Field Evolved Resistance in *Helicoverpa armigera* (Lepidoptera: Noctuidae) to *Bacillus thuringiensis* Toxin Cry1Ac in Pakistan

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Abstract

*Helicoverpa armigera* (Hübner) is one of the most destructive pests of several field and vegetable crops, with indiscriminate use of insecticides contributing to multiple instances of resistance. In the present study we assessed whether *H. armigera* had developed resistance to Bt cotton and compared the results with several conventional insecticides. Furthermore, the genetics of resistance was also investigated to determine the inheritance to Cry1Ac resistance. To investigate the development of resistance to Bt cotton, and selected foliar insecticides, *H. armigera* populations were sampled in 2010 and 2011 in several cotton production regions in Pakistan. The resistance ratios (RR) for Cry1Ac, chlorpyrifos, profenofos, cypermethrin, spinosad, indoxacarb, abamectin and deltamethrin were 580-fold, 320-, 1110-, 1950-, 200-, 380, 690, and 40-fold, respectively, compared with the laboratory susceptible (Lab-PK) population. Selection of the field collected population with Cry1Ac in 2010 for five generations increased RR to 5440-fold. The selection also increased RR for deltamethrin, chlorpyrifos, profenofos, cypermethrin, spinosad, indoxacarb, abamectin to 125-folds, 650-, 2840-, 9830-, 370-, 3090-, 1330-fold. The estimated LC₅₀ for reciprocal crosses were 105 µg/ml (Cry1Ac-SEL female × Lab-PK male) and 81 g µg/ml (Lab-PK female × Cry1Ac-SEL male) suggesting that the resistance to Cry1Ac was autosomal; the degree of dominance (D₅₀) was 0.60 and 0.57 respectively. Mixing of enzyme inhibitors significantly decreased resistance to Cry1Ac suggesting that the resistance to Cry1Ac and other insecticides tested in the present study was primarily metabolic. Resistance to Cry1Ac was probably due to a single but unstable factor suggesting that crop rotation with non-Bt cotton or other crops could reduce the selection pressure for *H. armigera* and improve the sustainability of Bt cotton.

Introduction

The cotton bollworm, *Helicoverpa armigera* (Hübner) (Noctuidae), is one of the most damaging and cosmopolitan pests causing significant economic loss to a wide range of field and vegetable crops [1]. Due to its wider host range, high fecundity, multiple generations, migratory behavior and insecticide resistance, it has become a much more difficult pest to manage [2]. The frequent use, distribution, and reproduction in any medium, provided the original author and source are credited.

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treated with a non-Bt foliar insecticide, or as a 4% refuge of non-transgenic plants that are left untreated.

Strategies for delaying insect pest resistance to cotton and maize expressing Bt Cry toxins were implemented from the introduction of these transgenes in 1996 and have so far proven to be effective in the US and other developed countries. In contrast, *Pectinophora gossypiella* from India and China has been shown to develop resistance to Bt transgenic cotton [9,10]. We were therefore interested in examining a similar trend in Pakistan, in *H. armigera*, as most of the growers in Pakistan do not follow the HRD strategy. We therefore surveyed the primary cotton growing areas of Pakistan to investigate whether *H. armigera* has developed resistance to the Bt toxin Cry1Ac after exposure to Bt cotton in the field. We further examined the number genes involved in resistance to Cry1Ac and mechanisms involved in resistance to Cry1Ac and chemical insecticides.

**Results**

**Toxicity of Insecticides to a Laboratory Susceptible Population and Field Population**

Toxicity of chlorpyrifos, profenofos Cry1Ac, indoxacarb and deltamethrin to the laboratory susceptible, Lab-PK was similar (overlapping of 95% FL; *P* > 0.05), but higher for cypermethrin and abamectin (Table 1). In contrast, the toxicity of spinosad was significantly lower (non-overlapping of 95% FL; *P* < 0.05) than cypermethrin and abamectin but was similar to other insecticides tested (Table 1). The slopes for all insecticides tested against Lab-PK were similar, but more shallow indicating that the response in the laboratory susceptible population to tested insecticides was heterogeneous.

The toxicity of all insecticides tested against a field collected population was significantly lower (*P* < 0.05) at G1 compared with Lab-PK. The resistance ratios for chlorpyrifos, profenofos, Cry1Ac, cypermethrin, spinosad, indoxacarb, abamectin and deltamethrin were 320-fold, 1110-, 580-, 1950-, 200-, 380, 690, and 40-fold respectively (Table 1). The slopes of the regression lines for the insecticides tested against field population at G1 were significantly steeper for chlorpyrifos, profenofos, Cry1Ac, cypermethrin, indoxacarb, abamectin and deltamethrin compared with Lab-PK, suggesting a homogenous response in the field collected population to these insecticides. The slope of spinosad however was more shallower than other insecticides but it was similar to Lab-PK (Table 1).

The response of *H. armigera* to Cry1Ac collected from various locations was similar; however the highest resistance ratio was obtained for the population collected from Multan (Fig. 1).

**Table 1.** Toxicity of various insecticides to laboratory susceptible (Lab-PK) and field collected populations of *H. armigera*.

| Population | Insecticides | LC50 (95% FL) (μg/ml) | Slope ± SE | RR1 | DR2 | n3 |
|------------|--------------|------------------------|------------|-----|-----|----|
| Susceptible | Cry1Ac | 0.58 (0.28–1.20) | 1.07±0.16 | – | – | 240 |
| Susceptible | Chlorpyrifos | 0.46 (0.18–0.96) | 1.03±0.17 | – | – | 240 |
| Susceptible | Profenofos | 0.50 (0.20–1.03) | 1.06±0.17 | – | – | 240 |
| Susceptible | Cypermethrin | 0.26 (0.12–0.50) | 1.20±0.17 | – | – | 240 |
| Susceptible | Spinosad | 1.45 (0.69–2.79) | 1.13±0.18 | – | – | 240 |
| Susceptible | Indoxacarb | 0.90 (0.48–1.71) | 1.27±0.17 | – | – | 240 |
| Susceptible | Abamectin | 0.23 (0.08–0.54) | 0.85±0.14 | – | – | 240 |
| Susceptible | Deltamethrin | 0.42 (0.17–0.78) | 1.27±0.22 | – | – | 240 |
| Field | Cry1Ac | 335.7 (244.2–477.6) | 2.43±0.31 | 579 | – | 240 |
| Field | Chlorpyrifos | 148.7 (106.3–217.0) | 2.29±0.27 | 323 | – | 240 |
| Field | Profenofos | 557.4 (420.3–734.5) | 3.20±0.46 | 1115 | – | 240 |
| Field | Cypermethrin | 506.8 (387.7–680.7) | 3.28±0.44 | 1949 | – | 240 |
| Field | Spinosad | 284.7 (180.0–484.7) | 1.74±0.30 | 196 | – | 240 |
| Field | Indoxacarb | 341.2 (260.8–459.7) | 3.08±0.34 | 379 | – | 240 |
| Field | Abamectin | 159.2 (121.3–208.8) | 3.15±0.53 | 692 | – | 240 |
| Field | Deltamethrin | 16.02 (12.74–20.32) | 3.49±0.45 | 38 | – | 240 |
| Field-UNSEL (G6) | Cry1Ac | 19.65 (11.56–33.04) | 1.50±0.22 | 34 | –0.21 | 240 |
| Field-UNSEL (G6) | Chlorpyrifos | 11.75 (7.96–17.83) | 2.04±0.25 | 26 | –0.18 | 240 |
| Field-UNSEL (G6) | Profenofos | 18.48 (12.30–27.83) | 2.05±0.25 | 37 | –0.25 | 240 |
| Field-UNSEL (G6) | Cypermethrin | 20.07 (12.71–31.48) | 1.83±0.24 | 77 | –0.23 | 240 |
| Field-UNSEL (G6) | Spinosad | 9.22 (5.76–14.81) | 1.66±0.23 | 6 | –0.25 | 240 |
| Field-UNSEL (G6) | Indoxacarb | 8.07 (4.04–14.37) | 1.29±0.20 | 9 | –0.27 | 240 |
| Field-UNSEL (G6) | Abamectin | 9.27 (5.88–14.79) | 1.77±0.24 | 40 | –0.21 | 240 |
| Field-UNSEL (G6) | Deltamethrin | 1.76 (1.14–2.61) | 1.90±0.30 | 4 | –0.16 | 240 |

1. Resistance ratio = LC50 of field and unselected (UNSEL) population / LC50 of Lab-PK.
2. Decline in resistance = Log10 (initial LC50–final LC50) / number of generation the population of unexposed to toxicant.
3. The number of larvae exposed to toxins in bioassays including controls.
4. Generations.

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Response to Selection and Cross-resistance in Cry1Ac-SEL Population

Mortality at different selection doses of 300, 500 and 1000 µg AI mL⁻¹, determined after 7 days exposure to Cry1Ac were 59, 35, 15 and 40% respectively. Selection of the field population with Cry1Ac from G1 to G5 increased the resistance ratio (RR) to 160-fold for Cry1Ac compared with the Unselected field population. However, when it was compared with Lab-PK, the RR increased from 580-fold to 5440-fold (just five generations of selection).

Similarly, selection with Cry1Ac also increased RR for deltamethrin, chlorpyrifos, profenofos, cypermethrin, spinosad, indoxacarb, abamectin, and abamectin.

Table 2. Cross-resistance and instability pattern in a Cry1Ac-selected (Cry1Ac-SEL) population of H. armigera.

| Population                  | Insecticides | LC50 (95% FL) (µg/ml) | Slope ± SE | RR¹ | RR² | DR³ | n⁴ |
|-----------------------------|--------------|-----------------------|------------|-----|-----|-----|-----|
| Cry1Ac-SEL (G6)⁵             | Cry1Ac       | 3154 (2548–4000)      | 3.65 ± 0.47| 5438 | 160 | –   | 240 |
| Cry1Ac-SEL (G6)              | Chlorpyrifos | 1220 (957.3–1585)     | 3.28 ± 0.39| 2652 | 104 | –   | 240 |
| Cry1Ac-SEL (G6)              | Profenofos   | 1421 (998.8–2022)     | 2.31 ± 0.28| 2842 | 77  | –   | 240 |
| Cry1Ac-SEL (G6)              | Cypermethrin | 2557 (2093–3164)      | 4.06 ± 0.50| 9834 | 127 | –   | 240 |
| Cry1Ac-SEL (G6)              | Spinosad     | 533.3 (322.2–836.6)   | 1.88 ± 0.30| 368  | 58  | –   | 240 |
| Cry1Ac-SEL (G6)              | Indoxacarb   | 2779 (2293–3424)      | 4.24 ± 0.51| 3088 | 344 | –   | 240 |
| Cry1Ac-SEL (G6)              | Abamectin    | 306.4 (222.1–430.4)   | 2.52 ± 0.31| 1332 | 33  | –   | 240 |
| Cry1Ac-SEL (G6)              | Deltamethrin | 52.86 (36.71–89.46)   | 2.79 ± 0.41| 125  | 18  | –   | 240 |
| Cry1Ac-UNSEL (G10)           | Cry1Ac       | 199.2 (141.6–275.7)   | 2.36 ± 0.35| 343  | 10  | −0.30| 240 |
| Cry1Ac-UNSEL (G10)           | Chlorpyrifos | 93.20 (64.45–135.9)   | 2.17 ± 0.28| 203  | 8   | −0.28| 240 |
| Cry1Ac-UNSEL (G10)           | Profenofos   | 98.61 (65.18–147.4)   | 2.03 ± 0.25| 197  | 5   | −0.29| 240 |
| Cry1Ac-UNSEL (G10)           | Cypermethrin | 204.6 (135.5–268.9)   | 2.96 ± 0.40| 787  | 10  | −0.27| 240 |
| Cry1Ac-UNSEL (G10)           | Spinosad     | 69.11 (44.33–105.4)   | 1.81 ± 0.27| 48   | 7   | −0.22| 240 |
| Cry1Ac-UNSEL (G10)           | Indoxacarb   | 111.8 (70.27–162.5)   | 1.99 ± 0.32| 124  | 14  | −0.35| 240 |
| Cry1Ac-UNSEL (G10)           | Abamectin    | 50.38 (34.99–71.88)   | 2.26 ± 0.31| 219  | 5   | −0.20| 240 |
| Cry1Ac-UNSEL (G10)           | Deltamethrin | 10.04 (7.85–12.89)    | 3.59 ± 0.44| 24   |     |     |     |

¹Resistance ratio = LC50 of Cry1Ac-SEL and unselected (UNSEL) populations/LC50 of Lab-PK.
²Resistance ratio = LC50 of Cry1Ac-SEL and Cry1Ac unselected populations/LC50 of UNSEL (Table 1).
³Decline in resistance = Log10(initial LC50–final LC50)/number of generation the population of unexposed to toxicant.
⁴The number of larvae exposed to toxins in bioassays including controls.
⁵Generations.

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carb, abamectin to 125-fold, 650-, 2840-, 9830-, 370-, 3090- and 1330-fold, compared with Lab-PK (Table 2).

Selection with Cry1Ac also increased the slope for the probit line for the Cry1Ac-SEL population compared with the field population at G1. The slope for the insecticides tested against Cry1Ac-SEL also increased significantly indicating increase in homogeneity in the selected population.

The bioassays on various Bt cotton varieties revealed that the survival of Cry1Ac-SEL population was almost 100% on all varieties which was significantly higher than any other strain tested (Fig. 2). In contrast, 100% mortality was obtained with the Lab-PK population while the UNSEL and field population response was similar among varieties (Fig. 2).

Reversion of Resistance to Cry1Ac and Other Insecticides in Field and Cry1Ac-SEL Populations

In order to investigate the stability of resistance to Cry1Ac and other insecticides in field populations and Cry1Ac-SEL, Cry1AC-SEL was maintained for 6 generations without exposure to insecticides. Bioassays of field populations at G5 showed a significant reduction (P<0.05) in resistance ratio with a reversion rate of 0.21. Similarly, rearing field population without exposure to insecticides also reduced ratios for chlorpyrifos, profenofos, cypermethrin, spinosad, indoxacarb and abamectin (Table 1). The reversion rate of resistance to deltamethrin in the field population was the least 0.16 while it was the highest for indoxacarb (0.27; Table 1).

The Cry1Ac-SEL population was also monitored for reversion of resistance to deltamethrin and other insecticides for four generations (G7–G10). Bioassays carried out at G11 indicated that resistance to Cry1Ac declined significantly (P<0.05) in four generations, near the level of field evolved resistance. Likewise, resistance to chlorpyrifos, profenofos, cypermethrin, spinosad, indoxacar, abamectin and deltamethrin was also reduced significantly from G6 to G10. The rate of decline of resistance in Cry1Ac-SEL population was similar to the reversion rate for the field selected population.

The parameter R is used to estimate response to selection (Falconer, 1989) which can also be applied to determine the number of generations required to a 10-fold change in resistance. The inverse of R is the number of generations required for a 10-fold change in LC50. The R value for Cry1Ac is 0.21 suggesting that only five generations are required to increase 10-fold resistance to Cry1Ac.

Inheritance of Cry1Ac Resistance

The LC50 of Cry1Ac for Cry1Ac-SEL population was over 3000-fold which was significantly higher than LC50 of Cry1Ac for Lab-PK. Estimated LC50 for F1 female progeny from reciprocal crosses were 105 μg/ml (Cry1Ac-SEL female × Lab-PK male) and 81g μg/ml (Lab-PK female × Cry1Ac-SEL male). The LC50 values and mean slopes for the concentration mortality line did not differ

| population | LC50 (95% FL) (μg/ml) | Slope ± SE | Dc | 
|------------|-----------------------|------------|----|
| Cry1Ac-SEL (G6) | 3154 (2548–4000) | 3.65±0.47 | – |
| Lab-PK | 0.58 (0.28–1.20) | 1.07±0.16 | – |
| Cry1Ac-SEL6 × Lab-PK | 105.5(70.69–163.0) | 1.76±0.21 | 0.60 |
| Lab-PK6 × Cry1Ac-SEL | 81.21(52.19–129.3) | 1.56±0.20 | 0.57 |
| F1 × Cry1Ac-SEL | 2.68(1.86–4.10) | 0.98±0.23 | – |

1Degree of dominance at LC50, which was calculated as described previously [24].
2Generations.
significantly between F₁ progeny of the reciprocal crosses between Lab-PK and Cry1Ac-SEL, suggesting that the inheritance of resistance to Cry1Ac was autosomal; neither maternal effects nor sex linkages were evident (Table 3). The degree of dominance (D₁₄C) for the reciprocal crosses was 0.60 for F₁ (Cry1Ac-SEL female × Lab-PK male) and 0.57 for F₁ (Lab-PK female × Cry1Ac-SEL male), indicating incomplete dominance of resistance to Cry1Ac in Cry1Ac-SEL population.

Pooled F₁ progeny were backcrossed to the Lab-PK colony, resulting in a slope of 0.98; this is similar to Lab-PK but about 4-fold less than the estimated slope for Cry1Ac-SEL, about 2-fold less than F₁ progeny indicate decreased genetic variance in the backcrossed progeny compared with F₁ progeny. The decreased genetic variance indicates the number of loci with major effect on resistance to Cry1Ac was very low. Similarly the direct test for a monogenic mode of inheritance of resistance, which is based on the goodness of fit between observed and expected mortality at seven (Table 3). Likewise, calculation of the minimum number of independently segregating loci with equal and additive contributions to resistance had given an estimate of 0.34, which also supported the conclusion that resistance was conferred by a single locus.

Resistance to Insecticides is Inhibited by Synergists

The synergistic effects of PBO and DEF on Cry1Ac and other insecticides were determined in Lab-PK and Cry1Ac-SEL populations of *H. armigera*. The monooxygenase specific inhibitor PBO showed 273-fold synergism to Cry1Ac in the Cry1Ac-SEL population at G6; however, no synergism was observed for Lab-PK (Table 4). Only a 2-fold level of resistance remained after the application of PBO and Cry1Ac together, suggesting that the major mechanism was associated with mono-oxidases or esterases since PBO has also been shown to inhibit the activity of esterases [12]. When the esterase specific inhibitor DEF was used, a high level of synergism (73-fold) was observed against the Cry1Ac-SEL population but no effect of DEF was detected in Lab-PK (Table 4). The occurrence of synergism for both inhibitors suggests that enhanced activities of esterases, or probably mono-oxidases, are involved in resistance to Cry1Ac in *H. armigera*.

Similarly, when insecticides like profenofos and indoxacarb were used in a mixture with either PBO or DEF against Cry1Ac-SEL population, a high level of synergism was observed but there was no effect of inhibitors on Cry1Ac toxicity against Lab-PK. The most intriguing observation was synergism of PBO or DEF with Cry1Ac against Cry1Ac-SEL population (Table 4).

### Discussion

Our data suggest that *H. armigera* has developed resistance to Cry1Ac in the field in Pakistan. In Pakistan illegal planting of Bt cotton has occurred since 1999, without following the HRD strategy. Previous studies from India and China had reported field evolved resistance to Cry1Ac in *Pectinophora gossypiella* [9,13]. In the present study, our data show a high level of resistance, not only to Cry1Ac, but also to conventional insecticides such as pyrethroids and organophosphates. To confirm whether the resistance to Cry1Ac and conventional insecticides was associated with the same mechanism of resistance, selection experiments were performed in the laboratory with Cry1Ac on a field collected population. After six generations of selection, resistance to Cry1Ac increased significantly (non-overlapping of 95% FL). Cross-resistance patterns in Cry1Ac-SEL could result from enzymes such as metabolic enzymes [8,12] and mutation at an insecticidal target site [3]. The high level of resistance shown by the Cry1Ac-SEL population suggests either a common mechanism affecting these insecticides or genetically linked independent mechanisms for Bt toxin Cry1Ac and deltamethrin. The findings of the present study are similar to previously reported results of *Plutella xylostella* resistance to deltamethrin, which showed a high level of reciprocal cross-resistance to Cry1Ac [8]. Similarly, our data also suggest that the resistance to Cry1Ac in the selected population was due to involvement of metabolic enzymes as was previously shown for *P. xylostella* [8] or *H. armigera* [12]. The metabolic enzymes have several isoenzymes that can act on different insecticides; if an insecticide selects some isoenzymes that can affect different insecticides then cross-resistance is possible [14]. When PBO, mono-oxidase or esterases inhibitor or DEF esterases inhibitors were used in combination with Cry1Ac or deltamethrin, the resistance to both toxicants was reduced significantly suggesting that the major mechanism of resistance to Cry1Ac or deltamethrin was associated with esterases. We also carried out bioassays with profenofos and indoxacarb in the presence of PBO or DEF and the data suggest that the resistance to the insecticides was

### Table 4. Susceptibility of Cry1Ac-selected (Cry-SEL) populations of *H. armigera* to Cry1Ac and other insecticides tested in the presence or absence of a PBO or DEF.

| Population | Treatment | LC₅₀ (95% FL) | Slope ± SE | RR¹ | SR² |
|------------|-----------|---------------|-----------|-----|-----|
| Lab-PK     | Cry1Ac    | 0.58 (0.28–1.20) | 1.07 ± 0.16 | –   | –   |
|            | Cry1Ac+ PBO | 0.50 (0.39–1.50) | 3.09 ± 0.46 | –   | 1   |
|            | Cry1Ac+ DEF | 0.48 (0.12–1.34) | 1.44 ± 0.23 | –   | 1   |
|            | Profenofos | 0.50 (0.20–1.03) | 1.06 ± 0.17 | –   | –   |
|            | Profenofos+ PBO | 0.98 (0.52–1.60) | 1.58 ± 0.24 | –   | 1   |
|            | Indoxacarb | 0.90 (0.48–1.71) | 1.27 ± 0.17 | –   | –   |
|            | Indoxacarb+ PBO | 1.05 (0.62–3.10) | 3.54 ± 0.49 | –   | –   |
|            | Indoxacarb+ DEF | 1.29 (0.81–2.44) | 1.79 ± 0.27 | –   | –   |
|            | Deltamethrin | 0.42 (0.17–0.78) | 1.27 ± 0.22 | –   | –   |
|            | Deltamethrin + PBO | 0.14 (0.08–0.43) | 1.56 ± 0.23 | –   | –   |
|            | Deltamethrin+ DEF | 0.09 (0.03–0.23) | 1.23 ± 0.13 | –   | –   |
| Cry1Ac-SEL | Cry1Ac    | 315.4 (254.8–540.3) | 1.65 ± 0.37 | 544 | –   |
|            | Cry1Ac+ PBO | 1.15 (0.56–2.44) | 1.07 ± 0.15 | 2   | 274 |
|            | Cry1Ac+ DEF | 4.31 (2.80–6.66) | 1.82 ± 0.27 | 9   | 73  |
|            | Profenofos | 421 (221.8–602.7) | 2.17 ± 0.29 | 842 | –   |
|            | Profenofos+ PBO | 1.27 (0.76–2.06) | 1.79 ± 0.27 | 1   | 331 |
|            | Indoxacarb | 277.9 (122.2–436.7) | 2.24 ± 0.41 | 309 | –   |
|            | Indoxacarb+ PBO | 1.25 (0.70–2.47) | 1.40 ± 0.18 | 1   | 222 |
|            | Indoxacarb+ DEF | 4.54 (3.14–6.52) | 2.26 ± 0.31 | 4   | 61  |
|            | Deltamethrin | 52.3 (36.7–89.5) | 2.79 ± 0.41 | 125 | –   |
|            | Deltamethrin + PBO | 5.12 (3.12–15.6) | 1.93 ± 0.23 | 37  | 10  |
|            | Deltamethrin+ DEF | 3.25 (1.11–11.3) | 2.12 ± 0.14 | 36  | 16  |

¹ Resistance ratio was LC₅₀ of Cry1Ac or insecticides for Cry1Ac-SEL/LC₅₀ of Cry1Ac or insecticides for Lab-PK.
²The synergism ratio (SR) was calculated from LC₅₀ of Cry1Ac or other insecticides tested/LC₅₀ of Cry1Ac+inhibitor or insecticides+inhibitor.

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associated with metabolic enzyme, esterases. The Cry1Ac-SEL population was derived from a field population which was collected from an area where Bt cotton has been grown since 1999. However pyrethroids and neonicotinoids are also being sprayed to control sucking insect pests such as *Bemisia tabaci*, suggesting that *H. armigera* in Pakistan is not only exposed to Bt toxin Cry1Ac but also to conventional insecticides. The common mechanism of resistance to several insecticides as shown by Cry1Ac-SEL, is literally a result of field exposure. Although laboratory experiments cannot completely reflect field conditions, our results are consistent with the fact that the rapid development of resistance to Cry1Ac and deltamethrin observed in field populations was due to exposure to insecticides in the field.

A field collected population was also selected for susceptibility to different insecticides. It was found that after one year of selection in the laboratory for 14 generations, the susceptibility to insecticides increased significantly. The resulting Lab-PK was significantly more susceptible to insecticides than another laboratory population of *H. armigera* from Pakistan with identical bioassay systems to the present studies [5]. The most probable reason for high susceptibility in the Lab-PK population was due to its collection from a non-cotton growing (Islamabad). Resistance alleles, if present at the time of collection, were likely lost due to rearing the pest without exposure to insecticides.

Data obtained on the stability of the mechanism of resistance in the selected or the field collected population suggest instability of resistance as the LC90 decreased significantly in the absence of selection. Rapid reversion of resistance in the Cry1Ac-SEL population and field collected population suggest that high fitness costs may be associated with resistance. The decline in resistance could also be due to the presence of heterozygotes in the selected population. A high level of resistance to conventional insecticides has been reported to decline rapidly in populations selected in the laboratory or in the field [15]. The rapid decline in resistance in the Cry1Ac-SEL population indicates that if the toxin is removed from the field, the resistance could also decline quickly. However, this is unlikely in Pakistan as farmers are maintaining Bt cotton throughout the year as a ratoon crop. This is a serious concern and will likely increase the selection pressure, allowing resistance to rapidly spread to other areas of Pakistan.

Reciprocal crosses between resistant and susceptible populations can provide information on dominance of resistance genes, sex linkage and the number of genes involved in resistance to insecticides. The results of these crosses between Cry1Ac-SEL and Lab-PK showed no significant difference in LC50s of reciprocal crosses, suggesting that the resistance was autosomal and no sex linkage was observed. Like resistance to deltamethrin and indoxacarb in *Spodoptera litura*, and Cry1Ac, deltamethrin and spinosad resistance in *P. xylostella* from Pakistan, resistance in *H. armigera* was incompletely dominant. The resemblance of genetics of resistance among various compounds could be due to similarity in selection protocols as the selection is carried out with the aim of having about 30% survival of exposed larvae, which could lead to an incompletely dominant mode of inheritance [14]. However the mode of inheritance at a given concentration of insecticide also depends upon the life stage of an insect especially first instars are generally more susceptible than later instars [8,16]. Although we have not carried out assays at first instar but our observations suggest that F1 progeny of Cry1Ac-SEL was more susceptible at neonate stage than the second instar (the stage which was tested in bioassays). Resistance to insecticides is generally monogenic [17]. The backcrossing of F1 progeny to parents usually support the estimate the number of genes involved in resistance [18] and the data of the present study suggest that resistance to Cry1Ac in the Cry1Ac-SEL population is controlled by single gene. The minimum number of effective factors estimated in the present study are less than 1 (nE <1), which also suggests that resistance is controlled primarily by one locus [19].

Stability of resistance and cross-resistance of Cry1Ac to deltamethrin suggest that common resistance mechanism was linked with the failure of two distant pest management control agents against *H. armigera* from Pakistan. Our data show a significant survival of the Cry1Ac-SEL, UNSEL and field collected *H. armigera* on Bt cotton (Fig. 2) suggesting that even small decreases in susceptibility to Cry1Ac could reduce the efficacy of Bt cotton in the field. Based on the field data described above and our bioassay results, we hypothesize that the magnitude of resistance documented here reduces the efficacy of Cry1Ac producing Bt cotton against *H. armigera* in the field. The results on *Pectinophora gossypella* resistance to Cry1Ac from India and China suggest that the refuge strategy has helped to delay resistance [9]. In contrast our study suggests that field collected population of *H. armigera* survived on Bt cotton. The most probable reasons for the survival are that in Pakistan cotton growers are using Bt cotton varieties which do not have sufficiently high expression of Cry1Ac. Similarly, small land-holding farmers are not using the refuge of non-Bt cotton to increase the population of hybrid progeny. Recently due to high cotton prices in the market, growers in Pakistan are keeping the cotton plants throughout the year as a ratoon crop which is also exposing *H. armigera* a year around to Bt toxin and thus increasing the chances of resistance development.

Most of the Bt cotton varieties being planted in Pakistan were locally developed. Varieties developed by Monsanto were not approved to grow in Pakistan. While in other countries such as Australia and the US key conditions of the HDR strategy are being met and susceptibility to Cry1Ac has not decreased substantially in the target pests, despite a relatively high initial frequency of resistance [12,13]. The most important option to counter resistance in *H. armigera* is to switch to Bt cotton expressing two toxins, Cry2Ab and Cry1Ac [13]. However, for long-term sustainable IRM, cotton with two or more toxins other than Cry1Ac would be better for countering resistance to Cry1Ac [13]. A second option, which is very unlikely in Pakistani agriculture, is to increase plantings of non-Bt cotton. Finally using other control tactics, such as cultural practices in combination with Bt cotton, could be another option to suppress resistance to Bt and conventional insecticides.

**Materials and Methods**

**Insects**

*Helicoverpa armigera* larvae were collected from non-Bt cotton fields, as the treatment regimes used provide a greater chance for the generation of resistance than the regimes used in vegetables. By pest scouting of fields from five districts, namely Multan, Khanewal, Lodhran, Bahawalpur and Rahim Yar Khan approximately 400 larvae were collected in 2010. No specific permit was required to collect insects from the field as the fields were privately owned and merely by speaking to private owners, collection was made. Since the collection was not involved endangered species therefore no such permission was required from any concerned authority in Pakistan. The areas are in Punjab Province and under multiple cropping systems, with several cultivated crops such as cotton, maize, sorghum, millet, rice, sugarcane, wheat, potato, vegetables, fodder crops and orchards (Fig. 3). These crops are grown side by side, depending on the season. An insecticide-susceptible population, labeled as Lab-PK, was collected from...
Multan, Punjab province from non Bt cotton field and selected for susceptibility in the laboratory as described previously [14].

Larvae were reared on semi-synthetic wheatgerm-based diet in the laboratory at 25±2°C and 60-65% relative humidity with a 14:10 h light:dark photoperiod. Diet was replaced after 24 h, and pupae were collected on alternate days. The adults that emerged were kept in Perspex oviposition cages (30×30×30 cm) with two sides sealed with muslin to maintain ventilation and fed on a solution containing sucrose (100 g L⁻¹), vitamin solution (20 mL L⁻¹) and methyl 4-hydroxybenzoate (2 g L⁻¹) presented on a soaked cotton wool ball.

Insecticides

Commercial formulations of the different insecticides used in bioassays included spinosad (Tracers 24SC, Dow Agro-Sciences, UK), indoxacarb (Stewards15SC, DuPont, Pakistan), abamectin (Agrimec TM 1.8EC, Syngenta, UK), cypermethrin 100 g L⁻¹ EC (Arrivo® 10EC; FM, Philadelphia, PA), deltamethrin 105 g L⁻¹ EC (Decis Super® 10.5EC; Bayer Crop Science, France), Profenofos 500 g L⁻¹ EC (Curacron® 50EC; Syngenta Crop Protection, Switzerland), chlorpyrifos 400 g L⁻¹ EC (Lorsban® 40EC; Dow AgroSciences, UK) and Cry1Ac. The source of Cry1Ac was a lyophilized (freeze-dried) formulation of MVP II containing≈20% Cry1Ac protoxin of B. thuringensis variety kurstaki encapsulated by transgenic Pseudomonas fluorescens Migula (Mycogen Corporation, San Diego, CA). It was stored at −80°C until used and before use it was allowed to warm at room temperature. Appropriate amounts of the lyophilized material were weighed for each concentration and suspended in distilled water.

Bioassays

Second instar H. armigera larvae were used for all bioassays, in which the insecticides or Cry1Ac was incorporated into an artificial wheatgerm diet [20]. The Cry1Ac or insecticides were serially diluted with distilled water and then mixed with diet at an appropriate temperature of diet. Toxins incorporated freshly prepared diet was poured into 140-ml plastic Petri-dishes. For
controls, distilled water was mixed with the diet. All assays included seven to eight toxin doses (concentrations) each with three to eight replicates and 30 larvae were placed on each replicate. The Petri-dishes were wrapped black paper to avoid cannibalism [21] and incubated at 27 °C, 70% RH, and a photoperiod of 14:10 (LD) h for 7 d. The Helicoverpa armigera are known to cannibalized however if they are placed in dark place, this behavior can be avoided [21]. Larval mortality and stunting (larvae that failed to molt to third instars) were recorded as response data. Dosage mortality data were analyzed by probit analysis [22].

We used five Bt cotton varieties viz. Bt121, Bt856, Bt456, Bt802 and Bt703, which expressed Cry1Ac to determine survival of field collected, Cry1Ac-SEL, UNSEL and Lab-PK populations on these varieties. These varieties were grown in pots of 45 cm to 30 cm using clay loam soil with farm yard manure as organic fertilizers. The pots were kept in an open field to avoid damage to Cry1Ac as it is an established fact that the expression of toxin declined with the age and also the plants will be less toxic if they are grown in the greenhouse. The five first instar larvae per plant were released on eight weeks old 10 plants of each variety and the larvae were allowed to complete their larval stage. The plants with larvae were placed at 27 °C, 70% RH, and a photoperiod of 14:10 (LD) h.

Selection with Cry1Ac

The field collected population was divided into two sub-populations. One population was selected with Cry1Ac (Cry1Ac-SEL) while the second population was left unselected (UNSEL) for five generations. The selection was done using three concentrations of MIVP viz. 300, 500 and 100 μg Al mL⁻¹ and about 300 larvae were used in each round of selection. The larvae were exposed to the toxin for seven days and after exposing the larvae to toxin the survived larvae were reared on freshly prepared diet without toxin until they pupated. Mean survival of larvae after exposure to Cry1Ac concentrations over four generations was 35% for Cry1Ac-SEL.

Effect of Inhibitors on Pesticides Activities

The toxicities of Cry1Ac, profenofos, indoxacarb and deltamethrin to Cry1Ac-SEL, UNSEL, and Lab-PK populations were determined in the presence of two inhibitors, piperonyl butoxide (PBO; Sigma Ltd, UK), an inhibitor of cytochrome P450 monoxygenases (microsomal oxidases) and of esterases, and S,S,S-tri-n-butyl phosphorothioate (DEF; Sigma Ltd, UK), an esterase-specific inhibitor as described previously [8]. The 10 mg L⁻¹ of the inhibitor was added to various concentrations of the pesticides and larvae were exposed as described above. The mortality was recorded after seven days exposure to the pesticides. The synergism ratio (SR) was calculated by dividing the LC₅₀ of the population treated with pesticide alone by the LC₅₀ of the strain treated with pesticide plus the inhibitor.

Genetics of Resistance to Cry1Ac

The response of F₁ progeny to Cry1Ac was evaluated in mass reciprocal crosses between Cry1Ac-SEL and laboratory susceptible (Lab-PK) populations. To produce F₁, mass crosses using 50 adults of each sex provided enough offspring for multiple-concentration testing and calculation of the 50% lethal concentration (LC₅₀). The degree of dominance for LC₅₀ (DLC) was calculated as described by Sayer et al [23] and Bourguet et al. [24] Backcrossed offspring were obtained from F₁ × Lab-PK. This backcross was preferred to F₁ × Cry1Ac-SEL because the resistance was incompletely dominant and differed more from Lab-PK than from Cry1Ac-SEL.

Data Analysis

Mortality data were corrected by Abbott’s formula [25] where necessary and analysed by probit analysis [26] using the software POLO-PC [22]. The estimates of LC₅₀ values and their 95% fiducial limits (FL) were obtained by probit analysis using Polo. Because of the inherent variability of bioassays, pair-wise comparisons of LC₅₀ values were made at the 1% significance level where individual 95% FL for two treatments do not overlap [27]. Resistance ratios were determined by dividing the LC₅₀ values of field populations by the LC₅₀ of Lab-PK. Cross-resistance pattern among insecticides was studied with pair-wise correlation co-efficient of LC₅₀ values of the field populations for each insecticide.

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Author Contributions

Conceived and designed the experiments: AHKA AHS. Performed the experiments: AHKA. Analyzed the data: AHS. Contributed reagents/materials/analysis tools: MN MA. Wrote the paper: AHS.

References

1. King ABS (1994) Heliothia/Helicoverpa armigera (Lepidoptera: Noctuidae), In: G. Matthews and J. Tunstall (eds), Insect Pests of Cotton. CAB International, Wallingford, UK. 445–446.
2. McCaffery AR (1998) Resistance to insecticides in heliothine Lepidoptera: a global view; 1998 Apr 08–09; London, England. 1735–1750.
3. Ferre J, Van Rie J (2002) Biochemistry and genetics of insect resistance to Bacillus thuringiensis. Ann Rev Entomol 47: 501–533.
4. Sayer AH, Wright DJ (2006) Genetics and evidence for an esterase-associated mechanism of resistance to indoxacarb in a field population of diamondback moth. Pest Manag Sci 62: 1045–1051.
5. Ahmad M, Arif MI, Ahmad Z (1995) Monitoring insecticide resistance Helicoverpa armigera (Lepidoptera: Noctuidae) in Pakistan. J Econ Entomol 88: 771–776.
6. furchen-Constant RH, Daborn PJ, Goff GL (2004) The genetics and genomics of insecticide resistance. Trend Gen 20: 163–170.
7. Ahmad M, Arif MI, Ahmad Z, Denholm I (2002) Cotton whitefly [Bemisia tabaci] resistance to organophosphate and pyrethroid insecticides in Pakistan. Pest Manag Sci 58: 203–208.
8. Sayer AH, Moores G, Crickmore N, Wright DJ (2008) Cross-resistance between a Bacillus thuringiensis Cry toxin and non-Bt insecticides in the diamondback moth. Pest Manag Sci 64: 813–819.
9. Dhurma S, Gujar GT (2011) Field-evolved resistance to Bt toxin Cry1Ac in the pink bollworm, Pectinophora gossypiella (Saunders) (Lepidoptera: Gelechiidae), from India. Pest Manag Sci 67: 891–903.
10. Zhang HN, Yin W, Zhao J, Jin L, Yang YH, et al. (2011) Early warning of cotton bollworm resistance associated with intensive planting of Bt cotton in China. PLoS ONE 6: e22874.
11. Sokal RR, Rohlf FJ (1981) Biometry, 3rd edition. WH Freeman, San Francisco, CA, USA.
12. Gunning RV, Dang HT, Kemp FC, Nicholson IC, Moores GD (2005) New resistance mechanism in Helicoverpa armigera threatens transgenic crops expressing Bacillus thuringiensis Cry1Ac toxin. Appl Environ Microbiol 71: 2558–2563.
13. Wan P, Huang Y, Wu H, Huang M, Cong S, et al. (2012) Increased frequency of pink bollworm resistance to Bt toxin Cry1Ac in China. PLoS ONE 7: e29975.
14. Ahmad M, Sayer AH, Crickmore N, Saleem MA (2007) Genetics and mechanism of resistance to deltamethrin in a field population of Spodoptera litura (Lepidoptera : Noctuidae). Pest Manag Sci 63: 1002–1010.
15. Carriere Y, Dennehy TJ, Pedersen B, Haller S, Ellers-Kirk C, et al. (2001) Large-scale management of insect resistance to transgenic cotton in Arizona: Can transgenic insecticidal crops be sustained? J Econ Entomol 94: 315–325.

16. Shad SA, Sayyed AH, Saleem MA (2010) Cross-resistance, mode of inheritance and stability of resistance to emamectin in Spodoptera litura (Lepidoptera: Noctuidae) Pest ManagSci 66: 839–846.

17. Horowitz AR, Gorman K, Ross G, Denholm I (2003) Inheritance of pyriproxyfen resistance in the whitefly, Bemisia tabaci (Q biotype). Arch Insect Biochem Physiol 54: 177–186.

18. Tabashnik BE (1991) Determining the mode of inheritance of pesticide resistance with backcross experiments. J Econ Entomol 84: 703–712.

19. Lande R (1981) The minimum number of genes contributing to quantitative variation between and within populations Genetics 99: 541–553.

20. Sayyed AH, Ahmad M, Crickmore N (2000) Fitness costs limit the development of resistance to indoxacarb and deltamethrin in Helicoverpa armigera (Lepidoptera: Noctuidae). J Econ Entomol 103: 1927–1933.

21. Ahmad M, Arif MI, Ahmad Z (1995) Monitoring insecticide resistance of Helicoverpa armigera (Lepidoptera, Noctuidae) in Pakistan. J Econ Entomol 88: 771–776.

22. LeOra S (2003) Polo Plus a users guide to probit or logit analysis. LeOra Software, Berkeley, CA.

23. Sayyed AH, Haward R, Herrero S, Ferre J, Wright DJ (2000) Genetic and biochemical approach for characterization of resistance to Bacillus thuringiensis toxin Cry1Ac in a field population of the diamondback moth, Plutella xylostella. Appl Environ Microbiol 66: 1509–1516.

24. Bourguet D, Genissel A, Raymond M (2000) Insecticide resistance and dominance levels. J Econ Entomol 93: 1588–1595.

25. Abbott SW (1925) A method of computing the effectiveness of an insecticide. J Econ Entomol 18: 265–267.

26. Finney D (1971) Probit analysis, 3rd ed. Cambridge University Press, Cambridge, United Kingdom.

27. Litchfield JT, Wilcoxon F (1949) A simplified method of evaluating dose-effect experiments. J Pharmacol Exp Ther 99: 99–103.