Growth and Bioactive Compound Synthesis in Cultivated Lettuce Subject to Light-quality Changes

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Abstract. This study aimed to determine the effect of changes in light quality on the improvement of growth and bioactive compound synthesis in red-leaf lettuce (Lactuca sativa L. ‘Sunmang’) grown in a plant factory with electrical lighting. Lettuce seedlings were subjected to 12 light treatments combining five lighting sources: red (R; 655 nm), blue (B; 456 nm), and different ratios of red and blue light combined with three light-emitting diodes [LEDs (R9B1, R8B2, and R6B4)]. Treatments were divided into control (continuous irradiation of each light source for 4 weeks), monochromatic (changing from R to B at 1, 2, or 3 weeks after the onset of the experiments), and combined (changing from R9B1 to R8B2 or R6B4 at 2 or 3 weeks after the onset of the experiments). Growth and photosynthetic rates of lettuce increased with increasing ratios of red light, whereas chlorophyll and antioxidant phenolic content decreased with increasing ratios of red light. Individual phenolic compounds, including chlorogenic, caffeic, chicoric, and ferulic acids, and kaempferol, showed a similar trend to that of total phenolics. Moreover, transcript levels of phenylalanine ammonia-lyase (PAL) and chalcone synthase (CHS) genes were rapidly upregulated by changing light quality from red to blue. Although the concentration of bioactive compounds in lettuce leaves enhanced with blue light, their contents per lettuce plant were more directly affected by red light, suggesting that biomass as well as bioactive compounds’ accumulation should be considered to enhance phytochemical production. In addition, results suggested that growth and antioxidant phenolic compound synthesis were more sensitive to monochromatic light than to combined light variations. In conclusion, the adjustment of light quality at a specific growth stage should be considered as a strategic tool for improving crop yield, nutritional quality, or both in a plant factory with electrical lighting.

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Closed-type plant production systems incur additional costs when using electrical light sources instead of sunlight, which requires an increase in crop yield and quality to obtain economic feasibility for crop cultivation (Piovene et al., 2015). Therefore, a systematic approach is necessary for developing technology that controls light conditions as these influence crop yield and quality factors. Light-emitting diodes, commonly used as electrical light sources for closed-type plant production systems, can generate optimal light conditions for improving crop yield because of their reduced energy consumption and control of light quality, intensity, and period (Massa et al., 2008). The production of phytochemicals, which are considered major indicators of crop quality, can be enhanced through LED irradiation (Bian et al., 2014; Lefsrud et al., 2008; Son et al., 2012). However, most closed-type plant production systems have only been performing a passive light control, such as fixed light quality and intensity, rather than changing light quality using LEDs.

Recently, Bian et al. (2014) reported that the accumulation of phytochemicals in vegetable crops, such as lettuce, cucumber, tomato, radish, and spinach, depended on light quality and delineated the effects of phytochemicals under different light quality and intensity. Light quality, in particular, had a major impact on phytochemicals: carotenoids (β-carotene, lutein), phenolics (anthocyanins, flavonoids), and vitamin C respond differently to LED quality in terms of ultraviolet-A (315–380 nm), blue (425–490 nm), green (490–550 nm), red (625–700 nm), and far red (700–740 nm) radiation in green vegetables, tomato, cucumber, and sweet pepper (Olle and Viršilė, 2013). Ultraviolet, which has high energy and short wavelength, is known to promote biosynthesis of phytochemicals by inducing plants’ defense mechanisms as this radiation usually stresses the plant (Gartia et al., 2003). Among the visible spectrums, red and blue light are important for photosynthesis and have often been used in plant research and commercial production. According to our previous studies, red and blue LEDs effectively enhanced lettuce (L. sativa L.) plant growth and the synthesis of secondary metabolites, respectively, although the response to light quality depended on lettuce variety (cultivar) (Son et al., 2012; Son and Oh, 2013, 2015).

In general, plants allocate or distribute their resources between growth and development, which was described in the so-called carbon-nutrient balance model (Bryant et al., 1983) and growth-differentiation balance framework (Hermes and Mattson, 1992). This resource partitioning might affect the synthesis of secondary metabolites differently during all growth stages. Therefore, determining a reasonable harvest time and controlling the growth environment are necessary to obtain mature plants with maximal concentrations of secondary metabolites. In this aspect, the narrow-handwidth light provided by LEDs may affect the marketable value of crops, and the irradiation protocol might be a strategy in crops’ production technology. Carvalho and Folkta (2014) suggested that light change during sprouting of kale seeds affected growth and development and could be a method to produce value-added crops. Similarly, light shift using blue and red LEDs in the production of lettuce and basil was effective in promoting plant growth and phenolic content (Jishi et al., 2016; Taulavuori et al., 2016). Based on these premises, the present study aims to determine the effect of changing light quality on growth and secondary metabolites of lettuce at a particular stage, providing basic information for improving the content of secondary metabolites. Changes in growth rate and secondary metabolite contents of lettuces were monitored to test if the content of bioactive compounds per lettuce plant would be higher in plants irradiated with a light quality that improved growth rate (mainly red light) followed by a light quality that improved secondary metabolites production (mainly blue light) than it would be in plants continuously irradiated by red or blue light.

Materials and Methods

Plant material and growing conditions
Red-leaf lettuce seeds (‘Sunmang’; Nongwoo Bio, Suwon, Korea) were sown in a plug tray (32 mL/cell) with horticultural growing medium (Myung-Moon; Dongbu Hannong, Seoul, Korea). The plug tray was placed on a shelf and the following growth conditions were maintained for 18 d: air temperature, 20 °C; relative humidity (RH), 60%; CO₂ concentration, 1000 μmol·mol⁻¹; photosynthetic photon flux density (PPFD) generated by fluorescent lamps, 119 ± 5 μmol·m⁻²·s⁻¹;
and light period, 12 h. Twenty-five seedlings were allocated to each light-quality treatment (one seedling per pot; pots 7 cm L × 7 cm W × 7.4 cm H) and transferred to other shelves in the same growing room. Lettuce plants were subirrigated with distilled water every 2–3 d and with a nutrient solution (17.3 N: 4.0 P: 8.0 K, pH 5.5, electrical conductivity 1.16 dS·m⁻¹) once a week for 4 weeks after transplanting. All lettuce plants were grown in a plant factory (4 m L × 2 m W × 3 m H) under controlled environmental conditions (air temperature, 20 °C; RH, 60%; CO₂ concentration, 1000 mol·mol⁻¹; and light period, 12 h) for 4 weeks after transplanting. Idetical LED irradiation conditions (PPFD, 151 ± 4 μmol·m⁻²·s⁻¹) and light period (12 h) were applied to each treatment. Lettuce plants were systematically rearranged every day to avoid a disproportionate distribution of irradiation.

Light-quality treatment

Five plate-type (48 cm L × 48 cm W) monochromatic (blue and red) and combined (red/blue ratios based on the number of the LED chips) R9B1, R8B2, and R6B4) LEDs were used as lighting sources in this study. The wavelengths of red (R, Bright LED Electronics, Seoul, Korea) and blue (B, ITSwell, Incheon, Korea) LEDs were 655 and 456 nm, respectively. The spectral characteristics of each lighting source were measured and adjusted at nine points (one in the center and eight on the edges of the tray) using an LI-1800 spectroradiometer and an LI-1900 portable photosynthesis system (Li-COR Inc., Lincoln, NE) for 3–4 weeks after the onset of the treatments. The leaf area of the lettuce plants grown under combined-type LED irradiation changed from red to blue at 1 (M1), 2 (M2), or 3 (M3) weeks after the onset of the experiment. The combined-type LED irradiation changed from R9B1 to R8B2 or R6B4 at 2 (C1 and C3) or 3 (C2 and C4) weeks after the onset of the treatment. All treatments were applied to lettuce plants within the same area of the plant factory and at the same time.

Growth characteristics

Biomass. Growth characteristics, such as the fresh and dry weight (DW) of shoots and roots, leaf number, and leaf area, were measured at 1-week intervals after the onset of the treatments. The fresh weight of shoots and roots was determined using a Si-234 electronic scale (Denver Instruments, Bohemia, NY). To obtain shoot and root DW, these were placed for 3 d at 70 °C in a VS-120203 oven (Vision Scientific, Daejeon, Korea) and then weighed. Leaf area was measured using a leaf area meter (LI-3000A; Li-COR Inc., Lincoln, NE).

Chlorophyll content and photosynthetic rate. Chlorophyll content was measured every week as the SPAD value determined by a portable chlorophyll meter (SPAD-502; KONICA MINOLTA, Osaka, Japan). The net photosynthesis of the lettuces grown under 12 treatments was measured using the LI-6400 portable photosynthesis system (Li-COR Inc., Lincoln, NE) for 3–4 weeks after the onset of the treatments. To precisely determine the effect of light quality, a 6400-08 clear chamber (Li-COR Inc., Lincoln, NE) was used as this chamber can transmit LED light into the leaf samples placed within it. The conditions of flow rate, CO₂ levels, PPFD, and leaf temperature within each leaf sample cuvette were maintained at 350 μmol·s⁻¹, 1000 μmol·mol⁻¹, 150 ± 5 μmol·m⁻²·s⁻¹, and 20 °C, respectively. SPAD values and photosynthetic rates were determined using the third leaf from the top of each plant.

Projected leaf area (PLA). To determine the leaf area index of the lettuces grown under each treatment, PLAs were obtained in the image analysis software LIA32 (K. Yamamoto, Nagoya University, Nagoya, Japan).

Table 1. The 12 light treatments applied, based on monochromatic (M) red (R) and blue (B) LEDs, or on combined (C) LEDs, and the fraction of blue and red light within each treatment.

| Treatment | 0 | 1 | 2 | 3 | 4 | Red | Blue |
|-----------|---|---|---|---|---|-----|------|
| R         | R | R | R | R | R | 100 | 0    |
| B         | B | R | B | B | B | 0   | 100  |
| M1        | R | B | B | B | B | 23  | 77   |
| M2        | R | R | B | B | B | 47  | 53   |
| M3        | R | R | R | R | B | 73  | 27   |
| R9B1      | R9B1 | R9B1 | R9B1 | R9B1 | R9B1 | 87  | 13   |
| R8B2      | R8B2 | R8B2 | R8B2 | R8B2 | R8B2 | 76  | 24   |
| R6B4      | R6B4 | R6B4 | R6B4 | R6B4 | R6B4 | 52  | 48   |
| C1        | R9B1 | R9B1 | R9B1 | R9B1 | R9B1 | 81  | 19   |
| C2        | R9B1 | R9B1 | R9B1 | R9B1 | R9B1 | 73  | 27   |
| C3        | R9B1 | R9B1 | R9B1 | R9B1 | R9B1 | 55  | 45   |
| C4        | R9B1 | R9B1 | R9B1 | R9B1 | R9B1 | 55  | 45   |

LED = light-emitting diode.

* Fractions of integrated blue and red wavelengths in terms of photosynthetic photon flux density at each treatment during all stages.

Table 2. Primers and cycle conditions used in the quantitative real-time PCR and their target gene.

| Gene⁷ | Accession no. | Primer sequence | Length (mer) | Cycle conditions |
|-------|---------------|-----------------|--------------|-----------------|
| LsPAL | AF299330.1     | F: CAAGGGAAAGCAGGAGATTTAC R: CTGGAAACGTCGATCAATGG | 20/20 | Tm° (°C) | No. of cycles |
| LsCHS | AB525909.1     | F: CTCACTAAGCTCTGGAGGCCT R: TTGTCCAACGAGGGAATCAA | 20/20 | 55 | 40 |
| Lsactin | AY260165.1   | F: AGCAACTGGGATGACATGGA R: GGGTGGAGGAGGTGCTCAGT | 20/20 | 52 | 40 |

⁷ LsPAL (Lactuca sativa) phenylalanine ammonia-lyase mRNA, LsCHS (Lactuca sativa) chalcone synthase, and Lsactin (Lactuca sativa) actin mRNA.

³ Annealing temperature.

HortScience Vol. 52(4) April 2017 585
Japan) for 4 weeks after the onset of LED treatments. The PLAs obtained in each treatment were used to calculate the planting density and the total phenolic content per unit area (square meters).

Secondary metabolites

Lyophilized shoots were ground using a blender (MFM-002H; Hibell, Hwaseong, Korea) and stored at 4 °C until they were used to determine the amount of total phenolics, antioxidants, and individual phenolics.

Total phenolic compounds and antioxidant capacity. Phenolic compound concentrations were determined using the Folin–Ciocalteu reagent method (Ainsworth and Gillespie, 2007), which are the key genes for the biosynthesis of phenolic compounds and flavonoids, respectively, were determined. Total RNA was isolated using the RNeasy Plant Mini Kit (QIAGEN, Dusseldorf, Germany), and its concentration in each sample was determined using a DS-11 NanoDrop spectrophotometer (DeNovix, Wilmington, DE). Complementary DNA was synthesized from the RNA isolated from each sample using the QuantiTect Reverse Transcription Kit (QIAGEN). Quantitative real-time polymerase chain reaction (PCR) was performed in a Rotor-gene 6000 (Corbett Research, Mortlake, Australia) using 2x QuantiMix SYBR Kit (PhileKorea, Seoul, Korea) with minor modifications. A portion of about 40 mg of each ground sample was extracted using 4 mL of 80% acetone for 15 min with ultrasonication (Sk5210HP; Hangzhou Nade Scientific Instrument, Zhejiang, China). Each sample extract was then centrifuged at 13,000 g, for 5 min at room temperature, and the supernatant was used in the subsequent analyses. Total phenolic concentrations and antioxidant capacity were determined as described in Son and Oh (2013, 2015) and expressed as the gallic acid–equivalent (GAE; mg) and the trolox-equivalent antioxidant capacity (TEAC; μmol) per shoot DW (mg GAE·g–1 shoot DW and μmol TEAC·g–1 DW), respectively.

Individual phenolic compounds. About 100 mg of each ground sample was mixed with 1 mL of acidified acetonitrile (0.5% v/v HCl). This mixture was hydrolyzed in a water bath at 80 °C and sonicated for 30 min (Sk5210HP ultrasonicator; Hangzhou Nade Scientific Instrument). After centrifuging at 3000 g, for 20 min and filtering through a 0.22-μm syringe filter (Noble Bio, Hwaseong, Korea), each sample’s supernatant was used in the analysis of polyphenol compounds. These were characterized using a YL9100 high-performance liquid chromatography system (HPLC; Younglin, Anyang, Korea) and separated on an ACE AQ column (4.6 mm × 250 mm, 5 μm; Advanced Chromatography Technologies, Aberdeen, UK) equipped with a guard column. The column temperature and injection volume were set to 30 °C and 10 μL, respectively, and 100% acetonitrile and 0.5% acetic acid in water were used as solvents A and B, respectively. The elution gradient was 0% to 10% A for 10 min, 10% to 20% A for 20 min, 20% to 30% A for 10 min, 30% to 40% A for 10 min, 40% to 80% A for 10 min, 80% to 0% A for 1 min, and 0% to 0% A for 9 min. The flow rate was 0.8 mL·min–1 and absorbance was recorded at 320 nm. Calibration curves, expressed as milligrams of analyte per gram of DW (mg·g–1 DW), were generated using standard chlorogenic, caffeic, chicoric, and ferulic acids, and kaempferol (all from Sigma-Aldrich, St. Louis, MO).

Gene expression. The youngest leaves of lettuce exposed to different treatments were collected every week and frozen with liquid nitrogen. Differences in the transcript levels of phenylalanine ammonia-lyase (PAL) and chalcone synthase (CHS), which are the key genes for the biosynthesis of phenolic compounds and flavonoids, respectively, were determined. Total RNA was isolated using the RNEasy Plant Mini Kit (Qiagen, Dusseldorf, Germany), and its concentration in each sample was determined using a DS-11 NanoDrop spectrophotometer (DeNovix, Wilmington, DE). Complementary DNA was synthesized from the RNA isolated from each sample using the QuantiTect Reverse Transcription Kit (Qiagen). Quantitative real-time polymerase chain reaction (PCR) was performed in a Rotor-gene 6000 (Corbett Research, Mortlake, Australia) using 2x QuantiMix SYBR Kit (PhileKorea, Seoul, Korea)

Fig. 1. Growth of shoots (A and B), roots (C and D), and leaves (E and F) of the lettuce plants grown under several light treatments using monochromatic (M) light-emitting diodes (LEDs) (red, R; blue, B) at different stages. R and B indicate the continuous irradiation of each of these lights for 4 weeks, and M1, M2, and M3 indicate changing from red to blue light at 1, 2, or 3 weeks after the onset of the treatments, respectively. The data are means ± se (n = 4). Different small caps indicate significant differences at P = 0.05 (*), P = 0.01 (**), and P = 0.001 (***).

Fig. 2. SPAD (A) and net photosynthesis (B) of the lettuce plants grown under the monochromatic (M) light treatments (red, R; blue, B) at different stages. R and B indicate the continuous irradiation of each of these lights during 4 weeks of the treatment, and M1, M2, and M3 indicate changes from red to blue light at 1, 2, or 3 weeks after the onset of the treatments, respectively. The data are means ± se (n = 4). Different small caps indicate significant differences at P = 0.05 (*) and P = 0.001 (***).
of the data presented here is supported by expression (three times). Reproducibility the measurements of growth and secondary metabolite parameters showed no significant difference irrespective of B. In all treatments, a rapid decline in lettuce growth was observed immediately after changing from red to blue light. During all growth stages, gradual significant differences were observed as blue irradiation increased.

**Chlorophyll content and photosynthetic rate.** Changing from red to blue light resulted in significant differences in lettuce’s chlorophyll content and photosynthetic rate (Fig. 2). After 4 weeks of treatment, SPAD, an indirect index of chlorophyll content, was the lowest in the R treatment (Fig. 2A). Treatments using blue light (B, M1, M2, and M3) showed a significantly higher SPAD value compared with R. A rapid increase in SPAD was observed in the week after the change from red to blue light (M1, M2, and M3). After 3 weeks of treatment, the photosynthetic rate was significantly higher (about 1.2 times) in plants exposed to red light (R and M3) than it was in other treatments (Fig. 2B). At the fourth week, R continued to show the highest photosynthetic rate among treatments, but the value in M3 decreased to a level similar to that of other treatments as it implied changing from red to blue light.

**Total phenolic compounds and antioxidant capacity.** Changing light quality from red to blue had a significant effect on total phenolic concentration and antioxidant capacity of lettuce leaves (Fig. 3). Total phenolic concentra-

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**Results**

**Growth characteristics.** Lettuces irradiated with red light and then with blue light showed significant differences in growth characteristics (Fig. 1). All growth characteristics showed the highest values after irradiation with red light for 4 weeks and a pronounced growth inhibition after irradiation with blue light. Lettuce growth after 4 weeks of treatment decreased as the time of irradiation with blue light increased (R > M3 > M2 > M1 > B), although M1 and M2 showed no significant difference irrespective of B. In all treatments, a rapid decline in lettuce growth was observed immediately after changing from red to blue light.

**Monochromatic LEDs**

**Total phenolic compounds.** Changing light quality from red to blue affected the concentration of the phenolic compounds and antioxidant capacity more sensitively in lettuce than that of contents. Individual phenolic compounds. Changing light quality from red to blue induced the highest contents of caffeic and chicoric acids, whereas M3 induced the highest

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**Table 3.** Concentrations and contents of individual phenolic compounds in the lettuce plants grown under five light treatments using monochromatic (M) LEDs (red, R; blue, B). R and B indicate the continuous irradiation of each of these lights for 4 weeks, and M1, M2, and M3 indicate changing from red to blue light at 1, 2, or 3 weeks after the onset of the treatments, respectively (n = 4).

| Treatment | Chlorogenic acid (mg g−1 shoot DW) | Caffeic acid (mg g−1 shoot DW) | Chicoric acid (mg g−1 shoot DW) | Ferulic acid (mg g−1 shoot DW) | Kaempferol (mg g−1 shoot DW) |
|-----------|----------------------------------|-------------------------------|--------------------------------|-------------------------------|-------------------------------|
| R         | 3.79 ± 0.32                      | 0.32                          | 1.40 ± 0.32                    | 0.03 ± 0.03                   | 6.38 ± 0.53                   |
| B         | 6.65 ± 0.24                      | 0.24                          | 1.08 ± 0.26                    | 0.03 ± 0.03                   | 4.27 ± 0.15                   |
| M1        | 9.42 ± 0.27                      | 0.27                          | 1.02 ± 0.28                    | 0.06 ± 0.03                   | 6.27 ± 0.18                   |
| M2        | 9.28 ± 0.28                      | 0.28                          | 1.02 ± 0.32                    | 0.01 ± 0.03                   | 6.81 ± 0.21                   |
| M3        | 7.06 ± 0.31                      | 0.31                          | 1.67 ± 0.33                    | 0.11 ± 0.03                   | 7.81 ± 0.34                   |

**Significance**

- **NS** = nonsignificant, P = 0.05 (*), P = 0.01 (**), and P = 0.001 (**).
contents of chlorogenic acid and the significantly highest ferulic acid values.

Gene expression. To assess the effect of changing light quality on the biosynthetic pathways for secondary metabolites, \( P_{\text{AL}} \) and \( \text{CHS} \) gene expression was measured by the quantitative real-time PCR in lettuce leaves at each stage of light quality change from red to blue (Fig. 4). Changing from red to blue light increased the transcript levels of \( P_{\text{AL}} \) and \( \text{CHS} \) at each stage, with the highest upregulations of these genes being observed at M1 (8.1 and 4.7 times, respectively).

Combined LEDs

Growth characteristics. Changing light quality in combined LEDs affected growth performance (Fig. 5). Among the control groups, i.e., plants continuously irradiated with R9B1, R8B2, and R6B4, the treatment comprising the highest percentage of blue light (R6B4) showed the lowest values in most growth characteristics, including shoot fresh weight, whereas R9B1 generally showed the highest values. Among the C treatments, C3 (from R9B1 to R6B4 after 2 weeks) showed the lowest values in all growth characteristics except for leaf number. Although growth characteristics under C1, C2, and C4 treatments were slightly distinct, a significant difference was not observed between these treatments. On the other hand, leaf number was similar to all treatments except R6B4.

Chlorophyll content and photosynthetic rate. Changing light quality in combined LEDs affected SPAD and photosynthetic rate during several growth stages of lettuces (Fig. 6). At 3 and 4 weeks of treatment, SPAD was not significantly different among treatments; however, the low SPAD value seen in R9B1 significantly increased after irradiation with R8B2 (C1) and R6B4 (C3) LEDs, which contained a larger proportion of blue (Fig. 6A). After 3 weeks, net photosynthesis was significantly higher under R9B1, C2, and C4 than it was in other treatments. All treatments showed lower net photosynthesis than R9B1 after 4 weeks of LED irradiation, with R6B4 presenting the lowest value (Fig. 6B).

Total phenolic compounds and antioxidant capacity. The effect of changing light quality in combined LED treatments on the total phenolic concentration and antioxidant capacity is shown in Fig. 7. Total phenolic concentration showed the lowest level in the R9B1 treatment, which contained the smallest fraction of blue light. On the other hand, changing light quality from R9B1 to R8B2 or R6B4 after 2 or 3 weeks of treatment resulted in the rapid increase of the total phenolic concentration, with the increasing rate being higher in R6B4 (41%) than in R8B2 (24%). However, when comparing the phenolic contents per shoot (plant), all treatments except R6B4 showed no significantly different alteration after 4 weeks (Fig. 7C). Antioxidant capacity exhibited a similar trend (Fig. 7B and D).

**Individual phenolic compounds.** Changing light quality from R9B1 to R8B2 or R6B4 affected lettuce’s concentration of chlorogenic, caffeic, chicoric, and ferulic acids, and kaempferol (Table 4). Chlorogenic acid concentration was not significantly different among treatments, although higher values were obtained with larger proportions of blue-light irradiation (R6B4 > R8B2 > R9B1). In addition, regardless of the time at which light-quality change was performed, increasing blue light irradiation (changing from R9B1 to R8B2 or R6B4; C1–C4) led to an increase in individual...
phenolic compounds. For chicoric acid, all C treatments showed a significantly higher value than R9B1 (>1.2 times). However, when considering the phenolic contents per unit of shoot dry weight (DW) (A and B) and per plant (C and D) obtained for the lettuce plants grown under several combined (C) light treatments at different stages. R9B1, R8B2, and R6B4 indicate the continuous irradiation of these combined lights for 4 weeks. C1, C2, C3, and C4 indicate changing from R9B1 to R8B2 or R6B4 at 2 or 3 weeks after the onset of the treatments, respectively. The data are means ± se (n = 4). Different small caps indicate significant differences at P = 0.01 (***) and P = 0.001 (****).

Discussion

Growth characteristics. Previous studies have shown that monochromatic red light was efficient for the growth of shoots and roots (Son et al., 2012) and that increased ratios of red in combined red and blue irradiation also enhanced lettuce growth (Son and Oh, 2013). The present results support these conclusions, although no significant difference was registered among M1, M2, and B treatments in lettuce irradiated with monochromatic light (Fig. 1A). M1, M2, and B contained 23%, 47%, and 0% of integrated red light, respectively, and these results are, therefore, consistent with nonsignificant differences in lettuce growth found in our previous study for treatments with <50% of red light irradiation (Son and Oh, 2013).

The nonsignificant differences in shoot growth among all treatments except R6B4 and C3 (Fig. 4A), where the fraction of integrated red light was >76%, also corroborated the results of Son and Oh (2013), where no significant differences were found between treatments with >70% of red light irradiation (87R/13B and 74R/26B). These results suggest that the fraction of red light integrated during growth stages influences growth, with differences in growth performances depending on the stage at which red light is irradiated (Table 1).

Chlorophyll content and photosynthetic rate. An increasing fraction of blue light effectively increased SPAD, the index of chlorophyll content. This is in agreement with the Banaś et al.'s (2012) report on blue light leading to the formation of chlorophyll, with the effective increase in chlorophyll formation resulting from monochromatic blue LED irradiation, and with the increase in SPAD resulting from increased ratios of blue light in combined LEDs (Son et al., 2012; Son and Oh, 2013). However, SPAD values after 4 weeks of irradiation were similar for all monochromatic treatments except B (Fig. 2A) and showed no significant differences among the combined treatments including blue light (Fig. 6A). These results imply that the qualitative effects of blue light are larger than its quantitative effects and that blue light is important for chlorophyll biosynthesis (Hogewoning et al., 2010).

Red light, on the other hand, had an effect on the photosynthetic rates similar to that registered for growth, suggesting a correlation between these two parameters (Kumagai et al., 2009). However, Kim et al. (2004) and Son and Oh (2015) found no correlation between the photosynthetic rate and growth values of lettuce leaves irradiated with an electrical lighting source installed in a leaf cuvette. This discrepancy might be explained by the setup used in the present study to measure photosynthesis: lettuce leaves were maintained under each lighting conditions for 4 weeks after transplanting, and photosynthesis was measured in a transparent leaf chamber (model 6400-08; LI-COR). Thus, studies evaluating light quality and photosynthesis under similar light conditions are necessary to obtain accurate data on photosynthetic rates in the plants’ growth environment.

Secondary metabolites. In contrast to the growth results, the fraction of blue light irradiated was effective on the accumulation of secondary metabolites. Son et al. (2012) reported an enhancement in phenolic concentration and antioxidant capacity resulting from the PAL gateway enzyme activation in the biosynthesis of phenolics induced by monochromatic blue LEDs. In addition, the increase in phenolic concentration and antioxidant capacity accompanying the increase in the fraction of blue light in combined LEDs suggested that blue light within the visible wavelength is important for the production of secondary metabolites in lettuce (Son and Oh, 2013). The results obtained here for individual phenolic compounds were consistent with those of Taulavuo et al. (2016), who observed that enhancing blue light during light periods increased the concentrations of many bioactive compounds in lettuce, including chicoric acid. This effect has been reported in several lettuce cultivars (Li and Kubota, 2009; Ouzounis et al., 2015). Thus, blue light has been referred to
in Table 4. Concentrations and contents of individual phenolic compounds in the lettuce plants grown under seven light treatments using combined (C) LEDs. R9B1, R8B2, and R6B4 indicate the continuous irradiation of each of these lights for 4 weeks. C1, C2, C3, and C4 indicate changing from R9B1 to R8B2 or R6B4 at 2 or 3 weeks after the onset of treatments, respectively (n = 4).

| Treatment | Concentration (mg g−1 shoot DW−1) | Content (mg/shoot) |
|-----------|---------------------------------|-------------------|
|           | Chlorogenic acid | Caffeic acid | Chicoric acid | Ferulic acid | Kaempferol | Chlorogenic acid | Caffeic acid | Chicoric acid | Ferulic acid | Kaempferol |
| R9B1      | 7.59               | 0.58         | 1.74           | 0.09         | 0.03       | 12.61          | 0.95          | 0.15         | 0.05         |           |
| R8B2      | 8.71               | 0.47         | 1.94           | 0.11         | 0.03       | 10.84          | 0.58          | 0.14         | 0.04         |           |
| R6B4      | 9.35               | 0.45         | 2.00           | 0.10         | 0.03       | 7.22           | 0.35          | 0.14         | 0.03         |           |
| C1        | 9.24               | 0.49         | 1.97           | 0.10         | 0.04       | 13.04          | 0.57          | 0.14         | 0.05         |           |
| C2        | 9.35               | 0.47         | 2.25           | 0.11         | 0.03       | 11.59          | 0.58          | 0.14         | 0.03         |           |
| C3        | 8.52               | 0.42         | 2.03           | 0.07         | 0.03       | 8.90           | 0.44          | 0.07         | 0.04         |           |
| C4        | 8.44               | 0.44         | 2.11           | 0.08         | 0.03       | 10.66          | 0.55          | 0.10         | 0.04         |           |

Significance: NS = nonsignificant, NS* = different lowercase letters within the columns indicate significant differences according to Duncan’s multiple range test. **P = 0.01 (***), and ***P = 0.001 (***)

Generally, it is well known that changing the light quality may be associated with metabolic changes in growth and secondary metabolites in optimum plant densities within a limited area. Hence, the light-quality changes performed in this study were effective to maximize the production of secondary metabolites in optimum plant densities within a limited area.

Conclusions. The growth rate and biosynthesis of secondary metabolites observed in lettuce at different stages clearly responded to changes in light quality, with red light being the most effective on growth and blue light being the most effective on the

Fig. 8. Lettuce plants (A), projected leaf area (B), planting density (C), and total phenolic content (D) of the lettuce plants grown under several treatments using monochromatic and combined lights. R and B indicate the continuous irradiation of red and blue light, respectively, for 4 weeks. M1, M2, and M3 indicate changes from red to blue light at 1, 2, or 3 weeks after the onset of treatments, respectively. R9B1, R8B2, and R6B4 indicate the continuous irradiation of combined lights for 4 weeks. C1, C2, C3, and C4 indicate changing from R9B1 to R8B2 or R6B4 at 2 or 3 weeks after the onset of treatments, respectively. The data are means ± SE (n = 4). Different small caps indicate significant differences at P = 0.01 (**) and P = 0.001 (***)
biosynthesis of secondary metabolites. As this study was conducted using monochromatic and combined LEDs under the same environmental conditions and during the same period, it enabled comparisons between these two light sources, which were not possible in our previous studies (Son et al., 2012; Son and Oh, 2013). Thus, it was possible to verify that combined light induced larger rates of growth and accumulation of secondary metabolites than monochromatic light. This clear difference was also corroborated by physiological (plant size), morphological (leaf shape), and accumulation of secondary metabolites (leaf pigment) data. These results suggest that light changes can be used in commercial plant cultivation areas that target bioactive compounds. This study also suggested that changing light quality using different LEDs or a combination of LEDs to enhance the productivity (growth) and quality (secondary metabolites) of crops might be a meaningful technique to produce higher quality and less-expensive crops.

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