Moderately enhancing cytokinin level by down-regulation of \textit{GhCKX} expression in cotton concurrently increases fiber and seed yield

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Received: 13 January 2014 / Accepted: 29 August 2014 / Published online: 29 January 2015
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Abstract  Cotton is the leading natural fiber crop in the world. Cotton seeds are also an important oil and protein source. However, enhancement of fiber abundance usually leads to a smaller seed. Thus, it has become a challenge for cotton breeding to concurrently increase fiber yield and seed yield. To improve cotton yield, we elevated the endogenous cytokinin level in transgenic cotton by constitutive suppression of cytokinin dehydrogenase (CKX), a key negative regulator controlling endogenous cytokinin in plants. The slightly and moderately suppressed transgenic cotton plants showed normal growth and development, while the severely suppressed plants exhibited a typical cytokinin-overproduction alteration. The suppression of CKX led to an enhancement of endogenous cytokinins in transgenic cotton plants. Total cytokinins in moderately suppressed lines, CR-3 and CR-6, increased by 20.4 and 55.5 % respectively, and that in the severely suppressed line (CR-13) increased by 134.2 % compared to the wild type. The moderately suppressed lines showed a delay in leaf senescence, higher photosynthesis, more fruiting branches and bolls, and bigger seed size. Field trials showed that seed yield and lint yield of the moderately suppressed CR-6 line increased by 15.4 and 20.0 %, respectively. Meanwhile, the enhanced cytokinin level in transgenic cottons did not show significant influence on fiber qualities. Our data demonstrated that \textit{CKX} is a promising gene for crop yield improvement.

Keywords  CKX · Cytokinin · Cotton fiber · Seed · Yield

Introduction

Cotton is one of the most valuable commercial crops in the world. In addition to the fiber used for textile manufacturing, cotton seed is an important source of oil and protein, and the seed hulls are used for cattle feed and mushroom production (Sunilkumar et al. 2006; Wilkins et al. 2000). Cotton fiber is derived from single epidermal cells of seeds. Enhancing the abundance of fiber usually leads to smaller seeds (Miller and Rawlings 1967; Zhang et al. 2005). Thus, it is a challenge for cotton breeding to concurrently increase both fiber yield and seed yield.

Cytokinins are a group of phytohormones that regulate cell division and influence numerous
developmental and physiological processes of plants (Werner and Schmülling 2009), including leaf senescence (Gan and Amasino 1995), vascular development (Mähönen et al. 2006), cell differentiation at shoot and root apical meristem (Wolters and Jürgens 2009), nutrient uptake and allocation (Séguêla et al. 2008), abiotic (Peleg et al. 2011; Rivero et al. 2007) and biotic stress responses (Choi et al. 2010; Siemens et al. 2006), and regulation of source–sink relationships (Roitsch and Ehneß 2000). Importantly, recent studies have revealed that cytokinin is a key regulator for seed yield (Ashikari et al. 2005; Bartrina et al. 2011).

Exogenous application of kinetin (6-furfurylaminopurine), one of the cytokinin compounds, was able to improve the yield of seed cotton and lint fiber (Sawan et al. 2000). However, large-scale commercial applications of cytokinins in crops are unfeasible in practice due to high costs and time consumption (Li et al. 2004). Genetic modification provides a means to manipulate hormone concentrations in plants. Promoting cytokinin biosynthesis is an effective method of overproducing cytokinin in plants. The ipt gene encoding isopentenyl transferase, which catalyses the rate-limiting step in cytokinin biosynthesis, has been widely used for the enhancement of cytokinins in transgenic plants (Klee et al. 1987; Smigocki and Owens 1988; Smigocki 1991; Wang et al. 1997; Geng et al. 2001, 2002; van der Graaff et al. 2001; Kuppu et al. 2013; Reguera et al. 2013; Rupp et al. 1999; Synkova et al. 1999). However, constitutive overexpression of the gene is usually associated with adverse effects on growth and development of the plants (Smart et al. 1991; Medford et al. 1989; McKenzie et al. 1994; van der Graaff et al. 2001; Geng et al. 2001; Guo et al. 2005; Synková et al. 1999). Another approach to elevating endogenous cytokinins is the suppression of cytokinin deactivation. Cytokinin oxidase/dehydrogenase (CKX), which catalyzes the catabolism of cytokinins to inactive products that lack the N6-unsaturated side chain (Jones and Schreiber 1997), is a crucial negative regulator controlling endogenous cytokinin contents in the plant kingdom (Mok and Mok 2001; Schmülling et al. 2003; Werner et al. 2001, 2003; Kowalska et al. 2010). Decrease of CKX expression level results in more cytokinins in inflorescence meristems, and thus more grains (Ashikari et al. 2005).

To increase endogenous cytokinins in cotton, we had tried to generate transgenic cottons using a constitutive promoter, CaMV35S, to control the expression of ipt. However, as happened in cucumber transformed with 35S::ipt reported by Smigocki and Owens (1989), the transformed calli never developed further into plant regenerants (unpublished data). In this paper, by using GhCKXRNAi, we successfully generated cytokinin-overproduction cottons. Our data demonstrated that moderate suppression of GhCKX can significantly improve both seed and fiber yield of cotton.

Materials and methods

Plant transformation and growth conditions

The 35S::GhCKXRNAi construct (Zeng et al. 2012) was introduced into cotton using the method of Agrobacterium-mediated transformation as described previously (Luo et al. 2007). Kanamycin-resistant and GUS-positive plants were screened out for further study. Cotton plants were grown in greenhouse in 30×928 cm (diameter×height) pots under a 16-h light/8-h dark cycle, at 26–32 °C. Pindstrup Substrate (Pindstrup Mosebrug A/S, Denmark) was used as potting mixture. The plants were watered once every 2 days, and fed once every month with compound fertilizer (XIYANG 17-17-17, China).

RNA isolation and RT-PCR analyses

Total RNA was isolated from stem shoots according to the manual of EASY Spin Plant RNA Kit (Aidlab, China). cDNA was synthesized by RevertAid First Strand cDNA Synthesis Kit (Fermentas, Canada). For RT-PCR analysis, the cDNA sample was used to amplify a GhCKX segment with primers GhCKX1 (5′-ccaaagggcacggtcattctg-3′) and GhCKX2 (5′-cccccttcggctgctgtg-3′). The HISTONE3 gene (AF024716) was used as the internal control with primers GhHIS1 (5′-gaagcctcatcgataccgtc-3′) and GhHIS2 (5′-ctaccatatcaattg-3′). Each reaction contained 10 μL SsoFast EvaGreen Supermix (Bio-Rad, USA), 500 nM each of forward and reverse primers and 1 μL of cDNA. PCR was carried out on a CFX96™ Real-Time System with following conditions: a pretreatment (95 °C, 2 min) followed by 40 amplification cycles (95 °C, 15 s; 56 °C, 15 s; 72 °C, 20 s).
Quantification of endogenous cytokinins

The concentration of endogenous cytokinins was determined in young leaves (the first main-stem leaf from the apex at 130 days after sowing [DAS]), mature leaves (the fifth main-stem leaf from the apex at 130 DAS), stem shoots (the first internode from the apex at 90 DAS), flower buds (3 days before anthesis) and 35-DPA (days post-anthesis) ovules. Tissues (100 mg) were pooled for each sample. Three independent biological replicates were analyzed for the tissue of each line. Extraction, purification, and quantification of endogenous cytokinins by high-performance liquid chromatography linked to a 4000 Q TRAP LC/MS/MS system (ABsciex, USA) were performed according to the method described by Zeng et al. (2012). The product ion pairs of each cytokinin component and corresponding deuterated internal standards, as well as their acquisition parameters, are summarized in Table S4.

Determination of chlorophyll contents

Leaves (100 mg fresh weight (FW)) were placed in a 10 mL tube containing 5 mL 80 % acetone and incubated in the dark at room temperature until the tissues became white. Total chlorophyll was determined using absorbance at 645 nm and 663 nm according to the equation: 20.2 A645 + 8.02 A663 (Chory et al. 1994).

Photosynthesis measurements

Photosynthetic rate was measured by the Li-6400 portable photosynthesis system (LI-COR, Lincoln, NE, USA). The fourth main-stem leaf from the apex (functional leaf) of 10 plants per line at 120 DAS was used for the analysis. The instrument was set at saturating light of 1,200 μmol m⁻² s⁻¹ and a CO₂ concentration of 400 ppm.

Determination of soluble sugar and soluble protein

Ovules and boll shells of 35-DPA bolls at third fruit branch (base to top) were frozen in liquid nitrogen. Samples (100 mg FW of each) were homogenized and extracted in 5 mL 80 % ethanol, and heated for 45 min in boiling water. After cooling to room temperature, the extraction was centrifuged at 3,000 rpm for 10 min. The supernatant was collected to determine the amount of soluble sugars by the colorimetric method with anthrone-sulfuric acid at 620 nm with glucose as a standard.

The protein content was determined by the Coomassie Brilliant Blue G-250 method with bovine serum albumin as the standard. Soluble proteins were extracted from 0.2 g of fresh tissue at 4 °C with 5 mL of extracting solution containing 93.7 % 0.2 M Na₂ HPO₄, 5.2 % 0.2 M NaH₂PO₄, 2 % PVP, and 0.1 % (v/v) β-mercaptoethanol for 60 min. After centrifugation at 4,000 rpm for 10 min, the supernatant was used for protein determination.

Ovule culture

Ovules at 0 DPA were harvested and cultured in BT medium according to the method of Beasley and Ting (1974). As well as 0.5 μM gibberellic acid and 5 μM indole-3-acetic acid, trans-Zeatin with concentrations of 0, 0.5, 1.0, 5.0, 8.0, and 15 μM was added to the medium. After 2-week culture in the dark at 32 °C, ovules were observed by an MVX-10 microscope (Olympus, Japan).

Scanning electron microscopy (SEM)

The 0-DPA ovules were imaged by an S-3400 N SEM (Hitachi, Japan). From the similar region of each SEM image, an area of 250 × 250 μm² was selected for fiber initial counting.

Field experiment

Homozygous T₂ generation lines of 35S::CKXRNAi were planted in 2010 in an experimental farm at Southwest University (Chongqing, China) with 15 plants per row. Line CR-6 (T₄) was selected as the best line of the moderately down-regulated GhCKX cotton.

To assess the agronomic performance of the line, plants were grown in field conditions for randomized comparative trial in 2012 with three replications. Each block was 20 m² and contained 60 plants in four rows 1.0 m apart. The space between two neighboring plants in a row was 0.33 m. The plant height (from soil surface to top), fruit branch number, and square number were measured at 105 DAS. The boll number was counted at 130 DAS. After harvest, seed cotton was ginned. Fibers and seeds were weighed separately, and the number of seeds per boll, lint percentage (fiber
weight/seed cotton weight), seed index (the weight of 100 seeds), lint index (the lint weight of 100 seeds), fiber yield and seed yield were determined. Fibers (3 × ~ 15 g) from each line were sampled randomly and sent to the National Center for Evaluation of Fiber Quality (Anyang, China) for measuring their qualities.

Results

Down-regulation of GhCKX in cotton

To increase the cytokinin level in cotton by down-regulation of the expression of GhCKX, we generated transgenic CaMV35S::GhCKXRNAi (CR) cottons. 17 independent transgenic cottons were obtained. Based on GhCKX expression levels detected by real-time RT-PCR, transgenic cottons were divided into three groups: slightly suppressed (gene expression level was decreased by less than 30 %; CR-1, -2, and -7), moderately suppressed (decreased by 30–70 %; CR-3, -4, -5, -6, -8, -9, -10, and -15), and severely suppressed (over 70 %; CR-11, -12, -13, -14, -16, and -17) (Fig. 1). The severely suppressed GhCKX cottons showed a typical cytokinin overproduction phenotype, e.g. stunted shoots, shortened internode, smaller leaves and floral organs, and even sterility (CR-17, Fig. S1). In contrast, no developmental abnormality was observed in the slightly and moderately suppressed cottons.

Fig. 1 Relative expression levels of GhCKX in T1 transgenic cotton plants. RNAs were isolated from young stems of transgenic cotton plants and analyzed by real-time RT-PCR. WT wild type, NT nontransgenic plants segregated from selfed progenies of transgenic cottons. Error bars indicate SD of three replicates.

Down-regulation of GhCKX-enhanced endogenous cytokinins

To confirm that regulation of GhCKX changes the level of endogenous cytokinins in transgenic cottons, we determined the content of 13 different cytokinins in the leaves, stem shoots, flower buds, and ovules of two GhCKX moderately suppressed lines (CR-3 and -6) and one severely suppressed line (CR-13) by LC/MS (Tables 1 and S1). Total cytokinins in young leaves of transgenic cotton lines CR-3, -6 and -13 increased 13.6, 28.2, and 79.7 %, respectively, compared to that of the wild type. In mature leaves, the cytokinin level in transgenic lines increased 99.4, 106.7, and 194.5 %, respectively. For stem shoots, flower buds, and ovules, the averaged levels in CR-3 and CR-6 rose 39.8, 59.9, and 36.0 %, respectively, and the content in severely suppressed line CR-13 was rather high, increasing by 61.1, 315.6, and 139.9 %, respectively (Table 1). With the decline in gene expression, the content of cytokinins increased significantly, confirming that down-regulation of GhCKX elevated the level of endogenous cytokinins in cotton.

Moderate down-regulation of GhCKX delayed leaf senescence

Chlorophyll degradation is a sign of leaf senescence (Kusaba et al. 2007). To investigate whether down-regulation of GhCKX in cotton delays the leaf
senescence, we compared the chlorophyll content in the leaves of GhCKX moderately suppressed lines (CR-3 and -6), severely suppressed line (CR-13), and wild type at 90, 105, and 120 DAS in field conditions. For upper leaves, there was a steady increase in chlorophyll content in transgenic lines and the wild type from 90 DAS to 105 and 120 DAS (Fig. 2). The content in the three transgenic lines was higher than that in the wild type. For the middle leaves, along with aging, the content in 105- and 120-DAS leaves of transgenic lines and wild-type plants declined. However, the content in transgenic leaves was higher than the wild type. In the aged leaf, i.e. the 120-DAS lower leaves, the content decreased dramatically compared with the upper leaves (Fig. 2). The content in the three transgenic lines was higher than that in the wild type. For the middle leaves, along with aging, the content in 105- and 120-DAS leaves of transgenic lines and wild-type plants declined. However, the content in transgenic leaves was higher than the wild type. In the aged leaf, i.e. the 120-DAS lower leaves, the content decreased dramatically compared with the upper leaves (Fig. 2). The content in the three transgenic lines was higher than that in the wild type. For the middle leaves, along with aging, the content in 105- and 120-DAS leaves of transgenic lines and wild-type plants declined. However, the content in transgenic leaves was higher than the wild type. In the aged leaf, i.e. the 120-DAS lower leaves, the content decreased dramatically compared with the upper leaves (Fig. 2). The content in the three transgenic lines was higher than that in the wild type. For the middle leaves, along with aging, the content in 105- and 120-DAS leaves of transgenic lines and wild-type plants declined. However, the content in transgenic leaves was higher than the wild type. In the aged leaf, i.e. the 120-DAS lower leaves, the content decreased dramatically compared with the upper leaves (Fig. 2). The content in the three transgenic lines was higher than that in the wild type. For the middle leaves, along with aging, the content in 105- and 120-DAS leaves of transgenic lines and wild-type plants declined. However, the content in transgenic leaves was higher than the wild type. In the aged leaf, i.e. the 120-DAS lower leaves, the content decreased dramatically compared with the upper leaves (Fig. 2).

Moderate down-regulation of GhCKX increased leaf photosynthesis and enhanced soluble sugar and protein in the boll

We measured photosynthesis of the fourth leaf from the apex at 120 DAS. The photosynthetic rate of CR-3 and CR-6 leaves was higher than the wild type, showing a 10.7 and 12.2 % increase, respectively. However, there was no such enhancement in the leaves of the severely suppressed line CR-13 (Fig. 3a). In addition, the concentrations of soluble sugar and protein in the boll shell and the ovule at 35 DPA (days post-anthesis) of CR-6 and CR-13 were significantly higher than those of the wild type (Fig. 3b, c).

Table 1 Total content of cytokinins in transgenic lines and wild type (ng/g fresh weight)

| Line   | Cytokinins | Young leaf | Mature leaf | Stem shoot | Flower bud | Ovule | Total | Increase level (%) |
|--------|------------|------------|-------------|------------|------------|-------|-------|-------------------|
| WT     |            | 75.5       | 32.6        | 142.5      | 38.5       | 983.3 | 1,272.4 | –                 |
| CR-3   |            | 85.8       | 65.0        | 190.3      | 53.0       | 1,138.2 | 1,532.3 | 20.4              |
| CR-6   |            | 96.8       | 67.4        | 208.0      | 70.1       | 1,536.9 | 1,979.2 | 55.5              |
| CR-13  |            | 135.7      | 96.0        | 229.6      | 160.0      | 2,359.1 | 2,980.4 | 134.2             |

Well-ground samples were extracted in cold 80 % (v/v) methanol. Cytokinins were purified by a Sep-Pak® Plus C18 cartridge (Waters, Ireland) and determined by HPLC–MS/MS systems (ABsciex, USA). The total values were the sum of cytokinin contents in young leaf, mature leaf, stem shoot, flower bud, and ovule, respectively. WT wild type; CR-3, CR-6 and CR-13, 35S::GhCKXRNAi transgenic plants

Based on the expression level of GhCKX (Fig. 1), coupled with phenotype observation in transgenic cottons (T2 generations), one slightly suppressed (the level decreased by less than 30 %) GhCKX line, CR-7, five moderately down-regulated lines, CR-3, -4, -5, -6, and -8 (decreased by 30–70 %), and two severely suppressed (decreased by over 70 %) lines, CR-11 and CR-13, were planted in field conditions to observe their agronomic performance. At the flower stage, the plant height of severely suppressed GhCKX lines CR-11 and -13 was 82.5 ± 13.7 cm and 68.7 ± 6.0 cm respectively, much lower than the control (100.5 ± 3.9 cm; Table S2). At the same time, the height of most of the moderately down-regulated lines was higher, while the square (flower bud) number at 105 DAS of most of the moderately down-regulated lines was higher, while the square number of the two severely
suppressed lines was lower than that of the wild type. After harvest, we counted the boll number per plant and calculated the seed and lint fiber yields. The agronomic behavior of the slightly down-regulated line, CR-7, was very close to the wild type (Table S2). The moderately down-regulated cottons produced more bolls than the wild type. The averaged boll number of the five moderately down-regulated lines was 30.7 ± 1.7, 19.0% more than the wild type. Moreover, bolls produced by these lines were bigger than the wild type, as indicated by the seed cotton weight per boll and seed index (the weight in grams of 100 seeds, Table S2). Consequently, seed cotton yield per plant of the lines increased significantly. In contrast, the number of bolls and the seed size in the severely suppressed cottons, CR-11 and CR-13, were lower and smaller than the wild type.

Fig. 2 Chlorophyll contents of upper, middle, and lower leaves in 90-, 105- and 120-DAS plants. Results are presented as mean ± SD (n = 3). Asterisks (*) indicate a significant difference (P < 0.05). FW fresh weight

Fig. 3 Comparison of photosynthetic rate in leaves, and soluble sugar and protein concentrations in bolls between transgenic lines and wild type. a Photosynthetic rate of the fourth leaf from the apex at 120 DAS. Error bars indicate SD of 10 randomly selected plants of each line. b Soluble sugar content of 35-DPA ovules and boll shells. c Soluble protein content of 35-DPA ovules and boll shells. Error bars indicate SD of data of three replicates for each line. For each column, means that are not followed by the same letter are significantly different according to Tukey’s range test at the 0.05 level.
Interestingly, lint index (the weight in grams of lint in 100 seeds), a measure of the abundance of fiber on the seed, and the lint yield per plant increased in the moderately down-regulated lines, demonstrating that the enhancement of seeds was not at the cost of lint fibers; instead, both seed and lint yield were concurrently improved. However, boll number per plant, seed number and seed cotton per boll, and seed cotton yield and lint yield per plant of the severely suppressed lines CR-11 and CR-13 were dramatically lower than the wild type (Table S2), suggesting that over-dosage of endogenous cytokinins reduced both seed and lint yield. With regard to the quality of the fiber, we did not find notable alterations in fiber length, fiber strength, or micronaire in transgenic cottons (Table S3), suggesting that the increase of endogenous cytokinins in cotton had little influence on the qualities of transgenic fibers.

To further assess the performance of the moderately down-regulated GhCKX cotton in yield improvement, we selected CR-6 (T2) as the best line for field trials. As in T2 generation, yield components, including boll number per plant, seed cotton per boll, lint index and seed index, of the transgenic line were higher than that of the wild type. The seed yield and lint yield increased 15.4 and 20.0 %, respectively, as compared to the wild type (Table 2). The field trial results confirmed that moderate down-regulation of GhCKX enhanced both fiber and seed yield.

**Discussion**

Cotton fibers are derived from single cells of ovule epidermis (Stewart 1975) and fiber cell development is tightly influenced by seed (Ruan 2013). Phytohormones play important roles in the growth and development of seed and fiber cells of cotton (Beasley and Ting 1973; Basra and Saha 1999; Lee et al. 2007). Indole-3-acetic acid and gibberellins were able to promote fiber cell initiation and elongation (Chen et al. 1988, 1996; Zhang et al. 2011; Xiao et al. 2010). Ethylene plays a major role in promoting fiber elongation (Shi et al. 2006), whereas abscisic acid (ABA) inhibits fiber growth (Dhindsa et al. 1976; Gokani et al. 1998; Lee et al. 2007). brassinosteroids (BRs) participate in both fiber initiation and elongation (Sun et al. 2005; Luo et al. 2007). The inhibitory effect of BRZ on fiber cells could be overcome by

| Line | Boll number | Seed cotton yield (kg/plot) | Lint yield (kg/plot) | Seed number per boll | Seed index (g) | Seed cotton index (g) | Lint index (g) | Micronaire (Fiber strength) | Fiber length (mm) |
|------|-------------|-----------------------------|---------------------|---------------------|----------------|----------------------|---------------|---------------------------|------------------|
| WT   | 23.5 ± 1.3  | 4.7 ± 0.1                   | 39.5 ± 0.2          | 6.9 ± 0.2           | 10.6 ± 0.3     | 11.2 ± 0.1           | 7.5 ± 0.4     | 5.1 ± 0.1                 | 30.8 ± 0.2 |
| CR-6 | 25.6 ± 1.4  | 4.9 ± 0.1*                  | 39.8 ± 0.5          | 7.4 ± 0.1*          | 11.2 ± 0.1*    | 7.5 ± 0.4*           | 4.5 ± 0.2*    | 4.5 ± 0.2*                | 31.0 ± 0.2 |

Plants were grown in the farm at Southwest University (Chongqing, China) in a randomized comparative trial with three replications. The plot size was 25 m². Results are presented as mean ± SD (n = 3). Asterisk (*) indicates significant differences at P < 0.05 according to Student’s t-test.
ethylene, indicating the cross-talk between BRs and ethylene (Shi et al. 2006). Cytokins have an important role in shoot and seed development, but fiber elongation is inhibited by high concentrations of cytokins in ovule culture (Beasley and Ting 1974). However, the effects of cytokins on the yield of cotton fiber and seeds remained to be evaluated.

To elevate the endogenous level of cytokins, we successfully generated 35S::GhCKXRNAi cottons. A dose-dependent effect of the gene expression on phenotype alterations was observed. The slightly suppressed cottons did not show distinct morphological and agronomic changes relative to the wild type, while moderate suppression resulted in a delay in leaf senescence, increase in photosynthesis, more fruiting branches and bolls, bigger size of seed, and consequently higher yield of seed and fiber. Along with the decrease in GhCKX expression, the negative impact of cytokinin overproduction on plant growth and development became progressively more severe, and the yield of seed and fiber of the severely suppressed cottons dramatically declined. The cytokinin content assay was well consistent with the dosage change in gene expression. That is to say, along with the decline in gene expression, the endogenous level of cytokins increased and the enhancement of endogenous cytokinins creates the alterations taking place in the transgenic cottons.

The increased cytokinin content caused an enhancement of seed yield in rice (Ashikari et al. 2005) and Arabidopsis (Bartrina et al. 2011). Our ovule culture results demonstrated that application of ZT (trans-Zeatin) at appropriate concentrations (5.0–8.0 μM) in media promoted seed development. However, a high concentration of the cytokinin (15 μM) inhibited seed development (Fig. S3). Results in transgenic 35S::GhCKXRNAi cottons supported this observation. Ovules of the severely suppressed cotton contained high concentration of cytokinins (Tables 1 and S1) and the cotton plants produced less seeds with smaller size, whereas the seed number, seed size, and seed cotton yield were significantly enhanced in the moderately suppressed cottons (Table S2). These results indicated that a moderate enhancement of cytokinins promotes the development of seeds, but overdosage of them inhibits it.

In the ovule culture experiment, fiber elongation was inhibited in cytokinin-overproduction transgenic cottons (Yu et al. 2000). In our experiments, no distinct inhibition of fiber initiation was observed in moderately suppressed transgenic cottons (Fig. S4). Meanwhile, there was no distinct difference in fiber length between transgenic lines and the wild type (Table S3), implying that moderate overexpression of GhCKXRNAi has little negative effect on fiber development. Instead, lint index and lint yield of the moderately suppressed GhCKX lines were significantly enhanced. There are two explanations for this enhancement. First, the number of fibers per seed is limited by the surface area of the ovule epidermis. Thus, enhancement of cotton seed development could achieve cotton seed and fiber yield improvement (Ruan 2013). In our experiments, a proper increase of cytokinins promoted seed development and enlarge the seed size (Fig. S3). Bigger seed means larger area of seed coat that bears more fibers. More importantly, early leaf senescence could decrease the assimilation supply for boll and fiber development (Bauer et al. 2000; Peng and Krieg 1991), thus reducing lint yield and fiber properties (Wright 1999; Dong et al. 2006). The delay of leaf senescence in transgenic cottons leads to an increase in chlorophyll concentration and photosynthetic rate of the leaves. Our results are in agreement with previous findings that early-fruit removal significantly increased cytokinins in both main-stem leaves and xylem sap and delayed the main-stem leaf senescence characterized by increases in chlorophyll concentration and photosynthetic activity of leaves (Dong et al. 2009). The enhanced photosynthesis in the leaves of transgenic cotton plants and the increased soluble sugar and protein of the boll (Fig. 3) provide more carbohydrates for the development of seed and fiber. As a result, seed yield and fiber yield concurrently improve.

Although seed-specific expression of ipt can alleviate developmental abnormalities resulting from constitutive overexpression of the gene (Ma et al. 2002, 2008; Daskalova et al. 2007), the shortcoming of this strategy is that the advantages associated with increase in cytokinins, such as delay in leaf senescence, increase in photosynthesis of leaves, more branches, tolerance to abiotic and biotic stress (Choi et al. 2010; Jeon et al. 2010; Nishiyama et al. 2011; Peleg et al. 2011; Rivero et al., 2007), and resistance to pathogens (Choi et al. 2011), may not benefit the whole plant. In our previous experiments, we failed to generate transgenic 35S::ipt cotton. In contrast, in this paper, using the strategy of suppressing cytokinin
deactivation, we successfully obtained transgenic 35S::GhCKXRNAi cottons in which endogenous cytokinin levels were globally and constitutively enhanced (Tables 1 and S1). In terms of manipulation of cytokinin levels in plants, CKX is more likely a softer regulator than ipt. Taken together, our data demonstrated that moderately enhancing endogenous cytokinins by suppressing CKX is a feasible and effective strategy for yield improvement, not only for cotton but also for other seed crops, such as canola, soybean, maize, and rice.

Acknowledgments This work was supported by the National Natural Science Foundation of China (31130039 to Y. P) and Chinese Ministry of Science and Technology (2010CB16000 to Y. P), and Ph.D. Programs Foundation of Southwest University, China (Rb20100005 to J. Z).

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