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The role of gene splicing, gene amplification and regulation in mosquito insecticide resistance

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The primary routes of insecticide resistance in all insects are alterations in the insecticide target sites or changes in the rate at which the insecticide is detoxified. Three enzyme systems, glutathione S-transferases, esterases and monooxygenases, are involved in the detoxification of the four major insecticide classes. These enzymes act by rapidly metabolizing the insecticide to non-toxic products, or by rapidly binding and very slowly turning over the insecticide (sequestration). In Culex mosquitoes, the most common organophosphate insecticide resistance mechanism is caused by co-amplification of two The amplified esterases are differentially regulated, with three times more $Est\beta 2^1$ being produced than Estα2¹. Cis-acting regulatory sequences associated with these esterases are under investigation. All the amplified esterases in different Culex species act through sequestration. The rates at which they bind with insecticides are more rapid than those for their non-amplified counterparts in the insecticide-susceptible insects. In contrast, esterase-based organophosphate resistance in Anopheles is invariably based on changes in substrate specificities and increased turnover rates of a small subset of insecticides. The up-regulation of both glutathione S-transferases and monooxygenases in resistant mosquitoes is due to the effects of a single major gene in each case. The products of these major genes upregulate a broad range of enzymes. The diversity of glutathione S-transferases produced by Anopheles mosquitoes is increased by the splicing of different 5' ends of genes, with a single 3' end, within one class of this enzyme family. The trans-acting regulatory factors responsible for the up-regulation of both the monooxygenase and glutathione S-transferases still need to be identified, but the recent development of molecular tools for positional cloning in *Anopheles gambiae* now makes this possible.

Keywords: mosquitoes; insecticide; gene amplification; gene splicing; *Anopheles*; *Culex*

1. INTRODUCTION

Major mechanisms of insecticide resistance in insects involve either mutation within the target site of the insecticide, or an alteration in the rate of insecticide detoxification. The enzymes involved in this detoxification may be quantitatively and/or qualitatively altered. There are three enzyme groups, esterases, glutathione Stransferases and monooxygenases, involved in metabolic resistance to the four major groups of insecticides. Esterase-based resistance has been reported from more than 30 different medical, veterinary or agricultural insect pests (Hemingway & Karunaratne 1998). mosquitoes it is the primary mechanism for organophosphorus (OP) insecticide resistance (Bisset et al. 1991; Herath et al. 1987; Karunaratne et al. 1993), and in some cases a secondary mechanism for carbamate resistance (Peiris & Hemingway 1993). Esterases produce a broad spectrum of resistance in many Culex species, but in Anopheles esterase-based resistance is usually specific to the OP malathion (Hemingway 1983, 1985; Herath et al. 1987).

Glutathione S-transferases in mosquitoes commonly confer resistance to the organochlorine insecticide DDT (Prapanthadara et al. 1993, 1996), and can act as a secondary OP resistance mechanism (Hemingway et al. 1991). In house flies, their role in OP resistance is more widely documented (Clark et al. 1984, 1986). DDT resistance in mosquitoes has generally been attributed to a single major-gene effect (Davidson 1963, 1956), although multigenic effects have been suggested in some instances (Lines & Nassor 1991). Reports of monooxygenase-based resistance are relatively rare in mosquitoes, and many of these are based on synergistic effects with piperonyl butoxide, which is not absolutely diagnostic. Pyrethroid resistance in Anopheles gambiae in East and West Africa appears to be linked to increased monooxygenase titres, in the latter case combined with an altered target-site mechanism (Brogdon et al. 1997). OP resistance in A. subpictus, a vector of malaria in Sri Lanka, is also linked to increased monooxygenase titres and higher insecticide metabolic rates (Hemingway et al. 1991; Martinez-Torres et al. 1998).

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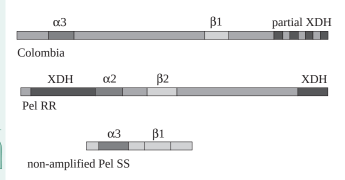


Figure 1. Structure of the amplicons associated with insecticide resistance in *Culex quinquefasciatus* strains from Colombia and Sri Lanka (PelRR) compared with the non-amplified esterase gene arrangement in the insecticide-susceptible strain PelSS. The $\text{Est}\alpha 2^1/\beta 2^1$ amplicon also has a complete gene with high homology to xanthine dehydrogenase (XDH) (Coleman & Hemingway 1997).

2. THE MOLECULAR BASIS OF METABOLIC RESISTANCE IN MOSQUITOES

(a) Gene amplification

The development of resistance to xenobiotics by amplification of the genes involved in their detoxication is common in several organisms. Gene amplification in the insecticide-resistant TEMR strain of the mosquito Culex quinquefasciatus was first shown for the Estβ1¹ esterase. It was originally estimated that there were up to 250 copies of this esterase gene per cell in resistant insects (Mouches et al. 1990), but this estimate has recently been revised to approximately 20 copies, i.e. similar to the estimates for Myzus esterases (Guillemaud et al. 1997). The Est β l genes are clustered between the centromere and the apex of chromosome II (Nance et al. 1990), and are inherited in a pseudo-monofactorial manner (Peiris & Hemingway 1993). The most common amplified esterases in *Culex* are Est $\alpha 2^1$ and Est $\beta 2^1$, which occur in ca. 90% of all the OPinsecticide-resistant C. quinquefasciatus strains analysed (Hemingway & Karunaratne 1998). The TEMR Estβl¹ and common Estβ2¹ from numerous strains have 97% identity at the amino-acid level. The high identity suggests that the Est \betas are an allelic series from a single locus. The approximately 47% deduced amino-acid Est $\alpha 2^1$ has homology with all the Estβs. This level of homology, along with conserved intron-exon boundaries and the close proximity (1.7 kb) of the two genes in a head-to-head arrangement in the susceptible insects (Vaughan et al. 1997), suggests that the two genes arose through an ancient duplication. In resistant insects Estα2¹ and Estβ2¹ are also in a head-to-head arrangement, but they are ca. 2.7 kb apart (Vaughan et al. 1997). The increase in the intergenic DNA between the two genes is accounted for by two large (ca. 500 bp) and one small insertion in the resistant insects. These insertions introduce DNA motifs that have high homologies to BARBIE boxes, ARE elements and Zeste elements (Vaughan et al. 1997; N. Hawkes, unpublished data). The structure of the Estα2¹/ESTβ2¹ amplicon and the related EstBl amplicons compared with the esterase gene arrangements in susceptible insects are given in figure 1.

It has been suggested, on the basis of the identical nature of the amplified $Est\alpha 2^l$ and $Est\beta 2^l$ restriction

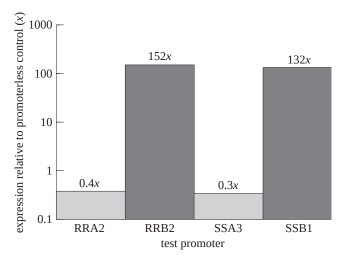


Figure 2. Expression rates in a luciferase reporter assay relative to a promoterless control when the intergenic spacer region from resistant (RR) and susceptible (SS) insects is cloned in the orientation of either the Est β gene or the Est α gene (labelled A and B, respectively).

digest patterns from resistant *Culex* populations worldwide, that amplification is a rare or unique event that occurs primarily through migration (Raymond *et al.* 1991). We now know that amplification of these genes has appeared independently at least five times (Hemingway & Karunaratne 1998), and that resistance is occurring through gene amplification and rapidly spreading by migration. Further evidence that the chromosomal region containing these esterases represents an amplification 'hot-spot' comes from other *Culex* species. In *C. tritaeniorhynchus* the homologous Estβ gene CtrEstβl¹ has been amplified and there is no possibility of this having occurred through gene flow between the species (Karunaratne *et al.* 1998).

In Anopheles stephensi there are at least three enzymes that are able to metabolize the OP malathion in resistant insects. None of these esterase genes are amplified and they are all present in very low quantities in the resistant and susceptible insects, conferring resistance through efficient insecticide metabolism (K. G. I. Jayawardena and J. Hemingway, unpublished data).

(b) Gene expression

Increased gene expression, rather than gene amplification, is the primary molecular basis of glutathione S-transferase and monooxygenase-based resistance in mosquitoes. However, gene amplification and elevated expression are not mutually exclusive. The Estα21 and Est $\alpha 2^1$ genes from *Culex* appear to be both amplified and increased in expression. The two genes are present in a 1:1 stoichiometry, being co-amplified, but approximately three times more $\operatorname{Est}\beta 2^1$ than $\operatorname{Est}\alpha 2^1$ is obtained from protein purifications of resistant insect homogenates (Karunaratne 1994). This difference may reflect differential protein or mRNA stability, or result from variations in the efficiencies of the two promoters, which are both contained within the intergenic spacer (figure 1). We are currently characterizing the Estα2¹ and Estβ2¹ promoters, and have cloned the intergenic spacer in both orientations upstream of the reporter gene luciferase. The

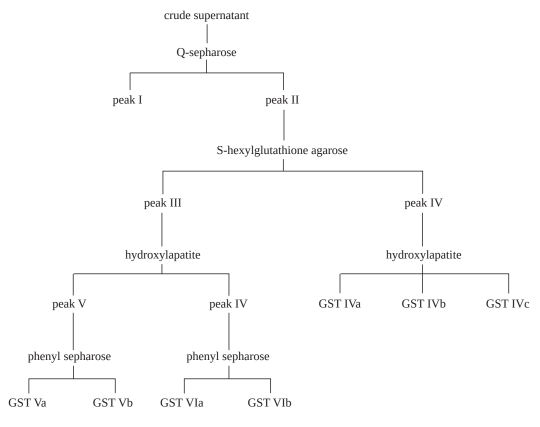


Figure 3. Schematic representation of the purification of glutathione S-transferases (GST) from *Anopheles gambiae*. The final peaks of GST activity still all contained multiple protein bands.

Table 1. DDT dehydrochlorinase activity^a exhibited by various peaks of glutathione S-transferase activity partially purified from A. gambiae, as described in figure 3

(S, susceptible strain enzymes; R, resistant strain enzymes.)

GSTs	$\rm nmole\ DDE\ mg^{-1}$		nmole/unit GSTb DDE activity		$nmole\ DDE\ g^{-1}\ larvae$	
	S	R	S	R	S	R
IVa	_	278.0	< 0.3	6.1	_	14.6
IVb	_	912.5	< 0.3	22.6	_	15.6
IVc	_	22.0	< 0.3	2.2	_	1.5
Va	173.3	564.7	144.5	241.3	48.4	344.9
Vb	65.6	765.6	121.5	243.1	7.6	68.1
VIa	27.4	50.5	94.6	112.2	4.2	39.6
VIb	9.8	178.6	98.3	235.1	0.9	52.4

^a DDT-dehydrochlorinase activity is defined as nmole DDE formation per two hours.

resultant constructs have been transfected into a range of insect and mammalian cell lines. Inserting the spacer at the same site, but in different orientations, reproducibly generates luciferase expression from the $Est\beta 2^l$ promoter many times greater than from the $Est\alpha 2^l$ orientation. This is true from both the resistant (2.7 kb) and susceptible (1.7 kb) spacers (figure 2). The differences in promoter strength may reflect differences in the relative locations of significant regulatory elements. Further studies are in progress.

In the malaria vector, *Anopheles gambiae*, resistance to DDT in both larvae and adults is conferred by increased levels of many glutathione S-transferases (GSTs). Resistance in the two life stages is conferred by different

genes, although the end result of both is a measurable increase in GST activity and DDT dehydrochlorination. Resistance has been studied in greatest detail in larvae. Resistant insects have increased levels of DDT dehydrochlorinase activity associated with seven partially purified peaks of GST activity (figure 3; and table 1) (Prapanthadara *et al.* 1993, 1995). There is an enormous diversity of GSTs found in both resistant and susceptible insects.

At present two broad classes of GSTs have been cloned from insects. Representatives from both classes have been cloned from A. gambiae (although the single insect class II GST cloned from A. gambiae is not expressed in larvae). All three class I GSTs cloned are able to use DDT as a

 $[^]b\,A$ unit of GST activity is defined as $\mu mole\,min^{-1}\,mg^{-1}$ with CDNB as the substrate.

substrate (Ranson et al. 1997a,b). Antisera raised to these expressed GSTs indicates that all of them belong to the peak IV GSTs represented in figure 3. We do not have, as yet, have any molecular data on the GSTs from peaks V and VI, but our current information suggests that they belong to GST classes that have not so far been characterized from any insect. The simplest hypothesis for the molecular basis of this GST-based resistance and for the similar organization of the monooxygenase-based pyrethroid resistance is that trans-acting regulators are involved in the up-regulation of these enzyme families. We are currently employing a positional cloning approach, in collaboration with Professor F. Collins, USA, to identify these regulator genes. The positional cloning takes advantage of the high-density microsatellite marker genetic map, in situ hybridization, polytene chromosome microdissection, and a BAC library, which have all been recently developed for A. gambiae.

(c) Gene splicing

Initial biochemical work on GSTs from A. gambiae demonstrated the diversity of GST enzymes present in this insect. The molecular work undertaken subsequent to this showed that the class I GSTs were all at a single chromosome location. Extensive sequencing of stretches of DNA at this location revealed a full-length intron-less gene, as occurs in Drosophila, and numerous apparent 5' truncated pseudogenes. Reverse transcriptase-polymerase chain reaction and Southern blot analysis has demonstrated that these are not pseudogenes, but are actively transcribed with the splicing of different 5' exons to a single 3' exon, occurring to produce a diversity of GSTs in both resistant and susceptible A. gambiae (Ranson et al. 1998).

The next decade should see extensive progress in our understanding of metabolically based insecticide resistance in insect pests, allowing for the development of new control methods to allow us to counteract this rapidly changing evolutionary phenomenon.

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