

The evolution of insecticide resistance in the peach–potato aphid, *Myzus persicae*

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The peach–potato aphid *Myzus persicae* (Sulzer) can resist a wide range of insecticides, but until recently (1990) the only mechanism identified was the increased production of carboxylesterases (E4 or FE4), which cause enhanced degradation and sequestration of insecticidal esters. We have now identified two forms of target-site resistance involving changes in the acetylcholinesterase (*AChE*) and sodium channel (*kdr*) genes. Biochemical and DNA diagnostic methods can be used to identify all three mechanisms in individual aphids, and thereby establish their spatial distributions and temporal dynamics. Amplified genes underlie the increased production of esterases, but their expression is modulated by DNA methylation. Amplification of the *E4* gene is in strong linkage disequilibrium with the *kdr* mechanism. This may reflect strong insecticidal selection favouring aphids with multiple mechanisms, tight chromosomal linkage and/or the prominence of parthenogenesis in many *M. persicae* populations. The decreased fitness of resistant aphids under winter conditions may be a consequence of the altered sodium-channel gene affecting behaviour and/or the perception of external stimuli.

Keywords: aphids; carboxylesterases; chromosomal linkage; gene amplification; target-site resistance

1. GENETIC CONSIDERATIONS

Any consideration of the evolution of resistance in peach–potato aphids (*Myzus persicae*), must take into account their complex life cycle, which can vary according to the different environments in which they occur (Blackman 1974). They have herbaceous summer (secondary) hosts, which include many annual crops such as potatoes, sugar beet, chrysanthemums, tobacco and various brassicas, on which they reproduce by parthenogenesis. Parthenogenetic populations develop as a mixture of clones, with the most favoured ones as potential dominators of the population. They can then overwinter either asexually (through continued parthenogenesis on protected crops, winter weeds or field crops such as oil seed rape) or as sexually-produced eggs on a woody (primary) host. Because *M. persicae* has a strict requirement for peach (*Prunus persica*) as its primary host, sexual reproduction and meiotic recombination are confined to those areas where this tree is present. The variation in reproductive mode is not only due to the environment but is also dependent on genotype (Blackman 1974). Thus, some clonal lines respond to autumn conditions by producing males and sexual egg-laying females (holocycly), whereas others can only reproduce through continuous parthenogenesis (anholocycly), and yet others are able to produce males but not females (androcycly). Clearly, this will have a large impact on population structure and gene flow.

In addition to showing variation in their reproductive mode, *Myzus* can display preferences between summer hosts. This can be very marked, to the extent that the

aphids become recognizable morphometrically, as with the tobacco-feeding form, which has been classified as a distinct species, *M. nicotianae* (Blackman 1987). Such segregation would again be expected to restrict gene flow. However, in areas where peach is grown close to tobacco crops, for example in Greece, there is the opportunity for genetic mixing because the two forms retain their ability to interbreed.

A third genetic feature relevant to the evolution of resistance in *M. persicae* is their karyotype. It has long been recognized that a heterozygous translocation between autosomes 1 and 3 is associated with one type of esterase-based resistance (Blackman *et al.* 1978). The translocation appears to reduce the aphid's ability to reproduce sexually, again having implications for the spread of resistance genes. Aphids with the A 1,3 translocation are widely distributed in warm temperate and tropical regions of the world, and in protected (glasshouse) crops in northern Europe, where they are mostly either anholocyclic or androcyclic.

2. INSECTICIDE DETOXIFICATION BY ESTERASES

(a) Biochemistry

For many years, the only resistance mechanism identified in *M. persicae* was the overproduction of insecticide-detoxifying esterases. This form of resistance was first implicated in the late 1960s by the demonstration that all resistant strains showed an increased ability to hydrolyse the model esterase substrate, 1-naphthyl acetate (Needham & Sawicki 1971). It was subsequently shown to

arise from the increased production of one of two forms of esterase, E4 or FE4, that were distinguishable electrophoretically (reviewed by Devonshire 1989). These esterases can account for as much as one per cent of the aphid's total body protein, and give a broad spectrum of resistance to organophosphorus, carbamate and pyrethroid insecticides as a consequence of both ester hydrolysis and sequestration (Devonshire & Moores 1982). Although FE4 hydrolyses insecticides slightly faster than E4, the form of esterase overproduced has little effect on the level of resistance (Devonshire *et al.* 1983).

The biochemical evidence for the role of these esterases in resistance was supported by selection experiments: spraying *M. persicae* populations with pyrethroid insecticides selected strongly for aphids carrying esterase-based resistance to organophosphates and carbamates, and vice versa. This selection has been observed both in the open field (French-Constant *et al.* 1988a) and in field cages colonized by known initial proportions of susceptible and resistant aphid clones (French-Constant *et al.* 1987). However, for pyrethroids the contribution of esterases to resistance has recently been shown to be secondary to that conferred by target-site insensitivity, with the two mechanisms being co-selected as a consequence of a strong linkage disequilibrium between them (see below).

(b) *Amplification of esterase genes*

The observation that aphid clones with increased amounts of esterase fell into a series with a progressive doubling of enzyme activity, led to the hypothesis that the underlying cause of resistance was amplification of their structural genes, rather than transcriptional control (Devonshire & Sawicki 1979). This was confirmed in later work, but the phenomenon was shown to be more complex than a simple series of successive duplications (Field *et al.* 1988). The genes encoding the two esterase forms, E4 and FE4, each span approximately 5 kb and are very similar (Field *et al.* 1993). They differ by only nine amino-acid substitutions; and FE4 has an additional 12 amino acids at the C-terminus as a consequence of a mutation that changes the stop codon present in the E4 gene. All clones studied so far have had identical E4 or FE4 genes, including the introns that have been sequenced. This suggests that a single amplification event occurred for each gene, and that they then became dispersed. Likewise, the tobacco-feeding form, *M. nicotianae*, has been shown to have exactly the same amplified genes, indicating that there is gene flow between this and *M. persicae* (Field *et al.* 1994).

In both *M. persicae* and *M. nicotianae*, amplification of E4 genes always occurs in conjunction with the A 1,3 translocation, whereas aphids that have only the amplified FE4 genes are apparently of normal karyotype. It is very rare to find both forms of the esterase genes amplified in a single individual, pointing to some reproductive isolation between aphids carrying the different genes. The two amplified genes were first seen together in the progeny of laboratory crosses between E4 and FE4 clones (Blackman *et al.* 1996); and the combination has recently been found in a small number of aphids collected in the field in Greece (Blackman *et al.* 1998). In the latter case, aphids with both genes amplified have also had the translocation.

The organization of the amplified genes has been studied by analysing large restriction fragments by means of pulsed-field gel electrophoresis (PFGE), which gave very different results for E4 and FE4 genes. Rare-cutting enzymes gave a single DNA fragment of 300–350 kb for amplified E4 genes in R₃ aphids, whereas those producing comparable amounts of the FE4 enzyme gave a family of fragments (Field *et al.* 1996). The complexity of the FE4 pattern precluded detailed analysis, but partial digests of the amplified E4 genes indicated that they were arranged in a tandem array of 12 copies, with the esterase gene being part of a 24 kb amplicon (Field & Devonshire 1997). However, recent work (Field *et al.*, unpublished data) has shown that the number of gene copies for both E4 and FE4 genes is higher, and more in line with the level of esterase production.

Further analysis of the E4 and FE4 genes has provided evidence that in susceptible aphids they are in a head-to-tail arrangement with E4 upstream of FE4, with about 19 kb of intervening sequence (Field & Devonshire 1998). Their close similarity suggests a recent duplication event, followed by limited divergence of the two forms. The subsequent amplifications of the E4 or FE4 genes must have involved separate events, each occurring once and then being selected by insecticide treatment and spread by migration (figure 1). The 5' ends of the genes differ, suggesting that they are regulated in different ways. FE4 appears to have a conventional promoter region, whereas E4 has a CpG island (associated with a 1.7 kb insertion that is absent from the FE4 gene) that may influence expression through changes in DNA methylation (Field & Devonshire 1998). This would be in accord with the observation that only E4 genes appear to undergo reversion through changes in transcription (see below). The novel joint (NJ) in the DNA sequence generated during the E4 amplification is just upstream of the transcription start site, and the FE4 joint (FJ) generated by the initial duplication event leading to the divergence of FE4 is only 3 bp away from the NJ. This suggests that this part of the chromosome might be prone to recombination, perhaps as a consequence of the initial duplication. The position of the NJ generated by the FE4 amplification has not yet been identified.

(c) *Chromosomal location and inheritance of esterase genes*

Fluorescence *in situ* hybridization (FISH) has enabled the sites of esterase gene amplification to be identified. The situation is simplest for amplified E4 genes, which, with one exception amongst those clones studied, have always been found at a single locus on the truncated autosome 3, close to the translocation break-point. In the one exceptional clone, the amplified E4 genes were also found at two other loci on autosomes 2 and 5 (Blackman *et al.* 1995). In contrast, FE4 genes have been found widely dispersed around the genome, with interclonal variation in the number of loci, some apparently being homozygous and others heterozygous in any one clone (Blackman *et al.* 1995, 1996). These results are in line with the PFGE data, which indicated a single block of E4 amplicons, but several differently sized groups of FE4 genes (Field *et al.* 1996). Crossing experiments, some coupled with FISH analysis, have shown that inheritance patterns are consistent with a single heterozygous locus of E4 amplification

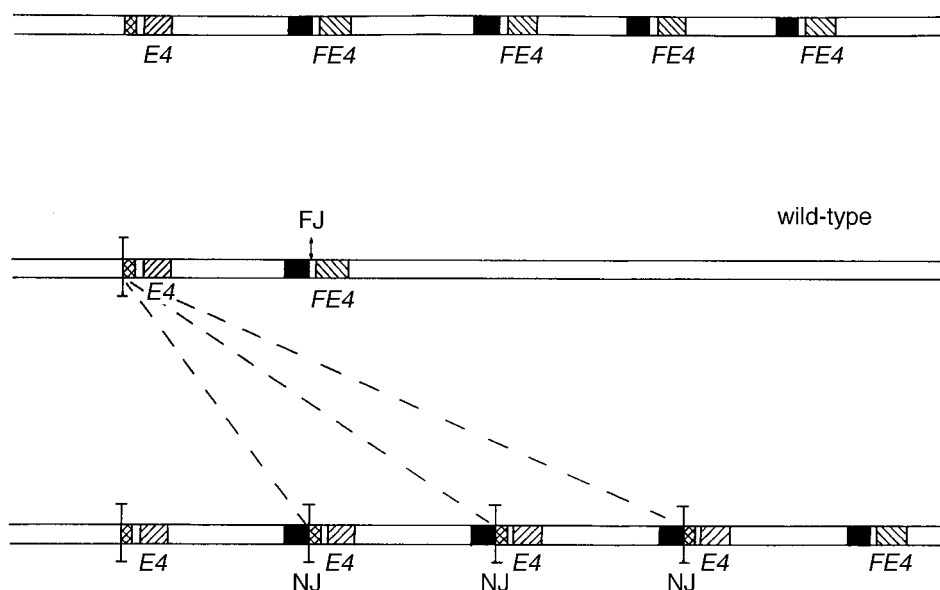


Figure 1. Model for the amplification and organization of *E4* and *FE4* genes (based on Field & Devonshire 1998).

in R_3 clones (Blackman *et al.* 1978, 1996). In contrast, two- or three-locus models were required for clones showing different degrees of *FE4* amplification (Blackman *et al.* 1996).

Despite their widespread dispersion around the genome, restriction analysis of the amplified genes indicates that all *FE4* copies are in the same immediate genetic background, i.e. the amplicon structures are maintained. The same conservation applies to the one aphid clone where the *E4* amplicons occur at three loci (Field & Devonshire 1998). This suggests that in each case an original amplification event occurred recently and the amplicon was then moved, intact, around the genome through rearrangements such as translocations and inversions, or perhaps mediated by transposable elements. The position and sequences around the site of initial duplication (FJ in figure 1) are identical in all of the aphid clones examined, reinforcing the likelihood of a single event that has, subsequently, become widely spread in populations around the world.

(d) *Transcriptional control of amplified esterase genes*

Some clonal cultures of aphids with amplified *E4*, but apparently not *FE4*, genes lose resistance spontaneously, concomitant with ceasing to over produce the esterase (Sawicki *et al.* 1980). This 'reversion' phenomenon occurs infrequently and stochastically in laboratory cultures in the absence of selection. However, recent evidence indicates that revertant aphids might be more common in the field than originally considered (L. M. Field and A. L. Devonshire, unpublished data). This loss of esterase-gene expression seems analogous to that seen in cell cultures resistant to cytotoxic drugs, which arises from loss of the amplified genes that confer resistance (Stark 1993). However, in *M. persicae*, revertant aphids retain their full complement of amplified esterase genes, suggesting that decreased transcription is responsible for the loss of enzyme production.

In aphids, the loss of expression was associated with changes in the methylation of the amplified esterase genes (Field *et al.* 1989); further work demonstrated a tight

correlation between the two events (Hick *et al.* 1996). The amplified genes are expressed when 5-methylcytosine is present in and around the gene, and they are silenced when the methylation is lost in revertants. This positive correlation is contrary to the situation in vertebrates, where methylation is usually associated with gene silencing (Holliday *et al.* 1996). Revertant aphid clones spontaneously produce a small proportion of offspring with higher esterase levels, which are selected when exposed to insecticides (French-Constant *et al.* 1988*b*). This plasticity confers a potential advantage to these aphid clones by enabling them to avoid producing the large amount of esterase protein when it is not needed.

(e) *Origin and spread of amplified esterase genes*

The molecular studies of amplified esterase genes, as in mosquitoes (Raymond *et al.*, this issue), points to a small number of isolated mutation events (Field & Devonshire 1997, 1998) which then spread rapidly through migration. The mode of aphid reproduction, coupled with their potential for rapid growth as asexual populations, mean that the parthenogenetic offspring of an individual with a marked advantage could become dominant very quickly. Furthermore, the migration of aphids, through both short- and long-distance flight, will lead to the dispersal of resistance genes. However, another way in which insects can become dispersed around the world is as a result of human activity. This has been invoked for mosquitoes, in which their movement through long-distance transport in boats and aircraft appears to have played a part in the rapid spread of resistance genes (Pasteur & Raymond 1996). Likewise, the international trade in plants and produce offers considerable potential for the widespread distribution of phytophagous insect pests (Frey 1993; Denholm *et al.*, this issue).

3. TARGET-SITE RESISTANCE

(a) *Insecticide-insensitive acetylcholinesterase*

Insecticide-insensitive acetylcholinesterase (AChE), the target for organophosphorus and carbamate insecticides, is an important resistance mechanism in many insect

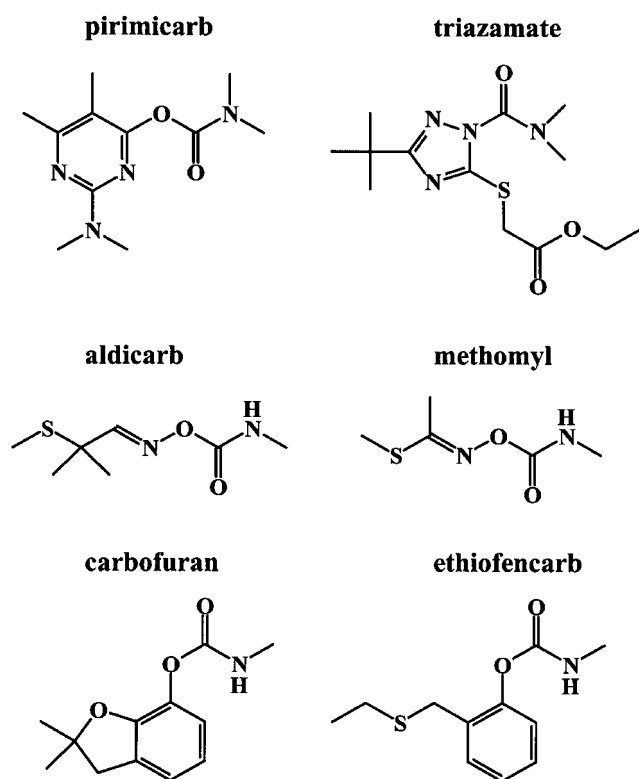


Figure 2. Chemical structures of *N,N*-dimethylcarbamates (pirimicarb and triazamate) to which the *Myzus* AChE is insensitive, and some *N*-monomethylcarbamates that are unaffected by this resistance mechanism.

species, but was not found in the *M. persicae*–*M. nicotianae* complex until 1990 (Moores *et al.* 1994a). It was first detected in populations from Greece, followed by those from Japan and South America, and has recently shown a northward expansion in its European distribution (Moores *et al.* 1994b). Aphids with insensitive AChE were first identified in the UK in 1995 from samples which were collected in aerial suction traps but did not cause any control problems in that year. However, in 1996, insensitive AChE occurred commonly in aphids from eastern England (Foster *et al.* 1998) where there were control failures with pirimicarb, the favoured insecticide for controlling the *M. persicae* that are very resistant (R_2 and R_3) owing to esterase overproduction.

The fact that insensitive AChE was only found recently (1990), and only in association with the esterase mechanism (both *E4* and *FE4*), suggests that it evolved after the amplified esterases. The incidence of what appears to be the same *AChE* gene alongside both forms of esterase suggests that it originated in a holocyclic individual and then spread through the *M. persicae*–*M. nicotianae* complex. If it originated as a single mutation event, its introgression into aphids with the A 1,3 translocation (which are anholocyclic or androcyclic) must have been through a female mating with a male from an androcyclic *E4*-producing clone. The continuous parthenogenetic reproduction of the offspring of such a cross could then have enabled it to be stably maintained in a heterozygous condition, when its phenotypic expression (resistance to pirimicarb) is semi-dominant (Moores *et al.* 1994b). However, it has also been found in a homozygous

condition in combination with amplified *E4* as well as *FE4*, indicating further mating, presumably through *E4* males crossing with females of normal karyotype also with the insensitive *AChE* gene. In England, aphids with the insensitive *AChE* gene have all been red and with amplified *E4*, whereas in Greece, where *FE4* predominates, the insensitive form has been found in both the red and green colour morphs.

The insensitive AChE confers strong resistance specifically to pirimicarb and triazamate, the latter being a novel and otherwise very effective triazole aphicide (Dewar *et al.* 1994) which also inhibits AChE. Both are *N,N*-dimethylcarbamates; there is no insensitivity to a range of organophosphorus compounds nor to aryl and oxime *N*-monomethylcarbamates (figure 2), suggesting that the bulkier substituents on the carbamate nitrogen are critical for the expression of resistance in this particular species. The insensitivity appears to arise at the inhibition stage, rather than from differential reactivation. Reactivation rates (k_3 , measured as described by Devonshire & Moores (1982)) were too slow to be determined for the sensitive AChE; whilst the insensitive form gave values of only $0.0029 \pm 0.0001 \text{ min}^{-1}$ and $0.00047 \pm 0.00007 \text{ min}^{-1}$ for mono- and dimethylcarbamylated enzyme, respectively.

Although the molecular basis of AChE insensitivity in other species involves one or more point mutations that change amino-acid residues close to the catalytic site of the enzyme (Mutero *et al.* 1994; Devonshire *et al.* 1998), the changes in *M. persicae* have not yet been established.

(b) *Modified sodium channels (knockdown resistance or kdr)*

We have recently shown that the esterase-based resistance to pyrethroids in *M. persicae* is of secondary importance compared with a *kdr*-type mechanism not previously identified in this species (Martinez-Torres *et al.* 1997, 1998), and that their co-selection appears to have been a consequence of their close association in some clones (Field *et al.* 1997). Aphids lacking the *kdr* mechanism, but with R_3 levels of esterase, only show approximately five-fold resistance to deltamethrin. The *kdr* mechanism alone (as found only in revertant aphids) confers 35-fold resistance and this is enhanced a further 15-fold by R_3 levels of esterase (figure 3). Analysis of aphid clones collected over many years, and reared in the laboratory, indicated that this mechanism had long been present in populations, but was only identified once a molecular diagnostic method became available. The cross-resistance it confers to DDT has since enabled the use of a discriminating dose bioassay for identifying its presence. The *kdr* mechanism involves a mutation causing the replacement of a leucine by phenylalanine in the domain IIS6 transmembrane region of the insect *para*-type sodium-channel gene. This mutation was first identified in house flies (Williamson *et al.* 1996) and cockroaches (Miyazaki *et al.* 1996), and the same change has since been found in several insect species including *M. persicae* (Martinez-Torres *et al.* 1997; French-Constant *et al.*, this issue).

In UK populations, the *kdr* mechanism shows strong linkage disequilibrium with the amplification of *E4*, but not *FE4*, genes (table 1). This perhaps reflects the almost

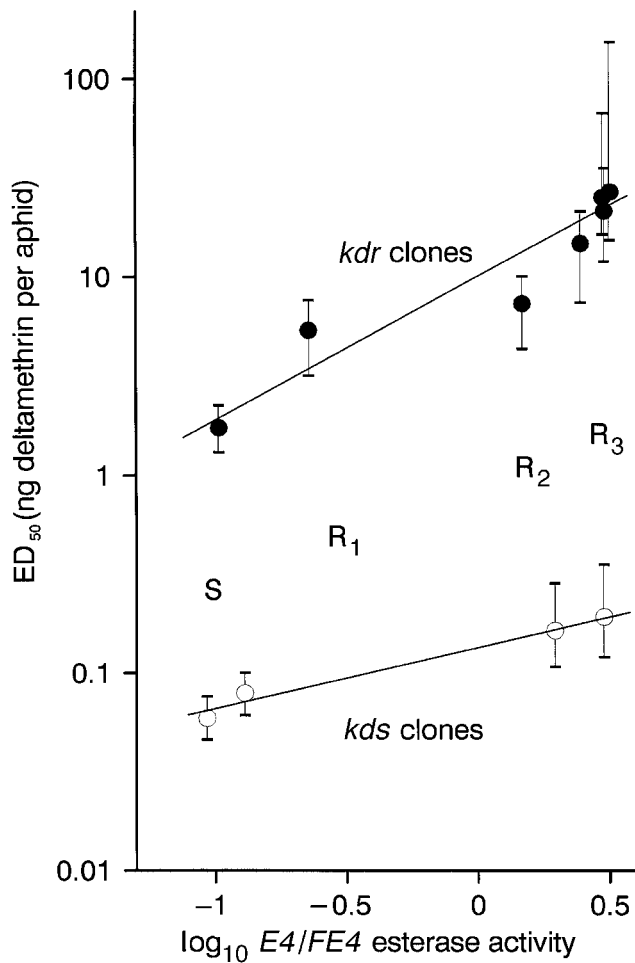


Figure 3. Relationship between toxicity of deltamethrin (ED_{50}) and mean esterase activity, measured by immunoassay, in *kdr* and *kds* (i.e. lacking the *kdr* mechanism) aphid clones. Limits of bars indicate 95% confidence intervals given by probit analysis. All of the *kdr* clones have highly amplified *E4* genes, but the clone with S esterase activity is a revertant, and those with R_1 and R_2 esterase levels appear to be partial revertants, based on analysis of DNA methylation patterns (Field *et al.* 1989).

exclusive chromosomal location of the amplified *E4* genes at a single locus on the truncated chromosome 3 close to the translocation break-point, whereas amplified *FE4* genes are more dispersed around the genome (Blackman *et al.* 1995, 1996, 1998). However, this linkage of the *kdr* mechanism with *E4* gene amplification does not hold fully outside the UK: in a broader survey of aphids from 15 countries, 5 of the 30 clones with amplified *E4* genes lacked the *kdr* mechanism (Field *et al.* 1997). It is not known whether the close association of the *E4* and *kdr* mechanisms reflects a tight chromosomal linkage. It seems more likely that the *kdr* mutation first occurred in a single *E4*-overproducing, translocated aphid, and gave its progeny such a large selective advantage when exposed to a wide variety of insecticides that they survived preferentially and have become widely dispersed as a clone. Following this hypothesis, males from this (androcyclic) clone would then have passed the *kdr* mutation via the sexual phase to some *FE4*-overproducing genotypes.

Table 1. Incidence of the *kdr* mechanism^a in UK aphids with different esterase levels and amplification of *E4* and *FE4* genes

esterase level	number of clones with:			
	amplified <i>E4</i>		amplified <i>FE4</i>	
	<i>kdr</i>	<i>kds</i> ^b	<i>kdr</i>	<i>kds</i>
R_1	8	0	1	0
R_2	26	0	3	11
R_3	67	0	1	1

^a Based on a diagnostic bioassay with DDT (Field *et al.* 1997).
^b The corresponding susceptible phenotype is referred to as *kds* for simplicity.

4. RELATIVE FITNESS OF SUSCEPTIBLE AND RESISTANT APHIDS

One of the tenets of resistance management is that, in the absence of insecticides, resistant insects are less 'fit' than their susceptible counterparts. Early studies of *M. persicae* provided evidence for some resistant clones moving less readily between different host species in the laboratory (Eggers-Schumacher 1983). Evidence of a more marked fitness deficit under field conditions began to emerge from the results of regular monitoring of populations throughout and between growing seasons in England (Smith *et al.* 1990). The proportion of aphids with esterase-based resistance built up during the summer in populations sampled from crops and, more randomly, in the aerial suction traps of the Rothamsted Insect Survey, presumably reflected selection pressure from insecticide use on the various crop hosts of *M. persicae*. However, their proportions declined over several winters, so that the overall situation remained relatively stable from 1988 to 1995 (Muggleton *et al.* 1996), suggesting that the more resistant forms were less able to survive the winter climate.

The poorer overwintering performance of *E4*-overproducing resistant aphids, compared with that of susceptible aphids, was clearly demonstrated in four out of nine field experiments over the winters of 1992–3 and 1993–4 (Foster *et al.* 1996). Several distinct clones from each of the resistance categories, S, R_1 , R_2 , R_3 and revertants, were colonized as early-instar nymphs on to oilseed rape crops, and their survival determined a month later; one such trial is shown in figure 4. These predominantly showed a negative correlation between survival and resistance level, with revertants behaving similarly to their R_3 counterparts (indicating that overproduction of esterase *per se* is not responsible for the differences observed). The relationship correlated strongly with three meteorological variables: day temperature below 2 °C, mean rainfall and mean wind speed. One of the factors implicated in this maladaptive behaviour was the tendency for the more resistant aphids to remain on deteriorating leaves, whereas the more susceptible clones moved more readily. This was apparent in both field trials and laboratory experiments (Foster *et al.* 1997), and would be expected to occur to increase the risk of the resistant aphids being separated from the plant after leaf-fall. Another characteristic was that some resistant

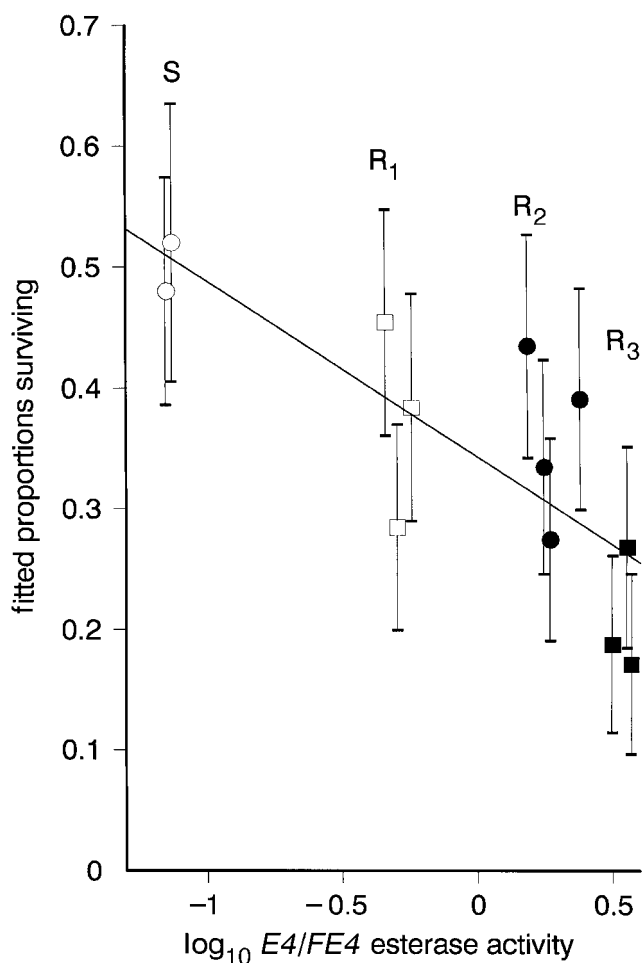


Figure 4. Relationship between esterase activity and the proportion of aphids surviving on oilseed rape crops in the field during a three week period in February 1994 (based on Foster *et al.* 1996). Symbols: open circles, S clones; open squares, R₁ clones; filled circles, R₂ clones; filled squares, R₃ clones. Limits of bars indicate \pm 1 s.e.

clones of *M. persicae* were shown to be less responsive to their own alarm pheromone (Dawson *et al.* 1983); and more recent work (Foster *et al.*, unpublished data) has shown this to be so in a wider range of clones.

At the time of these experiments, classification of resistant aphids was based on their esterase content. We now know that the *kdr* factor commonly occurs in combination with high E4 production, in UK clones. Because the mutation underlying *kdr* is in the sodium-channel gene, this would seem to be a more likely factor mediating the modified behaviour responsible for the reduced overwintering success, e.g. by reducing the sensitivity of the nervous system to stimuli. The *smellblind* mutant in *Drosophila melanogaster* provides a precedent for such an effect of a sodium-channel mutation (Lilly *et al.* 1994).

5. THE FUTURE

The way in which resistance will evolve in the future will depend on the pest control measures adopted. Our increasing knowledge of the underlying mechanisms, and the availability of sensitive and rapid diagnostic methods for their identification, opens the way to make rational

choices of insecticides to minimize selection pressure in particular circumstances. Increasing pressure to minimize pesticide use is also likely to help reduce the selection for resistance. However, one of the immediate needs is for novel chemicals circumventing, or even exploiting (Hedley *et al.* 1998), existing mechanisms. The recent introduction of the chloronicotinyl class of aphicides, with its distinctive chemistry and mode of action, holds great promise for addressing this need (Elbert *et al.* 1990). However, there is already evidence for a low level of resistance to these compounds in some *M. persicae* populations (Devine *et al.* 1996), and if their promise is to be fulfilled, it will be essential to manage their use carefully from the outset. This issue is addressed by others in this volume.

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