Modification of Petunia Seedling Carbohydrate Partitioning by Irradiance

David F. Graper and Will Healy
Department of Horticulture, University of Maryland, College Park MD 20742

Abstract. Petunia × hybrida Vill. ‘Red Flash’ plants received either 10 or 20 mol·day⁻¹ photosynthetic photon flux (PPF) in growth chambers at: 175 µmol·m⁻²·s⁻¹ for 16 hours, 350 µmol·m⁻²·s⁻¹ for 8 or 16 hours, or 350 µmol·m⁻²·s⁻¹ for 8 hours plus 8 hours of incandescent photoperiod extension (5 µmol·m⁻²·s⁻¹ PPF). The irradiation components of peak, total, and duration were examined. Doubling total PPF increased total carbohydrate (CHO) production by 60%, seedling dry weight (DW) by 30%, rate of seedling growth by 25%, and acid invertase activity by 50% compared to the other treatments, once the seedlings had reached the two-leaf stage. Seedlings receiving 20 mol·day⁻¹ PPF partitioned 14% more CHO into ethanol soluble sugars rather than starch, which may explain the increase in relative growth rate observed with supplemental irradiance treatments. Extending the photoperiod for 8 hours with 5 µmol·m⁻²·s⁻¹ PPF reduced total CHO production by 50% compared to the same treatment without photoperiodic lighting. Treatment with 350 µmol·m⁻²·s⁻¹ for 8 hours resulted in the highest O₂ evolution (8.8 µmol O₂/min per dm²). Increasing the photoperiod from 8 to 16 hours gave the lowest rate of O₂ evolution (4.5 µmol O₂/min per dm²). Previous reports of the importance of photosynthetic period in controlling partitioning between starch and sugars may have simply observed a decreasing rate of starch accumulation due to increased total PPF.

Peak PPF and the photosynthetic (Pn) period are two components of irradiance that are influential in CHO metabolism and transport (Chatterton and Silvius, 1980b; Grange, 1985; Silvius et al., 1979). Elevating peak PPF increased total net photosynthesis 25% and resulted in a higher rate of soluble CHO translocation within Glycine max L. (Silvius et al., 1979). Transferring G. max plants from 600 to 950 µmol·m⁻²·s⁻¹ initially increased starch production by 60%, but after a 2-day lag period, starch production decreased and translocation increased by 20% (Silvius et al., 1979). Grange (1985) reported higher levels of starch in Capsicum annum L. as peak PPF was increased. The translocation rate also increased by doubling the Pn period from 7 to 14 h (Chatterton and Silvius, 1980b). In each of these experiments, peak PPF or Pn period was increased, which resulted in increased net photosynthesis, starch production, or translocation. While peak PPF or Pn period increased, total integrated PPF also increased. Therefore, the reported increases in net photosynthesis, starch production, and translocation may have actually been due, in part, to changes in total PPF rather than peak PPF or Pn period.

Petunias have been the focus of considerable research due to their importance as a bedding plant crop. Much of this earlier research focused on irradiance treatments imposed for extended periods between sowing and flowering (Carpenter and Beck, 1973; Krizek et al., 1974; Merritt and Kohl, 1982; Seeley, 1960). Petunia is a C3 plant reported to be day-neutral before the six-leaf stage (Carpenter and Carlson, 1974), at which time it responds as a long-day plant.

We used Petunia × hybrida Villm. ‘Red Flash’ seedlings as a model plant to determine the effects of photoperiod, peak PPF, and total PPF on seedling growth, Pn capacity, CHO metabolism, and invertase activity of seedlings at the two-leaf stage of development. Supplemental irradiance treatments during periods of low ambient insolation in the greenhouse simultaneously increase total PPF, peak PPF, photoperiod, and Pn period. Determining how these factors interact to accelerate photosynthesis and growth rate and alter CHO partitioning should indicate possible mechanisms involved in hastening seedling growth.

Materials and Methods

General protocol. ‘Red Flash’ petunia seed was sown in 288 plug flats (cells, 2 x 2 x 3-cm tall; cell, 6.25 ml; 288 cells per flat). Before sowing, the flats were filled about one-third full with Redi-Earth peat-lite mix (W.R. Grace and Co., Cambridge, Mass.) and saturated with tap water. About 1.5 g flat of 17N–9P–13K (17N-3.9P–10.8K) Osmocote (9.370 ammonium; Sierra Chemical Co., Milpitas, Calif.) was spread uniformly over each flat using an Old Mill seeder (Old Mill Seeder Co., Savage, Md.). The flats were then completely filled with the medium and again saturated with tap water. One seed was then sown per cell with the Old Mill seeder.

The seeded flats were placed in plastic bags and germinated in darkness near 27°C until the first cotyledons merged. After removing the bags, two flats, representing two sampling units, were randomly assigned to each of four specially designed PGC chambers (Parameter Generation and Control, Black Mountain, N. C.) continuously at 21 ± 1°C. Flats were rotated within each chamber daily to minimize the effects of light and water distribution patterns.

Plants were irradiated using eight or 16 F48PG17 cool-white fluorescent lamps per chamber, suspended 30 to 50 cm above the plants, to provide either 175 or 350 µmol·m⁻²·s⁻¹ at plant level. The lamps were isolated from the plants by a plexiglas barrier in a ventilated light cap. Plants were supplied with either 10 or 20 mol·day⁻¹ PPF using one of the following treatments: 175 µmol·m⁻²·s⁻¹ for 16 h (175/16), 350 µmol·m⁻²·s⁻¹ for 8 h (350/8) or 16 h (350/16), or 350 µmol·m⁻²·s⁻¹ for 8-h plus
Photoperiod

The light cycle within each chamber began at staggered 15-min intervals to facilitate the sampling of plants. Sufficient seedling shoots (30 to 50 shoots, cut at the medium surface) at the two-leaf stage were sampled to comprise two 500-mg fresh weight (FWT) subsamples from each flat for CHO content and 2.5 g FWT for acid invertase activity analysis. Samples for analysis were collected 2 h into the light cycle. Plants were designated at the two-leaf stage once the third true leaf had become microscopically visible. Samples for CHO analysis were frozen at –70°C. Tissue for invertase analysis was placed in cold extraction buffer on ice until all samples were ground. The number of days for the majority of the plants to reach the two-leaf stage and mean seedling DW were recorded.

The experiment was replicated in time with treatments randomly assigned to chambers before each experiment commenced. Treatment means were separated using SAS General Linear Models procedure and single degree-of-freedom comparisons testing for main effects and interactions of the four treatment effects of peak PPF, Pn period, photoperiod, and total PPF as described in Table 1 (SAS Institute, Inc., Cary, N.C.).

Carbohydrate extraction. Tissue was homogenized in 80% EtOH using a polytron (Brinkmann Instruments, Westbury, N. Y.) with a β-phenyl-D-glucose internal standard. Homogenates were placed in a boiling water bath for 15 min and centrifuged at 1100 × g for 10 min. The supernatant was decanted and saved for soluble sugar analysis. The pellet was rinsed three times with 80% EtOH and given a final rinsing with distilled, deionized H2O (Huber and Israel, 1982). The combined supernatants were evaporated and all volumes were equalized to 2 ml with H2O.

Sugar analysis. The 2 ml of CHO extracts were filtered through a 2-cm column of polyvinylpolypyrrolidone to remove phenols, placed into microtubes, frozen in liquid N2, and stored at –20°C overnight. Samples were thawed, centrifuged at 20,850 × g for 15 min, and filtered through a 0.2-µm filter disk. Soluble sugars were measured using a gas chromatograph–mass spectrophotometer following TMS (tetramethylsilane) derivatization (Li and Schuman, 1980).

Starch extraction. The pellet from EtOH soluble sugar extraction was resuspended in 1 ml of 2 N KOH and boiled for 15 min before the solution was neutralized with 200 µl of 1 N acetic acid. Starch was digested by 1 ml of amyloglucosidase (400 units/ml) in 100 mM citric acid buffer at pH 5.5. Samples were incubated for 4 h at 37°C with periodic agitation. Tubes were boiled for 1 min and centrifuged at 1100 × g for 5 min (Beaudry et al., 1989).

Starch content was calculated by measuring glucose content. The glucose oxidase reagent consisted of 60 ml of 200 mM Heps-NaOH (pH 7), 40 ml glycerol, 6 mg peroxidase, 14 mg dianisidine, and 26 mg glucose oxidase. The glucose content of a 100-µl sample of supernatant was measured. Glucose determination was made by adding 2 ml of glucose oxidase reagent plus 900 µl of 50 mM citric acid buffer at pH 5.5. Samples were incubated at 37°C for 15 min. Reactions were stopped by adding 4 ml of 5 N HCl. Glucose was measured by absorbance at 540 nm (Chatterson and Silvius, 1979).

Invertase analysis. The 2.5-g FWT sample was ground in 50 mM Heps-NaOH (pH 6.9) using a Polytron. The suspension was filtered through Miracloth, rinsed with additional buffer, and centrifuged for 15 min at 26,000 × g. Supernatant was dialyzed for 12 h in 10 mM Heps-NaOH (pH 6.9) at 4°C. The solution was replaced and allowed to dialyze an additional 6 h. Enzyme activity was measured by using 50 µl Na-acetate buffer at pH 5.5, 50 µl H2O, and 50 µl enzyme extract. Samples were incubated at 30°C for 5 min before 50 µl of 400 mM sucrose was added; they then were incubated for 15 min at 30°C. Reactions were stopped by placing samples into an ice bath and adding 1 ml of 100 mM borate buffer at pH 9.0. Reducing sugar content was then measured at 276 nm after reaction with 200 µl of 1% 2-cyanoacetamide (Gross, 1982).

Oxygen evolution measurement. Due to the small size and limited leaf expansion of the seedlings, measurement of Pn rate by CO2 uptake using infrared-gas analysis was not feasible. Therefore, plant Pn capacity was estimated by measuring shoot 0, evolution in a Hansatech LD2 leaf disk electrode (Decagon Devices, Pullman, Wash.).

When a large proportion of the seedlings in a treatment reached the two-leaf stage, a section of the flat, with seedlings intact, was removed from the chamber for 0, evolution measurement in the laboratory. Representative plants were used for 0, evolution measurements. Measurements were made between 2 and 7 h into the photoperiod, repeating measurements of each of the treatments each 60 to 75 min. The leaves of four entire petunia shoots were carefully oriented in the chamber to allow for uniform irradiation of all leaves by the xenon light source. The general procedures, described by Delieu and Walker (1983) and Walker (1985) were followed with the slight modification of using whole leaves rather than 10-cm leaf disks for 0, evolution measurement. Two 4.25-cm Whatman #1 filter paper disks, separated by a layer of cheesecloth, were cut to fit inside the LD2 chamber. This “sandwich” was then saturated with 0.5 ml of a 1 mM KHCO3 solution to supply the CO2 necessary for photosynthesis. Seedlings were held in the chamber with the light source in place for 4 to 5 min before calibration and 0, measurement. Readings were taken each minute until a steady state of 0, evolution was observed, usually requiring 8 to 12 min. The xenon light source was operated at 1000 µmol m–2 s–1 PPF during the measurement. Preliminary observations of O2 evolution revealed that evolution was fairly linear from ≈600 to 1250 µmol m–2 s–1 (r = 0.83) in ‘Red Flash’ petunia seedlings during the sampling period used (data not presented).

Results and Discussion

Plants in the 350/16 treatment reached the two-leaf stage in 25% fewer days with 30% greater DW than those receiving half the total PPF or a similar Pn period (Table 2). Data obtained under 350/16 conditions supported the results from earlier work (Carpenter and Carlson, 1974; Graper and Healy, 1987, 1989; Lang and Healy, 1985) where increased total PPF yielded larger plants in reduced time. Increasing the photoperiod (350/PE) by incandescent day extension failed to decrease the days to reach

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Peak PPF (µmol·m–2·s–1)</th>
<th>Pn period (h·day–1)</th>
<th>Photoperiod (h·day–1)</th>
<th>Total PPF (mol·m–2·day–1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>175/16</td>
<td>175</td>
<td>16</td>
<td>16</td>
<td>10</td>
</tr>
<tr>
<td>350/8</td>
<td>350</td>
<td>8</td>
<td>8</td>
<td>10</td>
</tr>
<tr>
<td>350/16</td>
<td>350</td>
<td>16</td>
<td>16</td>
<td>20</td>
</tr>
<tr>
<td>350/PE</td>
<td>350</td>
<td>8</td>
<td>16</td>
<td>10</td>
</tr>
</tbody>
</table>

Table 1. Irradiance conditions in the growth chambers: peak PPF (µmol·m–2·s–1), Pn period (h·day–1), photoperiod (h·day–1), and total PPF (mol·m–2·day–1).
the two-leaf stage, but also decreased seedling DW by an average of 55% compared to 350/16 (Table 2). Lang and Healy (1985) reported that a similar low-level incandescent PE treatment for 7 days decreased seedling DW by 25% compared to PE using high-level irradiance at $\approx 85 \mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$.

In all treatments, leaf areas ranged from 0.7 to 0.9 cm²/plant. However, O₂ evolution was 31% higher under an 8-h photoperiod at 350 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ PPF than under the 175/16 treatment and was reduced by 48% in the 350/PE treatment (Table 2). Specific leaf weight (SLW) followed trends similar to O₂ evolution, with the 350/PE treatment having the lowest value recorded (Table 2).

Chabot et al. (1979) found *Fragaria virginiana* (Duch.) SLW increased as peak or total PPF increased. Also, net photosynthesis was increased when the total PPF was increased from 10.09 to 19.98 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{day}^{-1}$, but not when peak PPF was increased from 105 to 560 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$. Our results show similar trends, with O₂ evolution and SLW increasing as total PPF increased (Table 2). However, single degree-of-freedom comparisons of O₂ evolution show a significant difference due to photoperiod and Pn period as well (Table 2). This result is due to the effect of the high rate of O₂ evolution in the 350/8 treatment, which caused these tests to be significant.

As total PPF was increased from 10 to 20 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{day}^{-1}$, Chabot et al. (1979) found a linear slope of 0.23 ($\alpha = 0.0001$, $r' = 0.96$) with SLW. Although this type of correlation cannot be made from our data due to the few treatments under 350 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$, increasing total PPF from 10 to 20 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{day}^{-1}$ increased SLW by 30% (Table 2). Increasing the peak PPF from 175 to 350 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ also resulted in a 34% increase in SLW.

Reports in the literature of Pn rates for petunia seedlings and other genera are extremely limited. However, photosynthesis measurements made with mature leaves, which may not be directly comparable to the whole-plant measurements used in this study, have been reported. In *F. virginiana*, Chabot et al. (1979) found that Pn rate was positively correlated with peak and total PPF. Other investigators (Bunce, 1983; Chatterton and Silvius, 1981) obtained higher photosynthetic rates in *G. max* by lengthening the Pn period while maintaining constant total PPF. *Helianthus annuus* L. and *Amaranthus hypochondriacus* L. responded similarly to changes in Pn period (Bunce, 1983). The expanding leaves used in our study responded more strongly to increasing peak PPF in conjunction with the Pn period (Table 2).

Shoot CHO content at the two-leaf stage, like the morphological and Pn measurements, was significantly modified by total PPF. Plants grown at 350/16, which had the highest DW–30% higher than the mean of the other treatments (Table 2), often contained significantly more fructose, glucose, sucrose, starch, and total soluble sugars, resulting in a 30% to 90% higher total CHO content than in plants of the other treatments (Table 3). Acid invertase activity in these plants was increased by 50% compared to other treatments. Growing plants under the 350/PE reduced levels of all sugars an average of 80% and reduced invertase activity an average of 100%. In general, acid invertase activity and the amount of total CHO was positively correlated with seedling DW at the two-leaf stage (Tables 2 and 3).

Chatterton and Silvius (1979, 1981) investigated the effect of growing *G. max* at either 7- or 14-h photoperiods and Pn periods using treatments corresponding to our three treatments: 350/8, 350/16, and 350/PE. *Glycine max* plants grown at the same peak PPF while doubling the Pn period from 7 to 14 h had a greatly reduced rate of starch production. Our data indicate that increasing the Pn period while holding the total PPF constant did not decrease the amount of starch but did decrease the total sugar content (Table 3). When the Pn period and total PPF were doubled, the sugar content was further increased. However, little consideration was given by Chatterton and Silvius (1979, 1981) to the fact that while doubling the Pn period, the total PPF was doubled as well.

*Hordeum vulgare* L. preferentially partitioned CHO into sugars when the total PPF was increased (Farrar and Farrar, 1987). Similar results were observed in our studies when sugar and starch content was expressed as a percentage of the total CHO produced by the plant at the two-leaf stage (Table 4). Doubling the peak PPF of a 16-h photoperiod from 175 to 350 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ increased partitioning into sugars by 25% and decreased partitioning into starch by 32%. Increasing the photoperiod with an 8-h PE treatment decreased partitioning into sugar compared to the additional 8 h of 350 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ (Table 4).

Britz et al. (1985) reported that doubling the Pn period and total PPF or doubling only the photoperiod reduced the rate of starch accumulation in *Digitaria decumbens* Stent. by 200% to 300%. However, petunias grown continually in the 350/PE treatment, which had twice the photoperiod as those in 350/8, partitioned similar proportions of total CHO into sugars and starch (Table 4).

Increased acid invertase activity is closely associated with developing or rapidly expanding leaves (Claussen et al., 1985; Morris and Arthur, 1984, 1985). Seedlings grown with 350/16, which were developing at a faster rate than plants in other treatments, had 50% higher invertase activity (Table 3). This increase may result in more rapid breakdown of sucrose into its constituent sugars, fructose and glucose, promoting rapid growth. Earlier findings, using supplemental high-pressure sodium, showed more rapid leaf unfolding in treatments with increased relative growth rate (Graper, 1990; Graper and Healy, 1991).

Total PPF was, in most cases, the dominant factor in increasing petunia seedling growth rate and the production and partitioning of CHO into sugars vs. starch in seedlings at the two-leaf stage. Increasing peak PPF increased SLW as previously reported (Chabot et al., 1979). However, the greatest increase in Pn was observed when peak PPF increased from 175 to 350 $\mu\text{mol}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$ while Pn was maintained at 8 h (Table 2). Most reports have indicated that total PPF was the dominant factor influencing Pn rate, but often these researchers failed to hold the Pn period or peak PPF constant, which could lead to erroneous conclusions (Bunce, 1983; Chabot et al., 1979; Chatterton and Silvius, 1980a, 1980b).

Increasing the photoperiod with low-level incandescent PE decreased total concentration of sugars and starch within the plant by 80% and 20%, respectively, and decreased acid invertase activity by 50% and seedling DW, SLW, and O₂ evolution at the two-leaf stage by 55%, 85%, and 65%, respectively, as compared to the other treatments. However, CHO partitioning

---

Table 2. *Petunia x hybrida* ‘Red Flash’ seedling characteristics at the two-leaf stage for the four irradiance treatments. (See Table 1 for treatment explanation.)

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Days to two-leaf stage</th>
<th>DW (mg)</th>
<th>SLW (mg·cm⁻²)</th>
<th>O₂ evolution (µmol O₂·min⁻¹·dm⁻²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>175/16</td>
<td>16.3 ab</td>
<td>2.1 a</td>
<td>2.6 b</td>
<td>6.1 b</td>
</tr>
<tr>
<td>350/8</td>
<td>18.2 a</td>
<td>2.0 ab</td>
<td>2.8 b</td>
<td>8.8 a</td>
</tr>
<tr>
<td>350/16</td>
<td>14.0 b</td>
<td>2.4 a</td>
<td>3.5 a</td>
<td>7.5 ab</td>
</tr>
<tr>
<td>350/PE</td>
<td>17.3 a</td>
<td>1.4 b</td>
<td>1.6 c</td>
<td>4.5 c</td>
</tr>
</tbody>
</table>

*Mean separation in columns by LSD, *P* = 0.05.*
between starch and sugar was consistent with the other two treatments receiving 10 mol·m⁻²·day⁻¹ PPF. Increasing the Pn period while holding the total PPF constant did not increase total CHO content or decrease total starch content. Only when total PPF was increased in conjunction with Pn period was total CHO content increased by 60% (Table 3) and seedling DW increased by 13% (Table 2).

Our research indicates that increasing the total PPF to seedlings in the greenhouse with supplemental lighting may in fact increase the seedling growth rate by increasing CHO partitioning into sugars rather than starch. Although total PPF is the dominant factor, increasing the peak PPF and Pn periods also play a critical role.

**Literature Cited**


