Effects of different light quality on growth, photosynthetic characteristic and chloroplast ultrastructure of upland cotton (Gossypium hirsutum L.) seedlings
Effects of different light quality on growth, photosynthetic characteristic and chloroplast ultrastructure of upland cotton (Gossypium hirsutum L.) seedlings

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ABSTRACT
The objective of the present study was to evaluate the effects of different light emitting diodes (LEDs) on growth, photosynthetic characteristic and chloroplast ultrastructure of upland cotton (Gossypium hirsutum L.) seedlings. Seedlings of the cultivar Sumian 22 were grown under seven different lights including blue (B) plus red (R) LEDs BR (1:8, 1:3, 1:1, 3:1), B LEDs, R LEDs and fluorescent lamps (FL) with the 100 μmol m⁻² s⁻¹ photosynthesis photon flux (PPF) and a 12 hour photoperiod for 40 days. Compared with FL, fresh mass, dry mass, root length, stem width, root activity, length of palisade tissue and stomatal frequencies were significantly greater as well as the thickest grana lamella in chloroplast in seedlings grown under BR1:8 LEDs; The chlorophyll a, chlorophyll b, total chlorophyll contents, photosynthetic rate, leaf thickness, spongy tissue length and stomatal areas were highest in seedlings under B LEDs; The stem length, leaf area, sucrose, soluble sugar and starch concentrations were highest as well as the greatest number and volume of starch grains in chloroplast in seedlings under R LEDs. It was suggested that BR1:8 LEDs might be used as a primary light for the cultivation of upland cotton seedlings under controlled
conditions.

Keywords: Light emitting diodes; Photosynthetic characteristic; Chloroplast ultrastructure; Spongy tissue length; Stomatal area; Upland cotton (Gossypium hirsutum L.)

INTRODUCTION

Light affected the growth and development of plants (Gong et al., 2015). Two main light qualities were detected by pigment systems in plants, phytochrome and blue-absorbing pigments (BAPs). Phytochrome was most sensitive to red light and far-red light, while BAPs are influenced by B and ultraviolet-A (UV-A) light spectrum (Moe and Heins, 1990). Both light quantity of photon flux and wavelength were very important for plant growth and development (Moshe and Dalia, 2007). R LEDs light might increase starch accumulation by inhibiting the translocation of photosynthates out of leaves (Saebo et al., 1995). In addition, B LEDs light were important for the formation of chlorophyll, chloroplast development and stomatal opening (Senger et al., 1982). Light from B plus R LEDs influenced anatomical features, photosynthesis, growth and development in pepper, lily, strawberry, cherry tomato, rapeseed, non-heading Chinese cabbage and chrysanthemum (Schuerger et al., 1997; Lian et al., 2002; Nhut et al., 2003; Kim et al., 2004; Kurilcik et al., 2008; Liu et al., 2011a, 2011b; Fan et al., 2013a; 2013b; Li et al., 2013).

LEDs were solid-state, long-lasting and durable sources of narrow-band light which can be used in a range of horticultural and photo-biological applications. LEDs provided an opportunity to optimize the spectra for a given plant response and have been used as primary light sources for space-based plant research chambers and bio-regenerative life-support systems, such as plant tissue culture, establishment horticulture, seedling production and zoological experiments (Guo et al., 2008; Stutte, 2009). LEDs had been successfully used to cultivate several plant species, including lettuce (Kim et al., 2004; Kim et al., 2006; Li and Kubota, 2009; Stutte et al., 2009), pepper (Schuerger et al., 1997), spinach (Yorio et al., 2001), Chinese cabbage (Avercheva et al., 2009), non-heading Chinese cabbage (Li et al., 2012; Fan et al., 2013), cucumber (Sander et al., 2010), potato (Jiao and Fang, 2004), tomato (Liu et al., 2011a; 2011b; Fan et al., 2013), upland cotton (Li et al., 2010), maize (Felker et al., 1995), wheat (Goins et al., 1997), strawberry (Nhut et al., 2003), grape (Poudel et al., 2008), cymbidium (Tanaka et al., 1998), lilium (Lian et al., 2002), marigold and salvia (Heo et al., 2002), chrysanthemum (Kim et al., 2004; Lund et al., 2007; Kurilcik et al., 2008) and rehmanniae (Hahn et al., 2000).
Although the previous studies had identified various physiological and morphological effects of light quality on many plant species, few reports had addressed the effects of LEDs and FL on growth, photosynthesis and chloroplast ultrastructure of upland cotton seedlings. Plant leaves were the main organ of photosynthesis, and structure and function of chloroplasts are important for the growth of plants and influence physiological and ecological responses (Peng and Zhou, 2009). Upland cotton was the primary species of cotton cultivated for fibers to be used in textile industry (Kouakou et al., 2007). Seedling culture and transplantation was major system for Chinese cotton production. The system of cotton seedling culture and transplantation was an important breakthrough for cotton production and cultivation technology, helping to the double yields of seeds and fiber in China. It is necessary to study the most effective lights for the industrial cultivation of upland cotton seedlings. The major objective of the present study was to examine the effects of BR (1:8, 1:3, 1:1, 3:1) LEDs, B LEDs, R LEDs and FL (control) on growth, photosynthesis and chloroplast ultrastructure of upland cotton seedlings and select a better light source for upland cotton seedlings grown under the controlled conditions. The present study should provide a theoretical basis and technical support for the reasonable selection of light source for new LEDs lights in cotton factory seedling and facility cultivation.

MATERIALS AND METHODS

Plant materials

The experiments were conducted in growth chamber at Nanjing Agricultural University. Upland cotton cultivar, Sumian 22, was used as the plant material. Seeds of similar size were selected for sowing. Seeds were sown in cells filled with vermiculite and peat (1:1 v/v) for hydroponic cultivation, with one seed per cell. Culture conditions were maintained at 25°C-26°C (day and night) and 40%-50% (relative humidity) in greenhouse. After 7 days, seedlings with two expanded cotyledons were transplanted into plastic tanks (8 cm × 15 cm) which containing vermiculite and peat (1:1 v/v). The seedlings were cultured in incubator which was designed and made by Nanjing Agricultural University.

Experimental design

Seedlings were treated by BR (1:8, 1:3, 1:1, 3:1) LEDs, B LEDs, R LEDs and FL (control). Cultures were maintained at 25°C-26°C (day and night) and 40%-50% (relative humidity) with 100 μmol s⁻¹m⁻² photosynthesis photon flux (PPF) for a 12 hour photoperiod. The positions of
seedlings were assigned to each light treatment according to the same light intensity; The experiment was completely randomized design with three replications. Seedlings were treated by the type’s lights for 40 days. The spectral-energy distribution of the BR (1:8, 1:3, 1:1, 3:1, 11W) LEDs, B LEDs (11W), R LEDs (11W), and FL (T5, 28W) was measured using an spectral photometer (OPT-2000, Optpeco Inc., Beijing, China; Fig 1). The BR LEDs was determined according to the proportion of total light intensity.

Assessment of morphological index
The plants were harvested 40 days after applying treatment. Seedlings were dried at 85°C until a constant mass was reached to determine dry mass. The mass of each seedling was measured using an electronic balance. Stem length was measured from the main stem base to the top of a seedling using a ruler, and stem diameter was measured at the internode nearest to the root using a Venire caliper. The leaf area (in cm²) of each seedling was measured using a Leaf Area Meter (LI-3000, LI-COR Inc., USA).

Assessment of chlorophyll content
Chlorophyll was extracted from the leaves of fifteen seedlings at a similar position within each treatment to examine chlorophyll content. The fresh leaves were weighed to 0.1 g. 0.1 g leaf samples were placed into a mortar with quartz sand, and 10 mL of 80% acetone was added. The chlorophyll was then extracted until the leaf turned white. The optical density (OD) was measured using a spectrophotometer (UV-1200, Jin Peng Inc., Shanghai, China) at 663 nm for chlorophyll a (Chl. a, OD663) and at 645 nm for chlorophyll b (Chl. b, OD645). The concentrations of chlorophyll a and chlorophyll b were determined from the following equations (Lichtenthaler and Wellburn, 1983):

\[
\text{Chlorophyll a (mg·g}^{-1}) = (12.72 \text{ OD663} - 2.59 \text{ OD645}) \frac{V}{1000 W}
\]

\[
\text{Chlorophyll b (mg·g}^{-1}) = (22.88 \text{ OD645} - 4.67 \text{ OD663}) \frac{V}{1000 W}
\]

\[
\text{Total chlorophyll (mg·g}^{-1}) = (8.05 \text{ OD663} + 20.29 \text{ OD645}) \frac{V}{1000 W}
\]

Where V is the total volume of acetone extract (mL), W is the fresh weight (g) of the sample, Ca and Cb is respectively the concentration of chlorophyll a and chlorophyll b.

Assessment of leaf photosynthesis
When plants were grown under different lights for 40 days, measurements were taken on the functional leaves (the third leaves from the top) of the main stems of three seedlings in triplicate.
The photosynthetic rate was performed with a portable photosynthesis system (LI-6400, LI-COR Inc., Lincoln, USA) from 9:30-10:30 am. PPF was set to measure at 100 μmol m⁻²·s⁻¹, and the experimental conditions such as leaf temperature, CO₂ concentration and relative humidity were 24 ± 2°C, 380 ± 5 μL/L, and 40 ± 5%, respectively (Zeng et al., 2012).

Assessment of root activity

A root sample was excised from the lateral roots of fifteen seedlings at a similar position within each treatment. The 0.5 g fresh sample was treated in 5 mL of 0.1% 2, 3, 5-triphenyltetrazolium chloride (TTC) and 5 mL of 0.067 M potassium phosphate buffer, mixed thoroughly and kept for two hours at 37°C. The reaction was terminated with 2 mL of 1 M H₂SO₄. Then the root was removed, rinsed two to three times with distilled water, placed into a mortar with quartz sand and 10 mL of acetone and ground until it turned white. To make a standard curve, 50, 100, 150, 200 or 250 μL of 0.1% TTC was added to five volumetric flasks, and Na₂S₂O₄ and distilled water were added to reach a volume of 10 mL. The optical density was measured using a UV-1200 spectrophotometer (UV-1200, Jin Peng Inc., Shanghai, China) at 490 nm. Root activity was determined using the following equation:

Root activity (mg·g⁻¹·h⁻¹) = ρ · V / W · T (Li et al., 2010). Where ρ is optical density, V is volume (mL) of acetone extract, T is the time (h) of reactions, and W is fresh mass (g) of the sample.

Assessment of sugar and starch contents

Total sugar content was extracted using the method of Martin (Martin et al., 2000) with slight modifications. Leaves (0.5 g) were ground in a mortar with liquid nitrogen. Then 1 mL of 80% ethanol was added, and the mixture was filtered through filter paper. The filtrates were recovered, and the residues were washed again with 70% ethanol and filtered. Both filtrates were mixed, and then 3 mL of distilled water was added. The extract was centrifuged at 12, 000r for 15 min, and 1 mL of supernatant was collected. Soluble sugar was determined by the sulfuric acid-anthrone method and measured at 620 nm. Sucrose was determined by the phloroglucinol method and measured a UV-1200 spectrophotometer (UV-1200, Jin Peng Inc., Shanghai, China) at 480 nm. The method of Takahashi (Takahashi et al., 1995) was used for starch extraction. The residue obtained after ethanol extraction was re-suspended with 0.1 M sodium acetate buffer (pH 4.8) and boiled for 20 min. The gelatinized starch was digested with amyloglucosidase for four hours at 37°C and boiled again to stop the enzymatic reaction. After cooling, the mixture was centrifuged,
and the amount of soluble sugar in the supernatant was determined by anthrone colorimetry. The starch content was estimated by converting glucose to starch equivalents using a factor of 0.9 (Li et al., 2010).

Assessment of leaf anatomical features

Leaf sections (1 cm²; 1 cm × 1 cm) including small lateral veins from fully expanded leaves (from the second or third nodes) of fifteen seedlings were selected and fixed for two days in 50 mL of formaldehyde-based fixative solution containing 95% ethanol, glacial acetic acid and 37% formaldehyde (95:5:5, v/v/v). The leaf samples were dehydrated in a graded ethanol series (75%, 85%, 95%, 100%, and 100%), embedded in paraffin, sectioned, mounted on glass slides, and treated with safranin and fast green stain (Li et al., 2010). The stained sections of leaf tissues were analyzed using a microscope (DP71, Olympus Inc., Japan). Images were viewed on a monitor and analyzed using Motic Images Plus 2.0. Leaf cross-sections were measured for leaf thickness, lengths of palisade tissue and spongy tissue.

Assessment of stomatal features

Samples of fifteen seedlings were collected from fully expanded leaves (from the second or third nodes) of each seedling for stomatal observation. Absorbent cotton fiber was wetted with water, and the abaxial and adaxial surfaces of the leaves were swabbed. When the leaf was dry, transparent nail polish was brushed onto both sides. When the nail polish had air dried and formed a membrane, transparent adhesive tape was pressed onto both sides of each leaf, stripped, and then pressed on a slide. The slide was treated with a neutral plastic seal to make a temporary slide. Epidermal fingerprints were observed by using an optical microscope (Li et al., 2010). The slides were analyzed using an Olympus microscope (DP71, Olympus Inc., Japan). The area and frequency of stomata were measured using Motic Images Plus 2.0.

Assessment of chloroplast ultrastructure

Fully expanded leaves from fifteen seedlings at a similar position within each treatment were destructively sampled. Leaf sections (0.25 cm²; 0.5 cm × 0.5 cm) including small lateral veins were excised from the second leaf from the top and about 1 cm away from the petiole base. The samples were fixed in 2.5% glutaraldehyde (pH 7.0) at 4°C for 12 hours and bledd by vacuum pump. The leaf sections were rinsed three times with 0.1 M phosphate buffer for 20 min, fixed in 1% osmic acid (pH 7.0) for two hours, rinsed three times with 0.1 M phosphate buffer for 20 min,
and then dehydrated in an ethanol series (30%, 50%, 70%, 80%, 90%) for 20 min each. The dehydrated samples were imbued with acetone and embedded in Epon-812 epoxy resin, polymerized, sectioned, and stained with 2% uranyl acetate solution (pH 4.2) for 30 min followed by lead citrate solution (pH 12) for 30 min. The stained samples were then rinsed with distilled water 30 times. After the sections were dry, the samples were observed using a transmission electron microscope (H-7650, Hitachi Inc., Japan).

Statistical Analyses

Statistical analyses were conducted using SPSS statistical product and service solutions for windows version 18.0 (SPSS, Chicago, USA). Each treatment was replicated three times. The data were analyzed using an analysis of variance (ANOVA), and the differences between means were tested using Duncan’s multiple range test (P < 0.05).

RESULTS

Changes in growth and chlorophyll contents under different blue plus red LEDs lights

The measured parameters of the growth and morphogenesis of seedlings, including fresh weight, dry weight, root length, stem length, stem width and leaf area, showed differences under the four mixtures of blue plus red LEDs. Fresh weight, dry weight, root length, stem length, stem width and leaf area were the greatest in the seedlings grown under the BR1:8 LEDs compared with the other lights (Table 1). The present results demonstrate that the BR1:8 LEDs provided more suitable light for the growth of upland cotton seedlings than the other ratios of blue plus red LEDs (BR1:3, 1:1 or 3:1). The chlorophyll a, chlorophyll b and total chlorophyll contents were the greatest in the seedlings grown under BR1:8, which showed significantly higher than the BR3:1, 1:1 or 1:3; However, the BR3:1, 1:1 or 1:3 lights had no significant differences (Table 2).

Changes in growth parameters under different lights

Different lights had variable effects on the growth of upland cotton seedlings. Compared with FL, fresh mass, dry mass, root length and stem width were significantly greater in seedlings grown under BR1:8 LEDs; Stem length and leaf area were significantly greater in seedlings grown under R LEDs (Table 3).

Changes in root activity under different lights

Different lights varied significantly in their effects on root activity in upland cotton seedlings. Root activity of seedlings grown under BR1:8 LEDs was 95.26 mg · g⁻¹h⁻¹, a value 28.21% higher...
than those of the seedlings grown under B LEDs. Root activity was the lowest in seedlings grown under FL (Fig. 2).

**Changes in leaf chlorophyll contents under different lights**

The trend of chlorophyll a, chlorophyll b and total chlorophyll contents under four treatments were the same, the values were highest in seedlings under B LEDs, followed by those grown under BR1:8 LEDs, and the lowest in those grown under FL (Fig. 3). However, contents of chlorophyll a, chlorophyll b and total chlorophyll of seedlings under BR1:8 LEDs and R LEDs were not statistically different.

**Changes in photosynthetic rate under different lights**

Different lights exhibited obvious differences in their effects on photosynthetic rate of upland cotton seedlings. Photosynthetic rate of seedlings grown under BR1:8 LEDs was 8.19 μ mol m⁻² s⁻¹, a value 44.9% higher than those of the seedlings grown under FL. Compared with FL, photosynthetic rate of seedlings grown under B LEDs were the greatest. However, photosynthetic rate of seedlings grown under BR1:8 LEDs and B LEDs were not statistically different (Fig. 4).

**Changes in photosynthetic production under different lights**

The trend of sucrose, soluble sugar and starch concentrations under four treatments were the same, the values were highest in seedlings under R LEDs, followed by BR1:8 LEDs and the lowest in those grown under FL (Table 4).

**Changes in leaf anatomy under different lights**

Leaf thickness and spongy tissue length were the greatest in seedlings grown under B LEDs, followed by those grown under BR1:8 LEDs, and the smallest in seedlings grown under FL. Leaf thickness of seedlings under R LEDs and FL were not statistically different. Length of palisade tissue was the greatest in seedlings grown under BR1:8 LEDs. However, leaf thickness of seedlings under B and R LEDs were not statistically different. (Table 5).

**Changes in chloroplast ultrastructure under different lights**

Seedlings grown under BR1:8 LEDs exhibited a high integrity of the chloroplast ultrastructure with well-developed lamellar structure, elliptical and well-developed starch grains, and thick grana and lamellae (Fig. 5 B and 6 B). Seedlings grown under B LEDs also exhibited a high integrity of the chloroplast ultrastructure with a clearly visible lamellar structure and elliptical starch grains (Fig. 5 C and 6 C). When seedlings grown under R LEDs, the most chloroplasts exhibited a
disrupted ultrastructure with disjoint and ruptured grana and lamellae, but the number and volume
of starch grains were greater than in chloroplasts of seedlings grown under the other light sources
(Fig. 5 D and 6 D). When seedlings were grown under FL, the chloroplast ultrastructure was
substantially modified, the chloroplast lamellae were distorted, and the lamellar structure was faint
(Fig. 5 A and Fig. 6 A). The present results demonstrated that chloroplast structures were well
developed in seedlings grown under BR1:8 LEDs and B LEDs.

Changes in leaf stomata features under different lights
Stomatal areas on the adaxial and abaxial surfaces of leaves were the greatest in seedlings grown
under B LEDs, followed by those grown under BR1:8 LEDs, and the smallest in seedlings grown
under FL (Table 6 and Fig. 7). Stomatal frequencies on the adaxial and abaxial surfaces of leaves
were the greatest in seedlings grown under BR1:8 LEDs. Stomatal frequencies on the abaxial
surfaces of leaves were twice as high as those on the adaxial surfaces in seedlings grown under all
light sources, but the stomatal area did not differ significantly between the adaxial and abaxial
surfaces.

DISCUSSION
Blue plus red LEDs provide a better light for plant growth
The spectrum of sunlight ranges from 380 to 2600 nm. Wavelengths from 390 to 760 nm (visible
light), the physiologically effective wavelengths in sunlight, were known as photo-synthetically
available radiation. Blue-violet light (430 to 450 nm) and red light (from 640 to 660 nm) had the
highest effective photosynthetic rates (Pan et al., 2008). The light sources generally used for plant
growth under controlled conditions include fluorescent lamps, metal halides, high-pressure sodium
lamps, and incandescent lamps (Hahn et al., 2000). The shortcomings of these lights included high
energy consumption and low effectiveness toward plant seedlings. Several plants grew well under
different blue plus red LEDs lights, including non-heading Chinese Cabbage of B:R (1:8,1:6),
Lilium and chrysanthemum of B:R (1:1), strawberry of B:R (3:7) and rapeseed of B:R (3:1) (Hahn
et al., 2000; Lian et al., 2002; Nhut et al., 2003; Kim et al., 2004; Li et al., 2012; Fan et al., 2013;
Li et al., 2013). A certain mixture of blue plus red LEDs might combine the advantages of
monochromatic red LEDs and monochromatic blue LEDs and overcome their disadvantages. The
optimal proportion of blue plus red LEDs light varied with the plant species or cultivars. Therefore,
identifying the optimal proportion of blue plus red LEDs was critical for promoting the seedlings
growth of different plant species. However, few previous papers had examined effects of LEDs, sunlight and fluorescent lamps on growth of upland cotton seedlings. The present results demonstrated that B:R=1:8 LEDs had the significant advantages over sunlight and fluorescent lamps for growth of upland cotton seedlings (Table 1 and 3), so BR1:8 LEDs could be used as a light source for the industrial cultivation of upland cotton seedlings under controlled conditions.

**How lights affect leaf structure and its function?**

Plants were able to respond to changes in irradiance environment by modifying structural and physiological traits of their leaves (Wyka et al., 2008; Dunbar-Co et al., 2009; Guan et al., 2011). The structure and function of chloroplasts were important for the growth of plants and influence their physiological responses (Peng and Zhou, 2009). In addition, chloroplast development depended on light, and different wavelengths of light affected chloroplast structure and chemical changes in plants (Deng, 2007). Chloroplasts developed normally under blue plus red light in tomato leaves, and lamella structure was stacked densely (Zhang et al., 2010). Chloroplasts seemed to show the best development under blue, blue plus red, and blue, red plus green light treatments (Liu et al., 2011b). Our results showed that seedlings grown under B:R=1:8 LEDs exhibited a high integrity of the chloroplast ultrastructure with well-developed lamellar structure and thick grana and lamellae (Fig. 5 B and 6 B), seedlings grown under B LEDs also exhibited a high integrity of the chloroplast ultrastructure with a clearly visible lamellar structure (Fig. 5 C and 6 C). Christopher and Mullet (1994) reported that the expression of a number of chloroplast-encoded genes requires high irradiance B light. Our result was consistent with the previous studies. It might because that cryptochromes (CRYs) and phototropins were specifically sensitive to B light, and phytochromes were specifically sensitive to R light (Whitelam and Halliday, 2007).

The effects of spectral quality on anatomical changes in leaf tissues of pepper plants were generally correlated with the amount of B LEDs light (Schuerger et al., 1997). The largest areas of palisade cells had been observed in birch leaves exposed to B LEDs light (Saebo et al., 1995). Palisade tissue cells in tomato leaves under R plus B LEDs were especially well-developed and spongy tissue cells under the same treatment were localized in an orderly fashion (Liu et al., 2011b). The present results demonstrated that leaf thickness and length of spongy tissue were greatest under B LEDs and that the length of palisade tissue was greatest under B:R=1:8 LEDs.
The present results are inconsistent with those of Schuerger (1997) and Saebo (1995). The palisade tissues contained more chloroplasts than spongy tissues, and many photosynthetic pigments and enzymes were distributed in the grana and lamella of chloroplasts (Pan et al., 2008). Our results also showed that the seedlings grown under the B plus R light and B light were grown well. Longer palisade tissues and thicker grana lamella of the chloroplast in leaves might be beneficial for the growth of upland cotton seedlings.

Phytochrome affected photosynthesis by affecting chlorophyll content (Casal, 2000). Our results showed that the photosynthetic rate and the pigments was the highest under B LEDs, followed by B:R=1:8 LEDs (Fig. 3 and 4). Meanwhile, the net photosynthetic rate of chrysanthemum was the lowest under B LEDs, but with the highest pigments (Kim et al., 2004). However, the highest photosynthetic pigments were in tomato leaves with R, B plus green LEDs treatment, but net photosynthesis was increased significantly under the R plus B LEDs and R, B plus green LEDs (Liu et al., 2011a; 2011b). Our results were inconsistent with the study of Liu (2011) and Kim (2004). The capacity of leaves photosynthesis was related to the numbers of grana and the dense lamella structure by the different light quality (Liu et al., 2011b). Our result showed that the alteration of the leaf structure seems to relate with the photosynthesis. Stomata had dramatic эффект на fotosynthesis (Pan et al., 2008). The present results demonstrated that stomatal area of the seedlings grown under B LEDs was greater than that of the seedlings grown under the other light sources (Table 6 and Fig. 7). A direct effect of photochrome on stomatal development and higher SPAD values along with higher numbers of stomata have been recorded under B LEDs (Farquhar and Sharkey, 1982). In the present study, stomatal development might be affected by the chlorophyll content, which is related to the stomatal area in upland cotton seedlings grown under B LEDs.

**How LEDs on photosynthetic production metabolism?**

Light decreased the chlorophyll content of cymbidium (Tanaka et al., 1998). The lowest pigments in tomato leaves of seedlings were found in those with R LEDs treatment (Liu et al., 2011a). The present results demonstrated that the chlorophyll content was lower (Fig. 3) in upland cotton seedlings grown under R LEDs than those grown under B LEDs or B plus R LEDs. The results are consistent with those of Tanaka and Liu. Therefore, R LEDs may decrease the chlorophyll content of leaves in upland cotton.
Starch was the major storage carbohydrate in plants and has many important functions (Geiger et al., 1995). R LEDs light enhanced starch accumulation in Glycine and Sorghum (Britz and Sager, 1990). Light quality regulated the carbohydrate metabolism of higher plants, the carbohydrate content was high under R LEDs light (Kowallik, 1982). The accumulation of starch in chloroplasts, which was enhanced by R LEDs light, may inhibit photosynthesis. Thus, R LEDs light appeared to inhibit the translocation process (Saebo et al., 1995). Excess starch accumulation inhibited photosynthesis in leaves (Bondada and Syvertsen, 2005). Chloroplast of cherry tomato leaves under R LEDs was relatively rich in starch granules (Liu et al., 2011b). The present results demonstrated that contents of sucrose, soluble sugar and starch were greatest (Table 4) in seedlings grown under R LEDs and that the number and volume of starch grains were significantly increased in chloroplasts of seedlings grown under R LEDs (Fig. 6 D), the photosynthetic rate was lower in seedlings grown under R LEDs compared with those grown under blue LEDs or B plus R LEDs (Fig. 4). The present results were consistent with those of the previous studies (Kowallik, 1982; Britz and Sager, 1990; Saebo et al., 1995; Bondada and Syvertsen, 2005; Liu et al., 2011b). The photosynthetic carbon metabolic pathway was not static but was influenced by environmental conditions (Zhang et al., 2015). R LEDs promoted the production of photosynthetic products but might inhibit the transportation of photosynthetic products out of the leaves, starch accumulation in the leaves and leaf photosynthesis was prohibited in upland cotton seedlings.

CONCLUSIONS

The present study might be the first paper of determining different light qualities on growth, photosynthetic characteristic and chloroplast ultra-structure of upland cotton cultivar Sumian 22. The mixture blue plus red (BR1:8) LEDs light might be propitious and necessary to upland cotton seedling growth and can be used as a primary lights for cotton seedling cultivation.

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Author's contributions

C-M.T and Z-G. X design the study. H-M.L did the data analysis and wrote the paper. C-M.T corrected the article.

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| Light treatment | Fresh weight (g) | Dry weight (g) | Root length (cm) | Stem length (cm) | Stem width (cm) | Leaf area (cm²) |
|-----------------|-----------------|----------------|------------------|------------------|----------------|----------------|
| B:R=1:8         | 5.46±0.03a      | 0.72±0.02a     | 12.40±0.04a      | 21.35±0.56a      | 2.33±0.03a      | 53.70±0.45a    |
| B:R=1:3         | 4.19±0.06c      | 0.60±0.03b     | 11.15±0.12b      | 18.60±0.45b      | 2.01±0.06b      | 40.06±0.96b    |
| B:R=1:1         | 4.74±0.05b      | 0.62±0.04b     | 10.10±0.09c      | 19.00±0.52b      | 1.75±0.12c      | 41.52±0.87b    |
| B:R=3:1         | 4.84±0.06b      | 0.61±0.05b     | 10.95±0.08b      | 18.30±0.78b      | 2.04±0.06b      | 42.80±0.56b    |

B:R=1:8, 11.1% Blue plus 88.9% red light-emitting diodes; B:R=1:3, 25% Blue plus 75% red light-emitting diodes; B:R=1:1, 50% Blue plus 50% red light-emitting diodes; B:R=3:1, 75% Blue plus 25% red light-emitting diodes;

Values are the mean ± standard deviation; The different letters within the columns indicate significant differences at P<0.05 by Duncan’s Multiple Range Test.
Table 2: The effects of different blue plus red LEDs lights on the chlorophyll contents of upland cotton seedlings

| Light treatment | Chl. a (mg · g⁻¹) | Chl. b (mg · g⁻¹) | Chl.(a+b) (mg · g⁻¹) |
|----------------|--------------------|--------------------|----------------------|
| B:R=1:8        | 1.63±0.02a         | 0.79±0.03a         | 2.41±0.03a           |
| B:R=1:3        | 1.55±0.03b         | 0.67±0.02b         | 2.22±0.05b           |
| B:R=1:1        | 1.59±0.01b         | 0.72±0.03b         | 2.28±0.06b           |
| B:R=3:1        | 1.58±0.02b         | 0.70±0.02b         | 2.33±0.02b           |

B:R=1:8, 11.1% Blue plus 88.9% red light-emitting diodes; B:R=1:3, 25% Blue plus 75% red light-emitting diodes; B:R=1:1, 50% Blue plus 50% red light-emitting diodes; B:R=3:1, 75% Blue plus 25% red light-emitting diodes;

Values are the mean ± standard deviation; The different letters within the columns indicate significant differences at P<0.05 by Duncan’s Multiple Range Test.
### Table 3 The effects of different lights on growth of upland cotton seedlings

| Light treatment | Fresh mass (g) | Dry mass (g) | Root length (cm) | Stem length (cm) | Stem width (cm) | Leaf area (cm²) |
|-----------------|----------------|--------------|------------------|------------------|-----------------|-----------------|
| FL              | 1.79±0.12d     | 0.27±0.01c   | 4.86±0.23c       | 12.50±0.62c      | 1.67±0.15c      | 28.11±3.27d     |
| B:R=1:8         | 3.89±0.15a     | 0.53±0.03a   | 8.06±0.18a       | 17.1±1.12ab      | 2.31±0.16a      | 49.27±2.24b     |
| B               | 2.67±0.18c     | 0.35±0.04b   | 6.44±0.32b       | 15.2±1.50b       | 2.12±0.14ab     | 32.29±1.45c     |
| R               | 3.40±0.08b     | 0.40±0.02b   | 4.94±0.27c       | 18.32±1.11a      | 1.92±0.18bc     | 63.56±4.31a     |

FL, Fluorescent lamps; B:R=1:8, Blue plus red light emitting diodes; B, Blue light emitting diodes; R, Red light emitting diodes; Data represent mean ±standard deviation of three replicates (n = 3); The different letters within the columns indicate significant differences at P<0.05 by Duncan’s Test.
Table 4 The effects of different lights on photosynthetic products of upland cotton seedlings

| Light treatment | Sucrose (mg·g⁻¹) | Soluble sugar (mg·g⁻¹) | Starch (mg·g⁻¹) |
|-----------------|------------------|------------------------|-----------------|
| FL              | 10.93±0.14d      | 48.01±5.32d            | 6.17±0.87d      |
| B:R=1:8         | 16.92±0.20b      | 96.30±8.90b            | 18.45±1.28b     |
| B               | 15.37±0.16c      | 66.61±3.08c            | 10.40±0.38c     |
| R               | 18.72±0.60a      | 139.06±10.56a          | 25.01±2.07a     |

FL, Fluorescent lamps; B:R=1:8, Blue plus red light emitting diodes; B, Blue light emitting diodes; R, Red light emitting diodes. Data represent mean ± standard deviation of three replicates (n = 3). The different letters within the columns indicate significant differences at P<0.05 by Duncan’s Test.
Table 5 The effects of different lights on leaf anatomy of upland cotton seedlings

| Light treatment | Leaf thickness (μm) | Palisade tissue Length (μm) | Spongy tissue Length (μm) |
|-----------------|---------------------|-----------------------------|--------------------------|
| FL              | 189.68±8.62c        | 54.56±3.35c                 | 80.41±2.92d              |
| B:R=1:8         | 250.89±12.32b       | 89.90±1.53a                 | 100.56±2.30b             |
| B               | 280.26±10.02a       | 80.40±3.36b                 | 120.17±2.53a             |
| R               | 194.44±9.12c        | 75.06±3.39b                 | 86.11±2.13c              |

FL, Fluorescent lamps; B:R=1:8, Blue plus red light emitting diodes; B, Blue light emitting diodes; R, Red light emitting diodes; Data represent mean ±standard deviation of three replicates (n = 3); The different letters within the columns indicate significant differences at P<0.05 by Duncan’s Test.
Table 6 The effects of different lights on leaf stomata of upland cotton seedlings

| Light treatment  | Area of a stomata | Stomata frequency |
|------------------|-------------------|-------------------|
|                  | (μm²)             | (number / mm²)    |
|                  | Adaxial surface   | Abaxial surface   |
|                  | Adaxial surface   | Abaxial surface   |
| FL               | 5.75±0.12d        | 5.95±0.09d        |
|                  | 330.18±12.62bc    | 1006.70±22.34bc   |
| B:R=1:8          | 8.04±0.17b        | 8.05±0.21b        |
|                  | 402.60±12.08a     | 1191.70±22.42a    |
| B                | 8.75±0.19a        | 8.85±0.12a        |
|                  | 350.58±21.35b     | 1049.40±32.54b    |
| R                | 6.05±0.22c        | 6.66±0.16c        |
|                  | 315.26±10.12c     | 955.13±32.36c     |

FL, Fluorescent lamps; B:R=1:8, Blue plus red light emitting diodes; B, Blue light emitting diodes; R, Red light emitting diodes; Data represent mean ±standard deviation of three replicates (n = 3); The different letters within the columns indicate significant differences at P<0.05 by Duncan’s Test.
Fig. 1 The spectral distribution of light treatments

FL: Fluorescent lamp; B: 100% blue light; R: 100% red light.
Fig. 2 Effects of different lights on root activity of upland cotton seedlings

FL, fluorescent lamps; B:R=1:8, Blue plus red light emitting diodes; B, Blue light emitting diodes; R, Red light emitting diodes; The different letters indicate significant differences at P<0.05 by Duncan’s Test.
Fig. 3 Effects of different lights on chlorophyll content of upland cotton seedlings

FL, fluorescent lamps; B:R=1:8, Blue plus red light emitting diodes; B, Blue light emitting diodes; R, Red light emitting diodes; The different letters indicate significant differences at P<0.05 by Duncan’s Test.
Fig. 4 Effects of different lights on the photosynthetic rates of upland cotton seedlings

FL, fluorescent lamps; B:R=1:8, Blue plus red light emitting diodes; B, Blue light emitting diodes; R, Red light emitting diodes; The different letters indicate significant differences at P<0.05 by Duncan’s Test.
**Fig. 5** Effects of different lights on starch grains in chloroplast of upland cotton seedlings

A. fluorescent lamps; B. Blue plus red light emitting diodes (B:R=1:8); C. Blue light emitting diodes; D. Red light emitting diodes; S, Starch gains; Bar = 10 μm.
**Fig. 6** Effects of different lights on the lamella in chloroplasts of upland cotton seedlings

- A. fluorescent lamps; B. Blue plus red light emitting diodes (B:R=1:8); C. Blue light emitting diodes; D. Red light emitting diodes; S: Starch gains; GL: Grana lamellae; Bar=5 μm.
**Fig. 7** Effects of different lights on the abaxial surface stomata of leaves in upland cotton seedlings

- A, fluorescent lamps; B, Blue plus red light emitting diodes (B:R=1:8); C, Blue light emitting diodes; D, Red light emitting diodes; Sto: Stomata; Ep: Epidermis; Bar=25μm.