Salicylic acid remodeling of the rhizosphere microbiome induces watermelon root resistance against *Fusarium oxysporum* f. sp. *niveum* infection

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**Introduction**

Watermelon (*Citrullus lanatus*) is an important horticultural crop worldwide (Everts and Himmelstein, 2015). However, the commercial cultivation of this fruit is severely threatened by the soil-borne fungus *Fusarium oxysporum* f. sp. *niveum* (FON), which causes *Fusarium* wilt, a disease that leads to a significant decline in crop quality and yield (Xu et al.,...
Salicylic acid (SA), a phytohormone present in plants and some microbes, is reported to be an important signaling molecule that induces plant resistance to diseases (Yang et al., 2015; Zhang and Li, 2019). With the deciphering of the watermelon genome, Lü et al. found via gene ChiP transcription analysis that the phenylalanine ammonia lyase (PAL) gene plays an important role in the lignin metabolic pathway of resistance to Fusarium wilt (Lü et al., 2011). Lv reported that the intercropping of wheat and watermelon with FON induced SA synthesis in the watermelon plant to enhance its resistance to Fusarium wilt (Lv et al., 2018). Furthermore, our research demonstrated the important role of SA in regulating watermelon resistance against Fusarium wilt, with results indicating that the significantly expressed C. lanatus PAL (CIPAL) and non-pathogen-related (NPR) genes play key roles in SA synthesis and signal transduction in this plant species (Zhu et al., 2022a,b).

Recently, increasing evidence has suggested that the rhizosphere microbiome plays critical roles in promoting plant growth and health, such as enhancing nutrient uptake by the host plant and increasing its resistance against pathogen attack (Levy et al., 2017; Berendsen et al., 2018). The establishment of plant–rhizosphere microbiome interaction is a highly coordinated event influenced by the host plant and soil. For instance, the plant immune system shapes the microbiome, which, in turn, can increase the plant’s immune capacity (Heintz and Mair, 2014; Levy et al., 2017; Aleklett et al., 2018). Notably, our previous studies have demonstrated the important role of the soil microbial community structure in controlling the occurrence of Fusarium wilt in watermelons (Zhu et al., 2018, 2019). For instance, our results indicated that the presence of beneficial microbes, such as Rhodanobacter, Pseudomonas, Sphingomonas, and Herbaspirillum, is important for the prevention of watermelon disease (Zhu et al., 2020).

Consistently, more researchers have begun to notice that rhizosphere microbiome composition is influenced by an array of plant-derived metabolic substances (Trivedi et al., 2020; Liu et al., 2021). Additionally, the essential role of SA in recruiting specific rhizosphere microbiomes has been previously reported. For instance, Trivedi et al. showed that SA affected the abundance of specific bacterial groups in the roots via a combination of direct and indirect effects (Trivedi et al., 2020). However, the mechanism by which watermelon plants affect microbiome assembly and the impact of this interconnectedness on plant and microbiota functions remains unclear. Chen et al. found that systemic accumulation of SA could affect microbiome assembly in the rhizosphere of Arabidopsis plants after foliar infection by pathogens (Chen et al., 2020). Therefore, through metagenomic sequencing and relevant physiological and biochemical analyses, we aimed to explore the effect of exogenous SA in remodeling the watermelon rhizosphere microbiome to induce resistance against FON infection. Our research findings provide a new strategy for controlling watermelon Fusarium wilt from the perspective of environmental ecology and have significant value for promoting sustainable agricultural development.

Materials and methods

Experimental process and sampling

The experiment was conducted in the city of Changsha (112°58′42″E, 28°11′49″N), Hunan Province, China. The soil used for planting was sandy loam, collected from our field experiment at the Gaoqiao Scientific Research Base of the Hunan Academy of Agricultural Sciences in Changsha (Zhu et al., 2020). Our previous studies found that the high content of pathogens in the soil under a continuous cropping system leads to SA accumulation in the plant (Zhu et al., 2022a,b). Therefore, we sterilized the soil to make them as same background before use (LDZM-80KCS-3 vertical pressure steam sterilizer, ZHONGANG, Shanghai, China) to avoid errors and study how SA can improve watermelon immune resistance at the early stage after FON infection. The watermelon variety used was Zaojia 8,424 (Xinjiang Farmer Seed Technology Co., Ltd., Urumqi, China), which is the main cultivar on the Chinese market. The watermelon seedlings were cultivated in seedling pots containing peat, perlite, and vermiculite (6:3:1) and grown in a biochemical incubator (LRH-300, ZHUJIANG, Taihong, Shaoguan, China) that was set at 25°C during 16 h of light and 18°C during 8 h of darkness. After 30 days, each plant was transplanted into separate pots. When the seedlings were at the two-leaf stage, 5 ml of 100 μM exogenous SA (Sigma-Aldrich LLC., Merck KGaA, Darmstadt, Germany) was incorporated into the root zone of each plant, and another 5 ml was added 24 h later. The Fusarium strain FON was firstly incubated in the dark for 7 days on a PDA plate at 28°C. Then, a bam plug was selected from the PDA plate and placed into 300 ml of potato dextrose broth in a flask, before propagation on a rotary shaker at 200rmp at 26–30°C. Two days later, 5 ml FON (1 × 10^6 conidia/mL) was added to the root zone of each plant (Li et al., 2019).

The plants were divided into the following three groups: S, mock-inoculation control (H₂O); SA, 100 μM exogenous SA + FON; and SF, FON only. We set six different sampling times before and after the different treatments: 0 days post-inoculation (dpi) (before treatment), 12 h post-inoculation (hpi), 1 dpi, 3 dpi, 5 dpi, and 7 dpi. Sampling was stopped at 7 dpi when the disease symptoms (yellowing and wilting) started to appear. We selected 10 watermelon plants as one replicate and set three independent replicates (30 plants) for each sample group, at six different sampling times with every three treatments (16 groups). Therefore, 480 plant samples were collected from 480 pots.

The soil samples were designated S0 (mock-inoculation control, before treatment), S3 (mock-inoculation control, 3 dpi), SF3 (FON inoculation control, 3 dpi), S (exogenous SA inoculation control, before treatment), S3 (exogenous SA inoculation control, 3 dpi), SF3 (FON inoculation control, 3 dpi).

Abbreviations: ARO, antibiotic resistance ontology; dpi, days post-inoculation; FON, Fusarium oxysporum f. sp. niveum; hpi, hours post-inoculation; MDA, malondialdehyde; PAL, phenylalanine ammonia lyase; POD, peroxidase; SA, salicylic acid.

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Determination of the watermelon plant root morphology and physiological and biochemical indexes

First, the root morphology of each plant was captured using a digital camera. The roots were then cut with sterilized scissors, and their fresh weights were measured using an electronic balance. Thereafter, the roots were placed in sterilized 5 ml centrifuge tubes to test their peroxidase (POD) and phenylalanine ammonia lyase (PAL) activities and malondialdehyde (MDA) content. The POD and PAL activities were analyzed using a BC0095 peroxidase assay kit and BC0215 PAL test kit (Beijing Solarbio Science & Technology Co., Ltd., Beijing, China), respectively, whereas the MDA content was determined using the thiobarbituric acid method using a Rongsheng BC0025 MDA assay kit (Beijing Solar-bio Science & Technology Co., Ltd., Beijing, China), whereas the MDA content was determined using the thiobarbituric acid method using a Rongsheng BC0025 MDA assay kit (Beijing Solar-bio Science & Technology Co., Ltd., Beijing, China), respectively, whereas the MDA content was determined using the thiobarbituric acid method using a Rongsheng BC0025 MDA assay kit (Beijing Solar-bio Science & Technology Co., Ltd.).

The disease incidence was calculated as follows:

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\text{Disease incidence (\%)} = \frac{\text{no. of infected plants}}{\text{total number of plants surveyed}} \times 100.
\]

Soil DNA library preparation

A total of 1 μg DNA per sample was used for sequencing. In brief, DNA sequences 350bp in size were fragmented by sonication, and the fragments were then end-polished, A-tailed, and ligated with the full-length adaptor for Illumina sequencing followed by PCR amplification. The PCR products were purified (AMPure XP system), and libraries were prepared and analyzed for size distribution using the Agilent2100 Bioanalyzer. Three biological replicates per sample were analyzed. Raw data were obtained using the Illumina NovaSeq 6000 sequencing platform.

Metagenomic sequencing

Readfq V8 was used to acquire the clean data for subsequent analysis against the host database using the Basic Local Alignment Search Tool (BLAST), which uses Bowtie 2.2.4 as default software to filter the reads that are of host origin. After all the reads that were not used in the forward step were combined, SOAPdenovo was used to generate the mixed assembly using the same parameters as those applied for the single assembly. The Scaffigs (≥500bp) assembled from both the single and mixed assemblies were used to predict the open reading frames (ORFs) using MetaGeneMark software, and sequences shorter than 100nt were filtered from the predicted result with default parameters. For ORF prediction, sequence redundancy was reduced using the Cluster Database at High Identity with Tolerance software, and a unique initial gene catalog was obtained. The clean data of each sample were mapped to the initial gene catalog using Bowtie 2.2.4 with the following parameter settings: --end-to-end, -sensitive, -I 200, and -X 400. Genes with two or fewer reads in each sample were filtered, and the gene catalog (UniGene database) obtained was used for subsequent analysis.

The abundance of each gene in each sample was statistically analyzed based on the number of mapped reads and their lengths. The basic information statistics, core- and pan-genomic analyses, correlation analysis of the samples, and Venn diagram analysis of the number of genes were all based on the abundance of each gene in each sample in the gene catalog. DIAMOND software was used to blast the UniGene database to the sequences of bacteria, fungi, archaea, and viruses, which were all extracted from the non-redundant database of the National Center for Biotechnology Information (NCBI). Finally, the clean reads were deposited in the NCBI Sequence Read Archive database (Accession number: PRJNA707127).

Assembly of the Core rhizosphere communities, common functional databases used, and resistance gene annotation

To determine the dynamic changes in the dominant soil microbial communities during all three sampling times in the different treatment groups, we used community bar-plot analysis to identify the 10 most abundant soil microbial communities at both the phylum and genus levels. Moreover, we used multiple t-tests to compare significant differences in soil microbial communities at 3 and 7 dpi, both at the phylum and genus levels, after SA application. To elucidate the molecular mechanism by which the rhizosphere microbial community cooperates to induce plant resistance against *Fusarium* wilt, we blasted the unique genes against various functional annotation databases such as the Kyoto Encyclopedia of Genes and Genomes (KEGG), Evolutionary Genealogy of Genes: Non-supervised Orthologous Groups (eggNOG), Carbohydrate-Active Enzymes (CAZY), and the Comprehensive Antibiotic Resistance Database (CARD) to determine their abundance and analyzed them statistically with a visual display.
KEGG, eggNOG, and CAZy databases were used for functional annotation of the resistance genes. The Unigenes were aligned to the CARD database using Resistance Gene Identifier software with default parameter settings. The relative abundance of the antibiotic resistance ontology (ARO) cluster, abundance bar charts, abundance cluster heatmaps, and differences in the number of resistance genes between soil groups were displayed according to the alignment results. Similarly, the distribution of resistance genes in each sample and analyses of the species attribution of those genes and their resistance mechanisms were also investigated.

**Statistical analysis**

Statistical analysis was performed using GraphPad Prism 9 (GraphPad Software, San Diego, CA, United States). All values are expressed as mean ± standard error ($n=3$). The differences...
between the groups were tested using an analysis of similarities. Figures were constructed using Microsoft Office 2010 (Microsoft Corporation, Redmond, WA, United States).

**Results**

**Effectiveness of salicylic acid treatment in controlling *Fusarium* wilt of watermelon**

To clarify whether exogenous SA treatment can control *Fusarium* wilt disease in watermelon, we first compared the root phenotypes after exposure to exogenous SA + FON or FON alone at five different sampling times. The SA group had more fibrous roots than the SF group did. The roots in the SF group began to turn yellow at 3 dpi, and obvious plant yellowing and wilting symptoms were observed at 7 dpi (Figure 1). Furthermore, there was a significant difference in the fresh weight of the roots between the samples at 1, 3, and 7 dpi. For instance, the root fresh weight of the SA group was 2.5 times heavier than that of the SF group at 7 dpi (Figure 2A). The incidence of *Fusarium* wilt after SA application was significantly lower (by ~20%) than that in plants not treated with phytohormones (Figure 2B). Moreover, the MDA content in the SF group first increased from 12 hpi onward, then declined at 3 dpi and finally increased significantly again at 7 dpi. However, we noticed an increase in MDA content at 3 dpi in both SA and S groups (Figure 2C). Similarly, POD activity was significantly increased at 3 dpi in the SA and S groups, which indicated that this sampling time might be a key time point for activating resistance defenses in watermelon plants (Figure 2D). Moreover, there were significant enhancements in PAL activity in the SA group at 3 dpi and 7 dpi, whereas the enzyme activity was significantly lower in the SF group at 7 dpi (Figure 2E), which confirmed the symptoms and disease incidence results. Therefore, the results from plant root physiological and biochemical studies indicate that exogenous SA treatment can effectively reduce the incidence of *Fusarium* wilt in watermelon.

**Sequencing and metagenome assembly**

We selected three sampling times to compare the dynamic changes in the dominant soil microbial communities between the different treatments. Gene sequencing analysis revealed that the ORFs were approximately 200–600 nt in length (Supplementary Figure S1A). According to the results of the quality control analysis, the evenness of the number of ORFs observed in the samples tended to be consistent (Supplementary Figure S1B). The Venn diagram showed that there were 624,730 overlapping genes among all groups.
of samples, and there were unique genes in each group (Supplementary Figure S1C).

Dynamic changes in the soil microbial community structure

The top 10 phyla in all samples were Proteobacteria, Actinobacteria, Bacteroidetes, Gemmatimonadetes, Firmicutes, Acidobacteria, Verrucomicrobia, Deinococcus-Thermus, Elusimicrobiota, and Chloroflexi (Figure 3A). The top 10 genera were Rhodanobacter, Micromonaspora, Massilia, Flavobacterium, Cellvibrio, Frateuria, Stenotrophomonas, Streptomyces, Pseudomonas, and Solimonas (Figure 3B). Additionally, the heatmap showed dynamic changes in the rhizosphere communities among the groups at different sampling times; that is, there was an obvious enrichment of Proteobacteria and Firmicutes in the SA3 sample but a higher accumulation of Actinobacteria, Candidatus Saccharibacteria, Candidatus Woesebacteria, Chlamydiae, and Mucoromycota in the SF3 sample (Figure 4A). In agreement with previous research, our results indicated that Proteobacteria is the second largest phylum of hydrogenogenic CO oxidizers, which may play a significant role in helping plants against FON infection (Badger and Bek, 2008; Wang and Sugiyama, 2020). Overall, our results indicated that the application of SA changed the watermelon rhizosphere soil microbial communities.

For instance, at the phylum level, the abundances of Actinobacteria, Chlamydiae, Chloroflexi, Cyanobacteria, Dictyoglomy, Proteobacteria, Thermotogae, and Thaumarchaeota differed significantly between the SA3 and SF3 samples (Figure 4B). Moreover, the SA3 sample showed a significant abundance of Proteobacteria, whereas the SF3 sample was significantly enriched with Actinobacteria. In contrast, the SA7 sample had a significantly higher abundance of Blastocladiomycota, Dictyoglomy, and Elusimicrobiota, whereas the SF7 sample had a high abundance of Actinobacteria, Dictyoglomy, Gemmatimonadetes, Deinococcus-Thermus, Ascomycota, and Planctomycetes (Figure 4B). At the genus level, the abundance of Rhodanobacter increased significantly in the SA3 sample, whereas that of Actinoplanes, Burkholderia, Chryseolinea, Luteimonas, Micromonaspora, Nitrolancea, Ohtaekwangia, Sorangium, Sphingomonas, and Xanthomonas decreased significantly compared to the levels in the SF3 sample (Figure 4C). However, the abundances of Stenotrophomonas, Sphingomonas, Ramibacter, Herbaspirillum, Polaromonas, and Azospirillum were significantly higher, whereas those of Dokdonella, Gemmatirosa, Castellaniella, Gemmatimonas, Chitinophaga, Myxococcus, Mizugakibacter, Trichoderma, and Moritella were significantly lower in the SA7 soil than in the SF7 sample (Figure 4C). Collectively, these results suggest that although the structure of the main soil microbial community did not change, the abundance of specific species was significantly altered at different time points after pathogen injection.

Functional annotations and gene taxonomy predictions for the different rhizosphere microbiomes

For the SA3 sample, KEGG analysis of the cluster at level 1 revealed the significant activation of four major pathways: environmental information processing, cellular processes, metabolism, and human diseases (Figure 5A). Detailed information on the 43 pathways at cluster level 2 is displayed in Figure 5B. The heatmap of the KEGG ortholog groups (Figure 5C) showed that K06042 (precorrin-8X/cobalt-precorrin-8 methyl mutase), K02055 (putative spermidine/putrescine transport system substrate-binding protein), K16164 (acyl pyruvate mutase), K02055 (putative spermidine/putrescine transport system substrate-binding protein), K16164 (acyl pyruvate mutase), K02014 (iron complex outer membrane receptor protein) were more highly enriched in the SF3 sample than in the SA3 soil. Furthermore, the distribution of linear discriminant analysis scores between the SA3 and SF3 KEGG ortholog groups indicated that K02014 (iron complex outer membrane receptor protein) was increased in SA3, whereas K03088 (RNA polymerase sigma-70 factor, ECF subfamily) was increased in SF3 (Figure 5C).

The heatmap of the eggNOG classification, where the significantly expressed genes were merged into 12 groups, is shown
Figure 6A. Histograms of the distribution of linear discriminant analysis scores for resistance genes with statistically significant differences between the groups are shown in Figure 6B. For instance, there was a significantly higher expression level of genes related to transcription, carbohydrate transport, and metabolism in SF3 than in S3 plants. The genes significantly expressed in SA3 belonged mainly to post-translational; intracellular trafficking, secretion; cell motility; and cell wall: membrane envelope biogenesis. In contrast, those in SF3 belonged primarily to transcription and secondary metabolite biosynthesis. However, SF7 had a significantly higher number of genes related to translation, ribosomal structure, and biogenesis; cell cycle control and cell division; carbohydrate transport and metabolism; intracellular trafficking, secretion; energy production and conversion; defense mechanisms; and replication, recombination, and repair.

A Venn diagram of the distribution of resistance genes among the five selected samples is shown in Figure 7A. Heatmap analysis of the six CAZy classes indicated that the highly enriched genes in SA3 were in the classes GH: glycoside hydrolases and PL: polysaccharide lyases (Figure 7B). Analysis of the relative abundance of Unigenes showed that the major facilitator superfamily antibiotic efflux pump (bcr_1), drug class of aminoglycoside antibiotics, resistance mechanism of antibiotic inactivation (AAC6_IIC), and Serratia metallo-beta-lactamase (SMB_1) genes were highly expressed in SA3, whereas the ATP-binding cassette antibiotic efflux pump (msrC), drug class of aminoglycoside antibiotic, resistance mechanism of antibiotic inactivation (AAC6_IIC), ATP-binding cassette antibiotic efflux pump, major facilitator superfamily antibiotic efflux pump, resistance-nodulation-cell division antibiotic efflux pump (Pseudomonas_aeruginosa_soxR), and defense-resistant mprF (Listeria_monocytogenes_mprF) genes were increased in SA7 (Figure 7C). Finally, two-circle graphs (Figure 8) of the main phyla associated with the resistance genes expressed in the soil samples indicated that Proteobacteria were the predominant source in SA3, whereas Chloroflexi, Acidobacteria, and Cyanobacteria expressed most of the resistance genes in SA7. Interestingly, some resistance genes from Ascomycota were observed only in the SF groups and those from Firmicutes only in the SF7 group.
FIGURE 5
Functional diversity of the microbiomes among different soil samples based on KEGG pathways analysis (A) Heatmap of enrichment differences in six major metabolic pathways on level 1 among the samples. (B) Heatmap of enrichment differences in metabolic pathways on level 2 among the samples. The X-axis represents the sample group names; the Y-axis represents the KEGG metabolic pathway annotation information. (C) Distribution of linear discriminant analysis (LDA) scores between the SA3 and SF3 KEGG ortholog groups. S0, Mock-inoculation control, before treatment; S3, mock-inoculation control, 3 dpi; S7, mock-inoculation control, 7 dpi; SA3, SA + FON treatment, 3 dpi; SA7, SA + FON treatment, 7 dpi; SF3, FON treatment, 3 dpi; SF7, FON treatment, 7 dpi; FON, Fusarium oxysporum f. sp. niveum; SA, salicylic acid; dpi, days post-inoculation.
Discussion

Exogenous salicylic acid application leads to the enrichment of beneficial microbes in the watermelon rhizosphere

The rhizosphere plays a fundamental role in microbe–microbe and plant–soil–microbe interactions. Microbes and their interactions can extend the capacity of plants for disease resistance and improve their nutrient use efficiency (Berendsen et al., 2018). Emerging evidence indicates that some microbial symbionts communicate with the plant immune system through multiple feedback mechanisms, giving the plant the ability to resist pathogens and to maintain growth and development (Liu et al., 2021). Many studies have emphasized the role of soil microbial communities in enhancing plant growth and health (Alekkett et al., 2018; Toju et al., 2018). For example, Xin et al. found that the plant immune system is required to maintain the normal growth of commensal bacteria in Arabidopsis (Xin et al., 2016). Other studies have reported that assemblages of host-specific microbiomes in the rhizosphere are vital for disease resistance. For instance, some disease-resistant crop varieties are enriched in specific sets of bacterial species in the rhizosphere, which contribute to the suppression of pathogens (Delgado-Baquerizo et al., 2018). Another study showed that SA causes changes in the microbiome through allelopathy in wheat (Kong et al., 2020). It is not known how plant roots normally select and maintain a healthy rhizosphere microbiota. SA plays an important role in regulating plant immunity, which is necessary for systemically acquired resistance (Trivedi et al., 2020). In agreement with this theory, our results indicated that although the structure of the main soil microbial community did not change, specific microbes were significantly altered at different time points after pathogen injection. Likewise, we observed that Proteobacteria, the second largest phylum of hydrogenogenic CO oxidizers (Badger and Bek, 2008; Wang and Sugiyama, 2020), accumulated significantly in SA3. Lebeis et al. found that SA could modulate the root microbiome of A. thaliana. Specifically, plants with altered SA signaling had root microbiomes that differed from each other in their relative abundance of Proteobacteria as one of the core microbiomes when compared with those of wild-type plants (Lebeis et al., 2015). Furthermore, at the genus level, SA3 had a significantly higher abundance of...
Rhodanobacter than SF3 and SA7 had a higher abundance of Azospirillum, Herbaspirillum, Stenotrophomonas, and Sphingomonas than SF7. Rhodanobacter is capable of oxidizing ammonia and of denitrification (De Cercq et al., 2006; Huo et al., 2018). Some members of the genus Azospirillum exhibit biocontrol activity against phytopathogens and have been used as biofertilizers because of their plant growth-promoting activities, such as biological nitrogen fixation, hormone production, phosphate solubilization, and siderophore production (Mendez-Gomez et al., 2020). Sphingomonas is considered an abundant microbial resource for the biodegradation of aromatic compounds, thus showing great potential for environmental protection and industrial production applications because of its high metabolic capacity and multifunctional physiological characteristics (Vorholt et al., 2017). Stenotrophomonas maltophilia, which exists widely in water, soil, and animals, is a multidrug-resistant opportunistic pathogen that causes life-threatening infections in immunocompromised individuals (Messila et al., 2007; Mendes et al., 2013; Hassan and Bano, 2016). Herbaspirillum species have been described as closely associated with plants, both endophytically and epiphytically, because their nitrogenase activity promotes plant growth (Wang et al., 2014). For instance, using histochemical analysis, Herbaspirillum seropedicae has been shown to colonize the root surfaces and inner tissues of maize, sorghum, wheat, and rice seedlings grown in vermiculite (Ramos et al., 2020).

Therefore, these significantly altered microbial communities confirmed our hypothesis that SA can recruit beneficial microorganisms (such as Rhodanobacter, Sphingomonas, and Micromonospora) in the watermelon rhizosphere. The next question that needs to be answered is whether SA-influenced changes in rhizosphere microflora are beneficial to the antagonism of plants against pathogens, that is, is the increased resistance of plants to pathogens induced directly by SA alone, or by the rhizosphere microbiota, or by both?

Significantly expressed microbiome genes may interact with salicylic acid signaling to regulate watermelon resistance to disease

Many researchers have found that SA and plant-associated beneficial microorganisms are key candidates for systemic acquired resistance (SAR) and induced systemic resistance (ISR), respectively (Berendsen et al., 2018; Teixeira et al., 2019). Therefore, to elucidate the molecular mechanism through which the rhizosphere microbial community cooperates with the watermelon plant to induce
resistance against *Fusarium* wilt, we blasted unique genes to annotate their functions. Surprisingly, genes enriched in environmental information processing, cellular processes, and metabolism pathways were significantly more highly expressed in SA3 than in SF3. Furthermore, K02014 (iron complex outer membrane receptor protein) was expressed at significantly higher levels in SA3 than in SF3. Simultaneously, the genes enriched in the process groups of cell motility and cell wall: membrane envelope biogenesis were expressed more significantly in SA3 than in SF3. These results confirm our hypothesis that the rhizosphere microbiome assemblage is affected by SA signal transduction pathways (Mendes et al., 2011; Chaparro et al., 2014; Vorholt et al., 2017; Delgado-Baquerizo et al., 2018). Notably, the resistance genes from GHs and PLs were highly accumulated in SA3 compared with those in the other soil groups. In particular, AAC6_IIC was consistently more significantly accumulated in the SA groups, indicating that it may play an important role in interplay with plant lipid membranes. Furthermore, we noticed that the relative abundance of Proteobacteria corresponding to resistance genes was the highest post-inoculation, which means that more of such genes may come from species in this phylum. Our results confirm the conclusions of other studies that plant genetics and agricultural practices can potentially impose selective pressures on specific microbes and microbial communities. Thus, our findings not only show that SA treatment is beneficial to the antagonism of plants against pathogens but also suggest that, during pathogen infection, the rhizosphere microbiome may play a key role in activating a series of defensive feedback mechanisms in the plant through SA signal transduction pathways.

**Conclusion**

In conclusion, our results indicate that exogenous SA treatment can specifically increase some beneficial rhizosphere species that can confer resistance against FON inoculation. The glycoside hydrolase and polysaccharide lyase genes in the microbiome, specifically from a group of beneficial microbes (such as *Rhodanobacter*, *Sphingomonas*, and *Micromonospora*), may potentially induce activation of the plant immune system against *Fusarium* wilt disease and promote plant growth. Our results provide a novel strategy for controlling *Fusarium* wilt in watermelon by manipulating the
rhizosphere microbiome through phytohormones, such as exogenous SA treatment. Furthermore, we aim to elucidate the mechanisms underlying the interplay between lipid membrane signaling and SA signal transduction pathways in future studies.

Data availability statement

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary material.

Author contributions

FZ and RW: conceptualization. YF, ZW, and FZ: methodology. PW, KY, and FZ: investigation. FZ: writing—original draft preparation. EY, YF, and RW: writing—review and editing and funding acquisition. LX and RW: supervision. All authors contributed to the article and approved the submitted version.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.1015038/full#supplementary-material

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Supplementary material

The Supplementary material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.1015038/full#supplementary-material
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