Role of the Type VI Secretion System in the Pathogenicity of *Pseudomonas syringae* pv. *actinidiae*, the Causative Agent of Kiwifruit Bacterial Canker

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The type VI secretion system (T6SS), a macromolecular machine, plays an important role in the pathogenicity of many Gram-negative bacteria. However, the role of T6SS in the pathogenicity of *Pseudomonas syringae* pv. *actinidiae* (*Psa*), the pathogen of kiwifruit bacterial canker, is yet to be studied. Here, we found a T6SS gene cluster consisting of 13 core genes (A-J) in the genome of *Psa* M228 based on a genome-wide analysis. To determine whether the T6SS gene cluster affects the pathogenicity of *Psa* M228, T6SS and its 13 core gene deletion mutants were constructed and their pathogenicity was determined. The deletion mutants showed different degrees of reduction in pathogenicity compared with the wild-type strain M228; in *tssM* and *tssJ* mutants, pathogenicity was significantly reduced by 78.7 and 71.3%, respectively. The pathogenicity results were also confirmed by electron microscopy. To further confirm that the reduction in pathogenicity is related to the function of T6SS, we selected the T6SS gene cluster, comprising *tssM* and *tssJ*, for further analyses. Western blot results revealed that *tssM* and *tssJ* were necessary for hemolytic co-regulatory protein secretion, indicating that they encode a functional T6SS. Further, we explored the mechanism by which T6SS affects the pathogenicity of *Psa* M228. The ability of bacterial competition, biofilm formation, hydrogen peroxide tolerance, and proteolytic activity were all weakened in the deletion mutants M228ΔT6SS, M228Δ*tssM*, and M228Δ*tssJ*. All these properties of the two gene complementation mutants were restored to the same levels as those of the wild-type strain, M228. Quantitative real-time results showed that during the interaction between the deletion mutant M228ΔT6SS and the host, expression levels of T3SS transcriptional regulatory gene *hrpR*, structural genes *hrpZ*, *hrcC*, *hopP1*, and effector genes *hopH1* and *hopM1* were down-regulated at different levels. Taken together, our data provide evidence for the first time that the T6SS plays an important role in the pathogenicity of *Psa*, probably via effects on bacterial competition, biofilm formation, and environmental adaptability. Moreover, a complicated relationship exists between T6SS and T3SS.

**Keywords:** pathogenicity, type VI secretion system, Kiwifruit bacterial canker, type III secretion system, competition, biofilm formation, environmental adaptability
INTRODUCTION

The bacterial canker of kiwifruit is caused by the virulent form of *Pseudomonas syringae* pv. *actinidiae* (*Psa*) and is the most prevalent disease in the kiwifruit industry (Pinheiro et al., 2020). *Psa* can enter the plant through stomata, water holes, lenticels, wounds (caused by birds, insects, and/or human contact) and can colonize the plant (including branches, leaves, buds, leaf marks, and pruned diseased branches) for a long time without inducing external symptoms, until a suitable environmental condition is developed (Donati et al., 2020). Systemic infection occurs through various pathways (Stefani and Giovanardi, 2011; Tontou et al., 2014). *Psa* can infect young twigs systemically within a few minutes (Michelotti et al., 2018). Therefore, pathogenic bacteria can infect kiwifruits repeatedly, breed in cortex, expand up and down, and even move to the xylem and central column of the plant, causing severe pathogenicity under appropriate conditions (Gao et al., 2016). Due to the latent and short-term outbreak characteristics of *Psa*, bacterial canker has been identified as a destructive disease of kiwifruit, which results in production losses worldwide. Recently, this disease has received more attention in the main kiwifruit planting areas owing to the huge economic losses (Vanneste, 2017; Hang et al., 2018; Kim et al., 2019; Williams et al., 2020). Therefore, identifying the pathogenic mechanism will play a vital role in effective prevention and control of the disease.

Bacterial phytopathogens usually infect their host plants by various extracellular proteins or secretion of effectors through secretory systems (Khan et al., 2018; Deb et al., 2019). These secretion systems are a class of complex nanomolecular machines that can transport the virulence proteins to the external environment or host cells directly or indirectly (Costa et al., 2015; Wen et al., 2018). To date, at least six different types of secretion systems (I–VI) have been identified in Gram-negative pathogenic bacteria (Mcquade and Stock, 2018). Among them, a newly discovered secretion system, the type VI secretion system (T6SS), has been shown to play an important role in pathogenicity (Agnetti et al., 2019; Zhu et al., 2020) and has also been implicated in bacterial interactions or environmental adaptations such as colonization, biofilm formation, resistance, and survival (Mathias et al., 2017; Wu et al., 2018; Ben-aakov and Salomon, 2019; Asolkar and Ramesh, 2020). T6SS comprises 13 core genes (*tssA* to *tssM*) usually encoded within the same gene cluster (Leiman et al., 2009; Catarina et al., 2011; Kapitein et al., 2013; Kudryashev et al., 2015; Gianfanelli et al., 2016). Among them, hemolytic co-regulatory protein (Hcp) of bacteriophage T4 (Pell et al., 2009; Nguyen et al., 2018) is one of the major substrates secreted by T6SS. Since the secretion of Hcp is T6SS-dependent, it is often a reliable indicator of whether T6SS functions appropriately (Pukatzki et al., 2009).

Multiple studies have found that T6SS was closely related to the pathogenicity in many pathogenic bacteria strains. Mattinen et al. (2010) found that there was four H in the extract from pathogenic bacterium *Pectobacterium atrosepticum*, when it was cultured in minimal medium supplemented with host extract. Overexpression mutants of the *hcp* genes were constructed. The pathogenicity of the mutant strains was enhanced compared with that of the wild-type strain. Hcp may act as a new virulence factor in *P. atrosepticum* (Mattinen et al., 2010). Since the secretion of Hcp is dependent on the T6SS secretion system, the results indicates that T6SS plays an important role in *P. atrosepticum*. The crown gall disease caused by *Agrobacterium tumefaciens* is a worldwide tumor forming disease of the plants and is a major problem for plant nursery industries. It can cause great economic loss in fruit plants (Habbadi et al., 2017). Wu et al. (2018) found that the pathogenicity of *A. tumefaciens* is closely related to T6SS. *Pantoea ananatis* LMG 2665 is the most prevalent pathogenic bacterium found in pineapple fruit and onions. Shyntum et al. (2015) found that two sets of T6SS in its genome, namely T6SS-1 and T6SS-3. T6SS-1 is closely related to the pathogenicity and competitiveness of the strain *P. ananatis* LMG 2665, while T6SS-3 has no effect on its pathogenicity. This indicates that T6SS-1 is an important virulence factor in *P. ananatis*. In addition, the core genes *tssM* (Zhang et al., 2012) and *tssB* (Zhang et al., 2014) of T6SS are essential for the pathogenicity of *Ralstonia solanacearum*. This indicates that T6SS plays an important role in an increasing number of plant bacterial diseases, where the core genes of T6SS have different functions. The bacterial canker of kiwifruit, caused by *Psa*, is the most prevalent disease in the kiwifruit industry; however, the function of T6SS in the pathogenicity of *Psa* is still unclear. In this study, we found a T6SS gene cluster consisting of 13 core genes (A-J) in the genome of *Psa* M228 based on a genome-wide analysis. We studied the role of T6SS in the pathogenicity of *Psa* by mutant construction through homologous recombination technology and conducted a preliminary exploration of its mechanism. The results showed that T6SS is an important pathogenic determinant in *Psa* M228 and plays a role in bacterial competition, biofilm formation, hydrogen peroxide tolerance, proteolytic ability, and T3SS function. Our results provide insights into the functions of T6SS in the pathogenic mechanism of *Psa*.

MATERIALS AND METHODS

**Bacterial Strains, Plasmids, and Reagents**

*Pseudomonas syringae* pv. *actinidiae* strain M228 (*Psa* M228) was isolated from kiwi leaves in the Meixian, Shaanxi Province of China in 2010 (Gao et al., 2016). Deletion and complementation mutants were constructed in this study. The strain *Psa* M228 and mutants were grown on Luria-Bertani (LB) agar plates or cultured in LB liquid medium at 25 ± 1°C. *Escherichia coli* DH5α was grown in LB medium at 37°C. For gene cloning, *E. coli* S17-1::pir was cultured under the same conditions for bacterial gene transfer. When necessary, ampicillin (Amp, 10 µg/mL), kanamycin (Km, 10 µg/mL), tetracycline (Tcr, 10 µg/mL), erythromycin (Em, 10 µg/mL), and nalidixic acid (NAL, 10 µg/mL) were used. All strains used in this study were provided by the Plant Pathology Laboratory at Northwest A&F University, Yangling, and PRC. All reagents and solvents were of analytical grade. Antibiotics were purchased from Sigma (United States). Plasmid pK18mobSacB was used for gene deletion, and purchased from Biovector Science Lab, Inc.
(Beijing, China); pDSK-GFPuV was used for fluorescently labeled strains construction, and was provide by Ph.D. Mysore (The Samuel Roberts Noble Foundation, United States); pMarA was used for Bacillus transformation, and provided by Professor Qi Wang (China Agricultural University, Beijing, China); pDSK was used for gene complementation, pBR322 was used for E. coli DH5α transformation, and these two plasmids were provided by Professor Xi-Hui Shen (Northwest A&F University, Yangling, China).

T6SS Bioinformatics Analysis of Psa M228

The whole genome of Psa M228 has been sequenced by our laboratory, and the genome data have been registered on NCBI (ANJ10000000.2). The gene tssC sequence of Psa M228 was obtained through bioinformatics analysis, and an evolutionary tree was constructed from the tssC sequence of 68 species in 38 genera that contain T6SS. There are three representative types of T6SS and from the evolutionary tree; the species containing them were selected. Subsequently, a phylogenetic tree was constructed to determine the type of T6SS Psa M228 belonged to. The evolutionary history was inferred using the Maximum Likelihood method based on the Le_Gascuel 2008 model (Le and Gascuel, 1993). The bootstrap consensus tree inferred from 1000 replicates was used to show the evolutionary history of the taxa analyzed (Felsenstein, 1985). Branches corresponding to partitions reproduced in less than 60% bootstrap replicates were collapsed. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) is shown next to the branches (Felsenstein, 1985). The initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and the topology with a superior log likelihood value was selected. A discrete Gamma distribution was used to model evolutionary rate differences among sites (five categories (+G, parameter = 2.4316]). The analysis involved 34 amino acid sequences. All positions containing gaps and missing data were eliminated. A total of 443 positions in the final dataset was obtained. Evolutionary analyses were conducted in MEGA6 (Tamura et al., 2013).

According to the phylogenetic tree, the T6SS structural genes of Psa M228 were annotated. The database of Clusters of Orthologous Groups of proteins (COGs) was obtained from the National Center of Biotechnology Information1. A schematic diagram of the T6SS structure was constructed using the Pseudomonas aeruginosa PA01 gene structure map as a reference (Sana et al., 2016).

Generation of Psa M228 Mutant Strains

Mutant strains with deletions of T6SS and its 13 core genes were constructed by the homologous recombination method (Kvitko and Collmer, 2011). In brief, PCR primer synthesis and DNA sequencing were performed by TsingKe Biotech Co., Ltd. (Beijing, China). Psa genomic and plasmid DNAs were isolated using an E.Z.N.A. TM Bacterial DNA Kit (OMEGA, United States) and Plasmid Mini Kit I (OMEGA, United States), respectively. Restriction enzymes were purchased from TaKaRa (Dalian, China). Deletion mutants were constructed using the suicide vector pK18mobsacB (Schafer et al., 1994). The gene fragments including their left and right arms were amplified using the corresponding primers (Supplementary Table S1). The DNA fragments were digested with EcoRI/HindIII and ligated to pK18mobsacB. The vector pK18mobsacB carrying each target gene fragment was inserted by the Km resistance gene and deletion vectors were constructed. Then, the vectors were introduced into the M228 strain by conjugal mating experiments. The first recombinant mutants were screened on LB (Km10-NAL10-Amp 10 μg/mL) and PCR detected using primers SacB-F/R (Supplementary Table S2). The mutant candidates were then screened to remove the suicide plasmid, using LB agar plates containing 20% sucrose without sodium chloride. The results of target gene deletion were confirmed by PCR amplification and DNA sequencing.

Complementation of Psa M228ΔtssM and M228ΔtssJ in Deletion Mutants

The complement mutants were constructed as follows: primers TssM-C/F/R and TssJ-C/F/R (Supplementary Table 1) were used to amplify the full-length tssM (3849 bp) and tssJ (474 bp) genes from the Psa M228 genome, as well as the upstream putative promoters. The PCR products were cloned into the BamHI and NdeI sites of plasmid PDSK, and the plasmids were transformed into E. coli DH5α. The complementation plasmids PDSK-tssM and PDSK-tssJ were extracted from E. coli DH5α according to the manufacturer's instructions in the Plasmid Mini Kit I (OMEGA, United States), and then electroporated into mutants M228ΔtssM and M228ΔtssJ, respectively. The complement mutants were screened by kanamycin resistance on LB agar.

Spread Assays

The spreading ability of the strain Psa M228 and its mutant M228ΔT6SS was determined as previously described (Gao et al., 2016). In brief, the strains Psa M228 and its mutant M228ΔT6SS were successfully transformed with pDSK-GFPuV by electroporation (Huang et al., 2013). Kiwi branches (cv. “Hongyang”) were surface-disinfected by rinsing with running water, soaked in 0.6% sodium hypochlorite solution for 5 min, and rinsed with sterile water again. The base of the petiole was wrapped with absorbent cotton to avoid evaporation. Punctures were made at the main leaf vein 1–2 cm away from the petiole, and GFPuV labeled bacterial culture of 10 μL (diluted to OD600 = 0.1, concentration = 1 × 10⁸ CFU/mL) were dripped onto the wound. The inoculated materials were placed in an artificial climate incubator (photoperiod L/D: 16/8 h; day and night temperature: 16/4°C, relative humidity 95%). The results were observed using a stereo fluorescence microscope (Leica MZ10F, Leica Microsystems, Germany) after 2 days.

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1 ftp://ftp.ncbi.nih.gov/pub/COG/COG2014/static/lists/listAciave.html
Pathogenicity Assays
Pathogenicity of the strain \textit{Psa} M228 and its mutants was determined in kiwi (cv. “Hongyang”), as described previously (Zhao et al., 2019). Healthy branches of kiwi (cv. “Hongyang”) were cut into short branches of about 15 cm each and rinsed with sterile water; the ends were sealed with paraffin to avoid evaporation. In each test, a wound deep to the phloem within with sterile water; the ends were sealed with paraffin to avoid evaporation. Each branch was maintained in an artificial climate incubator with five branches, and each experiment was repeated three times. The branches were maintained in an artificial climate incubator under natural day and night cycles. The lesion lengths of the branches were measured after 30 days. The lesion length is the average of three replicates. Statistical significance was determined by Student’s \( t \)-test.

Electron Microscopic Observation
Healthy leaves from kiwifruit (cv. “Hongyang”) were inoculated separately with bacterial suspensions of \textit{Psa} M228 and its mutant strains M228ΔtssM and M228ΔtssJ, using the same method as described in section “Pathogenicity Assays.” Scanning electron microscopy (SEM) samples were prepared by taking the leaf blade (5–7 mm) from the edge of sick and healthy junctions after 48 and 96 h. Carefully processed as described by Kang (1995), the colonization of bacterial pathogens in leaf tissue was observed using a JSH 6360 SEM (JEOL Ltd., Tokyo, Japan) at 15 kV. Transmission electron microscope (TEM) samples from 96 h culture were prepared in the same way as described by Kang (1995), and alterations in the ultrastructure of the leaf tissue infested by pathogenic bacteria were observed using an HT7700 TEM (HITACHI Company, Tokyo, Japan).

Western Blot Assays
To explore whether the T6SS gene cluster encodes a functioning T6SS, Hcp secretion was assayed by western blot. The wild-type strains \textit{Psa} M228 and mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ were cultured as described in section “Electron Microscopic Observation.” The supernatant and cells were obtained by centrifugation at 13,000 \( \times \) g for 10 min at 4°C. The protein content was determined using a Micro BCA™ Protein Assay Kit (Thermo Fisher Scientific, United States). Extracellular and intracellular proteins were subjected to 12% SDS PAGE and then transferred to PVDF membranes. The immunoblot analysis was performed using anti-VSV-G-tag monoclonal antibody (VSVG) (CB100151, Cali-Bio, United States) and anti-RNA polymerase (RNAP) antibody (W0023, NeoClone, Beijing, China) at a dilution of 1:1000 as the primary and secondary antibodies were goat anti-mouse horseradish peroxidase (HRP) (DY60203) (DIYI BIO TECHNOLOGY, Shanghai, China) and goat anti-rabbit HRP (DY60202) (DIYI BIO TECHNOLOGY, Shanghai, China), respectively, at a dilution of 1:10,000. The differences between the bands of intracellular and extracellular proteins were compared between M228 and the mutants, using RNAP as a control.

Bacterial Competition Assays
\textit{Escherichia coli} DH5\( \alpha \) was transformed with plasmid pBR322 to confer tetracycline resistance and \textit{Bacillus} was transformed with plasmid pMarA to confer erythromycin resistance used for the competition studies. The competition assays were carried out as per the protocol described by MacIntyre et al. (2010). Competitor strains were grown overnight in LB broth supplemented with tetracycline (10 \( \mu \)g/mL) or erythromycin (10 \( \mu \)g/mL). The cells were centrifuged and washed twice with fresh sterile LB broth (10,000 \( \times \) g, 1 min). The washed cells were resuspended in LB broth (OD600 = 0.1, concentration = 1 \( \times \) 108 CFU/mL) and combined with the \textit{Psa} M228 wild-type or mutant strain M228ΔT6SS at a ratio of 1:1. The mixture was cultured at 25°C for 24 h, and then spotted on LB agar plates and incubated at 30°C for 12 h for the competition strain bacterial analysis.

Biofilm Assays
Biofilm assays were performed using the method described by Stepanovic et al. (2000). The results were determined by measuring the absorbance at 570 nm (OD570) using a Thermo Multiskan EX Micro plate Photometer (Thermo Fisher Scientific Inc., United States). The experiments were repeated three times with six replicates per treatment. The absorbance at 570 nm was the average of six replicates. Statistical significance was determined by Student’s \( t \)-test.

Environmental Adaptability Assays
The hydrogen peroxide (H\(_2\)O\(_2\)) tolerance and proteolytic ability were evaluated in accordance with previously described procedures (Molina et al., 2005). The bacterial solutions (diluted to OD600 = 0.1, concentration = 1 \( \times \) 108 CFU/mL) were inoculated into LB broth medium containing H\(_2\)O\(_2\) solution (10 mL, 0.06 mol/L) at a ratio of 1:100. Bacterial growth was measured at 600 nm absorbance every 12 h. A total of 2 \( \mu \)L of the bacterial solution (diluted to OD600 = 0.1, concentration = 1 \( \times \) 108 CFU/mL) was inoculated in the acid hydrolyzed case in a medium plate (1%w/v). Colony diameter was measured after culturing at 25°C for 14 days. All assays were repeated three times, and each treatment was performed in triplicate. Statistical significance was determined by Student’s \( t \)-test.

Quantitative Real-Time Assays
Healthy branches from kiwifruit (cv. “Hongyang”) were prepared as described in section “Pathogenicity Assays.” Suspension of \textit{Psa} M228 and mutant M228ΔT6SS (diluted to OD600 = 0.5, concentration = 5 \( \times \) 108 CFU/mL, 10 \( \mu \)L) were inoculated into the wound made in advance, and cultivated in an artificial climate incubator (the illumination time was 16 h and the dark time was 8 h). The bark of the inoculated wound (deep to phloem) was taken at 2 and 16 h after inoculation. Operations for bacterial RNA extraction, cDNA synthesis, and qPCR were performed as previously described (Broms et al., 2009). The reference genes were gyrB (DNA gyrase subunit B), dusA (tRNA dihydrouridine synthase), and ftrA (transcriptional regulator) (Hirose et al., 2020). Data were analyzed following the protocol...
of the comparative critical threshold (CT) method (Zhang et al., 2014). The expression levels of the target genes (hrpR, hrpZ, hrcC, hopP1, hopH1, and hopM1) at different inoculation times were calculated relative to endogenous control genes using the relative quantification method (Lopez and Pardo, 2005). Quantitative data were presented as fold change mutant strain to the relative quantification method (Lopez and Pardo, 2005). Error bars represent the standard deviation of the ΔΔC value. Statistical significance was determined by Student's t-test. The quantitative real-time PCR (qRT-PCR) was performed in triplicate for technical replicates.

RESULTS

T6SS Gene Structure Diagram of Psa M228

According to the results of the phylogenetic tree constructed from the tssC sequence of 68 species in 38 genera (Supplementary Figure 1), and its simplified phylogenetic tree (Figure 1A), one complete T6SS was found in Psa M228 and belonged to the same branch as the P. aeruginosa PA01 H3-T6SS (HIS-III) gene cluster. Then, the T6SS gene structure diagram in M228 was obtained.

As shown in Figure 1B, there are 13 conserved structural genes such as tssa-tssM necessary for T6SS in M228. The COGs of the cluster are impA, impB, impC, hcp, impE, impH, clpV, vgrG, lip, impJ, impK, and impL, all of which are located in the M228 genome sequence contig054. Among them, tssB and tssC are the needle-sheath structural genes, tssD is the needle-tube structural gene, and tssM, tssJ, and tssL are transmembrane structural genes. However, there is a frameshift mutation in the transmembrane structural protein gene tssM (1272 codons) and the ATP hydrolase gene tssH (854 codons) (Figure 1B).

Construction of T6SS Gene Cluster and 13 Core Genes Deletion Mutants in Psa M228

The previous sequence analysis showed that there is a T6SS gene cluster in the Psa M228 genome, which consists of 13 core genes (tssa-tssM). The function of the gene cluster and core genes in Psa has not been studied yet. Thus, in this study, we began by deleting the T6SS gene cluster and its individual 13 core genes of Psa M228 using Red-recombineering technique (Kvitko and Collmer, 2011), yielding strains M228ΔT6SS and 13 mutants from M228ΔtssA to M228ΔtssM.

T6SS of Psa M228 Is Required for Spreading

The spreading ability of mutant M228ΔT6SS in leaf veins was significantly different from that of wild-type M228. The spreading range of M228 in leaf veins increased with an increase in incubation time, and fluorescence began to weaken after 12 days. However, the spreading ability of M228ΔT6SS in leaf veins was significantly weakened; it failed to expand rapidly in the leaf veins and indicated only weak fluorescence near the inoculation point (Figure 2B).

T6SS of Psa M228 Is Required for Pathogenicity

To clarify the role of T6SS in the pathogenicity of M228, the pathogenicity of mutants lacking the T6SS gene cluster or 13 core genes was assessed by conducting pathogenicity tests on healthy branches of kiwi (cv. “Hongyang”), as described previously (Zhao et al., 2019). The pathogenicity of the mutant M228ΔT6SS was reduced by 69.65% compared with that of the wild-type strain M228 (Figure 2A). All the deletion mutants of 13 core genes of T6SS showed different degrees of pathogenicity reduction compared with that of the wild-type strain M228 (Figure 3A), especially gene tssM and tssJ (Figure 3B). The pathogenicity of mutants M228ΔtssM and M228ΔtssJ mutants were significantly reduced by 78.7 and 71.3%, respectively. However, the complemented strain M228ΔtssM-R and M228ΔtssJ-R restored the pathogenicity to wild-type levels (Figure 3B).

The difference in pathogenicity between M228 and the mutants was also confirmed by SEM (Figure 4) and TEM (Figure 5). After 4 days of inoculation of the host kiwi fruit leaves using wild-type strain M228, a large number of pathogens could be observed in the host tissue (Figure 4B). Serious plasmolysis occurred in the host cells of the pathogenic bacteria colonized area, and electron density of the host cell wall decreased and its chloroplast was degraded (Figures 5B,C). After inoculation of the mutants M228ΔtssM (Figure 4C) and M228ΔtssJ (Figure 4E), only a small amount of bacterial colonization was found in the host tissue at 4 days. The host cell was intact, and the organelles were not significantly degraded due to pathogen colonization (Figures 5D,F). However, there was no significant difference in the colonization and infection ability between the wild-type strain M228 and the complemented strain M228ΔtssM-R (Figures 4D, 5E) and M228ΔtssJ-R (Figures 4F, 5G). The mutants can cause a similar phenomenon to the host cell, indicating that its infection ability is restored to the wild-type level. The results showed that the T6SS gene cluster played an important role in the pathogenicity of Psa in kiwi fruit, especially the core genes tssM and tssJ.

T6SS Gene Cluster Encodes a Functioning T6SS in Psa M228

To investigate the function of T6SS, M228 and the deletion mutant strain, M228ΔT6SS, M228ΔtssM, and M228ΔtssJ were examined for the secretion of Hcp, which is an indication of Functional T6SS in many bacterial species (Ma and Mekalanos, 2010). As shown in Figure 6, Hcp was detected by western blot from both cells and supernatants for the wild-type strain Psa M228, but only detected in the cells and not in the supernatant for mutant strains M228ΔT6SS, M228ΔtssM, and M228ΔtssJ. This result demonstrated that T6SS is essential for the secretion of Hcp in Psa M228. The results also showed that T6SS in the pathogenic strain PsaM228 is functional, and its core genes...
FIGURE 1 | Structure of T6SS gene cluster analysis of the pathogenic strain Psa M228. (A) Molecular phylogenetic analysis by Maximum Likelihood method. The evolutionary history was inferred using the Maximum Likelihood method based on the Le_Gascuel_2008 model (Le and Gascuel, 1993). The bootstrap consensus tree inferred from 1000 replicates was used to represent the evolutionary history of the taxa analyzed (Felsenstein, 1985). Branches corresponding to partitions reproduced in less than 60% bootstrap replicates were collapsed. The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (1000 replicates) is shown next to the branches (Felsenstein, 1985). The initial tree(s) for the heuristic search were obtained automatically by applying (Continued)
Neiher-Join and BioNJ algorithms to a matrix of pairwise distances estimated using a JTT model, and the topology with a superior log likelihood value was selected. A discrete Gamma distribution was used to model evolutionary rate differences among sites (five categories (+G, parameter = 2.4316)). The analysis involved 34 amino acid sequences. All positions containing gaps and missing data were eliminated. A total of 443 positions in the final dataset was obtained. Evolutionary analyses were conducted in MEGA6 (Tamura et al., 2013). According to the results of the phylogenetic tree, one complete T6SS was found in Psa M228 and belonged to the same branch as that of the P. aeruginosa PAO1 H3-T6SS (HIS-III) gene cluster. * indicates the functional T6SS that has been reported. ** indicates the presence of frameshift mutation. The gene products are shown below the arrows. The database of Clusters of Orthologous Groups of proteins (COGs) was obtained from the National Center of Biotechnology Information (see text footnote 1).

**FIGURE 2** Pathogenicity and spreading ability analysis of the pathogenic strain M228 and its mutants M228ΔT6SS. (A) Pathogenicity analysis. The pathogenicity of the mutant M228ΔT6SS mutant was reduced by 69.65% compared with that of the wild-type strain. The lesion length is the average of five replicates. Statistical significance was determined by Student’s t-test. ** Indicates the difference is significant at the 0.01 level. (B) Spreading ability analysis+. The spreading ability of mutant M228ΔT6SS in leaf veins was significantly different from that of wild-type M228. The spreading range of M228 in leaf veins increased with an increase in incubation time, and fluorescence began to weaken after 12 days. However, the spreading ability of M228ΔT6SS in leaf veins was significantly weakened, and it failed to expand rapidly in leaf veins, and only showed weak fluorescence near the inoculation point.

**tssM** and **tssJ** are critical for Hcp secretion and pathogenicity. The internal control protein RNAP was stably expressed in the cell, but was not detected in the supernatant, confirming the reliability of the results.

**The T6SS of Psa M228 Is Used to Compete Against Bacteria**

In order to determine whether the T6SS is functional in competition between bacterial species, a competitive test was...
FIGURE 3 | Pathogenicity analysis of the deletion mutant strains of 13 core genes constitute T6SS. (A) The lesion lengths analysis of the deletion mutants infecting on branches of kiwi. All the deletion mutants of 13 core genes of T6SS showed different degrees of pathogenicity reduction compared with that of the wild-type strain M228, especially gene *tssM* and *tssJ*. The pathogenicity of mutants M228Δ*tssM* and M228Δ*tssJ* mutants were significantly reduced by 78.7% and 71.3%, respectively. However, the complemented strain M228Δ*tssM*-R and M228Δ*tssJ*-R restored the pathogenicity to wild-type levels. The lesion lengths are the average of three replicate. Statistical significance was determined by Student’s *t*-test. **“** Means the difference is significant at the 0.05 level; ****“** means the difference is significant at the 0.01 level. (B) The lesion observation of mutant strain M228Δ*tssM* and M228Δ*tssJ* infecting on branches of kiwi.

FIGURE 4 | Histological observation of the pathogenic strain M228 and its mutants infecting leaves of kiwi by SEM. (A) Healthy kiwi leaves; (B) after 4 days of inoculation of the host kiwifruit leaves using wild-type strain M228, a large number of pathogens could be observed in the host tissue. (C,E) After 4 days of inoculation of the host kiwifruit leaves using the mutants M228Δ*tssM* and M228Δ*tssJ*, only a small amount of bacterial colonization was found in the host tissue; (D,F) after 4 days of inoculation of the host kiwifruit leaves using the mutants M228Δ*tssM*-R and M228Δ*tssJ*-R, a large number of pathogens could be observed in the host tissue, and there was no significant difference in the colonization and infection ability between the wild-type strain M228 and the complemented strains. UE, upper epidermis; PT, palisade tissue; ST, sponge tissue; PH, phloem; XY, xylem; LE, lower epidermis; LVB, lateral vein; B, pathogenic bacteria.
FIGURE 5 | Ultrastructure analysis of kiwi leaves infected by the pathogenic strain M228 and its mutants by TEM. (A) Healthy kiwi leaves; (B,C) after 4 days of inoculation of the host kiwifruit leaves using wild-type strain M228, serious plasmolysis occurred in the host cells of the pathogenic bacteria colonized area, and electron density of the host cell wall decreased and its chloroplast was degraded; (D,F) after 4 days of inoculation of the host kiwifruit leaves using the mutants M228ΔtssM and M228ΔtssJ, the host cell was intact, and organelles were not significantly degraded due to pathogen colonization. (E,G) After 4 days of inoculation of the host kiwifruit leaves using the mutants M228ΔtssM-R and M228ΔtssJ-R, serious plasmolysis occurred in the host cells of the pathogenic bacteria colonized area, and electron density of the host cell wall decreased and its chloroplast was degraded; the mutants could cause a similar phenomenon to the host cell with wild-type strain M228. HCW, host cell wall; N, nucleus; V, vacuole; Chl, chloroplast; M, mitochondria; B, pathogenic bacteria; CW, bacterial cell wall; ‡, host cell plasmolysis.

FIGURE 6 | Immunoblot analysis of Psa M228 and mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ Hcp export. Hcp was detected by western blot from both cells and supernatant for the wild-type strain Psa M228, but was only detected in the cells and not in the supernatant for mutant strain M228ΔT6SS, M228ΔtssM, and M228ΔtssJ. The internal control protein RNAP was stably expressed in the cell, but was not detected in the supernatant, confirming the reliability of the results. Lane M, protein marker.
FIGURE 7 | Competitive, biofilm formation, and environmental adaptability analysis of wild-type strain M228 and mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ. (A) Bacterial competition assay. When the competitor strains were cocultured with wild-type Psa M228 or the complementation mutants M228ΔtssM-R and M228ΔtssJ-R, the viable cell numbers showed an obvious drop from 46.5, 58.0 to 31.0, and 35.2%, respectively. In contrast, when the competitor strains were cocultured with deletion mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ, the survival of competitor bacteria increased up to the same (Continued)
conducted in order to examine whether the mutants lacking the T6SS gene clusters could reduce the number of *E. coli* DH5α and *Bacillus* strains on agar plates. When the competitor strains were cocultured with wild-type *Psa* M228 or the complementation mutants M228ΔtssM-R and M228ΔtssJ-R, viable cell numbers showed an obvious drop from 46.5, 58.0 to 31.0, and 35.2, respectively. In contrast, when the competitor strains were cocultured with deletion mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ, the survival of competitor bacteria was increased up to the same level as that of the control (Figure 7A). Overall, the results showed that the T6SS gene cluster is related to bacterial competition in *Psa* M228.

**T6SS of *Psa* M228 Is Required for Biofilm Formation**

To investigate the function of T6SS, M228, and the deletion mutant strains, M228ΔT6SS, M228ΔtssM, and M228ΔtssJ were examined for biofilm formation. As shown in Figure 7B, the biofilm formation ability of the three mutants was significantly lower than that of the wild-type strain M228 (P < 0.05): M228ΔT6SS decreased by 53.8%, M228ΔtssM by 29.5%, and M228ΔtssJ by 38.7%. After the deletion genes were restored, the biofilm formation ability of the mutants was restored to the same level as that of M228.

**The T6SS of *Psa* M228 Is Required for H$_2$O$_2$ Tolerance and Proteolytic Ability**

In order to determine whether the T6SS is functional in environmental adaptability of *Psa* M228, we examined whether the mutants lacking the T6SS gene clusters could weaken the H$_2$O$_2$ tolerance and proteolytic ability.

The H$_2$O$_2$ tolerance results of mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ was significantly lower than that of the wild-type strain M228, decreased by 59.1, 26.1, and 46.6% at 12 h, respectively, and decreased by 5.0, 8.5, and 7.2% at 24 h, respectively (Figure 7C). The hydrogen peroxide tolerance was significantly affected by T6SS in the early stages. The proteolytic ability of mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ was significantly lower than that of the wild-type strain M228 (P < 0.01), and decreased by 34.8, 25.0, and 20.66%, respectively (Figure 7D). The H$_2$O$_2$ tolerance and proteolytic ability were all restored when the deletion genes were restored.

**T3SS-Related Genes Show Down-Regulated Expression in *Psa* M228ΔT6SS**

The qRT-PCR technique was used to analyze the relationship between the T6SS cluster gene and T3SS-related gene expression. During the interaction between the mutant strain M228ΔT6SS and the host, expression levels of T3SS transcriptional regulatory genes *hrpR*, structural genes *hrpZ*, *hrcC*, *hopP1*, and effector genes *hopM1* and *hopH1* were regulated. Following interaction for 2 h, T3SS-related genes *hrpR*, *hrpZ*, *hrcC*, *hopP1*, *hopH1*, and *hopM1* of the mutant M228ΔT6SS, were down-regulated by 2.01, 1.48, 1.46, 1.14, 1.05, and 1.2 times compared to the wild-type strain; following interaction for 16 h, the expression of the genes was down-regulated by 2.71, 1.76, 1.70, 1.48, 1.78, and 1.83 times, using *gyrB* as an internal reference. A similar down-regulation result was obtained using *dusA* and *ftrA* as internal reference genes (Figure 8). The expression levels of the T3SS related genes were significantly affected by T6SS in the later stage.

**DISCUSSION**

The protein secretion system plays an important role in the pathogenicity and host infection process of many bacterial pathogens (Gerlach and Hensel, 2007). Among them, T3SS plays a vital role in the pathogenicity of many pathogenic bacteria and can directly transfer effectors from pathogenic bacteria to host cells and exerts pathogenic functions (Cornelis et al., 2006; Zhang et al., 2018). Many studies have confirmed that T3SS plays an important role in the phytopathogenic bacteria of a variety of economic crops. Our previous studies also indicated that T3SS promoters are important pathogenic factors (Zhao et al., 2019) by the differential proteome data of strong and weak pathogenic strains. However, in this study, a T6SS cluster consisting of 13 core genes (A–J) was found in *Psa* M228. Although there is a frameshift mutation in the transmembrane structural protein gene *tssM* (1272 codons) and the ATP hydrolyase gene *tssH* (854 codons), the pathogenicity was weaker in all deletion mutants, including mutants M228ΔtssM and M228ΔtssH, than in the wild-type strain M228. It seems that the frameshift mutations of these two genes did not affect their function. This mechanism requires further research.

We further explored whether the gene cluster in *Psa* 228 is a functional T6SS and plays a similar function in pathogenicity as it does in other pathogenic bacteria. Western blot results...
**FIGURE 8** | Differential expression of T3SS-related genes analysis in the process of pathogenic strain *Psa* M228 and its mutant M228ΔT6SS infecting kiwi branches. During the interaction between the mutant strain M228ΔT6SS and the host, the expression levels of the T3SS transcriptional regulatory genes *hrpR*, structural genes *hrpZ, hrcC, hopP1*, and effector genes *hopM1* and *hopH1* were regulated. Following interaction for 2 h, T3SS-related genes *hrpR, hrcC, hopP1, hopH1*, and *hopM1* of the mutant M228ΔT6SS, were down-regulated by 2.01, 1.48, 1.46, 1.14, 1.05, and 1.2 times compared to the wild-type strain; following interaction for 16 h, the expression of the genes was down-regulated by 2.71, 1.76, 1.70, 1.48, 1.78, and 1.83 times, using *gyrB* as internal reference; a similar downregulation result was obtained using *dusA* and *ftrA* as internal reference genes.

showed that the T6SS was necessary for Hcp secretion, showing that the T6SS gene cluster in M228 is functional. To determine whether the observed decrease in pathogenicity is related to a functioning T6SS, we generated 13 core genes of T6SS gene cluster deletion mutants. These mutations all led to attenuated pathogenesis and weakened the spread ability of kiwi plants.
at different levels, especially tssM and tssJ. All these properties of the two gene complementation mutants were restored to the same level as those of the wild-type strain M228. The results indicate that the pathogenicity reduction is related to T6SS. To find out whether the effect of T6SS on pathogenicity is related to T3SS, we also conducted a preliminary study. Quantitative real-time results showed that T3SS genes related to pathogenicity were all down regulated in mutant M228ΔT6SS compared to those in the wild-type strain Psa M228, especially in the later stages of infection. Further research is needed to understand whether T6SS affects pathogenicity by regulating the T3SS-related genes expressed indirectly or by secreting virulence factors directly. The study results are consistent with the experimental results of a single T6SS gene mutation in R. solanacearum (Luo et al., 2016) and the relationship between T3SS and T6SS in the pathogenicity of Edwardsiella tarda (Zhang et al., 2016).

In nature or in the host, microorganisms usually exist in complex environmental conditions. For their own survival, they must compete with other organisms for limited nutrients and space. Studies have shown that many Gram-negative bacteria can kill or inhibit other bacteria through T6SS (Alcoforado et al., 2015). For example, the human pathogen Vibrio cholerae kills E. coli by secreting virulence factors through T6SS (MacIntyre et al., 2010). In addition, recent studies have shown that T6SS plays an important role in P. aeruginosa and E. coli cells. A variety of antibacterial toxins that plays an important role in bacterial competition can be delivered to the target cell through T6SS (Basler et al., 2013; Brunet et al., 2013; Durand et al., 2014; Alcoforado et al., 2015). It has been reported that E. coli is susceptible to T6SS and can be killed by the antibacterial toxin secreted by T6SS. Therefore, E. coli is often used as an ideal competitor (MacIntyre et al., 2010; Zheng et al., 2011; Weber et al., 2013). Bacillus is one of the most abundant species of plant endophytes (Gadhave et al., 2018; Gordon et al., 2020). Therefore, in this work, we used E. coli and Bacillus as competitors. Our results showed that T6SS is required for efficient bacterial competition. Moreover, how the competition is carried out in vitro requires further research and needs to be confirmed using in vivo survival assays. In addition, we investigated the role of the T6SS in bacterial biofilm formation, which is essential during infection and pathogenic processes (Yu et al., 2016). Our research indicated that T6SS plays an important role in the pathogenicity of Psa M228. These results present novel insights regarding the pathogenesis of Psa.

The adaptability of plants to various pressures from the natural environment needs to be achieved by regulating the pressure signal transduction pathways. When plants are attacked by pathogenic bacteria, it produces a series of stress responses, including the oxidative burst (Nicaisse and Candresse, 2017; Kanwar and Jha, 2019; Guo and Li, 2000). Previous studies have suggested that these reactive oxygen species have pathological functions such as inhibiting the growth of bacterial and fungal pathogens (Jiang et al., 2011). Our study showed that compared to the wild-type strain M228, the growth of the deletion mutants M228ΔT6SS, M228ΔtssM, and M228ΔtssJ was significantly inhibited under H2O2 stress. The deletion of T6SS, tssM, and tssJ from Psa M228 may lead to reduced tolerance to H2O2, thereby affecting the survival of the pathogenic bacteria. The results indicate that T6SS may be associated with resistance to environmental stress in pathogenic bacteria. However, this requires further investigation.

CONCLUSION

Here, we found a T6SS gene cluster consisting of 13 core genes (A-M) in the genome of Psa M228 based on a genome-wide analysis. We provide evidence that T6SS plays an important role in pathogenesis, bacterial competition, biofilm formation, and environmental adaptability. The next step is to clarify the assembly and secretion mechanism of T6SS and identify the virulence factors secreted by T6SS in Psa. This research has deepened our understanding of the pathogenic mechanism of Psa.

DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

AUTHOR CONTRIBUTIONS

NW, NH, and LH designed the research. NW, NH, RT, XG, JC, ZW, and YL performed the experiments. NW, NH, RT, ZW, and YL analyzed the data. NW and NH wrote the manuscript. 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.2021.627785/full#supplementary-material
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