Comparison of genetically modified insect-resistant maize and non-transgenic maize revealed changes in soil metabolomes but not in rhizosphere bacterial community

Yanjun Chen, Libo Pan, Mengyun Ren, Junsheng Li, Xiao Guan, and Jun Tao

ABSTRACT
The deliberate introduction of the beneficial gene in crop plants through transgenic technology can provide enormous agricultural and economic benefits. However, the impact of commercialization of these crops on the ecosystem particularly on belowground soil biodiversity is still uncertain. Here, we examined and compared the effects of a non-transgenic maize cultivar and an insect-resistant transgenic maize cultivar genetically engineered with cry1Ah gene from Bacillus thuringiensis, on the rhizosphere bacterial community using 16S rDNA amplicon sequencing and soil metabolome profile using UPLC/MS analysis at six different growth stages. We found no significant differences in bacterial community composition and diversity at all growth stages between the two cultivars. The analysis of bacterial beta-diversity showed an evident difference in community structure attributed to plant different growth stages but not to the plant type. In contrast, the soil metabolic profile of transgenic maize differed from that of the non-transgenic plant at some growth stages, and most of the altered metabolites were usually related to the metabolism but not to the plant-microbe interaction related pathways. These results suggest that genetic modification with the cry1Ah gene- altered maize soil metabolism but had no obvious effect on the rhizosphere bacterial community.

Introduction
With the ever-growing human population, food security is the major concern of this century. Thus, the agriculture sector has been revolutionized to obtain higher crop yield per capita through various sustainable approaches. Genetically modified (GM) crops, as a result of the introduction of beneficial genes in a crop plant, can provide sustainable agronomic and economic benefits. For instance, when a gene from Bacillus thuringiensis (Bt) responsible for the production of insecticidal toxin is introduced into certain crops, it allows protection of GM plants against a specific group of insect pests through the direct action of insecticidal toxin. Since their first commercialization in 1996, several GM crops such as insect-resistant, herbicide-resistant, combined insect- and herbicide-resistant and viral disease-resistant crops are being cultivated in different parts of the world. Although the global cultivable area of GM is increasing every year, their effects on ecosystem biodiversity remain controversial.

Some serious agricultural concerns associated with the commercial cultivation of GM crops have been reported so far. In Bt crops, for example, the target insect pests may develop Bt toxin resistance over time (Guan, Hou, Dai, Liu, Liu, Gu, Jin, Yang, Fabrick, Wu, 2021). Another concern is the transgene flow from Bt crops to surrounding plant diversity, and the potential development of ‘super weed’ is one of its examples. Similar to aboveground, the study of consequences of commercial cultivation of Bt crops on belowground biodiversity is of equal importance, as the health and performance of crop plants is highly dependent on soil biological processes.

Soil microbial community is a principal component of soil ecosystem functioning. Plant root harbors a mesmerizing diversity of microbes...
mainly dominated by the bacterial community in their ambient environment that is called the rhizosphere.\textsuperscript{13,14} These rhizosphere bacteria, ranging from plant pathogenic to beneficial ones, have great influences on host plant health as some may regulate plant nutrient acquisition ability or modulate host immunity.\textsuperscript{15–17} Plant actively secretes photosynthetically fixed carbon in the shape of root exudates, usually consisting of primary and secondary metabolites, into the soil, which acts as the energy source for the rhizosphere microbial community\textsuperscript{18} (Rahman et al.,\textsuperscript{19} 2021). This phenomenon gives hosts to preferentially select and shape their rhizosphere microbiota through modification in root exudation.\textsuperscript{20–23}

GM plants also secrete these metabolites to determine root-associated microbiota. The expression of specific proteins from GM plants may alter the composition of root metabolites, thereby the composition of root-associated microbiota, which may, in turn, affect the soil biological environment.\textsuperscript{24} So far, contrasting effects of GM crops on rhizosphere microbial community have been reported with a plethora of studies suggesting no obvious effect,\textsuperscript{25–28} while some studies showing significant changes in the rhizosphere microbial community as compared to non-GM crops\textsuperscript{29} (Guan, Wei, Stewart, Tang, 2021). Studies also indicate that other factors such as the plant growth stage could be a major indicator of changes in the rhizosphere microbial community.\textsuperscript{30,31} These studies suggest the necessity of assessing the effects of a given genetically modified crop on its rhizosphere microbial community before its commercialization.

The Cry1Ah protein of cry1Ah, a novel insecticidal gene from Bt subspecies, exhibits great toxicity to Lepidopteran insects, and its efficiency has been found higher than other insecticidal genes such as cry1Ab and cry1Ac.\textsuperscript{32,33} The expression of Cry1Ah might lead to altering plant root metabolites composition to induce changes in ambient soil biodiversity.\textsuperscript{24} Although we found no significant effects on weeds, nematodes and other invertebrate occurrences in cry1Ah genetically engineered maize HGK60 surroundings as compared to its non-transgenic control,\textsuperscript{34,35} its effects on the rhizosphere microbial community remain to be explored. Here, we used 16s rDNA amplicon sequencing to study the changes in rhizosphere bacterial community composition, and non-targeted metabolomics for soil metabolite profiling of transgenic and non-transgenic maize.

**Materials and Methods**

**Plant Material and Field Experiment**

The seeds of GM maize HGK60 with insecticidal gene cry1Ah and non-transgenic maize ZHENG58 (control) were provided by the Chinese Academy of Agricultural Sciences, China. The field experiment was carried out from May to September 2020 in an open field at the experimental station (116°36’34”E, 39°36’10’’N) of the Chinese Academy of Agricultural Sciences located in Langfang, China, with a tropical monsoon climate, an average lowest-highest temperature of 24.3–28.8°C, and average monthly precipitation about 122 mm for the plant growth period. The experimental site has been cultivated with HGK60 maize for more than 10 years.

Both maize types were planted in separate plots (10 m × 10 m) in a randomized block design with six replicates for each type (~12 plots). Plots were separated from each other with a 1 m wide uncultivated zone. The sowing of seeds in each plot was carried out with inter-plant spacing of 25 cm and inter-row spacing of 60 cm. The plots were managed conventionally.

**Sample Collection**

Rhizosphere soil samples from both maize types were collected at six different growth stages, i.e., pre-planting stage (April 28), seedling stage (May 10), bell stage (June 10), heading stage (June 30), fully ripe stage (August 10), and after-harvest stage (September 10). Briefly, the maize plant roots were carefully removed from the soil and shaken by hand to remove loosely attached soil (not rhizosphere soil). Then, soils tightly adhering to roots were removed by a sterile brush (rhizosphere soils). To limit the influence of soil heterogeneity, the preparation of composite soil samples for DNA extraction is recommended (Vestergaard et al.,\textsuperscript{36} 2017). Therefore, a composite sample was prepared from random ten maize plants in each replicate of the individual treatment. This resulted in six composite rhizosphere soil samples for each
treatment. After sieving (2 mm mesh), these fresh rhizosphere soil samples were stored at −80°C for DNA extraction.

**DNA Extraction, Illumina MiSeq Sequencing and Raw Data Processing**

Soil DNA was extracted from 0.5 g of rhizosphere soil using a FastDNA ® SPIN Kit for soil (MP Biomedicals, Santa Ana, CA) following the manufacturer’s instructions. The concentration and purity of DNA was confirmed by an ultra-micro spectrophotometer.

The V1-V9 region of the bacterial gene was amplified in PCR assay using the universal primer set of 8 F (5′-AGAGTTTGATCCTGGCTCAG-3′) and 1509 R (5′-GNTACCTGTTACGACTT-3′). The PCR reaction (50 μL) was: Trans Fastpfbu 1 μL, 5 x Buffer 10 μL, 5 x Stimulate 5 μL, dNTPs (2.5 × 10⁻³ mol/L each) 5 μL, Primer Mix (1 μmol/L) 2 μL, gDNA 1 μL, NFW 26 μL. The conditions for PCR reactions were: Pre-denaturation temperature 98°C (2 min), denaturation temperature 95°C (30 s), annealing temperature 60°C (45 s), extension temperature 72°C (90 s), after 35 cycles, the extension was terminated at 72°C for 10 min. The product of the triplicate reaction was pooled and purified using an Agarose Gel DNA purification kit (TaKaRa). Then, a TBA-380 micro-fluorometer with PicoGreen reagent was used to quantify the purified amplicons. The third generation full-length 16S amplicon sequencing was then performed on PacBio platform at Novogene Co. Ltd., Beijing, China.

The obtained raw reads were demultiplexed, quality filtered and further processed using FLASH. Operational taxonomic units (OTUs) were generated by binning the unique sequences at 97% sequence similarity with the help of an agglomerative clustering algorithm using UPRASE. The classification of sequences of OTUs was carried out with SILVA database. The identification and removal of chimeric sequences were done with the help of UPRASE 6.1 in QIIME. The data of all the sequences was uploaded in NCBI Sequence Archive with the submission accession number (PRJNA75527).

**Metabolites Extraction from Maize Soil, Untargeted Metabolomics Analysis and Raw Data Processing**

The procedure described by was used for the collection of maize soil metabolites with some modifications. Briefly, maize plants with whole root system were collected and made sure that most of the root-soil was retained, and placed in a pot (size varied with the plant age). For pre- and post-harvest stages, only soil (100 g) for each sample was collected and used. 50% methanol solution (0.05% formic acid, v/v) was applied (15 mL for pre-harvest, seedling and post-harvest growth stage for 1 min each, and 30 mL for the remaining three growth stages for 2 min each) and flushed through the pot with pressure using a syringe. After that, 10 mL of extract was collected in a centrifuge tube, and centrifuged to pellet soil residues (5 min, 3500 g). The supernatant (4 mL) was collected and transferred to a new centrifuge tube and initially frozen in liquid nitrogen. Then, the samples were freeze-dried for 48 hours and stored at −80°C for metabolites analysis.

A total of 72 samples (2 treatments × 6 growth stages × 6 replicates) were re-suspended in a 100 μL of solution (50% methanol, 49.9% water, 0.1% formic acid; v/v), sonicated at 4°C, and centrifuged at 14000 g for 15 min at 4°C. The supernatant (80 μL) was then transferred into glass vials containing a glass insert before the analysis on a UPLC system (Thermo, Ultimate 3000LC). Hyper gold C18 (1.9 μm internal diameter) column eluted with a multistep gradient throughout 0.3 mL/min at 40°C was used. The gradient used was consisted of A (94.9% water, 5% acetonitrile, and 0.1% formic acid), and B (99.9% acetonitrile, and 0.1% formic acid). The UPLC system was coupled with a mass spectrometer (Triple-TOF 5600) coupled with an electrospray ionization source. Data acquisition was carried out in full scan mode along with IDA mode. The analysis conditions set for mass spectrometry were as follows: Ion source temperature, 550°C (for both + and − ion); ion spray voltage 5500 V (+) and 4500 V (−); collision energy 10 ev (for both + and − ion) interface heater temperature 550°C (+) and 600°C (−), and curtain gas 35 PSI. The QCs were injected at regular time intervals throughout the run to provide data that can be assessed repeatedly.
The raw data acquired from UPLC-MS were processed with the help of progenesis QI software (Waters Corp., USA). The threshold values set were as: precursor tolerance, 5 ppm; fragment tolerance, 10 ppm; and retention time, 0.02 min. The data were obtained with m/z, peak retention time and peak intensities. The retention time m/z was used as the ion identifier. Any peak with a missing value was removed. The targeted Peak Finding function in Master View 1.0 software was used to match the molecular formula with the published known compounds in the database. For non-targeted peak findings: we imported the data into the Marker View 1.2.1 software, matched and picked the chromatographic peak using peak finding options. The taxonomic identification of metabolites was carried out online on The Human Metabolome Database and LIPID MAPS using the matched formula. Data from both negative and positive ions were combined for bioinformatics analysis in R (version 4.1.1). Finally, the KEGG pathways were identified and constructed at KEGG online (http://www.genome.jp/kegg/).

Statistical Analysis

The alpha diversity indices of the rhizosphere bacterial community including species richness and Shannon index were calculated using QIIME. The bacterial beta diversity analysis, the weighted UniFrac distances, were calculated using QIIME. For estimation of dissimilarity in bacterial community structure, principal component analysis (PCA), principal coordinate analysis (PCoA) and Bray-Curtis dissimilarity matrix calculations were conducted in the vegan package and visualized in ggplot2 package in R.

The data of alpha diversity and bacterial differential relative abundance were analyzed by analysis of variance (ANOVA), and means were compared based on student t-test. False discovery rate (FDR) was calculated for bacterial and soil metabolites differential analysis using edgeR package in R. The significance of data based on FDR value (=0.05) was considered with a fold change threshold (log-fold-change >1 or <-1). The differences in soil metabolites between two maize cultivars were compared using partial least square discriminant analysis (PLS-DA). Graphs were visualized using the ggplot2 package in R.

Results

Alpha and Beta Diversity of Maize Rhizosphere Bacterial Community

The transgenic maize did not affect the rhizosphere bacterial alpha diversity as the values of alpha diversity indices were not different between transgenic and control maize, except the Shannon index which was higher in control maize at the first stage (P > .05) (Fig. 1a, b). PCA and PCoA analysis based on UniFrac distance showed that the bacterial community distinctly differed among all the six growth stages in both maize cultivars; however, no difference was observed between transgenic and control maize when compared at each stage (Fig. 1c, d). On average, the explained Bray-Curtis dissimilarity ranged from 37.80% to 49.48% between samples of transgenic and control at all six stages (Fig. 1e). However, the analysis of similarities (ANOSIM) revealed that the dissimilarity was not strong (R-value close to zero) and statistically non-significant (P > .05) at all six stages (Fig. 1e). The results of adonis analysis of comparison of two groups at each growth stage are provided in supplementary file 1.

Composition of Maize Rhizosphere Bacterial Community

The rarefaction curve of bacterial OTUs at 97% of sequence similarity tended to reach the saturation plateau (Fig. 2a), indicating that large diversity of bacterial community was covered by the present analysis. The Good’s coverage, which reflects the captured microbial diversity in samples, for transgenic and control maize was 81.99% and 82.04%, respectively (Fig. 2b). A total of 50 classified bacterial phyla were detected in all the samples while 2.42% of sequences were unclassified. Proteobacteria, Acidobacteria and Bacteroidetes were the dominant groups, together accounting for 72.13% of the total sequences (Fig. 2c). At the class level, 34 classified groups were found in all the samples whereas 4.44% of total sequences could not be assigned to any class group. Major classes were Gammaproteobacteria, Bacteroidia, Alphaproteobacteria and Deltaproteobacteria, together accounting for 46.14% of total sequences (Fig. 2d).
Taxonomic assignment of identified sequences at the genus level resulted in 558 identified and unidentified groups. Among prominent classified genera (relative abundance >0.1% in at least one treatment), the relative abundances of *Azohydromonas* at stage one, *Mesorhizobium* at stage two, *Bryobacter*, *Ohtaekwangia* and *Paenisporosarcina* at stage three, *Dongia*, *Nordella*, *Paenibacillus* and *Gaiella* at stage four, *Mesorhizobium* at stage five, and *Pedomicrobium* at stage six were higher, whereas that of *Gemmata* and *Segetibacter* at stage two, *Archangium* at stage three, *Cupriavidus* at stage five and *Opitutus* at stage six were lower in transgenic maize rhizosphere than control (*P* < .05) (Fig. 2e). The FDR value obtained from the differential analysis of bacterial community at all taxonomic levels was more than 0.05 (data not shown), indicating that the bacterial composition was not different in transgenic maize and control.

**Difference in Maize Soil Metabolomics Profile of Transgenic and Control Maize**

Soil metabolomics profiling of two maize cultivars generated a total of 1730 compounds. PLS-DA analysis revealed complete separation between the metabolites of transgenic and control at each
growth stage (Supplementary file 2; Fig. S1). Moreover, the degree of change in metabolites increased up to the fourth growth stage and decreased afterward (Fig. 3). In total, 246 metabolites were altered based on threshold of FDR < 0.05; log2foldchange>1 or <-1, with 37 metabolites overlapping at different growth stages (Supplementary file 3). As compared to control, the transgenic maize increased the concentration of 3, 3, 12, 126, 14 and 14 compounds while decreased that of 7, 12, 42, 29, 6 and 9 compounds at stages 1, 2, 3, 4, 5 and 6, respectively (Fig. 3). The results suggest an obvious effect of transgenic maize on soil metabolite profile as compared to control.

The altered metabolites were further assigned to their respective KEGG pathways resulting in a total of 59 KEGG pathways at level III, 43 of those belonging to metabolism at KEGG level 1 (Supplementary file 2; Table S1). The detailed assignment of altered metabolites in both maize cultivars is provided in Table 1. In addition, most of the altered metabolites related to KEGG pathways were observed at growth stage four followed by three.

**Relationship between Maize Soil Metabolites and Rhizosphere Bacterial Community**

Since the relative abundances of bacterial OTUs in transgenic and control maize rhizosphere at each growth stage were not different (FDR > 0.5), we only selected the top 10 most abundant OTUs and 10 most abundant altered metabolites at each growth stage, and their relationships were tested using Spearman’s rank correlation coefficient (ρ) (threshold, ρ > 0.7 or ρ < −0.7, and P < .05) (Fig. 4).

At growth stage one, the relative abundances of OTU1 belonging to family Pyrinomonadaceae, and OTU3 belonging to genus *Stenotrophobacter* were positively correlated with altered metabolites identified as D-gluconic acid, 2-[amino(3-chloroanilino)
Figure 3. Volcano plot illustrating the differentially altered soil metabolites of transgenic maize as compared to control cultivar at different growth stages. The threshold level for FDR < 0.05 and fold change (2) > 1 or < -1 was set.

Table 1. The number of differentially altered metabolites belonging to KEGG pathways (level II). The detailed relationship between these metabolites and KEGG pathways are provided in Supplementary file 2; Table S1.

| KEGG pathway level II                           | Stage 1 | Stage 2 | Stage 3 | Stage 4 | Stage 5 | Stage 6 |
|------------------------------------------------|---------|---------|---------|---------|---------|---------|
| Up                                             | Down    | Up      | Down    | Up      | Down    | Up      |
| Skeleton-based classification                   | –       | –       | –       | –       | –       | –       |
| Membrane transport                             | –       | –       | 1       | 2       | 1       | –       |
| Signal transduction                            | –       | –       | –       | 1       | –       | –       |
| Signaling molecules and interaction            | –       | –       | –       | –       | 1       | –       |
| Folding, sorting and degradation               | –       | –       | 1       | –       | –       | –       |
| Amino acid metabolism                          | –       | –       | 1       | 8       | –       | –       |
| Biosynthesis of other secondary metabolites    | –       | –       | 1       | 3       | 6       | –       |
| Carbohydrate metabolism                        | –       | –       | –       | 6       | 1       | –       |
| Chemical structure transformation maps          | –       | –       | –       | 4       | 1       | –       |
| Global and overview maps                       | –       | 4       | 2       | 3       | 13      | 37      |
| Lipid metabolism                               | –       | –       | 2       | 13      | 3       | 1       |
| Metabolism of cofactors and vitamins           | –       | –       | –       | 4       | 2       | 3       |
| Metabolism of terpenoids and polyketides       | –       | –       | –       | 1       | 1       | –       |
| Nucleotide metabolism                          | –       | –       | –       | 2       | –       | –       |
| Nucleotide metabolism                          | –       | –       | –       | –       | 2       | –       |
| Xenobiotics biodegradation and metabolism       | –       | –       | 1       | 1       | 11      | –       |
| Circulatory system                             | –       | –       | –       | –       | –       | –       |
| Digestive system                               | –       | –       | 1       | 1       | 2       | –       |
| Endocrine system                               | –       | –       | –       | 1       | –       | –       |
| Environmental adaptation                       | –       | –       | –       | 2       | –       | –       |
| Nervous system                                 | –       | –       | –       | –       | 1       | –       |
| Sensory system                                 | –       | –       | –       | –       | –       | –       |
methylene]malononitrile, Glycyl-L-prolyl-L-lysine (only OTU3) and Prolylleucine, whereas OTU2 belonging to family Nitrosomonadaceae was negatively correlated with Albiflorin and GPK, and all were decreased in transgenic maize soil. At stage three, OTU2 and OTU14 belonging to the family Nitrosomonadaceae were positively correlated with the metabolite Rufloxacin, and OTU3 was negatively correlated with 7-Methylguanosine; all were increased in transgenic maize soil. At stage four, OTU21 from order Nostocales was positively correlated to a Triazole compound while OTU14 was negatively correlated with Norfentanyl, both increased in transgenic maize. At stage five, the OTU18 from Gemmatimonadaceae was positively correlated to an unidentified benzodioxin containing compound. The OTU18 along with OTU11 from the family Nitrosomonadaceae and OUT10 from genus Steroidobacter were positively correlated with Methyl palmitate at stage six which was increased in transgenic maize. Another correlation analysis revealed that the metabolite inventory in transgenic and control maize soils were non significantly ($P > .05$) correlated with the diversity of the bacterial community at all six different growth stages (Fig. 5).

**Discussion**

The transfer of transgene flow from GM crops into the soil via crop stubbles or root exudates may cause changes in plant ambient soil environment, potentially impacting the soil biodiversity and ultimately posing a threat to the soil ecosystem. Therefore, the impact of transgenic crops on soil fauna has been extensively studied in different GM crops, largely
resulting in minimal to no effects on biodiversity. In this context, the variable response of soil microbial community to Bt crops has triggered a controversial debate over the last two decades mainly due to the different approaches used to study the rhizosphere microbial community.

Using amplicon sequencing, we found that the alpha and beta diversities of rhizosphere bacterial community of transgenic and control maize did not differ, except the Shannon index at earlier stage which could be the usual transient impact that Bt crops exert on soil microbes. However, obvious differences were found in bacterial beta diversity at different growth stages of both maize cultivars. Likewise, the differences in bacterial relative abundances based on ANOVA were not supported by FDR analysis, which reduces the probability of false-positive in statistical significance level. These results revealed that the rhizosphere bacterial community varies with growth stages but rather than plant genotypes. Corroborating with these results, several studies have found that GM crops do not pose significant effects on soil microbial community composition and diversity (Sun et al., 2016) Fan et al. (2019) found in a 2-year-long trial that insect-resistant transgenic maize carrying cry1le gene did not affect the soil fauna. A recent study found that the maize genetically engineered with mcry1Ab and mcry2Ab genes had minimal to no effect on the rhizosphere bacterial community, and that the different development stages could induce those changes. When required, a plant can shape and modify its root microbiota through alteration in root exudation. The difference in bacterial community structure at different growth stages in our study is certain because of the difference in root exudation chemistry due to variable root biomass and hormonal changes at different stages of growth and development.

Although the rhizosphere bacterial community was not altered in the transgenic maize in our study, we yet analyzed the changes in soil metabolomic profile due to the potential transgene flow. As predicted earlier, the metabolomic profile greatly differed between transgenic and non-transgenic maize cultivars at all stages, and the difference was more prominent at mid-stages. It is inevitable because the Bt toxin of GM crops has been shown to impact the root exudation profile. The novel insecticidal protein from Bt crops can be directly released into the soil through root exudation, or their expression can trigger
changes in various metabolic pathways resulting in a change in concentration of certain metabolites in the soil surrounding the root system. Moreover, the prominent difference in soil metabolites during mid-growth stages as compared to other stages could be attributed to the difference in plant physiology. During the transition from vegetative growth to reproductive growth, plants undergo several hormonal and metabolic changes, which may greatly alter the root metabolism as well as the metabolites diversity in the root ambient soil.

Plant roots exude a proportion of photosynthetically fixed carbon into the soil in the form of different organic compounds, which serve as a major energy reservoir for the soil microbial community. Therefore, the composition of root-associated microbial community (i.e., in the rhizosphere) is strictly linked with the composition of root exudates of a plant, and any change in root exudation could potentially alter the soil biodiversity in the close vicinity. Despite the obvious differences in metabolic profile in the soils of two maize cultivars at each growth stage, similar to the results of most of the previous studies conducted on cry1Ah insecticidal maize, no difference in rhizosphere bacterial community was found in our study. Interestingly, the relative abundances of some bacterial OTUs were yet correlated with the altered metabolites in transgenic maize. Even though the different soil metabolic profiles could not alter the bacterial community composition, the association of some bacterial taxa to altered metabolites is not surprising but intuitive.

Based on Spearman’s rank correlation coefficient, we also found that the bacterial diversity was not related to the changes in soil metabolite assortment. Nevertheless, the effects of GM crops on the rhizosphere microbial community are instinct. However, it depends on the spectrum of activity of the transgene protein, the change in root exudation profile and the nature and toxicity of altered metabolites. Since examining the nature of all the altered compounds and their level of toxicity on the soil bacterial community was quite troublesome, we screened out the KEGG pathways involving those altered metabolites at different stages. We found that almost third quarter of KEGG pathways with altered metabolites was related to metabolism only, and none of them was found involved in plant–microbe interaction-related pathways such as bacterial chemotaxis, two-component system, biofilm formation (Bonlan, 2001), quorum sensing, MAPK signaling and plant hormone signal transduction. These results suggested that although the metabolite inventory in transgenic maize soil differed from that of non-transgenic maize, the altered metabolic profile was not quite related to the bacterial recruitment process. This may explain why the rhizosphere bacterial community composition, diversity and structure in the cry1Ah transgenic maize were not different from that of the non-transgenic maize even with a different soil metabolite composition. However, we strongly recommend investigating the specificity of altered compounds in GM crops and their effects on soil microbial community both in vivo and in vitro to deepen our understandings of the potential effects of transgenic crops on soil biodiversity.

Conclusion

The environmental biosafety of transgenic crops has always been a big concern since their commercialization. In this study, we assessed the effects of insect-resistant transgenic maize genetically engineered with cry1Ah gene on rhizosphere bacterial community and changes in soil metabolites at six different growth stages. We found that the change in rhizosphere bacterial community was related to the plant developmental stages but not to the plant genetic modification, while metabolic profile greatly differed in transgenic and non-transgenic maize. This study revealed that the insect-resistant transgenic maize genetically engineered with cry1Ah gene has no obvious effect on the rhizosphere bacterial community. However, the potential role of recognized and unrecognized metabolites altered in transgenic maize soil cannot be overlooked and should be assessed to evaluate the biosafety of GM maize.

Disclosure Statement

No potential conflict of interest was reported by the authors.
**Funding**

This work was funded by the National Science and Technology Major Project of the Ministry of Science and Technology of China (National Science and Technology Major Project 2016ZX08012005).

**ORCID**

Yanjun Chen [http://orcid.org/0000-0002-5618-4086](http://orcid.org/0000-0002-5618-4086)

**Author Contributions**

Conceptualization, Y C, X G, J T; methodology, Y C; software, M R; validation, Y C, L P, M R, J L, X G, J T; formal analysis, Y C, L P, M R, J L, X G, J T; investigation, Y C, L P, M R, J L, X G, J T; resources, X G, J T; data curation, Y C; writing—original draft preparation, Y C; writing—review and editing, X G, J T; visualization, Y C; supervision, X G, J T; project administration, X G, J T; funding acquisition, X G, J T. All authors have read and approved the final version of the manuscript.

**References**

1. Ramankutty N, Mehrabi Z, Waha K, Jarvis L, Kremen C, Herrero M, Rieseberg LH. Trends in global agricultural land use: implications for environmental health and food security. Annu Rev Plant Biol. 2018 Apr 29;69(1):789–815. doi:10.1146/annurev-arplant-042817-040256.

2. Catacora-Vargas G, Binimelis R, Myhr AI, Wynne B. Socio-economic research on genetically modified crops: a study of the literature. Agric Human Values. 2018 Jun;35(2):489–513. doi:10.1007/s10460-017-9842-4.

3. Palma L, Muñoz D, Berry C, Murillo J, Caballero P. Bacillus thuringiensis toxins: an overview of their biocidal activity. Toxins. 2014 Dec;6(12):3296–325. doi:10.3390/toxins6123296.

4. Anderson J, Ellsworth PC, Faria JC, Head GP, Owen MD, Pilcher CD, Shelton AM, Meissle M. Genetically engineered crops: importance of diversified integrated pest management for agricultural sustainability. Front Bioeng Biotechnol. 2019 Feb 20;7:24. doi:10.3389/fbioe.2019.00024.

5. Kumar K, Gambhir G, Dass A, Tripathi AK, Singh A, Jha AK, Yadava P, Choudhary M, Rakshit S. Genetically modified crops: current status and future prospects. Planta. 2020 Apr;251(4):1–27. doi:10.1007/s00425-020-03372-8.

6. Nawaz MA, Mesnage R, Tsatsakis AM, Golokhvast KS, Yang SH, Antoniou MN, Chung G. Addressing concerns over the fate of DNA derived from genetically modified food in the human body: a review. Food Chem Toxicol. 2019 Feb 1;124:423–30. doi:10.1016/j.fct.2018.12.030.

7. Campos SO, Santana IV, Silva C, Santos-Amaya OF, Guedes RN, Pereira EF. Bt-induced hormesis in Bt-resistant insects: theoretical possibility or factual concern?. Ecotoxicol Environ Saf. 2019 Nov 15;183:109577. doi:10.1016/j.ecoenv.2019.109577.

8. Tabashnik BE, Carrière Y. Surge in insect resistance to transgenic crops and prospects for sustainability. Nat Biotechnol. 2017 Oct;35(10):926–35. doi:10.1038/nbt.3974.

9. Mandal A, Sarkar B, Owens G, Thakur JK, Manna MC, Niazi NK, Jayaraman S, Patra AK. Impact of genetically modified crops on rhizosphere microorganisms and processes: a review focusing on Bt cotton. Appl Soil Ecol. 2020 Apr 1;148:103492. doi:10.1016/j.apsoil.2019.103492.

10. Berendse RL, Pieterse CM, Bakker PA. The rhizosphere microbiome and plant health. Trends Plant Sci. 2012 Aug 1;17(8):478–86. doi:10.1016/j.tplants.2012.04.001.

11. Gao D, Pan X, Khashi U Rahman M, Zhou X, Wu F. Common mycorrhizal networks benefit to the asymmetric interspecific facilitation via K exchange in an agricultural intercropping system. Biol Fertil Soils. 2021;57(7):959–71. doi:10.1007/s00374-021-01561-5.

12. Guan F, Hou B, Dai X, Liu S, Liu J, Gu Y, Jin L, Yang Y, Fabrick JA, Wu Y. Multiple origins of a single point mutation in the cotton bollworm tetraspanin gene confers dominant resistance to Bt cotton. Pest Manag Sci. 2021 Mar;77(3):1169–77. doi:10.1002/ps.6192.

13. Guan ZJ, Wei W, Stewart CN Jr, Tang ZX. Effects of transgenic oilseed rape harboring the Cry1Ac gene on microbial communities in the rhizosphere soil. Eur J Soil Biol. 2021 Mar 1;103:103277. doi:10.1016/j.ejsobi.2021.103277.

14. Khashi U Rahman M, Zhou X, Wu F. The role of root exudates, CMNs, and VOCs in plant–plant interaction. J Plant Interact. 2019 Jan 1;14(1):630–36. doi:10.1080/17429145.2019.1689581.

15. Bai YC, Chang YY, Hussain M, Lu B, Zhang JP, Song XB, Lei XS, Pei D. Soil chemical and microbiological properties are changed by long-term chemical fertilizers that limit ecosystem functioning. Microorganisms. 2020 May;8(5):694. doi:10.3390/microorganisms8050694.

16. Hussain M, Hamid MI, Tian J, Hu J, Zhang X, Chen J, Xiang M, Liu X. Bacterial community assemblages in the rhizosphere soil, root endosphere and cyst of soybean cyst nematode-suppressive soil challenged with nematodes. FEMS Microbiol Ecol. 2018 Oct;94(10):fiy142. doi:10.1093/femsec/fiy142.

17. Trivedi P, Leach JE, Tringe SG, Sa T, Singh BK. Plant–microbiome interactions: from community assembly to plant health. Nat Rev Microbiol. 2020 Nov;18 (11):607–21. doi:10.1038/s41579-020-0412-1.

18. Vives-Peris V, de Ollas C, Gómez-Cadenas A, Pérez-Clemente RM. Root exudates: from plant to rhizosphere and beyond. Plant Cell Rep. 2020 Jan;39(1):3–17. doi:10.1007/s00299-019-02447-5.
19. Rahman MK, Wang X, Gao D, Zhou X, and Wu F. Root exudates increase phosphorus availability in the tomato/potato onion intercropping system. Plant and Soil. 2021 Apr 7:1–8
20. Badri DV, Vivanco JM. Regulation and function of root exudates. Plant Cell Environ. 2009 Jun;32(6):666–81. doi:10.1111/j.1365-3040.2009.01926.x
21. Baetz U, Martinoia E. Root exudates: the hidden part of plant defense. Trends Plant Sci. 2014 Feb 1;19(2):90–98. doi:10.1016/j.tplants.2013.11.006
22. Topalović O, Hussain M, Heuer H. Plants and associated soil microbiota cooperatively suppress plant-parasitic nematodes. Front Microbiol. 2020 Feb 28;11:313. doi:10.3389/fmicb.2020.00313
23. Zhou X, Khashi U Rahman M, Liu J, Wu F. Soil acidification mediates changes in soil bacterial community assembly processes in response to agricultural intensification. Environ Microbiol. 2021;23(8):4741–55. doi:10.1111/1462-2900.15675.
24. Singh AK, Dubey SK. Current trends in Bt crops and their fate on associated microbial community dynamics: a review. Protoplasma. 2016 May 1;253(3):663–81. doi:10.1007/s00709-015-0903-5
25. Kapur M, Bhatia R, Pandey G, Pandey J, Paul D, Jain RK. A case study for assessment of microbial community dynamics in genetically modified Bt cotton crop fields.Curr Microbiol. 2010 Aug;61(2):118–24. doi:10.1007/s00284-010-9585-6.
26. Li P, Li Y, Shi J, Yu Z, Pan A, Tang X, Ming F. Impact of transgenic Cry1Ac+ CpTI cotton on diversity and dynamics of rhizosphere bacterial community of different root environments. Sci Total Environ. 2018 Oct 1;637:233–43. doi:10.1016/j.scitotenv.2018.05.013
27. Qiao Q, Wang F, Zhang J, Chen Y, Zhang C, Liu G, Zhang H, Ma C, Zhang J. The variation in the rhizosphere microbiome of cotton with soil type, genotype and developmental stage. Sci Rep. 2017 Jun 21;7(1):1–0. doi:10.1038/s41598-017-04213-7.
28. Yaqoob A, Shahid AA, Salisu IB, Azam S, Ahmed M, Rao AQ. Effects of cry toxins on non-target soil bacteria during a 2-year follow up study. Span J Agric Res. 2019;17(2):303. doi:10.5424/sjar/2019172-14605.
29. Zhang M, Feng M, Xiao L, Song X, Ding G, Yang W. Persistence of Cry1Ac protein from transgenic Bt cotton cultivation and residue returning in fields and its effect on functional diversity of soil microbial communities. Pedosphere. 2019 Feb 1;29(1):114–22. doi:10.1016/S1002-0160(17)60475-2.
30. Fazal A, Wen ZL, Lu YT, Hua XM, Yang MK, Yin TM, Han HW, Lin HY, Wang XM, Lu GH, et al. Assembly and shifts of the bacterial rhizobiome of field grown transgenic maize line carrying mcry1Ab and mcry2Ab genes at different developmental stages. Plant Growth Regul. 2020 May;91(1):113–26. doi:10.1007/s10725-020-00591-7.
31. Pan J, Lv X, Jin D, Bai Z, Qi H, Zhang H, Zhuang G. Developmental stage has a greater effect than Cry1Ac expression in transgenic cotton on the phyllosphere mycobiome. Can J Microbiol. 2019;65(2):116–25. doi:10.1139/cjm-2018-0309.
32. Li X-Y, Lang Z-H, Zhang J, He K-L, ZHU L, Huang D-F. Acquisition of insect-resistant transgenic maize harboring a truncated cry1Ah gene via Agrobacterium-mediated transformation. J Integr Agric. 2014 May 1;13(5):937–44. doi:10.1007/s13693-013-0531-6.
33. Xue J, Liang G, Crickmore N, Li H, He K, Song F, Feng X, Huang D, Zhang J. Cloning and characterization of a novel Cry1A toxin from Bacillus thuringiensis with high toxicity to the Asian corn borer and other lepidopteran insects. FEMS Microbiol Lett. 2008 Mar 1;280(1):95–101. doi:10.1111/j.1574-6968.2007.01053.x.
34. Chen YJ, Guan X, Ren MY. Impacts of transgenic insect-resistant maize HGK60 with Cry1Ah gene on biodiversity of arthropods and weeds in the field. Ying Yong Sheng Tai Xue Bao J Appl Ecol. 2020 Dec 1;31(12):4180–88. doi:10.13287/j.1001-9332.202012.031.
35. Ren M, Chen Y, Guan X. Effects of transgenic insect-resistant maize harboring Cry1Ah on invertebrate biodiversity. J Agro-Environ Sci. 2019;38:2287–96.
36. Vestergaard, G., Schulz, S., and Schöler, A. et al. Making big data smart—how to use metagenomics to understand soil quality. Biol Fertil Soils 53, 479–484 (2017). 10.1007/s00374-017-1191-3
37. Magoč T, Salzberg SL. FLASH: fast length adjustment of short reads to improve genome assemblies. Bioinformatics. 2011 Nov 1;27(21):2957–63. doi:10.1093/bioinformatics/btr507.
38. Edgar RC. UPARSE: highly accurate OTU sequences from microbial amplicon reads. Nat Methods. 2013 Oct;10(10):996–98. doi:10.1038/nmeth.2604.
39. Quast C, Pruesse E, Yilmaz P, Gerken J, Schweer T, Yarza P, Peplies J, Göcker F. The SILVA ribosomal RNA gene database project: improved data processing and web-based tools. Nucleic Acids Res. 2012 Nov 27;41(D1):D590–6. doi:10.1093/nar/gks1219.
40. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, Fierer N, Peña AG, Goodrich JK, Gordon JI, et al. QIIME allows analysis of high-throughput community sequencing data. Nat Methods. 2010 May;7(5):335–36. doi:10.1038/nmeth.f.303.
41. Pétiracq P, Williams A, Cotton A, McFarlane AE, Rolfe SA, Ton J. Metabolite profiling of non-sterile rhizosphere soil. Plant J. 2017 Oct;92(1):147–62. doi:10.1111/tpj.13639.
42. Wang Y, Liu J, Yang F, Zhou W, Mao S, Lin J, Yan X. Untargeted LC-MS-based metabolomics revealed specific metabolic changes in cotyledons and roots of Ricinus communis during early seedling establishment under salt stress. Plant Physiol Biochem. 2021 Jun 1;163:108–18. doi:10.1016/j.plaphy.2021.03.019.
43. Liu B, Zeng Q, Yan F, Xu H, Xu C. Effects of transgenic plants on soil microorganisms. Plant Soil. 2005 Apr;271(1):1–3. doi:10.1007/s11104-004-1610-8.

44. Widmer F. Assessing effects of transgenic crops on soil microbial communities. Green Gene Technol. 2007;22, 207–34.

45. Benjamini Y, Hochberg Y. Controlling the false discovery rate: a practical and powerful approach to multiple testing. J Royal Stat Soc Series B (Methodol). 1995 Jan;57(1):289–300.

46. Filion M. Do transgenic plants affect rhizobacteria populations?. Microb Biotechnol. 2008 Nov;1(6):463–75. doi:10.1111/j.1751-7915.2008.00047.x.

47. Sun, C., Geng, L., Wang, M., Shao, G., Liu, Y., Shu, C., and Zhang, J., 2016. No adverse effects of transgenic maize on population dynamics of endophytic Bacillus subtilis strain B916-gfp. MicrobiologyOpen, 6(1), p. e00404.

48. Saxena D, Stotzky G. Bacillus thuringiensis (Bt) toxin released from root exudates and biomass of Bt corn has no apparent effect on earthworms, nematodes, protozoa, bacteria, and fungi in soil. Soil Biol Biochem. 2001 Jul 1;33(9):1225–30. doi:10.1016/S0038-0717(01)00027-X.

49. Fan, C., Wu, F., Dong, J., Wang, B., Yin, J., and Song, X., 2019. No impact of transgenic cry1le maize on the diversity, abundance and composition of soil fauna in a 2-year field trial. Scientific reports, 9(1), pp.1–9

50. Hamid MI, Hussain M, Wu Y, Zhang X, Xiang M, Liu X, Sessitsch A. Successive soybean-monoculture cropping assembles rhizosphere microbial communities for the soil suppression of soybean cyst nematode. FEMS Microbiol Ecol. 2017 Jan;93(1):fiw222. doi:10.1093/femsec/fiw222.

51. Li C, Tian Q, U Rahman MK, Wu F. Effect of anti-fungal compound phytosphingosine in wheat root exudates on the rhizosphere soil microbial community of watermelon. Plant Soil. 2020 Nov;456(1–2):223–40. doi:10.1007/s11104-020-04702-1.

52. Houlden A, Timms-Wilson TM, Day MJ, Bailey MJ. Influence of plant developmental stage on microbial community structure and activity in the rhizosphere of three field crops. FEMS Microbiol Ecol. 2008 Aug 1;65(2):193–201. doi:10.1111/j.1574-6941.2008.00535.x.

53. Andow DA, Zwahlen C. Assessing environmental risks of transgenic plants. Ecol Lett. 2006 Feb;9(2):196–214. doi:10.1111/j.1461-0248.2005.00846.x.

54. Saxena D, Flores S, Stotzky G. Insecticidal toxin in root exudates from Bt corn. Nature. 1999 Dec;402(6761):480. doi:10.1038/49997.

55. Busov V, Meilan R, Pearce DW, Rood SB, Ma C, Tschaplinski TJ, Strauss SH. Transgenic modification of gai or rgl1 causes dwarfing and alters gibberellins, root growth, and metabolite profiles in Populus. Planta. 2006 Jul;224(2):288–99. doi:10.1007/s00425-005-0213-9.

56. Liu Y, Sun G, Zhong Z, Ji L, Zhang Y, Zhou J, Zheng X, Deng K. Overexpression of AtEDT1 promotes root elongation and affects medicinal secondary metabolite biosynthesis in roots of transgenic Salvia miltiorrhiza. Protoplasma. 2017 Jul;254(4):1617–25. doi:10.1007/s00709-016-1045-0.

57. Raubuch M, Roose K, Warnstorff K, Wichern F, Joergensen RG. Respiration pattern and microbial use of field-grown transgenic Bt-maize residues. Soil Biol Biochem. 2007 Sep 1;39(9):2380–89. doi:10.1016/j.soilbio.2007.04.012.

58. Yan B, Hou J, Cui J, He C, Li W, Chen X, Li M, Wang W. The effects of endogenous hormones on the flowering and fruiting of Glycyrrhiza uralensis. Plants. 2019 Nov;8(11):519. doi:10.3390/plants8110519.

59. Huang XF, Chaparro JM, Reardon KF, Zhang R, Shen Q, Vivanco JM. Rhizosphere interactions: root exudates, microbes, and microbial communities. Botany. 2014;92(4):267–75. doi:10.1139/cjb-2013-0225.

60. Shi J, Gong X, U Rahman MK, Tian Q, Zhou X, Wu F. Effects of wheat root exudates on bacterial communities in the rhizosphere of watermelon. Plant Soil Environ. 2021;67(No. 12):271–28. doi:10.17221/419/2021-PSE.

61. Tan SC, Liu JY, Rahman M, Ma CL, Wu FZ, Zhou XG. Effects of selected root exudates components on soil Pseudomonas spp. community structures and abundances. Allolepathy J. 2020;50:85–93.

62. Velasco AG, Kowalchuk GA, Mañero FJG, Ramos B, Yergeau E, García JAL. Increased microbial activity and nitrogen mineralization coupled to changes in microbial community structure in the rhizosphere of Bt corn. Appl Soil Ecol. 2013 Jun 1;68:46–56. doi:10.1016/j.apsoil.2013.03.010.

63. Cui H, Shu C, Song F, Gao J, Zhang J. Effect of cry1Ah-transgenic maize on community structure of microorganism in rhizosphere soil. Dongbei Nongye Daxue Xuebao. 2011;42:30–38.

64. Donlan RM. Biofilm formation: a clinically relevant microbiological process. Clin Infect Dis. 2001 Oct 15;33(8):1387–92. doi:10.1086/322972.

65. Sun C, Geng L, Wang M, Shao G, Liu Y, Shu C, Zhang J. No adverse effects of transgenic maize on population dynamics of endophytic Bacillus subtilis strain B916-gfp. MicrobiologyOpen. 2017 Feb;6(1):e00404. doi:10.1002/mbo3.404.

66. Ma K, Kou J, Rahman MK, Du W, Liang X, Wu F, Li W, Pan K. Palmitic acid mediated change of rhizosphere and alleviation of Fusarium wilt disease in watermelon. Saudi J Biol Sci. 2021;28(6):3616–23. doi:10.1016/j.sjbs.2021.03.040.

67. Rahman MK, Tan S, Ma C, Wu F, Zhou X. Exogenously applied ferulic acid and p-coumaric acid differentially affect cucumber rhizosphere Trichoderma spp. Commun Struct Abundance Plant Soil Environ. 2020;66:461–67.
68. Sourjik V, Wingreen NS. Responding to chemical gradients: bacterial chemotaxis. Curr Opin Cell Biol. 2012 Apr 1;24(2):262–68. doi:10.1016/j.ceb.2011.11.008.

69. Chang C, Stewart RC. The two-component system: regulation of diverse signaling pathways in prokaryotes and eukaryotes. Plant Physiol. 1998 Jul 1;117(3):723–31. doi:10.1104/pp.117.3.723.

70. Donlan RM. Biofilm formation: a clinically relevant microbiological process. Clinical infectious diseases. 2001 Oct 15;33(8):1387–92

71. Waters CM, Bassler BL. Quorum sensing: cell-to-cell communication in bacteria. Annu Rev Cell Dev Biol. 2005 Nov 10;21(1):319–46. doi:10.1146/annurev.cellbio.21.012704.131001.

72. Seger R, Krebs EG. The MAPK signaling cascade. FASEB J. 1995 Jun;9(9):726–35. doi:10.1096/fasebj.9.9.7601337.

73. Chory J, Wu D. Weaving the complex web of signal transduction. Plant Physiol. 2001 Jan 1;125(1):77–80. doi:10.1104/pp.125.1.77.