  values for all PHT parametrizations of NET remain unchanged 501 between the hot year (2020) and the normal year (2021). These  $\Psi_{root}$  values exhibit a negative 502 503 correlation with TWD, indicating that more negative  $\Psi_{root}$  values are associated with higher TWD (Figure 5). This negative correlation is statistically significant only for the GMP 504 parameterization. It is worth mentioning that the lower  $\Psi_{root}$  values produced by the GMP 505 506 parameterization result from the associated  $\Psi_{p50}$  parameter that is almost 2.0 MPa lower than that 507 of the PAP and SSP parameterizations (Table 2). It's important to highlight a notable difference in the relationship between the simulated  $\Psi_{root}$  and measured TWD between the hot and normal 508 509 years for BDT species. In 2020, all three parameterizations (GMP, PAP, SSP) show a significant negative correlation between TWD and  $\Psi_{root}$ , with a p-value less than 0.001. However, in 2021, 510 they are uncorrelated. Finally, it is interesting to highlight the different ranges of water potential 511 values simulated for the roots of NET and BDT trees, driven by the different  $\Psi_{p50}$  values as 512

513 defined by the PHT parameterization (Table 2).

514



515

516 **<u>Figure 5</u>**. Correlation between simulated root water potentials ( $\Psi_{root}$ ) and measured tree water

517 deficit (TWD) of the standalone forest stands (Experiment 1) using the generic model

518 parameterization (GMP), PFT adjusted parameterization (PAP), and Species-Specific

519 Parameterization (SSP). The canopy stress factor ( $\beta$ ) is used to visually characterize the stress at

520 leaf level. The p-values (p) of the Pearson correlation coefficient (r) are not significant when

521 marked with n.s.

#### 522 3.3 Mixed Patched Forest experiment

The three PHT parameterizations exhibit minimal differences in  $E_{\rm T}$  magnitude and 523 canopy stress ( $\beta$ ). These minor differences are reflected in the values of the index of agreement 524 (I<sub>na</sub>), ranging from 0.798 (SSP PHT) to 0.824 (GMP PHT). Despite these nuances, the 525 relationship between  $\Psi_{root}$  and TWD is not significant in 2021 for all model parameterizations 526 (i.e., GMP, PAP, and SSP). However, a significant (p < 0.001) and negative correlation is 527 observed for 2020 (Figure 6). It's noteworthy to highlight the remarkably similar range of 528  $\Psi_{root}$  values (0 - 1.5 MPa) simulated by the model despite the distinct  $\Psi_{p50}$  values used by NET 529 (i.e., -5.2 MPa) and BDT (i.e., -2.65 MPa) in the GMP parameterization. This suggests that BDT 530 531 forest types play a crucial role as the main driver of the ecosystem status at the grid cell level.



534 **Figure 6**. modeled ( $E_{T-mod}$ ) and measured ( $E_{T-mea}$ ) transpiration and the correlation between root 535 matric potentials ( $\Psi_{root}$ ) and tree water deficit (TWD) of a mixed patched forest (Experiment 2) 536 using the generic model parameterization (GMP), the PFT adjusted parameterization (PAP), and 537 the Species-Specific Parameterization (SSP). The canopy stress factor ( $\beta$ ) is used to visually 538 characterize the stress at the leaf level of each parameterization. The p-values of the Pearson 539 correlation coefficient (r) are not significant when marked with n.s.

540 The performance of the simulated  $E_{\rm T}$  of the individual fractions of the mixed-patched 541 forest under all three parameterizations (i.e., GMP, PAP, SSP) is illustrated in Figure 7. It is 542 crucial to emphasize that differences in  $E_{\rm T}$  between the fractions arise from their 543 proportional contribution to the overall forest composition. Additionally, the sum of the 544 individual contributions always equals the grid cell values (grey background dots in each 545 subplot). The simulated  $E_{\rm T}$  values are associated, even in this experiment, with mild canopy 546 stress conditions, reflected by  $\beta$  values ranging between 0.7 and 0.8.

For the NET forest fraction, both GMP and PAP parameterizations tend to overestimate  $E_{T}$ . However, the level of agreement (i.e.,  $I_{oa}$ ) between modelled and measured transpiration values improved significantly (from 0.517 to 0.763) when the PHT parameterization at the PFT level is refined. In contrast, the modeled  $E_{T}$  for the BDT fraction underestimates the actual  $E_{T}$ , with negligible differences between the GMP and PAP parameterization. However, when the

- 552 SSP parameterization is applied in the mixed-patched forest configuration, the results for NET
- and BDT species are remarkably different. NET species reach an  $I_{oa}$  of 0.723 and 0.616 for
- 554 Douglas fir and Spruce, respectively, while BDT species have values of 0.656 and 0.672 for Oak
- and Beech, respectively. The low performance of BDT species is due to an overestimation of  $\sim 50\%$  for Oak and an underestimation of one-third for Beech compared to measured  $E_{\rm T}$ .
- $\sim 50\%$  for Oak and an underestimation of one-third for Beech compared to measured  $E_{\rm T}$ . Refining the PHT parameterization does not substantially affect the performance of the model
- when analyzing its integrated response at the grid cell level ( $I_{oa-G}$  values range between 0.79 and
- 559 0.82). However, it has significant implications in the analysis of the transpiration response of
- 560 individual fractions. The use of a refined PHT plays an important role in improving the simulated
- 561 model response for those needleleaf evergreen species growing as isolated patches within the
- 562 catchment. Conversely, the improved PHT parameterization does not enhance the simulation of
- the transpiration response for those broadleaf species growing in fully-mixed forest stands.
- 564 Interestingly, there is an increment in I<sub>oa</sub> when moving from standalone conditions (Experiment
- 1) to mixed patched forest conditions (Experiment 2) for all species. The improvement in  $I_{oa}$  for
- 566 Douglas fir (from 0.692 to 0.723) is slightly lower compared to that of Spruce (from 0.563 to
- 567 0.616), Oak (from 0.58 to 0.66), and Beech (from 0.55 to 0.67).





- 570 **Figure 7**. Simulated ( $E_{\text{T-mod}}$ ) and measured ( $E_{\text{T-mea}}$ ) transpiration rates of the individual fractions
- of a mixed patched forest (Experiment 2) using the generic model parameterization (GMP), the PFT-adjusted parameterization (PAP), and Species-Specific Parameterization (SSP). The canopy
- 572 stress factor ( $\beta$ ) is used to visually characterize the stress at the leaf level of each
- parameterization. The grey dots represent the  $E_{T-mod}$  of the grid cell for each PHT
- 575 parameterization according to the results in Figure 6. Each plot has two indices of agreement
- <sup>576</sup> representing the grid cell ( $I_{oa-G}$ ) and the individual fractions within the grid cell ( $I_{oa-F}$ ).
- Figure 8 illustrates the relationship between the  $\Psi_{root}$  and TWD for the mixed patched 577 forest condition (Experiment 2). In this model configuration, there is a significant negative 578 correlation (p<0.001) between these two variables for both BDT and NET PFTs during the hot 579 year (i.e., 2020). This significant relationship persists across PHT parameterizations for both 580 PFTs. A clearer signal is obtained when using the SSP parameters for both Oak and Beech in the 581 case of BDT and for Spruce in the case of NET. Note that this significant relationship did not 582 emerge for NET in Experiment 1 when this ecosystem was simulated as a standalone forest 583 stand. Similarly, the relationship between  $\Psi_{root}$  and TWD for BDT differs between Experiment 1 584 and Experiment 2. In mixed-patched conditions, trees share the same soil water resources, the 585 signal emerging for Oak species is highly influenced by Beech species due to their larger water 586 use that drives the soil water changes in the catchment. Lastly, it is interesting to highlight that 587  $\Psi_{root}$  remained close to zero in the wetter year (i.e., 2021) and when TWD is less than 300-400 588 μm during the hot year (i.e., 2020). However, when TWD exceeds 300-400 μm in the hot year, 589 there is a significant negative correlation between  $\Psi_{root}$  and observed TWD. 590





- 593 <u>Figure 8</u>. Correlation between the root matric potentials ( $\Psi_{root}$ ) and tree water deficit (TWD) of 594 the mixed patched forest (Experiment 2) using the generic model parameterization (GMP), PFT 595 adjusted parameterization (PAP), and Species-Specific Parameterization (SSP). The canopy
- 596 stress factor ( $\beta$ ) is used to visually characterize stress at the leaf level. The p-values of the
- 597 Pearson correlation coefficient ( $\mathbf{r}$ ) are not significant when marked with n.s.
- 598 3.4 Fully-mixed Forest Experiment

Figure 9 compares the results of Experiment 3, where fully-mixed conditions are 599 represented only by the BDT PFT at the Weierbach catchment. The I<sub>oa</sub> values of the three PHT 600 parameterizations (i.e., GMP, PAP, SSP) do not differ substantially at the grid cell level, with a 601 602 minor increase from 0.757 to 0.778 when moving from GMP to SSP (see Figure S2 and Table 603 2). The SSP parameterization enhances modeled  $E_{\rm T}$  for individual species when compared with the standalone condition of Experiment 1. The  $I_{oa}$  increases for both Oak (from 0.582 to 0.674) 604 and Beech (from 0.555 to 0.664). The shared resources at the grid cell (i.e., the soil 605 606 compartment) and a similar PHT influence the strong similarities in  $\Psi_{root}$  among parameterizations, where the GMP shows a slight difference with PAP and SSP at the grid cell 607

- level. The differentiation between the hot year (2020) and normal year (2021) year follows the
- same trend as in Experiment 1 for GMP-BDT, PAP-BDT, and SSP-Beech. However, when Oak
- shares the soil water resources with Beech (i.e., Experiments 2 and 3), the Oak fraction depicts
- 611 more negative  $\Psi_{\text{root}}$  than in Experiment 1.





- 616 using the generic model parameterization (GMP), the PFT-adjusted parameterization (PAP), and
- 617 the Species-Specific Parameterization (SSP). The canopy stress factor ( $\beta$ ) is used to visually
- characterize stress at the leaf level of each parameterization. The p-values of the Pearson
- 619 correlation coefficient (r) are not significant when marked with n.s.
- 620 3.5 Identifying the stress periods at Weierbach

In the 11-years of simulations, the mean daily air temperatures of the four hot years (i.e., 621 622 2015, 2018, 2019, 2020) exceeded 27 °C for prolonged periods in spring and summer, with big differences in frequency and timing among the years (see Figure S12). The response of the 623 624 canopy stress factor ( $\beta$ ) and leaf water potential ( $\Psi_{\text{leaf}}$ ) to high air temperatures is similar between normal and hot years, hampering a characterization of the stress periods based on these metrics. 625 Conversely, the root water potential ( $\Psi_{root}$ ) provides a good differentiation between hot and 626 normal years for broadleaf (i.e., BDT, Oak, Beech) forest fractions, but not for the needleleaf 627 (i.e., NET, Douglas fir, Spruce) forest fractions (Figure 10). This contrast showed by broadleaf 628 and needleleaf forest fractions for the  $\Psi_{root}$  is diluted at grid cell level, where the differentiation 629 630 between normal and hot years shows a mixed response from both land covers (see Figure S14). By selecting the minimum  $\Psi_{root}$  value (1<sup>st</sup> quantile) of normal years as an assumed limit for stress 631 conditions (dotted blue line in Figures 10 and S14), it is possible to identify the periods were the 632 vegetation experiences more stress in broadleaf forest and at grid cell level but not for the 633 needleleaf forest. 634





637 Figure 10. Multiannual variability of the root ( $\Psi_{root}$ ) water potentials per forest fraction simulated

- 638 for the period between 2011 and 2021 at the Weierbach catchment. The distinction between
- normal and hot years is made according to the European Drought Observatory (EDO, 2023)
- 640 classification and based on the meteorological observations in Luxembourg. Each line

corresponds to an individual year modeled between 2011 and 2021, differentiated between
 normal (blue lines) and hot (red lines) years.

#### 643 4 Discussion

4.1 Importance of a refined plant hydraulic representation of mixed forests

The data from experimental forest stands are crucial in benchmarking physiological 645 processes in vegetation (e.g., Chaney et al., 2016; Sabot et al., 2020). Most datasets describing 646 PHTs are based on forest stands of different ages, management practices, and plant organs (e.g., 647 Choat et al., 2012), so caution is advised when using them widely to describe vegetation (Chu et 648 al., 2021). The characterization of PHTs per individual PFT lumps species together by 649 considering only the latitude and plant physiognomy (e.g., Lin et al., 2015), neglecting regional 650 and current distribution of tree species. Physiological characterization is fundamental to 651 reproducing vegetation water use strategies in CLM5 (Vidale et al., 2021). Li et al. (2021) 652 stressed the importance of implementing a better hydraulic trait-based representation to improve 653 the flux simulations in land surface models (LSMs) during drought events. This is crucial 654 because CLM5's generic parameterization is highly sensitive to dry conditions, affecting gross 655 primary productivity (GPP) and E<sub>T</sub> fluxes (e.g., Jiménez-Rodríguez et al., 2022; Wu et al., 2020). 656

While some assume that CLM5's PHT parameterization represents ecosystems correctly 657 (e.g., Raczka et al., 2021; Wu et al., 2020), present study shows differences in PHT leading to an 658 overestimation of  $E_{\rm T}$  in the GMP compared to PAP or the SSP. Some have improved the PHT 659 parameterization of CLM5, enhancing the representation of the tree species in several forest 660 stands dominated by a single tree species (e.g., Ali et al., 2022; Jiménez-Rodríguez et al., 2024). 661 In the present study, refining PHT parameterization for the individual needleleaf species (i.e., 662 SSP-Spruce, SSP-Douglas fir) drastically improved modeled  $E_{T}$ . However, uncertainties in PHT 663 for individual tree species complicate parameter selection, even at the species level (e.g., Lobo et 664 al., 2018; Rosner et al., 2019). Refinement efforts for two broadleaf species (Oak and Beech) did 665 not improve simulated  $E_{\rm T}$  when considered as standalone forest patches, possibly due to different 666 age structures and age-dependent PHT parameters (Lutz et al., 2013; Zeller & Pretzsch, 2019). 667 Hence, by including the SSP we largely improved the  $E_{\rm T}$  simulations of individual forest stands 668 despite the small differences at grid cell level. This condition enforces the need for an individual 669 evaluation of model performances for each forest fraction (Figure 7), rather than relying solely 670 on aggregated values at the grid cell level (Figures 6). 671

672 To date, few studies have investigated the effect of refining the PHTs in CLM5 for fullymixed forests. For example, Birch et al. (2021) and Kennedy et al. (2019), addressed the PHT 673 characterization from a lumped perspective focusing on the overall fluxes (e.g., eddy-covariance 674 data) and not on the individual performance at the species level. In the current study, LAI was 675 segregated between Oak and Beech in a fully-mixed stand, and simulated  $E_{\rm T}$  for both species 676 was compared to sap flow data separated by species. Surprisingly, no improvement in  $E_{\rm T}$ 677 simulations resulted from species-specific PHT parameterization. This suggests potential 678 deficiencies in other aspects of the model, such as the representation of water use strategies 679 (Jiménez-Rodríguez et al., 2024), or competition for light and water resources between the 680 species. 681

#### 683 4.2 The stressed vegetation and the hidden roots

Extreme temperatures and drier soil conditions are the main factors causing a decline in 684 gross primary productivity in forest ecosystems (J. Li et al., 2023). These conditions bring about 685 significant changes in water potentials within the plant, helping identify periods of high stress 686 when vegetation could suffer irreversible damage. Predawn water potentials ( $\Psi_{\text{predawn}}$ ) have been 687 used in the past to assess stress in forest ecosystems (e.g., Lavoir et al., 2009; Peiffer et al., 688 2014), even though continuous monitoring of  $\Psi_{predawn}$  is challenging. Although there are no 689 direct measurements of plant water potentials in the Weierbach catchment, insights into tree 690 water status can be deduced from TWD, strongly linked to midday canopy water potentials 691 (Steppe, 2018). Our PCA analysis shows that TWD measurements at Weierbach heavily depend 692 on soil water content ( $\theta$ ), while remaining unaffected by atmospheric drivers (Figure 3D). Tree 693 water storage has been identified as an effective mechanism that prevents trees from dehydrating 694 695 when the air around them becomes dry (Preisler et al., 2022). The capacity of trees to store water depends on  $\theta$  and their sapwood area (Hartzell et al., 2017). This helps them avoid stress caused 696 by water shortage if the water supply is enough to meet atmospheric dryness. Trees in Weierbach 697 have a considerably large sapwood area (see Table S3), which helps them buffer against 698 atmospheric drought through their water storage capacity. This also strengthens the relationship 699 between changes in  $\theta$  and stem diameter (i.e., TWD). 700

The simulated root water potentials ( $\Psi_{root}$ ) for all mixed forest configurations at the grid 701 cell level and for all broadleaf parameterizations (i.e. BDT, SSP-Oak and SSP-Beech) show a 702 703 strong negative correlation with observed TWD in the hot year, which is not observed in the normal year (Figures 5, 6, 8 and 9). In contrast, simulated leaf water potentials ( $\Psi_{\text{leaf}}$ ) and  $\Psi_{\text{root}}$ 704 705 for needleleaf species in the standalone simulations (Figure 5) are very noisy and do not show a 706 clear difference between the years. However, when the needleleaf fractions are in a mixed forest 707 configuration, their  $\Psi_{\text{root}}$  values do show a significant negative correlation with TWD (Figure 8), likely because of the strong influence of the co-occurring broadleaf species on the soil water 708 709 resources. Considering that the measured TWD does reach higher values in the hot year compared to the normal year for all species and the expected link between TWD and canopy 710 water potentials (Steppe, 2018), a stronger correlation between simulated water potentials and 711 TWD could suggest a more realistic model simulation. Since the needleleaf species cover a 712 713 separate part of the catchment in the Weierbach, it was not expected that they really do compete for water with the broadleaf species. However, the standalone model configuration resulted in 714 715 worse reproduction of observed sap flux for the needleleaf trees (Figure 4 vs. Figure 7) and a lower correlation between TWD and  $\Psi_{root}$  than the mixed patched forest configuration (Figure 5 716 vs. Figure 8), suggesting that the latter configuration produced more realistic results. 717

When modeling mixed forest ecosystems, CLM5 uses a single lower boundary condition 718 shared by multiple PFTs in each grid cell (Lawrence et al., 2019). This forces all PFTs to share 719 the same soil moisture pool, meaning that the dominant PFT drives the transpiration response 720 721 and water potentials of the other PFTs in a mixed forest configuration. The plant water uptake module allows the vegetation to use deeper water sources when the upper layers are too dry to 722 supply the canopy water demand (Kennedy et al., 2019; Lawrence et al., 2019). This 723 724 characteristic enables representation of the adaptability of adult Beech trees (Brinkmann et al., 2019) and deep water use by Oaks (Aranda et al., 2005; Bréda et al., 1993). However, it also 725 allows species that usually rely on shallower soil water to use deeper resources, forcing an 726 unrealistic response, as in the case of Spruce and Douglas Fir in Weierbach, where the vegetation 727

appears to be unaffected by water stress in the model. Therefore, when multiple PFTs share the

same grid cell in CLM5, the dominant cover in the landscape will mask or dilute important

aspects related to the water use of the other PFTs. This is the case in the Weierbach catchment,

<sup>731</sup> where BDT (for GMC and PAC) and Beech (for SSP) are the dominant forest fractions driving

the  $E_{\rm T}$  and hydraulic stress at the grid-cell level. The simplicity of the exponential root density profile used in most LSMs (e.g., JULES, CLM, ORCHIDEE) (Druel et al., 2017; Harper et al.,

2021; Lawrence et al., 2019) does not represent recent evidence of rooting strategies where the

735 fractional root allocation depends on the accessibility (e.g., depth) of the groundwater and

capillary fringe by the vegetation (Perona et al., 2022; Wang et al., 2018). Further, the

ran exponential root profile used in LSMs overlooks the fact that vertical root distributions can be

quite variable and dynamically follow changes in water availability (Fukuzawa et al., 2007;

739López et al., 2001; Zwetsloot & Bauerle, 2021).

4.3 The challenge of representing mixed forest ecosystems

The representation of PFTs in mixed forests is critical in LSMs because it may introduce 741 742 significant uncertainties concerning carbon, energy, and water fluxes (Hartley et al., 2017). In CLM5, applying it to complex ecosystems can lead to biases in flux and water stress estimation 743 (e.g., Song et al., 2020). Challenges arise in representing mixed patches and fully-mixed forests, 744 depending on the spatial distribution and contribution of different tree species. In this context, 745 patched mixed forests composed of well-spaced fractions that do not strongly interact 746 hydrologically can easily be represented by separate grid cells due to their physiognomic 747 748 differences (e.g., needleleaf versus broadleaf). In fully-mixed forest ecosystems (Experiment 3), where different tree species coexist and share soil resources, the random spatial distribution 749 750 challenges accurate representation of individual species in the model, impacting the precise calculation of transpiration fluxes (e.g.,  $E_{\rm T}$ ). This might explain why a more detailed 751 characterization of PHT per species did not enhance  $E_{\rm T}$  estimates (see Section 3.4). In the 752 753 Weierbach catchment, the heterogeneous distribution of Beech and Oak trees leads to varied LAI per species across the landscape, influencing how the ecosystem responds to water-limited 754 conditions (Forrester et al., 2022). The interactions among tree species and environmental drivers 755 may induce the trees to express different PHT within the range of each species (Drew et al., 756 2011; Lobo et al., 2018). For example, it has been found that some vulnerable tree species are 757 more resilient to drought conditions growing in a mixed forest stand than as a mono-specific 758

stands (Schäfer et al., 2019).

760 4.4 Challenges and opportunities in identifying vegetation stress in land surface models

In previous CLM versions, vegetation stress metrics were tied to a soil stress factor ( $\beta_{soil}$ ), connecting the soil compartment directly to the photosynthetic apparatus (Shrestha et al., 2018). CLM5 replaced this with the plant hydraulic system (PHS), using the plant vulnerability curve (PVC) for water transport and a canopy stress factor ( $\beta$ ) for leaf-level gas exchange (Kennedy et

767 (1 VC) for water transport and a canopy sites ratio (p) for real revergas exenange (Reinedy e al., 2019; Lawrence et al., 2019). However, using  $\beta$  alone as an indicator of tree stress in the

766 Weierbach catchment is misleading (see Figure S13), as it does not capture the hydraulic stress

conditions experienced by the species during the study period. To accurately identify stress

periods in CLM5, it is necessary to go beyond  $\beta$  and consider other metrics. The static

parameterization of PHTs in CLM5 ignores the temporal changes in plant physiology (Wieder et

al., 2019), allowing the vegetation to fully recover after a stress period without accounting for its

duration. This contrasts with evidence suggesting slow recovery after extreme hydraulic stress in
vegetation responses (Brodersen et al., 2019; Klein et al., 2018; Meinzer & McCulloh, 2013).

Using tree water deficit (TWD) as a proxy for canopy water potentials (Dietrich et al., 773 2018; Schäfer et al., 2019) suggests the possibility of using this variable as an indicator of stress. 774 In the Weierbach catchment, TWD clearly indicates the periods when most tree species (i.e., 775 776 Oak, Beech, and Dougas fir) are affected by hydraulic stress due to stem shrinkage. This stress is reflected in the model by the root water potential ( $\Psi_{root}$ ) at the grid cell level, particularly for 777 broadleaf covers, where  $\Psi_{root}$  is directly linked to soil water potential ( $\Psi_{soil}$ ), driving the root 778 system's response. The Weierbach catchment has experienced three consecutive hot years from 779 2018 to 2020 with limited water input. These frequent extreme weather events are likely to affect 780 the vegetation's response to hydraulic stress, resulting in legacy effects. However, it's difficult to 781 differentiate between normal and hot years using  $\Psi_{\text{leaf}}$  and  $\beta$  when compared to TWD. This could 782 be due to the model's PHS not accounting for tree capacitance, leading to an unintended lack of 783 legacy effects. The PHS includes the changes in  $\theta$  as the drivers of legacy effects, which better 784 represents the hydraulic stress response shown by TWD in Weierbach since the root system 785 depends on  $\Psi_{soil}$ . 786

#### 787 5 Conclusions

The forest's response to extreme weather events such as heatwaves is highly dependent 788 on forest structure and species composition, where mixed forests are presumed to be more 789 resilient to droughts. However, the representation of mixed forests using the plant functional type 790 791 (PFT) classification system can misrepresent the stress response of such complex ecosystems. This work addressed these relevant scientific issues by implementing the Community Land 792 Model version 5.0, at the experimental Weierbach catchment in Luxembourg. Several numerical 793 experiments were carried out, assuming three forest conditions (i.e., standalone, mixed-patched, 794 and fully-mixed forests). For each set of experiments, different plant hydraulic parameterizations 795 were used to characterize the PHS of the model. 796

The forest transpiration ( $E_{\rm T}$ ) based on sap flow measurements did not exceed 4.0 mm d<sup>-1</sup>, 797 with minor differences between normal and hot years. Catchment  $E_{\rm T}$  was strongly influence by 798 solar radiation  $(R_s)$  and relative humidity (RH). The estimations of tree water deficit (TWD) 799 800 based on incremental stem diameter recordings showed a strong difference between years, with 801 the hot year (i.e., 2020) having much larger values than the normal year (i.e., 2021) across species and PFTs. The interannual differences of TWD were primarily affected by the soil water 802 content ( $\theta$ ) suggesting that soil drought rather than atmospheric drought was the primary driver 803 for vegetation stress. 804

The three numerical experiments showed that the characterization of plant hydraulic traits 805 (PHTs) based on species rather than a "PFT-typical" parameterization improved the  $E_{\rm T}$  estimates 806 of homogeneous forest stands and to a lesser extent that of mixed forests. Also, the use of a 807 808 refined parameterization improved the capability of the model in reproducing the vegetation water stress at both coarse (i.e., PFT) and fine (i.e., species) levels for the broadleaf stands. 809 Representing fully-mixed forests as individual species showed limitations on the  $E_{\rm T}$  estimates as 810 even though the grid-cell level do not differ among parameterizations. These results highlight the 811 importance of refining the characterization of the PHT assigned to conventional PFT, where the 812 improvement should consider the spatial distribution of the species present in a region. The 813 814 results of the numerical experiments showed the constraints to use the simulated root water

- potential ( $\Psi_{root}$ ) to identify hydraulic stress periods in needleleaf covers. Whereas the
- 816 performance of  $\Psi_{root}$  in broadleaf covers highlighted its utility to identify such periods of
- 817 hydraulic stress.
- 818

#### 819 Acronyms

Acronym	Meaning
BDF	Broadleaf deciduous forest
BDT	Broadleaf deciduous temperate trees
EDO	European drought observatory
GMP	Generic model parameterization
LAI	Leaf area index
PAP	PFT adjusted parameterization
PCA	Principal component cnalysis
PFT	Plant functional type
Ioa	Index of agreement
	-f forest fraction
	-g grid cell level
NEF	Needleleaf evergreen forest
NET	Needleleaf evergreen temperate trees
PHS	Plant hydraulic system
PHT	Plant hydraulic traits
PLC	% loss of hydraulic conductance
PVC	Plant vulnerability curve
SSP	Species specific parametrization
TWD	Tree water deficit

820

#### 821 Acknowledgments

- 822 This work is supported by the Luxembourg National Research Fund (FNR) CORE program
- 823 (grant no. FNR/CORE/C19/SR/13652816/CAPACITY). GF and RS acknowledges the financial
- support of the FNR (grant no. AFR/STEP-UP ID 12546983, grant no.
- 825 FNR/CORE/C17/SR/11702136/EFFECT). The authors would like to thank Jérôme Juilleret for
- his invaluable field assistance in the collection of sap flow and dendrometer data and Lindsey
- 827 Stokes for proofreading the manuscript.

#### 828 **Open Research**

- 829 The COSMO-REA6 data used to complete the missing data for the atmospheric forcing in the
- study are available from the opendata-FTP server at DWD
- 831 (<u>https://opendata.dwd.de/climate\_environment/REA/COSMO\_REA6/</u>) (COSMO-REA6, 2019).

- CLM5.0 is publicly available through the Community Terrestrial System Model (CTSM) git
- repository (Tag name: release-clm5.0.34) via <u>https://github.com/ESCOMP/ctsm</u> (CTSM,
- 834 2017/2022). All analyses were done with the open source software R (R Core Team, 2022). The
- atmospheric forcing, daily averages of sap flow and dendrometer data, the results of the
- 836 numerical experiments, and the R scripts for post-processing the data are available from
- 837 ZENODO repository via <u>https://doi.org/10.5281/zenodo.7306884</u> with a Creative Commons
- Attribution 4.0 International license for the files (Jiménez-Rodríguez, 2023).
- 839

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Figure 1.



## Broadleaf Deciduous Forest (BDF)

Area: Tree Density: Species:

77.6 % 302 tree ha<sup>-1</sup> Fagus sylvatica *Quercus petraea x robur* 





Species:

# Needleleaf Evergreen Forest (NEF) Area:

22.4 % 319 tree ha<sup>-1</sup> Tree Density: Picea abies Pseudotsuga menziesii



Figure 2.

Standalone Forests Experiment 1 Mixed Patched Forests Experiment 2 Fully Mixed Forests Experiment 3









GMP

PAP







SSP





Figure 3.



Figure 4.



Figure 5.



Figure 6.



Figure 7.



Figure 8.



Figure 9.



▲ 2020 × 2021



Figure 10.

