Reduced caterpillar damage can benefit plant bugs in Bt cotton

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Bt cotton was genetically modified to produce insecticidal proteins targeting Lepidopteran pests and is therefore only minimally affected by caterpillar damage. This could lead to reduced levels of inherent, systemically inducible defensive compounds in Bt cotton which might benefit other important cotton herbivores such as plant bugs. We studied the effects of plant defense induction on the performance of the plant bug *Lygus hesperus* by caging nymphs on different food sources (bolls/squares) of Bt and non-Bt cotton which were either undamaged, damaged by Bt tolerant caterpillars, or treated with jasmonic acid (JA). Terpenoid induction patterns of JA-treated and *L. hesperus*-damaged plants were characterized for different plant structures and artificial diet assays using purified terpenoids (gossypol/heliocide H1/4) were conducted. Nymphs were negatively affected if kept on plants damaged by caterpillars or sprayed with JA. Performance of nymphs was increased if they fed on squares and by the Bt-trait which had a positive effect on boll quality as food. In general, JA-sprayed plants (but not *L. hesperus* infested plants) showed increased levels of terpenoids in the plant structures analyzed, which was especially pronounced in Bt cotton. Nymphs were not negatively affected by terpenoids in artificial diet assays indicating that other inducible cotton responses are responsible for the found negative effects on *L. hesperus*. Overall, genetically engineered plant defenses can benefit plant bugs by releasing them from plant-mediated indirect competition with lepidopterans which might contribute to increasing numbers of hemipterans in Bt cotton.

The cultivation of insect-resistant genetically engineered crops producing Cry proteins from *Bacillus thuringiensis* (Bt crops) helps to control a range of key lepidopteran and coleopteran pest species while reducing the amount of chemical insecticide applications1–2. The area-wide use of Bt-transgenic crops, mainly maize (*Zea mays*) and cotton (*Gossypium hirsutum*), has led to significant population declines in target pests3–6. However, increasing numbers of herbivorous pests not targeted by the Cry proteins have been reported from Bt crops. This is the case in particular for Bt cotton where sucking bugs, such as plant bugs (Heteroptera: Miridae) and stink bugs (Heteroptera: Pentatomidae), have become problematic pests in some cropping systems7–14. Increased issues with non-target pests in maize, on the other hand, are mainly limited to the western bean cutworm (*Striacosta albicosta*) (Lepidoptera: Noctuidae) 12,13. The reasons causing the increase in *S. albicosta* numbers are not fully understood but may include a variety of factors, such as reduced direct competition with herbivores targeted by the Bt trait, a reduction in insecticide use, as well as other ecological, agronomic and climatic causes8. In the case of Bt-transgenic cotton, several factors predisposing non-target pest to become more problematic have been suggested. Increases of sucking bugs in Bt cotton can mainly be attributed to a reduction in broad-spectrum insecticide applications as many insecticides against pest Lepidoptera also incidentally control other herbivore species7,14. In addition, there is mounting evidence that the strong reduction of lepidopteran populations in Bt cotton and, associated therewith, altered interspecific interactions among species also benefits non-target herbivores15. Stink bugs as well as cotton aphids, *Aphis gossypii* (Hemiptera: Aphididae) can benefit from the release of either direct interference competition or plant-mediated indirect competition with Bt-sensitive Lepidoptera16–18. There is evidence that plant-mediated indirect competition in cotton is partly driven by inducible defensive compounds. Best studied is a set of biosynthetically related non-volatile terpenoids (e.g. gossypol, heliocides, hemigossypolone) that are stored in subepidermal pigment glands20,21. These terpenoids are systemically induced in response to plant damage by tissue feeders22–24. They provide resistance primarily against lepidopterans, but may also be toxic to a range of other herbivores25. Reduced caterpillar damage on Bt cotton lowers the levels of...
inducible cotton defensive compounds, which in turn might improve the performance of non-target herbivores, as has been demonstrated for *A. gossypii* [17]. Plant-mediated indirect competition accounts for a major part of all interspecific herbivore interactions in natural ecosystems and can affect whole arthropod communities [25–27]. Thus, a release from indirect competition with caterpillars could also contribute to increasing numbers of sucking bugs in Bt cotton.

The western tarnished plant bug, *Lygus hesperus* (Hemiptera: Miridae) is a key herbivore in Bt cotton in the southwestern United States [11,28]. *L. hesperus* attacks mainly young cotton flower buds (squares), young bolls and growing points, where it feeds on enzymatically liquefied plant tissue [29,30]. This often leads to localized tissue necrosis and abortion of the attacked structure [29]. Literature documenting the role of cotton terpenoids as a defense mechanism against *L. hesperus* is limited. However, Tingey *et al.* [31] and Ellington *et al.* [32] showed that cotton varieties with low densities of gossypol-containing pigment glands positively affect *Lygus* spp. performance and population size.

Using *L. hesperus* as a model species we hypothesized that plant bugs benefit from reduced caterpillar damage in Bt cotton as they might profit from reduced levels of caterpillar-induced cotton defenses. This hypothesis was tested in the greenhouse, were we studied *L. hesperus* performance on Bt and non-Bt cotton subjected to different induction treatments. In further greenhouse and laboratory experiments we elucidated induction patterns of defensive cotton terpenoids in different plant structures that *L. hesperus* feeds on and studied their potential as explanatory factors affecting *L. hesperus* performance.

### Results

**Experiment 1: Effect of defense induction, food source and plant type on *L. hesperus* performance.**

On average, mortality of *L. hesperus* on Bt and non-Bt plants was 25% and 20% higher when nymphs were kept on caterpillar- and JA-induced plants, respectively, in comparison to control plants (Fig. 1). Likewise, 30% and 10% fewer nymphs developed into adults on caterpillar- and JA-induced plants compared to controls, respectively (Fig. 1). The number of nymphs that died during the experiment and the number of nymphs that developed to adults (development rate) (Table 1) was significantly reduced on plants damaged by *Spodoptera exigua* (Lepidoptera: Noctuidae) caterpillars when compared to undamaged control plants. Similarly, plants sprayed with JA had a significant negative impact on nymph survival whereas the development rate was reduced, albeit not significantly, by JA (Table 1). Mortality and development rate was not only affected by damage treatments but was strongly dependent on food sources (Fig. 1). Feeding on squares had a strong positive effect on nymph survival whereas the development rate was reduced, albeit not significantly, by JA (Table 1). Mortality and development rate was not only affected by damage treatments but was strongly dependent on food sources (Fig. 1). Feeding on squares had a strong positive effect on nymph survival whereas the development rate was reduced, albeit not significantly, by JA (Table 1). Mortality and development rate was not only affected by damage treatments but was strongly dependent on food sources (Fig. 1). Feeding on squares had a strong positive effect on nymph survival whereas the development rate was reduced, albeit not significantly, by JA (Table 1). Mortality and development rate was not only affected by damage treatments but was strongly dependent on food sources (Fig. 1). Feeding on squares had a strong positive effect on nymph survival whereas the development rate was reduced, albeit not significantly, by JA (Table 1). 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In subsequent analyses, neither terpenoid concentrations nor C:N ratios measured in youngest leaves of experimental plants were significantly correlated with *L. hesperus* mortality or development success (all variables $p > 0.05$).

Weight gain of adult *L. hesperus* (max. 24 h old) was significantly negatively affected by bolls as a food source when compared with squares (estimate $= -0.49 \pm 0.18$, $z = 2.73$, $p = 0.006$) whereas plant type (Bt-trait) and induction treatments had no significant effect on *L. hesperus* weight (all variables $p > 0.05$).
Experiment 2: Cotton defense induction in fruiting structures and leaves. Terpenoid production in the youngest leaves, and to a lesser degree, also in squares, and the boll endo- and mesocarp was significantly affected by induction treatment and plant type (Table 2, Fig. 3). Post-hoc tests (not shown) revealed that the induction treatment effects can be attributed to JA. In leaves and squares, JA-sprayed plants had significantly higher terpenoid levels than controls or *L. hesperus*-damaged plants. In the boll endo- and mesocarp, terpenoid levels of JA-sprayed plants were significantly higher than levels of *L. hesperus*-damaged plants whereas control plants showed intermediate levels. Furthermore, Bt plants showed on average higher terpenoid levels in leaves, squares and the boll endo- and mesocarp than non-Bt plants (Table 2). The boll exocarp showed no signs of induction by any treatment. Levels of different terpenoids varied strongly among different plant structures (Fig. 3). JA-induced levels of gossypol were about 4 times higher in squares when compared to leaves, 5 times higher than in the boll exocarp and 500 times higher when compared to boll endo- and mesocarp. In contrast, induced levels of hemigossypolone in young leaves were 7–10 times higher when compared to square levels and 3–550 times higher than in boll structures. JA-induced as well as constitutive heliocide H1/H4 levels were highest in the boll exocarp followed by the JA-induced levels measured in leaves. Heliocide H1/H4 levels measured in squares and boll meso/endocarp were 1–2 orders of magnitude lower compared with levels measured in leaves or the exocarp.

**Table 1.** Parameter estimates of the averaged candidate models explaining factors affecting survival or successful development of 3rd instar *Lygus hesperus* nymphs into adults. Nymphs were kept for 14 days on squares, bolls or both of Bt or non-Bt-cotton plants that had been induced by *Spodoptera exigua* caterpillars, treated with jasmonic acid (JA), or remained uninduced.

**Figure 2.** Percentage of *Lygus hesperus* nymphs (n = 341) that successfully developed into adults during the experiment. Nymphs were kept on Bt (n = 161) or non-Bt (n = 180) cotton (*Gossypium hirsutum*) and either fed on squares during their development (solid line) or exclusively fed on bolls (dashed line).

| Survival Parameters | Estimate | SE | z value | p value |
|---------------------|----------|----|---------|---------|
| Induction treatment: |          |    |         |         |
| Caterpillar         | −1.03    | 0.37| 2.79    | 0.005   |
| JA                  | −0.71    | 0.35| 2.01    | 0.04    |
| Food source: Square | 1.29     | 0.32| 4.10    | < 0.001 |

Variables with no relevant explanatory power: Plant type, terpenoids, C:N ratio

| Development rate Parameters | Estimate | SE | z value | p value |
|-----------------------------|----------|----|---------|---------|
| Induction treatment:        |          |    |         |         |
| Caterpillar                 | −1.31    | 0.42| 3.11    | 0.002   |
| JA                          | −0.68    | 0.41| 1.66    | 0.10    |
| Plant type: Bt              | 1.54     | 0.70| 2.20    | 0.03    |
| Food source: Square         | 2.75     | 0.71| 4.89    | < 0.001 |
| Interactions: Square × Bt   | −2.13    | 0.76| 2.78    | 0.005   |

Variables with no relevant explanatory power: Terpenoids, C:N ratio
**Experiment 3: Effect of terpenoids on *L. hesperus* performance.** Different diet treatments had no significant effect on the total number of days that *L. hesperus* nymphs needed to develop into adults (7.97 ± 0.07 days) or the number of days they spent in the third, fourth, or fifth instar (Kruskal-Wallis; df = 3, all p > 0.18). Likewise, neither survival (Chi-square = 2.18, df = 3, p = 0.54) nor weight gain (fw = 7.68 ± 0.15 mg) was affected by different diets (ANOVA; F3, 62 = 0.137, p = 0.94).

**Discussion**

The area-wide use of Bt-crops has significantly affected species composition in agro-ecosystems. In Bt-cotton, hemipterans such as plant and stink bugs have in many cases increased in relevance as pests. While this is mainly due to the fact that the use of broad-spectrum insecticides is reduced in Bt cotton, our study indicates that a release from plant-mediated indirect competition with caterpillars could have additionally contributed to the increasing numbers of plant bugs in some Bt cotton systems.
Impact of cotton defense induction and cotton terpenoids on *L. hesperus* performance.  We found that cotton defense induction by *S. exigua* caterpillars had a negative effect on *L. hesperus* survival and development rate (but not weight gain) (Table 1). Similarly, plants sprayed with JA negatively affected *L. hesperus* performance (Table 1). This can most likely be attributed to JA-mediated cotton defense induction. However, we cannot rule out that also EtOH, which was used to dissolve JA, negatively affected *L. hesperus* performance since we did not control for any potential negative effects of EtOH. However, the low concentration of EtOH applied to the plants (100 μl/60 ml), the high volatility of EtOH as well as the finding that artificial diet containing EtOH did not negatively affect *L. hesperus* nymphs (experiment 3) speak against a negative impact of EtOH on *L. hesperus* in our experiment.

Plant-mediated interactions among herbivores, which are linked to alterations in plant defense levels, are well documented for a range of plant-herbivore systems. In agreement with our results, previous studies reported that caterpillar-damaged cotton can adversely affect hemipteran species, i.e. cotton aphids, *A. gossypii* and the stink bug, *N. viridula* and *E. servus* (both: Hemiptera: Pentatomidae) [15,30]. There is evidence to suggest that these effects are caused by the induction of defensive terpenoids as a response to caterpillar damage or treatment with JA [25,28]. This is also supported by studies documenting that *L. hesperus* nymphs have higher survival rates and occur in higher densities on glandless cotton varieties with comparatively low levels of terpenoids [31,32].

However, when we exposed *L. hesperus* nymphs to pure gossypol or heliocide H1/H4 mixed into artificial diet, we observed no negative effects on their performance. In the case of heliocides H1/H4, the results need to be interpreted with caution since the purity of the compound used in our study was only ca. 50% and the concentration was about two orders of magnitude lower than concentrations found in squares or leaves of induced cotton plants. We can thus not rule out that heliocide H1/H4 in higher concentrations and purity might affect *L. hesperus*. In contrast, the purity of the gossypol used was high and concentrations in the artificial diet study were on the upper limit of concentrations measured in induced cotton leaves. Gossypol is known to be light-sensitive.

In order to minimize gossypol degradation in our study, terpenoid containing diet packs were exchanged every 4–5 days. Other artificial diet studies using similar concentrations of gossypol, showed that it remains bioactive during this period of time [34,39]. That gossypol has no adverse effect on *L. hesperus* is also supported by the results of our greenhouse experiment. Squares as a food source had a positive effect on *L. hesperus* performance when compared with bolls despite the fact that squares contained much higher concentrations of gossypol (Fig. 3). That cotton terpenoids might not be responsible for plant-mediated indirect competition between caterpillars and sucking bugs was also suggested by Zelinger et al. [19]. They found that the boll-feeding stink bug *E. servus* avoided cotton plants damaged by caterpillars of *Helioverpa zea* (Boddie) (Lepidoptera: Noctuidae) while it was attracted to plants damaged by *Heliotris virens* Fabricius (Lepidoptera: Noctuidae). This, despite the fact that the concentration of terpenoids was significantly greater in seeds of *H. virens*-damaged plants compared to *H. zea*-damaged plants.

Given the large array of different cotton defenses against caterpillars, it is most likely that other potentially inducible defense mechanisms, such as chlorogenic acid, condensed tannins, or other phenolic compounds might explain the negative effects of cotton induction on *L. hesperus* performance [15,21,40]. Although C:N ratios in plants had no effect on *L. hesperus* performance, we cannot rule out that other changes in cottons nutritional quality affected *L. hesperus* as it has been reported that caterpillar damage can affect amino acid composition, water content or the oxidative status in cotton [20,21].

Cotton terpenoid induction in different plant structures. Terpenoid production in the boll exocarp was not inducible by any treatment, but JA-sprayed plants showed higher terpenoid levels in leaves and squares and to a lesser degree also in the boll endo/mesocarp. In contrast, *L. hesperus* feeding led to no terpenoid induction in any of the plant structures, indicating that nymphs did not trigger the induction of terpenoids in cotton. While chewing herbivores generally induce JA-related defenses it has been found that many sucking hemipterans like aphids or whiteflies can bypass such plant defense responses [42,43]. However, to what degree *L. hesperus* can manipulate the array of other cotton defenses is not well understood. Studies by Rodriguez-Saona et al. [14] and Williams et al. [45] show that *L. hesperus* feeding can induce volatiles comparable to caterpillar-induced volatile blends which might allow the plant to respond to *L. hesperus* infestations by, for example, attracting natural enemies.

Impact of food source and plant type on *L. hesperus* performance. *L. hesperus* mortality and development, as well as weight gain, was strongly affected by food source (Table 1, Fig. 1). Feeding on squares had a positive effect on *L. hesperus* development and survival compared with individuals that had only access to bolls. Furthermore, adults that developed only on bolls where significantly lighter than adults that had fed on squares. Likewise, Chen and Parajulee [46] showed that adults that developed only on bolls where significantly lighter than adults that had fed on squares. Furthermore, adults that developed only on bolls where significantly lighter than adults that had fed on squares. In the field, however, nymphs are less restricted in their food choice and likely prefer to feed on squares over bolls (Table 1, Fig. 2). Therefore, when compared with our results from the greenhouse, the food source parameter is probably less relevant in affecting *L. hesperus* performance under field conditions.
Herbivory can have profound effects on a host plants’ inducible responses which may entail a higher plant resistance to subsequent attacks by conspecifics or other herbivores. We demonstrated that the absence of such plant-mediated competition between herbivores can be an important additional factor contributing to the increased populations of plant bugs in Bt cotton in some regions. While the observed effects seem to be related to one of the many inducible mechanisms in cotton, it appears that the defensive terpenoid gossypol does not play a role in the cotton–L. hesperus interaction. Besides plant induction, L. hesperus performance was also affected by the Bt-trait and the food source. The latter might, however, be less relevant in the field where plant bugs can freely choose between different plant structures as food sources.

Materials and Methods

Plants and Insects. Commercial cotton plants (G. hirsutum), i.e., Bt cotton (Bollgard II Roundup Ready Flex cotton, DP1359B2RF; event MON15985 × MON1445, Monsanto, St. Louis, USA) and the genetically closest non-Bt cotton cultivar (Sure grow 125, Monsanto, St. Louis, USA) were individually grown under greenhouse conditions (25 ± 4°C, av. 30% RH) in 3.8 l plastic pots containing a soil-sand mixture (9:5). Plants were watered daily and fertilized weekly using 100 ml of a 20% N, 20% P2O5, 20% K2O at 1 g l–1 (= 200 PPM N) Nutricleur General Purpose soluble fertilizer solution (Plant Marvel Laboratories, Chicago, USA). For all experiments 7–8 week old plants were used that possessed squares (flower buds) and young bolls (approx. 1 cm diameter). L. hespe- rus was reared under laboratory conditions (av. 27°C, 30% RH, 14:10 LD cycle). The first two instars were reared on green beans. Later instars were kept in the colony was founded from collections in alfalfa and cotton fields in Maricopa, AZ USA in 2013. Bt-tolerant, fourth instar S. exigua were obtained from Frontier Agricultural Sciences (http://www.insecttreated.com).

Experiment 1: Effect of cotton defense induction on L. hesperus performance. The aim of this experiment was to study the effect of cotton defense induction on L. hesperus performance. Bt and non-Bt cotton plants were exposed to one of the following induction treatments in the greenhouse: (i) plants were exposed to four instar S. exigua. A total of six larvae per plant were individually caged on single leaves equally distributed among top, medium, and lower node regions using organdy cloth bags with Velcro that fastened around the pet- irole. Larvae were transferred to new leaves every 2–3 days. After 1 week all larvae and bags were removed; (ii) plants were induced with the plant hormone jasmonic acid (JA) (Sigma-Aldrich, St. Louis, USA), which is known to induce cotton defense responses. Each plant was sprayed with a solution containing 60 ml distilled water and 5 mg of JA dissolved in 100 µl EtOH using a vaporizer; (iii) plants remained untreated (control). This resulted in a total of six treatments (three induction treatments each for Bt and non-Bt plants). A total of 20–24 plants were subjected to each treatment. One week after application of the induction treatments, three freshly molted and weighed third instar L. hesperus nymphs were caged in individual organdy cloth bags with a Velcro fastener on each plant (experimental unit). Nymphs were randomly caged on branchlets with either a single young boll or a square as food source. Preliminary feeding studies using artificial diet indicated that most third instar nymphs successfully developed into adults within 7–8 days (not shown). To make sure that the nymphs had enough time to develop into adults under greenhouse conditions they were kept on plants for a maximum of 14 days. After an initial period of five days, nymphs were checked daily during the experiment. Individuals that developed into adults were removed from plants and weighed again to calculate the weight gain.

L. hesperus nymphs were transferred to new squares or young bolls of the same plant if the plants aborted these structures due to L. hesperus feeding damage or if young bolls or squares developed into older bolls or flowers. Therefore, nymphs fed either on squares, only on bolls or on both structures during the experiment. Mortality and the number of nymphs that survived but did not manage to develop into adults was recorded. The youngest leaf of each plant in all treatments was sampled and stored at −80°C for further biochemical analyses (see measured terpenoids and plant nutrients) at the end of the experiment.

Experiment 2: Cotton defense induction in different plant structures. To study to what degree JA and L. hesperus feeding induces defense responses in different cotton plant structures and if these induction patterns are in agreement with the results from experiment 1, Bt and non-Bt cotton plants were exposed to one of the following induction treatments in the greenhouse: (i) three freshly molted third instar L. hesperus nymphs were randomly caged in organdy cloth bags with a Velcro fastener on young bolls or squares of each plant for 7 days. Nymphs were transferred to new squares or bolls of the same plant if cotton plants dropped these structures or if young bolls or squares developed into old bolls or flowers. Infested plants where more than one nymph died during the experiment were discarded; (ii) plants were induced with JA (positive control, described above); (iii) plants remained untreated (negative control). This resulted in a total of six treatments (three induction treatments each for Bt and non-Bt plants). A total of 10–14 individual plants were subjected to each treatment. After 7 days, all nymphs were removed and a young, not yet fully expanded leaf from the top of each plant was collected. A random square and both the exocarp and the locule (endo-and mesocarp) of a random young boll that was not previously infested (approx. 1 cm diameter) were sampled if available on the plant, and stored at −80°C for further biochemical analyses.

Experiment 3: Effect of terpenoids on L. hesperus performance. An artificial diet study was conducted with freshly molted third instar nymphs to assess the sensitivity of L. hesperus to two terpenoids, gossypol and heliocide H1/H4. Nymphs were fed with artificial diet described by Debolt, which was spiked with either: (i) 2 mg gossypol dissolved in 4.5 µl EtOH (95%)/g diet; (ii) artificial diet containing 0.9 mg heliocide H1/H4 dissolved in 4.5 µl EtOH (95%)/g diet; (iii) artificial diet containing just the EtOH solvent (4.5 µl EtOH (95%)/g
et al. 17. Gossypol was identified by comparing selection58. For each response variable, models were fitted with all possible combinations of the explanatory variables plant type (Bt and non-Bt cotton), induction treatment (caterpillar damage, JA treated, and control), plant type (Bt and non-Bt cotton) and food source (squares and bolls) as explanatory variables. In a second step, the same response variables were analyzed using a GLMM with terpenoids (gossypol, heliocide H1/ H4, hemigossypolone) and C:N ratios in the youngest leaves of the experimental plants as explanatory variables.

The two response variables, nymph weight gain of individuals that developed into adults was analyzed using a linear mixed model (LMM) (“lmer” function of the lme4 package, version 1.1–12)57 with induction treatment and plant type (see above) as explanatory variables. In all models individual plants were used as random effects.

Factors affecting L. hesperus performance were identified with an information theoretic framework of model selection59. For each response model, models were fitted with all possible combinations of the explanatory variables as well as their 2-way interactions using the “dredge” function of the R-package MuMln (version 1.15.6)39. Models were then ranked according to Akaike information criterion corrected for finite sample sizes (AICc).

To determine the explanatory variables that best explained variation in L. hesperus performance, all models were selected that conformed to two rules: First, only models with a Δ AICc value of ≤6 were selected, i.e., all models whose AICc value was at most 6 higher than the lowest AICc obtained. Second, a model was only selected if its AICc value was at most 6 higher than the lowest AICc obtained. The identity of the terpenoids was furthermore confirmed by mass spectrometry. We were unable to confirm the identity of a distinct peak assigned to heliocide H2/H3 with massspectrometrical analyses. We therefore did not include heliocide H2/H3 in this study. Terpenoid concentrations were quantified in terms of gossypol equivalents44.

To quantify C:N ratios from leaves of greenhouse experiment 1, 4–5 mg of lyophilized tissue from each leaf was individually filled in tin capsules and the C:N ratios for each sample was subsequently measured by elemental analysis (Hekatech Euro EA 3000, Wegberg Germany) as described in Leifeld et al.39.

Statistical analyses. For all statistical analyses, the Software R (version 3.2.3) was used56. The standard error of the mean is provided for all mean values (mean ± SE).

Experiment 1: Effect of cotton defense induction on L. hesperus performance. The two response variables, nymphs that died within 14 days (binomial distribution) and nymphs that developed within 14 days into adults (binomial distribution), were analyzed using generalized linear mixed models (GLMM) (“glm” function of the R-package lme4, version 1.1–12)57 with induction treatments (caterpillar damage, JA treated, and control), plant type (Bt and non-Bt cotton) and food source (squares and bolls) as explanatory variables. In a second step, the same response variables were analyzed using a GLMM with terpenoids (gossypol, heliocide H1/ H4, hemigossypolone) and C:N ratios in the youngest leaves of the experimental plants as explanatory variables.

The two response variables were analyzed using separate two-way ANOVAs using the “lm” function. In the case of significant effects of plant type or induction treatment, Tukey HSD post-hoc tests were conducted (package agricolae, version 1.2–4)39. Data were square root transformed prior to analysis to meet the assumptions for normality and homoscedasticity.

Experiment 2: Cotton defense induction in different plant structures. The effect of the explanatory variables plant type (Bt and non-Bt cotton), induction treatment (L. hesperus infested, JA treated, and control) and their interaction on terpenoid concentrations in youngest leaves, squares, and developing bolls were analyzed using separate two-way ANOVAs using the “lm” function. In the case of significant effects of plant type or induction treatment, Tukey HSD post-hoc tests were conducted (package agricolae, version 1.2–4)39. Data were square root transformed prior to analysis to meet the assumptions for normality and homoscedasticity.

Experiment 3: Effect of terpenoids on L. hesperus performance. Kruskal-Wallis tests followed by Holm-Bonferroni post-hoc tests (package agricolae) were used to test the effect of different diet treatments on total nymphal development time. Furthermore, the effect of the treatment on development times for each instar was tested separately. The effect of the treatment on net weight gain was analyzed using ANOVA followed by
Tukey HSD post-hoc tests. Survival among different diets was compared using a Chi-square test (package gmodels version 2.16.2). Nymphs fed with acephate spiked diet (positive control) died within 24 h after the experiment started and were therefore not included in the analyses.

References
1. ISAAA. Global Status of Commercialized Biotech/GM Crops: 2016. ISAAA Brief No. 52. (ISAAA, Ithaca, USA, 2016).
2. NASEM. National Academies of Sciences, Engineering, Medicine—Genetically Engineered Crops: Experiences and Prospects (National Academies Press, Washington, USA, 2017).
3. Carrière, Y. et al. Long-term regional suppression of pink bollworm by Bacillus thuringiensis cotton. Proc. Natl. Acad. Sci. USA 100, 1519–1523 (2003).
4. Adamczyk, J. J. & Hubbard, D. Changes in populations of Heliothis virescens (F.) (Lepidoptera: Noctuidae) and Helicoverpa zea (Boddie) (Lepidoptera: Noctuidae) in the Mississippi Delta from 1986 to 2005 as indicated by adult male pheromone traps. J. Cotton Sci. 10, 829–836 (2006).
5. Wu, K.-M., Lu, Y.-H., Feng, H.-Q., Jiang, Y.-Y. & Zhao, J.-Z. Suppression of cotton bollworm in multiple crops in China with Bt toxin-containing cotton. Science 321, 1676–1678 (2008).
6. Hutchison, W. D. et al. Genetically engineered Bt corn and range expansion of the western bean cutworm (Lepidoptera: Noctuidae) in the United States: a response to Greenpeace Germany. J. Integr. Pest Manage. 2, B1–B8 (2011).
7. Lu, Y. et al. Mirid bug outbreaks in multiple crops correlated with wide-scale adoption of Bt cotton in China. Science 328, 1115–1118 (2010).
8. Naranjo, S. E. Impacts of Bt transgenic cotton on integrated pest management. J. Agrie. Food Chem. 59, 5842–5851 (2011).
9. Choquette, N. F. & Bonning, B. C. Toxins for transgenic resistance to hemipteran pests. Toxins 4, 405–429 (2012).
10. Wilson, L. et al. IPM in the transgenic era: a review of the challenges from emerging pests in Australian cotton systems. Crop. Pasture Sci. 64, 737–749 (2013).
11. Williams, M. Cotton insect losses-2016; http://www.entomology.msstate.edu/resources/cottoncrop.asp Accessed 27 February (2017).
12. Catangui, M. A. & Berg, R. K. Western bean cutworm, S. elysia littoralis (Smith) (Lepidoptera: Noctuidae), as a potential pest of transgenic Cry1Ab Bacillus thuringiensis corn hybrids in South Dakota. Environ. Entomol. 35, 1439–1452 (2006).
13. Eichenseer, H. & Stroehlein, R. Frequency and severity of western bean cutworm (Lepidoptera: Noctuidae) ear damage in transgenic corn hybrids expressing different Bacillus thuringiensis cry toxins. J. Econ. Entomol. 101, 555–563 (2008).
14. Wu, K., Li, W., Feng, H. & Guo, Y. Seasonal abundance of the mirids, Lygus lucorum and Adelphocoris spp. (Hemiptera: Miridae) on Bt cotton in northern China. Crop Protect. 21, 997–1002 (2002).
15. Zeilinger, A. R., Olson, D. M. & Andow, D. A. Competitive release and outbreaks of non-target pests associated with transgenic Bt cotton. Ecol. Appl. 26, 1047–1054 (2016).
16. Zeilinger, A. R., Olson, D. M. & Andow, D. A. Competition between stink bug and heliothine caterpillar pests on cotton at within-plant spatial scales. Entomol. Exp. Appl. 141, 59–70 (2011).
17. Hagenbucher, S. et al. Pest trade-offs in technology: reduced damage by caterpillars in Bt cotton benefits aphids. Proc. Roy. Soc. B 280, 20130042 (2013).
18. Whitehouse, M. et al. Target and nontarget effects of novel “triple-stacked” Bt-transgenic cotton 1: Canopy arthropod communities. Environ. Entomol. 43, 218–241 (2014).
19. Zeilinger, A. R. et al. Behavioural and chemical mechanisms of plant-mediated deterrence and attraction among frugivorous insects. Ecol. Entomol. 40, 532–542 (2015).
20. Altman, D. W., Stipanovic, R. D. & Bell, A. A. Terpenoids in foliar pigment glands of A, D and AD genome cottons: introgression potential for pest resistance. J. Hered. 81, 447–454 (1990).
21. Hagenbucher, S., Olson, D. M., Ruberson, J. R., Wackers, F. L. & Romeis, J. Resistance mechanisms against arthropod herbivores in cotton and their interactions with natural enemies. Crit. Rev. Plant Sci. 32, 458–482 (2013).
22. Alborn, H. T., Röse, U. S. & McAuslane, H. J. Systemic induction of feeding deterrents in cotton plants by feeding of Spodoptera spp. Larvae. J. Chem. Ecol. 22, 919–932 (1996).
23. Bezemer, T. M., Wagenaar, R., Van Dam, N. M. & Wackers, F. L. Interactions between above-and belowground insect herbivores as mediated by the plant defense system. Oikos 101, 555–562 (2003).
24. Eiserring, M. et al. Cotton defense induction patterns under spatially, temporally and quantitatively varying herbivory levels. Front. Plant Sci. 8, 234 (2017).
25. Kaplan, I. & Denno, R. F. Interspecific interactions in phytophagous insects revisited: a quantitative assessment of competition theory. Ecol. Lett. 10, 977–994 (2007).
26. Ohgushi, T., Craig, T. P. & Price, P. W. Ecological communities: plant mediation in indirect interaction webs (Cambridge University Press, Cambridge, UK, 2007).
27. Uraizmi, S., Ando, Y. & Miki, T. Linkages among trait-mediated indirect effects: a new framework for the indirect interaction web. Popul. Ecol. 52, 485–497 (2010).
28. Ellsworth, P. C. & Barkley, V. Cost-effective Lygus management in Arizona cotton. Cotton: A College of Agriculture Report (2001).
29. Tingey, W. M. & Pillmer, E. A. Lygus bugs: crop resistance and physiological nature of feeding injury. Bull. Entomol. Soc. Am. 23, 277–287 (1977).
30. Leigh, T., Roach, S. & Watson, T. Biology and ecology of important insect and mite pests of cotton. In Cotton Insects and Mites: Characterization and Management (eds E. King, J. Phillips & R. J. Coleman) 17–69 (The Cotton Foundation Publisher, Memphis, USA, 1996).
31. Tingey, W. M., Leigh, T. F. & Hyer, A. H. Lygus hesperus: growth, survival, and egg laying resistance of cotton genotypes. J. Econ. Entomol. 68, 28–30 (1975).
32. Ellington, J. et al. Approach to the evaluation of some factors affecting insect resistance in one ‘Acala’ and seven sister genotypes of Stoneville cotton in New Mexico. J. Econ. Entomol. 77, 612–618 (1984).
33. Soler, R. et al. Impact of foliar herbivory on the development of a root-feeding insect and its parasitoid. Oecologia 152, 257–264 (2007).
34. Erb, M. et al. Signal signature of aboveground-induced resistance upon belowground herbivory in maize. Plant J. 59, 292–302 (2009).
35. Wang, M., Bieiere, A., Van Der Putten, W. H. & Bezemer, T. M. Sequential effects of root and foliar herbivory on aboveground and belowground induced plant defense responses and insect performance. Oecologia 175, 187–198 (2014).
36. Hagenbucher, S., Wackers, F. L. & Romeis, J. Indirect multi-trophic interactions mediated by induced plant resistance: impact of caterpillar feeding on aphid parasitoids. Biol. Lett. 10, 20130795 (2014).
37. Nomeir, A. & Abou-Donia, M. Photodecomposition of gossypol by ultraviolet irradiation. J. Am. Oil Chem. Soc. 62, 87–89 (1985).
38. Bottger, G. & Patana, R. Growth, development, and survival of certain Lepidoptera fed gossypol in the diet. J. Econ. Entomol. 59, 1166–1168 (1966).
39. Hagenbucher, S., Eiserring, M., Meisle, M. & Romeis, J. Interaction of transgenic and natural insect resistance mechanisms against Spodoptera littoralis in cotton. Pest Manage. Sci. 73, 1670–1678 (2017).
40. Bi, J. L., Murphy, J. B. & Felton, G. W. Antinutritive and oxidative components as mechanisms of induced resistance in cotton to Helicoverpa zea. J. Chem. Ecol. 23, 97–117 (1997).
41. Schmidt, L., Schurr, U. & Roese, U. S. Local and systemic effects of two herbivores with different feeding mechanisms on primary metabolism of cotton leaves. Plant, Cell Environ. 32, 893–903 (2009).
42. Walling, L. I. Avoiding effective defenses: strategies employed by phloem-feeding insects. Plant Physiol. 146, 859–866 (2008).
43. Eisenring, M., Glauser, G., Meissle, M. & Romeis, J. Differential impact of herbivores from three feeding guilds on systemic secondary metabolite induction, phytohormone levels and plant-mediated herbivore interactions. J. Chem. Ecol. 44, 1178–1189 (2018).
44. Rodriguez-Saona, C., Crafts-Brandner, S. J., Williams, L. III & Paré, P. W. Lygus hesperus feeding and salivary gland extracts induce volatile emissions in plants. J. Chem. Ecol. 28, 1733–1747 (2002).
45. Williams, L., Rodriguez-Saona, C., Paré, P. W. & Crafts-Brandner, S. J. The piercing-sucking herbivores Lygus hesperus and Nezara viridula induce volatile emissions in plants. Arch. Insect Biochem. Physiol. 58, 84–96 (2005).
46. Chen, C. & Parajulee, M. N. Development and population growth of Lygus hesperus on selected weed hosts, artificial diet and cotton in the laboratory. J. Econ. Entomol. 103, 2009–2018 (2010).
47. Tarpley, L. & Sassenrath, G. Carbohydrate profiles during cotton floral bud (square) development. J. Agr. Crop Sci. 192, 363–372 (2006).
48. Eisenring, M., Romeis, J., Naranjo, S. E. & Meissle, M. Multitrophic Cry-protein flow in a dual-gene Bt-cotton field. Agric., Ecosyst. Environ. 247, 283–289 (2017).
49. Debolt, J. W. Meridic diet for rearing successive generations of Lygus hesperus. Ann. Entomol. Soc. Am. 75, 119–122 (1982).
50. Rodriguez-Saona, C., Crafts-Brandner, S. J., Paré, P. W. & Henneberry, T. J. Exogenous methyl jasmonate induces volatile emissions in cotton plants. J. Chem. Ecol. 27, 679–695 (2001).
51. Stipanovic, R. D., Bell, A. A., O’Brien, D. H. & Lukefahr, M. J. Heliciode H1. A new insecticidal C25 terpenoid from cotton (Gossypium hirsutum). J. Agric. Food Chem. 26, 115–118 (1978).
52. Benson, C. G., Wyllie, S. G., Leach, D. N., Mares, C. L. & Fitt, G. P. Improved method for the rapid determination of terpenoid aldehydes in cotton. J. Agric. Food Chem. 49, 2181–2184 (2001).
53. Stipanovic, R. D., Altman, D. W., Begin, D. L., Greenblatt, G. A. & Benedict, J. H. Terpenoid aldehydes in upland cotton: analysis by aniline and HPLC methods. J. Agric. Food Chem. 36, 509–515 (1988).
54. McAuslane, H. J., Alborn, H. T. & Toth, J. P. Systemic induction of terpenoid aldehydes in cotton pigment glands by feeding of larval Spodoptera exigua. J. Chem. Ecol. 23, 2861–2879 (1997).
55. Leifeld, J., Ammann, C., Nefel, A. & Fuhrer, J. A comparison of repeated soil inventory and carbon flux budget to detect soil carbon stock changes after conversion from cropland to grasslands. Global Change Biol. 17, 3366–3375 (2011).
56. R Core Team. R: a language and environment for statistical computing (R Foundation for Statistical Computing, Vienna, Austria, http://www.r-project.org/, 2017).
57. Bates, D., Mächler, M., Bolker, B. & Walker, S. Fitting linear mixed-effects models using lme4. J. Stat. Softw. 67, 1–48 (2015).
58. Burnham, K. P. & Anderson, P. Model Selection and Multimodel Inference: A Practical Information-Theoretic Approach. (Springer, New York, USA, 2002).
59. Barton, K. MuMln: Multi-Model Inference. R package version 1.15.6, https://CRAN.R-project.org/package=MuMIn (2016).
60. Richards, S. A. Dealing with overdispersed count data in applied ecology. J. Appl. Ecol. 45, 218–227 (2008).
61. De Mendiburu, F. Agricolae: Statistical Procedures for Agricultural Research. R package version 1.2–4, https://CRAN.R-project.org/package=agricolae (2016).
62. Warnes, G. R., Bolker, B., Lumley, T. & Johnson, R. C. Contributions from Randall C. Johnson are Copyright SAIC-Frederick, Inc. Funded by the Intramural Research Program, of the NIH, National Cancer Institute and Center for Cancer Research under NCI Contract NO1-CO-12400. gmodels: Various R Programming Tools for Model Fitting. R package version 2.16.2, https://CRAN.R-project.org/package=gmodels (2013).

Acknowledgements
We thank Felix Wettstein and Jiwon Park for technical support as well as Christian Bochet for the heliocide H1/H4 extraction. This work was supported by the Swiss National Science Foundation [SNF grant No. 31003A-149794].

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M.E., M.M., J.R. and S.N. planned and designed the experiments, M.E., S.N. and A.A. conducted the experiments, M.E. performed the chemical analyses, M.E. and S.B. analyzed the data statistically, M.E., M.M., S.B., S.N. and J.R. wrote the manuscript.

Additional Information
Competing Interests: The authors declare no competing interests.

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