RESEARCH ARTICLE

Methoxyfenozide tolerance in *Chrysoperla carnea*: Inheritance, dominance and preliminary detoxification mechanisms

Muhammad Mudassir Mansoor1,2*, Sarfraz Ali Shad2*

1 Fatima Sugar Research & Development Centre, Fatima Sugar Mills Ltd, Muzaffargarh, Punjab, Pakistan,
2 Department of Entomology, Faculty of Agricultural Sciences and Technology, Bahauddin Zakariya University, Multan, Punjab, Pakistan

* honeybeepak@gmail.com (MMM); sarfrazshad@bzu.edu.pk (SAS)

Abstract

Lacewings exist in insecticide-dominant cropping systems. They are prime biological control agents due to outstanding ability of insecticide resistance development. This study examines occurrence of methoxyfenozide resistance and its subsequent effects on cross-resistance to other insecticides, inheritance and mechanism of resistance in *C. carnea*. Methoxy-SEL strain of *C. carnea* selected for 15 generations developed 3531.67-fold resistance to methoxyfenozide. Overlapping fiducial limits of LC50s of F1 and F1' (reciprocal crosses) suggested an autosomal and incompletely dominant mode of inheritance. Resistance to methoxyfenozide was polygenic and its realized heritability value was high (h2 = 0.62). Both PBO and DEF significantly changed LC50s indicating cytochrome P450-dependent monooxygenases and esterases detoxifying the resistance in Methoxy-SEL strain. Resistance to all tested insecticide was unstable but decrease rate was very negligible. These results have implications for preservation of biological control and effective use in insecticide-dominant cropping systems.

Introduction

A key element of pest management strategies in agro-ecosystems is to develop basic understanding of the impact on non-target natural enemies [1]. Insecticides prevail as a dominant tool for pest management regardless of having potential consequences on non-target arthropods [2]. Biological control is one of the constituents of different Integrated Pest Management (IPM) programs in multiple cropping situations [3]. Although, developing compatibility between insecticides and natural enemies has been portrayed as debatable and complicated issue [4], but both are still being used in agriculture.

*Chrysoperla carnea* (Stephens) (Neuroptera:Chrsopidae) is an effective voracious predator. It is widely used against several insect pests including whiteflies, aphids, jassids [5], thrips, mites, mealybugs and lepidoptera eggs. Due to its effectiveness, this predator is commercially available worldwide making it a cosmopolitan species. It has developed remarkable resistance
to several groups of insecticides including conventional, pyrethroids [5, 6], novel mode of action [7–9] and insect growth regulators (IGRs) [10–12].

To make an IPM program successful, it is important to use selective insecticides in tandem with natural enemies and other management tactics. This may deliver comprehensive management of target pests better than either approach singly [13]. Selectivity of insecticides with respect to biological control agents can be evaluated in the field and laboratory, but most studies have been done in controlled laboratory conditions. This is due to the probability of uncontrollable biotic and abiotic pressures in field studies [14].

Characterization of different aspects of insecticide resistance is essential to recognize the phenomenon of resistance development. For this purpose, studying insecticide resistance, genetics, mechanism, and other relevant features is important [15, 16]. It includes knowledgebase for designing effective IPM strategy including natural enemies while eliminating target pests. Studies reporting insecticide resistance, cross-resistance, inheritance, realized heritability mechanism and stability in C. carnea have been conducted [5, 7–10, 17]. However, there is no report of methoxyfenozide resistance and its characterization for this natural enemy.

Methoxyfenozide is an insect growth regulator (IGR) with ecdysone receptor agonist action [18]. This IGR is quite selective in action, posses very low acute toxicity on mammals, harmful to target pest species [19], and compatible with natural enemies [20]. Recently, several resistance aspects of this IGR have been documented from several countries such as selection [21], resistance risk assessment and baseline monitoring [22], toxicity and kinetics in Spodoptera exigua (Hübner) [23], mechanism and stability in Spodoptera litura (Fabricius) [24], cross-resistance and fitness costs [25] and genetics in Musca domestica L. [26]. Information about the effects of methoxyfenozide on natural enemies is very limited. Hewa-Kapuge et al., 2003 [27] studied the effects of this IGR on an egg parasitoid Trichogramma barassicae Bez. (Hymenoptera: Trichogrammatidae) and concluded that it is potentially suitable to control pests in the presence of this natural enemy. As C. carnea is an admired and dominant predator existing in multiple cropping systems, it is important to study methoxyfenozide resistance and its aspects on this natural enemy.

Materials and methods

C. carnea collection and rearing

About 300 adults of C. carnea were collected with a ventilated plastic vial (15 x 45 mm) from cotton, sugarcane and vegetable fields in District Muzaffargarh ((30.0703° N, 71.1933° E) in early spring. Verbal consent of local growers was obtained before collections from different fields so no legal permits were necessary in this regard. These adults were shifted to the Biological Control Laboratory, Fatima Sugar Research & Development Centre, District Muzaffargarh, in plastic cages (23 x 38 x 38 cm) and reared on an artificial diet mixture of water, honey and yeast (4:2:1 ratio) [6, 7]. Black glossy papers were instantly lined with ceiling of rearing cares for egg laying and replaced within 24 hours. At least 5–10 eggs removed from this black glossy paper by a sharp knife were placed inside transparent gelatin capsule (500 mg). A culture of Angoumois grain moth, Sitotroga cerealella Oliver was taken from IPM Station PARC, Faculty of Agricultural Sciences and Technology Bahauddin Zakariya University, Multan in 2007 and reared without any insecticide exposure [17]. Eggs of S. cerealella were placed in the transparent capsule to feed the C. carnea larvae. Temperature, relative humidity and photoperiod were maintained as previously reported in recent publications [8, 9].
Insecticides
Formulated insecticides used for bioassays include methoxyfenozide (Runner® 240SC, Arysta Life Sciences, Pakistan) 100–800 μg ml⁻¹, acetamiprid (Mospilon® 20 WP, Dow Agro-Sciences) 250–2000 μg ml⁻¹, profenofos (Curacron® 500 EC, Syngenta, Pakistan) 25–200 μg ml⁻¹ and lambda-cyhalothrin, (Karate® 2.5 EC, Syngenta, Pakistan) 50–400 μg ml⁻¹. For synergism tests, Piperonyl butoxide (PBO; Sigma Ltd, UK), inhibitors of cytochrome P450 monooxygenases (microsomal oxidases) & esterases, (DEF; Sigma Ltd, UK), an another esterase specific inhibitor, S,S,S-tributylphosphorothriothioate were used.

Concentration-response bioassays
Topical bioassays were performed to evaluate toxicity of tested insecticides on 1st instar larvae of *C. carnea* (Field). A droplet of 0.5 μl was applied from 1-ml glass syringe of Micro-applicator (Burkard Manufacturing Co. Ltd., Hertfordshire, England) directly on thorax of larvae [6]. Four serial concentrations of each tested insecticide were made and each concentration was replicated four times. There were 20 larvae in each replication making a total of 320 larvae treated per insecticide. However, only 30 larvae of *C. carnea* were treated with distilled water as control [28].

A susceptible strain of *C. carnea* (Susceptible) which was obtained in 2007 and reared at Biological Control Laboratory, Fatima Sugar Research & Development Centre, District Muzaffargarh was also used in bioassays. Larvae exposed to distilled water or the treatments were kept in transparent capsules with eggs of *S. cerealella* till pupation [29].

Selection with methoxyfenozide
Larvae of *C. carnea* (Field) were grouped into two sub-groups. At least 100 adults were included in each group. One sub-group (UNSEL) was reared without any insecticide treatment while second sub-group (Methoxy-SEL) was treated with varying levels (800 to 12800 μg ml⁻¹) of methoxyfenozide from G1 to G15 [9, 10]. Larvae were treated as reported in concentration-response bioassay section.

Genetic crosses
At least 30 male adults from Methoxy-SEL and 30 female adults from Susceptible strain were crossed to obtain F₁. Similarly, 30 female adults from Methoxy-SEL and 30 male adults from Susceptible strain were mated to get F₁. Another strain, F₂ was developed by crossing 30 males and females of F₁ strain. A Backcross (BC₁) was also developed using 30 females from F₁ strain and 30 males from Susceptible strain.

Degree of dominance (*D_{LC}*). *D_{LC}* of methoxyfenozide resistance was estimated with the following formula. Resistance was assumed completely dominant (if *D_{LC}* = 1) and completely recessive (if *D_{LC}* = 0) [30].

\[
D_{LC} = \frac{\log LC_{RS} - \log LC_{S}}{\log LC_{R} - \log LC_{S}}.
\]

Where Log LC₉₅ = LC₀ of Methoxy-SEL, LC₉₅ = LC₀ of F₁ and LC₅₀ = LC₀ of Susceptible strains [31].

Effective dominance (*D_{ML}*). of resistance to methoxyfenozide was estimated as

\[
D_{ML} = \frac{MT_{RS} - MT_{SS}}{MT_{RR} - MT_{SS}}.
\]

Where MT₉₅ (F₁), MT₉₅ (Methoxy-SEL) and MT₅₀ (Susceptible) shows percent mortality on any single tested dose of insecticide. Resistance to methoxyfenozide was completely dominant (if *D_{ML}* = 1) and completely recessive (if *D_{ML}* = 0) [32].
**Calculation of gene frequency.** Monogenic resistance hypothesis was tested using Goodness of fit (Chi-square) test [33]. Equation for testing null hypothesis of monogenic resistance is given below.

$$\chi^2 = (F - pn)^2 / pqn.$$  

Where F is mortality in BC$_1$, p is expected mortality, n is total larvae exposed to any dose and q is 1-p. Null hypothesis of monogenic resistance get rejected if 50% expected and observed mortalities show significant difference ($p > 0.05$).

Gene frequency responsible for methoxyfenozide resistance can be calculated as

$$\eta_E = (X_{RR} - X_{SS})^2 / (8\sigma^2 S).$$

Where $X_{RR} = \text{Log of LC}_{50}$ of Methoxy-SEL and $X_{SS} = \text{Log of LC}_{50}$ of Susceptible strain [34]. The $\sigma^2 S$ was estimated as follow:

$$\sigma^2 S = \sigma^2 B_1 - [\sigma^2 F_1 + 0.5 \sigma^2 X_{SS} + \sigma^2 X_{RR}]$$

Where $\sigma^2 B_1$, $\sigma^2 F_1$, $\sigma^2 X_{SS}$, and $\sigma^2 X_{RR}$ are variances of BC$_1$, F$_1$, Susceptible and Methoxy-SEL strains.

**Realized heritability ($h^2$)**

Realized heritability of methoxyfenozide resistance was calculated as follows.

$$h^2 = \frac{\text{Response to selection (R)}}{\text{Selection differential (S)}}.$$  

[35].

Response to selection was calculated as

$$R = \frac{\text{Log final LC}_{50} \text{ of Methoxy - SEL} - \text{Log initial LC}_{50} \text{ of Field Pop}}{n},$$

Here, n shows total number of generations exposed to methoxyfenozide.

Selection differential (S) was calculated with given formula:

$$S = \text{Intensity of selection (i) \times Phenotypic standard deviation (}\sigma p).$$

Intensity of selection was as,

$$i = 1.583 - 0.0193336p + 0.0000428p^2 + 3.65194 / p,$$

Where p is average survival of Methoxy-SEL strain.

Phenotypic standard deviation ($\sigma p$) was determined as

$$\sigma p = \left[1/2(\text{final slope} + \text{initial slope})\right]^{-1}.$$  

**Biochemical mechanism**

PBO and DEF (5 mg ml$^{-1}$) were diluted in acetone and mixed with insecticide concentrations. Experiments to determine the resistance mechanism were conducted as previously reported by Mansoor et al., 2017 [7].

**Stability and decrease rate of resistance**

The sub-group (UNSEL) left unexposed to any insecticide was used for this study. Resistance stability and its decrease rate (DR) to tested insecticides was measured using given
formula [36].

\[ DR = \frac{\log (\text{final } LC_{50}) - \log (\text{initial } LC_{50})}{n} \]

Total number of \textit{C. carnea} generations required for a 10-fold decrease in resistance to methoxyfenozide and other tested insecticides was also calculated as follows.

\[ GR = \frac{1}{R} \]

where \( R \) is response to selection.

**Data collection and statistical analysis**

Treated larvae were tapped softly by a needle-like brush after 72 hours of bioassays. These were considered dead if there was no movement. Mortality data was corrected [37] if necessary. Mortality data was further analyzed with Probit Analysis [38]. These analyses produced Median Lethal Concentration (\( LC_{50} \)), 95% Fiducial Limits (FLs), standard errors and slope values. \( LC_{50} \)s were regarded significantly different if FLs had no overlapping [39, 40].

**Results**

**Toxicity of various insecticides to Susceptible, field, UNSEL and Methoxy-SEL strains**

Methoxyfenozide was significantly less toxic to Susceptible strain followed by profenofos and lambda-cyhalothrin (95% FLs didn’t overlap). Toxicity of profenofos and lambda-cyhalothrin was significantly similar (95% FLs overlapping). Acetamiprid was the most toxic insecticide than all other tested insecticides (95% FLs didn’t overlap) (Fig 1).

For field strain, methoxyfenozide was significantly less toxic than profenofos (95% FLs didn’t overlap). However, it was relatively similar to that of acetamiprid and lambda-cyhalothrin (95% FLs overlapping). Profenofos and lambda-cyhalothrin were the most toxic

![Graph showing toxicity of various insecticides](https://doi.org/10.1371/journal.pone.0265304.g001)
insecticides. Toxicity to acetamiprid was significantly lower than profenofos and lambda-cyhalothrin (95% FLs didn’t overlap) (Fig 2).

The toxicity of methoxyfenozide was significantly low compared to other tested insecticides (95% FLs didn’t overlap). Acetamiprid was less toxic than lambda-cyhalothrin (95% FLs didn’t overlap). However, toxicity of profenofos and lambda-cyhalothrin was significantly high and similar (95% FLs overlapping) (Table 1).

Methoxy-SEL was 3531.67-fold and 35.91-fold more resistant than susceptible and field strains, respectively (Table 1). Average response of selection to methoxyfenozide was 82% after 72 hours of exposure (Table 3).

**Cross-resistance pattern**

Cross-resistance testing revealed that selection to methoxyfenozide didn’t increase resistance to any tested insecticides compared to field strain (Table 1). However, there was a slight change of 2.09-fold resistance to profenofos (95% FLs overlapping).

**Maternal effects and sex linkage**

Resistance in Methoxy-SEL was 3531.67-fold higher than susceptible strain. Resistance to methoxyfenozide dropped from 3531.67-fold to 1115.84-fold and 783.11-fold for F1 and F1' reciprocal crosses, respectively, compared to Susceptible (Fig 3). LC50s of these reciprocal crosses were not significantly different suggesting an autosomal and no sex linkage was involved in resistance development.

**Degree of dominance.** The D_{LC} values for F1, F1' and F2 strains were 0.71, 0.63 and 0.69, respectively (Fig 3). These results suggest an incompletely dominant inheritance of methoxyfenozide resistance in C. carnea. The results for D_{ML} show that level of methoxyfenozide resistance dominance decreased when methoxyfenozide dose was increased from 1600 to 12800 μg ml⁻¹. Resistance was likely completely dominant at lowest concentration (D_{ML} = 0.94) and incompletely dominant at highest concentration (D_{ML} = 0.52) tested (Fig 4).

![Fig 2. Toxicity of various insecticides to field strain (G1) of Chrysoperla carnea and resistance ratios. Error Bars show fiducial limits (FLs) of LC50s.](https://doi.org/10.1371/journal.pone.0265304.g002)
Number of genes controlling resistance. Monogenic test revealed that observed mortalities were significantly different ($P < 0.05$) compared to expected results of mortality (Fig 5). These significant differences advocated that resistance to methoxyfenozide is polygenic.

Realized heritability

The LC$_{50}$ value increased from 731.51 to 26275.66 μg ml$^{-1}$ in Methoxy-SEL strain after selecting 15 generations with methoxyfenozide. The realized heritability value of methoxyfenozide resistance was 0.62 (Table 2). The $h^2$ value of methoxyfenozide resistance was high while expected number of generations required to gain a 10-fold increase in LC$_{50}$ was only 10 (reciprocal of R: Table 2).

Synergism tests

Both PBO and DEF didn’t synergize the toxicity of methoxyfenozide against susceptible strain (95% FLs overlapping). However, toxicity of methoxyfenozide was significantly synergized by both PBO and DEF against Methoxy-SEL strain (95% FLs didn’t overlap). PBO and DEF decreased the LC$_{50}$ values from 26275.66 to 3713.24 μg ml$^{-1}$ and 26275.66 to 5901.98 μg ml$^{-1}$, respectively, in Methoxy-SEL strain (Fig 6).

Stability and decrease rate of resistance

Resistance to all tested insecticides dropped significantly from G1 to G15. This suggested that resistance to methoxyfenozide and other sampled insecticides was unstable (95% FLs not overlapping). The decline rates for methoxyfenozide, acetamiprid, profenofos and lambda-cyhalothrin were -0.06, -0.08, -0.07, and -0.05, respectively. *C. carnea* would require at least 16, 13, 14 and 18 generations for a 10-fold decrease in resistance to methoxyfenozide, acetamiprid, profenofos and lambda-cyhalothrin, respectively (Table 3).

---

### Table 1. Toxicity of various insecticides to Methox-SEL population of *Chrysoperla carnea*.

| Strain     | Insecticide    | LC$_{50}$ (95% FLs) μg ml$^{-1}$ | Fit of probit line | N$^a$ | RR$^b$ | RR$^c$ |
|------------|----------------|---------------------------------|--------------------|-------|-------|-------|
|            |                |                                 | Slope (±SE)        | $\chi^2$ | Df | $P$ |                                 |                                 |                                 |
| Methox-SEL (G15) | Methoxyfenozide | 26275.66(16686.46–67071.95) | 1.56(0.30)         | 0.30   | 3    | 0.96  | 350 | 3531.67 | 35.91 |
|             | Acetamiprid    | 1289.21(1025.58–1772.77)        | 1.57(0.23)         | 0.08   | 3    | 0.99  | 350 | 2432.47 | 1.17  |
|             | Profenofos     | 413.47(251.26–1210.58)         | 1.34(0.27)         | 0.02   | 3    | 0.99  | 350 | 135.56  | 2.09   |
|             | Lambda-cyhalothrin | 533.82(382.90–960.01) | 1.63(0.27)         | 0.58   | 3    | 0.90  | 350 | 196.98  | 1.40   |

$^a$N Total larvae in a bioassay including control.  
$^b$RR resistance ratio, LC$_{50}$ of field population and Methox-SEL strains/LC$_{50}$ of Susceptible strain.  
$^c$RR resistance ratio, LC$_{50}$ of Methox-SEL strain/LC$_{50}$ of field population.

https://doi.org/10.1371/journal.pone.0265304.t001

### Table 2. Estimation of the realized heritability of resistance in Methox-SEL strain of *Chrysoperla carnea*.

| N | Insecticide     | Initial log LC$_{50}$ (μg ml$^{-1}$) | Final log LC$_{50}$ (μg ml$^{-1}$) | Response to selection (R) | Initial slope | Final slope | $\sigma_p$ | Selection differential (S) | $h^2$ |
|---|----------------|-------------------------------------|-------------------------------------|---------------------------|---------------|-------------|-----------|---------------------------|-------|
| 15 | Methoxyfenozide | 2.86                                | 4.42                                | 0.10                      | 0.31          | 2.13        | 1.56      | 0.54                      | 0.17  | 0.62                      |

$N$ is the number of generations exposed with Methoxyfenozide.  
$P$ is the average survival% of green lacewing larvae throughout the selection.  
i is the intensity of selection.  
$\sigma_p$ is the phenotypic variation.

https://doi.org/10.1371/journal.pone.0265304.t002
**Discussion**

Methoxyfenozide is a bio-rational insecticide and it acts as an ecdysone agonist. It is used to control numerous insect pests, especially pests from lepidoptera and diptera. After 15 generations of regular selection with methoxyfenozide, Methoxy-SEL strain of *C. carnea* developed 3531.67-fold resistance compared with susceptible strain. This showed immense potential of...
Fig 5. Direct test of monogenic inheritance of resistance to methoxyfenozide by comparing expected and observed mortality of the backcross (F1♀ x Susceptible ♂) of Chrysoperla carnea.

https://doi.org/10.1371/journal.pone.0265304.g005

Fig 6. Toxicity of methoxyfenozide alone and with PBO or DEF to the susceptible and Methox-SEL Chrysoperla carnea strains.

https://doi.org/10.1371/journal.pone.0265304.g006
this natural enemy to develop resistance to this insecticide under selection pressure. There are numerous insecticide resistance reports for *C. carnea* [5, 7–10, 29, 41] indicating potential of this natural enemy in diverse field conditions.

Current results showed that Methoxy-SEL developed no cross-resistance to acetamiprid (1.17-fold), lambda-cyhalothrin (1.40-fold) and profenofos (2.09-fold) compared to field strain (95% FLs overlapped). Previously, no cross-resistance to other insecticides in methoxyfenozide selected populations has been reported. For example, a population of *S. litura* (Fabricius) selected with methoxyfenozide showed no cross-resistance to profenofos, spinosad, fipronil, lufenuron, methoioxynil or thiocarb [24]. A methoxyfenozide-selected strain of *M. domestica* showed no cross-resistance to bifenthrin and spinosad but very low cross-resistance to chlorpyrifos and fipronil [25]. Obliquebanded leafroller, *Choristoneura rosacea* (Harris) collected from various locations showed little cross-resistance between benzoylhydrazine, tebufenozide, methoxyfenozide and azinphosmethyl [42]. Lack of cross-resistance to different insecticides in Methoxy-SEL strain could be due to different resistance mechanisms [43, 44]. These results are in agreement with our previous findings of no, or little, cross-resistance exhibited by different insecticide-selected strains of *C. carnea* [7, 9, 10, 17].

Genetic crosses between insecticide-resistance and susceptible populations of natural enemies are a valuable tool to understand mode of inheritance, effective dominance, degree of dominance and number of genes supporting resistance development. Foreseeing how a natural enemy resistant to methoxyfenozide such as *C. carnea* may pass resistance to succeeding or susceptible populations and its possible impacts on insecticide resistance stability is only possible by studying genetics in laboratory. Reciprocal crosses F₁, F₁', and F₂ strains of *C. carnea* showed degree of dominance (D₁,c) values of 0.71, 0.63, and 0.69, respectively. These results indicated that LC₅₀s were not significantly different in reciprocal crosses suggesting an autosomal mode of inheritance. These outcomes are similar to previous studies on inheritance of resistance to acetamiprid, pyriproxyfen and cyromazine [9, 11, 12]. These results are in agreement to a recent study concluding autosomal, and no sex linkage in methoxyfenozide resistance in *M. domestica* [26].

Studying effective dominance showed that methoxyfenozide resistance is incompletely dominant in Methoxy-SEL strain of *C. carnea*. Interestingly, resistance was likely completely dominant (D₅₀ = 0.94) at the lowest dose (1600 μg ml⁻¹) tested. Changing concentration of any insecticide may affect dominance level [45]. Heritability of resistance increases due to increase in resistance dominance level, thus amplifying development of resistance. Dominance level for any particular factor is generally considered fixed but it may be affected by the genetic history and environmental circumstances [32]. Susceptible alleles may remain for a prolonged period even if resistance alleles show completely or incompletely dominant inheritance [46] and this situation supports interaction between dominant and recessive genes [47]. Current
findings indicated that methoxyfenozide resistance increased rapidly due to selection pressure in *C. carnea* while selection levels, diverse environments, and population structures may cause dissimilar responses between field and laboratory populations. However, it also suggested that methoxyfenozide would not kill the heterozygotes easily as resistance to this insecticide is associated with dominant genes.

Monogenic resistance depended on chi-square (Goodness of fit test) and estimation of total number of generations showed that methoxyfenozide resistance is controlled by more than one gene. This suggests that *C. carnea* has polygenic resistance for methoxyfenozide. Polygenic resistance is common among field collected populations selected under laboratory conditions due to natural selection variations but monogenic resistance takes place in natural populations only [48]. However, Sayyed and Wright [49] reported that multiple genes controlling resistance could be evenly spread between natural and laboratory-selected populations. Our results concurred with previous studies reporting polygenic resistance to deltamethrin [17], buprofezin [10], acetamiprid [9], pyriproxyfen [11] and cyromazine [12] in *C. carnea*.

Realized heritability can be used effectively in assessing potential of increase in resistance and fate of an insecticide in laboratory-selected populations [35]. A high realized heritability value (0.62) suggests high genetic variation and quick increase in methoxyfenozide resistance in Methoxy-SEL strain of *C. carnea* after only 15 generations of selection. Furthermore, this strain would require only 10 generations (Reciprocal of R, Table 2) for a 10-fold increase in resistance. Even though laboratory conditions are not a true match of field conditions, the estimated $h^2$ and predictable rate of methoxyfenozide resistance through laboratory selection have implications for biological control management. Therefore, this insecticide can be employed wisely in field.

Significant synergism by PBO and DEF on methoxyfenozide in Methoxy-SEL strain showed that cytochrome P450 monooxygenase and esterase might have a significant effect on detoxification of this insecticide. These results are in accordance with several reports of synergistic effects of PBO and DEF in methoxyfenozide-selected *S. exigua* [23], *S. litura* [24] and *M. domestica* [26]. Previously, Sayyed et al., [17], Mansoor et al., [7], Mansoor and Shad [10] and Mansoor and Shad [12] showed significant effect of PBO and DEF in synergizing the impact of deltamethrin, nitenpyram, buprofezin and cyromazine resistance in *C. carnea*, respectively.

Stability of insecticide resistance is a remarkable tool for result-oriented field utilization of natural enemies with idea of conservation. Natural enemies, such as *C. carnea* possessing insecticide resistance, which is stable, can be of benefit in IPM systems. This feature also ensures survival of non-targets especially when insecticide selection pressure is removed [7, 11]. Bioassays on UNSSEL population of *C. carnea* concluded that resistance to methoxyfenozide and other insecticides was not stable. However, the decrease rate was minimal suggesting that this population will take at least 16 generations to show a 10-fold decrease in resistance. In contrast, the higher realized heritability ($h^2$) and predictable rate of resistance development (reciprocal of R, Table 2) suggested that the same would need only 10 generations to acquire 10-fold resistance. This fact, accompanied with negligible decrease rates, also indicates the possibility of maintaining resistance for longer in field conditions if *C. carnea* receives periodic selection pressure of selected insecticides.

In conclusion, methoxyfenozide resistance in *C. carnea* is autosomal, incompletely dominant, controlled by multiple genes and settled by cytochrome P450-dependent monooxygenases and esterases. Although, hybrid individuals demonstrated incompletely dominant resistance, they also confer potential boost in resistance, suggesting that *C. carnea* has natural potential to acquire resistance to methoxyfenozide. Instable resistance in Methoxy-SEL and no cross-resistance to acetamiprid, profenofos and lambda-cyhalothrin with high to very high resistance in field populations suggested that these insecticides could be used in integration with *C. carnea* keeping in view the current resistance status of different pests.
Acknowledgments
Authors are grateful to Dr. Robert J. Whitworth, Associate Professor, Department of Entomology, Kansas State University, USA for improvement in English language and sense of this article.

Author Contributions
Conceptualization: Muhammad Mudassir Mansoor, Sarfraz Ali Shad.
Data curation: Muhammad Mudassir Mansoor.
Formal analysis: Muhammad Mudassir Mansoor, Sarfraz Ali Shad.
Investigation: Muhammad Mudassir Mansoor.
Methodology: Muhammad Mudassir Mansoor.
Supervision: Sarfraz Ali Shad.
Writing – original draft: Muhammad Mudassir Mansoor.
Writing – review & editing: Muhammad Mudassir Mansoor.

References
1. Desneux N, Decourten A, Delpeuch J-M. The sublethal effects of pesticides on beneficial arthropods. Annu Rev Entomol. 2007; 52:81–106. https://doi.org/10.1146/annurev.ento.52.110405.091440 PMID: 16842032
2. Roubos CR, Rodriguez-Saona C, Isacs R. Mitigating the effects of insecticides on arthropod biological control at field and landscape scales. Biol Control. 2014; 75:28–38.
3. Bale J, Van Lenteren J, Bigler F. Biological control and sustainable food production. P Roy Soc B-Biol Sci. 2008; 363(1492):761–76. https://doi.org/10.1098/rstb.2007.2182 PMID: 17827110
4. Devine GJ, Furlong MJ. Insecticide use: Contexts and ecological consequences. Agri Hum Values. 2007; 24(3):281–306.
5. Pathan AK, Sayyed AH, Aslam M, Liu T-X, Razzaq M, Gillani WA. Resistance to pyrethroids and organophosphates increased fitness and predation potential of Chrysoperla carnea (Neuroptera: Chrysopidae). J Econ Entomol. 2010; 103(3):823–34. https://doi.org/10.1603/ec09260 PMID: 20568629
6. Pathan AK, Sayyed AH, Aslam M, Razaq M, Jilani G, Saleem MA. Evidence of field-evolved resistance to organophosphates and pyrethroids in Chrysoperla carnea (Neuroptera: Chrysopidae). J Econ Entomol. 2008; 101(5):1676–84. https://doi.org/10.1603/0022-0493(2008)101[1676:eofrto]2.0.co;2 PMID: 18950051
7. Mansoor MM, Raza ABM, Abbas N, Aqeel MA, Afzal M. Resistance of green lacewing, Chrysoperla carnea Stephens to nitenpyram: cross-resistance patterns, mechanism, stability, and realized heritability. Pestic Biochem Physiol. 2017; 135:59–63. https://doi.org/10.1016/j.pestbp.2016.06.004 PMID: 28043332
8. Mansoor MM, Shad SA. Resistance, its stability and reversion rate of resistance to imidacloprid, indoxacarb and chlorfenapyr in a field population of green lacewing Chrysoperla carnea (Stephens)(Neuroptera: Chrysopidae). Arch Phytopathol Plant Protect. 2019; 52(9):1–11. https://doi.org/10.1080/03235408.2019.1660561
9. Mansoor MM, Shad SA. Genetics, cross-resistance and realized heritability of resistance to acetamiprid in generalist predator, Chrysoperla carnea (Steph.) (Neuroptera: Chrysopidae). Egypt J Biol Pest Co. 2020; 30(1):1–8.
10. Mansoor MM, Shad SA. Resistance of green lacewing, Chrysoperla carnea (Stephens), to buprofezin: Cross resistance patterns, preliminary mechanism and realized heritability. Biol Control. 2019; 129:123–7.
11. Mansoor MM, Shad SA. Inheritance of polygenic but stable pyriproxyfen resistance in a bio-control agent Chrysoperla carnea (Neuroptera: Chrysopidae): Cross-resistance and realized heritability. Pest Manage Sci. 2020;doi.org/10.1002/ps.5952.
12. Mansoor MM, Shad SA. Biochemical mechanism, inheritance and cross-resistance to cyromazine in a non-target Chrysoperla carnea: a potential predator of whiteflies and aphids. Chemosphere. 2020; 260:127620. https://doi.org/10.1016/j.chemosphere.2020.127620 PMID: 32758770
13. Gentz MC, Murdoch G, King GF. Tandem use of selective insecticides and natural enemies for effective, reduced-risk pest management. Biol Control. 2010; 52(3):208–15.

14. Franz J, Bogenschütz H, Hassan S, Huang P, Naton E, Suter H, et al. Results of a joint pesticide test programme by the working group: pesticides and beneficial arthropods. Entomophaga. 1980; 25(3):231–6.

15. Afzal M, Ijaz M, Farooq Z, Shad S, Abbas N. Genetics and preliminary mechanism of chlorpyrifos resistance in Phenacoccus solenopsis Tinsley (Homoptera: Pseudococcidae). Pestic Biochem Physiol. 2015; 119:42–7. https://doi.org/10.1016/j.pestbp.2015.02.008 PMID: 25868815

16. Ejaz M, Afzal MBS, Shabbir G, Serrão JE, Shad SA, Muhammad W. Laboratory selection of chlorpyrifos resistance in an Invasive Pest, Phenacoccus solenopsis (Homoptera: Pseudococcidae): cross-resistance, stability and fitness cost. Pestic Biochem Physiol. 2017; 137:8–14. https://doi.org/10.1016/j.pestbp.2016.09.001 PMID: 28364807

17. Sayyed AH, Pathan AK, Faheem U. Cross-resistance, genetics and stability of resistance to deltamethrin in a population of Chrysoperla carnea from Multan, Pakistan. Pestic Biochem Physiol. 2010; 98(3):325–32.

18. IRAC. IRAC mode of action classification (version 8.1). 2020:pp, 1–26.

19. Dhadialla TS, Carlson GR, Le DP. New insecticides with ecdysteroidal and juvenile hormone activity. Annu Rev Entomol. 1998; 43(1):545–69. https://doi.org/10.1146/annurev.ento.43.1.545 PMID: 9444757

20. Schneider M, Smagghge G, Pineda S, Vinuela E. Action of insect growth regulator insecticides and spinosad on life history parameters and absorption in third-instar larvae of the endoparasitoid Hyposoter didymator. Biol Control. 2004; 31:189–98.

21. Mosallanejad H, Soin T, Smagghge G. Selection for resistance to methoxyfenozide and 20-hydroxyecdysone in cells of the beet armyworm, Spodoptera exigua. Arch Insect Biochem Physiol. 2008; 67(1):42–7. https://doi.org/10.1002/arch.20220 PMID: 18044724

22. Moulton JK, Pepper DA, Jansson RK, Dennehy TJ. Pro-active management of beet armyworm (Lepidoptera: Noctuidae) resistance to tebufenozide and methoxyfenozide: baseline monitoring, risk assessment, and isolation of resistance. J Econ Entomol. 2002; 95(2):414–24. https://doi.org/10.1603/0022-0493-95.2.414 PMID: 12020022

23. Smagghge G, Pineda S, Carton B, Estal PD, Budia F, Vinuela E. Toxicity and kinetics of methoxyfenozide in greenhouse-selected Spodoptera exigua (Lepidoptera: Noctuidae). Pest Manage Sci. 2003; 59(11):1203–9.

24. Rehan A, Freed S. Resistance selection, mechanism and stability of Spodoptera littoralis (Lepidoptera: Noctuidae) to methoxyfenozide. Pestic Biochem Physiol. 2014; 110:7–12. https://doi.org/10.1016/j.pestbp.2014.02.001 PMID: 24759045

25. Shah RM, Shad SA, Abbas N. Methoxyfenozide resistance of the housefly, Musca domestica L.(Diptera: Muscidae): cross-resistance patterns, stability and associated fitness costs. Pest Manage Sci. 2017; 73(1):254–61. https://doi.org/10.1002/ps.4296 PMID: 27098995

26. Shah R, Abbas N, Shad S, Binyamin M. Determination of the Genetic and Synergistic Suppression of a Methoxyfenozide-Resistant Strain of the House Fly Musca domestica L.(Diptera: Muscidae). Neotrop Entomol. 2018; 47(5):709–15. https://doi.org/10.1007/s13744-018-0604-9 PMID: 29654414

27. Hewa-Kapuge S, McDougall S, Hoffmann AA. Effects of methoxyfenozide, indoxacarb, and other insecticides on the beneficial egg parasitoid Trichogramma nr. brassicae (Hymenoptera: Trichogrammatidae) under laboratory and field conditions. J Econ Entomol. 2003; 96(4):1083–90. https://doi.org/10.1093/jee/96.4.1083 PMID: 14503578

28. Robertson J, Preisler H. Pesticide Bioassays With Arthropods. CRC Boca Raton, FL. 1992.

29. Abbas N, Mansoor MM, Shad SA, Pathan AK, Waheed A, Ejaz M, et al. Fitness cost and realized heritability of resistance to spinosad in Chrysoperla carnea (Neuroptera: Chrysopidae). Bull Entomol Res. 2014; 104(6):707–15. https://doi.org/10.1017/S0007485314000522 PMID: 25033090

30. Bourguet D, Raymond M. The molecular basis of dominance relationships: the case of some recent adaptive genes. J Evol Biol. 1998; 11(1):103–22.

31. Stone B. A formula for determining degree of dominance in cases of monofactorial inheritance of resistance to chemicals. Bull WHO. 1968; 38(2):325. PMID: 5302309

32. Bourguet D, Genissel A, Raymond M. Insecticide resistance and dominance levels. J Econ Entomol. 2000; 93(6):1588–95. https://doi.org/10.1603/0022-0493-93.6.1588 PMID: 11142285

33. Sokal R, Rohlf F. Biometry, third ed. WH Freeman & Co., San Francisco, USA; 1981.

34. Lande R. The minimum number of genes contributing to quantitative variation between and within populations. Genetics. 1981; 99(3–4):541–53. https://doi.org/10.1093/genetics/99.3-4.541 PMID: 7343418
35. Tabashnik BE. Resistance risk assessment: realized heritability of resistance to Bacillus thuringiensis in diamondback moth (Lepidoptera: Plutellidae), tobacco budworm (Lepidoptera: Noctuidae), and Colorado potato beetle (Coleoptera: Chrysomelidae). J Econ Entomol. 1992; 85(5):1551–9.

36. Attique M, Khalq A, Sayyed A. Could resistance to insecticides in Plutella xylostella (Lep., Plutellidae) be overcome by insecticide mixtures? J Appl Entomol. 2006; 130(2):122–7.

37. Abbott W. A method of computing the effectiveness of an insecticide. J Econ Entomol. 1925; 18(2):265–7.

38. Finney D. Probit Analysis-A statistical Analysis of the Sigmoid Response Curve. Cambridge University Press; 1971.

39. Litchfield JJ, Wilcoxon F. A simplified method of evaluating dose-effect experiments. J Pharmacol Exp Ther. 1949; 96(2):99–113. PMID: 18152921

40. Robertson JL, Jones MM, Olguin E, Alberts B. Bioassays with arthropods: CRC press; 2017.

41. Mansoor MM, Abbas N, Shad SA, Pathan AK, Razaq M. Increased fitness and realized heritability in emamectin benzoate-resistant Chrysoperla carnea (Neuroptera: Chrysopidae). Ecotoxicology. 2013; 22(8):1232–40. https://doi.org/10.1007/s10646-013-1111-8 PMID: 23975538

42. Smirle MJ, Thomas Lowery D, Zurowski CL. Resistance and cross-resistance to four insecticides in populations of obliquebanded leafroller (Lepidoptera: Tortricidae). J Econ Entomol. 2002; 95(4):820–5. https://doi.org/10.1603/0022-0493-95.4.820 PMID: 12216826

43. Abbas NS, Ali S, Razaq MV, Abdul, Aslam M. Resistance of Spodoptera litura (Lepidoptera: Noctuidae) to profenofos: relative fitness and cross resistance. Crop Protect. 2014; 58:49–54.

44. Zewen L, Zhaojun H, Yinchang W, Lingchun Z, Hongwei Z, Chengjun L. Selection for imidaclopid resistance in Nilaparvata lugens: cross-resistance patterns and possible mechanisms. Pest Manage Sci. 2003; 59(12):1355–9. https://doi.org/10.1002/ps.768 PMID: 14667059

45. Georgiou GP. Management of resistance in arthropods. Pest Resistance to Pesticides: Springer; 1983. p. 769–92.

46. Falconer D. Introduction to Quantitative Genetics, 3rd edn Longmans Green. John Wiley & Sons, Harlow, Essex, UK/New York; 1989.

47. Sayyed AH, Haward R, Herrero S, Ferré J, Wright DJ. 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. 2000; 66(4):1509–16. https://doi.org/10.1128/AEM.66.4.1509-1516.2000 PMID: 10742234

48. McKenzie JA, Parker A, Yen J. Polygenic and single gene responses to selection for resistance to diazinon in Lucilia cuprina. Genetics. 1992; 130(3):613–20. https://doi.org/10.1093/genetics/130.3.613 PMID: 1551581

49. Sayyed AH, Wright DJ. Cross-resistance and inheritance of resistance to Bacillus thuringiensis toxin Cry1Ac in diamondback moth (Plutella xylostella L) from lowland Malaysia. Pest Manage Sci. 2001; 57(5):413–21. https://doi.org/10.1002/ps.313 PMID: 11374157