Cloning and characterization of NYD-OP7, a novel deltamethrin resistance associated gene from Culex pipiens pallens

Xiaobang Hu a,1, Yan Sun a,1, Weijie Wang a, Mingxia Yang a, Lixin Sun a, Wenbin Tan a, Jing Sun a, Jin Qian a, Lei Ma a, Donghui Zhang a, Guiyun Yan b, Changliang Zhu a,*

a Department of Pathogen Biology, Nanjing Medical University, Jiangsu Province Key Laboratory of Modern Pathogen Biology, Nanjing 210029, PR China
b Program in Public Health, College of Health Sciences, University of California at Irvine, Irvine, CA 92697-4050, USA

Received 27 June 2006; accepted 20 September 2006
Available online 26 September 2006

Abstract

One mosquito opsin gene, NYD-OP7, has been cloned from Culex pipiens pallens. An open reading frame (ORF) of 1116 bp was found to encode a putative 371 amino acids protein which exhibits high identity with opsins from Aedes and Anopheles mosquitoes. Transcript expression of NYD-OP7 was determined by real-time PCR in all life stages of deltamethrin-susceptible and -resistant strains of the Culex mosquito. The results demonstrated that this gene is expressed at all developmental stages, and it is expressed predominantly at the pupae and adult stages. Meanwhile, in pupae and adults, NYD-OP7 is overexpressed in deltamethrin-resistant strain than in -susceptible strain. Importantly, stable expression of NYD-OP7 in the mosquito C6/36 cells can confer moderate deltamethrin resistance. Our study provides the first direct evidence that increased expression of an opsin gene may play some role in the development of deltamethrin resistance in Cx. pipiens pallens.

© 2006 Elsevier Inc. All rights reserved.

Keywords: Culex pipiens pallens; Opsin; Reverse transcription PCR; Transfection; Deltamethrin resistance

1. Introduction

Opsins, the visual pigments of animal photoreceptors, belong to the G-protein coupled receptor family which is characterized by a seven-helix transmembrane topology and by the ability to activate heterotrimeric G-proteins [1–3]. Since the first sequence of an opsin, bovine rhodopsin, was determined by conventional protein sequencing and cDNA sequencing, more than 1000 opsins have been identified [4,5]. In invertebrates, some photoreceptor opsins have been well-characterized, both in terms of their sequences and cellular functions [1]. When sequence alignments are made of the invertebrate and vertebrate opsins, it is evident that individual sequence may be quite different, but the tertiary structure appears conserved [5,6].

Insecticide resistance is a major obstacle to the control of vector-borne diseases [7–9]. Many studies have suggested that the insecticide resistance phenotype evolves rapidly based on the selection of major effect genes [10,11]. However, recent genome-wide transcription profiling indicated that a broader range of genes may be involved and that insecticide resistance may be more complex than previously considered [12–14].

As part of an ongoing study of insecticide resistance in Cx. pipiens pallens, we have employed suppression subtractive hybridization (SSH) and cDNA microarray to identify differentially expressed genes between deltamethrin-susceptible and -resistant strains of Cx. pipiens pallens. Opsin is one of only ten genes that showed over-expression in deltamethrin-resistant strain [15]. In the present study, we isolated and sequenced the complete sequence of this gene and
termed it NYD-OP7. We then characterized the expression pattern of NYD-OP7 gene in different developmental stages of Cx. pipiens pallens. In addition, we investigated the role of NYD-OP7 in deltamethrin resistance by transfecting this gene into the mosquito C6/36 cells and tried to determine if over-expression of NYD-OP7 can change the sensitivity of the cells to deltamethrin.

2. Materials and methods

2.1. Mosquito strains

The deltamethrin-susceptible strain of Cx. pipiens pallens was obtained from the Shanghai Insect Institute of the Chinese Academy of Sciences and was maintained in our laboratory. The mosquitoes had never been exposed to any insecticides and the median lethal concentrations (LC50) of deltamethrin (Roussel Uclaf, France) was 0.0008 mg/L. The strain was reared and bred at 25–27 °C in a 16 h light/8 h dark photoperiod. The deltamethrin-resistant strain was derived from the susceptible early fourth instar larvae by selection with deltamethrin for 10 generations until the resistant ratio (LC50 of filial generation/LC50 of parental generation) reached 400 [16].

2.2. Construction of cDNA library

Total RNA were extracted from 300 adult females of resistant strain using the RNeasy maxi kit (Qiagen, Germany) according to the manufacturer’s instructions. The integrity of total RNA was determined by denaturing agarose gel electrophoresis and the yield and purity of total RNA was estimated by spectrophotometry. The poly(A) mRNA was purified from total RNA with the PolyA-tract mRNA isolation system (Promega, USA), and 2 μg polyA mRNA was used for cDNA library construction with λExcell/NotI/EcoRI/CIP vector system (Pharmacia, USA) following the manufacturer’s protocols. The cDNA library contained approximately 5 × 104 clones and was amplified.

2.3. Cloning and sequencing of NYD-OP7 cDNA

To isolate the full-length of culex opsins gene, we used the standard SP6 and T7 library vector primers and the primers designed based on the sequence of the SSH fragment we reported previously (GenBank Accession No. BE247812). The sequence of the oligonucleotide primers was: forward 5′-GGCGTCTGGTTGACCCATTCT-3′; reverse 5′-CACG ATCGGGTTGTAGACAG-3′. PCR conditions were: initial denaturation at 94 °C for 5 min, followed by 32 cycles of 94 °C for 30 s, 60 °C for 30 s and 72 °C for 90 s with final 10 min extension at 72 °C. The PCR products were separated by 1% agarose gel electrophoresis and purified using a QIA quick Gel extraction kit (Qiagen, Germany). Products were then cloned into the pGEM-T easy vector (Promega, USA) and sequenced at Shanghai Invitrogen Biotechnology Co., Ltd. (Shanghai, China). Then the sequences of above three fragments were assembled to generate a putative full-length cDNA of opsins. A pair of PCR primers: forward 5′-TTAAGTCTGAAGTGATGTC-3′ and reverse 5′-CTAGCAAAAATGCAATC-3′ were designed for the amplification of full-length cDNA according to the assembled sequence of opsins. PCR was carried out using LA Taq polymerase (TaKaRa, Japan) using the following protocol: 94 °C for 5 min, followed by 32 cycles of 94 °C for 30 s, 60 °C for 30 s and 72 °C for 90 s with final 10 min extension at 72 °C. Then the PCR product was purified, cloned into the pGEM-T easy vector and sequenced.

2.4. Sequence alignment and phylogenetic tree

The standard protein/-protein BLAST sequence comparison (blastp; www.ncbi.nlm.nih.gov BLAST/) and PSI-BLAST programs were used to search for sequences in the GenBank and SWISS-PROT databases with similarities to the translated sequences of NYD-OP7 [17]. Sequence alignment and phylogenetic tree analysis using the neighbour-joining method was carried out by the CLUSTAL W program [18]. The opsins included in our analysis were: twelve sequences from An. gambiae genome [2], six sequences from Drosophila melanogaster, three sequences from Manduca sexta, six sequences from Papilio glaucus and two sequences from Aedes aegypti [6].

2.5. RNA extraction and cDNA synthesis

Total RNA were extracted from embryos, first, second, third and fourth instar larvae, pupae, and adults of both susceptible and resistant strains using the RNeasy mini kit (Qiagen, Germany) according to the manufacturer’s protocol, and contaminant genomic DNA was removed by DNase I treatment. cDNA was synthesized from 2 μg of total RNA with M-MLV reverse transcriptase (Promega, USA) and random oligonucleotide primers according to the manufacturer’s protocol.

2.6. Real-time PCR of NYD-OP7 in mosquito developmental stages

Aliquots of cDNA were amplified on an ABI PRISM 7300 (Applied Biosystems) using the SYBR green PCR kit (PE Applied Biosystems) with 25 μl reaction mixtures. The PCR mixture contained 1× SYBR PCR buffer, 20 mM MgCl2, 0.2 mM (each) dNTP, 0.2 μM primers, and 0.3 μl of Taq polymerase (2.5 U/μl). The primers used were: NYD-OP7: forward 5′-GGGCGCTTCGTGACTCTTT-3′, reverse 5′-GGTTCACGCACTGCGTCCT-3′; β-actin: forward 5′-GCTTCCCTGTCATCACTGG-3′, reverse 5′-GTTGTGGCGAACAGATCTC-3′. For amplification, the following program was employed: 40 cycles of 95 °C for 15 s, 60 °C for
1 min. A melting curve program was run immediately after the PCR program. Control measurement was done in the absence of reverse transcriptase to make sure that the level of genomic DNA contaminants was negligible.

Real-time PCR data were analyzed with a 7300 System SDS Software v1.2.1 (Applied Biosystems) to estimate transcript copy numbers for each sample. The raw threshold cycle (Ct) values were normalized against β-actin standard to obtain normalized Ct values, which were then used to calculate relative expression levels in the samples. The expression level of NYD-OP7 in susceptible strain egg was considered as background level, or 1. To verify reproducibility, the real-time PCR analysis was repeated twice in independent experiments, with three replicates for each sample.

2.7. Recombinant insect expression vector construction

The 1116 bp NYD-OP7 coding sequence, from the putative translational ATG to the end of the ORF, was amplified by PCR using the cDNA library as the template. Forward and reverse primers, containing at their ends, respectively, EcoRI and NotI restriction sites, were: forward 5'-CGGAATTCCTGAGCTTTCTGAGCC-3' and reverse 5'-ACTAGCGGCGCTGACTTAAACCTTCCTTG-3'. The PCR product was ligated into the pGEM-T easy vector. Then the recombinant plasmid was digested with SacII and NotI and the purified fragment was ligated into the SacII–NotI linearized insect expression vector pIE1-3 (Novagen, USA), resulting in the pIE1-3/OP7 construct. The integrity of the construct was verified by restriction enzyme digestion and sequencing. The pIE1-3 vector is designed for constitutive expression of recombinant protein from the baculovirus iel promoter in stably transfected insect cells [19]. To obtain stably transfected cells, a pIE1-3 recombinant should be cotransfected with pIE1-neo vector (Novagen, USA), which contains the Neo selectable marker under iel control.

2.8. Cell culture and transfection

Mosquito C6/36 cells were obtained from China Center for Type Culture Collection (CCTCC, China). The cells were maintained in Eagle's minimum essential media (EMEM) (Invitrogen, USA) supplemented with non-essential amino acids, fetal bovine serum (10%) (Sijiqing, China), and antibiotics. The cells were grown in a 5% CO2 humidified incubator at 28 °C. Transfection was performed using Cellfectin reagent (Invitrogen, USA) according to the manufacturer's protocol. Briefly, 4 × 10^5 cells/well were plated into a 6-well plate in growth medium without antibiotics and cultured overnight. Then the pIE1-3/OP7 construct and the pIE1-3 vector only, were each separately cotransfected with pIE1-neo construct at a ratio 4:1 into C6/36 cells according to the manufacturer's protocol. After 24 h, the cells were selected with 300 μg/ml G418 (Calbiochem, Germany) for 2 weeks and the medium was changed every 4 days. Following selection, G418-resistant colonies were isolated and expanded as individual clonal cell lines for subsequent analysis.

2.9. RT-PCR analysis

Total RNA was extracted from cells with TRIzol reagent (Invitrogen, USA) according to the manufacturer's protocol and the reverse transcript reaction was performed as above. The resulting cDNA preparation was subjected to PCR amplification using the forward primer specific to NYD-OP7 (5'-GTGGGCTTCTCGACTTTT-3') and the pIE1-3 vector specific HRAS reverse primer (5'-AAGATTAGCGAC GCTGCT-3'). Reaction conditions were as follows: 94 °C for 5 min; 30 cycles (25 cycles for β-actin) of 94 °C for 30 s, 58 °C for 30 s, 72 °C for 40 s; 5 min extension at 72 °C for 10 min. To confirm equal loading, PCR amplification of the β-actin gene was also done in parallel. The primers used for β-actin PCR amplification were: forward 5'-CACCAGGTT GTGATTG GTCGG-3', reverse 5'-CCACCAGATCCAGAC GGAGT-3'. The PCR products were analyzed by electrophoresis on 1% agarose gel containing ethidium bromide, and photographed under UV light.

2.10. Cell proliferation ([3H]-TdR) assay

[3H]Thymidine (1 mCi/ml, Shanghai Institute of Applied Physics, Chinese Academy of Sciences) was used to determine the proliferation activity of stable C6/36 transfected cells. Cells in exponential growth phase were trypsinized and resuspended in growth media. Cells were plated in 96-well plates (5000 cells/well) and cultured 4 h to allow for cell attachment. Cells were then treated with various concentrations of deltamethrin (0, 10, 15, 20, 25, 30, 40, 80, and 120 μM). After incubation for 72 h, [3H]thymidine (0.5 μCi/well) was added, and cultures were incubated for a further 16 h. Incorporation of [3H]thymidine into DNA was measured using an Wallac 1414 liquid scintillation counter (WALLAC, Finland). Data were gathered as counts/min, representing the fraction of cells that contained newly synthesized DNA. Inhibition of proliferation was determined by comparing the level of [3H]thymidine uptake with controls. Three independent experiments were done in quadruplicate wells. Deltamethrin was dissolved in DMSO and the final concentration of DMSO in the medium is 0.1%.

2.11. Statistics

Data were analyzed by Student’s t test. The level of significance was set at p < 0.05.

3. Results

3.1. Isolation full-length of the Cx. pipiens pallens NYD-OP7 gene

To isolate opsin gene from the Cx. pipiens pallens, we performed a RT-PCR screen of cDNA library using the
standard SP6 and T7 library vector primers and primers designed based on the sequences we reported previously (GenBank Accession No. BE247812). The sequences we got were assembled to generate a putative full-length cDNA of opsin. Then a pair of PCR primers was designed for the amplification of full-length cDNA. Sequencing of the amplified transcript led to the identification of a novel opsin gene which we termed NYD-OP7 (GenBank Accession No. AY749413). The NYD-OP7 cDNA has an open reading frame of 1116 bp, encoding 371 amino acids (Fig. 1).

3.2. Sequence and phylogenetic analysis

Alignment of the deduced NYD-OP7 amino acid sequence with Ae. aegypti and An. gambiae opsins revealed sequence identities of 87% with Ops1, 86% with GPRop1-4, 83% with Ops2, 79% with GPRop6, and 63% with GPRop5 (Fig. 2). The newly isolated opsin displays typical hallmarks of rhodopsins, such as a highly conserved lysine residue in the seventh transmembrane region, to which the retinal chromophore is attached; putative phosphorylation

---

Fig. 1. The nucleotide and deduced amino acid sequences of Cx. pipiens pallens NYD-OP7. The deduced amino acid sequence is presented below the nucleotide sequence in single letter code. The predicted seven transmembrane segments are underlined. The putative polyadenylation signals in the 3'-untranslated region are also underlined (thick lines). The two cysteine residues in the first and second extracellular loops are boxed. The 317th K-residue is shown in bold. Potential sequences for G-protein binding sites, DRY and QAKKMNV, as mentioned in Section 3, are indicated by open boxes. Potential glycosylation sites in the N terminals and HEE motif are indicated by shaded boxes. The asterisk indicates the stop codon.
sites (Ser, Thr) near the C-terminus; putative N-glycosylation sites (Asn-Xaa-Ser/Thr) near the N-terminus, and two cysteine residues (Cys-121 and Cys-198) in the first and second extracellular loop, which may form a disulfide bond. NYD-OP7 contains a DRY sequence at the junction of helix III and the second intracellular loop that is very similar to the ERY sequence found in bovine rhodopsin. This sequence contains a charged pair that is required for transducin activation [20,21]. Although dissimilar from the vertebrate opsins, the sequence QAKKMNV found in the N-terminal half of cytoplasmic loop 3 is very highly conserved among the invertebrate opsins and is located within a region of the protein that is also required for transducin activation [22]. Most insect rhodopsins have a His-Glu-Lys (HEK) motif at the beginning of a conserved region in the third cytoplasmatic loop, and it is implicated in rhodopsin–G-protein interaction. Interestingly, the HEK motif is modified to His-Glu-Glu in NYD-OP7. Other insect rhodopsins in which the lysine residue of the HEK motif is substituted include two *M. sexta* rhodopsins [23], two *An. gambiae* rhodopsins [2], one *calliphora* rhodopsin [24] and one *Ae. aegypti* opsin [6].

Fig. 3 shows the molecular phylogenetic relationship between NYD-OP7 and some other insect opsins. Results showed that the NYD-OP7 sequence is most closely related to the *Ae. aegypti* OpsI, and it clearly belongs to a group comprising the known arthropod opsins.

### 3.3. Expression profile of the NYD-OP7 gene

The expression of the NYD-OP7 gene was analyzed by real-time PCR amplification of the corresponding mRNA from different *Cx. pipiens pallens* developmental stages. As shown in Fig. 4, NYD-OP7 is expressed at significantly high levels in pupae and adults, whereas expression is very low in the egg, first, second, third and fourth instar larval stages. Interestingly, in pupae and adults, the expression of NYD-OP7 is significantly higher in resistant strain than in susceptible strain.

### 3.4. NYD-OP7 expression and deltamethrin sensitivity analysis of pIE1-3/OP7 transfected C6/36 cells

To elucidate the contribution of the NYD-OP7 gene product to deltamethrin resistance, the mosquito C6/36 cells were transfected with the pIE1-3/OP7 insect expression vector. After G418 selection, one stably transfected NYD-OP7 subclone was obtained. Analysis of this transfectant demonstrates a higher expression of NYD-OP7 gene as compared with the empty vector transfectant control (Fig. 5). These subclones did not show any growth morphological changes and then they were examined for deltamethrin sensitivity by [3H]-TdR assay. As shown in Fig. 6, NYD-OP7 transfected C6/36 cells are relatively resistant to deltamethrin at the concentrations of 15, 20 and 25 µM (*p* < 0.05). Meanwhile, a slight increase of cell viability was also observed at 30 µM deltamethrin but without statistical significance (*p* = 0.0744).

### 4. Discussion

Opsins are membrane proteins with molecular masses of 30–50 kDa that are related to the protein moiety of the photoreceptive molecule rhodopsin; typically they are considered to act as light sensors in animals, but there are still many unanswered questions concerning the functions of different opsins [5,25–28]. Opin gene has been cloned in various invertebrates, such as *D. melanogaster* [29],...
An. gambiae [2], Ae. aegypti [6], M. sexta [23], P. glaucus [30] and aphid [31].

In this experiment, we studied a novel cDNA encoding opsin (NYD-OP7) isolated from Cx. pipiens pallens. Considering the amino acid sequence of this gene, the translated protein is expected to have a seven-transmembrane structure, and it has all the conserved residues required for a functional opsin protein. This feature and the amino acid identity of NYD-OP7 with known opsins suggest that it encodes a functional opsin protein. We have compared the amino acid sequence of the novel opsin with those of other insect opsins. The phylogenetic analysis indicates that NYD-OP7 is a member of the group including Ae. aegypti Ops1 and An. gambiae GPRop1-3. Real-time PCR analysis revealed the expression of NYD-OP7 at all developmental stages both in deltamethrin-susceptible and -resistant...
strains and it appeared that \textit{NYD-OP7} mRNA levels are significantly lower in \textit{C. pipiens pallens} embryos and larvae than in older developmental stages. Whether the expression pattern reflects a function for \textit{NYD-OP7} in eye development remains to be seen.

In previous study, we identified 16 genes that were differentially transcribed among the deltamethrin-susceptible and -resistant strains of \textit{C. pipiens pallens} and as shown by microarray, one \textit{opsin} gene is 3.06-fold overexpressed in the resistant strain [15]. In this paper, we also found that in pupae and adults, \textit{NYD-OP7} is overexpressed in resistant strain than in susceptible strain, which shows that \textit{NYD-OP7} may play a role in the deltamethrin resistance phenotype. However, it remains possible that \textit{NYD-OP7} over-expression is an indirect consequence of acquired deltamethrin resistance. To determine whether \textit{NYD-OP7} directly participates in the establishment of the deltamethrin resistance phenotype, we transfected this gene into the mosquito \textit{C6/36} cells and to determine if over-expression of \textit{NYD-OP7} can change the deltamethrin sensitivity in \textit{C6/36} cells. The results showed that \textit{NYD-OP7} transfected cells are relatively resistant to deltamethrin at low drug concentrations which suggests that \textit{NYD-OP7} is directly involved in the development of deltamethrin resistance.

Previously, by using a small-scale microarray representing all \textit{Drosophila} P450 genes, Daborn et al. and Le Goff et al. suggested that resistance to DDT in the field is monogenic and is due to the over-expression of a single P450 gene, \textit{Cyp6g1} [10,32]. However, by using a detox chip and a genome-wide microarray of \textit{An. gambiae}, David et al. and Vontas et al. found that many genes that belong to families not usually associated with insecticide resistance (such as peptidases, sodium/calcium exchangers and genes implicated in lipid and carbohydrate metabolism) were over-transcribed in the resistant strain [13,14]. Meanwhile, by using genome-wide microarray, Pedra et al. demonstrated that DDT-metabolic resistance in \textit{Drosophila} is associated with over-expression of detoxification related genes but also over-expression of ion transport, signal transduction, RNA transcription, and lipid and sugar metabolism pathways[12]. Interestingly, they also reported that \textit{Drosophila} UV-sensitive opsin (Rh3 and Rh4) and Blue-sensitive opsin (NinaE) were overexpressed in the DDT-resistant strain, suggesting that \textit{opsin} gene may play some role in DDT-resistance [12].
In the case of NYD-OP7, we hypothesized that as one member of G-protein coupled receptor family, NYD-OP7 may contribute to the phenotype of insecticide resistance by activating downstream G-protein and signal transduction pathway. It is also possible that the deltamethrin-resistant phenotype requires the coordinated expression of many different genes so that over-expression of individual gene is necessary, but maybe not sufficient, to generate a high resistance phenotype. It should also be mentioned that because of lack of a proper antibody, we cannot do western blot to confirm the over-expression of NYD-OP7 in stably transfected cells at the protein level. So there is a possibility that the expression of NYD-OP7 at the protein level is not as high as the mRNA level which may be the reason for why high NYD-OP7 transcriptional level expression only causes a modest increase in resistance. In a recent study, we have proved that the cotransfection of two genes (NYD-Tr and NYD-Ch) into C6/36 cells can induce deltamethrin resistance [33]. All these data suggest that insecticide resistance is more complex than we considered and we believe that further investigation should be done to fully elucidate the role that NYD-OP7 has in insecticide resistance.

In summary, we have reported the cloning and identification of a novel opsin gene (NYD-OP7) from Cx. pipiens pallens. Based on the expression profiles and previous SSH and microarray results, this gene appears to have some role in deltamethrin resistance. Over-expression of NYD-OP7 in mosquito C6/36 cells can confer moderate resistance to
deltamethrin, suggesting that NYD-OP7 directly participates in the development of deltamethrin resistance and the precise definition of its role awaits further investigation.

Acknowledgments

We thank Dr. Grace Jones (Department of Biology, University of Kentucky, USA) for his generous gift of plasmid pIE1-neo. We thank Mrs. Jianmei Xu for her help about data analysis. We also thank the anonymous reviewers for their helpful comments and suggestions. This work was supported by the National Natural Science Foundation of China (Nos. 30170835 and 30371255), the Natural Science Foundation of Jiangsu Province (No. BK2004147) and the Academic Natural Science Foundation of Jiangsu Province (No. 03KJB180082).

References
