The Proportion of Blue Light from Light-emitting Diodes Alters Microgreen Phytochemical Profiles in a Species-specific Manner

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Abstract. Microgreens are specialty vegetables that contain human health-promoting phytochemicals. Typically, microgreens are cultivated in controlled environments under red and blue light-emitting diodes (LEDs). However, the impact of varying the proportions of these light qualities on the composition of diverse phytochemicals in indoor-grown microgreens is unclear. To address this problem, the levels of chlorophylls, carotenoids, ascorbates, phenolics, anthocyanins, and nitrate were examined in arugula (Eruca sativa L.), ‘Red Russian’ kale [Brassica napus L. subsp. napus var. pabularia (DC.) Alef.], ‘Mizuna’ mustard (Brassica juncea L.), and red cabbage (Brassica oleracea L. var. capitata f. rubra) microgreens following cultivation under LEDs supplying varying proportions of blue light (5% to 30%) and red light (70% to 95%). Varying the proportion of blue light did not affect the extractable levels of total chlorophyll, total carotenoids, or nitrate in all four microgreen species. Generally, the levels of reduced and total ascorbate were greatest in arugula, kale, and mustard microgreens at 20% blue light, and a minor decrease was apparent at 30% blue light. These metabolite profiles were not impacted by the blue light percentage in red cabbage. Kale and mustard accumulated more total phenolics at 30% blue light than all other blue light regimens; however, this phytochemical attribute was unaffected in arugula and red cabbage. The total anthocyanin concentration increased proportionally with the percentage of supplied blue light up to 30% in all microgreens, with the exception of mustard. Our research showed that 20% blue light supplied from LED arrays is ideal for achieving optimal levels of both reduced and total ascorbate in all microgreens except red cabbage, and that 30% blue light promotes the greatest accumulation of total anthocyanin in indoor-grown Brassicaceae microgreens, with the exception of mustard.

Microgreens are a group of specialty crops that are normally harvested when both cotyledons have expanded and the first true leaves start to emerge (Treadwell et al., 2016). Microgreens are a rich source of health-promoting phytochemicals compared with their corresponding mature plants (Xiao et al., 2012). Phytochemicals that are present in microgreens, specifically in those of the Brassicaceae family, include ascorbate (i.e., vitamin C), chlorophylls, carotenoids, phenolics, and anthocyanins (Xiao et al., 2019). Although light harvesting pigments such as chlorophylls are not well-known for human health benefits, the consumption of the other aforementioned phytochemicals can potentially reduce the risk of chronic and degenerative diseases (Zhang et al., 2015). Carotenoids include carotenes and xanthophylls that function as accessory photosynthetic pigments in plants, some of which are converted in humans to vitamin A, which is crucial for vision, or as antioxidants that can prevent cancer and other chronic diseases (Rao and Rao, 2007; Thoma et al., 2020). Another well-known plant-derived antioxidant is ascorbate. In humans, ascorbate deficiency is linked to scurvy, and dietary ascorbate regulates collagen formation and iron uptake and is involved in epigenetic control of various diseases (Macknight et al., 2017). Phenolic compounds that occur in plants include simple aromatic compounds such as hydroxycinnamic acids and complex polyphenols like anthocyanin pigments. Apart from their antioxidant benefits, some of these phenolic molecules have anti-inflammatory effects in humans (Zhang and Tsao, 2016). In addition, microgreens contain lower levels of nitrate compared with mature leafy vegetables (Pinto et al., 2015). From the standpoint of human health, the effect of dietary nitrate has been the subject of much debate. There is evidence that nitrate derived from vegetables in the diet can provide cardioprotective benefits, including the capacity to lower blood pressure (Machha and Schechter, 2012). There is some concern that an excess of food-derived nitrate can be problematic because it is sequentially converted to nitrite and nitrosamine in the body; furthermore, the accumulation of the latter is associated with an increased risk of certain types of cancer (Bian et al., 2015; Mensinga et al., 2003).

Microgreens can be grown in indoor environments with electrical lighting as the sole light source, which circumvents the restrictions of ambient light on plant growth, such as biomass accumulation (Darko et al., 2014). LED technology has many advantages over traditional light sources (e.g., high-pressure sodium light and fluorescent light), including energy efficiency, low heat output, long operating lifetime, and lower environmental cost (Bourget, 2008). Moreover, LEDs afford the opportunity to modulate spectral composition to optimize the morphology and phytochemical distribution of indoor-grown plants (Bian et al., 2015; Yeh and Chung, 2009). It is known that blue light (400–500 nm) and red light (600–700 nm) are readily absorbed by leaves and used for photosynthesis (McCree, 1972). Red light supplemented with blue light increases photosynthetic capacity and biomass accumulation in plant seedlings (Hogewoning et al., 2010; Tanaka et al., 1998). The combination of blue and red LEDs is widely used for horticultural crop production, including microgreens (Jones-Baumgardt et al., 2019; Massa et al., 2008; Ying et al., 2020).

Typically, the photosynthetic photon flux density (PPFD) that is supplied to leafy green vegetables that are cultivated indoors can be as high as 300 μmol·m⁻²·s⁻¹ (Thoma et al., 2020). For Brassicaceae microgreens grown under blue and red LEDs, PPFDs of 300 to 400 μmol·m⁻²·s⁻¹ are optimal for yield and aesthetic quality (Jones-Baumgardt et al., 2019). However, high blue:red light ratios are known to decrease growth and yield of herbs (Dou et al., 2017). A recent study found that LEDs supplying red:blue light ratios of 95% red:5% blue to 85% red:15% blue are optimal for hypocotyl length and cotyledon area in kale and mustard microgreens; however, a reduction in these morphological.
attributes is apparent at higher blue light percentages (Ying et al., 2020). In addition, microgreens amass more reddish than green surface coloration when the proportion of blue light in the LED array is increased from 5% to 30% (Ying et al., 2020). These morphological changes may be associated with modifications in plant vitamins and secondary metabolites under increasing proportions of blue light supplied by LEDs and potentially affect the nutritional quality.

Light quality has a pronounced effect on the levels of phytochemicals that accumulate within plants under controlled environments (Bian et al., 2015). Previous studies indicated that some phytochemical profiles vary with the proportion of blue light supplied by LEDs, albeit in a manner that is dependent on the plant species and/or developmental stage, including in their microgreen form. For example, the amount of chlorophyll in cucumber (Cucumis sativus L.) leaves is elevated with increases in blue light proportions from 0% to 50% supplied from red:blue LEDs (Hogewoning et al., 2010). In contrast, there is no difference in the chlorophyll concentrations in tomato (Solanum lycopersicum L.), salvia (Salvia splendens Sellow ex Schult.), or impatiens (Impatiens walleriana Hook.f.) seedlings cultivated under LEDs (Hogewoning et al., 2010). In contrast, increasing the percentage of blue light from 0% to 33% enhances the total carotenoid concentrations in beet (Beta vulgaris L.) plantlets (Li et al., 2013); however, ascorbate tends to be enriched at 25 separate canopy level positions, as random placement in six separate compartments within the growth chamber under different lighting treatments. Opaque white vertical blinds separated the light quality treatment compartments within the growth chamber. Each lighting treatment compartment contained two Heliospectra LED arrays (LX602C; Heliospectra AB, Gothenburg, Sweden) that were positioned 36 cm apart.

Materials and Methods

Experimental materials and design. The experiment was conducted in a walk-in growth chamber with a controlled environment [7.3 m x 4.0 m x 2.5 m (length x width x height)] at the University of Guelph. Details related to the experimental design and environmental growth conditions are described in a previous report (Ying et al., 2020). Briefly, seeds of arugula (Eruca sativa L.), ‘Red Russian’ kale [Brassica napus L. subsp. napus var. palubaria (DC.) Afor], ‘Mizuna’ mustard (Brassica juncea L.), and red cabbage (Brassica oleracea L. var. capitata f. Rubra) were separately sown on organic substrates supplied by Greenbelt Microgreens Inc. (Lynden, ON, Canada) in fiber trays (23.5 cm x 48.5 cm x 3.5 cm). After watering, the sown microgreen trays were randomly placed in six separate compartments within the growth chamber under different lighting treatments. Opaque white vertical blinds separated the light quality treatment compartments within the growth chamber. Each lighting treatment compartment contained two Heliospectra LED arrays (LX602C; Heliospectra AB, Gothenburg, Sweden) that were positioned 36 cm apart in the center of the plot area and 56.5 cm above the substrate in the seed trays. The experiment comprised six light quality treatments with varying proportions of blue (445 nm peak) and red (660 nm peak) light supplied from the aforementioned LED arrays. These six LED regimens were as follows: 5% blue light and 95% red light; 10% blue light and 90% red light; 15% blue light and 85% red light; 20% blue light and 80% red light; 25% blue light and 75% red light; and 30% blue light and 70% red light. All lighting treatments had a fixed PPFD of 300 μmol·m⁻²·s⁻¹ and were applied for a 16-h photoperiod. For each light quality treatment, PPFD and spectral distribution were measured at 25 separate canopy level positions, as described previously (Ying et al., 2020). For simplicity, henceforth, all LED quality treatments are referred to as the percentage of supplied blue light. For each lighting treatment, there were four trays of sown seeds, with each tray containing only one of the aforementioned species. The experiment was a randomized complete block design with six light treatments within a single chamber with a controlled environment. The temperature in the walk-in chamber with a controlled environment was set and maintained at 21 °C during the light period (10:00 AM to 2:00 AM) and 17 °C during the dark period. The relative humidity was maintained at ~80% over the course of the experiment. The temperature and relative humidity were monitored by an automated system (Argus Controls Systems Ltd., Surrey, BC, Canada). All growing substrates were top-irrigated at least once each day using tap water until the appearance of visible drainage. A total of three separate light quality experiments were performed.

Tissue sampling for phytochemical analyses. For each microgreen species and light quality treatment within each experiment, three subsamples (~5 g per subsample) of aboveground plant parts, including cotyledons, stems, and first true leaves, were collected 10 d after sowing for kale and red cabbage and 11 d after sowing for arugula and mustard. The harvested microgreens were immediately flash-frozen in liquid N₂, pulverized with a prechilled mortar and pestle. The flash-frozen microgreen powders were stored at –80 °C for up to 8 months before phytochemical analyses. Unless otherwise mentioned, all chemicals required were purchased from Sigma-Aldrich Inc. (Oakville, ON, Canada). Glacial acetic acid, methanol, and ortho-phosphoric acid were obtained from Thermo-Fisher Scientific (Mississauga, ON, Canada).

Chlorophyll and carotenoids. To analyze chlorophyll and carotenoids from each microgreen LED regimen treatment replicate, cryogenic microgreen powder (20 mg) was transferred to a prechilled Eppendorf tube and resuspended in 1 mL of ice-cold 100% methanol. The sample was vortexed for 1 min and incubated on ice under darkness for ~10 min, followed by a second mixing by vortexing before centrifugation at 21,000 g for 5 min at 4 °C. The supernatant was collected and serial dilutions (up to three times) in a final volume of 200 μL were prepared in 100% ice-cold methanol. The methanolic extract and its dilutions were transferred to a 96-well microplate reader (SpectraMax 384 Plus; Molecular Devices, Sunnyvale, CA) and absorbance readings were performed at 665, 652, and 476 nm for chlorophyll a, chlorophyll b, and carotenoids, respectively. The chlorophyll a, chlorophyll b, and total carotenoid concentrations were calculated using the equations for methanolic solutions described by Lichtenthaler and Buschmann (2001).

Ascorbate metabolites. An established method reported by Flaherty et al. (2018) was used to assess reduced ascorbate, dehydroascorbate,
and total ascorbate concentrations. These metabolites were extracted from frozen microgreen powders (200 mg) after grinding the tissue with an ice-cold mortar and pestle with 500 μL of 6% (w/v) meta-phosphoric acid in the presence of acid-washed silica sand and then centrifuged at 13,000 g, for 10 min at 4 °C. The supernatant was filtered through a 0.45-μm polytetrafluoroethylene filter (diameter, 13 mm; Mandel Scientific Inc., Guelph, ON, Canada). The preparation containing reduced ascorbate was analyzed by HPLC as described herein. To analyze total ascorbate concentrations, the supernatant was subjected to a chemical reduction to promote the conversion of dehydroascorbate to reduced ascorbate. A 50-μL aliquot of the supernatant was combined with 25 μL of 400 mM dithiothreitol (DTT) in a 2-M Tris base. After incubating for 15 min at 24 °C, 25 μL of 8.5% (v/v) ortho-phosphoric acid was added to end the reduction step.

The total and reduced ascorbate preparations were analyzed by HPLC; 1 μL of the non-DTT-treated extract (reduced ascorbate concentration) and 2 μL of the DTT-treated extract (total ascorbate concentration) were separately analyzed using the HPLC procedure. The aforementioned extracts were injected on a Restek Ultra Aqueous C18 column (150 × 4.6 mm, 5 μm particle; Chromatographies Specialties Inc, ON, Canada) attached to an Agilent 1200 HPLC (Agilent Technologies, Mississauga, ON, Canada) system and thermostatted at 20 °C. Ascorbate was eluted isocratically with 20 mM ortho-phosphoric acid at a flow rate of 1 mL min⁻¹ over 10 min. The absorbance of the eluate was monitored at 254 nm and ascorbate peaks were compared with a known range of authentic L-ascorbate (Sigma-Aldrich Inc.) standard for quantification of the reduced ascorbate concentration in the DTT-free extract and the total ascorbate concentration in the DTT-treated extract. The dehydroascorbate concentration was calculated by subtracting the ascorbate detected in the DTT-free extract from that apparent in the DTT-treated extract. The levels of ascorbate in both DTT and non-DTT containing extracts were corrected by assessing the recovery of a known amount of authentic reduced ascorbate (377–778 nmol) that was added to a representative duplicate extract. The HPLC analysis revealed that the recovery of the reduced ascorbate that was spiked into the ascorbate extract was in the range of 70% to 97%.

Total phenolic concentration. The total phenolic concentration was evaluated according to the procedure described by Ainsworth and Gillespie (2007), with some modifications. Approximately 20 mg of frozen microgreen powder was transferred to a 2-mL Eppendorf tube, resuspended in 1 mL ice-cold 100% methanol, and vortexed twice for 1 min. The sample was then centrifuged at 13,000 g for 5 min at 4 °C. A 25-μL aliquot of the supernatant and dilutions prepared in 100% methanol were separately dispensed into wells of a 96-well microplate. Thereafter, 125 μL of 10% (v/v) Folin-Ciocalteau reagent was added to each well, and the plate was incubated at room temperature for 10 min, followed by the addition of 125 μL of 7.5% (w/v) Na2CO3. The absorbance was measured at 765 nm using a SpectraMax 384 Plus microplate reader, and the total phenolic concentration was compared with an authentic gallic acid standard curve ranging from 0.018 to 0.6 mg mL⁻¹.

Total anthocyanin concentration. For each microgreen and its light quality treatment replicate subsample, anthocyanins were extracted with acidified methanol according to the method of Roepke and Bozzo (2015). During the extraction process, 100 mg of frozen microgreen powder was combined with 500 μL of methanol:acetic acid:Milli-Q water (9:1:10, v/v/v). The homogenate was vortexed for 1 min, agitated on a nutator (Adams Nutator; Becton, Dickinson, and Company, Franklin Lakes, NJ) for 20 min, and then centrifuged at 16,000 g, for 10 min. The supernatant was collected and the residue was re-extracted twice as described previously. The supernatants from each successive extraction were pooled and partitioned against an equivalent volume of chloroform for the removal of chlorophyll. The 1.5-mL acidified methanol phase was collected and dried in a speedvac (Savant SC100 Speed Vac Concentrator; Thermo Fisher Scientific). The anthocyanin residue was dissolved in 200 μL of methanol containing 0.1% HCL (v/v). Thereafter, the absorption was measured at 530 nm with a SpectraMax 384 Plus microplate reader. The total anthocyanin content was expressed
salicylic acid in sulfuric acid, followed by incubation at room temperature for 20 min. Thereafter, 1 mL of 8% (w/v) NaOH was added to each sample. The absorbance was measured at 410 nm with a SpectraMax 384 Plus microplate reader and compared with a nitrate standard curve ranging from 12.65 to 1600 µg·mL⁻¹. For each sample, nonspecific absorbance was estimated by incubating the aqueous extract with 40 µL of sulfuric acid. To determine the amount of nitrate per microgreen sample, the nonspecific absorbance was subtracted from the absorbance of an assay performed in the presence of salicylic acid.

**Statistical analysis.** The relationships between alterations in phytochemical concentrations in each microgreen as a function of blue light percentage treatment were determined using regression analyses. To this end, linear and quadratic regressions for phytochemical data corresponding to experimental treatment replicates and their subsamples were established with SPSS software (version 25.0; IBM, Armonk, NY). Data were analyzed for normality of residuals and homoscedasticity by using the Shapiro-Wilk and Levene test, respectively. When applicable, homoscedasticity of the data was verified with residual plots.

**Results**

**Chlorophyll and carotenoid concentrations.** Altering the percentage of blue light supplied from LEDs delivering a PPFD of 300 µmol·m⁻²·s⁻¹ had no effect on the concentrations of total chlorophyll and chlorophyll a for all four microgreen species (Fig. 1). The average total chlorophyll contents across all six blue light treatments were 0.71, 0.51, 0.62, and 0.69 mg·g⁻¹ fresh weight (FW) for arugula, kale, mustard, and red cabbage, respectively. Similarly, the blue light percentage supplied from LEDs had no effect on the chlorophyll b, with the exception of mustard. Therefore, the chlorophyll b concentration in mustard microgreens cultivated under 30% blue light was 14% smaller than that in those cultivated under 5% blue light (Fig. 1). The total carotenoid concentration was similar across all LED light treatments, irrespective of species (Fig. 2). The average concentration of this phytochemical was 0.1 mg·g⁻¹ FW across all four microgreens and their light quality treatments.

**Ascorbate metabolite concentrations.** The effect of altering the blue light percentage supplied from sole-source LEDs on ascorbate metabolite composition in microgreens varied with the species (Fig. 3). For red cabbage, the reduced ascorbate concentration was comparable across all light quality treatments. For the other microgreens, the concentrations of reduced and total ascorbate concentrations increased 13% to 29% in microgreens cultivated under 20% blue light relative to 5% blue light. Thereafter, 9% to 11% lower concentrations of reduced ascorbate and 7% to 15% lower total ascorbate levels were apparent at 30% blue light relative to these maximal levels (Fig. 3). Dehydroascorbate levels were unaffected by the percentage of blue light with the exception of arugula, which was 200% to 344% greater under blue light percentages of 15% to 20% relative to the smallest levels detected at 5% and 30% blue light. Dehydroascorbate accounted for 3% to 13% of the total ascorbate in the tested microgreen species.

**Total phenolic and total anthocyanin concentrations.** The total phenolic concentration of microgreens was affected by light quality, although in a species-dependent manner (Fig. 4). For kale, an increased proportion of blue light culminated in more total phenolics; for example, cultivation under 30% blue light yielded an 18% greater total phenolic concentration than that apparent under 5% blue light. Similarly, the concentration of total phenolics in mustard microgreens increased linearly with elevated proportions of blue light; levels detected at 30% blue light were 24% greater than those apparent in the lowest blue light percentage treatment. The concentrations of total phenolics in red cabbage and arugula were not impacted by varying the blue light percentage supplied from LED arrays during cultivation in a controlled environment.

The total anthocyanin concentration was enhanced in arugula, kale, and red cabbage in response to an increased proportion of blue light (Fig. 5). In arugula, kale, and red cabbage, the total anthocyanin levels were, respectively, 70%, 65%, and 75% greater in microgreens cultivated under 30% blue light than under the 5% blue light treatment. Total anthocyanin concentrations in mustard were similar for all LED treatments and were unaffected by the percentage of blue light. At its maximum, the absolute anthocyanin concentration in mustard microgreens was 1.7- to 18.6-fold that of the highest levels detected in the other three microgreens.

**Nitrate concentration.** The concentration of extractable nitrate was unaffected by the percentage of blue light supplied by the LED array, regardless of microgreen species (Table 1). The nitrate levels detected across all microgreens and their LED light quality treatments were between 0.10 and 0.72 mg·g⁻¹ FW. The average level was greatest in red cabbage and least in arugula, with regard to the extractable nitrate concentration.

**Discussion**

Effect of varying blue light percentages in red:blue LED lighting on the concentrations of total chlorophyll, carotenoid, and nitrate concentrations in microgreens. The percentage of blue light supplied by LEDs did not affect the total chlorophyll in the four microgreen species. The impact of blue light on chlorophyll levels in plants during early development seems to be species-dependent. Similarly, chlorophyll levels in imatiops, salvia, and tomato are unaffected by blue light percentages in the range of 6% to 50% supplied by red:blue LEDs (Wollaeger and
The reduction in red light availability may have been instrumental in the lack of effect on chlorophyll production in all four microgreen species investigated because there is a precedence that red light at 660 nm promotes the activation of phytochrome B and, consequently, the involvement of this protein in chlorophyll biosynthesis (Kreslavski et al., 2018). Therefore, lowering the proportion of red light at the expense of enhancing that of blue light may have reduced the activation of phytochrome-mediated regulation of chlorophyll biosynthesis in the Brassicaceae microgreens investigated during this study.

The total carotenoid concentration was unaffected by the increased blue light percentage in this study, which is in accordance with the results of previous research of basal microgreens cultivated under LEDs supplying varying proportions of blue light in the range of 33% to 67% (Lobie et al., 2017). However, the total carotenoid concentration increases in beet microgreens when blue light supplied from red:blue LEDs is increased from 0% to 33%, but not in parsley or mustard microgreens (Samuolienë et al., 2017). As photosynthetic accessory pigments, carotenoids dissipate excess light energy by combining with singlet oxygen to prevent oxidative damage under high light (Jahns and Holzwarth, 2012). The possibility remains that, during the present study, there was no impact of blue light because microgreens were grown under the same PPFD of 300 μmol·m⁻²·s⁻¹. This PPFD has been shown to be optimal for microgreen quality, including the detection of surface green coloration, and, hence, the lack of yellowing in microgreens (Jones-Baumgardt et al., 2019).

Nitrate is the main source of nitrogen that plant roots acquire from the soil and is involved, for example, in the synthesis of proteins and chlorophyll in plants (Marschner, 1995). However, an increased amount of dietary nitrate could have a negative impact on human health. In the present study, extractable nitrate levels were unaffected by LEDs generating varying proportions of blue light percentages, regardless of species. This is consistent with the results of a previous study of lettuce in which nitrate was found to be equivalent in 10% and 25% blue light delivered from red:blue LEDs at a PPFD of 200 μmol·m⁻²·s⁻¹ (Zhang et al., 2018). The nitrate levels in Brassicaceae microgreens are lower than the maximum nitrate levels that are allowable for foodstuffs in some global markets (European Union, 2011). The lack of the blue light effect on nitrate composition is in accordance with the observation that chlorophyll levels in all four Brassicaceae microgreens were not impacted by the proportion of blue light in our study. This association is likely because nitrate availability in leaves affects the biosynthesis of chlorophyll and its precursors (Wen et al., 2019). Previous research indicated that ascorbate:nitrate ratios >1 occur in some greenhouse-grown vegetables, and this comparison is postulated to serve as a measure of the nutritional value of horticultural products (Pokluda, 2006). In this study, the total ascorbate:nitrate ratio ranged between 0.8 and 6.3 across all four microgreen species and their blue light percentage treatments. In general, the average total ascorbate:nitrate ratios were 1.1 and 1.2 in red cabbage and kale, respectively; however, these ratios were higher in arugula and mustard, with averages of 2.2 and 4.1, respectively. It is worth mentioning that the greatest total ascorbate:nitrate ratio was apparent at 20% blue light for all four microgreens assessed during this study. Therefore, the Brassicaceae microgreens cultivated during
the present study can be considered nutritious according to the Pokluda (2006) rationale.  

**Effect of blue light in red:blue LEDs on the concentrations of ascorbate, total phenolics, and total anthocyanins in microgreens.** With the exception of red cabbage, the total and reduced ascorbate concentrations were elevated to a maximum in all microgreens with an increase in blue light (i.e., 20% blue light) relative to the lowest proportion of 5%. Our results mirror those of a previous study of lamb’s lettuce (*Valerianella locusta*) in which ascorbate levels are greatest at 10% to 20% blue light supplied by LEDs in a greenhouse (Długosz-Grochowska et al., 2017). The biosynthesis of ascorbate is dependent on the presence of light and its related biochemical processes, such as the photosynthetic electron transport pathway in leaf chloroplasts (Yabuta et al., 2007). It is known that increased proportions of blue light generated by LEDs than plants cultivated under 0% to 13% and 0% to 35% blue light for ‘Summag’ cultivar and ‘Grand Rapid TBR’ cultivar, respectively (Son and Oh, 2013). However, total phenolics were unaffected by blue light in red cabbage and arugula microgreens in this study. These findings are consistent with the variable effect that red:blue LED ratios have on the concentration of total phenolics in acyanic and cyanic basil microgreens (Lobiuc et al., 2017). The increase in total phenolics with an elevation in the blue light percentage could be due to increased activity of phenylalanine ammonia-lyase, which is the initial step in the phenylpropanoid pathway leading to the production of flavonoids, including anthocyanins. In fact, high anthocyanin levels and phenylalanine ammonia-lyase activity are evident in the leaves of red curly lettuce grown under red LEDs supplemented with blue light relative to plants exposed to red or blue light alone (Heo et al., 2012).  

An increased percentage of blue light in blue:red LED arrays proportionally increased the total anthocyanin concentration in arugula, kale, and red cabbage. Similarly, basil microgreens accumulate more anthocyanins under 50% and 66.7% blue light than under 33.3% blue light (Lobiuc et al., 2017). Overall, the accumulation of anthocyanins under an increased blue light percentage in the microgreens investigated during this study could be associated with more red coloration of microgreen cotyledons, which is a trait that was reported by a previous study (Ying et al., 2020). The increased levels of anthocyanins under a high percentage of blue light could be due to enhanced activities of key anthocyanin biosynthesis enzymes (e.g., anthocyanidin synthase) (Xu et al., 2014). Blue light promotes anthocyanin accumulation primarily through the action of crytochromes, but it also requires active phytochrome proteins (Ahmad and Cashmore, 1997; Bouly et al., 2007; Kadomura-Ishikawa et al., 2013). It is
Table 1. The impact of light quality on total nitrate concentrations of arugula (Eruca sativa L.), ‘Red Russian’ kale (Brassica napus L. subsp. napus var. palubaria (DC.) Aef.), ‘Mizuna’ mustard (Brassica juncea L.), and red cabbage (Brassica oleracea L. var. capitata f. rubra).

<table>
<thead>
<tr>
<th>Blue light percentage</th>
<th>Red cabbage</th>
<th>Kale</th>
<th>Arugula</th>
<th>Mustard</th>
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<tr>
<td>5%</td>
<td>0.53 ± 0.30</td>
<td>0.34 ± 0.21</td>
<td>0.41 ± 0.38</td>
<td>0.12 ± 0.05</td>
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<td>15%</td>
<td>0.47 ± 0.28</td>
<td>0.39 ± 0.33</td>
<td>0.29 ± 0.26</td>
<td>0.27 ± 0.10</td>
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<tr>
<td>25%</td>
<td>0.60 ± 0.30</td>
<td>0.38 ± 0.26</td>
<td>0.41 ± 0.37</td>
<td>0.15 ± 0.05</td>
</tr>
<tr>
<td>30%</td>
<td>0.44 ± 0.28</td>
<td>0.26 ± 0.12</td>
<td>0.41 ± 0.37</td>
<td>0.10 ± 0.05</td>
</tr>
<tr>
<td>35%</td>
<td>0.72 ± 0.41</td>
<td>0.29 ± 0.17</td>
<td>0.30 ± 0.24</td>
<td>0.22 ± 0.11</td>
</tr>
<tr>
<td>40%</td>
<td>0.61 ± 0.33</td>
<td>0.40 ± 0.20</td>
<td>0.31 ± 0.25</td>
<td>0.16 ± 0.07</td>
</tr>
</tbody>
</table>

*Microgreens were cultivated under a photosynthetic photon flux density of 300 μmol·m⁻²·s⁻¹ with varying proportions of red light and blue light.  
*Data represent the mean ± SE of three separate experiments.


