Pseudomonas cannabina pv. alisalensis TrpA Is Required for Virulence in Multiple Host Plants

Nanami Sakata, Takako Ishiga and Yasuhiro Ishiga*

Faculty of Life and Environmental Sciences, University of Tsukuba, Tsukuba, Japan

Pseudomonas cannabina pv. alisalensis (Pca) causes bacterial leaf spot and blight of Brassicaceae and Poaceae. We previously identified several potential Pca virulence factors with transposon mutagenesis. Among these a trpA mutant disrupted the tryptophan synthase alpha chain, and had an effect on disease symptom development and bacterial multiplication. To assess the importance of TrpA in Pca virulence, we characterized the trpA mutant based on inoculation test and Pca gene expression profiles. The trpA mutant showed reduced virulence when dip- and syringe-inoculated on cabbage and oat. Moreover, epiphytic bacterial populations of the trpA mutant were also reduced compared to the wild-type (WT). These results suggest that TrpA contributes to bacterial multiplication on the leaf surface and in the apoplast, and disease development. Additionally, several Brassicaceae (including Japanese radish, broccoli, and Chinese cabbage) also exhibited reduced symptom development when inoculated with the trpA mutant. Moreover, trpA disruption led to downregulation of bacterial virulence genes, including type three effectors (T3Es) and the phytotoxin coronatine (COR), and to upregulation of tryptophan biosynthesis genes. These results indicate that a trade-off between virulence factor production and Pca multiplication with tryptophan might be regulated in the infection processes.

Keywords: Pseudomonas cannabina pv. alisalensis, tryptophan, type three secretion system, coronatine, cabbage, oat

INTRODUCTION

The foliar plant bacterial pathogen Pseudomonas syringae causes economically important diseases in a wide range of plants (Agrios, 2005). P. syringae colonizes leaf surfaces (epiphytic) of host plants, enters natural opening sites, including stomata, and then multiplies in the leaf interior (apoplast) (Xin and He, 2013). During infection processes, P. syringae suppresses plant basal defense by using virulence factors, such as specialized protein secretion systems, toxins, plant hormones, bacterial surface attachment factors, flagella, and siderophores (Xin and He, 2013). P. syringae pv. tomato (Pst) DC3000 also infects Arabidopsis (Whalen et al., 1991), and is used as a model pathogen to study plant-bacterial interactions. Pst DC3000 is a highly aggressive pathogen once inside host tissue, and uses many type three effectors (T3Es) and the phytotoxin coronatine (COR; Buell et al., 2003; Feil et al., 2005). The function of these two Pst DC3000 virulence factors have been well characterized at the molecular level. However, Boch et al. (2002) demonstrated that a wide range of plant-induced loci in Pst DC3000 included not only virulence associated genes such as...
hrp genes and COR biosynthesis genes, but also genes involved in stress tolerance, polysaccharide synthesis, nutrient uptake, amino acid assimilation, and carbon metabolism. Although virulence associated genes related to type three secretion system (T3SS) and COR have been investigated well in the P. syringae virulence, functions of other genes including amino acid metabolism remain largely unclear.

_Pseudomonas cannabina pv. alisalensis_ (Pcal) causes bacterial leaf spot and blight of Brassicaceae and Poaceae (Takikawa and Takahashi, 2014). _Pcal_ was formally classified as _P. syringae pv. maculicola_ (Psm). Although _Pcal_ and _Psm_ have similar characteristics, these two pathogens are defined by some bacteriological characteristics, genetic traits, and their ability to infect monocot plants such as oat (_Avena sativa_ and timothy (_Phleum pretense_)) (Cintas et al., 2002; Bull et al., 2010; Takikawa and Takahashi, 2014). Our recent study identified several potential _Pcal_ virulence factors, including T3SS, membrane transporters, transcriptional factors, and amino acid metabolism (Sakata et al., 2019). Among these, the mutants which were disrupted in amino acid metabolism showed no pathogenicity, similar to a T3SS mutant (Sakata et al., 2019). A _trpA_ (encoding tryptophan synthase alpha chain) mutant exhibited reduced disease symptom development and bacterial multiplication (Sakata et al., 2019). Helmann et al. (2019) conducted a genome wide screening to identify _P. syringae pv. syringae_ (Pss) B728a virulence factors, and identified that _trpA_ contributes to bacterial fitness on both the leaf surface and in the apoplast. Tryptophan is the least abundant amino acid in leaf exudates from many plant species (Morgan and Tukey, 1964; Rico and Preston, 2008). Although amino acid metabolism is essential for bacterial growth, the function of amino acid metabolism in plant bacterial virulence has not been investigated.

Here, we showed that TrpA contributes to _Pcal_ virulence in successful infection processes. Moreover, _trpA_ mutant expression profiles analysis showed downregulation of virulence related genes, including T3Es and COR, and upregulation of tryptophan biosynthesis related genes compared to WT. These results suggest that trade-off between virulence factor production and tryptophan biosynthesis might be present in bacterial infection processes.

**MATERIALS AND METHODS**

**Bacterial Strains, Plasmids, and Growth Conditions**

The bacterial strains and plasmids used in this study are described in Supplementary Table 1. _P. cannabina pv. alisalensis_ strain KB211 (_Pcal_ KB211) was used as the pathogenic strain to inoculate cabbage, oat, Japanese radish, broccoli, and Chinese cabbage. _Pcal_ wild-type (WT) was grown on King’s B (KB; King et al., 1954) medium at 28°C. NB35, NF2, NF34, NI13, NH11, NM37, and NN31 were grown on KB containing kanamycin (10 µg/ml) (Km). _trpA_ mutant complemented with pDSK-trpA was grown on KB containing Km (10 µg/ml) and gentamicin (Gen) (25 µg/ml) (Supplementary Table 1). Before _Pcal_ inoculation, bacteria were suspended in sterile distilled H₂O, and the bacterial cell densities at 600 nm (OD₆₀₀) were measured using a Biowave CO8000 Cell Density Meter (Funakoshi, Tokyo, Japan).

**Bacterial in vitro Growth Measurements**

Wild-type, the _trpA_ mutant, and the _trpA_ mutant complemented with pDSK-trpA were grown at 28°C on Luria-Bertani (LB; Sambrook et al., 1989) medium. The bacterial suspensions were standardized to an OD₆₀₀ of 0.01 with LB, and bacterial growth was measured at OD₆₀₀ after 24 h. WT and the _trpA_ mutant were also grown at 28°C in mannitol-glutamate (MG; Keane et al., 1970) medium. The bacterial suspensions were standardized to an OD₆₀₀ of 0.1. L-tryptophan (FUJIFILM Wako Pure Chemical Corporation, Osaka, Japan) was used to co-incubate with MG medium at 1, 10, and 100 µM, respectively. Bacterial growth was measured at OD₆₀₀ for 24 h.

**Plant Materials**

Plants used for _Pcal_ virulence assays include cabbage (_Brassica oleracea var. capitata_) cv. Kinkei 201, oat (_Avena strigosa_) cv. Hayoat, Japanese radish (_Raphanus sativus var. longipinnatus_) cv. Natsutsukasa, Chinese cabbage (_Brassica rapa var. pekinensis_) cv. Akimeki, and broccoli (_Brassica oleracea var. Italica_). Midoribue. All plants were grown from seed at 23–25°C with a light intensity of 200 µE/m²/s and a 16 h light/8 h dark photoperiod. Plants were used for dip- and spray-inoculation assays around two weeks after germination, and for syringe-inoculation assays around 3 weeks after germination. For flood-inoculation, cabbage seeds were germinated and grown on 1/2 strength Murashige and Skoog (MS) medium (0.3% phytagel) with Gamborg vitamins (Sigma-Aldrich, St. Louis, MO, United States). Cabbage seedlings were incubated in a growth chamber at 24°C with a light intensity of 200 µE/m²/s and a 12 h light/12 h dark photoperiod, and used for the inoculation assays 2 weeks after germination.

**Bacterial Inoculation**

To assay for disease on cabbage, oat, Japanese radish, broccoli, and Chinese cabbage plants, dip-inoculations were conducted by soaking seedlings in bacterial suspensions (5 × 10⁷ CFU/ml) containing 0.025% Silwet L-77 (OSI Specialities, Danbury, CT, United States). The seedlings were then incubated in growth chambers at 85–95% RH for the first 24 h, then at 80–85% RH for the rest of the experimental period. Disease symptoms were photographed at 5 days post-inoculation (dpi). To assess bacterial growth in all plants, the internal bacterial population was measured after dip-inoculation. Inoculated seedlings were collected, and two inoculated leaves were measured chronologically. The leaves were surface-sterilized with 10% H₂O₂ for 3 min. After washing three times with sterile distilled water, the leaves were homogenized in sterile distilled water, and diluted samples were plated onto solid KB agar medium. Two or three days after dilution sample plating, the bacterial colony forming units (CFUs) were counted and normalized as CFU per gram, using the total leaf weight. The bacterial populations at 0 dpi were estimated using
leaves harvested 1-hour post-inoculation (hpi) without surface-sterilization. The bacterial populations were evaluated in at least three independent experiments.

For syringe-inoculation, cabbage and oat leaves were syringe-inoculated with bacterial suspensions (5 \times 10^6 or 5 \times 10^7 CFU/ml) with a 1-ml blunt syringe. L-tryptophan was co-inoculated with the \textit{trpA} mutant at 100 \mu M, 1 mM, 10 mM, and 50 mM, respectively. The plants were then incubated at 70–80% RH for the rest of the experimental period. Leaves were removed and photographed at 5 dpi. To assess bacterial growth in cabbage, the internal bacterial population was measured after syringe-inoculation. Leaf disks were harvested using a 3.5 mm-diameter cork-borer from syringe-infiltrated leaf zones. To assess bacterial growth in oat, leaf pieces were cut from syringe-infiltrated leaf zones and the area (cm²) measured. The bacterial CFUs were counted and normalized as CFU per cm² of leaf tissue. The bacterial populations were evaluated in at least three independent experiments.

**Epiphytic Bacterial Growth Assay**

Flood-inoculation was conducted as described previously (Ishiga et al., 2011). Briefly, 50 ml of bacterial suspension (1 \times 10^8 CFU/ml) made in sterile distilled H₂O containing 0.025% Silwet L-77 (OSI Specialties, Danbury, CT, United States) was dispensed onto a plate containing 2-week-old cabbage seedlings grown on 1/2 strength MS medium for uniform contamination. The expression profiles were evaluated in at least three independent experiments.

**Monitoring Bacterial Gene Expression in planta**

To analyze \textit{Pcal} gene expression profiles during infection, we syringe-inoculated cabbage plants with \textit{Pcal} at 5 \times 10^7 CFU/ml, and at 3 and 6 hrs the total RNAs including plant and bacterial RNAs were extracted from infected leaves and purified. Total RNA extraction and real-time quantitative RT-PCR (RT-qPCR) were done as described previously (Ishiga and Ichinose, 2016). Two micrograms of total RNA were treated with gDNA Remover (TOYOBO) to eliminate genomic DNA, and the DNase-treated RNA was reverse transcribed using the ReverTra Ace qPCR RT Master Mix (TOYOBO). The cDNA (1:10) was then used for RT-qPCR using the primers shown in Supplementary Table 2 with THUNDERBIRD SYBR qPCR Mix (TOYOBO) on a Thermal Cycler Dice Real Time System (Takara Bio). \textit{Pcal} KB211 outer membrane porin \textit{F} (\textit{opfF}) and \textit{recombinase A} (\textit{recA}) were used to normalize gene expression. Gene expression of the \textit{trpA} mutant was calculated as a relative value of WT expression. The reagent blank (no-template) controls were used to detect contamination. The expression profiles were evaluated in four independent samples.

**RESULTS**

**TrpA Is Involved in \textit{Pcal} Pathogenicity**

To investigate functions of amino acid metabolism in \textit{Pcal} virulence, we conducted inoculation assay using six amino acid metabolism mutants (Sakata et al., 2019). These six mutants (NF2, NF34, NH11, NI13, NM37, and NN31; see Supplementary Table 1) and a T3SS mutant (NB35) were syringe-inoculated into cabbage to investigate which mutants were most involved in \textit{Pcal} virulence. NH11, NM37, and NN31 showed necrosis similar to WT. NB35, NF34, and NI13 only showed chlorosis. Only the NF2 mutant, which is mutated in the \textit{trpA} (encoding tryptophan synthase alpha chain), showed no symptoms (Figure 1A). Moreover, \textit{trpA} mutant bacterial populations were severely reduced among all mutants, including the T3SS mutant (Figure 1B).

**TrpA Contributes to \textit{Pcal} Multiplication in the Apoplast to Cause Disease**

\textit{trpA} is apparently dispensable for \textit{Pcal} growth in rich LB medium, since no growth difference was observed between the WT and the \textit{trpA} mutant (Supplementary Figure 1). Cabbage
inoculated with WT showed severe chlorosis, but the trpA mutant induced no symptoms (Figure 2A). Although bacterial populations of the WT and the trpA mutant complemented with pDSK-trpA reached around $1 \times 10^8$ CFU/ml when dip-inoculated onto cabbage, trpA mutant populations were $10^4$ times less (Figure 2B). Moreover, the trpA mutant multiplication defect was observed within 1 dpi (Figure 2B). To further investigate whether the trpA mutant is impaired in apoplastic growth and pathogenicity, syringe-infiltration was used to bypass stomatal defense. When infiltrated directly into the cabbage apoplast, trpA mutant disease development and bacterial populations were decreased compared to the WT (Figures 2C,D).

Importantly, Pcal is pathogenic on monocot plants, such as oat, as well as cruciferous species (Ishiyama et al., 2013). Oat plants inoculated with the trpA mutant also showed reduced symptoms as well as bacterial populations for both dip- and syringe-inoculation methods (Figures 2E–H).

### TrpA Contributes to Multiplication on the Leaf Surface and Disease

Globally, the microbiota on plant leaf surfaces is estimated to around $10^{26}$ bacterial cells (Lindow and Brandl, 2003). The plant surface is generally considered to be suboptimal for microbes (Lindow and Brandl, 2003). Because the trpA mutant is auxotrophic, and does not grow on minimal medium (MG medium) (Sakata et al., 2019), the trpA mutant is likely to be affected by other plant microbiomes, which compete for limited nutrients. Thus, to rule out the possibility that the trpA mutant showed reduced virulence because of a microbiome effect, we conducted flood-inoculation onto cabbage, which can assay the plant-bacterial interaction with a sterility test (Ishiga et al., 2011), and investigated epiphytic bacterial populations. The trpA mutant showed reduced population via flood-inoculation (Figure 3).

### Tryptophan Is an Essential Growth Resource

To investigate whether exogenous tryptophan application restores this virulence impairment, we first measured bacterial growth in MG medium with tryptophan. The trpA mutant was growth deficient in MG medium (Figure 4A). Co-inoculation with only 1 $\mu$M tryptophan restored trpA mutant bacterial growth to the WT level in vitro (Figure 4A).

We further tested whether exogenous tryptophan application restores trpA mutant virulence on cabbage. Co-inoculation of the trpA mutant with tryptophan restored virulence in a dose-dependent manner (Figures 4B,C). Although only 10 $\mu$M tryptophan restored trpA mutant bacterial growth to the WT level in vitro, at least 50 mM tryptophan was required for the trpA mutant to multiply to a similar level as WT in cabbage by 5 dpi (Figure 4B). Although bacterial multiplications were recovered, 50 mM exogenous application could not fully recover disease symptoms (Figure 4C and Supplementary Figure 2A). To further investigate symptom development and bacterial populations, we conducted inoculation with or without tryptophan (50 mM) in a time-dependent manner. Exogenous tryptophan application partially recovered trpA mutant multiplication at 1 and 3 dpi (Supplementary Figure 2B).

### trpA Mutant Virulence Genes Show Reduced Expression During Infection

Ten $\mu$M exogenous tryptophan application recovered bacterial growth in vitro, while at least 50 mM of tryptophan was required for growth recovery in planta (Figures 4A,B). Therefore, we assumed that tryptophan is not only a nutrient but also involved in regulating bacterial virulence gene expression. We firstly examined Pcal gene expression profiles in culture medium. As we expected, T3Es genes (including avrPto, hopM1,
FIGURE 2 | Disease phenotypes and bacterial populations of *Pseudomonas cannabina* pv. *alisaensis* KB211 WT and the *trpA* mutant in cabbage and oat after dip- and syringe-inoculation. Disease symptoms (A) and bacterial populations (B) in cabbage dip-inoculated with WT, the *trpA* mutant, and the *trpA* mutant complemented with pDSK-trpA. Disease symptoms (C) and bacterial populations (D) in cabbage syringe-inoculated with WT, the *trpA* mutant, and the *trpA* mutant complemented with pDSK-trpA. Disease symptoms (E) and bacterial populations (F) in oat dip-inoculated with WT, the *trpA* mutant, and the *trpA* mutant complemented with pDSK-trpA. Disease symptoms (G) and bacterial populations (H) in oat syringe-inoculated with WT, the *trpA* mutant, and the *trpA* mutant complemented with pDSK-trpA. Cabbage and oat were dip-inoculated with 5 × 10^7 CFU/ml of inoculum containing 0.025% Silwet-L-77 and syringe-inoculated with 5 × 10^5 CFU/ml of inoculum, respectively. Bacterial concentrations in the plant leaves were evaluated at 0, 1, 3, and 5 dpi. The leaves were photographed 5 dpi. Vertical bars indicate the standard error for at least three independent experiments. Different letters indicate a significant difference among treatments based on a Tukey's HSD test (p < 0.05). Scale bar shows 2 cm.
Since the trpA Host Plants TrpA contributes to disease on multiple expression in the trpA trpA during infection. The trpB gene expression profiles. × 5 conducted syringe-inoculation into cabbage with these strains at 5 × 10^7 CFU/ml and examined earlier time points including 3, 6, and 9 hpi. (Supplementary Figure 4). Therefore, we conducted syringe-inoculation into cabbage with these strains at 5 × 10^7 CFU/ml and extracted the total RNAs at 3 and 6 hpi. T3Es genes (Figures 5A–C) and COR biosynthesis related genes (Figures 5D–F) exhibited reduced expression in the trpA mutant compared to WT. We also investigated tryptophan biosynthesis gene expression profiles. trpB, trpE, trpG, and trpI showed greater expression in the trpA mutant than in the WT (Figures 5G–J).

**TrpA Contributes to Disease on Multiple Host Plants**

Since the trpA mutant showed no pathogenicity on both cabbage and oat, we hypothesized that TrpA was required for pathogenesis on multiple host plants. We therefore inoculated the WT and trpA mutant onto several host plants, including Japanese radish, broccoli, and Chinese cabbage, and measured bacterial populations. The trpA mutant showed reduced symptom development and bacterial multiplication in all host plants (Figure 6 and Supplementary Figure 5).

**DISCUSSION**

Plant pathogens deploy multiple virulence factors for successful infection. This study was performed to understand the role of TrpA in Pcal virulence. Overall, this study showed that TrpA contributes to Pcal pathogenesis and is important for successful infection in multiple host plants. Importantly, trpA disruption leads to downregulation of bacterial virulence genes, including T3Es and COR, and to upregulation of tryptophan biosynthesis related genes, demonstrating the importance of tryptophan biosynthesis in bacterial pathogenesis.

Among six amino acid metabolism mutants we identified (Sakata et al., 2019), the trpA mutant showed no pathogenesis (Figure 1). These results indicate that tryptophan biosynthesis is critical for Pcal virulence. Schreiber et al. (2012) conducted a high-throughput forward genetic screen and demonstrated the nutritional requirements of Psm ES4326 (Pcal ES4326) in Arabidopsis. Amino acid auxotrophs showed dramatically reduced bacterial populations following spray inoculation (Schreiber et al., 2012), suggesting that free amino acids are limited on leaf surfaces and are important for growth. Together, these results underscore the importance of amino acid metabolism in plant bacterial pathogen virulence.

Our data showed that TrpA contributes to multiplication on the leaf surface and in the apoplast, causing disease (Figures 2, 3). Helmann et al. (2019) conducted RB-TnSeq to define the fitness contributions of Pss B728a genes. Consistent with our results, they demonstrated that genes within the tryptophan biosynthetic pathway had the greatest effect on fitness both on the leaf surface and in the apoplast (Helmann et al., 2019). Generally, the plant surface is considered suboptimal for microbes, which provides limited nutrient resources to bacterial colonists (Lindow and Brandl, 2003). Morgan and Tukey (1964) demonstrated that tryptophan was not detected in several plant species among 20 amino acids detected in plant leachates, suggesting a pressing need for its synthesis by bacterial colonists. Moreover, the ability of Pcal to synthesize tryptophan strongly influences bacterial proliferation in multiple host plants (Figure 6). Together, tryptophan biosynthesis is an essential process for successful Pcal infection on multiple host plants.

Bacterial multiplication of the trpA mutant was restored to WT levels with only 10 µM tryptophan in vitro (Figure 4A). Exogenous tryptophan application rescued the trpA mutant bacterial multiplication defect in planta, but at least 50 mM tryptophan was required (Figure 4B). Therefore, we assumed that the reduced virulence of the trpA mutant is involved in plant-bacterial interactions, as well as lack of nutrition. Transcripts of T3Es and COR related genes were reduced during trpA mutant infection compared to WT (Figure 5). Conversely, tryptophan
biosynthesis related genes showed greater expression during trpA mutant infection than WT (Figure 5). The tryptophan biosynthesis pathway and genes have been proposed to be highly conserved in proteobacteria (Bae et al., 1989; Essar et al., 1990; Gussin, 2004). In P. syringae, the trpBA operon is regulated by TrpI, a LysR-type transcriptional activator whose gene is transcribed divergently (Auerbach et al., 1993). When tryptophan concentration is low, indoleglycerol phosphate (InGP) accumulates and TrpI assumes its active conformation, where it is able to bind at two operator sites in the trpI-trpBA intergenic region (Chang and Crawford, 1990; Merino et al., 2008). Our results demonstrated that these genes showed greater expression in the trpA mutant than in the WT (Figure 5).

Taken together, our results suggest that trpA mutation directly affects tryptophan biosynthesis genes, and indirectly affects virulence factor related genes. One possible explanation how mutation in trpA affects T3Es and COR related genes is trade-off between nutrition acquisition and virulence. Several studies have demonstrated that nutrient assimilation during host infection is critical for pathogenesis (Brown et al., 2008; Eisenreich et al., 2010; Barbier et al., 2011; Schoen et al., 2014). In the plant pathogen Ralstonia solanacearum, exopolysaccharide (EPS) (which are critical for disease symptom production) biosynthesis and secretion represent a significant energetic cost for the pathogen, resulting in reduced bacterial growth (Peyraud et al., 2016). Furthermore, the animal pathogen Salmonella

**FIGURE 4** | Effect of exogenous tryptophan application on trpA mutant growth. (A) Growth of Pseudomonas cannabina pv. alisalensis KB211 WT and the trpA mutant in MG medium. WT and the trpA mutant were adjusted to an OD600 of 0.1 in MG medium. Additionally, the trpA mutant was co-incubated with tryptophan; 1, 10, and 100 μM, respectively. Then, cultures were incubated with shaking at 28°C. Bacterial populations were quantified at 24 h. (B) Bacterial populations in cabbage syringe-inoculated with WT and the trpA mutant. Cabbage plants were syringe inoculated with 5 × 10^5 CFU/ml of WT and the trpA mutant. The trpA mutant was co-inoculated with tryptophan; 100 μM, 1 mM, 10 mM, and 50 mM, respectively. Bacterial concentrations in the plant leaves were evaluated at 5 dpi. (C) Disease symptoms on cabbage co-inoculated with tryptophan. Cabbage plants were syringe-inoculated with 5 × 10^5 CFU/ml of WT and the trpA mutant. The trpA mutant was co-inoculated with 100 μM, 1, 10, and 50 mM tryptophan, respectively. The leaves were photographed at 5 dpi. Scale bar shows 2 cm.
FIGURE 5 | Expression profiles of bacterial virulence genes and tryptophan biosynthesis genes during Pseudomonas cannabina pv. alisalensis KB211 WT and trpA mutant infection. Cabbage plants were syringe-inoculated with $5 \times 10^7$ CFU/ml of WT and the trpA mutant and total RNAs were collected 3 and 6 h after inoculation. Expression profiles of type three effector (T3Es) genes [including $avrPto$ (A), $avrE1$ (B), and $hopM1$ (C)] and COR biosynthesis related genes [including $cmaA$ (D), $cfl$ (E), and $corR$ (F)] were investigated. Additionally, expression profiles of tryptophan biosynthesis related genes [including $trpB$ (G), $trpE$ (H), $trpG$ (I), and $trpI$ (J)] were also investigated. Total RNA was extracted for use in real-time quantitative reverse transcription-polymerase chain reaction (RT-qPCR) with gene-specific primer sets (Supplementary Table 2). Expression was normalized using oprE and recA. Vertical bars indicate the standard error for at least six biological replicates. Different letters indicate a significant difference among treatments based on a Tukey’s HSD test ($p < 0.05$).
also balances the trade-off between fast growth and T3SS production (Ackermann et al., 2008; Diard et al., 2013). Taken together, the trade-off between growth and virulence should be considered in plant and bacterial pathogen interactions. Although further investigation will be necessary to understand how mutation in trpA affect the bacterial virulence factors, trade-off between virulence factor production and bacterial proliferation with nutrients such as tryptophan might be present in Pcal infection processes.

In conclusion, these data strongly suggest that TrpA contributes to bacterial multiplication on the leaf surface and in the apoplast, and contributes to disease in multiple host plants. Furthermore, trpA mutation leads to downregulation of virulence genes related to T3Es and COR. Since most amino acids are apparently present at relatively low apoplastic concentrations (O’Leary et al., 2016), it is expected that lack of amino acid metabolites function causes reduced virulence in addition to growth defects. Our findings suggest that amino acid metabolites can be targeted for developing new disease control strategies.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repositoryrepositories and accession number(s) can be found below: https://figshare.com/s/753ec4995e4086b047da.

AUTHOR CONTRIBUTIONS

NS and YI designed the experiments and wrote the manuscript. All authors performed the experiments.

FUNDING

This work was supported, in part, by Japan Society for the Promotion of Science (JSPS), Grant Number: 19K06045 (YI), by Sustainable Food Security Research Project in the form of an operational grant from the National University Corporation (YI), and by Japan Science and technology Agency (JST), Exploratory Research for Advanced Technology (ERATO) NOMURA Microbial Community Control Project, Grant Number: JPMJER1502 (YI).

ACKNOWLEDGMENTS

We thank Dr. Christina Baker for editing the manuscript. Pcal was kindly given from the Nagano vegetable and ornamental crops experiment station, Nagano, Japan.

SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.659734/full#supplementary-material
REFERENCES

Ackermann, M., Stecher, B., Freed, N. E., Songhet, P., Hardt, W.-D., and Doebele, M. (2008). Self-destructive cooperation mediated by phenotypic noise. *Nature* **454**, 987–990. doi: 10.1038/nature06706

Agrios, G. N. (2005). *Plant Pathology*, 5th Edn. Amsterdam: Elsevier Academic Press.

Auerbach, S., Gao, J., and Gussin, G. N. (1993). Nucleotide sequences of the trp, trpA, and trpA genes of *Pseudomonas syringae*: positive control unique to fluorescent pseudomonads. *Gene* **123**, 25–32. doi: 10.1016/0378-1119(93)90534-a

Bae, Y. M., Holmgren, E., and Crawford, I. P. (1989). Rhizobium meliloti anthranilate synthase gene: cloning, sequence, and expression in *Escherichia coli*. *J. Bacteriol.* **171**, 3471–3478. doi: 10.1128/jb.171.6.3471-3478.1989

Barbier, T., Nicolas, C., and Letesson, J. J. (2011). *Brucella* adaptation and survival at the crossroad of metabolism and virulence. *FEBS Lett.* **585**, 2929–2934. doi: 10.1016/j.febslet.2011.08.011

Boch, J., Joardar, V., Gao, L., Robertson, T. L., Lim, M., and Kunkel, B. N. (2002). Identification of *Pseudomonas syringae* pv. tomato genes induced during infection of *Arabidopsis thaliana*. *Mol. Microbiol.* **44**, 73–88. doi: 10.1046/j.1365-2958.2002.02877.x

Brown, S. A., Palmer, K. L., and Whiteley, M. (2008). Revisiting the host as a growth medium. *Nat. Rev. Microbiol.* **6**, 657–666. doi: 10.1038/nrmicro1953

Buell, C. R., Joardar, V., Lindeberg, M., Selengut, J., Paulsen, I. T., Gwinn, M. L., et al. (2003). The complete genome sequence of the *Arabidopsis* and *tomato* pathogen *Pseudomonas syringae* pv. *tomato* DC3000. *Proc. Natl. Acad. Sci. U.S.A.* **100**, 10181–10186. doi: 10.1073/pnas.1731982100

Bull, C. T., Manceau, C., Lydon, J., Kong, H., Vinatorz, B. A., and Fischer-Le Saux, M. (2010). *Pseudomonas cannabina* pv. *cannabina* pv. *nov.* and *Pseudomonas cannabina* pv. *alissalensis* (Cintas Koike and Bull, 2000) comb. nov., are members of the emended species *Pseudomonas cannabina* (ex Šutić & Dowson 1959) Gardan, Shaflk, Belouin, Brosch, Grimont & Grimont 1999. *Syst. Appl. Microbiol.* **33**, 105–115. doi: 10.1016/j.syapm.2010.02.001

Chang, M., and Crawford, I. P. (1990). The roles of indoleglycerol phosphate and the Trp1 protein in the expression of trpBA from *Pseudomonas aeruginosa*. *Nucleic Acids Res.* **18**, 979–988. doi: 10.1093/nar/18.4.979

Cintas, N. A., Koike, S. T., and Bull, C. T. (2002). A new pathovar, *Pseudomonas* *Brucella* *syringae* pv. *Brucella* *syringae* pv. *Brucella* *syringae* pv. *alissalensis*. *J. Gen. Plant Pathol.* **79**, 155–157. doi: 10.1007/s10327-013-0427-9

Keene, P. J., Kerr, A., and New, P. B. (1970). Crown gall of stone fruit ii. identification and nomenclature of agrobacterium isolates. *Aust. J. Bio. Sci.* **23**, 301–307.

Lindow, S. E., and Brandl, M. T. (2003). Microbiology of the phyllosphere. *Appl. Environ. Microbiol.* **69**, 1875–1885. doi: 10.1128/aem.69.6.1875-1883.2003

Merino, E., Jensen, R. A., and Yanofsky, C. (2008). Evolution of bacterial trp operons and their regulation. *Curr. Opin. Microbiol.* **11**, 78–86. doi: 10.1016/j.copmi.2008.02.005

Morgan, J. V., and Tukey, H. B. (1964). Characterization of leachate from plant foliage. *Plant Physiol.* **39**, 590–593. doi: 10.1104/pp.39.4.590

O’Leary, B. M., Neale, H. C., and Gelisius, C. M. (2016). Early changes in apoplast composition associated with defense and disease in interactions between *Phaseolus vulgaris* and the halo blight pathogen *Pseudomonas syringae* pv. *phaseolicola*. *Plant Cell Environ.* **39**, 2172–2184. doi: 10.1111/pce.12770

Palmer, D. A., and Bender, C. L. (1993). Effects of environmental and nutritional factors on production of the polyketide phytotoxin coronatine by *Pseudomonas syringae* pv. *glycinea*. *Appl. Environ. Microbiol.* **59**, 1619–1626. doi: 10.1128/AEM.59.5.1619-1626.1993

Peyraud, R., Cottret, L., Marmiesse, L., Gouzy, J., and Genin, S. (2016). A resource allocation trade-off between virulence and proliferation drives metabolic versatility in the plant pathogen *Ralstonia solanacearum*. *PLoS Pathog.* **12**(e1005939). doi: 10.1371/journal.ppat.1005939

Rico, A., and Preston, G. M. (2008). *Pseudomonas syringae* pv. *tomato* DC3000 uses constitutive and apoplast-induced nutrient assimilation pathways to catalyze nutrients that are abundant in the tomato apoplast. *Mol. Plant Microbe. Interact.* **21**, 269–282. doi: 10.1099/mpmi.2008-2-0289

Sakata, N., Ishiga, T., Masuo, S., Hashimoto, Y., and Ishiga, Y. (2021). *Coronatine* contributes to *Pseudomonas cannabina* pv. *alissalensis* virulence by overcoming both stomatal and apoplastic defenses in dicot and monocot plants. *Mol. Plant Microbe Interact.* doi: 10.1099/mpmi.0.0261-R [Epub ahead of print].

Sakata, N., Ishiga, T., Saito, H., Nguyen, V. T., and Ishiga, Y. (2019). Transposon mutagenesis reveals *Pseudomonas cannabina* pv. *alissalensis* optimizes its virulence factors for pathogenicity on different hosts. *PeerJ*. 7:e7698. doi: 10.7717/peerj.7698

Sambrook, J., Fritsch, E. F., and Maniatis, T. (1989). *Molecular Cloning: A Laboratory Manual*, 2nd Edn. Cold spring harbor, NY: Cold spring harbor laboratory press.

Sawada, T., Eguchi, M., Asaki, S., Kashiwagi, M., Takahashi, F., et al. (2018). MexEF-OprN multidrug efflux pump transporter negatively links to virulence factors for pathogenicity on different hosts. *PeerJ*. 7:e7698.

Sakata, N., Ishiga, T., Masuo, S., Hashimoto, Y., and Ishiga, Y. (2021). *Coronatine* contributes to *Pseudomonas cannabina* pv. *alissalensis* virulence by overcoming both stomatal and apoplastic defenses in dicot and monocot plants. *Mol. Plant Microbe Interact.* doi: 10.1099/mpmi.0.0261-R [Epub ahead of print].

Schoen, C., Kischkies, L., Elias, J., and Ampattu, B. J. (2014). Metabolism and virulence in *Neisseria meningitidis*. *Front. Cell. Infect. Microbiol.* **4**:114. doi: 10.3389/fcimb.2014.00114

Schreiber, K. J., Ye, D., Fisch, E., Jian, A., Lo, T., and Desveaux, D. (2012). A high-throughput forward genetic screen identifies genes required for virulence of *Pseudomonas syringae* pv. *maculicola* ES4326 on *Arabidopsis*. *PLoS One* **7**(e41461). doi: 10.1371/journal.pone.0041461

Takikawa, Y., and Takahashi, F. (2014). Bacterial leaf spot and blight of crucifer species caused by *Pseudomonas syringae* pv. *maculicola* and...
Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Copyright © 2021 Sakata, Ishiga and Ishiga. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.