Stearoyl-ACP Δ⁹ Desaturase 6 and 8 (GhA-SAD6 and GhD-SAD8) Are Responsible for Biosynthesis of Palmitoleic Acid Specifically in Developing Endosperm of Upland Cotton Seeds

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Palmitoleic acid (16:1 Δ⁹) is one kind of ω-7 fatty acids (ω-7 FAs) widely used in food, nutraceutical and industry. However, such high-valued ω-7 FA only has a trace level in mature seeds of cotton and other common oil crops. We found that palmitoleic acid (>10.58 Mol%) was specially enriched in developing cotton endosperm which is disappeared in its mature seed. The present study was conducted to investigate the mechanism underlying high accumulation of palmitoleic acid in developing endosperm but not in embryo of upland cotton (Gossypium hirsutum L.) seed. Of 17 stearoyl-ACP Δ⁹ desaturases (SAD) gene family members identified in upland cotton, six GhSADs may specifically work in the desaturation of palmitic acid (16:0-ACP) to produce palmitoleic acid (16:1 Δ⁹-ACP), which were revealed by examining the key amino acids in the catalytic center and their cis-elements. Gene expression analysis showed that spatial patterns of these GhSADs were different in developing ovules, with GhA-SAD6 and GhD-SAD8 preferentially expressed in developing endosperms. Functional analysis by transient expression in Nicotiana benthamiana leaves and genetic complementary assay using yeast mutant BY4389 strain unable to synthesize unsaturated fatty acids demonstrated that GhA-SAD6 and GhD-SAD8 have strong substrate specificity for 16:0-ACP. In contrast, GhA-SAD5 and GhA-SAD7 exhibited high specific activity on 18:0-ACP. Taken together, these data evidence that GhA-SAD6 and GhD-SAD8 are responsible for making palmitoleic acid in developing cotton endosperms, and provide endogenous gene targets for genetic modification to enrich ω-7 FAs in cotton seed oil required for sustainable production of functionality-valued products.

Keywords: cotton, ACP-Δ⁹ desaturase, palmitoleic acid, endosperm, substrate specificity

INTRODUCTION

Cotton (Gossypium sp.) is widely cultivated in the world as an important industrial crop, providing excellent fiber and protein plus oil. Particularly, cottonseed oil is an important vegetable oil and broadly used in food/catering, clean biofuel and other industries (Jiao et al., 2013; Sturtevant et al., 2017). Fatty acid composition and its content determine the value and utilization
of vegetable oil. The conventional cottonseed oil mainly consists of 26% palmitic acid (16:0), 58% linoleic acid (18:2\(\Delta 9,12\)), 13% oleic acid (18:1\(\Delta 9\)) and 2% stearic acid (18:0) (Zhao et al., 2018). Since linoleic acid is oxidatively unstable, cottonseed oil is easily oxidized and rancid, and thus it normally needs partial hydrogenation before food applications. However, byproducts of this process are trans-fatty acids, which are harmful for human health (Ginter and Simko, 2016). Saturated palmitic acid is not recommended for direct edible use because of its detrimental effects on human health despite of its highly oxidative stability (Shang et al., 2017). Monounsaturated fatty acids (MUFRs) such as oleic acid and palmitoleic acid (16:1\(\Delta 9\)) are health-promoting fatty acids, but they are in tiny content in cottonseed oil. Therefore, the development of designer cottonseed oil with nutritionally desirable fatty acids has been an attractive object to meet oil marketing demand.

Omega-7 fatty acids (\(\omega-7\) FAs), are monounsaturated lipids with a double-bond seven atoms away from the methyl end of the acyl chain, including palmitoleic acid (16:1\(\Delta 9\)), and its elongation products, vaccenic acid (cis-18:1\(\Delta 11\)) and paullinic acid (cis-20:1\(\Delta 12\)). \(\omega-7\) FAs are high-valued in human healthcare, but also in multiple industrial applications, ranging from the precursor of 1-octene for polyethylene to the feedstock of biodegradable lubricant and biodiesel (Curb et al., 2000; Misra et al., 2010; Mozaffarian et al., 2010; Nguyen et al., 2010, 2015). Unfortunately, such valued \(\omega-7\) FAs are very low abundance in commercialized field oilseeds like cotton and soybean (Glycine max). Only a few of uncommercialized plants can synthesize and accumulate much high level of \(\omega-7\) FAs in their seeds (Wu et al., 2012). For example, contents of \(\omega-7\) FAs are high up to 64%, 30%, and 32% in the mature seeds of cat’s claw (Doxantha unguis-cati L.), macadamia (Macadamia integrifolia), and sea buckthorn (Hippophae rhamnoides), respectively (Cahoon et al., 1998; Giummeson et al., 2000; Kallio et al., 2002). Interestingly, \(\omega-7\) FAs were found to be highly concentrated in the endosperm (aleurone) but not in the embryo in the seeds of Arabidopsis thaliana and rapeseed (Brassica napus), with 15 mol% in Arabidopsis and 35% in rapeseed, respectively (Bryant et al., 2016). It is noteworthy that the endosperm is reduced to a single cell layer of residues surrounding the embryo in these mature seeds (Penfield et al., 2004; Li et al., 2006). Such a thin peripheral aleurone and seed coat are tightly stuck together, and consequently are stripped off the embryo during the dehulling process before oil extraction. The detailed molecular mechanism responsible for such high biosynthesis of \(\omega-7\) FAs in the resource plants but not in the embryo of common oilseeds remains to be elucidated despite a number of enzymes and transcriptional factors were found to play important roles in formation of \(\omega-7\) FAs (Bryant et al., 2016; Troncoso-Ponce et al., 2016).

In plants, fatty acyl-ACPs of different chain lengths and saturation levels are initially biosynthesized in plastid. Then, fatty acyl-ACPs are catalyzed by fatty acyl-ACP thioesterases to release free fatty acids chains, which are transported from plastid via cytosol into endoplasmic reticulum (ER) for functional modifications, such as acyl chain elongation and further desaturation. Finally, fatty acids are channeled into triacylglycerol (TAG) by cascade acylation reactions. Alternatively, fatty acids could also be integrated into phosphatidylcholine (PC) and other acyl carrier molecules.

It is believed that \(\omega-7\) FAs are produced by \(\Delta 9\) desaturation of palmitic acid (16:0) to form palmitoleic acid (16:1\(\Delta 9\)), which can then be elongated (Cahoon et al., 1998). Such \(\Delta 9\) desaturation is catalyzed by acyl-ACP \(\Delta 9\) desaturase (AAD, EC 1.14.99.6) family (Parvini et al., 2016). The archetype for this family is the stearoyl-ACP \(\Delta 9\) desaturase (named shortly as SAD) which normally desaturates stearic acid (18:0) to form oleic acid (18:1\(\Delta 9\), cis-\(\omega-9\)) at high efficiency in plants. However, various AAD isoforms also exhibit different levels of substrate specificity on 16:0-ACP (McKeon and Stumpf, 1982). Those AAD isoforms with preference on 16:0-ACP instead of 18:0-ACP as substrate are also named as palmitoyl-ACP \(\Delta 9\) desaturases (PADs), and some of which were isolated from several high accumulators of \(\omega-7\) FAs (Cahoon et al., 1997, 1998).

It is therefore recognized that the specific activity of SAD enzymes for substrates determines the ratio of 16:1\(\Delta 9\) to 18:1\(\Delta 9\) and their derivatives (\(\omega-7\) and \(\omega-9\) FAs) in plant seeds. The key amino acids located in the catalytic domain were reported to play important roles for the substrate specificity of SAD enzymes (Whittle and Shanklin, 2001; Zhang Y. et al., 2015), but knowledge is still limited for the molecular determination of such substrate specificity. Recently, Arabidopsis AtAAD2 and AtAAD3 were identified as the major contributors for the synthesis of \(\omega-7\) FAs in the aleurone, though seven AtAAD family members were detected in Arabidopsis genome (Bryant et al., 2016; Troncoso-Ponce et al., 2016).

As described above, cottonseeds contain high level of 16:0 (\(\sim 25\%\)) and 18:2\(\Delta 9,12\) (\(>50\%\)), but little 16:1\(\Delta 9\) (\(<1\%\)) and 18:1\(\Delta 11\) (\(<1\%\)), showing that SAD enzymes responsible for \(\Delta 9\) desaturation in upland cotton may have higher selectivity for 18:0-ACP than for 16:0-ACP. However, RNAi down-regulation of \(\text{KASII}\) resulted in remarkable increase of 16:1\(\Delta 9\) and 18:1\(\Delta 11\) in cotton seeds by 10% and 1.5%, respectively, indicating the existence of SAD isoforms having high specific activity to catalyze 16:0-ACP to 16:1-ACP (Liu et al., 2017). More importantly, we examined fatty acid profiles in separate tissues from cotton seeds, revealing that a certain amount of 16:1\(\Delta 9\) were concentrated in developing endosperm but not in embryo. Overall, there should be existed some SAD members responsible for biosynthesis of 16:1\(\Delta 9\) in upland cotton. How many members of SAD family are in allotetraploid upland cotton? Is there any SAD isoform having a higher substrate specificity toward 16:0-ACP or 18:0-ACP? If so, what are the distinct structure features between 16:0- and 18:0-specific SAD isoforms? How they work coordinately in controlling the biosynthesis of \(\omega-7\) and \(\omega-9\) FAs in cotton seeds? Particularly, which members of SAD family in cotton are responsible for \(\omega-7\) FA enrichment in developing endosperm but not in embryo? Since endosperm would be vanished from the mature seeds, what physiological roles are ascribed to \(\omega-7\) FAs exclusively accumulated in developing endosperm? All these questions need to be addressed.

Here, a genome-wide characterization was performed to identify candidate SAD family members in upland cotton (Gossypium hirsutum L.), a naturally occurring allotetraploid (2n = 4x = (AD)\(_2\) = 52) produced by interspecific hybridization
of A- and D-genome diploid progenitor species (Zhang T. et al., 2015). Analysis on the key amino acids in the functional domain of GhSAD members and their cis-elements as well as 3D structure were conducted to identify the putative 16:0-specific GhSADs. And then, the temporal-spatial expression patterns of these GhSAD genes were examined by both RNA-seq data and qRT-PCR in various cotton tissues and developing seeds. Subsequently, for functional analysis of the target GhSADs, we employed the transient expression of these GhSAD genes in tobacco (Nicotiana benthamiana) leaf tissues and the heterologous expression in yeast (Saccharomyces cerevisiae) mutant BY4389 lacking the synthesis of unsaturated fatty acids, followed by detecting fatty acid profiles in the target samples. In the whole, the present data evidence that Gha-SAD6 and GhD-SAD8 are responsible for making palmitoleic acid in developing cotton endosperm, providing novel insight to understand oil and ω-7 FA biosynthesis in plant seeds and scientific reference for lipid metabolic engineering in commercial oilseeds.

MATERIALS AND METHODS

Plant Materials

Upland cotton (G. hirsutum cv. Zhong Mian 21) was planted in the experimental station at Shanxi Agricultural University, Taigu, China. Roots, stems and leaves were sampled from 6-week-old seedlings. Whole flower at 0 day after flowering (DAF) and developing ovules were sampled at 5, 10, 15, 20, 25, 35, and 45 DAF. Then ovules at 15, 25, 35 and 45 DAF were dissected manually to separate the embryos from the endosperms. All samples were instantly frozen in liquid nitrogen and stored at −70°C for subsequent experiment. Tobacco (N. benthamiana) was grown in an artificial incubator keeping at 26°C, 60% relative humidity and 14 h light/10 h dark photoperiod (Zhang et al., 2014).

Gene Identification, Functional Domain, Three-Dimensional Model and Promoter Analysis

The conserved FA_desaturase_2 domain (PF03405) was retrieved from Pfam database and used as the query to perform BLAST against proteins of G. hirsutum from the CottonGen database. The sequence features were identified by HMMER version 3.0 with default parameters, and their conserved motifs were further verified by SMART and CDD tool. Only the sequence containing conserved FA_desaturase_2 domain was employed for further analysis. All candidate GhSAD genes were renamed according to their chromosomal distributions, denoting GhA-SAD1~GhA-SAD9 and GhD-SAD1~GhD-SAD8. Transcriptional data were downloaded from cotton research institute of Nanjing Agricultural University.

For cis-element analysis, the extracted 2-kb upstream sequences from the initiation codon of all GhSADs were searched in the PlantCARE database. The data of AtSAD proteins were downloaded from TAIR database. Castor SAD sequences and SAD in cat’s claw (Muc-PAD) were downloaded from NCBI. To investigate the key amino acid residues that influence the catalytic properties of SAD enzyme, amino acid sequence comparisons of all GhSAD proteins and the reference protein sequences (RcSAD1, AtFAB2, AtAAD3, AtAAD2, and Muc-PAD) were conducted. Then, three-dimensional structures of GhSADs were modeled using Swiss-model, with the crystal structure of RcSAD1 (NP_001310659.1) as the template. In addition, the key amino acids and ligands in catalytic activity center were marked using Discovery Studio 4.1 software, with the default parameters.

RNA Isolation and Real-Time Quantitative PCR

EASYspin Plus Plant RNA Kit (Aidlab, Beijing, China) was used for the isolation of total RNA from all sample types. First-strand cDNA was reversely transcripted using PrimeScriptTM II 1st Strand cDNA Synthesis Kit (Takara, Kyoto, Japan) according to the manufacturer’s instruction. Gene-specific primers used were listed in Supplementary Table S1. The experiment was carried out on a Bio-Rad CFX96 Real-Time PCR system (Bio-Rad, United States) with three replicates. Following reaction conditions were applied: 95°C for 10 min; 40 cycles of 95°C for 15 s, 60°C for 1 min; 72°C for 20 s. The melting curve was analyzed after the final cycle, ramping from 65 to 95°C with increment 0.5°C/5 s. The normalized relative expression was calculated using the \( 2^{-\Delta \Delta Cq} \) method (Livak and Schmittgen, 2001). Cotton EF-1α (DQ174251) and histone 3 (AF024716.1) genes were employed as the internal references (Jiang et al., 2012; Shang et al., 2015).

Construction of Plant and Yeast Expression Vectors of the GhSAD Members

To test their substrate preferences, the full-length open reading frames (ORFs) of four representative GhSAD genes (GhA-SAD5, GhA-SAD6, GhA-SAD7, and GhD-SAD8) were amplified by PCR with oligonucleotides (Supplementary Table S2) and then cloned into pCAMBIA1303 vector under the drive of cauliflower mosaic virus (CaMV) 35S promoter. The resulting vectors were transferred into Agrobacterium strain of GV3101 using the freeze-thaw method for further transient expression in tobacco leaves by infiltration. The empty vector pCAMBIA1303 was used as blank control. For functional analysis using yeast expression system, the target GhSAD genes with optimized codons were then cloned into the yeast expression vector pYES2 under the control of the GAL1 promoter. Additionally, AtFAB2 and AtAAD3 from A. thaliana were cloned according to the published sequences.
and inserted into pYES2 as positive controls. Empty vector of pYES2 was set as blank control. Related primers were listed in Supplementary Table S2.

**Transient Expression Assays in *N. benthamiana* Leaves**

The constructed plant expression vectors were transferred into *Agrobacterium tumefaciens* strain GV3101. Agroinfiltration was employed for transient expression of the target genes in tobacco leaves. Briefly, the bacteria bearing the vector were cultured in LB medium and incubated at 28°C overnight with 250 rpm. Bacterial cultures were then centrifuged at 8,000 rpm for 3 min and resuspended into fresh infiltration medium supplemented with acetosyringone, MgSO$_4$ and MES to an optical density of 0.5 at 600 nm. Agroinfiltration was performed using a needle-free syringe in the leaves of six-week-old *N. benthamiana* plants. The treated leaves were harvested 5 days later, and then ground up into powder for the extraction of total lipids.

**Yeast Culture and Transformation**

The mutant yeast *S. cerevisiae* strain BY4389 (MATa ole1 Δ::LEU2 ura3-52 his4) that cannot synthesize unsaturated fatty acids was purchased from Osaka University, Japan (Xue et al., 2016). Before transformation, the cells were cultivated at 28°C in modified yeast-extract peptone dextrose (YPD) medium (0.01% sodium oleate, 1% yeast extract, 2% peptone and 2% glucose) with shaking (200 rpm). When the concentration was at OD600≈0.5, the cells were harvested by centrifugation, washed three times with precooling sterilized water. Then, the cells were resuspended in 100 mM LiAc solution. After removing LiAc, we got the competent cells of yeast with high transformation efficiency. The constructed yeast expression vector and empty pYES2 were separately transformed into *S. cerevisiae* mutant strain BY4389 using Yeast Transformation Kit (Coolaber, Beijing, China). Transformed cells were selected on the modified YPD medium lacking uracil (SC-Ura). After cultured 3 days at 28°C, PCR confirmation was used to select positive clones. The correct transformed cells were then transferred in the corresponding selective medium to increase the yeast biomass for 24 h. For induction, glucose was replaced by galactose (2%, w/v) and sodium oleate was removed, and cells were further cultivated for another 72 h. At last, transformed cells were harvested and freeze-dried into powder to for fatty acid analysis.

**Lipid Extraction and Fatty Acid Methyl Ester Analysis**

Total lipids from developing cotton embryo and endosperm (15, 25, 35 and 45 DAF), the wild and transgenic *N. benthamiana* leaves and yeast samples were extracted, and then converted into fatty acid methyl esters (FAMEs) as previously described (Bhattacharya et al., 2015). Briefly, FAMEs were transmethylated using 2 mL of methanol that contained 2.5% sulfuric acid (v/v) and then incubated at 85°C for 2 h. For quantification of the FAs, an appropriate amount of C17:0 FAME (Sigma) was added as internal standard. FAMEs were analyzed by Agilent gas chromatograph equipped with a flame ionization detector and a capillary column (HP-88 100 m × 0.25 mm × 0.2 μm). Nitrogen was used as carrier gas with a flow rate of 2.64 ml/min. 1 μL of the sample was injected into the column with three replicates. The injection and detector temperatures were both set as 250°C. The gas chromatograph oven was programmed at an initial temperature of 140°C for 5 min, and then increased up to 250°C at a rate of 15°C/min. Fatty acid component was estimated according to the retention time of fatty acid standard and data was collected by peak area normalization with C17:0 internal standard.

**RESULTS**

**Differential Distribution of Fatty Acids Between Embryos and Endosperms in Developing Cottonseeds, Especially for Palmitoleic Acid**

Angiosperm seed formation is characterized by the double fertilization of the embryo sac. During this process, one sperm cell fertilizes the egg cell to produce a zygote, which develops into an embryo, whereas the second sperm cell targets the central cell polar nuclei, leading to the endosperm formation. Both the diploid embryo and the triploid endosperm are enclosed by the maternally derived ovular integument layers that form seed coat. As a typical exalbuminous seed, cotton seed has no endosperm left at mature stage despite abundant endosperms surrounding the embryo are generated during the early seed development, providing nutrition for the enlarging embryo as it develops (Figure 1A).

To investigate whether fatty acid profiles are different between embryos and endosperms during development, we harvested embryo and endosperm samples, respectively, at four stages of seed development denoted as 15 DAF (days after flowering), 25, 35, and 45 DAF (Figure 1A). At the early stage of 15 DAF, the endosperm forms a thick milky layer surrounding the embryo. During seed development and maturation, the endosperm volume is gradually reduced into a thin layer closely to seed shell (25–35 DAF), accompanied by the embryo growing. At the end of seed development (45 DAF), the endosperm is disappeared in cotton seed.

The examination of FA contents in developing cotton seeds (Figure 1B) showed that FA profiles were significant different between endosperm and embryo. The most difference is that 16:1Δ9, a kind of ω-7 FAs, was largely accumulated in endosperm with the peak at 25 DAF (16.95 Mol% of total FAs), but only trace level (<1 Mol%) in embryo. However, content of 18:1Δ9, a kind of ω-9 FAs, was higher in embryo than in endosperm by 1.2–1.7 folds, with the maximum level (15.04 Mol%) in embryo at 25 DAF. Palmitic acid (16:0) was the major FA in endosperm at 15 DAF, and it reduced gradually during seed development, with lower level in endosperm than in embryo from 25 to 35 DAF. Linoleic acid (18:2Δ9,12), the major FA in embryo, is increased up to ~60% in embryo at 25 DAF, with lower level in developing endosperm. The contents of 18:0 and 18:3Δ9,12,15 were not significant between embryo and endosperm from 25 to 35 DAF.
despite these two fatty acid contents were higher in embryo than in endosperm at 15 DAF of the seed development. Notably, abundant palmitoleic acid (16:1Δ9) were found to be selectively accumulated in developing endosperm, suggesting that some unknown mechanism may control this differential accumulation of palmitoleic acid between the endosperm and embryo.

Identification of Seventeen Stearoyl-ACP Δ9 Desaturases (GhSADs) and Six Putative 16:0-ACP Specific GhSADs From Upland Cotton

As described in the introduction, some members (if not all) of SAD family can catalyze palmitic acid (16:0)-ACP to form palmitoleic acid (16:1Δ9)-ACP in plant seeds. The preferred accumulation of 16:1Δ9 in the developing cotton endosperm observed above indicates that SAD enzyme working in endosperm may have high substrate specificity for palmitic acid (16:0). To investigate such SAD members with high 16:0-ACP specificity, we performed a genome-wide characterization of SAD family members in upland cotton (G. hirsutum) genome. Total of 17 putative GhSADs were identified according to the conserved domain and their integrity. These predicted GhSADs were designated as GhA-SAD1~GhA-SAD9 and GhD-SAD1~GhD-SAD8 based on chromosome number of A- or D-genome where they located (Supplementary Table S3).
The known reports have established that RcSAD1 from castor (*Ricinus communis*) (Lindqvist et al., 1996) and AtFAB2 from Arabidopsis (Lightner et al., 1994) are characterized as the 18:0-ACP specific SADs while AtAAD2 and AtAAD3 from Arabidopsis (Troncoso-Ponce et al., 2016) as well as Muc-PAD from cat's claw (*Macfadyena unguis-cati*) (Cahoon et al., 1998) are identified as the 16:0-ACP specific enzymes. Moreover, the substrate specificity of these enzymes is determined by a group of eight amino acid residues comprising the region of the catalytic sites (also referred as the lower portion of the hydrophobic binding pocket for FA). To identify which GhSAD may be the candidate of 16:0-ACP specific enzyme, these well-characterized SADs were used as the references to perform in silico analyses of 17 GhSAD sequences including sequence comparisons and modeling of 3D protein structures (Arnold et al., 2006; Bordoli et al., 2009).

Table 1 summarized the comparison of the eight amino acid (AA) residues lining the lower portion of the hydrophobic binding pocket for FA in those enzymes tested. Seventeen GhSADs were clearly divided into two groups. Group 1 with ten GhSAD members had the identical AA residues in the conserved functional domains as RcSAD1 and AtFAB2 (Table 1), which are ω-9 producing SAD isoforms. Group 2 consisting of GhA-SAD5, 6, 7, and GhD-SAD4, 5, 7, 8 were gathered together with 16:0-ACP specific Muc-PAD, AtAAD2 and AtAAD3, although divergent residues existed in the eight-AA domain of those enzymes. The 3D model revealed that the eight AA residues were located near the bottom of the substrate-binding channel close to diiron center of the enzymes examined (Figure 2I). The identical eight-AA domain in the 18:0-ACP specific enzymes is predicted to shape a deep/long substrate cavity (pocket/channel) to accommodate 18:0-ACP substrate (Figures 2A). However, the divergent eight-AA domain in the 16:0-ACP specific enzymes is predicted to make the substrate channel shorter, which is more adaptable for 16:0-ACP substrate (Figures 2B,C,D).

To further test whether these GhSADs in Group 2 are the candidate enzymes responsible for 16:1<sup>ω-9</sup> biosynthesis in developing cotton endosperm, *cis*-elements analyses were conducted on the promoter sequences of the corresponding genes. The promoters of *GhA-SAD5*, *GhA-SAD6*, *GhA-SAD7*, *GhA-SAD9*, *GhD-SAD1*, *GhD-SAD2*, *GhD-SAD3*, and *GhD-SAD6* were used as references to perform in silico analyses of 17 GhSAD sequences including sequence comparisons and modeling of 3D protein structures (Arnold et al., 2006; Bordoli et al., 2009).

### Table 1 | Comparison of the eight amino acid residues comprising the region of the catalytic sites of GhSADs.

| Ricinus communis | Arabidopsis thaliana | Gossypium hirsutum |
|------------------|----------------------|--------------------|
| RcSAD1 | AtFAB2 | GhA-SAD1 | GhA-SAD2 | GhA-SAD3 | GhA-SAD4 | GhA-SAD8 | GhA-SAD9 | GhD-SAD1 | GhD-SAD2 | GhD-SAD3 | GhD-SAD6 |
| L118 | L156 | L141 | L150 | L140 | L151 | L150 | L150 | L150 | L141 | L151 | L150 |
| F189 | F227 | F212 | F211 | F211 | F212 | F221 | F221 | F221 | F212 | F222 | F221 |
| L115 | L153 | L138 | L147 | L137 | L148 | L147 | L147 | L147 | L138 | L149 | L147 |
| T117 | T155 | T140 | T149 | T139 | T150 | T149 | T149 | T149 | T149 | T150 | T149 |
| G188 | G226 | G211 | G210 | G210 | G211 | G220 | G220 | G220 | G220 | G211 | G220 |
| T181 | T219 | T204 | T213 | T203 | T214 | T213 | T213 | T213 | T204 | T214 | T213 |
| P179 | P217 | P202 | P211 | P201 | P212 | P211 | P211 | P211 | P202 | P212 | P211 |
| M114 | M152 | M137 | M146 | M136 | M147 | M146 | M146 | M146 | M137 | M148 | M146 |

| Arabidopsis thaliana | Dauricum unguis-cati | Gossypium hirsutum |
|----------------------|---------------------|--------------------|
| AtAAD3 | AtAAD2 | Muc-PAD | GhD-SAD4 | GhD-SAD5 | GhD-SAD6 | GhD-SAD7 | GhD-SAD8 |
| L153 | L163 | W152 | L151 | L31 | L150 | L152 | M148 | M148 | L151 |
| F224 | F234 | F223 | F222 | F102 | F221 | F223 | A219 | A219 | Y222 |
| L150 | L160 | L149 | L149 | L28 | L147 | L149 | L145 | L145 | L148 |
| T152 | T162 | T151 | T150 | R30 | R149 | T151 | T147 | T147 | T150 |
| G223 | G233 | G222 | G221 | T101 | T220 | T222 | G218 | G218 | A221 |
| F218 | F236 | F215 | F214 | I94 | I213 | A215 | S211 | S211 | A214 |
| S214 | S224 | P213 | P212 | V192 | V211 | P213 | S209 | S209 | M212 |
| T149 | T159 | M148 | M148 | I48 | I27 | M146 | M148 | L144 | M147 |

Sequences were conducted by GeneDoc software using RcSAD1, AtFAB2, AtAAD3 as well as Muc-PAD as templates. The GenBank or TAIR accession numbers of the temple sequences listed as the follows: RcSAD1 (NP_001310659.1) from *R. communis*; AtFAB2 (At2g43710), AtAAD2 (At3g02610), and AtAAD3 (At5g16230) from *A. thaliana*; Muc-PAD (AFV61670.1) from *D. unguis-cati*. Orange and pink color represent varied amino acids residues in the domain of 16:0-ACP specific isoforms and the putative 16:0-ACP specific GhSADs, respectively, compared to the 18:0-ACP specific isoforms characterized.
GhD-SAD5, GhD-SAD7 and GhD-SAD8 contained multiple skn-1 motifs with the core sequences of GTCAT which is necessary for gene expression in endosperm (Zhang et al., 2013). In addition, GhD-SAD5 and GhD-SAD8 have another GCN4 motif (the core sequences is TGAGTCA) involved in gene expression in endosperm (Supplementary Table S4) (Song et al., 2012). However, the promoter of GhD-SAD4 lacked such cis-element.

Overall, sequence analysis indicate that GhA-SAD5, 6, 7, and GhD-SAD5, 7, 8 may be the putative 16:0-ACP specific SADs responsible for palmitoleic acid biosynthesis in developing cotton endosperm.

**Expression Patterns of GhSAD Genes in Different Cotton Tissues**

If any of GhSADs identified above functions in endosperm palmitoleic acid production, it must be expressed in developing cotton endosperm. To determine this, we firstly examined their transcript abundance during developing ovules using cotton RNA-seq profiles publically available (see text footnote 5). As shown in Figure 3, these 17 GhSADs exhibited different expression patterns in developing ovules. High expressions of GhD-SAD7, GhA-SAD3, and GhD-SAD4, occurred at the early stages of ovules. GhA-SAD2, GhD-SAD1 and GhD-SAD3 abundantly appeared at 5 DAF. Seven GhSAD (GhA-SAD5, GhA-SAD6, GhD-SAD5, GhA-SAD-7, GhD-SAD8, GhA-SAD1, and GhD-SAD2) transcripts greatly accumulated at the middle stage of ovules (from 10 DAF to 25 DAF), while GhA-SAD4 and GhD-SAD6 largely expressed at the late stage of ovules (35 DAF).

To further verify the expression patterns of these GhSADs, we employed qRT-PCR to investigate their mRNA levels in various cotton tissues (Figure 4). Six GhSADs with high expressions were selected for this verification. Specific primers for qRT-PCR were designed at 3′ untranslated regions of the GhSADs because of their highly conserved ORF sequences (Supplementary Table S1). As shown in Figure 4, GhA-SAD5, GhA-SAD6, and GhD-SAD8 were highly expressed in the early middle developing ovules (5 and 10 DAF) while GhA-SAD7 enriched at the late developing stage (20 DAF). GhA-SAD7 also had the highest expression level in root compared to other tissues, indicating that GhA-SAD7 may function importantly in root. For flower tissue, both GhD-SAD5 and GhD-SAD7 had abundant expression levels, implying that GhD-SAD5 and GhD-SAD7 play roles in flower development. It is known that the fastest period of oil/FA biosynthesis in cotton endosperm was the middle stage (10–15 DAF) of the developing ovule. Combination of RNA-seq and qRT-PCR data suggests that GhA-SAD6, GhD-SAD8, GhA-SAD5, and GhA-SAD7 may play roles in FA/oil biosynthesis in developing cotton endosperm.

**FIGURE 2** 3D structure models of GhSAD protein monomers (C, D, upper panel I) and C18:0/C16:0-ACP chains as well as side chains of key amino acids close to catalytic center of diiron ions (lower panel II). All 3D models were predicted on Swiss-model (https://swissmodel.expasy.org/) with RcSAD1 (NP_001310659.1) and AtAAD3 as temples (A, B). Files of 3D structure of GhSADs were showed using Discovery Studio 4.1 software. Yellow bolls in upper panel I refer to diiron ions. Blue bolls/spheres refer to common amino acids as same as RcSAD1 while pink bolls refer to the varied amino acids. The fatty-acyl chains and side chains of amino acids were drawn by ChemDraw software (lower panel II). Black color represents C18:0-ACP and its common amino acids. Red indicates C16:0-ACP and varied amino acids.
**GhA-SAD6 and GhA-SAD8 Mostly Expressed in Developing Endosperm, Positively Correlating With Palmitoleic Acid Accumulation in Cotton Seeds**

To explore whether those GhSADs observed above are associated with 16:1Δ⁹ production in endosperm, developing cotton seeds at 15, 25, 35, and 45 DAF were then dissected and the two fractions, embryo and endosperm, were separately obtained to examine FA profiles and the expression patterns of the four GhSAD genes (GhA-SAD5, GhA-SAD6, GhA-SAD7, and GhD-SAD8) were selected according to the above characterization.

As shown in Figure 5, the four GhSAD genes exhibited differential expression patterns between embryo and endosperm of cotton seeds. The mRNA levels of GhA-SAD5, GhA-SAD6, and GhD-SAD8 were significantly high in the endosperm but very low in the embryo during seed development from 15 to 25 DAF, a period of palmitoleic acid level increased rapidly in the endosperm (Figures 5A, B, D). However, GhA-SAD7 expression was no obvious difference between embryo and endosperm (Figure 5C). Only a trace level of the four gene transcripts was detected in the embryo at the stage of 45 DAF. By this time, no endosperm was left in the seed. In addition, all these four genes had extremely low expression levels in cotton embryos and endosperms at 35 DAF compared to the two early stages (15 and 25 DAF). All these data suggest that GhA-SAD6 and GhD-SAD8 may be the main players for fast oil and palmitoleic acid accumulation in developing cotton endosperm although GhA-SAD5 has a higher expression in endosperm at 15 DAF.

To further confirm the association between GhA-SAD6 and GhD-SAD8 expressions and palmitoleic acid level in the endosperm of developing cotton seeds, two set of data, MUFA (16:1Δ⁹ and 18:1Δ⁹) contents and expression levels of GhA-SAD6 and GhD-SAD8 were integrated into one figure (Figure 6). In the embryo, neither 16:1Δ⁹ nor 18:1Δ⁹ content change was obviously correlated with the expression level of either GhA-SAD6 or GhD-SAD8 gene (Figures 6A, C). In the endosperm, no correlation was also detected between 18:1Δ⁹ content and the two gene expressions (Figure 6D). However, a significantly positive correlation was detected between 16:1Δ⁹ content and GhA-SAD6 or GhD-SAD8 expression in the endosperm fractions (Figure 6B). Moreover, linear regression analysis also showed that the level of palmitoleic acid in the endosperm is positively correlated with the expression of GhA-SAD6 (r = 0.752) and GhD-SAD8 (r = 0.886), respectively, again indicating that GhA-SAD6 and GhD-SAD8 may be the key...
Liu et al. Function of GhA-SAD6 and GhD-SAD8

FIGURE 4 | Deduced GhSAD expression patterns of GhA-SAD5 (A), GhD-SAD5 (B), GhA-SAD6 (C), GhA-SAD7 (D), GhD-SAD7 (E), and GhD-SAD8 (F) in various cotton tissues. Bar charts show the relative expression levels of GhSAD genes normalized to that of Histone3 and EF-1α measured by qRT-PCR. The analysis was performed with three biological samples for each tissue. Ovu-5d, 10d and 20d represent 5, 10, and 20 DAF of developing cottonseed ovules. The method of $2^{-\Delta\Delta Ct}$ was used in this analysis. ** represents statistically significant difference values of $P < 0.05$ and *** for $P < 0.01$ based on two-tailed Student’s t-tests.

contributors for biosynthesis and accumulation of palmitoleic acid in cotton endosperm.

GhA-SAD6 and GhD-SAD8 Boosted Accumulation of Palmitoleic Acid When Transient Expression in N. benthamiana Leaves

In order to identify the roles of GhA-SAD6 and GhD-SAD8 for biosynthesis of palmitoleic acid (16:1$ \Delta^9$), we transiently overexpressed them separately in leaves of N. benthamiana by Agrobacterium-mediated infiltrating. GhA-SAD5 and GhA-SAD7 were also transiently expressed, respectively, in tobacco leaves for comparison. The transformed leaves were harvested at 5 days after infiltration, and used for lipid extraction. FA analysis by GC (Figure 7) revealed that compared to the wild-type and empty-vector controls, either GhA-SAD6 or GhD-SAD8 overexpression led to a significant increase of palmitoleic acid by at least 4–5 folds. However, GhA-SAD5 or GhA-SAD7 expression alone resulted in a slight enhance of oleic acid (18:1$ \Delta^9$) by 2 folds around, with no effects on palmitoleic acid biosynthesis. These results indicate that GhA-SAD6 and GhD-SAD8 are specific for the ectopic enrichment of palmitoleic acid (e.g., to catalyze 16:0-ACP into 16:1$ \Delta^9$-ACP) in the leaves whereas GhA-SAD5 and GhA-SAD7 are specific for oleic acid biosynthesis (e.g., to catalyze 18:0-ACP into 18:1$ \Delta^9$-ACP). Other changes of FA profiles were the increase of linoleic acid (18:2$ \Delta^9,12$) but reduction of linolenic acid (18:3$ \Delta^9,12,15$) in the GhA-SAD5- or GhA-SAD7-expressed leaves. Such change in polyunsaturated fatty acid levels maybe due to the activity changes of the endogenous NbFAD2 and NbFAD3 enzymes in tobacco leaves.

GhA-SAD6 and GhD-SAD8 Accounted for Palmitoleic Acid Production When Expressed in Yeast Mutant BY4389

To further confirm the function of GhA-SAD6 and GhD-SAD8 for biosynthesis of palmitoleic acid (16:1$ \Delta^9$), we overexpressed
FIGURE 5 | Expression patterns of GhA-SAD6 (A), GhA-SAD6 (B), GhA-SAD7 (C), and GhD-SAD8 (D) in developing cotton embryos and endosperms at 15, 25, 35, and 45 DAF. The same experimental and statistical methods are performed as in Figure 4.

the four GhSAD genes (GhA-SAD5/6/7 and GhD-SAD8), respectively, in yeast mutant BY4389 disable in synthesis of unsaturated fatty acids. For this genetic complementary assays, Arabidopsis AtFAB2 (18:0-ACP specific) and AtAAD3 (16:0-ACP specific) were employed as the positive controls, and an empty pYES2 vector was used as the negative control. FA composition was examined in the transformed BY4389 strains.

Notably, all transformed yeast mutants except for empty pYES2 vector restored their desaturase function, and thus survived in SC-Ura medium without any unsaturated fatty acid. FA profiles detected (Figure 8) showed that the transgenic yeasts synthesized new but unbalanced-level unsaturated fatty acids (UFAs). As expected, AtAAD3-transgenic yeast generated 16.9 Mol% of C16:1Δ9 while AtFAB2-transgenics produced 19.03 Mol% of C18:1Δ9. Like in the AtAAD3-transgenics, both GhA-SAD6- and GhD-SAD8-expressed yeast accumulated high levels of C16:1Δ9 by 13.83 and 9.88 Mol%, respectively, but almost no detectable of C18:1Δ9. In contrast, the yeast mutants harboring GhA-SAD5 or GhA-SAD7 produced large amount of C18:1Δ9 by approximately 10 Mol%, with <3.5 Mol% of C16:1Δ9, which was similar to FA profiles in the AtFAB2-transgenic yeast. Overall, the present data obtained by in vivo assays using yeast mutant transformation again revealed that GhA-SAD6 and GhD-SAD8 both had palmitoyl-ACP substrate specificity, with the former stronger. Whereas, GhA-SAD5 and GhA-SAD7 showed to prefer stearoyl-ACP substrate over palmitoyl-ACP despite of low enzymatic activities.

DISCUSSION

Enrichment of ω-7 FAs in Developing Cotton Seeds May Benefit the Embryo Development

For the albuminous seeds such as cereal grains and castor beans, the embryo is embedded in the endosperm which persists to the mature seed stage as a storage tissue supplying nutrients for the seedling that will grow from the embryo (Huh et al., 2007). However, for exalbuminous seeds such as soybean, peanut and cotton, the endosperm is absorbed by the embryo during seed development, and the cotyledons...
become filled with stored food (Yan et al., 2014). And thus, no endosperm is left in maturity seeds. A differential partitioning of reserve compounds between different tissues in Arabidopsis seeds was examined, with a dramatically increased accumulation of the economically important ω-7 FAs in the endosperm, but not in the embryo (Penfield et al., 2004; Barthole et al., 2014). Here, we demonstrate that palmitoleic acid (16:1 Δ9) is enriched in developing cotton endosperm but not in the embryo. This ω-7 FA accumulation is different in Arabidopsis and rapeseed where this unusual fatty acid is highly concentrated in a single cell layer (aleurone) of the residual endosperm surrounding the embryo in the mature seed, with the major species being cis-vaccenic acid (18:1 Δ11) and paullinic acid (20:1 Δ13) instead of palmitoleic acid (16:1 Δ9) (Li et al., 2006). Such difference suggests that there may be different temporal-spatial regulation mechanisms in different plant seeds. And it also raises the question of what is the biological function of these ω-7 FAs enriched in the developing endosperm or the residual endosperm.

In Arabidopsis, transcriptional factors MYB115 and MYB118 coordinately upregulate the transcription of AtAAD2 and AtAAD3 which catalyze the synthesis of ω-7 FAs, leading to high levels of ω-7 FA accumulation exclusively in the endosperm of early maturing seeds (Bryant et al., 2016; Troncoso-Ponce et al., 2016). Arabidopsis double mutant of MYB115 and MYB118 resulted in significant decrease of ω-7 FAs in the endosperm, concurrently with a slight delay in the elongation and enlargement of the embryo associated with a moderate

**FIGURE 6** | The correlation of the expression of GhA-SAD6 and GhD-SAD8 with molar percentage of C16:1 Δ9 or C18:1 Δ9 in developing cotton embryos and endosperms. Columns showed fatty acid content (Mol% of total fatty acids) and lines for relative expression (Red for GhA-SAD6, blue for GhD-SAD8). (A, C) represented the correlation between the fatty acid content (C16:1 Δ9 or C18:1 Δ9) and relative expression levels of GhA-SAD6 or GhD-SAD8 in developing embryos, respectively. (B, D) were the correlation of C16:1 Δ9 or C18:1 Δ9 content with GhA-SAD6 or GhD-SAD8 in developing endosperms, respectively. Six biological replicates were analyzed in each tissue.
decrease of mature seed dry weight. This indicates that ω-7 FAs may function for seed development. However, seeds from the double mutant of AtAAD3 and AtAAD2 appear normal and can germinate readily under standard growth conditions. Even so, function of ω-7 FAs for seed germination cannot be excluded because the low level of remained ω-7 FAs in the endosperm of this double mutant may impair the examination of possible phenotypes.

To clarify the physiological roles of ω-7 FAs enriched in developing cotton endosperm, we conducted an in vitro cultivation of young embryos isolated from early developing seeds (data not shown). Addition of palmitoleic acid into the medium promoted the embryo development, with the embryo growing more health compared to the embryo cultured in the medium without this ω-7 FA. Moreover, this beneficial effect of ω-7 FA addition exhibited much stronger under cold stress. This demonstrates that ω-7 FA accumulation in the developing endosperm may function importantly for cotton seed development. Further lab work such as CRISPR-Cas9 mediated 16:0-ACP Δ⁹ desaturation (Kachroo et al., 2007) is under way to investigate the biological functions of ω-7 FA, particularly for seed development and germination.

Members of Cotton GhSAD Family May Function Differentially, With GhA-SAD6 and GhD-SAD8 as the Major Players for Palmitoleic Acid Enrichment in Developing Cotton Endosperm

Generally, soluble acyl-ACP Δ⁹ desaturases (AAD) catalyze the formation of first unsaturated fatty acid in FA biosynthesis pathway in plant plastid. As a polygenic family, AAD contains various isoforms. For example, multiple members of AAD family were identified in several plant genomes, with 7 members in Arabidopsis (Kachroo et al., 2007), 8 members in cacao (Theobroma cacao) (Zhang Y. et al., 2015) and 3 members in olive (Olea europaea) (Parvini et al., 2016), respectively. Moreover, different AAD isoforms showed functional diversity and distinct expression patterns. Among 7 AtSAD members, SSI2/FAB is the predominant and the best characterized isoform, with expression in all seed tissues and 18:0-ACP Δ⁹ desaturation (Lightner et al., 1994) despite it was also active on 16:0-ACP (Kachroo et al., 2007). Microarray analysis by Le et al. (2010) showed that AtAAD2 expression was mainly in the peripheral endosperm, whereas AtAAD3 largely expressed in the chalazal region of the endosperm (Le et al., 2010; Bryant et al., 2016).
Further analysis on the single or double mutant of the genes by Bryant et al. (2016) evidenced that AtAAD2 and AtAAD3 function in 16:0-ACP Δ⁹ desaturation and contribute to the biosynthesis of ω-7 FAs specifically in the aleurone. Possibly, the accumulation of ω-7 FAs in rapeseed aleurone plus seed coat results from the homologs of AtAAD3 since rapeseed has the relatively close phylogenetic relationship with Arabidopsis (Bryant et al., 2016). Although such findings are obtained from the characterization of the multigene family of AAD, the functions of most isoforms of this family remain to be elucidated.

Previous study showed that total of 17 stearoyl-ACP Δ⁹ desaturases (SAD) were identified in upland cotton genome, and their expression were developmentally regulated (Shang et al., 2017). However, no functional analysis of these 17 GhSADs is available, hence raising the question whether these GhSADs are redundant to the monounsaturated fatty acid formation or non-functional enzyme, or function separately from one another.

In the light of the present study, it demonstrates that the identical GhSAD family members were identified in upland cotton (Supplementary Table S3). Four GhSAD members including GhA-SAD5, GhA-SAD6, GhA-SAD7 and GhD-SAD8 showed higher expression levels at early stage of developing ovules than in other vegetative organs (Figure 4). Detailed expression analysis in dissection cottonseeds of different developmental stages revealed that the four genes above were highly expressed in early developing endosperm (15–25 DAF) (Figure 5), especially for GhA-SAD6 and GhD-SAD8 (Figure 5). However, the transcripts of GhA-SAD2, GhA-SAD3, and GhD-SAD6 mostly accumulated in the embryo (data not shown). Such diverse spatiotemporal expression patterns indicate that these GhSADs tested may function differentially. In order to investigate their functions, particularly those GhSADs selectively expressed in developing cotton endosperm, we conduct in vivo functional assays including transient expression in tobacco leaves and heterologous expressing in yeast strain BY4389.
unable to synthesize unsaturated fatty acids. The overexpression of both GhA-SAD6 and GhD-SAD8 substantially increase palmitoleic acid (16:1<sub>Δ9</sub>) synthesis in leaves, accompanied by a concomitant decrease in palmitic acid (16:0) (Figure 7). Unlike GhA-SAD6 and GhD-SAD8, the transient expression of either GhA-SAD5 or GhA-SAD7 slightly enhance oleic acid (18:1<sub>Δ9</sub>) in the leaves at the expense of stearic acid (18:0) (Figure 7). Similar results were also obtained by functional complementation assay in the yeast system (Figure 8). These reveal that in the developing cotton endosperm, GhA-SAD6 and GhD-SAD8 are responsible for the synthesis of palmitoleic acid (16:1<sub>Δ9</sub>) by 16:0-ACP desaturation while GhA-SAD5 and GhA-SAD7 are responsible for the synthesis of oleic acid (18:1<sub>Δ9</sub>) by 18:0-ACP desaturation. Together, our data demonstrate for the first time that the four GhSAD members (GhD-SAD8, GhA-SAD5, GhA-SAD6 and GhA-SAD7) are functional acyl-ACP Δ<sup>9</sup> desaturase enzymes but with different substrate preferences. GhA-SAD6 and GhD-SAD8 can preferentially catalyze 16:0-ACP to 16:1<sub>Δ9</sub>-ACP, whereas GhA-SAD5 and GhA-SAD7 are able to selectively catalyze 18:0-ACP to 18:1<sub>Δ9</sub>-ACP although <i>in vitro</i> assay has not been conducted yet for identifying the substrate specificity of those GhSAD members. Evolutionally, the occurrence of multiple SAD members in upland cotton and other plant species tested likely provide diverse enzymes with selective expression so as to achieve fine spatiotemporal accumulation of different monounsaturated fatty acids as in cases of the seeds from cotton, Arabidopsis and rapeseed where fatty acid profiles were different between embryo and endosperm. Further functional characterization is required for every member of this SAD family. Nevertheless, our study brings new perspectives in this regard.

**Key Amino Acid Residues Lining the Bottom Part of the Substrate-Binding Channels of SADs Are Crucial Determinants of the Substrate Specificity of the Enzymes, but Not the Only Factor**

As described above, plant SAD family composed of soluble plastidial proteins contains multiple members with differential expressions in different tissues and distinct substrate specificities. Some isoforms of the enzyme like castor RcSAD1 and Arabidopsis AtFAB2 show high preference for 18:0-ACP, yielding 18:1<sub>Δ9</sub>-ACP, while others such as cat’s claw Muc-PAD, Arabidopsis AtAAD2 and AtAAD3 exhibit strong activity on 16:0-ACP, producing 16:1<sub>Δ9</sub>-ACP. It have been reported that Δ<sup>9</sup> desaturases can recognize 14-, 16-, and 18-carbon chain lengths, and others perform desaturation with Δ<sup>6</sup> and Δ<sup>9</sup> regiospecificities (Whittle et al., 2005; Guy et al., 2007). In view of evolution, such diverse substrate specificities of SADs likely provide a competitive advantage for host plants to produce the required monounsaturated FAs for life activities including maintaining membrane fluidity and channeling into storage triacylglycerols (TAGs) in storage tissues (Shanklin and Cahoon, 1998), benefiting the evolution of novel metabolic diversity (Dyer et al., 2002).

Previous studies on the molecular mechanism underlying SAD substrate specificity were mostly conducted using the crystal structure of castor 18:0-ACP Δ<sup>9</sup> desaturase (RcSAD1) as a model (Lindqvist et al., 1996; Cahoon et al., 1997), showing that a group of eight amino acid residues lining the bottom part of the substrate channels of SADs is crucial for determining the substrate specificity of the enzymes (Cahoon et al., 1998). For example, RcSAD1 could change its binding substrate for 16:0-ACP from the wild type with 18:0-ACP when two key residues (T117 and G188) were substituted by Arg and Leu. Moreover, the mutant RcSAD1 activity for 16:0-ACP was higher by 82 folds than the wild enzyme for 18:0-ACP (Whittle and Shanklin, 2001). The side chain sizes of these 8 amino acids influence the depth and width of the substrate channel of the enzyme (Cahoon et al., 1997). The substrate channel of the archetype Δ<sup>9</sup> SAD is deep enough to accommodate 18:0-acyl chain, thus producing Δ<sup>9</sup> monoenes (Figure 2A). However, the shorter substrate channel of the enzyme is more adapted to 16:0-acyl chain, forming a ω-7 FA (e.g., palmitoleic acid, 16:1<sub>Δ9</sub>) (Figures 2A–C). These eight amino acids appear to be well conserved among the SAD isoforms characterized so far in higher plants. The amino acid sequences of the most SAD isoforms are highly homology to that of the castor 18:0-ACP Δ<sup>9</sup> desaturase (RcSAD1), implying that SADs share a common architecture able to accommodate different substrate binding models. Like in Arabidopsis, AtSADs share a high degree of amino acid sequence similarity and a common structural fold (Kachroo et al., 2007). Moreover, the three divergent residues (F226/216, S224/214, and T159/149) of the eight amino acids lining the bottom part of the substrate channels of AtAAD2 and AtAAD3 conferred their strong activity for 16:0-ACP, but not 18:0-ACP catalyzed by the SAD archetype (Bryant et al., 2016).

Base on such conservation, the corresponding amino acid residues in 17 cotton SADs were identified by a polypeptide sequence alignment of the eight amino acid residues lining the bottom part of the substrate channels of the GhSAD proteins using the typical 18:0-ACP specific SADs (RcSAD1 and AtFAB2) and the 16:0-ACP specific SADs (Muc-PAD, AtFAB2, AtAAD2, and AtAAD3) as references (Table 1). Interestingly, the eight amino acids are all identical for the group of 18:0-ACP specific SADs. However, they are divergent for the group of 16:0-ACP specific SADs. Only one amino acid residue of L115 in RcSAD1 is identical for all SADs tested here. This alignment indicates that 10 GhSADs may be 18:0-ACP specific while the other seven GhSADs possibly be 16:0-ACP specific. Further 3D modeling of GhSAD protein monomers showed that GhA-SAD6 and GhD-SAD8 had the similar structure of the substrate-bound cavity adjacent to the active diiron center with AtAAD3, but different from RcSAD1, suggesting that these two GhSADs have 16:0-ACP activity. Their substrate specificities were confirmed by our functional assay using tobacco transient expression and yeast expression system (Figures 7, 8). In contrast, GhA-SAD5 and GhA-SAD7 exhibited 18:0-ACP preference (Figures 7, 8) although they were classed into the same group of 16:0-ACP specificity by amino acid
sequence alignment (Table 1). On the other hand, GhA-SAD6 and GhD-SAD8 only have 50% identity for these eight amino acid residues despite of the similar substrate specificity, just like cases of Muc-PAD, AtAAD2, and AtAAD3 with less identity but the same substrate activity. Even for RcSAD1 and Muc-PAD, only one residue was diverged for the eight amino acids between them. Taken together, these indicate that other factors may also affect the substrate specificity of SADs beyond the predominant role played by the eight amino acid residues.

In consistence with this, increasing characterization of more SADs then revealed that for substrate preference of a given chain length, some isoforms can also desaturated slightly shorter- or longer-acyl chains, despite of a reduced affinity (Kachroo et al., 2007; Rodriguez et al., 2015). Some AADs then revealed that for substrate preference of a given chain length, some isoforms can also desaturated slightly shorter- or longer-acyl chains, despite of a reduced affinity. For instance, ivy (Hedera helix) AAD desaturated at the Δ⁹ position when presented with 18:0-ACP substrate while it catalyzed Δ⁴ desaturation when supplied with 16:0-ACP (Whittle et al., 2005). Moreover, this AAD also converted 16:1Δ⁹-ACP and 18:1Δ⁹-ACP to the corresponding Δ⁵,⁹ dienes. In contrast to ivy AAD, Thunbergia AAD produced 16:1Δ⁶ with 16:0-ACP substrate but when presented with 18:0-ACP substrate generated a mixture of Δ⁶ and Δ⁸ desaturated products (Cahoon et al., 1997). Possibly, the mechanism determining the substrate specificity of AADs may evolve independently among different plants species.

CONCLUSION

The present study revealed that palmitoleic acid (16:1Δ⁹), a kind of ω-7 FAs, is specifically enriched in developing cotton endosperm, possibly benefiting the embryo development. Total of 17 GhSADs are identified in upland cotton genome and their expressions are spatiotemporally diverse, indicating that they may function differentially. Particularly, GhA-SAD6 and GhD-SAD8 are preferentially expressed in the developing endosperm whereas GhA-SAD2 and GhD-SAD6 are restricted to the embryo. Our functional characterizations using transient expression in tobacco leaves and heterologous expression in yeast mutant BY4389 provide the first evidence showing that GhA-SAD6 and GhD-SAD8 are 16:0-ACP Δ⁹ desaturases responsible for production of 16:1Δ⁹ in the developing cotton endosperm while GhA-SAD5 and GhA-SAD7 act as 18:0-ACP Δ⁹ desaturases accounting for 18:1Δ⁹ biosynthesis in this tissue. In addition, polypeptide sequence alignment and 3D modeling of the protein monomers of these GhA-SAD6 and GhD-SAD8 suggested that their 16:0-ACP substrate specificities may be determined by the eight amino acid residues lining the bottom part of their substrate channels which distinguish them from the archetype 18:0-ACP Δ⁹ desaturase. Taken together, this study brought new insights into the SAD-mediated biosynthesis and differential accumulation of special lipid reserve in developing cotton seeds. The identified GhA-SAD6 and GhD-SAD8 could be good endogenous genes for engineering to produce higher oil enriched with ω-7 FAs in cotton seeds or other oil crops for food and industrial applications.

DATA AVAILABILITY

No datasets were generated or analyzed for this study.

AUTHOR CONTRIBUTIONS

BL, JX, XJ, and RL designed the experiments and drafted the manuscript. BL and YS did the vector construction, fatty acid analysis and material plantation. BL and XM carried out the expression and RNA-seq analysis. RL and XJ revised the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2019.00703/full#supplementary-material

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**Conflict of Interest Statement:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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