Testing Pollen of Single and Stacked Insect-Resistant Bt-Maize on *In vitro* Reared Honey Bee Larvae

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**Abstract**

The ecologically and economically important honey bee (*Apis mellifera*) is a key non-target arthropod species in environmental risk assessment (ERA) of genetically modified (GM) crops. Honey bee larvae are directly exposed to transgenic products by the consumption of GM pollen. But most ERA studies only consider responses of adult bees, although Bt-proteins primarily affect the larval phases of target organisms. We adopted an *in vitro* larvae rearing system, to assess lethal and sublethal effects of Bt-pollen consumption in a standardized eco-toxicological bioassay. The effects of pollen from two Bt-maize cultivars, one expressing a single and the other a total of three Bt-proteins, on the survival and prepupa/pupal weight of honey bee larvae were analyzed. The control treatments included pollen from three non-transgenic maize varieties and of *Heliconia rostrata*. Three days old larvae were fed the realistic exposure dose of 2 mg pollen within the semi-artificial diet. The larvae were monitored over 120 h, until the prepupal stage, where larvae terminate feeding and growing. Neither single nor stacked Bt-maize pollen showed an adverse effect on larval survival and the prepupal weight. In contrast, feeding of *H. rostrata* pollen caused significant toxic effects. The results of this study indicate that pollen of the tested Bt-varieties does not harm the development of *in vitro* reared *A. mellifera* larvae. To sustain the ecosystem service of pollination, Bt-impact on *A. mellifera* should always be a crucial part of regulatory biosafety assessments. We suggest that our approach of feeding GM pollen on *in vitro* reared honey bee larvae is well suited of becoming a standard bioassay in regulatory risk assessment schemes of GM crops.

**Introduction**

Pollinators provide key ecosystem services by maintaining both the biodiversity of wild plants and agricultural productivity [1,2] at an estimated value of US $217 billion yearly [3]. The most important pollinator species worldwide is the honey bee *Apis mellifera* [4], with populations present in all countries growing genetically modified (GM) crops [5,6]. Hence, honey bees are a key non-target test species for assessing the potential adverse impacts of GM crops on pollinators [7,8].

Crops expressing insecticidal proteins derived from the bacterium Bacillus thuringiensis (Bt [Bt-proteins]) are among the most widely cultivated GM crops worldwide [6]. A recent meta-analysis showed no adverse effects of Bt-crops on *A. mellifera* [7]. All of the re-analyzed studies tested only the effect of single Bt-proteins. However, one future trend in plant biotechnology is the stacking of multiple resistance traits. An example is Bt-maize SmartStax™, released in 2010 in the USA with six different insect resistance genes for above- and below-ground insect control, with two additional herbicide tolerance genes [6]. Hence, regulatory authorities are in need of up to date test-standards, to guide robust first-Tier laboratory experiments to assess the risks of new GM plants to non-target organisms [9].

Floral pollen is the sole protein source of *A. mellifera* colonies [10] and pollen of a variety of important crops is collected by bee foragers [8]. Adults and larvae of *A. mellifera* are directly exposed to transgenic material via pollen consumption of GM-crops, as planted in mass monocultures. On average, a worker consumes 3.4 to 4.3 mg of pollen per day [10], with colonies accumulating up to 55 kg per year [11]. Bees exposed to Mon810 maize pollen did not transmit quantifiable amounts of the Bt-proteins via their hypopharyngeal glands into the larval food they secrete [12]. Nevertheless, pollen is also straightforwardly added by nurse bees to the larval food [13]. It was reported that larvae consumed 1720–2310 maize pollen grains under semi-field exposure conditions, which is reflecting a worst case maize pollen exposure of 1.52–2.04 mg [14]. In comparison, European butterfly larvae fed with pollen grains from the transgenic maize variety Bt-176 were lethally affected at much lower exposure doses: LD$_{50}$ value of only 8 pollen grains per Diamond-back moth larva, and 32–39 pollen grains for Small tortoiseshells, Peacocks, European corn borers and Cabbage white larvae [15].

Bt-proteins confer plant-protection against herbivorous insects, with immature holometabolous pest insects showing a high susceptibility by a lethal damage to the gut [16]. This considering, especially young honey bee larva are amenable as non-target test organisms for GM crop pollen, because they represent a potentially sensitive life stage. In addition to larvae, young hive bees consume the most pollen within colonies [17], thus young bees are also amenable for precautionary tests on biosafety.
Nonetheless, Bt susceptibility in target insect adults is considered limited [18], in comparison to the lethal effects on larval stages [19,20]. To date, only minor fractions of peer-reviewed pollen feeding studies assess the risks on honey bee larvae [9]. Studies on Bt-pollen feeding to larvae have solely been performed within colonies [7,21]. In general, studies on the colony level are confounded by environmental influences and by nurse bees which remove the dietary treatments of the larvae. Thus, to be robust, laboratory bioassays need to exclude such uncontrolled factors as far as possible [9]. In this paper, to assess possible lethal and sublethal effects of GM crop pollen on the survival and prepupal weight of individual _A. mellifera_ larvae, we adopted a controlled _in vitro_ rearing bioassay [22,23]. The test larvae were exposed by adding fresh Bt-maize pollen directly in their artificial diet. This approach simulates the natural way of pollen consumption, whereby pollen is digested within the gut and Bt-proteins get exposed. Mechanistically, this is of key importance as the lethality among target-organisms is caused by the disruption of the gut epithelium by Bt-protein-receptor interactions [24]. This study fills an important gap in ERA’s on bees, as disruption of the gut epithelium by Bt-protein-receptor interactions [24].

**Materials and Methods**

**Pollen**

Multiple pollen types were collected for the _in vitro_ pollen feeding experiment (Table 1). Pollen of field grown maize varieties were collected by shaking flowering maize tassels in paper bags. The experiment (Table 1). Pollen of field grown maize varieties were lacking. 

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**Materials and Methods**

**Pollen**

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Pollen of the single transgenic Bt-maize event Mon810 (DKc7565, cultivar Novelis, Monsanto Co.) was collected on July 4th 2008 near Kitzingen (Lower Franconia, Germany). This maize variety expresses Cry1Ab proteins for the control of _Ostrinia nubilalis_ (Hubner) (Lepidoptera: Crambidae) at concentrations of 1–97 ng/g fresh weight (fwt) in pollen (Sauer and Jehle, pers. comm.). Cry1Ac at levels of mean 4.24 μg/g (n = 16, fwt in pollen, Sauer and Jehle, pers. comm.) and Cry2Ab2 at a level of mean 1.19 μg/g (n = 16, fwt in pollen, Sauer and Jehle, pers. comm.). Cry1A.105 is a chimeric gene synthesized by combining 4 native Bt-gene domains of cry1Ab, cry1F and cry1Ac [26]. This chimeric protein provides an increased activity against lepidopteran species compared to the original Cry1Ab protein as expressed in Mon810. The other parental line, Mon80017 (DKc3143), confers resistance to coleopteran pests, the Western, Northern and Mexican corn rootworms _Diabrotica spp._ (Coleoptera: Chrysomelidae) by the expression of the Bt-protein Cry3Bb1 at levels of mean 6.95 μg/g (n = 16, fwt in pollen, Sauer and Jehle, pers. comm.) (trademark YieldGard® Rootworm). Mon80017 also expresses an _Agrobacterium_ _sp_. CP4 derived 3-enolpyruvylshikimate-3-phosphate synthase (CPS) conferring tolerance against glyphosate, the active ingredient of the herbicide Roundup (trademark Roundup Ready®) at an expression level of 170 μg/g (fwt in pollen; www.gmo-compass.org/pdf/regulation/maize/MON89034xMON88017_application.pdf).

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Staked Bt-maize pollen and also control pollen of three conventional maize varieties was collected in the week of August 4th 2008 near Braunschweig (Germany). These maize varieties were grown on an experimental field in a randomized block-design with eight replications. Samples were collected from all 30×40 m subplots, pooling the pollen into one representative sample per variety. The non-GM variety DKc5340 (Monsanto Co.) is near-isogenic to the tested stacked Bt-maize variety, DKc4250 (Monsanto Co.) is more distantly related and Benicia (Pioneer Hi-Bred, Johnston, Iowa, USA) is totally unrelated to the stacked event (Table 1).

Table 1. Feeding treatments of _in vitro_ reared honey bee larvae for the Bt-pollen bioassay.

| Treatment | Plant variety | n Larvae | Colonies | Pollen/2 mg |
|-----------|---------------|----------|----------|-------------|
| 1 Transgenic maize | Stacked Bt; Mon89034 × Mon88017 | 20 | 5 | 1701 |
| 2 Transgenic maize | Single Bt; DKc7565 | 20 | 5 | 1750 |
| 3 Control maize | Near isogenic line; DKc5340 | 19 | 5 | 1784 |
| 4 Control maize | Distant related; DKc4250 | 20 | 5 | 1753 |
| 5 Control maize | Unrelated; Benicia | 20 | 5 | 1722 |
| 6 No pollen control | - | 12 | 6 | 0 |
| 7 Positive toxic control | _Heliconia rostrata_ (H) | 10 | 5 | 1600 |
| 1,2 Pooled Bt-maize | Transgenic maize (Bt) | 40 | 5 | 1726 |
| 3,4,5 Pooled control maize | Control maize (C) | 59 | 5 | 1753 |

*Treatment maize 1 expresses three Bt-proteins encoded by the genes cry1A.105, cry2Ab2 and cry3Bb1 from _Bacillus thuringiensis_ that confer resistance against certain lepidopteran and coleopteran pests and additionally expresses the _CP4 epsps_ gene for glyphosate-tolerance. Treatment maize 2 expresses a single lepidopteran specific Bt-toxin encoded by the gene cry1Ab. In addition, control treatments, tested plant varieties, number of larvae, colonies and counted pollen grains per 2 mg pollen treatment are indicated.

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In vitro larvae rearing and treatment applications

The rearing of larvae upon hatchment under laboratory conditions was performed following the protocols by Aupinel et al. and Hendriksma et al. [22,23]. Six donor honey bee colonies were selected from different Upper-Franconian apiaries, choosing naturally mated non-sibling queens (Apis mellifera carnica). By means of an excluder lid, the queens were trapped within their colonies on artificial combs (Nicoplast®) (day 1; D1, 25th June 2009). After 91 hours, without grafting manipulation, larvae within plastic queen cups were collected from the combs. A 72 hours development time of the embryos until the hatchment of eggs, the larvae had a mean chronological age of 9:30 hours (D4; min. 0 to max. 19 hours old) and were typically first instars [25].

The subsequent laboratory rearing was performed with larvae in queen-cups mounted in culture plates, placed in a hermetic plexiglass desiccator within an incubator at 35 °Celsius. The larvae were fed once a day over D4 to D9 with a 10-, 10-, 20-, 30-, 40-, microbalance to measure the weight to the nearest 0.001 g. with soft metal tweezers into a new clean cell on an analytical balance. Prepupae were transplanted only measured after defecation. Every prepupa was transplanted are physically terminated. Hence, the effective gain in weight can be assessed possible lethal effects of Bt-maize pollen during the exposure and the potential Bt-protein-receptor based mechanism

larvae defecate and molt their intestine at this stage, both the remaining total exposure time, until the diet was completely finished at the non-feeding days D10/D11. The maize pollen were consumed over 2011 [23].

The identity of the replicate donor colonies was included as a random factor in the models to take the non-independence of larvae from individual colonies into account [23].

Prepupae weights were analyzed with linear mixed effects models using the package nlme [31]. The survival dynamics of larvae were analyzed with Cox proportional hazards regression models [32] using the R packages survival and survnet [33,34]. A dynamic survival analysis is not applicable when all individuals of a group survive; in that case a Chi-square analysis was used.

Three test levels were considered. An overall sort-effect was tested over the five maize varieties. All treatments were also tested individually, paired to one another, to indicate sort effects. The significance of P values (α = 0.05) of multiple comparison were determined with an α-correction using the sequential Holm-Bonferroni procedure [35]. In case of no detectable difference, the treatment comparisons were summarized by evaluating the pooled data on Bt-maize pollen with control maize pollen data, also pooled.

Results

Survival

All 40 larvae fed with B-maize pollen survived the 120 hours of dietary exposure upon the prepupal phase (Fig. 1). The survival rate of the conventional maize pollen fed larvae did not differ significantly from Bt-maize pollen fed larvae {C: 56 out of 59: 95%}, (Chisq = 0.72, df = 1, P = 0.40). Of all the maize pollen fed larvae (N = 99), in total 97% survived until the prepupal phase. Specific survival rates were: for stacked Bt-maize 100%, near-isogenic line 100%, Mon810 100%, DKe1250 95%, and for Benicia 90%. Thus, no significant difference among the five maize pollen varieties was found (Chisq = 5.41, df = 4, P = 0.28).

![Figure 1. Survival analysis of honey bee larvae treated with pollen enriched diets.](image)

The dashed curve “Bt” indicates the 100% survival rate for Bt-pollen treated larvae (stacked Bt-maize expressing Cry1A.105, Cry2Ab2 and Cry3Bb1 and single Bt-maize expressing Cry1A.B were pooled; n = 40 larvae). Curve “C” indicates survival for three conventional (control) maize pollen treatments (pooled n = 59 larvae). No significant differences in survival rates were found among maize pollen treatments (neither individually, nor pooled). Compared to the other treatments, the larvae fed with the toxic Heliconia rostrata pollen (H; n = 10) had a significantly lowered survival rate. doi:10.1371/journal.pone.0028174.g001

Statistics

The data were analyzed with mixed models using different packages of the open source statistic software R version 2.11.1 [30]. The identity of the replicate donor colonies was included as a

In vitro Honey Bee Larvae Assay for GM Crop Pollen

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Effects of pollen from single and multiple Bt-maize varieties on honey bee larvae

Among the larvae fed with diets without pollen, the individual survival dynamics and the survival rate of 92% did not differ compared to larvae fed with maize pollen enriched diets (all P values≤0.64). In contrast, significantly fewer larvae survived the larval phase when they were fed with *Heliconia rostrata* pollen compared to the other six treatments (P values≤0.01, all significant with an α/6 sequential Holm-Bonferroni correction) (Fig. 1).

Sublethal effects on the prepupae weight

With a mean of 142.3 mg, prepupae weights of Bt-maize pollen fed larvae were almost identical to the mean weight of conventional maize pollen fed larvae (142.6 mg; t = −0.20, df = 1, P = 0.82) (Table 2). A general variety-effect, considering possible differences between the five maize varieties, was not found (F = 0.26, df = 4, P = 0.90) thus the weight distributions of the transgenic and non-transgenic maize pollen treatments were all alike (Fig. 2).

Individual comparison shows that mean prepupae weights differed neither between stacked Bt-pollen and pollen from the near-isogenic line (t = 0.83, df = 33, P = 0.41), nor between the stacked Bt-variety and the single Bt-variety (t = 0.81, df = 34, P = 0.42) (Table 2). In contrast, *H. rostrata* pollen fed larvae showed a significantly lower mean prepupae weight compared to all the other treatments (mean 87.7 mg±21.0 SD; P values≤0.001) (Table 2).

Discussion

Honey bees are the most important pollinators in agricultural ecosystems. In order to minimize the environmental risks of cultivating GM crops and their discussed contribution of being an underlying factor of the globally observed bee losses, robust and highly standardized risk assessment methods for honey bees are imperative. Here we present an effective pollen based method to test the direct effects of GM crops on *in vitro* reared larvae. Our test system reflects the natural exposure under field conditions and is therefore highly recommended for regulatory studies.

Effects of pollen from single and multiple Bt-maize varieties on honey bee larvae

One recent trend in plant biotechnology is stacking of multiple insect resistance traits in a single cultivar [6]. Honey bees are exposed to mass flowering GM crops and not a single published study deals with the effect of stacked Bt-cultivars on bees. The results of this study did not indicate adverse effects of the consumption of single and stacked Bt-maize pollen on the survival and prepupae weight of *in vitro* reared *A. mellifera* larvae. At a realistic exposure dose, the 120 h survivorship of Bt-pollen treated larvae was 100% until the prepupae phase (Fig. 1). At the prepupae stage, where larvae had terminated feeding, digesting...
and growing, were no indications of a sublethal Bt-pollen effect on the weight of the prepupae (Fig. 2).

The outcome of our data on stacked Bt effects are in line with earlier brood tests under colony conditions on single insect resistant Bt-maize pollen [36] or single purified Bt-proteins [7,37]. In contrast to these colony level studies, the current results are achieved by testing under controlled laboratory conditions, with minimum control mortality. Compared to single Bt-proteins in pollen or in purified form, our plant produced stacked Bt-proteins, with the chimeric B-t protein Cry1A.105, indicate a similar level of safety. In accordance, a stacked maize variety, expressing Bt proteins VIP3A and Cry1Ab, also caused no adverse effects on the biodiversity of arthropods during a 3 year ERA field experiment [38]. A stacked cotton cultivar, expressing cowpea trypsin inhibitor (CpTI) and Cry1Ac in pollen carried no lethal risk for honey bees, though a worst case feeding regime did cause feeding inhibition [39]. However, in studies comparing Cry1 with transgenic protease inhibitors, it was found that only the latter was causing reduced feeding effects [12,40,41,42].

The stacking of insect resistance traits in one crop aims to enhance the effectiveness towards target pest insects, to cause an additive or synergistic toxicity. Among target pest insects, synergistic effects between e.g. Cry1Ab, Cry1Ac, Cry1F and/or Cry2Ab2 have been reported [43,44]. Involved in toxicant synergies are mostly uptake, transportation or degradation pathways [45], causing a higher toxicity and a lower selectivity. Hence, potential synergistic effects on non-target insect also deserve consideration. The honey bee, a key non-target insect, has never shown lethality to Bt-proteins [7] and our data support the notion that, synergistic effects by stacking Bt-proteins at plant produced levels are unlikely a risk to bees. However, sublethal effects [46] on feeding, learning performance and foraging behavior might occur [47,48]. Indeed, the in vitro approach covers the opportunity of testing of potential sublethal effects, by a subsequent behavioral tests on hatched bees [49].

In order to examine a potential effect of increased protein expression levels, two Bt-maize varieties with different expression levels were compared. Bt-maize variety Mon89034×Mon88017 has compared to Mon810 a 10² to 10⁴ times increased Bt-protein expression level in pollen (see material and methods). Hypothetically, Mon810 could have had Bt-protein levels under a toxic threshold, but the larvae remain unharmed by the multifold Bt-protein of stacked Bt-maize pollen.

Pollen bioassays

The current bioassay tests GM plant material directly and realistically, by reflecting a natural consumption and digestion of pollen by A. mellifera larvae. It closes an important knowledge gap between in vivo colony experiments [7,14,36,37,38] and in vitro experiments with purified transgenic proteins [50,51,52]. Although purified proteins are ideal to test worst case exposure scenarios [9,48], the E. coli produced purified Bt-substances do not represent a field situation. And although field experiments have realistic pollen exposure conditions, a down-side is a variety of uncontrolled environmental factors. In addition, pollinator field studies have to be synchronized to the flowering period and they are space and time consuming and therefore relatively costly [53]. Finally, within a bee colony many factors such as colony size, diseases, and nutrition could have an influence on the brood development. The presented robust bioassay minimizes any environmental effect on larval development and allows a good control of dietary pollen amounts (Table 1).

The conventional non GM maize cultivars (Table 1) allow a secure assessment of the impact of the introduced transgenic traits [54]. It makes assessments comprehensive, since it enables a reliable estimate of naturally occurring variation within the crop species. Though having tested the total of five maize varieties, no maize-sort related differences were found. Nevertheless, the toxic control treatment and the power analysis indicated that monitoring discernible effects of pollen on honey bee larvae was effective (Fig. S1, Fig. S2, Power analysis S1). At the given sample sizes, the test was able to distinguish effects on both the survival and the weight endpoint.

The functionality of the pollen bioassay is proven by the feeding dose of 1600 H. rostrata pollen, which caused significant lethal and sublethal effects on larvae. This dose caused 50% of the larvae to die in 72 hours (LT50) and 100% to die in 7 days (LT100). The results demonstrate the usefulness of positive controls in order to i) validate the ingestion of pollen treatments, ii) to demonstrate the capacity of detecting treatment effects and iii) to allow comparisons with other studies [9].

Precise and robust ERA methods are needed for honey bees [55]. Our bioassay is well suited to monitor environmental pollution of pollen or natural pollen toxicity (Pictures S1). Of genuine concern are systemic, lipophilic chemicals (e.g. neonicotinoids) as used in agriculture, because the plant pollen are a carrier of pesticides into honeybee colonies. Such pesticides may cause (sub-)lethal effects and can be extremely persistent [46]. Our pollen test is widely applicable and it fits international tiered risk assessment schemes for regulatory biosafety assessments of any new transgenic trait. Hence, we propose the in vitro bioassay for consideration as a standard pre-release test for all polleniferous transgenic crops.

Supporting Information

Figure S1 Statistical power analysis for survival data of honey bee larvae on maize pollen enriched diets. The survival power analysis was based on a one-tailed 2-proportions test on mortality rate differences, comparing a control and a treatment group with a same sample size. Determining treatment effects more sensitively at higher sample sizes, the curves indicate the level of power with dotted lines for 0.4, striped lines for 0.6 and a continuous line for 0.8 power at analysis (significance level of α = 0.05). (Power analysis S1). (TIF)

Figure S2 Statistical power analysis for prepupae weight data of honey bee larvae on maize pollen enriched diets. The weight difference power analysis was based on a two-tailed t-test on weight differences between the treatment group and the control (with same sample sizes). The sensitivity to measure the mg weight differences is relating to the general variance in weight of all maize pollen fed larvae (142 mg±8.5 SD, n = 96). The significance level of α = 0.05 at 0.4, 0.6 and 0.8 power determined which sample sizes were needed to indicate effects. (Power analysis S1). (TIF)

Pictures S1 A honey bee larvae in vitro bioassay for testing pollen toxicity, considering GM-maize pollen (Zea mays) and pollen of Heliconia rostrata. Pictures by Harmen P. Hendriksma (legends are embedded in the pictures). (PDF)

Power Analysis S1 An analysis of statistical power to indicate mortality and weight differences within the experimental data. Supplementary information in addition to Figure S1 and Figure S2 on power analysis. (DOC)
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Author Contributions

Conceived and designed the experiments: HPH SH. Performed the experiments: HPH SH. Analyzed the data: HPH. Contributed reagents/materials/analysis tools: ISD. Wrote the paper: HPH SH ISD.

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