The Phylogeny and Functional Characterization of Peanut Acyl-ACP Thioesterases

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Abstract
Fatty acyl-acyl thioesterases (FATs), which hydrolyze the thioester bond linking acyl chains to an acyl carrier protein, thereby terminating their elongation, contribute significantly to the fatty acid (FA) content and composition of seed storage lipids. The peanut (Arachis hypogaea L.) genome was found to harbor 21 FAT (AhFAT) genes, distributed over 12 of the 20 chromosomes. The length of their predicted translation products varied from 74 to 415 residues, and all but one included the 1–2 Acyl-ACP_TE conserved domains. All of the coding sequences were interrupted by at least one intron, with the exon number ranging from two to 12, and five of the genes were liable to alternative splicing. When the RNA-Seq platform was used to assess the transcriptional behavior of the 21 AhFAT genes, transcription of only 13 was detectable in samples of root, leaves, and developing seed; among these, six were transcribed throughout the plant, three were root-specific and one was leaf-specific. A detailed analysis of a pair of homologous AhFATs showed that the coding region of each was split into six exons and that both were transcribed in all of the plant organs surveyed (although the intensity of their transcription was not the same in immature seed). The product of both genes was deposited in the chloroplast outer membrane. The constitutive expression of these genes in either yeast or Arabidopsis thaliana increased the FA content, especially that of saturated FAs. In peanut genome, 21 AhFAT genes were found and two of them were transformed into yeast and Arabidopsis for function identification. Results showed that overexpression of these two genes could increase the FA content, especially the saturated FAs content.

Keywords Fatty acid · Transcriptional level · Gene structure · Sub-cellular localization · Saturated fatty acids content · Genetic transformation

Abbreviations
AS Alternative splicing
C14:0 Myristic acid
C16:0 Palmitic acid
C16:1 Palmitoleic acid

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C18:0  Stearic acid  
C18:1  Oleic acid  
C18:2  Linoleic acid  
C18:3  Linolenic acid  
C20:0  Arachidic acid  
C20:1  Eicosenoic acid  
C20:2  Eicosadienoic acid  
C22:1  Erucic acid  
FA  Fatty acid  
FAT  Fatty acyl-acyl thioesterase  
RT-PCR  Real-time quantitative polymerase chain reaction  
Taqman-PCR  Taqman-polymerase chain reaction

Introduction

Fatty acyl-acyl carrier protein thioesterases (FATs) are pivotal enzymes in the synthesis of fatty acids (FAs). Following the termination of acyl chain elongation, FAs are hydrolyzed by FATs and transported to the cytosol (Beisson et al. 2003; Li-Beisson et al. 2010; Sandelius and Aronsson 2008). As a result, FATs determine the acyl chain length of FAs. Two groups of FAT have been recognized: while FATAs prefer unsaturated fatty acids as a substrate, FATBs prefer unsaturated fatty acids (Jones 1995). Within the FATBs, subgroup 1 enzymes act on long-chain fatty acids (Dörmann et al. 1995; Filichkin et al. 2006; Jha et al. 2010; Moreno-Pérez et al. 2011; Sánchezgarcía et al. 2010), while subgroup 2 enzymes target medium-length chain fatty acids (Filichkin et al. 2006; Jing et al. 2011; Voelker 1996; Voelker et al. 1997). The genetic manipulation of various plant FAT genes has been shown to influence FA composition: for example, the overexpression of the Arabidopsis thaliana gene AtFATB1 results in the accumulation of C16:0 (palmitic acid) (Dörmann et al. 2000). The expression in Chlamydomonas reinhardtii of a FATA gene isolated from the algal species Dunaliella tertiolecta has the effect of producing 63% and 94% more neutral lipids than the wild type, without compromising growth (Tan and Yuan 2017). When a FATB gene isolated from the cigar plant Cuphea lanceolata was expressed in C. reinhardtii, the production of C14:0 (Myristic acid)-containing triacylglycerols increased by up to 1.6-fold (Inaba et al. 2017). The heterologous expression of a Brassica napus FATB gene in yeast raised the content of the saturated C16:0 and C18:0 (stearic acid) by, respectively, 46% and 22%, at the same time reducing that of both the unsaturated C16:1 (palmitoleic acid) and C18:1 (oleic acid) by, respectively, 15% and 31% (Tan et al. 2015).

The peanut (Arachis hypogaea L.) is a leading oil seed crop. Several genes influencing its seed oil content have been identified, among which are a number of FATs. The expression of an AhFATA in both Escherichia coli and certain algal species has been shown to be effective in both altering the FA profile and increasing the content of a number of FAs (Chen et al. 2017). Manipulation of AhFATB1 has been effective in increasing the content of both saturated and unsaturated FAs (Chen et al. 2012, 2017; Wen et al. 2012). With the acquisition of the genome sequence of peanut (Bertioli et al. 2016; Bertioli et al. 2019; Zhuang et al. 2019), it is now possible to identify the species’ full complement of AhFAT genes and to characterize the functionality of each of them. The aim of the current research was document the family of AhFAT genes, to clarify their phylogeny and gene structure, and to profile their transcription in different organs of the plant. Moreover, two not previously isolated FATs have been transformed and functionally analyzed in Saccharomyces cerevisiae and A. thaliana hosts.

Materials and Methods

Plant and Yeast Materials

The peanut cultivar (cv.) ‘Fenghua 1’ was used for gene isolation. ‘Fenghua 1’ was planted in greenhouse with 16-h light/8-h dark photoperiod and 30 °C day/22 °C night temperature cycle. When it got to 12 days, the young mainstem, mainstem leaves, and root tips of ten plants were taken down and frozen in liquid nitrogen. In the full podding bearing stage, seeds of different developmental stages were taken down and frozen in liquid nitrogen. These materials were kept at − 80 °C until use. Genetic transformation experiments involved A. thaliana ecotype Col-0 and yeast (S. cerevisiae) strain W303. Arabidopsis plants were grown in flower pots in a temperature controlled incubator, with 16-h light/8-h dark photoperiod and 21 °C day/16 °C night temperature cycle. W303 yeast strains were kept on YPD medium (Qingdao Hope Bio-Technology Co. Ltd, China) and grown at 30 °C.

Sequence Analysis

The query sequences used to reveal the family of AhFAT genes represented in the species’ genome sequence (https://www.peanutbase.org/) were AhFATA (GenBank accession GU324446.1) and AhFATB (EF117305.1). Gene structures were analyzed using GSDS software (gsds.cbi.pku.edu.cn) and transmembrane domain analysis was conducted using TMHMM software (http://www.cbs.dtu.dk/services/TMHMM/). Sub-cellular localization was predicted using the ProtComp 9.0 package (https://linux1.softberry.com/berry.phtml?topic=protcomppl&group=programs&subgroup=proloc). Conserved domains were identified using SMART software (https://smart.embl-heidelberg.de/).
Phylogenetic Analysis

The AhFATA and AhFATB1 sequences were used to scan the Phytozome database (https://phytozome.jgi.doe.gov/pz/portal.html) for plant FATs (Supplemental Table S1). The resulting sequences were used to construct a FAT phylogeny based on ClustalW software (https://www.genome.jp/tools-bin/clustalw), with parameters set to default values. Mega6 software (https://www.megasoftware.net/) was used to construct a plant FAT phylogeny, based on the neighbor-joining method; statistical confidence in the clade branching points was obtained by running 1,000 bootstrap replicates (Tamura et al. 2013).

Transcription Profiling Analysis

RNA was extracted from the all roots and leaves of 12-day old cv. ‘Fenghua 1’ seedlings and from developing seeds harvested either 30 or 50 days after flowering (respectively “seed1” and “seed2”). RNA was extracted using the TRIzol reagent (Invitrogen, Carlsbad, CA, USA) then treated with RNase-free DNase I (New England Biolabs, USA) for 30 min at 37 °C to degrade any contaminating DNA present. The concentration and purity of the resulting RNA preparations were assessed using a NanoDrop 2000 spectrophotometer (Thermo Fisher Scientific, Wilmington, DE, USA) and its integrity was checked using an RNA Nano 6000 Assay Kit (Agilent Technologies, CA, USA). A 1.5 μg aliquot of RNA was processed with a Ribo-Zero rRNA Removal Kit (Epicentre, Madison, WI, USA) to remove the rRNA component, and the subsequently prepared sequencing libraries based on the residual RNA, following treatment with an NEBNext® UltraTM Directional RNA Library Prep Kit for Illumina® (New England Biolabs, USA). Index codes were added to enable each sequence to be attributed its organ provenance. Paired-end sequences were generated by an Illumina HiSeq2500 platform. The resulting sequence data, stripped of adapter sequence and low-quality reads, have been deposited with NCBI (SRA ID: SRP093901). The resulting sequence data, stripped of adapter sequence and low-quality reads, have been deposited with NCBI (SRA ID: SRP093901). The sequences were aligned with the genome sequence of the peanut cv. ‘Tifrunner’ (https://www.peanutbase.org/peanut_genome) using TopHat2 software (Kim et al. 2013). Aligned reads were assembled into a full transcriptome using the Cufflinks v2.2.1 program (Trapnell et al. 2010). FPKM values for each transcript were used to estimate transcript abundance. The expression levels of FATBs from Physcomitrella patens, Populus trichocarpa, and Glycine max are obtained from Phytozome v12.1 (https://phytozome.jgi.doe.gov/pz/portal.html).

AhFATB2 Isolation

Seed1 cDNA was provided as the template for PCRs driven by either primer pair AhFATB2-1-F/-R or AhFATB2-2-F/-R (Supplemental Table S2). Each 25 μL PCR contained 1 μL cDNA (100 ng/μL), 2 μL dNTP (10 mM each), 2.5 μL 10× TransTaq HiFi buffer (https://www.transgenbiotech.com), 1 μL TransTaq HiFi DNA polymerase (https://www.transgenbiotech.com), 1 μL of each primer (10 μM), and 16.5 μL ddH2O. The amplification program comprised an initial denaturation (95 °C/5 min), followed by 35 cycles of 95 °C/30 s, 60 °C/30 s, 72 °C/35 s, with a final elongation of 72 °C/10 min. The amplicons were resolved by electrophoresis through a 1% agarose gel, and sequenced by Sangon Biotech (Shanghai, China).

Isolation of RNA and Semi-quantitative PCR

Total RNA was extracted from the root, stem, leaf, flower, and seeds at 10, 20, 30, 40, 50, and 60 days after flowering of cv. ‘Fenghua 1’ plants, using a DP441 RNAprep Pure Plant Kit (Tiangen, Beijing, China), and was reverse transcribed using a Thermo Scientific RevertAid First Strand cDNA Synthesis kit (https://www.thermofisher.com/cn/zh/home.html). A Taqman-PCR assay (Agilent Technologies, California, USA) was used to determine transcript abundance, based on the primer pairs FATB2F/R (Supplemental Table S2) for the amplification of a fragment of AhFATB2 and the fluoresently labeled pairs FATB2-1-P/2-P (Supplemental Table S2) for the detection of, respectively, AhFATB2-1 and AhFATB2-2. The reference sequence was Actin (GenBank number GO264911), assayed with the primer pair AhActin-S/-A (Supplemental Table S2). Each 20 μL PCR contained 10 μL TaqMan Fast qPCR Master Mix, 0.4 μL of each of AhFATB2F/R (10 μM each), 0.4 μL AhFATB2-1-P or AhFATB2-2-P (10 μM), 2 μL cDNA (100 ng/μL), and 6.8 μL ddH2O. The amplification program comprised an initial denaturation (94 °C/3 min), followed by 45 cycles of 95 °C/5 s, 57 °C/15 s, 72 °C/30 s, with a final elongation of 72 °C/10 min. The relative transcript abundance of each FATB2 was calculated using the $2^{-\Delta\Delta CT}$ method (Livak and Schmittgen 2001). Each experiment was run in triplicate.

Sub-cellular Localization

The open reading frames (lacking the stop codon) of AhFATB2-1 and AhFATB2-2 were PCR-amplified using the primer pairs FATB2-1-GFP-F/-R and FATB2-2-GFP-F/-R (Supplemental Table S2), after which the amplicons were treated with Hind III and BamH I, then cloned into the multiple cloning site of the pBSK+-35S-EGFP plant transient expression vector (kindly provided by Weicai Yang, Institute
of Genetics and Developmental Biology, Chinese Academy of Science). The constructs were transformed into A. thaliana mesophyll protoplasts using the PEG-calcium transfection method (Yoo et al. 2007). Fluorescence generated by the expression of GFP was observed by confocal laser scanning microscopy.

**Yeast Transformation**

AhFATB2 fragments were PCR-amplified using the primer pairs FATB2-1-BamHI-F/-KpnI-R or FATB2-2-BamHI-F/-KpnI-R (Supplemental Table S2), after which the amplicons were treated with Kpn I and BamH I, then inserted into the multiple cloning site of the galactose-inducible yeast expression vector pESC-URA (Agilent Technologies). The resulting constructs plasmids were transformed into S. cerevisiae strain W303 competent cells which were grown on a selective SD-URA medium. Pooled colonies from each strain were grown in a selective liquid medium, then diluted into a liquid medium containing galactose and raffinose to induce protein expression. Cells harboring an empty plasmid served as the negative control. The cells were dried, then analyzed for their fatty acid content, following a gas chromatography protocol described by Zheng et al. (2017).

**A. thaliana Transformation**

The AhFATB2 open reading frames were PCR-amplified using the primer pairs FATB2-1-KpnI-F/-SacI-R or FATB2-2-KpnI-F/-SacI-R (Supplemental Table S2) and the resulting amplicons inserted first into the binary vector PRI101 (kindly provided by Minqin Wang, Shandong University, China) and thence into Agrobacterium tumefaciens strain LBA4404 (Poirier et al. 2000). A. thaliana plants were transformed using inflorescence infection method (Clough and Bent 1998). Progeny of putative transgenics were plated on half-strength MS medium (Murashige and Skoog 1962) containing 100 mg/L kanamycin in order to select for transgene-positive T1 individuals, and this procedure was repeated to obtain the T2 and T3 generations. The progeny of T3 plants were analyzed for their fatty acid content, following a gas chromatography protocol described by Zheng et al. (2017).

**Results**

**The AhFAT Gene Content of the Cultivated Peanut Genome**

In total, 21 potential AhFATs were uncovered from A. hypogaea genome (Supplemental Table S3); four loci mapped to each of chromosomes Ahrahy.12 and Ahrahy.13, three to Ahrahy.01, two to Ahrahy.02, and the other eight each mapped to different chromosomes. Two of the 21 genes were FATAs, namely Arahy.YI95N7.1 (on chromosome Ahrahy.04) and Arahy.QR9F0J.1 (Ahrahy.14); the 19 AhFATB genes were distributed across ten chromosomes. On the basis of their sequences, Arahy.YI95N7.1 was deemed to be a synonym of the previously described gene AhFATA, as was Arahy.KT88MX.1 of AhFATB1 (Chen et al. 2012; Wen et al. 2012). The length of the predicted translation products ranged from 74 (Ahrahy.L512HM) to 415 (Ahrahy.SJ2WML) residues. By sequence alignment in NCBI, we found Arahy.L512HM appeared to be an incomplete protein. By comparing the gene sequences with their genomic sequences, we found five of the genes produced more than one isoform: while both Arahy.QR9F0J and Arahy.L8EKIA produced three splicing variants, Arahy.H8W5PF, Arahy.LCII34, and Arahy.3BV5AA each produced two. Alternative splicing (AS) led to the formation of one transmembrane domain of Arahy.QR9F0J.3. Arahy.QR9F0J.3 was the only one which had transmembrane domain, whereas the other AhFATs had none. It had no effect on the products’ sub-cellular localization, with all 21 products predicted to be deposited in the chloroplast. With the exception of the very short Arahy.L512HM.1, all of the proteins possessed a 1–2 AcylACP_TE domain, with six also possessing an Acyl-thio_N domain.

There was extensive variation with respect to the gene structures of the AhFATs, with the number of exons ranging from two to twelve (Fig. 1). The shorter genes Arahy.L512HM.1, Arahy.C2SFPD.1, Arahy.0VM2VP.1, Arahy.QR9F0J.3, and Arahy.SL0JPW.1 each featured only two or three exons, while the others’ coding sequences were split into between six and 12 fragments. The Arahy.L8EKIA.I coding sequence was interrupted by 11 introns and featured three alternative splicing variants (one complete, one comprising eight exons, and one of seven exons); these isoforms derived from exon skipping, alternative transcription start sites and alternative transcription termination sites. The three isoforms generated by Arahy.QR9F0J were of length eight, six, and four exons. Intron retention was observed for both Arahy.QR9F0J.1 and Arahy.QR9F0J.2.

**Phylogeny of the AhFATs**

After removing gene fragments and incomplete sequences, the scan of plant FATs delivered a set of the 179 sequences from 26 species (Supplemental Table S1) used to construct a FAT phylogeny (Fig. 2). The representative from C. reinhardtii was located closest to the root of the tree, while the remainder formed two large groups (the FATAs and the FATBs). The two subgroups FATB1 and FATB2 were recognized: the former consisted entirely of genes harbored by angiosperm species, and comprised four branches (I through IV). The dicotyledoneae FATBs (including
some of the peanut FATBs) clustered within branch I, whereas branch II consisted entirely of monocotyledoneae FATBs, branch III only of peanut FATBs, and branch IV of FATBs from *Prunus persica* and *Populus trichocarpa*. With respect to the FATB2 subgroup, genes harbored by primitive land plant species located close to the root, and the remainder clustered into two groups (branches V and VI); and branch V contained all of the other primitive land plant genes, and branch VI the angiosperm genes. Peanut FATBs were distributed over the four branches I, III, V, and VI, each branch consisted of four, seven, two, and two AhFATBs, respectively. The FATAs produced by *Marchantia polymorpha* (a liverwort), *Selaginella moellendorffii* (a lycophyte), and *Physcomitrella patens* (a bryophyte) were located close to the root, and the remainder formed branches VII and VIII of the phylogenetic tree. Branch VII members were from monocotyledoneae and some dicotyledoneae species, while branch VIII clustered genes exclusively from dicotyledoneae (including the AhFATAs). The most closely related sequences to the AhFATs were from other Leguminaceae, such as soybean and *Medicago truncatula*.

**Transcription Analysis**

When the abundance of the various AhFAT transcripts was assessed using RNA-Seq data, only 13 of the 21 genes were represented (Fig. 3). Both AhFATAs (*Arah.y195N7.1* and *Arah.YR9F0J.1*) were transcribed in all four samples, and especially in the seed1 and leaf samples. Four of the AhFATB genes (*Arah.YT8BFX.1*, *Arah.L8EKA1.1*, *Arah.L4EP3N.1*, and *Arah.YR9F0J.1*) were expressed. *Arah.YT8BFX.1* and *Arah.L8EKA1.1* were root-specific, whereas *Arah.YR9F0J.1* had a low expression level (with FPKM = 0.81) specifically in the leaf. Branch V had two members, namely, *Arah.YR9F0J.1* and *Arah.75YJ3Z.1*. *Arah.YR9F0J.1* was leaf-specific, whereas *Arah.75YJ3Z.1* was root-specific, whereas *Arah.SJ2WML.1* had a low expression level in...
seed1, root, and leaf (with FPKM values of 0.5, 0.24, and 0.95, respectively) but no expression in seed2. Combined with RNA-Seq data (Fig. 3) and phylogenetic tree analysis (Fig. 2), the results demonstrated that the AhFATB genes in branch I were ubiquitously transcribed in all tissues, whereas the FATB genes in branches III–VI were expressed with tissue-specific patterns.

To verify the universality of this phenomenon, we analyzed the expression patterns of FATBs from P. patens, P. trichocarpa, and G. max (Fig. S1A–C). The three species have many FAB genes in their genomes, similar to peanut. Moreover, they exhibit different levels of evolution. The P. patens genome harbored eight FATBs, two of which (i.e., Pp3c15_18050V3.1 and Pp3c17_23790V3.1) were located closest to the root of the FATB group, while the others were located in branch V (Fig. 2). Pp3c19_20600V3.1 and Pp3c18_15010V3.1 were ubiquitously transcribed in all tissues at high levels, and the others were expressed in different tissues (Fig. S1A). The P. trichocarpa genome harbored 11 FATBs, which were distributed over branches I, IV, V, and VI, with each branch consisting of four, two, two, and three FATBs, respectively (Figs. 2, S1B). Four members in branch

Fig. 2 Phylogenetic tree analysis of plant FATs. The tree was conducted with Mega6 software based on the neighbor-joining method. Statistical confidence in the clade branching points was obtained by running 1,000 bootstrap replicates.
AhFATB2 Isolation and Analysis of Its Sequence

When cDNA prepared from seeds harvested 30 days after flowering was amplified using the primer pairs FATB2-1/F/-R and FATB2-2-F/-R (Supplemental Table S2), the two sequences thereby generated proved to be highly similar (99.4%) to those of Arahy.L4EP3N.1 and Arahy.4E7QKU.1. (Their sequences, designated AhFATB2-1 and AhFATB2-2, have been archived in GenBank under accession numbers MH105081 and MH105082). The two sequences, differing from one another at just eight nucleotides, comprised a 1,245 nt open reading frame, encoding the identical 414 residue polypeptide. Both shared the three conserved catalytic residues Asp311, Asp313, and Cys348, as well as the Acyl-ACP_TE domain. Their level of similarity with the sequence of AhFATB (GenBank number EF117305.1) was 91.4%. AhFATB2-1 mapped to chromosome Arahy.10 and AhFATB2-2 to chromosome Arahy.20, a pair of chromosomes recognized as homologous; they shared a different gene structure (six and seven exons, respectively) and variation in their intronic sequence (Fig. 1).

Semi-quantitative RT-PCR Analyses of AhFATB2-1 and AhFATB2-2

The very high similarity between the AhFATB2-1 and AhFATB2-2 sequences made it impossible to distinguish their transcripts from one another using conventional RT-PCR. To achieve the necessary discrimination, a Taqman-PCR method was employed. A comparison of their transcript abundance in the root, stem, leaf, and flower, sampled at 10, 20, 30, 40, 50, and 60 days after flowering, showed that they shared a very similar transcriptional profile. Both were more abundantly transcribed in the flower than elsewhere (Fig. 4a). In seeds, the expression level of AhFATB2-1 and AhFATB2-2 increased with the seeds development from 10 to 30 d, then peaked at 30 days, and decreased along with the seeds maturation. The period from 20 to 40 d was the most important stage of peanut seed development, the high expression level of AhFATBs was in accordance with the rapid oil accumulation in this stage (Fig. 4b).

Sub-cellular Localization of AhFATB2-1 and AhFATB2-2

When the two genes were expressed in A. thaliana protoplasts, the transgene was expressed most strongly in the vicinity of the chloroplasts, specifically in their outer membranes (Fig. 5).

FA Analysis of Transgenic Yeast

An analysis of the FA content of non-transformed yeast cells highlighted four major fatty acid types, namely C18:1, C16:1, C16:0, and C18:0; these accounted for, respectively, 37.5%, 34.6%, 15.1%, and 8.9% of the total FA content. An equivalent analysis of the transgenic yeast cells showed that cells expressing AhFATB2-1 accumulated 35.0% of the wild type level of C18:1, 50.1% of C16:1, 197.9% of C16:0, and 319.6%...
of C18:0, while the AhFATB2-2 transformants behaved similarly (40.1% of the wild type level of C18:1, 45.5% of C16:1, 170.0% of C16:0, and 277.4% of C18:0). The overall increase in FA content was 92.1% for the AhFATB2-1 transformants and 85.4% for the AhFATB2-2 transformants (Fig. 6a). The constitutive expression of both AhFATB2s in yeast lines clearly altered the composition of FAs (Fig. 6b): the content of saturated FAs (C16:0 and C18:0) increased, whereas that of unsaturated FAs (C16:1 and C18:1) decreased.

The effect of AhFATB2 constitutive expression in A. thaliana

When the two AhFATB2 genes were constitutively expressed in A. thaliana, the FA content of the seeds was raised from 167.6 mg/g dry weight to, respectively, 283.3 mg/g dry weight and 281.2 mg/g dry weight (Fig. 7a). In the seed of non-transgenic A. thaliana, nine fatty acid types were detected, namely C16:0, C18:0, C18:1, C18:2 (linoleic acid), C18:3 (linolenic acid), C20:0 (arachidic acid), C20:1 (eicosenic acid), C20:2 (eicosadienoic acid), and C22:1 (erucic acid). The effect of the transgene was to raise the content of six of these, and in particular that of C16:0 and C18:0; meanwhile, the content of both C18:1 and C20:1 was reduced. In non-transgenic A. thaliana seeds, the predominant fatty acids were unsaturated (C18:2, C18:3, C20:1, C18:1), accounting for 67.5% of the total fatty acid content (Fig. 7b). However, in the seed of the transgenic A. thaliana lines, C16:0 became the most abundant fatty acid, representing around 31% of the total fatty acid content; at the
same time, the content of the saturated fatty acids C18:0 and C20:0 was also significantly increased.

**Discussion**

Thioesterases are ubiquitous in both prokaryotic and eukaryotic organisms. Based on their catalytically active site, their oligomerization, and their substrate specificity, they have been allocated EC numbers from 3.1.2.1 to 3.1.2.27, with many remaining unclassified (EC 3.1.2.-) (Cantu et al. 2010). The FATs (EC 3.1.2.14) hydrolyze FAs synthesized by the FA synthase complex. The FATs found in bacteria and plants have been classified into ten subfamilies (Jing et al. 2011); here, a collection of 179 documented plant FATs, which included both FATAs and FATBs, fell into eight clades (Fig. 2). Peanut FATBs were distributed over the four branches I, III, V, and VI. Of particular interest
was the finding that one of these (branch III) was composed entirely of FATBs synthesized by *Arachis* spp. (seven by *A. hypogaea*, five by *A. ipaensis* and four by *A. duranensis*) (Fig. 2). A likely explanation for this rather surprising outcome is that these genes have evolved from a single ancestral copy by duplication. Of the seven *A. hypogaea* branch III FATBs, only two (*Arahy.3BV5AA.1* and *Arahy.BC15R8.1*) appeared to be active (Fig. 3): transcripts of the former gene were particularly abundant in the root, whereas the latter was transcribed—at a rather low level—exclusively in the leaf.

The large number of *FAT* genes present in the genome varies widely between species. This variation cannot be accounted for by suggesting that their number simply accumulates during evolution, since although the algal species *C. reinhardtii* encodes just one FAT, the genomes of the primitive land species *M. polymorpha* (a liverwort), *S. melloendorfii* (a lycophyte), and *P. patens* (a bryophyte) harbor, respectively, four, six, and ten *FAT* genes, a greater number than is harbored by most angiosperm species (Supplemental Table S1). Nor can the large number of *FAT* genes harbored by peanut (21 *AhFATs*) does not be explained by its being a tetraploid, since the hexaploid species bread wheat harbors just ten genes. Finally, there is no discernible relationship between the number of *FAT* genes and genome size, as can be seen from a comparison between the small genome species *A. thaliana* and the large genome species bread wheat. A possibility which remains to be tested is that selection for high oil content in the seed tends to favor the accumulation of *FAT* genes.

Based on RNA-Seq data, only 13 of the 21 peanut *FATs* were transcribed in root, leaf, and seed tissue (Fig. 3), and only six of these were ubiquitously transcribed. This observation implies a degree of sub-functionalization among the genes, as well as suggesting a high rate of gene silencing. However, the present experiment did not cover many other *FAT* genes and genome size, as can be seen from a comparison between the small genome species *A. thaliana* and the large genome species bread wheat. A possibility which remains to be tested is that selection for high oil content in the seed tends to favor the accumulation of *FAT* genes.

In response to the world energy crisis and global climate change, biodiesel has attracted renewed interest in the exploration for sustainable development (Zinoviev et al. 2010). Free fatty acids (FFAs) have been extensively applied in the manufacture of biofuels, cosmetics, and pharmaceutical drugs (Shin et al. 2016). Given that the overexpression of *FATs* has considerably increased the production of FFAs in prokaryotes and eukaryotes (Chen et al. 2017; Li et al. 2017; Lin and Lee 2017; Shin et al. 2016; Tan and Yuan 2017), *FATs* remain the first choice in the production of oleochemicals. The substrate specificity of *FATs* often determines the composition of oils. Thus, many plant *FATs* have been isolated and remolded to produce oleochemicals with desired chain lengths (Lozada et al. 2018). Peanut is an important oil crop (with seed oil content up to 50%), and its genome possesses 21 *FATs*, which are far more than any other species (Supplemental Tables S1, S3). Not all *AhFATs* are suitable for oleochemical production. At least four *AhFATs*, namely, *Arahy.KT8BMX.1*, *Arahy.L8EKIA.1*, *Arahy.LAEP3N.1*, and *Arahy.4E7QKU.1*, with high expression levels in branch I are good choices for biofuel production. Studies have demonstrated that all these *AhFATs* can increase the oil content of the transformed microalgae (Chen et al. 2012, 2017; Li et al. 2017; Wen et al. 2012), yeast, and plants (Figs. 6, 7). In addition, the *FATs* in branch I from other plants are good candidates for biofuel production because of their high expression level (Supplemental Fig. S1).

**Conclusions**

In summary, we analyzed the peanut *FAT* gene family in peanut genome, and analyzed their phylogeny relationship with other plant *FATs*. Here, two new *AhFATB* genes from cultivated peanut were cloned and transformed into *S.
cerevisiae and A. thaliana for function verification. Results showed that both of them could increase the FA content significantly, especially that of saturated FAs.

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Compliance with Ethical Standards

Conflict of interest The authors declare that they have no conflict of interest.

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