Light-limited photosynthesis under energy-saving film decreases eggplant yield

Sachin G. Chavan¹ | Chelsea Maier¹ | Yagiz Alagoz¹ | Joao C. Filipe² | Charles R. Warren³ | Han Lin⁴ | Baohua Jia⁴ | Michael E. Loik⁵ | Christopher I. Cazzonelli¹ | Zhonghua H. Chen¹ | Oula Ghannoum¹ | David T. Tissue¹

¹National Vegetable Protected Cropping Centre, Hawkesbury Institute for the Environment, Western Sydney University, Penrith, NSW, Australia
²Environmental and Conservation Sciences, Murdoch University, Murdoch, WA, Australia
³University of Sydney, Camperdown, NSW, Australia
⁴Faculty of Science, Engineering and Technology, Centre for Translational Atomaterials, Swinburne University of Technology, Hawthorn, Vic., Australia
⁵Department of Environmental Studies, University of California, Santa Cruz, CA, USA

Correspondence
David T. Tissue, National Vegetable Protected Cropping Centre, Hawkesbury Institute for the Environment, Western Sydney University, Locked Bag 1797, Penrith, NSW 2751, Australia. Email: D.Tissue@westernsydney.edu.au

Funding information
Horticulture Innovation Australia projects, Grant/Award Number: VG16070 and VG17003; Australian Indian Institute (AII) New Generation Network (NGN)

Abstract
Glasshouse films with adjustable light transmittance and energy-efficient designs have the potential to reduce (up to 80%) the high energy cost for greenhouse horticulture operations. Whether these films compromise the quantity and quality of light transmission for photosynthesis and crop yield remains unclear. A “Smart Glass” film ULR-80 (SG) was applied to a high-tech greenhouse horticulture facility, and two experimental trials were conducted by growing eggplant (Solanum melongena) using commercial vertical cultivation and management practices. SG blocked 85% of ultraviolet (UV), 58% of far-red, and 26% of red light, leading to an overall reduction of 19% in photosynthetically active radiation (PAR, 380–699 nm) and a 25% reduction in total season fruit yield. There was a 53% (season mean) reduction in net short-wave radiation (radiometer range, 385–2,105 nm upward; 295–2,685 nm downward) that generated a net reduction of 8% in heat load and reduced water and nutrient consumption by 18%, leading to improved energy and resource use efficiency. Eggplant adjusted to the altered SG light environment via decreased maximum light-saturated photosynthetic rates ($A_{max}$) and lower xanthophyll de-epoxidation state. The shift in light characteristics under SG led to reduced photosynthesis, which may have reduced source (leaf) to sink (fruit) carbon distribution, increased fruit abortion and decreased fruit yield, but did not affect nutritional quality. We conclude that SG increases energy and resource use efficiency, without affecting fruit quality, but the reduction in photosynthesis and eggplant yield is high. The solution is to re-engineer the SG to increase penetration of UV and PAR, while maintaining blockage of greenhouse heat gain.

KEYWORDS
eggplant, energy, light, photosynthesis, protected cropping, smart glass
most of the wavelengths required by plants for photosynthesis. Glazing and/or the application of films can change light intensity and spectral quality, thereby having an adverse effect on plant growth, photosynthesis, biomass partitioning, yield, and quality (Hao & Papadopoulos, 1999; Loik et al., 2017). Theoretically, blocking radiation not required for photosynthesis can decrease heat build-up in the glasshouse and hence reduce the energy cost required to maintain cooling in summer. However, this theory of photonics and material science still has not been properly tested in a high-tech greenhouse with a commercial horticultural crop over two seasons.

Plants respond to light intensity, spectral quality, and photoperiod (Babla et al., 2019; Ballaré & Pierik, 2017; Cazzonelli et al., 2020; Poorter et al., 2019). At the leaf level, blue photons are used less efficiently than orange and red photons in photosynthesis (Bugbee, 2016; Inada, 1976; McCree, 1971). The change in spectral quality, especially the ratio of red to far-red light, can affect plant phenology and development of buds, flowers, and fruits (Ballaré & Pierik, 2017; Cerdán & Chory, 2003). Plants cope with light fluctuations via adjustments at the whole organism, cellular, biochemical and molecular levels (Ruban, 2009). The light energy absorbed by pigments in the photosystems is used to drive chemical reactions for photosynthesis, and dissipate excessive light energy from photosystem II (PSII) via chlorophyll $a$ fluorescence and by several other thermal dissipation mechanisms (Baker, 2008; Logan, Adams, & Demmig-Adams, 2007). Photosystems I and II are composed of varying amounts of Chl $a$, Chl $b$, $\beta$-carotene, and xanthophylls (lutein, antheraxanthin, violaxanthin, and neoxanthin) which facilitate quenching of excess PSII energy. Carotenoid pigments play an important function in facilitating photosynthesis and photoprotection, thereby contributing to an optimal carbon balance from source (leaf) to sink (fruit) (Baranski & Cazzonelli, 2016; Demmig-Adams, Garab, Adams, and Govindjee, 2014). A reduction in photosynthesis will lower the supply of carbon in source leaves and carbohydrate translocation to sinks such as fruits, thereby affecting fruit set (Aloni, Karni, Zaidman, & Schafer, 1996; Turner & Wien, 1994). Limitations in photosynthesis can decrease crop yield and quality (Hao & Papadopoulos, 1999), depending on the light environment.

The primary objective of this study was to determine the impact of SG on light quality and quantity, and subsequently on photosynthetic carbon assimilation, leaf biochemistry, yield, and nutritional quality of eggplant (*Solanum melongena*) using a high-tech glasshouse facility. We used standard management practices during two greenhouse trials on a commercial eggplant cultivar (cv Tracey) to assess the efficacy of SG on reducing resource use while minimizing and growth. In addition, novel materials with insulation properties trap heat during winter and save energy on heating. SG could significantly contribute to reducing the energy costs in greenhouse operations.
negative impacts on crop yield and quality with the following key hypotheses that SG will: (a) block a significant amount of biologically nonuseful radiation that contributes to heat generation, consequently saving energy on cooling the greenhouse; (b) decrease PAR by a small amount (5%–10%) which will not significantly affect yield; and (c) affect physiological and biochemical characteristics of leaves and fruits.

2 | MATERIALS AND METHODS

2.1 | Facility description and glass specifications

The first SG trial was conducted in the state-of-the-art glasshouse facility designed for research and commercial production of horticultural crops at Western Sydney University, NSW, Australia (Figure S1). The 1,800 m² advanced glasshouse facility established in late 2017 is equipped with Priva software and hardware (Priva) to monitor and control temperature, humidity, nutrients, CO₂, and irrigation. Glasshouse air temperature is controlled by chilled air blowers, curtains, and opening vents. Relative humidity (RH) is controlled using a humidification system, and air temperature is partially controlled using hot water circulation through radiant pipes. We used four 105-m² research compartments with precise environmental control of atmospheric CO₂, air temperature, RH, and hydroponic nutrient and water delivery. Each research compartment included 6 gutters, used to deliver nutrients and water, which support 120–150 plants.

Two research compartments were fitted with HD1AR diffuse glass (70% haze; control compartments) and two research compartments had HD1AR diffuse glass, but were also coated with ULR-80 window film (Solar Gard; Saint-Gobain Performance Plastics). The SG film ULR-80 (Table S1) is a potentially suitable glazing material for greenhouse crop production. It has low thermal emissivity (0.87) which blocks the light that mainly contributes to heat, but transmits most of the wavelengths of light used by plants for growth in the PAR region. According to the manufacturer specifications, SG blocks ~88% light in the infrared (IR) and far-infrared (FIR) region between 780 and 2,500 nm; and >99% light in the ultraviolet (UV) region between 300 and 400 nm. In addition, SG blocks 43% of total solar energy with 40% transmission, 54% absorption, and 6% reflectance. The two control research compartments consist of roof glass (70% diffuse light) and wall glass (5% diffuse light) (Table S1).

2.2 | Plant growth and management

*Solanum melongena* (cv. Tracey eggplant grafted on tomato cv. Kaiser stems) was the first horticulture crop tested under the SG for two experiments. The first experiment was conducted from January 2018 (Australian mid-summer) through autumn to July 2018 (winter), and the second experiment was conducted from September 2018 (early spring) through summer to March 2019 (early autumn). For each experiment, 6-week-old nursery-grown seedlings were transplanted in Rockwool slabs and transferred into two control hazed glass (Control) and two SG (Treatment) compartments. Each compartment had six gutters (length 10.8 m, width 25 cm; AIS Greenworks, Castle Hill) with 10 Rockwool slabs (90 × 15 × 10 cm; Grodan) per gutter. Three plants per slab were planted in the four middle gutters, and two plants per slab were planted in the two side gutters and served as buffer plants. A total of 160 plants were grown in each chamber, but all measurements were performed on the 120 plants grown in the four middle gutters to avoid edge effects. Plants were grown at standard growth conditions under natural light (as described in Table S2 and Figure S2) and were provided nonlimiting nutrients and water by the Priva computer-programmed fertigation (nutrients and water) system. Three stems were selected to grow from each plant with weekly pruning and cutting according to commercial practices of eggplant production for vertical protected cultivation. Each stem was considered as an individual plant for replication, and all measurements were performed per stem.

2.3 | Light environment measurements

Light quality and quantity were measured using a portable spectroradiometer (PS300; Apogee Instruments, Inc.) and a PAR sensor (LI-190SZ Quantum Sensor; LI-COR) at the roof level during both experimental trials. Except for the spectroradiometer, all other sensors continually logged data providing output as 5-min averages. Additional sensors including hobo pendant temp/light data logger (UA-002-08; Instrument Choice), PAR (LI-190R-SMV-50 Quantum Sensor; LI-COR), net radiometer (SN-500; Apogee Instruments), and diffuse light sensor (BF5 sunshine sensor, Delta T Devices) were deployed to measure detailed light profiles during the second experimental trial. Three hobo pendant temp/light data loggers (at the base, middle, and top positions of the canopy), 5 PAR sensors (at canopy level), and a net radiometer were installed in each chamber. The diffuse light sensors were installed in one control and one SG chamber.

2.4 | Energy and nutrient savings calculations

The Priva system continuously records energy expenditure on cooling (kW) using water flow, the temperature of the water before entering the chiller, and after exiting the chiller. Each of the research compartments was cooled via two 1.2 kW
Fan Coil Units (FCUs). Chilled water, from one of the two 75 kW chillers, is supplied in a closed loop to each of the two FCUs in each room. The chilled water flows through these two units and is then returned to the 200,000 L storage tank. Priva records the supply and the return temperature of chilled water in each room. The meters do not measure the actual energy in kWh, unlike a meter for electricity, but can be used to calculate an energy value based on three variables: (a) water flowing through the flow meter; (b) temperature of the supply chilled water; and (c) temperature of the return chilled water. It does not record the ON/OFF of the FCUs, but if it reads a significant difference in the temperature of the supply and return, it sends a pulse to Priva. All data are based on the same reading, which allows us to directly use these numbers to determine the energy consumption. The Priva system also continuously records fertilizer and water supply to the irrigation system, and the irrigation water that subsequently enters the drainage system. The net consumption of fertilizer and water is determined using supply and drain values.

2.5 | Plant growth and productivity measurements

Plant growth and yield parameters were measured periodically in both experimental trials. Replication \( n \) refers to the total number of plants in two control or two SG chambers. Height was measured 79, 95, 109, 121, and 137 days after planting (DAP) during Experiment 1 \( (n = 120, 60 \text{ stems per chamber}) \) and 111, 125, 140, and 155 DAP in Experiment 2 \( (n = 24, 12 \text{ stems per chamber}) \). Bud, flower, and fruit number were measured 164, 171, and 178 DAP during Experiment 1 \( (n = 72, 36 \text{ stems per chamber}) \) and 84, 98, 110, 117, 131, and 146 DAP during Experiment 2 \( (n = 36, 18 \text{ per chamber, respectively}) \). Bud, flower, and fruit development were tracked weekly to test the rate of development of selected tagged buds until plants attained full development to the fruit stage and harvest. Twelve weeks after planting, eggplant fruits (only those between 350–450 g, representing commercial harvest mass) were harvested weekly for 18 and 16 weeks during Experiment 1 \( (n = 360, 180 \text{ stems per chamber}) \) and Experiment 2 \( (n = 240, 120 \text{ stems per chamber}) \), respectively. The weight of individual eggplant fruit (between 350–450 g) and the number of fruits per stem were recorded. Pruned biomass per chamber was weighed at 5 time points (62, 75, 85, 90, and 96 DAP) in Experiment 2.

2.6 | Leaf gas exchange measurements

Instantaneous steady-state leaf gas exchange measurements \( (n > 15) \) were performed using a portable, open-mode gas exchange system (LI-6400XT; LI-COR). Measurements were performed at 1,500 \( \mu \text{mol m}^{-2} \text{s}^{-1} \) PAR with two \( \text{CO}_2 \) concentrations (400 \( \mu \text{l/L} \) during Experiment 1 and 500 \( \mu \text{l/L} \) during Experiment 2) and 25°C leaf temperature. The response of \( A_{\text{sat}} \) to light \( (Q) \) (A–Q curve) was measured at 25°C leaf temperature at eight light levels \( (0, 50, 100, 250, 500, 1,000, 1,500, \text{and } 2,000 \mu \text{mol m}^{-2} \text{s}^{-1}) \) in Experiment 1 \( (n > 8) \), and 11 steps of light levels \( (0, 25, 50, 100, 200, 300, 400, 500, 1,000, 1,500, \text{and } 2,000 \mu \text{mol m}^{-2} \text{s}^{-1}) \) in Experiment 2 \( (n > 18) \). The response of \( A_{\text{sat}} \) to substomatal \( \text{CO}_2 \) mole fraction \( (C_i) \) (A–C\(_i\) response curve) was measured in eight steps of \( \text{CO}_2 \) concentrations \( (50, 100, 230, 330, 420, 650, 1,200, \text{and } 1,800 \mu \text{l/L}) \) at 25°C leaf temperature during Experiment 1. Spot measurements at 25°C leaf temperature and 500 \( \mu \text{l/L} \) \( \text{CO}_2 \) were also performed during Experiment 2 \( (n > 4) \) using the clear leaf cuvette under natural light conditions. The light response curve means were fitted using the following equation (Ögren & Evans, 1993; Xu et al., 2019).

\[
A = \frac{(\phi_{\text{max}} \cdot I + A_{\text{max}}) - \sqrt{(\phi_{\text{max}} \cdot I + A_{\text{max}})^2 - 4 \cdot \theta \cdot \phi_{\text{max}} \cdot I \cdot A_{\text{max}}}}{2 \cdot \theta} - R_d
\]

where, \( I = \) absorbed irradiance, we assumed absorptance = 0.85; \( A = \text{CO}_2 \) assimilation rate at given light; \( R_d = \) dark respiration; \( \phi_{\text{max}} = \) maximum quantum yield of PSII; \( A_{\text{max}} = \) maximum light-saturated \( \text{CO}_2 \) assimilation rate; and \( \theta = \) curvature factor of the light response curve.

2.7 | Spectral analysis of leaves using a spectroradiometer

Leaf reflectance was collected using an ASD spectroradiometer (FieldSpec 4, Malvern Panalytical Ltd) with a spectral range of 350–2,500 nm. The sensor has a sampling interval of 1.4 and 1.1 nm for 350–1,000 nm and 1,001–2,500 nm regions, respectively. Fully expanded leaves of eggplants were collected from the plant’s middle canopy from the four chambers of the glasshouse; measurements were taken with the aid of a leaf clip attached to a plant probe over a 3-hr period (9 a.m. to noon). The leaf clip allows the leaf to touch plant probe and keep the light beam at an angle of 45°. Reflectance spectral values were developed from the conversion of spectra by referencing a 99% Spectralon calibration panel (Labsphere, Inc.). A reference measurement of the calibration panel was taken before the first measurement and every 30 min onwards. For each leaf, four measurements were taken from six different spots. Spectral index values were estimated for each leaf using the mean of these 24 measurements. Spectral indices, including Water Band Index (WBI) for leaf water content (Peñuelas, Llusia, Piñol, & Filella, 1997), modified Normalized Difference Vegetation Index (mNDVI) for chlorophyll content (Fuentes, Gamon, Qui, Sims, & Roberts, 2001), Photochemical Reflectance Index (PRI) for xanthophyll cycle pigments (Gamon, Peñuelas, & Field, 1992), red–green ratio (RGR) for anthocyanin content (Fuentes et al., 2001), Structure
Intensive Pigment Index (SIPI) for carotenoid-to-chlorophyll a ratio (Peñuelas, Baret, & Filella, 1995), red–far-red ratio (RFR) (Mascarini, Lorenzo, & Vilella, 2006), and Normalized Phaeophytinization Index (NPQI) for chlorophyll degradation (Barnes, Balaguer, Manrique, Elvira, & Davison, 1992), were calculated as follows,

\[ WBI = \frac{R_{500}}{R_{700}} \]

\[ mNDVI = \frac{R_{750} - R_{305}}{R_{750} + R_{305}} \]

\[ PRI = \frac{R_{531} - R_{570}}{R_{531} + R_{570}} \]

\[ RGR = \frac{\sum_{n=699}^{n=600} R_n}{\sum_{n=599}^{n=500} R_n} \]

\[ SIPI = \frac{R_{800} - R_{445}}{R_{800} - R_{680}} \]

\[ RFR = \frac{R_{680}}{R_{730}} \]

\[ NPQI = \frac{R_{415} - R_{435}}{R_{415} + R_{435}} \]

2.8 | SPAD measurements and leaf pigment analysis using high-performance liquid chromatography

One leaf per plant from five different plants per chamber was used for SPAD measurements and then for pigment analysis using high-performance liquid chromatography (HPLC) and gas chromatography coupled with mass spectrometry (GCMS). Three leaf disks were punched from the top, middle and bottom position of a fully expanded mature leaf using a size 10 (2.54 cm² leaf area) cork borer in the morning hours between 10 a.m. and 12 p.m. Samples were snap-frozen using liquid nitrogen and kept at −80°C until further analysis. Fresh leaf weight was measured to calculate leaf mass per unit area (LMA) and for quantification of carotenoids and pigments. For both control and treatment, ten biological replicates were collected, frozen in the liquid N₂, and ground to a fine powder with TissueLyser (Qiagen). Carotenoids were extracted under low-light conditions with 500 µl extraction buffer (60% v/v ethyl acetate:40% v/v acetone and 0.1% BHT) and partitioned into the ethyl acetate layer by adding 500 µl of H₂O. The carotenoid-containing organic phase was separated via centrifugation and analyzed by reverse-phase HPLC (Agilent 1200 Series) using GraceSmart-C18 (4-µm, 4.6 × 250-mm column; Alltech) column. HPLC runs were performed as previously described (Alagoz, Dhami, Mitchell, & Cazzonelli, 2020). Pigments were identified based upon their specific retention time (RT) relative to known standards and their spectral characteristics at 440 nm (lutein—L, β-carotene—β, antheraxanthin—A, zeaxanthin—Z, neoxanthin—N, violaxanthin—V, and chlorophylls) and 286 nm (phytoene). Carotenoid quantification was performed as previously described except cis-carotene phytoene (Pogson, McDonald, Truong, Britton, & DellaPenna, 1996). Phytoene is quantified by using its molar extinction coefficient and molecular weight to convert the peak area in micrograms per gram fresh weight (µg/g FW) as previously described (Britton, Liaab-Jensen, & Pfander, 1995). All pigments were quantified at absorption wavelengths with maximum detection. The de-epoxidation state (DPS) of the xanthophyll cycle was calculated as DPS = (A + Z)/(A + Z + V).

2.9 | Leaf metabolite analysis using gas chromatography-mass spectrometry

One leaf disk (from the middle position per leaf) was used for metabolite profiling using gas chromatography–mass spectrometry (GCMS). Each leaf disk was extracted using methanol/chloroform/water (700/400/800 = 1,100 µl aqueous phase) by grinding with sand, followed by phase separation. A 200 µl aliquot of the aqueous phase of the extract was dried, and 50.0 µl 20 µg/ml ribitol was added followed by redrying for 3 hr. Finally, the extract was derivatized with 40 µl MOX followed by 60 µl MSTFA before analysis by GC-MS as described previously (Lisec, Schauer, Kopka, Willmitzer, & Fernie, 2006). Peaks were aligned and retention indices calculated against alkanes (Kovat’s RI). Peak picking, deconvolution, and ID were performed with MS-DIAL (Tsugawa et al., 2015) using generic GC-MS parameters, and MSP file for 15,302 entries of metabolites with Kovat’s RI. The data matrix was manually edited to verify IDs and remove deconvolution errors. Ions with m/z 73 or 147 were not used as quantification ions.

2.10 | Statistics and data analysis

Data analyses and plotting were performed using R computer software (R Core Team, 2019). The treatment effect was analyzed using one-way analysis of variance (ANOVA). The linear model involved testing of each parameter over two treatment conditions, SG and Control glass, using measurements from two SG and two control glass rooms. Replication, for example, n = 10, refers to 10 plants/stems per treatment or 5 plants/stems from each chamber. The homogeneity of variance was tested using Levene’s test from the car package. The parameters showing unequal variance (with less than .05
probability for Levene's test) were corrected using Welch's t-test for unequal variances using the oneway.test function in R. Other packages were also used, including (but not limited to) lubridate (for effective use of dates in plots), sciplot (for plotting), and doby (for calculating means and standard errors). For GCMS data analysis, unpaired t tests were used for univariate comparisons of metabolite concentrations, with p-values corrected to account for the false discovery rate due to multiple comparisons (Benjamini and Hochberg, 1995) using an Excel spreadsheet (Pike, 2011). The significance levels for ANOVA were $p > .05 = \text{ns}$, $* p < .05$, $** p < .01$, and $*** p < .001$.

3 | RESULTS

3.1 | SG blocks UV and light wavelengths $>800$ nm and significantly reduces PAR

Spectroradiometer measurements validated manufacturer SG specifications, including blockage of UV and infrared (Table S1 and Figure 1). Although a modest reduction ($-5\%$ to $-10\%$) in overall light transmission was expected, a considerable amount of PAR was blocked by SG with higher reduction in red light (600–750 nm) relative to blue or green light (Table 1, Figure 1). SG blocked most of the UV (221–279 nm, $-85\%$), and a considerable amount of red (600–699 nm, $-26\%$) and far-red (710–850 nm, $-58\%$), with an overall reduction of $-19\%$ PAR integrated from 280 to 799 nm (Table 1 and Figure 1). Thus, SG changed both the quantity and quality of the light spectrum.

Daily light integral (DLI) measured using a PAR sensor at roof level in each room was significantly reduced ($-24\%$ and $-28\%$ during experiments 1 and 2, respectively) under SG relative to control (Table 1 and Figure 1). The reduction in DLI measured at canopy level ($-21\%$ in SG relative to Control) was relatively lower than roof level reduction ($-28\%$ in SG relative to Control) in DLI during Experiment 2 (experiment mean). In addition, the proportion of diffuse light measured using a diffuse light sensor was $-25\%$ lower in SG relative to Control (Table 1). Short-wave radiation (385–2,105 nm

![Figure 1](https://example.com/fig1.png)
upward; 295–2,685 nm downward), which mostly contributes to heat generation in the glasshouse, was measured during Experiment 2 using a net radiometer and was reduced by −53% under SG (Table 1, Figure 2). The blocked short-wave radiation consequently reduced energy expenditure on cooling (−8%) by chillers and net fertigation (fertilizer + water) consumption (−18%) under SG relative to Control (Figure 2). In addition, the visible light intensity measured in lux by the hobo pendant temp/light data logger showed significant reduction in daily average light measured at the top (−56%), middle (−70%), and bottom (−67%) of the canopy (Table S3 and Figure 3). Thus, SG blocks most of the heat-generating

### TABLE 1 Summary of radiation, light intensity, canopy temperature, leaf gas exchange, and leaf mass area (LMA) measurements

| Parameter (mean) | Exp | Treatment | Change (%) | p-Value |
|------------------|-----|-----------|------------|---------|
| **Radiation parameters** | | | | |
| SW radiation (kWh m⁻² day⁻¹) | 2 | Control | 10,576 ± 302 | 4,970 ± 142 | −53 | 2.2 × 10⁻¹⁶ |
| Diffuse light (kWh m⁻² day⁻¹) | 2 | Control | 6,743 ± 149 | 5,037 ± 112 | −25 | 2.2 × 10⁻¹⁶ |
| DLI at roof (mol⁻¹ m⁻² day⁻¹) | 1 | Control | 11.3 ± 0.2 | 8.5 ± 0.1 | −24 | 2.2 × 10⁻¹⁶ |
| DLI at roof (mol⁻¹ m⁻² day⁻¹) | 2 | Control | 23.5 ± 0.6 | 16.9 ± 0.4 | −28 | 2.2 × 10⁻¹⁶ |
| DLI at canopy (mol⁻¹ m⁻² day⁻¹) | 2 | Control | 14.7 ± 0.4 | 11.6 ± 0.3 | −21 | 6.9 × 10⁻⁹ |
| **Light spectrum measurements using spectroradiometer** | | | | |
| UV (221–279 nm, µmol m⁻² s⁻¹) | 2 | Control | 21.7 ± 2.3 | 3.3 ± 0.6 | −84 | 3.2 × 10⁻⁷ |
| Blue (280–499 nm, µmol m⁻² s⁻¹) | 2 | Control | 212 ± 15 | 179 ± 4 | −15 | .052 |
| Green (500–599 nm, µmol m⁻² s⁻¹) | 2 | Control | 279 ± 20 | 236 ± 6 | −15 | .056 |
| Red (600–699 nm, µmol m⁻² s⁻¹) | 2 | Control | 298 ± 27 | 219 ± 7 | −26 | .01 |
| PAR (380–699 nm, µmol m⁻² s⁻¹) | 2 | Control | 792 ± 63 | 638 ± 17 | −19 | .02 |
| Far-red (719–850 nm, µmol m⁻² s⁻¹) | 2 | Control | 309 ± 39 | 128 ± 9 | −58 | .0002 |
| **Gas exchange parameters under saturated light** | | | | |
| A sat (µmol m⁻² s⁻¹) | 1 | Control | 25.2 ± 0.5 | 22.1 ± 0.6 | −12 | .0009 |
| 2 | Control | 29.4 ± 0.6 | 24.1 ± 0.5 | −18 | 2.7 × 10⁻⁷ |
| g s (mol m⁻² s⁻¹) | 1 | Control | 0.66 ± 0.05 | 0.50 ± 0.04 | −24 | .01 |
| 2 | Control | 0.49 ± 0.03 | 0.48 ± 0.04 | −2 | .9 |
| C i (µl/L) | 1 | Control | 285 ± 3 | 276 ± 4 | −3 | .1 |
| 2 | Control | 335 ± 5 | 357 ± 6 | +6 | .02 |
| PWUE (A/g s) | 1 | Control | 40 ± 3 | 47 ± 3 | +10 | .1 |
| 2 | Control | 64 ± 3 | 56 ± 4 | −12 | .1 |
| R d (µmol m⁻² s⁻¹) | 2 | Control | −2.46 ± 0.07 | −2.11 ± 0.08 | −14 | .003 |
| **Gas exchange parameters under natural growth light** | | | | |
| PARi (µmol m⁻² s⁻¹) | 2 | Control | 1,406 ± 16 | 1,168 ± 16 | −17 | 2.2 × 10⁻¹⁶ |
| A gl (µmol m⁻² s⁻¹) | 2 | Control | 29.3 ± 0.3 | 23.1 ± 0.3 | −21 | 2.2 × 10⁻¹⁶ |
| g sgl (mol m⁻² s⁻¹) | 2 | Control | 0.38 ± 0.01 | 0.31 ± 0.01 | −18 | .0003 |
| PWUE gl (A gl/g sgl) | 2 | Control | 79 ± 2 | 77 ± 2 | −2 | .39 |
| **Light response curve modeled parameters** | | | | |
| A max (µmol m⁻² s⁻¹) | 2 | Control | 31.5 ± 0.7 | 24.6 ± 0.6 | −22 | 1.2 × 10⁻⁷ |
| Φ max (mol CO₂ mol⁻¹ quanta⁻¹) | 2 | Control | 0.039 ± 0.001 | 0.037 ± 0.002 | −5 | .3 |
| Θ (Dimensionless) | 2 | Control | 0.87 ± 0.01 | 0.85 ± 0.01 | −2 | .08 |

Note: One-way analysis of variance for the Smart Glass effect on radiation (n = 318) during two experiments (Exp) including daily total means for short wave (SW), diffused light, and daily light integral (DLI); light spectrum (n = 18, spectroradiometer measurements at nine locations per chamber) including UV, blue, green, red, PAR, and far-red light wavelengths; instantaneous leaf gas exchange (n > 15) including light-saturated CO₂ assimilation rates (A sat), stomatal conductance (g s) and intercellular CO₂ concentration (C i) and photosynthetic water use efficiency (PWUE); instantaneous gas exchange at natural growth light (n = 5) including average PAR measured using LI-6400 (PARi), CO₂ assimilation rates at growth light (A gl), stomatal conductance and growth light (g sgl) and photosynthetic water use efficiency at growth light (PWUE gl); and light response curve modeled parameters (n > 18) including maximum light-saturated CO₂ assimilation rates (A max), maximum quantum yield (Φ max), and curvature factor (Θ).
energy not required by plants, thereby saving energy on cooling and resource use. However, SG also considerably reduces PAR required for photosynthesis and growth.

3.2 | **SG reduces eggplant photosynthesis due to light limitation**

The impact of an altered light environment on photosynthesis was investigated by measuring instantaneous leaf gas exchange and light response curves. Altered light quality and quantity, including reduction in PAR, decreased light-saturated CO₂ assimilation rates (Aₘₐₓ) (−12% and −18% in experiments 1 and 2, respectively). However, stomatal conductance (gₛ) decreased (−24%) only in Experiment 1 (Table 1 and Figure 4). In Experiment 2, instantaneous leaf gas exchange measured at natural growth light levels (1,406 and 1,168 μmol m⁻² s⁻¹ mean PAR in control and SG, respectively) showed reductions in CO₂ assimilation rates (A) (−21%) and gₛ (−18%) under SG (Table 1 and Figure 4). Leaf level photosynthetic water use efficiency (PWUE) did not differ, either during light-saturated or ambient growth light conditions (Table 1). In addition, average daily canopy temperature measured at the top, middle, and bottom position of the canopy was reduced by 0.5–0.9°C (±0.05) under SG during Experiment 2 (Table S2 and Figure 3).

Based on AQ curves, photosynthetic rates were generally reduced under SG at higher light intensities in both experiments. Maximum light-saturated CO₂ assimilation rates (Aₘₐₓ) (−22%) were significantly reduced under SG in Experiment 2, while maximum quantum yield (Φₘₐₓ) and curvature factor (θ) were similar under both control and SG (Table 1 and Figure 4). The dark respiration (R_d) measured during light response curves was decreased by 14% under SG relative to control (Table 1). Therefore, reductions in PAR under SG caused light limitation and decreased photosynthesis, particularly at higher light levels, suggesting adaptive changes in the photosynthetic apparatus without changes in the photosynthetic efficiency.

3.3 | **Eggplant leaves grown under SG have an altered xanthophyll composition**

The composition and abundance of carotenoid pigments was quantified in top canopy leaves from control and SG grown plants. Downregulation of photosynthesis and Aₘₐₓ in low-light
conditions in SG was correlated with an altered pigment composition and spectral indices. There was a significant reduction in specific xanthophyll pigments (A, Z, V, and N), yet no change in lutein or β-carotene. Altered light under SG significantly reduced pool sizes of A (−26%), Z (−45%), and V (−18%). De-epoxidation state (DPS) was consequently lowered (−14%) in leaves from plants grown under SG. In addition, the photochemical reflectance index (PRI) was significantly increased (+8%) under SG, which is inversely proportional to the DPS (Gamon et al., 1992; Peñuelas, Filella, Lloret, Munoz, & Vilajeliu, 1995) (Table 2 and Figure 5). A lower structure intensive pigment index (SIPI), a measure of carotenoid-to-chlorophyll a ratio (Peñuelas, Baret, et al., 1995), was consistent with a lower carotenoid/chlorophyll ratio quantified by HPLC (−0.3% and −8%) in SG relative to Control leaves. There was a reduction in mNDVI (−2%) and SPAD values (−6%) that suggest that leaf chlorophyll content was slightly lower in SG grown plants. However, HPLC data showed no significant difference in chlorophyll content when measured per unit fresh weight (Table 2). Rather, a lower leaf water content evident from reduced WBI (−1%) (Peñuelas, Gamon, Griffin, & Field, 1993) and LMA (−9%) indicated that chlorophyll content was reduced per unit leaf area, but not per unit fresh weight (Table 2 and Figure 5). It is worth noting that the spectral indices and physical measurements do not usually commensurate with each other, given the different way they are measured and that the indices are “indicators” rather than direct estimates. Untargeted GCMS of polar metabolites resolved >200 features in leaves (Table S4). After FDR correction, peaks areas (i.e., concentration) of 13 metabolites differed significantly between SG and control (FDR-corrected t test). However, all of the significantly different metabolites were present at low concentrations and in no cases were fold-differences large. Therefore, leaves from plants grown under SG acclimated with an altered xanthophyll composition and DPS with minimal alteration in metabolite, total carotenoid, or chlorophyll levels (Table 2 and Figure 5).

3.4 | SG does not affect morphological features or fruit quality, but a high fruit abortion rate reduces yield

Plant morphological traits including height, bud, flower, and fruit number were analyzed in response to altered light environment under SG. Plants grown under SG had similar height, number of flowers, and number of buds (Table 3). However, mean fruit number (−28%, p-value < .001 and −23%, p-value < .001) and fruit weight (−32%, p-value < .001 and −24%, p-value < .001) were significantly reduced leading to decreased productivity under SG relative to the control.
A reduction in fruit number was attributed to increased abortion of flowers or fertilized young fruits (chi-square test, p-value < .01) under SG (Figure 6). In addition, the biomass harvested after pruning was lower (−17%) under SG relative to control (Table 3).

Fruit quality parameters, including pH, titratable acidity, moisture, total soluble solids (brix), mineral content (ash), elemental composition (AGVITA, Table S5), metabolites (GCMS, Table S6), sugar content (HPLC), fat (ANKOM), and nitrogen (DUMAS) content, were assessed. None of the >400 metabolites resolved by untargeted GC-MS differed significantly between SG and control (FDR-corrected t test). We found increases in total sugars (+8%), sucrose (+29%), and Fe (+28%), and decreases in mineral content (−9% and −28% in experiments 1 and 2, respectively), but otherwise parameters were unchanged (Table 3 and Table S3). In summary, SG did not affect eggplant morphological traits, but increased abortion rate in fertilized young fruits, thereby decreasing fruit yield without major changes in fruit quality.

4 | DISCUSSION

SG film ULR-80 blocked 85% of UV (221–279 nm), 26% of red (600–699 nm), and 58% far-red (710–850 nm) light with an overall reduction of 19% PAR (280–799 nm) and 53% reduction (season mean) in short-wave radiation (385–2,105 nm upward; 295–2,685 nm downward) measured using spectroradiometer. This consequently reduced energy expenditure for cooling and water and nutrient consumption. However, SG also reduced mean season PAR (DLI: −24% and −28% in experiments 1 and 2, respectively) leading to reductions in photosynthesis and hence productivity (mean fruit weight: −32 and −24% in experiments 1 and 2, respectively), and generally did not affect fruit quality except for significantly increasing the sweetness of the fruits. Growth under SG reduced $A_{\text{max}}$ and the xanthophyll cycle pigments (A, V, and Z) and DPS, thereby highlighting that SG grown plants may have partially acclimated to low-light conditions. Novel glazing materials with low thermal emissivity can be applied to greenhouses to reduce energy expenditure and resource use (water and nutrients), but specifically SG film ULR-80 will require spectral compositional modification to maximize PAR transmission to avoid compromising plant productivity.

4.1 | SG blocks radiation and decreases energy use for cooling, water use, and nutrient consumption

According to manufacturer specifications, SG film (ULR-80) was anticipated to block UV and mostly higher wavelengths of light with marginal reductions (−5 to −10%) in light transmission. However, SG blocked a considerable amount of PAR at the canopy level (−25%, season mean), leading to a light limitation for plant growth and photosynthesis. Significant reductions (−53%, season mean) in short-wave radiation under SG blocked radiation contributing to heat, ultimately
decreasing energy used by chillers for cooling (−8%) and irrigation (water + nutrient) consumption (−18%). A previous study with tomatoes reported energy saving up to 25%–33% using glass with an antireflection coating with some near-infrared (NIR) reflective properties (Hemming et al., 2011). Another study with tomatoes grown under wavelength-selective photovoltaic systems (WSPVs) found small water savings due to reduced (−25%) stomatal conductance (Loik et al., 2017). WSPVs absorbed some of the blue and green wavelengths of the solar spectrum for electricity generation, but transmitted remaining wavelengths including most of the red light (Loik et al., 2017). In contrast, SG reduced the intensity of light mainly in the red-light region of the visible light spectrum, which suggested differences in the quality of light in our study relative to Loik et al. (2017). The reduction in water and nutrient consumption of the eggplant crop in our study can be attributed to a reduction in radiation load, as well as decreased photosynthesis and productivity. The

### TABLE 2 Summary of reflectance-based spectral indices, SPAD measurements, leaf mass per area (LMA), and pigment analysis using HPLC

| Parameter (mean)                                      | Treatment                      | Exp | Control     | Smart glass | Change (%) | p-Value |
|-------------------------------------------------------|--------------------------------|-----|-------------|-------------|------------|---------|
| Spectral Index parameters                             |                                |     |             |             |            |         |
| Leaf water content (WBI)                              |                                |     | 1.0456 ± 0.0005 | 1.0364 ± 0.001 | −1         | 1.4 × 10⁻⁸ |
| Chlorophyll content (mNDVI)                           |                                |     | 0.640 ± 0.002  | 0.625 ± 0.004  | −2         | .002 |
| Xanthophyll cycle (PRI)                               |                                |     | 0.0437 ± 0.0003 | 0.0475 ± 0.0005 | +8         | 9.3 × 10⁻⁷ |
| Carotenoid/chl-a (SIPI)                               |                                |     | 1.0142 ± 0.0003 | 1.011 ± 0.0002 | −0.3       | 1.9 × 10⁻⁷ |
| Red–Green ratio (RGR)                                 |                                |     | 0.671 ± 0.002  | 0.631 ± 0.004  | −6         | 2.9 × 10⁻⁹ |
| Chlorophyll degradation (NPQI)                         |                                |     | 0.010 ± 0.001  | 0.021 ± 0.001  | +50        | 2.8 × 10⁻⁶ |
| Red–Far-red ratio (RFR)                               |                                |     | 0.0901 ± 0.0004 | 0.088 ± 0.0003 | −2         | .001 |
| Chlorophyll and carotenoid absolute levels measured by HPLC |                                |     |             |             |            |         |
| Chlorophyll a (µg/gfw)                                |                                | 2   | 1,543 ± 37  | 1,569 ± 29  | +1         | .5 |
| Chlorophyll b (µg/gfw)                                |                                | 2   | 624 ± 17   | 642 ± 14   | +3         | .4 |
| Phytoene (µg/gfw)                                     |                                | 2   | 137 ± 12  | 69 ± 6  | −49        | .0001 |
| B-Carotene (µg/gfw)                                   |                                | 2   | 105 ± 2   | 101 ± 2   | −4         | .1 |
| Lutein (µg/gfw)                                       |                                | 2   | 190 ± 4   | 194 ± 5   | +2         | .5 |
| Neoxanthin (µg/gfw)                                   |                                | 2   | 54 ± 1   | 44 ± 1   | −18        | .0005 |
| Xanthophyll cycle pigments using HPLC                 |                                |     |             |             |            |         |
| Violaxanthin (µg/gfw)                                 |                                | 2   | 73 ± 2   | 60 ± 2   | −18        | .0009 |
| Antheraxanthin (µg/gfw)                               |                                | 2   | 3.5 ± 0.1 | 2.6 ± 0.1 | −26        | .0004 |
| Zeaxanthin (µg/gfw)                                   |                                | 2   | 2.2 ± 0.1 | 1.2 ± 0.1 | −45        | 3.6 × 10⁻⁵ |
| De-epoxidation (DPS)                                  |                                | 2   | 0.073 ± 0.003 | 0.063 ± 0.002 | −14        | .03 |
| Tot carotenoid (µg/gfw)                               |                                | 2   | 430 ± 8 | 404 ± 11  | −6         | .09 |
| Tot chlorophyll (µg/gfw)                              |                                | 2   | 2,168 ± 54 | 2,212 ± 44 | +2         | .5 |
| Xanthophyll/chlorophyll                               |                                | 2   | 0.0365 ± 0.0009 | 0.0291 ± 0.0007 | −20        | 1.1 × 10⁻⁵ |
| Carotenoid/chlorophyll                                |                                | 2   | 0.198 ± 0.003 | 0.182 ± 0.002 | −8         | .0007 |
| SPAD values at different leaf positions               |                                |     |             |             |            |         |
| Leaf-Top                                              |                                | 2   | 56.1 ± 0.7 | 52.6 ± 1.1 | −6         | .02 |
| Leaf-Middle                                           |                                | 2   | 56.7 ± 0.9 | 53.8 ± 1.1 | −5         | .06 |
| Leaf-Bottom                                           |                                | 2   | 57.4 ± 0.9 | 52.9 ± 1.2 | −7         | .01 |
| Leaf mass area (LMA) using leaf fresh weight per unit area |                                |     |             |             |            |         |
| Leaf-Top (mg/cm²)                                     |                                | 2   | 19.4 ± 0.3 | 17.9 ± 0.4 | −7         | .01 |
| Leaf-Middle (mg/cm²)                                  |                                | 2   | 20.5 ± 0.5 | 18.4 ± 0.4 | −10        | .007 |
| Leaf-Bottom (mg/cm²)                                  |                                | 2   | 20.4 ± 0.6 | 17.8 ± 0.4 | −12        | .003 |

Note: Summary of statistical analysis using one-way analysis of variance (ANOVA) for the Smart Glass effect on spectral indices (n = 20), leaf pigment parameters, SPAD measurements, and LMA (n = 10).
electrical power used by chillers is an indirect measurement of energy use (kWh) calculated using water flow and temperatures, before and after cooling. Hence, the actual total energy savings could be different, and a detailed, certified accounting of energy usage is required in future investigations.

4.2 | Plants acclimated to low light by reducing $A_{\text{max}}$ and xanthophyll composition

SG changed light quantity and quality and this was reflected in the responses of photosynthetic activity and pigments. Light-limited reduction in photosynthetic rates is consistent with tomatoes (~20%) grown under WSPVs when measured at higher light levels (Loik et al., 2017). Interestingly, photosynthetic light saturation was observed at ~500 μmol m$^{-2}$ s$^{-1}$ in tomato (Loik et al., 2017) relative to ~1,000 μmol m$^{-2}$ s$^{-1}$ in eggplants (current study) which can be due to differences in growth CO$_2$, temperature (Xin, Li, Zhang, & Hu, 2019), and species. Stomatal conductance was decreased by SG in one of the two eggplant experiments. Loik et al. (2017) also found a higher reduction in $g_s$ than $A_{\text{sat}}$ which was linked to reduced blue light under WSPVs which plays an important role in stomatal functioning. Light intensity and quality both affect stomata (O’Carrigan et al., 2014) and in dynamic light environments, stomata have been found to respond more slowly than photosynthesis, resulting in noncoordination between $A$ and $g_s$ (McAusland et al., 2016). Reduced stomatal conductance, similar to reduced photosynthesis, is partly in response to lower light intensity under SG. However, altered light quality, particularly vastly reduced red and far-red light, may have modified the stomatal response in Experiment 2 and further work is required to understand the impact of SG on light quality and stomata.

In the current study, $A_{\text{sat}}$ and $A_{\text{max}}$ were reduced without significant changes in $\Phi_{\text{max}}$ and $\theta$, suggesting that the photosynthetic apparatus acclimated in response to reduced light intensity (Evans & Poorter, 2001). Acclimation was generally uniform across light-dependent and light-independent reactions of photosynthesis, including photosynthetic efficiency and the electron or photon cost of CO$_2$ fixation, which aligns with unchanged total chlorophyll or carotenoid content. In contrast, chlorophyll $a/b$ ratios and electron transport components decreased at lower light levels in spinach and pea (Evans, 1987; Terashima & Evans, 1988) which can be attributed to the stronger light treatment (>70% lower light for spinach and >80% lower light for pea) compared to relatively modest light treatment (~26% lower light for eggplant) in our study. Unchanged total chlorophyll and carotenoid content suggests that the light treatment in our study was not strong enough to induce changes in total pigment levels, but the shifted light environment could alter pigment composition associated with light capture and photoprotection.

Carotenoid pigments such as the xanthophylls facilitate nonphotochemical quenching (NPQ) and light capture (Demmig-Adams et al., 2014; Niyogi, 1999). Selective synthesis and degradation of chloroplast components during acclimation have been shown to modulate the composition and function of the photosynthetic apparatus (Bailey, Walters, Jansson, & Horton, 2001). Under high light, violaxanthin undergoes de-epoxidation (DPS) via an antheraxanthin intermediate back to zeaxanthin in the thylakoid pigment bed to help dissipate excess light-induced excitation energy as heat and minimize photo-oxidative stress (Demmig-Adams & Adams, 2006; Demmig-Adams et al., 2014; Havaux, Dall’Osto, Cuiné, Giuliano, & Bassi, 2004; Marin et al., 1996). The DPS
was lower (~0.07) in eggplant leaves due to the markedly low abundance of A and Z, keeping consistent with tomato (Ding, Wang, Liu, & Zhang, 2017) and rice leaves (Yin et al., 2010), in comparison to eucalyptus tree leaves that have a considerably higher DPS (~0.7) and similar abundances of V, A, and Z (Dhami, Drake, Tjoelker, Tissue, & Cazzonelli, 2020). Lower light levels are linked to lower DPS values, which can rise during the midday in response to higher light levels (Ding et al., 2006). Our results suggest a limited capacity for eggplant to use the xanthophyll cycle for photoprotection, perhaps relying instead on the production of antioxidants (Logan, Kornyeyev, Hardison, & Holaday, 2006). Spectral indices (e.g., SIPI and PRI) provide additional evidence to support the lower DPS, and these indices have been successfully used for quantifying biophysical characteristics of agricultural crops (Peñuelas, Baret, et al., 1995; Peñuelas, Filella, et al., 1995; Thenkabail, Smith, & De Pauw, 2000). Plants grown under SG appear to have acclimated by lowering their xanthophyll composition, without affecting lutein, β-carotene, or chlorophyll levels. This is consistent with the recent meta-analysis on plant responses to light (Poorter et al., 2019), where the xanthophyll-to-chlorophyll ratio

**TABLE 3** Summary of plant morphology, yield, and fruit quality parameters

| Parameter (mean) | Exp | Control | Smart glass | Change (%) | p-Value |
|------------------|-----|---------|-------------|------------|---------|
| **Productivity and development parameters** | | | | | |
| Mean height (cm/stem) | 1 | 236 ± 2 | 234 ± 2 | −1 | .6 |
| | | 2 | 276 ± 3 | 279 ± 4 | +1 | .6 |
| Mean bud number (n/stem) | 1 | 6.4 ± 0.2 | 6.4 ± 0.3 | 0 | .9 |
| | | 2 | 6.2 ± 0.2 | 6.2 ± 0.2 | 0 | .9 |
| Mean flower number (n/stem) | 1 | 0.6 ± 0.1 | 0.6 ± 0.1 | 0 | ns |
| | | 2 | 1.6 ± 0.1 | 1.7 ± 0.1 | +5 | .3 |
| Mean fruit number (g/stem) | 1 | 0.39 ± 0.01 | 0.28 ± 0.01 | −28 | 2.2 × 10⁻¹⁶ |
| | | 2 | 0.53 ± 0.01 | 0.41 ± 0.01 | −23 | 1.1 × 10⁻¹³ |
| Mean fruit weight (g/stem) | 1 | 155 ± 2 | 105 ± 2 | −32 | 2.2 × 10⁻¹⁶ |
| | | 2 | 192 ± 4 | 145 ± 3 | −24 | 2.2 × 10⁻¹⁶ |
| Total yield (kg m⁻² year⁻¹) | 1 + 2 | 41.3 | 31.8 | −23 | NA |
| Pruned biomass (kg) | 2 | 6.3 | 5.2 | −17 | NA |
| **Eggplant fruit quality parameters** | | | | | |
| Mineral (ash) (g/100 g) | 1 | 0.43 ± 0.01 | 0.39 ± 0.01 | −9 | .02 |
| | | 2 | 0.35 ± 0.01 | 0.25 ± 0.01 | −28 | 3.5 × 10⁻⁷ |
| pH | 1 | 5.46 ± 0.02 | 5.53 ± 0.02 | +1 | .03 |
| | | 2 | 5.09 ± 0.02 | 5.04 ± 0.02 | −1 | .1 |
| Titratable acidity (mq NaOH/kg) | 1 | 9.6 ± 0.4 | 9.3 ± 0.6 | −3 | .7 |
| | | 2 | 9.9 ± 0.4 | 9.4 ± 0.2 | −5 | .3 |
| Moisture (%) | 1 | 93.9 ± 0.2 | 94.4 ± 0.2 | +1 | .2 |
| | | 2 | 94.7 ± 0.2 | 95.3 ± 0.1 | +1 | .07 |
| Total soluble solids (Brix) | 1 | 3.6 ± 0.1 | 3.1 ± 0.1 | −12 | .02 |
| | | 2 | 2.91 ± 0.07 | 2.65 ± 0.05 | −8 | .07 |
| Glucose (g/100 g) | 1 | 1.02 ± 0.01 | 1.09 ± 0.01 | +6 | .004 |
| Fructose (g/100 g) | 1 | 1.07 ± 0.01 | 1.13 ± 0.01 | +5 | .01 |
| Sucrose (g/100 g) | 1 | 0.17 ± 0.01 | 0.24 ± 0.02 | +29 | .03 |
| Total sugars (g/100 g) | 1 | 2.27 ± 0.03 | 2.47 ± 0.04 | +8 | .002 |
| Fat (%) | 1 | 0.072 ± 0.008 | 0.059 ± 0.005 | −18 | .1 |
| N (%) | 1 | 0.112 ± 0.004 | 0.107 ± 0.002 | −4 | .3 |
| Protein (%) | 1 | 0.702 ± 0.026 | 0.674 ± 0.013 | −4 | .3 |

Note: Summary of statistical analysis using one-way analysis of variance (ANOVA) for the Smart Glass effect on plant height (n > 24), bud/flower/fruit number (n > 36), pruned total biomass (per chamber), yield (experiment total), fruit weight (n > 240), and fruit quality parameters (n = 6–10).
correlates with the quantity of light. Lower DPS and zeaxanthin levels under SG also suggest plants may have reduced NPQ based on the curvilinear relationship between zeaxanthin and NPQ (Cheng et al., 2003). However, photosynthesis was limited by electron transport rate under WSPVs, yet no differences were found in NPQ (Loik et al., 2017). Taken together, a reduction in $A_{\text{max}}$ was associated with a reduction in xanthophyll composition and DPS in SG leaves, thereby revealing a reduced photosynthetic capacity for plants acclimated to the SG environment.

4.3 Reduced photosynthesis and high abortion rate under SG decreases yield without changing fruit quality

In crop plants, the average yield is generally reduced by 0.8%–1% for every 1% reduction in light intensity (Marcelis, Broekhuijsen, Meinen, Nijs, & Raaphorst, 2006). In accordence, we found that light-limited (~ −26% DLI) photosynthesis under SG reduced fruit yield (~ −28%) without significantly affecting fruit quality and plant morphological traits, including plant height, bud number, and flower number. One of the few changes in fruit quality (e.g., 29% increase in sucrose content) was a positive impact of SG, while the decrease in mineral (−9% and −28% in experiments 1 and 2, respectively) content was a negative impact. The reduction in fruit yield was driven by reduced fruit number due to a high flower abortion rate under SG relative to control. A very high rate of flower abortion (56.2% in cv Emi and 93.4% in cv Long Negro) has been reported for eggplant cultivars (Passam & Khah, 1992). However, a previous study found a decrease in flowers, flower buds, fresh fruit weight, and fruit growth period under reduced light intensity in eggplants (Uzun, 2007). One cultivar of tomato (cv. Clarence) grown under WSPVs also showed a significant decrease in fruit number and mass due to lower light and photosynthesis (Loik et al., 2017). Poorter et al. (2019) also showed a strong relationship between light intensity and fruit number. The yield reduction in our study could be related to the control of carbon from source to sink. Limited availability of carbon due to reduced photosynthesis may have triggered plants to decrease the number of fruits developed to full maturation, which was evident from high abortion rates in SG. Source-sink regulation is known to control fruit load depending on the availability of photosynthate for translocation during fruit development (Marcelis, Heuvelink, Baan Hofman-Eijer, Den Bakker, & Xue, 2004), which allows plants to produce fewer, but fully developed and better quality fruits (Pallas et al., 2013). Fruit set is related to assimilate supply (source strength) in pepper and low light decreased fruit set due to lower capacity to accumulate sugars and starch during the day (Aloni et al., 1996). Turner and Wien, (1994) suggested that the low-light stress-induced abscission in pepper associated with reduced assimilate partitioning to flower buds could be related to the high assimilate consumption in the maintenance of expanded leaves. However, light quality was
also altered in SG, which may have induced fruit abortion and decreased yield (Cerdán & Chory, 2003). Hence, further investigation is required to understand if light quality, light quantity or both are driving the reduction in fruit yield.

5 | CONCLUSIONS

SG blocked UV and light wavelengths >780 nm, but also a significant proportion of PAR mainly in the red-light region of the spectrum, contributing to decreased energy, water, and nutrient consumption. Reductions in PAR reduced photosynthesis in leaves from SG grown plants, which was associated with a decrease in yield due mainly to higher fruit abortion rates, without affecting fruit quality. SG did not affect morphological features, including plant height, floral bud number or the number of open flowers. Further investigation into whether light quality and/or quantity primarily reduce fruit yield will shed light on how to engineer a new generation of SG for protected cropping industries. It should be noted that SG is likely to have different effects in a crop-specific manner (e.g., vegetative crops such as leafy vegetables may have a different response because leaves, and not reproductive structures, are harvested for yield) and during periods of primarily low light, including winter growing periods. Thus, additional SG trials with different crop types and primarily low-light conditions are required to identify the most appropriate SG characteristics for use with a wide variety of crop plants. Overall, this research shows that novel glass technologies can provide significant energy savings for commercial vegetable greenhouses and may benefit growers who seek to develop sustainable food production with lower resource use in the future. Our study also demonstrates that spectral changes in light may generate biochemical changes, which can be used in biofortification of vegetables and fruits. In addition, SG and similar glazing materials can be further advanced to redirect biologically unused radiation into generation of electricity through solar technologies in the future. The current study demonstrates a comprehensive investigation of potential glazing materials and its impact on plant growth and productivity for commercial use in greenhouse horticulture, which will be useful for future protected cropping research.

ACKNOWLEDGMENTS

We thank Dr. Wei Liang for crop growth and management, Mr Goran Lopaticki for technical operation and maintenance of the glasshouse, Dr. Craig Barton for support in sensor and data logger installation, Dr. Rosalie Durham for providing fruit quality analysis methods, and Dr Margaret Andrew for the use of the spectroradiometer and advice on the spectral indices. This work was supported by Horticulture Innovation Australia projects VG16070 and VG17003. YA was supported by the Australian Indian Institute (AII) New Generation Network (NGN) fellowship. Data are available at the Mendeley research data repository (http://dx.doi.org/10.17632/h225w9kvmer.1).

REFERENCES

Ahamed, S., Guo, H., & Tanini, K. (2019). Energy saving techniques for reducing the heating cost of conventional greenhouses. Biosystems Engineering, 178, 9–33. https://doi.org/10.1016/jbiosystemseng.2018.10.017
Alagoz, Y., Dhami, N., Mitchell, C., & Cazzonelli, C. I. 2020). cis/trans carotenoid extraction, purification, detection, quantification, and profiling in plant tissues. In M. Rodriguez-Concepción, & R. Welsch (Eds.), Plant and food carotenoids (pp. 145–163). New York, NY: Springer US. https://doi.org/10.1007/978-1-4939-9952-1_11
Aloni, B., Karni, L., Zaidman, Z., & Schaffer, A. A. (1996). Changes of carbohydrates in pepper (Capsicum annum L.) flowers in relation to their abscission under different shading regimes. Annals of Botany, 78(2), 163–168. https://doi.org/10.1006/anbo.1996.0109
Andersson, N. E., & Nielsen, O. F. (2000). Energy consumption, light distribution and plant growth in greenhouse partly insulated with non-transparent material, 5.
Babla, M., Cai, S., Chen, G., Tissue, D. T., Cazzonelli, C. I., & Chen, Z.-H. (2019). Molecular evolution and interaction of membrane transport and photoreception in plants. Frontiers in Genetics, 10, 956. https://doi.org/10.3389/fgen.2019.00956
Bailey, S., Walters, R. G., Jansson, S., & Horton, P. (2001). Acclimation of Arabidopsis thaliana to the light environment: The existence of separate low light and high light responses. Planta, 213(5), 794–801. https://doi.org/10.1007/s004250100556
Baker, N. R. (2008). Chlorophyll fluorescence: A probe of photosynthesis in vivo. Annual Review of Plant Biology, 59(1), 89–113. https://doi.org/10.1146/annurev.arplant.59.032607.092759
Bakker, J. C., Adams, S. R., Boulard, T., & Montero, J. I. (2016). Innovative technologies for an efficient use of energy. Acta Horticulturae, 801, 49–62. https://doi.org/10.17660/ActaHortic.2008.801.1
Ballaré, C. L., & Pierik, R. (2017). The shade-avoidance syndrome: Multiple signals and ecological consequences. Plant, Cell and Environment, 40(11), 2530–2543. https://doi.org/10.1111/pce.12914
Baranski, R., & Cazzonelli, C. I. (2016). Carotenoid biosynthesis and regulation in plants. https://doi.org/10.1002/9781118622223.ch10
Barnes, J. D., Balaguer, L., Manrique, E., Elvira, S., & Davison, A. W. (1992). A reappraisal of the use of DMSO for the extraction and determination of chlorophylls a and b in lichens and higher plants. Environmental and Experimental Botany, 32(2), 85–100. https://doi.org/10.1016/0098-8472(92)90034-Y
Benjamini, Y., & Hochberg, Y. (1995). Controlling the False Discovery Rate: A Practical and Powerful Approach to Multiple Testing. Journal of the Royal Statistical Society. Series B (Methodological), 57(1), 289–300. https://www.jstor.org/stable/2346101
Britton, G., Liaaeb-Jensen, S., & Pfander, H. (1995). Carotenoids. Volume 1B: Spectroscopy. Basel, Switzerland: Birkhäuser Verlag AG.
Bugbee, B. (2016). Toward an optimal spectral quality for plant growth and development: The importance of radiation capture. *Acta Horticulturae*, 1134, 1–12. https://doi.org/10.17660/ActaHortic.2016.1134.1

Cazzonelli, C. I., Hou, X., Alagöz, Y., Rivers, J., Dhami, N., Lee, J., … Poong, B. J. (2020). A cis-carotene derived apocarotenoid regulates etioplast and chloroplast development. *Elife*, 9, e45310. https://doi.org/10.7554/eLife.45310

Cerdán, P. D., & Chory, J. (2003). Regulation of flowering time by light. *Science*, 297(5583), 2126–2129. https://doi.org/10.1126/science.1074401

Cuce, E., Harjunowibowo, D., & Cuce, P. M. (2016). Renewable and sustainable energy saving strategies for greenhouse systems: A comprehensive review. *Renewable and Sustainable Energy Reviews*, 64, 34–59. https://doi.org/10.1016/j.rser.2016.05.077

Demmig-Adams, B., & Adams, W. W. (2006). Photoprotection in an ecological context: The remarkable complexity of thermal energy dissipation. *New Phytologist*, 172(1), 11–21. https://doi.org/10.1111/j.1469-8137.2006.01835.x

Demmig-Adams, B., Garab, G., Adams III, W., & Govindjee (Eds.). (2014). *Non-photothermal quenching and energy dissipation in plants, algae and cyanobacteria*. Dordrecht, The Netherlands: Springer Netherlands. https://doi.org/10.1007/978-94-017-9302-1

Dhami, N., Drake, J. E., Tjoelker, M. G., Tissue, D. T., & Cazzonelli, C. I. (2020). An extreme heatwave enhanced the xanthophyll de-epoxidation state in leaves of Eucalyptus trees grown in the field. *Physiology and Molecular Biology of Plants*, 26(2), 211–218. https://doi.org/10.1007/s12298-019-00729-6

Ding, F., Wang, M., Liu, B., & Zhang, S. (2017). Exogenous melatonin mitigates photoinhibition by accelerating non-photothermal quenching in tomato seedlings exposed to moderate light during chilling. *Frontiers in Plant Science*, 8, 244. https://doi.org/10.3389/fpls.2017.00244

Ding, L., Wang, K. J., Jiang, G. M., Li, Y. G., Jiang, C. D., Liu, M. Z., … Peng, Y. (2006). Diurnal variation of gas exchange, chlorophyll fluorescence, and xanthophyll cycle components of maize hybrids released in different years. *Photosynthetica*, 44(1), 26–31. https://doi.org/10.1007/s11099-005-0154-3

Evans, J. (1987). The relationship between electron transport components and photostatic capacity in pea leaves grown at different irradiances. *Australian Journal of Plant Physiology*, 14(2), 157. https://doi.org/10.1071/PP970157

Evans, J., & Poorter, H. (2001). Photosynthetic acclimation of plants to growth irradiance: The relative importance of specific leaf area and nitrogen partitioning in maximizing carbon gain. *Plant, Cell and Environment*, 24(8), 755–767. https://doi.org/10.1046/j.1354-307X.2001.00724.x

Fuentes, D. A., Gamon, J. A., Qiu, H., Sims, D. A., & Roberts, D. A. (2001). Mapping Canadian boreal forest vegetation using pigment and water absorption features derived from the AVIRIS sensor. *Journal of Geophysical Research: Atmospheres*, 106(D24), 33565–33577. https://doi.org/10.1029/2001JD900110

Gamon, J. A., Peñuelas, J., & Field, C. B. (1992). A narrow-waveband spectral index that tracks diurnal changes in photosynthetic efficiency. *Remote Sensing of Environment*, 41(1), 35–44. https://doi.org/10.1016/0034-4257(92)90059-S

Goto, E. (2003). Effects of light quality on growth of crop plants under artificial lighting. *Environment Control in Biology (Japan)*, 41(2), 121–132. https://doi.org/10.2525/ecb1963.41.121

Hao, X., & Papadopoulos, A. P. (1999). Effects of supplemental lighting and cover materials on growth, photosynthesis, biomass partitioning, early yield and quality of greenhouse cucumber. *Scientia Horticulturae*, 80(1), 1–18. https://doi.org/10.1016/S0304-4268(98)00217-9

Havaux, M., & Dall’Osto, L., Ciui, S., Giuliano, G., & Bassi, R., (2004). The effect of zeaxanthin as the only xanthophyll on the structure and function of the photosynthetic apparatus in *Arabidopsis thaliana*. *Journal of Biological Chemistry*, 279(14), 13878–13888. https://doi.org/10.1074/jbc.M311154200

Hemming, S., Kempkes, F. L. K., & Janse, J. (2012). New greenhouse concept with high insulating double glass and new climate control strategies – Modelling and first results from a cucumber experiment. *Acta Horticulturae*, 952, 231–239. https://doi.org/10.17660/ActaHortic.2012.952.28

Hemming, S., Kempkes, F. L. K., & Mohammadkhani, V. (2011). New glass coatings for high insulating greenhouses without light losses – Energy saving, crop production and economic potentials. *Acta Horticulturae*, 893, 217–226. https://doi.org/10.17660/ActaHortic.2011.893.15

Inada, K. (1976). Action spectra for photosynthesis in higher plants. *Plant Cell Physiology*, 17, 355–365.

Kwon, J. K., Khoshimkhujaev, B., Lee, J. H., Yu, I. H., Park, K. S., & Choi, H. G. (2017). Growth and yield of tomato and cucumber plants in polycarbonate or glass greenhouses. *Horticultural Science and Technology*, 35(1), 79–87. https://doi.org/10.7255/HORT.20170009

Lisec, J., Schauer, N., Kopka, J., Willmitzer, L., & Fernie, A. R. (2006). Gas chromatography mass spectrometry–based metabolite profiling in plants. *Nature Protocols*, 1(1), 387–396. https://doi.org/10.1038/nprot.2006.59

Logan, B. A., Adams, W. W., & Demmig-Adams, B. (2007). Avoiding common pitfalls of chlorophyll fluorescence analysis under field conditions. *Functional Plant Biology*, 34(9), 853–859. https://doi.org/10.1071/FP07113

Logan, B. A., Kornyeyev, D., Hardison, J., & Holaday, A. S. (2006). The role of antioxidant enzymes in photoprotection. *Photosynthesis Research*, 88(2), 119–132. https://doi.org/10.1007/s11192-006-9043-2

Loik, M. E., Carter, S. A., Alers, G., Wade, C. E., Shugar, D., Corrado, C., … Kitayama, C. (2017). Wavelength-selective solar photovoltaic systems: Powering greenhouses for plant growth at the food-energy-water nexus. *Earth’s Future*, 5(10), 1044–1053. https://doi.org/10.1002/2016EF000531

Marcelis, L. F. M., Broekhuisjen, A. G. M., Meinen, E., Nijs, E. M. F. M., & Raaphorst, M. G. M. (2006). Quantification of the growth response to light quality of greenhouse grown crops. *Acta Horticulturae*, 711, 97–104. https://doi.org/10.17660/ActaHortic.2006.711.9

Marcelis, L. F. M., Heuvelink, E., Baan Hofman-Eijer, L. R., Den Bakker, J., & Xue, L. B. (2004). Flower and fruit abortion in sweet pepper in relation to source and sink strength. *Journal of Experimental Botany*, 55(406), 2261–2268. https://doi.org/10.1093/jxb/erh245

Marin, E., Nussaume, L., Quesada, A., Gonneau, M., Sotta, B., Hugueney, P., … Marion-Poll, A. (1996). Molecular identification of zeaxanthin epoxidase of Nicotiana plumaginifolia, a gene involved in asbiscic acid biosynthesis and corresponding to the ABA
locus of Arabidopsis thaliana. *The EMBO Journal*, 15(10), 2331–2342. https://doi.org/10.1002/1460-2075.1996.tb00589.x

Marucci, A., & Cappuccini, A. (2016). Dynamic photovoltaic greenhouse: Energy efficiency in clear sky conditions. *Applied Energy*, 170, 362–376. https://doi.org/10.1016/j.apenergy.2016.02.138

Mascarenas, L., Lorenzo, G. A., & Villella, F. (2006). Leaf area index, water index, and red: Far red ratio calculated by spectral reflectance and its relation to plant architecture and cut rose production. *Journal of the American Society for Horticultural Science*, 131(3), 313–319.

McAusland, L., Vialet-Chabrand, S., Davey, P., Baker, N. R., Brendel, O., & Lawson, T. (2016). Effects of kinetics of light-induced stomatal responses on photosynthesis and water-use efficiency. *New Phytologist*, 211(4), 1209–1220. https://doi.org/10.1111/nph.14000

McCree, K. J. (1971). The action spectrum, absorptance and quantum yield of photosynthesis in crop plants. *Agricultural Meteorology*, 9, 191–216. https://doi.org/10.1016/0002-1571(71)90022-7

McCree, K. J. (1971). The action spectrum, absorptance and quantum yield of photosynthesis in crop plants. *Agricultural Meteorology*, 9, 191–216. https://doi.org/10.1016/0002-1571(71)90022-7

Miyagi, K. K. (1999). Photoprotection revisited: Genetic and molecular approaches. *Annual Review of Plant Physiology and Plant Molecular Biology*, 50, 333–359. https://doi.org/10.1146/annurev.arplant.50.1.333

O’Carrigan, A., Babla, M., Wang, F., Liu, X., Mak, M., Thomas, R., … Chen, Z.-H. (2014). Analysis of gas exchange, stomatal behaviour and micronutrients uncovers dynamic response and adaptation of tomato plants to monochromatic light treatments. *Plant Physiology and Biochemistry*, 82, 105–115. https://doi.org/10.1016/j.plaphy.2014.05.012

Ögren, E., & Evans, J. R. (1993). Photosynthetic light-response curves. *Planta*, 189(2), 182–190. https://doi.org/10.1007/BF00195075

Ouzounis, T., Rosenqvist, E., & Ottosen, C.-O. (2015). Spectral effects of light-emitting diodes on plant growth, visual color quality, and photosynthetic photon efficacy. *White versus blue plus red radiation*.

Ouzounis, T., Rosenqvist, E., & Ottosen, C.-O. (2015). Spectral effects of light-emitting diodes on plant growth, visual color quality, and photosynthetic photon efficacy. *White versus blue plus red radiation*.

Peñuelas, J., Gamon, J. A., Griffin, K. L., & Field, C. B. (1993). Assessing community type, plant biomass, pigment composition, and photosynthetic efficiency of aquatic vegetation from spectral reflectance. *Remote Sensing of Environment*, 46(2), 110–118. https://doi.org/10.1016/0034-4257(93)90088-F

Peñuelas, J., Llusia, J., Piiuel, J., & Filella, I. (1997). Photochemical reflectance index and leaf photosynthetic radiation-use-efficiency assessment in Mediterranean trees. *International Journal of Remote Sensing*, 18(13), 2863–2868. https://doi.org/10.1080/014311697217387

Pike, N. (2011). Using false discovery rates for multiple comparisons in ecology and evolution. *Methods in Ecology and Evolution*, 2(3), 278–282. https://doi.org/10.1111/j.2041-210X.2010.00061.x

Pogson, B., McDonald, K. A., Truong, M., Britton, G., & DellaPenna, D. (1996). Arabidopsis carotenoid mutants demonstrate that lutein is not essential for photosynthesis in higher plants. *The Plant Cell*, 8(9), 1627–1639. https://doi.org/10.1105/tpc.8.9.1627

Pooter, H., Niinemets, Ü., Niagkas, N., Siebenkäs, A., Mienenää, M., Matsubara, S., & Pons, T. (2019). A meta-analysis of plant responses to light intensity for 70 traits ranging from molecules to whole plant performance. *New Phytologist*, 223(3), 1073–1105. https://doi.org/10.1111/nph.15754

Price, K. E., electronically mailed to the journal.

R Core Team (2019). *R: A language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing. Retrieved from https://www.R-project.org/

Rigby, E. (2019). *Protected cropping in subtropical climates. A report for Nuffield Australia Farming Solutions*. Retrieved from https://nuffield.com.au/energy-rigby-2/ [Online Resource]

Roser, M., & Ritchie, H. (2019). “Yields and land use in agriculture”. Published online at OurWorldInData.org. Retrieved from https://ourworldindata.org/yields-and-land-use-in-agriculture [Online Resource].

Ruban, A. V. (2009). Plants in light. *Communicative and Integrative Biology*, 2(1), 50–55. https://doi.org/10.4161/cib.2.1.17504

Takii, M., Rohani, A., & Rahmati-Joneidabad, M. (2018). Solar thermal simulation and applications in greenhouse. *Information Processing in Agriculture*, 5(1), 83–113. https://doi.org/10.1016/j.ipa.2017.10.003

Terashima, I., & Evans, J. R. (1988). Effects of light and nitrogen nutrition on the organization of the photosynthetic apparatus in spinach. *Plant and Cell Physiology*, 29(1), 143–155. https://doi.org/10.1093/oxfordjournals.pcp.a077461

Thenkabail, P. S., Smith, R. B., & De Pauw, E. (2000). Hyperspectral vegetation indices and their relationships with agricultural crop characteristics. *Remote Sensing of Environment*, 71(2), 158–182. https://doi.org/10.1016/S0034-4257(99)00067-X

Tsugawa, H., Cajka, T., Kind, T., Ma, Y., Higgins, B., Ikeda, K., … Arita, M. (2015). *MS-DIAL: Data-independent MS/MS deconvolution for comprehensive metabolome analysis*. *Nature Methods*, 12(6), 523–526. https://doi.org/10.1038/nmeth.3393

Turner, A. D., & Wien, H. C. (1994). Photosynthesis, dark respiration and bud sugar concentrations in pepper cultivars differing in susceptibility to stress-induced bud abscission. *Annals of Botany*, 73(6), 623–628. https://doi.org/10.1006/anbo.1994.1078

Uzun, S. (2007). Effect of light and temperature on the phenology and maturation of the fruit of eggplant (*Solanum melongena*) grown in greenhouses. *New Zealand Journal of Crop and Horticultural Science*, 35(1), 51–59. https://doi.org/10.1080/01140670709510167

Xin, P., Li, B., Zhang, H., & Hu, J. (2019). Optimization and control of the light environment for greenhouse crop production. *Scientific Reports*, 9(1), 1–13. https://doi.org/10.1038/s41598-019-44980-z

Xu, J., Lv, Y., Liu, X., Wei, Q., Qi, Z., Yang, S., & Liao, L. (2019). A general non-rectangular hyperbola equation for photosynthetic light response curve of rice at various leaf ages. *Scientific Reports*, 9(1), 1–8. https://doi.org/10.1038/s41598-019-46248-y
Yang, L. Y., Wang, L. T., Ma, J. H., Ma, E. D., Li, J. Y., & Gong, M. (2017). Effects of light quality on growth and development, photosynthetic characteristics and content of carbohydrates in tobacco (Nicotiana tabacum L.) plants. *Photosynthetica, 55*(3), 467–477. https://doi.org/10.1007/s11099-016-0668-x

Yin, Y., Li, S., Liao, W., Lu, Q., Wen, X., & Lu, C. (2010). Photosystem II photochemistry, photoinhibition, and the xanthophyll cycle in heat-stressed rice leaves. *Journal of Plant Physiology, 167*(12), 959–966. https://doi.org/10.1016/j.jplph.2009.12.021

Zhu, X.-G., Long, S. P., & Ort, D. R. (2010). Improving photosynthetic efficiency for greater yield. *Annual Review of Plant Biology, 61*(1), 235–261. https://doi.org/10.1146/annurev-plant-042809-112206

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

**How to cite this article:** Chavan SG, Maier C, Alagoz Y, et al. Light-limited photosynthesis under energy-saving film decreases eggplant yield. *Food Energy Secur*. 2020;00:e245. https://doi.org/10.1002/fes3.245