Biochemical Characterization of Acetamiprid Resistance in Laboratory-Bred Population of Aedes aegypti L. Larvae

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ABSTRACT
The constant rise in cases of Zika, Dengue and Chikungunya worldwide has made control of Aedes aegypti a principal concern. The most recommended plan to control mosquito-borne diseases primarily lies on vector management and disturbing their disease-transmission cycle. Wide-ranging use of different classes of organic insecticides for mosquito control has led to the development of high levels of resistance making them less operative at safe dosages imposing us to explore novel insecticides. Present study investigates the bioefficacy of a neonicotinoid, acetamiprid on the Ae. aegypti larvae, development of resistance after subjecting acetamiprid selection pressure for 10 successive generations and biochemical characterization of the resistance developed. Acetamiprid exposure of the parent population of Ae. aegypti early fourth instars resulted in respective LC50 and LC90 values of 0.188 ppm and 1.315 ppm. Selection with acetamiprid for 10 successive generations (ACSF-10) reduced its efficacy by 20-fold. Involvement of four enzymes; alpha-esterases, beta-esterases, glutathione-S-transferases and acetylcholinesterases in development of acetamiprid resistance was investigated to uncover mode of action of acetamiprid. An elevation of 1.4-fold and 2.1-fold was observed in alpha-esterases and beta-esterases activity in ACSF-10 as compared to ACSF-5. However, activity of glutathione-S-transferases decreased in ACSF-5 which rose to 12-fold in ACSF-10. Similarly, the activity of acetylcholinesterases was found to be much higher in resistant generations as compared to the parental strains. The results indicated individual/synergistic contribution of different enzymes leading to acetamiprid detoxification. Further research is being conducted to identify the role of target site mutations in resistance development.

Keywords: Aedes aegypti, acetamiprid, esterases, glutathione-S-transferase, acetylcholinesterases

1. INTRODUCTION
The WHO global strategy for dengue prevention and control by 2020 is based on the fact that almost half of the world population resides in dengue-prone area; resulting in 50 to 100 million estimated annual dengue infections. Currently, approximately 75% population of Asia-Pacific region is exposed to dengue. The actual number of cases of dengue is even worse, because of severe underreporting and misclassification [1, 2]. In addition, most countries in Southeast Asia with endemic malaria are experiencing increased Aedes-borne diseases due to ecological changes arising from poorly controlled population movement and extensive exploitation of natural environments.

The prevention of dengue transmission primarily depends on reduction of the human-vector contact using residual chemical compounds; while dengue infection can be kept under check by accurate diagnosis and prompt effective treatment [3, 4]. For decades, DDT was used in mosquito vector control programs as an inter-domiciliary spray; which was gradually discontinued because of social and environmental concerns. The organochlorines were then replaced with organophosphates followed by synthetic pyrethroids. Since then, pyrethroids have been used extensively for the insecticide treatment of bed nets (ITN) and as indoor residual sprays (IRS) in many parts of the country. However, repeated contact with these insecticides has led, in some cases, to high levels of resistance in vector populations. The increased development of mosquito resistance to pyrethroids is of particular concern for many integrated mosquito control programs that utilize insecticides for vector control [5].

Despite extensive research in the field, management of insecticide resistance remains the major challenge in mosquito control leading to the need of developing strategies which could ensure long-term efficacy of toxicants and; delay or prevent the development of resistance. Identification and introduction of new insecticides in the fields with mosquito control potential and novel mode of action has become a major task in front of the researchers. Neonicotinoids are among such
insecticides which are being explored by the researchers as mosquito control agents. These are the synthetic analogues of natural insecticide nicotine, an active component of tobacco and are considered agonist neurotoxins targeting various sensitive sites in insect nervous system [6-8]. Initially discovered in late 1980s, these are now used for various purposes ranging from plant protection, biocides, and veterinary products to the control of invertebrate pests in fish farming [9]. Reports reveal high pest control efficacy of neonicotinoids as they have been effective at much lower concentration and quantity in comparison to the traditionally used insecticides [10]. In addition, their low mammalian toxicity, negligible cross resistance and unique mode of action suggest the possible use of neonicotinoids as ideal alternatives to various synthetic organic insecticides [11]. The newest major group of neonicotinoids includes Acetamiprid, Imidaclorpid, Clothianidin, Dinotefuran, Nitenpyram, Thiacloprid and Thiamethoxam. The half-life and residual nature of these compounds vary widely from a few months to much longer durations [12-14]. These chemicals have been reported effective against a variety of economically important crop pests including whitefly, aphids, leafhoppers, wireworms, plant hoppers and mealybugs [15] and also showed efficacy against mosquito larvae as compared to other classes [16].

Acetamiprid, an odourless neonicotinoid insecticide manufactured under the trade names Assail, and Chipco by Aventis CropScience India, is a nicotinic agonist that reacts with nicotinic acetylcholine receptors (nAChRs) [17] located in the post-synaptic neural dendrites of brain, spinal cord, ganglia and muscular junctions; and mediates fast cholinergic transmission, a unique mode of action [18,19]. Acetamiprid is also known to possess both acute contact and stomach poisoning activity [18, 20]. This insecticide, therefore, has become an important component of integrated pest management since it was registered commercially in 1995.

In this study, the larvae of *Ae. aegypti* were exposed to acetamiprid for 10 successive generations to assess the probable larval resistance development. Thereafter, we conducted a series of biochemical assays integrated with dose-mortality bioassays for detection of resistance and to outline the underlined mechanisms involved in neonicotinoid resistance in *Ae. aegypti* L.

### 1.1. Materials and Methods

#### 1.1.1. Culture of Aedes aegypti L.

Dengue fever mosquitoes, *Ae. aegypti*, at adult and larval stages, were obtained from ICGEB (International Centre for Genetic Engineering and Biotechnology), New Delhi, India. The colony was maintained in an insect rearing unit of Insect Pest and Vector Laboratory, Acharya Narendra Dev College, University of Delhi, India under controlled conditions of 28 °C ± 1 °C, 80% ± 5% RH, 14 hr of light and 10 hr of darkness [21]. Adults kept in clothed cages were fed on sugary juice of water-soaked raisins while female *Ae. aegypti* were provided with occasional blood meals for egg maturation. Eggs collected in an ovitrap were transferred into the enamel trays filled with at least 1.5 – 2.0 L of dechlorinated water. The hatched larvae were fed on powdered dog biscuits and yeast in a ratio of 3:1 till they transformed into pupae [22, 23]. Trays were kept clean and water was changed every day to prevent formation of scum on the water surface. The pupae collected on regular basis were kept in clothed cages for adult emergence.

### 1.1.2. Insecticide susceptibility test

The parent strain (PS) of *Ae. aegypti* L. was exposed to acetamiprid to estimate its larvicidal efficacy [22, 23]. The larvae were then subjected to selection pressure of acetamiprid at LC₀ level till 10 successive generations to assess the development of resistance. The acetamiprid-selected 5th generation (ACSF-5) and 10th generation (ACSF-10) were employed in the current experiments.

#### 1.1.3. Biochemical assay

The larvae of parent generation of *Aedes aegypti* L. and those obtained after subject to selection pressure with acetamiprid for successive 5 and 10 generations were characterized biochemically. The standard method to identify the probable biochemical mechanism involved in insecticides resistance provided by WHO [24] was used with few minor changes. Individual larva was homogenized in 200µL of autoclaved water on ice. The 25µl × 2 (replicate) of homogenized mixture was segregated for the acetylcholinesterase assay and the remaining homogenate was spun in a refrigerated microfuge at maximum speed for 30 s. Supernatant separated after centrifugation was used for estimation of proteins, alpha-esterases, beta-esterases and glutathione-s-transferases.

#### 1.1.3.1. Protein estimation

The 10µL × 2 (replicate) supernatant from all the larval strains was pipetted in the microtiter plate. The 300µL of BIORAD protein reagent was added to the homogenate. A blank was run simultaneously replacing homogenate with the water. The absorbance of the mixture was read at 570 nm with the help of ELISA plate reader. The standard curve was plotted and the concentration of the protein of individual larvae from PS, ACSF-5 and ACSF-10 was calculated in mg/mL and in n moles.
1.3.2. Non-specific esterases estimation

The 10 µL × 2 (replicate) of the supernatant of each strain was pipetted in the microtiter plate. The 200 µL of Alpha/Beta naphthyl acetate was added in each titer and incubated for 15 mins. Visual changes were interpreted by adding 50 µL of fast blue stain (fresly prepared) in each well. The absorbance was measured at 570 nm to evaluate the elevation in the level of esterases in each strain.

1.3.3. Glutathione-S-transferases bioassay

The 20 µL × 2 (replicate) of the supernatant of each strain was pipetted in the microtiter plate to which 50 µL of 2mM GSH (Reduced glutathione) and 50 µL of 1mM CDNB (1-Chloro-2,4-dinitrobenzene) was added. The ELISA plates were then read continuously for 5 mins at 340 nm to study the enzyme kinetics.

1.3.4. Acetylcholinesterase’s inhibition bioassay

The 25 µL × 2 (replicate) of crude insect homogenate of each strain was placed in separate wells of a microtiter plate. The ASCHI (Acetylthiocholine iodide) solution was divided into two 10mL aliquots. One aliquot was added with 20 µL of 0.1M propoxur. The 145 µL of 1% solution of Triton X-100 was added to the homogenate to solubilize acetylcholinesterases, followed by addition of 10 µL of DTNB (Dithiobis 2-nitrobenzoic acid) solution. Thereafter, 25 µL of ASCHI+propoxur was added to the other replicate. The reaction mixture was then incubated for 1 hr after which end point reading was taken by reading the absorbance at 405 nm.

1.2. Our Contribution

This paper discusses the possible use of acetamiprid for the control of dengue vector, Ae. aegypti. The development of acetamiprid resistance in the vector and characterization of the underlying biochemical and molecular mechanisms that may potentially play a role in the resistance are elucidated. The variations in the enzyme levels in the resistant Ae. aegypti population are determined while the role of target site mutations in resistance development is under exploration. These investigations could help in formulation of resistance management strategies to eliminate heterozygotes and resistant homozygotes, the efficacy of which can only be established after field trials.

1.3. Paper Structure

The rest of the paper is organized as follows. Section 2 explains the insecticide resistance, its mechanism the biochemical characterization of resistance. The acetamiprid susceptibility in Ae. aegypti and development of resistance are presented and analysed. This section tells about the correlation of three major classes of detoxifying enzymes: Non-specific esterases, Glutathione-S-transferases and acetylcholinesterase with increase in level of acetamiprid resistance in Ae. aegypti. These results are discussed with that conducted elsewhere. Finally, Section 3 concludes the paper and presents direction for future research.

2. RESULTS AND DISCUSSION

Employment of chemical insecticides is the prime strategy in control of mosquito vectors. Resistance to different classes of insecticides; organochlorines, organophosphates and pyrethroids has been reported in different mosquito vectors which may be attributed to constant selection pressure of insecticide used in agriculture as well as in residential areas [25-27]. The forced paths available for the supervision of insecticide resistance have created an urgent necessity to identify novel insecticides for effective control of mosquito species which have developed multiple resistances to the insecticides belonging to different classes [28]. Present study involves, neonicotinoids, a class of neurotoxic insecticides with highly systemic toxicity and long-term persistence. They bind to the nicotinic acetylcholine receptors in the nervous system and block the route of nerve impulses, the binding potency conferred by a particular confirmation of the molecule [29].

The insecticidal potential of different neonicotinoids has been reported earlier [28, 30]. Most of these studies are focused on the toxicity assays of imidacloprid. In the present study, an attempt was made to study the probable use of a neonicotinoid, acetamiprid, for Ae. aegypti management. The early fourth instars of Ae. aegypti were tested for their susceptibility and tolerance levels against the insecticide. The parent strains and those subjected to continuous selection pressure of acetamiprid (ACSF-5 and ACSF-10) were assessed for elevation in the xenobiotic metabolizing enzymes. The study displayed high susceptibility of the Ae. aegypti larva to the acetamiprid. Exposure of the early fourth instars of parent generation resulted in LC50 and LC90 values of 0.188 ppm and 1.31 ppm, respectively. Selection of these larvae with acetamiprid at LC50 levels resulted in the development of considerable level of resistance. The early fourth instars of Ae. aegypti subjected to continuous selection pressure of acetamiprid for 10 successive generation developed 19.74-fold resistance to acetamiprid (Table 1). The respective LC50 and LC90 values recorded in the parent strain rose to 3.711 and 10.088 ppm in ACSF-10. Similar results were reported by Urugayala et al. [28] in an Indian strain of Ae. aegypti susceptible to organochlorines, organophosphates and pyrethroids. They obtained a little higher LC50 value (0.558 ppm) on larval bioassay with imidacloprid.
Identification of biochemical-based resistance mechanisms using microplate enzyme assays in a single mosquito is more informative and could be of value in early detection of insecticide resistance in field population [13]. Target site insensitive AChE assay and detoxification enzymes, a- and β-esterases and GST assays were conducted to identify the probable biochemical mechanism involved in acetamiprid resistance.

The larval selection with acetamiprid for 10 successive generations resulted in accountable biochemical alterations in of *Ae. aegypti* larvae. Alpha-esterase activity increased by 1.27-fold in ACSF-10 as compared to the susceptible generation. However, the results showed an insignificantly reduced esterase activity by 0.93-fold in ACSF-5 as compared to the parent strain (PS) (Table 2; Figure 1). The elevated levels of beta-esterase activity in *Ae. aegypti* was evidenced by the appearance of pink colour in the solution indicating the formation of 2-naphthol.

Beta-esterase activity oscillated a lot between the generations tested. Subjection of acetamiprid selection pressure for 5 generations decreased beta-esterase activity by 0.63-fold which later increased significantly by 1.33-fold in ACSF-10 (Table 2). The esterase activity in parent strain and ACSF-5 did not differ significantly (p>0.05), whereas the activity was statistically different from that observed in ACSF-10 (Figure 2). Similar results were reported by Hemingway [27] who found quantitative increase of esterases in malathion-resistant *An. stephensi* from Pakistan.

The Glutathione-S-transferase assay is based on the conjugation of reduced glutathione with CDNB (3,4-dichloro-4-nitrophenyl (1-glutathione) in presence of Glutathione-S-transferase as visual interpretation is not possible due to lack of any colour appearance in this reaction. The conjugation is calculated by noting the enzyme kinetics through scanning the activity continuously for at least 5 mins, which is then calculated in nmoles/min/mL. The results showed that ACSF-5 exhibited significantly reduced GST activity as compared to PS (p<0.05). However, the activity shot up in ACSF-10 by 1.50 (Table 2; Figure 3). In addition, an increased AChE activity was observed in the average population of ACSF-5 and ACSF-10; resulting in 0.07% decreased AChE inhibition in ACSF-5 which further decreased to 2.94% in ACSF-10. Safi et al. [31] also reported metabolic-based

### Table 1 Larval LC$_{50}$ and LC$_{90}$ (in ppm) of parent susceptible (PS) and acetamiprid-larval selected strain of *Aedes aegypti* selected for 10 successive generations

| Strain  | LC$_{50}$ (ppm) | LC$_{90}$ (ppm) | Heterogeneity $\chi^2$ (DF) | Slope ± SEM | RR |
|---------|----------------|----------------|-----------------------------|-------------|----|
| PS      | 0.188 (0.119-0.248)** | 1.315 (0.759-5.241)** | 3.395 (3) | 1.517 ± 0.295 | ----- |
| ACSF-5* | 1.659 (0.727-2.065) | 4.509 (3.284-20.236) | 2.285 (3) | 2.951 ± 0.146 | 3.427 |
| ACSF-10 | 3.711 (2.915-4.356) | 10.888 (7.120-32.184) | 0.959 (6) | 2.975 ± 0.119 | 19.74 |

ACSF* - Acetamiprid selected filial RR - Resistance Ratio
**Figures in parentheses represent lower and upper 95% Confidence Limit
LC$_{50}$ - Lethal Concentration at which 50% larvae are killed, LC$_{90}$ - Lethal Concentration at which 90% larvae are killed; SEM=Standard Error of Mean, DF = degree of freedom

### Table 2 Mean Protein, α- and β-esterase, GST activities and % AChE inhibition in laboratory-bred population of *Aedes aegypti* L. after exposure to acetamiprid for 10 successive generations

| Strain  | Protein Concentration (nmoles) ± SEM* | α-Esterase (nmoles/min/mg protein) ± SEM* | β-Esterase (nmoles/min/mg protein) ± SEM* | GST activity (nmoles/min/ml) ± SEM* | % AChE inhibition ± SEM* |
|---------|--------------------------------------|---------------------------------------------|---------------------------------------------|----------------------------------|------------------------|
| PS      | 8.06 ± 0.27 a                         | 1.32 ± 0.05 a$_1$                          | 2.23 ± 0.22 a$_2$                          | 1.80 ± 0.18 b$_3$                | 62.00 ± 2.95 a$_4$     |
| ACSF-5* | 7.95 ± 0.27 a                         | 1.29 ± 0.03 a$_1$                          | 1.40 ± 0.42 a$_2$                          | 0.23 ± 0.02 c$_3$                | 61.93 ± 2.33 a$_4$     |
| ACSF-10 | 8.30 ± 0.29 a                         | 1.77 ± 0.05 b$_1$                          | 2.98 ± 0.17 b$_2$                          | 2.71 ± 0.19 a$_3$                | 59.06 ± 4.12 a$_4$     |

*Figures in each column followed by different letters are significantly different p < 0.05, one-way ANOVA followed by Tukey’s all pair wise multiple Comparison test; SEM: Standard error of Mean
mechanisms, including esterases, P450s and glutathione S-transferase (GSTs) combined with insensitive AChE in An. stephensi from Kunar and Nangarhar provinces of Afghanistan. They further stated that the high level of resistance was found in the Nangarhar population compared to the Kunar population due to selection of different pesticides in agriculture.

The results indicate that the neonicotinoid resistance in Ae. aegypti larvae can be ascribed to the enhanced metabolism by detoxification enzymes; and elevated levels of esterases and glutathione-S-transferases (Table 2; Figure 4). It is suggested that combined or any one of these mechanisms may possibly be involved in acetamiprid resistance development in Ae. aegypti. These are well known and established mechanisms for resistance to pyrethroids and OPs in mosquitoes [26]. The involvement of P450 mediated monooxygenases, elevated non-specific esterases, and reduced sensitivity of sodium ion channels in insects has been reported as common insecticide resistance strategy against pyrethroids [32-35]. Moreover, increased level of glutathione-S-transferases (GSTs) has been associated with conferring pyrethroid inhibition in different mosquito species; Ae. aegypti, An. gambiae and An. dirus B [36].
The elevated GSTs have been found to bind to molecules of various pyrethroid insecticides compromising effectiveness and toxicity by a sequestering mechanism. Although the spread of pyrethroid resistance has increased in disease vectors worldwide, the actual operational impact of resistance in control of disease vectors and transmission remains limited.

Though the results suggest differential involvement of metabolic detoxification, esterases, glutathione-S-transferases and AChE inhibition in the development of acetamiprid resistance in *Ae. aegypti*; yet increase in insecticide resistance is a multidimensional and dynamic process that depends upon many factors. The increase in the frequency of resistant *Ae. aegypti* population demands the use of insecticides with different modes of action.

3. CONCLUSION

In the present study, susceptible population of *Ae. aegypti* L. was subjected to acetamiprid selection pressure and resistance was developed. The results suggest the probable role of metabolic detoxifying enzymes, such as elevated esterase levels and glutathione-S-transferases in the development of resistance. The results also indicate individual or synergistic contribution of different enzymes leading to acetamiprid detoxification. However, increase in insecticide resistance is a multifaceted and vigorous process and depends upon many factors. The increase in the frequency of resistant *Ae. aegypti* population demands the proper insecticide resistance management. The investigations could help in formulation of resistance management strategies to eliminate mosquito as well as in development of new insecticides from the already existing one.

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