Short-term Temperature Change Affects the Carbon Exchange Characteristics and Growth of Four Bedding Plant Species

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ABSTRACT. Bedding plants are exposed to a wide range of environmental conditions, both during production and in the landscape. This research compared the effect of short-term temperature changes on the CO₂ exchange rates of four popular bedding plant species. Net photosynthesis (Pₚₑₙ) and dark respiration (Rₚₐ₃) of geranium (Pelargonium ×hortorum L.H. Bail.), marigold (Tagetes patula L.), pansy (Viola ×wittrockiana Gams.), and petunia (Petunia ×hybrida Hort. Vilm.-Andr.) were measured at temperatures ranging from 8 to 38 °C (for Pₚₑₙ) and 6 to 36 °C (for Rₚₐ₃). Net photosynthesis of all species was maximal at 14 to 15 °C, while Rₚₐ₃ of all four species increased exponentially with increasing temperature. Gross photosynthesis (Pₕᵣₑₓ) was estimated as the sum of Pₚₑₙ and Rₚₐ₃, and was greater for petunia than for the other three species. Gross photosynthesis was less sensitive to temperature than either Pₚₑₙ or Rₚₐ₃, suggesting that temperature effects on Pₚₑₙ were caused mainly by increased respiration at higher temperatures. Gas exchange–temperature response curves were not useful in determining the heat tolerance of these species. There were significant differences among species in the estimated Rₚₐ₃ at 0 °C and the Q₁₀ for Rₚₐ₃. Differences in the Q₁₀ for Rₚₐ₃ were related to growth rate and plant size. Large plants had a greater Q₁₀ for Rₚₐ₃, apparently because these plants had a higher ratio of maintenance to growth respiration than small plants. The Q₁₀ of the maintenance respiration coefficient was estimated from the correlation between the Q₁₀ and relative growth rate, and was found to be 2.5 to 2.6.

Temperature affects growth and physiology of plants, both during production and in the landscape. A thorough understanding of temperature effects on plant growth is needed to develop guidelines for optimal temperatures during production, and can help to predict plant performance during production and in landscapes. Since almost all dry weight accumulation of plants is the result of carbon fixation, CO₂ exchange rate measurements are particularly useful in predicting plant growth. Photosynthesis and respiration temperature-response curves of various species have been used to predict plant performance and heat tolerance (Higgins et al., 1992; Leonardos et al., 1994; Ranney and Peet, 1994; Ranney and Ruter, 1997).

Most commonly, CO₂ exchange rates of individual leaves have been measured (Higgins et al., 1992; Larigauderie and Körner, 1995; Martindale and Leegood, 1997; Ranney and Peet, 1994; Ranney and Ruter, 1997). Leaf photosynthesis measurements can provide valuable physiological information (e.g., stomatal conductance, CO₂ concentrations in the stomatal cavity) and allow for relatively easy measurements of the environmental effects (e.g., light intensity, temperature, CO₂ concentration) on gas exchange rates. However, leaf gas exchange measurements often are poorly correlated with growth or yield (Elmore, 1980; Evans, 1993). Leaf measurements ignore differences in total leaf area, spatial variability within a canopy, and respiratory CO₂ efflux from roots, stems, flowers, and fruits (van Iersel and Bugbee, 2000). Therefore, whole plant measurements can give a better indication of environmental effects on plant growth.

Zelitch (1982) has shown that there generally is a good correlation between canopy photosynthesis and dry matter production or yield. However, reports of whole-plant CO₂ exchange rate measurements to determine how floricultural crops respond to changing environmental conditions appear rare. Leonardos et al. (1994) measured whole-plant CO₂ exchange rates of Alstroemeria at different light intensities, atmospheric CO₂ concentrations, and temperatures, and Miller et al. (2001) measured the effects of light intensity and temperature on CO₂ exchange rates of two angelonia (Angelonia angustifolia Benth.) cultivars.

The objectives of this study were to determine whether species from different genera respond differently to short-term temperature changes, and whether whole-plant CO₂ exchange responses to temperature can be used as an indicator of the heat tolerance of bedding plants. Photosynthesis–temperature response curves commonly are used to compare the heat tolerance of species, but these data normally are collected at the single leaf level (e.g., Al-Khatib and Paulsen, 1999; Higgins et al., 1992; Ranney and Ruter, 1997). No studies which compare the responses of whole plant CO₂ exchange rates of different species to temperature could be identified. The four species used in this study were geranium (Pelargonium ×hortorum), marigold (Tagetes patula), and petunia (Petunia ×hybrida); marigold (Tagetes patula), and pansy (Viola ×wittrockiana). These species were selected because they are among the most popular bedding plant species in the United States and differ in heat tolerance. Pansy performs well at cool temperatures, but is particularly heat-sensitive and generally does not survive summers in a warm climate like that in the southeastern United States. The other three species are better adapted to high temperatures and generally perform well at higher temperatures than pansy. Initial analysis of the data revealed differences in the response of dark respiration (Rₚₐ₃) to temperature among species. Therefore, another objective of this study was to explain these differences.

Materials and Methods

PLANT MATERIAL. Marigold ‘Antigua orange’, geranium ‘Pinto violet’, petunia ‘Dreams red’, and pansy ‘Scarlet bronze’ seedlings were received from a commercial grower (Speedling, Inc., Blairsville, Ga.) on 17 Sept. 1999 and transplanted into 10-cm
(510-mL) square pots filled with a peat-based growing medium (MetroMix 300; Scotts Co., Marysville, Ohio). The plants were placed in a double-layer polyethylene-covered greenhouse and watered as needed. Plants were fertilized twice weekly with a water-soluble fertilizer solution containing 200 mg L⁻¹ N, made with a 20N–4.3P–16.6K fertilizer (20–10–20 Peat-lite Special, Scotts Co.). Greenhouse temperature and relative humidity averaged 25 °C and 60%, respectively.

**Measurements.** CO₂ exchange measurements were started on 18 October. Data were collected with a semicontinuous, 10-chamber, whole-plant gas exchange system (van Iersel and Bugbee, 2000). Multiple plants (15 in this case) can be measured together inside one gas exchange chamber simultaneously, thus reducing experimental error caused by plant-to-plant variability. The gas exchange system consisted of acrylic chambers (0.32 × 0.5 × 0.6 m³), eight of which were placed in two growth chambers (model E-15; Conviron, Winnipeg, Canada), and the other two chambers were placed outside of the growth chambers and were measured without any plants in them. Gas exchange of eight groups of 15 plants (two groups from each species) was measured simultaneously. Air flow through each gas exchange chamber (0.5 L s⁻¹) was measured with mass flow meters (GFM37-32; Aalborg Instruments and Controls, Monsey, N.Y.) and the difference in CO₂ concentration between the air entering and exiting the gas exchange chambers was measured with an infrared gas analyzer (IRGA) in differential mode (LI-6262; LI-COR, Lincoln, Nebr.). Whole-chamber CO₂ exchange rates (µmol s⁻¹) were calculated as the product of mass flow (mol s⁻¹) and the difference between the CO₂ concentration of the air entering and exiting the chamber (µmol mol⁻¹). Each chamber was measured for 30 s, once every 10 min. Gas exchange data were corrected for zero drift of the IRGA, by subtracting the CO₂ exchange rate of the empty chambers from that of the chambers with plants in them.

Fifteen plants from a single species were placed in a gas exchange chamber to determine carbon exchange rate—temperature response curves. Since there were eight gas exchange chambers available, two replications of the four species could be measured at one time. Since only two replications could be measured simultaneously, the experiment was repeated in time to obtain a total of four replications of each species. The first two replications were measured 32 and 33 d after transplanting and the last two replications at 39 and 40 d after transplanting.

Plants were placed inside the gas exchange chambers the evening before the start of the measurements and kept in the dark for 16 h, while the growth chamber temperature was gradually decreased from 23 to 10 °C. The growth chamber temperature then was decreased to 1 °C at the start of the photosynthesis measurements. Net photosynthesis—temperature response curves were determined during a 24-h period, during which the growth chamber temperature was increased stepwise from 1 to 36 °C in 5 °C steps of 3 h each. Due to radiative heating by the lights, actual air temperatures inside the gas exchange chambers were higher than the rest of the growth chamber and ranged from 8 to 38 °C. Each temperature was maintained for 3 h, but only data from the last hour were used for analysis. This allowed the gas exchange measurements to stabilize after the temperature was changed. Since whole plants, and the pots in which they were grown, were placed in the gas exchange chambers, the measurements represent the combined effects of shoots, roots, and microbial respiration in the growing medium. Microbial respiration is negligible compared to plant CO₂ exchange, unless small plants are grown in relatively large pots. Thus, microbial respiration likely had little effect on the measured CO₂ exchange rates.

Air temperatures inside the gas exchange chambers were measured with shielded, aspirated, type T thermocouples. Light was provided by fluorescent lamps and photosynthetic photon flux density at the top of the plant canopies was 650 µmol·m⁻²·s⁻¹. Relative humidity inside the gas exchange chambers was not controlled and ranged from ≈85% at 8 °C to 70% at 38 °C as measured with capacitance sensors (HTO-45, Rotronic, Huntington, N.Y.). Therefore, the vapor pressure deficit increased from 0.16 to 2.0 kPa with increasing temperature. Since an increase in vapor pressure deficit with increasing temperature would be expected to occur under natural conditions as well, no attempt was made to control it. The CO₂ concentrations inside the gas exchange chambers ranged from 300 to 400 µmol mol⁻¹, depending on the photosynthetic CO₂ uptake by the plants.

Dark respiration measurements were collected during the following 24-h period, during which the growth chamber temperature was decreased stepwise from 38 to 3 °C, again in 5 °C steps of 3 h each. This resulted in temperatures ranging from 36 to 6 °C inside the gas exchange chambers. Relative humidity ranged from 100% at 6 °C to 70% at 38 °C and the vapor pressure deficit increased from 0 to 2.0 kPa. CO₂ concentrations ranged from 410 to 470 µmol·mol⁻¹, depending on the respiratory CO₂ efflux from the plants. Shoot and root dry weights (DW) of the plants were determined after finishing the respiration measurements.

**Data analysis.** Net photosynthesis—temperature response curves were fitted using quadratic models (R² > 0.98). Results of the regression analysis were then used to estimate the temperature at which P net reached a maximum.

Temperature response coefficients (Q10) for dark respiration were determined by fitting an exponential curve to the data (R² > 0.97)

\[ R_{dark} = R_0 \times Q_{10}^{T/T_0} \]  \[ \text{[Eq. 1]} \]

where \( R_0 \) is the estimated \( R_{dark} \) at 0 °C (µmol s⁻¹), \( Q_{10} \) is the relative increase in \( R_{dark} \) with a 10 °C increase in temperature, and \( T \) is the temperature (°C).

A temperature response curve for gross photosynthesis (\( P_{gross} \)) was constructed by adding the estimated rates of \( P_{net} \) and \( R_{dark} \), assuming a 12-h light and 12-h dark period, and the same rate at T + 5 °C divided by the rate at T – 5 °C.

To predict the effect of temperature on plant growth rate, daily carbon gain of the plants (DCG, µmol·d⁻¹) was estimated at different temperatures (T) using the regression results for \( P_{net} \) and \( R_{dark} \), assuming a 12-h light and 12-h dark period, and the same temperature in the light and dark:

\[ DCG_T = 12 \times (P_{net,T} - R_{dark,T}) \times 3600 \]  \[ \text{[Eq. 2]} \]

A photoperiod of 12 h was used for these calculations, because it is close to the natural photoperiod that the plants were exposed to in the greenhouse.

Carbon use efficiency at different temperatures (CUEₜ, mol·mol⁻¹, the ratio of photosynthates incorporated into dry matter to carbohydrates fixed in photosynthesis) at different temperatures was estimated as (again assuming a 12 h photoperiod):

\[ \text{CUE}_T = \frac{DCG_T}{P_{gross,T} \times 12 \times 3600} \]  \[ \text{[Eq. 3]} \]

For a more detailed evaluation of CO₂ exchange differences among the species, the respiration rate at 25 °C (R₂₅) was esti-
Table 1. Dry weight and calculated optimal temperature (T_{opt}) for net photosynthesis, Q_{10} for dark respiration, estimated whole plant dark respiration (R_{dark}), specific dark respiration at 0 and 25 °C (R_{0} and R_{25}), specific respiration rates at 0 °C (R_{s}; mol·g^{-1}·s^{-1}) was calculated as the ratio between DCG and total plant dry weight:

\[ \text{RGR}_{25} = \text{DCG}_{25}/\text{DW} \]  

All CO₂ exchange rates are expressed on a per plant basis (i.e., measured whole-chamber CO₂ exchange rates divided by 15 plants per chamber).

The experiment was designed as a randomized complete block with four treatments (species) and four replications. The experimental unit was a group of 15 plants within a gas exchange chamber, and four groups of 15 plants (one group of each species) inside a single growth chamber was a replication. Data were analyzed by regression and analysis of variance. Due to differences in plant size, there were significant differences between the first and last two replications for many of the measured and calculated parameters. This was accounted for in the statistical analysis by including the block (i.e., replication) effect in the analysis of variance. Means were separated with Tukey’s multiple comparison procedure.

**Results and Discussion**

Net photosynthesis of all four species increased as temperatures were increased above 8 °C, reached a maximum between 13.4 and 15.5 °C, and decreased again at higher temperatures (Fig. 1, Table 1). This photosynthetic response to temperature is typical for many C_{3} species (Bednarz and van Iersel, 2001; Higgins et al., 1992; Leonards et al., 1994; Miller et al., 2001; van Iersel and Lindstrom, 1999) and is consistent with the model for Rubisco-limited photosynthesis by Bernacchi et al. (2001). Although the optimal temperature for P_{net} may be species-dependent (Higgins et al., 1992), there were no significant differences in optimal temperature for P_{net} among the species used in this study (Table 1). Petunia had a higher P_{net} than the other three species at most temperatures (<34 °C), but its P_{net} declined more rapidly at supraoptimal temperatures, and all species had similar P_{net} at 34 to 38 °C. Higgins et al. (1992) also reported that P_{net} of different species respond differently to short-term temperature changes, while van Iersel and Lindstrom (1999) reported differences in photosynthesis responses to temperature among three magnolia (Magnolia grandiflora L.) cultivars.

Dark respiration increased exponentially for all species, with significant differences in R_{0} and Q_{10} among the species (Table 1). Geranium and petunia had the highest R_{0}, while marigold and petunia had the highest Q_{10} for R_{dark}. The combination of a high R_{0} and Q_{10} of petunia resulted in higher R_{dark} of petunia than any of the other species at temperatures above 20 °C. Since R_{0} is the whole-plant R_{dark} at 0 °C, and thus depends on plant dry weight, specific respiration rates at 0 °C (R_{s}/DW) were calculated to correct for this. The specific respiration rate at 0 °C was more than three times as high for geranium than for marigold, with intermediate values for pansy and petunia (Table 1).

Gross photosynthesis of petunia was ≈0.03 μmol·s^{-1} higher than that of the other three species, independent of temperature (P
< 0.05), while there were no differences in P_{gros} among the other three species. The higher P_{gros} of petunia most likely was the result of the relatively large leaf area of these plants, as compared to that of the other species (data not shown). At low temperatures, P_{gros} may be limited by enzyme activity (Björkman et al., 1980), which explains the increase in P_{gros} with increasing temperatures from 8 to 22 °C. Photorespiration also increases with increasing temperature (Keys et al., 1977), and since photorespiration is not included in R_{dark} measurements, estimates of P_{gros} are not corrected for photorespiration. Increasing photorespiration with increasing temperature can explain the decrease in P_{gros} at temperatures above 22 °C. Stomatal effects may have contributed as well. The leaf-to-air vapor pressure deficit increased with increasing temperature, which may result in partial stomatal closure (Day, 2000).

Gross photosynthesis of all four species was less sensitive to temperature than either P_{net} or R_{dark}, with Q_{10} values ranging from 0.9 to 1.2, compared to 0.4 to 1.1 for P_{net} and 2.0 to 2.4 for R_{dark} (a Q_{10} value close to one indicates little response to temperature) (Table 1, Fig. 2). The low sensitivity of P_{gros} to temperature suggests that temperature effects on P_{net} were caused mainly by the increase in respiration with increasing temperature, rather than by temperature effects on the actual photosynthetic activity of the plants.

The Q_{10} for P_{gros} and P_{net} generally decreased with increasing temperature, although there was little change in the Q_{10} for P_{gros} of marigold, pansy, and petunia at temperatures above 22 °C. At low temperatures (13 to 15 °C), the Q_{10} for P_{gros} of marigold was higher than that of pansy, indicating that P_{gros} of marigold increased more rapidly than that of pansy with increasing temperature in this range. At temperatures >26 °C, geranium had a lower Q_{10} for P_{gros} than the other species, indicating that its P_{gros} declined more rapidly at high temperatures.

At high temperatures, marigold and petunia had a lower Q_{10} for P_{net} than pansy (>24 °C for marigold and >30 °C for petunia). Since there were no differences in Q_{10} for P_{gros} among marigold, pansy, and petunia at high temperatures, the higher Q_{10} for P_{net} of pansy apparently was the result of the high Q_{10} for R_{dark} of petunia and marigold. Dark respiration of these two species increased more rapidly with increasing temperature, resulting in a more rapid decrease in P_{net} of petunia and marigold than that of pansy.

Although temperature effects on the CO2 exchange rate give a good indication how these physiological processes are affected by temperature, they are not a direct indicator of plant growth, since growth is affected by both photosynthesis and respiration. Among species, petunia had the highest DCG at temperatures ≤22 °C, while geranium and pansy had the highest DCG at 36 °C (P < 0.05) (Fig. 3A). Daily carbon gain of all four species increased slightly from 8 to ≈12 °C and decreased at higher temperatures. Daily carbon gain of marigold and petunia was negative (i.e., the plants would be expected lose carbon) at temperatures >32 °C, while DCG of geranium and pansy was negative at temperatures >34 °C. Many plant species have the ability to acclimate to

![Fig. 2. The Q_{10} of A) net photosynthesis (P_{net}) and B) gross photosynthesis (P_{gros}) of four bedding plant species at different air temperatures. Data points are the mean ± 1 standard error (n = 4). Error bars not shown are within the limit of the symbol.](image)

![Fig. 3. The effect of short-term changes in air temperature on (A) the daily carbon gain (DCG), a measure of plant growth rate, and (B) carbon use efficiency (CUE), the ratio of carbon incorporated into the plant to the amount of carbon fixed in gross photosynthesis, of four bedding plant species. Data points are the mean (per plant) ± 1 standard error (n = 4). Error bars not shown are within the limit of the symbol.](image)
different environmental conditions, including temperature. Thus, it is important to realize that these DCG values predict short-term effects of temperature on growth, and do not imply that these species are unable to grow at temperatures >34 °C. Species with the ability to acclimate to changing temperatures have higher optimal temperatures for photosynthesis when grown under warmer conditions than under cool conditions (Björkman et al., 1980), while Rdark at a given temperature decreases (Hurry et al., 1996; Pearcy, 1977; Rook, 1969; Tranquillini et al., 1986). Therefore, if the plants in this study had been grown at higher temperatures, the temperature at which DCG became negative likely would have likewise been higher.

Growth of plants is a function of how many photosynthates are fixed in Pgross and the efficiency with which these photosynthates are converted into structural carbon (i.e., CUE). Carbon use efficiency did not differ among species at any temperature and differences in plant size among species, correlated between plant dry weight and Q10 for Rdark were determined. The Q10 was correlated with plant size (r = 0.88, P < 0.0001; Fig. 4D), suggesting that species differences in Q10 may have been an indirect effect of differences in plant dry weight.

To better understand species and dry weight effects on Rdark, growth and respiration-related parameters were estimated for a temperature of 25 °C (Table 1, Fig. 4). Although there were large differences in dry weight among species, estimated DCG25 did not differ, while there were significant differences in RGR25, R25, and R25/DW (Table 1). There was a positive correlation between R25 and plant dry weight, but this correlation was poor (r = 0.53, P = 0.04; Fig. 4A) and there was no correlation between dry weight and DCG25 (r = 0.00, results not shown). This indicates that large plants do not necessarily grow faster or respire more than small plants.

Dry weight was negatively correlated with both R25/DW (r = -0.89, P < 0.0001) and with RGR25 (r = -0.93, P < 0.0001) (Fig. 4B and C), indicating that the metabolic activity of small plants was higher than that of large plants (per unit dry weight). These findings are similar to those of Bednarz and van Iersel (2001), who found a positive correlation between Rdark of whole cotton

Pansy, Geranium, Marigold, and Petunia
(Gossypium hirsutum L.) plants and dry weight, but a stronger and negative correlation between Rdark/DW and dry weight. The higher relative physiological activity of smaller plants (i.e. RGR25 and R25/DW) may explain species differences in Q10 for Rnet, as well as the correlation between the Q10 for Rnet and plant dry weight. These effects apparently were caused by different effects of temperature on maintenance and growth respiration. The maintenance respiration coefficient (maintenance respiration per unit dry weight) generally is considered to be temperature-sensitive, partly because of increased protein turnover at higher temperatures (de Wit et al., 1970; Thornley and Johnson, 1990).

In contrast, the growth respiration coefficient (growth respiration per unit growth) generally is considered to be temperature-insensitive, unless temperature influences the chemical composition or metabolic pathways of the plants (Penning de Vries et al., 1974). Therefore, respiration of plants with a high ratio of maintenance to growth respiration would be expected to be more temperature sensitive than those with a low ratio, and thus to have a higher Q10 for Rdark. The decrease in RGR25 and R25/DW with increasing plant size (Fig. 4B and C) suggests that larger plants allocated a larger fraction of their total respiration to maintenance. This is consistent with the finding that large plants had a higher Q10 than small plants (Fig. 4D).

Assuming that growth respiration is insensitive to temperature and that maintenance respiration of the different species increases similarly to increasing temperatures, the temperature sensitivity of maintenance respiration can be estimated from the regression of Q10 versus RGR25 (Fig. 5). When RGR25 = 0, there is no growth and therefore no growth respiration, so the Y-intercept of the regression line (2.58) is an estimate of the Q10 for maintenance respiration. However, the calculated value for RGR25 (using [Eq. 4]) depends on the duration of the photoperiod used in the calculations of DCG25 [Eq. 2]. Thus, photoperiod duration also affects the Y-intercept of the regression line of Q10 vs. RGR25, and this effect was quantified by using different photoperiod durations for the calculation of RGR25. Estimated values of the Q10 for the maintenance coefficient were 2.20 ± 0.16 (r = −0.43), 2.62 ± 0.09 (r = −0.84), 2.58 ± 0.08 (r = −0.87), 2.61 ± 0.08 (r = −0.88), and 2.62 ± 0.08 (r = −0.88) for light periods of 6, 9, 12, 15, and 18 h, respectively (estimate ± standard error, followed by the correlation coefficient of the regression line). Thus, the effects of photoperiod duration on the estimate of the Q10 for maintenance respiration were negligible, except when a light period of 6 h was used for the calculations. Using a 6-h photoperiod, however, not only resulted in a low estimate for Q10 with a relatively high standard error, but also in a low r2 value for the regression line, indicating that this estimate was less accurate than those at longer photoperiods. Thus, an estimate of 2.5 to 2.6 for the Q10 for maintenance respiration of the four species in this study seems reasonable. Q10 estimates for the maintenance respiration of whole plants appear rare. Winzeler et al. (1976) estimated the maintenance respiration coefficient of whole unicum barley (Hordeum vulgare L.) from Rdark and dry weight measurements and found that it had a Q10 of ≈3. In contrast, Gifford (1995) reported that the maintenance respiration coefficient of wheat (Triticum aestivum L.) increased with an increase in temperature from 15 to 20 °C, but was insensitive to a further increase in temperature, while the growth respiration coefficient decreased from 15 to 20 °C and increased with a further increase in temperature. This appears to be the only report of such a complex relation of growth and maintenance respiration to temperature. Others have estimated the Q10 for maintenance respiration of specific organs, and reported a Q10 of ≈2 (Marcelis and Baan Hofman-Eijer, 1995; Szaniawski and Kielkiewicz, 1982).

These results indicate that the response of Rdark to temperature depends on the ratio of maintenance to growth respiration in plants. Since growth and maintenance respiration depend on growth rate and plant size, respectively (Hesketh et al., 1980; McCree, 1974), these factors may affect respiration responses to temperature. Since RGR is the ratio between growth and weight, it may be especially important in interpreting respiratory responses to temperature. Unless differences in growth rate, plant size, and/or RGR are taken into account, it will be difficult to make meaningful comparisons of temperature effects on respiration among experimental treatments or species.

This problem may be especially important in whole plant measurements, because responses of all different organs are integrated in the measurements. Plants with a high RGR will have a higher proportion of young tissue than plants with a low RGR. Young, expanding leaves generally have higher respiration rates (per unit leaf area or dry weight) than older leaves (Yemm, 1965), presumably at least partly because most of the growth respiration occurs in young, growing tissues. Thus, the age distribution of the leaves will affect whole-plant Rnet, as well as the ratio between growth and maintenance respiration. Since much of the response of Pnet to temperature appears to be caused by changes in Rdark, rather than Pnet, RGR is likely to affect the temperature response of Pnet as well as that of Rdark. This problem can be circumvented in leaf gas exchange measurements by measuring leaves of similar age. This may explain why leaf photosynthesis measurements have been used successfully to study differences in temperature responses among species (e.g., Al-Khatib and Paulsen, 1999; Higgins et al., 1992; Ranney and Ruter, 1997), while the current whole-plant measurements did not detect any species differences.

Conclusions

Although there were differences in carbon exchange rate among the four species used in this study, gas exchange—temperature response curves were not useful in determining the heat tolerance of different bedding plant species. Gross photosynthesis was not very sensitive to changes in temperature from 8 to 36 °C (Q10 between 0.9 and 1.2), while the respiration rate of all four species increased exponentially (Q10 of 2.0 to 2.4, dependent on species). The rapid increase in Rdark with increasing temperature resulted in a decrease in Pnet at high temperatures. All four species reached their maximal Pnet at 14 to 15 °C. There were differences in the Rdark and Q10 for Rdark among the species. However, differences in Rdark and therefore indirectly Pnet responses to temperature appeared to be caused by differences in plant dry weight and RGR, rather than true physiological differences among species. The Q10 for Rdark increased with increasing plant dry weight. This apparently occurred because large plants had a lower RGR and presumably a higher maintenance to growth respiration ratio than small plants. Since the maintenance respiration coefficient is temperature sensitive, while the growth respiration coefficient is not, plants with a high ratio of maintenance to growth respiration had a higher Q10 for Rdark. The Q10 for the maintenance respiration was estimated from the correlation between RGR and Q10 for Rdark and found to be ≈2.5 to 2.6. These results suggest that it is important to consider the ratio of maintenance to growth respiration when studying temperature effects on whole-plant respiration.
Literature Cited


