



# Heat Waves Alter Carbon Allocation and Increase Mortality of Aleppo Pine Under Dry Conditions

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Birami B, Gattmann M, Heyer AG, Grote R, Arneth A and Ruehr NK (2018) Heat Waves Alter Carbon Allocation and Increase Mortality of Aleppo Pine Under Dry Conditions. Front. For. Glob. Change 1:8. doi: 10.3389/ffgc.2018.00008 Climate extremes are likely to occur more frequently in the future, including a combination of heat waves and drought. However, the responses of trees to combined stress and their post-stress recovery are not fully understood yet. Therefore, this study investigated the responses of semi-arid Pinus halepensis seedlings to moderate drought, heat and combined heat-drought stress, as well as post-stress recovery. The seedlings were grown under controlled conditions and exposed to two 4-days-long heat periods, reaching air temperature maxima of 42°C and vapor pressure deficit (VPD) of 7 kPa. Day- and nighttime canopy gas exchange was measured and differences in shoot and root allocation of non-structural carbohydrate (NSC) compounds (soluble sugars, starch, cyclitols, and carboxylic acids) assessed. Fluorescence parameters, nitrate levels, proline content and shoot water potential ( $\psi$ ) provided additional indicators for stress severity and recovery performance. During the heat periods, net photosynthesis and stomatal conductance decreased immediately. This decline was modest under well-watered conditions, with transpiration and dark respiration rates remaining high and despite reductions in root NSC content, trees recovered following heat release. This was not the case in the heat-drought treatment, where stress resulted in high mortality rates and the few surviving seedlings showed reduced gas exchange rates and low root NSC content, while leaf nitrate and proline remained elevated even 3 weeks after heat release. Shoot  $\psi$  indicated that hydraulic failure was not the reason for mortality in the heat-drought seedlings. Instead, we argue that low transpiration rates, which resulted in needle temperatures >47°C during heat stress (6°C above air temperature) have caused irreversible damage. In summary, it could be demonstrated that heat waves in combination with moderate drought can either result in increased mortality or, if the seedlings survive, in delayed recovery. This highlights the potential of an increase in heat wave temperatures to trigger forest decline in semi-arid regions.

Keywords: carbon allocation, combined stress, cyclitols, extreme heat wave, nitrate, non-structural carbohydrates, proline, recovery

# INTRODUCTION

Forest dieback related to climate extremes has been observed in most regions of the world (Allen et al., 2010; Anderegg et al., 2012, 2015). In particular heat waves combined with drought, which are increasing in frequency and duration (Meehl and Tebaldi, 2004; Schär, 2015), could be a major trigger of tree mortality (Williams et al., 2012; Allen et al., 2015), yet our understanding of physiological processes within trees to such extremes is scarce.

Trees tightly regulate stomatal conductance  $(g_s)$  to balance water supply and water loss. The close coordination between leaf water potential ( $\psi$ ) and  $g_s$  during drought conditions has been reported manifold (Jarvis, 1976; Burghardt and Riederer, 2003; Ripullone et al., 2007; Klein and Niu, 2014; Anderegg et al., 2018), while the responses of  $g_s$  to changes in evaporative demand, particular in combination with high temperatures is less clear. During experimental heat wave conditions,  $g_s$  has been found to decline (Tatarinov et al., 2015; Duarte et al., 2016; Ruehr et al., 2016; Drake et al., 2018) or to be not affected (Ameye et al., 2012) or even to increase (Urban et al., 2017a,b). Despite such differences in stomatal responses, most heat wave studies have found transpiration (*E*) to increase under high soil water content. At first, this seems counterintuitive, but can be explained by increases in evaporative demand, which stimulates E and thus is cooling of the leaf.

Increased E causes water-use efficiency (WUE) to sharply decrease, because photosynthesis typically declines at high temperatures (Ameye et al., 2012; Tatarinov et al., 2015; Duarte et al., 2016; Ruehr et al., 2016; Urban et al., 2017b; Drake et al., 2018), which results in a pronounced carbon-water decoupling during heat waves (Drake et al., 2018). The underlying reasons for the strong decline of photosynthesis under heat stress are manifold and include decreases in gs (Ruehr et al., 2016), reduced enzymatic activity (Haldimann and Feller, 2004; Rashid et al., 2018), inactivation of integral proteins and increased membrane leakage (Quinn, 1988; Hays et al., 2001; Zhang and Sharkey, 2009; Watson, 2015), or a combination of influences (Duarte et al., 2016). In addition, increased photorespiration (Teskey et al., 2015) and mitochondrial respiration decreases photosynthetic efficiency. Temperature thresholds resulting in permanent photosynthetic and whole-leaf damage are reported to range between 40 and 50°C (Colombo and Timmer, 1992; Hüve et al., 2011; O'sullivan et al., 2017; Niinemets, 2018; Rätsep et al., 2018), whereas the exact threshold depends on species, the duration of exposure, water availability and adaptive metabolic responses.

Besides direct tissues damage, temperature increments affect tree C allocation dynamics (Zhao et al., 2013; Ruehr et al., 2016). Because respiration typically increases with temperature,

C loss can become larger than C uptake (Zhao et al., 2013) and NSC reserves in trees may deplete. In addition, shoot-toroot allocation patterns are sensitive to stress. While reduced C transport from shoots to roots under drought conditions has been observed (Ruehr et al., 2009; Zang et al., 2014; Blessing et al., 2015), C allocation toward roots can increase under high temperatures (Blessing et al., 2015). However, it remains unclear how such changes in C allocation may affect the resilience of trees. For example, the formation of specific C compounds, such as proline or sugar alcohols can prevent protein denaturation at high temperatures (Hamilton, 2001; Jaindl and Popp, 2006). The amino acid proline can also act as osmolyte, antioxidative defense and membrane stabilizing molecule (Verbruggen and Hermans, 2008; Szabados and Savouré, 2010). In addition, large NSC reserves will be critical to sustain high respiration rates. This indicates a potentially important role of the primary C metabolism in mitigating long-term damages and hence could influence post-stress recovery. It can be speculated that besides actual tissue damage, the amount of C maintained during stress could also be a driving force of repair mechanisms (Galiano et al., 2017).

Following stress release, the ability of trees to recover from heat waves depends on stress severity, which is directly influenced by heat exposure (Hüve et al., 2011). If heat waves are additionally combined with drought, evaporative cooling is diminished and canopy temperature increases (Scherrer et al., 2011). This should then accelerate stress-induced damage and potentially delays post-stress recovery. Semi-arid regions are at particular risk because air temperatures are already high so that even a slight warming can surpass critical levels, especially when evaporative cooling is low (Rotenberg and Yakir, 2010; Liu et al., 2013). Regeneration in semi-arid forests might especially be jeopardized because seedlings are most vulnerable to desiccation due to low rooting depth (limited water supply) and exposure to particularly high temperatures (Kolb and Robberecht, 1996).

The present study investigates the impacts of high temperatures under well-watered and dry conditions on seedlings of Pinus halepensis, originating from one of the driest pine forest plantations in the world, located in the Negev desert (Rotenberg and Yakir, 2010). In this region, short heat waves frequently occur that already affects the forest's carbon and water cycling (Tatarinov et al., 2015). For future conditions, it is expected that temperatures are increasing further and precipitation during winter is decreasing (Giorgi and Lionello, 2008; Tabari and Willems, 2018). Thus, the trees might be pushed beyond their stress tolerance limits and the survival of this unique forest might be at risk. In order to increase understanding of Aleppo pine's responses to expected extreme heat waves we investigated shoot gas exchange, fluorescence parameters, and some primary metabolites in shoots and roots in a greenhouse experiment under controlled environmental conditions. In particular the following research questions were addressed: (1) How does shoot gas exchange and water-use efficiency respond to a simulated heat waves with or without drought and do  $g_s$ , photosynthesis and E also depend on vapor pressure deficit? (2) What are the main drivers of metabolite dynamics and do they affect the allocation between shoots and

**Abbreviations:**  $A_{net}$ , light-phase carbon assimilation/net photosynthesis; C, fixed carbon; *E*, transpiration;  $E_{day}$ , light-phase transpiration;  $g_s$ , stomatal conductance; NSC, non-structural carbon;  $\Pi$ , osmotic potential; PAR, photosynthetic active radiation;  $\psi$ , leaf water potential;  $R_{dark}$ , dark respiration; RH, relative air humidity; RSW, relative soil water content; TCA, Tri-carboxylic acid cycle; VPD, vapor pressure deficit; WUE, water-use efficiency; WUEa, apparent water-use efficiency; WUEi, intrinsic water-use efficiency.

roots? (3) Are heat and heat-drought stress responses fully reversible post-stress?

# MATERIAL AND METHODS

# Plant Cultivation and Initial Pre-treatment Conditions

Pinus halepensis (Miller) seedlings were grown from seeds in a scientific greenhouse facility in Garmisch-Partenkirchen, Germany (732 m a.s.l., 47°28′32.87″N, 11°3′44.03″E). The origin of the seed material is a 50-years-old Aleppo pine plantation in Israel (Yatir forest). Cones of trees were sampled growing in close-proximity to a meteorological station and flux tower (IL-Yat, 650 m a.s.l., 31°20′49.2″N, 35°03′07.2″E). About 2-4 weeks after germination of seedlings in Germany, the seedlings were transferred to 1 L pots each, containing a mixture (2:8) of quartz sand (0.7 and 1-2 mm) and vermiculite (ca. 3 mm) with 2 g of slow-release fertilizer (Osmocote<sup>®</sup> Exact + TE 3-4months fertilizer 16-9-12 + 2MgO + TE, Everis International B.V., Heerlen, The Netherlands). Three months before starting the experiment, 7 months-old seedlings were planted in larger pots (2.5 L) containing a mixture of 1:2:1 quartz sand (1–2 mm): quartz sand (Dorsolit 4-6 mm): vermiculite (3 mm) with 6 g of slow-release fertilizer added (Osmocote® Exact + TE 5-6 months fertilizer 15-9-12 + 2MgO + TE). Potted seedlings were irrigated regularly, starting with 0.1 L per week, which was later adapted to meet increased water demand of the growing seedlings to 0.15 L per week.

Seedlings were grown under average light intensity of 693  $\pm$  324  $\mu$ mol m $^{-2}s^{-1}$  (supplemented by sodium vapor greenhouse lamps, T-agro 400 W, Philips) and temperatures and humidity adapted to 10-years monthly-averaged day and night air temperature and relative humidity measured at the Yatir forest site (see **Table S1**). One month before the start of our experiment, seedlings were gradually acclimated from average daytime temperatures of  $18^{\circ}$ C (night:  $12^{\circ}$ C) to  $25^{\circ}$ C (night:  $15^{\circ}$ C). This is close to the 10-years average measured at Yatir forest during May, when the occurrence of short heat waves is typically observed (Tatarinov et al., 2015).

Seedlings were assigned randomly either to a control, drought, heat or heat-drought treatment with 30 seedlings per group. To maintain ambient air temperature and relative humidity in the control and drought treatment, while conducting heat waves in the heat and heat-drought treatment, the seedlings were placed in two adjacent, but individually controllable compartments of the greenhouse facility.

## **Environmental Variables**

Air temperature and relative humidity sensors (CS215, Campbell Scientific Inc., Logan, Utah, US, enclosed in aspirated radiation shields type 43502, Young, Traverse City, MI, USA) and photosynthetic active radiation (PAR) (PQS 1, Kipp & Zonen, Delft, The Netherlands) were measured continuously at canopy height in each greenhouse compartment. Carbon dioxide concentration was monitored with a CO<sub>2</sub> probe (GMT 222, Vaisala, Helsinki, Finland). Soil water content was measured automatically in 10 pots per treatment using substrate-specific

calibrated probes (10HS, Decagon Devices, Inc., WA, USA). All environmental sensors produced half-hourly data that were recorded with data loggers (CR1000, Campbell Scientific, Inc. USA).

Soil moisture is given as relative soil water content (RSW) as follows:

$$RSW = 100*\frac{(SWC_{\text{sample}} - SWC_{\text{min}})}{(SWC_{\text{max}} - SWC_{\text{min}})}$$
(1)

where SWC<sub>sample</sub> is the actual volumetric soil water content, SWC<sub>min</sub> is the minimum volumetric soil water content after drying the soil for 48 h at 60°C. SWC<sub>max</sub> is the soil water content at maximum water holding capacity, which was about 200 cm<sup>3</sup> L<sup>-1</sup> (20% v/v) for our substrate.

## **Experimental Conditions**

Aleppo pine seedlings exposed to heat and heat-drought treatments were subjected to two heat periods of 4 days each (April 27th-April 30th and May 7th-May 10th, 2016). We simulated naturally occurring heat waves by a gradual increasing temperature throughout 2 days, reaching a maximum during the following 2 days (Tatarinov et al., 2015). In order to test how projected further temperature increases of 2-6°C in the Mediterranean region (Seneviratne et al., 2012) might affect metabolism and survival of tree seedlings, we chose to set 5°C higher air temperature maxima than have been reported so far (Tatarinov et al., 2015). This resulted in a maximum of 43.1°C during the first heat wave and 42.4°C during the second heat wave. Relative humidity (RH) during the heat waves was kept between 20 and 40%, which is similar to air water content under ambient temperature conditions. This resulted in a pronounced increase of VPD up to 7.5 kPa during the heat periods (Figures 1A,B) which corresponds to observations in the Yatir forest (Tatarinov et al., 2015). In total, extreme air temperatures above 40°C and VPD > 6 kPa were maintained for 10h during the first heat wave and 12h during the second heat wave. Dark-phase (PAR < 10  $\mu$ mol m<sup>-2</sup>s<sup>-1</sup>) temperatures reached 33.0°C during the first- and 32.9°C during the second heat wave with a total exposure time of 13 h above 30°C for each heat wave. Outside light was supplemented by sodium vapor lamps, resulting in an average PAR of 416  $\pm$  105  $\mu$ mol m<sup>-2</sup>s<sup>-1</sup> during the course of the experiment (Figure 1C). Irrigation of the seedlings was adapted to maintain targeted soil water availability. During the initial control conditions and during the recovery at the end of the experiment, seedlings were watered to a RSW of 40-50% (about 0.15 L three-times per week), which is similar to the average wet season condition of the Yatir forest site (pers. com. Yakir Preisler). One week before the start of the first heat wave, irrigation was withheld in the drought and heat-drought treatment (for 4 days) until a RSW of about 15% was reached (Figure 1D). This RSW was maintained by watering with 0.05 L three-times per week. One week after the end of the second heat period, all plants were irrigated to reach similar RSW conditions of 40-50%.

In order to avoid a possible influence on the seedlings from the location within the greenhouse, the position of the



seedlings were changed randomly every 2 weeks during the initial growth phase. During the experimental phase which took place in two separate greenhouse compartments, the seedling position was unchanged because automated shoot cuvettes for gas exchange measurements (see section Gas Exchange Measurements) and soil moisture sensors were permanently installed. During this phase the seedlings were placed spatially interspersed in a randomized block design. Differences in daily-averaged RH between the greenhouse compartments were below 3% ( $\Delta$ PAR <6%) when heat waves were not applied. Air temperature differences were  $\sim$ 1°C before the first heat wave and 0.2°C during the remainder of the experiment.

## **Gas Exchange Measurements**

Net photosynthesis  $(A_{net})$ , night respiration  $(R_{dark})$  and light phase transpiration  $(E_{day})$  of *Pinus halepensis* shoots were measured with an automated cuvette system (Duarte et al., 2016; Bamberger et al., 2017). Highly UV-transmissive acrylic glass tubes (PMMA Saalberg, 30L: 18W) were placed around tree shoots (n = 4 per treatment). The bottom side of each cuvette was sealed by an acrylic glass cap, which allowed the insertion of the seedling. Remaining gaps between tree stem and cap were sealed using plastic putty (Teroson, Düsseldorf, Germany). Each cuvette was supplied with a photodiode for PAR spectrum (G1118, Hamamatsu Photonics, Hamamatsu, Japan) and cross-calibrated with high-precision PAR sensors (PQS 1, Kipp & Zonen, Delft, the Netherlands). For temperature measurements, each cuvette was equipped with a previously calibrated thermocouple (5SC-TTTI-36-2M, Newport Electronics GmbH, Deckenpfronn, Germany). Air mixing within the cuvettes was maintained by a fan (412 FM, ebm-papst GmbH & Co. KG, Mulfingen, Germany). The cuvettes were measured sequentially for 10 min each. After the distal cap had closed automatically, reference air of known CO2 and H2O concentration at a rate of 5L min<sup>-1</sup> was supplied to the cuvettes. The flow rate was adjusted by a digital mass flow controller (F-201CZ-10K, Bronkhorst, Ruurlo, Netherlands). The measurement air (reference) was generated by an oil-free compressor (SLP-07E-S73, Anest Iwata, Yokohama, Japan) with an Ultra Zero Air generator (Ultra Zero Air GT, LNI Schmidlin SA, Geneva, Switzerland). CO2 and water vapor (nebulizing evaporation pump, LCU Liquid Calibration Unit, Ionicon, Innsbruck, Austria) was supplied at a constant rate to the air stream, resulting in a CO<sub>2</sub> concentration of 438  $\pm$  3  $\mu$ mol mol<sup>-1</sup> and a water vapor concentration of 6.5  $\pm$  0.1 mmol mol<sup>-1</sup> on average during the experiment.

The slight overpressure that was deliberately generated during each measurement prevented outside air from entering the system. The resulting sample air stream was  $1-3 \text{ L} \text{ min}^{-1}$ . Concentration changes between reference and sample air (each provided with  $0.5 \text{ L} \text{ min}^{-1}$ ) were measured with a

LI-7000, which was connected to a LI-840 (both LI-COR Inc., Lincoln, NE, USA) for absolute concentration measurements of the reference air stream. The sample cells of the two instruments were all supplied with an air stream of  $0.5 \,\mathrm{Lmin^{-1}}$  each. The LI-840 and LI-7000 were zero and span calibrated before the start of the experiment and the two measurement cells of the LI-7000 were matched on a weekly basis.

As an additional quality check, one cuvette per greenhouse compartment was left empty and allowed detecting any offset between the reference and sample air not caused by plant gas exchange. The concentration differences were small (CO<sub>2</sub>: 0.4 [-0.2, 1.2]  $\mu$ mol mol<sup>-1</sup> in median with lower and upper quartiles; H<sub>2</sub>O 0.07 [0.05, 0.12] mmol mol<sup>-1</sup> with lower and upper quartiles) and were removed by subtraction upon data analysis. After each measurement cycle, the system was flushed with reference air for 1 min. For flux calculation the last 180 s per measurement were used if the following criterion for stability was met: change in [CO<sub>2</sub>] < 0.5  $\mu$ mol s<sup>-1</sup> and change in [H<sub>2</sub>O] < 0.5 mmol s<sup>-1</sup>. In total, 94% of the measurements were used for flux calculations.

Gas exchange rates (**Figure 2**) were calculated following an open chamber approach. In brief,  $(E_{dav})$  was derived as follows:

$$E_{\text{day}} = \frac{\dot{m} \Delta W}{\text{Area}_{\text{leaf}} * \left(1 - \frac{W_{\text{sample}}}{1,000}\right)}$$
(2)

Where  $\dot{m}$  is mass flow [mol s<sup>-1</sup>] into the cuvettes,  $\Delta W$  the difference of water vapor in reference- and sample air stream [mol mol<sup>-1</sup>],  $W_{\text{sample}}$  the water vapor concentration of the sample air [mol mol<sup>-1</sup>] and Area<sub>leaf</sub> [m<sup>2</sup>] the half-sided needle area of the shoot.

 $\rm CO_2$  gas exchange fluxes separated into  $A_{\rm net}$  and  $R_{\rm dark}$  were calculated as:

$$A_{\text{net}}(R_{\text{dark}}) = -\frac{\dot{m}*\Delta \text{CO}_2}{\text{Area}_{\text{leaf}}} - \frac{\text{CO}_2 \text{ sample}*\text{E}}{1,000}$$
(3)

With  $\Delta CO_2$  as the difference in [CO<sub>2</sub>] between reference and sample air stream [mol mol<sup>-1</sup>], CO<sub>2sample</sub> carbon dioxide concentration of the sample air stream [mol mol<sup>-1</sup>].

 $g_s$  was calculated as:

$$g_{\rm s} = \frac{\mathrm{E}\left(1,000 - \frac{W_{\rm leaf} + W_{\rm sample}}{2}\right)}{W_{\rm leaf} - W_{\rm sample}} \tag{4}$$

$$W_{\text{leaf}} = \frac{e_{\text{sat leaf}}}{p} *1,000 \tag{5}$$

With stomatal conductance  $g_s$  in mol m<sup>-2</sup>s<sup>-1</sup>, saturated vapor pressure of the leaf with  $e_{sat} = e_{sat}$  in bar and atmospheric pressure p in kPa. Duarte et al. (2016) showed that the ventilation generated by the fans in the cuvettes allowed well air mixing and thus boundary layer for the calculation of  $g_s$  could be neglected. In order to determine changes in WUE during heat stress, apparent water-use efficiency  $WUE_a$  and intrinsic wateruse efficiency  $WUE_i$  were derived using the following equations:

$$WUE_{a} = \frac{A_{net}}{E_{day}}$$
(6)

$$WUE_{i} = \frac{A_{net}}{gs}$$
(7)

 $WUE_a$  is driven by  $CO_2$  assimilation and water loss at the leaf level, and thus provides a measure for the water cost of carbon fixation.  $WUE_i$  provides a measure for leaf physiological changes and is used to characterize photosynthetic efficiency independently of environmental drivers (Bonan et al., 2014; Wieser et al., 2018).

#### Chlorophyll Fluorescence

A portable photosynthesis system (LI-6400XT, LiCor Bioscience, Lincoln, NE, USA) supplemented with a fluorescence head (6400-40 Leaf Chamber Fluorometer, LiCor Bioscience, Lincoln, NE, USA) was used to measure chlorophyll fluorescence on 4 seedlings per treatment before the start of the treatments, just 1 day after the second heat wave had ended and 3 weeks later during recovery. Measurements were conducted between 9 a.m. and 1 p.m. at a leaf temperature of 25°C and reference  $[CO_2]$  of 400  $\mu$ mol mol<sup>-1</sup>. Needles were clamped into the leaf cuvette fully covering the cuvette area (2 cm<sup>2</sup>). Needles were acclimated to actinic light with optimal PAR (1,200 µmol  $m^{-2}s^{-1}$ , pre-determined from light response curves) with a blue light proportion of 10% for several min. As soon as changes in fluorescence intensity (F) ceased to values <5, F was considered stable and the measuring sequence was initiated. First, chlorophyll fluorescence at actinic light (F<sub>c</sub>) was measured as a steady state value, then needles were briefly exposed to a saturating flash of >7,000  $\mu$ mol m<sup>-2</sup>s<sup>-1</sup> and maximum fluorescence (F'm) was measured. This was followed by a fast switch from actinic light to far red radiation. This socalled dark pulse allowed measurement of minimum chlorophyll fluorescence  $(F'_0)$ . The photochemical parameters effective photosystem II quantum yield ( $\Phi_{PSII}$ ), maximum light adapted quantum yield of the photosystem II  $(F'_v/F'_m)$ , coefficients of photochemical fluorescence quenching (qP) and relative electron transfer rate (ETR) were calculated using standard procedures.

#### **Biomass Sampling and Sample Preparation**

Entire seedlings were sampled on three occasions during the course of the experiment: pre-stress (April 20th), at the last day of the second heat wave (Mai 10th) and 3 weeks later at the end of the recovery period (Mai 30th). During each sampling campaign, a minimum of six seedlings per treatment (including dead seedlings, see below) were sampled according to a randomized block design. All plant material containing needles above the former cotyledons was assigned as shoot, all plant material below the first roots as root material. Stems were collected but were not used in this study. Additionally, all plant samples were weighted, photographed, and immediately frozen



in liquid nitrogen. Time of harvest was between 1 and 2 p.m. Prior to freezing, water potential of shoots ( $\psi_{shoot}$ ) was measured on some seedlings using a pressure chamber (Model, 600 PMS-Instruments, Albany, OR, USA). Roots and shoots were ground to fine powder in liquid nitrogen with and samples were divided into aliquots and stored at  $-80^{\circ}$ C for further analysis. Dry weight was assessed gravimetrically by drying samples for 48 h at 60°C. In total, 23 control, 22 drought, 28 heat, and 28 heat-drought treated seedlings were sampled. For each sampling campaign metabolites were analyzed in root and shoot tissues and averaged per treatment.

# **Seedling Mortality**

Mortality was determined by two criteria. The first was that foliage appeared dry and had lost most of their greenness (for example see **Figure S1**) while the second criterion was based on absolute shoot water content. In seedlings that looked quite healthy and green, absolute shoot water content was between 58 and 70%, while shoot water content in the other seedlings ranged from 8.2 to 46.7% (**Figure S2**) However, we like to highlight that the decline in shoot water content was unlikely the reason of death (e.g., seedlings of the well-watered heat treatment died as well, but  $\psi_{\text{shoot}}$  did not indicate water stress in the surviving seedlings, see **Figure 3**), but rather a consequence: after the shoots were permanently damaged and the majority of needles

and/or roots no longer functional, water supply ceased and the shoots began to desiccate. The stem desiccation phenomenon was supported by dendrometer data (**Figure S3**). Here, stem dehydration indicated by diameter shrinkage of dying seedlings was observed to occur after the heat waves. Separating between living and dead seedlings was a necessary prerequisite before analyzing the gas exchange and metabolite data.

# Analysis of Carbohydrates

Analysis of NSC in shoot and root samples was done following the approach reported by Brauner et al. (2014) based on extraction using boiling ethanol (80%v/v), which was recently defined as standard method for sugar extraction (Quentin et al., 2015). Extracts were analyzed by HPLC (Dionex DX-500 HPLC system, Thermo Scientific) using pulsed amperometric detection by a gold electrode (Dionex ED Au, Thermo Scientific). Soluble sugar compounds (inositol, pinitol, glucose, fructose, sucrose) were separated by a PA1 column (Dionex CarboPac PA-1, Thermo Scientific). Myo-inositol and D-pinitol were co-eluted by our chromatographic method and therefore are referred to cumulative as cyclitols. Glucose, fructose and sucrose are treated together as soluble NSC because single components responded in the same direction to the treatments applied. Remaining extracts were stored at  $-20^{\circ}$ C.



**FIGURE 3** | Shoot water potential ( $\psi_{shoot}$ ) in control, drought, heat, heat-drought trees is given before stress (before), after the second heat wave (stress) and 3 weeks after stress (recovery). The error bars are  $\pm 1$  SE (n = 6, but heat-drought treatment during stress and recovery n = 3). Significant treatment effects are given \*(ANOVA,  $P \le 0.05$ ).

Starch extraction was carried out with a modified approach suggested by Hoch et al. (2003). Remaining pellets from soluble sugar extraction were suspended in 1 ml of deionized water (H<sub>2</sub>Odd), heated to 95°C for 45 min and then cooled to 30°C before 1 ml of amyloglucosidase reagent (10 mM acetate Buffer pH 4.5, 1 mg ml<sup>-1</sup> amyloglucosidase) was added. Glucose-oxidase reagent was added containing 2 U ml<sup>-1</sup> horse radish peroxidase, 5 U ml<sup>-1</sup> glucose oxidase and 0.1 mg ml<sup>-1</sup> o-Dianisidin. Samples were kept at 30°C for 2 h, then the reaction was stopped with 5 N HCL and absorption was measured at 540 nm photometrically (Ultrospec 2000 UV/VIS Spectrophotometer, Pharmacia Biotech).

Analysis of carboxylic acids (malic-, fumaric-, citric acid), extracted into hot water, was done as reported by Brauner et al. (2014), again using HPLC (Dionex DX-500 HPLC system, Dionex IonPac AS11-HC, Thermo Scientific) coupled to a suppressor (Dionex AERS 500 Carbonate Electrolytically Regenerated Suppressor, Thermo Scientific) with detection by conductivity (Dionex ED, Thermo Scientific). On the same chromatogram, nitrate, phosphate, sulfate, and chloride were detected as well.

#### Proline

Proline in shoot and root tissues was analyzed following Rienth et al. (2014). In brief, 25 mg of frozen sample powder was extracted for 10 min in 1 ml H<sub>2</sub>Odd at 4°C and centrifuged (13,000 rpm for 15 min). 500  $\mu$ l of supernatant was added to 500  $\mu$ l of concentrated Formic acid and vortexed for 5 min. 500  $\mu$ l of Ninhydrin reagent was added (3% Ninhydrin in Dimethylsulfoxide). Samples were heated (100°C for 15 min) and immediately cooled on ice and then centrifuged (at 13,000 rpm for 1 min). Absorption of the supernatant was measured at 520 nm (Ultrospec 2000 UV/VIS Spectrophotometer, Pharmacia Biotech).

# **Needle Surface Temperature**

Needle surface temperature was measured optically by an infrared camera system (PI 450, Optris, Germany) and analyzed using the manufacturer's software. Recordings were taken at the last day of the second heat wave between 1 and 4 p.m. of at least four individuals per treatment. Air temperature during the measurements was used to correct for background radiation. Emissivity of *P. halepensis* needles was set to 0.97 according to Monod et al. (2009).

## **Calculation of Osmotic Pressure Potential**

Osmotic pressure potential was calculated using the van't Hoff osmotic pressure equation for aqueous mixed electrolyte solvents:

$$\Pi = M * i * R * T \tag{8}$$

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where  $\Pi$  is the osmotic pressure potential Pa (kg m<sup>-1</sup> s<sup>-1</sup>), M is the molar fraction of all measured osmotic active components (cumulative) including total C, proline, nitrate, sulfate, and phosphate concentrations. R is the ideal gas constant (8.31445 kg m<sup>2</sup> mol<sup>-1</sup>K<sup>-1</sup>s<sup>2</sup> at standard temperature of 25°C). The van't Hoff factor i represents the degree of dissociation of each solute component. We calculated with fully dissociated solute components for the case that a salt would dissociate into two ions (e.g., i<sub>glucose</sub> = 1, i<sub>nitrate</sub> = 2).

#### **Statistics and Data-Analysis**

Treatment effects on metabolite concentrations and differences between treatments were analyzed separately per sampling period (pre-stress, stress and recovery). Analysis of variance (ANOVA) was used to detect treatment effects and a subsequent *posthoc* test (Tukey's honest significant difference test) was used to reveale differences between treatment groups. Samples that were considered as "dead" were excluded from the analysis. The effect size of treatments compared to the control was calculated as a percent treatment effect (*D*):

$$D = 100*\frac{(\text{mean}_{\text{t}} - \text{mean}_{\text{c}})}{\text{mean}_{\text{c}}}$$
(9)

where mean<sub>t</sub> is the treatment average and mean<sub>c</sub> is the average of the control, the standard error (SE) of the treatment effect ( $D_{SE}$ ) was calculated as follows:

$$D_{\rm SE} = 100* \sqrt{\left(\left(\frac{1}{\rm mean_c}\right)*SE_t\right)^2 + \left(\left(\frac{\rm mean_t}{\rm mean_c^2}\right)*SE_C\right)^2}$$
(10)

with SE<sub>t</sub> and SE<sub>c</sub> are the SE of treatments or control. Because shoot gas exchange was measured on the same seedlings throughout the experiments, treatment effects were assessed by application of linear mixed-effects models, which account for the repeated sampling-in-time design. All data processing and statistical analysis was done using R version 3.2.2 (R Core Team, 2015), extended with the "lme4" package (Bates et al., 2015) for linear mixed-effects models.

**TABLE 1** | Comparison of needle temperature (T<sub>needle</sub>) between treatments.

T <sub>needle</sub> [°C]	SE	$\Delta^{\circ}$ C (T <sub>needle</sub> – T <sub>air</sub> )
28.6	0.63 [ <i>n</i> = 6]	3.4
30.7	0.33 [ <i>n</i> = 4]	5.2
45.3	0.75 [n = 6]	3.1
47.2	0.57 [ <i>n</i> = 4]	5.7
	28.6 30.7 45.3	$\begin{array}{c} 28.6 \\ 30.7 \\ 45.3 \end{array} \begin{array}{c} 0.63 \ [n=6] \\ 0.33 \ [n=4] \\ 0.75 \ [n=6] \end{array}$

Measurements were done using an infrared camera at the end of the second heat wave when air temperature was 40.8–42.1°C in the heat and heat-drought treatment and 25.1–25.4°C in the control and drought treatment. Shown are treatment averages, standard error ( $\pm 1$  SE) and temperature difference between needle and air temperature.

# RESULTS

## **Stress Intensity and Mortality**

Heat waves resulted in pronounced death and caused 5 seedlings (23%) in the heat treatment and 15 seedlings (68%) in the heat-drought treatment to die. The surviving seedlings showed only a moderate reduction in  $\psi_{shoot}$  (Figure 3). Thus, it is unlikely that hydraulic failure was a cause of mortality. However, needle temperatures were affected differently by the treatments. While needle temperatures in the drought and heat-drought treatment were on average 2.6°C higher than in the control and heat treatment (control: +3.4°C above ambient, heat:  $+3.1^{\circ}$ C, drought:  $+5.2^{\circ}$ C, heat-drought:  $+5.7^{\circ}$ C; Table 1). Thus, shoots under heat combined with drought had absolute needle temperature >47°C. Although the exact time point of mortality was not investigated in each of the seedlings, the continuous gas exchange measurements pointed to the end of the first heat wave (Figure 2 intermitted lines) as daytime gas exchange rates did not recover, but declined even further. Shoot Rdark reached values close to zero  $\sim 1$  week after the end of the last heat wave (Figure 2C), which coincides with a halt of the continuous diameter development (Figure S3) and indicates that the seedlings were dead. Since the seedlings continued to dry out after death, the shoot water content was below 49% at biomass sampling 2 weeks later, which was in agreement with the death criterion (see Methods section).

## Shoot Gas Exchange and Chlorophyll Fluorescence During Stress and Recovery

In the surviving seedlings, large differences in shoot gas exchange were observed between the treatments (see **Table S3** for linear mixed effect model results). While  $A_{net}$  and  $R_{dark}$  in drought-treated seedlings decreased proportionally (**Figures 2A,B**), we found contrasting responses under high temperature stress. Most markedly, in both the heat and heat-drought treatment a sharp decline in  $A_{net}$  was contrasted by a pronounced increase in  $R_{dark}$  (heat: +101%, heat-drought: +35%,  $P \leq 0.001$ ) during the first heat wave. In the surviving heat-drought seedling, the increase in  $R_{dark}$  during the first heat wave and  $R_{dark}$  did not reach control levels throughout the remainder of the experiment (-49.8%,  $P \leq 0.001$ ). In contrast, the heat-treated seedlings maintained high  $R_{dark}$  rates and  $A_{net}$  recovered instantaneously after the second heat wave had ended.

Canopy transpiration was strongly affected by the two heat waves and increased markedly in the heat treatment, while water deficit resulted in a reduction of  $E_{day}$  in the heat-drought treatment (**Figure 2C**). In both treatments the changes in  $E_{day}$ were accompanied by decreasing  $g_s$  (**Figure 2D**). Because  $A_{net}$ appeared coupled to  $g_s$  but uncoupled from  $E_{day}$  during heat wave conditions, WUE<sub>a</sub> declined sharply while WUE<sub>i</sub> remained relatively unchanged, but slightly increased in the surviving heatdrought seedling post-stress ( $P \le 0.001$ ; **Figures 2E,F**).

The relationships of shoot gas exchange with VPD (**Figure 4**) revealed striking differences between drought and well-watered seedlings. In the heat treatment, a decline in  $g_s$  was particularly visible during the initial increase in VPD (0.5–3 kPa), while  $g_s$  remained surprisingly constant at higher VPD. This resulted in increasing  $E_{day}$ , while photosynthesis remained constant over a temperature range between 32 and 42°C. The decline of  $g_s$  in the heat-drought treatment indicated stomatal closure at a VPD of about 3.5 kPa, tightly limiting  $E_{day}$  and  $A_{net}$ .

The impairment of the photosynthetic apparatus under heatdrought stress was clearly visible in all measured chlorophyll fluorescence parameters 1 day after the last heat wave (**Figure 5**), where  $\Phi_{PSII}$  showed a treatment effect (ANOVA,  $P \leq 0.001$ ) and the strongest decline by 78% compared to the control (*posthoc*,  $P \leq 0.001$ ) and the drought treatment (*post-hoc*,  $P \leq$ 0.05). The parameters also reflected the reduced, but continuing photosynthetic activity in the heat and drought treatment, in which small but not significant reductions were observed. The apparent recovery of all chlorophyll fluorescence parameters in the heat-drought treatment should be interpreted with caution, because only data of intact needles of the three surviving seedlings are reported.

## Non-structural Carbohydrates

We found no distinct stress responses of NSC concentrations in shoots (**Figures 6A,C,E,G**). In contrast, soluble NSC and starch contents in roots declined in response to heat and heat-drought stress (**Figures 6B,D**; see **Tables S4, S5**). Heat-drought caused a steep depletion of soluble sugars (-97%, 2.4 µmol gDW<sup>-1</sup>) and starch (-98%, 0.8 µmol gDW<sup>-1</sup>). Carboxylic acids and cyclitols were also significantly affected by heat-drought stress, causing an overall treatment effect (ANOVA,  $P \le 0.001$ ) and reduction in total C of 83% compared to the control. The Heat-drought treatment differed from all other treatments (*post-hoc*,  $P \le 0.05$ ).

During recovery, carboxylic acids (toward control and drought) and cyclitols (toward all treatments) increased significantly in shoots of heat-drought treated seedlings (**Figures 6E,G**; *post-hoc*,  $P \le 0.05$ ). While in the heat and in the heat-drought treatment root soluble NSC and starch increased during recovery close to control values (**Figures 6B,D**), total C remained reduced in the heat-drought treatment (-85%, ANOVA,  $P \le 0.001$ ). In dying seedlings, NSC contents did not recover (**Figure 6**) and starch storage in shoots and roots was close to depletion.

## Proline, Nitrate, and Osmotic Potential

Concentrations increased in shoots of heat and heat-drought treated seedlings during stress and recovery (Figures 7A,B)



bin-averaged in VPD classes of 0.5 kPa (10 a.m. -7 p.m.).

with a treatment effect after recovery (ANOVA,  $P \leq 0.001$ ). Dead seedlings showed accumulation of proline during both sampling campaigns. Proline content in roots was much smaller and no clear response to the treatments was found.

Shoot nitrate content, proposed here as an indirect measure for nitrogen (N) assimilation, was 2 to 4 times higher during stress in heat and heat-drought seedlings (*post-hoc*,  $P \le 0.05$ ) than the control or drought treatment. Following recovery, nitrate content in the heat treatment decreased to control levels, while it showed a lasting effect in the heat-drought treatment (ANOVA,  $P \le 0.05$ ) and remained elevated toward drought (*post-hoc*,  $P \le 0.05$ ). No obvious effect of the treatments on root nitrate was visible (**Figure 7D**). Again, the dead seedlings showed highest concentrations and no decline during recovery.

The osmotic potential ( $\Pi$ ) of shoots remained relatively constant in all treatments throughout the experiment.  $\Pi$  in roots showed a tendency to decrease in the drought (treatment effect; ANOVA,  $P \leq 0.05$ ) and heat-drought treatment although root total C declined (**Figure 6J**) because root water content was reduced (**Table S2**).





# DISCUSSION

#### **High Temperature and Mortality**

Although the exact cause of seedling mortality is not clear, it likely relates to the high shoot surface temperatures that developed in particular when transpiration ceased. As indicated by the highresolution dendrometer data (Figure S3) seedlings in the heatdrought treatment died right after the last day of the first heat wave and desiccated rapidly afterwards. In contrast, seedlings in the heat treatment showed strong stem diameter declines, indicating death, only after the second heat wave had ended. Such a delay in stress response might indicate that cellular damage (e.g., in needles, roots and/or cambium cells) progresses over time, as has been observed after high temperature stress, e.g., Hüve et al. (2011), until critical levels are reached beyond which tree functioning cannot longer be maintained. This was also described as indirect damage by Colombo and Timmer (1992). The differences in the occurrence of mortality between heat and heat-drought treatments might reflect dosage effects of the experienced heat stress. The reduced transpiration in the heatdrought treatment (c. 0.5 mmol  $m^{-2}s^{-1}$ ) vs. the heat treatment (c. 2 mmol  $m^{-2}s^{-1}$ ) at a VPD of 4 kPa and 42°C air temperature resulted in 2.6°C higher needle temperatures at the end of the second heat wave (Table 1). Such an increase in leaf temperature has also been observed in mature forests when drought-exposed and well-watered trees were compared (Scherrer et al., 2011). With regard to the pronounced mortality under heat-drought as observed in our experiment, leaf cooling could have been crucial for survival as leaf temperatures had reached 47°C, a critical value for tissue damage in conifers (Colombo and Timmer, 1992; Bigras, 2000). The importance of evaporative cooling at lower VPD was examined recently for Pinus taeda (Urban et al., 2017b). The study showed that seedlings were able to keep stomata open to maintain high transpiration rates. This resulted in lower leaf temperatures. According to a field experiment on *Eucalyptus parramattensis* by Drake et al. (2018), leaf surface heating was reduced by  $2.8^{\circ}$ C on average because transpiration was maintained at high rates even at temperatures >43°C.

The ability of the pine seedlings to cool via transpiration was found to fail under the water limiting conditions applied here. Under field conditions, leaf cooling can be supported by transport of sensible heat via high wind speed or convective air flow. Such upward flow conditions and cooling of forest canopy have been shown at the Yatir forest (Rotenberg and Yakir, 2011; Eder et al., 2015). However, wind speed and convective exchange for small seedlings under canopy is lower as for mature trees. Also, tissue temperature is strongly affected by the closeness of bare and hot soil surfaces (Kolb and Robberecht, 1996). The authors found that under such conditions, seedlings with higher stomatal conductance were more likely to survive. This compares well to the findings that trees with low  $E_{day}$  and  $g_s$  died, while seedlings that maintained generally higher Eday rates survived (Figure 2). In summary, the high temperatures experienced by the seedlings in the heat and heat-drought treatment caused pronounced stress, likely resulting in membrane and protein damage causing higher mortality in trees with lower evaporative cooling capacity.

# Stress Severity and Impairment of N Assimilation

The impact of heat and heat-drought stress was not only reflected in high mortality rates but also by highly elevated proline



**FIGURE 6** | Changes in non-structural carbohydrate (NSC) concentrations for shoot and root tissues during stress and recovery relative to before stress conditions. Shown are (**A**,**B**) soluble non-structural carbohydrates (soluble NSC: glucose, fructose, sucrose), (**C**,**D**) starch (**E**,**F**) carboxylic acids (malate, fumarate, citrate), (**G**,**H**) cyclitols (myo-inositol, D-pinitol), and (**I**,**J**) total carbohydrates (given as C6 equivalents of the shown metabolites) for control, drought, heat, heat-drought and dead seedlings. The error bars are  $\pm 1$  SE (n = 6, but heat-drought during stress and recovery n = 3, dead heat samples n = 5, dead heat-drought samples n = 15). The intermitted lines provide comparison to before stress conditions. Significant treatment effects are given \*(ANOVA,  $P \le 0.05$ ). See **Table S2** for absolute concentrations.

contents. Proline was not upregulated in response to the applied drought alone. However, drought stress can be viewed as rather mild, as stomata did not close completely and  $\psi_{shoot}$  remained well above critical values for xylem embolism formation (Oliveras

et al., 2003; Delzon et al., 2010; Klein et al., 2011; David-Schwartz et al., 2016). Thus, function of proline as an osmoprotectant was of minor importance, which is in line with an unchanged  $\Pi$  of shoots during drought compared with control. It has been found



stress conditions for control, drought, heat, heat-drought and dead seedlings. Osmotic potential was calculated from measured solutes and tissue water content using the van't Hoff equation. The error bars are  $\pm 1$  SE (n = 6, but heat-drought treatment during stress and recovery n = 3). The intermitted line provides comparison to before stress conditions. Significant treatment effects are given \*(ANOVA,  $P \le 0.05$ ). In dead seedlings, when changes exceed axis scaling, the mean value was placed below the SE.

that proline accumulates under salinity stress (Hamilton, 2001; Hayat et al., 2012). Although not directly relevant it is interesting to note that a quadratic relationship between shoot chloride content and proline concentrations was found (**Figure S4**). This might have been caused by increased uptake of chloride by roots at high temperatures (Turner and Lahav, 1985).

Proline upregulation under heat stress is reported as beneficial for plants due to its role as protein stabilizer (e.g., complex II of mitochondrial electron transport chain; Hamilton, 2001). Moreover, proline synthesis serves as an efficient NADPH scavenger to regenerate NADP<sup>+</sup> (Szabados and Savouré, 2010; Hayat et al., 2012; Zhang et al., 2015) and to prevent radical production in the thylakoid electron transfer chain (Kramer et al., 2004). This mechanism might enhance ETR by regenerating electron acceptors, and thus reducing oxidative stress. Indeed, we found that ETR was unaffected in the heat treatment, while it was strongly reduced in the heat-drought treatment. Thus, the membrane and protein stabilizing ability of proline could not prevent photo-inhibition at the time of highest needle temperatures in the heat-drought treatment (>47°C). Levels of nitrate in shoots of heat and heat-drought stressed Aleppo pines were elevated, while they did not change in roots, thus indicating heat-induced inactivation of leaf nitrogen assimilation. Decreased nitrate reductase activity has been observed to occur at temperatures above 40°C in cereals (Onwueme et al., 1971; Pal et al., 1976) and nitrate reductase activity recovers quickly after heat release. A fast recovery of N assimilation in Aleppo pine 2 days after stress release may also be concluded from the drop in leaf nitrate content (see **Figure 7**), while in the dead seedlings nitrate levels remained highly elevated (>900%).

# Metabolic Responses Toward Heat-Drought Stress

The degree of stress and reduction in gas exchange were reflected in root NSC concentrations. In the heat-drought treatment, soluble NSC, starch, carboxylic acids and cyclitols declined to a larger extend than in the heat treatment although root zone temperatures were similar in both treatments (data not shown). Supposedly, the smaller C uptake in the heatdrought treatment decreased phloem transport (Ruehr et al., 2009; Sevanto, 2014, 2018), which caused a strong depletion of soluble sugars and starch storage in roots. Total NSC levels were close to zero in the roots of the heat-drought seedlings, which may indicate critical C shortage in roots, limiting the otherwise high maintenance respiration in roots at high temperatures (Jarvi and Burton, 2018; Tjoelker, 2018). This could have affected root functionality and hence contributed to the observed large mortality rates in the heat-drought treatment.

Changes in lipids, which can be critical for C supply during stress, were not measured. In particular in pine trees, lipids (triglycerides) together with cyclitols are reported to be a large carbon storage pool (Piispanen and Saranpää, 2002; Hoch et al., 2003) and this pool can easily be channeled into the Tricarboxylic-acid cycle especially during extreme stress (Fischer et al., 2015). This partly explains the weak responses of carboxylic acids under heat and heat-drought, whereas root NSC was depleted. Furthermore, carboxylic acids together with cyclitols, did surpass or at least equal the concentration of the typically measured NSC (starch, sucrose, glucose, and fructose; **Table S2**), and hence should be considered as important plant carbohydrate pools.

Carboxylic acids exert key functions in the regulatory network of plants. For example, intermediates from the tri-carboxylicacid cycle (TCA) are important for the generation of ATP and thus key in respiratory processes. Additionally, they can provide carbon skeletons for amino acids and carbohydrates (Fernie et al., 2004). An increase in carboxylic acids during recovery might indicate enhanced investment into provision of carbon skeletons for maintenance and repair mechanisms. For heat-drought treated seedlings the accumulation of carboxylic acids can be explained by the reduction of  $R_{dark}$ . A similar upregulation of TCA intermediates was recently reported for eucalypts in response to heat and drought (Correia et al., 2018).

While shoot soluble NSC did not decline, shoot  $R_{dark}$  was much lower during the second than in the first heat wave and during the period of recovery. Presumably, available C for respiration was not limited in the heat-drought treatment at this stage because of actively maintained high sugar levels. This can be concluded from the upregulation of compatible solutes, including cyclitols during recovery. A similar pattern has also been observed in Scots pine where newly assimilated carbohydrates were allocated to cyclitols during recovery from drought (Galiano et al., 2017). Only heatdrought was associated with an increase in cyclitols during recovery, which might be explained by the very high needle temperatures that were found in this treatment and the protective function of cyclitols against heat denaturation (Jaindl and Popp, 2006).

Reports on NSC concentrations during drought have often been found to be inconclusive and to differ between species. Although hydraulic failure is often lethal, drought-induced lethality is not always connected to NSC depletion (Adams et al., 2017). In fact, NSC concentrations during drought were found to increase, decrease or remain constant in a wide range of tree species (Hartmann and Trumbore, 2016). Information on

isolated heat effects on NSC of tree seedlings is scarce and might depend on heat dosage rather than temperature maxima. For example, it has been postulated that high temperatures lead to an increase in shoot NSC (Sevanto and Dickman, 2015). Also, Marias et al. (2017) have found an increase of sugars in shoots, when coffee plants have been exposed to a heat pulse (49°C for 45 min). This is in stark contrast to our observations of relatively unchanged NSC levels in shoots and declining NSC in roots during the heat and heat-drought treatment. We suggest that the key for explaining the results is the duration of the experiment, which is corroborated by findings that longer stress periods did also not change NSC concentrations in leaves of Eucalyptus globulus, while NSC concentrations declined when treated with additional drought (Correia et al., 2018). To conclude this section, the primary metabolism of Aleppo pine shoots appears to be highly buffered against deviations even under extraordinary stressful conditions. In contrast, the root metabolism decouples from the source supply under mild drought and/or extreme heat waves and is clearly affected by combined stressors. These changes in allocation patterns might indicate an approach to preserve source functionality at the cost of sink integrity.

# Photosynthetic Inhibition and Recovery From Acute Heat Stress

Water availability during heat waves seems to be a key factor of survival (Bauweraerts et al., 2014; Ruehr et al., 2016) since it translates in evaporative cooling capacity (Drake et al., 2018) and protects the photosynthetic apparatus. This is supported by leaf fluorescence that showed only a moderate decline under heat stress (only ETR; *post-hoc*,  $P \leq 0.05$ ) while in heat-drought trees chlorophyll fluorescence parameters (**PSII**,  $F'_v/F'_m$ , qP, and ETR) decreased strongly. A decline in  $\Phi$ PSII indicates a saturation of the electron transfer chain, bearing the risk of oxidative stress (Murchie and Lawson, 2013). A reduction of  $\Phi$ PSII due to pigment degeneration can be neglected because of their stability above 60°C (Rätsep et al., 2018) and only temperatures above 50°C are considered critical for the disaggregation of the harvesting complexes (Tang et al., 2007; Nellaepalli et al., 2014). ETR, on the other hand, largely depends on thylakoid membrane integrity, which is disturbed at much lower temperatures of about 38°C (Havaux et al., 1996; Bukhov et al., 1999). Dark-adapted maximum operating efficiency F<sub>v</sub>/F<sub>m</sub> is widely used for the estimation of stress severity (Murchie and Lawson, 2013). In this experiment,  $F'_v/F'_m$ (light-adapted) was recorded instead. This parameter decreases with increasing non-photochemical quenching (Murchie and Lawson, 2013), which means the dissipation of excess energy by heat. Together with the hypothesis of a saturated photosystem and reduced ETR, this provides evidence for stress-induced limitations of processes downstream of the light reaction of photosynthesis. There is increasing evidence that thresholds for irreparable/slowly recoverable damage are typically reached at temperatures >45°C in temperate regions (Hüve et al., 2011).

The apparent small decline of chlorophyll fluorescence parameters in the heat treatment, measured 1 day after the last heat wave had ended, may either reflect a relative mild impairment of the photosynthetic apparatus, or indicate a fast recovery of chlorophyll fluorescence parameters 1 day after the acute stress. Fast recovery responses of photosynthetic parameters after acute heat waves (when water was not limiting) were observed in herbaceous plants, as well as in some tree species and typically relate to temperatures <45°C (Hüve et al., 2011; Ameye et al., 2012; Guha et al., 2018). However, complete recovery of the few surviving seedlings in the heat-drought treatment seemed to be impaired.

# Canopy Gas Exchange Affected by High Temperatures, Atmospheric Demand and Soil Drought

Heat stress, which induces high atmospheric evaporative demand has been reported to result in (partial) stomatal closure (Ameye et al., 2012; Bauweraerts et al., 2013; Duarte et al., 2016; Garcia-Forner et al., 2016; Ruehr et al., 2016). In our study, we found that stomata did not fully close in the heat treatment even under extreme vapor pressure deficits of >6 kPa. Indeed, partial stomatal closure was enough to maintain midday stem WP at moderate levels of -1.2 MPa (measured at the last day of the second heat wave), indicating that the xylem water transport was operating under non-harmful conditions. Under the same atmospheric conditions but reduced irrigation, the stress on the hydraulic system of the seedlings was larger (-1.8 MPa)and stomata were almost fully closed preventing any further drop in  $\psi_{shoot}$  and hence xylem embolism. Separating the effects of high temperature and VPD on stomatal responses is challenging. In experiments with temperature rise independent of VPD,  $g_s$  has been shown to increase (Urban et al., 2017a,b). A temperature-driven increase in  $g_s$  might simply reflect enhanced water loss along with rising temperatures without any further opening of stomata aperture, or could reflect an active cooling mechanism that prevents further closing of stomata. While the mechanism has not yet been explained in detail, various results indicate an active role of leaf cooling as a reasonable option (Urban et al., 2017a,b; Drake et al., 2018).

 $A_{\text{net}}$  and  $E_{\text{day}}$  were uncoupled during the heat waves, resulting in a strong decline in WUE<sub>a</sub>. Both, fluxes showed slightly different response pattern during the first and the second heat wave with a less pronounced decrease in  $A_{\text{net}}$  in the former and a less pronounced increase in  $E_{\text{day}}$  during the latter. This resulted in similar amplitudes of WUE<sub>a</sub> decrease in both heat waves.  $R_{\text{dark}}$  was similarly dampened during the second heat wave, especially in the surviving heat-drought seedling. A similar decline in the amplitudes of gas exchange in response to repeated heat waves had been reported before for various other tree species (Duarte et al., 2016; Guha et al., 2018) and had been linked to cellular damage. Also WUE<sub>i</sub> remained elevated during recovery from combined heat and drought stress. This was at the cost of reduced transpiration, which resulted in higher needle surface temperatures along with elevated levels of compatible solutes and dramatically changed shoot-to-root C allocation patterns.

# CONCLUSION

The combination of heat and drought stress affected Aleppo pine seedlings differently than drought or heat alone. This was not restricted to the stress periods per se, but became more pronounced during post-stress since high mortality rates and delayed recovery of the surviving seedlings occurred. The observed delay in recovery and pronounced mortality in the heat-drought treatment clearly demonstrated that physiological stress responses can continue after environmental stress has been released. Moreover, it showed that a tight regulation of a plant's water balance via stomatal closure at the cost of evaporative cooling can result in excessive needle temperatures under the experimental conditions applied here. Although the exact cause of seedling death is difficult to depict, it is likely that observed needle temperatures (>47°C) either directly damaged shoot tissues or indirectly affected root vitality via reduced C translocation. In summary, we conclude that increases in heat wave temperatures, as predicted to occur during the next decades, can have disastrous effects in dry environments. It seems that in semi-arid forests where drought is a common phenomenon, even a single heat wave that surpasses a given threshold (probably above 47°C needle temperature) may have widely detrimental effects.

# DATA AVAILABILITY STATEMENT

All relevant data is contained within the manuscript and the supplementary files. Shoot gas exchange data will be made available by the authors upon request.

# **AUTHOR CONTRIBUTIONS**

AA, BB, NR, and RG designed the study. BB, MG, and NR conducted experiments. AH, BB, and MG conducted measurements. AH, BB, and NR contributed to data evaluation. BB and NR wrote the manuscript. AA, AH, MG, and RG contributed to revision.

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#### SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/ffgc.2018. 00008/full#supplementary-material

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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