Light Effects on Wax Begonia: Photosynthesis, Growth Respiration, Maintenance Respiration, and Carbon Use Efficiency

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Abstract. The effect of increasing daily light integral (DLI; 5.3, 9.5, 14.4, and 19.4 mol·m⁻²·d⁻¹) on photosynthesis and respiration of wax begonia (Begonia semperflorens-cultorum Hort.) was examined by measuring CO₂ exchange rates (CER) for a period of 25 d in a whole-plant gas exchange system. Although plant growth rate (GR, increase in dry weight per day) increased linearly with increasing DLI, plants grown at low DLI (5.3 or 9.5 mol·m⁻²·d⁻¹) respired more carbohydrates than were fixed in photosynthesis during the early growth period (13 and 4 d, respectively), resulting in a negative daily carbon gain (DCG) and GR. Carbon use efficiency [CUE, the ratio of carbon incorporated into the plant to C fixed in gross photosynthesis (Pc)] of plants grown at low DLI was low, since these plants used most of the C fixed in Pc for maintenance respiration (Rm), leaving few, if any, C for growth and growth respiration (Rg). Maintenance respiration accounted for a smaller fraction of the total respiration with increasing DLI. In addition, the importance of Rg in the carbon balance of the plants decreased over time, resulting in an increase in CUE. At harvest, crop dry weight (DWcrop) increased linearly with increasing DLI, due to the increased photosynthesis and CUE at high PPF.

Dry matter production and crop growth rate are strongly correlated to the amount of light intercepted by plants (Lawlor, 1995). To capture the maximum amount of radiation and optimize light use, shade-grown plants may undergo various physiological and morphological modifications. For example, a decrease in leaf number and an increase in specific leaf area (leaf area per unit leaf weight) with decreasing DLI have been reported in tall fescue [Festuca arundinacea Schreb. (Allard et al., 1991)] and dracaena [Dracaena sanderana Hort. Sander ex Mast (Vladimirova et al., 1997)], while a decrease in leaf number, leaf thickness, and stomatal number have been reported in mangosteen [Garcinia mangostana L. (Weibel et al., 1994)]. Other studies have reported modifications in photosynthetic (Nocini et al., 1991b) and respiration rates (Noguchi et al., 2001) in response to varying levels of DLI (Nocini et al., 1991b; Noguchi et al., 2001). However, the ability of shade grown plants to increase their photosynthesis when exposed to high photosynthetic photon flux (PPF) is limited, because of relatively low levels of carboxylation enzymes and components of the electron transport chain (Björkman, 1981). Indeed, research has indicated a decrease in the quantum yield of shade plants grown under high DLI (Funnell et al., 2002).

Plant growth results when carbohydrates synthesized in photosynthesis exceed those lost in respiration. Experimental evidence indicates that 30% to 50% of carbohydrates synthesized in photosynthesis are lost in respiration during crop growth (Amthor, 1984; Lawlor, 1995; van Irsel and Seymour, 2000). Therefore, the significance of respiration in plant growth should not be underestimated. Research on whole-plant respiration has increased since Hesketh et al. (1971) and McCree (1974) related respiration to Pp, dry weight of plants, and separated it into maintenance and growth components. According to Penning de Vries (1975), Rm involves release of energy (ATP) only for subcellular maintenance processes, whereas Rg involves release of energy for all processes that result in a net increase in biomass. To understand the physiological basis of growth, both photosynthesis and respiration have to be studied. Although a large amount of literature is available on the effects of light on photosynthesis in shade plants (e.g., Callan and Kennedy, 1995; Funnell et al., 2002; Greenway and Lieffers, 1997; Nocini et al., 1991a, 1991b), few studies have focused on respiration, and Rg and Rm, in particular, in shade plants grown at varying DLI levels.

Most of the commercially available equipment for measuring photosynthesis and respiration in plants is designed for leaf measurements. Traditional techniques of measuring leaf photosynthesis do not accurately determine the dry matter production and growth rate of plants (van Irsel and Bugbee, 2000). Continuous measurements of the CER of whole plants (for weeks) can depict the growth rate of plants more accurately than individual leaf measurements, as they directly measure accumulated C (a measure of growth rate) in plants (van Irsel and Kang, 2002). Although there have been other whole-plant photosynthesis studies in the genus Begonia (Giaglaras et al., 1995; Ehler and Hansen, 1998), these studies focused on short-term responses of photosynthesis to PPF. Short-term responses may differ greatly from long-term responses (weeks as opposed to minutes or hours), since many species have the ability to acclimate to different light levels. Effects of acclimation to light on plant growth and CER can only be determined by exposing plants to different light levels over a prolonged period. Here we measured whole-crop CER of wax begonia continuously for a period of 25 d in a 10-chamber, whole-plant gas exchange system. The objective was to quantify the effects of various DLI levels on plant growth, photosynthesis, growth respiration, maintenance respiration, and CUE of wax begonia.

Materials and Methods

Plant Material. Wax begonia ‘Cocktail Vodka’ plug seedlings

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were procured in cell flats (288 cells/flat) from a commercial grower (Speedling Inc., Blairsville, Ga.) on 13 June 2001. Seedlings were transplanted into 36-cell flats (160 mL/cell, Jumbo 606; TLC polyform, Plymouth, Minn.) filled with a soilless growing medium (Fafard 2P; Fafard Co., Anderson, S.C.). About one gram of a 14N–6.1P–11.6K slow-release fertilizer (14–14–14 Osmocote, The Scotts Co., Marysville, Ohio) was added to the growing medium in each cell prior to transplanting. One cell in each flat was left empty for inserting an irrigation pipe, which was inserted through a hole in the side of the gas exchange chamber. Flats were placed in a watertight tray, lined with a double-layer of capillary mat (Vattex F capillary watering system, OS plastics, Norcross, Ga.). A portion of the capillary mat was hung over one side of the tray to allow excess water to drain. About 2 L of water was added to the trays at approximately weekly intervals. Immediately after transplanting, the plants were placed inside whole-plant gas exchange chambers (0.32 × 0.5 × 0.6 m³) (van Iersel and Bugbee, 2000) arranged inside two growth chambers (model E-15; Conviron, Winnipeg, Canada). Ten gas exchange chambers were used in the experiment, two of which were controls (Van Iersel and Bugbee, 2000) arranged inside two growth chambers

To obtain the required DLI levels, gas exchange chambers were covered with shade cloth of varying thickness. Before placing plants inside the chambers, the DLI at canopy height was measured simultaneously in eight chambers using quantum sensors (QSO; Apogee Instruments Inc., Logan, Utah) connected to a datalogger (CR10X; Campbell Sci.). The average DLI measured in different treatments were 5.3, 9.5, 14.4, and 19.4 mol·m⁻²·d⁻¹ [corresponding to instantaneous PPF of 106, 189, 286, and 385 µmol·m⁻²·s⁻¹, respectively]. The light source was a mix of fluorescent and incandescent lamps. A photoperiod of 14 h was maintained inside the growth chambers.

**CO₂ Exchange Rate Measurements.** Data were collected using a previously described whole-plant gas exchange system (van Iersel and Bugbee, 2000), which directly measures net-photosynthesis (Pn, µmol·s⁻¹) and dark respiration (Rd, µmol·s⁻¹) of each crop semi-continuously for 30 s, once every 10 min (see Table 1 for an explanation and units of the measured and calculated variables). Measurement of CER in each chamber took 1 min per chamber, because an additional 30 s were needed to purge air from the previously measured chamber from the tubing, followed by a 30-s period of actual CER measurement. The CER was measured for 25 d. About 0.4 L·s⁻¹ of ambient air was blown into each gas exchange chamber by a rotary vane blower (DT 3.40; Becker, Cuyahoga Falls, Ohio). The air flow into each gas exchange chamber was measured using mass flow meters (GF M37-32; Aalborg Instruments and Controls, Monsey, N.Y.) and the CO₂ concentration of air from the blower was measured with an IRGA (SBA-1; PP-systems, Haverhill, Mass.). The difference in the CO₂ concentration of the incoming and exiting air from gas exchange chambers was measured with an IRGA in differential mode (LI-6262; LI-COR, Lincoln, Nebr.).

The CER (µmol·s⁻¹) was calculated as the product of mass flow (moles of air per second) and the difference in CO₂ concentration of the air entering and exiting the chambers (µmol·mol⁻¹). Since the two empty control chambers did not contain plants, CER measured from these chambers is an estimate of the error due to zero drift of the IRGA. These errors were corrected by subtracting.

**ENVIRONMENTAL CONTROL.** Shielded, aspirated, type-T-thermocouples were used to measure the air temperature inside all gas exchange chambers. The chamber temperature was maintained at 25 °C with electric resistance heaters (100 W) controlled by a datalogger (CR10T; Campbell Sci.). The chamber temperature was maintained at 25 °C with electric resistance heaters (100 W) controlled by a datalogger (CR10T; Campbell Sci.). Humidity probes (HTO-45R; Rotronic, Huntington, N.Y.) were placed in four of the gas exchange chambers.**
the average CER of the two empty gas exchange chambers from that of chambers containing plants. Although CER measurements were taken every 10 min, only average values for $P_n$ (averaged over the entire light period) and $R_g$ (dark-period average) are used here. Hence, daily average gross photosynthesis ($P_g$, $\mu$mol·s$^{-1}$) was calculated as

$$P_g = P_n - R_d \quad [\text{Eq. 1}].$$

Since $R_d$ was measured as the CO$_2$ exchange rate, it is expressed as a negative value, hence a negative sign in the above equation. This equation assumes that respiration rates during the light and dark periods were equal (van Iersel and Bugbee, 2000). Since one mole of either CO$_2$ or CH$_2$O contain an equal amount of C, CER data can be directly converted in glucose equivalents. Thus, the total amount of respired carbon per day ($R_{\text{CH2O}}$, g of CH$_2$O per day) was calculated as $-R_d \times 30 \times 86,400/10^6$ (30 is the molecular weight of CH$_2$O, 86,400 the number of seconds in a day, and $10^6$ converts $\mu$mol to mol).

Daily carbon gain (µmol of C per day or the net amount of C fixed by the 35 plants in a gas exchange chamber per day) was calculated as

$$DCG = (P_n \times t_{\text{light}}) + (R_d \times t_{\text{dark}}) \quad [\text{Eq. 2}]$$

where $t_{\text{light}}$ and $t_{\text{dark}}$ are the duration of the light and dark periods (s), respectively.

Integrating DCG over time (t) provides an estimate of cumulative carbon gain (CCG, mol of C; the total amount of carbon accumulated in plants since the start of experiment, a measure of plant mass):

$$CCG = \int DCG \, dt \quad [\text{Eq. 3}]$$

Cumulative carbon gain at the end of the experiment was plotted against $DWCROP$ (g) in each treatment, and a linear equation was fitted to describe the relationship:

$$DWCROP = DWO_0 + 12 \times CCG/f_c \quad [\text{Eq. 4}]$$

where $DWO_0$ is the estimated initial dry weight of the plants before starting the experiment, 12 converts moles of carbon to grams of carbon, and $f_c$ is the estimated carbon content of the plants (g of C/g of dry mass). Dry weight of plants at the end of each day ($DWCROP$) in different treatments was estimated from this equation, based on the CCG of the plants.

To look at the effects of DLI on CER on the efficiency of light used to fix C, CCG/m$^2$ ground area was determined by dividing CCG by the ground area of the gas exchange chamber (0.184 m$^2$). Cumulative DLI (DLI$_{\text{cum}}$, mol·m$^{-2}$) was determined by multiplying DLI by 25 d (duration of the gas exchange study), and CCG/DLI$_{\text{cum}}$ (mol of carbon fixed during the growing period per mol of incident light) was calculated from these data.

Growth rate (g of dry mass per day) of plants in different treatments can be estimated from DCG (mold$^{-1}$) as

$$GR = 12 \times DCG/f_c \quad [\text{Eq. 5}]$$

dividing CCG by the ground area of the gas exchange chamber.

Relative growth rate ($RGR$, g of new dry mass per g of existing dry mass per day) of the plants was estimated from GR and $DWCROP$:

$$RGR = GR/DWCROP \quad [\text{Eq. 6}]$$

Carbon use efficiency (mol of C incorporated per mol of C fixed in gross photosynthesis) was calculated as follows:

$$\text{CUE} = P_n/(P_n \times t_{\text{light}}) \quad [\text{Eq. 7}]$$

Growth ($g_r$) and maintenance ($m_r$) respiration coefficients can be estimated from the specific respiration rate ($R_{\text{spec}}$, g of CH$_2$O per g of dry mass per day) and $RGR$ (Hesketh et al., 1971):

$$R_{\text{spec}} = R_{\text{CH2O}}/DWCROP = m_r + g_r \times RGR \quad [\text{Eq. 8a}]$$

This equation was modified for potential DLI effects on $g_r$ and $m_r$:

$$R_{\text{spec}} = a_0 + a_1 \times DLI + (b_0 + b_1 \times DLI) \times RGR \quad [\text{Eq. 8b}]$$

where $m_r = a_0 + a_1 \times DLI$ (g of CH$_2$O per g of dry mass per day) and $g_r = b_0 + b_1 \times DLI$ (g of CH$_2$O per g of dry mass).

Growth and maintenance respiration rates (g of CH$_2$O per day) of plants were estimated as

$$R_g = g_r \times GR \quad [\text{Eq. 9}]$$

and

$$R_m = m_r \times DW_{\text{day}} \quad [\text{Eq. 10}]$$

To determine the relationship between CUE and RGR or $R_{\text{CH2O}}/R_{\text{CDEO}}$, we fitted rectangular hyperbolas [CUE = $x/(a \times x + b)$], since van Iersel (2003) has shown that

$$1/CUE = 1 + g_r + m_r/RGR \quad [\text{Eq. 11a}]$$

can be rearranged as

$$\text{CUE} = RGR/[(1 + g_r) \times RGR + m_r] \quad [\text{Eq. 11b}]$$

where $g_r$ is in units of moles of C respired per mol of C incorporated into the plants, and $m_r$ is in mol of C respired per mol of C in the plants per day.

**Final Measurements.** At harvest (25 d after transplanting), leaf chlorophyll concentration, leaf area, and shoot dry weight of 23 plants were determined. The remaining 12 plants were left in the gas exchange chambers for an additional day (for light response curves; Nemali, 2002), after which their root and shoot dry weights were determined. Leaf chlorophyll concentration (of 4 leaves from separate plants) was assessed using a chlorophyll meter (SPAD-502; Minolta Co., Japan) and leaf areas were measured using a leaf area meter (LI-3100; LI-COR). Total leaf area was estimated from the leaf area of the measured 23 plants. Root dry weight of the harvested 23 plants was estimated as the product of shoot dry weight of 23 plants and root to shoot ratio measured for the remaining 12 plants. Leaf area ratio (LAR) was estimated as the ratio of leaf area to $DWCROP$.

**Experimental Design and Data Analysis.** The experimental layout was a randomized complete block with two replications. Each experimental block (growth chamber) consisted of four DLI treatments (four gas exchange chambers), and each experimental unit (each gas exchange chamber) consisted of 35 plants. The effects of DLI on CER and other calculated variables were tested separately for each measurement day with linear and quadratic regression analysis. Other relationships among response variables were analyzed with both linear and nonlinear regressions, with $P < 0.05$ considered to be statistically significant.

**Results and Discussion**

**Plant Growth—DLI Relationships.** At harvest, $DWCROP$ and leaf area increased linearly, while leaf chlorophyll concentration decreased linearly with increasing DLI (Table 2). Leaf area ratio and shoot to root ratio were not affected by DLI. Increases in LAR with decreasing DLI have been reported in tall fescue (Allard et al., 1991) and dracaena (Vladimirova et al., 1997). To capture more light, plants grown in shade tend to develop larger and thinner leaves than those grown in full sunlight (Allard et al., 1991; Weibel et al., 1994). Since neither shoot-root ratio nor LAR was affected, wax begonia did not preferentially allocate a larger fraction of the available carbohydrates to leaf area production in response to a decrease in DLI from 19.4 to 5.3 mol·m$^{-2}$·d$^{-1}$. Thus, the increase in leaf area with decreasing DLI have been reported in response to shading (Adams and Demmig-Adams, 1992; Close et al., 2001), and may help plants maximize light interception under low DLI.
There was a strong correlation between DW\textsubscript{crop} and CCG ($r = 0.96$), which indicates that gas exchange data were a realistic measure of crop growth (Fig. 1). Crop dry weight increased by 21.4 g for every mole of C incorporated by the plants. Carbon content in the plants ($f_c$) was estimated from the slope of the regression line as 0.56 g·g$^{-1}$ (12 g·mol$^{-1}$/21.4 g·mol$^{-1}$). The estimated C content in wax begonia was higher than most other reported values, 0.465 g·g$^{-1}$ for pansy ($Viola \times wittrockiana$ Gams.) (van Iersel and Kang, 2002), 0.45 g·g$^{-1}$ [pumpkin leaves ($Cucurbita pepo$ L.) (Turgeon and Webb, 1975)], 0.396 g·g$^{-1}$ [white clover ($Trifolium repens$ L.) (McCree and Troughton, 1966a)], and 0.421 g·g$^{-1}$ [sugar beet leaves ($Beta vulgaris$ L.) (Terry and Mortimer, 2002)].

As expected, C fixation of the plants was closely related to DLI. There was a quadratic relationship between CCG and DLI\textsubscript{cum} (Fig. 2, top). The intercept of the regression line with the x-axis (125 mol·m$^{-2}$, equivalent to 5.0 mol·m$^{-2}$·d$^{-1}$, or 100 µmol·m$^{-2}$·s$^{-1}$ with a 14-h photoperiod) is the estimated DLI at which there would have been no net gain of C, i.e., the functional light compensation point. The functional light compensation point is higher than the light compensation point determined from photosynthesis–PPF response curves, which was 22 µmol·m$^{-2}$·s$^{-1}$ for these plants (Nemali, 2002). Note that the light compensation point as determined from photosynthesis–PPF response curves is an indication of the

| DLI (mol·m$^{-2}$·d$^{-1}$) | Chlorophyll concn (spad units) | Leaf area (m$^2$) | DW\textsubscript{CROP} (g) | DW\textsubscript{SHOOT} (g) | DW\textsubscript{ROOT} (g) | Leaf area ratio (m$^2$·kg$^{-1}$) | Shoot to root ratio |
|--------------------------|-----------------------------|----------------|----------------|----------------|----------------|------------------|------------------|
| 3.3                      | 30.3                        | 0.8            | 23.9          | 16.4          | 7.6           | 31.5             | 2.2              |
| 9.5                      | 29.3                        | 1.1            | 30.5          | 19.8          | 10.7          | 36.1             | 1.9              |
| 14.4                     | 29.2                        | 1.3            | 36.3          | 26.3          | 10.1          | 35.8             | 2.6              |
| 19.4                     | 28.5                        | 1.4            | 38.4          | 27.2          | 11.2          | 37.5             | 2.4              |
| $r$                      | -0.84                       | 0.89           | 0.94          | 0.94          | ---           | ---              | ---              |
| intercept                | 30.6$^{***}$                | 0.58$^{**}$    | 19.4$^{***}$  | 12.2$^{***}$  | 9.9           | 35.2             | 2.3              |
| DLI$\textsubscript{coef}$| -0.11$^*$                   | 0.05$^{**}$    | 1.07$^{***}$  | 0.85$^{***}$  | NS            | NS               | NS               |

NS,*,**,***Nonsignificant or significant at $P < 0.05$, or 0.005, or 0.0005, respectively.

DAILY LIGHT INTEGRAL AND CO$_2$ EXCHANGE RATES. There was a strong correlation between DW\textsubscript{crop} and CCG ($r = 0.96$), which indicates that gas exchange data were a realistic measure of crop growth (Fig. 1). Crop dry weight increased by 21.4 g for every mole of C incorporated by the plants. Carbon content in the plants ($f_c$) was estimated from the slope of the regression line as 0.56 g·g$^{-1}$ (12 g·mol$^{-1}$/21.4 g·mol$^{-1}$). The estimated C content in wax begonia was higher than most other reported values, 0.465 g·g$^{-1}$ for pansy ($Viola \times wittrockiana$ Gams.) (van Iersel and Kang, 2002), 0.45 g·g$^{-1}$ [pumpkin leaves ($Cucurbita pepo$ L.) (Turgeon and Webb, 1975)], 0.396 g·g$^{-1}$ [white clover ($Trifolium repens$ L.) (McCree and Troughton, 1966a)], and 0.421 g·g$^{-1}$ [sugar beet leaves ($Beta vulgaris$ L.) (Terry and Mortimer, 2002)].

As expected, C fixation of the plants was closely related to DLI. There was a quadratic relationship between CCG and DLI\textsubscript{cum} (Fig. 2, top). The intercept of the regression line with the x-axis (125 mol·m$^{-2}$, equivalent to 5.0 mol·m$^{-2}$·d$^{-1}$, or 100 µmol·m$^{-2}$·s$^{-1}$ with a 14-h photoperiod) is the estimated DLI at which there would have been no net gain of C, i.e., the functional light compensation point. The functional light compensation point is higher than the light compensation point determined from photosynthesis–PPF response curves, which was 22 µmol·m$^{-2}$·s$^{-1}$ for these plants (Nemali, 2002). Note that the light compensation point as determined from photosynthesis–PPF response curves is an indication of the
PPF at which \( P_n \) is zero. Plants grown at a PPF level at which \( P_n \) is zero would have a negative DCG, since there would be no net C exchange during the light period and a C loss during the dark period. Plants grown at the functional light compensation point would have a net C gain during the light period that equals the C loss in the dark.

There was a quadratic relationship between CCG/DLI\(\text{cum} \) and DLI (Fig. 2 bottom). The ratio of CCG to DLI\(\text{cum} \) is a measure of the efficiency with which plants used the incident light to incorporate C. This efficiency was very low (\( \approx 0.93 \) mmol·mol\(^{-1} \)) at 5 mol·m\(^{-2} \)·d\(^{-1} \), reached a maximum of 9.43 mmol·mol\(^{-1} \) at 15.6 mol·m\(^{-2} \)·d\(^{-1} \), and decreased again slightly at higher DLIs. Treatment effects on CCG/DLI\(\text{cum} \) can be due to differences in light absorption, \( P_g \) per unit absorbed PPF, and the conversion of photosynthates into structural dry matter (CUE). The low ratio of CCG to DLI\(\text{cum} \) at low light intensity was due at least partly to low light interception, since leaf area increased linearly with increasing DLI (Table 2). In addition, CUE increased with increasing DLI (see below), while \( P_g \) per unit absorbed PPF could not be determined, since only incident, not absorbed, PPF was measured. However, the photosynthetic efficiency of leaves (carbon fixed per unit absorbed PPF) decreases, when photosynthesis becomes light saturated (Jones, 1992). This can explain the decrease in CCG/DLI\(\text{cum} \) when DLI was greater than 15.2 mol·m\(^{-2} \)·d\(^{-1} \). As in Fig. 2 top, the x-intercept in Fig. 2 bottom (4.5 mol·m\(^{-2} \)·d\(^{-1} \) or 88 \( \mu \)mol·m\(^{-2} \)·s\(^{-1} \) with 14-h photoperiod) is an estimate of the functional light compensation point. The small difference between the two estimated functional light compensation points can be attributed to error in the data points used for the regressions.

Gross photosynthesis, \( P_g \), and \( R_d \) of the plants increased linearly with increasing DLI and increased steadily throughout the experiment (Fig. 3). This continuing increase in CO\(_2\) assimilation suggests that even near the end of the experiment, the plants did not use all incoming \( PPF \) for \( \text{CO}_2 \) assimilation, presumably due to an incomplete absorption of the incoming \( PPF \). The continued increase in photosynthesis throughout the experiment is consistent with the low efficiency at low light intensity. However, the photosynthetic efficiency of leaves (carbon fixed per unit absorbed PPF) decreases, when photosynthesis becomes light saturated (Jones, 1992). This can explain the decrease in CCG/DLI\(\text{cum} \) when DLI was greater than 15.2 mol·m\(^{-2} \)·d\(^{-1} \). As in Fig. 2 top, the x-intercept in Fig. 2 bottom (4.5 mol·m\(^{-2} \)·d\(^{-1} \) or 88 \( \mu \)mol·m\(^{-2} \)·s\(^{-1} \) with 14-h photoperiod) is an estimate of the functional light compensation point. The small difference between the two estimated functional light compensation points can be attributed to error in the data points used for the regressions.
with the finding that the growth of white clover keeps increasing with increasing leaf area index (McCree and Troughton, 1966b). The leaf area index of wax begonias grown at 19.4 mol·m–2·d–1 was 8.8 at the end of the experiment.

Although Rg in all treatments generally increased throughout the experiment, there was a distinct decrease in Rg after plants were watered on day 13. This was accompanied by an increase in DCG (Fig. 4), which indicates that growth of the plants was limited by a mild drought stress prior to this irrigation. There were no visible symptoms of drought (e.g., wilting) at any time.

Negative DCG values (i.e., net carbon losses) were seen during the initial 13 and 4 d for plants grown at a DLI of 5.3 and 9.5 mol·m–2·d–1, respectively (Fig. 4). This may be partly due to microbial breakdown of the organic matter in the growing medium, which increases total respiration, and thus decreases DCG. However, even under a high DLI of 21.4 mol·m–2·d–1, negative DCG values were reported after transplanting vinca (Catharanthus roseus L.) (van Iersel, 1999) into an inorganic growing medium, where microbial breakdown of organic components of the growing medium did not affect the measurements. This indicates that plants can respire more carbohydrates than are synthesized in photosynthesis soon after transplanting (possibly from storage forms like starch). This may be related to the need to repair damage caused by transplanting.

Since CCG is DCG integrated over time, treatment effects on DCG resulted in differences in CCG as well. Thus, plants grown at 5.3 and 9.5 mol·m–2·d–1 had a negative CCG during the initial 20 and 6 d of crop growth, respectively (Fig. 4). Both DCG and CCG were positively correlated with DLI throughout the experiment (P < 0.05), which would be expected based on PPF effects on photosynthesis (Fig. 2).

Relative growth rate increased linearly with DLI until day 10, and from day 10 onwards, quadratic responses of RGR to DLI were seen (Fig. 4). There was a large increase in RGR with an increase in DLI from 5.3 to 14.4 mol·m–2·d–1, followed by only a small or no increase in RGR as DLI increased from 14.4 to 19.4 mol·m–2·d–1. From day 21 onwards, RGR was similar for plants grown at 14.4 and 19.4 mol·m–2·d–1 (Fig. 4). Somewhat surprisingly, RGR of the plants increased throughout the first 18 d, after which it remained fairly steady in all treatments. Normally, RGR decreases with increasing plant size because of increased intra- and inter-plant competition for light (Kvet et al., 1971). The low RGR during the early part of this experiment may be due to transplant shock and acclimation of the plants to the environmental conditions in the gas exchange chambers. At harvest, the RGR for plants grown at 5.3, 9.5, 14.4, and 19.4 mol·m–2·d–1 was 0.013, 0.027, 0.036, and 0.035 g·g–1·d–1, respectively. Similar to DCG, RGR was negative during the initial 4 and 13 d for plants grown at 5.3 and 9.5 mol·m–2·d–1. These estimated RGR values...
Increased gr, but had no significant effect on m, indicating that gr remained constant and m decreased from 65 to 39 mg·g⁻¹·d⁻¹ when plants were shifted from high to low irradiance. In contrast to that experiment, plants in our study were grown at lower, constant DLI levels and probably better adapted to low DLI due to the shade tolerance of wax begonia.

Literature values for m differ widely. In their review, Hesketh and Jones (1980) reported m values for many species ranging from 6 to 91 mg·g⁻¹·d⁻¹, while Amthor (2000) states that m falls in the range of 3.3 to 138 mg·g⁻¹·d⁻¹. The estimated m of wax begonia (31 mg·g⁻¹·d⁻¹) is well within these ranges, and similar to that of Italian ryegrass tops [Lolium multiflorum L. (37 mg·g⁻¹·d⁻¹)], higher than that of chrysanthemum [Chrysanthemum morifolium L. (17 mg·g⁻¹·d⁻¹)], perennial ryegrass [Lolium perenne L. (14 mg·g⁻¹·d⁻¹)], and tomato [Lycopersicon esculentum L. (12 mg·g⁻¹·d⁻¹)], but lower than that of roots of Italian ryegrass (91 mg·g⁻¹·d⁻¹) (Hesketh and Jones, 1980).

Our g estimates (0.40 to 0.62 g·g⁻¹) were slightly higher than those of maize [Zea mays L. (0.34 g·g⁻¹)] and cotton [Gossypium hirsutum L. (0.33 to 0.39 g·g⁻¹)], similar to that of chrysanthemum (0.56 g·g⁻¹), and lower than that of roots of Italian ryegrass (0.67 g·g⁻¹) (reviewed by Hesketh and Jones, 1980). Experimental estimates from a variety of studies suggest that g is similar (generally differing by <10%) among plant parts, species, and growing conditions (Cannell and Thornley, 2000), and normally close to 0.4 g·g⁻¹. However, g depends on which chemical compounds are synthesized by the plants (Penning de Vries et al., 1974), which in turn may depend on DLI. Plants grown under a high DLI generally have higher concentrations of photosynthetic enzymes (Hikosaka, 1996), and the production of enzymes requires a relatively much energy (Penning de Vries et al., 1974). Therefore, an increase in DLI is likely to result in an increase in g, as well.

Since Rg and Rm were calculated as g × GR and m × DWcrop, respectively, trends in Rg and Rm were similar to those for DCG and CCG. Both Rg and Rm increased with increasing DLI throughout the experiment (results not shown). Throughout the entire experiment, and at all DLI levels, Rm was greater than Rg (Fig. 6). At the start of the experiment, Rm accounted for almost all of Rg (97 to 100%). At harvest, the percentage of Rm was 91, 79, 69, and 66% of Rg for plants grown at 5.3, 9.5, 14.4, and 19.4 mol·m⁻²·d⁻¹, respectively. The fraction of Rg that was used for growth increased throughout most of the experiment and with increasing DLI (Fig. 6). Growth respiration accounted for a higher fraction of Rg with increasing DLI because both DCG and g increased with increasing DLI. Usually, Rg accounts for most (<80%) of the total respiration during the vegetative stage, and decreases during the reproductive stage (to ~20%; Lawlor, 1995). The high ratio of Rm to Rg of wax begonia may be an indirect effect of the low photosynthetic rate of these plants. At light saturation, Pₚ per unit leaf area of these plants ranged from 4.5 mmol·m⁻²·s⁻¹ for plants grown at a DLI level of 5.3 mol·m⁻²·d⁻¹ to 8.1 mmol·m⁻²·s⁻¹ for plants grown at 19.4 mol·m⁻²·d⁻¹ (Nemali, 2002), while typical maximum leaf Pₚ rates of C₃ plants range from 14 to 40 mmol·m⁻²·s⁻¹ (Jones, 1992). Thus, wax begonias produce relatively low levels of carbohydrates, and most of these are used for Rg, leaving a small carbohydrate pool for growth and Rm. This also can explain the slow growth habit of wax begonias.

**Carbon Use Efficiency.** At the lowest DLI level (5.3 mol·m⁻²·d⁻¹), this equation, the estimated m for plants was 0.031 g·g⁻¹·d⁻¹ and g for plants grown at 5.3, 9.5, 14.4, and 19.4 mol·m⁻²·d⁻¹ were 0.40, 0.47, 0.54, and 0.62 g·g⁻¹, respectively. An earlier study on white clover (McCree, 1982) grown initially at high PPF (1750 mmol·m⁻²·s⁻¹) and subsequently at a lower PPF (350 mmol·m⁻²·s⁻¹) indicated that g remained constant and m decreased from 65 to 39 mg·g⁻¹·d⁻¹ when plants were shifted from high to low irradiance.
CUE increased from -0.48 to 0.21 mol·mol⁻¹ during the experiment, while CUE increased from 0.07 to 0.45 mol·mol⁻¹ at a DLI level of 19.4 mol·m⁻²·d⁻¹ (Fig. 7A). Carbon use efficiency generally ranges from 0.5 to 0.7 mol·mol⁻¹ (Bednarz and van Iersel, 1999; Gifford 1995). The low CUE of wax begonia can be explained from its low photosynthetic rate, and the large fraction of carbohydrates used for Rₚ (Fig. 6).

Carbon use efficiency of plants responded quadratically to an increase in DLI. There was an increase in CUE from 5.3 to 14.4 mol·m⁻²·d⁻¹, while CUE was similar at 14.4 and 19.4 mol·m⁻²·d⁻¹ (Fig. 7A). There was a sudden increase in CUE after watering on day 14, due to a decrease in Rₚ and an increase in DCG (Figs. 3 and 4). In all DLI treatments, CUE increased during the first 14 d of the experiment. At the two lowest light levels, CUE was negative early in the experiment, a direct effect of the negative DCG as plants did not produce enough carbohydrates to meet maintenance requirements during the first 13 and 4 d, respectively (Fig. 6). A negative DCG indicates that plants were catabolizing reserves or structural dry matter to meet the respiratory demand.

Although CO₂ production from microbial breakdown of organic components of the growing medium is negligible compared to the metabolic activity of rapidly growing plants (van Iersel and Bugbee, 2000), it may be significant when the CO₂ exchange rate of the plants is low. This would result in an underestimation of DCG, not affect Pₙ estimates (unless microbial breakdown of the growing medium differs in the light and in darkness), and therefore result in an underestimation of CUE. This may have been a factor contributing to the low CUE of plants grown at low DLI during the early part of the growing period. However, previous work has shown that a negative CUE may occur after transplanting, even when there is no microbial breakdown of the growing medium. After vinca was transplanted into an inorganic growing medium, Rₚ increased rapidly and CUE became negative (van Iersel, 1999).

There was a close relationship between CUE and the ratio of Rₚ to Rₙ₋₋ₐₐ₆ (R² = 0.96) and between CUE and RGR (R² = 0.99) (Fig. 7B, C). The regression results from the hyperbolic relationship between CUE and RGR can be used to estimate gₚ and mₙ. Based on Eq. 11b, estimates of gₚ and mₙ are 0.383 mol·mol⁻¹ and 0.0222 mol·mol⁻¹·d⁻¹, respectively. These estimates can be converted to 0.54 g·g⁻¹ (grams of glucose per gram dry matter) for gₚ and 0.031 g·g⁻¹·d⁻¹ for mₙ, based on an fc of 0.56 for the plant material and 0.40 for glucose. Since data from all DLI treatments were combined in this regression analysis, these estimates ignore DLI effects on gₚ and thus represent an average for the different treatments. Nonetheless, the regression resulted in an excellent fit (R² = 0.99), so apparently any differences in gₚ among DLI treatments had little or no effect on CUE. Presumably, this was the case because Rₚ only accounted for a small fraction of total respiration (Fig. 6).

The asymptotes of the regressions of CUE versus the ratio of Rₚ to Rₙ₋₋ₐₐ₆ (0.74 mol·mol⁻¹) and versus RGR (0.72 mol·mol⁻¹) (Fig. 7B, C) are estimates of the theoretical maximum for CUE (i.e., CUE in the absence of any Rₚ) for wax begonia. These values are similar to estimated maximum CUEs of pansy (0.67 mol·mol⁻¹; van Iersel and Kang, 2002) and lettuce (Lactuca sativa L.) (0.68 mol·mol⁻¹; van Iersel, 2003). The actual CUE of the plants was much lower (<0.5) than the theoretical maximum because a large fraction of photosynthates was used for maintenance.

Conclusions

Plants grown at 5.3 and 9.5 mol·m⁻²·d⁻¹ had a low growth rate due to inadequate incident DLI, insufficient light interception (at least partially due to small leaf areas), and low photosynthetic rates. Growth rate of plants increased with increasing DLI because both Pₚ and CUE increased with increasing DLI. At the lower DLI (5.3 and 9.5 mol·m⁻²·d⁻¹), photosynthesis was initially insufficient to meet maintenance requirements for carbohydrates, resulting in a negative DCG, RGR, and CUE. The importance of Rₚ in the carbon balance, and thereby on the growth rate of the plants, decreased with increasing DLI. At low DLI, plants produce relatively few carbohydrates, and only a small fraction of this carbohydrate pool is available for growth after maintenance requirements are met. This explains the higher Rₚ/Rₙ₋₋ₐₐ₆ and lower growth rate of plants grown at 5.3 and 9.5 mol·m⁻²·d⁻¹. Wax begonia had lower CUE and RGR than most reported values for other crops. This too may be attributed to the low photosynthetic capacity of wax begonia. Although considered to be shade tolerant, growth of wax begonias is inhibited at DLI levels of 9.5 mol·m⁻²·d⁻¹, which is considered a moderate amount of light inside a greenhouse. Growth and CUE of wax begonia were constrained because they had more Rₚ than Rₚ. Photosynthetic rates were much lower than those of other C₃ species even at the highest DLI. A future issue would be to identify the physiological reason for these low photosynthetic rates.

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