Rhizobacterial community structure in grafted tomato plants infected by *Ralstonia solanacearum*

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Abstract. Navitasari L, Joko T, Murti RH, Arwiyanto T. 2020. Rhizobacterial community structure in grafted tomato plants infected by *Ralstonia solanacearum*. Biodiversitas 21: 4888-4895. Bacterial wilt disease caused by *Ralstonia solanacearum* is a devastating soil-borne vascular disease of tomato leading to a 100% yield loss. One of the alternatives to suppress the infestation of *R. solanacearum* infestation is the application of grafting techniques, which has been studied and successfully practiced by tomato growers. However, the infestation mode of *R. solanacearum* and the rhizobacterial community structure in grafted tomato plants are poorly reported. In this study, the rhizobacterial community structure in grafted tomato plants infected by *R. solanacearum* was investigated. The experiment was conducted on tomato germplasms with the implementation of tube grafting using resistant rootstocks (Amelia from Indonesia, H.7996 from Asian Vegetable Research Development Center/AVRDC) and susceptible scion (Servo from Indonesia). The rhizobacterial community structure was analyzed by metagenomic study under 16S rRNA genes sequencing with a distinct region (16SV3-V4) that was amplified using a specific primer (16SV4: 515F-806R) 5’-GTGCCAGCMGCCGCGGTAA and 5’GGACTACHVHHRHTCTAAAT. The results indicated that the grafted tomato plants and resistant rootstocks that were infected by *R. solanacearum* showed significantly lower intensity of bacterial wilt disease compared to the susceptible scion. The rhizobacterial community structure in the grafted tomato plants infected by *R. solanacearum* was indicated by predominant phyla of Proteobacteria, Firmicutes, and Actinobacteria with dominant genera of *Pseudomonas* and *Bacillus*. Besides, significant difference was also indicated by species of *Geitlerinema* sp. in the grafted tomatoes infected by *R. solanacearum*.

Keywords: Bacterial wilt, disease intensity, grafting, predominant rhizobacteria

INTRODUCTION

Bacterial wilt disease caused by *Ralstonia solanacearum* is a devastating soil-borne vascular disease that severely affects tomato (*Solanum lycopersicum* L.), leading to a 100% yield loss (Genova et al. 2013; Hemelda et al. 2019). *R. solanacearum* infected tomato plants by colonizing the plant rhizosphere, where the pathogen must compete with the other bacterial taxa (Hibbing et al. 2010). After reaching a threshold density, pathogen switches on its virulence gene expression and invades plant roots (Schell 2000). Once within xylem vessels, *R. solanacearum* rapidly spreads to aerial plant parts throughout the vascular system (Dalsing et al. 2015) and blocks the water flow via excessive production of extracellular polysaccharides (Genin and Denny 2012). Control strategies of bacterial wilt disease caused by *R. solanacearum* is difficult to perform due to the high variability capacity of the pathogen to survive in diverse environments as well as its wide host range (Yanti et al. 2017). One of the alternatives to suppress the development of bacterial wilt disease is grafting method using rootstock from resistant varieties.

Grafting has been utilized to manage bacterial wilt disease in tomato plants worldwide (Rivard et al. 2012). The method can provide an effective management strategy to reduce bacterial wilt disease incidence and subsequent crop loss (Louws et al. 2010). Grafting method using rootstock from resistant varieties is the most effective method in disease control of bacterial wilt disease in the fields. The grafting method using resistant rootstock significantly reduces bacterial wilt incidence in tomato plants (Louws et al. 2010; McAvoy et al. 2012). Resistant rootstock in grafting also influences and changes microbial community in the rhizosphere (Gilardi et al. 2013; Perez-Alfoceae et al. 2015), thereby changing the rhizobacterial diversity and affecting the pathogens in the rhizosphere. Poudel et al. (2019) reported that grafting between tomato BHN589 and Maxifort or RST-04-106 without infection of *R. solanacearum* indicated that the diversity and abundance bacterial communities in rhizosphere were higher than in endosphere, and the abundance of bacteria in the grafted tomato plants was higher than in the non-grafted ones. However, there is little information about how the infection of *R. solanacearum* in grafted tomato plants suppresses bacterial wilt disease and influences the diversity of rhizobacterial community structure. Therefore, the effects of tomato grafting using resistant rootstock on the rhizobacterial community structure and bacterial wilt disease intensity were investigated in this study. The evidence that the rhizobacterial community structure was influenced by resistant rootstock, grafting, and infections of *R. solanacearum* were provided.
MATERIALS AND METHODS

Preparation of bacterial inoculum

*Ralstonia solanacearum* was isolated from the rhizosphere of tomato plants showing bacterial wilt disease symptoms in Yogyakarta, Indonesia. The phylotype, race, and biovar of isolated *R. solanacearum* were also analyzed. *R. solanacearum* was incubated at a temperature of 28°C and grown on Yeast Peptone Glucose agar (Yeast extract 5 g/L, peptone 10 g/L. Glucose 20 g/L, agar 15 g/L, and aquadest 1 L) (Laeshita and Arwiyanto 2007). *R. solanacearum* grown on YPGA medium for 48 hours were added to 10 ml sterile aquadest and then shaken, and the suspension of *R. solanacearum* was measured with Spectrophotometer (UV-vis Genesys 10S, Thermo Scientific) to obtain the OD 0.1 at λ 580nm. The suspension of *R. solanacearum* having OD 0.1 was used for the infection treatment in the grafted tomato plants.

Grafting and planting in the field

The rootstocks from two resistant tomato cultivars were used for grafting. The resistant rootstocks used were Amelia (Matahari Seed Company) and H7996 (Asian vegetable research development Center/AVRDC), while the susceptible scion used was Servo (East-West Company). The grafting made were GrftAmlS (Amelia and Servo) and GrftH7996S (H7996 and Servo) using tube graft method. The grafts were held in plastic tubes with a diameter of 1 cm, and grafting performed when rootstock and scion have same diameter (1 cm) (21 days old). The grafts were then moved into a healing chamber and maintained for ten days at a temperature of 22-25°C and relative humidity of 90-95%. After 10 days in healing chamber, the rooted grafted and non-grafted tomato plants (control plants) were removed and then planted in field under plastic mulch (Arwiyanto et al. 2015). A 30 ml of *R. solanacearum* inoculum was splashed close to the roots of plants and the bacterial wilt disease intensity was measured every week by calculating the disease intensity (Widyaningsih et al. 2019; Solekha et al. 2019). The experiment was arranged in a randomized complete block design consisting of 5 treatments and 4 replications with 10 plants within each treatment, and the total number of experimental units was 200 plants (Arwiyanto et al. 2018).

\[ I = \frac{\sum_{n=1}^{N} X \times v}{N \times Z} \times 100\% \]

Where:

- **I**: Disease intensity (%)
- **n**: number of damaged plants
- **v**: Score of the damage level (1: healthy; score 3: < 15% wilt on leaf; score 5: 3 >15%-45% wilt on leaf; score 7: > 45%-85% wilt on leaf; and Score 9: > 85% wilt on leaf and death)
- **Z**: the highest damage level value
- **N**: Total sample plants observed

Preparation of rhizosphere soil samples for metagenomic analysis

Sample analysis of rhizobacteria taken from grafted (GrftAmlS and GrftH7996S) and non-grafted tomato plants (Amelia, H7996, Servo) infected by *R. solanacearum* was performed at 30 days after planting. Total samples for analysis were 15 samples of 50 g of rhizosphere soil. The rhizosphere soil was collected as previously described (Riley and Barber, 1969).

Extraction of genomic DNA and Amplicon Generation

Total genomic DNA from each sample was extracted using Phusion® High-Fidelity PCR Master Mix (New England Biolabs). The DNA concentration and purity were monitored on 1% agarose gels. According to the concentration, DNA was diluted to 1ng/μL using sterile water. Meanwhile, the amplicon generation, 16S rRNA (genes of distinct regions/16SV3-V4), was amplified using specific primer (16SV4: 515F-806R) 5'-GTGCCAGCMGCCGCGGTAA and 5'GGACTACHV1HHHHTWTCTAAAT. Samples with a bright main strip between 400-450bp were chosen for further experiments. Then, the libraries generated with Ion® Fragment Library Kit 48 rxns for Thermo fisher and quantified via Qubit and Q-PCR were sequenced by IonS5™XL (Thermo fisher) (Wang et al. 2018).

Metagenomic analysis

Metagenomic analysis was performed at Novogene Bioinformatics Technology Co., Ltd (Singapore). A 5 μg DNA was used to construct 400-450bp sequencing libraries performed by Illumina Hiseq PE150 (Illumina, San Diego, USA). The libraries generated with Ion Plus Fragment Library Kit 48 rxns for Thermofisher and quantified via Qubit and Q-PCR were sequenced by IonS5™XL (Thermofisher). Single-end read was assigned to samples based on their unique barcode and truncated by cutting off the barcode and primer sequence. The reads were compared with the reference database (Gold database, http://drive5.com/uchime/uchime_download.html) using UCHIME algorithm (UCHIME Algorithm, http://www.drive5.com/usearch/manual/uchime_algo.html) to detect chimera sequences (report template life.pl), and then the chimera sequences were removed to get effective reads final (Caporaso et al. 2010, Edgar et al. 2011). Sequences analyses were performed with Uparse software (Uparse v7.0.1001 http://drive5.com/uparse/) using all the effective reads. Sequences with ≥97% similarity were assigned to the same OTUs (Haas et al. 2011). Analysis using Mothur software was performed on the SSUrRNA database of SILVA Database (http://www.arb-silva.de/) used for species annotation at each taxonomic rank (Threshold: 0.8–1) kingdom, phylum, class, order, family, genus, and species. Meanwhile, the phylogenetic relationship of all OTUs representative sequences was analyzed using the MUSCLE (Version 3.8.31, http://www.drive5.com/muscle/) (Quast et al. 2012). Analysis of diversities and species in each sample was carried out based on all effective reads by 97% DNA sequence with operational Taxonomy Units (OTUs) generating 97%
sequence identity homologous in species, and then the rank of relative abundance and diversities of rhizobacteria were drawn by Ternary Plot.

**Statistical analysis**

The data of bacterial wilt disease intensity were statistically analyzed with ANOVA, and the data showing significant differences were further analyzed with Duncan Multiple Range Test (DMRT) at alpha 5%. Meanwhile, the rhizobacterial community structure in within-group and among groups was determined using T-test, and the variation of rhizobacteria species in groups was analyzed with Metastats.

**RESULTS AND DISCUSSION**

**Analysis of Ralstonia solanacearum and measurement of bacterial wilt disease intensity**

*Ralstonia solanacearum* observed in this study belongs to phylotype 1, race 1, and biovar 3. There was no significant difference in the bacterial wilt disease intensity between the grafted tomato plants and resistant rootstocks (Amelia and H7996). On the contrary, significant difference in the bacterial wilt disease intensity was observed in the susceptible scion (Servo) (Table 1). Generally, bacterial wilt disease intensity in the grafted tomato plants, resistant rootstocks, and susceptible scion increased from 7 to 28 days after transplanting (Figure 1). The increase of bacterial wilt diseases intensity is related to the resistance level of the rootstock, susceptible scion, and the combination (grafting) (Table 1). The intensity of bacterial wilt disease in the grafted tomato plants (GrfAmIS and GrfH7996S) was lower compared to that in the susceptible scion (Servo), but higher than that in the resistant rootstocks (Amelia and H.7996) (Figure 1). The intensity of bacterial wilt disease in the resistant rootstocks (Amelia and H.7996) and grafted tomato plants (GrfAmIS and GrfH7996S) was significantly lower than that in the susceptible scion (Servo) (Table 1).

Table 1. Bacterial wilt disease intensity and resistance level

| Group       | Bacterial wilt disease intensity (%) | Resistance level       |
|-------------|--------------------------------------|------------------------|
| Amelia      | 21.8750 a                           | Moderate resistant     |
| GrfAMIS     | 38.0000 a                           | Moderately susceptible |
| H.7996      | 30.3750 a                           | Moderately resistant   |
| GrfH.7996S  | 40.6875 a                           | Moderately susceptible |
| Servo       | 69.3750 b                           | Susceptible            |

Note: Means followed by the same letters are not significantly different based on the Duncan’s Multiple Range Test (DMRT) (p<0.05). Resistant level based on Janaki and Putturaja (2012). GrfAMIS (grafting Amelia and Servo), GrfH7996S (Grafting H7996 and Servo)

Figure 1. Percentage of bacterial wilt disease intensity in resistant rootstocks (Amelia and H.7996), susceptible scion (Servo), and grafted tomato plants using resistant rootstock and susceptible scion

The resistant rootstocks of Amelia and grafting between resistant rootstocks of Amelia and susceptible scions of Servo were indicated by dominant rhizobacterial species of *Fictibacillus arsenicus*. Meanwhile, the resistant rootstocks of H7996 were indicated by predominant rhizobacterial species of *Pseudomonas resinovorans*. The different result was found in the grafting between resistant rootstocks of H7996 and susceptible scions of Servo that showed similar dominant rhizobacterial species (*Sphingomonas jaspsi*) to the dominant rhizobacterial species in the susceptible scion (Figure 5). Besides, the relative abundance of *R. solanacearum* was also found in the rhizosphere. The resistant rootstocks of Amelia and the grafting between resistant rootstocks of Amelia and susceptible scion showed relative abundance of *R. solanacearum* that was higher compared to the relative abundance in the resistant rootstocks of H.7996, the grafting between resistant rootstocks of H.7996 and susceptible scion, and the susceptible scion. The relative abundance of *R. solanacearum* in the grafted tomato plants was higher than in the resistant rootstocks but smaller than in the susceptible scion (Figure 5).
The relative abundance of *R. solanacearum* in rhizosphere affected the relative abundance of others rhizobacteria as shown in Table 2 that indicated the relative abundance of *R. solanacearum* affected the difference of rhizobacteria in phylum, class, order, and species. Significant difference in rhizobacterial phylum was shown by Phylum Latescibacteria. The phylum Latescibacteria was significantly different in GrftAmlS, GrH7996S, resistant rootstocks of H7996, and susceptible scion of Servo but not in the resistant rootstocks of Amelia. The significant differences in class and order of rhizobacteria were shown by class of unidentified Acidobacteria of the phylum Acidobacteria and order of unidentified Alphaproteobacteria and Acidobacteriales. The relative abundance of *R. solanacearum* in rhizosphere also influenced significant difference of rhizobacteria in species of *Geiternema* sp. in the grafted tomato plants (GrftAmlS and GrH7996S) and susceptible scion of servo.

Figure 3. Diversity of rhizobacterial community structure
Figure 4. Predominant species of rhizobacteria in resistant rootstocks, susceptible scions, and grafted tomato plants. The resistant rootstocks are Amelia and H.7996; the grafting of Amelia-Servo is GrftAmlS, the grafting of H.7996-Servo is GrH.7996S, and the susceptible scion is Servo.

Figure 5. Top ten dominant relative abundant rhizobacterial species.

Table 2. The difference in the abundance of the rhizobacteria in grafted tomato plants, resistant rootstocks, and susceptible scions.

| Group    | Phylum           | The difference in the abundance of rhizobacteria between group | Class          | Order                          | Species               |
|----------|------------------|---------------------------------------------------------------|----------------|-------------------------------|-----------------------|
| Amelia   | NS               | Unidentified Acidobacteria*                                   | Acidobacteriales**, unidentified Alphaproteobacteria* | NS               |                      |
| GrftAmlS | Latiscibacteria  | Unidentified Acidobacteria*                                   | Acidobacteriales**, unidentified Alphaproteobacteria* | NS               |                      |
| H.7996   | Latiscibacteria  | Unidentified Acidobacteria*                                   | Acidobacteriales**, unidentified Alphaproteobacteria* | NS               |                      |
| GrH7996S | Latiscibacteria  | Unidentified Acidobacteria*                                   | Acidobacteriales**, unidentified Alphaproteobacteria* | Geitlerinema sp.** |                      |
| Servo    | Latiscibacteria  | Unidentified Acidobacteria*                                   | Acidobacteriales**, unidentified Alphaproteobacteria* | Geitlerinema sp.** |                      |

Note: * significantly different at P < 0.05; ** significantly different at P < 0.01; NS: not significantly different; grafted tomato plants are GrftAmlS and GrH7996S; resistant rootstocks are Amelia and H7996; and susceptible scion is Servo.
The use of resistant cultivar in the management of bacterial wilt disease caused by *R. solanacearum* is one of the highest priorities. For instance, resistant rootstocks in grafting have been used to suppress bacterial wilt disease. Huet (2014) reported that the most effective method of disease control of bacterial wilt disease is through the use of resistant cultivar. McAvoy et al. (2012) also reported that a commonly used grafting method using resistant rootstock significantly decreased bacterial wilt incidence in fields. The result in this study showed that bacterial wilt disease intensity in grafted tomato plants was not significantly different compared to that in the resistant rootstocks, but significantly lower than that in the susceptible scion. It indicates that grafting using resistant rootstocks reduced bacterial wilt diseases intensity caused by *R. solanacearum* in fields. The use of resistant rootstocks in grafting can restrict bacterial spread in the roots. Furthermore, resistant plants can also limit the bacterial spread in the vascular stem tissue. However, combination grafting of resistant rootstocks with susceptible scion changes the resistance level from moderately resistant to moderately susceptible. The resistance level of Amelia and H7996 are moderately resistant, making them recommended as rootstocks that are resistant to *R. solanacearum* (Genova et al. 2013, Laeshita and Arwiyanto, 2017).

The change in resistance level from resistant to moderately resistant is influenced by the new race of *R. solanacearum* and the environment. A new pathogen race destroys the resistance of a plant variety. Plant varieties with vertical resistance (resistance is determined by a single gene or a few genes and only effective against some strains of pathogen) need replacement of resistance every few years (3, 5, or 10 years), and this is depending on the pathogen genetic plasticity, a particular gene, or a genetic combination, as well as environmental conditions on the development of the disease (Laeshita and Arwiyanto, 2017; Sutrisno et al. 2018). Deberdt et al. (2014) also reported that tomato cultivar H-7996 is known to be resistant to *R. solanacearum* but changing to be susceptible after being inoculated with *R. solanacearum* strains IIB, which is an emerging ecotype. The resistance and susceptibility of plant varieties are influenced by environmental factors, such as changes in the physical environment and pathogens race (Semangun, 2006; Wei et al. 2017). Besides, the change in the resistance level is also influenced by relative abundance of *R. solanacearum*. Generally, all grafted tomato plants, resistant rootstock, and susceptible scion showed relative abundance of *R. solanacearum* in rhizosphere. However, the relative abundance of *R. solanacearum* in resistant rootstock was higher than that in the grafted tomato plants and susceptible scion. It indicates that resistant rootstock can limit bacterial spread in plants, and the effect is indicated by the low intensity of bacterial wilt disease compared to the grafted tomato plants and susceptible scion. However, grafted tomato plants with a combination of resistant rootstock and susceptible scion showed lower intensity of bacterial wilt disease intensity compared to the susceptible scion. It was influenced by relative abundance of *R. solanacearum* in the grafted tomato that was higher than in the resistant rootstock but lower than in the susceptible scion. Besides, the intensity of bacterial wilt disease was also influenced by diversity of rhizobacteria in the rhizosphere. When attacked by pathogen, plant select and recruit bacteria in rhizosphere by root exudates, and then root exudates influence community structure of the rhizosphere microbes by recruiting beneficial microbes and repelling pathogens. The recruitment and selection of rhizobacteria influenced the rhizobacterial community structure (Bakker et al. 2013, Wei et al. 2015).

The rhizobacterial diversity and community structure associated with the relative abundance of *R. solanacearum* were shown by dominant rhizobacterial phylum of rhizobacteria. The dominant rhizobacterial phylum was shown by phylum Proteobacteria (alpha, beta, and gamma proteobacteria), Firmicutes (Bacilli), and Actinobacteria (Nocardioides). Mendes et al. (2011) reported that infection of pathogen in plant was influenced by predominant phylum of Proteobacteria, Firmicutes, and Actinobacteria. Proteobacteria (alpha, beta, and gamma proteobacteria) and firmicutes (Bacilli) were identified as the most dynamic taxa associated with disease suppression. Actinobacteria have been observed to exhibit uniquely increased survival rates through periods of environmental stress (Legget et al. 2012). When attacked by pathogen, plants seem to actively select specific elements of their bacterial rhizosphere microflora and then select and recruit specific soil microbes into the rhizosphere microbiome. Plant roots also influence specific functions of the microbiome, and a plant seems to respond to pathogen infection by systemic signaling (Joko et al. 2018), leading to enhanced biocontrol activity in the microbiome (Bakker et al. 2013; Massart et al. 2015). Infection of pathogen in the rhizosphere also affects plant defense and recruitment of beneficial microbe by root exudate. Plant exudation can change the rhizosphere community composition by recruiting certain beneficial microbes and directly repelling the pathogen (Joko et al. 2012).

Infection of *R. solanacearum* pathogen showed significant effect on rhizobacterial phylum. The significant difference in the rhizobacterial phylum related to relative abundance of *R. solanacearum* was shown by phylum Latescibacteria. Latescibacteria has SAGs harbor extensive uptake systems for sugars and amino acids/oligopeptides (Youssef et al. 2015). Uronic acids and uronic acid derivatives are potentially imported using a single common transporter, a sugar-phosphate permease transporter of the major facilitator superfamily (Joko et al. 2007) similar to the ExuT transporter of *R. solanacearum* (Gonzalez and Allen 2003). Besides, the significant difference in rhizobacteria was also shown by *Geitlerinema* sp. Patel et al. (2018) reported that the Cyanobacterium *Geitlerinema* sp. has the characteristic of phycocyanin that correlated with antioxidant activity and therapeutic agent against oxidative stress. Infection of *R. solanacearum* causes oxidative stress on plant, and this response of plants is used to adapt and suppress pathogen. Colonization of *R.*
solanacearum is exposed to host-derived ROS, which triggers a bacterial oxidative stress response that adapts the pathogen to the xylem environment (Flores and Allen, 2009). Infection of R. solanacearum was also related to the dominant rhizobacterial species in genus Bacillus (Fibicicillus arsenicus and Bacillus fuscus), Pseudomonas (Pseudomonas resinovorans), and Sphingomonas (Sphingomonas jaspsi). The genus of Bacillus and Pseudomonas are known as antagonistic microorganisms, which are mainly isolated to control pathogens from the rhizosphere and plant microflora (Lamsal et al. 2012). P. resinovorans displays a wide range of antimicrobial and antifungal activity (Fikri et al. 2018), and S. jaspsi produces carotenoids, including zeaxanthin. Zeaxanthin is one of the carotenoids that are a stronger ROS scavenger. When bacterial cells are exposed to the oxidative stress as a result of cellular metabolism, zeaxanthin accumulated in the bacterial cell membrane might quench the ROS, thereby decreasing their lethal effects (Tian et al. 2007).

In conclusion, grafting using resistant rootstock and infection of R. solanacearum influences the intensity of bacterial wilt disease and rhizobacterial community structure. The intensity of bacterial wilt disease intensity in grafted tomato plants was not significantly different compared to that in the resistant rootstock, but significantly different from that in the susceptible scion. The grafted tomato plants showed lower intensity of bacterial wilt diseases compared to the susceptible scion, but showed higher intensity compared to the resistant rootstocks. The intensity of bacterial wilt diseases is related to the rhizobacterial community structure indicated by dominant phyla in Proteobacteria, Firmicutes, and Actinobacteria with dominant genera of Pseudomonas and Bacillus. Besides, significant difference is also indicated by species of Geitlerinema sp. on tomatoes grafted infected by R. solanacearum.

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