The Molecular Cloning and Expression Analysis of a CYP71 Gene in *Ginkgo biloba* L.

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Abstract

Cytochrome P450 monooxygenases (CYPs) are a group of redox proteins that catalyze various oxidative reactions in plant secondary metabolism. To explore the function of the CYP71 gene in *Ginkgo biloba* under biotic and abiotic stresses, a full-length CYP gene, designated GbCYP71, was first isolated and characterized from leaves of *G. biloba*. It contained a 1512-bp open reading frame (ORF) encoding 503 amino-acid-deduced polypeptide whose theoretical molecular weight was 56.9 kDa. The genomic DNA sequence of GbCYP71 contained two exons and one intron. The cDNA of GbCYP71 was subcloned in a pET-32a vector and then transformed into *E. coli* strain BL21 (DE3). A protein with a molecular weight of 76.4 kDa was subsequently identified and found to be consistent with the above theoretical value. Transient expression analysis revealed that the GbCYP71 protein may be located in the *G. biloba* cell cytoplasm. GbCYP71 was expressed in almost all ginkgo tissues, including leaves, stamens, gynoecia, stems and, preferentially, roots. Expression-profiling analyses revealed that GbCYP71 can be induced by salinity stress and phytohormone signals, including salicylic acid, abscisic acid, methyl jasmonate and ethephon, but is repressed by heat and cold stresses. These results indicate that GbCYP71 mainly functions in responding to biotic and abiotic stresses.

Keywords: abiotic stress, expression pattern, *Ginkgo biloba*, GbCYP71, phytohormone induction

Introduction

Cytochrome P450 monooxygenases (CYPs) form the largest enzyme family and present in various tissues (Werck-Reichhart *et al.*, 2000) in higher-order plants. The CYPs are a group of redox proteins that catalyze the oxidation steps of various substrates using oxygen and NAD(P)H (Guttikonda *et al.*, 2010; Isin and Güengerich, 2007). The first reported plant CYP gene, *CYP71A1*, was a ripening-related gene from the mesocarp of an avocado (Bozak *et al.*, 1990). Since its discovery, substantial works have been performed on cloning and characterizing a large array of CYPs. At present, more than 127 families and 5100 sequences of plant CYPs have been annotated and named (Nelson and Werck-Reichhart, 2011); however, the biological functions of most of these proteins in plants remain unknown. Plant CYPs have been shown to participate in many metabolic pathways, especially those related to plant secondary metabolism, which mainly involves the synthesis of endogenous compounds, such as structural macromolecules, pigments, and defense compounds; the synthesis or catabolism of all types of hormones or signaling molecules; and the metabolism of xenobiotics (Morant *et al.*, 2003; Werck-Reichhart *et al.*, 2002). CYPs catalyze many oxidation reactions, including epoxidations, hydroxylations, dealkylations, isomerizations, dehydrations, aryl migrations, decarboxylations, ring formations, ring extensions and carbon-carbon bond cleavages (Rupasinghe and Schuler, 2006). The ability of CYPs to catalyze such diverse reactions in plants contributes greatly to understanding the challenge posed by abiotic stresses and pathogens in nature.

An ancient gene superfamily, CYPs are classified into various families and subfamilies on the basis of the protein sequence identity and phylogeny (Nelson, 2006; Nelson *et al.*, 1996). CYPs with at least 40% similarity belong to the same family, and CYPs with at least 55% similarity belong to the same subfamily. CYPs with at least 97% similarity are assumed to be allelic variants unless otherwise demonstrated. Plant CYP families have been assigned names, including CYP51, CYP71 to CYP99 and CYP701 to CYP772, as depicted on the Cytochrome P450 webpage (http://drnelson.uthsc.edu/CytochromeP450.html). The nomenclature of a CYP gene is regulated the same way as the number following the CYP: the prefix is for the family, the letter is for the subfamily, and the number is for the particular gene. Based on evolutionary relationships in phylogenetic trees, the CYPs of land plants are grouped into 11 clans, and each clan is derived from a different gene ancestor (Nelson and Werck-Reichhart, 2011).

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Most CYP members contain approximately 500 amino acids and have a molecular weight of 45 kD to 65 kD (Donaldson and Luster, 1991; Schuler, 1996). Some conserved domains have been well identified in the primary CYP amino acid sequence; the most highly conserved CYP regions are the proline-rich region, PERF, K-helix, and I-helix regions and the heme-binding domains. Following the CYP N-terminal hydrophobic helix, the highly conserved proline-rich protein region is thought to act as a membrane hinge, usually obeying the consensus P/I-P-G-P-X/G/P-X-P (Yamazaki et al., 1993). The I-helix contains a conserved region A/G-X-G-X-D-T-T/S corresponding to the proton transfer groove on the distal side of the heme (Wercz-Reichhart and Feyereisen, 2000). There is a K-E-T-L-R consensus sequence with conserved Glu (E) and Arg (R) in the K-helix; its E and R form an E-R-R triad with R from P-E-R-F. The triad is considered to be involved in locking the heme pocket into position and assures heme association with the protein (Paquette et al., 2000). The heme-binding domain in the C-terminal region of the protein contains the consensus sequence F-X-X-G-X-D-E/T, which is the main feature in CYP identification. The conserved cysteine (C) of the heme-binding domain serves as the fifth ligand for the heme iron (Chapple, 1998). Only the E-R-R triad and the C in the heme-binding domain are conserved in all plant CYP sequences among these conserved domains (Wercz-Reichhart et al., 2002). In addition, plant CYPs vary in their subcellular locations, and most of them are endoplasmic reticulum (ER) binding proteins; the N-terminal hydrophobic helix anchors to the ER membrane, and most of the rest of the protein is located on the cytosolic face of the membrane (Chapple, 1998; Wercz-Reichhart and Feyereisen, 2000).

Ginkgo biloba is an ancient plant that originated in the Carboniferous period and thrived in the Jurassic period. It is the only remaining species of Ginkgoaceae and is often called a “living fossil” (Shen et al., 2006). G. biloba is one of the most popular functional plants used worldwide today, especially for medicinal purposes. Plants in nature frequently experience complicated climate changes and have to develop a broad range of complex defense systems to adapt. Geological records show that G. biloba exhibits minor changes in its morphology, indicating its strong adaptability in changing environments and tolerance of harsh conditions (Royer et al., 2003). In addition, Ginkgo is widely distributed around the world, thanks to characteristics such as resistance to disease, pests, drought, low temperatures, barren landscapes, and nuclear radiation (Cheng et al., 2013). To date, CYP genes have been reported in 66 species, including algae, planus, pteridophytes, needle-leaved plants, monocotyledons, and dicotyledons. However, the CYP gene has not been reported in G. biloba. In this study, we isolated the cDNA of a CYP gene from G. biloba and characterized the physical-chemical properties of its putative protein using bioinformatics tools. The expression patterns of the GbCYP were analyzed by real-time PCR. Additionally, GbCYP was subcloned into an expression vector and underwent heterologous expression in E. coli and N. benthamiana to further elucidate its functional characterization in G. biloba.

Materials and Methods

Plant materials and growth conditions

Different tissues, including leaves, stamens, gynoecia, stems and roots, were collected from several 16-year-old grafted Ginkgo biloba plants that were grown in the outdoor ginkgo garden of Nanjing Forestry University in Jiangsu Province, China. These tissues were immediately frozen in liquid nitrogen, transferred to an ultra-low-temperature freezer and kept at -80°C prior to total RNA extraction.

Mature zygotic embryos of G. biloba were incubated in an MS induction medium to initiate callus cultures; the MS basal medium was supplemented with 2 mg/L 6-benzyladenine (6-BA) and 1.5 mg/L naphthaleneacetic acid (NAA). After 4 weeks of incubation, the original seed embryos generated calluses, which were transferred to fresh MS induction media at 2-week intervals. The calluses, having a similar growth rate, were transferred to an MS liquid medium supplemented with 6-BA (2 mg/L) and NAA (1.5 mg/L) after two subcultures and then were cultured on a rotary shaker at 100 rpm, in the light, at 25 ± 1 °C. The suspension cultures were subcultured three times at 2-week intervals, and the calluses that appeared pale yellow-white, translucent, or browning were omitted.

The tobacco (Nicotiana benthamiana) used for the transient expression of fluorescent fusion proteins was approximately 4 to 5 weeks old and was planted in an artificial climate chamber at 21°C with 70% relative humidity, with 14 hours of light (500 μmol/m²/s) and 10 hours of dark.

Hormones and stress treatments

To investigate induction elicitors, the callus lines were treated with 100 μmol/L AsA, 100 μmol/L ascobic acid (ABA), 100 μmol/L methyl jasmonate (MeJA), 40 μmol/L ethephon (ETH) and 200 mmol/L sodium chloride (NaCl), separately, while the calluses without treatment were used as a control (CK) (Gong et al., 2006; Liao et al., 2015; Novo-Uzal et al., 2014). To explore the response to temperature stress, the callus lines were treated at 4°C and 40°C separately, while the control group was treated at 25°C. Samples were collected at 0, 3, 6, 12, 24 and 48 h after the beginning of treatments, immediately frozen in liquid nitrogen and stored at -80°C. Each sample at each time point included three biological replicates.

Molecular cloning

Total RNA was extracted from different tissues subjected to different treatments using the MiniBEST Plant RNA Extraction Kit (Takara, Japan), according to the manufacturer’s instructions. The RNA sample was subsequently treated with DNase I according to the extraction kit instructions to remove contaminating genomic DNA. Genomic DNA was extracted from fresh leaves following use of the Plant DNA Isolation Reagent (Takara, Japan). The quality of the RNA and genomic DNA was determined by agarose gel electrophoresis. The concentration was detected by UV spectrophotometry. Reverse transcription-polymerase chain reaction was performed using a Transcriptor First Stand cDNA Synthesis Kit (Roche, Germany).

The specific primers CYPU and CYPD (Table 1) were designed and synthesized (Shanghai Generay Biotech, China) based on the transcriptome sequencing. To obtain the cDNA
and genomic sequences of GbCYP71, a touchdown thermal cycle program was utilized under the following conditions: 94 °C for 4 min, 94 °C for 30 s, 58 °C for 30 s and 72 °C for 1 min 30 s, followed by 9 cycles at decreasing annealing temperatures in decrements of 0.5 °C per cycle, and then 25 cycles of 30 s at 94 °C, 30 s at 94 °C and 1 min 30 s at 72 °C, as well as a final extension at 72 °C for 10 min. The PCR products were purified and cloned into a pMD19-T vector (Takara, Japan) for sequencing.

Bioinformatics analysis and phylogenetic tree analysis

The obtained sequences were analyzed using online bioinformatics tools (http://www.ncbi.nlm.nih.gov and http://web.expasy.org/protparam/). The software Vector NTI Advance 11 was used for sequence multi-alignment. Sequences of homologous CYP71 proteins present in the NCBI database were retrieved using BLAST network services. The protein signal sequence was predicted using the SignalP 4.1 Server (http://www.cbs.dtu.dk/services/SignalP/), and the transmembrane domain was analyzed by the TMHMM server (http://www.cbs.dtu.dk/services/TMHMM/). Subcellular localization presumption was performed using TargetP 1.1 (http://www.cbs.dtu.dk/services/TargetP/). Phylogenetic and molecular evolutionary analyses of the amino acid sequences from other plant species were constructed with MEGA 6.06 software by the neighbor joining method and measured by bootstrap analysis with 1000 replicates.

RNA extraction and qPCR

A real-time quantitative PCR detection system (qPCR) was used to investigate the transcription profile of GbCYP71 in different tissues and in the callus samples collected at different time points after treatment. qPCR was performed using an ABI PRISM 7500 Sequence Detection System (Applied Biosystems, USA), using the SYBR Green PCR Master Mix (Roche, Germany) PCR system. PCR was performed with a 20 µL reaction consisting of 50 ng of template cDNA, 2×SYBR Green Mastermix, 4 µL of forward and reverse primers and 2.5 mM of each primer. The PCR product was separated by electrophoresis on 1% (w/v) agarose gels and then cloned into pET-32a, which was digested by BamH I at 37 °C for 4 h. The recombinant plasmid was introduced into E. coli BL21 (DE3) by the heat shock method. A single colony of E. coli BL21 (DE3) cells harboring the expression plasmid was grown in LB medium containing ampicillin (50 µg/mL) with 160 rpm shaking at 37 °C. For induction, IPTG was added to the E. coli culture at a final concentration of 1.0 mM when the optical density (OD₆₀₀) reached approximately 0.6. The cells continued growing for 6 h at 37 °C before being collected by centrifugation. The collected cells with pET32a-GbCYP71 or pET-32a plasmid were lysed by sonication in PBS buffer (20 mM, pH 7.4) and centrifuged at 12,000 ×g for 15 min at 4 °C. The pellets were re-suspended and then boiled for 5 min after adding 5×SDS loading buffer (4:1, v/v). SDS-PAGE analysis was used to examine 10 µg of each of the samples, followed by a staining procedure with Coomassie brilliant blue R250. Western blot analysis was performed to verify the expression of a GbCYP71 protein recombinating a His-tag in the N-terminus. After electrophoresis, the proteins were electro-transferred onto a PVDF membrane (0.45 µm) and detected with TRUE-Tag Anti-His mAb (Vazyme, China) and a secondary antibody, HRP-conjugated Goat Anti-Mouse IgG (H+L) (Vazyme, China). Enhanced ECL (Vazyme, China) was used in the color reaction.

Transient expression in N. benthamiana

For transient expression of GbCYP71 in tobacco, all plasmids were constructed using Gateway technology (Invitrogen, USA). To generate the entry vector, pENTR/D/TOPO vector (Invitrogen) was used to connect with the gene fragment encoding GbCYP71. The gene fragment was amplified from cDNA with the primer pair CYPS2/CYPA1 and was designed according to the guidelines of TOPO Cloning instruction (Table 1). The expression vectors (CYP71-GFP) for tobacco injection, including the promoter of 35S and enhanced green fluorescent protein (EGFP), were generated by LR reactions using the LR Clonase II enzyme (Invitrogen, USA) with the corresponding entry vector and the vector destination.

### Table 1. Primers for GbCYP71

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5’-3’)</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>CYPS1</td>
<td>GCCATGCTTATCAGGAAGTGCAGTGCA</td>
<td>Gene-specific primer, forward</td>
</tr>
<tr>
<td>CYPS2</td>
<td>CACCATGTTGAAGGAGCTGCA</td>
<td>Gene-specific primer, reverse</td>
</tr>
<tr>
<td>CYPA1</td>
<td>CTCTGGTGGGTCCTGACGCTTGTCTGAGTGGATGAGTCAATGCTGCTG</td>
<td>Primer for ORF subclone, forward</td>
</tr>
<tr>
<td>CYPA2</td>
<td>CCACTCTGAGGAGGAGCTGCA</td>
<td>Primer for ORF subclone, reverse</td>
</tr>
</tbody>
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pB7WG2.0 (Karimi et al., 2002). The reference vector (35S-RFP) for the expression indicator was constructed with 35S and red fluorescent protein (RFP) and was located in the cytoplasm. The plasmid encoding p19 protein was used as a viral silencing suppressor. Each constructed plasmid was introduced into Agrobacterium strain harboring the plasmid encoding GbCYP71, RFP and p19 protein was mixed in a 1:1:1 ratio before being transfected into the underside of tobacco leaves. The leaves were agroinfiltrated according to Sparkes et al., (2006). Three leaves per plant were infiltrated with the same constructs to yield experimental triplicates. After 3 to 4 days of infiltration (the period when fluorescence was optimal), fluorescent protein localization was monitored in the lower epidermal cells of the leaves using a fluorescence microscope (Leica DM500B).

### Statistical analysis

Statistical analysis was performed by one-way analyses of variance (ANOVA), and mean differences were compared by Tukey’s multiple range test using SPSS 17.0.

### Results

**Characterization of GbCYP71 in Ginkgo biloba**

The full-length cDNA sequence of the GbCYP71 gene was obtained from *G. biloba*. It is a 1568-bp sequence containing a 1512-bp open reading frame (ORF). A 1630-bp genomic DNA sequence of the GbCYP71 gene was amplified by a pair of specific primers derived from the cDNA sequence containing the start and stop codon region. Comparison with the full-length cDNA sequence showed that the genomic DNA and cDNA were 100% identical in the coding region, although the genomic DNA contained an intron. Exon 1 (894 bp) and exon 2 (618 bp) were separated by intron 1 (118 bp). The putative splicing site obeyed the GU/AG rule (Fig. 1).

The putative GbCYP71 protein contains 503 amino acids. Analysis revealed that the isolectric point (pI) and molecular weight of this peptide were 6.88 and 56.9 kDa, respectively (ProtParam tool). The putative formula of GbCYP71 was C_{259}H_{489}O_{25}N_{30}S_{50}. The instability index (II) was computed to be 39.70 (which classified the protein as stable), and its grand average of hydropathicity (GRAVY) was predicted to be -0.165. SignalP 4.1 analysis showed that GbCYP71 had a signal peptide sequence with a cleavage site between Gly and Ser. Transmembrane analysis by the TMHMM Server v. 2.0 showed that GbCYP71 had one transmembrane domain located between Gly 3 and Phe 21 along the polypeptide chain. The TargetP 1.1 Server predicted that the subcellular localization of GbCYP71 was in the secretory pathway.

As shown in Fig. 1, the deduced amino acid sequences of this gene showed the typical features of membrane-bound P450s. Alignment by NCBI’s Blastx tool showed that GbCYP71 shared moderate homology with CYPs from *Picea sitchensis* (59%, ABR18406.1), *Cucumis melo* (56%, XP_004406681.1), *Vitis arizonica* × *Vitis rupestris* (55%, ACM98788.1), *Cicer arietinum* (54%, XP_004504031.1) and *Populus euphratica* (53%, XP_011025622.1). However, the functions of the above proteins have not been reported. Nevertheless, this predicted protein has high homology (40-49%) with known CYP71 proteins of other higher-order plants, which indicates that the protein encoded by the cDNA obtained in this study may belong to the CYP71 protein family.

**Molecular evolution analysis**

A phylogenetic tree was constructed using the neighbor-joining method by MEGA 6.06 software to investigate the evolutionary relationships between GbCYP71 and other CYPs from different species. An unrooted neighbor-joining cladogram is shown in Fig. 2; the numbers at the nodes show bootstrap support values (1000 replicates), and the scale line below the tree represents relative evolutionary distance. In this phylogenetic tree, the relationships between CYPs from analyzed plants
generally reflect standard relationships pertaining to gymnosperms, pteridophytes, monocotyledons and dicotyledons. GbCYP71 belongs to the gymnosperm group and has a relatively close relationship with the conifer *Picea sitchensis*.

Expression of recombinant GbCYP71 in *E. coli*

GbCYP71 was expressed in *E. coli* by the pET32a expression vector. SDS-PAGE showed that the molecular weight of the expressed recombinant protein was estimated as 76.4 kDa, which is the sum of the target gene (56.9 kDa) and a histidine marker (approximately 20 kDa) (Fig. 3 a), and was consistent with the predicted value. Western blotting of purified recombinant GbCYP71 protein confirmed its specific immune reactivity to anti-His antibodies (Fig. 3 b).

Subcellular localization of GbCYP71 Proteins in *N. benthamiana*

To investigate the biological activity of the GbCYP71 protein, its subcellular localization was analyzed using the transient expression system in *N. benthamiana*. As expected, 35S-RFP showed cytosolic localization when expressed in an *N. benthamiana* epidermal cell (Fig. 4 b). The recombinant fusion protein CYP71-GFP showed similar accumulation in the cell, which indicated that the GbCYP71 protein was located in the cytoplasm (Fig. 4).

Expression of GbCYP71 in different tissues

To explore the expression pattern of GbCYP71 in *G. biloba*, qPCR was performed using the primers CYP-F and CYP-R. As shown in Fig 5, the expression of GbCYP71 could be detected in all tissues, but at varying levels. GbCYP71 was preferentially expressed in roots and leaves, and the expression level in roots was significantly higher (P<0.05) than that in leaves and in other tissues. Subsequently, the expression level of GbCYP71 in leaves was significantly higher (P<0.05) than that in stems. The lowest level of GbCYP71 was in the stamens.

Expression of GbCYP71 under salt and different temperature stresses

To determine if GbCYP71 was involved in the responses to abiotic stresses, the gene’s transcript levels were measured by qPCR after treating the callus lines with high salinity, cold (4 °C) and heat (40 °C) stress. GbCYP71 expression was significantly affected by NaCl and extreme temperature (Fig. 6). The expression level of GbCYP71 increased quickly after NaCl...
treatment and reached a peak (10.3-fold) at 6 h. Then, it decreased sharply to a basal level (0.63-fold) at 12 h, followed by a slow decline with a 12 h duration, and finally maintained a 0.33-fold expression level relative to the control until 48 h (Fig. 6 b). Compared with room temperature, GbCYP71 expression was suppressed under extreme temperatures (Fig. 6 a, c and d). Under high temperature (40 °C), GbCYP71 expression decreased quickly and finally dropped to 0.17-fold that of the control (0 h) at 24 h (Fig. 6 c). However, at low temperature (4 °C), GbCYP71 expression decreased to a minimum of 0.42-fold of the control after 3 hours of treatment and then increased slightly and recovered to the same level as the control (P<0.01) at 12-48 h (Fig. 6 d).

Expression of GbCYP71 under hormone treatments

Expression profiles for GbCYP71 at different stages of SA, ABA, ETH and MeJA treatment were determined in ginkgo calluses to assess the possible involvement of GbCYP71 in signaling pathways used by phytohormones. GbCYP71 transcript levels increased significantly with 100 µmol/L SA, reached an expression peak (8.64-fold) at 24 h, and then decreased sharply to their basal level (Fig. 7 a). GbCYP71 transcript levels were also stimulated by 100 µmol/L ABA; the strongest response to ABA (20.7-fold induction) was observed at 12 h (Fig. 7 b). Afterward, GbCYP71 transcript levels decreased as treatment time increased, reaching a 2.95-fold difference from the control at 48 h. GbCYP71 transcription levels decreased slightly at 3 h after 40 µmol/L ETH treatment, then increased significantly between 3 and 6 h, and finally decreased until 48 h (Fig. 7 c). In response to 100 µmol/L MeJA, GbCYP71 transcript levels were elevated significantly between 0 and 6 h post treatment and reached an expression peak (8.76-fold) at 3 h, after which they decreased to their basal level (Fig. 7 d).

Discussion

CYPs constitute a superfamily of hemoproteins with diverse functions in plants. They have a systematic nomenclature based on their amino acid sequence identity. Due to limitations in purifying CYP from higher-order plant sources, studies of CYPs in higher plants have lagged behind those in animals and microorganisms (Toguri and Tokugawa, 1994). Members of the same family encode homologous proteins and have similar structures and properties. Similarly, it is possible for genes from different families to participate in similar pathways. In this study, the deduced protein sequence of the CYP71 gene in G. biloba contains the classically conserved domains of plant CYP proteins, such as the proline-rich protein region, the C-helix involved in oxygen binding, the E-R-R triad, and the cysteine in the heme-binding region. Blastx also indicated that GbCYP71 belonged to the CYP71 family. The CYP phylogenetic tree shows that GbCYP71 was in the gymnosperm cluster and diverged earlier than CYPs from other plants. Some studies have been performed to investigate whether CYP genes (CYP73A1 through CYP73A41) from different species were generally encoding cinnamate 4-hydroxylase (Ehlting et al., 2006). In another study, low amino acid identity (13%) was noted among four CYPs (CYP73A45, CYP75B1, CYP84A1, CYP98A3); similar mechanisms for substrate binding were predicted as they mediate the phenylpropanoid pathway of Arabidopsis (Rupasinghe et al., 2003). By comparatively analyzing the three CYP crystal structures, Hasemann et al. (1995) suggested that although the primary sequences of CYPs were highly diverse, their folding and architecture were remarkably conserved throughout evolution. CYP71 forms the largest CYP family in plants with various functions, of which most are related to plant
secondary metabolism (Gierl, 2009; Nelson and Werck-Reichhart, 2011). CYPs in plants appear to be differentially expressed in various tissues and organs (Guttikonda et al., 2010; Yamaguchi et al., 2014). In this study, the expression level of GbCYP71 varied greatly in different tissues; the expression level was highest in the root and lowest in the male flower. Because plants are sessile organisms, they are directly exposed to common adverse environmental factors that are known to induce oxidative stress, such as salinity, drought and low temperature (Bohnert et al., 1995; Mittova et al., 2004). Elucidating the stress perception, signaling, and response mechanisms of plants is crucial to understanding how they improve their stress tolerance in environmentally demanding conditions. In this study, we found that GbCYP71 expression was significantly induced under salt stress and decreased under extreme temperature stress (4 °C, 40 °C). Our results suggest that GbCYP71 is highly expressed in the root, which may strengthen ginkgo’s resistance to salt stress.

Additionally, GbCYP71 expression was not induced by high or low temperature; its suppressed expression may be due to the slow growth of callus, which was caused by extreme temperature. Salinity stress is one of the most severe factors limiting plant productivity in abiotic stress; high salt conditions can affect the entire physiological process from seed germination to plant development. Salinity causes osmotic stress in plants in the root zone through decreases in the osmotic potential of surrounding soil (Pasternak, 1987). Temperature stresses are another common factor that can restrict plant growth and development.

ABA, SA and MeJA are hormones in higher-order plants that play important roles in activating plant defense mechanisms in response to biotic and abiotic stresses, such as drought, low temperature, salinity and pathogen attack. Furthermore, ABA, SA and MeJA are important cellular regulators participating in various developmental processes, such as embryo and seed development, root growth, seedling establishment and vegetative development (Cutler et al., 2010; Vlot et al., 2009; Wasternack and Hause, 2002; Wasternack and Parther, 1997). Osmotic stress elicited by salinity or drought is a very important environmental constraint for land plant survival and distribution. Phytohormones ABA, SA and MeJA can increase in response to osmotic stress. Genes responsive to osmotic stress can also be induced by these exogenous hormones (Borsani et al., 2001; Ishitani et al., 1997; Xu et al., 1994). In this study, GbCYP71 was induced by all exogenous hormones. ABA, SA and MeJA elicited similar significant increases in gene expression. These results revealed that the GbCYP71 gene may play a role in resistance to biotic and abiotic stresses, as well as in osmotic regulation.

At present, because an efficient ginkgo regeneration system has not been established with explants, it is difficult to study ginkgo gene functions except via heterologous expression. New technologies could be applied in this field, such as the plant protoplast system, which has been widely used in popular research (Lin et al., 2013). In this study, the GbCYP71 protein was expressed successfully in transgenic strains. Additional studies could purify recombinant proteins from E. coli cells and analyze their enzyme activity. Further analysis of the GbCYP71 gene and its promoter sequence is required to study the signaling pathways involved in biotic and abiotic stress regulation and transgenic technology should be applied to study the functions of these cloned genes.

Conclusions

This work characterized the molecular structure of the CYP71 gene from Ginkgo biloba and analyzed its expression pattern under various treatments. Bioinformatics analyses demonstrated that GbCYP71 is a membrane-bound P450s protein with five classically conservative domains. Expression pattern analysis showed that GbCYP71 is preferentially expressed in roots and that its expression is induced by salinity stress and phytohormones but repressed by cold and heat stresses. This study revealed that GbCYP71 may participate in plant defense responses to abiotic stress.

Acknowledgments

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References


Gong YF, Zao ZH, Guo BH, Sun XF, Tang KX (2006). Molecular cloning and expression profile analysis of Ginkgo biloba DKS gene encoding 1-deoxy-D-xylulose 5-phosphate synthase, the first committed enzyme of...


