A Novel L-ascorbate Peroxidase 6 Gene, ScAPX6, Plays an Important Role in the Regulation of Response to Biotic and Abiotic Stresses in Sugarcane

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The L-ascorbate peroxidase 6 gene (APX6) is one of the most important genes for scavenging H$_2$O$_2$ and plays a vital role in plant resistance to environmental stresses. In this study, a novel ScAPX6 gene (GenBank Accession No. KT907352) was obtained from a sugarcane variety (ROC22). Bioinformatics analysis showed that ScAPX6 has a cDNA length of 1,086 bp and encoded 333 amino acid residues. Subcellular localization confirmed that ScAPX6 was located in the chloroplast. Enhanced growth of Escherichia coli BL21 cells that expressed ScAPX6 showed high tolerance under copper (Cu) stress. Real-time quantitative PCR analysis revealed that ScAPX6 was constitutively expressed wherein with the highest expression levels in sugarcane pith and leaf and the lowest in the root. ScAPX6 was down-regulated by salicylic acid (SA), hydrogen peroxide (H$_2$O$_2$), polyethylene glycol (PEG) and sodium chloride (NaCl) stimuli. Interestingly, it was significantly up-regulated under the stresses of abscisic acid (ABA) and methyl jasmonate (MeJA) wherein with the highest inducible expression levels at 6 h at 6.0- and 70.0-times higher, respectively than that of control. Overexpression of ScAPX6 in Nicotiana benthamiana leaves enhanced the resistance to the infection of tobacco pathogens Pseudomonas solanacearum and Fusarium solani var. coeruleum. These results implied that ScAPX6 might positively respond to ABA, MeJA, and Cu, but might negatively respond to the stresses of SA, H$_2$O$_2$, PEG, and NaCl. Keeping in view the current investigation, ScAPX6 could be associated with the hypersensitive response (HR) or immunity of sugarcane, which will provide a baseline for the function identification of sugarcane ScAPX6.

Keywords: sugarcane, L-ascorbate peroxidase 6 gene, subcellular localization, biotic and abiotic stresses, transient overexpression
INTRODUCTION

In addition to *Oryza sativa*, *Triticum aestivum*, and *Ze a mays*, sugarcane is the fourth largest staple food for the people of China. Sugarcane planting and production are of great significance in sugar supply (Li, 2000). However, the growth and development of sugarcane is severely affected by various abiotic and biotic stresses, such as drought, cold, salinity, heavy metals, high temperature, viruses, fungi, and so on (Li, 2000; Xu et al., 2008). As reported, environmental stimuli can induce active oxygen system which may cause injury to plant cells (Mittler et al., 2004, 2011). Peroxidases (EC number 1.11.1.1.), including glutathione peroxidase (GPX), catalase (CAT), and ascorbate peroxidase (APX), are widespread in organisms and can remove the reactive oxygen (Shigeoka et al., 2002; Apel and Hirt, 2014). APX, belonging to type I heme peroxidase and copper oxidase family, is widely employed in plants and can rapidly scavenge hydrogen peroxide (H₂O₂) in the ascorbic acid (ASA) and glutathione (GSH) cycle (Shigeoka et al., 2002). The role of APX is highly specific to ascorbic acid, that is, to help electron donor to oxidation (Chen and Asada, 1989; Mittler, 2002; Foyer and Noctor, 2005).

According to the orientation characteristic, there are three mainly APX subfamilies in plants, such as cytoplasm APX (cAPX), thylakoid APX (tAPX), and APX-R (Apx-Related) (Mano et al., 1997; Shigeoka et al., 2002; Chew et al., 2003; Dunand et al., 2011). APXs gene have been reported in several plants such as *Nelumbo nucifera* (Chen et al., 2011), *Hordeum vulgare* (Shi et al., 2001), *Solanum tuberosum* (Kawakami et al., 2002), *Z. mays* (Breusegem et al., 1995), and *Vitis pseudoreticulata* (Lin et al., 2006). The expression of APX can be regulated by environmental stimuli, such as salt (Badawi et al., 2004), temperature (Kawakami et al., 2002), high light (Maruta et al., 2010), and heavy metal stresses (Pallavi and Dubey, 2007). Research has also shown that APX induced by adversity stress can regulate the content of H₂O₂ in the cell and redox signaling, and then affect plant tolerance to the osmotic stress (Andréia et al., 2012). This characteristic of APX enzyme activity may be treated as one of the physiological and biochemical indexes measuring crop resistance to biotic and abiotic stresses, for instance, water deficit (Nayyar and Gupta, 2006) and high temperature (Almeselmani et al., 2006). Kornyeyev et al. (2001) transferred the chloroplast APX gene into *Gossypium spp.*, and it was found that APX activity in the transgenic cotton leaf was higher than that of the wild type. Overexpression of tAPX genes increased the resistance of *Nicotiana tabacum* and *Arabidopsis thaliana* to the oxidative stress induced by methyl violet essence (Yabuta et al., 2002; Murgia et al., 2004).

Until now, there are four nucleotide sequences of APX genes, which have been identified in sugarcane. Wang Z. Q. et al. (2015) indicated that the APX enzyme activity in sugarcane smut resistant variety Yacheng05-179 was significantly higher than the susceptible variety Luicheng03-182 after inoculated with *Sporisorium scitamineum*. As a result the expression level of one sugarcane ScAPX gene (GenBank Acc. No. KJ7565501) increased under the stresses of salicylic acid (SA), methyl jasmonate (MeJA), abscisic acid (ABA), H₂O₂, sodium chloride (NaCl), and polyethylene glycol (PEG). Wang S. et al. (2015) demonstrated that TAPX gene (GenBank Acc. No. IQ958327) played a part in sugarcane resistance to osmotic stress. Huang et al. (2013) found that sugarcane S-APX2 gene showed highly homologous with rice APX (GenBank Acc. No. XP_002463451.1) and mazie APX (GenBank Acc. No. DAA41857.1). Another sugarcane APX gene (GenBank Acc. No. KX235995) was found in *Saccharum arundinaceum*, but its function was unclear.

From all the above, cloning the APXs gene of different isozymes is necessary to better understand the APX gene family and know more about their expression levels under different stress conditions. In the present study, a putative APX6 unigene, named as ScAPX6, was cloned and identified based on our previous transcriptome data of sugarcane in response to sorghum mosaic virus (SrMV) infection (Bioproject number: PRJNA379719). The sequence characters of ScAPX6 was analyzed by bioinformatics analysis, and the expression patterns of ScAPX6 gene after exposure to various stresses, such as ABA, MeJA, SA, H₂O₂, PEG, NaCl, and copper (Cu), were detected by real-time quantitative polymerase chain reaction (qRT-PCR). Furthermore, its expression in *Escherichia coli*, subcellular localization and transient overexpression in *Nicotiana benthamiana* were also investigated. This study will be helpful to understand the gene function of ScAPX6 in sugarcane.

MATERIALS AND METHODS

Plant Material and Treatments

For the analysis of the tissue-specific expression of ScAPX6, six healthy plants of 10 months old of ROC22 were used, and then +1 leaf, root, bud, skin and pith were collected. The samples were fixed in liquid nitrogen and stored at −80°C until the extraction of total RNA.

For the abiotic treatment, uniform tissue cultured plantlets of ROC22 at the five or six leaf stage were transferred to water for one week and then treated by the following six stress conditions with root dipping of 5 mmol L⁻¹ SA, 100 μmol L⁻¹ MeJA, 100 μmol L⁻¹ ABA, 10 μmol L⁻¹ H₂O₂, 25.0% PEG 8000, 250 mmol L⁻¹ NaCl, and 100 mmol L⁻¹ copper chloride (CuCl₂), at 28°C with 16 h light and 8 h darkness (Su et al., 2014a). The whole plantlets under SA, MeJA, H₂O₂, PEG, and NaCl stresses were harvested at 0, 6, 12, and 24 h, respectively. Another set of plantlets under Cu stress was harvested at 0, 12, 24, and 48 h, respectively. Three plantlets per time point were gathered and immediately fixed in liquid nitrogen, and stored at −80°C until the extraction of total RNA.

Total RNA Extraction and the First-strand cDNA Synthesis

Total RNA of the treated samples was extracted by Trizol® Reagent (Invitrogen, Carlsbad, CA, USA) according to the manufacturer's instructions. The first-strand cDNA synthesis was performed using Prime-Script™ RT Reagent Kit (TaKaRa, Dalian, China) following manufacturer's instructions and tested by 1% agarose gel electrophoresis.
Sugarcane ScAPX6 Gene Isolation and Gateway Entry Vector Construction

The sequence of a putative APX6 unigene (ScAPX6) from our previous transcriptome data of sugarcane in response to SrMV infection was used to design the cloning primer APX6-1F/1R (Table 1). The first-strand cDNA of ROC22 was used as amplification template. The reverse transcription–polymerase chain reaction (RT-PCR) procedure was 94°C for 4 min; 94°C for 30 s, 55°C for 30 s, 72°C for 2 min, 35 cycles; and 72°C for 10 min. RT-PCR products were gel-purified and cloned into pMD19-T vector (TaKaRa, Dalian, China), and then transformed into E. coli DH5α competent cells and sequenced (Sangon, Shanghai, China).

The open reading frame (ORF) of ScAPX6 with Gateway entry adapters attB1 and attB2 was amplified from the plasmid of pMD19-T-ScAPX6 by the primers (APX6-3F/3R) (Table 1). The touchdown PCR procedure was 94°C for 4 min; 94°C for 30 s, 70°C for 30 s and then each loop drop 0.5°C for 1 min and 30 s, 35 cycles; and 72°C for 10 min. The PCR amplification products were gel-purified and transformed into the Gateway® donor vector of pDONR221 (Invitrogen, USA) following the manufacturer’s instructions of Gateway® BP Clonase™ II Enzyme Mix (Invitrogen, USA). The mixture of BP reaction was transformed into DH5α competent cells and sequenced (Sangon, Shanghai, China). The positive plasmid pDONR221-ScAPX6 was achieved and then used for the constructions of prokaryotic expression vector and eukaryotic expression vector.

Bioinformatics Analysis

The ORF was translated and analyzed by ORF Finder (https://www.ncbi.nlm.nih.gov/orffinder/). Conserved domain of ScAPX6 was predicted by the SMART program (http://smart.embl-heidelberg.de/) and NCBI Conserved Domains Database (CDD) (http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi). The ExPaSy tool (http://us.expasy.org/tools) was used to predict the basic physical and chemical properties of ScAPX6. The cleavage sites of the signal peptides were predicted by SignalP 4.1 Server (http://www.cbs.dtu.dk/services/SignalP/). Prediction of transmembrane helices in ScAPX6 protein was performed by TMHMM Server v. 2.0 (http://www.cbs.dtu.dk/services/TMHMM-2.0/). Psort software was used to predict the subcellular localization of ScAPX6. GOR IV software (https://npsa-prabi.ibcp.fr/cgi-bin/npsa_automat.pl?page=/NPSA/npsa_gor4.html) was used to analyze the secondary structure of ScAPX6. The protein 3D model was predicted by SWISSMODEL software (http://swissmodel.expasy.org/). The homologous sequences of ScAPX6 were obtained using Blastp in NCBI. DNAMAN software was used for the multiple sequence alignment. The phylogenetic tree of ScAPX6 was constructed with amino acid sequences from other species by the neighbor-joining (NJ) method (1,000 bootstrap replicates) using the MEGA 6.06 (Saitou and Nei, 1987).

Subcellular Colocalization Assay

The ORF of ScAPX6 was amplified by the primer APX6–4F/4R, and then was inserted into the Bsal and Eco311I restriction sites of the pBWA(V)HS-cddb-GLosgfpl vector. Then the recombinant vector pBWA(V)HS-ScAPX6-GLosgfpl and the chloroplast marker vector were co-transformed in rice protoplasts with PEG solution (40% W/V PEG 4000, 0.2 mol-L⁻¹ mannitol and 0.1 mol-L⁻¹ calcium chloride). The mixture was cultured in dark for 30 min, and then the protoplasts was gathered and cultured in dark for 16–24 h. The method of transformation of rice protoplasts was followed by Datta and Datta (1999). The subcellular localization of the fusion protein was observed by a confocal laser scanning microscope Leica TCS SP5 (Germany).

Expression of ScAPX6 in E. coli BL21 (DE3) Strain

The plasmid of pDONR221-ScAPX6 was digested with Aval and then gel-purified for LR reaction with prokaryotic expressive vector of pEZYHb according to the manufacturer’s instructions of LR Clonase™ II Enzyme Mix (Invitrogen, USA). The recombinant plasmid of pEZYHb-ScAPX6 was transformed into the competent cells E. coli BL21 (DE3) and then induced by 1.0 mmol-L⁻¹ isopropyl β-D-thiogalactoside (IPTG) at 28°C for 0, 2, 4, and 8 h (Guo et al., 2008). LB medium with E. coli BL21 (blank) and BL21+pEZYHb (control) were induced by 1.0 mmol-L⁻¹ IPTG for 0 and 8 h, respectively. The collected bacterial protein was analyzed by 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE).

Spot assay was conducted to study the responses of E. coli BL21 cells expressing the ScAPX6 gene under abiotic stress, such as NaCl, Cu and PEG. When OD₆₀₀ of E. coli BL21 cells in LB medium (containing 80 µg·mL⁻¹ ampicillin) reached to 0.6, IPTG with a concentration of 1.0 mmol·L⁻¹ was added, and then the cells were grown at 37°C for 12 h. The cultures were diluted to OD₆₀₀ = 0.6, and then diluted to two levels of 10⁻³ and 10⁻⁴ (Guo et al., 2012). Ten microliters from each level was spotted on LB plates containing NaCl (250, 500, and 750 mmol·L⁻¹), CuCl₂ (250, 500, and 750 µmol·L⁻¹) and PEG (15, 30, and 45%), respectively (Su et al., 2013). All plates were cultured in 37°C overnight and photographed.

Gene Expression Patterns of ScAPX6

SYBR Green Master (ROX) (Roche, China) and a 7500 qRT-PCR system (Applied Biosystems, South San Francisco, CA, USA) were applied to analyze gene expression levels of ScAPX6 in different tissues and in response to various stresses. The sequence-specific primer of ScAPX6 (APX6-2F/2R) (Table 1) was designed by Premier 5.0 software. The primer combination of clathrin adaptor complex (CAC) and cullin (CUL) (Table 1) was regarded as the internal control (Guo et al., 2014). The 20 µL reaction system containing 10 µL SYBR Green Master Mix, 0.8 µL each of 10 µmol·L⁻¹ upstream and downstream primers, 2 µL cDNA templates (20 x diluted cDNA) and 6.4 µL double distilled water. Each qRT-PCR was conducted in triplicate. The qRT-PCR procedure was 50°C for 2 min; 95°C for 10 min; 35 cycles of 95°C for 15 s, and 60°C for 1 min. The 2⁻ΔΔCt method (Livak and
TABLE 1 | Primers used in this study.

| Primer       | Sequence information (5′–3′)                           | Strategy                                      |
|--------------|--------------------------------------------------------|-----------------------------------------------|
| APX6-1F      | CTTGAGAAGGCAAGCCAGGA                                    | Gene cloning                                 |
| APX6-1R      | CGAGACACTGAGTACAGGGGAG                                  | Gene cloning                                 |
| APX6-2F      | QTGTGATTTGGCTGCTGTCG                                    | qRT-PCR analysis                             |
| APX6-2R      | TGCTGACCGCTGCTGCTG                                      | qRT-PCR analysis                             |
| CUL-F        | TGTGACTGGTACAGGGGAG                                     | qRT-PCR analysis                             |
| CUL-R        | ACACGTGAGCCAAAGGCA                                      | qRT-PCR analysis                             |
| CAC-F        | AGATCACTCCACCTCCTGNTAC                                  | qRT-PCR analysis                             |
| CAC-R        | AGATCACTCCACCTCCTGNTAC                                  | qRT-PCR analysis                             |
| APX6-3F      | GGGGACAAGTTTGTACAAAAAAGCAGGCTTCGAGCTGCCAAAGCCTC        | Gateway entry vector construction and RT-PCR analysis |
| APX6-3R      | GGGGACCACTTTGTACAAAAAAGCAGGCTTCGAGCTGCCAAAGCCTC        | Gateway entry vector construction and RT-PCR analysis |
| APX6-4F      | CGGTGCTCACACCTCCCAAGGACAGGACG                          | Subcellular localization vector construction  |
| APX6-4R      | CGGTGCTCACACCTCCCAAGGACAGGACG                          | Subcellular localization vector construction  |
| NtHSR201-F   | CAGCAGTCCTTTGGCGTTGTC                                   | qRT-PCR analysis                             |
| NtHSR201-R   | GCTCAGTTGACCGCTGATG                                     | qRT-PCR analysis                             |
| NtHSR203-F   | TGCCACGGCTGATGGGATA                                     | qRT-PCR analysis                             |
| NtHSR203-R   | GCACGAAACCTCCCAAGGACAGGACG                              | qRT-PCR analysis                             |
| NtHSR515-F   | TTGGGACGATGAGTTGTC                                      | qRT-PCR analysis                             |
| NtHSR515-R   | TTGGGACGATGAGTTGTC                                      | qRT-PCR analysis                             |
| NtPR-1a-c-F  | AACCTTGAGCTTGCAGGACGAGGAC                              | qRT-PCR analysis                             |
| NtPR-1a-c-R  | AACCTTGAGCTTGCAGGACGAGGAC                              | qRT-PCR analysis                             |
| NtPR2-F      | TGAGTGTCGCTCTGAGGTCATTGATG                              | qRT-PCR analysis                             |
| NtPR2-R      | AGCTCTCGCCCCCGGTTC                                     | qRT-PCR analysis                             |
| NtPR3-F      | CAGGAGGTATTGCTGCTGCTGCTG                               | qRT-PCR analysis                             |
| NtPR3-R      | CAGGAGGTATTGCTGCTGCTGCTG                               | qRT-PCR analysis                             |
| NtEF1-α-F    | CAGGAGGTATTGCTGCTGCTGCTG                               | qRT-PCR analysis                             |
| NtEF1-α-R    | CAGGAGGTATTGCTGCTGCTGCTG                               | qRT-PCR analysis                             |
| NtAcdeaminase-F | TCTGAGGTTACTGTCTGACTTGGATG                              | qRT-PCR analysis                             |
| NtAcdeaminase-R | TGAGGTTACTGTCTGACTTGGATG                              | qRT-PCR analysis                             |
| NIEF1-α-F    | TGAGGTTACTGTCTGACTTGGATG                                | qRT-PCR analysis                             |
| NIEF1-α-R    | TGAGGTTACTGTCTGACTTGGATG                                | qRT-PCR analysis                             |

attB1 and attB2 adapters were underlined in the forward primer APX6–3F and in the reverse primer APX6–3R, respectively.

Schmittgen, 2001) was employed to analyze the qRT-PCR data.

Transient Overexpression of ScAPX6 in N. benthamiana

To study the role of ScAPX6 in response to pathogen infection and its hypersensitive reaction in plant, an overexpressed vector pEarleyGate 203-ScAPX6 was constructed by Gateway cloning technique according to the manufacturer’s instructions of LR Clonase™ II Enzyme Mix (Invitrogen, USA). N. benthamiana leaves was inoculated with the vector of pEarleyGate 203-ScAPX6 by an Agrobacterium-mediated transient expression method conducted by Su et al. (2014b). Two important tobacco pathogens, Pseudomonas solanacearum and Fusarium solani var. coeruleum, were cultured in potato dextrose water (PDW) liquid medium at 28°C. When the two pathogens cells were cultured to an OD600 of 0.8, they were separately infected into the treated leaves that were agroinfiltrated with pEarleyGate 203-ScAPX6 for 24 h. Then the 3,3′-diaminobenzidine (DAB) staining, trypan blue staining and transcript analysis of the eight tobacco immunity-associated marker genes (Table 1), were conducted by the treated N. benthamiana leaves according to Su et al. (2016). RT-PCR was used to detect whether ScAPX6 has been overexpressed in N. benthamiana, with the RNA of treated leaves and ScAPX6 specific primer (APX6-3F/3R, Table 1), the NIEF1-α was treated as control. RT-PCR procedure was 94°C for 4 min; 94°C for 30 s, 72°C for 30 s, 72°C for 2 min, 35 cycles; and 72°C for 10 min. All treatment materials were cultured at 24°C (16 h light/8 h darkness) and then photographed at 1 and 7 day (d) separately. Each test was repeated three times.

DAB and trypan blue staining were used for histochemical analysis of Agrobacterium-infiltrated leaves. The leaves was soaked in DAB-HCl solution (1.0 mg·mL⁻¹, pH 5.8), and then cultured in the dark for 12 h. The leaves were destained with 95% ethanol at 100°C for 10 min (Su et al., 2014a), and then was imaged for H₂O₂ detection with a stereoscopic microscope (Nikon, Tokyo, Japan) and a light microscope (Leica, Wetzlar, Germany). Three biological replications were prepared. The leaves were also deal with...
trypan blue mixture, containing 10 mL lactic acid, 10 g phenol, 10 mL glycerol, 30 mL absolute ethanol, 10 mg trypan blue, and 10 mL distilled water, and then was boiled for 5 min. After staining, the leaves were soaked in a chloral hydrate solution (2.5 g·mL\(^{-1}\)) for decoloring (Dang et al., 2013). The blue color of the leaves for the cell death was also imaged with a stereoscopic microscope (Nikon, Tokyo, Japan) and a light microscope (Leica, Wetzlar, Germany).

## RESULTS

### Cloning and Bioinformatics Analysis of ScAPX6

In the present study, a full-length cDNA of APX6 unigene, which was named as ScAPX6 (GenBank Acc. No. KT907352), was isolated from ROC22. The cDNA sequence length of ScAPX6 was 1,086 bp (Figure S1) with a complete ORF (1,002 bp, from position 27 to position 1,025), encoding 333 amino acid

![FIGURE 1](image1.png) Nucleotide acid sequence and deduced amino acid sequence of sugarcane ScAPX6 gene obtained by RT-PCR. The start codon and termination codon were underlined in black. The primer used in RT-PCR was underlined in red line. The peroxidase like superfamily domain contains 222 amino acids (from 103 to 324) was highlighted in red. The amino acids highlighted in yellow represented the heme binding site. *, Stop codon.

![FIGURE 2](image2.png) Predicted 3D structure of ScAPX6. The plant-peroxidase-like domain was in red. Saccharum spp. hybrids (AMQ80947.1), Sorghum bicolor (XP_002445876.1), Setaria italica (XP_004973913.1), Oryza sativa Japonica Group (EAZ43377.1).
residues. ScAPX6 had a molecular mass of 36.21 kDa and an isoelectric point (pI) of 6.91. CDD search of NCBI showed that ScAPX6 belonged to a member of the plant-peroxidase-like superfamily (Figure 1). The instability index of ScAPX6 protein was 48.10, suggesting that ScAPX6 might be an unstable acid hydrophilic protein (Walker, 2005). Secondary structure

![Phylogenetic analysis of deduced amino acid sequence from ScAPX6 and other ascorbate peroxidases proteins. The GenBank accession number of proteins were according to Teixeira et al. (2004) and downloaded from NCBI. The neighbor-joining method with 1,000 bootstrap replications was used.](image1)

![Subcellular localization analysis of ScAPX6 in rice protoplasts. (a,d) green fluorescence; (b,f) visible light; (e) red fluorescence from chloroplast marker; (c,g) merged light.](image2)
prediction of ScAPX6 predicted that the percentages of alpha-helix, random coil, and extended strand were 44.74, 42.94, and 12.31%, respectively.

Furthermore, SWISSMODEL program showed that the main spatial structures of ScAPX6 were alpha-helix and random coil (Figure 2). Comparing ScAPX6 with O. sativa Japonica Group APX6 (EAZ43377.1), S. italic APX6 (XP_004973913.1), and S. bicolor APX6 (XP_002445876.1), we found that the spatial structure of these four APX6 was basically in line with each other, suggesting that ScAPX6 owned high conservation of spatial structure with different plant species. Psort software predicted that ScAPX6 might be located in the chloroplast thylakoid membrane, plasma membrane, chloroplast stroma, and chloroplast thylakoid space with the probabilities of 71.9, 65.0, 56.1, and 56.1%, respectively.

According to the classification method by Teixeira et al. (2004), the phylogenetic tree was separated into three groups, including cytosolic isoforms, Apx-R isoforms and chloroplastic isoforms (Figure 3). ScAPX6 was clustered into group chloroplastic isoforms. Two APX proteins reported in Saccharum hybrid cultivar, which were ScAPX (AIG52216.1) and TAPX (AGD80596.1), were also clustered into the same clade as ScAPX6.

Subcellular Localization
The recombinant vector pBWA(V)HS-ScAPX6-GLosgfP was constructed to understand the subcellular location of ScAPX6. The results showed that ScAPX6 and the chloroplast marker were located in the same place, so it was confirmed that ScAPX6 was located in the chloroplast, which is in accordance with the results of prediction (Figure 4).

Expression of ScAPX6 in E. coli BL21 (DE3) Strain
The ScAPX6 gene was combined with the expression vector pEZYHb and then was transformed into E. coli BL21 cell. The SDS-PAGE analysis (Figure 5) showed that ScAPX6 was expressed as a recombinant protein in the BL21 cell. In Figure 5, after induced by 1.0 mmol.L⁻¹ IPTG at 28°C for 2, 4, and 8 h, an obvious accumulation protein (including the 6× His-tag) at approximate 55 kDa was observed.

Overexpression of ScAPX6 in E. coli Enhances Cell Growth under Cu Stress
It has been reported that the APX activity was up-regulated by abiotic stress, such as heavy metal, high salinity, drought, high temperature, and wounding (Shi et al., 2001). In this study, spot assay was performed to study the function of ScAPX6 in response to abiotic stress in vivo. The control (BL21+pEZY-Hb) and the gene-expressed cells (BL21+pEZY-Hb-ScAPX6) grew in LB plates containing NaCl, CuCl₂, and PEG were performed (Figure 6). It had been recorded that the recombinant ScAPX6 cells showed a more rapid growth than the control in LB plates with PEG and CuCl₂ supplement, but not with NaCl, suggesting that the overexpression of ScAPX6 in E. coli could enhance its tolerance to PEG and Cu stress.

Tissue-Specific Expression of ScAPX6
qRT-PCR analysis showed that ScAPX6 was constitutively expressed in all five kinds of sugarcane tissues, including root, bud, skin, leaf, and pith, but with different expression levels (Figure 7). ScAPX6 showed the highest expression levels in the pith and leaf, and then on the skin, while the transcript in root was at the lowest level.

Gene Expression Patterns of ScAPX6 in Response to Abiotic Stress
qRT-PCR analysis revealed that the ScAPX6 gene exhibited different expression characteristics in response to ABA, MeJA, SA, H₂O₂, PEG, NaCl, and Cu stimuli (Figures 8A,B). As shown in Figure 8A, the transcripts of ScAPX6 were remarkably up-regulated under the stresses of ABA and MeJA, and with the highest inducible expression levels at 6 h, which were 6.0- and 70.0-times higher than that of control, respectively. However, ScAPX6 was down-regulated during the SA treatment and rapidly decreased at 6 h. Under the stress of PEG, the expression of ScAPX6 showed no change at 6 h, and then decreased at 12 h. ScAPX6 was down-regulated after the treatment of H₂O₂ and NaCl, but was up-regulated by the CuCl₂. These results demonstrated that ScAPX6 might positively respond to ABA, MeJA, and Cu stresses but negatively respond to SA, H₂O₂, PEG, and NaCl stresses.

Transient Overexpression of ScAPX6 Induces a Defense Response in N. benthamiana
After transient overexpression of ScAPX6 in N. benthamiana leaves for 1 day (d), the transcripts of ScAPX6 were detected by qRT-PCR (Figure 9A). As shown in Figures 9A,B darker DAB staining color and more intense trypan blue staining cells were observed in ScAPX6 leaves than that in the control (35S::00) after infiltration for 2 and 6 d, respectively. The
Liu et al.  
ScAPX6 Functions in Sugarcane Defense

**FIGURE 6**  | Spot assays of BL21+pEZY-Hb-ScAPX6 (b) and BL21+pEZY-Hb (control) (a) on LB plates with NaCl, PEG and CuCl$_2$. Isopropyl $\beta$-D-thiogalactoside (IPTG) was added to the cultures of BL21+pEZY-Hb-ScAPX6 and BL21+pEZY-Hb to induce the expression of recombinant protein. The cultures were adjusted to $\text{OD}_{600} = 0.6$. Ten microliters from $10^{-3}$ (left side of the red line on the plate) to $10^{-4}$ (right side of the red line on the plate) dilutions were spotted onto LB plates without any supplement (CK) (A) or with NaCl (250, 500, and 750 mmol·L$^{-1}$) (B), PEG (15, 30, and 45%) (C) and CuCl$_2$ (250, 500, and 750 µmol·L$^{-1}$) (D), respectively. NaCl, sodium chloride; PEG, polyethylene glycol; CuCl$_2$, copper chloride.

**FIGURE 7**  | Tissue-specific expression analysis of ScAPX6 in sugarcane. The error bars represented the standard error of each treating group ($n = 3$). Data were normalized to the CAC and CUL expression level. All data points were means ± SE ($n = 3$). Different lowercase letters indicate a significant difference, as determined by the Duncan’s new multiple range test ($p < 0.05$).

Eight immunity-associated marker genes in *N. benthamiana* were induced by transient overexpression of ScAPX6 (Figure 9C). The hypersensitive response (HR) marker genes, *NhHSR201* and *NhHSR203*, showed no change in transcript, while *NhHSR515* was up-regulated. The expression level of SA-responsive gene *NtPR2* remained unchanged, while *NtPR-1a/c* and *NtPR3* and two ethylene synthesis dependent genes, *NtEFE26* and *NtAccdeaminase*, were all down-regulated. Compared with the control leaves, 35S::ScAPX6 exhibited darker color reflecting high levels of H$_2$O$_2$ accumulation and intense hypersensitivity response.

To further investigate the response of ScAPX6 to pathogen, two tobacco pathogens, *P. solanacearum* and *F. solani* var. *coeruleum*, were separately injected into *N. benthamiana* containing 35S::ScAPX6 or the control. After inoculation with *P. solanacearum*, no disease symptom was found between 35S::ScAPX6 and the control leaves for 1 d, while 35S::ScAPX6 exhibited darker color than the control at 1 d by DAB staining. With elongated treatment time, although the DAB staining showed no difference between 35S::ScAPX6 and the
control leaves, the leaves in the control showed slight yellow phenomenon and necrotic spot, while the 35S:ScAPX6 only exhibited the faint wilting symptom after inoculation at 7 d (Figure 9D). After challenging with *F. solani* for 1 and 7 d (Figure 9E), the expression levels of *NhHSR201*, *NhHSR203*, and *NhPR2*, were unchanged or down-regulated in the control and 35S:ScAPX6 leaves. The transcripts of *NhPR-1a/c* and *NhPR3* were significantly down-regulated at 1 d and increased at 7 d in the control leaves, but were down-regulated or remained unchanged in the 35S:ScAPX6 leaves. The expression levels of *NhHSR515* and *NieFE26* were unchanged at 1 and 7 d and *NhAccdeaminase* was up-regulated at 7 d in the control leaves, while these three genes in the 35S:ScAPX6 leaves were all unchanged at 1 d and reached the peak values at 7 d after inoculation.

Likewise, for DAB staining, the 35S:ScAPX6 leaves showed darker color than the control after inoculation with *F. solani* var. *coeruleum* for 1 d and exhibited no difference at 7 d. No apparent disease symptom differences between 35S:ScAPX6 and the control were found at 1 d after inoculation (Figure 9F). Some symptoms, such as wilting, decay phenomenon and necrotic spot, were observed in the control leaves at 7 d, while the leaves in the control showed slight yellow phenomenon and necrotic spot, which was consistent with the result of bioinformatics analysis (Figure S1), which was different from the other already reported sugarcane APX genes in NCBI (ScAPX: GenBank Acc. No. KT907352) was cloned (Figure S1), which was different from the result of bioinformatics analysis.

**DISCUSSION**

Plant APXs are a multi-gene family (Shigeoka et al., 2002). Many plants APXs genes have been cloned and identified, including eight in *O. sativa* (Teixeira et al., 2005, 2006), four in *Vigna unguiculata* (D’Arcymeta et al., 2006) and *Spinacia oleracea*, respectively (Ishikawa et al., 1995, 1996, 1998), six in *Eucalyptus grandis* (Teixeira et al., 2005), seven in *Lycopersicon esculentum* (Najami et al., 2008), and nine in *A. thaliana* (Panchuk et al., 2002; Mittler et al., 2004; Narendra et al., 2006). In the present study, based on a putative APX6 unigene sequence from our previous transcriptome data, a sugarcane ScAPX6 gene (GenBank Acc. No. KJ7565501; GenBank Acc. No. KT907352) was cloned (Figure S1), which was different from the other already reported sugarcane APX genes in NCBI (ScAPX: GenBank Acc. No. KJ7565501; TAPX: GenBank Acc. No. JQ958327; APX: GenBank Acc. No. KX235995), and shared only 19.79% identity at the amino acid sequence level. Najami et al. (2008) found that in *Solanum lycopersicum* three cytosolic *SIA* genes, *SIA*1, *SIA*2, and *SIA*3, showed a high sequence identity (>90%). Teixeira et al. (2004) found that APX contained two isoforms, chloroplastic and nonchloroplastic isoforms. Furthermore, APX-R was found as a new heme-containing protein functionally associated with ascorbate peroxidase (Lazzarotto et al., 2011). In this study, ScAPX6 was clustered in chloroplastic isoforms (Figure 3), which was consistent with the study conducted by Teixeira et al. (2004). Subcellular localization of ScAPX6 in rice protoplast showed that ScAPX6::GFP was targeted at chloroplast (Figure 4), which was consistent with the result of bioinformatics predicted localization. Similar to other plant species, such as the APXs from *Cucumis melo* (Cheng et al., 2009) and *A. andraeanum* (Liu et al., 2013), ScAPX6 also contained a plant peroxidase like superfAMILY and the heme binding site and
In this study, the expression of ScAPX6 was noted with NaCl and PEG treatment. Previous research on sugarcane showed that the transcripts of ScAPX6 were separated down regulated by 300 mmol L\(^{-1}\) H\(_2\)O\(_2\), SA, and NaCl treatments. Previous research on sugarcane showed that the transcripts of ScAPX6 increased under the treatment of ABA, MeJA, SA, H\(_2\)O\(_2\), PEG, and NaCl (Wang Z. Q. et al., 2015). In this study, the expression of ScAPX6 was also up-regulated by both ABA and MeJA, but down-regulated by SA and H\(_2\)O\(_2\) treatments (Figure 8A). As reported, the expression of TAPX gene in sugarcane was significantly induced by NaCl and PEG stresses (Wang S. et al., 2015). Previous investigations have identified the transcripts of OsAPX7 and OsAPX8, which were separately down regulated by 300 mmol L\(^{-1}\) NaCl in rice roots and leaves (Hong and Kao, 2007; Yamane et al., 2010). Similarly, in this study, down regulation of the transcripts of ScAPX6 were noted with NaCl and PEG treatment (Figure 8B), which was consistent with the results of the spot assay that the recombinant protein of ScAPX6 expressed in E. coli BL21 did not show better growth than the control under both two

**FIGURE 9** | Transient overexpression of ScAPX6 in Nicotiana benthamiana leaves. (A) RT-PCR analysis of ScAPX6 in the N. benthamiana leaves after 1 d infiltration by Agrobacterium strain GV3101 carrying pEarleyGate 203-ScAPX6 and the empty vector (3SS::00). (B) DAB (3,3′-diaminobenzidine) staining and trypan blue staining of N. benthamiana leaves at 48 h and 6 d after Agrobacterium strain infiltration, respectively, (1) represented a (Continued)
treatments (Figure 6). Previous reports have provided evidence that over-expressed plant stress tolerance genes in E. coli cells could enhance their growth under abiotic stress (Gupta et al., 2010; Guo et al., 2013). For example, Su et al. (2014b) have tested a chitinase gene ScChi in E. coli, which showed better growth under NaCl, Cu, CdCl₂ and ZnSO₄ treatments. Duan et al. (2006) have indicated that the transgenic O. sativa with HvAPX1 gene was more tolerant to cadmium stress when compared with the wild type. In this study, the transcript of ScAPX6 was also up-regulated by the treatment of Cu (Figure 8B), which was in line with the results that the recombinant protein of ScAPX6 expressed in E. coli BL21 resulted in a better growth under Cu stress (Figure 6). Therefore, it was predicted that ScAPX6 could be helpful for the tolerance of sugarcane to Cu. These findings suggested ScAPX6 might be a positive response to ABA, MeJA, and Cu stresses, while showed the negative response to SA, H₂O₂, PEG, and NaCl stresses. However, what should be pointed out here is that, we only use treatment with some hormones, such as SA, ABA, and MeJA with one concentration, and we cannot directly link effect of our treatment with specific hormone pathway because the specific concentration of the hormones was not determined. These points need to be considered in future.

Previous studies have revealed that the overexpression of tApx gene in tobacco enhanced tolerance to chilling, methylviologen, and high-intensity light (Yabuta et al., 2002). Transgenic potato with simultaneous overexpression of APX, choline oxidase (codA), and SOD, increased tolerance of SSAC plants and lower levels of H₂O₂ under methylviologen, drought and salt-mediated oxidative stresses (Ahmad et al., 2010). Investigations have revealed that cell death could induce R gene expression, ion fluxes, stimulation of ROS, and defense-related hormones, which can efficiently restrict pathogen growth and development (Li et al., 2010; Melech-Bonfil and Sessa, 2010; Du et al., 2012). Thordal-Christensen et al. (1997) proved that DAB-uptake method can serve H₂O₂ detection at a subcellular level. Although the DAB reactions reflect increases in local H₂O₂, this method has been adapted to many other plant species for in situ detection of H₂O₂. Thordal-Christensen et al. (1997) speculated that this could be related either to the fact that the optimal pH (5.5–6.0) for DAB precipitation coincides with the expected pH level in the leaf, or to the strong polymerization observed in the plant tissue. Lai et al. (2013) found that 35S::BrERF11 transgenic tobacco plants showed significantly increased HR and H₂O₂ accumulation compared with wild-type plants according to trypan blue and DAB staining. In the present study, a darker DAB staining color was found after overexpression of ScAPX6 in N. benthamiana leaves and after inoculation with P. solanacearum and F. solani var. coeruleum at 1 d compared with that in the leaves of control (Figures 9B,D,F), which was indicative of the accumulation of H₂O₂ and resulted in intense hypersensitivity response, but with elongated treatment time, the DAB staining color became lighter, which was consistent with the fact that APX can rapidly scavenge H₂O₂ in the plant by the ASA-GSH cycle (Shigeoka et al., 2002). However, since the DAB buffer is far from real physiological conditions, whether itself results in the accumulation of H₂O₂ remains an open question. Together, the antimicrobial action against the tobacco pathogens after overexpression of ScAPX6 in N. benthamiana, suggesting that ScAPX6 may enhance the resistance to P. solanacearum and F. solani var. coeruleum.

CONCLUSIONS

In this study, a novel ascorbate peroxidase gene, ScAPX6 (GenBank Acc. No. KT907352), was isolated and characterized. The cDNA of ScAPX6 gene was 1,086 bp long with a complete 1,002 bp ORF, encoding 333 amino acids. Subcellular localization revealed that ScAPX6 was targeted in chloroplast. After inducing by IPTG, the accumulation protein of pEZYHb-ScAPX6 at 55 KDa led to a better growth of E. coli BL21 under Cu stress. ScAPX6 was constitutively expressed in sugarcane tissues. Besides, ScAPX6 showed positive response to ABA, MeJA, and Cu stresses, but negative response to the stresses of SA, H₂O₂, PEG, and NaCl. The overexpression of ScAPX6 in N. benthamiana leaves showed positive response against the attack of P. solanacearum and F. solani var. coeruleum. These results suggested that ScAPX6 plays an important role in the HR or immunity of sugarcane.

AUTHOR CONTRIBUTIONS

FL, YS, and YQ: Conceived, designed, and initiated the project; NH, HL, and SG: Prepared materials; FL, NH, LW, TS, WA, and JG: Performed experiments and contributed to data analysis and validation; FL and YS: Drafted the manuscript; LX, KM, YS, and YQ: Revised the manuscript. All authors read and approved the final manuscript.

ACKNOWLEDGMENTS

This work was supported by Natural Science Foundation of Fujian province, China (2015J06006), the National Natural Science Foundation of China (31501363), the Research Funds for Distinguished Young Scientists in Fujian Agriculture and Forestry University (xjq201630), the Research Funds for Distinguished Young Scientists in Fujian Provincial Department of Education, the earmarked fund for China Agricultural Research System (CARS-17) and the Program for New Century Excellent Talents in Fujian Province University (JA14095).

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2017.02262/full#supplementary-material

Figure S1 | Amplification of ScAPX6 gene in sugarcane. M: DNA marker 2,000 bp; 1: RT-PCR product.
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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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