Natural Enemies Delay Insect Resistance to Bt Crops

Xiaoxia Liu1,2, Mao Chen2*, Hilda L. Collins2, David W. Onstad3,4, Richard T. Roush4, Qingwen Zhang1, Elizabeth D. Earle5, Anthony M. Shelton2*

1 Department of Entomology, China Agricultural University, Beijing, China, 2 Department of Entomology, Cornell University, New York State Agricultural Experiment Station, Geneva, New York, United States of America, 3 Department of Entomology, University of Illinois, Urbana, Illinois, United States of America, 4 Melbourne School of Land and Environment, University of Melbourne, Victoria, Australia, 5 Department of Plant Breeding and Genetics, Cornell University, Ithaca, New York, United States of America

Abstract

We investigated whether development of resistance to a Bt crop in the presence of a natural enemy would be slower than without the natural enemy and whether biological control, in conjunction with a Bt crop, could effectively suppress the pest population. Additionally, we investigated whether insecticide-sprayed refuges of non-Bt crops would delay or accelerate resistance to the Bt crop. We used a system of Bt broccoli expressing Cry1Ac, a population of the pest Plutella xylostella with a low frequency of individuals resistant to Cry1Ac and the insecticide spinosad, and a natural enemy, Coleomegilla maculata, to conduct experiments over multiple generations. The results demonstrated that after 6 generations P. xylostella populations were very low in the treatment containing C. maculata and unsprayed non-Bt refuge plants. Furthermore, resistance to Bt plants evolved significantly slower in this treatment. In contrast, Bt plants with no refuge were completely defoliated in treatments without C. maculata after 4–5 generations. In the treatment containing sprayed non-Bt refuge plants and C. maculata, the P. xylostella population was low, although the speed of resistance selection to Cry1Ac was significantly increased. These data demonstrate that natural enemies can delay resistance to Bt plants and have significant implications for integrated pest management (IPM) with Bt crops.

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* E-mail: ams@cornell.edu

¤a Current address: Monsanto Company, Chesterfield, Missouri, United States of America

¤b Current address: Dupont Agricultural Biotechnology, Wilmington, Delaware, United States of America

Introduction

The commercialization of plants expressing insecticidal crystal (Cry) proteins from Bacillus thuringiensis (Bt) for insect management has revolutionized agriculture and become a major tool for integrated pest management (IPM) programs [1–2]. In 2011, Bt crops were grown on nearly 70 million ha in 27 countries in 2012 [3]. Bt crops have provided economic benefits to growers and reduced the use of other insecticides [1,4–6], suppressed pest populations on a regional basis [7–9], conserved natural enemies [10] and promoted biological control services in agricultural landscapes [6]. However, the development of insect resistance is a major threat to the sustainable use of Bt crops [11–12].

Since Bt crops were first commercialized in 1996, there is evidence that three lepidopteran pests have evolved resistance to Bt crops in the open field [13–15] and one case of a coleopteran pest [16]. Resistance to Bt plants is a serious concern, but the relatively few number of cases is in stark contrast to many cases of resistance to conventional insecticides, which has occurred much more rapidly [17]. Commonly proposed reasons for the few confirmed cases of resistance to Bt plants are the high dose of Bt proteins expressed in plants and the use of refuges of non-Bt plants that can serve as a pool of Bt susceptible alleles in the population [18–19].

Another possible reason for the relatively few cases of resistance to Bt plants could be their safety to natural enemies that help suppress pest populations. Numerous studies have investigated the effects of Bt crops and Cry proteins on natural enemies (predators and parasitoids) in the laboratory and field [20–21]. A meta-analysis has confirmed the safety of Bt proteins, especially when compared to traditional insecticides [10]. When negative effects on natural enemies have been observed with Bt proteins, they appear to be due to the poor quality of the host and not the Cry protein [22], but see Desneux et al., 2010 [23]. The safety of several Bt proteins has been verified in tritrophic studies conducted with Bt-resistant or non-susceptible herbivores that avoided the problems of prey-quality in some previous studies [22]. Allowing Bt-resistant hosts to ingest Bt proteins and then feeding the hosts to natural enemies (both predators and parasitoids) has revealed no effects on the natural enemies [24–26]. However, some reports continue to suggest natural enemies may be harmed by Bt proteins [27], but these have been challenged [28].

The conservation of natural enemies by the use of Bt plants could also influence the development of resistance to Bt crops. This question was first studied by Gould et al. [29] in their conceptual and mathematical models on tritrophic interactions of a plant, an herbivore and a natural enemy. Their simplest conclusion was that natural enemies that increase differential fitness between
susceptible and resistant phenotypes on host plants will accelerate resistance; those that decrease the differential will delay resistance. Johnson et al. [30–31] carried out controlled studies of a parasitoid and a pathogenic fungus that attack *Heliothis virescens* on *Bt* tobacco and concluded that the parasitoid would likely delay the development of resistance to transgenic tobacco, while the pathogen would likely promote the development of resistance. Mallampalli et al. [32] discovered that different prey species of a generalist predator had different effects on the development of resistance by *Leptinotarsa decemlineata* to *Bt* potato: the presence of one prey species delayed resistance while the other accelerated resistance. Heimpel et al. [33] reported that the form of egg mortality could influence the rate of resistance, but the importance of egg mortality depended on other ecological processes in the pest population. Other simulation models reported that natural enemies could slow insect resistance to *Bt* crops or *Bt* pesticides [34–36]. As this summary indicates, the literature contains suggestions that natural enemies could delay or accelerate resistance, depending on whether there is a differential impact on susceptible or resistant phenotypes.

In the present study we used a unique system, composed of broccoli plants transformed to express Cry1Ac protein, a population of *Plutella xylostella*, a global pest of crucifers [37] with a low frequency of resistant individuals to Cry1Ac and the insecticide spinosad, and the predaceous ladybird beetle, *Coleomegilla maculata*, to conduct a multigenerational study in the greenhouse. Our objectives were to determine: (1) if a natural enemy can delay the development of insect resistance to a *Bt* crop; and (2) if biological control in conjunction with *Bt* crops can effectively suppress the pest population. In addition, to simulate field-realistic conditions for both the predator and prey, we sprayed refuges with insecticide in some treatments, but not others, and observed those effects on the development of insect resistance.

**Results**

**Population Density of *P. xylostella***

The predator, presence of a refuge, and use of a spray on the refuge each influenced the population dynamics of *P. xylostella* per *Bt* plant over the 6 generations of the experiment. A repeated-measures ANOVA, with generation and treatment as factors, yielded a significant effect for generations (*F*$_{5,164} = 77.101$, *P* < 0.001), for treatments (*F*$_{2,31} = 31.788$, *P* < 0.001), and the interaction term generation*treatments (*F*$_{10,326} = 16.250$, *P* < 0.001). During the 1st generation, few *P. xylostella* were found on *Bt* plants (Fig. 1A), and there were no significant differences between treatments using one-way ANOVA (*F*$_{1,258} = 0.900$). By the 2nd generation, there was an average of 7 *P. xylostella* larvae and pupae per *Bt* plant in the treatment with only *Bt* plants, but the number of *P. xylostella* was still about 1 per plant in the other treatments (*F*$_{1,376} = 3.767, P = 0.026$). *Bt* plants were completely defoliated and control failure was evident in the *Bt* plant-only treatment at the 3rd generation when the number of *P. xylostella* increased to 51 per *Bt* plant (*F*$_{1,667} = 8.667, P = 0.001$).

At the 4th generation, the number of *P. xylostella* increased to 12 per *Bt* plant in treatment *Bt*R+K (75% *Bt* plants +25% non-*Bt* refuge plants) and 29 per *Bt* plant in *Bt*+SR (75% *Bt* plants +25% spinosad-sprayed non-*Bt* refuge plants), significantly higher than those in treatments *Bt*R+Cm (75% *Bt* plants +25% non-*Bt* refuge plants + predator) and *Bt*+SR+Cm (75% *Bt* plants +25% spinosad-sprayed non-*Bt* refuge plants + predator) (*F*$_{1,21} = 21.294, P < 0.001$).

There was still only about 1 *P. xylostella* per *Bt* plant in treatment *Bt*R+Cm at the 5th and 6th generations, significantly lower than other treatments (5th generation: *F*$_{1,59} = 59.203, P < 0.001$; 6th generation: *F*$_{1,61} = 61.164, P < 0.001$). Most importantly, over the 6 generations of the test, only treatment *Bt*R+Cm maintained <2 *P. xylostella* per *Bt* plant at each generation, suggesting the important role that the predator played in maintaining a low pest population.

When spinosad was used (treatment *Bt*+SR+Cm), the pest population was also maintained at a low level, except for a flare-up in the 5th generation when the *P. xylostella* population was >40 per *Bt* plant. Without the use of the predator (*Bt*R+K), the pest population gradually rose and peaked at 61 per plant by the 6th generation, compared to being maintained at about 1 for all generations when the predator was used (*Bt*R+Cm). When the predator was replaced by the insecticide spinosad (*Bt*+SR), the pest population increased more rapidly and peaked at 107 per plant in the 6th generation, again showing the strong and lasting benefit of the predator in maintaining a low pest population.

Examing the pest population on the refuge plants in the cages provided another indication of the overall performance of the treatments. Using repeated measures analysis, there were significant differences for generations (*F*$_{5,283} = 29.341, P < 0.001$), for treatments (*F*$_{1,60} = 6.250, P < 0.001$), and the interaction term generation*treatments (*F*$_{14,960} = 10.071, P < 0.001$). On refuge plants, the density of *P. xylostella* varied between 5–120 *P. xylostella* per plant between generations and treatments (Fig 1B). Only the populations in *Bt*+SR+Cm, which combined *Bt* plants, spinosad and the predator, were consistently the lowest and did not exceed 20 per plant in any generation. Use of the predator alone (*Bt*R+Cm) maintained a relatively low and stable pest population over all 6 generations. In treatments with 100% non-*Bt* refuge plants, *P. xylostella* densities exceeded 100 per plant at each generation despite keeping only 3 defoliated plants with their larvae and pupae in each cage to reduce the overall population. Therefore, we did not include the treatment of only non-*Bt* plants in Figure 1B.

**Population Density of *C. maculata***

In treatments *Bt*R+Cm and *Bt*+SR+Cm, predator populations generally increased as the pest, *P. xylostella*, populations increased. A repeated-measures ANOVA yielded a significant effect for generations (*F*$_{5,383} = 11.873, P = 0.002$), for treatments (*F*$_{1,9} = 9.410, P = 0.022$), and the interaction term generation*treatments (*F*$_{5,383} = 11.225, P = 0.003$). Only a few *C. maculata* adults were found in the 1st generation because only 3 pairs were released at the start of the experiment (Fig. 2A). Predator populations on *Bt* plants in treatment *Bt*R+Cm remained about 1 per plant because of the low pest population, especially on *Bt* plants through the 6th generation (Fig. 1A). In treatment *Bt*+SR+Cm, predator populations remained about 1 per plant until the 5th generation when they increased to >4.5 in the 5th and 6th generations (Fig. 2A).

Mean values for the 5th and 6th generations by the independent-test between the two treatments, respectively, differed significantly (G5: *t*$_{05} = 4.562, P = 0.004$; G6: *t*$_{05} = 6.268, P = 0.001$). This likely maintained the pest population on the *Bt* plants at a low population (Fig. 1A), although resistance increased (Table 1).

Predator populations were generally higher on the refuge plants (Fig. 2B) than on the *Bt* plants (Fig. 2A), likely reflecting the higher pest density on these plants. For the three treatments *Bt*R+Cm, *Bt*+SR+Cm, and non-*Bt*+Cm, there were no significant differences in predator density in most generations. The repeated measures showed significant effects for generations (*F*$_{5,460} = 12.234, P < 0.001$) and the interaction term generation*treatments (*F*$_{6,920} = 3.241, P = 0.012$), but not for the treatments (*F*$_{2,101} = 0.175$).
Resistance to Cry1Ac

Survival of *P. xylostella* larvae indicates resistance and, on *Bt* broccoli, it was significantly affected when analyzed using repeated-measures for generations ($F_{2.714} = 35.792$, $P<0.001$), for treatments ($F_{6} = 81.199$, $P<0.001$), and the interaction term generation*treatments ($F_{16.282} = 6.719$, $P<0.001$). Resistance levels reached 74.9% after only 3 generations in the treatment with *Bt* plants only, significantly greater than the survival in other treatments ($F_{6} = 50.049$, $P<0.001$) and >2x the next highest treatment of *Bt*+SR (75% *Bt* plants + 25% spinosad-sprayed non-*Bt* refuge plants) (Table 1). The high rate of resistance in the treatment with only *Bt* plants was sustained through the 6th generation. In treatment *Bt*+SR, resistance also developed rapidly, with 56.0% *P. xylostella* surviving at the 4th generation, significantly greater than the treatments *Bt*+R, *Bt*+R+*Cm*, *Bt*+SR+*Cm* and the control cages (*Non-Bt* Only and *Non-Bt+Cm*) ($F_{6} = 33.319$, $P<0.001$). The survival rates were 4–7% in treatment *Bt*+SR (with predators) at all generations, but increased to 32.5% and 39.5% in treatment *Bt*+R (without predators) at the 5th and 6th generations, respectively. There were significant differences among treatments (5th generation: $F_{6} = 26.302$, $P<0.001$; 6th generation: $F_{6} = 38.850$, $P<0.001$) with <10% survival in

Figure 1. *Plutella xylostella* populations on *Bt* plants (A) and on non-*Bt* refuge plants (B) in greenhouse cages (Means ± SEM). *Bt*+R: 75% *Bt* and 25% non-*Bt* refuge plants; *Bt*+R+*Cm*: 75% *Bt* and 25% non-*Bt* refuge plants with *C. maculata*; *Bt*+SR: 75% *Bt* and 25% spinosad-sprayed non-*Bt* refuge plants; *Bt*+SR+*Cm*: 75% *Bt* and 25% spinosad-sprayed non-*Bt* refuge plants with *C. maculata*. Different letters within the same generations denote significant differences ($P<0.05$, one-way ANOVA, Games-Howell test for G3 and G6 in A, G1, G3 and G5 in B, other comparisons by Tukey HSD test).

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Bt+R+Cm, Non-Bt Only and Non-Bt+Cm, and >70% survival in Bt+SR, Bt+SR+Cm and Bt Only, while Bt+R had nearly 40% survival in the 6th generation. These significant differences at the 6th generation among Bt+R, Bt+R+Cm and Bt+SR+Cm highlight the ability of the predator to slow the rate of resistance to Bt plants. Treatment Bt+R+Cm, which included the predator, had the lowest survival (6.0%) while the treatment without the predator (Bt+R) had 39.5% survival. Including the predator but using spinosad (Bt+SR+Cm), which has some toxicity to Cm [38], increased the development of resistance to Bt plants as indicated by the 72.8% survival (Table 1).

Resistance to Spinosad
In the treatments in which spinosad was used (Bt+SR and Bt+SR+Cm), only low levels of survival of P. xylostella were detected. Survival of individuals removed from cages and fed spinosad-treated leaves was only about 1% at the 4th and 6th generations.

Discussion
A large body of literature has evaluated the potential ecological risks of Bt crops to non-target organisms including natural enemies of insect pests [1,2,22,39]. However, as is the case with conventional insecticides, the interaction of natural enemies and...
relatively non-toxic to natural enemies, it does have a toxic effect on pest populations, which could result in resistance to individuals emerging from cages. This is so far for many key pests, then matings between resistant insects will greatly reduce the possibility of creating homozygous resistant adults and larvae on "maculata" aggregation, a significant number of them occur in areas where food density is low. In the present study, results indicate that refuges slow resistance, but a sprayed refuge could accelerate it. These findings are in line with our previous field study [42]. In the treatment with only "Bt" plants without refuge plants, control failure and high insect densities were observed in the 3rd generation (two replications) and the 4th generation (the other two replications). Plants in the treatment of spinosad-sprayed non-"Bt" broccoli plants ("Bt+SR") were completely defoliated between the 4th and 5th generations.

In the treatment ("Bt+SR+Cm") in which the predator and spinosad were used, 36.8% of the pest population survived on "Bt" plants by the 5th generation and by the 6th generation that had increased to 72.3% (Table 1). Spinosad killed most susceptible "P. xylostella" on refuge plants, and not enough susceptible individuals mated with the resistant individuals on "Bt" plants. Therefore, resistance development was quick although there were "C. maculata" in the treatment ("Bt+SR+Cm"). However, the pest populations were very low in this treatment with 16.5 "P. xylostella" larvae per refuge plant (Fig. 1A) and 8.9 per "Bt" plant (Fig. 1B) at the 6th generation. In the 6th generation, we found >100 "C. maculata" in each cage. This indicates that the high predator population controlled "P. xylostella" density to a low level, despite the fact that the pest had developed resistance to the "Bt" toxin.

While farmers are concerned with reducing the likelihood of resistance, they are more immediately concerned with lowering the pest population to avoid crop injury. It is well worth noting that in our experiments we not only saw the lowest rate of resistance development in the prey when the predator was not decimated by the use of an insecticide, but also the lowest and least fluctuating pest population on "Bt" plants (Fig. 1A) and low and stable pest populations on non-"Bt" plants in the refuge (Fig. 1B). Thus, our data suggest that farmers can have sustainable management of pests if they combine "Bt" plants with biological control.

In conclusion, this study provides empirical evidence to confirm the theory that natural enemies can delay resistance development to "Bt" plants, but also demonstrates that it can do so while maintaining a low pest density and low crop damage. Non-"Bt" refuges are necessary to delay resistance to "Bt" plants, but spraying refuges could accelerate resistance if sprays reduce the function of important biological control agents of the pest. We suggest that host-plant resistance with "Bt" plants and biological control can be fully compatible within an overall integrated pest management (IPM) program. Our results have significant implications for IPM and insect resistance management (IRM) for "Bt" crops.

**Methods**

**Insects**

Three strains of "Plutella xylostella" were used to create a hybrid population for the cage tests: the susceptible Geneva 88 (SS), the Cry1Ac-resistant (Cry1Ac-RR), and the spinosad-resistant (Pearl-RR) [24,40,43] strains. The hybrid population was created by releasing 100 F1 RS1 (G88 female × Cry1Ac-RR male), 100 RS2 (G88 female × Pearl-RR male) and 300 G88 moths into a cage.

![Table 1. Survival on Cry1Ac leaf (%) (Means ± SEM) of *Plutella xylostella* larvae from adults taken from cages.](http://www.plosone.org/)

| Treatments       | Generation |
|------------------|------------|
|                  | G3         | G4         | G5         | G6         |
| Bt+R             | 3.3±1.86 A | 19.0±6.61 B| 32.5±14.55 BC| 39.5±14.06 B|
| Bt+R+Cm          | 4.7±3.18 A | 7.0±3.06 AB| 5.0±3.51 AB | 6.0±2.86 A  |
| Bt+SR            | 29.3±4.01 B| 56.0±4.74 C| 87.5±1.44 D | 83.8±1.07 D |
| Bt+SR+Cm         | 3.3±1.93 A | 20.7±2.19 B| 36.8±6.85 C | 72.8±3.28 D |
| Bt Only          | 74.9±5.46 C| 72.0±6.76 C| 90.5±3.97 D | 93.5±1.9 D  |
| Non-Bt Only      | 1.0±0.58 A | 1.3±0.33 A | 1.0±0.58 A  | 3.7±1.33 A  |
| Non-Bt+Cm       | 0.7±0.33 A | 1.0±0.58 A | 4.7±1.86 AB | 2.3±1.86 A  |

Bt+R: 75% "Bt" and 25% non-"Bt" refuge plants; Bt+R+Cm: 75% "Bt" and 25% non-"Bt" refuge plants with *C. maculata*; Bt+SR: 75% "Bt" and 25% spinosad-sprayed non-"Bt" refuge plants; Bt+SR+Cm: 75% "Bt" and 25% spinosad-sprayed non-"Bt" refuge plants with *C. maculata*; Bt Only: only "Bt" plants; Non-Bt Only: only non-"Bt" plants; Non-Bt+Cm: only non-"Bt" plants with *C. maculata*.

Different letters within the same column denote significant differences (*P*<0.05, one-way ANOVA, Tukey's test).

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Empirical data have demonstrated the usefulness of refuges for "Bt" crops. Tang et al. [40] reported that pest population growth was influenced by refuge size, with the highest populations occurring in treatments that had either no refuge plants or all refuge plants. That study also confirmed other models [41] that the development of resistance was inversely proportional to the size of the refuge. Most importantly, Tang et al. [40] demonstrated that a balance could be struck by refuge sizes that would delay resistance while at the same time limit the growth of the pest population and damage to the crop, as has been observed in the field [7,9].

In the present study, results indicate that refuges slow resistance, but a sprayed refuge could accelerate it. These findings are in line with our previous field study [42]. In the treatment with only "Bt" plants without refuge plants, control failure and high insect densities were observed in the 3rd generation (two replications) and the 4th generation (the other two replications). Plants in the treatment of spinosad-sprayed non-"Bt" broccoli plants ("Bt+SR") were completely defoliated between the 4th and 5th generations.

In the treatment ("Bt+SR+Cm") in which the predator and spinosad were used, 36.8% of the pest population survived on "Bt" plants by the 5th generation and by the 6th generation that had increased to 72.3% (Table 1). Spinosad killed most susceptible "P. xylostella" on refuge plants, and not enough susceptible individuals mated with the resistant individuals on "Bt" plants. Therefore, resistance development was quick although there were "C. maculata" in the treatment ("Bt+SR+Cm"). However, the pest populations were very low in this treatment with 16.5 "P. xylostella" larvae per refuge plant (Fig. 1A) and 8.9 per "Bt" plant (Fig. 1B) at the 6th generation. In the 6th generation, we found >100 "C. maculata" in each cage. This indicates that the high predator population controlled "P. xylostella" density to a low level, despite the fact that the pest had developed resistance to the "Bt" toxin.

While farmers are concerned with reducing the likelihood of resistance, they are more immediately concerned with lowering the pest population to avoid crop injury. It is well worth noting that in our experiments we not only saw the lowest rate of resistance development in the prey when the predator was not decimated by the use of an insecticide, but also the lowest and least fluctuating pest population on "Bt" plants (Fig. 1A) and low and stable pest populations on non-"Bt" plants in the refuge (Fig. 1B). Thus, our data suggest that farmers can have sustainable management of pests if they combine "Bt" plants with biological control.

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The total number of moths in the cage was 500 with a 1:1 ratio for female and male moths from each strain. Eggs were collected from the cage and put on artificial diet to rear F1 larvae. About 1,000 moths from F1-F4 were used to produce a synthetic population. F5 pupae were used in the selection experiments. The expected allele frequency of the synthetic population (square root of survival rate) was 0.1 for Cry1Ac and spinosad resistance. The mean survival of F3 larvae was 0.033 on Cry1Ac plants and 0.025 on spinosad-sprayed plants. Therefore, the actual initial allele frequency at the start of the experiment was estimated to be 0.057 (square root of 0.033) for Cry1Ac resistance and 0.050 (square root of 0.025) for spinosad resistance [43]. While this is higher than would be expected in the field initially when Bt crops are released (perhaps $10^{-3}$ or lower [41]), we expected that these initial frequencies would allow us to see differences among the treatments in a reasonable time frame.

Larvae and adult C. maculata were obtained from DuPont Pioneer (Johnston, IA) and maintained in a climatic chamber at 27±1°C, 65±5% RH and 16:8 L: D at Cornell University’s Department of Entomology at Geneva, NY. Both larvae and adults were reared on decapsulated eggs of brine shrimp, Artemia franciscana, (Brine Shrimp Direct, Ogden UT) and 1.5% agar solution provided separately as a water source.

Transgenic broccoli plants and insecticide

We used Brassica oleracea producing high levels of Cry1Ac for our Bt broccoli plants [44]. Bt broccoli plants with 8 true leaves were used, and analysis by ELISA indicated that the Cry1Ac protein level was 12.3±1.62 µg/g fresh leaf tissue (n = 7). To ensure the activity of the Bt broccoli, the plants were screened with P. xylostella neonates of F1 heterozygotes (G88 x Cry1Ac-RR) when plants were 4 to 5 wk old [45]. The Cry1Ac plants that killed 100% of neonates of F1 heterozygotes (SS x Cry1Ac-RR), indicating high levels of expression in the Bt plants, were used in the greenhouse experiments. Non-Bt broccoli (cv. Packman) was used in the refuge.

A commercial formulation of spinosad (SpinTor 2 SC, 240 g [AI]/Liter) was used. Refuge plants in the treatments B1 and B2 were sprayed in the cages using a small hand-held sprayer. The Cry1Ac plants were covered during the sprays to avoid drift. The concentration used was 90 ppm, which was the lowest field dose listed on the insecticide label. The insecticide-treated refuges were sprayed at the 1st, 3rd and 5th generations in order to keep the P. xylostella population in check in the cages.

Table 2. Experimental treatments.

| Group | Treatments | Replications |
|-------|------------|--------------|
| Bt+R  | 75% Bt plants and 25% non-Bt refuge plants | 4 |
| Bt+R+Cm | 75% Bt plants and 25% non-Bt refuge plants and C. maculata | 4 |
| Bt+SR | 75% Bt plants and 25% non-Bt refuge plants treated with spinosad | 4 |
| Bt+SR+Cm | 75% Bt plants and 25% non-Bt refuge plants treated with spinosad and C. maculata | 4 |
| Bt Only | 100% Bt plants only | 4 |
| Non-Bt Only | 100% Non-Bt refuge plants | 3 |
| Non-Bt+Cm | 100% Non-Bt refuge plants and C. maculata | 3 |

R: refuge.  
SR: refuge with spinosad.  
Cm: C. maculata  
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Experimental designs

The selection experiment was conducted in greenhouses at Cornell University’s New York State Agricultural Experiment Station and followed the general procedures used in previous studies [40, 43]. Each cage was 1.8 m long x 0.9 m wide x 1.7 m high and constructed of nylon netting. Adults, but not larvae, could easily move between the different broccoli types, which were separated by a nylon-netting barrier (0.9 m high). This arrangement simulated adjoining fields with frequent inter-field movement by adults but negligible movement of larvae.

Seven treatments were included in the experiment (Table 2). There were 16 plants total in each cage (12 Bt plants plus 4 non-Bt refuge plants), 250 F4 pupae of the synthetic P. xylostella population were released into each cage. Three pairs of 1-week old C. maculata were released into cages of the treatments Bt+R+Cm, Bt+SR+Cm and non-Bt+Cm when the P. xylostella larvae were 2nd instars. During the pupal period, all old plants were cut at the base of the stem, and plants and pots (which might have pupae on them) were kept in the cages for a week in order to allow adults to emerge. New plants were introduced into the cage to provide foliage for egg laying or existing larvae that had defoliated plants. There were four replications (cages) of Bt-plant treatments (Bt+R, Bt+R+Cm, Bt+SR, Bt+SR+Cm and Bt Only) and three of the controls (Non-Bt Only and non-Bt+Cm).

Data collection

Older larvae (primarily 3rd or 4th instars) and pupae of P. xylostella, and pupae and adults of C. maculata on broccoli plants were counted every generation when larval and pupal densities peaked. To test for resistance, 30–40 larvae from non-Bt refuge plants were collected from each cage at the 3rd, 4th and 5th generations. The larvae were reared on diet in the laboratory, and then adults were allowed to mate and eggs were harvested and tested as described below. At the 6th generation, we collected at least 60 pupae on refuge plants from most of the cages. The survival of 2nd instars derived from the pupae collected in each cage was tested on Cry1Ac broccoli leaf disks in 30-ml plastic cups [43]. For each cage, a total of 100 larvae in 10 replications were tested on Bt broccoli, and non-Bt broccoli was used as a control. We also tested survival of 2nd instars derived from B1 and B2 treatments on spinosad-dipped broccoli leaf disks in 30-ml plastic cups at the diagnostic dose of 10 ppm [45]. A total of 100 larvae were tested (10 replications, 10 larvae/rep) for each cage. For both Cry1Ac plants and spinosad treatments, survival was determined after 3 days at 27±1°C, 50±10% RH and 16:8 LD photoperiod.
Statistical analysis
All statistical analyses were conducted using SPSS 17.0 Windows [6]. Descriptive statistics are given as mean values and standard errors of the mean. Because the data fit the assumptions for parametric analysis, the repeated measures ANOVA was used with the factors of generations and treatments for analysis of the population of P. xylostella on Bt and non-Bt plants and the survival on Cry1Ac leaves of P. xylostella larvae from adults taken from cages. The survival rates were transformed by square root before analysis. For each generation, the differences of resistance development and the population of P. xylostella and C. maculata were analyzed by one-way ANOVA, and means were compared by Tukey HSD, if the data fit homoscedasticity, and Games-Howell test if not. Differences of the mean values in the population of C. maculata per Bt plants between the treatments of refuge + C. maculata and sprayed-refuge + C. maculata were examined by independent t-tests. In all tests P values <0.05 were considered significant.

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Author Contributions
Conceived and designed the experiments: XXL, MC, DO RR AMS. Performed the experiments: XXL, MC, AMS. Analyzed the data: XXL, MC, AMS. Contributed reagents/materials/analysis tools: HLC EDE. Wrote the paper: XXL, AMS. Discussed and helped analyze the results: QWZ.

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Author/s:
Liu, X; Chen, M; Collins, HL; Onstad, DW; Roush, RT; Zhang, Q; Earle, ED; Shelton, AM

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