High resilience of carbon transport in long-term drought-stressed mature Norway spruce trees within 2 weeks after drought release

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Abstract

Under ongoing global climate change, drought periods are predicted to increase in frequency and intensity in the future. Under these circumstances, it is crucial for tree’s survival to recover their restricted functionalities quickly after drought release. To elucidate the recovery of carbon (C) transport rates in c. 70-year-old Norway spruce (Picea abies [L.] KARST.) after 5 years of recurrent summer droughts, we conducted a continuous whole-tree $^{13}$C labeling experiment in parallel with watering. We determined the arrival time of current photoassimilates in major C sinks by tracing the $^{13}$C label in stem and soil CO$_2$ efflux, and tips of living fine roots. In the first week after watering, aboveground C transport rates (CTR) from crown to trunk base were still 50% lower in previously drought-stressed trees (0.16 ± 0.01 m h$^{-1}$) compared to controls (0.30 ± 0.06 m h$^{-1}$). Conversely, CTR below ground, that is, from the trunk base to soil CO$_2$ efflux were already similar between treatments (c. 0.03 m h$^{-1}$). Two weeks after watering, aboveground C transport of previously drought-stressed trees recovered to the level of the controls. Furthermore, regrowth of water-absorbing fine roots upon watering was supported by faster incorporation of $^{13}$C label in previously drought-stressed trees (0.16 ± 0.01 m h$^{-1}$) compared to controls (0.30 ± 0.06 m h$^{-1}$). Conversely, CTR below ground, that is, from the trunk base to soil CO$_2$ efflux were already similar between treatments (c. 0.03 m h$^{-1}$). Two weeks after watering, aboveground C transport of previously drought-stressed trees recovered to the level of the controls. Furthermore, regrowth of water-absorbing fine roots upon watering was supported by faster incorporation of $^{13}$C label in previously drought-stressed trees (0.16 ± 0.01 m h$^{-1}$) compared to controls (0.30 ± 0.06 m h$^{-1}$). Conversely, CTR below ground, that is, from the trunk base to soil CO$_2$ efflux were already similar between treatments (c. 0.03 m h$^{-1}$). Thus, the whole-tree C transport system from the crown to soil CO$_2$ efflux fully recovered within 2 weeks after drought release, and hence showed high resilience to recurrent summer droughts in mature Norway spruce forests. This high resilience of the C transport system is an important prerequisite for the recovery of other tree functionalities and productivity.
1 | INTRODUCTION

Global climate change has been causing significant and mostly negative impacts on forest ecosystem carbon (C) cycling such as reduced productivity (Ciais et al., 2005; Collins et al., 2013). Drought is one of the most influential drivers of tree mortality (Allen et al., 2010; 2015; McDowell et al., 2008; van Mantgem et al., 2009) and it is predicted to occur more frequently and for longer durations in the future (IPCC, 2007, 2014). Under these circumstances, tree survival primarily depends on the extent to which tree functionality is impaired by drought (i.e., resistance, Lloret et al., 2011). After drought release, it is then crucial that surviving trees recover their limited functionality back to pre-drought levels (i.e., resilience, Lloret et al., 2011). Since drought release typically causes a high C demand for repair and growth particularly in belowground sinks (Gao et al., 2021; Hagedorn et al., 2016; Joseph et al., 2020), C transport from leaves to sink organs is an important process for tree recovery (Ruehr et al., 2019). C assimilates are transported from the crown via the phloem to various above- and belowground C sinks (Lemoine et al., 2013; Salmon et al., 2019). Recent studies revealed that saplings (Barthel et al., 2011; Ruehr et al., 2009; Zang et al., 2014), young trees (Dannoura et al., 2019; Epron et al., 2016), and mature trees (Gao et al., 2021; Hesse et al., 2019; Wang et al., 2021) restricted transport of current photoassimilates under drought, thereby reducing the C supply to sinks. Upon drought release, C limitation in sink tissues can occur if the C transport would not recover fast enough to meet the sink demands (Hartmann et al., 2013; Hartmann et al., 2013; Sevanto, 2014; Winkler & Oberhuber, 2017), but knowledge on mature trees is scarce (Gao et al., 2021).

There are two main causes restricting transport of current photoassimilates from the crown along the stem to belowground C sinks under drought (Salmon et al., 2019). First, water limitation delays the export of sugars from leaves, increasing the mean residence time (MRT) of photoassimilates in leaves (Dannoura et al., 2019; Epron et al., 2012; Hesse et al., 2019; Ruehr et al., 2009). This is caused by accumulation of osmolytes, and/or production of secondary metabolites and volatile compounds (Epron & Dreyer, 1996; Ruehr et al., 2009; Salmon et al., 2019). Second, the phloem transport velocity can be reduced through increased phloem viscosity (Epron et al., 2016; Hesse et al., 2019; Sevanto, 2014, 2018; Woodruff, 2014), lower C source/sink strength (Lemoine et al., 2013; Ryan & Asao, 2014; Sevanto, 2014), and smaller phloem conduit diameter (Dannoura et al., 2019; Woodruff, 2014). Increased phloem viscosity is a result of water limitation in the xylem, as the xylem supplies the nearby phloem with water (Hölttä et al., 2006, 2009). Lower C source/sink strength (e.g., photosynthesis rates and stem/sink CO$_2$ efflux rates) limits sugar loading/unloading processes between C source/sink and phloem. This hinders the osmotic regulation in phloem and thus limits water exchange between phloem and xylem. Smaller phloem conduit diameter is caused by restricted cell expansion due to turgor reduction usually under severe drought (Hsiao, 1973), thereby reducing phloem conductivity.

Recovery of C transport depends on the restricting mechanisms. MRT of leaf sugars decreases after drought release within days (Zang et al., 2014). Drought release increases plant water potential and water availability in the xylem, typically followed by increased C source and sink strength (Gao et al., 2021; Hagedorn et al., 2016). Previous studies using young eucalypt trees (Epron et al., 2016) and a rainfall event in a naturally dry pine forest (Gao et al., 2021) reported that C transport velocity from crown to trunk base or soil was related to C source or sink strength, which typically decreases under drought and increases after drought release (Hagedorn et al., 2016; Joseph et al., 2020; Nikolova et al., 2009; Zang et al., 2014). Furthermore, there is increasing evidence that C source strength and C supply may be “sink controlled” (Faticchi et al., 2014; Gavito et al., 2019; Hagedorn et al., 2016; Körner, 2015). Conversely, drought-related reductions of phloem conduit diameter are expected to further restrict the phloem transport during the first weeks after stress release even if the phloem sap viscosity and C source/sink strength recover.

This present study was performed in the framework of the Kranzberg roof (KROOF) project, which was initiated to elucidate the drought responses of mature European beech (Fagus sylvatica L.) and Norway spruce (Picea abies [L.] KARST.; see details in Grams et al., 2021). Both tree species were exposed to recurrent summer droughts from 2014 to 2018 and leaf water potential reached values as low as −1.8 MPa, causing distinct drought effects such as reduced stem and fine root growth (Grams et al., 2021; Pretzsch et al., 2020; Zwetsloot & Bauerle, 2021) and acclimation in tree hydraulics (Tomasella et al., 2018). To predict the trajectories of forests under future climates, it is important to understand, to what extent tree functionality recovers after drought release and how fast. To answer this question, former drought plots were watered in early summer 2019 (Grams et al., 2021). In parallel with watering, we performed a whole-tree $^{13}$C labeling experiment on mature spruce trees to assess the resilience of their C transport processes, that is, the ability to recover to the level of control trees (Lloret et al., 2011).

We divided the C transport path from the crown to the soil CO$_2$ efflux into two parts (Figure 1), as drought release may affect them differently. (1) Aboveground transport from the crown (leaves) to the trunk base (aboveground transport hereafter), and (2) belowground transport from the trunk base to the soil CO$_2$ efflux (belowground transport hereafter). In addition, we also investigated a third process, (3) incorporation of current photoassimilates in living fine roots (Figure 1). The aboveground transport comprises sugar export from leaves and transport along the woody structures in the

**KEYWORDS**

$^{13}$C labeling, climate change, forest ecosystem, phloem, photosynthesis, *Picea abies*, recovery, soil CO$_2$ efflux, stem CO$_2$ efflux, watering
The aboveground C transport rates ($\text{CTR}_{\text{above}}$ in m h$^{-1}$) indicate how fast newly assimilated C can be supplied to belowground sinks. The belowground transport includes the phloem transport along roots and the CO$_2$ diffusion in the soil. The belowground C transport rates ($\text{CTR}_{\text{below}}$ in m h$^{-1}$) from trunk base to soil CO$_2$ efflux, and (3) Incorporation time (in h) of current photoassimilates from trunk base to fine root tips.

In a similar experiment by Gao et al. (2021) conducted in a naturally dry pine forest after a rainfall event is the only study to date investigating CTR of mature trees after drought release. We still lack knowledge on the recovery of highly productive forests under ongoing climate change. Furthermore, there is no study considering the effect of water availability on the above- and belowground transport individually. We show for the first time the resilience of the whole-tree C transport after repeated summer droughts in a highly productive Norway spruce forest stand of great ecological and economic relevance in central Europe (Caudullo et al., 2016).

2 | MATERIALS AND METHODS

2.1 | Experimental site

This study was conducted in a mixed forest with c. 90-year-old European beech and c. 70-year-old Norway spruce trees in Kranzberg Forest, located in southern Germany/Bavaria (11°39′42″E, 48°25′12″N; 490 m a.s.l.). The experimental site consists of 12 plots with three to seven beech and spruce trees each. At this site, a long-term throughfall exclusion (TE) and subsequent watering experiment was conducted as described in detail.
in Grams et al. (2021). Briefly, six plots were assigned to TE plots equipped with roofs and the other six plots without roofs to control plots (CO). All plots were trenched to 1 m of soil depth 4 years before the experiments started (Pretzsch et al., 2014). The mature beech and spruce trees in TE plots were then exposed to summer drought for five consecutive growing seasons (2014–2018). To investigate trees’ recovery processes, in early summer 2019, all TE plots were watered with c. 90 mm over 36 h and the soil water content increased to the level of the CO plots within 1 week (for further details see Grams et al., 2021). In parallel with the watering, we conducted a $^{13}$C labeling experiment on four CO and three TE spruce trees on neighboring plots (Figure 2a, for details see Table S1). In addition to the two labeled plots, we assessed three spruce trees each on additional CO and TE plots as non-labeled controls (Table S2). A canopy crane located next to these plots enabled the measurements of leaf photosynthesis, leaf water potential, and leaf osmotic potential in sun-lit canopy.

2.2 | Weather data

The mean photosynthetic photon flux density during the labeling period accounted to 788 ± 534 (SD) μmol m$^{-2}$ s$^{-1}$ (Figure 3a). During the daytime (from 5 am to 7 pm, CET) on the labeling days, mean temperature was 18.8 ± 4.3 (SD)°C (Figure 3b) and mean vapor pressure deficit was 0.6 ± 0.4 (SD) kPa. There were several rain periods during labeling on day 3, 7, and 9. Only on day 9, however, weak but continuous rainfall event with a high wind speed occurred throughout the daytime, accumulating to 7.8 mm (Figure 3b). Due to this weather conditions, a smaller $\delta^{13}$C shift in canopy air was achieved on day 9 (see below).

2.3 | CO$_2$ exposure and assessment of canopy air

The whole crowns of all spruce trees on the CO and TE plot, that is, four and three trees, respectively, were fumigated with $^{12}$C-depleted tank CO$_2$ ($\delta^{13}$C of −44.3 ± 0.2‰) using the isoFACE system described earlier (Grams et al., 2011; Kuptz et al., 2011). Depending on its crown size, each tree crown was equipped with 9-17 microperforated PVC tubes hanging vertically from a carrier structure (Figure 2b). These fumigation tubes were then connected to the CO$_2$ tank and the $^{13}$C-depleted CO$_2$ was released directly within the seven tree crowns.

The atmospheric CO$_2$ concentration and $\delta^{13}$C in tree canopy ($\delta^{13}$C$_{C}$) were continuously monitored during the labeling using a cavity ring-down spectroscopy (CRDS, ESP-1000; PICARRO). Two air measurement points were installed per tree c. 2 m inside the sun-lit crowns at 1 m distance from the stem (east and west orientation, one CO tree had only one measurement point, Figure 2a and Table S2). We took care that these sampling points had enough distances from the fumigation tubes (c. 1 m). A sample point above the canopy was used as a reference. The sample air was continuously transported to the CRDS by membrane pumps via PVC tubes. A computer-automated multiplexer system switched every 5 min between measurement positions and averages of the last 3 min were recorded by the CRDS. According to the mean CO$_2$ concentration of all 13 measurement points in canopy, which was measured continuously by an infra-red gas analyzer (BINOS 100 4P; Rosemount-Emerson Electric Co.), a mass flow controller regulated the amount of the CO$_2$ exposure through fumigation tubes. To calibrate the CRDS, two commercially available calibration gases were used (Ref.1: −9.7 ± 0.3‰ and Ref.2: −27.8 ± 0.3‰; Thermo Fisher Scientific). All $\delta^{13}$C values in this study were referenced to international standards (Vienna Pee Dee Belemnite).

The $^{13}$C labeling started in parallel with the watering and continued for 14 days, that is, from July 4, 2019 (day 0) to July 17, 2019 (day 13), from 5 am to 7 pm (CET). We targeted the mean CO$_2$ concentration in canopy air at +130 ppm relative to the ambient air above the canopy to create a shift of ~8.3‰. Due to variable wind exposition, however, each tree received different amounts of added CO$_2$. In CO trees, the mean canopy CO$_2$ concentration increased to 541 ± 16 ppm during labeling (Figure 3c, see values for individual trees in Table S1), shifting the $\delta^{13}$C$_{C}$ by...
−7.3 ± 0.5‰ on average (Figure 3d). In contrast, TE trees received less 13C-depleted CO2 with an increase in the mean canopy CO2 concentration to 495 ± 23 ppm (Figure 3e), causing smaller mean shift of δ13C by −5.1 ± 1.3‰ (Figure 3f) compared to CO trees. Furthermore, during the weak but longer rainfall event associated with a high wind speed on day 9, a smaller mean shift in δ13C was achieved in both CO and TE trees. Mean CO2 concentration and δ13C of the ambient air above the canopy were 413 ppm and −9.2‰ during labeling hours.

To track the current photoassimilates through the tree/soil system, we used the two experimentally induced changes in δ13C:
(1) Turn-on of CO2 exposure with 13C-depleted tank CO2 on day 0 of watering. This part of the experiment was used to calculate the arrival time of the 13C-depleted tracer in the observed C sinks in the first week after watering.
(2) Turn-off of the CO2 exposure system and subsequent increase in δ13C in the canopy air back to the initial, ambient level on day 13 of watering. In this part of the experiment, the arrival of unlabeled tracer (C with ambient δ13C)
in the studied C sinks was used to calculate CTR 2 weeks after watering.

2.4 | Measurement of phloem sugar

On day −1, 7, 13, and 21 around midday, phloem tissue samples were collected at the breast height of four labeled CO and three labeled TE trees using a cork borer (two disks with diameter of 5 mm for each tree, Tables S2 and S3). The dead bark was removed and the remaining phloem samples were immediately frozen on dry ice and subsequently freeze-dried. The dried material was milled to fine powder using a steel ball-mill (Retsch) and about 70 mg per sample were transferred into a 2 ml reaction vial and mixed with 1.5 ml deionized water. The fractions of water-soluble compounds were closed before isotopic analysis. The C isotopic composition of phloem sugars (\( \delta^{13}C_{\text{phloem}} \)) was analyzed with a thermal conversion elemental analyzer (PYRO cube; Elementar) that was coupled via a ConFlo III reference system to an isotope-ratio mass spectrometer (Finnigan Delta Plus XP, all supplied by Thermo Fisher Scientific). The typical measurement precision for in-house sugar standards was 0.3‰ (SD).

2.5 | Measurement of stem CO\(_2\) efflux

Rates of stem CO\(_2\) efflux and its stable C isotope composition (\( \delta^{13}C_{\text{stem}} \)) before and after watering were recorded using an isotope ratio infrared spectrometer (IRIS, DeltaRay, Thermo Fisher Scientific; Braden-Behrens et al., 2017). A total of 12 spruce trees were measured, three \( ^{13} \text{C} \)-labeled and three non-labeled trees in each treatment, that is, CO and TE (n = 3; Figure 2a; Tables S2 and S3). The non-labeled trees were used to correct for changes in \( ^{13} \text{C} \) discrimination caused by the watering and weather fluctuations. Plexiglas (Röhm GmbH) chambers (61–204 cm\(^2\)) were attached at ca. 1 m height on each stem after removing mosses, lichens, and algae. After a leak test using a slight overpressure (c. 2000 Pa), each chamber was supplied with reference air of a constant CO\(_2\) concentration of c. 413 ppm. Excess air was exhausted before entering the chamber to avoid an overpressure. The mixture of reference air plus stem-derived CO\(_2\) of each chamber was continuously pumped through PVC tubes to a computer-automated manifold with 16 channels, which changed the channel flowing to IRIS every 5 min. The CO\(_2\) concentration and the stable C isotope composition of the reference air were determined between measurement cycles (c. every 80 min). The same reference gases as for the CRDS system were used for calibration of the IRIS system (see above).

The rate of stem-derived CO\(_2\) efflux was calculated according to mass balance equation as described by Gamnitzer et al. (2009), using the mean values of the closest two measurements of the reference air.

\[
\text{Stem CO}_2 \text{ efflux (\mu mol m}^{-2}\text{s}^{-1}) = \frac{F_{\text{air}}}{V_{\text{mol A chamber}}} \left(\left[\text{CO}_2\right]_{\text{sample}} - \left[\text{CO}_2\right]_{\text{reference}}\right),
\]

where \( F_{\text{air}} \) gives the air flow through the chamber (L s\(^{-1}\)); \( V_{\text{mol}} \), the molar volume of gases (22.4 L mol\(^{-1}\)); \( A_{\text{chamber}} \), the chamber base area (m\(^2\)); \( \left[\text{CO}_2\right]_{\text{sample}} \) and \( \left[\text{CO}_2\right]_{\text{reference}} \), the CO\(_2\) concentration (ppm) of sample air from stem chambers and reference air, respectively.

\( \delta^{13}C_{\text{stem}} \) was calculated by the following equation using a two end-member mixing model (Dawson et al., 2002),

\[
\delta^{13}C_{\text{stem}}(\text{‰}) = \frac{([\text{CO}_2]_{\text{sample}} \times \delta^{13}C_{\text{sample}}) - ([\text{CO}_2]_{\text{reference}} \times \delta^{13}C_{\text{reference}})}{([\text{CO}_2]_{\text{sample}} - [\text{CO}_2]_{\text{reference}})},
\]

where \( \delta^{13}C_{\text{sample}} \) and \( \delta^{13}C_{\text{reference}} \) give the \( \delta^{13}C \) signature of sample air from stem chambers and that of reference air, respectively.

\( \delta^{13}C_{\text{stem}} \) can be affected by CO\(_2\) transported from belowground in xylem sap (Teskey et al., 2008). However, Kuptz et al. (2011) observed a positive correlation in \( \delta^{13}C \) between stem phloem and stem CO\(_2\) efflux in spruce trees at the same experimental site. In this study, we also found a positive linear correlation between \( \delta^{13}C_{\text{stem}} \) and \( \delta^{13}C_{\text{phloem}} \) (slope = 0.94, R\(^2\) = .30, p < .01; Figure S1). Likewise, \( \delta^{13}C_{\text{stem}} \) showed no significant difference between daytime and nighttime (data not shown). Therefore, as reported in previous studies (Kodama et al., 2008; Kuptz et al., 2011; Ubierna et al., 2009), we concluded that CO\(_2\) in xylem sap had negligible effect on \( \delta^{13}C_{\text{stem}} \).

Thus, we assessed the \( \delta^{13}C_{\text{stem}} \) as a surrogate of \( \delta^{13}C_{\text{phloem}} \).

2.6 | Measurement of soil CO\(_2\) efflux

Soil CO\(_2\) efflux rates and its isotopic C composition (\( \delta^{13}C_{\text{soil}} \)) were measured using a Li-8100 automated soil CO\(_2\) flux system with a Li-8150 multiplexer (Li-Cor Inc.), connected to an IRIS. The air stream leaving the Li-8100 was sampled by the IRIS at a flow rate of 80 ml min\(^{-1}\) and added back to the chamber air stream. Three automatically operating soil chambers (8100-104) per treatment, that is, CO and TE, were installed with 1 m distance from the spruce trees (Figure 2a; Table S2). Additionally, one chamber was installed close to the non-labeled beech trees in the TE plot (Figure 2a; Table S2). Measurement time per chamber was adapted based on the CO\(_2\) efflux rate: 5 min in the TE plot and 2:30 min in the CO plot. \( \delta^{13}C_{\text{soil}} \) was calculated using the Keeling plot approach (Keeling, 1958, 1961). Each single measurement was quality controlled based on the fit of the linear regressions. For soil CO\(_2\) efflux values were kept if \( R^2 \geq .8 \) and for
δ^{13}C_{soil} based on the Keeling plot approach if $R^2 \geq 0.9$. To calibrate the IRIS, two commercially available calibration gases were used (Ref.1: $-9.9 \pm 0.3\%$ and Ref.2: $-27.8 \pm 0.3\%$, Thermo Fisher Scientific).

During watering of the TE plots, the soil pores fill with water and the lighter $^{13}$C-depleted CO$_2$ gets pushed-out (Andersen et al., 2010; Subke et al., 2009; Unger et al., 2010). This interfered with our labeling experiment. Hence, we corrected for the δ$^{13}$C$_{soil}$ of the TE plot based on measurements of the additional chamber close to the non-labeled beech trees (see details in Figures S2 and S3). Due to a limitation in the number of soil chambers, a non-labeled chamber was not available for the CO plot. For purposes not related to this study, the CO plot was slightly watered (c. 12 mm over 12 h) in parallel to the TE plots. As we did not observe any significant effect of the watering on the δ$^{13}$C$_{soil}$ of the wet CO plot (Figure 4c), there was no need to apply this correction here.

2.7 Measurement of root tips

Fine roots were collected on day −7 and repeatedly after the watering with an interval of 1–2 days until day 25 (Table S3), from random sampling positions (17–18 samples per treatment and day, Table S2). The collected samples were carefully washed in petri dishes, and representative living root tips were cut off under a stereomicroscope. Individual root tips were placed in pre-weighed tin capsules and dried at 60°C. Their stable C isotope composition ($\delta^{13}$C) was determined with an isotope-ratio mass spectrometer (delta V Advantage; Thermo Fisher Scientific) coupled to an Elemental Analyzer (Euro EA; Eurovector). Due to the very small sample quantities (the smallest samples with c. 3 µg C), the C-blank (c. 0.6 µg C) of the tin capsules and their δ$^{13}$C were taken into account in the evaluation. As with δ$^{13}$C$_{stem}$, δ$^{13}$C$_{root}$ of non-labeled plots was assessed to correct for the effect of watering and weather fluctuations.

2.8 Calculation of arrival time and CTR

To determine the arrival time of the two tracers ($^{13}$C-depleted tracer after the start of labeling, and unlabeled tracer after the end of labeling) in stem/soil CO$_2$ efflux and living root tips, the courses of δ$^{13}$C were fitted by piecewise function (Figure 4). Since $^{13}$C-depleted CO$_2$ decreases δ$^{13}$C, the arrival time of the $^{13}$C-depleted tracer was defined as the point when δ$^{13}$C started to decrease. First, δ$^{13}$C data of each C sink were fitted to contain only two linear segments before and after the arrival of the tracers. Then, we performed a linear regression for the δ$^{13}$C data (lm function, R package “stats,” version: 3.6.1). Finally, the intersection of two linear fits was determined using “segmented” function (R package “segmented,” version: 1.3-0, red lines fitted to the δ$^{13}$C data). This function calculated a new regression model and this function described above (blue lines fitted to the δ$^{13}$C data) and was then defined as the arrival time of unlabeled C (blue vertical lines in Figure 4a–f). In the case of root tips, it was not possible to assign each root to the belonging tree. Therefore, all values were pooled for each treatment (Figure 4e,f), providing only one arrival time for each treatment.

Using the arrival time in stem and soil CO$_2$ efflux, the CTR$_{above}$ (aboveground C transport rates from crown to trunk base in m h$^{-1}$, Figure 1) and CTR$_{below}$ (belowground C transport rates from trunk base to soil CO$_2$ efflux in m h$^{-1}$, Figure 1) were calculated by:

$$\text{CTR(m h}^{-1}) = \frac{d}{t_f}$$

For CTR$_{below}$, t$_f$ (in h) gives the time lag between the start respectively end of labeling and the arrival time of the tracers at trunk base (stem CO$_2$ efflux). d (in m) represents the distance between the mean crown height (the middle of the crown, Table S1) of the tree and the height of the stem chamber. For CTR$_{above}$, t$_f$ (in h) gives the time lag between the arrival time of the tracers at trunk base and the arrival time at soil CO$_2$ efflux, with d (in m) representing the height of the stem chamber plus 1 m, since each soil chamber was placed at 1 m distance from each trunk. The real transport distance from trunk base to soil chamber can vary depending on the structure of roots. We assumed that there is no time lag between arrival of current photosynthesizes at trunk base/roots and the use of them in stem/root CO$_2$ efflux. We did not calculate CTR to living root tips, since the transport distance was unknown due to random sampling positions. Therefore, for the incorporation time of current photosynthesizes in fine roots, the time lags between the arrival time of the tracers at trunk base and the arrival time at root tips were compared between CO and TE trees instead (Figure 1).

The additional soil chamber in the TE plot enabled to correct for the effects of watering on δ$^{13}$C$_{soil}$ (see details in Figures S2 and S3). Due to stable weather conditions in the first week of the labeling with only few short and weak rain events, we were able to calculate the arrival time of $^{13}$C-depleted tracer in soil CO$_2$ efflux. However, unstable weather conditions during the second part of the experiment (day 7–13) did not allow to calculate the arrival time of unlabeled tracer in soil CO$_2$ efflux (they caused negative time lags). Reduced C gain on day 9 increased the δ$^{13}$C$_{soil}$ already before the unlabeled tracer arrived in soil CO$_2$ efflux, likely as more $^{13}$C-enriched old C was used (Steinmann et al., 2004; Wingate et al., 2010). Thus, we excluded the CTR$_{below}$ calculated using unlabeled tracer 2 weeks after watering.

2.9 Measurement of light-saturated CO$_2$ assimilation rates ($A_{sat}$), predawn leaf water potential ($\Psi_{pd}$), and leaf osmotic potential ($\pi_O$)

The light-saturated CO$_2$ assimilation rates at CO$_2$ concentration of 400 ppm ($A_{sat}$, expressed on the basis of total needle surface area)
FIGURE 4  Examples for the calculation of the arrival time of the $^{13}$C-tracers, using: (a, b) $\delta^{13}$C of stem CO$_2$ efflux ($\delta^{13}$C$_{stem}$) of one control (CO) and one previously drought-stressed (throughfall exclusion, TE) tree, (c, d) $\delta^{13}$C of soil CO$_2$ efflux ($\delta^{13}$C$_{soil}$) of one CO and one TE soil chamber, and (e, f) $\delta^{13}$C of living root tips ($\delta^{13}$C$_{root}$) of CO and TE trees. Dashed vertical lines are the start and the end of labeling. The red and blue lines fitted to the data show the results of the piecewise functions to estimate the arrival time of $^{13}$C-depleted and unlabeled tracer, respectively (see Section 2). The intersections of two lines, marked with solid red and blue vertical lines are the calculated arrival times in the first week and 2 weeks after the watering, respectively. These arrival times (displayed here with arrows) were then used to calculate the above- and belowground carbon transport rates (CTR$_{above}$, CTR$_{below}$) and the incorporation time in fine roots (see Section 2). The red and blue shaded area give the 95% confidence interval of the intersections. The data of the other trees are displayed in Figure S4 (stem CO$_2$ efflux) and in Figures S3 and S5 (soil CO$_2$ efflux). All the root samples were pooled for each plot (CO and TE)
were measured on fully sun-exposed 1-year-old needles using a LI-6800 gas exchange system (Li-Cor Inc.) between 8 am and 3 pm (CET), before (around day −14) and after watering on days 4 and 14 (Table S3). In TE trees, when annual branch growth was not sufficiently long to cover the measurement chamber, needles from the previous year(s) were also included. Because of the small number of replicates in the present labeling plots (access by the canopy crane was limited by the labeling infrastructure), we additionally measured four spruce trees of each treatment in other plots (in total n = 6; Table S2). During the measurements, we set the light intensity to 1500 μmol m−2 s−1 and kept the leaf temperature at 25°C. The relative humidity was set to 60–65%. After the measurements, the needles were harvested and scanned (Epson Perfection 4990 Photo; Epson Deutschland GmbH). The projected needle surface area was multiplied by the factor 3.2 to determine the total needle surface area (Goisser et al., 2016).

Pre-dawn leaf water potential (ΨPD) and leaf osmotic potential (xO) on fully sun-exposed twigs were determined on day −6, 2, 7, and 22 (n = 6, same trees used for ASat, Tables S2 and S3). ΨPD was measured using a Scholander pressure bomb (1505D; PMS Instrument Co.) before sunrise (3 am–5 am CET). xO was determined with pressure volume curves (PV curves), following Tomasella et al. (2018). Collected twigs (two needle age classes) were rehydrated, and subsequently, their weight and water potential were repeatedly measured.

2.10 Statistical analysis

We analyzed all data using R (version 4.0.3) in R studio (version 1.3.1093). The treatment effect on the CTR and the time lags were tested using a t test. Beforehand, we tested the homogeneity of variances (F-test) and the normality of the data (Shapiro test). Since the homogeneity of variances between CTRabove and CTRbelow was violated, we tested their difference with wilcox.test (package: stats, version: 3.6.1). The differences in these arrival times, was 0.16 ± 0.02 m h−1 and thus about half in TE spruce compared to CO spruce with 0.30 ± 0.05 m h−1 (Figure S4). Despite similar transport distance of 28.4 ± 0.3 and 27.0 ± 0.9 m in CO and TE trees, respectively (Table S1), the arrival of the 13C-depleted tracer in stem CO2 efflux was significantly delayed in TE trees compared to CO trees (p < .05). The 13C-depleted tracer was found in stem CO2 efflux of CO trees 95 ± 10 h after the start of labeling and watering, whereas in TE trees the tracer arrived after 163 ± 12 h. CTRabove, calculated from these arrival times, was 0.16 ± 0.01 m h−1 and thus about half in TE spruce compared to CO spruce with 0.30 ± 0.06 m h−1 (Figure 5a; p = .06). Already 2 weeks after watering, CTRabove determined with the arrival of unlabeled tracer did not differ between treatments anymore, because of a significant increase in CTRabove of TE trees to 0.39 ± 0.13 m h−1 (Figure 5b). CTRabove of CO trees remained almost constant during the study period (0.32 ± 0.05 m h−1 2 weeks after watering).

3 RESULTS

3.1 Aboveground transport rates (CTRabove) from crown to trunk base

The 13C-depleted CO2 was successfully taken up by tree crowns and transported downwards along the stem after the start of labeling. For example, δ13Cstem of one CO tree in Figure 4a was −26.1 ± 0.1‰ before the start of labeling and remained almost constant for 4 days after the start of labeling. Then, δ13Cstem suddenly decreased after the 13C-depleted tracer arrived. Similar courses of δ13Cstem were observed in all six labeled trees assessed in this study (Figure 4b; Figure S4). Despite similar transport distance of 28.4 ± 0.3 and 27.0 ± 0.9 m in CO and TE trees, respectively (Table S1), the arrival of the 13C-depleted tracer in stem CO2 efflux was significantly delayed in TE trees compared to CO trees (p < .05). The 13C-depleted tracer was found in stem CO2 efflux of CO trees 95 ± 10 h after the start of labeling and watering, whereas in TE trees the tracer arrived after 163 ± 12 h. CTRabove, calculated from these arrival times, was 0.16 ± 0.01 m h−1 and thus about half in TE spruce compared to CO spruce with 0.30 ± 0.06 m h−1 (Figure 5a; p = .06). Already 2 weeks after watering, CTRabove determined with the arrival of unlabeled tracer did not differ between treatments anymore, because of a significant increase in CTRabove of TE trees to 0.39 ± 0.13 m h−1 (Figure 5b). CTRabove of CO trees remained almost constant during the study period (0.32 ± 0.05 m h−1 2 weeks after watering).

3.2 Leaf osmotic potential (πO) and predawn water potential (ΨPD)

xO increased with ΨPD following a sigmoidal fit (Figure 6; p < .001). Before watering, πO of the TE trees was on average −0.93 ± 0.03 MPa, which was significantly lower than that of CO trees with −0.59 ± 0.02 MPa (p < .05). On day 7, ΨPD was then similar between treatments with −0.61 ± 0.02 and −0.69 ± 0.05 MPa in CO and TE trees, respectively (p > .6). The lowest xO of −2.44 ± 0.05 MPa was observed for TE trees before watering, which was significantly lower than in CO trees with −1.67 ± 0.04 MPa (p < .01). Correlated with ΨPD, xO of TE trees increased by 0.5 MPa until day 22 to −2.0 ± 0.04 MPa. Nevertheless, on day 22, xO of TE trees was still somewhat lower than in CO trees (p < .1) that stayed around −1.6 MPa throughout the study.

3.3 Belowground transport rates (CTRbelow) from trunk base to soil CO2 efflux

The labeling with 13C-depleted CO2 also caused a sudden decrease in δ13Csoil but with a smaller shift compared to δ13Cstem (Figure 4c, d). In the first week after watering, the 13C-depleted tracer was detected in soil CO2 efflux under CO trees 73 ± 22 h |
after the detection in the stem CO$_2$ efflux. The time lag was similar in TE trees with $62 \pm 37$ h ($p > .8$). CTR$_{\text{below}}$ calculated from these time lags, was not significantly different between CO and TE trees with $0.03 \pm 0.01$ and $0.11 \pm 0.08$ m h$^{-1}$, respectively (Figure 5a, $p > .7$). The large variance of TE trees was caused by one tree with a high CTR$_{\text{below}}$ (0.28 m h$^{-1}$). CTR$_{\text{below}}$ was significantly lower than CTR$_{\text{above}}$ ($p < .05$).

3.4 Incorporation of current photoassimilates in living fine roots

In the first week after watering, the $^{13}$C-depleted tracer was detected in the living root tips of TE trees within 12 ± 10 h after the detection in the stem CO$_2$ efflux, whereas CO trees incorporated the current photoassimilates much later, that is, within 73 ± 10 h ($p < .05$; Figure 7a). Two weeks after watering, the incorporation time of the unlabeled tracer significantly decreased in CO trees to 14 ± 8 h after the detection at the trunk base ($p < .05$), which was similar to that of TE trees (10 ± 5 h, $p > .7$; Figure 7b).
TABLE 1 Light-saturated CO\textsubscript{2} assimilation rates (\(A_{\text{sat}}\)) before (around day −14) and after (day 4 and 14) the watering (means ± SE, \(n = 6\), expressed on the basis of total needle area), and rates of stem and soil CO\textsubscript{2} efflux before (day −1) and after (day 4 and 14/15) watering (means ± SE, \(n = 3\)) in CO (control) and TE (previously drought-stressed, throughfall exclusion) trees. The lowercase letters indicate the significant differences among treatments and days, determined by a post-hoc test after applying a linear-mixed model. \(A_{\text{sat}}\), stem CO\textsubscript{2} efflux, and soil CO\textsubscript{2} efflux were tested separately.

|                      | Before | Day 4 | Day 14/15 |
|----------------------|--------|-------|-----------|
| \(A_{\text{sat}}\) (\(\mu\text{mol CO}_2 \cdot \text{m}^{-2} \cdot \text{s}^{-1}\)) |        |       |           |
| CO                   | 2.7 ± 0.2\textsuperscript{a} | 2.6 ± 0.3\textsuperscript{a} | 2.3 ± 0.3\textsuperscript{a} |
| TE                   | 2.1 ± 0.3\textsuperscript{a} | 2.4 ± 0.3\textsuperscript{a} | 2.1 ± 0.2\textsuperscript{a} |
| Stem CO\textsubscript{2} efflux (\(\mu\text{mol CO}_2 \cdot \text{m}^{-2} \cdot \text{s}^{-1}\)) |        |       |           |
| CO                   | 2.8 ± 0.8\textsuperscript{a} | 2.9 ± 0.7\textsuperscript{a} | 3.3 ± 0.8\textsuperscript{a} |
| TE                   | 3.3 ± 0.7\textsuperscript{a} | 2.8 ± 0.2\textsuperscript{a} | 3.3 ± 0.6\textsuperscript{a} |
| Soil CO\textsubscript{2} efflux (\(\mu\text{mol CO}_2 \cdot \text{m}^{-2} \cdot \text{s}^{-1}\)) |        |       |           |
| CO                   | 6.7 ± 0.4\textsuperscript{a} | 5.4 ± 0.3\textsuperscript{b} | 6.6 ± 0.4\textsuperscript{a} |
| TE                   | 1.7 ± 0.1\textsuperscript{c} | 1.8 ± 0.1\textsuperscript{c} | 2.1 ± 0.2\textsuperscript{c} |

3.5 | Changes in C source/sink relations upon watering

Before watering, light-saturated CO\textsubscript{2} assimilation rates (\(A_{\text{sat}}\)) were 2.7 ± 0.2 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) and thus hardly higher in CO compared to TE spruce with 2.1 ± 0.3 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) (Table 1; \(p > .6\)). Watering did not significantly affect the \(A_{\text{sat}}\) of TE spruce, which remained almost constant under both treatments until day 14 (on day 4: CO, 2.6 ± 0.3; TE, 2.4 ± 0.3; on day 14: CO, 2.3 ± 0.3; TE, 2.1 ± 0.2 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\)). Similarly, the rates of stem CO\textsubscript{2} efflux did not significantly differ between treatments before watering (Table 1; \(p > .9\)), although the CO\textsubscript{2} efflux was slightly higher in TE with 3.3 ± 0.7 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) compared to CO spruce with 2.8 ± 0.8 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\). Upon watering, the stem CO\textsubscript{2} efflux rates remained almost constant with 2.9 ± 0.7 and 3.3 ± 0.8 in CO, and 2.8 ± 0.2 and 3.3 ± 0.6 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) in TE trees on days 4 and 14.

Before watering, rates of soil CO\textsubscript{2} efflux were 1.7 ± 0.1 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) under TE trees, which were much lower than under CO trees with 6.7 ± 0.4 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) (\(p < .01\); Table 1). Soil CO\textsubscript{2} efflux rates under TE trees around 2.0 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) hardly increased after watering and remained significantly lower than those under CO trees with 5.4 ± 0.3 and 6.6 ± 0.4 \(\mu\text{mol m}^{-2} \cdot \text{s}^{-1}\) on days 4 and 15, respectively.

4 | DISCUSSION

This study aims to elucidate the whole-tree C transport in highly productive Norway spruce forests upon watering in a long-term climate-change experiment with repeated experimental summer droughts. In the last decades, Norway spruce forests have been showing immense dieback through severe drought (Arend et al., 2021; Boczoń et al., 2018; Hentschel et al., 2014; Rosner et al., 2016; Solberg, 2004). Also in our experimental site, we lost a couple of TE spruce trees during the drought period (Grams et al., 2021). In the present study, we ask whether surviving trees recover both the aboveground C transport, that is, from the crown to the trunk base, and the belowground C transport, that is, from the trunk base to the soil CO\textsubscript{2} efflux after drought release. As the third transport process, we show how fast the current photoassimilates are incorporated in fine roots after drought release.

4.1 | Aboveground transport from crown to trunk base recovered within 2 weeks after drought release

The observed CTR\textsubscript{above} of CO spruce (c. 0.30 \(\text{m h}^{-1}\)) is somewhat higher than the average of gymnosperm trees calculated in a meta-analysis (0.22 \(\text{m h}^{-1}\), Liesche et al., 2015), and corresponds to the values observed at the same site 10 years before (Kuptz et al., 2011). The repeated summer droughts restricted the CTR\textsubscript{above} of mature spruce. In the first week after drought release, the arrival of \(^{13}\text{C}\)-depleted tracer was still delayed by 2–3 days in TE trees, indicating a 46% reduction in CTR\textsubscript{above} compared to CO spruce (Figure 5a). In a counterpart experiment with pine trees growing on a naturally dry site (Gao et al., 2021), CTR from crown to rhizosphere doubled upon watering, similar to findings on mature spruce trees in the present study. This delay was likely to be caused by longer MRT of sugars in leaves (Dannoura et al., 2019; Epron et al., 2012; Hesse et al., 2019; Ruehr et al., 2009; Zang et al., 2014) and/or slower phloem transport (Hesse et al., 2019; Sevanto, 2014).

About 2 weeks after watering, CTR\textsubscript{above} of TE trees significantly increased to the level of CO trees, while CTR\textsubscript{above} of CO trees remained constant (Figure 5b). However, neither C source strength, that is, photosynthesis rates nor sink strength, assessed here as stem and soil CO\textsubscript{2} efflux, significantly increased within the first 2 weeks after watering (Table 1). Likewise, unaffected soil CO\textsubscript{2} efflux rates during 2 weeks after drought release were also observed in other Norway spruce forests, likely due to slow recovery of microbial activity (Muhr & Borken, 2009; Schindlbacher et al., 2012). Considering that ratio of autotrophic (root-derived) to heterotrophic (microbial) soil respiration under the present spruce trees is known to decrease during drought (Nikolova et al., 2009), autotrophic respiration also likely remained low after drought release. Thus, changes of C source/sink relations are unlikely to be a major cause for the impaired CTR\textsubscript{above}. This led to the rejection of H1 that CTR\textsubscript{above} would recover with C source and/or sink strength, which is different from the study of Gao et al. (2021) on pine trees.

Although not fully recovered to the rather constant level of CO trees, \(\pi_o\) of TE trees increased until day 22 after watering in parallel with \(\Psi_{pd}\) (Figure 6). This indicates a declined C demand for osmotic adjustments, implying a decrease in leaf sugar concentration and MRT after drought release. Therefore, the delayed
sugar export from leaves under drought was likely a component of slower C translocation from the crown to the trunk base. A quick recovery of MRT of sugars in leaves was also observed in beech saplings after drought release (Zang et al., 2014). In addition, since \( \Psi_{\text{soil}} \) of TE trees was significantly lower than that of CO trees before watering and increased to the control level by day 7 (Figure 6), increased phloem viscosity due to water limitation in the xylem might be another cause for the slower phloem transport under drought (Epron et al., 2016; Woodruff, 2014). In principle, CTR may be reduced by intensified leakage–retrieval of transported sugars in the phloem (van Bel, 2003; De Schepper et al., 2013; Epron et al., 2016), however, there is no evidence to date that this mechanism is enhanced under drought (Salmon et al., 2013). Considering the rapid increase in CTR above within 2 weeks, reduction in phloem conduit diameter is unlikely to have occurred, which is in line with unaffected branch phloem lumen area of the same TE spruce trees (Giai Petit, University of Padova, in preparation). Miller et al. (2020) also reported an unaffected sieve cell production of mature spruce under summer drought. Furthermore, phloem production of the present spruce likely peaked before watering under moderate water stress (c. −0.9 MPa), since it has been found to peak before mid-June in spruce trees (Grčar et al., 2014; Jyske et al., 2015; Miller et al., 2020). This explains the different results on other tree species including conifers, which decreased phloem growth and diameter under more severe water stress (Dannoura et al., 2019; Woodruff, 2014).

It is important to note that the xylem water potential and to some extent also \( \pi_x \), were continuously increasing after watering until the \(^{13}\text{C}\)-depleted tracer arrived in stem CO\(_2\) efflux around day 7 in TE trees. Therefore, the drought-induced reduction in CTR above might have been even more pronounced before the watering. Most importantly, the aboveground CTR from crown to trunk base of mature spruce fully recovered within 2 weeks after watering, hence showing high resilience to long-term and recurrent summer droughts.

4.2 | Belowground transport from trunk base to soil CO\(_2\) efflux was similar between treatments already in the first week after watering

The observed CTR below of CO trees (c. 0.03 m h\(^{-1}\)) was about 10 times lower than CTR above (Figure 5a), which is in line with the study of Mencuccini and Hölttä (2010) reporting on a slower belowground transport compared to transport along the stem phloem. The variance of CTR below in TE trees was high, likely due to the soil heterogeneities and unknown root structures from the trunk base to the soil of CO\(_2\) efflux assessments. Already in the first week after watering, CTR below was similar between CO and TE trees. However, conversely to our expectation, rates of soil CO\(_2\) efflux did not increase after watering (as discussed above), which led to the rejection of H2 that CTR below would recover in parallel with increasing C sink or source strength. Upon watering, water potential in leaves fully recovered within 1 week (Figure 6) and can be expected to have increased faster in roots in parallel with increasing soil water potential (Fiscus, 1972; Gleason et al., 2017; McCully, 1999). Therefore, we suggest a fast and full recovery of root phloem transport within few days, that is, even before the \(^{13}\text{C}\)-depleted tracer arrived at the trunk base (i.e., around day 7). Moreover, speed of soil CO\(_2\) diffusion was likely similar in soils of both treatments, as gas diffusion in soils is negatively correlated with soil water content (Kuzyakov & Gavrickova, 2010) that was very similar in TE and CO plots within few days after watering (Grams et al., 2021).

Since the distance of the aboveground transport is much longer than belowground in tall mature trees, particularly in shallow rooting spruce trees, the drought-reduced transport rates from crown to soil CO\(_2\) efflux are mainly caused by the restricted aboveground C transport from crown to trunk base. However, short young trees or deep rooting mature trees have higher ratio of belowground to total transport distance. Thus, the belowground C transport from trunk base to soil CO\(_2\) efflux might play a significant role for the whole-tree transport processes and forest C cycling (Gao et al., 2021), since CTR below is much lower than CTR above. Most importantly, not only aboveground but the whole-tree CTR from crown to soil CO\(_2\) efflux showed a full recovery within 2 weeks after watering, hence indicating high resilience to long-term and recurrent summer droughts.

4.3 | Incorporation of current photoassimilates in fine roots was faster in trees recovering from drought than in control trees

The \(^{13}\text{C}\)-depleted tracer was detected in living root tips of TE trees within 12 h after the arrival at the trunk base, but only 60 h later in CO trees (Figure 7a), confirming H3 that incorporation of current photoassimilates is faster in trees recovering from drought. The faster use of the tracers in living root tips of TE trees compared to controls coincided with the growth of new roots that started within few days after watering (personal observations on site), suggesting a higher C demand in fine roots of TE trees. However, the enhanced fine root growth upon watering in TE plots was not reflected in soil CO\(_2\) efflux, likely due to a small contribution of respiration of fine roots grown after watering to total soil CO\(_2\) efflux: that is, small biomass share of growing fine roots to total roots. Furthermore, Nikolova et al. (2020) found on the same spruce trees that respiration rates of fine roots and proportion of absorptive fine roots to the total root biomass were both small. A preferential investment of current photoassimilates following high C sink strength of growing fine roots has also been observed in young beech trees upon drought release (Hagedorn et al., 2016) and in naturally drought-stressed mature pine trees after a rainfall event (Gao et al., 2021; Joseph et al., 2020).

Not only previously drought-stressed spruce but also control trees responded with fast C incorporation in living fine roots after increase in soil water availability. During an intensive rain event on day 17 (following a short dry spell), fine root growth was likely induced in the shallow soil layers (Joseph et al., 2020; Meier & Leuschner, 2008). This may explain the fast arrival of unlabeled tracer in root
tips in both treatments 2 weeks after drought release (Figure 7b). Our results suggest, therefore, that the speed of incorporation of current photoassimilates in living root tips of mature spruce trees is strongly dependent on the C demand for root production, that is, “sink controlled” as suggested earlier (Fatchi et al., 2014; Gavito et al., 2019; Hagedorn et al., 2016; Körner, 2015). In contrast to previous studies (Gao et al., 2021; Hagedorn et al., 2016), the increased “sink demand” by stimulated fine root growth in mature spruce did not significantly affect the whole-tree CTR from crown to soil, since they were still reduced in the first week after drought release (Figure 5a). Above all, mature drought-stressed spruce trees respond to drought release by quickly supplying the growing root tips with current photoassimilates. In addition to the high resilience in whole-tree C transport, this response is essential to regenerate the water-absorbing root system.

5 | CONCLUSION

The present study reveals high resilience of the whole-tree C transport system in Norway spruce forests even after recurrent summer droughts. Once spruce trees manage to survive drought periods, their whole-tree C transport system may be expected to recover quickly after drought release. This ensures high resilience of C supply with current photoassimilates, in particular to belowground sinks such as growing fine roots. Once the water-absorbing root system is restored, long-term recovery of C uptake and supply to further sinks can be expected. However, recovery of the C transport is only one of the many important prerequisites for the recovery of tree productivity. Thus, long-term observations of C source and sink activities upon drought release are necessary to elucidate the recovery potential of productivity in central European forests dominated by Norway spruce stands.

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CONFLICT OF INTEREST

The authors declare that there is no conflict of interest.

AUTHOR CONTRIBUTIONS

TEEG and KP designed the experiment. KH, VR, and TEEG prepared and conducted the \(^{13}\)C labeling experiment. All authors contributed to the data collection. KH analyzed and interpreted the data with supports from TEEG, JD, and VR. KH wrote the manuscript with contributions of all authors.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study are openly available in mediaTUM at https://doi.org/10.14459/2021mp1637391.

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