RESEARCH PAPER

Whole-plant versus leaf-level regulation of photosynthetic responses after partial defoliation in *Eucalyptus globulus* saplings

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Abstract

Increases in photosynthetic capacity ($A_{1500}$) after defoliation have been attributed to changes in leaf-level biochemistry, water, and/or nutrient status. The hypothesis that transient photosynthetic responses to partial defoliation are regulated by whole-plant (e.g. source–sink relationships or changes in hydraulic conductance) rather than leaf-level mechanisms is tested here. Temporal variation in leaf-level gas exchange, chemistry, whole-plant soil-to-leaf hydraulic conductance ($K_p$), and aboveground biomass partitioning were determined to evaluate mechanisms responsible for increases in $A_{1500}$ of *Eucalyptus globulus* L. potted saplings. $A_{1500}$ increased in response to debudding (B), partial defoliation (D), and combined B&D treatments by up to 36% at 5 weeks after treatment. Changes in leaf-level factors partly explained increases in $A_{1500}$ of B and B&D treatments but not for D treatment. By week 5, saplings in B, B&D, and D treatments had similar leaf-specific $K_p$ to control trees by maintaining lower midday water potentials and higher transpiration rate per leaf area. Whole-plant source:sink ratios correlated strongly with $A_{1500}$. Further, unlike $K_p$, temporal changes in source:sink ratios tracked well with those observed for $A_{1500}$. The results indicate that increases in $A_{1500}$ after partial defoliation treatments were largely driven by an increased demand for assimilate by developing sinks rather than improvements in whole-plant water relations and changes in leaf-level factors. Three carbohydrates, galactosyl, stachyose, and, to a lesser extent, raffinose, correlated strongly with photosynthetic capacity, indicating that these sugars may function as signalling molecules in the regulation of longer term defoliation-induced gas exchange responses.

Key words: Carbohydrates, carbon limitation, defoliation, leaf water potential, photosynthesis, plant hydraulic conductance.

Introduction

After partial defoliation, a continuum of photosynthetic responses has been reported in the remaining foliage, with the direction, magnitude, and duration influenced by the type of damage, plant species, and environment (Sweet and...
There are two hypotheses that might explain photosynthetic responses to defoliation at a whole-plant level. The first, the source:sink (S:S) hypothesis, suggests that increases in photosynthetic rates after partial defoliation are regulated by changes in whole-plant carbon S:S relationships (Neales and Incoll, 1968; Layne and Flore, 1995; Pinkard et al., 2011b). The second implies that photosynthetic responses to defoliation are regulated by changes in whole-plant hydraulic conductance (Whitehead, 1998; Brodribb et al., 2007). Plant organs can be defined by their functional role as either net exporters (sources) or net consumers of carbohydrates (sinks) (Laporte and Delph, 1996; Neales and Incoll, 1968). The S:S hypothesis suggests that defoliation decreases source size while leaving sink demand relatively unchanged. Increased demand for carbohydrates from remaining leaves reduces end-product inhibition, which in turn increases the photosynthetic rates of the remaining leaves (Sweet and Wareing, 1966; Neales and Incoll, 1968; Layne and Flore, 1995). If whole-plant S:S relationships were driving photosynthetic responses to defoliation, increases in the rates of biochemical reactions of photosynthesis and changes in leaf-level carbohydrate concentrations might be expected. While increases in stomatal conductance ($g_s$) might also occur, this would not be a key driver of photosynthetic responses. Evidence to support the S:S hypothesis at the whole-plant scale is not well documented. While increases in the rates of biochemical reactions of photosynthesis are commonly reported following defoliation (von Caemmerer and Farquhar, 1984; Layne and Flore, 1995; Ozaki et al., 2004), these have not been linked explicitly to whole-plant S:S processes. Reductions in total non-structural carbohydrate (NSC) concentrations in the remaining leaves following defoliation have been observed (e.g. Layne and Flore, 1995; Myers et al., 1999; Zhou and Quebedeaux, 2003) and are consistent with an increased demand and export of NSC from source leaves to developing sinks. However, this response is not universal (see Lavigne et al., 2001; Turnbull et al., 2007; Eyles et al., 2009b). End-product inhibition may be related to a specific carbohydrate rather than total NSC concentration (Turnbull et al., 2007). In this study, the aim is to quantify not only total soluble sugar and starch concentrations but also up to eight specific carbohydrates in order to capture fully the effect of defoliation on carbohydrate metabolism.

Maximum rates of gas exchange and growth of plants are also regulated by plant hydraulic architecture (Whitehead, 1998; Brodribb et al., 2007). Leaves represent one of the largest sites of resistances to water flow in plants (Sack and Holbrook, 2006); thus, reductions in leaf area may result in a greater root-to-leaf hydraulic conductivity. Reductions in leaf area also represent reduced transpiring surface area and increased root:leaf ratio, which in turn improves the water and nutrient status of the remaining foliage per unit area (Welker and Menke, 1990). If hydraulic architecture was the key driver of photosynthetic responses to defoliation, a strong relationship between photosynthesis and $g_s$, irrespective of defoliation treatment, would be expected, along with increases in root-to-leaf hydraulic conductivity. The few studies that have examined the impact of defoliation on whole-plant water relations reported less negative midday leaf water potential ($\Psi_{mid}$) (Syvertsen, 1994; Singh and Thompson, 1995; Vanderklein and Reich, 2000; Quentin et al., 2011) and higher transpiration ($E_L$) per unit leaf area in response to defoliation (Meinzer and Grantz, 1990; Quentin et al., 2011; but see Oren et al., 1999). Meinzer and Grantz (1990) observed increased whole-plant soil-to-leaf hydraulic conductance ($K_p$) within 24h after defoliation in sugar cane and they attributed this response to increased $E_L$ and not to any changes in $\Psi$. A recent study noted higher $K_p$ 2 months after defoliation, and this was due to an increase in $E_L$ and a less negative $\Psi_{mid}$ in 4-year-old *Eucalyptus globulus* Labill. trees (Quentin et al., 2011).

The broad-leaved evergreen woody tree species, *E. globulus* was selected to examine the underlying mechanisms of photosynthetic up-regulation observed post-defoliation. This well-studied species exhibits an indeterminate growth habit and has a documented propensity for photosynthetic up-regulation (Pinkard et al., 2007; Eyles et al., 2009b). Temporal responses in leaf-level gas exchange, biochemical, N, and carbohydrates as well as whole-plant changes in $K_p$ and S:S ratios of well-watered and fertilized saplings were examined to test specific whole-plant- and leaf-level mechanisms that may explain the increased photosynthetic performance after partial defoliation and/or budding. In seeking to determine the contribution of whole-plant mechanisms, the focus is not only on the S:S ratio and hydraulic conductance, but leaf-level responses are also investigated to examine fully the effects of defoliation on carbon use and storage, and water relations during the refoliation process. It was hypothesized that photosynthetic responses to partial defoliation were determined by whole-plant processes related to changes in either the S:S ratio or hydraulic conductance. It was predicted that (i) if the demand for assimilates by sinks regulates photosynthesis of source leaves, then there will be an inverse relationship between photosynthesis and the S:S ratio, followed by a decline in photosynthesis as the S:S ratio is restored during crown recovery, with concomitant leaf-level increases in rates of biochemical reactions of photosynthesis and changes in NSCs; and (ii) if whole-plant water relations are responsible for photosynthetic up-regulation, then there will be a positive relationship between $A$ and $K_p$. 

Wareing, 1966; von Caemmerer and Farquhar, 1984; Lovett and Tobiesen, 1993; Reich et al., 1993; Pinkard et al., 2004; Turnbull et al., 2007; Eyles et al., 2009b, 2011). Explanations for photosynthetic responses have been largely focused on leaf-level mechanisms such as improvements in leaf N availability (Lavigne et al., 2001) and increases in the rates of biochemical reactions of photosynthesis (von Caemmerer and Farquhar, 1984; Layne and Flore, 1995; Ozaki et al., 2004; Turnbull et al., 2007). An alternative hypothesis is that whole-plant mechanisms explain the gas exchange responses. The scale at which photosynthetic responses are regulated affects the sorts of management strategies that may be effective in promoting recovery from defoliation, and influences the ways in which defoliation responses might be represented in productivity models (Pinkard et al., 2011a).
Materials and methods

Plant material

Fifty-three open-pollinated saplings of *E. globulus* were planted in 300 mm diameter pots (volume: 21 litres). In order to mimic natural growing conditions, saplings were supplied a low-phosphorus potting mix. They were grown outside for 5 months, watered until saturated daily, and fertilized with a slow-release pelletized fertilizer (Osmocote Native Gardens, N:P:K of 17.9:0.8:7.3, Scotts, Australia). The saplings were 68 ± 2 cm (mean ± SE) in height and had a diameter of 0.7 ± 0.3 cm at the start of the experiment (December 2010; aged 6 months). The saplings were exposed to ambient weather conditions for the full duration of the experiment.

Design

The experiment was established as a completely randomized design. Four foliage treatments were randomly applied to 12 saplings per treatment on 14 December, 2010. (i) Control (no defoliation) (C). (ii) Defoliated (remove leaves from 50% of crown length in the upper crown on December 2010 when saplings were 6 months old). The defoliation treatment removed leaves from the crown apex downwards (i.e. upper crown), excluding apical buds. Leaves were removed using long-nosed secateurs (D). (iii) A 100% removal of buds (remove all buds throughout the entire crown) (B). (iv) A combination of B and D (B&D).

These treatments were designed to manipulate S:S ratios (i.e. defoliation examined the effect of source leaf removal while budding examined the effect of sink removal). Previous studies have shown that 50% defoliation of the upper but not the lower crown elicits photosynthetic up-regulation (*Pinkard et al.*). All foliage material was collected and total foliage area removed was determined as follows. A subsample of 10 leaves per sapling was taken and fresh leaf area was measured using the scanning software WinRhizo (Regent Instruments, Quebec). Leaves were then dried at 65 °C for 3 d and weighed. The remaining leaf material per tree was also dried at 65 °C for 7 d. The area:dry mass ratio (specific leaf area, SLA) was used to calculate leaf area removed per tree.

Gas exchange measurements

Photosynthetic measurements were made immediately before treatment (week 0) and 2 (week 2), 5 (week 5), 6 (week 6), 8 (week 8), and 12 (week 12) weeks after imposition of treatments on four saplings per treatment per measurement week. The *A*$_{1500}$ and $g_c$ were measured with a portable open-path gas exchange system with CO$_2$ control (Li-Cor LI-6400 portable IRGA, Li-Cor, Lincoln, NE, USA). Measurements were undertaken with a standard 20 × 30 mm leaf chamber equipped with blue-red light-emitting diodes mounted on the top of the chamber (Model 6400-02B). Leaves of potted *E. globulus* saplings have been shown to be light saturated at a photosynthetic photon flux density (PPFD) of 1000 μmol m$^{-2}$ s$^{-1}$ (Eyles et al., 2009b); therefore, PPFD was set at 1500 μmol m$^{-2}$ s$^{-1}$. The ambient [CO$_2$] was maintained at 39 Pa. Three of the youngest fully expanded leaves located within the lower crown of each sapling were recorded after *A*$_{1500}$ and $g_c$ had stabilized (between 0.5 min and 2 min). In weeks 2, 5, and 6, it was not possible to follow this sampling protocol for the bud treatments (B&D and B treatments). In these two treatments, the leaf pair closest to the developing bud was selected for gas exchange measurements. Up to week 5, all leaves selected for measurement had been fully expanded before imposition of the defoliation/debudding treatments. To minimize the effects of time of measurement, a randomly chosen replicate from each treatment was measured.

The responses of *A* to varying [CO$_2$] were measured at week 5, 8, and 12. For the photosynthetic rate-intercellular CO$_2$ concentration (*A*/C) curves, PPFD was maintained at 2000 μmol m$^{-2}$ s$^{-1}$ and leaf temperature at 20 °C. Leaves were first equilibrated at a [CO$_2$] of 39 Pa, after which [CO$_2$] was reduced to 0 and then increased to 200 Pa in a total of 12 steps. The *A*/C curves were measured between 0900h and 1400h Eastern Standard Time (EST) over two consecutive days. Across all gas exchange measurements, leaf temperatures varied between 18 °C and 26 °C, vapour pressure deficit approximated ambient conditions, varying between 0.9 kPa and 1.3 kPa, while airflow through the chamber was 400 μmol s$^{-1}$. Each leaf used for photosynthetic measurements was sampled mid afternoon because maximum accumulation of sugars and starch occurs mid afternoon (1530–1630h EST) (*Zhou et al.*, 2001). Leaves were immediately frozen and stored at −20 °C pending chemical analyses.

Biomass harvests

At week 0, a biomass harvest of five saplings was conducted to develop an allometric relationship between basal diameter and leaf area (leaf area=0.0028×basal diameter$^{10.9}$, r$^2$=0.89), which were used to estimate percentage leaf area removed for each treatment in week 1. Further biomass harvests were conducted at week 5, 8, and 12. Four saplings per treatment (a total of 16 saplings per harvest date) were harvested for measurements of aboveground biomass. At each harvest, saplings were enclosed immediately in plastic bags and stored at −20 °C until processed. Each tree was divided into its three main biomass components: leaves, buds, and woody tissue, and oven-dried to constant mass at 65 °C. Prior to drying, the leaf areas of 10 leaves, representing a range of sizes, were measured using the scanning software WinRhizo for determination of SLA. Biomass was coarsely ground with a Thomas-Wiley mill model 4 (Thomas Scientific, Swedesboro, NJ, USA) and then subsampled for further grinding to a fine powder with a mixer mill MM200 grinder (Retsch, Haan, Germany) for N and carbohydrate analyses.

Plant hydraulic conductance

*K*$_W$ was measured 1, 5, 8, and 12 weeks after defoliation using the following equation:

$$K_W = \frac{E_L}{\Psi_{md} - \Psi_{pd}}$$

where *E*$_L$ is transpiration (mmol m$^{-2}$ s$^{-1}$) and $\Psi_{pd}$ and $\Psi_{md}$ (MPa) are pre-dawn and mid-day leaf water potentials, respectively. On the day prior to *K*$_W$ measurements, each pot was watered thoroughly and enclosed in a polypropylene bag sealed tightly around the base of the stem to prevent evaporation of soil water. *E*$_L$ was measured gravimetrically; that is, change in weight of the pot as measured over a 40–60 min period between 1130 h and 1230 h EST during maximal rates of transpiration on sunny days. *E*$_L$ was normalized by dividing all values by the total leaf area of the sapling (as determined from the biomass harvests, described above). $\Psi_{pd}$ and $\Psi_{md}$ were measured by collecting one leaf at 1300 h and 2400 h, respectively. They were placed immediately into plastic bags and kept in the dark until measurements were made using a 4.0 MPa pressure chamber (PMS Instruments Co., Corvallis, OR, USA).

Chemical analyses

Soluble carbohydrates were extracted from ~50mg of dried plant tissue in 10ml of 80% (v/v) ethanol in a 60 °C water bath for 30 min. The extraction solvent included an internal standard of 0.01% trehalose, a sugar which had not been previously detected in initial qualitative studies. After centrifugation (10 min, 2500 rpm), the soluble sugars were separated and quantified using ultra-performance liquid chromatography–mass spectrometry (UPLC-MS). Full details of the method used are detailed in the Supplementary Method S1 available at *JXB* online. Starch and complex sugars remaining in the undissolved pellet of plant material after ethanol extractions were...
enzymatically (amyloglucosidase; Fluka-10115, Sigma-Aldrich, St.
Louis, MO, USA) reduced to glucose using the method detailed in
Eyles et al. (2009a). It would have been preferable to have used freeze-
rather than oven-dried samples but, given that all samples were pro-
cessed in the same manner, and relative rather than absolute values
were used to compare treatment effects, it is considered that any
treatment effects, if present, would have continued to be evident
nonetheless. Although the surrace concentration in the samples was
lower than those found in other E. globulus studies (e.g. Merchant
et al., 2011), it was still possible to observe significant differences
across treatments (Table 1).
Leaf N concentration was determined on dried and ground mate-
rial with an elemental analyser (Thermo Finnigan EA 1112 Series
Flash Elemental Analyser, Thermo Scientific, MA, USA).

Data analyses
Photosynthetic responses to [CO₂] were fitted to the biochemical
model developed by Farquhar et al. (1980) and von Caemmerer and
Farquhar (1981), and presented in full by von Caemmerer (2000).
The values of the various parameters in the model are the standard
values tabulated by von Caemmerer (2000), but V_{\text{Cmax}} and J_{\text{max}} were
estimated from A/C curves. The fitting protocol used here was based
on an Excel workbook on the PhysEcol website (http://www.else-
vierdirect.com/companion.jsp?ISBN=9780123744609) that accom-
panies Landsberg and Sands (2011). The standard errors of, and cor-
relations between, the estimated parameters were obtained using the
NonlinXL software also available from the PhysEcol website.

The whole-plant aboveground S:S ratio (g:g) was calculated in
two ways. First, it was calculated as the ratio of biomass of pho-
tosynthetic tissues (leaves) to non-photosynthetic tissue (woody
tissue+buds) [hereafter referred to as source:sink (biomass)]. This
definition attempts to take into account stored NSC pools that may
occur in week 5; therefore, the effects of treatment on the fol-
lowing variables were explored: V_{\text{Cmax}}, J_{\text{max}}, leaf N, soluble sugars,
starch, and specific soluble sugars (fructose, galactinol, gentiobiose,
glucose, raffinose, stachyose, and sucrose) in week 5 only using one-
way ANOVA. The assumptions of ANOVA such as homogeneity of
variance and the Gaussian distribution were checked by the use of
qq plots and residual plots for all variables. Post-hoc comparisons
of means were made using Tukey’s method. Initially, the relation-
ships between A_{1500} and leaf- and whole-plant-level factors in week 5
were examined by linear regression. Associations between A_{1500}
and the explanatory variables were then explored using stepwise regres-
sion methods to examine whether increases in A_{1500} could be best
explained by (i) leaf-level factors only; (ii) whole-plant factors only;
or (iii) a combination of leaf- and whole-plant-level factors. A_{1500}
and SLA were analysed by SAS version 9.2. The other factors were
analysed by Genstat Version 10.1 (VSN International).

Results

Effect of defoliation treatments on leaf area
Leaf area at the start of the experiment in December aver-
age 0.586 ± 0.008 m². From the allometric relationships
developed between basal diameter and leaf area, it was esti-
mation that the percentage of leaf area removed in increas-
ing order of severity was as follows: C, 0%; B, 3.96 ± 0.2%;
D, 32.8 ± 1.3%; and B&D, 35.2 ± 1.3%. By week 5, leaf area
of the B&D treatment continued to be significantly reduced
by 50% compared with the C treatment (C, 1.32 ± 0.1; B,
0.97 ± 0.1; B&D, 0.66 ± 0.09, and D, 1.05 ± 0.1 m²; P < 0.05).

Table 1. Effect of treatments (B=bud damage; B&D=bud damage and defoliation;
C=control; D=50% defoliation) on leaf-level factors of Eucalyptus globulus in week 5 only.

| Parameter                      | B            | B&D          | C            | D            | P-value |
|-------------------------------|--------------|--------------|--------------|--------------|---------|
| V_{\text{Cmax}} [μmol m⁻² s⁻¹] | 90.7 (0.4)a  | 113.2 (5.9)b | 82.1 (1.4)a  | 94.9 (5.7)a  | ***     |
| J_{\text{max}} [μmol m⁻² s⁻¹]  | 163.8 (5.4)  | 184.8 (5.4)  | 137.2 (9.6)  | 137.5 (9.3)  | **      |
| Leaf Nₙₙ [g m⁻²]               | 1.24 (0.08)a | 1.70 (0.2)b  | 0.939 (0.07) | 1.163 (0.1)a | ***     |
| Leaf Nₙₙ [%]                   | 0.98 (0.07)  | 1.34 (0.1)   | 0.99 (0.08)  | 1.24 (0.09)  | P = 0.06 |
| SLA (cm² g⁻¹)                 | 78.3 (5.8)   | 78.6 (5.8)   | 106.6 (5.8)b | 106.9 (5.8)b | ***     |
| Soluble sugars (%)            | 5.9 (0.7)    | 6.0 (0.4)    | 5.4 (0.4)    | 6.5 (0.3)    | NS      |
| Starch (%)                    | 10.7 (1.6)   | 11.4 (1.1)   | 9.2 (0.7)    | 10.8 (1.2)   | NS      |
| Specific soluble sugars (mg g⁻¹) |            |              |              |              |         |
| Fructose                      | 22.7 (2.8)   | 19.7 (2.2)   | 21.9 (2.3)   | 24.0 (1.8)   | NS      |
| Galactinol                    | 2.89 (0.2)a  | 4.91 (0.2)b  | 2.31 (0.2)a  | 2.44 (0.4)a  | ***     |
| Gentiobiose                   | 4.54 (0.9)c  | 1.39 (0.2)a,b| 3.77 (0.7)b,c| 1.18 (0.3)a  | **      |
| Glucose                       | 19.5 (2.7)   | 18.3 (1.9)   | 17.8 (1.8)   | 19.8 (1.2)   | NS      |
| Raffinose                     | 1.91 (0.6)a  | 5.83 (0.6)b  | 1.43 (0.4)a  | 5.8 (0.2)b   | ***     |
| Stachyose                     | 0.31 (0.04)  | 0.62 (0.07)b | 0.24 (0.01)a | 0.38 (0.02)a | ***     |
| Sucrose                       | 0.79 (0.3)a  | 2.91 (0.5)b  | 0.57 (0.2)a  | 6.33 (0.6)c  | ***     |

Asterisks indicate significance of treatment (Trt), *P ≤ 0.05; **P ≤ 0.01; ***P ≤ 0.001; NS, non-significant. Different letters denote a significant treatment (α=0.05) effect within a measurement period. Error bars are 1 SE, n=4 for all treatments.
Defoliation treatments induce photosynthetic up-regulation

The \( A_{1500} \) and \( g_s \), measured at week 0 were not significantly different across treatments (i.e. mean values of \( A_{1500} = 13.6 \pm 0.2 \ \mu \text{mol m}^{-2} \ \text{s}^{-1} \) and \( g_s = 0.343 \pm 0.01 \ \text{mmol m}^{-2} \ \text{s}^{-1} \); \( P > 0.05 \)). Defoliation treatments significantly increased \( A_{1500} \) (Fig. 1A). By week 5, \( A_{1500} \) was 36, 18, and 17% higher in B&D, B, and D treatments, respectively, than in control saplings. These treatment differences were maintained through to week 6 but, by week 8, treatment effects were negligible. The changes in \( g_s \) followed a similar pattern to \( A_{1500} \) with the largest difference occurring in week 6 (Fig. 1B). At this sampling point, the \( g_s \) of both bud treatments was nearly double that measured in the C treatment. In week 8, it was only the \( g_s \) of \( B \) and not the B&D treatment that remained significantly higher than that of the control value. In week 5, the relationship between \( A_{1500} \) and \( g_s \) was significantly affected by treatment interactions (\( P < 0.05 \), Fig. 1C). In particular, significant regressions were found for the \( B \) and B&D treatments, and their respective slopes were different from those of the control saplings. This meant a higher \( A_{1500} \) per unit \( g_s \) was exhibited by saplings in the B&D and B treatments than in the C treatment.

Impact of defoliation treatment on whole-tree hydraulic conductivity per leaf area varies during defoliation

\( \Psi_{sd} \) was unaffected by defoliation treatment throughout the experiment (\( P > 0.05 \); data not shown). \( K_p \) was significantly affected by defoliation treatments (Fig. 2A). In week 1, the \( K_p \) in the B&D treatment was 78% higher than control values. The differences between the treatments and control saplings, however, decreased with time and, by week 5, the differences were <21%. In week 5, the reduction in total transpiring area brought about by the defoliation treatments was offset by a concomitant non-significant increase in \( E_l \) (Fig. 2B) and a decrease in \( \Psi_{md} \) (Fig. 2C). That is, \( E_l \) was 60% higher while \( \Psi_{md} \) was 31% lower for saplings in the B&D treatment compared with control values.

Whole-tree S:S ratios reflect defoliation treatments over time

The source: sink (biomass) of control trees was 1.8 (Fig. 2D). This declined with time to 1.05 by week 12, reflecting increased allocation of resources to woody tissue [i.e. woody tissue/total aboveground biomass \( \% = 32.6 \pm 0.02 \) (week 0), 42.9 ± 0.9 (week 5), 46.1 ± 0.9 (week 8), and 48 ± 1.6% (week 12), \( P < 0.05 \); Fig. 2D]. At week 1, defoliation treatments brought about a large reduction in the source:sink (biomass) of B&D and D treatments by 32% and 39%, respectively, compared with the C treatment. In contrast, there was an immediate reduction in sinks and therefore an increase in source: sink (biomass) in the B treatment. However, by week 5, the release of auxiliary buds, which replaced the single debudded apical bud with a new pair of buds (one arising from each leaf axil), created new sinks and this was reflected as a reduced source: sink (biomass) that was similar to that of the control saplings. The source: sink (biomass) of saplings in the B&D treatment remained significantly lower than the control value in week 5. Thereafter, the differences between treatments and control saplings decreased over time.

Leaf-level responses in week 5 vary with defoliation treatment

The largest contrast between treatments in \( A_{1500} \) occurred in week 5 (Fig. 1). Therefore, for clarity, the effects of defoliation treatments on leaf-level parameters in week 5 only are presented (see summary in Table 1). \( V_{Cmax} \) and \( J_{max} \) were significantly higher (~40%) than control values in the B&D treatment. SLA in both bud treatments was significantly lower (26%) than in the other two treatments. The leaf \( N_{mass} \) of saplings in the B&D treatment was almost twice as high as that of control saplings in week 5. Total soluble sugar concentrations showed a similar pattern to starch concentrations in week 5. The concentrations of five carbohydrates were significantly influenced by defoliation treatments in week 5 (Table 1). In contrast, no significant changes were observed for the inositol group, the unknown monosaccharide, and the two most dominant soluble sugars, glucose and fructose (data presented for fructose and glucose only, Table 1). Specifically, both defoliation treatments (D and B&D) resulted in significant increases in sucrose and raffinose but decreases in gentiobiose. For saplings in the B&D treatment, concentrations of stachyose and galactinol were 3- and 2-fold higher, respectively, than those in the C treatment.

Photosynthetic up-regulation primarily related to whole-plant responses

The relationships between \( A_{1500} \) and leaf- and whole-plant-level responses to defoliation treatments were explored in week 5 only. For leaf-level factors, \( A_{1500} \) was positively correlated with leaf \( N_{mass} \), \( V_{Cmax} \), and \( J_{max} \), and negatively correlated with SLA (Supplementary Fig. S1 at JXB online). \( A_{1500} \) was unrelated to starch and soluble sugar concentrations (Fig. 3A, B). These leaf-level relationships did not appear to be consistent across treatments (data not shown), suggesting that the changes in these values were not necessarily driving the photosynthetic up-regulation. Regression analyses between \( A_{1500} \) and specific soluble sugars showed that the direction and strength of these relationships varied with each soluble sugar (Fig. 3). Specifically, \( A_{1500} \) was positively correlated with stachyose, galactinol, and raffinose, but not sucrose and gentiobiose. Multiple regression analysis of all seven leaf-level factors that were found to be significantly related to \( A_{1500} \) (i.e. \( g_s \), \( V_{Cmax} \), \( J_{max} \), leaf \( N_{mass} \), stachyose, galactinol, and raffinose, Figs 1C, 3; Supplementary Fig. S1) showed that increases in \( A_{1500} \) were best explained by the predictor variables of galactinol and \( g_s \) (\( A_{1500} = 7.06 + 1.58 \text{Galactinol} + 11 \text{ } \times 0.76 \), \( r^2 = 0.76, P < 0.001 \)). However, there was considerable between-treatment variation in the relationship between \( A_{1500} \) and \( g_s \) (Fig. 1C), suggesting that galactinol had a large influence on the strength of this relationship.
Fig. 1. Effect of treatments (B=bud damage; B&D=bud damage and defoliation; C=control; D=50% defoliation) on $A_{1500}$ (A) and (B) $g_s$ on leaves of *Eucalyptus globulus*. Asterisks indicate significance of treatment (Trt), time, or their interaction (Int): **$P \leq 0.01$ and ***$P \leq 0.001$. Different letters denote a significant treatment ($\alpha=0.05$) effect within a measurement period. Error bars are 1 SE. (C) Relationship between $A_{1500}$ and $g_s$ of *E. globulus* in weeks 5 and 6. The significant regressions shown are described by the following equations: $A_{1500}(B)=18.63 \ g_s+8.70$ and $A_{1500}(B&D)=27.05 \ g_s+8.48$. 

$B&D: \ r^2 = 0.64, P < 0.05$ and $B: \ r^2 = 0.86, P < 0.01$.
At a whole-plant level, there was no significant relationship between $A_{1500}$ and $K_p$ ($P > 0.05$) (Fig. 4A). In contrast, there were significant negative relationships between $A_{1500}$ and source:sink (biomass) (Fig. 4B), source:sink (carbohydrate) (Fig. 4C), and leaf area (Fig. 4D). Multiple regression analysis at the whole-plant scale showed that increases in $A_{1500}$ were best explained by the predictor variable of source:sink (biomass) ($r^2=0.74$, $P < 0.001$, Fig. 4B) alone. The addition of $K_p$ or leaf area did not improve the correlation. A final multiple regression analysis of all leaf- and whole-plant-level variables showed that the best whole-plant model was not significantly improved by the addition of any of the leaf-level variables (data not shown).

**Discussion**

It has been demonstrated that whole-plant S:S relationships explain photosynthetic responses to defoliation and debudding. Part of the whole-plant response revolved around leaf-level changes that facilitate increases in photosynthetic rates, although the role of some leaf-level responses alone in determining photosynthetic responses (e.g. the relationship between $A_{1500}$ and galactinol+$g_s$) cannot be discounted conclusively. There was only weak evidence for hydraulically mediated photosynthetic responses. Following an initial discussion of the variable evidence that links increases in $A_{1500}$ with leaf-level responses, it is then proposed that the observed whole-plant responses improved understanding of photosynthetic responses to defoliation. In particular, it is argued that the temporal patterns of $A_{1500}$ are closely related to changes in S:S ratios during crown recovery.

**Leaf-level factors and photosynthetic up-regulation**

Leaf-level changes similar to those reported in other studies (Lavigne et al., 2001; Layne and Flore, 1995; Ozaki et al., 2004; Pinkard et al., 2011), and consistent with the whole-plant regulation of photosynthesis via S:S interactions, were observed. All treatments, and, in particular, the B&D and B treatments, increased $A_{1500}$ per unit $g_s$ (Fig. 1C), suggesting that these saplings became more efficient at fixing CO$_2$ per unit water lost, and that changes in $g_s$ were not driving photosynthetic responses to defoliation as would have been expected if photosynthetic changes were hydraulically mediated. Further, increases in $A_{1500}$ were accompanied by increases in $V_{C\text{max}}$ and $J_{\text{max}}$ (Table 1), suggestive of changes in the rates of biochemical reactions of photosynthesis and more rapid translocation of end-products. These responses, however, were only observed in the debudding and not in the D treatment. Decreases in SLA were also noted, but, once again, only in the debudding treatments (Table 1), providing some evidence that these treatments had affected the leaf

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Different letters denote a significant treatment ($\alpha=0.05$) effect within a measurement period. Error bars are 1 SE, $n=4$ for all treatments.
thickness and density of apparently ‘fully expanded’ leaves, though it remains unclear exactly how. In response to pathogen infection, the adaxial palisade layer of fully expanded leaves of *E. globulus* is capable of undergoing cell division to produce, for example, a necrophylactic periderm (Smith et al., 2007). Debudding treatments may similarly induce modifications in leaf anatomy that might affect leaf-level hydraulics (Brodribb et al., 2007).

In previous studies examining photosynthetic responses in *E. globulus* after defoliation, leaf $N_{\text{area}}$ has been reported either to increase (Turnbull et al., 2007; Pinkard et al., 2011b) or to not differ from the control value (Eyles et al., 2009b; Quentin et al., 2010). Turnbull et al. (2007) concluded that the immediate increases in $A_{1500}$ following defoliation were largely attributed to increases in $V_{\text{Cmax}}$ and $J_{\text{max}}$. In this study, a positive relationship between $A_{1500}$ and leaf $N_{\text{area}}$ was observed, but regression analysis suggested that $N_{\text{area}}$ did not explain the photosynthetic response to defoliation.

As reported in other studies, it was found that responses between $A_{1500}$ and various leaf-level variables were...
inconsistent across treatments, suggesting that the direction and magnitude of the leaf-level responses were dependent on the nature of the treatment imposed, and that the observed photosynthetic responses were not sufficiently explained by leaf-level mechanisms. The strong relationship between $A_{1500}$ and the combination of $g_a$ and galactinol does not reflect the observed strong among-treatment differences in the relationship between $A_{1500}$ and $g_a$, and may highlight the role of galactinol as a signalling molecule (see below).

Whole-plant water relations and photosynthetic up-regulation

Various measures of whole-plant responses to defoliation have arguably provided some support for hydraulically mediated control of gas exchange. In the current study, $E_L$ was significantly higher in both defoliation treatments (B&D and D) than control values in week 5 ($P < 0.01$). This result is consistent with previous studies (Meinzer and Grantz, 1990; Pataki et al., 1998; Oren et al., 1999; Quentin et al., 2011, 2012) and provides some evidence of improved water status. However, it was also found that a significantly more negative $\Psi_{md}$ enabled trees in the B&D, D, and B treatments to maintain similar $K_p$ values to that observed in the C treatment in week 5. While there was an immediate increase in $K_p$ 1 week after defoliation (especially in the B&D treatment) (Fig. 2), it is suggested that this initial increase in $K_p$ was not directly responsible for the increase in $A_{1500}$ observed in week 5 because the increase was not sustained whereas the increases in $A_{1500}$ were. The lack of correlation between $K_p$ and $A_{1500}$ across treatments in week 5 (Fig. 4A) provides good evidence that increases in $A_{1500}$ were not sufficiently explained by changes in $K_p$ alone. In additional support for this argument, it was found that $K_p$ was unrelated to $g_a$ in week 5 (data not shown), suggesting a transient disconnection between leaf-level water status and whole-plant hydraulic capacity.

The poor correlation between $K_p$ and $A_{1500}$ could be explained by an unchanging root:shoot ratio. While partial defoliation is assumed to increase root:shoot ratios (Reich et al., 1993; Lavigne et al., 2001; Ozaki et al., 2004), in the current study it is unclear how debudding would lead to substantial shifts in the root:shoot ratio where even the 100% removal of buds failed to cause any changes in leaf area compared with the C treatment. Studies that examined the effects of defoliation on biomass partitioning have consistently reported either a decrease (due to root death) or an unchanging root:shoot ratio (Vanderklein and Reich, 199; Ovaska et al., 1993; Reich et al., 1993; Syvertsen, 1994; Singh and Thompson, 1995). Although the root:shoot ratio was not explicitly examined in this study, a comparable defoliation study in young E. globulus field trees similarly observed a reduction, rather than an increase, in this ratio (Eyles et al., 2009a). This response most probably reflects the strategy of defoliated plants to increase aboveground allocation of biomass to new photosynthetic tissue, not necessarily at the expense of, but instead of, root biomass (Eyles et al., 2009a).

Source–sink interactions and photosynthetic up-regulation

In accordance with the S:S hypothesis, $A_{1500}$ was greater in the B&D, D, and B treatments than C treatment in weeks 5 and 6 (Fig. 1), which was followed by a decline in $A_{1500}$ as the source:sink (biomass) was restored by week 8 (Figs 1, 2D). As predicted, an inverse relationship between $A_{1500}$ and source:sink (biomass) was observed across treatments in week 5 (Fig. 4D), providing further support for S:S regulation of photosynthesis. The photosynthetic responses of the B treatment appears puzzling at first because debudding presumably should have increased source:sink (biomass). Previous studies examining photosynthetic responses to debudding suggest that the direction of the response is species specific. In herbaceous species, sink manipulation can decrease (Plaut et al., 1987), increase (von Caemmerer and Farquhar, 1984), or effect no change in A (Plaut et al., 1987; Evans, 1991). In woody species, debudding has mostly been found to increase A (Syvertsen, 1994; Lavigne et al., 2001; Ozaki et al., 2004; but see Myers et al., 1999). Consistent with the latter studies, significant increases in $A_{1500}$ were also found in the B treatment, but it is necessary to keep in mind that this measurement was made 5 weeks after defoliation (Fig. 1). At this measurement date, the source:sink (biomass) of the B treatment was similar to control values, most probably reflecting the rapid stimulation of auxiliary bud development, which created new sinks.

The refoliation process, which includes the production of new leaf and buds, is critical for the restoration of the S:S ratio and, for evergreen woody tree species with an indeterminate growth habit such as E. globulus, rapid refoliation after defoliation is possible. It is unclear how evergreen trees with a determinisitic growth habit would restore S:S ratios, particularly after a post-flush defoliation event. Interestingly, for some of these tree species, the increase in photosynthesis has been shown to be much more long-lived, up to 16 weeks rather than only 5 weeks (Reich et al., 1993; Eyles et al., 2011). Longer durations of photosynthetic up-regulation have also been observed for older trees, which potentially may require a longer time to restore S:S balance, as their sinks are relatively much larger than those of young trees. Alternatively, the demand for new resources by alternative sinks such as stem and/or roots may increase during periods when shoot growth ceases (Reich et al., 1993).

Reductions in NSC in source leaves have been suggested as evidence for a carbohydrate-mediated feedback between carbon sinks and sources (Zhou and Quebedeaux, 2003). Therefore, a negative relationship between carbohydrate and $A_{1500}$ would have been expected. While no such relationships were observed for either soluble sugar or starch (Fig. 3E, F), there was a negative relationship between $A_{1500}$ and sucrose for the two defoliation treatments ($A_{1500} = -1.02$Sucrose+23.12, $r^2 = 0.85$, $P < 0.001$, Fig. 3G), which supports the hypothesis. Sucrose is the major transport carbohydrate in plants (Koch, 2004).

Role of carbohydrates as signalling molecules in photosynthetic up-regulation

The present results suggest that increases in $A_{1500}$ were not related to variation in leaf-level bulk NSC, as similarly reported
in other studies (Lavigne et al., 2001; Turnbull et al., 2007; Quentin et al., 2010). Instead, the analysis of specific soluble sugars showed that increases in \( A_{1500} \) were not accompanied by the same changes in the carbohydrate profile across treatments (Table 1). In particular, strong positive correlations were found between galactinol, stachyose, and, to a lesser extent, raffinose and photosynthetic responses (Fig. 3). In addition to their critical roles as carbon and energy sources, carbohydrates are also recognized as signalling molecules in growth and development, and stress-related responses (Valluru and Van den Ende, 2011; Bolouri Moghaddam and Van den Ende, 2012). While sucrose is the major transport carbohydrate in plants (Koch, 2004), other small oligosaccharides such as raffinose and galactinol are also phloem mobile (Keller and Pharr, 1996). The present results provide some evidence that galactinol, stachyose, and raffinose may potentially function as signalling molecules in the regulation of gas exchange. Future experiments should focus on the putative signalling role of these sugars, monitoring their presence not only in leaves but also in phloem sap. It is interesting to note that the carbon content of phloem sap of \( E. \) globulus is dominated by sucrose and raffinose (Merchant et al., 2001). The latter has received little attention in the context of defoliation studies but, given that phloem is the main pathway for the movement of solutes and signalling among tissues of higher plants, examining the effects of defoliation on the composition of phloem sap will provide insights into the regulatory role of specific carbohydrates.

**Conclusion**

Although photosynthetic responses to defoliation treatments were influenced by both leaf- and whole-plant-level factors, by directly comparing these variables with multiple linear regression, strong evidence was provided that the direction and duration of these photosynthetic responses were regulated by changes in whole-plant S:S ratios in \( E. \) globulus saplings. There was little evidence that \( A_{1500} \) was related to \( K_p \) in week 5, suggesting that changes in \( A_{1500} \) were not hydraulically mediated. At a leaf level, only the interaction of \( g \) and galactinol explained photosynthetic responses to defoliation, and, given the between-treatment variation in the relationship between \( A_{1500} \) and \( g \) alone, there cannot be confidence in the capacity of this relationship to explain photosynthetic responses to defoliation without further examination of the role of galactinol. Strong positive correlations were observed between three specific carbohydrates: galactinol, stachyose, and raffinose, and photosynthetic responses, providing some evidence that these sugars may function as longer term signalling molecules in the regulation of gas exchange. Future studies examining whole-plant responses to defoliation treatments should be able to use the S:S hypothesis to predict the interactive effects of defoliation treatments with other stresses such as water or nutrient stress on photosynthetic responses (Pinkard et al., 2011b).

**Supplementary data**

Supplementary data are available at JXB online.

**Method S1.** Full description of the ultra-performance liquid chromatography–mass spectrometry (UPLC-MS) method used to analyse leaf soluble sugars.

**Figure S1.** Relationship between \( A_{1500} \) (measured directly) and leaf-level \( N_{mass} \) (a), SLA (b), \( V_{Cmax} \) (c), and \( J_{max} \) (d) of \( Eucalyptus \) globulus following treatments in week 5.

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