Effects of Disease Resistant Genetically Modified Rice on Soil Microbial Community Structure According to Growth Stage

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Abstract

BACKGROUND: This study investigated the effects of rice genetically modified to be resistant against rice blast and rice bacterial blight on the soil microbial community. A comparative analysis of the effects of rice genetically modified rice choline kinase (OsCK1) gene for disease resistance (GM rice) and the Nakdong parental cultivar (non-GM rice) on the soil microbial community at each stage was conducted using rhizosphere soil of the OsCK1 and Nakdong rice.

METHODS AND RESULTS: The soil chemistry at each growth stage and the bacterial and fungal population densities were analyzed. Soil DNA was extracted from the samples, and the microbial community structures of the two soils were analyzed by pyrosequencing. No significant differences were observed in the soil chemistry and microbial population density between the two soils. The taxonomic analysis showed that Chloroflexi, Proteobacteria, Firmicutes, Actinobacteria, and Acidobacteria were present in all soils as the major phyla. Although the source tracking analysis per phylogenetic rank revealed that there were differences in the bacteria between the GM and non-GM soil as well as among the cultivation stages, the GM and non-GM soil were grouped according to the growth stages in the UPGMA dendrogram analysis.

CONCLUSION: The difference in bacterial distributions between Nakdongs and OsCK1 rice soils at each phylogenetic level detected in microbial community analysis by pyrosequencing may be due to the genetic modification done on GM rice or due to heterogeneity of the soil environment. In order to clarify this, it is necessary to analyze changes in root exudates along with the expression of transgene. A more detailed study involving additional multilateral soil analyses is required.

Key words: Genetically modified rice, OsCK1, Soil microbial community
Introduction

The roots of plants secrete a wide variety of chemical compounds to attract microbes to the rhizosphere, resulting in the exclusive selection for the members of the soil microbial community (Huang et al., 2014). The roots release 5-21% of the photosynthate in the form of sugars, amino acids or metabolites, and these substances are used by the microbes in the rhizosphere (Badri et al., 2013b; Chaparro et al., 2013a). Specialized microbial communities are formed in the rhizosphere depending on the plant species, or even the variety, and plant developmental stage. They are also formed by the type and amounts of chemical compounds that comprise root exudates, which are determined by various environmental factors such as soil type, pH and temperature (Badri and Vivanco, 2009). Microbes associated with plants influence the health and growth of plants through a variety of mechanisms (Neumann et al., 2014; Haas and Défago, 2005; Menders et al., 2011; Weller et al., 2002). Numerous studies have been conducted on the interactions between plants and microbes at the plant-microbe level as well as the plant-microbiome level (Huang et al., 2014). Genetically modified (GM) crops might influence the soil microbial community in a direct way by the release of newly produced proteins in root exudates. In addition, these proteins might alter the metabolic pathways within the plant body, and the products of these pathways are released in the root exudates, thus imparting an indirect effect on the soil microbes. Therefore, a meticulous investigation of the potential effects of GM crops on the soil microbial community is required prior to their commercial cultivation.

GM crops resistant to diseases are needed to be developed to prepare for the effects of global warming due to climate change. To date, many GM crops resistant to diseases have been developed, and they are being widely cultivated. However, the cultivation of GM crops is causing many environmental concerns, one of which is the effects of GM crop cultivation on the soil microbial community, as mentioned above. The reason for this concern is that alterations of the soil microbial community can cause additional effects through interactions with plants. The effects of GM crops resistant to diseases on the soil microbial community have been analyzed and reported through various methods. The GM potato that produces exudates that are used by the microbes in the rhizosphere (Badri et al., 2013b; Chaparro et al., 2013a). Specialized microbial communities are formed in the rhizosphere depending on the plant species, or even the variety, and plant developmental stage. They are also formed by the type and amounts of chemical compounds that comprise root exudates, which are determined by various environmental factors such as soil type, pH and temperature (Badri and Vivanco, 2009). Microbes associated with plants influence the health and growth of plants through a variety of mechanisms (Neumann et al., 2014; Haas and Défago, 2005; Menders et al., 2011; Weller et al., 2002). Numerous studies have been conducted on the interactions between plants and microbes at the plant-microbe level as well as the plant-microbiome level (Huang et al., 2014). Genetically modified (GM) crops might influence the soil microbial community in a direct way by the release of newly produced proteins in root exudates. In addition, these proteins might alter the metabolic pathways within the plant body, and the products of these pathways are released in the root exudates, thus imparting an indirect effect on the soil microbes. Therefore, a meticulous investigation of the potential effects of GM crops on the soil microbial community is required prior to their commercial cultivation.

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Choline kinase is commonly distributed in eukaryotes, and it catalyzes the first stage of the Kennedy pathway, which produces phosphatidylcholine as a final product. Phosphatidylcholine mainly consists of phospholipid and is very important for the structure and functions of cell membranes (Gibellini and Smith, 2010; Lee et al., 2007). Many studies have reported that phosphatidylcholine is involved in adaptive responses to droughts, frost, high salinity, and insect damage in vascular plants (Gibellini and Smith, 2010; Lee et al., 2007). Kim et al. (2003) confirmed that the rice choline kinase (OsCK1) gene in rice plays a role in the cold response. Lee et al. (2007) developed GM rice resistant to rice blast and bacterial blight by overexpressing OsCK1 and is currently conducting an environmental risk assessment for its commercialization.

In this study, prior to its commercialization, the effects of GM rice resistant to rice blast and bacterial blight on the soil microbial community were studied. The GM rice and its parental cultivar Nakdong rice were planted in a rice paddy. Soil chemical properties and soil microbial population density at each growth stage were studied, and the microbial community structures of these two soils were comparatively analyzed using pyrosequencing.

Materials and Methods

Site and sampling

The experimental plot for OsCK1 (GM) and Nakdong (non-GM) rice cultivation was constructed in an isolated GMO experimental field at the National Institute of
Agricultural Sciences, Rural Development Administration located in Suwon, Korea. Seeds were sown in a seedling box and then transplanted in June 2012 after 3 weeks to three 4 x 4 m fields. Three replicates of rhizosphere soil samples were collected from each of the fields during the seedling, tillering, and maturity stages. In order to collect rhizosphere soil from the rice, we collected the rice plant with its roots, completely removed the bulk soil, and then collected as much soil as possible that was attached to the roots.

**Soil chemical analyses**

After collection, the soil samples were dried and then passed through a 2-mm sieve for chemical analyses. These analyses were performed according to the methods described by the National Institute of Agricultural Sciences (NIAST, 2000). A pH meter was used to measure the pH of soil suspensions produced by mixing soil and distilled water at a ratio of 1:5. Total nitrogen and carbon compositions were obtained via an elemental analyzer (vario Max CN, Elementar, Germany). Available phosphate was measured using the Lancaster method using a calorimetry assay. Exchangeable cations such as calcium, potassium, magnesium, and sodium were diffused in 1 N ammonium acetate (pH 7.0) and then analyzed using ICP (GBC Integra XL, Australia).

**Viable counts of bacteria and fungi**

The density of soil microbes was assessed by the enumeration of cultured total bacteria and fungi after inoculating soil samples in respective selective media. Ten grams of fresh soil was immersed in 90 mL of sterilized 0.85% NaCl solution and then suspended for 30 min using a shaking incubator (Vision Co., Korea) at 200 rpm. A series of dilutions was made using the suspension, and these dilutions were smeared onto three Petri dishes with R2A agar (Difco, Detroit, USA) containing cycloheximide (0.05 g/L) for bacterial culture and three Petri dishes with R2A agar containing chloramphenicol (0.02%) for fungal culture. The bacteria- and fungi-inoculated media were incubated at 28°C for 2 and 4 d, respectively, prior to counting the number of colonies. The number of microorganisms in each sample was calculated by counting the number of colonies in each of the three Petri dishes and using the average value as the colony-forming unit (cfu·g⁻¹ dry soil).

**Pyrosequencing**

Total DNA was extracted from microorganisms in the soil using a FastDNA Spin Kit (Qbiogen, USA) according to the manufacturer’s manual. The extracted DNA was amplified using primers targeting the V1 to V3 regions of the prokaryotic 16S rRNA gene. The primers used for bacteria were V1-9F (5'-CTATCCCTGGTGCCCTTG GCAGTC-3') and V3-541R (5'-CCATCTCATCCTGCATTCTCAGGC-3': underlining indicates the gene specific part) and V3-541R (5'-CCATCTCATCCTGCATTCTCAGGC-3': the X barcode is uniquely designed for each soil DNA samples, followed by common linker [AC]). PCR amplifications were performed using a C1000 Touch Thermal Cycler (Bio-Rad, CA, USA). A total of 100 ng template DNA was added to the PCR reaction (total volume of 50 μL), which contained Ex Taq buffer, 0.2 mM each dNTP, 0.5 μM each primer, and 2 units Ex Taq (Takara, Otsu, Japan). After an initial denaturation (94°C for 5 min), the PCR reaction was carried out using the touchdown program, which involved 10 cycles of denaturation (94°C for 30 s), annealing (60°C for 45 s), and extension (72°C for 90 s), where the annealing temperature was decreased by 0.5°C for each subsequent cycle. An additional 20 cycles of denaturation (94°C for 30 s), annealing (55°C for 45 s), and extension (72°C for 90 s) were carried out. The amplified products were confirmed by 2% agarose gel electrophoresis and visualized using the Gel Doc system (Bio-Rad). Amplicons were purified using a QIAquick® PCR Purification Kit (Qiagen, CA, USA) and quantified using a PicoGreen® dsDNA Assay Kit (Invitrogen, CA, USA). Equimolar concentrations of each amplicon from the different samples were pooled and purified using an AMPure bead kit (Agencourt Bioscience, MA, USA). Sequencing reactions were performed using a Roche GS FLX Titanium System at ChunLab Inc. (Seoul, Korea) according to the manufacturer’s instructions.

**Pyrosequencing data analyses**

The sequencing reads from the different samples were separated by unique barcodes. The sequences of the barcode, linker, and PCR primers were then removed from both sides of the original sequencing reads. The resultant sequences were subjected to a filtering process where only reads containing 0 to 1 ambiguous base calls (Ns) and 300 or more base pairs were selected for the final bioinformatics analyses. Nonspecific PCR amplicons that showed no match
with the 16S rRNA gene database in a BLASTN search (expectation value of $e^{-5}$) were also removed from the subsequent analyses. Chimeric sequences were detected by the analysis of differences in BLASTN based sequence similarity patterns between the first half and second half of an object sequence. When the first and second halves were differentially identified at the bacterial order level, the sequence was regarded as chimeric, thus eliminated. The obtained sequences were compared and classified using the EzTaxon Database (http://www.ezbiocloud.net) (Chun et al., 2007), which contains 16S rRNA gene sequences of type strains that have valid published names and representative species-level phylotypes of either cultured or uncultured entries in the GenBank public database with complete hierarchical taxonomic classification from the phylum to the species level. The cut-off values used for taxonomic assignments were as follows ($x =$ similarity): species ($x \geq 97\%$), genus ($97\% > x \geq 94\%$), family ($94\% > x \geq 90\%$), order ($90\% > x \geq 85\%$), class ($85\% > x \geq 80\%$) and phylum ($80 > x \geq 75\%$). If the similarity was lower than the specific cut-off value, the sequence was assigned as ‘unclassified’ (Chun et al., 2007; Unno et al., 2010; Park et al., 2012). For the taxonomic affiliation, if the scientific name for a taxon was unknown, the known name in the nomenclature was written first, and then, a suffix was added at the end of the name after an underscore (e.g. if the class name was unknown, a ‘c’ was written after the phylum name. ‘Acidobacteria_c’; c = class, o = order, f = family, g = genus, and s = species) (Kim et al., 2012).

A rarefaction curve that shows the increase in the ratio of operational taxonomic units (OTUs) to the analyzed sequence number was constructed based on the CD-HIT (Li et al., 2001; Fu et al., 2012) and Mothur software packages (Schloss et al., 2009). The number of OTUs was calculated from the sequence group that showed 97% sequence homology based on the taxonomy-based de novo clustering algorithm (Lee et al., 2012), and this number was used to calculate the Shannon diversity index ($H$), which is a measure of diversity and evenness, and the richness estimators ACE and Chao 1. In order to analyze the sharedness of the species shared between the soils, a single source tracking analysis was conducted using the CLcommunity™ for Microbial Community Analysis program (ChunLab, Seoul, Korea) by ChunLab. The Nakdong rice soil at the seedling stage was chosen for the sink, whereas the rest of the soil samples, that is, the all stages of OsCK1 rice soils and tillering and maturity stage Nakdong rice soils were the source. The sharedness between the sink and source was calculated using the following formula: Sharedness (%) $= [(s/a) \times 100 + (s/b) \times 100]/2$, where ‘s’ represents the number of sequences found in both samples, ‘a’ represents the total number of source samples, and ‘b’ represents the number of sink sample species. The similarity between each pair of communities was estimated using the Fast UniFrac web interface (Hamedy et al., 2010) and visualized using the unweighted pair group method with an arithmetic mean (UPGMA) dendrogram.

## Results

### Chemical characteristics of soil samples

Because differences in the soil chemical composition can affect soil microbial communities, soil pH, available phosphate, electrical conductivity, total nitrogen, organic matter, and cations were analyzed to identify possible differences between the rhizosphere soil of OsCK1 rice and that of Nakdong rice (Table S1).

| Growth Stage | Sample | pH  | P$^+$ (mg/kg) | EC$^b$ (dS/m) | T-N$^c$ (%) | OM$^d$ (g/kg) | K$^+$ | Ca$^{2+}$ | Mg$^{2+}$ | Na$^{+}$ |
|--------------|--------|-----|--------------|--------------|-------------|---------------|-------|----------|----------|---------|
| Seedling     | ND     | 6.7±0.02 | 96.0±0.07 | 0.3±0.01 | 0.2±0.01 | 20.6±0.2 | 0.7±0.01 | 7.8±0.25 | 2.1±0.01 | 0.3±0.00 |
|              | OsCK1  | 6.6±0.01 | 93.3±0.06 | 0.3±0.00 | 0.2±0.00 | 18.7±0.4 | 0.5±0.01 | 8.2±0.10 | 2.3±0.02 | 0.2±0.00 |
| Tillering    | ND     | 6.3±0.02 | 83.4±1.26 | 0.3±0.02 | 0.2±0.01 | 19.3±0.31 | 0.5±0.01 | 7.9±0.15 | 2.1±0.12 | 0.2±0.00 |
|              | OsCK1  | 6.3±0.03 | 80.2±1.66 | 0.3±0.01 | 0.2±0.01 | 18.7±0.09 | 0.4±0.01 | 8.1±0.34 | 2.2±0.04 | 0.2±0.00 |
| Maturity     | ND     | 6.4±0.01 | 69.1±1.66 | 0.2±0.00 | 0.2±0.00 | 19.2±0.26 | 0.4±0.00 | 7.2±0.20 | 2.1±0.12 | 0.2±0.01 |
|              | OsCK1  | 6.6±0.01 | 70.2±2.83 | 0.3±0.00 | 0.2±0.00 | 18.6±0.30 | 0.4±0.01 | 7.4±0.12 | 2.2±0.12 | 0.2±0.00 |

$^a$: Available phosphate; $^b$: Electrical conductivity; $^c$: Total nitrogen; $^d$: Soil organic matter; ND, Nakdong; OsCK1, disease-resistant transgenic rice. There was no significant difference between Nakdong and OsCK1 rice soils according to growth stage (t-test, $p < 0.05$).
There was no significant difference in soil pH between OsCK1 (pH 6.3–6.6) and Nakdong rice (pH 6.3–6.7). In this study, the available phosphate was 69–96 mg kg⁻¹ for OsCK1 rice and 70–93 mg kg⁻¹ for Nakdong rice soil, which was not significantly different. The soil electrical conductivity for OsCK1 was 0.2–0.3 dS m⁻¹ and that of Nakdong rice was 0.3 dS m⁻¹ throughout the growth stages, and the total nitrogen content was similar between the two soils. Moreover, there were no significant differences in either organic matter or cations, such as K⁺, Ca²⁺, Mg²⁺, or Na⁺, between OsCK1 and Nakdong rice soils.

Bacterial community comparative analysis

In this study, we did not observe significant difference in bacterial and fungal quantities between OsCK1 and Nakdong rice soil microbial communities (Table S2). The total number of pyrosequencing reads was 98,839, among which 51,927 were high-quality reads. The results for the OTU richness patterns from the Chao 1 and ACE analyses showed that the pattern for the Nakdong soil at the seedling stage was the highest, and ACE for the Nakdong rice soil and Chao 1 for the OsCK1 soil at the tillering stage were the lowest (Table 1). The results for the Shannon index values indicated that the diversity of the Nakdong rice soil at the seedling stage and OsCK1 soil at the maturity stage were the highest and those of the Nakdong and OsCK1 rice soils at the tillering stage were the lowest (Table 1).

| Growth Stage | Sample | Total No. of Reads | Observed phylotypes (OTUs) | Average Length (bp) | Ace | Chao1 | Shannon index | Coverage (%) |
|--------------|--------|--------------------|----------------------------|---------------------|-----|-------|---------------|--------------|
| Seedling     | ND     | 13,833             | 8,845                      | 463.4±0.5           | 18138.3±9284.2 | 9837.5±4440.1 | 7.6±0.4       | 51.5±7.2     |
|              | OsCK1  | 12,865             | 8,096                      | 464.2±0.4           | 16145.2±9334.4 | 9084.9±4876.0 | 7.5±0.4       | 52.7±10.0    |
| Tillering    | ND     | 12,298             | 6,837                      | 454.7±5.5           | 11669.9±273.9  | 6492.0±588.1  | 7.3±0.1       | 58.9±5.0     |
|              | OsCK1  | 14,943             | 7,103                      | 456.6±2.3           | 12257.3±1906.3 | 6411.9±57.1  | 7.3±0.1       | 63.1±12.4    |
| Maturity     | ND     | 22,588             | 10,586                     | 467.9±0.6           | 17304.1±9565.1 | 9764.9±5757.9 | 7.5±0.5       | 64.8±5.7     |
|              | OsCK1  | 22312              | 10,460                     | 467.7±0.4           | 17253.6±5527.5 | 9710.6±3927.7 | 7.6±0.4       | 63.6±8.5     |

Values indicate the colony-forming unit (cfu)/g wet weight ± standard deviation from three replications. There was no significant difference between Nakdong and OsCK1 rice soils according to growth stage (t-test, p < 0.05). ND, Nakdong; OsCK1, disease-resistant transgenic rice.

Estimates of the Shannon index were obtained based on 3% differences in DNA sequence alignments. Values of Ace, Chao1, Shannon and coverage indicate the mean ± standard deviation from three replications. There was no significant difference between Nakdong and OsCK1 rice soils (t-test, p < 0.05). ND, Nakdong; OsCK1, disease-resistant transgenic rice.

Table S2. Average number of colony-forming units in the rhizosphere soil

| Growth Stage | Sample | Total No. of Reads | Observed phylotypes (OTUs) | Average Length (bp) | Ace | Chao1 | Shannon index | Coverage (%) |
|--------------|--------|--------------------|----------------------------|---------------------|-----|-------|---------------|--------------|
| Seedling     | ND     | 13,833             | 8,845                      | 463.4±0.5           | 18138.3±9284.2 | 9837.5±4440.1 | 7.6±0.4       | 51.5±7.2     |
|              | OsCK1  | 12,865             | 8,096                      | 464.2±0.4           | 16145.2±9334.4 | 9084.9±4876.0 | 7.5±0.4       | 52.7±10.0    |
| Tillering    | ND     | 12,298             | 6,837                      | 454.7±5.5           | 11669.9±273.9  | 6492.0±588.1  | 7.3±0.1       | 58.9±5.0     |
|              | OsCK1  | 14,943             | 7,103                      | 456.6±2.3           | 12257.3±1906.3 | 6411.9±57.1  | 7.3±0.1       | 63.1±12.4    |
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|              | OsCK1  | 22312              | 10,460                     | 467.7±0.4           | 17253.6±5527.5 | 9710.6±3927.7 | 7.6±0.4       | 63.6±8.5     |
showed that Chloroflexi was present at about twice the level of Proteobacteria at the maturity stage. Among phyla showing a relative abundance over 1% in the seedling stage of Nakdong rice and OsCK1, Firmicutes had a significant difference of distribution rate between Nakdong and OsCK1 rice soil (p<0.05). In the tillering stage among phyla showing a relative abundance of over 1%, Chloroflexi, Proteobacteria, Nitrospira, Bacteroidetes, Planctomycetes, and Chlorobi had a significant difference between Nakdong and OsCK1 rice soil (p<0.05). In the maturity stage among phyla showing a relative abundance of over 1%, cyanobacteria had over 20% higher relative abundance in OsCK1 rice soil compared to that of Nakdong rice soil.

At the genus level, 2,442 genera were observed and 197 genera were found in all soils (Table 2).
Among genera showing a relative abundance of over 1%, the number of genus with significant difference in the relative abundance between Nakdong and OsCK1 rice soils was 2 in the seedling stage, 5 in the tillering stage, and 3 in the maturity stage. The genera with the highest relative abundance difference between Nakdong and OsCK1 rice soils were GU454901_g (Nakdong: 0.8%, OsCK1: 0.4) in the seedling stage, AM745150_f_uc (Nakdong: 1.51%, OsCK1: 2.35) in the tillering stage and Solibacter (Nakdong: 0.41%, OsCK1: 0.61) in the maturity stage.

The sharedness of the OTUs that comprise the microbial community structure was analyzed through single source tracking analysis (Table 3). When the Nakdong rice soil at the seedling stage was the sink and the rest were the source, the sharedness was 99.6%–99.9% at the phylum level, 96.9–99.8% at the class level, 91.9–99.6% at the order level, 85.8–98.8% at the family level, 79.8–96.4% at the genus level, and 45.3–76.7% at the species level. The UPGMA dendrogram analysis examining the overall similarity between the two soil bacterial communities showed that the soils grouped by time period (Fig. 2).

**Table 3. Sharedness analysis between sink and source soil**

| Phylogenetic Level | SS ND | SS ND | SS OsCK1 | SS OsCK1 | TS ND | TS ND | TS OsCK1 | TS OsCK1 | TS OsCK1 | TS OsCK1 | TS OsCK1 | TS OsCK1 | MS ND | MS ND | MS ND | MS ND | MS OsCK1 | MS OsCK1 | MS OsCK1 | MS OsCK1 | MS OsCK1 |
|-------------------|-------|-------|----------|----------|-------|-------|----------|----------|----------|----------|----------|----------|-------|-------|-------|-------|----------|----------|----------|----------|----------|
| Phylum            | 99.93 | 99.58 | 99.69    | 99.84    | 99.71 | 99.64 | 99.69    | 99.71    | 99.71    | 99.77    | 99.77    | 99.72    | 99.72   | 99.71   |
| Class             | 99.83 | 97.64 | 99.33    | 97.51    | 99.3  | 99.14 | 99.17    | 99.13    | 99.24    | 98.91    | 99.04    | 97.35   | 99.04   | 98.91   |
| Order             | 99.61 | 92.11 | 97.8     | 98.03    | 91.9  | 98.18 | 98.21    | 97.83    | 97.61    | 97.5    | 97.82    | 97.82   | 97.95   | 92.07   | 97.67   |
| Family            | 98.75 | 86.86 | 94.31    | 87.97    | 94.9  | 95.14 | 95.05    | 93.28    | 93.28    | 94.31    | 94.31   | 94.58   | 94.92   | 93.44   | 95.39   |
| Genus             | 96.35 | 79.97 | 85.62    | 87.35    | 97.7  | 86.24 | 87.26    | 87.77    | 86.03    | 86.03    | 85.88   | 87.38   | 86.71   | 81.44   | 88.33   |
| Species           | 76.67 | 51.38 | 51.08    | 57.37    | 45.28 | 52.68 | 62.29    | 53.22    | 50.95    | 50.95    | 52.37   | 64.25   | 55.92   | 50.8    | 55.49   | 65.17   |

* Sink: SSND#1 soil
* Source: SSND#2, SSND#3, SSOsCK1#1, SSOsCK1#2, SSOsCK1#3, TSND#1, TSND#2, TSND#3, TSOsCK1#1, TSOsCK1#2, TSOsCK1#3, MSND#1, MSND#2, MSND#3, MSOsCK1#1, MSOsCK1#2, and MSOsCK1#3 soils

SS, seedling stage; TS, tillering stage; MS, maturity stage; ND, Nakdong; OsCK1, disease-resistant transgenic rice

![Fig. 2. A UPGMA dendrogram based on the unweighted pair-wise Fast UniFrac distances between the bacterial communities in the ND and OsCK1 rice soil samples. SS, seedling stage; TS, tillering stage; MS, maturity stage; ND, Nakdong; OsCK1, disease-resistant transgenic rice.](image-url)
**Discussion**

**Analysis of the Soil Chemistry**

Soil chemistry can influence the soil microbial community structure, and many reports are available concerning the topic. The composition and diversity of the soil bacterial community are often closely related to soil pH (Jenkins et al., 2009; Lauber et al., 2009). A study that analyzed the differences in microbial community structures according to pH levels through pyrosequencing at the continental level found that the composition and diversity of the soil microbial community had a positive correlation with pH, and the overall difference in the community structure was caused by the prevalence of Acidobacteria, Actinobacteria, and Bacteroidetes (Lauber et al., 2009). In the present study, the pH of the OsCK1 and Nakdong rice soils ranged from 6.3–6.7 through all stages, which is higher than the average acidity (pH 5.6) of Korean soils for rice (Jung et al., 1998). One study reported that the addition of phosphate significantly increased the root biomass, shoot biomass, soil pH and microbial activity and caused noticeable changes in the fungal community and bacterial community structures as a result (Rooney and Clipson, 2009). In this study, the values for both soils ranged from 69–96 mg kg$^{-1}$ through all stages. Considering the average density of available phosphate of 95 mg•kg$^{-1}$ for Korean paddy soils, the available phosphate density decreases as development nears the maturity stage (Jung et al., 1998). In another study that investigated the effects of the electrical conductivity of the soil from a protected strawberry cultivation on the microbial ecology, it was found that the high electrical conductivity level soil had high values for soil microbial biomass, total bacteria, gram-negative and gram-positive bacteria, actinomycetes, fungi, and arbuscular mycorrhizal fungi (Lee et al., 2011). It was reported that changes in soil chemistry due to organic amendment caused changes in the total phospholipid fatty acid content and bacteria:fungi ratio (Ng et al., 2014a). In this study, no significant differences were observed between the soil chemistry results of the OsCK1 and Nakdong rice soils, indicating that the cultivation of OsCK1 rice did not alter the soil chemistry.

**Analysis of microbial community structure through pyrosequencing**

To date, many disease-resistant GM crops have been commercialized. Among these are transgenic potato resistant to potato virus Y (Solanum tuberosum, event name: RBMT15-101, RBMT21-129, RBMT21-350, RBMT22-82, SEMT15-02, and SEMT 15-15), transgenic squash resistant to zucchini yellow mosaic potyvirus and watermelon mosaic potyvirus (Cucurbita pepo, event name: ZW20), transgenic sweet pepper resistant to cucumber mosaic cucumovirus (Capsicum annuum, event name: PK-SP01), and transgenic papaya (Azad et al., 2014) resistant to PRSV. Disease-resistant transgenic crops can have a direct effect on the soil microbial community through the root exudates released into the rhizosphere, and thus, numerous studies have been conducted on these effects. For example, T4 lysozyme-producing transgenic potato (Heuer et al., 2002; Lottmann et al., 1999; Ahrenholtz et al., 2000; Lottmann and Berg, 2001), cecropin B-producing transgenic potato (Rasche et al., 2006; Sessitsch et al., 2003), transgenic plants that produce pathogenesis-related proteins (Vierheilig et al., 1993, 1995; Yang et al., 2002), and transgenic plants that cause a defense reaction by inducing systemic acquired resistance (Heuer et al., 2002; Lottmann et al., 1999; Sessitsch et al., 2003; Ahrenholtz et al., 2000; Lottmann and Berg, 2001; Vierheilig et al., 1995; Yang et al., 2002; Glandorf et al., 1997; Medina et al., 2003) were analyzed using amplified ribosomal DNA restriction analysis, ARISA, T-RFLP, CFU, CLPP, denaturing gradient gel electrophoresis (DGGE), fatty acid methyl ester, and repetitive extragenic palindromic PCR (rep-PCR) methods. It was found that could be effects, no effects, or minor effects on the soil microbial community structure in the rhizosphere. PRSV is a highly destructive disease for papaya production, and there have been various attempts to develop GM papaya resistant to PRSV. Coat protein-mediated protection, RNA-silencing, and replicase gene-mediated transformation have been used to produce GM papaya resistant to PRSV, which has been successfully commercialized and is now in production in many countries (Azad et al., 2014). A study was conducted on the effects of transgenic papayas expressing the replicase gene on the soil microbiota, and it was reported that the transgenic papayas can alter the soil chemistry, enzyme activity, and microbial communities (Wei et al., 2006). In addition to disease-resistant transgenic crops, transgenic crops resistant to harmful insects continue to be studied, including Cry1Ab GM maize, cotton, and rice;
transgenic Cry1Ab/Cry1Ac rice; transgenic Cry1Ac
eggplant and turnip; transgenic Cry1F maize;
transgenic Cry2Ab maize; transgenic Cry2Ab cotton;
transgenic Cry3Bb1 maize; and transgenic
Cry34/35Ab1 maize (Turrini et al., 2015). Their soil
microbial communities were analyzed using ARISA,
DGGE, quantitative PCR, CLPP, microarray, PCR-
RFLP, single-strand conformation polymorphism, T-
RFLP, RNA-stable isotope probing, phospholipid fatty
acid analysis, and pyrosequencing. The effects were
constant, transient, or none. Barriuso et al. (2012)
analyzed the soil microbial community structure of
the rhizosphere after 4 years of transgenic Cry1Ab Bt
maize production using pyrosequencing and
concluded that the alteration of the soil microbial
community structure in the rhizosphere was due to
climatic factors rather than the Bt gene. In the present
study, rhizosphere soils from transgenic OsCK1 rice
resistant to rice diseases and the parental cultivar
Nakdong rice were also analyzed by pyrosequencing,
and the similarity between the soils was investigated
using UPGMA dendrogram analysis. It was found
that the GM and non-GM soils were grouped together
at each stage and not grouped only by themselves.
The bacteria not present in both soils at each stage
were taxonomically analyzed, and the bacterial
populations from the two soils did not match 100%.
The similarity of the Nakdong rice at the seedling
stage as the sink for the rest of the soils was studied
by rank through single source tracking analysis. The
soil at the seedling stage showed over 90% similarity
with the other soils at the family level and over,
whereas it fell to over 80% similarity at the genus
level and to approximately 50% at the species level.
These results indicate differences not only in the soil
microbial community structure between the Nakdong
rice and OsCK1 soils but also in the soil microbial
community structure according to the stages. Among
the three stages, tillering stage showed the highest
difference in the relative abundance of microbial taxa
between Nakdong and OsCK1 rice soils.

As the tillering stage is the most active rice
growing phase, various products different from the
seedling or maturity stages are released through the
root exudation causing the difference in the microbial
distribution between Nakdong and OsCK1 rice soils
due to change in rhizosphere soil environment
(Aulakh et al., 2001). As root exudates, organic acids
and malic acid showed the highest concentrations
followed by tartaric, succinic, citric and lactic acids.
The exudation of organic acids has been analyzed to
replace the release of sugar according to the growth
of rice (Aulakh et al., 2001).

The difference in bacterial distribution rate
between Nakdong and OsCK1 rice soils at each
phylogenetic level detected in metagenome analysis
by pyrosequencing may be due to the difference
between GM and non-GM, or due to heterogeneity of
the soil environment. In order to clarify this, it is
necessary to analyze the changes of primary and
secondary metabolite pathways in the plant body
occurred by introducing transgene; to analyze
biochemical components of root exudates released
from plant roots; and to confirm whether or not
transgene derived proteins is contained in the root
exudates. If there is a difference in microbial
community composition or distribution rate between
GM and non-GM soils in the results, it is necessary to
analyze the function of microorganisms showing the
difference and the correlation with transgene.

Plants not only release primary and secondary
metabolites but also proteins as root exudates (Basu
et al., 1994, 1999; Charmont et al., 2005; Hu et al.,
2018). However, knowledge on how these released
proteins interact with rhizosphere microorganisms is
extremely limited. In the case of transgenic plants,
not only the function of target proteins but also the
changes in root exudates due to changes in plants
after expression of transgene should be compared
with root exudates of non-transgenic plants. To this
end, a combined approach of transcriptomics and
proteomics tools may be helpful in understanding
these interactions. In addition, long-term studies
should be conducted as to whether the changes in
root exudates due to the introduction of transgens
are sustained in soils and affect soil microorganisms.

Note

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