A Comprehensive Assessment of the Effects of Transgenic Cry1Ac/Cry1Ab Rice Huahui 1 on Adult *Micraspis discolor* (Fabricius) (Coleoptera: Coccinellidae)

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Abstract

*Micraspis discolor* (Fabricius) (Coleoptera: Coccinellidae) is a widely distributed coleoptera predator in southern Asia in rice ecosystem, and adult *M. discolor* feed on both rice pollen and soft-bodied arthropods. Bitrophic bioassay and tritrophic bioassay were conducted to evaluate the potential impact of Cry1Ac/Cry1Ab-expressing rice Huahui 1 and its non-transgenic counterpart Minghui 63 on fitness parameters of adult *M. discolor*. The results showed that the survival, and fecundity of this beetle's adults were not different when they fed on Bt rice or non-Bt rice pollen or *Nilaparvata lugens* (Stål) reared on Bt rice or non-Bt rice. Toxicity assessment to ensure *M. discolor* adults were not sensitive to Cry1Ab or Cry1Ac protein independent from the pollen background, *M. discolor* adults were fed with an artificial diet containing Cry1Ac, Cry1Ab or both protein approximately 10 times higher concentration than in Huahui 1 rice pollen. No difference was detected for any of the life-table parameters tested between Cry protein-containing and pure diet. Artificial diet containing E-64 (N-(trans-Epoxysuccinyl)-L-leucine 4-guanidinobutylamide) was included as a positive control. In contrast, the pre-oviposition and fecundity of *M. discolor* were significantly adversely affected by feeding on E-64-containing diet. In both bioassays, the uptakes of Cry protein by adult *M. discolor* were tested by ELISA measurements. These results indicated that adults of *M. discolor* are not affected by Cry1Ab- or Cry1Ac-expressing rice pollen and are not sensitive to Cry protein at concentrations exceeding the levels in rice pollen in Huahui1. This suggests that *M. discolor* adults would not be harmed by Cry1Ac/Cry1Ab rice if Bt rice Huahui 1 were commercialized.
Introduction

Rice, *Oryza sativa* L., is a main food crop for 3.5 billion people in the world [1]. Leaffolders such as *Chilo suppressalis* (Walker) (Lepidoptera: Pyralidae), *Scirpophaga incertulas* (Walker) (Lepidoptera: Pyralidae) and *Cnaphalocrocis medinalis* Guenée (Lepidoptera: Pyralidae) are three of the most serious pests of rice in temperate to tropical Asia [2–4]. Many synthetic *Bt* genes including *cry1Ab*, *cry1Ac* and *cry1B*, are derived from the bacterium *Bacillus thuringiensis* Berliner transferred into rice, conferring high resistance against lepidopteran pests have been developed since 1993 [4–6].

Huahui 1 is a transgenic *Bt* rice created by fusion a synthetic *cry1Ab/cry1Ac* gene into rice *MingHui63* [7–8], and it is a lepidopteran resistant variety. From 2009 to 2014, Huahui 1 and its hybrid line *Bt* Shanyou 63 were issued a safety certificate and approved for limited release in farmers’ fields in the Hubei Province in China [8–9].

*Micraspis discolor* is distributed from Southern China to Southern Asia [10–13]. It feeds on plants’ pollen, such as pollen of *Triticeae Dumort, Oryza sativa* L., as well as on a wide range of soft-bodied arthropods and eggs. It has been reported as both phytophagous and entomophagous. In rice ecosystems, adult *M. discolor* feed on rice pollen and *Nilaparvata lugens* (Stål) (BPH), *Sogatella furcifera* (Horváth) (WBPH), aphids and eggs or larvae of *C. suppressalis* and *T. incertulas* etc.

It has been recorded that planthoppers -BPH and WBPH are the predominant prey of *Micraspis* spp. [14–16], so *M. discolor* can play important roles in natural control of pest insects [10–14]. Feeding preference studies showed that adults of *M. discolor* have a significant preference for rice pollen over all insects from rice fields in paired-choice tests [14]. 1st and 2nd instar larvae mainly feed on rice pollen. *M. discolor* can complete its life on rice pollen or BPH alone [11].

Previous studies have demonstrated that *Bt* crop pollen expressing *Cry1Ab*, *Cry3A* and *Cry3Bb1*, poses little risk to predators [17–20]. As Huahui 1 may be commercialized in the future, the impact of *Bt* rice on *M. discolor* should be carefully evaluated before its commercialization. In this study, a comprehensive assessment of the effects of transgenic *Cry1Ac/Cry1Ab* rice Huahui 1 on adult *M. discolor* were investigated.

Results

The abundance of *Micraspis discolor* and other predatory Coleoptera in rice ecosystem in different rice growth periods in 3 rice-cropping seasons

Field studies showed that there were same trends of the abundance change of coleoptera predators in rice ecosystem in 3 rice-cropping seasons in Minghui 63 paddies. 3 species of coleoptera predators, *M. discolor*, *Ophionea indica* (Thunberg), *Paederus fuscipes* Curtis were main predators which feed on soft-bodied insects in whole rice growth period. In Sanya rice ecosystem, adult *M. discolor* was the most dominant coleoptera predator in whole rice growth period especially in blooming stage in 3 rice seasons (Fig 1) (S1 File). Adults *M. discolor* were 19.85±2.20, 28.35±0.89, 24.55±1.73 in each replicate in early-blooming stage in Spring, Summer and Autumn rice seasons, 2–6 fold the quantities in booting, tillering and filling stage of 3 rice-cropping seasons.

*Bt* rice pollen bioassay

Over 75% percent adult *M. discolor* survived the 26 days pollen feeding period, and the survival rates of males were slightly higher than that of females (Table 1) (S2 File). For both sexes,
survival did not differ between the Bt and non-Bt pollen treatments. Similarly, the pre-oviposition period, total fecundity and female and male dry weight were not affected by the Bt pollen treatment (Table 1). Additionally, the daily fecundity of females was not significantly affected by feeding on Cry1Ac/Cry1Ab -expressing rice pollen (RM-ANOVA; \(F_{1,76} = 0.12, P = 0.74\)) (Fig 2) (S3 File).

Table 1. Impact of the consumption of pollen from Cry1Ab/Cry1Ac-expressing Bt rice or the corresponding non-transformed rice plant on different life-table parameters of adult Micraspis discolor. Data are provided as mean (±SE).

| Fitness indicators                  | Minghui 63 pollen | Huahui 1 pollen | \(\chi^2\) | \(P\) |
|-------------------------------------|------------------|-----------------|------------|------|
| Female survival (%) \(a\)           | 76.7(30)         | 80.0(30)        | \(\chi^2 = 0.10\) | \(p = 0.75\) |
| Male survival (%) \(a\)             | 80.0(30)         | 83.3(30)        | \(\chi^2 = 0.11\) | \(p = 0.74\) |
| Pre-ovipos. period (days) \(b\)     | 13.17±0.08(30)   | 13.06±0.10(30)  | \(U = 73787.5\) | \(p = 0.51\) |
| Fecundity (eggs/female) \(c\)       | 19.97±0.37(30)   | 19.07±0.39(30)  | \(t = 1.68\) | \(p = 0.09\) |
| Female adult dry weight (mg) \(c\)  | 6.14±0.07(16)    | 6.26±0.06(17)   | \(t = -1.38\) | \(p = 0.17\) |
| Female adult dry weight (mg) \(c\)  | 3.97±0.07(17)    | 3.83±0.05(17)   | \(t = 1.59\) | \(p = 0.12\) |

The experiment was terminated after 26 days. \(N = 30\).

\(a\) Chi- square test
\(b\) Mann- Whitney U test
\(c\) Student’s t test

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The ELISA measurements showed that the mean (±SE) concentration of Cry1Ab, Cry1Ac protein in the pollen was 15.26 ± 0.02 μg/g and 11.92±0.01 μg/g dry weight. No Cry1Ac or Cry1Ab protein was detected in any sample of non-Bt rice Minghui63 pollen.

When the experiment was terminated after 26 days, ELISA results showed that adults fed on Bt rice pollen contained Cry1Ac/Cry1Ab concentrations with a mean (±SE) of 2.09±0.01 μg/g Cry1Ac and 1.55±0.02 μg/g Cry1Ab dry weight in female adults, 0.36±0.11 μg/g Cry1Ac and 0.25±0.01 μg/g Cry1Ab dry weight in male adults. No Cry toxin was detected in adults fed on non-Bt rice pollen.

**BPH reared on Bt rice or non-Bt rice bioassay**

Over 70% percent adult *M. discolor* fed BPH reared on Bt rice Huahui1 or Minghui 63 plants survived the 26 days period (Table 2) (S4 File). For both sexes, survival did not differ between BPH from Bt and non-Bt treatments. The pre-oviposition period, total fecundity and female and male dry weight were not different between 2 treatments (Table 2) (S4 File). The daily fecundity of females was not significantly affected by feeding on BPH from Huahui 1 rice (RM-ANOVA; F1,64 = 0.67, P = 0.42) (Fig 3) (S5 File). The experiment was terminated after 26 days. N = 30.

**Table 2. Impact of the consumption of BPH reared on Cry1Ab/Cry1Ac-expressing Bt rice or the corresponding non-transformed rice plants on different life-table parameters of adult *Micraspis discolor*.** Data are provided as mean (±SE).

| Fitness indicators                  | BPH on Minghui 63 | BPH on Huahui 1 |
|-------------------------------------|-------------------|-----------------|
| Female survival (%)                 | 76.7(30)          | 73.3(30)        |
| Male survival (%)                   | 70.0(30)          | 73.3(30)        |
| Pre-oviposit. period (days)         | 13.13±0.08(30)    | 13.09±0.09(30)  |
| Fecundity (eggs/female)             | 18.40±0.74(30)    | 19.71±0.73(30)  |
| Female adult dry weight (mg)        | 6.09±0.07(15)     | 6.25±0.06(16)   |
| Female adult dry weight (mg)        | 3.91±0.07(15)     | 3.88±0.06(17)   |

The experiment was terminated after 26 days. N = 30.

a Chi- square test  
b Mann- Whitney U test  
c Student’s t test
The ELISA measurements showed that the mean (±SE) concentration of Cry1Ab, Cry1Ac protein in the BPH reared on Huahui 1 was 3.23 ± 0.01 μg/g and 2.46±0.01 μg/g dry weight. No Cry1Ac or Cry1Ab protein was detected in any sample reared on non-Bt rice.

When the experiment was terminated after 26 days, ELISA results showed that female adults fed on BPH reared on Bt rice contained Cry1Ac/Cry1Ab concentrations with a mean (±SE) of 0.15±0.00 μg/g Cry1Ab and 0.11±0.00 μg/g Cry1Ac dry weight. Most male adults samples were detected no Cry1Ac/Cry1Ab concentration. No Cry toxin was detected in adults fed on BPH reared on non-Bt rice.

Purified Cry toxin bioassay

The survival rates of adult M. discolor were above 65% and did not differ between adults feeding on pure artificial diet or any Cry toxin treatment (Chi-square test, P > 0.05) (Table 3). While consumption of diet containing E-64 caused a decline in adult survival by more

| Fitness indicators                      | Cry1Ab          | Cry1Ac          | Cry1Ab/Cry1Ac | control | E-64          |
|-----------------------------------------|-----------------|-----------------|---------------|---------|---------------|
| Survival(%)a                           | 80.6(31)        | 79.7(32)        | 75.9(29)      | 78.4(37)| 65.5(29)      |
| Pre-oviposit. Period(days) b            | 13.87±0.70(31)  | 13.84±0.64(32)  | 13.62±0.62(29)| 13.92±0.48(37)| 16.79±0.51(29)** |
| Fecundity (eggs/female) c               | 18.77±1.52(31)  | 19.44±1.60(32)  | 19.31±1.55(29)| 20.86±1.41(37)| 6.79±0.61(29)** |
| Female adult dry weight (mg) e          | 6.30±0.18(14)   | 6.34±0.21(14)   | 6.16±0.28(13) | 6.43±0.17(14) | 6.09±0.19 (13)  |
| Male adult fresh weight (mg) f          | 4.03±0.16(14)   | 3.99±0.19(14)   | 3.97±0.16(13) | 3.99±0.18(14) | 3.81±0.15(13)   |

35–39 pairs of adult M. discolor were fed an artificial diet containing 150 mg/g Cry1Ab, 120 mg/g Cry1Ac, 150 mg/g Cry1Ab and 120 mg/g Cry1Ac or 25 mg/g E-64 (positive control) per g dry weight of artificial diet. Pure artificial diet served as a negative control. Pairs producing no eggs were removed from the analyses.

Statistical comparisons were made between each of the Cry1Ab, Cry1Ac, Cry1Ab/Cry1Ac and E-64 treatments with the control. Asterisks denote significant differences:

** p < 0.01

a Chi-square test
b Mann-Whitney U-test
c Dunnett test
than 10%, this difference also was not significant ($\chi^2 = 1.36, P = 0.24$) (Table 3) (S6 File). No significant differences were detected between the Cry1Ac/Cry1Ab treatment and the control for the pre-oviposition period, total fecundity and adult dry weight (all $P > 0.05$) (S6 File). In contrast, the pre-oviposition period was significantly prolonged (Mann—Whitney U test; $U = 58347.5, p = 0$), the total fecundity was significantly reduced (Dunnett test; $P < 0.05$) when \textit{M. discolor} adults were fed with the E-64-containing diet compared with those fed with pure artificial diet (Table 3) (S6 File). Likewise daily fecundity did not differ between the Cry1Ab, Cry1Ac or Cry1Ab/ Cry1Ac with control treatments ($P > 0.05$), while E-64 caused a significant reduction in daily fecundity (RM-ANOVA; $F_{1,49} = 54.364, P < 0.01$) (Fig 4) (S7 File).

After 26 days diet feeding, the ingestion of Cry1Ac/Cry1Ab by adult \textit{M. discolor} was confirmed by ELISA. Similar to Bt rice pollen feeding, the females contained much higher mean ($\pm$SE) concentrations of the protein (Cry1Ab $2.55 \pm 0.02 \, \mu g/g$, Cry1Ac $1.89\pm 0.03 \, \mu g/g$ dry weight) than males (Cry1Ab $0.41 \pm 0.01 \mu g/g$ and Cry1Ac $0.31 \pm 0.02 \mu g/g$ dry weight). No Bt protein was detected in adults fed pure artificial diet.

**Discussion**

New high yielding varieties, pesticides and nitrogen fertilizers have doubled the production of many crops including rice [21–22]. But the inputs of insecticides and fertilizers have resulted in some negative effects on environments [23–25]. Current methods for controlling insect pests mainly include good farming practices, biological control, breeding and growing resistant varieties, and the use of pesticides. However, the application of chemical insecticide is the major control measure for rice insect pests in Asia, which not only causes severe environmental pollution and the resurgence of herbivores but also reduces populations of the natural enemies of herbivores [23–25].

The Cry1Ab/Cry1Ac protein expressed \textit{Bt} rice Huahui 1 was issued a safety certificate in China in 2009. It is lepidopteran specific with high activity against \textit{C. suppressalis}, \textit{S. incertulas} and many other lepidopteran pests [5], [26–28]. Although predators were not the target insects,
M. discolor may have been altered due to the introduction of Cry1Ab/Cry1Ac gene. Such a potential effect requires proper assessment of the ecological risks posed by Bt rice, by carrying out a combination of laboratory tests, field experiments and longer-term monitoring, as suggested by Jeppson et al. [29].

In the laboratory study, ELISA measurements showed that Bt rice Huahui 1 pollen contained high concentrations of Cry1Ab/Cry1Ac, and Cry proteins also were detected in M. discolor when fed Bt rice pollen, although the Bt protein levels were 7 times lower for Cry1Ab and 6 times lower for Cry1Ac compared to the concentrations in the Bt rice pollen. The Bt protein levels of BPH were much lower than that in Bt rice pollen, the bitrophic results of BPH was the same as Huahui 1 pollen feeding bioassay. Same reports have been reported from other bitrophic and tritrophic experiments in which Bt plants pollen was used to expose predators to the insecticidal compounds [17, 20, 29–31].

The pollen/BPH feeding bioassays showed no adverse effect on the fitness of adult M. discolor after ingestion of Bt rice Huahui 1 pollen/BPH expressing Cry1Ab/Cry1Ac compared with pollen from the corresponding nontransformed rice Minghui 63. Similarly no negative effects on the survival and reproduction of adult M. discolor were observed when these beetles were provided with an artificial diet containing Cry1Ab, Cry1Ac or Cry1Ab/Cry1Ac at a concentration that was approximately 10 times higher than that measured in the Bt rice pollen. E-64 treatment (positive control) caused a significant prolongation of the pre-oviposition period and a decrease in fecundity. E-64 has been used as a positive control in insect bioassays for toxicity assessments, such as Coleomegilla maculata [32–33].

Our study indicated that Bt rice pollen or BPH reared on Bt rice has no adverse effect on M. discolor which fed on rice pollen or BPH alone. This result agrees with the studies of Wang et al., adults of Chrysoperla sinica are not affected by Cry2Aa-expressing rice pollen and are not sensitive to Cry2Aa protein at concentrations exceeding the levels in pollen [34]. Bai et al. also reported Cry1Ab toxin expressed in Bt rice pollen had no evident negative impacts on Propylea japonica fitness when the pollen was used as a food by this beetle [26], Other rice crop also showed the same results. Li et al. reported adults of Chrysoperla carnea (Stephens) are not affected by Bt maize pollen and Adalia bipunctata are not sensitive to Cry1Ab and Cry3Bb1 at concentrations exceeding the levels in pollen [35–36]. Li et al. also reported Cry1Ac and Cry2Ab Bt cotton demonstrates no detrimental effects on Coleomegilla maculata [32–33].

The E-64 treatment (positive control) caused a significant prolongation of the pre-oviposition period and a decrease in fecundity. Similar trend was observed for female M. discolor. These results demonstrate that the experimental system was able to detect treatment effects if present and is suitable for assessing the hazards of orally active insecticidal compounds on adult M. discolor. E-64 has been used as a positive control in insect bioassays for toxicity assessments, namely for Coleomegilla maculata [32–33] and Coccinella septempunctata [37].

In summary, this study using bitropic and tritropic bioassays and a direct feeding bioassay provides the most complete information so far on the potential direct and indirect effects of Cry1Ab and Cry1Ac on the ladybird beetle, M. discolor, a common and abundant predator in rice systems. ELISA measurements and sensitive-insect bioassays confirmed that the test insects were exposed to high concentrations of biologically active Cry proteins throughout the duration of the bioassays. This suggests that M. discolor adults would not be harmed by Cry1Ab/Cry1Ac rice if transgenic Bt rice Huahui 1 were commercialized.

Reports showed that Bt rice Shanyou 63 which was produced by crossbreeding Zhenshan 97A with Huahui 1 reduced pesticide applications 80% [38]. Meanwhile Bt rice Huahui 1 does not affect a important predator M. discolor adults. Consequently, Huahui 1 would not pose negative effects on both environment and predators if it were commercialized.
Materials and Methods

Ethics Statement

No specific permits were required for the described studies.

Plant materials

Transgenic Bt Huahui 1 and its non-transgenic counterpart Minghui 63 were evaluated.

Insects

*M. discolor* and *Nilaparvata lugens* (Stal) were collected from rice fields in the experimental farm of Sanya Science & Technology Academy for crop winter multiplication during May 2012 and since then maintained in the laboratory without introductions of field-collected insects. Eggs collected from the colony were kept in a climatic chamber at 16:8 h (L: D), 28 ± 1°C, and 75 ± 5%. After hatching, the larvae were kept individually in glass tubes (2 cm diameter, 9 cm long). 5–15 Drosophila adults were supplied daily to the tubes. After 6 generation newly emerged (<24 h after emergence) adults *M. discolor* were used for the experiments.

*N. lugens* maintained in rice Yuzhenxiang seedlings in the greenhouse in the experimental base of Tropical biological technology institute in Wenchang county. After more than 20 generation *N. lugens* were transferred to Huahui1 and Minghui 63 seedlings, 3rd instar *N. lugens* from Huahui1 and Minghui 63 were used for the experiments.

Purified proteins

Cry1Ab toxin and Cry1Ac toxin (99%) used in this study was provided by Envirologix Inc. E-64 (advanced pure) was provided by the Amresco Company.

The abundance of Coleoptera predators in rice ecosystem

There are three rice-cropping seasons according to sowing date in the studied area. The non-transgenic control Minghui 63 were sowed in early December in 2012, early April in 2013 and early July in 2013, and transplanted 1 month later. Field insect sampling by the D-vac method in minghui63 field were conducted at the experimental farm of Sanya Science & Technology Academy for crop winter multiplication in Hainan Province, China in 2013. Five samples were taken from each plot. For taking a sample, a plastic bin covered 20x20 cm² was placed over one hill and all arthropods inside the bin were carefully collected with a blower-vac machine sampler. There were 3 replicates, and each plot was 400 m². Rice seedlings were transplanted by hand at a density of 3 seedlings per hill. Spacing between rows was 20 cm and hills within rows were spaced 15 cm apart.

Field insect sampling were conducted in rice tillering, booting, early-blooming, blooming and filling stage with five point sampling method in every rice seasons. Each sampling site contained 9 rice hills. All treatments were managed by normal cultural practices except for pesticide sprays. Collected insects were put in 80% alcohol, and Coleoptera predators were counted in laboratory.

Rice pollen

Pollen was obtained from greenhouse-grown rice of the *Bt* rice lines Huahui 1 and the non-transgenic line Minghui 63 in the Wenchang experimental base of Tropical biological technology institute.
In the morning, inflorescences were cut from plants before anthesis. The obtained pollens was placed in sealed plastic vials and kept at $-20^\circ$C until use. In the laboratory, anthers enclosed within glumes were striped off by scissors.

**Bt rice pollen bioassay and BPH bioassay**

The experiment was conducted in the climatic carine with photoperiod 16:8 h (L: D), 28±1°C, and 75 ± 5% RH.

For each treatment, new emerged adults were transferred to a Petri dish (10.0 cm diameter, 2.6 cm tall). After 3 days, adults were paired and placed in glass tubes (2cm diameter, 9cm long) containing sufficient Bt or non-Bt rice pollens or BPH reared on Huahui1 or Minghui 63 and a water-saturated cotton ball with 5% leechee honey. Thereafter, daily fecundity and adult survival were recorded every 6 h. After 26 days, dry male and female adults were weighed.

When egg clutches were observed, adults *M. discolor* as well as the rice pollens/BPH were transferred to other clear glass tubes, and the numbers of egg clutches were recorded.

**Purified toxin bioassay**

The purified toxin bioassay was carried out as described above. The experimental system was used to assess the toxicity of Cry1Ac and Cry1Ab on *M. discolor*. Five diet treatments were tested: 1) artificial diet (negative control) containing of litchi honey, egg yolk powder (in proportions 6:3); 2) artificial diet containing Cry1Ab at 150 mg/g weight of diet; 3) artificial diet containing Cry1Ac at 120 mg/g dry weight of diet; 4) artificial diet containing Cry1Ab at 150 mg/g and Cry1Ac at 120 mg/g dry weight of diet; 5) artificial diet containing E-64 at 25 mg/g fresh weight of diet (positive control). Diets were replaced every day. The experiment was conducted with 30–33 pairs neonate *M. discolor* adults per treatment. The insects were observed every 6 h and their survival and development were recorded.

**ELISA measurements**

The concentrations of Cry1Ab and Cry1Ac in rice pollen, adult *M. discolor* and BPH were measured by enzyme-linked immunosorbent assays (ELISA) using the Cry1Ab/Cry1Ac detection kits from Envirologix (Portland Maine, USA). Prior to analysis, insects were washed in phosphate buffered saline with Tween-20 (PBST) buffer (provided in the kit) to remove any Bt toxin from their outer surface.

After adding PBST to the samples at a ratio of at least 1:10 (mg sample:ml buffer) in 1.5 ml centrifuge tubes, the samples were fully ground by hand using a plastic pestle. After centrifugation and appropriate dilution of the supernatants, ELISA was performed according to the kit’s instructions. The measured OD values were calibrated to a range of concentrations of Cry1Ab and Cry1Ac made from purified toxin solution.

**Data analysis**

For the pollen bioassay, statistical analyses were conducted between the Bt pollen and non-Bt pollen treatments. For the BPH bioassay, statistical analyses were conducted between BPH from Bt rice Huahui 1 and non-Bt rice Minghui 63 treatments. In the purified toxin experiment, comparisons were made between the Cry1Ac, Cry1Ab, Cry1Ac/Cry1Ab or E-64 treatment with control (pure artificial diet). For both experiments, Chi-square tests were carried out to compare adult survival (females and males separately). Mann—Whitney U-tests were used to compare the preoviposition. Data on the total fecundity and adult dry weight were compared.
using Student’s t-tests in the pollen feeding bioassay and by Dunnett tests in the purified protein bioassay. Daily fecundity data were analyzed using repeated measures (RM) ANOVA. All statistical analyses were conducted using the software package SPSS (version 16; SPSS, Inc., Chicago, IL).

Supporting Information

S1 File. Different abundance of 3 predators in booting(I), tilling(II), early-blooming(III), blooming(IV) and filling(V) phase in Spring, Summer and Autumn rice seasons in 2013 (Fig 1). (XLS)

S2 File. Impact of the consumption of pollen from Cry1Ab/Cry1Ac-expressing Bt rice or the corresponding non-transformed rice plant on different life-table parameters of adult Micraspis discolor (Table 1). (XLS)

S3 File. Daily fecundity of Micraspis discolor fed pollen from Cry1Ab/Cry1Ac Bt rice or the corresponding non-transformed rice plants (Fig 2). (XLS)

S4 File. Impact of the consumption of BPH from Cry1Ab/Cry1Ac Bt rice or the corresponding non-transformed rice plants on different life-table parameters of adult Micraspis discolor (Table 2). (XLS)

S5 File. Daily fecundity of Micraspis discolor fed BPH from Cry1Ab/Cry1Ac Bt rice or the corresponding non-transformed rice plant (Fig 3). (XLS)

S6 File. Effect of feeding a pure artificial diet or artificial diet with purified Cry1Ab, Cry1Ac and E-64 provided on different life-table parameters of adult Micraspis discolor (Table 3). (XLS)

S7 File. Daily fecundity of Micraspis discolor fed an artificial diet containing 150 mg/g Cry1Ab, 120 mg/g Cry1Ac, 150 mg/g Cry1Ab and 120 mg/g Cry1Ac or 25 mg/g E-64 (positive control) per g dry weight of artificial diet (Fig 4). (XLS)

Author Contributions

Conceived and designed the experiments: XZ APG. Performed the experiments: XZ HK JZ RZJ. Analyzed the data: XZ QXH YLG LX. Contributed reagents/materials/analysis tools: QXH. Wrote the paper: XZ RZJ APG.

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