The role of defoliation and root rot pathogen infection in driving the mode of drought-related physiological decline in Scots pine (Pinus sylvestris L.)

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Drought-related tree die-off episodes have been observed in all vegetated continents. Despite much research effort, however, the multiple interactions between carbon starvation, hydraulic failure and biotic agents in driving tree mortality under field conditions are still not well understood. We analysed the seasonal variability of non-structural carbohydrates (NSCs) in four organs (leaves, branches, trunk and roots), the vulnerability to embolism in roots and branches, native embolism (percentage loss of hydraulic conductivity (PLC)) in branches and the presence of root rot pathogens in defoliated and non-defoliated individuals in a declining Scots pine (Pinus sylvestris L.) population in the NE Iberian Peninsula in 2012, which included a particularly dry and warm summer. No differences were observed between defoliated and non-defoliated pines in hydraulic parameters, except for a higher vulnerability to embolism at pressures below −2 MPa in roots of defoliated pines. No differences were found between defoliation classes in branch PLC. Total NSC (TNSC, soluble sugars plus starch) values decreased during drought, particularly in leaves. Defoliation reduced TNSC levels across tree organs, especially just before (June) and during (August) drought. Root rot infection by the fungal pathogen Onnia P. Karst spp. was detected but it did not appear to be associated to tree defoliation. However, Onnia infection was associated with reduced leaf-specific hydraulic conductivity and sapwood depth, and thus contributed to hydraulic impairment, especially in defoliated pines. Infection was also associated with virtually depleted root starch reserves during and after drought in defoliated pines. Moreover, defoliated and infected trees tended to show lower basal area increment. Overall, our results show the intertwined nature of physiological mechanisms leading to drought-induced mortality and the inherent difficulty of isolating their contribution under field conditions.

Keywords: die-off, fungi pathogen, global change, hydraulic failure, non-structural carbohydrates.

Introduction

Drought-induced tree die-off is emerging as a global phenomenon, affecting a great variety of species and ecosystems in all vegetated continents of the world (Allen et al. 2010). Recent episodes of crown defoliation and tree mortality have been related to an increase of mean annual temperature and a decrease of annual rainfall in southern European forests (Carnicer et al. 2011) and with increasing severe droughts in the southwestern USA (van Mantgem et al. 2009, Williams et al. 2012). Extreme drought events are expected to become more frequent (IPCC 2013), which could accelerate drought-related tree mortality. These responses can be amplified in many regions by current trends towards increased stand basal area, associated with changes in forest management (Martínez-Vilalta et al. 2012). There are important feedback loops between forest dynamics and climate due to the key role of forests on the global climate.
water and carbon cycles (Bonan 2008) and thus widespread forest mortality can have rapid and drastic consequences for ecosystems (Anderegg et al. 2013a). Nonetheless, and despite these potential effects on ecosystem functioning, the mechanisms causing tree die-off are still poorly understood (Sala et al. 2010, McDowell 2011, McDowell et al. 2011). McDowell et al. (2008) introduced a framework with three main, non-exclusive mechanisms that could cause drought-induced mortality in trees: (i) carbon starvation, (ii) hydraulic failure and (iii) biotic agents. They also hypothesized that plants with a strict control of water loss through stomatal closure (isohydric species) would be more likely to die of carbon starvation during a long drought, whereas anisohydric species would more likely experience hydraulic failure during intense droughts, due to more negative xylem water potentials (McDowell et al. 2008). However, the evidence for or against these proposed mortality mechanisms is inconclusive, and recent reports have urged adoption of a more integrated approach focusing on the interrelations between plant hydraulics and the economy and transport of carbon in plants (McDowell and Sevanto 2010, Sala et al. 2010, 2012, McDowell 2011, McDowell et al. 2011).

Xylem vulnerability to embolism has been found to place a definitive limit on the physical tolerance of conifers (Brodribb and Cochard 2009) and angiosperms (Urli et al. 2013) to desiccation. A recent global synthesis has shown that many tree species operate within relatively narrow hydraulic safety margins in all major biomes of the world (Choat et al. 2012). Although structural and physiological acclimation to drought may result in large safety margins from hydraulic failure preceding death (Plaut et al. 2012), high levels of hydraulic dysfunction associated with drought-induced desiccation have been reported (Hoffmann et al. 2011, Nardini et al. 2013). In addition, evidence of hydraulic failure linked to canopy and root mortality has been found in declining trembling aspen (Populus tremuloides Michx.), without evidence of depletion of carbohydrate reserves (Anderegg et al. 2012), and in some Eucalyptus L’Hér. species (Mitchell et al. 2013).

The dynamics and role of non-structural carbohydrate (NSC) stores during drought are still under debate, but some agreement is emerging in that NSC concentrations tend to increase under moderate drought because growth ceases before photosynthesis, whereas they may decline sharply if drought conditions become extreme (McDowell 2011). Recent reports show that the same drought conditions can have contrasting effects on NSC concentrations even on species of the same genus (Nothofagus Blume) (Piper 2011). On the other hand, Galiano et al. (2011) reported extremely low NSC concentrations in the stem of defoliated Scots pine (Pinus sylvestris L.) trees and higher probability of drought-induced mortality in pines with lower NSC concentrations. Also, Adams et al. (2013) observed an association between tree mortality and reduced NSC levels in leaves in a drought simulation experiment on piñon pine (Pinus edulis Engelm.). In addition, drought-related reductions in stored NSC pools may affect tree organs differentially. For instance, Hartmann et al. (2013a) showed a reduction of NSC in Norway spruce (Picea abies (L.) Karst.) roots but not in leaves, which could be attributed to changes in carbon allocation.

Biotic agents and drought can interact to accelerate tree mortality. Drought-stressed trees may be more vulnerable to infection by fungal pathogens (Desprez-Loustau et al. 2006, La Porta et al. 2008) or to insect attacks (Matthias et al. 2007, Gaylord et al. 2013). Although pathogens can directly kill trees through the production of toxic metabolites, they can also induce hydraulic failure, e.g., via occlusion of the xylem, or carbon starvation by altering NSC demand or supply. The interactions between pathogen infection and the physiological mechanisms of drought-induced mortality depend on the trophic interaction (biotrophic, necrotrophic or vascular wilts) established with the tree (Oliva et al. 2014). For instance, blue-stain fungi infection can cause outright hydraulic failure via xylem occlusion (Hubbard et al. 2013) while root rot fungi may lead to a gradual tree decline and eventually to death because of chronic growth reductions and subsequent constraints on water transport (Oliva et al. 2012). In addition, root rot fungi can reduce carbon reserves by fungal consumption of stored carbohydrates or by the induction of carbon-expensive defences, causing tree growth reductions (Cruickshank et al. 2011).

Field studies on mature trees examining all three hypothesized mechanisms of drought-induced mortality are still scarce and rarely explore explicitly the interactions between different mechanisms and their occurrence in different plant organs. Here we compare the hydraulic properties and the dynamics of NSC and embolism as a function of defoliation and infection by fungal pathogens in Scots pine trees growing together in a site affected by drought-induced decline (Martínez-Vilalta and Piñol 2002, Hereş et al. 2012) and close to the dry limit of the distribution of the species. Previous studies have shown that defoliation precedes drought-induced mortality in Scots pine trees from the same or similar sites (Galíano et al. 2011, Poyatos et al. 2013), and that defoliation seems to be an inevitable consequence of drought in the most susceptible individuals rather than an (effective) strategy to cope with it (Poyatos et al. 2013). In this context, we address the following questions: (i) are defoliated Scots pines intrinsically more vulnerable to xylem embolism than non-defoliated ones, providing evidence in favour of hydraulic failure as an important component of the decline process? If so, do defoliated trees experience higher levels of native embolism under dry summer conditions? (i.e., Are leaf area reductions enough to compensate for their intrinsically higher vulnerability?) (ii) Do defoliated pines have lower seasonal NSC concentrations in all organs, supporting carbon starvation being involved in the mortality process? (iii) Is infection by fungal pathogens likely to enhance hydraulic dysfunction through increased levels of native embolism, or carbon depletion directly
or indirectly lowering NSC levels through increased consumption, reduced sapwood depth or low growth?

**Materials and methods**

**Study site**

The study was conducted in Tillar valley (41°19′N, 1°00′E; 990 m above sea level) within Poblet Forest Natural Reserve (Prades Mountains, NE Iberian Peninsula). The climate is typically Mediterranean, with a mean annual precipitation of 664 mm (spring and autumn being the rainiest seasons and with a marked summer dry period), and moderately warm temperatures (11.3 °C on average) (Poyatos et al. 2013). The soils are mostly Xerochrepts with fractional schist and clay loam texture, although outcrops of granitic sandy soils are also present (Hereter and Sánchez 1999). Our experimental area is mostly located on a NW-facing hillside with a very shallow and unstable soil due to the high stoniness and steep slopes (35° on average). More detailed information about the study area can be found in Hereter and Sánchez (1999).

The dominant canopy tree species in the study site is Scots pine and the understory consists mainly of the Mediterranean evergreen holm oak (*Quercus ilex* L.). Severe droughts have affected the study site since the 1990s (Martínez-Vilalta and Prado 2002, Hereš et al. 2012). Scots pine average standing mortality and crown defoliation in the Tillar valley are currently 12 and 52%, respectively. However, in some parts of the forest standing mortality is >20% and cumulative mortality is as high as 50% in the last 20 years (J. Martinez-Vilalta, unpublished data). The Scots pine population studied is more than 150 years old and has remained largely unmanaged for the past 30 years (Hereš et al. 2012). No major insect infestation episode associated with the forest decline in the area has been detected (M. Rojo, Catalan Forest Service, personal communication).

A mixed Holm oak–Scots pine stand with a predominantly northern aspect was selected for this study where defoliated and non-defoliated pines were living side by side. A total of 10 defoliated and 10 non-defoliated Scots pine trees were sampled (see Table S1 available as Supplementary Data at Tree Physiology Online). In order to minimize unwanted variation, all trees had a diameter at breast height (DBH) between 25 and 50 cm (average DBH 36.08 ± 1.53 cm; similar between defoliation classes \( P > 0.05 \); data not shown)) and the distance between sampled trees was always >5 m (the average minimum distance was 41.99 ± 13.74 m). In this study, defoliation was visually estimated relative to a completely healthy tree in the same population (cf. Galiano et al. 2011). A tree was considered as non-defoliated if the percentage of green needles was >60% (average green leaves of the sampled non-defoliated trees = 77%) and defoliated if the percentage of green needles was <40% (average green leaves of the sampled defoliated trees = 26%). The average height of Scots pines in the population studied was 14.1 ± 0.5 m (Poyatos et al. 2013). All measurements, in combination with a continuous monitoring of the main meteorological variables and soil moisture (cf. Poyatos et al. 2013 for details), were carried out during 2012. Sampling from defoliated trees avoided dead or dying branches (i.e., those with no green leaves), so our sampling can be considered representative only of the living part of the crown.

**Non-structural carbohydrates sampling and analysis**

Trees were sampled in March (late winter), June (late spring), August (mid-summer) and October (early autumn). Four organs (leaves, branches, trunk and coarse roots) were sampled from every tree to measure NSCs. Sun-exposed branches (0.5–1 cm of diameter with bark removed) were excised with a pole pruner around noon (12:00 pm–3:00 pm, local time) to minimize the impact of diurnal variation in NSC concentrations due to photosynthetic activity (Li et al. 2008, Gruber et al. 2012). One-year-old leaves were sampled from these branches. Trunk xylem at breast height and coarse roots at 5–10 cm soil depth were cored to obtain sapwood. Sapwood portion was visually estimated in situ from all the extracted cores. We also used these cores to measure the basal area increment (BAI) corresponding to the three most recent annual rings (see Table S1 available as Supplementary Data at Tree Physiology Online). All samples were placed immediately in paper bags and stored in a portable cooler containing cold accumulators. One of the defoliated Scots pine trees (tree 1637; see Table S1 available as Supplementary Data at Tree Physiology Online) died during the study period and leaves could not be sampled in August and October, whereas branches could be sampled in August but not in October.

At the end of every sampling day, samples were microwaved for 90 s in order to stop enzymatic activity and oven-dried for 72 h at 65 °C. Samples were then ground to fine powder in the laboratory. Total NSC (TNSC) was defined as including free sugars (glucose and fructose), sucrose and starch, and were analysed following the procedures described by Hoch et al. (2002) with some minor variations (cf. Galiano et al. 2011). Approximately 12–14 mg of sample powder was extracted with 1.6 ml distilled water at 100 °C for 60 min. After centrifugation, an aliquot of the extract was used for the determination of soluble sugars (glucose, fructose and sucrose), after enzymatic conversion of fructose and sucrose into glucose (invertase from *Saccharomyces cerevisiae* Meyen ex E.C. Hansen) and glucose hexokinase (*GHK* assay reagent, I4504 and G3293, Sigma-Aldrich, Madrid, Spain). Another aliquot was incubated with an amyloglucosidase from *Aspergillus niger* van Tieghem at 50 °C overnight, to break down all NSC (starch included) to glucose. The concentration of free glucose was determined photometrically in a 96-well microplate reader (Sunrise Basic Tecan, Männedorf, Switzerland) after enzymatic modification with some minor variations (cf. Galiano et al. 2011). Approximately 12–14 mg of sample powder was extracted with 1.6 ml distilled water at 100 °C for 60 min. After centrifugation, an aliquot of the extract was used for the determination of soluble sugars (glucose, fructose and sucrose), after enzymatic conversion of fructose and sucrose into glucose (invertase from *Saccharomyces cerevisiae* Meyen ex E.C. Hansen) and glucose hexokinase (*GHK* assay reagent) conversion of glucose to gluconate-6-phosphate. Starch was calculated as TNSC minus soluble sugars. All NSC values are
expressed as percent dry matter. Throughout the manuscript NSC is used to refer generically to NSCs, while TNSC is used to refer specifically to the total value of NSC (sum of starch and soluble sugars). The relationship between soluble sugars (SS) and TNSC will be expressed as SS : TNSC. Finally, we note that due to the current uncertainty in NSC quantification methodologies (Quentin et al., in review), our results should be considered valid in relative terms (as used here) but not necessarily comparable to the values obtained by other laboratories or using different methods.

Hydraulic conductivity and vulnerability to xylem embolism
Hydraulic measurements were taken on the same Scots pine individuals as NSCs, except for one defoliated tree that could not be sampled for hydraulics. Branches and roots were sampled in April and May 2012, respectively, for determining xylem vulnerability curves. We selected branches and roots containing internodal segments 0.4–1.1 cm in diameter and ~15 cm in length. Branches were always sampled from the exposed part of the canopy and were 3–5 years old. Root samples were taken at a soil depth of ~20 cm and always from the down slope side of the trunk, in order to control for differences in water availability or soil properties. Sampled branches and roots were always longer than 40 cm and were immediately wrapped in wet cloths and stored inside plastic bags until they were transported to the laboratory on the same day. Once in the laboratory, samples were stored at 4 °C until their vulnerability curves were established within <3 weeks. Right before that, all leaves distal to the measured branch segments were removed and their total area measured with a Li-Cor 3100 Area Meter (Li-Cor Inc., Lincoln, NE, USA).

Vulnerability curves relating the percentage loss of hydraulic conductivity (PLC) as a function of xylem water potential were established using the air injection method (Cochard et al. 1992). Hydraulic conductivity (water flow per unit pressure gradient) was measured using the XYL’EM embolism meter (Bronkhorst, Montigny-Les-Cormeilles, France) using deionized, degassed water (Liqui-Cel Mini-Module degassing membrane) and a pressure head of 4.5 kPa. The bark was removed from all measurement segments and their ends were cleanly shaved with a sharp razor blade before connecting to the XYL’EM apparatus.

Branch segments were rehydrated with deionized, degassed water prior to determining maximum hydraulic conductivity ($K_{max}$) (Espino and Schenk 2011), leading to stable and consistent $K_{max}$ measurements. Afterwards, all segments (four to six each time) were placed inside a multi-stem pressure chamber with both ends protruding. Then, we raised the pressure inside the chamber to 0.1 MPa (basal value) for 10 min, lowered the pressure, waited 10 min to allow the system to equilibrate and measured hydraulic conductivity under a low pressure head (4.5 kPa). Stable conductivity readings were usually achieved within 3 min. These measurements were considered to represent the maximum hydraulic conductivity ($K_{max}$) and were used as reference conductivity (PLC = 0) for the purpose of establishing vulnerability curves. We repeated this process, raising the injection pressure stepwise by 0.5 MPa (roots) or 1 MPa (branches), until the actual conductivity of the segment was <20% of $K_{max}$ or when we reached 4 MPa. Due to technical limitations, we could not increase the pressure in the chamber over 4 MPa, but only five samples (out of 38) had not reached 80% PLC at this pressure. We fitted vulnerability curves with the following function (Pammenter and Van der Willigen 1998):

$$PLC = \frac{100}{1 + \exp(\alpha(P - P_{50}))}$$

In this equation, PLC is the percentage loss of hydraulic conductivity, $P$ the applied pressure, $P_{50}$ the pressure (i.e., $-\psi$) causing a 50% PLC and $\alpha$ is related to the slope of the curve. Parameters were estimated with nonlinear least squares regression. Some vulnerability curves led to inconsistent results, due to technical problems, and were removed from the analyses. Final sample size was 15 roots (six from defoliated and nine from non-defoliated trees) and 15 branches (seven from defoliated and eight from non-defoliated trees).

Measurements of $K_{max}$ were used for calculating specific hydraulic conductivity ($K_s$, in m$^2$ MPa$^{-1}$ s$^{-1}$), as the ratio between maximum hydraulic conductivity and mean cross-sectional area of the segment (without bark), and leaf-specific conductivity ($K_L$, in m$^2$ MPa$^{-1}$ s$^{-1}$), as the quotient between maximum hydraulic conductivity and distal leaf area. Finally, we calculated the ratio between distal leaf area and cross-sectional area ($A_s : A_s$) of each branch segment.

Monitoring water potentials and native embolism
We sampled one exposed branch from each of the trees that had been sampled for vulnerability curves (9 defoliated and 10 non-defoliated individuals) monthly from May to August and in October 2012. Midday and predawn leaf water potentials were usually measured in situ on nearby defoliated and healthy pines on the same sampling dates. We selected branches at least 40 cm long and containing internodal segments of 4–10 cm length (0.3–0.9 cm in diameter). Immediately following excision, branch samples were placed in plastic bags with a small piece of damp paper towel, stored in a bigger bag containing cold accumulators and transported to the laboratory within 3 h. All branches were sampled before 8 a.m., solar time.

Once in the laboratory, we measured shoot water potential on one terminal shoot per sampled branch using a Scholander-type pressure chamber (PMS Instruments, Corvallis, OR, USA) (except for the initial, May sampling). To measure native embolism, we cut wood segments (4–10 cm in length) underwater from each sampled branch. The bark was removed from the measurement segments and their ends were cleanly shaved with a sharp razor blade before connecting them to the tubing system...
of the XYL’EM apparatus to measure native hydraulic conductivity ($K_i$) under a pressure head of $\sim 4.5$ kPa. Measurement samples were rehydrated overnight and their maximum hydraulic conductivity ($K_{\text{max}}$) was measured with the XYL’EM apparatus as described above. The PLC due to embolism was computed as:

$$\text{PLC} = 100 \left(1 - \frac{K_i}{K_{\text{max}}}\right)$$

(2)

**Root rot pathogens**

We focused on root and butt rot pathogens as possible contributors to the decline process observed in the forest studied. From each tree, we extracted two cores with an increment borer, one at stump height and the other in one of the large roots. Decay presence was noted and cores were kept under sterile conditions. Within 48 h, they were placed onto Hagem medium amended with benomyl ($10 \text{ mg} \text{l}^{-1}$) and cloramphenicol ($200 \text{ mg} \text{l}^{-1}$). Plates were checked weekly during the following 3 months, and any mycelia growth was sub-cultured. Identification of fungal isolates was first attempted by sequencing the ITS region. Some of our isolates showed an equally low similarity (95%) to other isolates classified as either *Onnia tomentosa* (Fr.) P. Karst or *Onnia circinata* (Fr.) P. Karst in reference databases such as GenBank or UNITE. We identified our cultures as *Onnia* P. Karst spp., as morphological characters in culture and comparison with reference cultures from CBS-KNAW Fungal Biodiversity Centre (CBS 246.30 *O. circinata*, CBS 278.55 *O. tomentosa*) did not yield a conclusive identification.

We found signs of fungi infection after the cultivation of the extracted cores in seven pines; three of them were non-defoliated and the other four were defoliated (see Table S1 available as Supplementary Data at *Tree Physiology* Online). One defoliated and one non-defoliated tree were infected by a heart rot fungi (*Porodaedalea pini* (Brot.) Murrill). These fungi mainly cause heartwood decay with no major known physiological effects (*Garbelotto 2004*), so we considered these two trees as ‘non-infected’ in our analyses. The other trees (two non-defoliated and three defoliated) were infected by *Onnia* sp., which causes root rot and physiological impairment, and we considered those trees as ‘infected’ in subsequent analyses.

**Data analysis**

We used different types of linear models to test the effects of defoliation and fungal infection on the measured response variables. In each case we included the main covariates (measured organ and sampling month, as appropriate) and the most biologically plausible interactions. Note that model complexity, in terms of the number of interactions that could be included, is subject to sample size limitations. Preliminary analyses showed that DBH effects were never significant in the models and therefore were not included in the final models reported here. General linear models were fitted to study the effects of defoliation class (two levels: defoliated or non-defoliated) and infection occurrence (two levels: infected or non-infected) on $K_i$, $A_i : A_S$, BAI, and root and trunk sapwood depths. Separate linear models were fitted for each of the five response variables (see Eq. (S1) available as Supplementary Data at *Tree Physiology* Online). Linear mixed models were used to analyse vulnerability-curve parameters ($\alpha$ and $P_{50}$) and $K_S$ as a function of defoliation class, organ (branch or root) and infection occurrence. Tree identity was included in the models as a random factor (see Eq. (S2) available as Supplementary Data at *Tree Physiology* Online). Similar mixed models were used to analyse shoot water potential, PLC, TNSC and SS : TNSC as a function of defoliation class, sampling date, organ (only for TNSC and SS : TNSC) and infection occurrence, including again tree identity.
as a random factor (see Eqs (S3) and (S4) available as Supplementary Data at *Tree Physiology* Online). Finally, starch concentration in roots was also analysed using similar models (without the organ effect) to study in detail the impact of root rot pathogens on this NSC fraction and how it developed over time (and hence in this model we included the interaction between defoliation class, infection occurrence and sampling date) (see Eq. (S5) available as Supplementary Data at *Tree Physiology* Online).

Variables $K_p$, $K_L$, $A_L$, $A_S$, TNSC, and root and trunk sapwood depth were log-transformed, and starch concentration in roots was square root-transformed, to achieve normality prior to all analyses. All analyses were carried out with R Statistical Software.
version 3.1.0 (R Core Team 2014) using the lm and lme functions.

Results

Meteorological conditions

The summer of 2012 was particularly dry and warm compared with the climatic average for the period 1951–2010: average air temperature (June–August) was 20.56 °C and total precipitation was only 56 mm (Figure 1), compared with climatic values (1951–2010) of 19.30 °C and 135.23 mm, respectively. High values of daytime-averaged temperatures and vapour pressure deficits (VPDs) occurred in mid-August, followed by very low soil water content (SWC) values (~0.08 m³ m⁻³) at the beginning of September, compared with a spring maximum soil moisture of 0.33 m³ m⁻³ (Figure 1).

Non-structural carbohydrates

Concentrations of total NSCs varied among tree organs, in the following order: TNSC (leaves) > TNSC (branches) > TNSC (roots) > TNSC (trunk) (Figure 2, Table 1). In general, non-defoliated Scots pine trees showed higher concentrations of TNSC throughout the study period and across all organs (Figure 2). Total non-structural carbohydrate values increased

<table>
<thead>
<tr>
<th>Parameter</th>
<th>log(TNSC)</th>
<th>SS:TNSC</th>
<th>sqrt(Starch)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>2.50 ± 0.19***</td>
<td>0.54 ± 0.06***</td>
<td>0.65 ± 0.13***</td>
</tr>
<tr>
<td>Branches</td>
<td>−1.31 ± 0.19***</td>
<td>0.27 ± 0.07***</td>
<td>ne</td>
</tr>
<tr>
<td>Trunk</td>
<td>−3.22 ± 0.19***</td>
<td>0.10 ± 0.07 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>Roots</td>
<td>−2.78 ± 0.19***</td>
<td>0.07 ± 0.07 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>June</td>
<td>−0.15 ± 0.19 (ns)</td>
<td>−0.08 ± 0.07 (ns)</td>
<td>−0.02 ± 0.18 (ns)</td>
</tr>
<tr>
<td>August</td>
<td>−1.17 ± 0.19***</td>
<td>0.41 ± 0.07***</td>
<td>−0.40 ± 0.18*</td>
</tr>
<tr>
<td>October</td>
<td>−1.28 ± 0.19***</td>
<td>0.37 ± 0.07***</td>
<td>−0.51 ± 0.18**</td>
</tr>
<tr>
<td>Non-defoliated</td>
<td>0.15 ± 0.17 (ns)</td>
<td>−0.10 ± 0.05*</td>
<td>0.21 ± 0.21</td>
</tr>
<tr>
<td>Non-infected</td>
<td>0.18 ± 0.20 (ns)</td>
<td>−0.09 ± 0.06 (ns)</td>
<td>−0.17 ± 0.16 (ns)</td>
</tr>
<tr>
<td>Branches:June</td>
<td>0.31 ± 0.19 (ns)</td>
<td>−0.13 ± 0.07*</td>
<td>ne</td>
</tr>
<tr>
<td>Trunk:June</td>
<td>0.31 ± 0.19 (ns)</td>
<td>−0.03 ± 0.07 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>Roots:June</td>
<td>0.21 ± 0.19 (ns)</td>
<td>0.01 ± 0.07 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>Branches:August</td>
<td>1.23 ± 0.19***</td>
<td>−0.37 ± 0.08***</td>
<td>ne</td>
</tr>
<tr>
<td>Trunk:August</td>
<td>1.21 ± 0.19***</td>
<td>−0.26 ± 0.08***</td>
<td>ne</td>
</tr>
<tr>
<td>Roots:August</td>
<td>1.26 ± 0.19***</td>
<td>−0.23 ± 0.08**</td>
<td>ne</td>
</tr>
<tr>
<td>Branches:October</td>
<td>0.88 ± 0.19***</td>
<td>−0.35 ± 0.08***</td>
<td>ne</td>
</tr>
<tr>
<td>Trunk:October</td>
<td>1.26 ± 0.19***</td>
<td>−0.33 ± 0.08***</td>
<td>ne</td>
</tr>
<tr>
<td>Roots:October</td>
<td>0.90 ± 0.19***</td>
<td>−0.33 ± 0.08***</td>
<td>ne</td>
</tr>
<tr>
<td>Branches:non-defoliated</td>
<td>0.05 ± 0.14 (ns)</td>
<td>0.10 ± 0.05*</td>
<td>ne</td>
</tr>
<tr>
<td>Trunk:non-defoliated</td>
<td>0.05 ± 0.14 (ns)</td>
<td>0.02 ± 0.05 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>Roots:non-defolated</td>
<td>0.29 ± 0.14*</td>
<td>0.04 ± 0.05 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>Branches:non-infected</td>
<td>−0.28 ± 0.16*</td>
<td>0.04 ± 0.06 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>Trunk:non-infected</td>
<td>−0.36 ± 0.16*</td>
<td>0.03 ± 0.06 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>Roots:non-infected</td>
<td>−0.39 ± 0.16*</td>
<td>0.03 ± 0.06 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>June:non-defolated</td>
<td>0.31 ± 0.14*</td>
<td>−0.07 ± 0.05 (ns)</td>
<td>0.28 ± 0.29 (ns)</td>
</tr>
<tr>
<td>August:non-defolated</td>
<td>0.14 ± 0.14 (ns)</td>
<td>0.11 ± 0.05*</td>
<td>0.04 ± 0.29 (ns)</td>
</tr>
<tr>
<td>October:non-defolated</td>
<td>0.27 ± 0.14 (ns)</td>
<td>0.01 ± 0.05 (ns)</td>
<td>0.47 ± 0.29 (ns)</td>
</tr>
<tr>
<td>June:non-infected</td>
<td>0.04 ± 0.16 (ns)</td>
<td>0.05 ± 0.06 (ns)</td>
<td>0.15 ± 0.22 (ns)</td>
</tr>
<tr>
<td>August:non-infected</td>
<td>−0.10 ± 0.16*</td>
<td>0.12 ± 0.06*</td>
<td>0.14 ± 0.22 (ns)</td>
</tr>
<tr>
<td>October:non-infected</td>
<td>0.16 ± 0.16 (ns)</td>
<td>0.07 ± 0.06 (ns)</td>
<td>0.59 ± 0.22**</td>
</tr>
<tr>
<td>Non-defoliated:non-infected</td>
<td>ne</td>
<td>ne</td>
<td>0 ± 0.24 (ns)</td>
</tr>
<tr>
<td>June:non-defolated:non-infected</td>
<td>ne</td>
<td>ne</td>
<td>−0.22 ± 0.33 (ns)</td>
</tr>
<tr>
<td>August:non-defolated:non-infected</td>
<td>ne</td>
<td>ne</td>
<td>−0.13 ± 0.33 (ns)</td>
</tr>
<tr>
<td>October:non-defolated:non-infected</td>
<td>ne</td>
<td>ne</td>
<td>−0.72 ± 0.33*</td>
</tr>
</tbody>
</table>
from March to June (except for leaves of defoliated pines) to reach a seasonal peak and then declined in August. Seasonal variation in TNSC was greater in leaves, especially for non-defoliated pines, which showed a more than fourfold decline in TNSC between the June peak and the minimum value in August. Post-drought October TNSC only slightly increased in trunks and leaves (Figure 2). Infection by fungal pathogens was not associated with significant differences in TNSC levels in any organ or month (Table 1).

The ratio of soluble sugars to TNSC (SS : TNSC) differed among tree organs but showed consistent seasonal dynamics across organs (Figure 2, Table 1). The value of SS : TNSC declined slightly from March to June, it increased in August, especially in leaves, and then declined in October. In August, SS : TNSC values were >0.8 for all organs and defoliation classes, indicating that most of the TNSC were in the form of soluble sugars (Figure 2). Starch levels at peak drought were virtually depleted in leaves (i.e., SS : TNSC was nearly 1). Moreover, there were significant differences in SS : TNSC between defoliation classes in some organs and months, whereby defoliated pines tended to show higher SS : TNSC values before the onset of drought but similar or lower values in August (Figure 2, Table 1). No differences in SS : TNSC were observed associated with fungi infection (Table 1).

Throughout the season, root starch tended to be lower in defoliated pines compared with non-defoliated ones (Figure 3). Root rot infection was associated with higher root starch levels in non-defoliated pines, but this effect was not observed in defoliated pines (Figure 3, Table 1). The triple interaction between month, defoliation and infection was statistically significant (Table 1). Interestingly, our results show that root starch concentration in October was virtually depleted in infected, defoliated pines, and it was much lower than in non-infected ones (Figure 3).

Vulnerability curves and hydraulic properties

Roots were more vulnerable to embolism than branches, reaching 50% PLC slightly below −2 MPa, compared with a value close to −3 MPa for branches (Figure 4). This effect was statistically significant (overall P-value for Organ = 0.0062 in a least significant means multiple comparison), although the difference was not significant for all the combinations of defoliation and infections classes (cf. Table 2). The vulnerability to embolism of branches was similar between defoliation classes (Figure 4b, Table 2). In roots, there was no difference between defoliation classes in $P_{50}$, but the slope of the vulnerability curve was steeper in defoliated trees (Table 2). This result implies that the roots of non-defoliated pines would begin to lose conductivity at higher (i.e., closer to zero) water potentials than those of defoliated individuals, but the latter would lose conductivity faster than non-defoliated pines as water potentials declines (Figure 4a). Infection did not affect vulnerability to xylem embolism (Table 2).

No differences in specific hydraulic conductivity ($K_s$) were found between roots and branches (Table 2), possibly due, at least in part, to the large variability in $K_s$ observed for roots.
Likewise, there were no significant differences in either $K_s$, $K_i$ or $A_s : A_i$ between defoliated and non-defoliated pines (Table 2). However, we found lower branch $K_i$ in infected trees (Table 2).

**Water potentials and native embolism**

Shoot water potentials measured in the morning, at the time of sampling for PLC measurements, increased from June to July (Figure 5b, Table 3), consistent with the rainfall events occurring in late June (Figure 1). Afterwards, water potentials declined sharply down to values around $-2$ MPa in August, coinciding with the driest part of the study period. These water potentials were usually between the corresponding predawn and midday shoot water potentials measured in simultaneous sampling campaigns in nearby Scots pine individuals (Figure 5c). We did not find any significant effect of defoliation or infection on water potential (Table 3).

Native PLC varied significantly among sampling dates, following the variation in shoot water potentials. Percentage loss of hydraulic conductivity was particularly low in July, with values around 20%, and it was highest in August, with average values around 65% (Figure 5a, Table 3). In August, six branches (four from non-defoliated, and two from defoliated pines) showed PLC values $>85\%$. Native embolism was not significantly different between defoliation and infection classes (Table 3), except for October when non-defoliated pines showed higher PLC (Figure 5b) ($P = 0.048$).

**Basal area increment and sapwood depth**

We found a marginally significant reduction of BAI associated with defoliation ($P = 0.064$) and this effect tended to be larger in infected trees as shown by the marginally significant ($P = 0.083$) interaction between defoliation and infection (Figure 6a).

Trunk sapwood depth showed patterns similar to those for BAI, but with significantly lower values for defoliated and infected individuals (Figure 6b). Finally, infected trees showed a significantly lower root sapwood depth compared with non-infected trees ($P = 0.039$), regardless of defoliation class, whereas defoliation had no significant effect on root sapwood depth (Figure 6c) ($P = 0.89$).

**Discussion**

Our results show that both high levels of embolism and low carbohydrate reserves occurred in the studied trees during a particularly dry summer. In addition, we show that defoliation was more associated with reduced carbohydrate reserves than with greater hydraulic impairment at the branch level. We also report that drought-induced defoliation and infection by a root rot fungus occur independently, but they both interact to determine the mode of physiological failure in Scots pine at the dry edge of its distribution. We note, however, that some of these results (particularly those regarding pathogen infection) should be interpreted with caution due to the low sample size.

**Defoliation is associated with lower carbon availability, but not with higher hydraulic impairment at the branch level**

We found no consistent differences between defoliated and non-defoliated pines in terms of their vulnerability to embolism according to $P_{50}$ values. This is consistent with previous studies comparing Scots pine populations suffering different levels of drought-induced decline in the same area (Martínez-Vilalta et al. 2012). Although we found steeper vulnerability curves for roots of defoliated pines, which would yield comparatively higher root PLC at water potentials lower than $-2$ MPa (Figure 4a), these conditions occur only during exceptional droughts (Poyatos et al. 2013). In addition, branch-level leaf $K_s$ and $A_s : A_i$ were similar for defoliated and non-defoliated pines, in contrast to different values observed across Scots pine populations within the same study area (Martínez-Vilalta and Piñol 2002) and to the lower $A_s : A_i$ reported at the tree level for defoliated pines (Poyatos et al. 2013). It should be emphasized, however, that our sampling design is representative at the branch level, but not necessarily at the whole-crown level, as we were forced to sample living (as opposed to random) branches in heavily defoliated crowns.
may be increasing over time as a result of repeated droughts (i.e., cavitation fatigue; Hacke et al. (2001)), as has been reported in the case of aspen die-off (Anderegg et al. 2013b).

The decline in PLC observed after a rainy period in July (Figure 5a) suggests some partial embolism reversal (e.g., McCulloh et al. 2011), but we cannot rule out the possibility that sampling artefacts may be causing an overestimation of this PLC recovery (e.g., Sperry 2013). Although our results imply that defoliation may be relatively effective in avoiding further decreases in branch PLC, maximum PLC values were still within a range (−50−90%) associated with canopy dieback in several angiosperms (Hoffmann et al. 2011, Anderegg et al. 2012, 2013b, Nardini et al. 2013) and gymnosperms (Plaut et al. 2012, Klein et al. 2013). Overall, these results also suggest that the steeper declines in whole-plant hydraulic conductance observed for defoliated pines during drought (Poyatos et al. 2013) may occur primarily belowground, as observed in other pine species prior to death (Plaut et al. 2013).

Defoliated pines showed consistently lower TNSC values than non-defoliated pines for most combinations of organ and season, as already observed in early autumn measurements of stem TNSC in other declining Scots pine populations (Galano et al. 2011), including the study site (Poyatos et al. 2013). Interestingly, we did not observe increased TNSC in leaves of defoliated pines, despite their higher assimilation rates (Y. Salmon, J.M. Torres-Ruiz, R. Poyatos, J. Martínez-Vilalta, P. Meir, H. Cochard, M. Mencuccini, unpublished data). Total NSC decreased in all organs during drought, and most dramatically

### Table 3. Summary of the fitted linear mixed models with the percentage loss of conductivity (PLC) and the water potential as response variables. The reference month was ‘May’ in PLC and ‘June’ in Water Potential, the reference defoliation class was ‘Defoliated’ and the reference infestation occurrence was ‘Infected’. The values are the estimate ± SE. Abbreviations: ns, no significant differences; *0.05 < P < 0.1; **0.01 < P < 0.05; ***P < 0.001; ne, not evaluated in the model. Conditional R² values are given for each model.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>PLC</th>
<th>Water potential</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>23.01 ± 10.98*</td>
<td>−1.57 ± 0.13***</td>
</tr>
<tr>
<td>June</td>
<td>10.95 ± 14.96 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>July</td>
<td>−9.95 ± 14.96 (ns)</td>
<td>−0.04 ± 0.19 (ns)</td>
</tr>
<tr>
<td>August</td>
<td>45.23 ± 14.96**</td>
<td>−0.69 ± 0.19***</td>
</tr>
<tr>
<td>October</td>
<td>−12.74 ± 15.00 (ns)</td>
<td>1.14 ± 0.19***</td>
</tr>
<tr>
<td>Non-defoliated</td>
<td>8.60 ± 10.47 (ns)</td>
<td>0.14 ± 0.13 (ns)</td>
</tr>
<tr>
<td>Non-infected</td>
<td>−0.52 ± 11.87 (ns)</td>
<td>0.02 ± 0.15 (ns)</td>
</tr>
<tr>
<td>June:non-defoliated</td>
<td>0.25 ± 14.33 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>July:non-defoliated</td>
<td>0.19 ± 14.33 (ns)</td>
<td>−0.07 ± 0.18 (ns)</td>
</tr>
<tr>
<td>August:non-defoliated</td>
<td>−8.15 ± 14.33 (ns)</td>
<td>0.01 ± 0.18 (ns)</td>
</tr>
<tr>
<td>October:non-defoliated</td>
<td>16.72 ± 14.64 (ns)</td>
<td>−0.30 ± 0.19 (ns)</td>
</tr>
<tr>
<td>June:non-infected</td>
<td>4.42 ± 16.23 (ns)</td>
<td>ne</td>
</tr>
<tr>
<td>July:non-infected</td>
<td>6.51 ± 16.23 (ns)</td>
<td>0.37 ± 0.21</td>
</tr>
<tr>
<td>August:non-infected</td>
<td>−8.16 ± 16.23 (ns)</td>
<td>0.20 ± 0.21 (ns)</td>
</tr>
<tr>
<td>October:non-infected</td>
<td>9.13 ± 16.38 (ns)</td>
<td>−0.05 ± 0.21 (ns)</td>
</tr>
</tbody>
</table>

*Figures 5. Seasonal variation in (a) native embolism expressed as PLC and (b) corresponding water potential, measured in the same branches, of defoliated and non-defoliated Scots pine trees. (c) Predawn and midday water potentials from nearby Scots pine trees from the same population measured on the same dates, where solid lines and symbols indicate defoliated trees and dashed lines and open symbols designate non-defoliated individuals. Error bars indicate ±1 SE. Different letters indicate significant differences (P < 0.05) between sampling dates.*
in leaves, tracking the seasonality of gas exchange (Poyatos et al. 2013).

The observed seasonal variation of the fraction of soluble sugars (SS : TNSC) (Figure 2) is consistent with starch concentration building up prior to bud-break and then decreasing during the summer period (Hoch and Körner 2003, Gruber et al. 2012). During August, the greater mobilization of starch to soluble sugars could be attributed to several causes, including osmoregulation (Sala et al. 2012), the de novo synthesis of carbon-rich compounds into defence against root rot pathogens (Oliva et al. 2012), the energy-dependent process of embolism repair (Brodersen and McElrone 2013) and the growth of new tissue. The two latter processes are consistent with the reduction of native embolism after August, reaching pre-drought values in October (Figure 5a).

Despite the combination of drought and defoliation, our results did not show a complete depletion of TNSC in any of the organs studied (Figure 2), even in the tree that died during the monitoring period (data not shown). Values of trunk TNSC as low as 0.1% were measured in Scots pine 1 year prior to death in a nearby population (Galiano et al. 2011) while our trunk TNSC values were always >0.4%. Experimental studies on conifer saplings (Hartmann et al. 2013b), including pine species (Mitchell et al. 2013), have found evidence for drastic reductions in TNSC (especially starch) in stems and/or roots at mortality, without actually showing completely depleted carbohydrate storage. Phloem impairment may preclude the translocation of NSC and its availability for the maintenance of xylem transport or for the production of defence compounds (Sala et al. 2010, Hartmann et al. 2013a, Sevanto et al. 2014). A recent modelling analysis in the same study system predicts slow, but not disrupted, phloem transport under extreme drought (M. Menecucini, F. Minunno, Y. Salmon, R. Poyatos, J. Martinez-Vilalta, T. Hölttä, unpublished data).

Root rot pathogens exacerbate hydraulic and carbon-related constraints at the tree level

We did not find a clear BAI reduction in Onnia-infected trees (Figure 6a), contrary to other studies reporting growth reductions caused by root rot pathogens from the Armillaria (Fr.) Staude (Cruickshank et al. 2011) and Heterobasidion Bref. genus (Oliva et al. 2012). Although the interaction between infection and defoliation was only marginally significant, the trend towards lower BAI in trees that were both defoliated and infected would be consistent with the reduced sapwood depth observed in these trees (Figure 6b). Hence, post-drought recovery of xylem-specific hydraulic conductivity would be severely constrained in the long-term. Lower sapwood depth in the roots of infected trees (Figure 6c) could be the ultimate consequence of the decay progress, due to the reduction in tree growth caused by the formation of a reaction zone (Oliva et al. 2012). Infection-driven reductions in sapwood depth in roots (and in the trunk of infected, defoliated trees) may also result in lower tissue capacitance and a decreased ability to buffer short-term variations in water potential under drought (McCulloh et al. 2014).

Even though Onnia infection did not affect the vulnerability to embolism, it was associated with reduced $K_L$, particularly in defoliated pines (Table 2). Hence, infection by a root rot pathogen likely exacerbated hydraulic constraints through effects on growth and sapwood depth but also more directly through its impact on hydraulic conductivity.

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Figure 6. Basal area increment (BAI) (a), trunk sapwood depth (b) and root sapwood depth (c) as a function of infection and defoliation classes. Different uppercase letters indicate significant differences ($P < 0.05$) between infestation occurrence, and different lowercase letters indicate significant differences between defoliation classes within a given infection class. Note that the interaction between defoliation and infection in the BAI model was marginally significant ($0.05 < P < 0.1$). Error bars ± 1 SE.
Our results show a complex picture of the root rot infection effects on NSC pools. We did not find any evidence for consistent reductions in TNSC across tree organs associated with *Onnia* infection. However, for non-defoliated pines, infection appeared to drive starch accumulation in roots, rather than depletion, possibly as a response to increased sink strength in the roots associated with higher C demand (de novo synthesis of defence compounds; Oliva et al. 2014). Presumably, defoliated pines were so severely carbon-limited that this starch sink in the roots could not be maintained, and they ended the year with extremely low levels of starch in roots (Figure 3). These findings are in line with a recently proposed framework which postulates that the net effect of the infection by a necrotrophic pathogen, such as *Onnia*, is strongly dependent on tree C availability and the timing of drought events (Oliva et al. 2014).

**A complex pathway to mortality**

In this final section we synthesize our current understanding on the process of drought-induced mortality in the study area, from the perspective of the comparison of coexisting defoliated and healthy individuals, using the information gathered here and in other studies conducted at the same site (Figure 7). Drought-induced defoliation is associated with drier microenvironments (Vilà-Cabrera et al. 2013) and with steeper reductions in whole-plant hydraulic conductance during seasonal drought (Poyatos et al. 2013). Hydraulic constraints may be related to higher vulnerability to embolism in roots (steeper vulnerability curves) and can be magnified by cavitation fatigue following repeated droughts. Defoliation and prolonged periods of near complete stomatal closure both contribute to reduce NSC in defoliated trees (Poyatos et al. 2013). Defoliated trees appear to enter a death spiral in which reduced C assimilation constrains radial growth (Heres et al. 2012) and crown development (Poyatos et al. 2013). Root rot fungi may further damage hydraulic function through direct effects on sapwood depth and cumulative growth reductions. Moreover, the demand for C-rich compounds for osmoregulation, hydraulic repair and defence against root rot infection may contribute to the depletion of C reserves in defoliated pines, possibly increasing the minimum C threshold for tree survival and hence accelerating tree mortality (Oliva et al. 2014). It remains to be established whether...
the framework outlined in Figure 7, which has been developed for only one species in a given region, can be applied to other species and study systems. Overall, our study reflects the intertwined nature of physiological mechanisms leading to drought-induced mortality (McDowell et al. 2011) and the inherent difficulty of isolating their contributions under field conditions.

Supplementary data
Supplementary data for this article are available at Tree Physiology Online.

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Conflict of interest
None declared.

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