Responses of Sweet Basil to Different Daily Light Integrals in Photosynthesis, Morphology, Yield, and Nutritional Quality

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Abstract. Consumption of basil (Ocimum basilicum) has been increasing worldwide in recent years because of its unique aromatic flavor and relatively high concentration of phenolics. To achieve a stable and reliable supply of basil, more growers are turning to indoor controlled-environment production with artificial lighting due to its high environmental controllability and sustainability. However, electricity cost for lighting is a major limiting factor to the commercial application of indoor vertical farming, and little information is available on the minimum light requirement to produce uniform and high-quality sweet basil. To determine the optimal daily light integral (DLI) for sweet basil production in indoor vertical farming, this study investigated the effects of five DLIs, namely, 9.3, 11.5, 12.9, 16.5, and 17.8 mol·m\textsuperscript{-2}·d\textsuperscript{-1} on basil growth and quality. ‘Improved Genovese Compact’ sweet basil was treated with five DLIs provided by white fluorescent lamps (FLs) for 21 d after germination, and gas exchange rate, growth, yield, and nutritional quality of basil plants were measured to evaluate the effects of the different DLIs on basil growth and quality. Results indicated that basil plants grown under higher DLIs of 12.9, 16.5, or 17.8 mol·m\textsuperscript{-2}·d\textsuperscript{-1} had higher net photosynthesis, transpiration, stomatal conductance (gs), transpiration, stomatal conductance (gs), compared with those under lower DLIs of 9.3 and 11.5 mol·m\textsuperscript{-2}·d\textsuperscript{-1}. High DLIs resulted in lower chlorophyll (Chl) a/b concentration per leaf fresh weight (FW), higher Chl a/b ratios, and larger and thicker leaves of basil plants. The shoot FW under DLIs of 12.9, 16.5, and 17.8 mol·m\textsuperscript{-2}·d\textsuperscript{-1} was 54.2%, 78.6%, and 77.9%, respectively, higher than that at a DLI of 9.3 mol·m\textsuperscript{-2}·d\textsuperscript{-1}. In addition, higher DLIs led to higher soluble sugar percent and dry matter percent than lower DLIs. The amounts of total anthocyanin, phenolics, and flavonoids per plant of sweet basil were also positively correlated to DLIs, and antioxidant capacity at a DLI of 17.8 mol·m\textsuperscript{-2}·d\textsuperscript{-1} was 73% higher than that at a DLI of 9.3 mol·m\textsuperscript{-2}·d\textsuperscript{-1}. Combining the results of growth, yield, and nutritional quality of sweet basil, we suggest a DLI of 12.9 mol·m\textsuperscript{-2}·d\textsuperscript{-1} for sweet basil commercial production in indoor vertical farming to minimize the energy cost while maintaining a high yield and nutritional quality.

Sweet basil (O. basilicum) is often referred as the “king of herbs” or the “royal herb” and is widely used in cooking and medicinal practices, as well as a fragrant, ornamental plant for gardens and containers because of its unique flavor and relatively high content of phenolic compounds (Chiang et al., 2005; Kruma et al., 2008; Makri and Kintzios, 2008). The United States is both the largest producer and importer of basil in the world, with most of its production in open fields (DAFF, 2012). However, the yield and quality of essential oils and phenolics of basil grown outdoors is hard to control and its phytochemical concentration varies widely with cultivation location, season, and cultivar (Fischer et al., 2011; Hassanpouraghdam et al., 2010; Pushpangadan and George, 2012). To achieve a stable and reliable supply of basil, more growers are turning to indoor controlled-environment production, which has proven to be a suitable alternative to open field and greenhouse production (Liaros et al., 2016; Saha et al., 2016).

Indoor vertical farming, also known as “plant factory,” is a highly controlled environmental system for plant production that uses multiple-layer culture shelves with artificial lighting (Despommier, 2010; Kozai et al., 2015). In consideration of global climate change and increasing urban populations, food security is an increasingly pressing matter, especially considering limited resources such as arable land, clean water, and fuel energy (Dunwoody, 2014; Liaros et al., 2016). Indoor vertical farming emerged as an environmentally sustainable form of plant production because of its high resource-use efficiency of both land and water (Despommier, 2013; Kozai, 2013; Kozai et al., 2015; Touliatos et al., 2016). The utilization efficiency of land, water, CO\textsubscript{2}, and light energy in indoor vertical farming were 100, 40, 2, and 1.7 times of those in greenhouses, respectively (Kozai, 2007; Ohyama et al., 2003; Yokoi et al., 2005). In recent years, the number of indoor vertical farming facilities has increased rapidly in Japan, China, and other Asian countries (Kozai et al., 2015). In North America, vertical farming has been built for commercial production of leafy greens, herbs, and transplants (Kozai et al., 2015). For example, AeroFarms, an enterprise specializing on indoor farming, built its ninth farm in Newark, NJ, and is the world’s largest indoor vertical farm based on annual output (AeroFarms, 2017). As one of the most popular herbs in the United States, sweet basil is a great candidate plant for indoor vertical farming because of its high value and demand (Liaros et al., 2016).

Light is one of the most important environmental factors that affects plant development and regulates plant behavior depending on light quantity, quality, direction, and duration (Chang et al., 2008; Dou et al., 2017; Figueiredo et al., 2008; Shafee-Hajjabad et al., 2016). Daily light integral [the product of photosynthetic photon flux density (PPFD) and photoperiod] represents the total PPF radiated by a light source in 24 h and usually has a linear relationship with crop yield and nutrient accumulation (Bochenek and Fallstrom, 2016; Colonna et al., 2016; Dai et al., 2009). Basil originates in tropical and subtropical regions and is adapted to moderately high PPFD and long-day irradiation (Pushpangadan and George, 2012). However, artificial lighting accounts for ≈80% of total electricity consumption in an indoor vertical farm, which makes energy conservation one of the biggest concerns for its commercial application (Ohyama et al., 2002). DLIs of 12–17 mol·m\textsuperscript{-2}·d\textsuperscript{-1} are recommended for vegetables and herbs in vertical farming in terms of energy savings (Albright et al., 2000; Kozai et al., 2015). A few studies explored the effects of DLIs from
13.5 to 34.6 mol·m⁻²·d⁻¹ on basil growth and development (Beaman et al., 2009; Chang et al., 2008), but no study has determined the optimum DLI between 12 and 17 mol·m⁻²·d⁻¹ for sweet basil production under indoor controlled environment. Between DLIs of 17.3 and 23.0 mol·m⁻²·d⁻¹, no differences in plant height, canopy diameter, or shoot yield among ‘Genovese’, ‘Italian Large Leaf’, and ‘Nufar’ basil were observed, which were lower than the basal growth under DLIs of 28.8 and 34.6 mol·m⁻²·d⁻¹ in a growth chamber, respectively (Beaman et al., 2009). In a glasshouse condition, there was no difference in photosynthesis of ‘Genovese’ basil between DLIs of 13.5 mol·m⁻²·d⁻¹ (light shading in a glasshouse) and 24.9 mol·m⁻²·d⁻¹ (full sunlight), whereas a DLI of 5.3 mol·m⁻²·d⁻¹ (heavy shading) significantly reduced the photosynthetic rate, leaf area per plant, shoot FW per plant, and total essential oils concentration (Chang et al., 2008). The total amount of essential oil of basil ‘Bageco’ increased significantly with supplemental radiation provided by high-pressure sodium-vapor lamp compared with plants grown under sunlight (Nitz and Schnitzler, 2004). Based on these circumstances, the objective of this article was to determine the minimum DLI for sweet basil production with comparable nutritional values in indoor vertical farming.

Materials and Methods

Plant materials and culture. The experiment was conducted in a large walk-in growth room with multiple “book-shelf stands” each with four vertical layers spaced 25 cm apart at the Texas AgriLife Research and Extension Center at El Paso, TX, from 7 Mar. to 26 Apr. 2017 and repeated from 17 Apr. to 29 May. ‘Improved Genovese Compact’ sweet basil (Johnny’s Selected Seeds, Winslow, ME) was used in both experiments. For both experiments, one basil seed per cell was sown in 72 square cell trays (length 3.86 cm, height 5.72 cm, and volume 59 mL) with all-purpose commercial mix Metro-Mix 360 (SunGro Horticulture, Bellevue, WA). All trays were put under mist in a greenhouse for germination. The seedlings were moved out from mist after germination and grown in a greenhouse for 2 weeks. The seedlings were then transplanted to 4” square pots (length 9.52 cm, height 8.26 cm, and volume 574 mL) with Metro-Mix 360 when roots were visible on the outside of the plug.

root ball, and uniform plants were selected and moved to the walk-in growth room for different DLI treatments for 21 d.

Treatments. The experiment was conducted as a completely randomized design with a single factor (DLI) at five levels, 9.3, 11.5, 12.9, 16.5, and 17.8 mol·m⁻²·d⁻¹ (hereafter, DLI 9.3, DLI 11.5, DLI 12.9, DLI 16.5, and DLI 17.8, respectively), created by growing basil plants under five different PPFD of 160, 200, 224, 290, or 310 µmol·m⁻²·s⁻¹, respectively, with the same 16-h photoperiod provided by Cool White Alto Linear FLs (Philips Lighting, Somerset, NJ). All treatments were randomly arranged in the growth room and 18 uniform plants were randomly assigned for each treatment (replications). For each growing layer (treatment), mechanical mini fans (LS1225A-X; AC Infinity, City of Industry, CA), temperature sensor, and reflective aluminum sheets were installed to keep a uniform growing environment among treatments. To minimize light distribution being disproportionate within each treatment, all plants were systematically rearranged every 3 d. The PPFD in each treatment was measured at 15 cm from FLs at nine points using PS-100 spectroradiometer (Apogee Instruments, Logan, UT). All plants were subirrigated with a nutrient solution containing 1.85 g L⁻¹ (277.5 ppm N) 15N–2.2P–12.5S (Peters 15–5–15 Ca–Mg Special; The Scotts Company, Marysville, O H) according to plants’ water requirement, maintaining an electrical conductivity of 2.0 dS·m⁻¹ and a pH of 6.0 as recommended (Kiferle et al., 2011; Park et al., 2016; Sgherri et al., 2010; Walters and Currey, 2015). Plant canopy temperatures in each treatment were recorded and maintained at 24.5 °C/21.3 °C day/night. The basil plants in the first experiment were transplanted 3 d later than the plants in the second experiment and basil plants that had a higher yield in the first experiment when harvest; however, both experiments showed a similar trend, so only data from the second experiment are presented.

Growth characteristics. The growth characteristics such as plant height, two perpendicular widths of the stems’ diameters and the number of internodes were recorded on day 1 (D1) of the treatment and then weekly. Six plants per treatment were randomly selected for measurement. Height and two perpendicular widths of the first branch of basil plants were measured on D21, the end of the experiment. Leaf area per plant was measured using a leaf area meter (LI-3100; LI-COR, Lincoln, NE), and shoot and root FW per plant were recorded on D21. The shoot and root tissues were dried at 80 °C in a drying oven (Grieve, Round Lake, IL) for 3 d to determine the dry weight (DW) per plant.

Gas exchange and Chl concentration analysis. A portable gas exchange analyzer (CIRAS-3; PP Systems International, Amesbury, MA) was used to measure the gas exchange rate of basil leaves on D20. A PLC3 leaf cuvette with an LED light unit was used, and PPFD, relative air humidity, and CO₂ concentration inside the leaf chamber were kept constant at 800 µmol·m⁻²·s⁻¹, 50%, and 390 µmol·m⁻¹, respectively. The soil plant analysis development (SPAD) index of basil was recorded weekly to quantify relative Chl concentration per leaf area in basil leaves using a Chl meter SPAD-502 (Konica-Minolta cooperation, Ltd., Osaka, Japan). On D21, ≈0.2 g of basil leaves were cut into small pieces and then extracted in 80% methanol (v/v) for 3 d. The absorbance of extracts was measured at 663 and 645 nm using a spectrophotometer (Genesys 10S ultraviolet/Vis; Thermo Fisher Scientific, Madison, WI), and the concentrations of Chl a and Chl b were calculated according to Porra et al. (1989) and were used to calculate Chl a/b concentration and Chl a/b ratio.

Nutritional quality measurement. Six plants per treatment were randomly selected for measurements of soluble sugar percent (%), anthocyanin concentration, total phenolic concentration, total flavonoid concentration, and antioxidant capacity of basil leaves on D21 to evaluate the effects of DLIs on basil nutritional quality. The soluble sugar percent of fresh basil leaves was measured using a Brix refractometer (Extech Instruments, Nashua, NH). Fresh basil leaves were collected and stored in a deep freezer at −80 °C (U1786A; Thermo Fisher Scientific, Marietta, OH) until phytochemical analysis. About 2 g of fresh basil leaves were ground in liquid nitrogen and extracted with 15 mL 1% acidified methanol in darkness. After overnight extraction, the mixture was centrifuged at 13,200 rpm for 15 min and the supernatant was collected for analysis. The absorbance of extracts was measured at 530 nm using a spectrophotometer mentioned previously and the anthocyanin concentration was expressed as milligram cyanidin-3-glucoside equivalents using a molar extinction coefficient of 29,600.

The total phenolic concentration of basil leaves was determined using the modified Folin–Ciocalteu reagent method (Xu and Mou, 2016) described as follows: a 100-µL extraction sample was added to a mixture of 150-µL distilled water and 750 µL 1/10 dilution Folin–Ciocalteu reagent; after 6-min reaction, 600 µL 7.5% Na₂CO₃ was added to the mixture. The mixture was incubated at 45 °C in a water bath for 10 min before the absorbance was measured at 725 nm using a microplate reader (ELX800; BioTek, Winooski, VT). Results were expressed as milligram gallic acid equivalent per gram FW of basil leaves. For total flavonoid concentration, a 20-µL extract was mixed with 85 µL distilled water and 5 µL 5% NaNO₂. After 6 min, 10 µL 10% AlCl₃·6H₂O was added. After another 5 min, 35 µL 1 M NaOH and 20 µL distilled water was added and then the absorbance was measured at 520 nm using the microplate reader mentioned previously. The results were expressed as milligram of (+) catechin hydrate equivalent per gram FW of basil leaves. The amounts of total anthocyanin, phenolic compound, and flavonoid per plant were calculated by

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multiplying the concentration of anthocyanin, phenolic compound, and flavonoid by leaf FW per plant.

The total antioxidant capacity of basil leaves was measured using the ferrous ion chelating activity (FICA) method (Xu and Mou, 2016) described as follows: a mixture of 24-μL extracts, 1.20 mL methanol, and 16 μL 2 mM ferrous chloride were vortexed vigorously. Thirty-two microliters of 5 mM ferrozine was then added and mixed vigorously, and the absorbance of mixture was measured at 562 nm after 4-min reaction using the spectrophotometer mentioned previously. The total antioxidant capacity was calculated as the absorbance difference between control (Acontrol) and sample (Asample): total antioxidant capacity (% FICA) = 100 × (Acontrol - Asample) / Acontrol.

Statistical analysis. One-way analysis of variance was conducted to test the effects of DLI on all measured parameters. Mean comparison among treatments was conducted using Student’s t method. Correlation test was conducted using the pairwise correlations method. All statistical analyses were performed using JMP (version 13; SAS Institute Inc., Cary, NC).

**Results**

Photosynthesis and Chl concentration of basil leaves under different DLIs. The relative Chl concentration per leaf area and SPAD readings increased significantly as basil growth stage developed and DLI increased (Fig. 1A). SPAD for treatments DLI 9.3, DLI 11.5, and DLI 12.9 increased from 30 to 37 after 21 d of treatment, whereas those in the DLI 16.5 and DLI 17.8 treatments increased to ≈41, which was 11% higher (Fig. 1A). In contrast, no difference in Chl a concentration per leaf FW was observed among the five different DLIs on D21, whereas Chl b concentration was higher for treatments DLI 9.3 and DLI 11.5, and lower for treatments DLI 12.9, DLI 16.5, and DLI 17.8 (Fig. 1B). Higher levels of Chl a/b ratio (Fig. 1C) and lower levels of Chl a+b concentration (Fig. 1B) were observed for treatments DLI 12.9, DLI 16.5, and DLI 17.8. The Chl a+b concentration per leaf FW for treatments DLI 9.3 and DLI 11.5 were ≈17% higher than that of basil grown under treatments DLI 12.9, DLI 16.5, and DLI 17.8 (Fig. 1B).

The leaf net photosynthetic rate per leaf area (Pnleaf), transpiration, and gS of basil leaves increased significantly as DLI increased and were the highest for treatments DLI 12.9, DLI 16.5, and DLI 17.8 (11.5, 10.6, and 10.4 μmol·m⁻²·s⁻¹), followed by treatments DLI 9.3 and DLI 11.5 (6.1 and 7.8 μmol·m⁻²·s⁻¹), respectively (Table 1). Pnleaf for treatments DLI 12.9 was 86% and 47% higher than that for treatments DLI 9.3 and DLI 11.5, respectively, and no difference among treatments DLI 12.9, DLI 16.5, or DLI 17.8 was observed (Table 1). Transpiration for treatment DLI 12.9 was 78% and 57% higher than that for treatments DLI 9.3 and DLI 11.5, whereas gS for treatments DLI 12.9 was 126% and 83% higher than that for treatments DLI 9.3 and DLI 11.5, respectively (Table 1).

Morphological differences of basil influenced by DLIs. Basil plants grown under higher DLIs had a larger canopy because of increased height and widths (Table 2) but had
similar number of internodes (data not presented). The plant widths responded faster to DLIs than plant height, with visible difference after 1-week DLI treatment, whereas it took 2 weeks for plant height to show difference among treatments. On D21, the plant height was the greatest for treatments DLI 12.9, DLI 16.5, and DLI 17.8 (22.1, 23.3, and 23.0 cm, respectively), followed by DLI 11.5 (20.2 cm), and was the lowest for DLI 9.3 (17.4 cm). Although the plant widths showed visual differences earlier than the height, the differences among five DLI treatments were small (Table 2).

Basil plants grown under higher DLIs had larger and thicker leaves, as well as greater branch height and widths (Table 3). With the same number of leaves, the leaf area per plant for treatment DLI 17.8 was 51% and 35% higher than that for treatments DLI 9.3 and DLI 11.5, whereas its specific leaf area (leaf area per unit leaf DW) was 30% and 21% lower, respectively. Lower specific leaf area under higher DLIs indicated that the thickness of basil leaves increased as DLIs increased. In addition to plant height and widths, the branching of basil was also positively correlated with DLIs. There were two pairs of fully expanded leaves for the first branch of basil plants grown under treatments DLI 12.9, DLI 16.5, and DLI 17.8, whereas only one pair of fully expanded leaves for treatment DLI 9.3 (data not presented), which contributed to increased branch height and widths under higher DLIs (Table 3).

Plant growth and yield of basil under different DLIs. The highest shoot FW per plant was observed in treatments DLI 12.9, DLI 16.5, and DLI 17.8 (20.2, 23.4, and 23.3 g, respectively), followed by DLI 11.5 (15.7 g), whereas DLI 9.3 (13.1 g) exhibited the lowest value (Fig. 2A). The fresh leaf and stem weight had the similar trend as fresh shoot yield, whereas the root FW per plant was the highest in treatments DLI 16.5 and DLI 17.8, followed by DLI 12.9, then DLI 11.5, and was the lowest in DLI 9.3. The leaf DW per plant was more sensitive to DLIs than leaf FW, and significant difference was observed among treatments DLI 12.9, DLI 16.5, and DLI 17.8 (1.22, 1.58, and 1.55 g, respectively) (Fig. 2B). The shoot DW per plant had a similar pattern with leaf DW, whereas shoot DW per plant in DLI 17.8 was more than 2-fold than that in DLI 9.3. The shoot FW and DW per plant were both positively correlated with DLIs at the time of harvest on D21, with correlation coefficients of 0.86 and 0.88, respectively (Fig. 3A). The shoot dry matter percent of basil was also positively influenced by DLIs, ranging from 6.7% to 9.2% (Fig. 3B).

Nutritional quality of basil leaves under different DLIs. The soluble sugar percent, total phenolic concentration, and total flavonoid concentration of basil leaves increased with DLIs and were 52%, 35%, and 85% higher in treatment DLI 17.8 compared with DLI 9.3, respectively (Table 4). There was no difference for anthocyanin concentration among different DLIs, ranging from 2.60 to 2.82 mg·100 g−1 leaf FW (Table 4).
increased phenolic compound and flavonoid concentration of basil leaves led to higher antioxidant capacities with increasing DLIs, which was 73% higher in treatment DLI 17.8 than DLI 9.3 (Table 4). Because of higher leaf FW per plant under higher DLIs, the amounts of total anthocyanin, phenolic, and flavonoid per plant were positively correlated with DLIs with correlation coefficients of 0.84, 0.96, and 0.89 respectively (Fig. 4).

**Discussion**

Photosynthetic capacity, Chl concentration, leaf morphology, growth, and yield of sweet basil. As the vital factor affecting plant photosynthesis, DLI or PPFD alters leaf Chl concentration to maximize photosynthetic efficiency and productivity (Retkute et al., 2015; Wittmann et al., 2001). In this study, the Pn_leaf of sweet basil increased from 6.1 μmol·m⁻²·s⁻¹ in treatment DLI 9.3 (relatively low PPFD of 160 μmol·m⁻²·s⁻¹) to 10.4 μmol·m⁻²·s⁻¹ in treatment DLI 17.8 (relatively high PPFD of 310 μmol·m⁻²·s⁻¹) (Table 1), inferring that the light saturation point of sweet basil is higher than 310 μmol·m⁻²·s⁻¹ under this environment. Similarly, Polyakova et al. (2015) reported that the Pn_leaf of ‘Ararat’ basil grown for 30 d under 240–260 μmol·m⁻²·s⁻¹ provided by induction lamps was more than twice higher than that of plants grown under 80–85 μmol·m⁻²·s⁻¹ provided by white LEDs. One reason for the increased Pn_leaf of high-light leaves is their generally higher Chl concentration per leaf area (Lichtenthaler et al., 2007). Pn_leaf represents the sum of individual cell CO₂ assimilation, and the thinner leaves under lower DLIs contain significantly less cells per leaf area as compared with thicker leaves under higher DLIs (Table 3), which consequently resulted in lower Chl concentration per leaf area (SPAD) and Pn_leaf (Fig. 1A; Table 1). SPAD reading of plants was mainly associated with a greater amount of nitrogen per leaf area, as well as higher concentration of Rubisco enzyme, and subsequently resulted in increased photosynthesis (Lichtenthaler, 1985). Increased SPAD reading also led to darker green leaves of basil plants under higher DLIs, which play an important role for consumers making purchasing decisions (Rouphael et al., 2012). Basil plants under higher DLIs exhibited higher Pn not only on leaf area basis but also on Chl basis and leaf DW basis (Fig. 5), which could be explained by the possession of chloroplasts adapted to higher PPFD under higher DLIs. The high-light–adapted chloroplasts had higher photosynthetic quantum conversion rate with adapted ultrastructure, biochemical organization and a special arrangement of Chls, and carotenoids in the thylakoids under higher DLIs, resulting in increased Pn on Chl basis and leaf DW basis (Lichtenthaler et al., 2007).

In contrast to Chl concentration on leaf area basis, basil leaves under lower DLIs had a significantly higher Chl a+b concentration per leaf FW, and treatment DLI 9.3 was up to 16% higher than treatment DLI 17.8 (Fig. 1B). This result was consistent with the Chl a+b concentration of ‘Ararat’ basil and Glycyrrhiza uralensis grown under different DLIs (Hou et al., 2010; Polyakova et al., 2015). The increased Chl a+b concentrations of basil leaves under lower DLIs resulted from increased Chl b concentration with similar Chl a concentration and consequently lower Chl a/b ratios (Fig. 1C). The difference in Chl a/b ratios is also a useful indicator of light conditions, with lower Chl a/b ratios in shade leaves and higher Chl a/b ratios in sun leaves (Sarijeva et al., 2007). Under lower DLIs, plants maximize light-harvesting capacity by increasing light-harvesting Chl–protein complex in photosystem II, which

![Graph](image)

**Fig. 3.** Correlation between shoot fresh weight per plant, shoot dry weight per plant (A), and dry matter percent (B) with daily light integrals of ‘Improved Genovese Compact’ sweet basil grown for 21 d at different DLIs in indoor controlled environment. Correlation test was conducted using pairwise correlations method.

**Table 4.** Brix, anthocyanin concentration, total phenolic concentration (gallic acid equivalent, GAE), total flavonoid concentration (+)-catechin hydrate equivalent, CHE), and antioxidant capacity (ferrous ion chelating activity, FICA) of ‘Improved Genovese Compact’ sweet basil leaves grown for 21 d at different daily light integrals (DLIs) in indoor controlled environment.

| Treatment | Brix (%) | Anthocyanin concn (mg·100 g⁻¹ FW) | Total phenolic concn (GAE mg·g⁻¹ FW) | Total flavonoid concn (CHE mg·g⁻¹ FW) | Antioxidant capacity (% FICA) |
|-----------|---------|----------------------------------|--------------------------------------|--------------------------------------|-------------------------------|
| DLI 9.3   | 2.3 c   | 2.60 a                           | 1.02 b                               | 0.34 c                               | 1.96 b                        |
| DLI 11.5  | 2.7 bc  | 2.76 a                           | 1.07 b                               | 0.47 b                               | 3.46 ab                       |
| DLI 12.9  | 2.9 b   | 2.82 a                           | 0.99 b                               | 0.40 bc                              | 3.80 ab                       |
| DLI 16.5  | 2.5 bc  | 2.82 a                           | 1.61 a                               | 0.90 a                               | 5.26 a                        |
| DLI 17.8  | 3.5 a   | 2.73 a                           | 1.38 a                               | 0.63 a                               | 3.37 ab                       |

*Means followed by the same letters are not significantly different within a column, according to Student’s t mean comparison (P < 0.05). FW = fresh weight.
contains most of the Chl b, and consequently a higher Chl a/b ratio (Kitajima and Hogan, 2003; Sarjieva et al., 2007). The increased Chl a/b concentration per leaf FW under lower DLIIs demonstrated the plants’ ability to maximize the light-harvesting capacity under lower light conditions (Dai et al., 2009). Accordingly, Chl a/b concentration was correlated with Chl a/b ratio negatively and Pnleaf positively (Fig. 6).

Plant photosynthetic rate per leaf area depends not only on photosynthetic biochemistry but also on the mesophyll structure of leaves (Retkute et al., 2015). Because resistance to CO2 diffusion from the submatal cavity to the stroma is substantial, the mesophyll structure affects Pnleaf by affecting the diffusion of CO2 (Terashima et al., 2001) and the penetration of light in leaves (Vogelmann and Martin, 1993). The increased gs under higher DLIIs indicated that basil leaves were able to open their stomata much wider under higher DLIIs, which increased the leaf transpiration accordingly (Table 1). This certainly appears to be an essential prerequisite for increased Pnleaf under higher DLIIs (Table 1; Fig. 5).

Basil leaves developed in lower DLIIs are thinner and smaller than those growing in higher DLIIs (Table 3), which reduced the respiratory cost of basil leaves to help compensate for the greatly decreased photosynthetic capacity (Dai et al., 2009). Meanwhile, the mesophyll cells of basil leaves under higher DLIIs are more compact (associated with higher dry matter percent) compared with lower DLIIs (Fig. 3B). Under lower DLIIs, decreased Pnleaf produced insufficient ATPs with low carbon fixation and carbohydrate biosynthesis, resulting in smaller plant canopy (Table 2) and decreased shoot and root FW/DW per plant (Fig. 2). Accordingly, the shortage of photo-assimilate supplies and inadequate sucrose synthesis led to a reduction in soluble sugar percent (Table 4) compared with plants grown under higher DLIIs.

Future research perspectives. This experiment was conducted at five DLIIs created by growing basil plants under five different PPFD with the same 16-h photoperiod. As one factor of the lighting conditions, photoperiod also influences leaf expansion, crop yield, and nutritional content accumulation of plants (Beaman et al., 2009). Few studies on responses of basil plants to different photoperiods in indoor controlled environment were published because it is believed that basil is a long-day plant, and a 16-h photoperiod was used in most studies on basil controlled-environment cultivation with artificial lighting (Beaman et al., 2009; Piovene et al., 2015). However, what is the response of basil plants to DLIIs created by different photoperiods with the same PPFD? Furthermore, what is the response of basil plants to different combinations of PPFD and photoperiod at the same optimal DLI? These might be the future research perspectives.
leaves, greater leaf and shoot yield, and higher dry matter percent under DLI’s of 12.9, 16.5, and 17.8 mol·m⁻²·d⁻¹, compared with 9.3 and 11.5 mol·m⁻²·d⁻¹. Meanwhile, the soluble sugar percent and amounts of total anthocyanin, phenolic compounds, and flavonoids per plant were positively correlated with DLIs, and the antioxidant capacity of basil leaves at a DLI of 17.8 mol·m⁻²·d⁻¹ was 73% higher than 9.3 mol·m⁻²·d⁻¹. Combining results in growth, yield, and nutritional quality of sweet basil, we suggest a DLI of 12.9 mol·m⁻²·d⁻¹ for basil commercial production in indoor vertical farming to minimize the energy cost while maintaining a high yield and nutritional quality.

Fig. 5. Net photosynthetic rate per chlorophyll weight per hour (A) and net photosynthetic rate per leaf dry weight per hour (B) of ‘Improved Genovese Compact’ sweet basil grown for 21 d at different daily light integrals in indoor controlled environment. Means with the same letters are not significantly different according to Student’s t mean comparison (P < 0.05).

Fig. 6. Correlation of chlorophyll (Chl) a+b concentration with Chl a/b ratio and net photosynthetic rate per leaf area with Chl a/b ratio of ‘Improved Genovese Compact’ sweet basil grown for 21 d at different daily light integrals in indoor controlled environment. Correlation test was conducted using pairwise correlations method.

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