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Source: Journal of Insect Science, 15(1) : 1-7

Published By: Entomological Society of America

URL: <https://doi.org/10.1093/jisesa/ieu174>

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## RESEARCH

# Identification and Expression of Two Novel Cytochrome P450 Genes, *CYP6CV1* and *CYP9A38*, in *Cnaphalocrocis medinalis* (Lepidoptera: Pyralidae)

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**Subject Editor:** Yu-Cheng Zhu

J. Insect Sci. (2015) 15(1): 50; DOI: 10.1093/jisesa/ieu174

**ABSTRACT.** *Cnaphalocrocis medinalis* Guénée can cause severe losses in rice. Cytochrome P450s play crucial roles in the metabolism of allelochemicals in herbivorous insects. Two novel P450 cDNAs, *CYP6CV1* and *CYP9A38*, were cloned from the midgut of *C. medinalis*. *CYP6CV1* encodes a protein of 500 amino acid residues, while *CYP9A38*-predicted protein has 531 amino acid residues. Both cDNA-predicted proteins contain the conserved functional domains for all P450s. Phylogenetic analyses showed that *CYP6CV1* is grouped in the cluster containing CYP6B members, while *CYP9A38* is in the cluster including CYP9 members. However, both clusters are contained in the same higher lineage. Homologous analysis revealed that *CYP6CV1* is most similar to CYP6B8, CYP6B7, CYP6B6, CYP6B2, and CYP6B4 with the highest amino acid identity of 41%. *CYP9A38* is closest to CYP9A17, CYP9A21, CYP9A20, and CYP9A19 with the highest amino acid identity of 66%. Studies of temporal expression profiles revealed that *CYP9A38* showed a steady increase in mRNA level during the five instar stages, but a low-expression level in pupae, and then presented at a high-expression level again in adults. Similar expression patterns were obtained with *CYP6CV1*. In the fifth instar larvae, *CYP6CV1* was mainly expressed in midgut and fat bodies, whereas *CYP9A38* was mainly expressed in midgut. Expression studies also revealed a 3.20-fold over-expression of *CYP6CV1* and 3.54-fold over-expression of *CYP9A38* after larval exposure to host rice resistance. Our results suggest that both *CYP6CV1* and *CYP9A38* may be involved in detoxification of rice phytochemicals.

**Key Words:** *Cnaphalocrocis medinalis*, *CYP6CV1*, *CYP9A38*, host rice resistance, expression profile

The rice leaffolder *Cnaphalocrocis medinalis* (Guénée) (Lepidoptera: Pyralidae) is a species of the Crambidae family. It is considered a major pest of rice (*Oryza sativa* L.) (Riley et al. 1995). The attacked rice plants dry up and the vigor of plants reduces. The yield loss may vary up to 10–50%. And it is more problematic at booting stage (Riley et al. 1995). Serious outbreaks of *C. medinalis* have been reported in many Asian countries including India, Korea, Japan, China, Malaysia, Sri Lanka, and Vietnam (Senthil et al. 2006). And *C. medinalis* can undertake migrations which makes the pest control difficult (Riley et al. 1995).

Some rice varieties such as TKM-6 (*O. sativa* L.) show resistance to *C. medinalis* to a certain extent, while some like Taichung Native 1 (TN1) (*O. sativa* L.) are susceptible to this pest. In Asia, TN1 and TKM-6 have been the most frequently used in crossing programs (Upadhyay et al. 2011). TN1 was developed in Taiwan. It was the first semidwarf indica to respond to nitrogen fertilizer up to ~100 kg/ha. TN1 is short-statured (83–85 cm) and high tillering. But the area planted gradually declined, because TN1s principal deficiency is susceptibility to several insects and diseases (Dalrymple 1978). TKM-6 was developed in India. It has a weak stem. It was used as a parent for several high yielding Indian and International Rice Research Institute (IRRI) varieties. The discovery of resistant gene *Bph 1* in TKM-6 is significant for rice breeders because the variety is resistant to several other diseases and insects (Dalrymple 1986, Khan and Joshi 1990). Therefore, both of TN1 and TKM-6 are important germ plasm resources in rice breeding, as well as vital materials in fundamental research. In this work, we use the two rice varieties as materials to rear or treat *C. medinalis* larvae.

The mechanism of resistance of TKM-6 to *C. medinalis* was attributed to a comparatively broader and thicker sclerenchymatous hypodermis in the stem, to the closer disposition of vascular bundles, the presence of more silicated cells in the leaf epidermis, more wax on leaf

surface and narrower, and more hairy leaves (Chandramani et al. 2009). One other factor responsible for resistance in TKM-6 seems to be of biochemical nature. The changes of various biochemical constituents such as leaf soluble protein, phenol, ortho-dihydroxy phenol, tannin, and enzymes viz., peroxidase, phenyl alanine ammonia lyase (PAL) were ever assessed spectrophotometrically in the rice genotypes before and after *C. medinalis* infestation. The protein profile was analyzed using sodium dodecyl sulphate-poly acrylamide gel electrophoresis (SDS-PAGE) method. A significant constituent of biochemical content such as tannin, phenol, and ortho-dihydroxy phenol was proved increased along with enzyme activities of peroxidase and PAL in the infested TKM-6 rice (Punithavalli et al. 2013). It is also evident that there are more biochemicals such as phenol, orthodihydroxy phenol, and tannin in TKM-6 than in TN1 plants, which were negatively correlated with *C. medinalis* damage. However, leaf protein content was less in TKM-6 than in TN1, which was positively correlated with the damage by *C. medinalis* (Punithavalli et al. 2013). Meanwhile, the increased death of early instars, the slower development of larvae, the reduced size or pupal stage, and abnormal behavior of *C. medinalis* were watched on TKM-6, when compared with that on the other rice varieties (Khan et al. 1989, Masoud et al. 1996).

It is well accepted that planting resistant rice varieties is an effective practice to control some of rice pests (Sogawa et al. 2003). But no rice varieties with sufficient resistance level have been developed to control *C. medinalis* in practical application to date (Rao et al. 2010). Why *C. medinalis* has so strong adaptation to its host rice remains unknown. Therefore, to research the molecular interaction of *C. medinalis* and its host rice resistance may obtain some crucial information to develop novel integrated pest managements (IPMs).

The P450s are important metabolic systems in insects because of their involvement in the syntheses of endogenous hormones, fatty acids

and steroids, and in the catabolism of xenobiotics, such as drugs, pesticides, and plant toxins (Scott 2008). Multiple P450 family genes were found in herbivorous insects, which represent an adaptation in the “animal–plant warfare” (Gonzalez and Nebert 1990). Ingestion of plant toxins frequently induces insects P450 genes responsible for catabolism of plant toxins. Some examples of insect P450 genes induced by phytochemicals are summarized in Feyereisen (2005), Schuler (2011), and Scott (2008). As noted previously, we found that *CYP6AE28* and *CYP6AE30* of *C. medinalis* are induced in response to resistant rice variety TKM-6 (Liu et al. 2010). In this study, we cloned another two P450 genes *CYP6CV1* and *CYP9A38* from *C. medinalis* and studied their expression profiles, in order to establish a foundation for further study of their functions.

## Materials and Methods

**Experimental Insects and RNA Isolation.** Insect cultures: *C. medinalis* larvae were collected from paddy fields in the Wuchang district, Wuhan City, China, and reared in containers on TN1 rice in booting stage, a leafhopper susceptible rice variety, at 25°C under a photoperiod of 16:8 (L:D) h, and 80% relative humidity.

Treatment with TKM-6 rice: For induction analyses of the P450 genes, the newly molted fifth instar larvae were transferred from TN1 plants to TKM-6 rice in booting stage and kept for 24 h. Whole insect bodies were collected and deeply frozen in liquid nitrogen then stored in –80°C refrigerator for further use.

RNA isolation and cDNA synthesis: For temporal expression analyses of the P450 genes, whole bodies of the newly developed larvae in the first, second, third, fourth, fifth instar stages, pupae, and adults were collected for total RNA isolation by using TRIzol reagent (Invitrogen, Carlsbad, CA). For spatial expression analyses, head, midgut, fat body, and carcass dissected from the fifth instar larvae treated with TKM-6 were used for RNA isolation. For induction analyses, the fifth instar larvae fed with TN1 and TKM-6 plants were used for RNA extraction. Half microgram of total RNA was used for cDNA synthesis using the iScript™ cDNA synthesis kit (BioRad, Hercules, CA).

**Cloning of the Full-Length cDNAs.** A pair of degenerate oligonucleotide primers (dCYP6-S and dCYP6-AS, Table 1) was designed for amplification of cytochrome P450 family six genes (Kasai et al. 2000). Another pair of degenerate primers (dCYP9-S and dCYP9-AS, Table 1) was used to amplify cytochrome P450 family nine genes (Stevens et al. 2000). The cDNAs transcribed from the midgut, RNA were used as template in the reverse transcription-polymerase chain reactions (RT-PCRs). Two P450 clones representing two novel P450 genes were selected after cloning, sequence and analyzing for CYP6 and nine homologies with the National Center for Biotechnology Information (NCBI) Basic Local Alignment Search Tool (BLAST)

Network Server. Gene-specific primers (GSPs) for 5' and 3' rapid-amplification of cDNA ends (RACE) were designed according to the two cDNA clones. The primers were rCYP6-S, rCYP6-AS, rCYP9-S, and rCYP9-AS (Table 1). The cDNA synthesis and RACE were performed exactly according to the instruction manual of the SMART RACE cDNA Amplification Kit (BD Bioscience Clontech, Palo Alto, CA).

Amplified fragments were routinely cloned into pGEM-T vector (Promega, Madison, WI) and sequenced. The nucleotide sequences of the 5' and 3' RACE products were aligned to form two full-length cDNA sequences and the cDNA-predicted proteins were called as CYP6CV1 and CYP9A38 by the P450 nomenclature committee (D.R. Nelson), respectively. Two pairs of primers for long distance-PCR (LD-PCR) were designed to amplify the internal sequences of the full cDNAs, respectively. The primers were lCYP6-S, lCYP6-AS, lCYP9-S, and lCYP9-AS (Table 1). The PCR system was heated at 95°C for 1 min and then amplified for 34 cycles (95°C for 30 s, 62°C for 30 s, and 72°C for 3 min). Amplified fragments were cloned and sequenced.

**Computer-Assisted Analysis of P450 cDNAs.** Molecular mass and isoelectric point were predicted by Compute pI/Mw tool ([http://us.expasy.org/tools/pi\\_tool.html](http://us.expasy.org/tools/pi_tool.html)). The transmembrane anchors of the deduced P450s were predicted by the TMPred server ([http://www.ch.embnet.org/software/TMPRED\\_form.html](http://www.ch.embnet.org/software/TMPRED_form.html)). ClustalX v1.8 (Thompson et al. 1997) was used to analyze the alignment. A molecular phylogenetic tree was constructed by the ClustalW Server (<http://crick.genesis.nig.ac.jp/homology/clustalw-e.shtml>) by using the bootstrap N-J tree option (number of bootstrap trials = 1,000; Page 1996).

**Real-Time Quantitative PCR Analysis.** The transcript levels of *CYP6CV1* and *CYP9A38* were determined by real-time quantitative PCR (qPCR), using iQ SYBR Green Supermix (Bio-Rad, Hercules, CA) in the iCycler iQ Real Time PCR Detection System (Bio-Rad, Hercules, CA). To standardize qPCR inputs, a master mix that contained iQ SYBR Green Supermix and forward and reverse primers was prepared (final concentration = 100 nM per reaction; primer sequences [qCYP6-S, qCYP6-AS, qCYP9-S, and qCYP9-AS] are listed in Table 1). The qPCRs were conducted with the same quantity of 10-fold diluted cDNA for each instar larva, pupa, and adults for analyses of temporal gene expression profiles. The reactions were also carried out with equivalent cDNA input for the organs, including head, midgut, fat body, and carcass from the fifth instar larvae for analyses of spatial gene expression profiles. In addition, equal cDNA input of the fifth instar larvae ingested with TN1 and TKM-6 plants were used for analyses of host resistance induction. PCR profiles were: 95°C for 5 min; 40 cycles of 95°C for 15 s, 55°C for 10 s, and 72°C for 10 s with a plate read at the end of each cycle. All reactions were performed in duplicates and three replicates were used to estimate variation.

**Table 1. Primers used in the present study**

| Primer set | Primer sequence (5'-3')   | Application type | Product length [bp] |
|------------|---|------------------|---------------------|
| dCYP6-S    | GA(A or G)AC(A or G or C or T)(A or C or T)(C or T)(A or G or C or T)(A or C)                       | RT-PCR           | 239                 |
| dCYP6-AS   | G(A or G or C or T)CC(A or G or C or T)(G or T)C  |                  |                     |
| dCYP9-S    | GG(A or G or C or T)CC(A or G or C or T)(G or T)C(A or G or C or T)CC(A or G)AA(A or G or C or T)GG | RT-PCR           | 404                 |
| dCYP9-AS   | TACGA(AG)(CT)IGCI(AG)(AT)IAA(CT)CC(CT)GA  |                  |                     |
| rCYP6-S    | CCIA(GT)(AG)CA(AG)TTIC(GT)IGGICC  | RACE             | 856                 |
| rCYP6-AS   | TGGCTGATTACAGTTTCCTGGAAGTGA   | RACE             | 1,258               |
| rCYP9-S    | TCGGAAATATTCTGGGTCGGCGTTA   | RACE             | 569                 |
| rCYP9-AS   | CTGGTATCGCGATGGACCGGATCTGC  | RACE             | 1,427               |
| lCYP6-S    | CTCGGAGAAGCGCTCGGGGTCAAAC   | LD-PCR           | 1,636               |
| lCYP6-AS   | CTCCAACATGGCGCTGCTCGTG  |                  |                     |
| lCYP9-S    | CTCAATTACGCCATCGCCTTC   | LD-PCR           | 1,554               |
| lCYP9-AS   | GCGCAAACCGGCTGAGCCATG   |                  |                     |
| qCYP6-S    | GGACAGCTGGGACGGGATGCTG  | qPCR             | 198                 |
| qCYP6-AS   | ACGTTTCCTGGAAGTATGTCA   |                  |                     |
| qCYP9-S    | CAAACCAATGCAATTCGAGGT   | qPCR             | 188                 |
| qCYP9-AS   | CAGATCCTCACCTTCTTCGCTT  |                  |                     |
|            | CGACCTTTCTTAGCCTCCATGA  |                  |                     |

S and AS indicate forward primer and reverse primer, respectively. Y = C or T, K = T or G.



Beta-actin was used as an internal reference. Relative accumulation of *CYP6CV1* and *CYP9A38* normalized against beta-actin was calculated from the formula  $2^{-\Delta\Delta Ct}$  where two is the reaction efficiency and  $\Delta\Delta Ct$  is the difference in beta-actin Ct values between a defined control and the rest samples in an assay, or the difference in *CYP6CV1* or *CYP9A38* Ct values between a defined control and the rest samples in an assay.

## Results

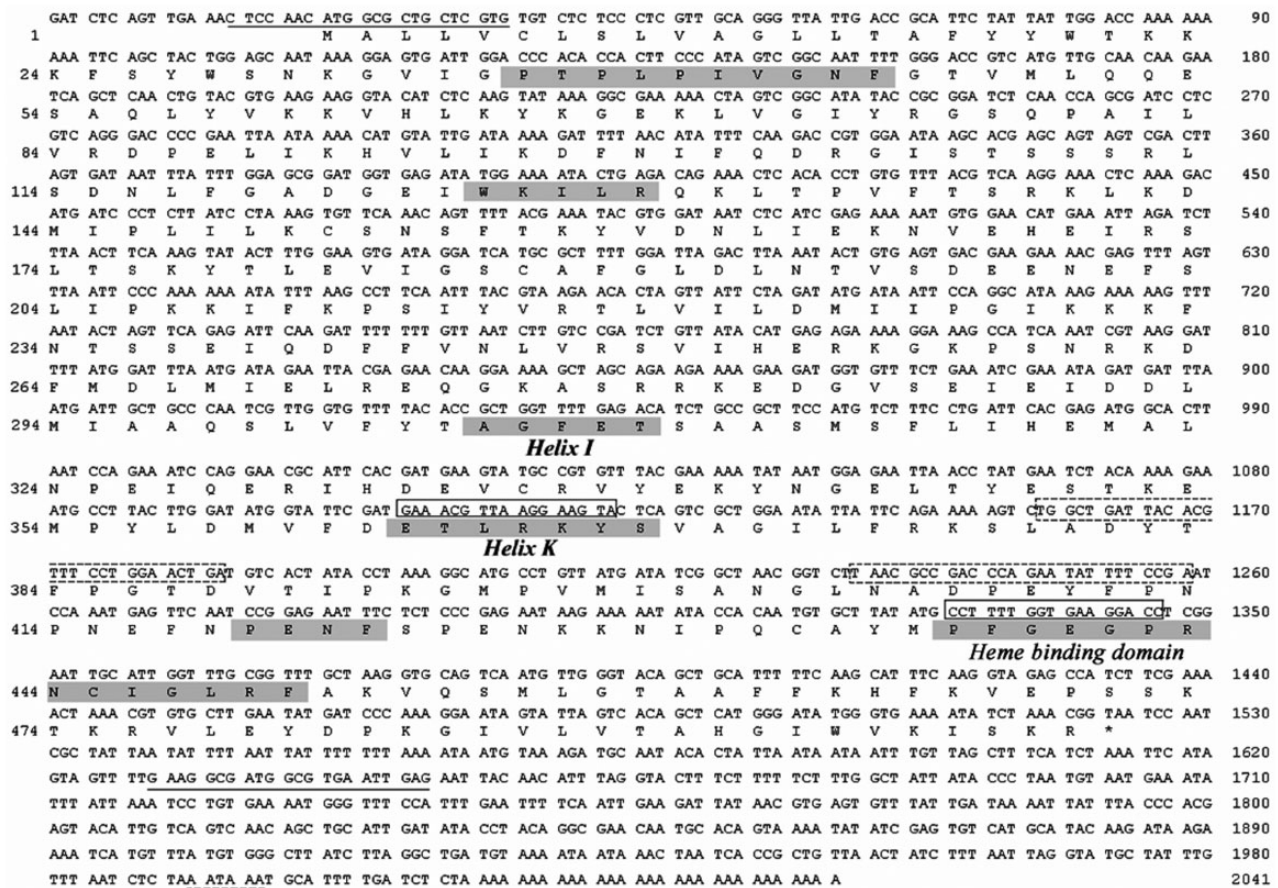
**cDNA Cloning and Characterization.** Two pairs of degenerate primers for the conserved regions of insect P450 proteins were used to amplify reverse-transcribed midgut mRNA of *C. medinalis*. The PCR products were cloned and sequenced. A clone with 239 bp in length encoding a reading frame of 79 amino acids was amplified by using dCYP6-S and dCYP6-AS; another clone with 404 bp encoding 134 amino acids was obtained by using dCYP9-S and dCYP9-AS as primers. Both clones scored highly with other P450 proteins by BLAST Network searches. Based on the two cDNA clones, GSPs were designed. Two P450 cDNAs with 2,041 and 1,853 bp in length were amplified by RACE and LD-PCR strategies. Close alignment with known insect CYP6 and CYP9 members showed that the two new P450s belonged to the two families, respectively. The former sequence (accession number FN421127) was called CYP6CV1, and the later (accession FN421128) was called CYP9A38.

CYP6CV1 nucleotide sequences analysis revealed that this cDNA contains a putative ORF of 1,503 bp, a 21 bp 5'-untranslated region (5'-UTR), and a 517 bp 3'-UTR, with a putative polyadenylation signal sequences (AATAAA) upstream of the poly(A) tract (Fig. 1). CYP9A38 cDNA is composed of a 1,593 bp ORF, with a 63 bp 5'-UTR

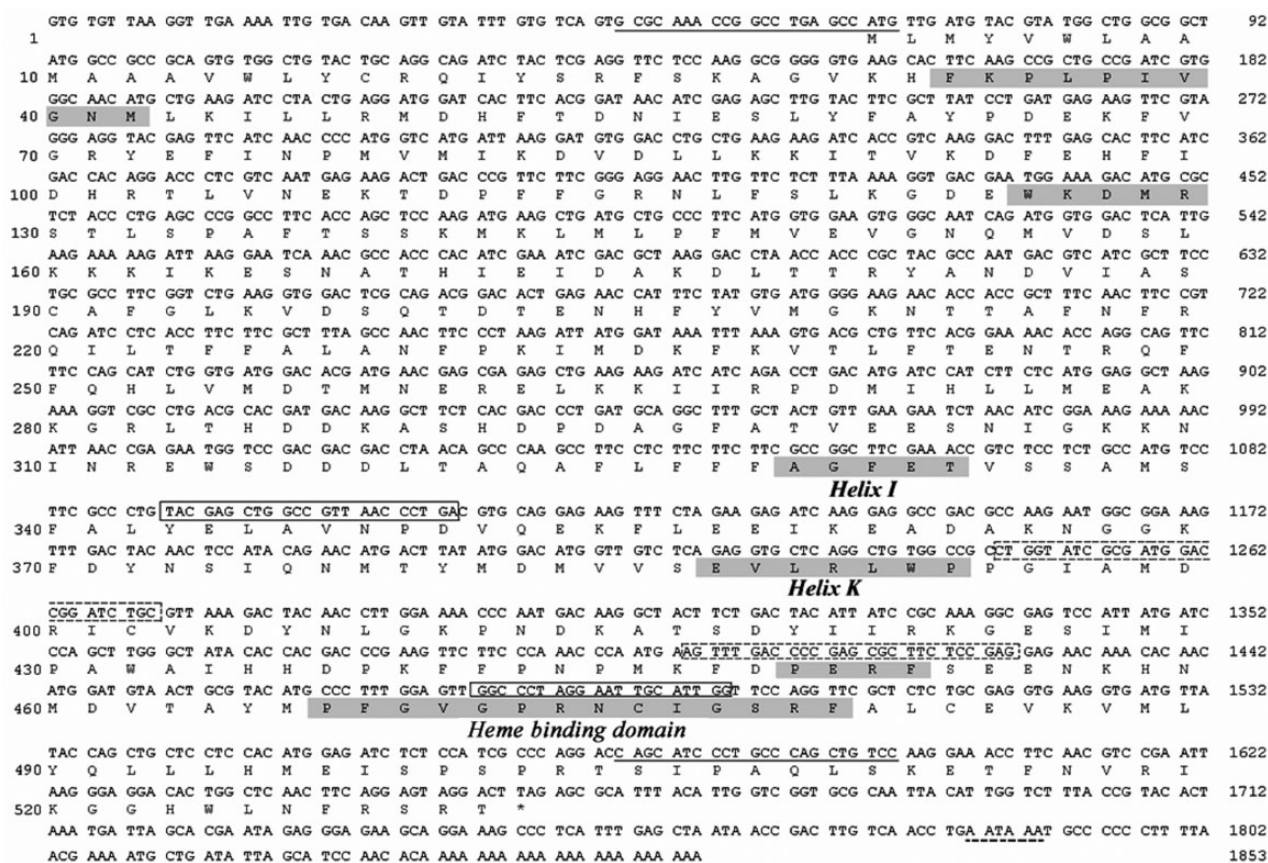
and a 197 bp 3'-UTR with a polyadenylation signal (AATAAA) upstream of the poly(A) tract (Fig. 2).

**Characterization of the cDNA-Predicted Proteins.** Based on the predicted amino acid sequence, CYP6CV1 has a theoretical *pI* value of 8.95 and molecular mass of 56,886, and CYP9A38 has a theoretical *pI* value of 7.67 and molecular mass of 61,454. Two strong inside-to-outside transmembrane helices from amino acid 2 to 20 and 293 to 315 in CYP6CV1 sequences, and three transmembrane helices from amino acid 1 to 18, 215 to 234 and 319 to 342 in CYP9A38 sequences were predicted, suggesting that both P450s are endoplasmic reticulum membrane-bound proteins. The putative proteins contain the typical motifs of an insect P450 protein (Nelson et al. 1993), including the proline and glycine rich hinge region (xxPxPxGxx), helix-C (WxxxR), helix-I (AGxE or DT), helix-K (ExxR), PExF (PxxFxxPxxF), and the heme-binding domain (FxxGxxxG) (Figs. 1 and 2). Figure 3 shows the alignment of CYP6CV1, CYP6AE28, CYP6AE30, and the other four CYP6 family members (CYP6A2, CYP6B2, CYP6AE1, and CYP6AE12). All of them contain the six residue sequence PExFxxP (PENFSP, position 419 to 424 in CYP6CV1) upstream of the heme-binding domain, specific to family six members (Nelson et al. 1993). Likewise, the SR(F or I or L)(A or G)xx(Q or E) sequence immediately following the heme-binding domain, specific to family 9, is found in CYP9A38 (SRFALCE, position 478 to 484; Maibèche-Coisne et al. 2005).

The relatedness of CYP6CV1, CYP6AE28, CYP6AE30, CYP9A38, and some CYP6 family P450s is revealed by the fact that CYP6CV1 and CYP6CV2 form an independent cluster contained in the clade including CYP6A2 and CYP6B members. CYP9A38 is in the cluster including other six CYP9 members. A higher lineage containing



**Fig. 1.** Nucleotide sequence and deduced amino acid sequence of *CYP6CV1* in *C. medinalis*. The proline and glycine rich hinge region, C-helix, I-helix, K-helix, PExF, and heme-binding domain are shaded. The consensus polyadenylation signal is indicated by a discrete underline. The degenerate primer corresponding regions are framed. The GSPs are framed with discrete line. The LD-PCR primers are underlined.



**Fig. 2.** Nucleotide sequence and deduced amino acid sequence of *CYP9A38* in *C. medinalis*. The conserved domains, polyadenylation signal, and primers are marked as in Fig. 1.

the above two clusters is formed in the phylogenetic tree. *CYP6AE28* and *CYP6AE1* form another independent cluster, while *CYP6AE30* is in the cluster containing *CYP6AE9* and *CYP6AE47* as shown in Fig. 4.

Homologous analysis revealed that *CYP6CV1* shows the highest percentage amino acid identities (41%) to *Helicoverpa zea* *CYP6B8* (AF285828), *Helicoverpa armigera* *CYP6B7* (ABE60887), *CYP6B6* (ABE60886), *CYP6B2* (ABE60885), and *Papilio glaucus* *CYP6B4* (AAB05892), respectively. *CYP9A38* is closest to members of the *CYP9A* subfamily. It shares the highest identity (66%) with *CYP9A17* (ACJ37388) from *H. armigera*, *CYP9A21* (ABN71369), *CYP9A20* (ABO07439), *CYP9A19* (ABQ18318) from *Bombyx mori*, respectively.

**Expression Analyses of *CYP6CV1* and *CYP9A38*.** We analyzed the mRNA levels of *CYP6CV1* and *CYP9A38* across insect life stages by quantitative RT-PCR (Fig. 5A). Both gene mRNA level exhibited a cyclic pattern. *CYP9A38* showed a steady increase in mRNA level during the five instar stages, but a low-expression level in pupae, and then presented at high-expression level again in adults. Similar expression patterns were obtained with *CYP6CV1*. *CYP6CV1* was significantly activated in the fourth, fifth instar larvae, and the adults. Namely, *CYP9A38* expression was 4.43-fold, 4.89-fold, 8.53-fold, 9.48-fold, 6.77-fold, and 4.50-fold comparatively to pupae for first, second, third, fourth, fifth larval stages and adults, respectively. *CYP6CV1* was expressed by 0.61-fold, 1.46-fold, 1.61-fold, 3.50-fold, 3.35-fold, and 3.44-fold comparatively to pupae for first, second, third, fourth, fifth larval stages and adults, respectively.

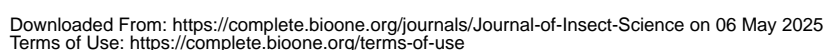
In the fifth instar larvae, *CYP6CV1* was mainly expressed in midgut and fat bodies, with 0.14-fold, 1.41-fold, and 0.27-fold comparatively to fat bodies for head, midgut, and carcass, respectively. Whereas *CYP9A38* was mainly expressed in the midgut, with 0.38-fold, 4.72-fold, and 0.32-fold comparatively to fat bodies for head, midgut, and carcass, respectively (Fig. 5B).

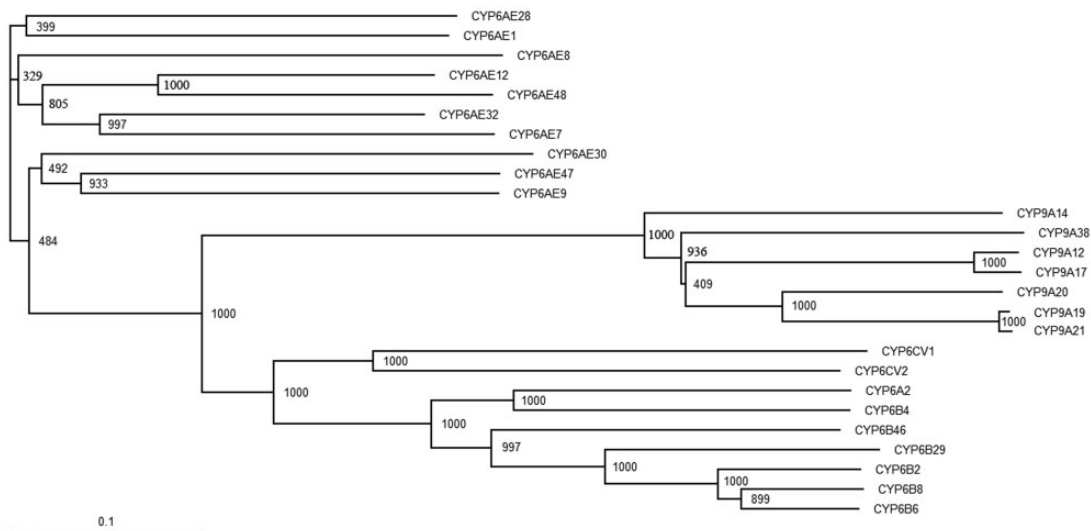
In the fifth instar larvae reared on the susceptible rice TN1 plants, both *CYP6CV1* and *CYP9A38* were constitutively expressed at low levels, but significantly induced by exposure to the resistant rice variety TKM-6 (Fig. 6). Expression studies revealed a 3.20-fold over-expression of *CYP6CV1* and 3.54-fold over-expression of *CYP9A38* after larval exposure to host rice resistance.

## Discussion

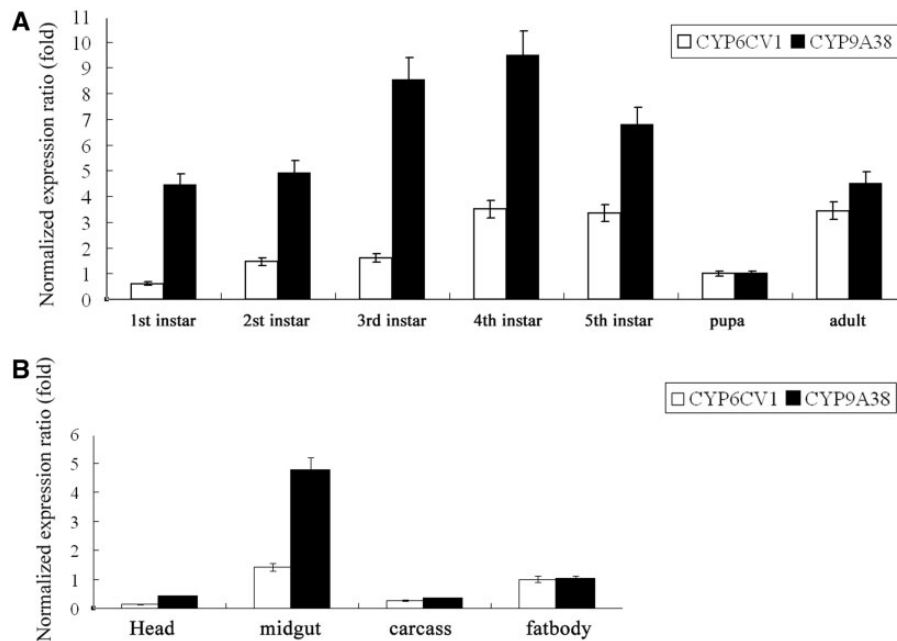
During the last three decades, people have focused interest in insect P450s on their role in fundamental physiological functions, such as growth, development or reproduction through the biosynthesis and the catabolism of key hormones, such as juvenile hormone (JH) or 20-hydroxyecdysone, in the oxidative metabolism of various xenobiotics including insecticides and plant phytochemicals (Feyereisen 2005). Most of the *CYP6* family members studied to date, especially in crop pests, are generally expressed in the digestive tract and fat body, and were found to be mainly responsible for insecticide metabolism and inactivation of phytochemicals (Li et al. 2002, Feyereisen 2005, Scott 2008). The insect *CYP9* family is most closely related to the *CYP6* family, and together with the *CYP28* family, they are grouped in the *CYP3* clan including some mammal P450s (Nelson 1998). *CYP9A1* from *Heliothis virescens*, the first member of this family, was found to be constitutively over-expressed in thiodicarb-selected tobacco budworms and may play a role in pesticide metabolism (Rose et al. 1997). *CYP9A2* was activated by the wild tomato compound 2-undecanone, indole-3-carbinol, phenobarbital, 2-tridecanone, and xanthotoxin. *CYP9A4* and *CYP9A5* were induced differentially by clofibrate and xanthotoxin (Stevens et al. 2000). *CYP9A13* may probably involve in the metabolism of odorant compounds and play a role in taste in the moth *Mamestra brassicae* (Maibèche-Coisne et al. 2005). *CYP9A12* and *CYP9A17* mRNA proved to be affected by deltamethrin, gossypol







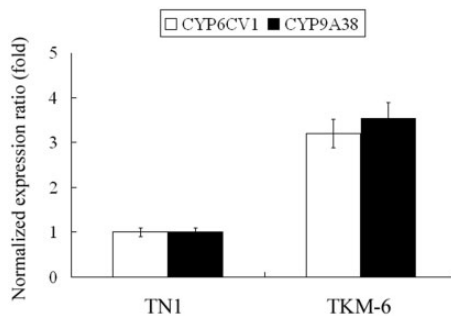
**Fig. 4.** Phylogenetic tree of full-length amino acid sequences of deduced CYP6CV1 and CYP9A38 and some members from CYP6 and nine families constructed by the neighbour-joining method. The tree was constructed with the full-length sequences of the P450s. The bootstrap values were indicated on each branch. The GenBank accession numbers of the P450s in a reduced version of the neighbour-joining tree are: *Plutella xylostella* CYP6CV2 ADW27429.1, *P. xuthus* CYP6A2 BAM18141.1, *Manduca sexta* CYP6B46 ADE05579.1, *Spodoptera litura* CYP6B29 ACY41036.1, *H. zea* CYP6B8 AAM90316.1, *P. glaucus* CYP6B4 AAB05892.1, *H. armigera* CYP6B6 AAY21920.1, *H. armigera* CYP6B2ABE60885.1, *C. medinalis* CYP6AE28 CAX94849.1, *C. medinalis* CYP6AE30 CBB07053.1, *M. sexta* CYP6AE32 ADE05581.1, *H. armigera* CYP6AE12 ABB69054.1, *D. pastinacella* CYP6AE1 AAP83689.1, *Spodoptera littoralis* CYP6AE48 AFP20589.1, *B. mori* CYP6AE9 NP\_001104004.1, *S. littoralis* CYP6AE47 AFP20588.1, *B. mori* CYP6AE7 NP\_001104006.1, *Bombyx mandarina* CYP6AE8 ABY40426.1, *B. mori* CYP9A20 NP\_001077079.1, *B. mori* CYP9A19 ABQ18318.1, *H. armigera* CYP9A12 ACB30273.2, *H. armigera* CYP9A17 AAY21809.1, *B. mori* CYP9A21 NP\_001103394.1, *H. armigera* CYP9A14 ABY47596.1.



**Fig. 5.** Expression levels of CYP6CV1 and CYP9A38 mRNA in different developmental stages of *C. medinalis* (A) and in different tissues of fifth instar larvae of *C. medinalis* (B). A. Expression levels of CYP6CV1 and CYP9A38 in *C. medinalis* larvae during larval-larval, larval-pupal, and pupal-adult transitions were detected by qRT-PCR and normalized against *actin* transcript, and then normalized to obtain an expression ratio of 1 in pupas. B. Detection of CYP6CV1 and CYP9A38 expression in head, midgut, carcass, and fat body of *C. medinalis*. Equal tissue equivalents were analyzed by qRT-PCR and normalized against *actin* transcript, and then normalized to obtain an expression ratio of 1 in fat bodies. mRNA profiles are representative of three independent replicates. Vertical bars indicate the SEM ( $n = 3$ ).

In our previous work, two P450 genes *CYP6AE28* and *CYP6AE30*, cloned from *C. medinalis* for the first time, were found to be induced by TKM-6 (Liu et al. 2010). The results together with the data in this study contribute to characterize the response of *C. medinalis* to rice dietary

phytochemicals and emphasize the role of P450 genes in the adaptation of *C. medinalis* larvae to resistant rice. More works including heterologous expression of the new P450 genes, reconstruction of heterologously expressed enzyme systems, studying the biochemical



**Fig. 6.** Expression levels of *CYP6CV1* and *CYP9A38* mRNA in *C. medinalis* larvae feeding on different rice varieties. mRNA profiles are representative of three independent replicates. Vertical bars indicate the SEM ( $n = 3$ ). TN1 and TKM-6 indicate *C. medinalis* larvae feeding on the rice plants of TN1 and TKM-6, respectively.

characteristics of the P450s and identification of their natural substances from rice, need to be carried out in the future to reveal the function of these P450 genes.

### Acknowledgments

We thank Dr. David R. Nelson and Dr. Daniel W. Nebert for naming the *C. medinalis* P450s. This work was supported by grants from the National Natural Science Foundation of China (No. 31301675, 30500328, 31071679, and 31272050), Educational Commission of Hubei Province of China (No. D20091007), Science and Technology Department of Hubei Province of China (No. 2010CDB04504 and 2011CDB064), Guangzhou Yangcheng Scholarship (10A042G).

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Received 21 December 2013; accepted 20 October 2014.