Monitoring Tritrophic Biocontrol Interactions Between *Bacillus* spp., *Fusarium oxysporum* f. sp. *cubense*, Tropical Race 4, and Banana Plants *in vivo* Based on Fluorescent Transformation System

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*Bacillus* spp. is effective biocontrol agents for Fusarium wilt of banana (FWB), tropical race 4 (TR4). This study explores the colonization by *Bacillus subtilis*, *Bacillus velezensis*, and *Bacillus amyloliquefaciens* of host banana plants and elucidates the mechanism of antagonistic TR4 biocontrol. The authors selected one *B. subtilis* strain, three *B. velezensis* strains, and three *B. amyloliquefaciens* strains that are proven to significantly inhibit TR4 *in vitro*, optimized the genetic transformation conditions and explored their colonization process in banana plants. The results showed that we successfully constructed an optimized fluorescent electro-transformation system (OD$_{600}$ of bacteria concentration = 0.7, plasmid concentration = 50 ng/μl, plasmid volume = 2 μl, transformation voltage = 1.8 kV, and transformation capacitance = 400 Ω) of TR4-inhibitory *Bacillus* spp. strains. The red fluorescent protein (RFP)-labeled strains were shown to have high stability with a plasmid-retention frequency above 98%, where bacterial growth rates and TR4 inhibition are unaffected by fluorescent plasmid insertion. *In vivo* colonizing observation by Laser Scanning Confocal Microscopy (LSCM) and Scanning Electron Microscopy (SEM) showed that *Bacillus* spp. can colonize the internal cells of banana plantlets roots. Further, fluorescent observation by LSCM showed these RFP-labeled bacteria exhibit chemotaxis (chemotaxis ratio was 1.85 ± 0.04) toward green fluorescent protein (GFP)-labeled TR4 hyphae in banana plants. We conclude that *B. subtilis*, *B. velezensis*, and *B. amyloliquefaciens* can successfully colonize banana plants and interact with TR4. Monitoring its dynamic interaction with TR4 and its biocontrol mechanism is under further study.

Keywords: *Bacillus* spp., biocontrol, electro-transformation, RFP-labeled *Bacillus*, *Bacillus* interaction with TR4
INTRODUCTION

Fusarium wilt of banana (FWB) caused by Fusarium oxysporum f. sp. cubense, especially Tropical Race 4 (TR4), is one of the most destructive diseases affecting the crop (Ghag et al., 2015; Presti et al., 2015; Carvalhais et al., 2019). Pathogen spores invade the vascular bundles of banana roots through wounds and then extend to the corms and pseudostems, causing the vascular bundles to become brown and necrotic, the leaves gradually wither and eventually the whole plant dies (Swarupa et al., 2014). Due to its characteristic of surviving in the soil for decades, once the pathogen is introduced into the soil, the infected banana orchard cannot be used for growing susceptible banana cultivars, which seriously affects the sustainable development of the banana industry, as there are few proven TR4-resistant cultivars (Ploetz, 2006, 2015).

Like all the other Foc strains, TR4 cannot be controlled using fungicides and cannot be eradicated from soil using fumigants (ProMusa, 2021). Crop rotation and intercropping have been used to reduce the infections and inoculum levels (Nadarajah et al., 2016). In China, farmers have been growing bananas in the presence of TR4 by rotating or intercropping with Chinese leek (Allium tuberosum; Nadarajah et al., 2016; Li et al., 2020a,b). The most effective solution supporting continued production of bananas in infested soils would be replacing susceptible cultivars with resistant ones. However, almost all important banana cultivars are susceptible to TR4 (Ploetz, 2015; Chen et al., 2019; Sun et al., 2019), and most commercial cultivars are triploid and sterile (non-seed bearing) makes banana breeding more difficult (Hwang and Ko, 2004; Heslop-Harrison and Schwarzacher, 2007). At present, few TR4-resistant banana cultivars have been bred and popularized (Daniells, 2011). Furthermore, even those cultivars still have to adapt to local cultivation practices and conditions. As a result, the spread of TR4 has led to an increase in research on biological control and biocontrol agents (BCAs) in suppressing the pathogen (Nadarajah et al., 2016; Bubici et al., 2019; Damodaran et al., 2020).

So far, many microbes such as Trichoderma spp., Pseudomonas spp., and Bacillus spp. have been widely used as BCAs (Bubici et al., 2019). The characteristics of spore-forming and rapid growth of Bacillus species confer them with an important advantage over other beneficial biological control microorganisms. In addition, many Bacillus species can synthesize a large number of secondary metabolites, which play a key role in antibiosis against detrimental microorganisms (Radhakrishnan et al., 2017; Fira et al., 2018). Bacillus subtilis is a representative Bacillus, which could produce a variety of antibiotics with different structures and activities; it also exhibits a wide range of antibacterial activities against different plant pathogens under in vitro conditions (Stein, 2005). Bacillus amyloliquefaciens, a type of Gram-positive bacterium (Priest et al., 1987), highly homologous with B. subtilis, has a single nutrient requirement and is harmless to the environment and human health. Due to these positive characteristics, many B. amyloliquefaciens strains have been isolated and identified as having significant inhibition and control effects on Ceitocybe bescens and Fusarium oxysporum (Guan and Jiang, 2013). Wang et al. (2015) and Xue et al. (2015) found that B. amyloliquefaciens combined with organic fertilizers can significantly reduce the incidence of Fusarium wilt. The B. amyloliquefaciens isolated by Zhang et al. (2014) showed a good biocontrol effect on banana wilt, and this strain can produce IAA and siderophore, promote the growth of banana plants, and has a high biocontrol potential. Bacillus velezensis is also a Gram-positive bacterium that is closely related to B. amyloliquefaciens. FZB42 is currently the most researched B. velezensis strain, which has been commercialized, and it is effective against various pathogens caused by bacteria and fungi (Borriess et al., 2011). Fan et al. (2011) integrated a plasmid carrying a gene encoding a fluorescent protein into the chromosome of the FZB42 strain and successfully observed the colonization and distribution in the roots of corn, Arabidopsis, and duckweed. However, so far there are few reports on the colonization and distribution of Bacillus on banana plants. Understanding how these beneficial microorganisms colonize and distribute in host banana plants will provide an important and favorable theoretical basis for using biocontrol strains to control FWB.

Most biocontrol Bacillus species are soil microorganisms that colonize the rhizosphere of plants and directly or indirectly promote plant growth through different mechanisms (Compant et al., 2005; Scherwinski et al., 2008; Malfanova et al., 2013). These play an important role in many fields, such as ecological restoration, biocatalysis, and biological control. The successful colonization of biocontrol strains is a prerequisite for the development of biocontrol promotion and disease prevention, and it is vital to explore their interaction processes with plants (Kang, 2019). Fluorescence transformation is currently the most successful approach for studying the colonization of Bacillus spp. and the interaction with plants in vivo (He, 2014). However, many wild-type Bacillus species with good bio-promoting and disease-preventing effects cannot easily form competent cells because of the unknown cellular restriction-repair system, which leads to constraints to a low efficiency of electric shock transformation (Alegre et al., 2004; Yasui et al., 2009). This seriously hinders banana research on horizontal manipulation or modification of Bacillus, as well as the further utilization and exploitation of the potential value of these biocontrol agents. Hence, there is a need to develop a reliable and efficient fluorescent-transformation system for monitoring the interactions between Bacillus, Foc TR4, and banana plants.

In this study, seven Bacillus species containing one B. subtilis, three B. velezensis, and three B. amyloliquefaciens strains with strong antagonistic effects on TR4 in vitro were selected. In order to obtain stable fluorescent-marked transformants and develop an efficient genetic transformation system, pYP69 carrying red fluorescent protein (RFP) was used as the fluorescent expression vector, which was successfully introduced into wild-type Bacillus strains according to the optimized experimental parameters. Furthermore, laser confocal observation confirmed that the fluorescent-transformed strains could be used for monitoring how Bacillus colonizes host banana plants (Figure 1).
MATERIALS AND METHODS

Source of Strains and Plasmid

*Bacillus subtilis* strain YN1419 (GenBank Accession No. MW647761) was isolated from banana cultivar Brazilian in Xishuangbanna, Yunnan, China. *Bacillus velezensis* strain YN1282-2 (GenBank Accession No. MW663765) was isolated from banana cultivar GCTCV-119. *Bacillus velezensis* N67 (GenBank Accession No. MW672323), WBN06 (GenBank Accession No. MW672324), and *B. amyloliquefaciens* G9R-3 (GenBank Accession No. MW674627; Zhou et al., 2020), HN04 (GenBank Accession No. MW674626) were isolated from Guangxi banana plantations. *Bacillus amyloliquefaciens* YN0904 (GenBank Accession No. MW647760) was isolated from Yunnan banana plantation. All experimental strains had proven strong *Foc* TR4 antagonistic effects in vitro (Li et al., 2021). *Foc* TR4 strain 15-1 (Zhang et al., 2018) was isolated from infected banana plants in Xishuangbanna, Yunnan, China. GFP-TR4 was constructed in our laboratory (Zhang et al., 2018) and was used in monitoring the interaction with bacteria. The plasmid pYP69 expressing RFP and the chloramphenicol-resistance gene (Supplementary Figure S1) were obtained as a gift from Dr. Yongmei Li (Plant Protection College of Yunnan Agricultural University, Kunming, China) and Dr. Yiyang Yu (Plant Protection College of Nanjing Agricultural University, Nanjing, China). pYP69 was constructed by the pYC127 as backbone and cloned with mKate2 coding sequence (Chen et al., 2012). The *Escherichia coli* DH5α was purchased from Beijing Biomed Biotechnology Co., Ltd. (Beijing, China). The strains were stored in 25% glycerol at −80°C. The isolates were reactivated on nutrition agar (NA) medium at 37°C for 24 h.

Culture Media

LB broth medium (LB; tryptone 10 g, yeast extract 5 g, NaCl 10 g, pH 7.0 for 11 with deionized water) was used to cultivate the bacteria and prepare the bacterial suspension. Potato Dextrose Agar (PDA) medium (200 g potato, 20 g glucose, 20 g agar, diluted to 1 l, natural pH) was used to activate the *Foc* TR4 and conduct the dual-culture experiments. Growth Medium [GM; LB with 3% glycine (Gly), 1% DL-threonine (DL-Thr), 0.03% Tween 80, and 9.1% sorbitol] was used to prepare competent cells. ETM buffer (40 ml glycerol, 360 ml deionized water, 36.4 g sorbitol, 36.4 g mannitol, 0.25 mM KH₂PO₄, 0.25 mM K₂HPO₄, and 0.5 mM MgCl₂) was used to wash away the ion components in competent cells. Recovery Medium (RM; LB with 9.1% sorbitol and 6.92% mannitol) was used for resuspension of competent cells after electroporation.

Detection of Plasmid pYP69 Expression in *Escherichia coli*

The *Escherichia coli* competent cells were thawed on ice. One microliter pYP69 was mixed gently into competent cells and put it on ice for 30 min stationary. Then, the heat shock was applied at 42°C for 60 s, and then, the culture was put on ice for 2 min. Five hundred microliter LB liquid medium was then added to the culture and resuscitated at 37°C, swirling at 180 rpm for 60 min, and then, the transformative culture was evenly spread on the 100 μg/ml ampicillin-resistant medium. The transformative plates were cultured at 37°C for 16 h, and then, single colonies were selected to observe fluorescence by the Fluorescence Microscope (Nikon 80I).

Plasmid Extraction

The *E. coli*-carrying pYP69 plasmid was inoculated into LB liquid medium containing ampicillin (100 μg/ml), and cultured
at 37°C, swirling at 220 rpm for 16 h. Plasmid extraction followed the instructions accompanying the OMEGA Plasmid DNA Extraction Kit (E.Z.N. A® Plasmid DNA Mini Kit I).

Electroporation
Preparation of Competent Cells
Competent cells preparation (Zhang et al., 2011) was carried out as follows. *Bacillus* strains stored at −80°C were inoculated in the LB solid medium overnight. Each single colony was individually selected and inoculated in 30 ml GM liquid medium by inoculating loop, cultivated at 37°C, 220 rpm for 16 h. Two milliliter cultured bacteria suspension was inoculated into 200 ml GM liquid medium, cultured at 37°C, and swirled at 220 rpm for 3 h. OD_{600} value was determined by spectrophotometer (Nanophotometer NP80 Touch). When OD_{600} value reached 0.5, 3% Gly, 1% D-glucosamine, and 0.03% Tween 80 were added into the culture for cell-wall weakening. The culture was arrested when the OD_{600} value reached 0.7. The bacterial suspension was put on ice for 10 min and centrifuged at 4°C, 1,300 g for 10 min; then, 50 ml precooled ETM solution was used to wash away the ion components in the GM medium for three times. About 500 μl ETM was used to suspend the competent cells, competent cells were divided into 100 μl per EP tubes and stored at −80°C.

Electroporation
The transformation (Kang, 2019) was carried out as follows. Two microliter plasmid pYP69 were added into 100 μl competent cells culture, mixed the plasmid and competent cells, and electroporation was carried out. The electroporation voltage was 1.8 kV, and the capacitance was 400 Ω. After electroporation, 1 ml RM medium was immediately added to the electroporation cup and then transferred to a 1.5 ml centrifuge tube, cultured at 37°C, shook at 220 rpm for 4 h. The culture was spread on the LB solid plate containing 10 μg/ml chloramphenicol, the positive transformants designated according to the format of RFP-*Bacillus*, e.g., RFP-N67, etc., were selected, and the fluorescence labeling was observed under the fluorescence microscope (Excitation wavelength: 555 nm/Emission wavelength: 584 nm).

Stability Determination of Plasmid pYP69 in *Bacillus*
The activated RFP-*Bacillus* strains were inoculated into liquid LB medium without antibiotics, cultured at 37°C, and shook at 180 rpm, with samples being taken every 5 h and spread on LB solid plates, and bacteria colonies were observed by fluorescence microscope. The percentage of red fluorescent colonies in total cells colonies used to calculate the stability of the plasmid in the RFP-strains. Three replicates were conducted.

Cell Growth Determination of RFP-Strains
Red fluorescent protein-*Bacillus* and wild-type (WT) strains were inoculated in the LB liquid medium and cultured to the concentration was OD_{600} reaching 1.0, and then, the bacteria suspension was transferred to the blank medium at a ratio of 1%. Bacteria were cultured at 37°C and shook at 180 rpm for 60 h. OD_{600} values of samples were measured every hour in the first 6 h. OD_{600} was measured every 2 h during 8–20th h, and OD_{600} was measured every 4 h during 22nd–60th h. Three replicates were made.

Antagonistic Activity of RFP-Strains on TR4
The dual-culture method was used to compare the antagonistic activity of the RFP-labeled *Bacillus* and WT strains against *Foc* TR4. TR4 was activated in PDA for 7 days at 28°C, and RFP-labeled bacteria and WT bacteria were activated at 37°C for 24 h. Individual 5 mm diameter disks of TR4 hyphae were placed in the center of each PDA plate. Plates were then inoculated with the RFP strain and the WT strain at 2.5 cm from the center by using inoculating loop. These were cultivated for 7 days, and the growth of TR4 hyphae was measured. The inhibition dual culture assay was carried out in three replicates.

Colonizing Observation of Biological Control *Bacillus*
A 10 ml red fluorescent bacterial suspension (cultivated 24 h at 37°C and 180 rpm, diluted to 1 × 10^6 cfu/ml) was poured into the tissue-culture bottle which cultured five banana plantlets by MS medium, gently shaking the bottle and placing in a culture incubator (30°C, 80% humidity, 12 h light/12 h dark), with the treatment adding just sterilized water as the control. Five replicates were made. Ten banana plant roots per treatment were randomly selected after inoculation for 7 days, and the roots were washed with flowing sterile water to remove the medium.

For fluorescent microscopy, tissue slices (thickness: 50 nm) were excised by freezing microtome and any bacteria in root tissues were observed by Laser Confocal Electron Microscope (Leica TCS-SP8). Excitation/emission wavelengths were 561 nm/570–640 nm for RFP (mKate2 protein).

For scanning microscopy, the critical point drying sample processing method was conducted. Tissue slices (thickness: 50 nm) were immersed in FAA fixative (5 ml 38% formaldehyde, 5 ml glacial acetic acid, 90 ml 70% ethyl alcohol, and 5 ml glycerol) at 4°C overnight. Then: the fixed samples were immersed in 50, 60, 70, 80, 90, and 95% alcohol successively to dehydrate 30 min was conducted in each alcohol concentration, then the samples were immersed in absolute ethanol for 30 min, and three replicates were conducted. Banana tissue samples were then put into Critical Point Dryer (K850) and Cressington Sputter Coater (108 AUTO) to spray gold coating. The fixed samples were put into Scanning Electron Microscope (ZEISS Sigma 300, Germany) to observe the bacterial colonization.

Laser Confocal Electron Microscope Observation of Interaction *in vivo*
In order to observe the interactions between strains and TR4 *in vivo*, 3 ml RFP-labeled bacterial culture in 9% physiological saline solution was inoculated into the leaf vascular bundles of the banana cultivar Brazilian by injection (Liu et al., 2020),
and a 5 mm agar disk with GFP-labeled TR4 mycelia (Zhang et al., 2018) was placed on the wound by inoculation loop. The latest fully expanded leaves were selected. Fluorescent observation by laser scanning confocal microscope (Leica TCS-SP8) was carried out after the leaves were cultured in 28 °C, 60% light, and 50% humidity for 7 days. Excitation/emission wavelengths were 561 nm/570–640 nm for RFP (mKate2 protein). Excitation/emission wavelengths were 488 nm/500–540 nm for GFP (AmCyan protein).

**Chemotaxis Assay of Bacteria to Pathogen**

Seven day-old Foc TR4 were inoculated in 50 ml diluted PDB (1: 50; v: v in H2O) at 28°C with shaking at 150 rpm for 24 h to obtain spores suspension of Foc TR4. Suspension was then washed twice with sterile ddH2O and incubated for 48 h in 5 ml sterile ddH2O at 28°C and 150 rpm. The supernatant was sterilized by filtration through a 0.22 μm membrane (Millex-GP) for use.

To obtain bacterial suspension, 1 × 10^7 colony ml^-1 RFP-N67 strains were grown in LB overnight at 37 °C, washed either with sterile ddH2O, and diluted in ddH2O to the OD_560 of 0.1.

Chemotaxis capillary assays (Palmieri et al., 2020) were carried out as follows. 250 μl bacterial suspension was added to the well (1.5 cm x 1.5 cm) in the glass slide together with a 10 μl capillary containing the test compound (Foc TR4 hyphal exudate or ddH2O). Slides were incubated for 60 min at 28°C, capillaries were carefully lifted, the content was serially diluted and plated onto LA medium with 10 μg/ml chloramphenicol, and CFUs were counted 24 h after incubation at 37°C. The chemotaxis ratio was calculated by dividing the number of bacteria in the tube containing the test compound (Foc TR4 hyphal exudate) by the number of bacteria in the tube containing the control (ddH2O). All experiments included four replicates and were performed three times with similar results.

**Statistical Analysis**

Data were analyzed by one-way ANOVA using the SPSS version 18.0 for Windows (Chicago, IL, United States). The figures and charts were drawn using ORIGIN 2018 (Massachusetts, United States).

**RESULTS**

**Plasmid pYP69 Expresses Fluorescence in Escherichia coli**

Before transforming *Bacillus*, we aimed to verify whether this plasmid could be expressed in *E. coli*, preserved, and extracted. Heat-shock transformation was used to transform the plasmid pYP69 into *E. coli* DH5α competent cells. We obtained positive transformants on plates containing ampicillin. Colonies glowed an obvious red color under the fluorescence microscope, and the red *E. coli* cells could still be visualized even after being cultured in the liquid. This verified that RFP-labeled cells can be observed under the fluorescence microscope (Figure 2A), indicating that the plasmid pYP69 can be expressed in *E. coli*.

**Plasmid pYP69 Expresses Fluorescence in *B. subtilis*, *B. velezensis*, and *B. amyloliquefaciens***

Fluorescence labeling is one of the best methods for tracing bacterial colonization (including *B. subtilis*, *B. velezensis*, and *B. amyloliquefaciens*) in plants. Here, electroporation with optimized conditions was used to transfer the plasmid pYP69 into competent cells of WT *Bacillus* strains. Positive transformants of *Bacillus* were obtained on the solid plate of 10 μg/ml chloramphenicol. These can be seen with the naked eye as the colonies of strains formed on the antibiotic plate gradually become light red. Fluorescence microscope observation results showed that the plasmid pYP69 has been successfully transformed into the WT strains and can express RFP in the *Bacillus* strains (Figure 2B; Supplementary Figures S2A–E). Several repeated experiments showed that although there was still relatively low trans-formants efficiency (1 × 10^10 cfu/μg of plasmid DNA); stable transformants from each *Bacillus* strain had already been generated.

**RFP-Labeled Strains Possess High Plasmid Stability**

The stability of plasmid expression in the strains is important for the control of fluorescent strains. Under condition of no antibiotic pressure, the red fluorescent strain was cultured in serial dilutions, and the samples were taken every 5 h. The dilution was evenly spread on a non-resistant plate. The proportion of colonies with fluorescence was counted under a fluorescence microscope to determine the frequency of plasmid retention. We found that the frequency of RFP plasmid cells is all greater than 98% after being cultured for 10 consecutive generations (Figure 3). Results confirmed the plasmid is rarely lost due to the proliferation of bacterial cells, indicating that the plasmid pYP69 can be stably expressed in these *Bacillus* strains, and it can be used for experiment tracing colonization and migration in plants.

**RFP-Labeled Strains Have the Similar Growth Condition of WT-Strains**

The growth of the RFP-*Bacillus* and WT strains was compared under the same inoculation and culture conditions. The results showed that the growth of RFP-YN0904, RFP-YN1282-2, RFP-WBN06, and RFP-N67 was consistent to WT strains, indicating that the plasmid pYP69 had no significant effect on the growth of these three *B. velezensis*. The wild-type strains of HN04 and G9R-3 grew slight faster in the log phase and reached the stable phase earlier than RFP strains, the wild-type strains YN1419 grew lower before the stable phase than the RFP strain, but they did not significantly affect normal growth of RFP bacteria (Figure 4; Supplementary Figures S3A–E). We speculate that it may be that the introduction of multicopy large plasmids affects the growth rate of *B. amyloliquefaciens* and *B. subtilis*, and good fluorescence performance of the plasmids in the *Bacillus* host may also create an additional metabolic burden on the strains.
RFP-Labeled Strains Retain Their Antagonistic Activity Against TR4

The metabolic burden caused by the introduction of exogenous plasmids sometimes affects other functions of the strain. In order to explore the effect on the biological characteristics of plasmid pYP69 insertion, the antagonistic activities of the RFP-Bacillus strain and the WT strain against the pathogenic fungus Foc TR4 were compared. The results showed that there was no significant difference in the antagonistic activity of the two types of strains against the tested pathogenic fungi (Figure 5; Supplementary Table S1). It shows that the expression of...
plasmid pYP69 does not affect the inhibitory activity of the Bacillus strains on the growth of the pathogenic fungus.

**Bacillus Can Successfully Colonize Banana Root Cells**

Biocontrol bacteria successfully colonizing plants are a necessary pre-condition for their biocontrol function. To explore the colonization capacity of Bacillus in banana plants, we inoculated tissue-cultured banana plantlets with B. velezensis RFP-N67. After 7 days of co-culture, we sampled the banana plantlets’ roots, and slice observation by Laser Confocal Microscopy (LCM) showed that RFP-N67 colonized the banana roots cells (Figure 6A) and Scanning Electron Microscopy (SEM) confirmed the bacteria could successfully colonized the roots xylem cells (Figures 6B, C). We also found that there were no bacteria in banana plants subject to low inoculation concentrations (1 × 10³–10⁵ cfu/ml), and that Bacillus can enter the root when the inoculation concentration reached a higher density (1 × 10⁶ cfu/ml). However, excessive concentrations could damage banana plantlets.

**Bacillus Exhibits Chemotaxis Toward Foc TR4 in Banana Plants**

The interactions between beneficial Bacillus strains and pathogens are the key to crop disease control. There are few studies on the interactions between biocontrol bacteria and TR4 in banana. Therefore, we selected RFP-labeled Bacillus RFP-N67 and used available green fluorescent labeled pathogens GFP-TR4, inoculating banana leaves simultaneously to observe whether they will interact in vivo. Leaf phenotypic observation showed that the lesion size in both the front and back sides of the leaves after the treatment inoculated with RFP-N67 and GFP-TR4 was significantly smaller than those found in leaves inoculated with GFP-TR4 only (Figure 7A). Then, we made ultrathin sections around the leaf lesions and successfully observed RFP-N67 and GFP-TR4 simultaneously under laser confocal microscopy, finding that RFP-N67 often appeared around GFP-TR4 mycelia (Figures 7B, C), and further chemotaxis assay, indicating that RFP-N67 exhibits a strong chemotaxis toward Foc TR4 (Figure 7D). Therefore, we can speculate that the biocontrol bacteria RFP-N67 we used can grow and reproduce normally in banana to exert their biocontrol functions, and that in the presence of pathogens, it can be quickly found to inhibit pathogen growth. However, this study only provides histological evidence. Quantitative analysis of specific pathogen growth inhibition needs to be carried out.

**DISCUSSION**

Over the last decade, many studies have focused on the interactions of Foc with beneficial microorganisms. Generally, researches have centered around “growth promotion,” “systemic resistance induction,” “secondary metabolites syntheses,” etc., which were facilitated by the rapid development of “-omics” tools (Bubici et al., 2019). However, as we know, there are no reliable currently available fluorescent labeled Bacillus strains for Banana–Endophyte–Pathogen tritrophic interaction studies. Currently, fluorescence labeling is the best approach for exploring interactions of Bacillus spp. with plants in vivo. Hence, the fluorescence transformation system we constructed is the first step and of great significance to study the interaction mechanisms associated with TR4 and biological control Bacillus. In the process of manipulation, the wild-type Bacillus generally has a low transformation efficiency, and some inert Bacillus strains cannot even be transformed at all, which seriously affects internal mechanism research of its beneficial properties, such as
disease prevention, growth promotion, efficient enzyme production, or antibiotics (He, 2014). Currently, the transformation of Bacillus mainly includes protoplast transformation, electric shock transformation, natural transformation, and protoplast electric shock transformation (Shen et al., 2013; He, 2014; Kang, 2019; Plucker et al., 2021; Wang, 2021). In this study, we have explored the different methods and conditions of fluorescent transformation of TR4-antagonistic strains. The results showed that we only got the positive transformants by the method of electric shock transformation. Through the natural transformation method, all seven Bacillus strains could not form naturally competent cells or positive transformants, even if we tested in the different transformation conditions, such as culture time, plasmid concentration, and recovery time. The low electro-transformation efficiency of wild-type Bacillus may be related to the restriction-repair system in the cell (Zhang et al., 2012). According to the statistics of REBASE (a professional database of restriction endonucleases), approximately 88% bacterial genomes contain restriction-repair systems, and 43% contain four or more restriction-repair systems (Roberts et al., 2007). The restriction-repair system is a barrier for bacteria to exclude external DNA, thus preventing the transformation of bacteriophages and external plasmids, thereby maintaining the integrity and functional stability of its own genetic material. At present, the restriction-repair systems have been found in a variety of bacteria and archaea (Roberts et al., 2003). Finding a way to prevent the wild-type bacteria from degrading the external DNA during the transformation process is the key to improving the efficiency of electric shock transformation.

In the process of cell-wall synthesis, glycine can replace D(L)-alanine in the peptidoglycan component of the bacterial
cell wall, which reduces the degree of cross-linking of peptidoglycan and interferes with the synthesis and assembly of the cell wall, thus loosening the cell wall (Anderson et al., 1966; Zhu et al., 2020). It has been reported that the addition of some compounds that inhibit cell-wall synthesis [such as glycine, threonine, penicillin, or Tween 80 (affecting cell membranes’ fluidity)] during the exponential growth stage of Bacillus can improve the transformation efficiency (Holo and Nes, 1989; Zhang et al., 2011). After numerous repeated experiments, we added glycine, sorbitol, threonine, and Tween

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**FIGURE 7** Bacillus velezensis RFP-N67 exhibits chemotaxis toward the GFP-TR4 pathogen. N67-RFP in 9% physiological saline solution was injected into the leaf vascular bundles, and the pathogen GFP-TR4 was placed on the injection wound. (A) Disease incidence of banana leaves after 7 days’ inoculation. The first two pictures show the front side of the leaves, and the last two pictures show the back side of the leaves. Three replicates were used. (B) Microscopy of the control treatment, which injected 9% physiological saline solution with no bacteria, inoculated the GFP-TR4 only. (C) Representative micrographs of N67-RFP showed significantly chemotropic toward GFP-TR4. BF, bright field; RFP, red fluorescent field; GFP, green fluorescent field; and MERGE, merge image with BF, RFP, and GFP fields. Three replicates were used. (D) Chemotaxis ratio of RFP-N67 toward Foc TR4. *Indicate the significance between treatments at the 0.05 level.
80 to the electro-transformation growth medium, finding this helps *Bacillus* more easily absorb external DNA to form positive transformants. Although transformation efficiency is between $1 \times 10^2$–$10^4$ cfu/µg of plasmid DNA in the repetitive experiment, indicating our transformation system is still needed to be further optimized, which means that there is still much space for improvement in the transformation of inert *Bacillus* strains. However, this electric shock transformation system is reproducible and stable because we have generated RFP-labeled *Bacillus* in all seven strains from two main banana producing areas, Yunnan and Guangxi provinces, China (Figure 2B; Supplementary Figures S2A–E).

There are other methods that can also improve transformation efficiency: a good way to temporarily inactivate restriction endonuclease in the host bacterium is by incubating at a certain temperature for a couple of times after electric shock. *Corynebacterium glutamicum* was cultured by shaking under low-temperature to prepare electro-transformation competent cells, and then rapidly heat shocked after the electric shock, the results showed that the electro-transformation efficiency was increased fourfold (Rest et al., 1999). Heat-shocking after electric shock transformation of *B. amyloliquefaciens*, increased transformation efficiency by 10-fold (Zhang et al., 2011). In addition, due to the restriction endonuclease and methylase in the wild-type bacteria restriction-repair systems often appearing in pairs, the in vivo methylation modification of the exogenous plasmid to be transformed in the same cell can evade restriction-enzyme digestion and degradation of the host wild-type bacteria during the transformation process, thereby improving the transformation efficiency. Through this strategy, the transformation efficiency of *Bifidobacterium adolescentis* ATCC15703 was increased fivefold (Yasui et al., 2009), and the transformation efficiency of *B. amyloliquefaciens* TA208 and *B. cereus* ATCC10987 was increased almost fourfold (Zhang et al., 2012).

A vital factor affecting the expression of fluorescent proteins in *Bacillus* is plasmid instability. In order to explore the stability of plasmid pYP69 in *Bacillus* strains, we detected RFP expression and the loss-ratio of plasmid in the transformed strains with regularly sampling. This showed that within 50h of culture, more than 98% of the cells expressed RFP, indicating that very few cells lost their plasmids (Figure 3), and *Bacillus* growth rate was not affected (Figure 4; Supplementary Figures S3A–E). According to previous reports (Bonfirate et al., 2013), the propagation rate of *Bacillus* in natural environments is 50–100 h/generation, and the rate of propagation under laboratory conditions is 20–30 min/generation. Based on this, it could be calculated that the number of fluorescent strain propagations during 50h is about 100–150 generations. In other words, the plasmid pYP69 still exists stably after 100–150 generations, indicating that it has a very strong compatibility with bacteria. Therefore, RFP-labeled *Bacillus* will be suitable for monitoring its colonizing activities in banana host plants in our next research step.

Nowadays, the biological control of banana Fusarium wilt is always focused on screening antagonistic strains, while ignoring the research on the colonization of antagonistic bacteria in soil or plants. Many of the selected antagonistic bacteria have obvious antibacterial effects in vitro or pot experiments, but they usually lose biocontrol effects in the field (Jing et al., 2020; Wei et al., 2020). Therefore, the successful colonization of biocontrol bacteria is an important pre-condition for its application and function in field. Some studies have shown that the colonization of plant growth promoting rhizobacteria (PGPR) is affected by abiotic factors, such as soil temperature, texture, water content, and oxygen content. Biotic factors including root exudates, plant growth conditions, bacterial chemotaxis, the nature of self-regulation mechanisms, and bacterial trophic type also affect colonization (Yajing, 2018). At present, a few studies have been carried out on the colonization of strains controlling banana wilt. Dai (Yajing, 2018) detected the quantity of different PGPR in banana soil rhizosphere by fluorescence quantitative PCR. The results indicated that the biomass of three antagonistic bacteria in banana rhizosphere soil was increased significantly, indicating that the three PGPR strains M8, C5, C14 can colonize banana roots (Lin, 2011). Their colonization determination studies used antibiotic-labeled strains and showed that labeled strains could be isolated by wound inoculation, irrigated inoculation, and axil inoculation. The control did not show any bacterial colonies, indicating that with three methods of inoculation, the labeled strains can be colonized in banana. Chao et al. (2010) used the scanning electron microscope to observe biocontrol bacteria FJAT-346-PA-K in tissue-cultured banana plantlets 10 days after inoculation. They successfully found that there were biocontrol bacteria in the roots’ internal tissues and banana stems. We inoculated the wild-type strain N67 into tissue-cultured banana plantlets, and it was also confirmed by scanning electron microscope that N67 could successfully colonize the banana root cells (Figures 6B,C). However, this method can only preliminarily judge the existence of biocontrol strains in the plant, and it cannot clearly figure out the internal activity in vivo. Therefore, we constructed red fluorescent-labeled strains to observe its dynamic migration in banana plants, and the results showed that the biocontrol strains can successfully colonize the roots but not in the corn or pseudostem due to short incubation period (Figure 6A). We also injected RFP-labeled bacteria into the detached leaf which is proved to be a reliable protocol (Liu et al., 2020). The results showed that the biocontrol strains can successfully colonize and grow in the leaf, as well as displaying a positive chemotaxis response toward TR4 hyphae, indicating that the biocontrol bacteria can effectively interact with and inhibit the pathogen in vivo (Figure 7). Other studies have shown that chemoattraction of bacteria could contribute to its root colonization (Palmieri et al., 2020). *Fusarium oxysporum* f. sp. *lycopersici* (Fol) is known to facilitate bacterial movement in search of nutrients which also exhibits chemotaxis toward plant roots (Furuno et al., 2009; Turra et al., 2015). We thus conclude that N67 not only directly inhibits TR4 hyphae growth, but it could also benefit from the capacity of pathogen hyphae trends to plant colonization and thus increases colonization efficiency. Significant progress has been made in other crops by using fluorescent-labeled strains for colonization observation. Kang (2019) observed that the endophytic *B. velezensis* CC09 Bv-GFP can not only effectively colonize wheat roots, but also
migrate to stem and leaf organs to achieve whole plant distribution. He (2014) observed the roots of maize seedlings inoculated with Y2-P43GFPmut3a by fluorescence microscope, showing that Y2 strain successfully colonized root surfaces and interiors. However, as far as we know, there is no research on the in vivo interaction of RFP-Bacillus for monitoring TR4 in banana, so dynamic migration is our next proposed research.

With the great progress of gene-sequencing technology, more and more Bacillus whole genomes have been sequenced. Using such tools, it will be important to explore the potential biocontrol mechanisms among Bacillus, pathogens, and plants (Bubici et al., 2019; Carrió et al., 2019; Jiang et al., 2019; Chen et al., 2020; Jing et al., 2020; Wei et al., 2020). The biocontrol mechanisms of Bacillus are usually considered to be based on one or more of: (1) Antagonism: Bacillus often secretes secretory secondary metabolites that inhibit pathogen growth. (2) Competition: Bacillus competes for niches with pathogens and other microorganisms to obtain nutrients and other resources. (3) Inducing systemic resistance: they can activate host defense responses by inducing systemic resistance. (4) Promoting growth: Bacillus can provide necessary mineral nutrition and plant hormones (e.g., IAA) for plant hosts to support their life activities. These mechanisms of Bacillus are interrelated and synergistic. Of course, the principal Bacillus biocontrol mechanism(s) could be different in different crops.

It is an innovative research direction to develop new biopesticides and new agricultural antibiotics by using the antagonistic behavior of microorganisms (Yajing, 2018). It has become an important measure for biological control of TR4 in organic farming by researching “biological fertilizer” (where organic fertilizers are inoculated with biocontrol agents) and banana growth-promoting bacterial agents to increase yield and disease resistance (Ling et al., 2014; Fu et al., 2016). Nowadays, biological control for sustainable banana production has attracted much attention. New biological microorganism agents are urgently needed to replace traditional chemical pesticides, which continue to cause pollution and damage the environment, and degrade soils. Microbial agents as recognized promising “pollution-free pesticides” will play a crucial role in managing agricultural and silvicultural diseases and protecting ecological balance.

CONCLUSION

We successfully developed an optimized fluorescent electro-transformation system of TR4-inhibitory Bacillus spp. strains (OD600 = 0.7, plasmid concentration = 50 ng/µl, volume = 2 µl, voltage = 1.8 kV, and capacitance = 400 Ω). The RFP-labeled Bacillus strains have high stability, and their growth rates and inhibition effects on TR4 are unaffected by fluorescent plasmid insertion. In vivo colonizing observation by Laser Scanning Confocal microscopy (LSCM) and SEM showed that Bacillus spp. can colonize the xylem cells of banana plantlets’ roots. Further fluorescent observation by LSCM showed these RFP-labeled bacteria exhibit chemotaxis toward the hyphae of the green fluorescent protein (GFP)-labeled TR4 pathogen in banana leaves.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

PH conceived, designed, and performed the experiment, analyzed the data, and wrote the paper. SL conceived, designed, and performed the experiment, and analyzed the data. S-JZ conceived and designed the experiment and prepared the manuscript. SX, HF, YW, and GH analyzed the data. WZ and GF provided the strains sources and analyzed the data. S-JZ and Y-YW supervised the research and provided funding support. All authors contributed to the article and approved the submitted version.

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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.2021.754918/full#supplementary-material
