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Effects of host plant and genetic background on the fitness costs of resistance to *Bacillus thuringiensis*

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Abstract

Novel resistance to pathogens and pesticides is commonly associated with a fitness cost. However, measurements of the fitness costs of insecticide resistance have used diverse methods to control for genetic background and rarely assess the effects of environmental variation. Here we explored how genetic background interacts with resource quality to affect the expression of the fitness costs associated with resistance. We used a serially backcrossed line of the diamondback moth, Plutella xylostella, resistant to the biopesticide Bacillus thuringiensis, to estimate the costs of resistance for insects feeding on two Brassica species. We found that fitness costs increased on the betterdefended Brassica oleracea cultivars. These data were included in two meta-analyses of fitness cost experiments that used standardized protocols (and a common resistant insect stock) but which varied in the methodology used to control for the effects of genetic background. The meta-analysis confirmed that fitness costs were higher on the low quality host (B. oleracea); and experimental methodology did not influence estimates of fitness costs on that plant species. In contrast, fitness costs were heterogeneous in the Brassica pekinensis studies: fitness costs in genetically homogenized lines were significantly higher than in studies using revertant insects. We hypothesize that fitness modifiers can moderate fitness costs on high quality plants but may not affect fitness when resource quality is low.

Keywords

Bacillus thuringiensis; compensation; gene X environment interaction; plant quality; resource quality

Introduction

Selection for resistance to pesticides, at least initially, produces a genetic correlation that results in a reduction in fitness in the absence of the pesticide (Crow, 1957; Foster *et al*, 2003; Gould, 1998). This is commonly referred to as a "fitness cost". These costs are not always detectable but environmental factors such as natural enemies, resource limitation and over-wintering have been observed to increase costs, although the mechanisms behind these increases remain unknown (Carrière *et al*, 2001; Janmaat and Myers, 2005; Raymond *et al*, 2007a; Raymond *et al*, 2005). Fitness costs can be important for the sustainable use of a pesticide: theoretically they are capable of inhibiting the evolution of resistance to pesticides

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in agro-ecosystems, especially when these costs are large or partially dominant (Carrière *et al*, 2001; Carrière and Tabashnik, 2001; Raymond *et al*, 2007a; Tabashnik, 1989).

Host plant defences are a major source of stress for herbivorous insects, and these defences have been known to alter the fitness costs associated with pesticide or pathogen resistance (Carrière *et al*, 2005; Carrière *et al*, 2004; Janmaat and Myers, 2005; Raymond *et al*, 2007b). Research into the effects of host plant defences or host plant varieties on the fitness costs of resistance has been primarily motivated by the ongoing need to improve resistance management. Since crop plant characteristics are, to some extent, under the control of growers or plant breeders it may be possible to choose plant crop varieties that maximize the fitness costs of resistance. However, the environment is not the only factor to significantly alter fitness costs. Genetic background can also have a major impact on the pleiotropic effects of any particular mutation. Average genetic heterogeneity may influence fitness costs (Paris *et al*, 2008) or the genetic background may include specific "modifier" genes which act to reduce fitness costs (Clarke and McKenzie, 1987; Maclean *et al*, 2004).

Different experimental methods used to estimate fitness costs can control for the effect of linked genes or genetic background. Estimation of fitness costs of pesticide resistance in insects has frequently used a simple "between population" approach that involves assessing the fitness of a recently isolated resistant field population against a standard laboratory population (Alyokhin and Ferro, 1999; Oliveira et al, 2007). However, this method makes no attempt to control for the effect of genetic background or genes linked to resistance loci, although genes linked with resistance alleles are known to have potentially deleterious consequences for fitness (Bergelson and Purrington, 1996; Chaplin and Mann, 1978; Parker, 1991). A more rigorous method is to use an experimental selection-based approach to generate matched pairs of resistant and susceptible lines. This can be done by repeatedly selecting for resistance in susceptible population, or allowing high levels of resistance to revert to susceptibility in an unselected line (Groeters et al, 1994; Luong and Polak, 2007). Identification of resistance genes also means that parents of paired resistant and susceptible populations with a common genetic background can be identified using PCR (Carrière et al, 2004; Gassmann et al., 2008; Yang et al., 2008). There is also "homogenization" or "introgression" approach in which near isogenic lines are produced by serially backcrossing resistant strain into a susceptible population and reselecting for resistance after each cross (Bergelson and Purrington, 1996). Finally genetic engineering approaches can be used to insert resistance genes and fully control for genetic background (Tian et al, 2003). Introgression-based protocols (and genetic engineering approaches) have been considered as the most rigourous means of producing matched pairs of resistant and susceptible lines since repeated rounds of recombination and selection ensures that observed fitness changes are due to mutations at the locus of interest, rather than due to variation in genes linked to resistance loci (Bergelson and Purrington, 1996). Introgression-based protocols have been commonly employed in plant studies but only rarely used in insect evolutionary ecological experiments (McCart et al, 2005; Yang et al, 2008).

Once paired resistant and susceptible lines have been created via reversion, selection, introgression or molecular methods, differences in fitness between these lines can be then assessed using either a fitness components approach (in which individual life history parameters are measured) or by using a selection based stability analysis (in which the decline in resistance in the absence of the selective agent is measured over a number of generations) (Gassmann *et al*, 2009; Roush and McKenzie, 1987). In this study we estimated the fitness costs associated with resistance to the entomopathogen *Bacillus thuringiensis* (*Bt*) using a fitness components analysis of serially backcrossed insects derived from the Karak strain of the diamondback moth *Plutella xylostella*. We investigated how the magnitude of these costs varies with different host plant species (*Brassica oleracea, Brassica pekinensis*),

which represent resources of very different quality for the larvae of this moth. We also explored how different experimental techniques that control for genetic background in a variety of ways could affect our estimation of fitness costs. Moreover, we investigated how controlling for genetic background would affect fitness cost estimates on both low and high quality host plants. This was carried out using a meta-analysis that explores the variation in fitness cost estimates derived from between-population, selection-based and homogenization approaches on the two host plant species (*B. oleracea, B. pekinensis*). Data for the meta-analysis was collated from both previously unpublished and published work and was produced using standardized protocols on insect populations with the same resistant progenitor (Raymond *et al,* 2007a; Raymond *et al,* 2005; Raymond *et al,* 2007b).

Materials and methods

Fitness experiment with serially backcrossed insects

The 'Karak' (*Bt* resistant) population of diamondback moth, *Plutella xylostella*, has monogenetic resistance to *Bt* based on reduced binding of toxins to the mid-gut (Sayyed *et al*, 2004). This population was backcrossed six times into a standard laboratory susceptible population "Lab-UK" (from Professor Juan Ferré, University of Valencia) to produce a hybrid population (KarUK₆). After each cross, hybrids were left unselected for at least one generation. Resistance was then re-selected for two generations as described previously using 50-100µg ml⁻¹ of DiPel DF® (*B. thuringiensis*, subsp. *kurstaki*, Valent Biosciences) (Raymond *et al*, 2005). In 2005 both the Lab-UK and KarUK₆ strains were crossed with the additional insect strain: "Newcastle" (from Dr Gordon Port, University of Newcastle) and the resistant population as above for one additional generation. All selection and routine culture of insects took place on Chinese cabbage, *Brassica pekinensis* (Lour.) Rupr. cv "One Kilo, S.B".

Relative resistance in the two resulting populations, denoted UK/Newc and KarUK₆/Newc, was measured with standard leaf-dip assays (Sayyed *et al*, 2000). The dominance of the fitness advantage conferred by resistance h was calculated according to the relative fitness of larvae at each dose of biopesticide- (D_{WT}) (Bourguet *et al*, 2000). Since there were zero counts for homozygous susceptible and resistant insects at some doses, the reference fitness in these calculations was the survival of insects in untreated controls.

KarUK₆/Newc resistant and UK/Newc susceptible populations, their F1 hybrids, and three Brassica cultivars, were used in a full-factorial experiment in the presence and absence of Bt (DiPel DF) in July 2006, nearly five years after the original collection of the Karak population (approximately 55 generations). The common cabbage, *B. oleracea* L. var. capitata, varieties "Wheelers Imperial" and "Red drumhead" were used at 16 weeks old and Chinese cabbage, B. pekinensis "One Kilo, S.B" at five weeks old. Individual plants (n = 8)were randomly allocated to treatments. DiPel was applied to plants with a hand sprayer with 0.1% v/v Tritox X-100. Spray adheres to the waxy B. oleracea less well than to B. pekinensis. We calibrated the spore density on plants by dilution plating of leaf washes onto agar plates (Bacillus cereus specific agar, Oxoid) and sprayed B. oleracea plants with correspondingly higher doses (270µg ml⁻¹ compared to 100µg ml⁻¹ on *B. pekinensis*) in order to produce similar spore densities. This dose is approximately 1/10th of the recommended full field dose; however, under laboratory conditions the Karak strain is not resistant to the full field dose. A lower dose, therefore, allows us to discriminate between the different resistant genotypes. After spray had dried freshly emerged neonate P. xylostella larvae were added to each plant (N=35 neonates on B. pekinensis, N=45 neonates on B. oleracea, N = 8 plants per treatment). After 7 days, pupae were removed regularly and adult emergence scored daily. Female fecundity was measured as described previously with up to four individual pairs of *P. xylostella* per plant (Raymond *et al*, 2007a).

Protocols of fitness experiments included in the meta-analysis

All fitness experiments were conducted according to the standard protocol described above with the same varieties of *B. pekinensis* ("One Kilo SB") and *B. oleracea* ("Wheelers Imperial") (Raymond et al, 2007a; Raymond et al, 2005; Raymond et al, 2007b) using insects derived from the Karak strain, which was collected in Malaysia in 2001 (Sayyed et al, 2004). In all studies resistant and susceptible insects were bio-assayed in the generation prior to fitness assays in order to confirm the efficacy of selection regimes. Prior to 2005 all experiments were conducted at Imperial College, Silwood Park campus and insects were selected with the purified Cry toxin Cry1Ac. Cry1Ab and Cry1Ac are the dominant virulence factors in B. thuringiensis kurstaki HD-1 (Liu et al, 1996). Both these toxins have similar toxicity to susceptible and Karak resistant P. xylostella (Sayyed et al, 2004). Subsequent experiments were carried out at the Department of Zoology, Oxford, where insects were selected with Bt as described above. In early experiments (Raymond et al., 2005; Raymond et al., 2006, between population experiments) survivorship was recorded as survival until successful pupation; subsequently survival was recorded as proportional adult emergence. Data from one published study, which used B. oleracea plants at eight weeks old, was excluded (Raymond et al, 2007b), since plants of this species were at least sixteen weeks old in all other experiments. Fitness costs on younger cabbage plants were lower than in experiments with older plants. Younger B. oleracea plants also represent a resource of higher quality to P. xylostella (Verkerk and Wright, 1994). Since one of our main aims was to explore the effects of low resource quality on fitness costs, analysing data from a higher quality version of our low quality species would have confounded that goal.

Data from three previously unpublished sets of experiments has been included in the metaanalysis presented here. This includes fitness data from experiments with KarUK₆ (ie backcrossed Karak prior to outcrossing with the Newcastle strain) carried out in 2005. This experiment used n = 8 host plants per genotype with 35 neonates per plant and was replicated twice in two independent blocks. Strong heterozygote advantage in the 2005 experiment suggested that the Lab-UK parent strain was inbred, fitness costs from that study were therefore conservatively estimated from the relative survivorship of resistant (KarUK₆) and heterozygote F1 insects (KarUK₆ X Lab-UK crosses). The "between-population" experiments carried out with the original Karak strain in 2003-4 comprise the second unpublished data set. These experiments used two laboratory susceptible strains: Lab-UK (described above) and Roth (a population maintained at Rothamsted Research for more than 100 generations). Between population experiments with B. pekinensis used the standard protocol with n = 8-10 host plants per genotype (Raymond et al, 2005). The final unpublished 2004 study used revertant and resistant Karak populations on B. pekinensis and used N=9 plants per genotype with 35 larvae per plant. Small differences in replication or numbers of larvae per plant in the published studies have been detailed in previous papers (Raymond et al, 2007a; Raymond et al, 2005; Raymond et al, 2007b). The total sample size for the B. pekinensis meta-analysis was n = 4535 insects and the total sample size in the B. *oleracea* meta-analysis was n = 2679 insects.

Statistical methods

Mean values from each individual plant were used in statistical analyses to avoid pseudoreplication. Fitness components were analysed with generalized linear models with insect population (genotype), plant variety and presence/absence of *Btk* as factors. Fecundity was square-root transformed prior to analysis. Proportional data were logit transformed and analysed with binomial errors; F-tests using the function "quasibinomial" were applied to correct for over-dispersion when appropriate (Crawley, 2005). Estimates of LC₅₀ and their 95% confidence limits (CLs) were calculated according to (Collett, 1991) from individually fitted analyses of deviance. Final model fitting and model assumptions were checked with

graphical analyses. Specific post hoc comparisons between levels within treatments were made using model treatment contrasts. The meta-analysis used a relative risk parameter calculated according to relative survivorship as an indicator of fitness costs (RR = p(survival resistant larvae)/p(survival susceptible larvae). Survival has been has been the most informative metric for fitness costs in the Karak strain (Raymond *et al*, 2007a; Raymond *et al*, 2005; Raymond *et al*, 2007b). We used a random effects meta-analysis with Woolf's test for heterogeneity between studies (DerSimonian and Laird, 1986). All data analysis was carried out in R v 2.6.2 (http://www.r-project.org).

Results

Fitness experiment with serially backcrossed insects

In the generation prior to the fitness experiment (June 2006) the resistance of the selected KarUK₆/Newc population, the susceptible UK/Newc population and both reciprocal F1 hybrid crosses was assayed with DiPel DF. These populations differed in their susceptibility to $Bt(\chi^2 = 16.2, d.f. = 3, P << 0.0001$, Figure 1). The LC₅₀ of the resistant KarUK₆/Newc population exceeded the maximum bioassay dose and was estimated at 728 µg ml⁻¹ (CLs 77.5, 6740 µg ml⁻¹) the LC₅₀ of the susceptible UK/Newc population was 0.325 µg ml⁻¹ (CLs 0.15, 0.70 µg ml⁻¹). The LC₅₀s of the F1 crosses were 4.36 µg ml⁻¹ (CLs 2.45, 7.75 µg ml⁻¹) (resistant mother) and 5.66 µg ml⁻¹ (CLs 2.75, 11.7 µg ml⁻¹) (resistant father). The dominance of resistance (D_{WT}) was 0.145 at a DiPel concentration of 125 µg ml⁻¹, D_{WT} increased to 0.37, 0.39, 0.52 and 0.875 as the biopesticide dose decreased (the doses were 25, 5, 1 and 0.2 µg ml⁻¹ respectively).

The two *B. oleracea* cvs were poorer hosts for diamondback moth than *B. pekinensis* in terms of survival (Figure 2, Table 1). There was, however, little difference in survival between the two *B. oleracea* cvs (Figure 2, Table 1). Simplification of the survival models by using plant species rather than plant variety as an explanatory factor resulted in no significant loss of deviance with respect to survivorship ($F_{6,127} = 0.80$, P = 0.58), fecundity ($F_{4.53} = 0.93$, P = 0.45), or development time ($F_{4.73} = 1.93$, P = 0.11).

Resistance benefited insects in the presence of sprays and imposed fitness costs in the absence of Bt primarily through survival (Figure 2) (full ANOVA in Table 1). Application of Bt reduced survivorship and this interacted with insect genotype (Figure 2). Fitness costs were recessive in this experiment since resistant insects had poorer survival relative to F1 heterozygotes on control plants (difference estimate 1.33, df = 143, t = -4.27, P < 0.001) whereas there was no difference between F1 crosses and susceptible insects in the absence of Bt (difference estimate -0.11, df = 142, t = 0.369, P = 0.71). In contrast, the net fitness benefit of resistance alleles (ie the benefit minus the cost of resistance) in the presence of Bt was effectively dominant on B. pekinensis ($D_{WT=} 1.13$, but was not significantly different from 1.0 as the fitness of RS and SS insects was not statistically distinguishable). On B. oleracea var. Wheeler's Imperial the net benefit of resistance was effectively overdominant ($D_{WT=} 2.55$), while on B. oleracea var. Red Drumhead no homozygous resistant insects survived Bt sprays, and survival of RS insects was 5.9%.

aside the impact of fitness costs, the impact of resistance was semi-dominant in this experiment.

Host plant species affected the strength of fitness costs and the fitness advantage of homozygous resistant insects in sprayed treatments. In both sprayed and control treatments homozygous resistant insects incurred additional reductions in fitness on B. oleracea plants relative to fully susceptible (difference estimate -1.46, df=142, t=-2.06, P=0.04) and heterozygote insects (difference estimate -1.91, df=141, t=-2.84 P=0.005) (Figure 2). There was a trend in which mean survivorship was highest for heterozygotes (RS) on B. oleracea (Figure 2) increasing from 11% for homozygous susceptibles to 18% for heterozygotes. However, post-hoc treatment comparisons indicated that the survivorship of these RS larvae was not significantly greater than fully susceptible (SS) larvae in the absence of sprays (difference estimate =0.45, se of difference =0.413, df=139, t=-1.08, P=0.28). The power of this test (ie the probability of rejecting a false null hypothesis) was estimated at 1.0 (Fleiss $et\ al$, 1980). It would therefore be unjustified to interpret these data as evidence of heterozygote advantage.

Time to emergence and fecundity of females from independent crosses was recorded in this experiment. However, low survivorship in some treatment combinations (e.g. sprayed susceptible insects or resistant insects on B. oleracea) led to low statistical power for investigating some interactions. There was however sufficient replication to explore main effects. The fecundity data, a total of 179 crosses from 66 independent replicates (plants) showed that exposure to Bt did not affect fecundity (number of fertile eggs laid- $F_{1,61} = 0.92$, P = 0.34), nor did host plant species ($F_{2,62} = 0.28$, P = 0.60). Only genotype significantly affected fecundity ($F_{2,62} = 7.36$, $F_{2,62} = 0.0014$). The mean fecundity (and se) of different genotypes was as follows: SS insect 158 fertile eggs (10.4), RS insects 159 fertile eggs (8.6), RR insects 85.7 fertile eggs (16.7). The development time of survivors was extended on $F_{2,82} = 5.35$, $F_{2,82}$

Meta-analysis of fitness cost experiments with the Karak strain

The meta-analysis of fitness cost experiments with *B. pekinensis* estimated the relative risk of survival for resistant (RR) insects at 0.72 (95% CLs 0.62, 0.83; 99.9% CLs 0.57, 0.92) indicating that the probability of resistant larvae surviving to adulthood is 0.72 that of susceptible larvae (Figure 3). We investigated whether different methods of controlling for genetic background affected estimates of fitness costs by conducting a test for homogeneity in estimates of relative risk between the different studies. We found strong evidence of heterogeneity between experiments (Woolf's $\chi^2 = 34.9$, df = 6, P << 0.001). The estimates of fitness costs were greatest in the backcrossed Kar/UK₆/Newc population and significantly higher in this study relative to the two studies using revertant susceptible strains, as indicated by the non-overlapping confidence intervals (Figure 3a). When we pooled studies according to estimation method there was still significant heterogeneity in the dataset (Woolf's $\chi^2 = 16.1$, df = 2, P << 0.001) and fitness costs estimates in the backcross studies (RR = 0.58; 95% CL 0.52-0.64) did not have overlapping confidence limits with the studies using revertant lines (RR = 0.79; 95%CL 0.71-0.89). Fitness costs in the B. oleracea experiments were much higher than in the *B. pekinensis* experiments (relative risk = 0.17, 95% CLs 0.11. 0.25, 99.9% CIs = 0.08, 0.33, Figure 3). There was, however, no significant between-study heterogeneity in estimates of fitness costs in the B. oleracea experiments (Woolf's $\chi^2 = 2.69$, df = 3, P = 0.442).

Discussion

Our experimental work with backcrossed lines showed that fitness costs are so high on *B. oleracea* that homozygous RR insects are effectively carrying a near-lethal recessive mutation. When exposed to the *Bt*-based biopesticide DiPel homozygous resistant insects on *B. oleracea* could not effectively express resistance as they rarely survived beyond the first instar. Heterozygous resistant insects, however, which do not carry fitness costs, did have significant resistance to a low dose of *Bt* (5-10% of field dose) on *B. oleracea*. The meta-analysis confirmed that fitness costs of resistance genes are much greater on mature *B. oleracea* than on the higher quality *B. pekinensis* plants. Fitness costs have been shown to vary with host plant in other insect species (Bird and Akhurst, 2007; Janmaat and Myers, 2005). However, costs do not always increase on better defended varieties (Carrière *et al*, 2005). Nevertheless the very high level of costs shown on *B. oleracea* plants confirms that variation within crop species may be exploitable as a means of improving resistance management strategies (Carrière *et al*, 2005; Carrière *et al*, 2004; Raymond *et al*, 2007a).

The genetics of resistance has a fundamental impact on the management and evolution of resistance to pesticides (Gould, 1998; Roush and McKenzie, 1987). A key aim of the highdose refuge strategy for the management of resistance to Bt is to ensure that the expression levels of Bt toxins in transgenic crops are high enough to kill resistant heterozygotes and ensure that resistance is effectively recessive (Tabashnik et al, 2004). In this study, bioassays with third instars insects showed that resistance became increasingly recessive as the dose of Bt increased (Sayyed et al, 2004) whereas in fitness experiments initiated with neonate larvae the survival of heterozgotes is roughly equal to homozygous resistant individuals, suggesting dominant resistance. Survivorship is however a complex trait affected by the genetics of resistance and the genetics of the fitness costs of resistance. In P. xylostella fitness costs are thought to predominantly act on neonate survival (Raymond et al, 2005) and therefore bioassays with third instar instars do not reveal the effects of fitness costs but only indicate the genetics of resistance itself. In contrast, neonate- adult survival incorporates both the genetics of resistance and the genetics of pleiotropic fitness costs. Therefore, in common with many resistance traits, the fitness costs of resistance to Bt in this study were entirely recessive (Gassmann et al, 2009) while the dominance of resistance itself was incomplete and dose dependent.

A meta-analysis of independent fitness cost studies with varying methodologies may not be as rigorous a means of hypothesis-testing as a designed experimental analysis of resistant and susceptible lines produced under different techniques. Nevertheless, fitness cost experiments with genetically homogenized insects are rare, and data sets that can examine the efficacy of different methodologies applied to the same population are even rarer. We therfore think it worthwhile to explore and discuss the patterns in our data sets with this broad caveat in mind (Bergelson and Purrington, 1996). With a repeatedly backcrossed population we can infer with increased confidence that the observed fitness costs are pleiotropic consequences of the resistance gene itself rather than the result of genes linked with this locus (Bergelson and Purrington, 1996). Thus, eliminating the effects of genetic background may allow a more precise estimation of the pleiotropic consequences of the resistance mutation and this method has been hailed as a gold standard for fitness cost studies (Bergelson and Purrington, 1996). One of the most surprising aspects of the two meta-analyses is the extent to which the studies which did not control for genetic background agreed with experiments using selection-based or homogenization protocols, a conclusion reached in an independent review of the Bt resistance literature (Gassmann et al, 2009). This may be because fitness costs are large and genes at other loci that have major effects on fitness in laboratory conditions are rare, or quickly lost in culture. Gassman et al. (2009) also showed that stability experiments, which measure the decline in resistance over

one or more generations tended to identify fitness costs more readily than fitness components experiments. Gassman *et al.* (2009) did not, however, distinguish between the results of studies that used different methods to control for genetic background.

In agreement with Bergelson & Purrington (1996) we found that observed fitness cost for insects feeding on one species (B. pekinensis) were elevated in studies using homogenized backcrosses compared to studies using selection based approaches, ie revertant susceptible lines. Studies with revertant lines may underestimate fitness costs if revertant lines harbour moderate frequencies of resistance genes. Alternatively, epistatic interactions between resistance genes and other loci can affect the strength of pleiotropic costs. We might expect that the genetic background with which the resistance genes were associated in the field to moderate fitness costs relative to a novel genetic background (Clarke and McKenzie, 1987; Colegrave and Buckling, 2005; Maclean et al. 2004). However, our estimate of the frequency of resistance alleles in the revertant Karak line was 0.375 prior to experiments (Raymond unpubl. dat. 2003). Given recessive fitness costs and a relative survivorship of 0.72 for RR homozygotes (Figure 2) a revertant susceptible population might have a relative fitness of approximately 0.96 that of SS homozygotes. This very modest decrease does not seem sufficient to explain the lower fitness costs observed in the revertant line studies, while epistatic interactions with the original genetic background remain a plausible cause of these lower costs.

A range of studies has indicated that the fitness costs of diverse classes of resistance tend to be elevated in stressful environments (Carrière et al, 2001; Gazave et al, 2001; Janmaat and Myers, 2005; Kraaijeveld and Godfray, 1997; Raymond et al, 2007a; Raymond et al, 2005) although there has not yet been a satisfactory explanation for this common phenomenon. In this study, near-complete replacement of the original genetic background of a field-selected population resulted in increased estimates of fitness costs when resource quality was high while replacement of genetic background had no effect on fitness costs when resource quality was low. These data imply that genetic background is more important for expression of fitness costs when resources are abundant or environmental quality high. One simple hypothesis that would explain this resource-dependent effect is that compensatory mutations or fitness modifiers are more effective at reducing fitness costs in P. xylostella when environmental quality is high. Compensation for the deleterious effects of new mutations may be widespread within insect genomes (Kulathinal et al, 2004). In some cases compensation can arise through interactions with genes already present in populations prior to selection for resistance (Cohan et al, 1994). If so, a long period of selection for resistance may not be required to produce compensatory mutations after resistance has increased to a high frequency.

One aim of this study was to compare the efficacy of different methodologies for measuring fitness costs. Our recommendation for the most suitable method is that it may depend upon (a) the research question and (b) whether the genetic background in the original insect is likely to be affecting the size of pleiotropic fitness costs. A wide range of methods for controlling for the effects of genetic background gave quite similar and robust results over a number of years suggesting that diverse methods can give reliable estimates of fitness costs. However, the replacement and homogenization of genetic background can lead to higher estimates of fitness costs relative to other techniques, as has been found in plants (Bergelson and Purrington, 1996). A homogenization approach will give the most reliable answer if the research question has the purely genetical aim of estimating the fitness consequences of a particular mutation. On the other hand, if the research question is to estimate the size of the fitness costs experienced by resistant insects in the field then other methods may be superior. If compensatory fitness modifiers in the original genetic background are reducing the fitness costs of resistance then homogenization methods may overestimate the costs of resistance

experienced in the field and it would be better to estimate fitness costs in the original genetic background. On the other hand, if observed fitness costs are inflated due to the action of genes linked to resistance loci that may have become fixed via inbreeding in the laboratory, genetic homogenization and repeated backcrossing provides extra assurance that any observed fitness costs are caused by the gene of interest.

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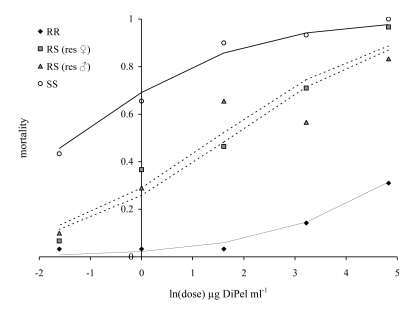


Figure 1.
Relative resistance of homozygous resistant (RR), heterozygous resistant (RS) and fully susceptible (SS) third instar larvae of diamondback moth larvae in leaf-dip bioassays. Data for the reciprocal F1 crosses used to produce RS heterozygotes have been plotted separately and the sex of the resistant parent indicated. Data are five-day mortality; lines represent fitted statistical models from a binomial glm.

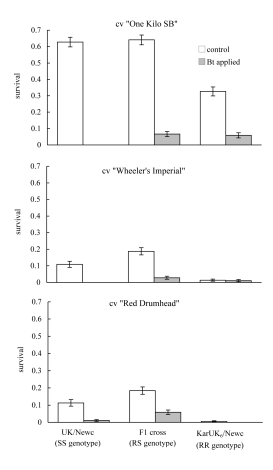


Figure 2. Survival (neonate – adult) of a Bt resistant population of diamondback moth (KarUK $_6$ /Newc), a susceptible population (UK/Newc) and their F1 hybrids on B. pekinensis (cv "One Kilo SB") and B. oleracea (cvs "Wheeler's Imperial" and "Red Drumhead"). Clear bars represent the spray-free treatment; shaded bars the Bt sprayed treatments. N=280 larvae on 8 host plants on B. pekinensis or 320 larvae on eight plants of B. oleracea. RR, RS and SS indicate the homozygous resistant, heterozygous resistant and homozygous susceptible genotypes respectively. Data are means and SEs (calculated using the normal approximation to the mean).

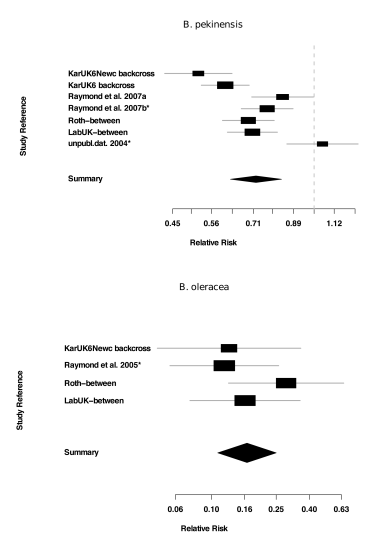


Figure 3.
Random effects meta-analyses of studies using diverse methods to estimate the fitness costs of resistance to *Bacillus thuringiensis* in diamondback moth feeding on two host plant species, *B. pekinensis* and *B. oleracea*. Fitness costs were calculated as the relative risk of survival to adulthood for resistant and susceptible larvae. The horizontal lines indicate the 95% confidence interval for each study and the point estimate is given by a square whose height is inversely proportional to the standard error of the estimate. The summary relative risk estimated from the whole data set is drawn as a diamond with horizontal limits at the confidence limits and width inversely proportional to its standard error. Experiments that used revertant susceptible lines to calculate fitness costs are denoted with an asterisk *; the studies denoted by "backcross" used genetic homogenization techniques as described in the methods; all other studies were between population designs that did not control for genetic background.

Table 1

ANOVA of larval survival for Bt resistant P. xylostella, Bt susceptible P. xylostella and their heterozygotes grown on two Brassica spp. of varying suitability. Data are logit-transformed survival (neonate-adult), the analysis used a generalized linear model with correction for over-dispersion.

	d.f.	Deviance	Resid. df	Resid. df Resid. Dev.	F
Null model			144	1817.5	
plant species	-	482.1	143	1335.4	160.4 ***
population	2	131.8	141	1203.6	22.0 ***
treatment	2	677.2	140	526.4	225.3 ***
plantXpopn	-	37.4	138	495.0	5.21 **
plantXtrtmt	4	35.7	137	461.0	11.36 ***
popnXtrtmt	2	51.8	135	410.9	8.33 ***
plantXpopnXtrtmnt 4	4	10.5	133	405.2	0.95

Significance terms:

p < 0.001p < 0.01