

# Ozone causes substantial reductions in the carbon sequestration of managed European forests

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## Abstract.

The annual, accumulated stomatal ozone uptake during the vegetation season, i.e. the species-specific Phytotoxic Ozone Dose  
15 above a threshold of  $1.0 \text{ nmol m}^{-2} \text{ s}^{-1}$ , POD<sub>1</sub>SPEC, was estimated for European forest tree plant functional types for the years  
2008 – 2012. These POD<sub>1</sub>SPEC estimates were based on ozone concentrations simulated with the EMEP CTM model in  
combination with stomatal ozone uptake estimated with the DO<sub>3</sub>SE (Deposition of Ozone for Stomatal Exchange) model.  
POD<sub>1</sub>SPEC -based dose-response relationships for impacts of ozone on forest growth rates were constructed based on results  
from multi-year experiments with young trees generated within the framework of the UNECE LRTAP Convention. Official  
20 information on forest gross growth rates and natural and harvest removals for different European countries for the years 2008  
– 2012 were used to estimate annual changes in forest living biomass carbon (C) stocks due to ozone. This was achieved using  
two different scenarios, with and without the negative impacts of ozone on forest gross growth rates, estimated using the  
POD<sub>1</sub>SPEC -based dose-response relationships for impacts. Results estimated the annual gap between forest gross growth and  
the total removals, i.e. the annual forest stock changes, both for current levels of ozone and associated impacts, as well as in  
25 the absence of ozone. Estimates were made by collating species-specific information into broad European coniferous and  
deciduous forest types for consistency with forest statistics. The default IPCC methodology was used to convert estimates of  
the impacts of ozone on the annual changes in forest living biomass C stocks. The results showed that the critical level for  
negative ozone impacts on forests suggested by the UNECE LRTAP Convention, based on POD<sub>1</sub>SPEC, was exceeded in large  
parts of Europe during 2008 - 2012, except for inland areas in the Mediterranean and for small parts of Continental Europe  
30 and the Fennoscandian mountain range. The highest POD<sub>1</sub>SPEC was estimated for the coastal regions of mid-latitude Europe  
including the UK, and extending northwards to mid-Sweden, south Norway and Finland. To the south, lower values for  
POD<sub>1</sub>SPEC were estimated for most of the Iberian Peninsula and parts of the Mediterranean coastal regions. It was estimated

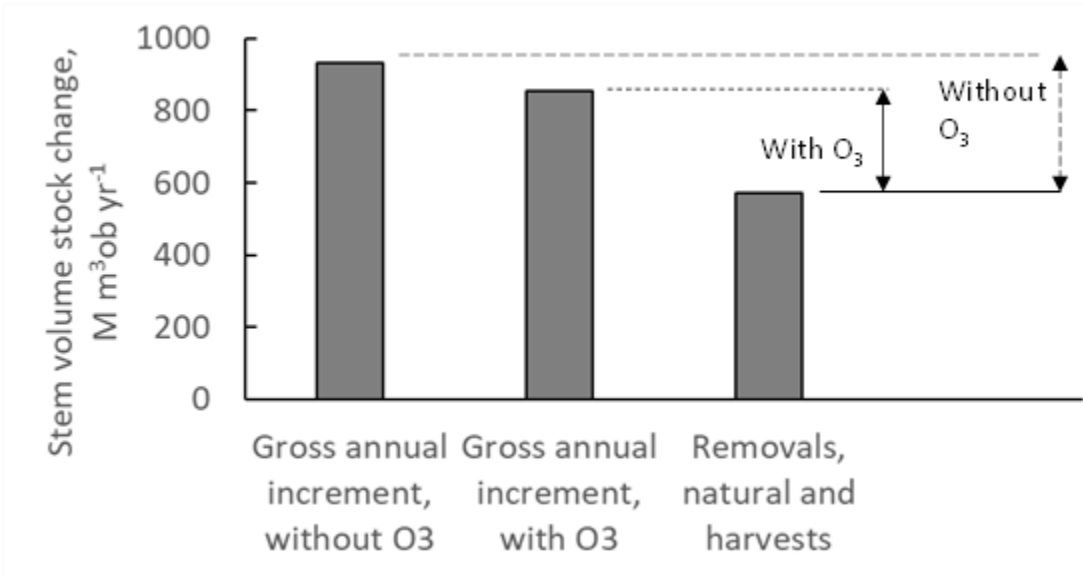
that reduced ozone exposure, similar to pre-industrial conditions, would increase European forest stem volume growth rates by 9%, but it would increase European forest annual net changes in standing stocks by 28%. The difference in gross forest stem volume growth with and without ozone impacts was relatively similar in, for example, Germany and France. However, since the gap between gross growth and total removals was much smaller for Germany, the enhanced growth in the absence of ozone had a much larger relative impact on the forest standing stock changes in Germany, compared to France. Summarized for all European forests, the C sequestration to the living biomass C stock was estimated to increase by 31% in the absence of ozone exposure. A thorough review of the literature resulted in the conclusion that mature trees under field conditions cannot be assumed to be less sensitive to ozone exposure compared to young trees under experimental conditions, strongly suggesting that these results are credible for European forest stands of different age classes.

### Short summary

Stomatal ozone uptake and the negative impacts on forest growth rates were estimated for European forests. This was translated to annual increments in the forest living biomass carbon stocks, with and without ozone exposure. In the absence of  $O_3$  exposure, European forest growth rates would on average increase by 9%, but the sequestration to the living biomass carbon stocks would increase by 31%, since the sequestration depends on the difference between growth and harvest rates.

Key figure

50 **Yearly changes in the European forest stem volume stocks, with and without the impacts of ozone pollution.**



## 1 Introduction

55 Forests are important for cost-effective, land-based climate mitigation (Griscome et al., 2017; Roe et al., 2021), including carbon (C) sequestration by the increases in the forest ecosystem C stocks (Pan et al., 2011). Furthermore, long-lived biogenic raw materials produced by forests can store C in harvested wood products and substitute for the use of fossil-based materials (Sathre and Gustavsson, 2009; Jasinevičius et al., 2015; Gustavsson et al., 2017). The yearly net C sink in global forests ecosystems has been estimated at 4 000 Mt CO<sub>2e</sub> (Pan et al., 2011) of which approximately 80% is stored in the living biomass  
60 C stocks. The importance of European forests in this capacity has been clearly demonstrated (Hyyrynen et al., 2023; Korosuo et al., 2023). European forests are likely to absorb ~10 % of all European anthropogenic CO<sub>2</sub> emissions (Janssens et al., 2003). The C stocks in European forests above and below ground biomass are in the order of 48 000 M t CO<sub>2e</sub> and it has been estimated to have increased annually over the time period 2010 – 2020 by 560 M t CO<sub>2e</sub>, i.e. by 1.4% annually (Forest Europe, 2020). Forest growth rates in many European countries are primarily determined by forest management (Etzold et al., 2020). In  
65 general, forests that are actively managed sequester C in the forest ecosystem carbon stocks at higher rates than non-managed forests (Nabuurs et al., 2008). The C sequestration of managed forests to a large extent depends on the balance between forest growth and removals, including harvests and natural losses (Soimakallio et al., 2021). Any measures that increase the productivity of temperate or boreal forest are likely to increase the forest C sequestration.

70 Forest growth rates are also affected by environmental conditions such as meteorological variables and air pollutants. Tropospheric ozone (O<sub>3</sub>) is an air pollutant that has been found to cause losses to tree biomass (Emberson, 2020). O<sub>3</sub> impacts on vegetation is estimated based on the accumulated amounts of O<sub>3</sub> that is taken up to the leaf interior through the stomata during the vegetation season, i.e. the Phytotoxic Ozone Dose, POD (Mills et al. 2011). Experimental studies fumigating young trees with O<sub>3</sub> for up to 10 years have been conducted across Europe for different tree species (Karlsson et al., 2007; Wittig et al., 2009). These experimental studies have been used within the UNECE Convention on Long-Range Transboundary Air  
75 Pollution (LRTAP) to develop POD-based dose-response relationships that can be used to identify areas across Europe where losses in whole tree biomass may occur due to elevated O<sub>3</sub> concentrations (LRTAP, 2017). Critical levels for forest trees were set to prevent an annual biomass loss under experimental conditions exceeding between 2 (coniferous forests) and 4% (deciduous forests), based on above ground or whole tree biomass. Maps of ozone uptake and exceedance of POD critical  
80 levels (Simpson et al., 2007, 2022, Franz et al., 2017) show where forests might be at risk of damage due to O<sub>3</sub> pollution. Identifying areas of such critical level exceedance can be used within LRTAP to apply an effects-based emission reduction policy (Emberson, 2020).

It is important to estimate actual O<sub>3</sub> impacts to forest trees under ambient O<sub>3</sub> pollution concentrations, since this would allow  
85 assessment of O<sub>3</sub> impacts on forest health (Marzuoli et al., 2019), productivity (Karlsson et al., 2005) and C sequestration rates, the latter becoming increasingly more important as nature-based solutions are promoted across Europe to increase

climate resilience (Calliari et al., 2022). Due to the multi-year lifetime of forest trees, O<sub>3</sub> impact studies involve a time component over which effect estimates are integrated. Many O<sub>3</sub> impact studies for trees report only the percent reduction of biomass caused by O<sub>3</sub> at the end of the experiment and information on the biomass at the start of the experimental fumigation with O<sub>3</sub> is often not provided. As a result, impacts on tree growth rates cannot be calculated directly. The significance of this problem increases at low growth rates in relation to the size of the O<sub>3</sub> effect (Karlsson, 2012). Information about O<sub>3</sub> impacts on forest growth rates is necessary for relevant assessments of long-term impacts of O<sub>3</sub> on forest growth and C sequestration (Korosuo et al., 2023). The use of the whole tree biomass, POD-based, dose-response relationship can then be applied for assessments of long-term impacts of O<sub>3</sub> on forest C sequestration in combination with methods (such as forest growth models) that are able to simulate O<sub>3</sub> effects on growth rates in a complex forest environment (Sitch et al., 2007; Franz et al., 2018; Subramanian et al., 2015; Otu-Larbi et al. 2020).

The first phase of the Tropospheric Ozone Assessment Report (TOAR; <https://igacproject.org/activities/TOAR/TOAR-I>) built the world's largest database of O<sub>3</sub> metrics to identify the global distribution of the pollutant and trends in O<sub>3</sub> concentrations over time. The second phase of TOAR (<https://igacproject.org/activities/TOAR/TOAR-II>), to which this paper contributes, has a broader scope, with the aims being to investigate the impact of tropospheric O<sub>3</sub> on human health and vegetation. The present work will address these goals by assessing the effects of ozone on C sequestration in forest living biomass. Hence, in this study we re-analyse existing whole tree biomass, POD-based, dose-response relationships based on multi-year experiments with young trees to provide estimates of O<sub>3</sub> impacts on the tree growth rates and changes in the forest living biomass C stocks. Consequently, O<sub>3</sub> impacts on European forest gross growth rates can be estimated using the LRTAP experimental information (LRTAP, 2017) in combination with readily available forest statistic information describing annual increments in forest stocks (Forest Europe, 2020, UNECE, 2011).

The overall aim of this study was to apply LRTAP air quality guidelines to estimate the impacts of the present O<sub>3</sub> exposure, based on POD, on the C sequestration to the living biomass C stocks of Europe's managed forests. This aim was achieved by addressing the following research questions:

1. How does POD vary spatially for key coniferous and deciduous forest tree species and functional types across Europe?
2. How can O<sub>3</sub> dose – response relationships for biomass reduction be converted into growth rate relationships for key European forest tree species and forest plant functional types?
3. How can impacts of O<sub>3</sub> exposure on the C sequestration of living biomass C stocks of forests be estimated, based on the difference between forest gross growth rates and total removals, the latter including both harvests and natural losses?
4. What is the magnitude of the negative impacts of present O<sub>3</sub> exposure on the C sequestration of the living biomass C stocks in forests, aggregated at both the European and national levels?

**2.1 Overview of methods**

In Figure 1 it is described how datasets and models are integrated and used to quantify the effect of O<sub>3</sub> on changes in annual living biomass C stock increments (Mt CO<sub>2e</sub> yr<sup>-1</sup>) of managed forests across Europe. Gridded surface meteorological data and O<sub>3</sub> concentration data at ca. 50 m height (assumed to be the top of the atmospheric surface or the constant flux layer, c.f. Stull  
125 (1988)) for Europe were provided from the EMEP MSC-W chemical transport model (CTM) (Simpson et al., 2007, 2012). Version rv4.35 of the EMEP model (Simpson et al., 2020), as used here, is driven by 3-hourly meteorological data from the European Centre for Medium Range Weather Forecasts Integrated Forecasting System (ECMWF-IFS, <https://www.ecmwf.int/en/research/modelling-and-prediction>). These meteorological data are interpolated to hourly resolution inside the EMEP model and variables such as cloud-cover and vapour pressure deficits (VPD) are calculated. Anthropogenic  
130 emissions data for the EMEP model are predominantly derived from official national estimates (Mathews et al., 2020), and biogenic emissions (isoprene, terpenes, etc) calculated as in Simpson et al. (2012). These calculations used 20 vertical layers, and O<sub>3</sub> data from the lowest vertical layer (which approximates to an average height of 50 m above the surface) were used in this study. For consistency with the land cover mapping data used in this project (Cinderby et al., 2007), all data were provided at a horizontal resolution of ~ 50 x 50 km<sup>2</sup> on the polar-stereographic which was used until 2016 by EMEP  
135 ([https://www.emep.int/mscw/emep\\_grid.html](https://www.emep.int/mscw/emep_grid.html)).

**Figure 1: Overview of the datasets and methods used to estimate the influence of O<sub>3</sub> on forest annual living biomass C stock increments (Mt CO<sub>2eq</sub> yr<sup>-1</sup>) for the time period 2008 - 2012. A negative sign was used for the removal of CO<sub>2</sub> from the atmosphere to the living biomass carbon stocks.**

140 The DO<sub>3</sub>SE (Deposition of Ozone for Stomatal Exchange, Emberson et al., 2000; 2007; Büker et al., 2012; 2015) dry deposition model estimates gridded O<sub>3</sub> concentrations down to the forest canopy height. From this, the stomatal O<sub>3</sub> flux values, i.e. the POD<sub>1</sub>SPEC (Phytotoxic Ozone Dose above a threshold of 1 nmol m<sup>-2</sup> Projected Leaf Area s<sup>-1</sup> for Specific tree species) metric recommended to assess O<sub>3</sub> risk to forests by the UNECE Mapping Manual (LRTAP, 2017), are calculated using the  
145 multiplicative stomatal conductance module, parameterised for bioregion-specific species (determined according to the biogeographic regions of Europe (EEA, 2016) and plant functional types (PFTs). POD<sub>1</sub>SPEC values are used with gross annual stem volume increment (GAI) dose-response relationships (DRRs) to estimate the effect of O<sub>3</sub> on GAI. These grid level estimates are aggregated to the national level using forest coverage data from the UNECE harmonized land cover map (Cinderby et al., 2007) scaled according to the mean 2008 to 2012 estimates of the total forest cover (Forest Europe, 2020, UNECE, 2011). This results in country level estimates of GAI, with and without O<sub>3</sub> pollution. Country level GAI values are  
150 converted to net annual stem volume increment (NAI) values by allowing for forest removals by both natural causes and fellings. Country values for annual net change in forest standing stocks (NCSS), both with and without O<sub>3</sub> pollution, are

converted to changes in living biomass C stocks using default methodologies of the IPCC (Penman et al., 2003), which include both above and below ground living biomass. This allows the effect of O<sub>3</sub> on annual C sequestration in the living biomass of managed European forests to be estimated.

## 2.2 Calculating POD<sub>1</sub>SPEC

The DO<sub>3</sub>SE deposition model was used in an off-line mode to calculate total deposition of O<sub>3</sub>. This allows the O<sub>3</sub> concentration at the top of the forest canopy (i.e. at the interface between the atmosphere and the canopy-influenced boundary layer) to be estimated for the different forest types across Europe. This off-line calculation uses the DO<sub>3</sub>SE stomatal flux model parameterisations provided in Table S1 to estimate the stomatal and non-stomatal deposition sink for O<sub>3</sub> from which the canopy height O<sub>3</sub> concentration  $c(z_{-1})$  in (nmol m<sup>-3</sup>) at height  $z_{-1}$ (m) can be calculated as described in Eq. 1.

$$c(z_1) = c(z_{Ref}) \cdot [1 - Ra(z_{Ref}, z_1) \cdot Vg_{Ref}] \quad [1]$$

where  $c(z_{Ref})$  is the concentration of O<sub>3</sub> from ca. 50m, assumed to represent the top of surface layer in the EMEP model,  $Ra(z_{Ref}, z_1)$  is the atmospheric resistance to O<sub>3</sub> transfer between  $z_{Ref}$  and  $z_1$ , and  $Vg_{Ref}$  is the deposition velocity of the tree canopy. It is important to note that this "big-leaf" calculation of  $c(z_1)$  makes use of big-leaf resistance and conductance terms, which differ from the leaf-level values used below (see Tuovinen et al., 2008, for more discussion of these distinctions).

We define stomatal O<sub>3</sub> flux model parameterisations by species, bioregions and PFTs since we know that stomatal O<sub>3</sub> uptake by forest trees is dependent upon tree physiology which varies by species and bioregion, the latter often causing different physiology in the same species (LRTAP, 2017). We define nine bioregions across Europe using the biogeographic regions of Europe map (EEA, 2016; see Table S2) and apply the appropriate stomatal conductance ( $g_{sto}$ ) model parameterisation (Table S1) in the estimation of  $g_{sto}$  following the components of Eq. [2] that are required for forest tree species as described in the UNECE Mapping Manual (LRTAP, 2017).

$$g_{sto} = g_{max} \cdot f_{phen} \cdot f_{light} \cdot \max\{f_{min}, (f_{temp} \cdot f_{VPD} \cdot f_{PAW})\} \quad [2]$$

Where  $g_{sto}$  is the actual stomatal conductance (mmol O<sub>3</sub> m<sup>-2</sup> PLA s<sup>-1</sup>, PLA, projected leaf area) and  $g_{max}$  is the species-specific maximum  $g_{sto}$  (mmol O<sub>3</sub> m<sup>-2</sup> PLA s<sup>-1</sup>) and the parameters  $f_{phen}$ ,  $f_{light}$ ,  $f_{temp}$ ,  $f_{VPD}$ ,  $f_{PAW}$  and  $f_{min}$  are all expressed in relative terms.  $f_{phen}$  allows for the variation in  $g_{sto}$  during the growing season,  $f_{min}$  defines a daytime minimum  $g_{sto}$  and the remaining factors represent the modifying influence of environmental variables (irradiance, temperature, atmospheric water vapour pressure deficit and plant available soil water respectively).

Stomatal conductance model parameterisations only exist for a limited number of forest tree species/ PFTs, namely: birch  
 185 (*Betula pendula*), beech (*Fagus sylvatica*), Norway spruce (*Picea abies*), Scots pine (*Pinus sylvestica*), Aleppo pine (*Pinus  
 halepensis*), temperate deciduous oak (*Quercus robur*), Mediterranean deciduous oak (*Quercus faginea*, *Quercus pyrenaica*  
 and *Quercus robur*) and Mediterranean evergreen oak (Holm oak, *Quercus ilex*). Table S2 describes the combination of  
 stomatal O<sub>3</sub> flux parameterisations (to estimate the POD<sub>1</sub>SPEC values) and GAI DRRs (to estimate the damage caused by  
 POD<sub>1</sub>SPEC that are used in each bioregion along with the number of EMEP grids these bioregions cover). Table S2 shows  
 190 that for those bioregions that comprise > 100 EMEP grids, between 29 and 85% of forest area is represented by these specific  
 species, the remainder being classified as other deciduous, other coniferous or mixed and represented by an appropriate species  
 of that bioregion and forest type.

The stomatal O<sub>3</sub> flux (F<sub>st</sub>) of sunlit leaves or needles of the upper canopy can then be calculated for forests across Europe using  
 195 Eq. [3].

$$F_{st} = c(z_1) \cdot g_{sto} \cdot \frac{r_c}{r_b + r_c} \quad [3]$$

where g<sub>sto</sub> is in m s<sup>-1</sup>, r<sub>b</sub> is the leaf quasi-laminar resistance and r<sub>c</sub> the leaf surface resistance, both given in s m<sup>-1</sup>. For further  
 200 details on the resistance scheme, see UNECE Mapping Manual (LRTAP, 2017).

The accumulated species-specific POD<sub>1</sub>SPEC (mmol O<sub>3</sub> m<sup>2</sup> PLA) was calculated according to Eq. [4] by summing modelled  
 hourly F<sub>st</sub> values (F<sub>sti</sub>) over the smaller of either a 6-month period (1 April to 30 Sept) or a species-specific period defined by  
 the forest latitude model (LRTAP, 2017).

$$POD_1SPEC = \sum_{i=1}^n [F_{sti} - y] \text{ for } F_{sti} \geq y \text{ nmol m}^{-2} \text{ projected leaf area (PLA) s}^{-1} \quad [4]$$

Where F<sub>sti</sub> is the hourly mean O<sub>3</sub> flux in nmol O<sub>3</sub> m<sup>-2</sup> PLA s<sup>-1</sup>, and n is the number of hours within the accumulation period.  
 The threshold of 1 nmol O<sub>3</sub> m<sup>-2</sup> PLA s<sup>-1</sup> (Mills et al., 2011; LRTAP, 2017) represents the ability of plants to detoxify a certain  
 210 amount of O<sub>3</sub>. POD<sub>1</sub>SPEC was calculated separately for coniferous and deciduous tree species and separately for scenarios  
 with (f\_PAW) and without (NSW) the influence of the estimated soil water deficit (Bueker et al., 2012).

### 2.3 Deriving POD<sub>1</sub>SPEC dose-response relationships for gross annual stem volume increment

POD<sub>1</sub>SPEC DRRs exist for a number of species/species groups and have been previously defined in B  ker et al. (2015). The  
 215 response parameter for these DRRs is a percentage reduction in total living biomass at the end of the experimental period,

corrected for the number of experimental years. As described in the introduction, this is not a suitable metric for use to estimate impact on long-term forest growth rates. To overcome this issue, we re-analysed the existing DRRs, converting the response variable from percentage reduction in total biomass to percentage reduction in GAI. This was achieved by defining a standard sigmoidal growth function for tree biomass based on the Richards equation [Eq. 5] (Richards, 1959).

$$Tree\ biomass = y_0 \left[ 1 - \exp(-y_1 \cdot tree\ age) \right]^{\left( \frac{1}{1-y_2} \right)} \quad [5]$$

Where  $y_0$ ,  $y_1$  and  $y_2$  are set to 1, 0.03 and 0.65 based on data collected for key European tree species from sites across Europe (Fellner & Rechberger, 2009; shown in Figure S1). This allows calculation of the tree biomass at the beginning of each  $O_3$  fumigation period (which usually starts when the tree is a few years old) and the GAI that would be expected each year until the end of the fumigation period. For each experiment, the  $O_3$  effect is estimated as the average percentage reduction on annual growth rate of each  $O_3$  treatment vs control, assuming that the relative  $O_3$  effect on growth rate is constant, irrespective of age of tree or GAI.

The GAI DRR is estimated by combining results of fumigation experiments conducted on a particular species or species group following the method of Fuhrer (1994). This estimates a regression for each individual experiment to define the biomass at zero  $O_3$  exposure. This is then used to scale the different treatment effects so that zero exposure is always associated with no effect at the individual experiment level. The percentage reductions in GAI for each experiment and treatment are pooled and a linear regression is drawn through the data to give a species- or PFT-specific DRR.

The UNECE harmonised land cover map (Cinderby et al., 2007), which describes coverage of individual forest tree species at a spatial scale of 1 x 1 km are aggregated to the ~50 x 50 km EMEP grids to provide an area-weighting for each species and PFT across the EMEP modelling domain. This area weighting is scaled according to the national forest area coverage data from the Forest Europe statistics (Forest Europe, 2020, UNECE, 2011) to ensure that national total forest cover is consistent with the GAI data.

## 2.4 Estimating the influence of $O_3$ on gross annual stem volume increments.

The species/ PFT specific DRRs are used to estimate the GAI under pre-industrial  $O_3$  concentrations (assumed constant at 10 ppb across the whole of Europe, after Volz & Kley, 1988) for each species/PFT in each grid. This allows an estimate of the reduction in GAI due to the occurrence of  $O_3$  pollution for each grid, which is then scaled to provide a country area weighted value for coniferous and deciduous (broadleaf) PFTs scaled, according to land cover.

## 2.5 Estimates of O<sub>3</sub> impacts on European forest carbon sequestration

Forest statistics for European countries are available at the national level from the Forest Europe statistics (<https://fra-data.fao.org/assessments/panEuropean/2020/FE/home/overview>), section “Increment and fellings, in forest available for wood supply”, indicator 3.1.). At the commencement of this study, information was available for forests at the national level for five-year periods until the period 2010, which is based on mean values for the period 2008 - 2012. In most cases information is presented separately for coniferous and broadleaf tree species. Hence, calculations of O<sub>3</sub> impact on European forest C sequestration were made separately for each country and separately for coniferous and broadleaf tree species. The assessment thus regards the annual values for the time period 2008-2012. Calculations were restricted for forests available for wood supply (FAWS). For most countries, the ratio of the area of total forest land to the area of FAWS did not exceed 1.3 (data not shown). Annual net changes in forest standing stocks, i.e. the summed stem volumes of living trees, were calculated according to the scheme outlined in Fig 2.

**Figure 2. An illustration of the scheme used to calculate the net changes in the forest standing stocks. Modified from Forest Europe (2020) and Korosuo et al. (2023). The arrow with the thickest line represents the final value calculated as the net forest standing stock change (NCSS).**

The dependent variable for O<sub>3</sub> impacts on net changes of forest C stocks was the GAI in FAWS. The annual rates of natural losses (ANL) are subtracted from the GAI to give the Net Annual stem volume Increment (NAI). It was assumed that O<sub>3</sub> exposure does not have an impact on the ANL, mainly since there are, as far as we are aware, no published studies on O<sub>3</sub> impacts on tree mortality. NAI is then subtracted by the annual rates of total fellings (ATF), including fellings of both living and dead trees and also including harvest residues, i.e. all biomass that is removed from the FAWS. The resulting values will be the annual net change in forest standing stocks (NCSS) in FAWS. Information was not available for all parameters listed above for all countries. Missing values were replaced based on the relations between the relevant parameters for nearby countries positioned within the same European forest region. The regions used were those used by Forest Europe (2020), i.e. North Europe, Central-West Europe, Central-East Europe, Southwest Europe, Southeast Europe and finally Russia with neighbouring countries.

The forest statistics for different European countries only provide information on the GAI, natural- and harvest removals of the total forests together with information for the different areas of coniferous and broadleaf forests. Hence, the share of the GAI, natural- and harvest removals between coniferous and broadleaf forests had to be assumed based on the differences in the forest areas. This assumption was tested and found to be relatively correct for Sweden, a country for which the relevant forest statistics were available separately for coniferous and broadleaf forests (data not shown).

280 The NCSS was converted to C stock changes as described in IPCC’s “Good Practice Guidance for Land Use, Land-Use Change and Forestry” (Penman et al., 2003).

$$\Delta C = I_v \cdot BEF \cdot (1+R) * D * CF \tag{6}$$

285 Where  $\Delta C$ , C sequestration to tree living biomass (tonnes C ha<sup>-1</sup> yr<sup>-1</sup>, a negative sign is used for the removal of CO<sub>2</sub> from the atmosphere to the living biomass carbon stocks;  $I_v$ , yearly increment of timber volume (m<sup>3</sup> ha<sup>-1</sup> yr<sup>-1</sup>);  $D$ , density stem (tonnes dry weight m<sup>-3</sup>);  $CF$ , “carbon fraction”, of dry matter (tonnes tonnes<sup>-1</sup>);  $BEF$ , biomass expansion factor, converts between stem biomass and total living biomass above ground;  $R$ , shoot/root ratio. The value of  $\Delta C$  was then converted to CO<sub>2</sub>-equivalents  
 290 (CO<sub>2e</sub>) by multiplying with 3.67. The constants used in the present study are shown in Table 1.

**Table 1. Constants used for the conversion from change in forest standing stock to changes in forest C stock, based on IPCC (Penman et al., 2003, Ågren et al., 2021).**

295 **3 Results**

The spatial distribution of the estimated POD<sub>1</sub>SPEC (mmol O<sub>3</sub> m<sup>-2</sup> PLA) across Europe is shown in Fig. 3, separately for coniferous and deciduous tree species, as an average value for the years 2008 to 2012. To give an indication of the variation in POD<sub>1</sub>SPEC between years, the differences between the minimum and maximum POD<sub>1</sub>SPEC values across 2008 to 2012 for conifers and deciduous species are also shown.

300 **Figure 3. The spatial distribution of the estimated POD<sub>1</sub>SPEC (mmol O<sub>3</sub> m<sup>-2</sup> PLA) across Europe accumulated during the growing season using the f<sub>PAW</sub> scenario, separately for coniferous (a) and broadleaf (c) tree species for annual mean values for the period 2008-2012 and as the difference in maximum and minimum POD<sub>1</sub>SPEC values across the years 2008-2012 (coniferous, b; broadleaf, d). Also shown is the mean annual value of AOT40 in ppb.hrs for the period 2008-2012 and the Continental Central Europe (CCE) growing season of Norway Spruce (e) and Beech (f).**

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The highest POD<sub>1</sub>SPEC values were estimated for the coastal regions of mid-latitude Europe including UK, for both coniferous (12 to 18 mmol O<sub>3</sub> (m<sup>2</sup> PLA)<sup>-1</sup>) and broadleaf (21 to 30 mmol O<sub>3</sub> (m<sup>2</sup> PLA)<sup>-1</sup>) forests, extending as far as mid-Sweden and south Norway and Finland. There were even high values estimated for the southern part of Iceland. To the south, lower values  
 310 for POD<sub>1</sub>SPEC were estimated for most of the Iberian Peninsula and all the Mediterranean coastal regions. There were high values estimated for POD<sub>1</sub>SPEC also in parts of the alpine region and some surrounding regions of inland continental Europe (Fig. 3a, c). There were low values (0 to 9 mmol O<sub>3</sub> (m<sup>2</sup> PLA)<sup>-1</sup>) for both coniferous and broadleaf deciduous estimated in

particular for the inland of the Iberian Peninsula but also for the eastern parts of continental Europe and the Balkan region. For coniferous forests, there were low values (around 6 to 9 mmol O<sub>3</sub> (m<sup>2</sup> PLA)<sup>-1</sup>) estimated for POD<sub>1</sub>SPEC also for large parts of Italy. In general, POD<sub>1</sub>SPEC values were considerable higher (by almost 10 mmol O<sub>3</sub> (m<sup>2</sup> PLA)<sup>-1</sup>) for deciduous as compared to coniferous trees (Fig. 3a, c). The inter annual differences (Fig. 3b, d) are also greater for broadleaf deciduous (often reaching up to 10 mmol O<sub>3</sub> (m<sup>2</sup> PLA)<sup>-1</sup>) compared to coniferous with inter annual variability over the five years of only up to around 5 mmol O<sub>3</sub> (m<sup>2</sup> PLA)<sup>-1</sup>. These differences are not overtly driven by soil water stress as might be expected since the NSW model runs also show a similar magnitude of variability, albeit over a slightly reduced extent across Europe (see Figure S2). Finally, it is clear from Fig. 3a, c, e and f that the AOT40 metric has a very different spatial pattern with a strong north to south gradient, identifying highest values (over 28 ppm.hrs) and hence risk in Mediterranean and mid to southern continental Europe.

Figure 4 gives an indication of which of growing season or key environmental variables (i.e. which of  $f_{temp}$ ,  $f_{VPD}$ ,  $f_{PAW}$ ) are most limiting of POD<sub>1</sub>SPEC across Europe. The length of the growing season may cause a limitation in the Northern latitudes when growing season length is below around 120 days (Figure 4a), however it is likely that temperature would also cause a limitation outside of the growing season. This is clear since the function used to describe the temperature limitation of stomatal conductance (Figure 4b) showed homogenously high values across most of Europe, except for the alpine region and northern Fennoscandia, Scotland and Iceland. The function used to describe the VPD limitation of stomatal conductance (Fig. 4c) showed a geographical variation that more resembled the pattern for the estimated POD<sub>1</sub>SPEC, with higher values for continental coastal regions and lower values for continental inland regions. The function used to describe the soil water deficit limitation of stomatal conductance,  $f_{PAW}$ , (Fig. 4d) showed low values only for the inland Iberian Peninsula.

**Figure 4. The limits to POD<sub>1</sub>SPEC from each of the key three environmental variables (a) growing season length (in days) for broadleaf deciduous forests, (b)  $f_{temp}$ , (c)  $f_{VPD}$  and (d)  $f_{PAW}$  averaged during the growing season for all species and all years (2008-2012) across Europe. Grid values are mean values across all plant functional types for the growing season and for daylight hours only. Plots for individual species are given in the supplementary (Figure S4 – S6).**

The GAI DRRs developed from the re-analysis of the experimental fumigation data are shown in Fig. 5b for coniferous species and 5d for broadleaf deciduous species. The percentage reduction in whole tree living biomass (taken from Büker et al. (2015)) are also shown (Fig. 5a for coniferous and 5c for broadleaf deciduous) for comparison. The GAI DRR regressions are given in Table S3. The GAI DRRs show small improvements in the statistical relationship of the regression between relative GAI and POD<sub>1</sub>SPEC, with R<sup>2</sup> values of 0.69 and 0.47 for coniferous and broadleaf deciduous respectively. The difference in the intercept at POD<sub>1</sub>SPEC is negligible between relative biomass and relative GAI but the slope for the negative impacts was slightly more negative for the DRRs based on GAI growth rate reductions. These results suggest the relative GAI DRRs are as robust as the relative biomass DRRs and hence are suitable for use in risk assessment.

**Figure 5. Dose-response relationships (DRRs) based on POD<sub>1</sub>SPEC for assessments based on impacts on the percent annual biomass loss (Relative biomass) (a, c) and the estimated impacts on gross annual increment (relative GAI) (b, d) separately for coniferous (a, b) and deciduous tree species (c, d). Other species and PFT dose-response relationships are given in the supplementary text (Table S3 and Figure S3).**

The estimated values for percent change in GAI, caused by the present O<sub>3</sub> compared to the absence of O<sub>3</sub> exposure, are shown in Table 2, calculated as mean values for the period 2008-2012 and mean values for different European countries. The reductions of GAI were calculated separately for coniferous and broadleaf deciduous tree species and separately for scenarios with (f<sub>PAW</sub>) and without (NSW) soil water deficits. The differences in reductions by O<sub>3</sub> exposure estimated by the two different O<sub>3</sub> scenario, NSW and f<sub>PAW</sub>, were very small (Table 2). Hence, only results based on the scenario f<sub>PAW</sub> will be used in the following.

The estimated reductions in GAI by O<sub>3</sub> exposure were substantial. The largest growth reductions, estimated by including f<sub>PAW</sub>, were found for deciduous tree species in Ireland (-32%), Poland (-28%) and Austria, (-28%) (Table 2). The mean reductions in GAI by O<sub>3</sub> for deciduous tree species across all countries was -17%. For coniferous tree species, the most negative values were estimated for Greece (-23%), Portugal (-19%) and Italy, (-17%). The mean reductions in GAI by O<sub>3</sub> for coniferous tree species was -7%.

**Table 2. Percent reductions in gross annual increments (GAI), caused by the present O<sub>3</sub> exposure, calculated as mean values for POD<sub>1</sub>SPEC for the period 2008-2012 and mean values for the area of different European countries. The estimated reduction of GAI was estimated based on O<sub>3</sub> dose-response relationships for impacts on growth rates (GAI DRRs) based on country-wide mean values for POD<sub>1</sub>SPEC calculated separately for coniferous and deciduous tree species and separately for scenarios with (f<sub>PAW</sub>) and without (NSW) the functions for soil water deficit included.**

The estimated total, gross annual stem volume stock changes for all forests in Europe are shown in Fig. 6, with and without the exposure to present O<sub>3</sub> doses, calculated as POD<sub>1</sub>SPEC. Also shown are the estimated total removals of forest stem volumes, including both natural and harvest removals. The rates of removals are assumed to be independent of O<sub>3</sub> regimes. Furthermore, in Fig. 6 the differences between the gross annual stock increments and the removals are indicated, separately for the presence of O<sub>3</sub> impacts (continuous line arrow) and in the absence of O<sub>3</sub> impacts (dashed line arrow).

**Figure 6. The rates of total, annual gross stem volume increment growth for forests in Europe as mean annual values for the time period 2008-2012, including the sum of coniferous and deciduous tree species, with and without the exposure to present O<sub>3</sub> doses, calculated as POD<sub>1</sub>SPEC with the scenario f<sub>PAW</sub>. Also shown are the estimated total removals of forest stem volumes, including both natural- and harvest removals. The rates of removals are assumed to be independent of O<sub>3</sub> regimes. Furthermore, the differences between the gross annual stock increments and the removals are indicated with vertical arrows, separately for the presence of O<sub>3</sub> impacts (continuous line arrow) and without the presence of O<sub>3</sub> impacts (dashed line arrow). ob, over bark.**

385 The rates of annual, gross stem volume increment of all European forests in the presence of O<sub>3</sub> were 854 M m<sup>3</sup> over bark (o.b.) yr<sup>-1</sup> and in the absence of O<sub>3</sub> 933 M m<sup>3</sup> o.b. yr<sup>-1</sup> (Fig. 6). This is an increase of 9% in the absence of O<sub>3</sub>. However, the annual net changes in forest standing stocks, NCSS, were 283 M m<sup>3</sup> o.b. yr<sup>-1</sup> in the presence of O<sub>3</sub> (continuous vertical line arrow in Fig. 6) and in the absence of O<sub>3</sub>, 363 M m<sup>3</sup> o.b. yr<sup>-1</sup> (dashed vertical line arrow in Fig. 6). This is an increase by 28% in the absence of O<sub>3</sub>. Hence, the removal of O<sub>3</sub> exposure would increase European forest stem volume growth rates by 9%, but it will increase European forest annual net changes in standing stocks by 28%. The results from this study, with modelling of negative ozone impacts on forest growth in combination with the application of forest statistics, illustrates the importance of not only considering O<sub>3</sub> impacts on forest gross growth rates but rather to consider the impacts on the gap between gross growth and the removals, i.e. net changes in forest standing stocks.

395 The annual gross stem volume increment for forests in the different European countries are shown in Fig. 7, with and without the exposure to present O<sub>3</sub> doses. Also shown are the natural and harvest removals, which are assumed to be independent of the scenarios for O<sub>3</sub> exposure. The gap between the current rates of gross growth and the total removals differs considerably in magnitude between different countries. For instance, Germany has a relatively small gap between GAI and the total (natural + harvest) removals, while France has a much larger gap. The absolute difference in GAI with and without O<sub>3</sub> exposure is relatively similar between these countries. However, this O<sub>3</sub> difference in GAI will have a much larger relative impact on the percent change in NCSS for Germany, compared to France, since the gap between current GAI and removals is smaller.

405 **Figure 7. The rates of annual gross stem volume increment growth for forests in different European countries, as annual values for the time period 2008-2012, including the sum of coniferous and deciduous tree species, with and without the exposure to present O<sub>3</sub> doses, calculated as POD<sub>1</sub>SPEC with the scenario f<sub>PAW</sub>. Also shown are the sum of the natural and harvest removals of forest stem volumes, i.e. the total removals. The rates of natural and harvest removals are assumed to be independent of ozone regimes. The countries are sorted left to right in alphabetical order.**

The change in the annual C sequestration in the forest living biomass C stock, in the absence compared to the presence of O<sub>3</sub> exposure is presented in Fig. 8. Results are presented for the individual European countries, as absolute values (Fig. 8a) and differences without and with O<sub>3</sub> (Fig. 8b), separately for coniferous and deciduous and for total forests. The absence of O<sub>3</sub> exposure would increase the C sequestration to the living biomass, in absolute values, the most in Germany, France, Italy and Poland. Summarized for all European forests, the C sequestration to the living biomass C stock was estimated to be 343 M t CO<sub>2e</sub> yr<sup>-1</sup> in the presence of O<sub>3</sub> and 449 M t CO<sub>2e</sub> yr<sup>-1</sup> in the absence of O<sub>3</sub>, i.e. an increase of 31%.

415 **Figure 8. The changes in the annual C sequestration in the living biomass C stocks, with and without the exposure to present O<sub>3</sub> doses, calculated as POD<sub>1</sub>SPEC with the scenario f<sub>PAW</sub>. (a). absolute values for total forests; (b). differences without versus with ozone, separately for coniferous, deciduous and total forests. Results are presented for the individual European countries, separately for coniferous and deciduous and for total forests. The countries are sorted left to right in alphabetical order. Removal of CO<sub>2</sub> from the atmosphere to the living biomass carbon stocks is shown with a negative sign.**

The percent changes in the annual C sequestration of forests in the different European countries, without the present O<sub>3</sub> exposure compared to with O<sub>3</sub> exposure, are shown in Fig. 9. Changes are shown separately for coniferous and deciduous forests and total forests. As an example, the percent increase in the annual C sequestration of forests in the absence of O<sub>3</sub> is much larger for Germany, compared to France. As mentioned above, this is because the annual C sequestration of forests in Germany in the presence of O<sub>3</sub> was so low that the increase due to the removal of the O<sub>3</sub> exposure will be relatively large.

As mentioned already above, the percent increase in the C sequestration to the forest living biomass C stock of all European forests in the absence of O<sub>3</sub> is 31%. In general, the impact of the absence of O<sub>3</sub> is larger for deciduous compared to coniferous forests. This is because in the O<sub>3</sub> GAI DRRs, deciduous forests are considered more sensitive to O<sub>3</sub> impacts compared to conifers. However, the estimates of the GAI of different tree functional types are somewhat uncertain since the forest statistics for different European countries only provides information on the GAI, natural- and harvest removals of the total forests together with information for the different areas of coniferous and deciduous forests.

**Figure 9. The percent changes in the annual C sequestration of forests in the different European countries, without the present O<sub>3</sub> exposure compared to with O<sub>3</sub> exposure. Changes are shown separately for coniferous and deciduous forests and for total forests.**

## **4 Discussion**

### **4.1 Spatial variation in POD<sub>1</sub>SPEC**

The UNECE Mapping Manual (LRTAP, 2017) sets the critical levels for O<sub>3</sub> impacts on trees in Europe, based on POD<sub>1</sub>SPEC, to 9.2 mmol m<sup>-2</sup> for Norway spruce, related to a 2 % annual biomass reduction, and to 5.2 mmol m<sup>-2</sup> for Beech and Birch, related to a 4 % annual biomass reduction. The results shown in this study (Fig. 3a, c) suggest that the critical level for deciduous forests was exceeded in large parts of Europe during the period 2008 - 2012, except for dry areas in the Iberian peninsula and for small parts of Continental Europe and the Fennoscandian mountain range. The exceedance of the critical levels for coniferous forests was somewhat less. This geographical distribution is very different from the exceedance of the critical levels based on the ozone exposure index AOT40 applied by the European Union (10 ppm.h), which shows a strong latitudinal gradient, with the highest values in the Mediterranean (Figure 3e and f).

The estimates of the POD<sub>1</sub>SPEC are the result of the combination of estimated O<sub>3</sub> concentrations and meteorological factors (f-functions) describing the limitation of the stomatal conductance (Emberson et al., 2000). The maps for the f-functions provide some explanations for the patterns for POD<sub>1</sub>SPEC across Europe (Fig. 4). The function used to describe the temperature limitation of stomatal conductance (Figure 4b) showed in general homogenously high values across most of Europe. The function used to describe the VPD limitation of stomatal conductance (Fig. 4c) showed a geographical variation

that more resembled the pattern for the estimated  $POD_1SPEC$ , with higher values for continental coastal regions and lower values for continental inland regions. Hence, this indicates that  $f_{VPD}$  was an important factor for the estimates of  $POD_1SPEC$  across Europe. The function used to describe the soil water deficit limitation of stomatal conductance,  $f_{PAW}$ , (Figure 4d) showed low values only for the inland Iberian Peninsula, explaining the low values for  $POD_1SPEC$  estimated for this region. The reduced geographical extent of the  $f_{PAW}$  limitation is likely due to the use of a rather insensitive  $f_{PAW}$  relationship (with limitation starting only once soil water had been reduced to a level where only 25% remains available to the plant). This rather insensitive response was selected to ensure the protective effect of soil water stress in limiting  $POD_1SPEC$  was not overestimated. Future studies could explore the effect of species-specific  $f_{PAW}$  in more detail especially since this was found to be an important variable determining  $POD_1SPEC$  and crown defoliation and visible foliar injury at specific study sites in France, Italy and Romania (Sicard et al., 2020). To give an indication of the effect of a more sensitive  $f_{PAW}$ , Figure S6 shows  $f_{PAW}$  values for each species for the year 2012 using an  $f_{PAW}$  relationship that reduces  $g_{sto}$  once 50% of available soil water is exceeded. This extends the limiting effect of soil water on  $POD_1SPEC$  further north to some parts of the Eastern continental Europe.

Southern Europe shows the highest variation of  $POD_1SPEC$  during the five years, most likely highlighting the influence of more pronounced changes in weather and hence ozone uptake conditions during those five years as compared to the rest of Europe. However, it should be noted that none of the five years targeted in this study was perceived as a meteorologically extreme, with none of the extreme European heat waves that characterised e.g. 2003 (Solberg et al., 2008, Lin et al., 2020).

The  $POD_1SPEC$  results in the present study are consistent with Anav et al (2022), who found that  $\sim 40\%$  of the forested area in the Northern Hemisphere exceeded the  $POD_1SPEC$  values of critical levels used in their study ( $5.2 \text{ mmol m}^{-2} \text{ y}^{-1}$  for boreal and continental deciduous forests,  $9.2 \text{ mmol m}^{-2} \text{ y}^{-1}$  for boreal and continental evergreen forests,  $14 \text{ mmol m}^{-2} \text{ y}^{-1}$  for temperate deciduous forests and  $47.3 \text{ mmol m}^{-2} \text{ y}^{-1}$  for temperate evergreen forests). Sicard et al (2020) found broad ranges of  $POD_1SPEC$  between 3 to  $>20 \text{ mmol O}_3 \text{ m}^{-2}$  for 15 locations in France, Italy and Romania during 2018 and 2019. A study at 30 rural sites in Switzerland found ranges in  $POD_1SPEC$  for beech of 12 to  $25 \text{ mmol O}_3 \text{ m}^{-2}$  (Braun et al., 2022) whilst our estimates are between 22 and  $28 \text{ mmol O}_3 \text{ m}^{-2}$ . For European wide studies, Simpson et al. (2022) calculated  $POD_1IAM_{DF}$  (the integrated assessment model recommendation of UNECE (LRTAP, 2017), which is largely derived from the parameterization for beech) with a more recent and higher resolution (Version rv4.45,  $0.3^\circ \text{ lon} \times 0.2^\circ \text{ lat}$ ) version of the EMEP CTM. They also found highest  $POD$  values (of around  $12\text{--}40 \text{ mmol O}_3 \text{ m}^{-2}$ ) in central Europe, Sweden and the UK, and lower values ( $0\text{--}6 \text{ mmol O}_3 \text{ m}^{-2}$ ) in many parts of Spain, rather similar to the results presented here. Finally, a recent study by Vlášáková et al. (2024) estimated  $POD_1SPEC$  for beech and Norway spruce according to the UNECE Mapping Manual functions and the CAMS ensemble forecast modelling which provided hourly  $O_3$  data. These results are consistent with our study findings, with  $POD_1SPEC$  for beech commonly ranging from 5 to  $30 \text{ mmol m}^{-2}$  in a similar spatial pattern to

that described by our broadleaf deciduous results in Figure 3c and for Norway spruce commonly ranging from 7 to 25 mmol m<sup>-2</sup> compared to our slightly lower values for coniferous forests of 5 to 18 mmol m<sup>-2</sup>.

## 4.2 The GAI dose-response relationships

490 We have developed GAI DRR so that the impacts of O<sub>3</sub> exposure on forest stand growth rates can be estimated annually and used with commonly available forest statistics to estimate annual changes in forest standing stocks and ultimately living biomass C stocks. A number of assumptions have been made in the derivation of these GAI DRRs. Firstly, we assume that O<sub>3</sub> effects on young forest trees will be equivalent to those effects on mature trees, at least in terms of the O<sub>3</sub> effect on growth rates. This is further discussed in section 4.4. Secondly, we assume that we can reliably estimate the growth rate for different  
495 tree species using the Richards equation [Eq. 5] (Richards, 1959). Given that the Richards equation is used to interpolate growth rates between very young trees (when biomass will be close to zero) and known biomass of trees at the end of a 2-to-10-year growth cycle, the uncertainties in this assumption are very small.

## 4.3. Impacts of ozone exposure on the forest living biomass C stock changes

Estimates of O<sub>3</sub> impacts on forest C sequestration for the different European countries, and for Europe in total, were in general  
500 much larger when based on the difference between GAI and total removals, compared with estimates based on ozone impacts on GAI itself. The removal of O<sub>3</sub> exposure was estimated to increase European forest stem volume growth rates by 9%, but it was estimated to increase European forest living biomass C stock changes by 31%, i.e. more than three times as much. This illustrated the importance of not only considering ozone impacts on forest growth rates but also to consider the impacts on the gap between gross growth rates and forest removals, i.e. the net changes in the forest standing stocks. Previous studies of the  
505 impacts of O<sub>3</sub> on vegetation C sequestration have in most cases focussed on the impacts of O<sub>3</sub> on photosynthesis (Felzer et al., 2005; Felzer et al., 2009; Sitch et al., 2007; Ren et al., 2007). However, direct impacts of environmental changes on C-sinks involved in the growth process has been shown to be more important for tree growth than indirect control via impacts on C-sources, such as photosynthesis (Millard et al., 2007; Fatichi et al., 2014; Körner, 2015; Eckes-Shephard et al., 2020; Kannenberg et al., 2022). This would suggest that approaches that estimate the negative impacts of O<sub>3</sub> on vegetation C  
510 sequestration only via impacts on instantaneous photosynthesis may not be correct and hence care should be taken when making estimates of ozone effects on forest growth using damage functions that only influence via photosynthetic based carbon assimilation (e.g. Sorrentino et al., 2025). The dose-response relationships used in our study were based on O<sub>3</sub> exposure experiments with the long-term biomass growth as the response parameter, thus mainly reflecting O<sub>3</sub> impacts on the C-sinks.

## 4.4. The applications for forest stands

515 The phytotoxicity of O<sub>3</sub> for young trees under experimental conditions is well documented (Karlsson et al., 2007; Wittig et al., 2009), but negative impacts of ozone on mature trees under field conditions have been more difficult to establish (Kolb and

Matyssek, 2001; Marzuoli et al., 2020). The most evident and well described case of negative ozone impacts on forest ecosystems is the ozone induced decline in the San Bernadino Forest in California (Miller and McBride, 1999). Negative impacts on forest growth were analysed mainly based on gradient studies (Miller et al., 1997; Arbaugh et al., 1999). It was estimated that the basal area increment growth rates of the Douglas fir forest were reduced by approximately 30%, when comparing the areas with daylight mean O<sub>3</sub> concentration of 64-67 and 76-83 ppb, respectively. These are high concentration values compared to the corresponding values that can be estimated in Europe today (Gaudel et al., 2018). On the other hand, the climate in San Bernadino Forest is a Mediterranean dry climate which might be expected to limit O<sub>3</sub> uptake.

The impacts of O<sub>3</sub> on the growth and vitality of different tree species have been studied extensively in Switzerland (Braun et al., 2014; 2017; 2022). These studies include both experimental exposure studies with young trees and epidemiological field studies where the stem growth and other vitality parameters have been correlated with environmental factors, including O<sub>3</sub> exposure. The agreement of the dose–response curve from the epidemiological studies with the experimental data with young Beech and Norway spruce trees suggests that the current dose–response curves for O<sub>3</sub> used within the LRTAP convention are also valid for mature forests (Braun et al., 2022). In a unique case study, a free-air ozone fumigation experiment was conducted at “Kranzberger Forst” (Freising, southern Germany), over 8 years on adult trees of European beech and Norway spruce. Pretzsch and Schütze (2018) concluded that in the twice ambient O<sub>3</sub> exposure the annual basal area growth of European Beech and Norway spruce decreased by 32 and 24%, respectively. Free air fumigation of different deciduous tree species in the Aspen-Face project (Karnosky et al., 2005; Talhem et al., 2014) demonstrated a negative impact of O<sub>3</sub> on the ecosystem C content that, however, had a transient development over time. Experimental studies in China (Feng et al., 2019) with poplar tree species demonstrated a consistent negative relationship between biomass accumulation and O<sub>3</sub> exposure both of small trees in open-top chambers and larger trees under chamber-less exposure. Oksanen et al. (2009) summarized the results from Finnish experiments with O<sub>3</sub> exposure of several different clones of Birch and concluded that an O<sub>3</sub> exposure of 15 ppm hours AOT40 may reduce the stem diameter growth of Birch by approximately 15 %. This is of the same magnitude as the estimates of AOT40 described for large parts of Europe (Figure 3F). Thus, based on literature results described above, mature trees under field conditions cannot be assumed to be less sensitive to ozone exposure compared to young trees under experimental conditions (Karlsson et al., 2024). Consequently, we suggest that our estimates of negative impacts of O<sub>3</sub> on forest growth rates and C stock changes, using DRRs from 2- to 10-year-old forest trees under experimental conditions, may provide reliable results for the different forests of variable age that exist across Europe.

**4.5. Assumptions**

The estimates of the impacts of O<sub>3</sub> exposure on the C sequestration of the managed forests were based on some additional assumptions. The first assumption was that the presence of O<sub>3</sub> did not affect the natural losses caused by storm fellings, drought, insect attacks etc. Free-air fumigation to elevated concentrations of O<sub>3</sub> were shown to increase aphid infestations in trembling

550 aspen, *Populus tremuloides* (Percy et al., 2003). If absence of O<sub>3</sub> exposure would reduce the risk for insect attacks or other diseases in trees, that would contribute to increase the C sequestration in European managed forests even further, since the gap between gross growth and removals would increase. Ozone has been shown to reduce the relative growth of below ground biomass (Gu et al., 2022). Reduced root growth can be expected to make trees more vulnerable to drought. Hence, the absence of O<sub>3</sub> exposure would have the potential to increase the C sequestration also from this aspect.

555 The second assumption was that harvest rates would not be affected by the changes in gross growth rates, i.e. that harvest rates are mainly dependent on the demand for forest raw materials. In countries that are heavily dependent on the supply of forest raw material to the forest industry, such as e.g. Sweden and Finland, the rates of harvests are carefully monitored by national forest inventories. For example, the Swedish Forest Agency regularly make recommendations about the rates of harvest in relation to the rates of forest gross growth, to make sure that harvest rates do not exceed growth rates. Levers et al. (2014)  
560 concluded that the harvest intensities of European forests were mainly explained by the share of plantation species, by terrain ruggedness, and by different country-specific characteristics. Forest growth rates were not considered as important for explaining forest harvest intensities in that study. Another important factor regulating the harvest rates is the forest age structure, i.e. when forests reach the economically optimal age for harvesting (Korosuo et al., 2023). An alternative assumption for the calculations would have been to keep harvest rates as a certain fraction of the growth rates. In this case increased growth  
565 rates in the absence of O<sub>3</sub> exposure would have been accompanied by a certain increase in harvest rates, that in absolute terms would have been lower than the increase in growth. Hence, also in this case, the absence of O<sub>3</sub> exposure would have resulted in increased C sequestration in European managed forests, but it would have been smaller than what has been presented in the current calculations in our study.

#### 4.6. Comparisons with other estimates of and policy for European forest carbon (C) sequestration

570 In the present study, the yearly C sequestration to the European forests living biomass C stocks, as a mean value 2008 – 2012, was estimated to be -343 M t CO<sub>2e</sub> yr<sup>-1</sup> in the presence of O<sub>3</sub>. Regarding the uncertainties and simplification of the calculations, this value is in the same order of magnitude as the value estimated in the Forest Europe assessment (Forest Europe, 2020), that the average annual sequestration of C in European forest biomass between 2010 and 2020 was around -560 M t CO<sub>2e</sub> yr<sup>-1</sup>. Pan et al (2011) estimated the yearly C sequestration to European forests, excluding Russia, to -570 M t CO<sub>2e</sub> for the period 2002-  
575 2007.

The European Union European Commission has issued an action plan to develop sustainable solutions to increase C sequestration by forests. The EU target is to increase the yearly uptake of CO<sub>2</sub> in the entire AFOLU sector (Agriculture, Forestry and Other Land Use) to -310 M t CO<sub>2e</sub> yr<sup>-1</sup> (Korosuo et al., 2023). It was estimated in the Forest Europe assessment  
580 (Forest Europe, 2020) that living woody biomass represents around 36% of the total C stocks in forests. Hence, the increase

in the yearly C sequestration to the European forests living biomass C stocks estimated in this study, by eliminating negative impacts of O<sub>3</sub> on forest gross growth rates, from -343 to -449 M t CO<sub>2e</sub> yr<sup>-1</sup>, clearly has the potential to substantially contribute to the European Commission action plan. The yearly C sequestration to the European forests is expected to be short of the so called “forest reference levels”, i.e. the forest C sequestration levels agreed by the EU member states during 2021–2025 (Hyrynen et al., 2023; Korosuo et al., 2023).

The estimates made in this study included only the O<sub>3</sub> influence of changes in the forest living biomass C stocks. However, there may be additional climate change abatement benefits caused by mitigating ozone concentrations, such as enhancing C held in the dead biomass and soil C stocks (Liski et al., 2002; Mäkipää et al., 2023). Both these C stock changes may be promoted by the increase litter production due to the increase in forest gross growth rates. Furthermore, C can be stored for some time in harvested wood products (HWP, Hyrynen et al., 2023) increasing C stocks. Finally, the forest raw materials produced may substitute the use of fossil-based raw materials (Leskinen et al., 2018; Korosuo et al., 2023).

**5 Conclusions**

- The annual, species-specific Phytotoxic Ozone Dose above a threshold of 1.0 nmol m<sup>-2</sup> s<sup>-1</sup>, POD<sub>1</sub>SPEC, was estimated for European forests plant functional types for the years 2008 – 2012.
- The critical level for negative impacts on forests suggested by the UN Air Convention, based on POD<sub>1</sub>SPEC, was exceeded in large parts of Europe.
- The highest POD<sub>1</sub>SPEC was estimated for the coastal regions of mid-latitude Europe including UK, for both coniferous and broadleaf forests.
- The annual forest standing stock changes were estimated based on POD<sub>1</sub>SPEC -based dose-response relationships for negative impacts of ozone on forest growth rates in combination with the estimated values for POD<sub>1</sub>SPEC and official forest statistics for forest growth and total removals. With the use of default IPCC methodology, this could be converted to estimates of changes in the forest living biomass C stocks.
- The absence of ozone exposure would increase European forest stem volume growth rates by 9%, but it would increase European forest living biomass C stocks increment rates by 31%.
- Mature trees under field conditions cannot be assumed to be less sensitive to ozone exposure compared to young trees under experimental conditions.

## 6 Author contribution

PEK, PB and LE designed the study. DS and SB performed the simulations. KS and FH provided important information. PEK,  
610 PB and LE prepared the manuscript with contributions from all co-authors.

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## 8 Data availability

Data will be available upon request.

## 9 Competing interests

625 The authors declare that they have no conflict of interest.

## 10 References

- Ågren, K., Högbom, L., Johansson, M., och Wilhelmsson, L.: Datainsamling till underlag för livscykelanalyser (LCA) av det svenska skogsbruket. Skogforsk ARBETSRAPPORT 1086–2021. ISSN 1404-305X. (In Swedish), 2021.
- 630 Anav, A., De Marco, A., Collalti, A., Emberson, L., Feng, Z., Lombardozzi, D., Sicard, P., Verbeke, T., Viovy, N., Vitale, M. and Paoletti, E.: Legislative and functional aspects of different metrics used for ozone risk assessment to forests. *Env. Poll.*, 295, 118690, <https://doi.org/10.1016/j.envpol.2021.118690>, 2022.

- Arbaugh, M.J., Peterson, D.L., Miller, P.J.: Air pollution effects on growth of Ponderosa pine, Jeffrey pine and Bigcone Douglas-Fir. In Miller, P.R & McBride, J.R., editors. Ecological Studies 134. Springer. ISBN 0-387-94493-3. 1999.
- 635 Büker, P., Morrissey, T., Briolat, A., Falk, R., Simpson, D., Tuovinen, J.-P., ... Emberson, L. D. (2012). DO3SE modelling of soil moisture to determine ozone flux to forest trees. *Atm. Chem. and Phys.*, 12(12), 5537–5562. <https://doi.org/10.5194/acp-12-5537-2012>, 2012.
- Büker, P., Feng, Z., Uddling, J., Briolat, a., Alonso, R., Braun, S., ... Emberson, L. D.: New flux based dose–response relationships for ozone for European forest tree species. *Env. Poll.*, 206, 163–174.
- 640 <https://doi.org/10.1016/j.envpol.2015.06.033>, 2015.
- Braun, S., Schindler, C., Rihm, B.: Growth losses in Swiss forests caused by ozone: Epidemiological data analysis of stem increment of *Fagus sylvatica* L. and *Picea abies* Karst. *Env. Poll.*, 192, 129–138. <https://doi.org/10.1016/j.envpol.2014.05.016>, 2014.
- Braun, S., Rihm, B., Schindler, C. : Growth trends of beech and Norway spruce in Switzerland: The role of nitrogen deposition, ozone, mineral nutrition and climate. *Sci. of the Tot. Env.*, 599–600, 637–646. <https://doi.org/10.1016/j.scitotenv.2017.04.230>,
- 645 2017.
- Braun, S.; Rihm, B.; Schindler, C.: Epidemiological Estimate of Growth Reduction by Ozone in *Fagus sylvatica* L. and *Picea abies* Karst.: Sensitivity Analysis and Comparison with Experimental Results. *Plants*, 11, 777. <https://doi.org/10.3390/plants11060777>, 2022.
- 650 Calliari, E., Castellari, S., Davis, M., Linnerooth-Bayer, J., Martin, J., Mysiak, J., Pastor, T., Ramieri, E., Scolobig, A., Sterk, M., Veerkamp, C.: Building climate resilience through nature-based solutions in Europe: A review of enabling knowledge, finance and governance frameworks. *Clim. Risk Man.*, 1, 37–100450. <https://doi.org/10.1016/j.crm.2022.100450>, 2022
- Cinderby, S., Emberson, L., Owen, A., & Ashmore, M.: LRTAP land cover map of Europe. In CCE Progress Report. Retrieved from [http://www.rivm.nl/bibliotheek/digitaaldepot/PBL\\_CCE\\_PR07PartI\\_5.pdf](http://www.rivm.nl/bibliotheek/digitaaldepot/PBL_CCE_PR07PartI_5.pdf), 2007.
- 655 Eckes-Shephard, A. H., Tiavlovsky, E., Chen, Y., Fonti, P., Friend, A.D.: Direct response of tree growth to soil water and its implications for terrestrial carbon cycle modelling. *Glob. Change Biol.* 27, 121–135. <https://doi.org/10.1111/gcb.15397>, 2021
- EEA.: Biogeographical Regions in Europe, <https://www.eea.europa.eu/en/analysis/maps-and-charts/biogeographical-regions-in-europe-2>, 2016.
- Emberson, L. D., Ashmore, M. R., Cambridge, H. M., Simpson, D., & Tuovinen, J. P.: Modelling stomatal ozone flux across
- 660 Europe, *Env. Poll.*, 109(3), 403–13. [https://doi.org/10.1016/S0269-7491\(00\)00043-9](https://doi.org/10.1016/S0269-7491(00)00043-9), 2000.
- Emberson, L. D., Büker, P., & Ashmore, M. R.: Assessing the risk caused by ground level ozone to European forest trees: a case study in pine, beech and oak across different climate regions, *Env. Poll.*, 147(3), 454–66. <https://doi.org/10.1016/j.envpol.2006.10.026>, 2007.
- Emberson, L., 2020. Effects of ozone on agriculture, forests and grasslands. *Phil. Trans. of the Roy. Soc. A*, 378(2183),
- 665 p.20190327, <https://doi.org/10.1098/rsta.2019.0327>, 2020.

- Etzold, S., Ferretti, M., Reinds, G. J., Solberg, S., Gessler, A., Waldner, P., Schaub, M., Simpson, D., Benham, S., Hansen, K., Ingerslev, M., Jonard, M., Karlsson, P.E., Lindroos, A.-J., Marchetto, A., Manninger, M., Meesenburg, H., Merilä, P., Nöjd, P., Rautio, P., Sanders, T.G.M., Seidling, W., Skudnik, M., Thimonier, A., Verstraeten, A., Vesterdal, L., Vejpustkova, M., de Vries, W.: Continental-scale forest growth in Europe is driven by management and further modulated by nitrogen deposition. *Forest Ecology and Management*, 458, 117762, <https://doi.org/10.1016/j.foreco.2019.117762>, 2020.
- Fatichi, S., Sebastian Leuzinger and Körner, C.: Moving beyond photosynthesis: from carbon source to sink-driven vegetation modelling. *New Phyt.*, 201, 1086–1095, <https://www.jstor.org/stable/newphytologist.201.4.1086>, 2014
- Fellner, J., & Rechberger, H.: Abundance of <sup>14</sup>C in biomass fractions of wastes and solid recovered fuels. *Waste Management*, 29(5), 1495–1503, <https://doi.org/10.1016/j.wasman.2008.11.023>, 2009.
- Felzer, B., Reilly, J., Melillo, J., Kicklighter, D., Sarofim, M., Wang, C., Prinn, R. and Zhuang, Q.: Future Effects of Ozone on Carbon Sequestration and Climate Change Policy Using a Global Biogeochemical Model. *Climatic Change*, 73(3), 345–373. <https://doi.org/10.1007/s10584-005-6776-4>, 2005
- Felzer, B. S., Cronin, T. W., Melillo, J. M., Kicklighter, D. W., & Schlosser, C. A.: Importance of carbon-nitrogen interactions and ozone on ecosystem hydrology during the 21st century. *Journal of Geophysical Research*, 114(G1), 1–10. <https://doi.org/10.1029/2008JG000826>, 2009.
- Feng, Z., Shang, B., Gao, F., Calatayud, V.: Current ambient and elevated ozone effects on poplar: A global meta-analysis and response relationships. *Sci. of the Tot. Env.* 654, 832-840. <https://doi.org/10.1016/j.scitotenv.2018.11.179>, 2019.
- Forest Europe.: State of Europe's Forests 2020. Prepared and published by: Ministerial Conference on the Protection of Forests in Europe - FOREST EUROPE. [www.foresteurope.org](http://www.foresteurope.org), 2020.
- Franz, M., Simpson, D., Arneth, A., and Zaehle, S.: Development and evaluation of an ozone deposition scheme for coupling to a terrestrial biosphere model, *Biogeosciences*, 14, 45–71, <https://doi.org/10.5194/bg-14-45-2017>, 2017.
- Franz, M., Alonso, R., Arneth, A., Büker, P., Elvira, S., Gerosa, G., Emberson, L., Feng, Z., Le Thiec, D., Marzuoli, R. and Oksanen, E.: Evaluation of simulated ozone effects in forest ecosystems against biomass damage estimates from fumigation experiments. *Biogeosciences*, 15(22), 6941-6957, <https://doi.org/10.5194/bg-15-6941-2018>, 2018.
- Fuhrer, J.: The critical level for ozone to protect agricultural crops – An assessment of data from European open-top chamber experiments. In: Fuhrer J, Achermann B, editors. Critical levels for ozone. A UN-ECE workshop report. Schriftenreihe der FAC Liebfeld 16, pp. 42-57, 1994.
- Gaudel, A., Cooper, O.R., Ancellet, G., Barret, B., Boynard, A., Burrows, J.P., Clerbaux, C., Coheur, P.F., Cuesta, J., Cuevas, E., Doniki, S.: Tropospheric Ozone Assessment Report: Present-day distribution and trends of tropospheric ozone relevant to climate and global atmospheric chemistry model evaluation. *Elem. Sci. Anth.*, 6, 39, <https://doi.org/10.1525/elementa.291>, 2018.

- Griscom, B.W., Adams, J., Ellis, P.W., Houghton, R.A., Lomax, G., Miteva, D.A., Schlesinger, W.H., Shoch, D., Siikamäki, J.V., Smith, P. and Woodbury, P.: Natural climate solutions. *Proc. Nat. Aca. Sci.*, 114, 11645–11650. [www.pnas.org/cgi/doi/10.1073/pnas.1710465114](http://www.pnas.org/cgi/doi/10.1073/pnas.1710465114), 2017.
- 700 Gu, X., Wang, T., & Li, C.: Elevated ozone decreases the multifunctionality of belowground ecosystems. *Global Change Biol.*, 29(3), 890-908. <https://doi.org/10.1111/gcb.16507>, (2023).
- Gustavsson, L., Haus, S., Lundblad, M. Lundström, A., Ortiz, C.A., Sathre, R., LeTruong, N., Wikberg, P.-E.: Climate change effects of forestry and substitution of carbon-intensive materials and fossil fuels. *Ren. and Sust. Energy Rev.*, 67, 612–624. <http://dx.doi.org/10.1016/j.rser.2016.09.056>, 2017.
- 705 Hyyrynen, M., Ollikainen, M., Seppälä, J.: European forest sinks and climate targets: past trends, main drivers, and future forecasts. *Eur. J. of For. Res.*, 142, 1207–1224. <https://doi.org/10.1007/s10342-023-01587-4>, 2023.
- Janssens, I.A., Freibauer, A., Ciais, P., Smith, P., Nabuurs, G.J., Folberth, G., Schlamadinger, B., Hutjes, R.W., Ceulemans, R., Schulze, E.D. and Valentini, R.: Europe's terrestrial biosphere absorbs 7 to 12% of European anthropogenic CO<sub>2</sub> emissions, *Science*, 300, 1538–1542, <https://www.science.org/doi/10.1126/science.1083592>, 2003.
- 710 Jasinevičius, G., Lindner, M., Pingoud K., Tykkyläinen, M.: Review of models for carbon accounting in harvested wood products, *Int. Wood Prod. J.*, 6(4), 198-212, <https://doi.org/10.1080/20426445.2015.1104078>, 2015.
- Kannenber, S.A., Cabon, A., Babst, F., Belmecheri, S., Delpierre, N., Guerrieri, R., Maxwell, J.T., Meinzer, F.C., Moore, D.J.P., Pappas, C., Ueyama, M., Ulrich, D.E.M., Voelker, S.L., Woodruff, D.R., Anderegg, W.R.L.: Drought-induced decoupling between carbon uptake and tree growth impacts forest carbon turnover time. *Agric. and For. Met.*, 322, 108996, <https://doi.org/10.1016/j.agrformet.2022.108996>, 2022.
- 715 Karlsson, P.E., Pleijel, H., Belhaj, M., Danielsson, H., Dahlin, B., Andersson, M., Hansson, M., Munthe, J., Grennfelt, P.: Economic assessment of the negative impacts of ozone on the crop yield and forest production. A case study of the Estate Östads Säteri in southwestern Sweden, *Ambio*, 34, 32-40, <https://doi.org/10.1579/0044-7447-34.1.32>, 2005.
- Karlsson, P.E., Braun, S., Broadmeadow, M., Elvira, S., Emberson, L., Gimeno, B.S., Le Thiec, D., Novak, K., Oksanen, E.,
- 720 Schaub, M., Uddling, J., Wilkinson, M.: Risk assessments for forest trees - the performance of the ozone flux versus the AOT concepts. *Env. Poll.*, 146, 608–616. <https://doi.org/10.1016/j.envpol.2006.06.012>, 2007.
- Karlsson, P.E.: Ozone Impacts on Carbon Sequestration in Northern and Central European Forests. IVL Rapport B 2065, 2012.
- Karlsson, P.E., Danielsson, H., Pleijel, H., Andersson, C.: Exceedance of critical levels for ozone impacts on Swedish forests - Evaluation of methodology for POD1SPEC calculations. IVL Report C 829, 2024.
- 725 Karnosky, D. F., Pregitzer, K.S., Zak, D.R., Kubiske, M.E., Hendrey, G.R., Weinstein, D., Nosal, M., Percy, K.E.: Scaling ozone responses of forest trees to the ecosystem level in a changing climate, *Plant, Cell & Env.* 28, 8, 965-981, <https://doi.org/10.1111/j.1365-3040.2005.01362.x>, 2005.
- Kolb, T. E., and R. Matyssek.: Limitations and perspectives about scaling ozone impacts in trees, *Env. Poll.*, 115(3), 373-393, [https://doi.org/10.1016/S0269-7491\(01\)00228-7](https://doi.org/10.1016/S0269-7491(01)00228-7), 2001.

- 730 Korosuo, A., Pilli, R., Abad Viñas, R., Blujdea, V., Colditz, R., Fiorese, G., Rossi, S., Vizzarri, M., Grassi, G.: The role of forests in the EU climate policy: are we on the right track? *Carbon Balance and Management*, 18(15), <https://doi.org/10.1186/s13021-023-00234-0>, 2023.
- Körner, C.: Paradigm shift in plant growth control. *Current Opinion in Plant Biology* 25, 107–114. <http://dx.doi.org/10.1016/j.pbi.2015.05.003>, 2015.
- 735 Leskinen, P., Cardellini, G., González-García, S., Hurmekoski, E., Sathre, R., Seppälä, J., Johannes Verkerk, P.: Substitution Effects of Wood-based Products in Climate Change Mitigation. From Science to Policy 7. European Forest Institute, 2018.
- Levers, C., Verkerk, P.J., Müller, D., Verburg, P.H., Butsic, V., Leitão, P.J., Lindner, M., Kuemmerle, T.: Drivers of forest harvesting intensity patterns in Europe. *Forest Ecology and Management*, 315, 160–172, <http://dx.doi.org/10.1016/j.foreco.2013.12.030>, 2014.
- 740 Lin, M., Horowitz, L. W., Xie, Y., Paulot, F., Malyshev, S., Shevliakova, E., Finco, A., Gerosa, G., Kubistin, D., and Pilegaard, K.: Vegetation feedbacks during drought exacerbate ozone air pollution extremes in Europe, *Nature Climate Change*, 1758–6798, <https://doi.org/10.1038/s41558-020-0743-y>, 2020.
- Liski, J. Daniel Perruchoud, Timo Karjalainen.: Increasing carbon stocks in the forest soils of western Europe. *Forest Ecology and Management*, 169, 159–175. [https://doi.org/10.1016/S0378-1127\(02\)00306-7](https://doi.org/10.1016/S0378-1127(02)00306-7), 2002.
- 745 LRTAP.: Manual on methodologies and criteria for modelling and mapping Critical Loads and Levels and air pollution effects, risks and trends. Chapter 3: Mapping critical levels for vegetation. Retrieved from <http://icpvegetation.ceh.ac.uk>., 2017.
- Marzuoli, R., Gerosa, G., Bussotti, F. and Pollastrini, M.: Assessing the impact of ozone on forest trees in an integrative perspective: Are foliar visible symptoms suitable predictors for growth reduction? A critical review. *Forests*, 10(12), 1144. <https://doi.org/10.3390/f10121144>, 2019
- 750 Matthews, B., Mareckova, K., Schindlbacher, S., Ullrich, B., and Wankmüller, R.: Emissions for 2018, in: Transboundary particulate matter, photo-oxidants, acidifying and eutrophying components. EMEP Status Report 1/2020, The Norwegian Meteorological Institute, Oslo, Norway, 37–57, <http://emep.int>, 2020.
- Millard, P., Sommerkorn, M. Grelet, G.-A.: Environmental change and carbon limitation in trees: a biochemical, ecophysiological and ecosystem appraisal. *New Phyt.*, 175, 11–28, <https://doi.org/10.1111/j.1469-8137.2007.02079.x>, 2007.
- 755 Miller, P.J., Arbaugh, M.J., Temple, P.J.: Ozone and its known potential effects on forests in western United States. In Sandermann, H., Wellburn, A.R., Heath, R.L., editors. *Ecological Studies* 127. Springer. ISBN 3-540-61321-8, 1997.
- Miller, P.J. and McBride, J.R.: Introduction. In *Oxidant air pollution impacts in the montane forests of southern California*. Miller, P.R & McBride, J.R., editors. *Ecological Studies* 134, Springer, ISBN 0-387-94493-3, 1999.
- Mills, G., Pleijel, H., Braun, S., Büker, P., Bermejo, V., Calvo, E., Danielsson, H., Emberson, L., Grünhage, L., González
- 760 Fernández, I., Harmens, H., Hayes, F., Karlsson, P.E., Simpson, D.: New stomatal flux based critical levels for ozone effects on vegetation. *Atmos. Env.*, 45, 5064–5068, <https://doi.org/10.1016/j.atmosenv.2011.06.009>, 2011.

- Mäkipää, R., Abramoff, R., Adamczyk, B., Baldy, V., Biryol, C., Bosela, M., Casals, P., Yuste, J.C., Dondini, M., Filipek, S., Garcia-Pausas, J., Gros, R., Gomoryov, E., Hashimoto, S., Hassegawa, M., Immonen, P., Laiho, R., Li, H., Li, Q., Luyssaert, S., Menival, C., Mori, T., Naudts, K., Santonja, M., Smolander, A., Toriyama, J., Tupek, B., Ubeda, X., Verkerk, P.J.,  
765 Lehtonen, A.: How does management affect soil C sequestration and greenhouse gas fluxes in boreal and temperate forests? – A review. *Forest Ecology and Management* 529, 120637, <https://doi.org/10.1016/j.foreco.2022.120637>, 2023
- Nabuurs, G.J., Paivinen, R., Sikkema, R., Mohren, G. M.: The role of European forests in the global carbon cycle - a review, 13(6), 345–358. [https://doi.org/10.1016/S0961-9534\(97\)00036-6](https://doi.org/10.1016/S0961-9534(97)00036-6), 1998.
- Oksanen, E., Manninen, S., Vapaavuori, E. and Holopainen, T.: Near-ambient ozone concentrations reduce the vigor of *Betula*  
770 and *Populus* species in Finland, *Ambio*, 38, 413–417. <https://doi.org/10.1579/0044-7447-38.8.413>, 2009.
- Otu-Larbi, F., Conte, A., Fares, S., Wild, O. and Ashworth, K.: Current and future impacts of drought and ozone stress on Northern Hemisphere forests. *Global Change Biology*, 26(11), 6218–6234, <https://doi.org/10.1111/gcb.15339>, 2020.
- Pan, Y., Birdsey, R., Fang, J., Houghton, R., Kauppi, P. E., Kurz, W., Hayes, D.: A large and persistent carbon sink in the world’s forests. *Science*, 333(6045), 988–993. <https://doi.org/10.1126/science.1201609>, 2011.
- 775 Penman, J., Gytarsky, M., Hiraishi, T., Kryg, T., Kruger, D., Pipatti, R., Buendia, L., Miwa, K., Ngara, T., Tanabe, K., Wagner, F. Good Practice Guidance for Land Use, Land-Use Change and Forestry. Published by the Institute for Global Environmental Strategies (IGES) for the IPCC. ISBN 4-88788-003-0, 2003.
- Percy, K.E., Caroline S., Awmack, R.L., Kubiske, M.E., Kopper, B.J., Isebrands, J.G., Pregitzer, K.S., Hendrey, G.R., Dickson, R.E., Zak, D.R., Oksanen, E., Sober, J., Harrington, R., Karnosky, D.F.: Altered performance of forest pests under atmospheres  
780 enriched by CO<sub>2</sub> and O<sub>3</sub>. *Nature*, 420, 403 – 407, <https://doi.org/10.1038/nature01028>, 2003.
- Pretzsch, H., Schütze, G.: Growth recovery of mature Norway spruce and European beech from chronic O<sub>3</sub> stress. *European J. of For. Res.*, 137, 251–263. <https://doi.org/10.1007/s10342-018-1106-3>, 2018.
- Ren, W., Tian, H., Chen, G., Liu, M., Zhang, C., Chappelka, A. H., & Pan, S. Influence of ozone pollution and climate variability on net primary productivity and carbon storage in China’s grassland ecosystems from 1961 to 2000. *Env. Poll.*,  
785 149(3), 327–35. <https://doi.org/10.1016/j.envpol.2007.05.029>, 2007.
- Richards, F. J.: A flexible growth constant for forests. *J. of Exp. Botany*, 10(39), 290–300. 1959.
- Roe, S., Streck, C., Beach, R., Busch, J., Chapman, M., Daioglou, V., Deppermann, A., Doelman, J., Emmet-Booth, J., Engelmann, J., Fricko, O., Frischmann, C., Funk, J., Grassi, G., Griscom, B., Havlik, P., Hanssen, S., Humpenöder, F., Landholm, D., Lomax, G., Lehmann, J., Mesnildrey, L., Nabuurs, G.-J., Popp, A., Rivard, R., Sanderman, J., Sohngen, B.:  
790 Land-based measures to mitigate climate change: Potential and feasibility by country. *Glob. Change Biol.*, 27, 6025–6058, <https://doi.org/10.1111/gcb.15873>, 2021.
- Sathre, R., Gustavsson, L.: Using wood products to mitigate climate change: External costs and structural change. *Applied Energy*, 86, 251–257, <https://doi.org/10.1016/j.apenergy.2008.04.007>, 2009.

Simpson, D., Ashmore, M. R., Emberson, L., & Tuovinen, J. P. A comparison of two different approaches for mapping potential ozone damage to vegetation. A model study, *Env. Poll.*, 146(3), 715-725, <https://doi.org/10.1016/j.envpol.2006.04.013>, 2007.

Simpson, D., Benedictow, A., Berge, H., Bergström, R., Emberson, L. D., Fagerli, H., Flechard, C. R., Hayman, G. D., Gauss, M., Jonson, J. E., Jenkin, M. E., Nyíri, A., Richter, C., Semeena, V. S., Tsyro, S., Tuovinen, J.-P., Valdebenito, Á., and Wind, P.: The EMEP MSC-W chemical transport model – technical description, *Atmos. Chem. Phys.*, 12, 7825–7865, <https://doi.org/10.5194/acp-12-7825-2012>, 2012.

Simpson, D., Gonzalez Fernandez, I. A., Segers, A., Tsyro, S., Valdebento, A., and Wind, P.: Updates to the EMEP/MSC-W model, 2021–2022, in: Transboundary particulate matter, photo-oxidants, acidifying and eutrophying components. EMEP Status Report 1/2022, The Norwegian Meteorological Institute, Oslo, Norway, 133–146, 2022.

Sicard, P., De Marco, A., Carrari, E., Dalstein-Richier, L., Hoshika, Y., Badea, O., Pitar, D., Fares, S., Conte, A., Popa, I. and Paoletti, E.: Epidemiological derivation of flux-based critical levels for visible ozone injury in European forests. *J. For. Res.* 31, 1509–1519, <https://doi.org/10.1007/s11676-020-01191-x>, 2020.

Sitch, S., Cox, P. M., Collins, W. J., & Huntingford, C.: Indirect radiative forcing of climate change through ozone effects on the land-carbon sink. *Nature*, 448(7155), 791–4. <https://doi.org/10.1038/nature06059>, 2007

Soimakallio, S., Kalliokoski, T., Lehtonen, A., Salminen, O.: On the trade-offs and synergies between forest carbon. *Mitigation and Adaptation Strategies for Global Change* 26, 4, <https://doi.org/10.1007/s11027-021-09942-9>, 2021.

Solberg, S., Hov, O., Sovde, A., Isaksen, I. S. A., Coddeville, P., De Backer, H., Forster, C., Orsolini, Y., and Uhse, K.: European surface ozone in the extreme summer 2003, *J. Geophys. Res.*, 113, D07307, <https://doi.org/10.1029/2007JD009098>, 2008.

Sorrentino, B., Anav, A., Calatayud, V., Collalti, A., Sicard, P., Leca, S., Fornasier, F., Paoletti, E. and De Marco, A.: Inconsistency between process-based model and dose-response function in estimating Biomass losses in Northern Hemisphere due to elevated O<sub>3</sub>. *Env. Poll.*, 125379, <https://doi.org/10.1016/j.envpol.2024.125379>, 2024.

Stull, R. B. An introduction to Atmospheric Boundary Layer Meteorology, Kluwer Academic Publishers, Dordrecht, 1988.

Subramanian, N., Karlsson, P.E., Bergh, J. and Nilsson, U., Impact of ozone on sequestration of carbon by Swedish Forests under a changing climate: a modeling study. *For. Sci.*, 61(3), 445-457, <https://doi.org/10.5849/forsci.14-026>, 2015.

alhelm, A.F., Pregitzer, K.S., Kubiske, M.E., Zak, D.R., Company, C.E., Burton, A.J., Dickson, R.E., Hendrey, G.R., Isebrands, J.G., Lewin, K.F. and Nagy, J.: Elevated carbon dioxide and ozone alter productivity and ecosystem carbon content in northern temperate forests. *Global Change Biology*, 20, 2492–2504. <https://doi.org/10.1111/gcb.12564>, 2014.

Tuovinen, J. P., & Simpson, D.: An aerodynamic correction for the European ozone risk assessment methodology. *Atmospheric Environment*, 42(36), 8371-8381, <https://doi.org/10.1016/j.atmosenv.2008.08.008>, 2008.

UNECE: <https://fra-data.fao.org/assessments/panEuropean/2020/FE/home/overview>, 2011.

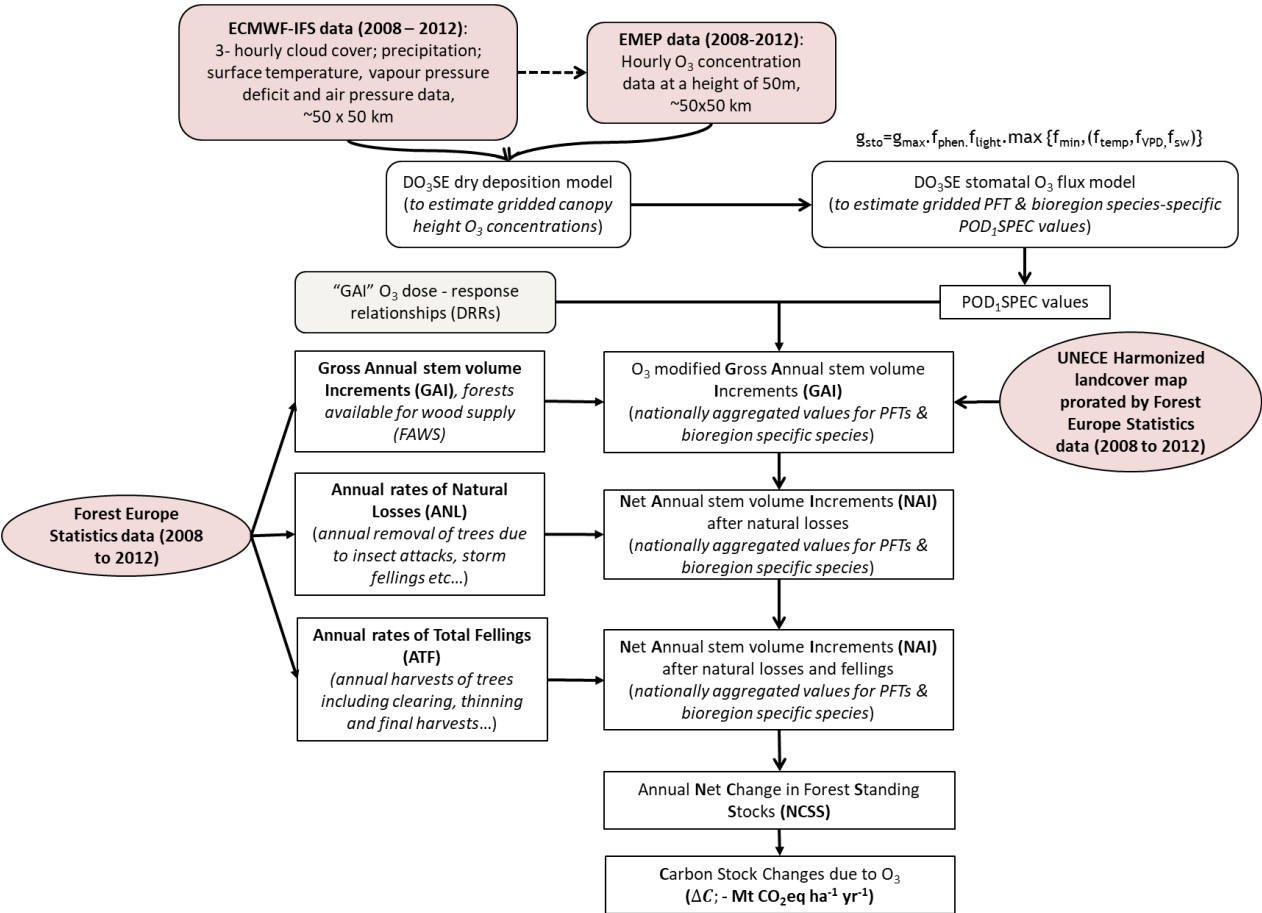
Vlasáková, L., Marková, J., Tognet, F., Horálek, J., Colette, A.: Evaluation of European-wide map creation of flux-based ozone indicator POD for selected tree species (Eionet Report – ETC HE 2022/23). European Topic Centre on Human Health and the Environment, 2022.

Volz, A., Kley, D.: Evaluation of the Montsouris series of ozone measurements made in the nineteenth century. *Nature*, 332, 240–242, <https://doi.org/10.1038/332240a0>, 1988.

Wittig, V. E., Ainsworth, E. a., Naidu, S. L., Karnosky, D. F., & Long, S. P.: Quantifying the impact of current and future tropospheric ozone on tree biomass, growth, physiology and biochemistry: a quantitative meta-analysis. *Global Change Biology*, 15(2), 396–424, <https://doi.org/10.1111/j.1365-2486.2008.01774.x>, 2009.

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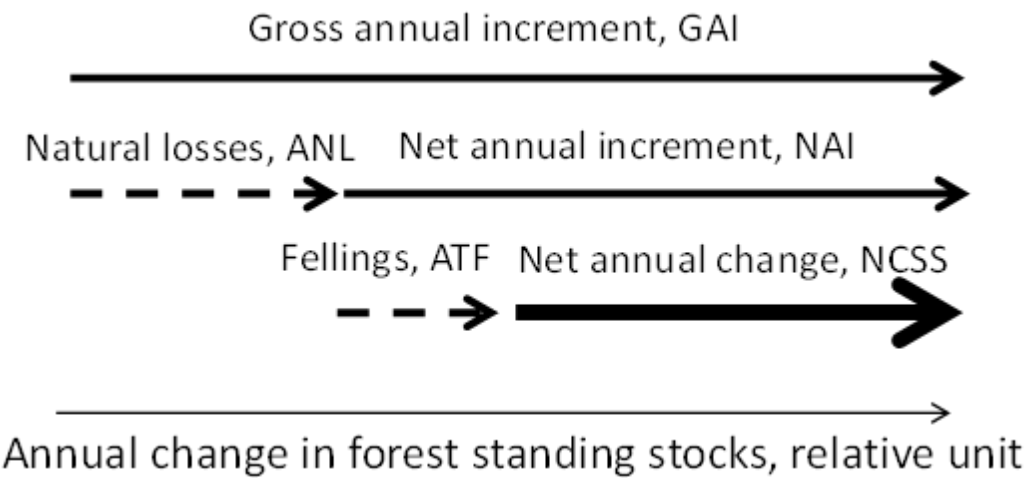
Figure 1: Overview of the datasets and methods used to estimate the influence of O<sub>3</sub> on forest annual living biomass C stock increments (Mt CO<sub>2eq</sub> yr<sup>-1</sup>) for the time period 2008 - 2012. A negative sign was used for the removal of CO<sub>2</sub> from the atmosphere to the living biomass carbon stocks.



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850 Figure 2. An illustration of the scheme used to calculate the net changes in the forest standing stocks. Modified from Forest Europe (2020) and Korosuo et al. (2023). The arrow with the thickest line represents the final value calculated as the net forest standing stock change (NCSS).



855 **Table 1. Constants used for the conversion from change in forest standing stock to changes in forest C stock, based on IPCC (Penman et al., 2003, Ågren et al., 2021).**

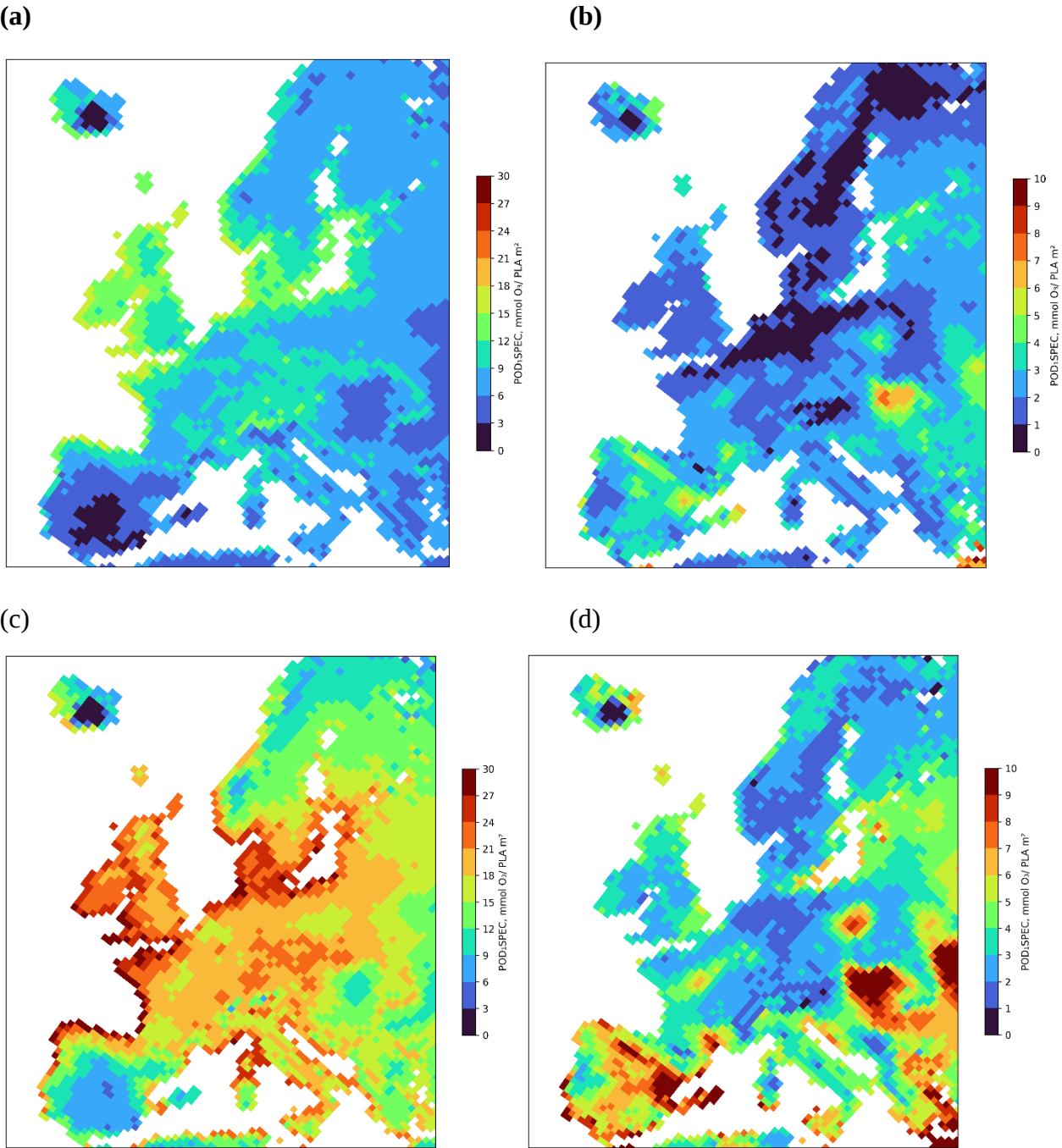
|                                                            | Coniferous tree species* | Broadleaf tree species* |
|------------------------------------------------------------|--------------------------|-------------------------|
| D, t dry mass m <sup>-3</sup>                              | 0.41**                   | 0.55***                 |
| BEF, biomass above ground expansion factor (dimensionless) | 1.125 <sup>#</sup>       | 1.15 <sup>##</sup>      |
| R, root/shoot ratio                                        | 0.32                     | 0.26                    |
| CF, carbon fraction of dry matter (t t <sup>-1</sup> ).    | 0.51 <sup>###</sup>      | 0.47 <sup>###</sup>     |

\* Valid for forest stands with standing biomass of 50-150 t ha<sup>-1</sup> for coniferous and 75-150 t ha<sup>-1</sup> for broadleaf species; \*\* mean value for *Picea abies* and *Pinus sylvestris*; \*\*\* mean value for *Betula* sp. and *Fagus sylvatica*; <sup>#</sup> mean value for *Picea abies* and *Pinus sylvestris*, boreal and temperate forests; <sup>##</sup> mean for broadleaf forests, boreal and temperate; <sup>###</sup> Ågren et al., 2021;

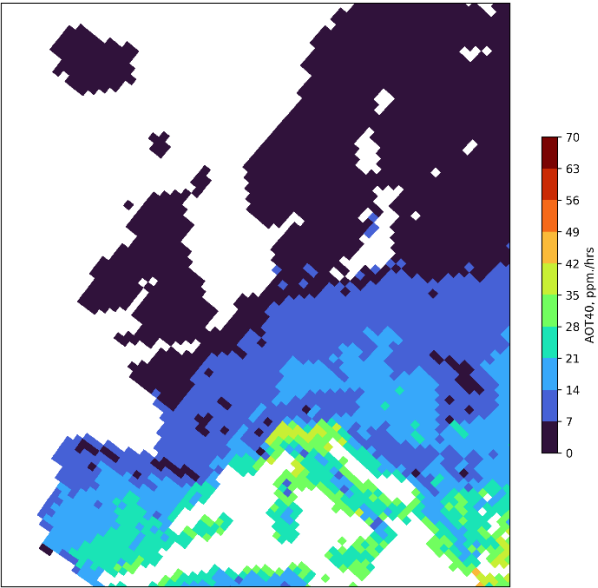
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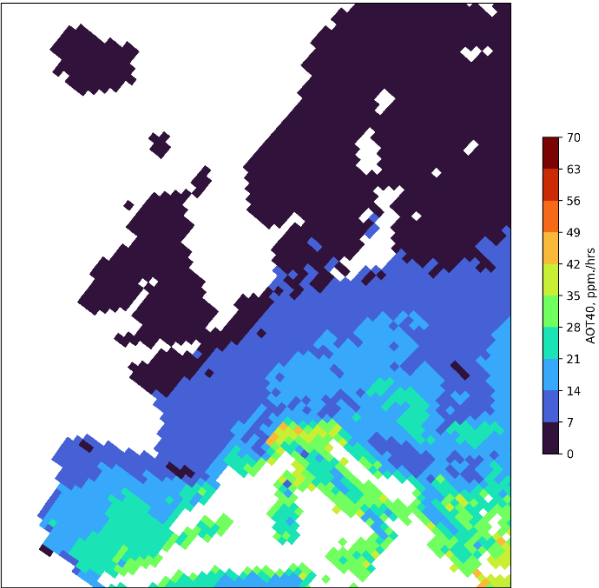
**Figure 3.** The spatial distribution of the estimated POD<sub>1</sub>SPEC (mmol O<sub>3</sub> m<sup>-2</sup> PLA) across Europe accumulated during the growing season using the f<sub>PAW</sub> scenario, separately for coniferous (a) and broadleaf (c) tree species for annual mean values for the period 2008-2012 and as the difference in maximum and minimum POD<sub>1</sub>SPEC values across the years 2008-2012 (coniferous, b; broadleaf, d). Also shown is the mean annual value of AOT40 in ppb.hrs for the period 2008-2012 and the Continental Central Europe (CCE) growing season of Norway Spruce (e) and Beech (f).



875 (e)



(f)

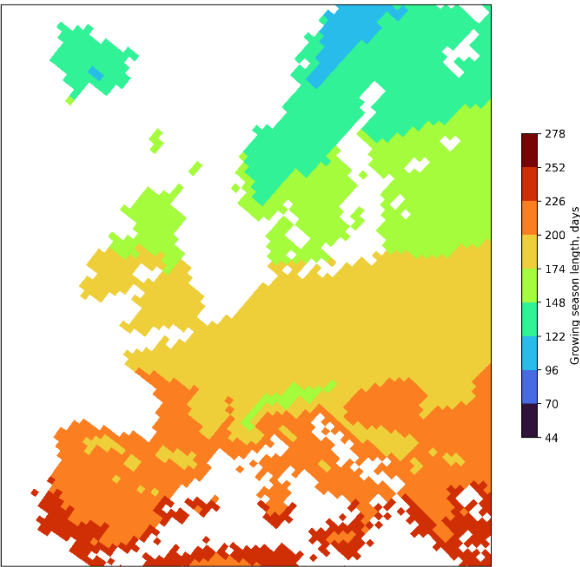


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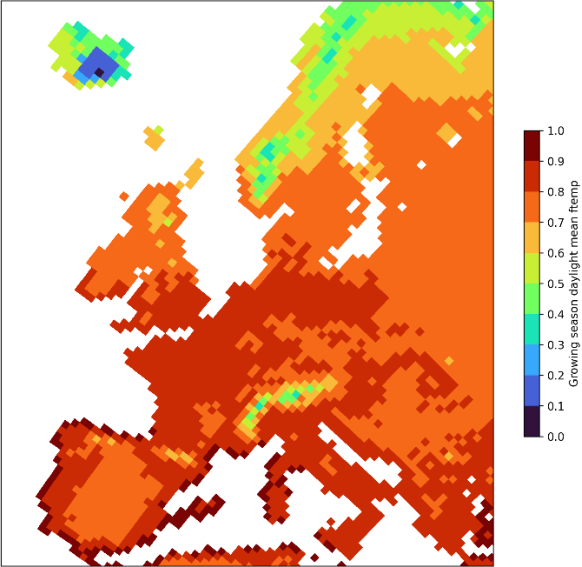
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Figure 4. The limits to POD<sub>1</sub>SPEC from each of the key three environmental variables (a) growing season length (in days) for broadleaf deciduous forests, (b)  $f_{temp}$ , (c)  $f_{VPD}$  and (d)  $f_{PAW}$  averaged during the growing season for all species and all years (2008-2012) across Europe. Grid values are mean values across all plant functional types for the growing season and for daylight hours only. Plots for individual species are given in the supplementary (Figure S4-S6).

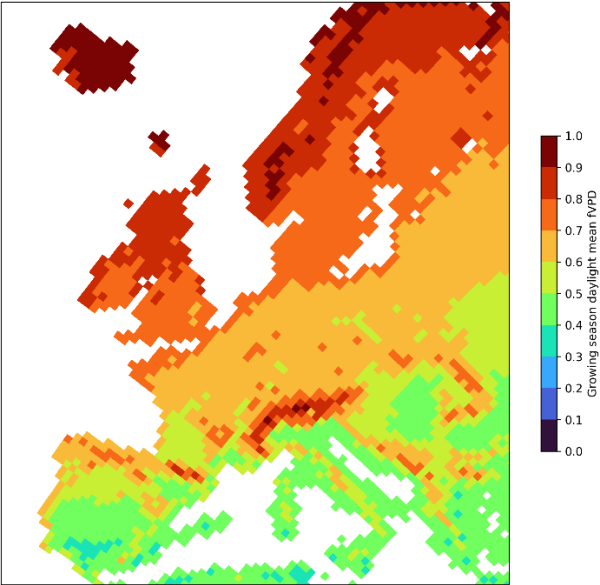
(a) Growing season length (days)



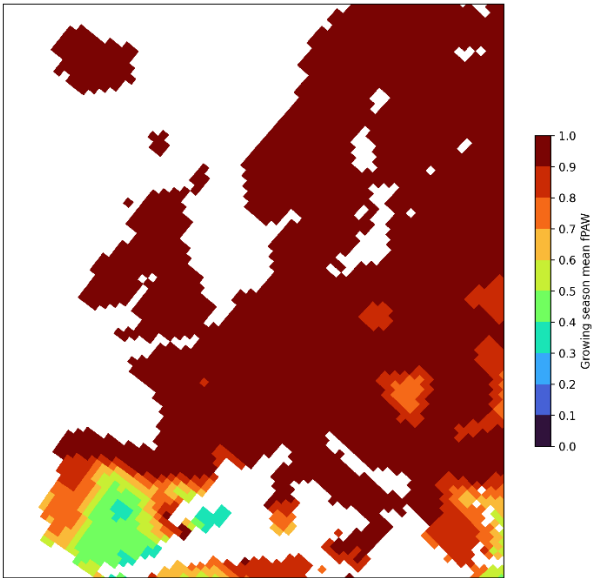
(b)  $f_{temp}$



(c)  $f_{VPD}$



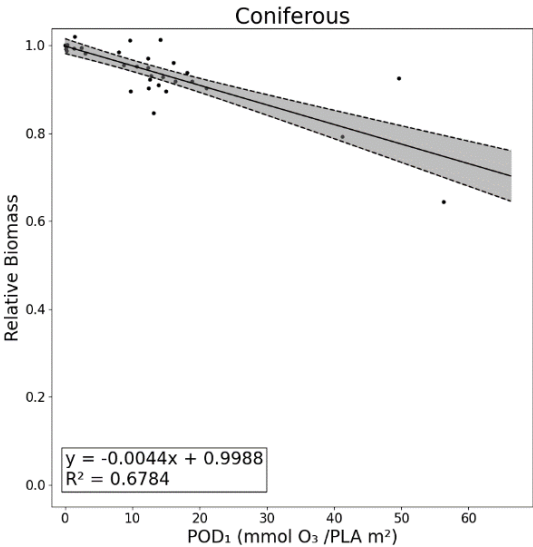
(d)  $f_{PAW}$



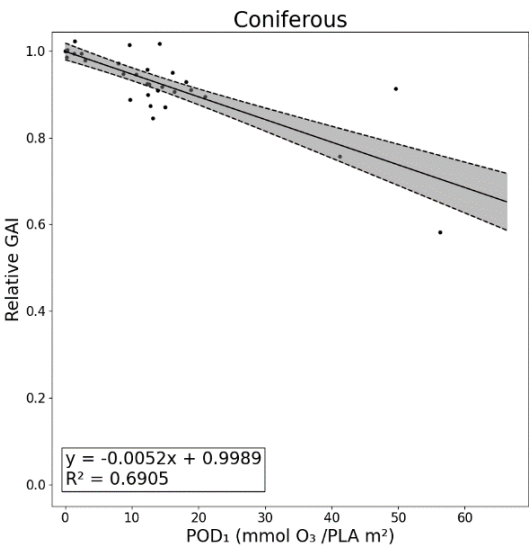
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**Figure 5. Dose-response relationships (DRRs) based on POD<sub>1</sub>SPEC for assessments based on impacts on the percent annual biomass loss (Relative biomass) (a, c) and the estimated impacts on gross annual increment (relative GAI) (b, d) separately for coniferous (a, b) and deciduous tree species (c, d). Other species and PFT dose-response relationships are given in the supplementary text (Table S3 and Figure S3).**

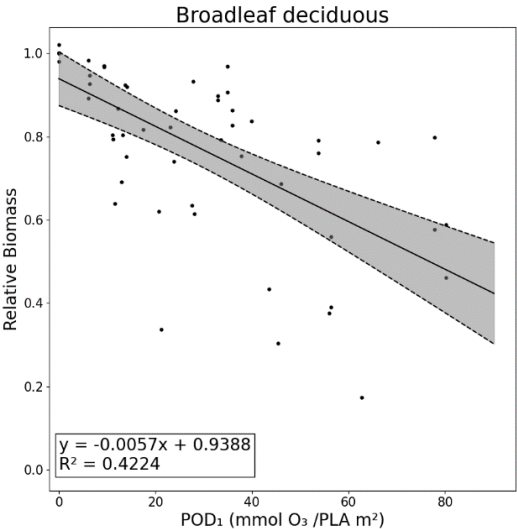
895 (a)



(b)



(c)



(d)

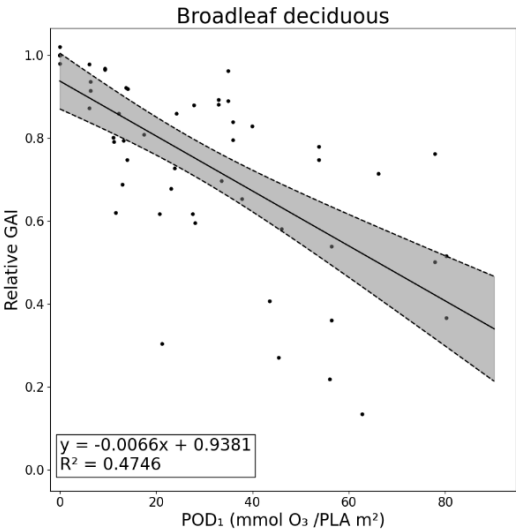
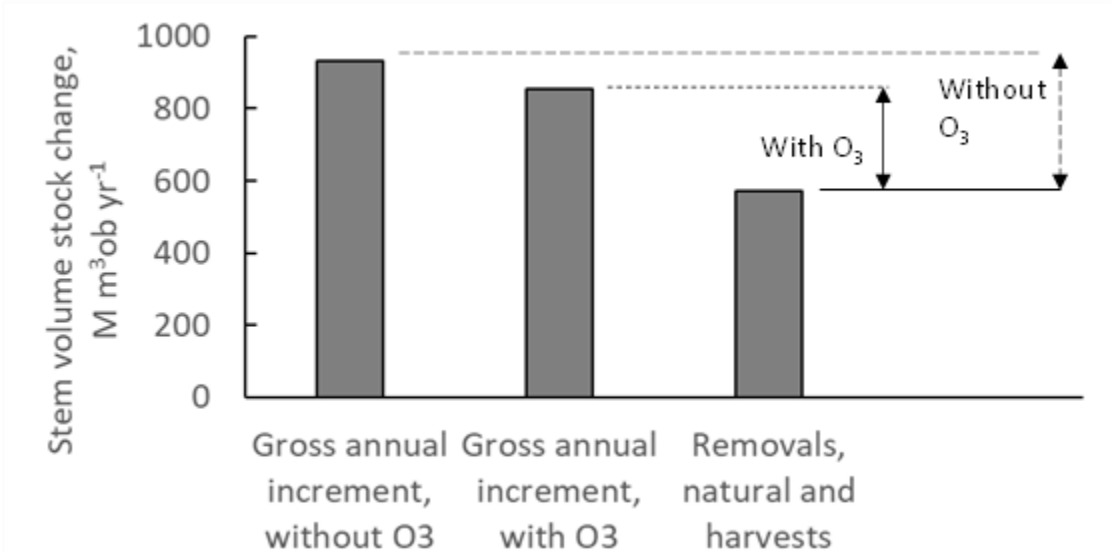


Table 2. Percent reductions in gross annual increments (GAI), caused by the present O<sub>3</sub> exposure, calculated as mean values for POD<sub>1</sub>SPEC for the period 2008-2012 and mean values for the area of different European countries. The estimated reduction of GAI was estimated based on O<sub>3</sub> dose-response relationships for impacts on growth rates (GAI DRRs) based on country-wide mean values for POD<sub>1</sub>SPEC calculated separately for coniferous and deciduous tree species and separately for scenarios with (fPAW) and without (NSW) the functions for soil water deficit included.

|                | Coniferous, NSW | Deciduous, NSW | Coniferous, fPAW | Deciduous, fPAW |
|----------------|-----------------|----------------|------------------|-----------------|
| Sweden         | -4              | -23            | -4               | -23             |
| Norway         | -2              | -17            | -2               | -17             |
| Finland        | -4              | -13            | -4               | -13             |
| Denmark        | -5              | -15            | -5               | -15             |
| Estonia        | -5              | -17            | -5               | -17             |
| Latvia         | -2              | -24            | -2               | -24             |
| Lithuania      | -4              | -14            | -4               | -14             |
| Germany        | -4              | -20            | -4               | -20             |
| Poland         | -1              | -29            | -0.5             | -28             |
| Netherlands    | -4              | -18            | -4               | -18             |
| Belgium        | -6              | -14            | -6               | -13             |
| United Kingdom | -4              | -23            | -4               | -23             |
| France         | -12             | -8             | -12              | -8              |
| Switzerland    | -1              | -26            | -1               | -26             |
| Austria        | -4              | -28            | -4               | -28             |
| Czech Republic | -4              | -26            | -4               | -26             |
| Hungary        | -10             | -5             | -10              | -4              |
| Slovakia       | -5              | -14            | -5               | -14             |
| Ireland        | -6              | -32            | -6               | -32             |
| Slovenia       | -8              | -18            | -8               | -18             |
| Bulgaria       | -7              | -17            | -7               | -17             |
| Romania        | -10             | -8             | -9               | -8              |
| Greece         | -26             | -13            | -23              | -11             |
| Italy          | -17             | -18            | -17              | -18             |
| Spain          | -13             | -15            | -9               | -10             |
| Portugal       | -22             | -13            | -19              | -9              |

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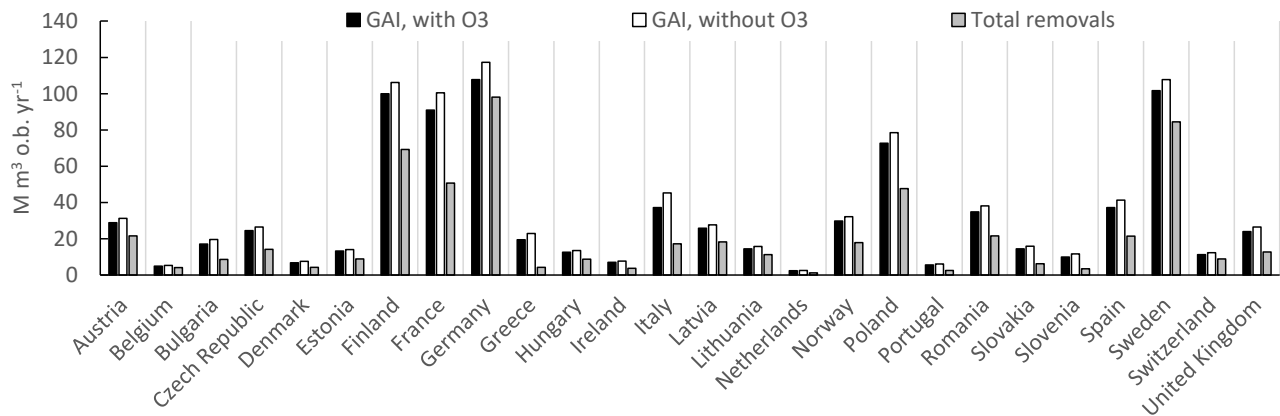
Figure 6. The rates of total, annual gross stem volume increment growth for forests in Europe as mean annual values for the time period 2008-2012, including the sum of coniferous and deciduous tree species, with and without the exposure to present O<sub>3</sub> doses, calculated as POD<sub>1</sub>SPEC with the scenario f<sub>PAW</sub>. Also shown are the estimated total removals of forest stem volumes, including both natural- and harvest removals. The rates of removals are assumed to be independent of O<sub>3</sub> regimes. Furthermore, the differences between the gross annual stock increments and the removals are indicated with vertical arrows, separately for the presence of O<sub>3</sub> impacts (continuous line arrow) and without the presence of O<sub>3</sub> impacts (dashed line arrow). ob, over bark.



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Figure 7. The rates of annual gross stem volume increment growth for forests in different European countries, as annual values for the time period 2008-2012, including the sum of coniferous and deciduous tree species, with and without the exposure to present O3 doses, calculated as POD<sub>1</sub>SPEC with the scenario f<sub>PAW</sub>. Also shown are the sum of the natural and harvest removals of forest stem volumes, i.e. the total removals. The rates of natural and harvest removals are assumed to be independent of ozone regimes. The countries are sorted left to right in alphabetical order.



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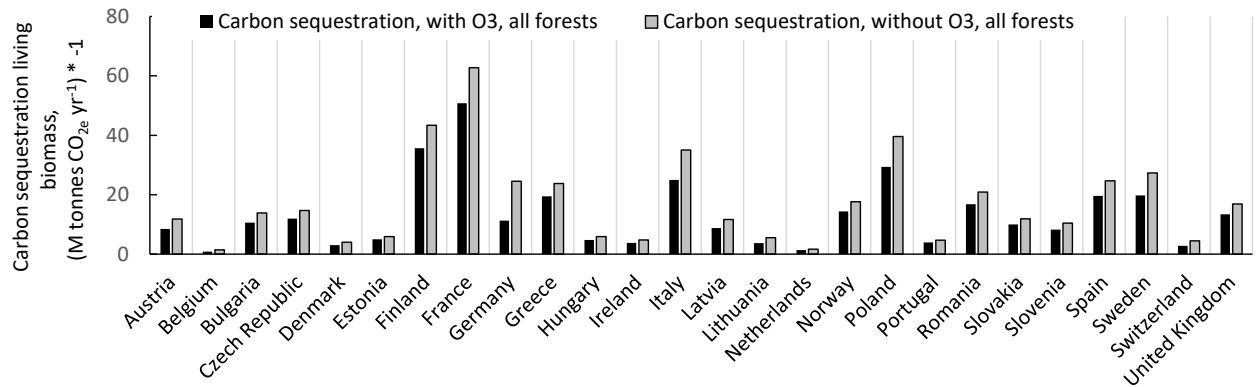
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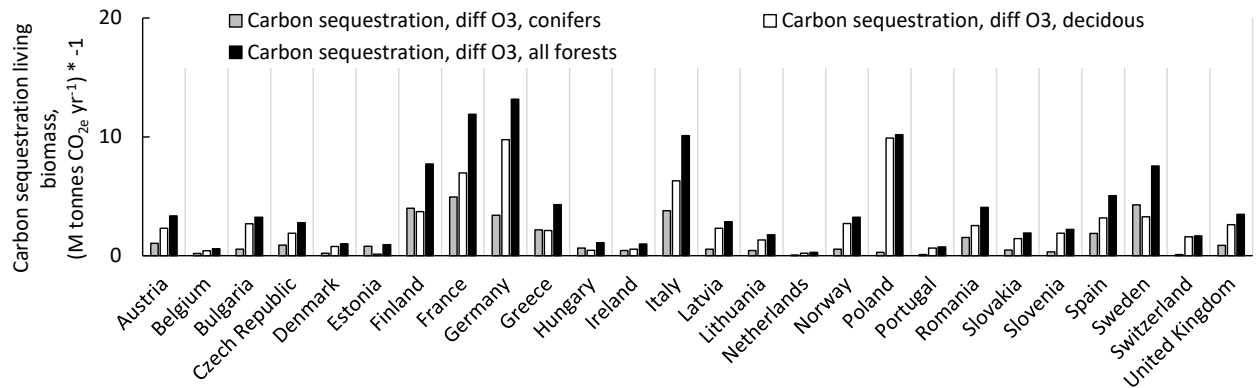
**Figure 8. The changes in the annual C sequestration in the living biomass C stocks, with and without the exposure to present O<sub>3</sub> doses, calculated as POD<sub>1</sub>SPEC with the scenario f<sub>PAW</sub>. (a). absolute values for total forests; (b). differences without versus with ozone, separately for coniferous, deciduous and total forests. Results are presented for the individual Europeans countries, separately for coniferous and deciduous and for total forests. The countries are sorted left to right in alphabetical order. Removal of CO<sub>2</sub> from the atmosphere to the living biomass carbon stocks is shown with a negative sign.**

a



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b



**Figure 9. The percent changes in the annual C sequestration of forests in the different European countries, without the present O<sub>3</sub> exposure compared to with O<sub>3</sub> exposure. Changes are shown separately for coniferous and deciduous forests and for total forests.**

