Effects of below-ground space limitation on performance of Eucalyptus seedlings: Does photosynthesis really control growth?

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Abstract

Interpreting limitations to plant growth requires understanding of the balance between carbon (C) source and sink activity in order to assess C allocation and biomass partitioning. This study used manipulations of soil volume to test how growth is coupled to physiology, allocation, and sink activity in Eucalyptus tereticornis seedlings. We grew seedlings in a large range of container sizes and planted containers flush to the soil alongside naturally sown seedlings (free). Reduced soil volume was expected to induce rapid negative effects on growth and physiology compared to free seedlings. It was hypothesized that soil volume effect would be largest in the smallest containers, resulting in physical constraints to growth independently of photosynthesis (A). Photosynthesis would then become sink-limited, resulting in the build-up of leaf nonstructural carbohydrates eventually leading to photosynthetic down regulation. We observed a negative container effect on aboveground growth soon after the experiment started. Although growth was consistently different across soil volumes mass, partitioning to leaves, stems, roots was conserved after 120 days. Photosynthetic capacity was also significantly reduced in containers, and was related to both leaf nitrogen content and starch accumulation. We developed a seedling growth model that utilized leaf A rates to allocate daily C uptake towards mass growth of stems, leaves and roots. We then asked whether the observed reductions in A explained the observed differences in seedling biomass. We found that although belowground sink limitation resulted in the down regulation of A, these reductions could not explain observed growth responses. Thus, as photosynthesis and growth were not coordinated a pool of excess C resulted in seedlings with soil volume restriction. This research highlights the need to further utilize mass balance approaches when evaluating plant C allocation and confirms that A and growth are not always directly related.

Key Words

photosynthesis, sink regulation, growth, carbon allocation, soil volume

Introduction

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Understanding plant growth requires knowledge of the mass balance that must be achieved between *C* uptake and subsequent allocation to growth, storage, and respiration. It is commonly
assumed plant growth is limited by *C* availability, yet it has long been demonstrated that correlations between *A* and growth are weak or seldom present. Körner (2013) argues that growth instead
controls *A*, as it is the norm for sink activity to feedback on source activity. This is supported by
evidence that growth of plants under environmental stress is not limited by the supply of photoassimilates (Palacio et al., 2014). As woody plants have highly integrated systems of competing
carbohydrate sinks (Kozlowski, 1992), growth should principally depend on the transport of these
photoassimilates between different tissue and organ sinks. Despite a wealth of studies, however,
large uncertainties still remain regarding the coordination of *C* supply, via *A*, and growth of woody
species.

In woody species, how A controls growth has been studied with manipulations of C source activity. Examples included elevated CO₂ experiments, for example FACE (reviewed in Ainsworth and Long (2005), and partial defoliation experiments. Elevated CO₂ has been shown to increase A rates (Drake et al., 1997; Ainsworth and Rogers, 2007) and across four FACE experiments this resulted in a conserved increase in forest production (Norby et al., 2005). Evidence from elevated CO₂ experiments, however, also reveals that the growth response tends to be much smaller than the photosynthetic enhancement (Kirschbaum, 2011). In defoliation experiments, compensatory increases in A are commonly shown yet are attributed to variable mechanisms, including reduction in end product inhibition (Iglesias et al., 2002; Zhou and Quebedeaux, 2003; Handa et al., 2005), enhanced biochemical activity (Ovaska et al., 1993a; Layne and Flore, 1995), increased stomatal conductance (Layne and Flore, 1995), leaf nutrient status (Turnbull et al., 2007), and regulatory sugar signaling (Eyles et al., 2013). However, increases in A did not always produce increased growth due to reductions in meristem sink strength (Palacio et al. 2012). C limitation to mycorrhizal colonization (Markkola et al. 2004), or an overall decrease in whole plant C gain (Ovaska et al., 1993b). These manipulations of C source activity expose unresolved issues with how changes in A do not always infer similar responses in growth.

Alternatively, tissue sink activity can restrict biomass production when limited by environmental or developmental constraints (Körner, 2003). This is because metabolic signaling networks, relaying information on C and nitrogen (N) status of different tissues, can down regulate photosynthetic activity (Paul and Fover, 2001). Whether this down regulation, via sink inhibition, exists in woody species has been tested through fruit removal, girdling, and low temperatures. In these studies, down regulation of A was frequently correlated to carbohydrate accumulation resulting from reduced tissue sink strength (Iglesias et al., 2002; Hoch et al., 2002; Urban and Alphonsout, 2007; Haouari et al., 2013). However, reductions in A were also attributed to biochemical limitations prior to carbohydrate accumulation (Nebauer et al., 2011), irreversible photo-oxidative damage (Duan et al., 2008), and stomatal limitation (Li et al., 2005). These mixed results are not surprising as we still know little about the pathways in which plants achieve balance between assimilation, storage, and growth across temporal scales (Smith and Stitt. 2007). As these manipulations likely impact source as well as sink activity simultaneously, affect water transport, are very extreme, or are specific to large annual fruiting sinks, they tell us little about source-sink coordination in normal field-grown woody species. This coordination is further confounded by the fact that under normal growing conditions, A is not always correlated to photosynthate accumulation, as with fir needles (Little and Loach, 1973). Unfortunately, there is still limited understating of the physiological roles of carbohydrates in photosynthetic regulation and the elements triggering the down-regulation process (Nebauer et al., 2011).

An alternative approach is to lower belowground *C* sink strength in tree seedlings by manipulating rooting volume, by varying the container size. The advantage of this approach is it allows a large range of manipulations, can be easily compared to naturally planted seedlings and mimics natural conditions as seedlings compete for space or reach bedrock. Seedlings undergo many physiological and morphological changes in response to rooting volume, including biomass partitioning, *A*, water relations, nutrient uptake and respiration (NeSmith and Duval, 1998, and refer-

ences therein). The rooting volume available to plants can decrease *C* sink strength by reducing root growth. Container size studies frequently have photosynthetic down-regulation, likely as a result of sink limitation (Arp, 1991; McConnaughay and Bazzaz, 1991; Gunderson and Wullschleger, 1994; Sage, 1994; Maina et al., 2002; Ronchi et al., 2006). A meta-analysis by Poorter et al. (2012a) concluded that *A* is the process likely to be the strongest affected by pot size and may best explain the observed effect on biomass seen in the large number of studies where containers are used. This conclusion arises because plants grown in small containers are shown to accumulate leaf starch while having lower *C* exchange and assimilate export rates (Robbins and Pharr, 1988). However, evidence in support for a trade-off between *C* storage and growth in trees is, to date, inconclusive (Palacio et al., 2014). Based on these previous studies, using container size as a sink-strength manipulation can be used to empirically test how growth and *A* are coordinated.

Another barrier in understanding the coordination between A and growth is a lack of knowledge 99 regarding the allocation of C in woody species. Understanding the fate of assimilated C is cru-100 cial when source or sink manipulations do not directly lead to equivalent changes in growth. As programmed plasticity in C allocation naturally occur as plants growth and develop, this must be accounted for when evaluating induced variation in growth (Wright and Mcconnaughay, 2002). 103 Thus, when C allocation is altered by a treatment, it should be shown whether these allocation patterns differ from common sized untreated plants (Reich et al., 2002; Poorter et al., 2012b). Only then can we test if functional balance preservation or optimal foraging occurs from changes in C uptake. For example, seedlings in containers may shift allocation to leaves if root restriction occurs or alter fine root morphology to increase nutrient uptake. In woody species, shifts in allo-108 metric relationships of leaves, stems and roots can reveal if growth is ontogenetically constrained or actively adjusted when A is affected. This then allows additional pools of C, such as root exudation or changes in tissue respiration, to be accounted for and evaluated.

This study utilizes a novel field design to investigate the coordination between growth and *A* in Eucalyptus tereticornis Sm. seedlings, by manipulating container size and thus rooting volume.

Seedlings were maintained under well watered conditions in order to evaluate only the effect of restricted soil volume and the limited nutrient resource pool. We used freely-rooted seedlings as a control for the container size treatments. Empirical results were combined with a simple plant growth model to simulate seedling growth with a *C* mass balance approach, which was then compared to observed harvested seedling mass. The model used whole-plant *C* gain, scaled from instantaneous rates of leaf *A*, to quantify seedling production over 120 days.

- 1). The manipulations of container size were hypothesized to induce a belowground sink limitation in these seedlings which was expected to be largest in the smallest containers, resulting in physical constraints to growth independently of *A*.
- 2). Reducing soil volume was expected to induce rapid negative effects on growth and physiology compared to free seedlings (container effect), with accumulation of leaf nonstructural carbohydrates triggering photosynthetic down regulation through time as a function of available soil volume.
- 3). Last, the growth model was expected to find agreement between simulated and observed seedling mass, through direct correlation of the effects of soil volume on rates of leaf *A*.

Methods

Experimental Design

This experiment was located on the Hawkesbury Forest Experiment site in Richmond, NSW, Australia. Plots were located in open cover with a site history that consists of a paddock that was converted from native pasture grasses. Top soils at this site, used for the study, are an alluvial formation of low-fertility sandy loam soils (380 and 108 mg kg⁻¹ total *N* and phosphorus respectively) with low organic matter (0.7 %) and low water holding capacity. At this site a soil hard layer exists

 $_{96}$ at \sim 1.0 m with a transition to heavy clay soils. The climate for the region is classified as sub-humid temperate.

Eucalyptus tereticornis seedlings, 20 weeks old and approximately 40 cm tall in tube stock, were chosen from a single local Cumberland plain cohort. Previous experiments have confirmed that species with tap roots (similar to *E. tereticornis*) use the center of the container as the medium for thick roots leaving the periphery of the soil as the most active sites for fine root proliferation (Biran and Eliassaf, 1980a,b). This is generally hypothesized to be a different response than seedlings with no taproot. By using a species with tap root growth and manipulations of container length rather than width, it is believed that a more realistic test of inhibition of growth through constrained soil volume would be achieved. Six seedlings were harvested before planting to measure initial leaf area and dry mass of leaves, stems and roots.

Six container volumes were used ranging from 5 l to 35 l, with a 22.5 cm diameter, and lengths ranging from 15 to 100 cm. Containers were constructed of PVC pipe and were filled with local top soil (described above). Soil in each container was packed to achieve a target soil bulk density of 1.7 g m⁻³. A Imidacloprid (BAYER CropScience) insecticide tablet was planted 5 cm below the roots of each seedling. Containers were planted flush with the soil surface inside metal sleeves, designed to minimize excess air space between the container and outside soil while also allowing for container removal. This allowed for soil temperatures in containers to reflect conditions of naturally sown (free) seedlings. Each experimental block (n=7) contained a complete replicate set of container volumes as well as one free seedling, with 1 m² spacing. For each free seedling, a 1 m² subplot was excavated to 0.5 m and replaced with the same soil used in each container. A border of root exclusion material was buried 0.25 m deep and extended 0.25 m above the ground surface around each subplot to exclude local vegetation.

Plants were watered weekly or when needed, accounting for natural precipitation, to maintain soil moisture at field capacity (13-15 %). Drain systems were built into each pot to prevent pooling of water in containers before root expansion, from reduced root uptake, or from large rainfall events. These conditions could lead to an anaerobic environment around the root that could hinder the uptake of water through reduced root conductance (Poorter et al., 2009), an undesired experimental artifact. A collection compartment in the bottom of containers, containing gravel covered by root exclusion mesh, was used to collect excess water for 20, 25, and 35 l containers. Plastic tubing (6 mm diameter) was inset into the gravel layer and extended through the top of the container. A lysimeter pump was then used to suction excess water, through the tubing, as needed. As small containers (5, 10, and 15 l) have a larger irradiation effect a simple bottom plug was used to drain excess water from the gravel compartment.

Growth and morphology metrics

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Seedlings were planted on January 21st 2013 and stem height, diameter at 15 cm and leaf count were measured weekly thereafter. Once the growth rate of individual plants had significantly declined a full biomass harvest was completed (May 21st 2013). Dry mass of leaves, stems, roots and cumulative leaf area (LI-3100C Area Meter; LI-COR, Lincoln, NE, USA) was measured for each seedling. Mean individual leaf area for each harvested seedling was calculated by dividing cumulative leaf area by total leaf count of only fully expanded leaves. This value was then used to interpolate cumulative leaf area through time with weekly leaf counts. Root mass was collected by passing soil from each container through a 1 mm sieve, washing, separating into fine and coarse roots (<2 mm and >2 mm diameter, respectively) and then drying to a constant mass. Roots from the free seedlings were collected by excavating each 1 m² subplot to 0.5 m depth. 25 g fresh weight subsamples of washed fine roots were analyzed, using WhinoRhizo software (Regent Instruments Inc.), for specific root length (SRL, cm m⁻¹).

Photosynthetic parameters

Leaf gas exchange measurements were performed bi-weekly at saturating light (Asat) and saturat-184 ing light and [CO₂] (A_{max}) on new fully expanded leaves. Measurements were initiated only after 185 sufficient new leaf growth occurred (March 17th, 2013), approximately 6 weeks following planting, and continued until the biomass harvest. Leaf level gas exchange was measured with a standard leaf chamber equipped with blue-red light emitting diodes using a portable gas exchange system (LI-6400, LI-COR, Lincoln, NE, USA). $A_{
m sat}$ measurements were made at PPFD of 1800 μ mol m $^{-1}$ s $^{-1}$ and $[CO_2]$ of 400 μ l l^{-1} and A_{max} with $[CO_2]$ of 1600 μ l l^{-1} and PPFD of 1800 μ mol photons m⁻¹ s⁻¹. This choice of light level to achieve light saturation is consistent with other studies on *Eucalyptus* species (Kallarackal and Somen, 1997; Pinkard et al., 1998; Crous et al., 2013; Drake et al., 2014). These measurements were conducted during midday (10:00-14:00 h) with leaf temperature maintained at 25°C. After leaves acclimated to the chamber environment, net CO_2 assimilation rate and stomatal conductance (g_s) were logged 5 times for both A_{sat} and A_{max} . Photosynthetic CO_2 195 response (AC_i) curves were also developed at 25°C on a random subset of each container size (n=3) after new leaves were first produced and immediately prior to the final harvest (May 23rd 2013). Each AC_i curve began at the reference [CO₂] of 400 μ l l⁻¹ and then consisted of 12 additional steps from [CO $_2$] of 50 to 1800 μ l l 1 at 25 $^\circ$ C at saturating light (above). From these curves the photosynthetic parameters, J_{max} and Vc_{max} , were quantified using the biochemical model of Farquhar 200 et al. (1980). Leaf dark respiration rates (R_d) was measured on each seedling during the same dates as AC_i curves using detached leaves inside a conifer chamber attached to the Licor 6400 at least 1 hour after sundown. Measurements were taken at a reference [CO2] of 400 µl l-1 while 203 leaf temperature was maintained at current ambient conditions. Specific leaf area (SLA, m² kg⁻¹) was calculated by measuring leaf area and dry mass for all individual leaves sampled during gas exchange campaigns.

107 Leaf water potential

Predawn (Ψ_{pd}) and midday (Ψ_{l}) leaf water potentials were measured for each seedling using a PMS 1505D pressure chamber (PMS Instruments, Albany, OR, USA) on fully expanded leaves during the same time period as AC_i and R_{d} . Leaves were detached and immediately stored inside foil covered bags before water potential measurements were performed. Ψ_{pd} was measured before sunrise and Ψ_{l} at midday 13:00-14:30 h. These measurements were used as a measure of static water stress on the seedlings (Sellin, 1999), and to ensure that the bulk soil water availability was high enough for plants as they became larger and roots filled the soil volume.

Leaf, root and soil chemistry

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Leaves used in each gas exchange measurements and subsamples of harvested roots were dried to a constant mass and milled for analysis of N content, δ^{13} C, and total non-structural carbohydrates (TNC). Pre-planting soil samples (n=6) and subsamples of soil from each container following harvest were sieved to remove organic material, air dried and milled for analysis. N concentrations of leaf and soil samples were determined using a Carlo Erba CE1110 elemental analyzer with thermal conductivity and mass spectromic detection (of N_2 and CO_2). The percentage of N in the sample was calculated by comparison with known standards. Leaf δ^{13} C was analyzed with an Delta V Advantage coupled to a Flash HT and Conflo IV isotope ratio mass spectrometer. Leaf samples were flash combusted at 1000°C to convert to to CO_2 , feed to the mass spectrometer and isotopic signatures are reported relative to the VPDP scale. Leaf TNC concentration was analyzed using a total starch assay kit (Megazyme International 303 Ireland Ltd., Wicklow, Ireland) and includes the starch (mg g⁻¹) and soluble sugar (mg g⁻¹) concentrations. Starch was quantified using a thermostable α -amylase and amyloglucosidase assay (McCleary et al., 1997) and soluble sugars were determined following the anthrone method (Ebell, 1969). Complete methods of the TNC assay are described in (Mitchell et al., 2013).

Seedling growth model

We developed a simple seedling growth model that utilized leaf *A* rates to allocate daily *C* assimilate towards biomass production of stems, leaves, fine roots and coarse roots. The model begins with mean initial tissue component biomass (leaf_i, stem_i and root_i) and a starting leaf area (LA_i) measured prior to planting. The initial biomass of roots was divided evenly between fine and coarse roots. The daily net biomass production of seedlings (P_i) is then given by

$$P_{\rm i} = L \, C_{\rm dav,i} \, \epsilon_{\rm c}^{-1} - R \tag{1}$$

where L is standing cumulative leaf area, $C_{day,i}$ is the predicted daily carbon gain, ϵ_c is a biomass conversion efficiency parameter and R is the total respiration of all tissue components. Total respiration was calculated as

$$R = \Sigma(R_{\mathsf{c}} M_{\mathsf{c}}) \tag{2}$$

where R_c is tissue respiration of fine roots, coarse roots or stems on a mass basis and M_c is the standing biomass of each component. R_{leaf} is represented in the calculation of C_{day} . The change in individual component biomass, here solved on a daily time step, is given by

$$\frac{dM_{c}}{dt} = A_{c} P_{i} - (\Lambda_{c} M_{c}) \tag{3}$$

where A_c is the component specific biomass partitioning to whole plant biomass (%) and Λ_c is component specific turnover rate. Due to the duration of the experiment Λ_{stem} was assumed to equal o. Total seedling biomass, per time step, was then equal to the sum of all biomass components; leaves, stems, fine roots and coarse roots.

 $C_{\rm day}$ was predicted by fitting a coupled photosynthesis - stomatal conductance model (Farquhar et al., 1980; Medlyn et al., 2011) in the 'plantecophys' package in R (Duursma, 2014) to the mean photosynthetic parameters (R_d, J_{max}, V_{cmax}, and g₁) for each treatment and meteorological data from an onsite weather station. Examples of the photosynthesis model are described in Medlyn et al. (2002) and the approach of the coupled leaf gas exchange model are described in Duursma et al. (2014). The g₁ parameter was generated by fitting observed g_s values into the optimal stomatal conductance model from (Medlyn et al., 2012). Combined with the meteorological parameters; PPFD, air temperature, and relative humidity, at 15 m intervals, leaf A rates (µmol CO₂ m⁻² s⁻¹) were predicted for each soil volume treatment. Rates were assumed to be representative of the entire 15 min meteorological interval. $C_{\rm day}$ was calculated by converting predicted rates to mass $C_{\rm day}$ for each soil volume treatment, one value for each day of the experiment. Thus, each daily time step for model runs included a value of $C_{\rm day}$ that represented both treatement specific photosynthetic parameters and meterological constraints across the duration of the experiment. All default parameters used in model simulations are reported in Table. 3.

We then utilized this model to test the hypothesis that the effects of belowground sink limitation on rates of leaf A where sufficient to accurately predict overall seedling production after 120 days. Each model was run with a sequence of C_{day} values (n=101) covering the entire range of predicted values of all soil volume treatments and the free seedling control. First, the model was simulated using only changes in A rates across treatments, via C_{day} , combined with mean values of mass partitioning and either published or local data of stem and root respiration rates (M_0). Next, the model was fit with with treatment specific mass partitioning from the final harvest (M_1) in order to determine how ontogenetic differences in seedling allometry improved biomass predictions. Finally, model sensitivity to 3 different C allocation scenarios, including changes to leaf C allocation, root respiration rates, and fine root C allocation ($M_{2,3,4}$ respectively), was tested to investigate possible sources of missing C from initial model simulations. This was accomplished by testing the sensitivity of the model to adjustments of C allocation to leaves, fine roots or root respiration rates by ± 50 % of default values. For all cases, biomass production was compared between model output and harvested seedlings.

₇₆ Data Analysis

Differences in experimental parameters with soil volume were analysed by one-way analysis of variance (ANOVA) in R with individual containers as random effects and soil volume as a categorical fixed effect. Tukeys post-hoc tests were performed in conjunction with ANOVA to determine which specific paired comparisons among soil volume treatments were different. Mixed model ANOVAs of A_{max} and leaf chemistry were performed using the 'nlme' package (Pinheiro et al., 2014) in R and r^2 values of mixed models were computed as in Nakagawa and Schielzeth (2013). Tests of allometric relationships between biomass components were implemented using major axis regression in the 'smatr' package in R (Warton et al., 2012). Results were considered significant at P < 0.05.

Sesults

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287 Growth and morphology metrics

In this open field study, colder temperatures and reductions in cumulative PPFD per day (Fig. 1) most likely lead to the reduced growth in the free seedlings in the final weeks of the experiment (Fig. 2). Combined with severe growth reductions in the smallest container volumes the experiment was chosen to be harvested after 120 days. Over this duration height, diameter, and leaf area diverged between container volumes (Fig. 2). First, seedling leaf area significantly diverged between soil volumes (P < 0.026) during the 5th week of the experiment. Following this period both height (8th week) and then diameter (9th week) significantly deviated across soil volumes (P < 0.002 & 0.001, respectively). Negative growth effects then manifested as severely reduced height gain and declining leaf area through time with small soil volumes across the final two months of the experiment. Seedlings maintained diameter growth throughout the experiment, although marginal with smaller soil volumes in the final month. Final seedling height significantly increased with increasing soil volume (P < 0.001). Increases in both final stem diameter (P < 0.001) and cumulative leaf area (both P < 0.001) were found with increasing soil volume and these differences were driven mainly by the largest container and the free seedling.

Harvested total seedling biomass at harvest was significantly different across container volumes (P<0.001) and with free seedlings (P<0.001, Table. 1). We analyzed the relationship between biomass growth with each fold increase in soil volume and found an increase of 34 % with a doubling of pot size, consistent with the meta-analysis of Poorter et al. (2012a). Additionally, plant biomass was highly correlated with total leaf area across all treatments (r^2 = .97, P<0.001). Differences in biomass partitioning to leaves, stems, and roots were not different across soil volumes when variation in seedling biomass within treatments was factored in the analysis. Across all treatments, the final harvested root:shoot was conserved in these seedlings, with a slightly higher shoot than root mass on average (\bar{x} =0.904).

SRL of harvested fine roots was not different across soil volumes (Table. 1). Over the duration of the experiment SLA was higher in free seedlings but was not different across containers sizes (Table. 1, P<0.001) and this pattern was evident in the first gas exchange measurement campaign (P<0.001).

Leaf and root chemistry

Leaf N % was significantly higher in free seedlings and the largest container volume at the onset of gas exchange measurements (6th week, P<0.001). Over the remaining duration of the experiment the smallest container volume had a significant reduction in leaf N % compared to other soil volumes, while free seedlings maintained a greater leaf N % (Table. 1, P<0.001). Additionally, mean leaf starch content in the smallest container was double that of free seedlings (P=0.042), while leaf

soluble sugars did not differ across treatments throughout the experiment (Table. 1). Differences in leaf starch between the free seedling and the smallest container were also evident during the first gas exchange campaign (*P*=0.0013).

Gas exchange and photosynthetic parameters

 $A_{\rm sat}$ (Fig. 3) and $A_{\rm max}$ (Table. 2) were both significantly higher in the largest container volume and the free seedling at the first measurement campaign (both P < 0.001). Across measurement campaigns $A_{\rm sat}$ was consistently higher in free seedlings than in containers (Figure 3, P < 0.001). The interaction between photosynthetic capacity, leaf starch, and leaf N on a mass basis was marginally significant (P = 0.0584) but $A_{\rm max}$ was highly correlated to both leaf N content and leaf starch (both P < 0.001). Across all measurement campaign $A_{\rm max}$ was higher when foliar N was also higher, usually associated with low levels of leaf starch (Fig. 3a). $A_{\rm max}$ was also lower when leaf starch was high as higher leaf N often did not coincide with high leaf starch (Fig. 3b)

The photosynthetic parameters J_{max} and Vc_{max} were not different across measurement campaigns, therefore the parameter means per treatment are reported here (Table. 2). Overall, both J_{max} and Vc_{max} were significantly higher in free seedlings with little variation between soil volume treatments (P=0.0012 & 0.0021, respectively). Leaf dark respiration rates were not significantly different across soil volumes (Table. 2). The g_1 parameter, generated for each seedling from the Medlyn et al. (2012) optimal stomatal conductance model, was not different across soil volumes (Table. 2). Predicted values of g_s , using the g_1 parameter, where highly correlated with observed values (r^2 = .74, P<0.001, data not shown).

Neither $\Psi_{\rm pd}$ nor $\Psi_{\rm l}$ were different across treatments, with mean values of -0.27 and -1.2 mPa across all seedlings, respectively. Although $g_{\rm s}$ in free seedlings was generally higher than those in containers (Table. 2, P=0.0023), the mean rates for all seedlings were high at 0.37 mol H $_{\rm 2}$ 0 m $_{\rm 2}$ s $_{\rm -1}$ 1 and did not decline significantly across the experiment duration. Additionally, leaf $\delta^{\rm 13}$ C at final harvest was not different across treatments (Table. 1). Combined these indices provide strong evidence that water stress was not apparent on these well-watered seedlings throughout the experiment. Soil N % at harvest was not different across soil volumes (\bar{x} =04.5 %) and decreased approximately 3 % across all containers over the experiment duration. This indicates that nutrient leaching from free seedlings or from draining of containers following natural rainfall events did not differ between treatments.

Modelling seedling biomass

Model M_0 , testing only treatment specific photosynthesis rates, was found to overestimate seedling production when compared to harvested seedling biomass. When both C_{day} and biomass of modeled and harvested seedlings were scaled to the control treatment this overestimation was greater for seedlings with soil volume restriction (Fig. 6a). Model M_1 , using treatment specific leaf, stem and root biomass partitioning, also overestimated seedling production for all soil volume treatments (Fig. 6b). With both M_0 and M_1 this overestimation resulted in a unexplained pool of missing carbon in seedlings that was not attributed to biomass growth. Models $M_{2,3,4}$ were then utilized to investigate how different C allocation scenarios, not explicitly tested within the framework of this experiment, might account for these pools of unexplained carbon. Both decreases in fine root C allocation and increases in leaf C allocation slightly improved biomass estimates but neither were sufficient to accurately predict the observed biomass response (Fig. S1a,c). Altering respiration rates of fine and coarse roots did not have a noticeable deviation from the original model (Fig. S1b).

- ₃₆₅ Discussion
- 366 Tables

Table 1: Responses of plant and leaf characterisitcs of Eucalyptus tereticornis seedlings to soil volume treatments. Each value reflects the mean(standard

error) of each treatment.	ո treatment.							
Volume (L)	/olume (L) Seedling mass (g) SLA (m² kg¹) Leaf Nitrogen (%) Leaf Sugars (%) Leaf Starch (%)	SLA (m ² kg ⁻¹)	Leaf Nitrogen (%)	Leaf Sugars (%)	Leaf Starch (%)	SRL (cm m ⁻¹)	SRL (cm m ⁻¹) Root Nitrogen (%) 5 ¹³ C (%o)	5¹³C (‰)
2	14.8 (1.82)	9.5 (0.23)	1.1 (0.02)	6.4 (0.28)	12.7 (0.97)	39.1 (5.47)	0.82 (0.05)	-30.1 (0.26)
10	20.0 (2.38)	9.8 (0.24)	1.3 (0.04)	6.7 (0.25)	9.4 (0.75)	34.2 (5.83)	0.75 (0.02)	-30.2 (0.25)
15	25.4 (2.49)	11.0 (0.47)	1.4 (0.06)	7.2 (0.28)	7.3 (0.73)	37.6 (4.63)	0.71 (0.02)	-30.3 (0.36)
20	23.4 (1.63)	9.8 (0.28)	1.4 (0.05)	6.6 (0.26)	9.5 (0.88)	45.3 (5.50)	0.76 (0.04)	-29.7 (0.28)
25	30.4 (5.49)	10.4 (0.37)	1.3 (0.06)	6.9 (0.24)	9.8 (0.71)	47.0 (7.10)	0.74 (0.02)	-29.7 (0.25)
35	52.2 (9.55)	11.3 (0.44)	1.5 (0.08)	6.8 (0.22)	9.8 (0.65)	50.6 (11.61)	0.77 (0.03)	-30.6 (0.38)
Free	174.5 (18.02)	13.0 (0.44)	2.4 (0.09)	7.4 (0.25)	6.8 (0.65)	43.7 (6.24)	0.87 (0.04)	-30.0 (0.34)

Table 2: Responses of leaf level gas exchange parameters of *Eucalyptus tereticornis* seedlings to soil volume treatments. Each value reflects the mean(standard error) of each treatment. Units for A_{max} and R_{dark} are μ and R_{s} are μ and R_{s} are μ and R_{s} are μ of each treatment.

Volume (L)	$A_{\sf max}$	R_{dark}	\int_{max}	V_{Cmax}	g _s	ğ
5	21.2 (0.9)	2.8 (0.3)	103.7 (5.8)	62.9 (2.8)	0.30 (0.01)	5.1 (0.1)
10	22.3 (1.4)	2.7 (0.4)	116.5 (5.9)	69.4 (2.9)	0.36 (0.01)	5.4 (0.1)
15	23.3 (1.2)	1.4 (0.1)		79.4 (10.0)	0.45 (0.01)	6.2 (0.2)
20	26.1 (0.7)	1.7 (0.1)		81.4 (6.0)	0.38 (0.01)	5.2 (0.2)
25	23.9 (0.9)	1.2 (0.1)		78.6 (2.6)	0.32 (0.01)	4.8 (0.2)
35	25.0 (1.0)	1.5 (0.2)		78.0 (3.5)	0.33 (0.02)	4.7 (0.2)
Free	33.1 (0.7)	1.3 (0.1)	171.7 (18.6)	101.2 (5.7)	0.49 (0.02)	5.0 (0.2)

Figures

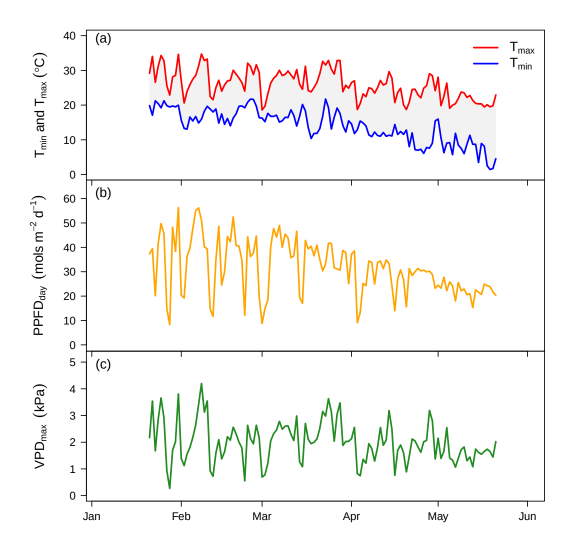


Figure 1: Daily maximum and minimum temperature (a), cumulative daily PPFD (b), and daily maximum vapour pressure deficit (c) across the experiment duration in 2013.

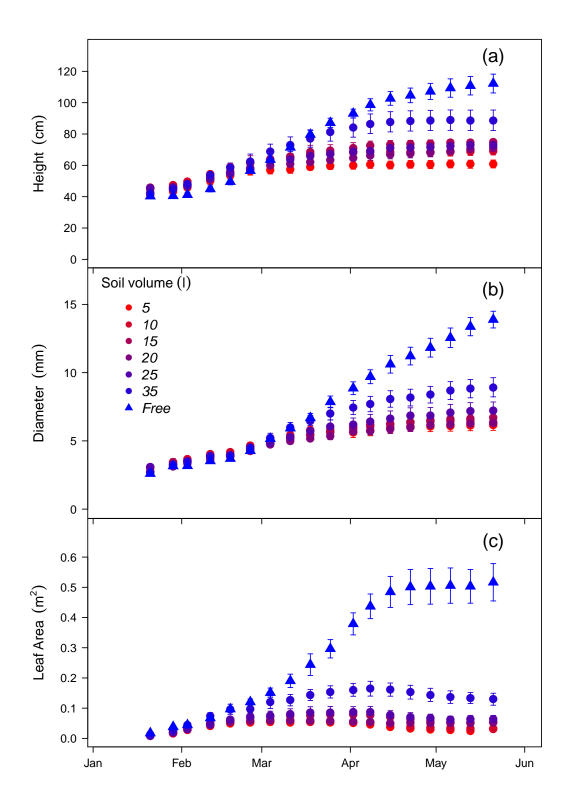


Figure 2: Soil volume treatment means \pm SE (n=7) of height growth (a), diameter growth (b), and interpolated seedling leaf area (c) measured weekly of *Eucalyptus tereticornis* seedlings across the experiment duration in 2013.

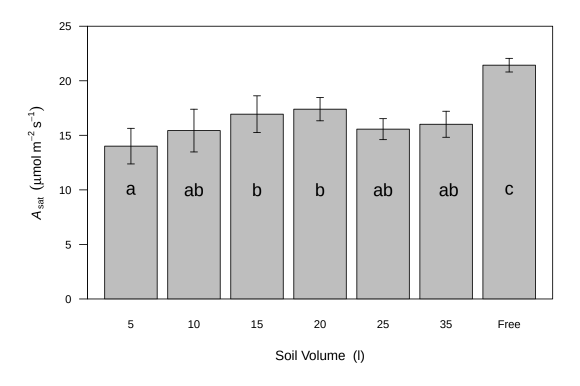


Figure 3: Soil volume treatment means \pm SE (n=7), across all measurement dates (n=6), of light saturated rates of photosynthesis at 25°C. Different letters represent significant differences between treatments.

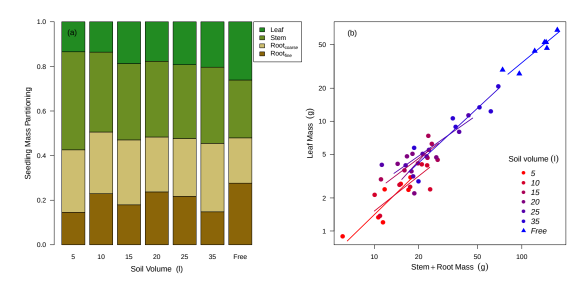


Figure 4: Soil volume treatment means (n=7) of mass partitioning to leaves, stems, and roots (a) and bi-variate relationships between mass allocation to leaves and stems + roots (b). Lines represent standardized major axis fitting of the log transformed allometric relationships of leaf mass fraction by treatment.



Figure 5: Photosynthetic capacity, on a leaf mass basis, as a function of accumulation of leaf starch (a) and leaf nitrogen content without TNC (b). Colors represent bins levels (n=5) of both leaf starch and nitrogen grouped from low to high . Lines represents predictions, for each bin level, from the linear mixed effects model equation of $A_{\rm max}$ as a function of starch and nitrogen. The marginal r^2 (fixed effects only) was 0.37 and the conditional r^2 (fixed and random effects) was 0.48 for the complete model.

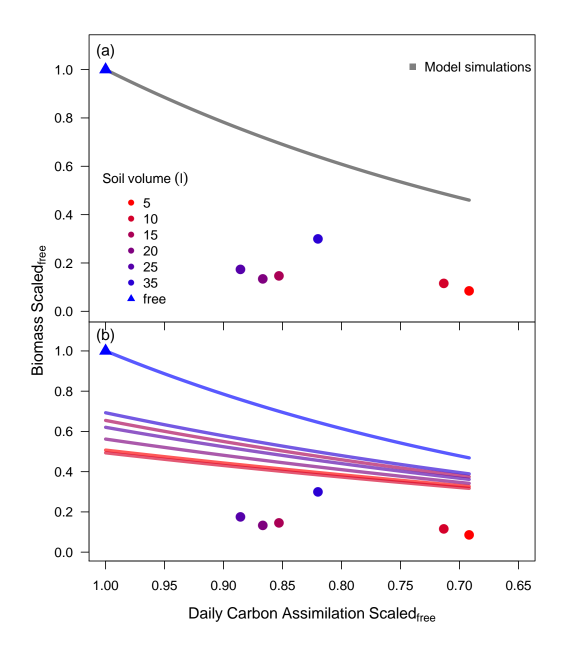


Figure 6: Modeled and harvested seedling biomass (g) versus the measure range of daily assimilated carbon gain (g m²) across all treatments (a) and then with treatment specific plant component mass partitioning included (b). Values of biomass and carbon gain are scaled to the free seedling with unlimited soil volume.

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Table 3: Seedling Growth Model Default Parameters

Variable	Default Value	Units	Source
Leaf area _i	0.035	m²	this study
Leaf mass _i	3.45	g	this study
Stem mass _i	1.51	g	this study
Root mass _i	0.99	g	this study
٤ _c	.65	g C g mass ⁻¹	Mäkelä (1997)
R _{coarse root}	0.00124	g C g root ⁻¹	Marsden et al. (2008)
$R_{fine\ root}$	0.010368	g C g root ⁻¹	Ryan et al. (2010)
R_{stem}	0.00187	g C g stem ⁻¹	Drake 2014 (unpublished)
C_{day}	4.7-6.8	g C m ⁻²	this study
Λ	1/365	yr ⁻¹	theoretical

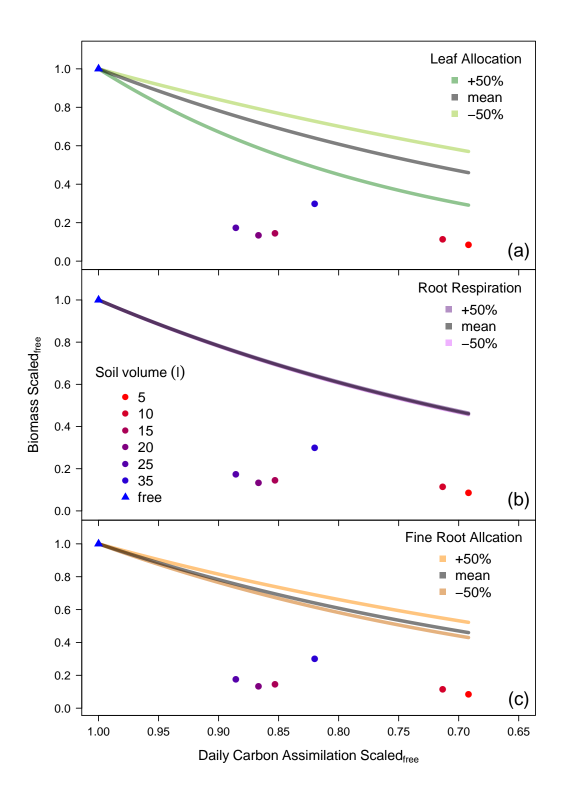


Figure S1: Modeled and harvested seedling biomass (g) versus daily assimilated carbon gain (g m^2) including sensitivity testing for unmeasured carbon allocation scenarios. Model parameters of leaf allocation (a), root respiration (b), and fine root allocation (c) were increased or decreased by 50%.

References

- Elizabeth A Ainsworth and Stephen P Long. What have we learned from 15 years of free-air CO2 enrichment (FACE)? A meta-analytic review of the responses of photosynthesis, canopy properties and plant production to rising CO2. *New Phytologist*, 165(2):351–372, 2005.
- Elizabeth A Ainsworth and Alistair Rogers. The response of photosynthesis and stomatal conductance to rising [CO2]: mechanisms and environmental interactions. *Plant, cell & environment*, 30 (3):258–270, 2007.
- W J Arp. Effects of source-sink relations on photosynthetic acclimation to elevated CO2. *Plant, Cell & Environment*, 14(8):869–875, 1991.
- I Biran and A Eliassaf. The effect of container size and aeration conditions on growth of roots and canopy of woody plants. *Scientia Horticulturae*, 12(4):385–394, 1980a.
- ³⁸⁰ I Biran and A Eliassaf. The effect of container shape on the development of roots and canopy of woody plants. *Scientia Horticulturae*, 12(2):183–193, 1980b.
- Kristine Y Crous, Audrey G Quentin, Yan-Shih Lin, Belinda E Medlyn, David G Williams, Craig V M
 Barton, and David S Ellsworth. Photosynthesis of temperate Eucalyptus globulus trees outside
 their native range has limited adjustment to elevated CO2 and climate warming. *Global change*biology, 19(12):3790–3807, 2013.
- Bert G Drake, Miquel A Gonzàlez-Meler, and Steve P Long. More efficient plants: a consequence of rising atmospheric CO2? *Annual review of plant biology*, 48(1):609–639, 1997.
- John E Drake, Michael J Aspinwall, Sebastian Pfautsch, Paul D Rymer, Peter B Reich, Renee A Smith, Kristine Y Crous, David T Tissue, Oula Ghannoum, and Mark G Tjoelker. The capacity to cope with climate warming declines from temperate to tropical latitudes in two widely distributed Eucalyptus species. *Global change biology*, 2014.
- Wei Duan, Pei G Fan, Li J Wang, Wei D Li, Shu T Yan, and Shao H Li. Photosynthetic response to low sink demand after fruit removal in relation to photoinhibition and photoprotection in peach trees. *Tree physiology*, 28(1):123–132, 2008.
- Remko Duursma. plantecophys: Modelling and analysis of leaf gas exchange data, 2014.
- Remko A Duursma, Craig V M Barton, Yan-Shih Lin, Belinda E Medlyn, Derek Eamus, David T Tissue, David S Ellsworth, and Ross E McMurtrie. The peaked response of transpiration rate to vapour pressure deficit in field conditions can be explained by the temperature optimum of photosynthesis. *Agricultural and Forest Meteorology*, 189:2–10, 2014.
- L F Ebell. Variation in total soluble sugars of conifer tissues with method of analysis. *Phytochem-istry*, 8(1):227–233, 1969.
- Alieta Eyles, Elizabeth A Pinkard, Noel W Davies, Ross Corkrey, Keith Churchill, Anthony P OGrady,
 Peter Sands, and Caroline Mohammed. Whole-plant versus leaf-level regulation of photosynthetic responses after partial defoliation in Eucalyptus globulus saplings. *Journal of experimental*botany, 64(6):1625–1636, 2013.
- G D Farquhar, S von von Caemmerer, and J A Berry. A biochemical model of photosynthetic CO2 assimilation in leaves of C3 species. *Planta*, 149(1):78–90, 1980.
- Carla A Gunderson and Stan D Wullschleger. Photosynthetic acclimation in trees to rising atmospheric CO2: a broader perspective. *Photosynthesis research*, 39(3):369–388, 1994.
- I Tanya Handa, Christian Körner, and Stephan Hättenschwiler. A test of the treeline carbon limitation hypothesis by in situ CO2 enrichment and defoliation. *Ecology*, 86(5):1288–1300, 2005.
- Afef Haouari, Marie-Christine Van Labeke, Kathy Steppe, Fethi Ben Mariem, Mohamed Braham, and Mohamed Chaieb. Fruit thinning affects photosynthetic activity, carbohydrate levels, and shoot and fruit development of olive trees grown under semiarid conditions. *Functional Plant Biology*, 40(11):1179–1186, 2013.

- Günter Hoch, Marianne Popp, and Christian Körner. Altitudinal increase of mobile carbon pools in Pinus cembra suggests sink limitation of growth at the Swiss treeline. *Oikos*, 98(3):361–374,
- Domingo J Iglesias, Ignacio Lliso, Francisco R Tadeo, and Manuel Talon. Regulation of photosynthesis through source: sink imbalance in citrus is mediated by carbohydrate content in leaves.

 Physiologia Plantarum, 116(4):563–572, 2002.
- J Kallarackal and C K Somen. An ecophysiological evaluation of the suitability of Eucalyptus grandis for planting in the tropics. *Forest Ecology and Management*, 95(1):53–61, 1997.
- Miko U F Kirschbaum. Does enhanced photosynthesis enhance growth? Lessons learned from CO2 enrichment studies. *Plant Physiology*, 155(1):117–124, 2011.
- ⁴²⁶ C Körner. Growth controls photosynthesis–mostly. *Nova Acta Leopoldina*, 114:273–283, 2013.
- ⁴²⁷ Christian Körner. Carbon limitation in trees. Journal of ecology, 91(1):4–17, 2003.
- T T Kozlowski. Carbohydrate sources and sinks in woody plants. *The Botanical Review*, 58(2):107–222, 1992.
- Desmond R Layne and J A Flore. End-product inhibition of photosynthesis in Prunus cerasus L. in response to whole-plant source-sink manipulation. *Journal of the American Society for Horticul-tural Science*, 120(4):583–599, 1995.
- W D Li, S H Li, S H Yang, J M Yang, X B Zheng, X D Li, and H M Yao. Photosynthesis in response to sink-source manipulations during different phenological stages of fruit development in peach trees: regulation by stomatal aperture and leaf temperature. *Journal of horticultural science & biotechnology*, 80(4):481–487, 2005.
- C H A Little and K Loach. Effect of changes in carbohydrate concentration on the rate of net photosynthesis in mature leaves of Abies balsamea. *Canadian Journal of Botany*, 51(4):751–758, 1973.
- Godfrey G Maina, Joel S Brown, and Mordechai Gersani. Intra-plant versus inter-plant root competition in beans: avoidance, resource matching or tragedy of the commons. *Plant Ecology*, 160 (2):235–247, 2002.
- Annikki Mäkelä. A carbon balance model of growth and self-pruning in trees based on structural relationships. *Forest Science*, 43(1):7–24, 1997.
- Claire Marsden, Yann Nouvellon, and Daniel Epron. Relating coarse root respiration to root diameter in clonal Eucalyptus stands in the Republic of the Congo. *Tree physiology*, 28(8):1245–1254,
 2008.
- B. V. McCleary, T. S. Gibson, and D. C. Mugford. Measurement of total starch in cereal products by amyloglucosidase- α -amylase method: Collaborative study. *Journal of AOAC International*, 80(3): 571–579, 1997. ISSN 1060-3271.
- K D M McConnaughay and F A Bazzaz. Is physical space a soil resource? *Ecology*, pages 94–103, 1991.
- B E Medlyn, Erwin Dreyer, D Ellsworth, M Forstreuter, P C Harley, M U F Kirschbaum, Xavier Le Roux, Pierre Montpied, J Strassemeyer, A Walcroft, and Others. Temperature response of parameters of a biochemically based model of photosynthesis. II. A review of experimental data. *Plant, Cell & Environment*, 25(9):1167–1179, 2002.
- Belinda E Medlyn, Remko A Duursma, Derek Eamus, David S Ellsworth, I Colin Prentice, Craig V M
 Barton, Kristine Y Crous, Paolo de Angelis, Michael Freeman, and Lisa Wingate. Reconciling the
 optimal and empirical approaches to modelling stomatal conductance. *Global Change Biology*,
 17(6):2134–2144, 2011.
- Belinda E Medlyn, Remko A Duursma, Derek Eamus, David S Ellsworth, I Colin Prentice, Craig V M Barton, Kristine Y Crous, Paolo Angelis, Michael Freeman, and Lisa Wingate. Reconciling the

- optimal and empirical approaches to modelling stomatal conductance. *Global Change Biology*, 18(11):3476, 2012.
- Patrick J Mitchell, Anthony P O'Grady, David T Tissue, Donald A White, Maria L Ottenschlaeger, and
 Elizabeth A Pinkard. Drought response strategies define the relative contributions of hydraulic
 dysfunction and carbohydrate depletion during tree mortality. New Phytologist, 197(3):862–872,
 2013.
- Shinichi Nakagawa and Holger Schielzeth. A general and simple method for obtaining R2 from generalized linear mixed-effects models. *Methods in Ecology and Evolution*, 4(2):133–142, 2013.
- Sergio G Nebauer, Begoña Renau-Morata, José Luis Guardiola, and Rosa-Victoria Molina. Photosynthesis down-regulation precedes carbohydrate accumulation under sink limitation in Citrus. *Tree Physiology*, page tpq103, 2011.
- D Scott NeSmith and John R Duval. The effect of container size. *HortTechnology*, 8(4):495–498, 1998.
- Richard J Norby, Evan H DeLucia, Birgit Gielen, Carlo Calfapietra, Christian P Giardina, John S King,
 Joanne Ledford, Heather R McCarthy, David J P Moore, Reinhart Ceulemans, and Others. Forest
 response to elevated CO2 is conserved across a broad range of productivity. *Proceedings of the*National Academy of Sciences of the United States of America, 102(50):18052–18056, 2005.
- Jari Ovaska, Ruuska Sari, Eevi Rintamäki, and Elina Vapaavuori. Combined effects of partial defoliation and nutrient availability on cloned Betula pendula saplings II. Changes in net photosynthesis and related biochemical properties. *Journal of Experimental Botany*, 44(8):1395–1402, 1993a.
- Jari Ovaska, Mari Walls, and Elina Vapaavuori. Combined Effects of Partial Defoliation and Nutrient
 Availability on Cloned Betula pendula Saplings I CHANGES IN GROWTH, PARTITIONING AND
 NITROGEN UPTAKE. Journal of Experimental Botany, 44(8):1385–1393, 1993b.
- Sara Palacio, Günter Hoch, Anna Sala, Christian Körner, and Pete Millard. Does carbon storage limit tree growth? *New Phytologist*, 201(4):1096–1100, 2014.
- Matthew J Paul and Christine H Foyer. Sink regulation of photosynthesis. *Journal of experimental* botany, 52(360):1383–1400, 2001.
- Jose Pinheiro, Douglas Bates, Saikat DebRoy, Deepayan Sarkar, and R Core Team. {nlme}: Lineαr and Nonlinear Mixed Effects Models, 2014. URL http://cran.r-project.org/package=nlme.
- E A Pinkard, C L Beadle, N J Davidson, and M Battaglia. Photosynthetic responses of Eucalyptus nitens (Deane and Maiden) Maiden to green pruning. *Trees*, 12(3):119–129, 1998.
- Hendrik Poorter, Ülo Niinemets, Lourens Poorter, Ian J Wright, and Rafael Villar. Causes and consequences of variation in leaf mass per area (LMA): a meta-analysis. *New Phytologist*, 182(3): 565–588, 2009.
- Hendrik Poorter, Jonas Bühler, Dagmar van Dusschoten, José Climent, and Johannes A Postma.
 Pot size matters: a meta-analysis of the effects of rooting volume on plant growth. Functional
 Plant Biology, 39(11):839–850, 2012a.
- Hendrik Poorter, Karl J Niklas, Peter B Reich, Jacek Oleksyn, Pieter Poot, and Liesje Mommer.
 Biomass allocation to leaves, stems and roots: meta-analyses of interspecific variation and environmental control. *New Phytologist*, 193(1):30–50, 2012b.
- Peter B Reich, Y Weisel, A Eshel, and U Kafkafi. Root-shoot relations: optimality in acclimation and adaptation or the Emperors New Clothes. *Plant roots: the hidden half*, pages 205–220, 2002.
- N Suzanne Robbins and David M Pharr. Effect of restricted root growth on carbohydrate metabolism and whole plant growth of Cucumis sativus L. *Plant physiology*, 87(2):409–413, 1988.

- Cláudio P Ronchi, Fábio M DaMatta, Karine D Batista, Gustavo A B K Moraes, Marcelo E Loureiro,
 and Carlos Ducatti. Growth and photosynthetic down-regulation in Coffea arabica in response
 to restricted root volume. Functional Plant Biology, 33(11):1013–1023, 2006.
- Michael G Ryan, Jose Luiz Stape, Dan Binkley, Sebastiao Fonseca, Rodolfo A Loos, Ernesto N Takahashi, Claudio R Silva, Sergio R Silva, Rodrigo E Hakamada, Jose Mario Ferreira, and Others. Factors controlling Eucalyptus productivity: How water availability and stand structure alter production and carbon allocation. *Forest ecology and management*, 259(9):1695–1703, 2010.
- Rowan F Sage. Acclimation of photosynthesis to increasing atmospheric CO2: the gas exchange perspective. *Photosynthesis research*, 39(3):351–368, 1994.
- Arne Sellin. Does pre-dawn water potential reflect conditions of equilibrium in plant and soil water status? *Acta Oecologica*, 20(1):51–59, 1999.
- Alison M Smith and Mark Stitt. Coordination of carbon supply and plant growth. *Plant, cell & environment*, 30(9):1126–1149, 2007.
- Tarryn L Turnbull, Mark A Adams, and Charles R Warren. Increased photosynthesis following partial defoliation of field-grown Eucalyptus globulus seedlings is not caused by increased leaf nitrogen. *Tree Physiology*, 27(10):1481–1492, 2007.
- L Urban and L Alphonsout. Girdling decreases photosynthetic electron fluxes and induces sustained photoprotection in mango leaves. *Tree Physiology*, 27(3):345–352, 2007.
- David I Warton, Remko A Duursma, Daniel S Falster, and Sara Taskinen. smatr 3–an R package for estimation and inference about allometric lines. *Methods in Ecology and Evolution*, 3(2):257–259, 2012.
- Sarah D Wright and Kelly D M Mcconnaughay. Interpreting phenotypic plasticity: the importance of ontogeny. *Plant Species Biology*, 17(2-3):119–131, 2002.
- Rui Zhou and Bruno Quebedeaux. Changes in photosynthesis and carbohydrate metabolism in mature apple leaves in response to whole plant source-sink manipulation. *Journal of the American Society for Horticultural Science*, 128(1):113–119, 2003.