Downsizing in plants—UV light induces pronounced morphological changes in the absence of stress

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Abstract

Ultraviolet (UV) light induces a stocky phenotype in many plant species. In this study, we investigate this effect with regard to specific UV wavebands (UV-A or UV-B) and the cause for this dwarfing. UV-A- or UV-B-enrichment of growth light both resulted in a smaller cucumber (Cucumis sativus L.) phenotype, exhibiting decreased stem and petiole lengths and leaf area (LA). Effects were larger in plants grown in UV-B- than in UV-A-enriched light. In plants grown in UV-A-enriched light, decreases in stem and petiole lengths were similar independent of tissue age. In the presence of UV-B radiation, stems and petioles were progressively shorter the younger the tissue. Also, plants grown under UV-A-enriched light significantly reallocated photosynthates from shoot to root and also had thicker leaves with decreased specific LA. Our data therefore imply different morphological plant regulatory mechanisms under UV-A and UV-B radiation. There was no evidence of stress in the UV-exposed plants, neither in photosynthetic parameters, total chlorophyll content, or in accumulation of damaged DNA (cyclobutane pyrimidine dimers). The abscisic acid content of the plants also was consistent with non-stress conditions. Parameters such as total leaf antioxidant activity, leaf adaxial epidermal flavonol content and foliar total UV-absorbing pigment levels revealed successful UV acclimation of the plants. Thus, the UV-induced dwarfing, which displayed different phenotypes depending on UV wavelengths, occurred in healthy cucumber plants, implying a regulatory adjustment as part of the UV acclimation processes involving UV-A and/or UV-B photoreceptors.
Introduction

The study of plant ultraviolet (UV) responses has gradually shifted from plant stress biology into the realm of plant regulatory responses (Jansen and Bornman, 2012). There is now a good, emerging understanding of the mechanism underly- ing the sensing of UV-B and UV-A wavelengths by a range of dedicated plant photoreceptors, including the phototropins, cryptochromes, and UV RESISTANCE LOCUS 8 (UVR8) (Paik and Huq, 2019). However, understanding of down- stream regulatory interactions which can substantially mod- ify UV-A and UV-B responses is only slowly emerging. Nevertheless, there is consensus that both UV-B and UV-A signaling pathways closely interact (Rai et al., 2019, 2020), with further crosstalk with, among others, phytochrome signaling. For example, UV-B through UVR8 accelerates degra- dation of PHYTOCHROME INTERACTING FACTORS (PIFs) that are part of the phytochrome-mediated elongation re- sponse to high far-red to red light ratio’s (Sharma et al., 2019). As a consequence of these interactions, plant responses under natural light conditions are not always identical to those observed under controlled, artificial lighting in the laboratory. For example, Morales et al. (2013) showed that under natural sunlight, UVR8 both positively and negatively affects UV-A-regulated gene expression and metabolite accumulation. Conversely, a high UV-A/blue light background radiation moderates UV-B-driven gene expression.

Understanding plant UV responses under natural condi- tions is particularly important in the context of climate change. Ongoing changes in the global climate, recovery of the stratospheric ozone layer, and interactions between these two processes are resulting in novel combinations of, among others, temperature, water availability, and solar UV radiation (Bornman et al., 2019). For example, plants in the Mediterranean are predicted to be exposed to higher UV levels, due to climate change-associated changes in cloud cover, together with increased spells of drought (Bornman et al., 2019). It has been hypothesized that UV-exposed plants will be more drought-protected. Indeed, Robson et al. (2015a) showed that when Betula pendula (silver birch) seedlings were exposed to a combination of natural UV and drought, wilting was less pronounced compared with that in plants which had just been exposed to drought. However, not all studies show such cross-tolerance (Rodriguez-Calzada et al., 2019), and there is still considerable uncertainty in the literature concerning the environmental relevance of cross- tolerance (Jansen et al., 2019). Nevertheless, it has been ar- gued that a key component of any putative cross-tolerance is the UV-induced change in plant architecture, and espe- cially a stockier phenotype. The UV-induced phenotype is characterized by shorter stems, internodes, and petioles, and a diminished leaf area (LA), often associated with an in- crease in leaf thickness (Jansen et al., 1998; Robson et al., 2015b), and some of these characteristics are shared with drought-acclimated plants. Yet, major questions remain concern- ing the stocky UV phenotype, and particularly the mechanism underlying the induction of such a phenotype. UV-induced stress, possibly involving reactive oxygen species (ROS; Hideg et al., 2013) and/or accumulation of damaged DNA (Kang et al., 1998), may affect plant architecture (Robson et al., 2015b). Conversely, a regulatory response mediated by a UV photoreceptor can drive architectural change. In the latter case, UV-B- and UV-A-induced responses may be different as these are driven by distinct photoreceptors.

The aim of the current study was to investigate whether morphological changes occurred in plants as a result of supplementing photosynthetically active radiation (PAR) with additional UV-A- or UV-B-enriched light, and to ascer- tain if such alteration is due to known UV-induced stress factors such as reduction of photosynthetic capacity (Jordan et al., 2016), ROS formation (Hideg et al., 2013), DNA damage (Kalbin et al., 2001), or changes in hormonal status (Hideg and Strid, 2017). The study was carried out in cu- cumber (Cucumis sativus L.), a model species representing broad-leaved, high biomass plants of considerable economic importance and which develops considerable phenotypic changes dependent on different wavelengths of UV light (Ballaré et al., 1991).

Results

A more dwarfed, UV-induced, plant architecture has been observed in many different plant species, following exposure to UV radiation. However, some of the strongest morpho- logical responses have been observed in cucumber (e.g. see Qian et al., 2020). In the present study, we analyzed the in- duction of a more dwarfed architecture in cucumber cv Hi Jack. Two-week-old cucumber plants were exposed to UV light for 14 days, during which the stem length of control plants increased from, on average, 4.1 to 47.9 cm. Plants exposed to UV-A- or UV-B-enriched radiation remained comparatively short (Figure 1, A) and reached just 79% and 68% of control stem length, respectively (Figure 1, B; P < 0.05; Table 1).

The UV-mediated decrease in elongation growth was not limited to plant height. A similar impediment of elongation could be observed for petioles. The typical petiole length for a control plant ranged between 1.5 and 11.9 cm for the seventh and the third leaf, respectively.

However, petioles of plants exposed to UV-A- or UV-B-enriched radiation remained considerably shorter (Figure 2). The relative effect of UV-A-enriched radiation was more-or- less constant across the range from older to younger leaves, not exceeding more than 17% inhibition. The decrease in petiole length was significant (P < 0.05) in UV-A-exposed plants for the first, second, and third petiole only (Table 1). In contrast, the effects of UV-B-enriched radiation were par- ticularly pronounced for the youngest leaves (fifth, sixth, and seventh petiole), with petiole length decreasing by more than 40% compared with control plants. This UV-B-induced decrease was significant for the first–seventh petioles (Table 1). Also, progressive decreases in petiole length under UV-B-
enriched radiation were found to be statistically significant when comparing adjoining petioles 1–6 (Table 2), that is a larger decrease the younger the tissue. However, there was no statistically significant difference in the extent of the decrease when comparing the small, developing petioles 6 and 7.

LA was also affected by UV light. Generally, only small changes in LA were obtained following treatment with UV-A-enriched light (Figure 3, A and B). Yet, exposure to UV-B-enriched light led to considerable decreases in LA. The UV-B-induced alteration was more pronounced for younger leaves, as is shown in Figure 3, A for true leaves 2–4 (15%, 24%, and 35% smaller leaves, respectively). For the UV-B treatment, these changes were all statistically significant (P < 0.05), whereas for UV-A only the 10% decrease in LA (Figure 3, A) of true leaf No. 1 was statistically significant.

Table 1 Significant differences between means of measured variables with regard to treatment and where “x” denotes P < 0.05

| Plant part     | Parameter | UV-A:control | UV-B:control | UV-A:UV-B | Figure number |
|----------------|-----------|--------------|--------------|-----------|---------------|
| Stem           | Length    | x            | x            | x         | 1             |
| First petiole  | Length    | x            | x            | x         | 2             |
| Second petiole | Length    | x            | x            | x         | 2             |
| Third petiole  | Length    | x            | x            | x         | 2             |
| Fourth petiole | Length    | x            | x            | x         | 2             |
| Fifth petiole  | Length    | x            | x            | x         | 2             |
| Sixth petiole  | Length    | x            | x            | x         | 2             |
| Seventh petiole| Length    | x            | x            | x         | 2             |
| Second true leaf| Area     | x            | x            | x         | 3A            |
| Third true leaf| Area      | x            | x            | x         | 3A            |
| Fourth true leaf| Area     | x            | x            | x         | 3A            |
| Total leaf     | Area      | x            | x            | x         | 3B            |
| Second true leaf| Dry mass | x            | x            | x         | 3C            |
| Third true leaf| Dry mass  | x            | x            | x         | 3C            |
| Fourth true leaf| Dry mass | x            | x            | x         | 3C            |
| Second true leaf| SLA      | x            | x            | x         | 4A            |
| Third true leaf| SLA       | x            | x            | x         | 4A            |
| Fourth true leaf| SLA     | x            | x            | x         | 4A            |
| Total leaf     | SLA       | x            | x            | x         | 4B            |
| Total plant    | Dry mass  | x            | x            | x         | 5A            |
| Shoot/root ratio|          | x            | x            | x         | 5B            |
| LMF            | x         | x            | x            | x         | 5C            |

T tests were used for total plant or leaf parameters, whereas paired T tests were used for petiole and true leaf parameters comparing developmental effects on same plant individuals. Petioles and leaves are numbered in order of appearance, with higher numbers for younger structures.
In addition, the progressive decrease in true LA under UV-B-enriched radiation was statistically significant for true leaf 2 compared with leaf 3, and leaf 3 compared with leaf 4 \( (P < 0.001; \text{Table 2}) \), confirming a larger decrease the younger the tissue. At the whole plant level, this resulted in a statistically significant decrease in LA by 5% for plants exposed to UV-A-enriched light and by 28% for plants grown under UV-B-enriched light \((\text{Figure 3, B})\). In parallel, UV-B exposure caused a statistically significant decrease in leaf dry weight for true leaves 2–4 \((\text{Figure 3, C and Table 1})\), while UV-A exposure caused a small increase.

The observed decrease in LA, together with a slightly larger decrease in leaf mass in plants exposed to UV-B-enriched light \((\text{Figure 3, A versus C})\), results in a statistically significant increase in specific LA (SLA) in the second and third leaves \((\text{Figure 4, A and Table 1})\). The older the leaves,

the larger the increase in SLA. A similar trend was also seen in the total plant SLA \((\text{Figure 4, B})\) which, however, was not statistically significant. In contrast, a clear and statistically significant negative effect of UV-A-enriched light on second, third, and fourth leaf SLA as well as on total plant SLA is discernible, with total SLA decreasing by as much as 18% compared with control leaves \((\text{Figure 4, B and Table 1})\).

There was a statistically significant difference in SLA between plants exposed to UV-A- and UV-B-enriched light in all cases \((\text{Figure 4, A and B and Table 1})\).

To explore whether the more compact architecture of plants exposed to UV-A- or UV-B-enriched light was related to an overall decrease in growth, both biomass and photosynthetic activity were measured. The dry weight of control plants ranged between 3.8 and 4.6 g (average 4.2 g) after 28 days of growth. Overall, UV-A-enriched radiation significantly stimulated plant biomass production by 14% relative to the control \((\text{Figure 5, A and Table 1})\).

UV-B-enriched radiation had a clear negative impact on biomass accumulation. The statistically significant decrease in biomass caused by UV-B was 28% \((\text{Figure 5, A and Table 1})\). UV-A also impacted on the shoot-to-root ratio, resulting in a 22% decrease \((\text{Figure 5, B and Table 1})\) due to the relatively high root biomass in plants exposed to UV-A-enriched light. No such effect was seen in plants exposed to UV-B-enriched light. A small but statistically significant increase in the leaf weight fraction, relative to the controls \((\text{Figure 5, C and Table 1})\), was induced by both UV-A and UV-B.

To understand the underlying cause of the observed alterations in plant morphology, we explored whether UV-exposed plants exhibited disrupted metabolism as a response to stress. Photosynthetic activities were monitored using chlorophyll fluorometry throughout the experiment for all treatments. The initial measurement of \(F_{v}/F_{m}\) on the four measuring days did not differ between treatment nor day and was 0.792 ± 0.012 \((\text{Supplemental Figure S1})\). Thus, the photosynthetic response did not change over time \((\text{Supplemental Figure S2})\) and just data from day 15, the first day after the end of the UV enrichment, are presented \((\text{Figure 6})\). The measurements were done on the youngest well-developed leaf and followed by exposure to actinic light at low \((302 \mu\text{mol m}^{-2} \text{s}^{-1})\) and high \((1860 \mu\text{mol m}^{-2} \text{s}^{-1})\) PAR. The lower PAR corresponded to a level in the range that the plants experienced during most of the day in the

![Figure 2](https://academic.oup.com/plphys/advance-article/doi/10.1093/plphys/kiab262/6294311)

**Table 2** Pairwise comparisons for statistical significance (paired t test) of petiole length and true LA in plants exposed to UV-B-enriched light and where ** is \(P < 0.01\); *** is \(P < 0.005\); and **** is \(P < 0.001\)

| Treatment       | Parameter | Plant part 1 | Plant part 2 | Level of significance | Figure number |
|-----------------|-----------|--------------|--------------|-----------------------|---------------|
| UV-B-enriched   | Petiole length | First petiole | Second petiole | **                     | 2             |
|                 |           | Second petiole | Third petiole | **                     | 2             |
|                 |           | Third petiole | Fourth petiole | **                     | 2             |
|                 |           | Fourth petiole | Fifth petiole | **                     | 2             |
|                 |           | Fifth petiole | Sixth petiole | **                     | 2             |
|                 |           | Sixth petiole | Seventh petiole | **                     | 2             |
| UV-B-enriched   | True LA   | Second true leaf | Third true leaf | ****                  | 3A            |
|                 |           | Third true leaf | Fourth true leaf | ****                  | 3A            |

Petioles and leaves are numbered in order of appearance, with higher numbers for younger structures.
greenhouse, while the high level corresponded to light saturation, where potential differences in light acclimation are most clearly shown. Despite the exposed position of the youngest fully developed leaf and exposure to the treatments for the entirety of its development, the operation efficiency of PSII ($F_0/F_m$; Figure 6, A) and fraction of open PSII ($q_L$; Figure 6, B) were both unaffected by UV treatments. The only treatment effect was a small but significant increase in heat dissipation through NPQ (Figure 6, C) in cucumbers grown in UV-A-enrichment. This was not a big enough increase to affect $F_0/F_m$ and $q_L$, indicating that photosynthesis was not affected by the UV enrichment.

In parallel to measurements of the photosynthetic activity, chlorophyll content was measured using a Dualex. For the duration of the experiment, the first true leaf slowly accumulated more chlorophyll per unit of LA (Figure 7, A). Treatment with UV-A- or UV-B-enriched radiation had no impact on this process. Likewise, in the youngest well-developed leaf, measured on day 15, one day after the end of UV treatment, there was no statistically significant effect of UV-A or UV-B on chlorophyll content (Figure 7, B).

A key component of plant UV protection is the accumulation of flavonols and related compounds. Here, we show a complex induction curve using two independent approaches that both peaked 5 days after commencement of UV treatment. LAEFC measurements (using the Dualex instrument) revealed that UV-B-enriched, and to a lesser extent UV-A-enriched, radiation-induced accumulation of flavonols in the second true leaf (Figure 8, A) compared with control plants. These changes were statistically significant for days 3, 5, and 10 of exposure (Supplemental Table S1). The model best describing the dependence between the results of the LAEFC and TUAP methods considered treatment (UV-A- or UV-B-enriched, or control) and leaf age:

$$\hat{y} = 0.42 + 0.044X - 0.0065X^3.$$  (1)

A small influence of leaf age ($X_3$) was found. $R^2$ of this fit was 0.84. The residual plots show a random pattern on both sides of 0 (Supplemental Figure S4, A and B), thus justifying the model assumptions that leaf age is also a determinant of the relationship between LAEFC and TUAP. Furthermore, and provided that the linear relationship is valid also for $X_3 < 0$, a zero level in the TUAP parameter (i.e. at the intercept of the axis of the LAEFC parameter) corresponded to a Dualex index of 0.42 in Equation 1, indicating the presence of flavonols in the epidermal cells when the flavonoid level in the bulk of the leaf is negligible.

Linked to the increase in flavonols, a significant increase in total antioxidant capacity (TAC) can be seen in leaves exposed for 3 or 5 days to UV-B-enriched light. To a small extent, leaves exposed to UV-A-enriched light also significantly increased their TAC (Figure 8, C and Supplemental Table S1). Interestingly, whereas both the LAEFC and TUAP parameters on day 14 returned to a level similar to the one...
before onset of UV exposure, or slightly below, the TAC parameter decreased significantly to approximately 50% of the initial value, independently of whether the plants had experienced any of the UV exposures or were controls.

To more accurately examine this biphasic nature of the TAC measurements, we studied the dependence between this assay and the two other methods applied in this study (LAEFC and TUAP). To describe the dependence between the TAC and the LAEFC assays, we assumed two simple linear relationships, without taking into account the type of treatment (UV-A- or UV-B-enriched), but dividing up the samples in those from younger leaves (≤ 5 days of exposure time, i.e. ≤ 19 days after sowing) and those from older leaves (≥ 10 days of exposure time, i.e. ≥ 24 days after sowing). In this case, the differences of the linear relationships between the TAC and LAEFC assays became apparent, as is shown in Figure 9, A. The model for the linear dependence with regard to young leaves was found to be:

\[
\hat{y} = 0.143 + 0.1276x,
\]

whereas for the linear dependence with regard to older leaves, the equation was

\[
\hat{y} = 0.124 + 0.2068x.
\]

\(R^2\) for the fits of data points to the two assumed linear relationships was 0.72 and 0.83, respectively. The intercept at zero TAC level was similar for both linearizations with a LAEFC value of approximately 0.13 (Eqs 2 and 3), indicating a low level of leaf epidermal flavonols that are not active as antioxidants.

Finally, we scrutinized the linear dependence between the TUAP (Figure 8, B) and the TAC methods (Figure 8, C), using the same model as applied above. In Figure 9, B, we again assumed two simple linear relationships, without taking into account the type of treatment (UV-A- or UV-B-enriched), but dividing up the samples in those for younger leaves (≤ 5 days of exposure time, i.e. ≤ 19 days after sowing) and those from older leaves (≥ 10 days of exposure time, i.e. ≥ 24 days after sowing). Two linear relationships between the TAC and TUAP data were estimated and the equation with regard to younger tissue was found to be

\[
\hat{y} = 2.37 + 0.29x.
\]

In this case, the equation explained variation in data with a factor \(R^2\) of 0.91. For older tissue, the equation became

\[
\hat{y} = 0.39 + 0.40x.
\]

\(R^2\) was even higher (0.98) for the older leaves and the difference between the intercepts at 0 absorbance for TUAP differed approximately six-fold (0.39–2.37).

To further understand the link between UV exposure and induced changes in plant morphology, accumulation of CPD dimers was measured in leaves of plants exposed to UV-A- or UV-B-enriched light and in control plants to see whether induced CPDs were associated with the smaller cucumber phenotype. Overall, the number of CPDs was low, and, with the exception of day 14, variability was limited. There were no statistically significant effects of UV-A or UV-B on CPD accumulation (Figure 10).

Finally, to explore whether changes in key plant hormones are associated with observed changes in plant morphology, leaf concentrations of abscisic acid (ABA), gibberellins (GA), and the auxins indole acetic acid (IAA) and indole-butyric acid (IBA) were quantified. Statistically significant decreases in ABA, IAA, and IBA were associated with leaf development.
No statistically significant effects of UV treatment were observed, although small decreases in the GA GA1, GA44, GA6, and GA15 were noted in plants exposed to either UV-A- or UV-B-enriched radiation (Table 3).

Discussion

Cucumber displays a strong morphological response when exposed to supplemental UV

Here, we explored the regulation of plant morphology by white light enriched with either UV-A or UV-B. The data show that enrichment of the spectrum with either UV wavelength results in a stocky cucumber phenotype. These data are in agreement with previous studies that show that cucumber is particularly responsive to UV-exposure (Krizek, 1978; Murali and Teramura, 1986; Ballaré et al., 1991; Adamse and Britz, 1992; Adamse et al., 1994; Krizek et al., 1994, 1997; Takeuchi et al., 1996; Fukuda et al., 2008; Shinkle et al., 2010; Yamasaki et al., 2010, 2014; Qian et al., 2019, 2020). Therefore, we consider cucumber a promising model species for the study of the stocky UV phenotype, a role that will be facilitated by the large leaf surface area which enables measurement of potential hormone gradients within organs. The data presented in this study show that especially elongation is affected by UV-exposure, with noted decreases in stem and petiole length, as well as in LA. These effects were larger in plants that had been grown in the UV-B-enriched than in the UV-A-enriched light environment (Figures 1, 2, and 3, A and B). Interestingly, whereas the petiole lengths and LA in plants treated with UV-A-enriched light were approximately the same independent of tissue age, there was a progressively larger decrease in these parameters the younger the tissue in the presence of UV-B radiation (Figures 2, 3, A and Table 2). Finally, in plants grown under the UV-A-enriched light regimen, a significant reallocation of photosynthate from shoot to root by more than 20% was seen (Figure 5, B). The differences between the effects of UV-A- and UV-B-enriched light on cucumber morphology indicate that there may be different developmental regulatory mechanisms involved in the two cases.

The overall results are thus consistent with earlier descriptions of the UV phenotype, which referred to a more "stocky" architecture (Barnes et al., 1996; Jansen et al., 1998; Robson et al., 2015b). Other aspects of the UV phenotype, such as leaf thickening are also apparent in the current study, and a decrease in SLA was noted in plants exposed to UV-A-enriched radiation. A similar UV-induced stocky phenotype has been observed in a substantial number of plant species (Robson et al., 2015b). Yet, this phenotype remains an enigma, in that major questions remain to be answered with respect to the wavelength specificity of its induction, the underlying mechanism of the response, and the functional importance of the induced architectural response for the plant.

The stocky UV phenotype is not associated with plant stress

Several hypotheses have, over the years, been proposed to explain the mechanism underlying UV-induced changes in plant morphology (Robson et al., 2015b). High UV intensities can drive the development of stress-induced morphogenetic responses (SIMRs) as first proposed by Potters et al. (2007). These responses are associated with the disruption of cellular metabolism (distress), resulting in a localized cessation of growth. Therefore, intensities of the UV light used are key and dose-dependent, as well as UV:PAR ratios (Rai et al,
Figure 6 Photosynthetic parameters measured in cucumber plants grown under different light conditions. A, The response of the operation efficiency of PSII ($F_{\text{q}0}/F_{\text{m}0}$). B, The fraction of open PSII ($q_L$). C, Heat dissipation measured as NPQ of fluorescence measured on the youngest well-developed leaf under an actinic PAR of 302 or 1860 μmol m$^{-2}$ s$^{-1}$ on day 15 after commencement of UV exposure (last day of UV exposure day 14) to UV-deficient control (white), UV-A-enriched (gray), and UV-B-enriched light (closed squares) treated plants. The data represent mean values ± SD with $n = 9$ for UV treatments and $n = 18$ for controls. T test was used for statistical analysis. ** were used to represent the significant difference for $P < 0.01$.

Figure 7 Chlorophyll content in cucumber plants grown under different light conditions. A, The adaxial surface chlorophyll levels expressed as "Dualex chlorophyll index" of the first true leaf of UV-deficient controls (open circles), UV-A-enriched light (gray triangles), and UV-B-enriched light (closed squares) treated plants. B, Adaxial surface chlorophyll levels of the youngest well-developed leaf measured on day 15 after commencement of treatment using UV-A-enriched (light gray) or UV-B-enriched (dark gray) growth light, compared with the corresponding UV-deficient control (white). Sampling was done as in Figure 1. The data represent mean values ± SD, $n = 9$ for the UV-enriched treatments and $n = 18$ for the control. Two-way ANOVA was used to test the effect of exposure time, and treatment on adaxial surface chlorophyll levels of the first true leaf. T test was used to test the significant difference of upper surface chlorophyll levels in youngest well-developed leaf between UV-treated and control samples.
In fact, UV dose–responses remain largely unknown, but those studies that have compared effects of multiple
doses on morphological parameters, suggest a complex re-
sponse curve (Brodfuehrer, 1955; Van de Staaij et al., 1997;
Qaderi et al., 2008). A key target for future research is deter-
mining the dose that causes the transition from physiologi-
cal regulation of morphology to SIMR, a transition that is of
interest from both a scientific and a horticultural perspec-
tive. The SIMR phenotype is characterized by decreased
elongation growth, which can result in a stockier phenotype,
as described in the current study. However, in the current
study, there is no evidence for disruptive stress. There was a
small increase in NPQ in plants grown under UV-A-enriched
light but no change was found in neither maximum photo-
chemical efficiency, redox state of PSII, nor the operation ef-
ciciency of PSII in any of the UV treatments measured using
chlorophyll a fluorometry, nor in total chlorophyll content.
Furthermore, there is no evidence for an increase in accu-
mulation of damaged DNA in the form of CPD dimers.
DNA damage may potentially result in a stocky phenotype
as UV-induced dimerization of DNA can impair DNA repli-
cation, and hence impede cell cycle progression, particu-
larly by slowing the G1-to-S phase (Jiang et al., 2011).
Indeed, several earlier reports refer to UV-mediated im-
pairment of cell division (Dickson and Caldwell, 1978;
Wargent et al., 2009). Others (Lake et al., 2009) refer to
larger cells in UV-exposed plants which can be explained
by endoreduplication, resulting in fewer, but bigger cells
(Radziejwoski et al., 2011). Plants showing symptoms of
UV stress would often have between 50 and 800 CPDs/Mb
(Kang et al., 1998; Kalbin et al., 2001; Pescheck et al.,
2014). Yet, in the current study, levels of CPDs/Mb were
one to two orders of magnitude less, indicating efficient
repair of damaged DNA. Therefore, the observed stocky
phenotype is not associated with accumulated DNA dam-
age. The data on the ABA content also are consistent with
non-stress conditions. Furthermore, the data show
increases in total antioxidant activity (TAC) and flavonol
concentrations in leaf adaxial epidermis (LAEFC) and the
entire leaf (TUAP). Taken together, these data reveal suc-
cessful UV acclimation. Thus, healthy plants display the
stocky UV phenotype, implying that the strong morpho-
logical response observed in UV-exposed cucumber seed-
lings is a regulatory adjustment that is part of the UV
acclimation processes involving UV-A and/or UV-B
photoreceptors.

Figure 8 Flavonoid content and antioxidative capacity in cucumber
plants grown under different light conditions. A, LAEFC measured
with Dualex. B, Total UV-absorbing pigments, TUAP, measured spec-
trophotometrically at 330 nm per leaf fresh weight. C, TAC measured
as nmol Trolox equivalents per mg leaf fresh weight. Measurements

Figure 8 (Continued)
were performed on the second true leaf of two-week-old cucumber
plants grown under UV-A-enriched (gray triangles) or UV-B-enriched
light (closed squares), respectively, and compared with the corre-
sponding controls (open circles). Data represent mean values ± se
with n = 15–18 for the UV-enriched treatments and n = 30–36 for the
control treatment in (A); n = 9 for the UV-enriched treatments and
n = 18 for the control treatment in (B); n = 3 for the UV-enriched
 treatments; and n = 6 for the control treatment in (C).
UV acclimation is accompanied by a decrease in biomass accumulation
Notwithstanding the apparent lack of plant stress under either UV-A- or UV-B-enriched light, a substantial decrease in produced total biomass was noted for plants exposed to UV-B-enriched light. We have interpreted this decrease in biomass production as a secondary consequence of UV acclimation. UV-B-exposed plants with shorter stems (including shorter internodes) and shorter petioles will condense the same number of leaves in a smaller area, hence increasing the likelihood of self-shading which, in turn, may decrease overall PAR capture, and hence photosynthetic productivity (Barnes et al., 1996). Consistently, UV-A had considerably smaller impacts on both stem and petiole length, and this was associated with a lack of impact on biomass production. UV-B-induced decreases in LA will further hamper PAR capture, an effect that is much smaller in plants raised under UV-A-enriched light. These UV-B-mediated decreases in light capture may also be accompanied by the often-reported UV-B-induced stomatal closure (He et al., 2013; Martínez-Lüscher et al., 2013; Tossi et al., 2014), which would similarly decrease photosynthesis in situ, and potentially decrease biomass production. In addition, the measured UV-A-induced decrease in SLA may potentially improve the water use efficiency of plants (Liu and Stützel, 2004), and although water use efficiency has not been measured in this study, several studies have reported UV-induced co-tolerance with drought (reviewed in Barnes et al., 2019).

Impact of UV-A- and UV-B-enriched light on flowering, fruit setting, and final yield
In a previous horticultural study carried out in two different nurseries over 3 years (Qian et al., 2020), it was found that...
Table 3 Main plant hormones, including ABA, GA, and the auxins IAA and IBA, present in the second true leaf on day 3, and 5 of UV treatments

| Hormone species | UV-A 3 day control | UV-A 3 day exposed | UV-A 5 day control | UV-A 5 day exposed | UV-B 3 day control | UV-B 3 day exposed | UV-B 5 day control | UV-B 5 day exposed |
|-----------------|-------------------|--------------------|--------------------|--------------------|-------------------|-------------------|-------------------|--------------------|
| ABA             | 85.5 ± 12.3       | 96.8 ± 9.0         | 48.3 ± 12.0        | 67.7 ± 16.0        | 71.9 ± 14.1       | 77.7 ± 3.4        | 54.9 ± 2.6        | 52.4 ± 13.4        |
| GA1             | 17.0 ± 7.4        | 9.7 ± 0.8          | 13.3 ± 1.2         | 11.7 ± 3.3         | 11.1 ± 2.6        | 10.0 ± 2.8        | 16.6 ± 5.7        | 11.7 ± 1.5         |
| GA4             | 2268 ± 1447       | 1123 ± 342         | 2104 ± 1214        | 1315 ± 186         | 1485 ± 312        | 1099 ± 298        | 4850 ± 3668       | 1184 ± 155         |
| GA9             | 468 ± 288         | 177 ± 55           | 424 ± 62           | 360 ± 120          | 268 ± 131         | 184 ± 89          | 832 ± 626         | 237 ± 57           |
| GA15            | 2367 ± 1403       | 839 ± 475          | 1825 ± 1112        | 1281 ± 713         | 1579 ± 944        | 1035 ± 617        | 4925 ± 3776       | 1109 ± 427         |
| IAA             | 84.4 ± 33.4       | 95.4 ± 3.0         | 60.2 ± 5.0         | 67.3 ± 6.6         | 88.4 ± 20.1       | 89.9 ± 3.7        | 67.2 ± 6.2        | 70.1 ± 7.5         |
| IBA             | 1362 ± 387        | 1242 ± 67          | 361 ± 139          | 522 ± 125          | 1197 ± 142        | 865 ± 368         | 531 ± 100         | 489 ± 193          |
| IBA-OH          | 1141 ± 357        | 934 ± 53           | 278 ± 108          | 427 ± 126          | 971 ± 133         | 696 ± 309         | 450 ± 90          | 422 ± 188          |

Notes: Plant tissue from one plant in each of three boxes per treatment was pooled together with three separate experiments giving n = 3 for the UV-enriched treatments and the control treatment. The data represent mean values ± SD and are given in pmol (g fresh weight)−1. For data of each plant hormone species, t test was used to test the significant difference between each treatment. The levels of the following hormone species were under the detection limit: GA3, GA4, GA5, GA8, GA12, GA19, GA20, IAA-OX, IAA-OH, IBA-OX, IBA-OH-C and IBA-OX-C. The levels of the following hormone species did not show any clear trend: GA7, IAA-C and IBA-C.

flowering, fruit setting, and total fruit yield were not influenced by pre-treatment of plants with UV-A-enriched light for 2 weeks. Two weeks pre-treatment with UV-B-enriched light led to an increased number of flowers in flowering nodes primarily in the third year (Supplemental Figure S3), whereas delayed fruit setting by 2 weeks was found in the experiment carried out in the second year but not in the first or the third year. The total yield of cucumber fruits was however not influenced by either UV-A- or UV-B-enriched light, compared with their corresponding controls, in any of the 3 years (Qian et al., 2020). Thus, apart from the fact that the total yield was not influenced by the treatments, flowering and fruit set was more variable in cucumber plants grown under UV-B-enriched conditions.

Regulatory mechanism(s) underlying the stocky UV-induced phenotype

Since the data in this study do not support an association between the stocky UV phenotype and plant stress, one or more different specific regulatory responses should be considered, as argued above, dependent on what part of the UV spectrum is supplementing the PAR. The UV-B photoreceptor UVR8 was discovered in Arabidopsis thaliana mutants that did not show UV-B-induced dwarfing of hypocotyls (Hayes et al., 2014). Thus, UVR8 has been strongly associated with control of plant architecture. It should also be noted that, in a recent study by Rai et al. (2020), there was a marked difference in regulation of gene expression between a UVR8-dominated effect at wavelengths below 335–350 nm and a cryptochrome-dominated regulatory mechanism at UV wavelengths above 350 nm. Notwithstanding, the UVR8- and CRY-regulatory mechanisms were interdependently influenced by each other.

Thus, the mechanism underlying the UV-B-induced stocky phenotype may relate to interactions with various cellular signaling pathways, including the phytochrome and cryptochrome pathways. In UV-exposed plants, UVR8 monomers bind CONSTITUTIVELY PHOTOMORPHOGENIC 1 (COP1), and the resulting UVR8-COP1 complexes enter the nucleus and promote UV-B signaling which inhibits auxin biosynthesis, signaling as well as hypocotyl elongation (Hectors et al., 2012). Parallel increases in the expression of the ELONGATED HYPOCOTYL 5 (HY5)/HY5-HOMOLOG (HYH) transcription factor may enhance transcription of polar auxin transport proteins PIN1 and PIN3, as well as several regulators of auxin signaling (Vanhaelewyn et al., 2016). Furthermore, sequestration of COP1 by UVR8 destabilizes PIF5, further interfering with auxin biosynthesis and signaling (Hayes et al., 2014; Vanhaelewyn et al., 2016). Yet, few studies have been able to demonstrate UV-induced changes in auxin levels. Hectors et al. (2012) reported (non-significant) UV-B-induced decreases in auxin levels in Arabidopsis leaves, as well as altered UV-B responses in auxin influx and biosynthesis mutants (Hectors et al., 2012). The current study does not present evidence for significant changes in auxin concentrations either, despite strong morphological responses, but a non-significant trend of decreasing GA concentrations is observed in both UV-A- and UV-B-exposed plants. UVR8 binding to COP1 can, via upregulation of transcription of HY5 and HYH, result in an increase in gibberellin 2-β-dioxigenase 1 (GA2OX1) levels, reducing GA concentrations (Hayes et al., 2014; Vanhaelewyn et al., 2016). The current study shows a trend of decreasing concentrations of GA1, GA44, GA9, and GA15, in both UV-A- and UV-B-exposed plants. Hormonal action may be important in cross-talk between UV light and other environmental factors affecting for instance plant water use efficiency and drought tolerance (Bernhard et al., 2020). However, the data in the current paper are not conclusive with respect to a role for UVR8, auxin, or gibberellic acid in mediating plant UV responses.
In fact, it is debatable whether the observed dwarfing response can simply be explained as UVR8 mediated, given that responses are induced by both UV-B and UV-A radiation, albeit with partly different outcomes (progressive decreases in petiole lengths and LA in plants grown in UV-B-enriched light; alteration in carbon allocation from shoots toward roots in UV-A-enriched light). Therefore, although the UVR8 action spectrum remains to be fully characterized in detail, particularly with respect to potential interactive responses to UV-A wavelengths, recent evidence suggests antagonistic effects whereby responsibility to UV-B is modulated by UV-A wavelengths and vice versa (Morales et al., 2013; Rai et al., 2020). Thus, although a UVR8-mediated mechanism appears to be the most likely candidate to explain observed decreases in both organ elongation and GA concentration in UV-B-exposed plants, major questions remain to be addressed concerning the regulation of the stocky phenotype under natural, solar light conditions where there is considerable scope for interactions between multiple wavelength bands, photoreceptors, and signaling pathways. In fact, brassinosteroids may also play a role in UV-regulation of gene expression (Sävenstrand et al., 2004) and development (Liang et al., 2018), a role which was further strengthened by the discovery of interaction of UVR8 with molecular regulators of brassinosteroids.

**Regulation of cucumber phenylpropanoid genes**

The regulation of all 12 different cucumber phenylalanine ammonia lyase (CsPAL) genes, all three cinnamate 4-hydroxylase (CsC4H) genes, and all three chalcone synthase (CsCHS) genes, as the result of exposure to UV-A- and UV-B-enriched light, has previously been reported (Qian et al., 2019). The proteins encoded by these genes encode the PAL, C4H, and CHS isoenzymes of the phenylpropanoid pathway that are all involved in the synthesis of flavonoids, including flavonols. Some of the genes were up-regulated by UV-A and others by UV-B. In particular, three of the latter genes (CsPAL4, CsPAL10, and CsCHS2) were up-regulated at least 30-fold by UV-B exposure. These three genes had in common an enrichment of upstream cis-acting elements (MREs, ACEs, and G-boxes) that may be of importance for amplification of expression of flavonoid biosynthesis genes under UV-B-containing light (Qian et al., 2019). Therefore, it is obvious that there are distinct regulatory effects by UV-B and UV-A on genes of importance for UV tolerance in cucumber.

**Relationships between flavonoid content and antioxidant capacity**

Protection against UV light was measured in three different ways reflecting the notion that UV-B-induced flavonoid compounds have both an antioxidative effect and a function as UV-absorbing compounds (Agati et al., 2012; Hideg and Strid, 2017). For estimation of leaf adaxial epidermal flavonol content (LAEFC), the Dualex method was used, for TUAP acidic methanol extraction and spectrophotometric detection at 330 nm was employed, and total anti-oxidative activity (TAC) was assayed using the Trolox method. Comparisons between LAEFC and TUAP measurements have been made previously (Barthod et al., 2007), and in line with published results the intercept of the plot of LAEFC versus TUAP measurements (Supplemental Figure S1) is inferring higher flavonol content in the epidermal cell layers compared with the average in the total foliar biomass.

The data of this paper show an increased flavonol level in leaves for up to 5 days after all three treatments (no UV control, UV-A-, or UV-B-enriched light). Thereafter, the flavonoid content declined (Figure 8, A and B). The dependence of the flavonoidal levels (LAEFC and TUAP), with a peak at Day 5 is likely to be due to leaf developmental processes. TAC also peaked on day 5 (Figure 8, C). That the peaks in all three parameters at Day 5 could be related to particular weather conditions seem highly unlikely given that all experiments were independently replicated three times during different weeks between February and June. For TUAP, readings after 14 days of UV exposure were similar to those measured at the onset of the experiment (day 0). For the LAEFC, the readout on day 14 returned to levels slightly lower than at the outset, whereas for TAC the antioxidant capacity decreased between day 0 and day 14 by about 50%, independently of whether the plants had experienced any UV exposures or were controls. Thus, as these trends were observed in both control and in UV-treated plants, a developmental process would be the likely cause. To improve understanding of developmental processes as well as the relationships between the three different parameters, further regression analyses were performed.

The slope of the TAC curve of antioxidant capacity when compared LAEFC or TUAP results indicate that the TAC parameter reflects two different physiological means of antioxidative activity, one being flavonoids and the second being another type of ROS scavenging of enzymatic or non-enzymatic nature, and which shows an age-dependent decrease by half through days 10–14. When estimating the linear dependence of the TAC and LAEFC experiments, we found that the intercept at a zero TAC level was similar for the two leaf-age-dependent linearizations used (Eqs 2 and 3), with a LAEFC flavonol index of approximately 0.13 (Figure 9, A), which indicates the presence of some flavonoidal compounds that lack antioxidant capacity, for example monohydroxylated species (Agati et al., 2012). Differences in the antioxidant capacity of different flavonol species have been extensively demonstrated (Csépegi et al., 2016). For example, some flavones, such as apigenin, have particularly low ferric ion reducing antioxidant power (Csépegi et al., 2016). Also, it should be considered that Trolox may not be a perfect proxy for antioxidant activity in general (Csépegi et al., 2016).

The difference in the slope of the two linear relationships in Supplemental Figure 9, A and the different shapes of the curves in Figure 8, A and B, indicate that, as the leaf tissue aged, the pool of leaf epidermal flavonols had lost its antioxidative capacity, either by accumulation of more oxidized...
forms of this type of compound, because of increased O-glycosylation, or due to another yet to be identified mechanism. The OH group on the 3-position on the A-ring of the flavonoid backbone is commonly glycosylated, which decreases antioxidant activity. Developmental changes in flavonol profiles have also previously been shown in for instance white mustard (Sinapis alba; Reifenrath and Müller, 2007) and grapevine (Vitis vinifera; Bouderias et al., 2020). Also, a study by Morgenstern et al. (2014) shows that the ratio of quercetin to kaempferol in buckthorn (Hippophaë rhamnoïdes) shows a strong developmental trend, rising from just over 100 at the beginning of the season, to over 400 by mid-summer, and this effect is paralleled by a drop in gallic acid and rutin levels (Morgenstern et al., 2014). Given the different antioxidant activities of different flavonols, this may underpin a change in antioxidant capacity.

In our study, the loss of anti-oxidative defense (i.e. TAC) was independent of whether the plants had been kept under control conditions or under supplementary UV-A- or UV-B-enriched light. Thus, this seems to be a true effect of leaf age. In old leaves, flavonoids constitute the bulk of the antioxidant capacity (intercept close to 0 in Figure 9, B), whereas in younger tissue half of the oxidant capacity (comparison of the curves in Figures 8, B and C and considering the intercept at 2.37 in Figure 9, B) is contributed by antioxidative systems other than flavonoids, be it enzymatic or non-enzymatic, that do not absorb light at 330 nm in methanol under acidic conditions. Leaf age-dependent changes in TAC have previously been shown in greenhouse-grown grapevine leaves (Majer and Hideg, 2012), using a several-fold higher biologically effective UV-B dose than we did. Four days of exposure led to a large increase in TAC in young leaves, similar to what we found in cucumber. However, in old leaves, 4 days of UV exposure led to decreased TAC. Clearly, there are strong interactions between UV acclimation and developmental processes that govern as disparate physiological parameters in plants as stem and petiole stretching, leaf expansion, flavonoid content, and TAC.

Conclusion
In this paper, we show that cucumber grown in UV-A- or UV-B-enriched light led to a stockier phenotype compared with non-UV-irradiated control plants. In plants grown in UV-A-enriched light, the decreases in stem and petiole lengths were similar independent of tissue age whereas in plants grown in UV-B-enriched light stems and petioles were progressively shorter the younger the tissue. In addition, plants grown under UV-A-enriched light significantly reallocated photosynthates from shoot to root, had thicker leaves, and decreased SLA. This infers different morphological plant regulatory mechanisms under UV-A and UV-B radiation, especially since there was no evidence of stress in any of the UV-exposed plants, as judged by the absence of effects on photosynthetic parameters, cyclobutane pyrimidine dimer levels, or ABA content. The total leaf antioxidant activity and UV-dependent accumulation patterns of flavonoidal compounds and leaf-age-dependent variation in these parameters also indicated successful acclimation of the plants to the two UV light regimens. Therefore, we conclude that the stocky UV phenotype developed in healthy plants, which in turn implies a strong regulatory adjustment and morphological response as part of a successful UV acclimation processes involving UV-A and/or UV-B photoreceptors.

Materials and methods
Plant material, growth conditions, and treatment conditions
Cucumber seeds (C. sativus L. cv “Hi Jack”) were sown one per 0.25 L pot in 14-7-15 NPK fortified peat (SE Horto AB, Hammenhög, Sweden), as described previously (Qian et al., 2019, 2020). Seedlings were grown in a greenhouse under natural daylight from the roof which was supplemented with 150–200 µmol m–2 s–1 PAR as measured 20 cm above the table using Vialox NAV-T Super 4Y high-pressure sodium lamps (Osram, Johannesburg, Sweden) for 16 h per day centered around solar noon, and only turned off when the natural irradiance reached 900 µmol m–2 s–1. The day/night temperature was 25°C/20°C and the relative humidity was set to 80%. Watering was done by adding water to the tray underneath the pots when the tray itself was completely dry. As soon as the cucumber seedlings had fully developed cotyledons, watering was commenced using a full nutrient solution (Svegro AB, Ekerö, Sweden).

Fourteen days after sowing, when the first true leaf of the cucumber seedlings was approximately 5 cm in diameter (about one-third of the diameter of a fully developed first true leaf), UV exposure commenced. The plants were then given either supplementary UV-A-enriched or UV-B-enriched irradiation for 4 h per day (centered around solar noon) in addition to the PAR described above. Controls were simultaneously exposed to PAR only (see below) in the same chamber as the corresponding UV-treated plants. The UV-A and UV-B exposures were carried out in separate greenhouse chambers and the treatments were alternated between the chambers when repeating the experiment (cf., Qian et al., 2019, 2020, for details).

Open top, front, and backside boxes (OTFB boxes), covered with Perspex on the left and right sides, were used for the different UV exposures. Each greenhouse compartment was equipped with up to six boxes, three being used for the UV treatments and three for the corresponding controls. Each OFTB box contained up to 48 plants per replicate. For the UV-A-enriched experiments, fluorescent UVA-340 tubes (Q-Lab, Cleveland, OH, USA) were used for exposure, whereas for the UV-B-enriched experiments, fluorescent Philips TL40/12 UV tubes (Eindhoven, The Netherlands) were employed. For the control OFTB boxes, UV-blocking Perspex was used to cover the top and all sides. For the UV-B-enriched experiment, 0.13-mm cellulose acetate (Nordbergs Tekniska AB, Vallentuna, Sweden) covered the top, front, and backside of the OFTB boxes with the purpose to remove any UV-C radiation emitted by the Philips TL40/12 UV tubes.
TL40/12 tubes. For the UV-A-enriched experiment, the OTFB boxes were similar to the boxes used in the control experiment but without any filtering material on top.

The spectral distribution of the light environments in the different treatments was measured using an OL756 double monochromator spectroradiometer (Optronic Laboratories, Orlando, FL, USA) 20 cm above the table. The details of doses were as described by Qian et al. (2019). Briefly, however, UV-A-enriched radiation contained 3.6 W UV-A m⁻² and a 45.5 mW m⁻² plant-weighted UV-B (calculated according to Yu and Björn, 1997), giving a total of plant-weighted UV-B of 0.6 kJ m⁻² day⁻¹ during the daily 4-h exposure. The UV-B-enriched irradiation had 83.4 mW m⁻² plant-weighted UV-B totaling 1.23 kJ m⁻² day⁻¹ plant-weighted UV-B. This exposure also contained 0.34 W UV-A m⁻². The daily irradiation outside in Lund, Sweden, under clear skies on a summer’s day is approximately 4.8 kJ m⁻² day⁻¹ of plant-weighted UV-B (Yu and Björn, 1997).

Morphological measurements
Between day 0 and day 14, morphological parameters were measured. A ruler was used to measure the lengths of stems and petioles. The dry matter (DM) of shoots (separated into stems, petioles, and leaves) and roots was measured using a digital balance (accuracy 0.001 g) following oven drying at 70°C for 20 h. The leaf mass fraction (LMF) was calculated as LMF = leaf DM/total shoot DM. The LA was determined from digitized photographs using ImageJ (https://imagej.nih.gov/ij/). As a measure of how much LA a plant builds with a given amount of leaf biomass, the SLA for true leaves 2–4 from the base of the plants, as well as for all leaves combined, was calculated as SLA = LA/leaf DM. For each experiment, six plants per treatment were measured, two from each of the three replicated treatment OTFB boxes and their corresponding controls. In total, three independent experiments were performed.

Chlorophyll fluorescence
Chlorophyll a fluorescence was measured with a MINI-PAM (Walz, Effeltrich, Germany) on the first true leaf from the bottom of the stem on days 1, 4, 8, and 14 of UV exposure, as well as on the youngest well-developed leaf (the top leaf which was fully developed, i.e. the diameter reached approximately 15 cm) on day 15 of UV exposure. In dark, the middle portion of attached leaves were fixed in a leaf clip holder (2030-B, Walz) fitted with a halogen lamp (2050-HB, Walz) and a heat absorbing glass filter (Callflex, Optic Balzers, Liechtenstein). The leaves were then dark adapted for 30 min using aluminium foil. \( F_0 \) and \( F_m \) were measured and the maximum photochemical efficiency of photosystem II (PSII) was calculated as \( F_v/F_m = (F_m - F_o)/F_m \). Subsequently, the leaf was exposed to actinic PAR of 302 or 1860 μmol m⁻² s⁻¹ for 10 min to achieve steady-state \( F_s \) and \( F_m' \), measured by 0.6 s saturating pulses. The operation efficiency of PSII (\( F_m' / F_m' \)), where \( F_m' = F_m - F_o \), the fraction of open PSII expressed as qL, and non-photochemical quenching (NPQ) were calculated as reviewed by Murchie and Lawson (2013). For each experiment, three plants per treatment were measured (one from each of the three replicated treatment OTFB boxes with accompanying control boxes) and in total three independent experiments were performed.

Biochemical analysis
Upper surface chlorophyll content in the first true leaf (on days 1, 4, 8, and 14 of UV exposure), and the youngest well-developed leaf on day 15 of UV exposure was measured using a DUALEX (Force-A, Orsay, France), following chlorophyll fluorescence measurements.

LAECF in the second true leaf (on days 0, 1, 3, 5, 10, and 14 of UV exposure) was measured using a DUALEX (Force-A) with \( n = 15–18 \) for the UV-enriched treatments and \( n = 30–36 \) for the control treatment. Additionally, total UV-absorbing pigments (TUAP; mostly flavonoids) were extracted from the second true leaf (on days 0, 1, 3, 5, 10, and 14 of UV exposure) for quantification. Leaves were snap frozen, then stored at -80°C until used. They were then ground in liquid nitrogen using a mortar and pestle and 0.1 g leaf material was placed into micro-tubes with 1 mL acidified methanol (1% HCl, 20% H₂O, 79% CH₃OH (v/v)) before incubation in the dark at 4°C for 4 days. Absorbance was recorded at 330 nm using a spectrophotometer (Shimadzu UV/VIS 1800). Absorbance was normalized per leaf fresh weight. For each experiment, three plants per treatment were measured (one from each of the three replicated treatment OTFB boxes with accompanying control boxes) and in total three independent experiments were performed, that is \( n = 9 \). For control samples, \( n = 18 \).

TAC was analyzed using a commercially available kit (TAC Assay kit, Sigma-MAK187). Ground leaf tissue (0.1 g; see above) from the second true leaf was extracted in 1 mL of ice cold 1 × phosphate-buffered saline (PBS) and following centrifugation, the supernatant was diluted 1:100 to bring absorbances of diluted extracts at 570 nm with Trolox standards and values normalized to tissue fresh weight. \( n = 3 \) for the UV-enriched treatments and \( n = 6 \) for the control treatment.

DNA damage detection
With replications as for flavonoid analysis, cyclobutane pyrimidine dimers were quantified using an immunoassay following the protocol from van de Poll et al. (2001). First, DNA was extracted from ground leaf tissue (see above) from the second true leaf from the base of the plant on days 1, 3, 5, 10, and 14 using the E.Z.N.A. Plant DNA Kit (Omega Bio-Tek, GA, USA), and dissolved in 100 μL TE buffer (pH 8.0). The DNA concentration was determined fluorometrically in a microplate reader (GENius, Tecan, Salzburg, Austria) using the Quantifluor dye (Promega, Madison, WI, USA). Of each sample, 50 ng DNA was used for the Southern blot. CPDs were subsequently labeled on the membrane by using a primary antibody against CPDs produced in mouse (H3 clone 4F6, Sigma Aldrich, St. Louis,
MO, USA). Detection was conducted by peroxidase coupled to the secondary antibody (Anti-Mouse IgG (whole molecule)-Peroxidase, Sigma Aldrich) using an enhanced chemiluminescent substrate (Pierce ECL, Thermo Fisher Scientific, Waltham, MA, USA). On each blot, a CPD calibration standard was included to allow absolute quantification of CPDs Mb\(^{-1}\) (Pescheck et al., 2014).

**Plant hormone analysis**

The second true leaves from the base of the plant were harvested on days 3 and 5. Leaves from three different plants within one experiment were pooled to obtain approximately 300 mg material. Leaf tissues were snap frozen in liquid nitrogen and kept at \(-80^\circ\text{C}\). All samples were then ground with a mortar and pestle in liquid nitrogen and again kept at \(-80^\circ\text{C}\) until used. In total, three independent experiments generated three replicates. Each pooled sample was subdivided in separate 100 mg fractions for the extraction of the different hormone groups.

**Auxin and ABA analysis**

Samples were extracted in 500 \(\mu\text{l}\) of 80% v/v methanol. [\(^{13}\)C]-IAA (100 pmol, (phenyl-\(^{13}\)C6)-indole-3-acetic acid, 99%, Cambridge Isotopes, Tewksbury, MA, USA) and D6-ABA (150 pmol, \(^{[4]}\text{H}_6\)+-cis, trans-abscisic acid, [(S)-5-\(^{[4]}\text{H}_6\)](1-hydroxy-2,6,6-trimethyl-4-oxocyclohex-2-en-1-yl)-3-methyl-(2Z,4E)-pentadienoic acid), Olchemim, Olomouc, Czech Republic) were added as internal tracers. After overnight extraction, samples were centrifuged (20 min, 15,000 g, 4\(^\circ\)C, in an Eppendorf 5810R centrifuge, Eppendorf, Hamburg, Germany) and the supernatants were aliquoted in two equal parts. One aliquot was acidified using 5.0 mL of 6.0% v/v formic acid and loaded on a reversed-phase (RP)-C18 cartridge (500 mg, BondElut Varian, Middleton, The Netherlands). The compounds of interest (IAA, ABA, and the oxidation products IAA-OX, IAA-OH, IBA-OX, and IBA-OH) were eluted with 5.0 mL of diethyl ether and dried under a nitrogen stream (TurboVap LV Evaporator, Zymark, New Boston, MA, USA). The remaining aliquot was hydrolyzed in 7.0 M NaOH for 3 h at 100\(^\circ\)C under a water-saturated nitrogen atmosphere. After hydrolysis, the samples were acidified using 2.0 M HCl, desalted on an RP-C18 cartridge (500 mg), and eluted with diethyl ether. All samples were methylated using ethereal diazomethane to improve analysis sensitivity. Samples were analyzed using an Acquity UPLC system linked to a TQD triple quadrupole detector (Waters, Milford, MA, USA) equipped with an electrospray interface in positive mode. Samples (6.0 \(\mu\text{l}\)) were injected on an Acquity UPLC BEH C18 RP column (1.7 \(\mu\text{m}, 2.1 \times 50 \text{ mm, Waters}) using a column temperature of 30\(^\circ\)C and eluted at 450 \(\mu\text{l}\) min\(^{-1}\) with the following gradient of 0.1% v/v formic acid in water (solvent A) and 0.1% v/v formic acid in acetoniitrile (solvent B): 0 – 0.8 min isocratic 92/8 A/B; 0.8 – 5 min linear gradient to 60/40 A/B; and 5 – 5.5 min linear gradient to 10/90 A/B. Quantitative analysis was performed by multiple reactant monitoring of selected transitions based on the MH\(^{+}\) ion (dwell time 0.02 s) and the most appropriate compound-specific product ions in combination with the compound-specific cone and collision settings. All data were processed using Masslynx/Quanlynx software V4.1 (Waters). Data are expressed in picomoles per gram fresh weight (pmol g\(^{-1}\) FW).

**Statistical analysis**

Statistical analysis was performed using either SPSS 19.0 (IBM, Armonk, NY, USA), STATA 14.0 or Wizard for Macintosh (App Store, Apple Inc., Cupertino, CA, USA). Morphological parameters (Figures 1–5) were analyzed using error propagation where the standard deviations of the ratios were approximated using Taylor linearization (Taylor, 1997) as further described in Qian et al. (2020), and tests of differences between means due to treatment (UV-A, UV-B, or control) were performed using analysis of variance (ANOVA). Paired \(T\) tests were used to test for changes in length of first–seventh petioles and parameters related to the second–fourth true leaves (Figures 2–4 and Tables 1, 2). Paired \(T\) test was also used for evaluating significant changes in each of LAEFC, TUAP, and TAC (Figure 8 and Supplementary Table S1). For data generated at different time periods of UV exposure, including analysis of chlorophyll fluorescence, chlorophyll content of the first true leaf,
and DNA dimer, two-way ANOVA was performed to test whether each variable was significantly affected by treatment or time of UV exposure. For data including chlorophyll fluorescence and chlorophyll content of the youngest well-developed leaf, and plant hormone concentration, T tests were performed to analyze if the differences between UV-exposed and control samples were significant or not.

To describe the relationship between the LAEFC and TUAP measurements, a simple regression model was first fitted. Thereafter, additional explanatory variables giving number of days of treatment/leaf age and two dummy variables taking into account treatment (UV-A-enriched, UV-B-enriched, and control) were included, to see if they would contribute in describing the dependent variable LAEFC. The final model included the TUAP variable and the variable days of treatment as explanatory variables. The residuals were analyzed and the assumptions behind the model seemed to be fulfilled, and showed no signs of systematic pattern, supporting the choice of model.

The same approach was used when TAC was the dependent variable and as explanatory variables in the full model; TUAP, days of treatment, two dummy variables for the treatment (UV-A, UV-B, and control). The final model included the same explanatory variables as the first model, TUAP and days of treatment, the other explanatory variables were not significant, thereby not contributing to the explanation of the values of TAC. The analysis of the residuals did not show any indication on deviations from the model assumptions.

Supplemental data
The following materials are available in the online version of this article.

Supplemental Table S1. Pairwise T test for significance of treatment of cucumber seedlings

Supplemental Figure S1. Maximum photochemical efficiency (Fv/Fm) measured in cucumber plants grown under different light conditions.

Supplemental Figure S2. Time courses of photosynthetic parameters measured in cucumber plants grown under different light conditions.

Supplemental Figure S3. The monophasic relationship between LAEFC and TUAP.

Supplemental Figure S4. Residual (e) plots for models of correlation between the three methods used to determine LAEFC, TUAP, and TAC.

Supplemental Figure S5. Average number of flowers in each leaf node in cucumber plants that had been exposed to UV-A-enriched, UV-B-enriched non-UV supplemented growth light as seedlings.

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