PgCYP76B93 docks on phenylurea herbicides and its expression enhances chlorotoluron tolerance in *Arabidopsis*

Jinhoon Jang1†, Sanjida Khanom1†, Youngkook Moon2†, Sooim Shin2 and Ok Ran Lee1*  

**Abstract**  
The phenylurea herbicides are used to control annual and perennial weeds on crop cultivating fields. The excessive usage of these agrochemicals increase many environmental problems. Thus, engineering transgenic plant for herbicide metabolism can provide efficient and eco-friendly means for enhanced phytoremediation capacity. Cytochrome P450 enzymes comprise one of the major plant enzyme families that mediate the oxidative degradation of xenobiotic chemicals, including herbicides. Considering these notions, phytoremediation properties of transgenic ginseng-derived *PgCYP76B93* in *Arabidopsis* were assessed. Phylogenetic tree of *PgCYP76B93* clustered in between close to the herbicide metabolism-related enzyme families and terpenoid biosynthesis-related. The expression of *PgCYP76B93* was considerably upregulated upon treatment with phenylurea herbicide, chlorotoluron. Simulated docking using Autodock program predicted possible interaction with chlorotoluron. Transgenic *Arabidopsis* plants overexpressing *PgCYP76B93* were resulted in slightly reduced plant height with relatively small leaves. The lower plant height in the *PgCYP76B93*-overexpressing line than in the control revealed that it was linked to the expression of gibberellin oxidases (GAOx). The bioassay of transgenic plants growing on herbicide-containing media revealed enhanced resistance against chlorotoluron.  

**Keywords:** *Panax ginseng*, Cytochrome P450, Herbicide, Chlorotoluron, Phytoremediation

**Introduction**  
Plant cytochrome P450 enzymes (CYPs) are involved in catalyzing various types of monooxygenation/hydroxylation reactions in primary and secondary metabolism [1]. One third of all known CYP enzymes are from the plant kingdom, and the number of enzymes is estimated to be up to 1% of the total number of annotated plant species. This implies that diversification within CYPs has led to the emergence of new metabolic pathways throughout land plant evolution.

A newly diversified species-specific secondary metabolic pathway in *Panax ginseng* Meyer involves ginseng saponins, also called ginsenosides [2, 3]. Considering the pharmaceutical efficacies of ginsenosides, studies on ginseng plants largely focused on ginsenoside biosynthesis [3, 4]. Among the 116 reported *P450* genes [5], two CYP genes have been reported to be involved in the dammarane-type of ginsenoside biosynthesis [6, 7] and one CYP, called β-amyrin 28-oxidase, is involved in the oleanane-type of ginsenoside biosynthesis [8]. However, besides their physiological roles in secondary metabolism, other important features of CYPs are their detoxification roles against herbicide [9]. The mechanism of acquiring increased herbicide metabolism has not yet been adequately demonstrated. A perennial plant, such as ginseng, has been grown commercially for up to 6 years or even more, and it can be exposed to herbicides over...
several years. Continuous exposure to herbicides might be involved in the diversification of more CYP family genes. Thus, taking the concept one step further, the functional characterization of other CYP family members from ginseng might shed light on herbicide metabolism. Modern herbicides contributed a lot to sustainable global food production by minimizing time to remove weeds and substituting destructive soil cultivation [10]. Thus understanding herbicides resistance mechanism is a major challenge for modern agriculture [10]. In higher plants, the herbicide detoxification genes have accumulated through the evolution of multigene families, like glutathione S-transferase [11] or glycosyl transferase [12] and cytochrome P450 enzyme families [9]. The CYPs are by far the largest family of enzymes that mediate the oxidative degradation of xenobiatic chemicals, including herbicides. The CYPs are heme protein that gains electrons from NADPH to catalyze the formation of a molecule of water and an oxygenated product. Xenobiotics such as aromatics, pesticides, and hydrocarbons are usually synthesized for industrial and agricultural purposes. Since some of them are harmful to living organisms, organisms have evolved efficient system such as CYPs to eliminate xenobiatic absorption. Genetically engineered plants for herbicide metabolism provide efficient and eco-friendly means for enhancement of detoxification of harmful substances [13].

The first evidence of the involvement of plant P450 in the metabolism of the herbicide monuron was reported in the microsomal fraction of cotton seedlings by Frear in 1969 [14]. To date, several CYP family members have been characterized as being involved in herbicide metabolism. CYP76B1 was isolated from Jerusalem artichoke (Helianthus tuberosus) and was found to metabolize herbicides belonging to the phenylurea class [15–17]. CYP71A10 was isolated from soybean and determined to metabolize the phenylurea herbicide, chlortoluron, when expressed in yeast, although with an efficiency 10 times lower than that of CYP76B1 [18]. The CYP76C1, CYP76C2, and CYP76C4 genes also provided suitable evidence for herbicide resistance in Arabidopsis [17]. When human-derived CYP1A2 was overexpressed in Arabidopsis, it also increased tolerance to the phenoylea herbicide linuron [19]. Very recently ginseng-derived CYP736A12 was reported to be involved in chlortoluron and isoproturon tolerance when overexpressed in Arabidopsis [20].

In this study, we report another ginseng-derived CYP gene annotated as PgCYP76B93 that are involved in herbicide tolerance. Homology based molecular docking of PgCYP76B93 shows the more possible interaction with chlortoluron rather than isoproturon. Transgenic plants overexpressing PgCYP76B93 displayed relatively reduced plant height and herbicide chlortoluron tolerance with respect to control lines. The observed functions of PgCYP76B93 provide insight into its use as a selectable marker for genetic engineering of multifunctional crops having the ability to detoxify agrochemicals and environmental contaminants.

Materials and methods
Plant materials and growth conditions
Panax ginseng Meyer ‘Chun-Poong’ was used as the plant material in this study. Arabidopsis thaliana (genotype Col-0) was used as a heterologous system. Mature Arabidopsis seeds were surface-sterilized for 5 min in 70% (v/v) ethanol and then rinsed three times with sterile water. The Arabidopsis seeds were sown on 1/2 MS medium (Duchefa Biochemie, Haarlem, Netherlands) containing 0.5 g/L MES (2-[N-morpholino] ethanesulfonic acid, 1% sucrose, and 0.8% phytoagar); the pH was adjusted to 5.7 using KOH. The seed-sown Petri plates were cold-treated for 2 days and then incubated under long-day photoperiod conditions—16 h light and 8 h dark—at 23°C in a growth chamber.

Sequence analysis
The ginseng EST clones were analyzed by using the BLAST (Basic Local Alignment Search Tool) program against a specialized cytochrome P450 database (https://dnelson.utmem.edu/CytochromeP450.html). The amino acid sequences were analyzed using the online programs (https://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi) and the ProtParam tool (https://web.expasy.org/protparam/). The amino acid sequences were aligned using the BioEdit program (version 7.1.9). A phylogenetic tree was generated according to the neighbor-joining method of the MEGA6 (version 6.06) program.

Chemical treatments in ginseng
To record gene expression levels with the treatment of herbicides, 10 μM of chlorotoluron (CAS Registry No. 15545-48-9, Cayman, USA) and isoproturon (CAS Registry No. 34123-59-6, Tokyo Chemical Industry, Japan) were treated to 4 weeks cultivated adventitious ginseng roots. The treated samples and their corresponding control were collected at different time intervals (0 h, 1 h, 4 h, 8 h, 12 h, 24 h, and 48 h) for qPCR.

RNA isolation and quantitative real-time PCR (qPCR)
Total RNA was isolated using the RNeasy Plant Mini Kit (Qiagen, Valencia, CA, USA) according to the manufacturer’s instructions with modifications. The DNase I (Takara, Japan) treatment was additionally performed before the washing step in a 100 μL reaction volume containing 1 μL of RNase inhibitor for 1 h. The concentration
of the extracted RNA was determined using the Nano- 
MD UV–Vis spectrophotometer (Scinco, Seoul, Korea). To synthesize the first-strand of cDNA, total RNA (up to 
5 μg maximum) was reverse transcribed using the Rever-
taID reverse transcriptase (Thermo, USA). A qPCR was 
performed using the Thermal Cycle Dice real-time PCR 
system (Takara, Shiga, Japan) according to the manufac-
turer’s instructions. The qPCR was performed in a 20 μL 
reaction volume using the following thermal cycling con-
ditions: initial denaturation at 95 °C for 30 s, followed 
by 40 cycles of 95 °C for 5 s and 60 °C for 30 s, an additional 
cycle of 95 °C for 15 s and 60 °C for 30 s, and a final dis-
sociation step at 95 °C for 15 s. At the end of the qPCR, a 
dissociation curve was generated to evaluate the gen-
eration of by-products. To determine the absolute and/or 
relative fold-differences in template abundance for each 
sample, the threshold cycle (Ct) value of each sample was 
normalized to that of β-actin, and the formula $2^{-\Delta\Delta Ct}$ 
(relative quantification) was used to calculate the gene 
expression levels. At least three independent experi-
ments were performed in order to validate the ΔΔCt method. Dissociation analysis at the end of each run was 
performed to confirm the specificity of the reaction. The gene-specific primers for $Pg$CYP76B93 were 5′-CAC 
AAG TCC CTG TCC ACC-3′ (forward) and 5′-GTG 
GTT GTG GAC GTG GAG-3′ (reverse). The control primers for ginseng β-actin (DC03005B05) were 5′-AGA 
GAT TCC GCT TGC AA-3′ (forward) and 5′-ATC 
AGC GAT ACC AGG GAA CA-3′ (reverse). Other primer sequences that were used are listed in Additional 
file 1: Table S1.

**Molecular docking study of $Pg$CYP76B93 with isoproturon and chlorotoluron**

The structure of CYP76B93 was predicted by ExPASy-
SWISS MODEL using Zebra Fish CYP-450 17A1 mutant 
Abitraterone complex (PDB code: 6B82) [21] as major 
template which exhibited 48% sequence similarity, 
because the crystal structure of CYP76B93 have not been 
defined yet. Molecules that are used to identify interac-
tion with CYP76B93 are isoproturon (CID: 36679) and 
chlorotoluron (CID: 27375) retrieved from the PubChem 
database. These molecules were exported as the standard 
3-dimentional XML format using the Open Babel 2.4.1. 
Docking of the two small molecules with CYP76B93 was 
processed by the AutoDock vina (The Scripps Research 
Institute, La Jolla, CA, USA) and set as 20 docking poses 
in the order of the highest negative ΔG. The flexibility of 
the ligands was kept as possible and placed at the inter-
face region. The binding possibility of isoproturon and 
chlorotoluron with CYP76B93 were examined accord-
ing to their binding energy, and then specific residues 
involved in interaction with CYP76B93 were determined 
by LigPlot+ [22] (The European Bioinformatics Institute).

**Vector map construction and in-planta transformation**

The full-length cDNA (1551 bp) of $Pg$CYP76B93 was 
amplified to functionally characterize in the Arabidop-
sis heterologous system. The in-planta gene transfer 
was carried out by using the constitutive 35S promoter-
driven pCAMBIA1390 [23]. $Pg$CYP76B93 was amplified 
using the following primers containing added restriction 
enzyme sites (SalI and MfeI): 5′-AA GTC GAC ATG GAT 
ATC TTA ACC ATG-3′ and 5′-CG CAA TTG AAT CAT 
AAC TGG AGT TG-3′. The purified PCR product was 
inserted into pCAMBIA1390 by performing overnight 
ligation at 16 °C. The $Pg$CYP76C9-ligated construct was 
confirmed by nucleotide sequencing and transformed 
into Arabidopsis using Agrobacterium tumefaciens strain 
C58C1 (pMP90) [24]. The transgenic transformants were 
selected on hygromycin-containing plates (50 μg/mL), 
and more than 15 T1 independent lines were selected. 
The homozygous transgenic lines carrying one copy of 
the insert and following a Mendelian segregation ratio 
were further characterized. For further data analysis, the 
Col-0 and empty vector lines were used as controls for the $Pg$CYP76B93-overexpressing lines.

**Phenylurea herbicide resistance test**

Surface-sterilized seeds were sown on the 1/2 MS media 
containing two phenylurea herbicide, chlorotoluron. After 2 days of cold stratification at 4 °C, the seedlings 
were grown at 23 °C under long-day photoperiod conditions 
of 16 h light/8 h dark. Chlorotoluron was added to 
the medium at different concentrations for preliminary 
tests, and a physiological concentration was chosen to 
show a clear-cut difference in tolerance.

**Statistical analysis**

Significance of the statistical analysis was measured 
based of the determination of Student’s t-test using MS 
excel software.

**Results and discussion**

**Isolation and identification of ginseng $Pg$CYP76B93 genes**

All cytochrome P450 (CYP) genes homologous with 
known CYP family genes were identified from the 
expressed sequence tag (EST) libraries constructed from 
the embryogenic calli, 4- and 14-year-old roots, leaves, 
bud, and methyl Jasmonic acid (MeJA)-treated adventi-
tious roots [25]. One EST clone showing similarity with 
CYP76B family was further chosen, and the recovery of 
its full-length cDNA was confirmed by rapid amplifi-
cation of cDNA ends (RACE) polymerase chain reac-
tion (PCR) [26]. The full-length cDNA of $CYP76B$ was
Fig. 1 Ginseng-derived cytochrome P450 (CYP) proteins are closely related to other CYP family proteins. a Phylogenetic tree of the PgCYP76B93 (marked in red box) protein with other closely related CYP proteins from Arabidopsis and several other crops. The phylogenetic tree was constructed using the ClustalX program (neighbor-joining method). At, Arabidopsis thaliana; Sm, Swertia mussotii; Panax ginseng; Cr, Catharanthus roseus; Pg, Ht, Helianthus tuberosus. The gene Accession numbers are: SmCYP76B10 (GU168041), CrCYP76B6 (AJ251269), HtCYP76B1 (Y10098), AtCYP76C7 (AT3G61040), AtCYP76C4 (AT2G45550), AtCYP76C1 (AT2G45560), HsCYP1A2 (GI73915100), PgCYP716A47 (JN604537), and PgCYP716A53v2 (JX036031). The bar represents 0.1 substitution per amino acid position. b Sequence alignment of PgCYP76B93 with other closest homologs. Black boxes indicate identical residues, and similar residues are shaded in grey. The four red box motifs represent the oxygen-binding domain (A/G) GX(D/E)T(T/S), K-helix region (ExxR), domain C (PER), and heme-binding region (FxxGxRxCG). Gaps were inserted to maximize homology.
annotated again as **PgCYP76B93** (CYP76B93 from *P. ginseng*) after consulting with the CYP nomenclature committee [27]. The original EST clone coding for **PgCYP76B93** was obtained from MeJA-treated adventitious roots, indicating that it can be induced by an elicitor and is expressed weakly in normal conditions. **PgCYP76B93** is a 1551 bp long gene encoding 517 amino acids. Other amino acid sequences of the characterized CYPs in plants were collected from various gene banks and were used to generate a phylogenetic tree (Fig. 1a). **PgCYP76B93** was clustered in between the CYP76B family enzymes involved in terpenoid biosynthesis [28, 29] and several CYP76B and CYP76C family enzymes [15, 16, 18], which were previously reported to be involved in phenylurea herbicide resistance (Fig. 1a). The enzymes belonging to the A-type subfamily of the CYP76 family enzymes were clustered relatively closer with **PgCYP76B93**, and three ginseng CYP enzymes involved in ginsenoside biosynthesis [6–8] were clustered far more distantly (Fig. 1a). This indicates that **PgCYP76B93** likely plays a role in herbicide resistance and/or terpenoid biosynthesis.

The **PgCYP76B93** enzyme was predicted to have a molecular weight of 58,390 kDa and a pl of 7.74 using the ProtParam program [30]. The CYP family enzymes were reported to share low similarity in their amino acid sequences, except for several conserved domains, which are necessary for their tertiary structure and enzymatic function [31]. The most well-conserved motif is the heme-binding region FxxGxRxRxG (also known as the CxG motif), in which C (cysteine) binds to the heme group. The ExxR and PER motifs form the E–R–R triad and play important roles in locking the structure of the heme pocket. The most conserved amino acids in the CYP family are glutamic acid (E) and arginine (R) in the ExxR motif and cysteine in the CxG motif, and these amino acids are also found in **PgCYP76B93** (Fig. 1b). The least conserved motif, (A/G)Gx(D/E)T(T/S) (also known as AGxDTT), contributes to oxygen binding and activation. The AGxDTT motif is also well conserved in **PgCYP76B93**, but the PER motif is replaced by the PKR motif (Fig. 1b). Understanding the significance of the substitution of K (lysine) for E requires further investigation.

Transcript levels of **PgCYP76B93** are gradually increased by chlorotoluron

In one step to understand the function of **PgCYP76B93**, modulation of transcripts against two phenylurea herbicides were evaluated according to the recent report [20]. Differential mRNA levels of **PgCYP76B93** was quantified in time-dependent manner by treatment of 10 μM of chlorotoluron and isoproturon (Fig. 2). The transcripts of

![Chlorotoluron](image1)

![Isoproturon](image2)
Fig. 3: Isoproturon and chlorotoluron docking near heme site of PgCYP76B93. **a** Five poses of isoproturon in green, and **b** six poses of chlorotoluron in blue are focused from all interactions on predicted PgCYP76B93 overlaying with Zebra Fish CYP-450 17A1 mutant Abiraterone complex. Heme is colored in red stick. Abiraterone is marked in light brown stick. **c, d** Residues of simulated PgCYP76B93 were expected to interact with isoproturon and chlorotoluron are presented by using LigPlot+ program. Hydrophobic interaction is indicated in spoked arcs labelling with residue, and hydrogen bond is marked in green dotted line with distance (Å) and residue. Distances from predicted heme of PgCYP76B93 to isoproturon and chlorotoluron are shown in **(e)**. All figures were produced using PyMol (https://pymol.org/2/).
Fig. 4  Heterologous overexpression of \textit{PgCYP76B93} reduced plant height and leaf size. \textbf{a} Transcript levels of \textit{PgCYP76B93} from 4-week-old leaves in control (Col-0 and empty vector control) and overexpression lines. Data represent the mean ± SE of three independent replicates at \( P < 0.05^{(*)} \) and \( P < 0.01^{(**)} \) by Student’s t-test. \textbf{b} Fully grown transgenic plants are smaller than the control plants. Scale bar = 2 cm. \textbf{c} Overall leaf size is lower in the overexpression lines than in the control lines. The leaves are arranged from cotyledons to the youngest leaves from left to right side. Scale bar = 1 cm. \textbf{d} Four-day-old seedlings display no significant difference in root length. Scale bar = 5 mm. \textbf{e} Statistical data displays the primary root length of each line from 4- and 7-day-old seedlings, \( n = 13–27 \).
*PgCYP76B93* were increased by chlorotoluron gradually from 1 to 24 h after treatments, and were initially upregulated by isoproturon up to 4 h (Fig. 2). It indicates that *PgCYP76B93* might play roles in herbicide detoxification, especially chlorotoluron.

**Molecular docking study shows putative phenylurea herbicide interaction with PgCYP76B93**

The major secondary structure of PgCYP76B93 is pretty identical to Zebra Fish CYP-450 17A1 mutant Abiraterone complex [21]. The heme site of simulated PgCYP76B93 was not appeared but it could be predicted by superimposing the simulated PgCYP76B93 on the Zebra Fish CYP-450 17A1 Abiraterone complex (Additional file 2: Fig. S1). When isoproturon and chlorotoluron were added to the AutoDock program to test the binding possibility with PgCYP76B93, possible docking simulation for the interaction was observed. Total 5 of 20 docking poses for isoproturon and 6 of 20 docking poses for chlorotoluron are resided around heme in active site (Fig. 3a, b). These positions are interestingly located on the similar site of abiraterone which is a substrate of the template protein, Zebra Fish CYP-450 17A1. Residues interacting with isoproturon and chlorotoluron around heme of CYP76B93 was identified by Ligplot program [22] (Fig. 3). Isoproturon dominantly forms hydrophobic interaction with Arg 102, Val119, Trp120, Met214, Asp315, Val318, Ala319, Thr323, Pro383, Ala384, Leu387, Ile388, Arg456, Pro459, Gly460 and Ile500 in 5 poses, and minor hydrogen bond with Ile388 and Arg390 in 1 poses near heme (Fig. 3c). Chlorotoluron majorly contacts with Val119, Trp120, Asp315, Val318, Ala319, Asp322, Thr323, Pro383, Ala384, Leu387, Ile388, Arg390, Arg456, Ile500 and Thr501 in 6 poses by hydrophobic interaction and minorly interacts with Ile388, Arg390 and Thr501 by hydrogen bond in 3 poses (Fig. 3d). Val119, Trp120, Val318, Ala319, Thr323, Thr384, Leu387, Ile388, Arg390, Arg456, Ile500 involved in interaction with isoproturon or chlorotoluron are frequently observed in each poses, so it can be further considered as a target for mutagenesis study. The closest distance from heme to isoproturon is 9.9 Å and to chlorotoluron is 4.3 Å (Fig. 3e). This result is believed to support the data that transgenic *Arabidopsis* possessing overexpression of CYP76B93 gene exhibits resistance against chlorotoluron other than isoproturon. Closely bound chlorotoluron to heme of PgCYP76B93 might be used as a substrate for undergoing detoxification mechanism.

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**Fig. 5** Gene expression patterns of gibberellin oxidases resulting from the overexpression of *PgCYP76B93*. Relative gene expression of four gibberellin oxidases (*GA2ox1*, *GA2ox2*, *GA3ox1*, and *GA20ox1*) in 14-a and 28-day-old seedlings b compared to Col-0. Data represent the mean ± standard error (SE) of three independent replicates at \( P < 0.05(*) \text{ and } P < 0.01(**) \) by Student’s t-test.
Overexpression of \( \text{PgCYP76B93} \) reduced plant height and leaf size

In order to understand the functional characteristics of \( \text{PgCYP76B93} \) in plant growth and development, its full-length coding sequence was overexpressed in heterologous \( \text{Arabidopsis} \) plants. Of the several T1 transgenic lines selected on the media containing an antibiotic, two were characterized further. These two transgenic lines showed on average 3.6 times more upregulation of \( \text{PgCYP76B93} \) (Fig. 4a) and displayed reduced plant height (Fig. 4b). In seedling stage, one strongly expressing line No.9 displayed reduced hypocotyl length (Additional...
tollerant phenotypes compared to the control plants (Fig. 6a). Statistical analysis also suggests that the survival rate (%) on 1/2 MS media containing herbicide is significant (Fig. 6b). To evaluate the resistance of already germinated seedlings, 4-day-old seedlings from Col-0 and PgCYP76B93 overexpression lines (OE lines) were analyzed in the presence and absence of two different concentrations of chlorotoluron, and found OE lines show apparent tolerant growth phenotype (Additional file 2: Fig. S3).

**Supplementary information**

**Supplementary information** accompanies this paper at https://doi.org/10.1186/s13765-020-00498-x.

Additional file 1: Table S1. PCR oligonucleotide primers used to confirm gene insertion.

Additional file 2: Fig. S1. Predicted 3D structure of CYP76B93. a The structure of Zebra Fish CYP-450 17A1 mutant Abiraterone complex. It is used as a major template to draw PgCYP76B93. Heme is colored in red, and Abiraterone is colored in light green. b The theoretical structure of PgCYP76B93. It was derived using EvPksy-SWISS MODEL. This figure was produced using PyMol (https://pymol.org/2/). Fig. S2. Heterologous overexpression of PgCYP76B93 reduced hydropocyt length. Statistical data of hyrocyclot length of each lines from 4-day-old seedlings. Data represent the mean ± standard error (SE) of three independent replicates at P < 0.05(*) by Student’s t-test. n = 23. Fig. S3. Overexpression of PgCYP76B93 confers herbicide tolerance. Four-day-old seedlings of Col-0 and OE lines were transferred on 1/2 MS media in the presence or absence of 4- and 7 μM of chlorotoluron for 18 days. Scale bars = 1 cm.

**Abbreviations**

CYP: Cytochrome P450 enzyme; qRT-PCR: Quantitative real-time reverse transcription-polymerase chain reaction; ABA: Abscisic acid; H2O2: Hydrogen peroxide; JA: Jasmonic acid; HAT: Hours after treatment.

**Authors’ contributions**

ORL conceived the project and designed the experiments. JHJ, and SK performed the experiments, except for the molecular docking experiment, which was performed by YKM and SS. ORL and JHJ analyzed the data and wrote the manuscript. All authors read and approved the final manuscript.

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**Availability of data and materials**

The datasets used and/or analysed during the current study are available from the corresponding author on reasonable request.

**Competing interests**

The authors declare that they have no competing interests.

**Author details**

1 Department of Applied Plant Science, College of Agriculture and Life Science, Chonnam National University, Gwangju 61186, Korea. 2 Interdisciplinary Program of Bioenergy and Biomaterials Graduate School, Department of Bioengineering and Biotechnology, College of Engineering, Chonnam National University, Gwangju 61186, Republic of Korea.

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Transcripts of gibberellin oxidases are altered by the overexpression of *PgCYP76B93*

Gibberellins (GAs) are well-known plant hormones that promote stem elongation as well as seed germination [32]. The *PgCYP76B93*-mediated reduced plant height (Fig. 4b) led us to analyze whether the transcript levels of gibberellin biosynthesis-related genes were changed. Gibberellins are biosynthesized from geranylgeranyl diphosphate (GGDP), which is a common C20 precursor for diterpenoids, using three different classes of enzymes [33]: (1) terpene synthases (2) cytochrome P450 monooxygenases, and (3) 2-oxoglutarate-dependent dioxygenases. To uncover a possible link with bioactive GA biosynthesis, four GA monooxidase genes were analyzed. Two GA oxidase enzymes, GA20oxI and GA3oxI, are involved in catalyzing the reaction of bioactive GA biosynthesis, whereas GA2oxI and GA2ox2 are involved in the conversion of bioactive gibberellins into an inactive form. In two different developmental stages, all transcripts of GA20oxI and GA3oxI were significantly downregulated, whereas GA20oxI and GA2ox2 were more upregulated (Fig. 5a, b). Though the mRNA levels of GA2ox2 were higher in the 28-day-old seedlings than in the 14-day-old seedlings, the general tendency was a decrease in the expression of the more bioactive form of GA oxidase gene, suggesting that the reduced height (Fig. 4b) of the *PgCYP76B93* overexpression lines is caused by reduction in the levels of the more bioactive forms of gibberellins.

**Constitutive overexpression of *PgCYP76B93* confers herbicide resistance**

*PgCYP76B93* is closely clustered with the CYP76B family of enzymes, which have been reported to be involved in herbicide metabolism [15–17]. Also upregulation of *PgCYP76B93* transcripts against herbicide sup -

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file 2: Fig. S2). It could be resulted in slightly reduced plant height in overexpression lines. Rosette leaves were smaller than those in the control lines (Fig. 4c). This implies that *PgCYP76B93* might be involved in plant growth and development via phytohormone regulation which is involved in stem and leaf development. However, the overexpression of *PgCYP76B93* did not alter the root growth phenotype (Fig. 4d, e) suggesting that the phytohormone-regulated *PgCYP76B93* gene is more functional in aerial parts than in underground parts.
References

1. Mizutani M, Ohra D (2010) Diversification of P450 genes during land plant evolution. Annu Rev Plant Biol 61:291–315
2. Kim YJ, Lee OR, Oh J, Jang MG, Yang DC (2014) Functional analysis of 3-hydroxy-3-methylglutaryl coenzyme A reductase encoding genes in intergeneric saponin-producing ginsengs. Plant Physiol 165:373–387
3. Nguyen NQ, Lee OR (2017) Overexpression of ginseng UGT72A1L1 causes organ fusion in the axillary leaf branch of Arabidopsis. J Ginseng Res 41:419–427
4. Nguyen NQ, Lee SC, Yang TJ, Lee OR (2017) cis-Prenyltransferase interacts with a Nogo-B receptor homolog for dolichol biosynthesis in Panax ginseng Meyer. J Ginseng Res 41:403–410
5. Devi BS, Kim YJ, Sathiyamoorthy S, Khorolragchaa A, Gayathri S, Parvin S, Yang DU, Selvi SK, Lee OR, Lee S, Yang DC (2011) Characterization of putative cytochrome P450 genes from Panax ginseng C. A. Meyer. Biochemistry 76:1347–1359
6. Han JY, Kim HJ, Kwon YS, Choi YE (2011) The Cyt P450 enzyme CYP716A47 catalyzes the formation of protopanaxadiol from dammarenediol-II during ginsenoside biosynthesis in Panax ginseng. Plant cell physiol 52:2062–2073
7. Han JY, Hwang HS, Choi SW, Kim HJ, Choi YE (2012) Cytochrome P450 CYP716A53v2 catalyzes the formation of protopanaxadiol from protopanaxadiol during ginsenoside biosynthesis in Panax ginseng. Plant Cell Physiol 53:1535–1545
8. Han JY, Kim MJ, Ban YW, Hwang HS, Choi YE (2013) The involvement of β-amyrin 28-oxidase (CYP716A52v2) in oleane-type ginsenoside biosynthesis in Panax ginseng. Plant Cell Physiol 54:2034–2046
9. Werck-Reichhart D, Hehn A, Didierjean L (2000) Cytochrome P450 for herbicide engineering tolerance. Trends Plant Sci 5:116–123
10. Powles SB, Yu Q (2010) Evolution in action: plant resistant to herbicides. Ann Rev Plant Biol 61:291–315
11. Brazier M, Cole DJ, Edwards R (2002) O-Glucosyltransferase activities toward phenolic natural products and xenobiotics in wheat and herbicide-resistant and herbicide-susceptible black-grass ( Alopecurus myosuroides). Phytochemistry 59:149–156
12. Kebeish R, Azab E, Peterhaensel C, El-Basheer R (2014) Engineering the metabolism of the phytoflavonoid chlortoluron in genetically modified Arabidopsis thaliana plants expressing the mammalian cytochrome P450 enzyme CYP1A2. Environ Sci Pollut Res Int 21(13):8224–8232
13. Frear DS, Swanson HR, Tanaka FS (1969) 3-(phenyl)-1-methylureas—isolation and characterization of a microsomal mixed function oxidase from cotton. Phytochemistry 8:2157–2169
14. Robinau T, Batard Y, Neldelkina S, Cabello-Hurtado F, LeFlet P, Sorokin O, Didierjean L, Werck-Reichhart D (1998) The chemically inducible plant cytochrome P450 CYP76B1 actively metabolizes phytolipids and other xenobiotics. Plant Physiol 118(3):1049–1056
15. Didierjean L, Gondet L, Perkins R, Lau SM, Schaller H, O’Keefe DP, Werck-Reichhart D (2002) Engineering herbicide metabolism in tobacco and Arabidopsis with CYP7681, a cytochrome P450 enzyme from Jerusalem artichoke. Plant Physiol 130:179–189
16. Hofier R, Boachon B, Renaud H, Gavira C, Miesch L, Iglesias J, Ginglinger JF, Allouche L, Miesch M, Grec S, Larbat R (2014) Dual function of the CYP76 family from Arabidopsis thaliana in the metabolism of monoterpenoids and phenylurea herbicides. Plant Physiol 166:1149–1161
17. Siminskiy B, Corbin FT, Ward ER, Fleischmann TJ, Dewey RE (1999) Expression of a soybean cytochrome P450 monooxygenase cDNA in yeast and tobacco enhances the metabolism of phenylurea herbicides. Proc Natl Acad Sci 96:1750–1755
18. Azab E, Kebeish R, Hegazy AK (2018) Expression of the human gene CYP1A2 tolerance and detoxification of the phytoflavonoid herbicide linuron in Arabidopsis thaliana plants and Escherichia coli. Environ Pollut 238:281–290
19. Brazier M, Cole DJ, Edwards R (2002) O-Glucosyltransferase activities toward phenolic natural products and xenobiotics in wheat and herbicide-resistant and herbicide-susceptible black-grass ( Alopecurus myosuroides). Phytochemistry 59:149–156
20. Allouche L, Miesch M, Grec S, Larbat R (2014) Dual function of the CYP76 family from Arabidopsis thaliana in the metabolism of monoterpenoids and phenylurea herbicides. Plant Physiol 166:1149–1161
21. González E, Johnson KM, Pallan PS, Pham TT, Zhang W, Lei L, Wawrzak Z, Yoshimoto F, Eslì M, Guengerich FP (2018) Inherent sterol 17α, 20-lyase activity in defunct cytochrome P450 17A enzymes. J Biol Chem 293(2):541–556
22. Lauskowska R, Swindells MB (2011) Ligand–protein interaction diagrams for drug discovery. J Chem Inf Model 51(10):2778–2786
23. Lee OR, Kim SJ, Kim HJ, Hong JK, Ryu SB, Lee SH, Cho HT (2010) Phospholipase A2 is required for PIN-FORMED protein trafficking to the plasma membrane in the Arabidopsis root. Plant Cell 22:1812–1825
24. Bechtold N, Pelletier G (1998) In planta Agrobacterium-mediated transformation of adult Arabidopsis thaliana plants by vacuum infiltration. Methods Mol Biol 82:259–266
25. Kim MK, Lee BS, In JG, Sun H, Yoon JH, Yang DC (2006) Comparative analysis of expressed sequence tags (ESTs) of ginseng leaf. Plant Cell Rep 25:599–606
26. Pulla RK, Lee OR, In JG, Parvin S, Kim YJ, Shim JS, Sun H, Kim YJ, Senthil K, Yang DC (2011) Identification and characterization of class I chitinase in Panax ginseng C. A. Meyer. Mol Biol Rep 38:95–102
27. Nielsen DR (2009) The cytochrome P450 homepage. Hum Genomics 325:68–74
28. Collu G, Unver N, Peltenburg-Loonan AM, van der Heijden R, Verpoorte R, Memelink J (2001) Genossil 10-hydroxylase, a cytochrome P450 enzyme involved in terpenoid indole alkaloid biosynthesis. FEBS Lett 508:215–220
29. Wang J, Liu Y, Cai Y, Zhang F, Xia G, Xiang F (2010) Cloning and functional analysis of geraniol 10-hydroxylase, a cytochrome P450 from Szentimuru satis Franch. Biosci Biotechnol Biochem 74:1583–1590
30. Wilkins MR, Gasteiger E, Bairoch A, Sanchez JC, Williams KL, Appel RD, Hochstrasser DF (1999) Protein identification and analysis tools in the ExPASy server. Methods Mol Biol 122:531–552
31. Chen W, Lee MK, Jefcoate C, Kim SC, Chen F, Yu JH (2014) Fungal cytochrome P450 enzymes: their distribution, structure, functions, family expansion, and evolutionary origin. Genome Biol Evol 6:1620–1634
32. Lester DR, Ross JJ, Davies PI, Reid JB (1997) Mendel’s stem length gene (le) encodes a gibberellin 3 β-hydroxylase. Plant Cell 9:1435–1443
33. Yamaguchi S (2008) Gibberellin metabolism and its regulation. Annu Rev Plant Biol 59:225–251

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