Pantoea ananatis Genetic Diversity Analysis Reveals Limited Genomic Diversity as Well as Accessory Genes Correlated with Onion Pathogenicity

Shaun P. Stice1†, Spencer D. Stumpf2†, Ron D. Gitaitis3, Brian H. Kvitko1,3* and Bhabesh Dutta2‡

1 Department of Plant Pathology, University of Georgia, Athens, GA, United States, 2 Department of Plant Pathology, University of Georgia, Tifton, GA, United States, 3 The Plant Center, University of Georgia, Athens, GA, United States

Pantoea ananatis is a member of the family Enterobacteriaceae and an enigmatic plant pathogen with a broad host range. Although P. ananatis strains can be aggressive on onion causing foliar necrosis and onion center rot, previous genomic analysis has shown that P. ananatis lacks the primary virulence secretion systems associated with other plant pathogens. We assessed a collection of fifty P. ananatis strains collected from Georgia over three decades to determine genetic factors that correlated with onion pathogenic potential. Previous genetic analysis studies have compared strains isolated from different hosts with varying diseases potential and isolation sources. Strains varied greatly in their pathogenic potential and aggressiveness on different cultivated Allium species like onion, leek, shallot, and chive. Using multi-locus sequence analysis (MLSA) and repetitive extragenic palindrome repeat (rep)-PCR techniques, we did not observe any correlation between onion pathogenic potential and genetic diversity among strains. Whole genome sequencing and pan-genomic analysis of a sub-set of 10 strains aided in the identification of a novel series of genetic regions, likely plasmid borne, and correlating with onion pathogenicity observed on single contigs of the genetic assemblies. We named these loci Onion Virulence Regions (OVR) A-D. The OVR loci contain genes involved in redox regulation as well as pectate lyase and rhamnogalacturonase genes. Previous studies have not identified distinct genetic loci or plasmids correlating with onion foliar pathogenicity or pathogenicity on a single host pathosystem. The lack of focus on a single host system for this phytopathogenic disease necessitates the pan-genomic analysis performed in this study.

Keywords: Pantoea ananatis, pangeneome, onion, CWDE, MLSA, bacterial secretion systems

INTRODUCTION

Pantoea ananatis is a member of the Enterobacteriaceae family and is a ubiquitous phytopathogenic bacterium capable of infecting diverse monocotyledonous hosts including maize, rice and pineapple as well as the woody host, Eucalyptus (Serrano, 1928; Tabei et al., 1988; Paccola-Meirelles et al., 2001; Coutinho et al., 2002; Coutinho and Venter, 2009). P. ananatis strains have also been
recovered from opportunistic infections in humans (De Baere et al., 2004). In the United States, \textit{P. ananatis} is of primary concern as a causative agent of center rot of onion (\textit{Allium cepa} L.). The first outbreak of center rot of onion in the United States occurred in Georgia in 1997 and since then it has been identified in different onion growing regions (Gitaitis and Gay, 1997; Schwartz and Otto, 2000; Carr et al., 2010). \textit{P. ananatis} outbreaks are a recurring challenge for onion production and outbreaks resulting in considerable crop loss have been documented (Walcott et al., 2002). \textit{P. ananatis} infection can be transmitted via seed but it is also a common epiphyte on weeds and is known to be transmitted transiently between plants by two different species of thrips (Gitaitis et al., 2002; Walcott et al., 2002; Dutta et al., 2014). Infection of onions is facilitated through wounds or thrips damage. \textit{P. ananatis} strains are known to express the ice-nucleation active (InaZ) cell surface protein which efficiently nucleates ice crystals and is associated with increased incidence of frost damage (Lindow, 1983; Abe et al., 1989). Upon leaf invasion, \textit{P. ananatis} reaches high levels in leaf blades and causes foliar necrosis. The pathogen can also progress from leaf blade to the corresponding onion bulb scale where, alongside colonization by secondary pathogens, it causes center rot symptoms in central scales (Carr et al., 2013).

Management and accurate diagnosis of \textit{P. ananatis} has been complicated due to the phenotypic variation among different strains in onion pathogenicity. \textit{P. ananatis} strains collected from non-onion sources (thrips, weeds) may either be highly aggressive on onions or completely onion-non-pathogenic. Currently no strain features have been described that can predict onion pathogenic potential.

\textit{Pantoea ananatis} is an enigma among bacterial phytopathogens. Whole genome sequencing has shown that phytopathogenic \textit{P. ananatis} strains lack Type II, Type III, and Type IV virulence-associated secretion systems as well as known phytotoxin synthesis genes (De Maayer et al., 2014; Sheibani-Tezerji et al., 2015; Weller-Stuart et al., 2017). In the absence of these pathogenicity factors, secretion systems used to deliver Cell Wall Degrading Enzymes (CWDEs) outside of the bacterium (i.e., T2SS), or virulence effector proteins directly into the plant cells (i.e., T3SS/T4SS); however, it remains an open question as to what host interaction mediated pathogenicity factors are associated with \textit{P. ananatis} infection of onion. Progressing leaf necrosis has been shown to be dependent on pathogen quorum sensing and swimming motility (Morohoshi et al., 2007). In addition, the Type VI Secretion System (T6SS), most commonly associated with inter-bacterial competition, was shown to play a role in foliar necrosis symptoms but with little effect on pathogen populations (Morohoshi et al., 2007; Shyntum et al., 2015; Weller-Stuart et al., 2017).

In this study we assessed a collection of fifty Georgia \textit{P. ananatis} strains collected over three decades (1997–2015) from diverse sources to determine what genetic factors correlated with onion pathogenic potential. Using multi-locus sequence analysis (MLSA) and repetitive extragenic palindrome (rep)-PCR techniques, we did not observe strong correlations between onion pathogenic potential and genetic diversity among strains. We found that strains varied greatly in their pathogenic potential and aggressiveness on different cultivated \textit{Allium} species like onion, shallot (\textit{Allium cepa} var. \textit{aggregatum}), chives (\textit{Allium schoenoprasum}), and leeks (\textit{Allium porrum}). Several strains that were non-pathogenic on onion displayed significant virulence on leek. Pan-genome analysis derived from whole genome sequencing of a sub-set of six onion pathogenic and four onion non-pathogenic Georgia strains identified four loci associated only with the six onion pathogenic strains. These four loci were present on a single contig. We named these loci as the Onion Virulence Regions A through D (OVR-A-D). The OVR loci contain pectate lyase and rhamnogalacturonase genes as well as genes that are involved in redox regulation.

**MATERIALS AND METHODS**

**Bacterial Strains, Identification, and Culture Preparation**

Fifty \textit{P. ananatis} strains used in this study were isolated from onion and other sources including weeds and thrips throughout the state of Georgia (Table 1). These strains were stored in sterile 15% aqueous glycerol solution at $-80\textdegree$C. The strains which were isolated from either symptomatic onion foliage, bulb, or from contaminated onion seeds were designated as “PNA” in our culture collection. Strains isolated from asymptomatic weeds or from the gut of thrips (\textit{Thrips tabaci} and \textit{Frankliniella fusca}) were designated as “PANS.” The source, year of isolation, and county of origin in Georgia for these strains are shown in Table 1. These strains were identified as \textit{P. ananatis} by their morphological and physiological characteristics such as Gram-negative, facultative anaerobic, positive for indole production, negative for nitrate reductase and phenylalanine deaminase and by PCR amplification of a 398 base pair (bp) fragment using \textit{P. ananatis} species-specific primers (Gitaitis et al., 2002).

Inoculum was prepared by transferring single colonies of each bacterial strain from 48-h old cultures on nutrient agar (NA) medium to nutrient broth or lysogeny broth (LB). The broth was shaken overnight on a rotary shaker (Inova; New Brunswick Scientific, Edison, NJ) at 150 rpm. After a 12-h of incubation, 10 ml of each bacterial suspension were centrifuged at 6,000 $\times$ g (Allegra 25R, Beckman Coulter, Fullerton, CA) for 5 min. The supernatant was discarded and the pellet was re-suspended in 0.1 M phosphate-buffered saline (PBS). Later, inoculum was adjusted using a spectrophotometer (Spectronic 20; Bausch and Lomb, Rochester, NY) to an optical density of 0.3 at 600 nm [$\approx 1 \times 10^{8}$ colony forming unit (CFU)/ml]. The bacterial suspension was serially diluted in PBS to obtain the desired concentration according to each experiment. PNA 97-1R is a spontaneous rifampicin (Rf) resistant clone of PNA 97-1 recovered after plating a dense suspension of PNA 97-1 onto an LB Rf 60 plate.

**Multi-locus Sequence Analysis (MLSA) of \textit{P. ananatis} Strains**

Total microbial genomic DNA was extracted from a 3 ml volume of overnight cultures grown at 28$\textdegree$C in nutrient broth using an UltraClean Microbial DNA Kit (MO BIO, Carlsbad, CA)
### TABLE 1 | Isolates of *Pantoea ananatis* and their associated phenotypic characteristics.

| Strain Name | Host | Place of isolation in Georgia (county) | Ice nucleation | Copper tolerance | Strain aggressiveness |
|-------------|------|----------------------------------------|----------------|------------------|---------------------|
| PANS 99-1   | *Richardia scabra* L. | Tift | − | − | ++ + |
| PANS 99-3   | *Richardia scabra* L. | Tift | − | − | ++ + |
| PANS 99-11  | *Digitaria sanguinalis* | Tift | + | − | ++ + |
| PANS 99-27  | *Desmodium tortuosum* | Toombs | + | − | ++ + |
| PANS 99-29  | *Digitaria sanguinalis* | Tift | + | − | ++ + |
| **PANS 01-2** | *Thrips tabaci* | Tift | + | − | ++ + |
| **PNA 97-1** | *Allium cepa* L. | Toombs | + | − | ++ + |
| PNA 98-3    | *Allium cepa* L. | Toombs | + | − | ++ + |
| PNA 99-3    | *Allium cepa* L. | Tift | + | − | ++ + |
| PNA 200-11  | *Allium cepa* L. seed | Tift | + | − | ++ + |
| PNA 02-18   | *Allium cepa* L. | Tattnall | + | − | ++ + |
| **PNA 06-1** | *Allium cepa* L. | Wayne | + | − | ++ + |
| **PNA 07-1** | *Allium cepa* L. | Tattnall | + | − | ++ + |
| **PNA 08-1** | *Allium cepa* L. | Tattnall | + | − | ++ + |
| **PNA 15-1** | *Allium cepa* L. | Tattnall | + | − | ++ + |
| PANS 99-22  | *Digitaria sanguinalis* | Tift | − | − | ++ + |
| **PANS 01-5** | *Thrips tabaci* | Tift | + | − | ++ + |
| **PANS 01-6** | *Thrips tabaci* | Tift | + | − | ++ + |
| PNA 07-5    | *Allium cepa* L. | Wayne | − | − | ++ + |
| PNA 13-1    | *Allium cepa* L. | Toombs | − | − | ++ + |
| PNA 14-1    | *Allium cepa* L. | Toombs | − | − | ++ + |
| **PNA 99-14** | *Allium cepa* L. | Tattnall | − | − | ++ + |
| PNA 14-4    | *Allium cepa* L. | Toombs | − | − | ++ + |
| **PNA 99-24** | *Vigna unguiculata* | Toombs | − | − | ++ + |
| **PNA 200-3** | *Allium cepa* L. seed | Tift | − | − | ++ + |
| PNAS 99-33  | *Richardia scabra* L. | Coffee | − | − | Y |
| PNAS 99-33  | *Richardia scabra* L. | Coffee | − | − | Y |
| **PANS 02-5** | *Frankliniella fusca* | Tift | − | − | Y |
| **PNA 97-11** | *Allium cepa* L. | Toombs | − | − | Y |
| **PNA 98-2** | *Allium cepa* L. | Tift | + | − | Y |
| **PNA 98-11** | *Allium cepa* L. | Tattnall | − | − | Y |
| **PNA 99-2** | *Allium cepa* L. | Tattnall | + | − | Y |
| **PNA 99-8** | *Allium cepa* L. | Wheeler | + | − | Y |
| **PANS 99-23** | *Cyperus esculentus* | Toombs | + | − | Y |
| **PANS 99-26** | *Chamaesyce hyssopifolia* | Toombs | + | − | Y |
| **PANS 99-6** | *Glandularia bipinnatifida* | Tift | − | − | Y |
| **PANS 200-1** | *Amaranthus viridis* | Toombs | − | − | Y |
| **PANS 99-7** | *Allium cepa* L. | Tattnall | − | − | N |
| **PANS 99-5** | *Glandularia bipinnatifida* | Tift | − | − | N |
| **PANS 99-32** | *Richardia scabra* L. | Toombs | + | − | N |
| **PANS 99-36** | *Richardia scabra* L. | Terrell | + | − | N |
| **PANS 02-1** | *Frankliniella fusca* | Tift | − | − | N |
| **PANS 04-2** | *Frankliniella fusca* | Tift | − | − | N |
| **PANS 99-9** | *Allium cepa* L. | Tattnall | + | − | N |
| **PANS 200-7** | *Allium cepa* L. seed | Tift | − | − | N |
| **PANS 200-8** | *Allium cepa* L. seed | Tift | − | − | N |
| **PANS 200-12** | *Allium cepa* L. seed | Tift | + | − | N |
| **PANS 05-1** | *Allium cepa* L. | Tattnall | + | − | N |

(Continued)
TABLE 1 | Continued

| Strain Name<sup>a</sup> | Host | Place of isolation in Georgia (county) | Ice nucleation<sup>b</sup> | Copper tolerance<sup>c</sup> | Strain aggressiveness<sup>d</sup> |
|-------------------------|------|--------------------------------------|--------------------------|--------------------------|--------------------------|
| PNA 07-10               | Allium cepa L. | Toombs | − | − | *N* ND ND ND |
| PNA 11-1                | Allium cepa L. | Toombs | − | − | *N* ND ND ND |

<sup>a</sup> First digits of the strain designation indicate year of isolation, e.g., 97 = 1997, 200 = 2000.

<sup>b</sup> Ice-nucleation test at −5°C.

<sup>c</sup> Strain aggressiveness on the foliage of onion, chive, leek, and shallot 5 dpi. Strains lacking pathogenicity are designated by a “−,” weakly aggressive strains on a host are represented by a “+,” moderately aggressive “++,” and highly aggressive “+++” Y/N denote previously recorded as pathogenic or non-pathogenic on onion.

<sup>d</sup> Bold names indicate whole genome sequencing (WGS) strains.

According to the manufacturer's instructions. Extracted genomic DNA was quantified on a NanoDrop 1000 spectrophotometer (Thermo Scientific, Wilmington, U.S.A.) and 50 ng of genomic DNA was used for each PCR reaction. Six primer pairs of housekeeping genes, *fusA*, *gyrB*, *leuS*, *pyrG*, *rpl*, and *rpoB*, were chosen based on a previous MLSA scheme (Salerno et al., 2007; Deletoile et al., 2009). The concatenation of these six genes represented a total of 2,961 bp. Primer sequences and PCR cycles are outlined in Supplementary Table 3. Each PCR reaction was performed in a 50 µl reaction consisted of 10 µl of 5x PCR buffer, 0.2 mM of dNTPs, 0.2 M of each primer, 1.25 U HotStart Taq Polymerase (Qiagen, Valencia, Ca) and 50 ng/µl of template DNA. PCR products were subjected to gel electrophoresis in Tris-acetate-EDTA buffer on a 1.5% agarose gel at 100 V for 45 min. Reactions resulting in positive amplifications and expected fragment size were purified with a DNA Clean and Concentrator Kit (Zymo Research Inc., Irvine, Ca), according to the manufacturer's instructions. Fragment sequencing was conducted by Eurofins, (Louisville, KY).

**Phylogenetic Analysis of Sequencing Data**

Sequences were manually trimmed and verified based on peak quality of forward and reverse chromatograms in Geneious 8.1.8 (Biomatters, Auckland, New Zealand) and were aligned using MUSCLE (http://www.ebi.ac.uk/Tools/maa/muscle/). The nucleotide sites presenting alignment gaps were excluded from analysis and were then concatenated. Consensus sequences were used to generate a maximum-likelihood (ML) tree based on the Tamura-Nei Model of MEGA 6.06 with bootstrapping repetitions of 1,000. Sequences from *P. agglomerans* 97-1 (accession no. FN434113.1) from GenBank were used as outgroups. MLST sequences generated from this study were deposited in GenBank under accession numbers MF925240–MF925289, MF939901–MF939950, MF939951–MF940000, MF940001–MF940050, MF940051–MF940100, and MF964629–MF964678.

**Repetitive Extragenic Palindrome (rep)-PCR Amplification and Analysis**

Genomic DNA from strains were isolated as described above and subjected to rep-PCR using REP1R-I (III ICG ICG ICA TCI GGC) and REP 2-I (ICG ICT TAT CIG GCC TAC) primers (Versalovic et al., 1991). The amplification was carried out in a total of 25 µl reactions containing 50 ng of DNA, 10 mM Tris-HCl, 50 mM KCl, 1.5 mM MgCl₂, 200 µM dNTPs, 2.5 U pureTaq DNA polymerase, and 0.5 µM of each primer pair. Amplifications were done using the following conditions: an initial denaturation at 95°C for 4 min, 45 cycles consisting of 94°C for 1 min, 45°C for 1 min, and 65°C for 8 min, followed by cooling for 4 min. Samples were separated by gel electrophoresis in Tris-acetate-EDTA buffer on a 1.5% agarose gel for 2 h at 100 V stained with GelGreen. DNA fingerprinting profile for each *P. ananatis* strain was compared using Dice’s (1945) coefficient of analysis with the aid of BioNumerics software package (Applied Math, Kortrijk, Belgium), and the unweighted pairwise group method with arithmetic mean algorithm was used to generate a dendrogram indicating strain relatedness.

**Aggressiveness of *P. ananatis* Strains on Onion, Chive, Leek, and Shallot**

The aggressiveness of 33 selected strains were tested on 6-week old shallots (cv. "Camelot F1"), chives (cv. "Dolores"), leeks (cv. "King Richard"), and onion (cv. "Century") under greenhouse conditions. Strains from 11 clades (*n* = 23) and from unresolved portion (*n* = 10) of the MLSA phylogenetic tree were selected for this assay. These strains represented 66% of the total *P. ananatis* strains from 11 clades and unresolved portion. Seedlings of each plant species were established in 10 × 8 cm (diameter × height) plastic pots (Hummert International, Earth City, MO) containing commercial potting mix. The seedlings were kept in greenhouse conditions and maintained at 25–28°C and 80–90% RH with a 12L:12D photoperiod. Seedlings were fertilized with osmocote smart-release plant food (14-14-14) (Scotts Miracle-Gro Company, Marysville, OH). Inoculum preparation and standardization were conducted as described above. Seedlings were inoculated by cutting the central leaf 1 cm from the apex with a sterile pair of scissors. Using a micropipette, a 10 µl drop of bacterial suspensions containing 1 × 10⁸ CFU/mL (≈1 × 10⁹ CFU) were placed at the cut end of the seedlings. Seedlings mechanically inoculated with PBS as described above were used as negative controls. Four replications per strain per host were used in a single experiment and this experiment was repeated one more time. The experiment was arranged in a randomized complete block design. Seedlings were checked...
visually for symptom development every day until 5 days after inoculation (DAI). At 5 DAI, lesion length for each seedling was measured and mean lesion length for each strain on each plant species were compared using Fisher’s protected least significant difference test at \( P < 0.05 \) level.

To confirm that symptoms observed were caused by \( P. ananatis \), bacteria from symptomatic tissue (\( n = 5 \) symptomatc plants per host) were isolated from a region adjoining the necrotic and healthy tissue and streaked onto Tryptic soy broth agar (TSBA) plates and incubated for 48 h at 28°C. After 48 h of incubation, yellow-pigmented colonies were isolated and tested for \( P. ananatis \) using physiological tests (as previously described) and species-specific TaqMan based polymerase-chain reaction (PCR) assays (Walcott et al., 2002). Briefly, presumptive \( P. ananatis \) colonies were picked using a sterile inoculation loop and suspended separately in a 2 ml micro-centrifuge tubes containing 25 \( \mu l \) of sterile deionized water. The bacterial suspension was heated (Modular Dry Block Heaters, Cole Parmer, IL) for 3 min at 95°C. A suspension (5 \( \mu l \)) was amplified in 20 \( \mu l \) of PCR mastermix containing 10 mM Tris-HCl (pH 9.0), 50 mM KCl, 0.1% Triton X-100, 1.5 mM MgCl\(_2\), and 0.2 mM of each nucleotide (dATP, dCTP, dGTP, and dTTP), 25 \( \mu M \) each of primer PanITS1 (5′-GTCTGATAGAAAAGATACG-3′) and EC5 (5′-CGGTTGATGCCCTGGGA-3′) and 10 \( \mu M \) of TaqMan probe 6-FAM TAGCGGTAGACTCCGCCCTTCA-BHQ. The PCR reaction was conducted in a Cepheid Smart Cycler (Sunnyvale, CA) using the following thermal profile: denaturation at 95°C for 180 s, 35 cycles each of denaturation at 95°C for 15 s, and annealing at 60°C for 40 s. Samples with cycle threshold (Ct) values <35 were considered positive for \( P. ananatis \).

### Ice Nucleation Test

To test the ice nucleating ability of \( P. ananatis \) strains, five separate drops of 10 \( \mu l \) aliquots of bacterial suspension (>1 \( \times \) \( 10^6 \) CFU/ml) were pipetted onto a sterile weigh boat in an ice-ethanol water bath adjusted to \(-5^\circ\)C. Sterile water was used as the negative control. Samples that froze concurrently with the positive control were considered ice nucleation positive and samples that remained in liquid form or with delayed ice-nucleation activity were considered ice nucleation negative. Fifty \( P. ananatis \) strains were used and the experiment was repeated twice.

### Copper Tolerance

Strains were grown and re-suspended to an optical density of 0.3 at 600 nm, as previously described, and suspensions were serially diluted to a concentration of \( 10^5 \) CFU/ml. Aliquots of 50 \( \mu l \) were spread onto nutrient agar (NA) and NA amended with 200 ppm CuSO\(_4\) \( \cdot \) 5H\(_2\)O. Plates were incubated at 28°C for 48–72 h and observed for bacterial growth.

### High Quality Draft Genome of PNA 97-1R

Genomic DNA from the prototype strain PNA 97-1R was obtained from overnight cultures started from single colonies.
using the Gentra Puregene Yeast/Bact kit (Qiagen Inc.), quantified using biospectrometer (Eppendorf), and visually assessed for quality on a 0.75% agarose gel. DNA quality, sequencing and assembly were carried out at the University of Delaware DNA Sequencing & Genotyping Center (Newark, DE). DNA was assessed for quality metrics via Fragment Analyzer (Advanced Analytical Technologies). Sequence reads were generated by a Pacific Biosciences RSII Single-Molecule DNA Sequencer following SMRT Bell library preparation. Basic de-novo assembly was carried out using PacBio HGAP assembler (v3) with seed reads 17 kilo base (kb) and above. Contigs were polished using Quiver and annotation was performed via Rapid Annotation using Subsystem Technology (RAST v2.0) or Prokaryote Genome Annotation Pipeline (PGAP v4.2.) (Zhao et al., 2012; Overbeek et al., 2014).

**Draft Genomes of 10 Diverse Strains**

Genomic DNA from 10 diverse strains (PNA 97-1R, PNA 06-01, PNA 200-3, PNA 15-1, PNA 99-7, PANS 04-02, PANS 01-02, PANS 99-23, PANS 99-36, and PANS 99-3) was obtained from overnight cultures started from single colonies using the Gentra Puregene Yeast/Bact Kit (Qiagen Inc.), quantified using biospectrometer (Eppendorf), and visually assessed as above on a 0.75% agarose gel. Sequencing, assembly, annotation, and variant calling were carried out at MicrobesNG (Birmingham, U.K.). Sequence reads were generated by an Illumina HiSeq using 250-bp paired-end libraries, and assembled using SPAdes (version 3.10.1). Reads were mapped to the reference genome PNA 97-1R and back to contigs using BWAMEM to ensure quality and 30X coverage. Annotation was performed via Prokka or RAST (Overbeek et al., 2014; Seemann, 2014).

**Pan-Genomic Analysis**

The Prokka annotated fragmented de novo assemblies for the ten draft genomes for strains (PNA 97-1R, PNA 06-01, PNA 200-3, PNA 15-1, PNA 99-7, PANS 04-02, PANS 01-02, PANS99-23, PANS 99-36, and PANS 99-3) were loaded in GFF3 format to the ROARY (v3.6.1) program using default parameters to calculate the pan-genome of the 10 strains (Page et al., 2015). ROARY was run on a windows computer through an emulated Linux machine as described by the authors (https://sanger-pathogens.github.io/Roary/). The resulting newick tree files were edited using Geneious (v10.1.3). The “gene_presence_absence.csv” output file was analyzed using Microsoft excel to identify gene that are present in the pathogenic strains and absent in the non-pathogenic strains (Supplementary File 1).

The average nucleotide identity of the strains (ANI) was calculated using the online ANI calculator server with default parameters to compare WGS strains to the genomic type strain LMG 20103 (http://envomics.ce.gatech.edu/ani/). Additional Pan-genomic analysis was conducted using RAST BLASTP pairwise alignment with PNA-97-1 listed as reference genome. The computationally intensive Pan Genomic Analysis Pipeline (PGAP) was run on the UGA Sapelo computer cluster (Zhao et al., 2012). The resulting “5.Orthologs_Cluster_Function” file was opened with Microsoft excel and genes clustering in pathogenic strains and non-pathogenic strains were cross referenced with the results from ROARY analysis. The resulting putative virulence genes were then identified in the contigs of the draft genomes using Geneious. The contigs containing putative virulence associated genes were extracted and aligned using the Geneious Mauve plugin (v2.3.1) (Darling et al., 2004). Putative secretion target signals were assessed by running genes through the Signal P (v4.1) Server and TMHMM Server (v2.0) for prediction of transmembrane helices in proteins (Darling et al., 2004; Nielsen, 2017). Genomes were also investigated for secondary metabolite non-ribosomal peptide synthetase (NRPS) biosynthesis gene clusters using the antiSMASH bacterial server. Whole genome MLST (wgMLST) was conducted using PGAdb-builder (Liu et al., 2016). The 95% occurrence allele scheme was selected to generate a wgMLST tree with P. agglomerans PC10 and P. stewartii subsp. indologens LMG 2632 selected as outgroups (Accession NZ_LIME0000000.1 and JPKO0000000.1). Genomic islands were identified using the IslandPath DIMOB function on IslandViewer4 server (Bertelli et al., 2017). Partial and complete phage regions were identified in the genome assemblies of the WGS strains using Phaster. Circular genome comparisons of WGS query strains and the OVR A-D locci to the reference strain PNA 97-1R were conducted using BRIG (GPLx3) (Alikhan et al., 2011).

**Nucleotide Sequence Accession Numbers**

The Whole Genome Shotgun project of *P. ananatis* strains (PNA 97-1R, PNA 06-1, PNA 200-3, PNA 15-1, PNA 99-7, PANS 04-2, PANS 01-2, PANS 99-23, PANS 99-36, and PANS 99-3) have been deposited at DDBJ/ENA/Genbank under the accessions (CP020943-CP020945), NMZY00000000, NMZX00000000, NMZZ00000000, NMZW00000000, NMZU00000000, NMZX00000000, NMZI00000000, and NMZR00000000, respectively. In addition, all bio-projects and accession numbers related to this sequencing project are linked under the NCBI umbrella project PRJNA 400632.

**RESULTS**

**Multi-locus Sequence Analysis**

Six housekeeping genes (*fusA, gyrB, leuS, pyrG, rpl*, and *rpoB*) from fifty *P. ananatis* strains were sequenced and a concatenated tree was developed. Sequence identity amongst the strains was >99% indicating low diversity. Overall, 11 small clades consisting of *P. ananatis* strains (*n = 35*) with bootstrap values >67 were observed (Figure 1). Among 11 clades, three clades contained strains, which were strictly isolated from onion or onion seed (designated PNA) (Figure 1). In addition, there were eight clades that contained strains isolated from diverse sources including onion or onion seed or weeds or thrips. None of the strains isolated from sources other than onion...
or onion seed (designated PANS) were exclusively resolved in a separate clade. Fifteen strains remain unresolved (Figure 1). There was no trend among strains with respect to their year of isolation, geographic location, source of isolation or ice-nucleation phenotype.

**Repetitive Extragenic Palindrome (rep)-PCR**

A dendrogram was constructed from the rep-PCR profiles of fifty *P. ananatis* strains. Strains that showed relatedness more than 60% were clustered into four separate clades containing PNA and PANS strains. Fingerprint patterns were not correlated with isolation source or phenotype. Clade I strains (*n* = 24) were 80% similar while strains in Clade II (*n* = 11) had similarity of 86% (Supplementary Figure 1). Five strains displayed identical fingerprint profile within Clade III and were unique from the strains in the clades I–IV (Supplementary Figure 1). Clade IV comprised of ten strains that showed fingerprint profiles different from other clades and displayed 61% genetic similarity (Supplementary Figure 1).

**Pathogenicity and Aggressiveness of *P. ananatis* Strains on Onion, Chive, Leek, and Shallot**

Thirty-three strains pathogenic or non-pathogenic on onion based on MLSA findings were selected for pathogenicity and aggressiveness study on onion, chive, leek, and shallot under controlled greenhouse conditions. PBS control plants displayed no foliar symptoms in any test. On onion, 100% of the previously determined pathogenic strains displayed typical foliar necrosis symptoms. However, there were significant differences among strains with respect to the aggressiveness (*P* < 0.0001; Table 1). Fifteen strains representing a mix of PNA and PANS strains were highly aggressive on onion with lesion length > 7.8 cm after 5-days of inoculation under greenhouse conditions. Moderate levels of aggressiveness were displayed by ten strains also representing a mix of PNA and PANS strains. All eight previously determined onion non-pathogenic PANS strains were again determined to be non-pathogenic on onions.

On chive, all strains except PNA 200-11 were non-pathogenic. The PNA 200-11 strain was weakly aggressive on chive with lesion length of 0.92 cm after 5-days of inoculation. Out of

---

**FIGURE 1 |** Maximum-likelihood tree based on the Tamura-Nei Model of *Pantoea ananatis* strains derived from concatenation of six housekeeping genes (*fusA*, *gyrB*, *leuS*, *pyrG*, *rplB*, *rpoB*). Bootstrap values are based on 1,000 repetitions and values >60% are shown in the figure.
25 onion pathogenic strains, 19 strains were also pathogenic on shallot and they varied significantly in aggressiveness ($P = 0.0007$). These strains were weakly to moderately aggressive and they produced lesions of $>0.6$ cm after 5 days of inoculation. Six strains, which were pathogenic on onion, were non-pathogenic on shallot; however, one onion nonpathogenic strain, PANS 200-1 was found to be mildly aggressive on shallot. On leek, of the 25 onion pathogenic strains, 12 strains were also pathogenic and again, they varied significantly in aggressiveness ($P < 0.0001$). These strains were either weakly or moderately aggressive on leek with lesion length $>0.8$ cm after 5 days of inoculation. Thirteen onion pathogenic strains were non-pathogenic on leek. However, four onion non-pathogenic strains were weakly to moderately aggressive on leek. All symptomatic samples ($n = 5$ per host type) assayed were determined as positive for $P.\ ananatis$ by a real-time PCR assay.

**Ice Nucleation and Copper Tolerance**

Ice nucleating ability was observed in 66% (33/50) of the strains tested. Of the 33 strains that were positive for ice nucleation, 18 were isolated from onion, three from onion seed, and 12 from other hosts. All 50 $P.\ ananatis$ strains were unable to grow on NA media amended with 200 ppm CuSO$_4$$\cdot$$5$H$_2$O. Confluent growth was observed on all non-amended NA plates (Table 1).

**Whole Genome Sequencing Summary**

The prototype strain PNA 97-1R was sequenced and assembled to near completion via PacBio sequencing. In addition, whole genome Illumina shotgun sequences (WGS) of PNA 97-1R and nine additional $P.\ ananatis$ strains from diverse primary and secondary sources associated with the center rot pathosystem were assembled and annotated to generate draft genomes for comparative pan-genomic analysis. The genomes of $P.\ ananatis$ strain ranged from 4.8 to 5.1 Mbp size which is consistent with the reported sizes $P.\ ananatis$ isolated from other locations (De Maayer et al., 2014). Genome features are reported in Table 3. The pathogenicity of each strain was assessed using both foliar and bulb scale assays (Figure 2). Annotated genomes were assessed for reported virulence elements including: Acyl homoserine lactone quorum sensing system, flagellar elements, and twitching motility type IV pili.

Universally, $P.\ ananatis$ strains lacked complete virulence-type T3SS and T4SS secretion systems as well as the T2SS commonly used by soft rot causing Enterobacteriaceae pathogens to deliver CWDEs and other virulence factors across the outer membrane. A scan of the genomes to identify NRPS/To delivery CWDEs and other virulence factors across the commonly used by soft rot causing Enterobacteriacea pathogens were assessed for reported virulence elements including: Acyl homoserine lactone quorum sensing system, flagellar elements, and twitching motility type IV pili.

Red Onion Scale Assay

The ability of $P.\ ananatis$ strains to clear red onion scales, and cause scale weakening was observed in five of the six onion pathogenic WGS strain (Figure 2, Table 2). The weakly aggressive PNA 200-3 did not exhibit scale clearing, but did cause traditional foliar necrotic symptoms (Figure 2, Table 2). The negative control did not exhibit clearing in the onion scale assay or blighting in the foliar assay. Thus, all six strains containing the OVR-A-D region exhibited blight symptoms in foliar assay and five of six strains containing the OVR-A-D region exhibited pigment clearing in the onion scale assay (Table 2).

**Draft Genomes of 10 $P.\ ananatis$ Strains from Georgia**

Quality assessment of extracted PNA 97-1R prototype strain DNA yielded fragments $>50$ kb. PacBio reads were assembled resulting in a high quality draft genome of prototype strain PNA 97-1R. The assembly consists of three contigs with sizes 4,571,292 bp; 295,591 bp; 180,232 bp, GC content 53.6, 52.4, and 49.6%, respectively; with an estimated 200X genome coverage (GenBank accession numbers CP020943-CP020945 respectively). The second largest contig 295,591 bp falls in the range of the large universal Pantoea plasmid LPP-1 280–789 kb (De Maayer et al., 2012). The presence of core factors such as thiamine biosynthesis proteins (thiaOSF), and MAUVE alignment with the AJ335 LPP-1 pEA320 (NC_017533.1) aided in characterizing this contig as homologous to the widely reported universal plasmid LPP-1 (Supplementary Figure 2; Weller-Stuart et al., 2017). MAUVE alignment yielded a single block with homology between the proposed PNA 97-1R LPP-1 plasmid (CP020944) and the AJ335 LPP-1 plasmid (Supplementary Figure 2).

MicrobesNG Illumina draft genome sequencing yielded 354,367–2,036,787 bp range in reads for assemblies. The average insert size ranged from 344 to 649 bp (Supplementary Table 1). Average coverage for draft-genome sequences ranges from 28.9 to 167.46%. The GC content ranged from 53.29 to 53.52%. The RAST annotation resulted in prediction of 4,681–4,992 protein coding sequences (CDS) among WGS strains, roughly 1,615–1,330, were predicted as hypothetical coding sequences (Supplementary Table 2). The number of predicted phages, prophages, and transposable elements varied widely among strains from 2 to 53 CDS (Supplementary Table 2). Most proteins coding sequences are devoted to carbohydrates, amino acids and derivatives; 455–630 CDS on average. No traditional complete virulence associated plant pathogen secretion systems were identified. Two proteins associated with the type II secretion system including gspE (cytosolic ATPase) and gspF (inner membrane protein) were identified in WGS genomes of all strains.
### TABLE 3 | Primers and conditions used for amplification and sequencing.

| Gene | Primer name | Primer sequence | Primer positions on coding sequences | PCR cycles | Template size (bp) | Template position on P. ananatis PA13 genome |
|------|-------------|-----------------|--------------------------------------|-------------|-------------------|--------------------------------------------|
| fusA | fusA3       | 5′-CAT CGG TAT CAG TGC KCA CAT Cga-3′ | 36–59 | 2 min 94°C; 1 min 94°C; 1 min 58°C; 1 min 72°C (31 cycles); 5 min 72°C | 639 | 4449773–450416 |
| fusA | fusA4       | 5′-GAG CAT CGG CTG AAC RCC TTT GTT-3′ | | | | |
| gyrB | gyrB3       | 5′-GGG TAA GGG CCG GGG TAT GTA-3′ | 57–77 | 2 min 94°C; 1 min 94°C; 1 min 58°C; 1 min 72°C (31 cycles); 5 min 72°C | 427 | 4108–4543 |
|       | gyrB4       | 5′-CGG TGG ACG TGG GCA TGG TGG AT-3′ | 1488–1508 | | | |
|       | gyrB3i      | 5′-AAC GCG ATC GAC GAA GGC-3′ | 136–152 | Primers used only for sequencing | | |
|       | gyrB4i      | 5′-TGG AAC COR TGC TGC CAG-3′ | 771–788 | | | |
| leuS | leuS3       | 5′-GAG ACC GTG CTG GGC AAC GAC CAR GT-3′ | 487–512 | 2 min 94°C; 1 min 94°C; 1 min 58°C; 1 min 72°C (31 cycles); 5 min 72°C | 640 | 3282463–3283102 |
|       | leuS4       | 5′-GGG CGG GGC CCA RTA RGCG CTG-3′ | 1274–1293 | | | |
| pyrG | pyrG3       | 5′-GGG GTC GTA TCC TCT GTG GGT AAA GGC-3′ | 31–56 | 2 min 94°C; 1 min 94°C; 1 min 58°C; 1 min 72°C (31 cycles); 5 min 72°C | 316 | 1019655–1019970 |
|       | pyrG4       | 5′-GGA AGG GCA GGG ATT CTA TAC CNC GKA-3′ | 434–460 | | | |
| rplB | rplB3       | 5′-GAG ATT AAC GTC TTG AGT AGC ATC-3′ | 227–254 | 2 min 94°C; 1 min 94°C; 1 min 58°C; 1 min 72°C (31 cycles); 5 min 72°C | 343 | 457307–457649 |
|       | rplB4       | 5′-GAC GAC CAC CAT GYG GGT CRT C-3′ | 665–706 | | | |
| rpoB | Vic2        | 5′-GAG TGC TGG AAG TGG TAA CC-3′ | 2469–2489 | 4 min 94°C; 30 s 94°C; 30 s 50°C; 30 s 72°C (31 cycles); 5 min 72°C | 596 | 4432827–4433422 |

*Size of the sequence template of the internal portion of PCR product used for sequence comparison.

*Location of each gene sequence analysis on P. ananatis PA13 reference genome (NCBI genome accession no. CP003085).
**Known Virulence Factors**

Virulence factors previously recorded as having importance in the pathogenicity of *P. ananatis* including: acyl-homoserine lactone mediated quorum sensing system, twitching motility, flagellar motility, two clusters of type VI secretion system (C1, C3) associated genes and effectors were present in WGS of all 10 *P. ananatis* strains. Among these, PANS 99-3 contained all three previously described type VI clusters (C1, C2, C3) (De Maayer et al., 2011).

**The Ice Nucleation Gene (inaZ) Is Present in All Strains**

Although only five of the 10 WGS strains have an ice nucleation positive phenotype, the ice-nucleation protein gene inaZ was found to be present in all strains sequenced in this study. The inaZ gene was observed to vary in lengths from 3,474 bp (PANS 04-2) to 4,209 bp (PANS 06-1). PNA 200-3 has three ORFs with homology to inaZs, which are severely truncated being, 750, 450, and 1,287 bp. The severely truncated nature of the inaZ loci in PNA 200-3 likely explains the ice nucleation minus phenotype in this strain. PANS 99-3, PNA 15-1, PNA 99-7, and PANS 04-2 encoded the inaZ gene but lacked an ice nucleation phenotype *in vitro*.

**Pan-Genomic Comparative Analysis of *P. ananatis* Strains from Georgia**

The ROARY pan genomic analysis identified a core genome consisting of 3,750 protein coding sequences (CDS) and an accessory genome consisting of 1,395 CDS. Dendrograms generated through pan-genomic pipelines based on presence or absence of genes, differentiated onion pathogenic strains from onion non-pathogenic strains (Figure 3). ROARY BLASTP comparisons clearly identified CDS with homology to the prototype strain PNA 97-1R while also indicating phage, and virulence associated regions absent from onion non-pathogenic strains, but present in onion pathogenic strains (Figure 4, Supplementary Figure 6). Comparative analysis identified 57 CDS on four contiguous gene clusters on one contig in all pathogenic strains, but absent in onion non-pathogenic strains (Supplementary File 1). wgMLST analysis resulted in several different clades with the outgroups, *P. agglomerans* and *P. stewartii* subsp. *indologens* clustering from the *P. ananatis* strains (Supplementary Figure 4). Strains PANS 99-36 and PANS 99-23 clustered from other strains and had similar years of isolation. The pathogenic strain PANS 99-3 and non-pathogenic strain PNA 99-7 clustered together. Genomic islands were identified in all *P. ananatis* strains. Strains had between 9 and 15 predicted genomic
FIGURE 3 | Comparison of two dendrograms of *Pantoea ananatis* strains generated through different pan-genome analysis programs (A) ROARY presence-absence based tree generated uses the first 4,000 genes in accessory genomes and Fast Tree to calculate distances (Page et al., 2015) (B) PGAP pan-genomic based UPGMA dendrogram is calculated with phylip (Zhao et al., 2012). Dotted box highlights foliar pathogen isolates. See NCBI accession numbers (Table 2).

FIGURE 4 | RAST Seed Viewer protein sequence-based comparison analysis of *Pantoea ananatis* strains to pathogenic proto-type strain PNA 97-1R. BLASTP analysis compares every protein in the concatenation of the reference genome (PNA 97-1R PacBio) to every protein in the comparison genome (9 other strains, Illumina HiSeq). Overlapping regions and heat map indicate level of homology between proteins. (P1) Phage related proteins including: hypothetical proteins, phage lytic proteins, phage tail proteins, phage replication proteins, phage DNA binding proteins, phage lysine and lysozymes (P2) Phage associated genes (OVR) onion virulence region loci including: cell wall degrading enzymes, metabolic related proteins, amino acid transporters, and sugar transporters. (P3) Hypothetical proteins, phage proteins, transporters, and transposable DNA elements. (P4) Hypothetical proteins, and putative transcriptional regulators. This plot should not be used to determine the location of genes in the comparison genome but highlights regions that may require further analysis.

islands. The size of the islands varied from 5,430 to 150,125 bp (Supplementary Table 4). The described OVR A-D regions were predicted as genomic islands in pathogenic strains. Islands unique to non-pathogenic strains were not identified. Partial and complete phage regions were identified in all WGS strains. The relative orientation and location of 5...
phage regions can be observed in the type strain PNA 97-1R (Supplementary Figure 5).

**Identification of Genomic Regions Associated with Onion Pathogenic Strains**

The 57 CDS shared between onion foliar pathogen strains were present on four contiguous regions, and were localized on a single contig among pathogenic strains (Table 2). The contiguous regions contained 30, 16, 2, and 9 CDS and were ordered based on orientation in the prototype strain PNA 97-1R PacBio genome (CP020945). Of the 57 CDS, 17 CDS are predicted, hypothetical proteins based on NCBI PGAP annotations (Table 4). The onion virulence associated regions were termed onion virulence regions A-D (OVR-A through OVR-D) (Table 4, Figure 5). All four regions are located on the smallest 180,232 bp contig from PNA 97-1R PacBio assembly (CP020945). PGAP pan-genomic analysis, which utilizes computationally intensive strategy to BLAST every gene to each genome confirmed, 20 of the pathogen associated genes were present only once in the onion pathogenic strains and entirely absent in the non-pathogenic strains (Table 4, Figure 5). MAUVE alignment of pathogenic strains along these regions indicated high levels of homology (Figure 5).
### TABLE 4 | ROARY presence-absence based comparison.

| Gene                          | Prokka annotation | NCBI locus | NCBI PGAP annotation | Length (bp) |
|-------------------------------|-------------------|------------|----------------------|-------------|
| **ONION VIRULENCE REGION A (OVR-A)** |                   |            |                      |             |
| xerC                          | Tyrosine recombinase XerC | B9Q16_23180 | Recombinase Cre      | 1,038       |
| Helix-turn-helix              |                   | B9Q16_23175 | Hypothetical protein CDS | 213         |
| fabG*                         | 3-oxoacyl-[acyl-carrier-protein] reductase FabG | B9Q16_23170 | Sugar dehydrogenase CDS | 774         |
| DSBA-like thioredoxin domain  |                   | B9Q16_23165 | Isomerase CDS        | 702         |
| nemA                          | N-ethylmaleimide reductase | B9Q16_23160 | Alkene reductase CDS | 1,116       |
| Thioredoxin-like protein srl0233 |                  | B9Q16_23155 | Thiol reductase thioredoxin CDS | 354         |
| Carboxymuconolactone decarboxylase family |           | B9Q16_23150 | Alkylhydroperoxidase CDS | 339         |
| nemR                          | HTH-type transcriptional repressor NemR | B9Q16_23145 | TetR family transcriptional regulator | 579         |
| Hypothetical protein          |                   | B9Q16_23140 | Hypothetical protein CDS | 528         |
| yjE                          | Uncharacterized inner membrane transporter yjE | B9Q16_23135 | EamA family transporter CDS | 864         |
| patB                          | Cystathionine beta-lyase PatB | B9Q16_23130 | Aminotransferase CDS | 1,212       |
| gor                           | Glutathione reductase | B9Q16_23125 | Glutathione-disulfide reductase CDS | 1,356       |
| osmC                          | OsmC-like protein | B9Q16_23120 | Osmotically inducible protein C CDS | 558         |
| ybiB                          | Xylose kinase      | B9Q16_23115 | Xyulokinase CDS      | 1,449       |
| rbsC                          | Ribose transport system permease protein RbsC | B9Q16_23110 | Sugar ABC transporter permease CDS | 1,026       |
| rbsB                          | D-ribose-binding periplasmic protein precursor | B9Q16_23105 | LacI family transcriptional regulator CDS | 981         |
| rbsA                          | Ribose import ATP-binding protein RbsA | B9Q16_23100 | ABS transporter CDS | 1,503       |
| Hypothetical protein          |                   | B9Q16_23095 | Hypothetical protein CDS | 285         |
| hxB                           | 3-hexulose-6-phosphate isomerase | B9Q16_23090 | SIS domain-containing protein | 597         |
| fabG                          | 3-oxoacyl-[acyl-carrier-protein] reductase FabG | B9Q16_23085 | Short-chain dehydrogenase CDS | 753         |
| Rpe                           | Ribulose-phosphate 3-epimerase | B9Q16_23080 | Ribulose-phosphate 3-epimerase CDS | 645         |
| garC                          | Galactitol permease IIC component | B9Q16_23075 | PTS galactitol transporter subunit IIC CDS | 1,254       |
| garB                          | Galactitol-specific phosphotransferase enzyme IIB component | B9Q16_23070 | PTS galactitol transporter subunit IIB CDS | 294         |
| fruA                          | PTS system fructose-specific EIIABC component | B9Q16_23065 | PTS galactitol transporter subunit IIA CDS | 450         |
| sorC                          | Sorbitol operon regulator | B9Q16_23060 | Hypothetical protein CDS | 969         |
| Major Facilitator Superfamily |                   | B9Q16_23905 | MFS transporter CDS | 1,143       |
| argT                          | Lysine-arginine-ornithine-binding periplasmic protein precursor | B9Q16_23895 | ABC transporter substrate-binding protein CDS | 783         |
| dapL                          | LL-diaminopimelate aminotransferase | B9Q16_23890 | Succinyldiaminopimelate transaminase CDS | 1,203       |
| peF                           | Pectate lyase E precursor | B9Q16_23040 | Pectate lyase CDS | 1,194       |
| * Rhamnogalacturonase B, N-terminal |           | B9Q16_23030 | Rhamnogalacturonase B CDS | 1,610       |
| **ONION VIRULENCE REGION B (OVR-B)** |                   |            |                      |             |
| csrR                          | HTH-type transcriptional repressor CsrR | B9Q16_23870 | Transcriptional regulator CDS | 693         |
| Major royal jelly protein     |                   | B9Q16_23865 | Hypothetical protein CDS | 930         |
| Ureidoglycylate lyase         |                   | B9Q16_23860 | 2-hydroxyhepta-2,4-diene-1,7-dioate isomerase CDS | 840         |
| ydeO                          | HTH-type transcriptional regulator YdeO | B9Q16_23855 | AraC family transcriptional regulator CDS | 828         |
| * putative kinase inhibitor   |                   | B9Q16_23850 | Phosphatidylethanolamine-binding protein CDS | 564         |
| T5orf172 domain               |                   | B9Q16_23845 | Chromosome partitioning protein ParA CDS | 1,584       |
| hypothetical protein          |                   | B9Q16_23840 | IS110 family transposase | 207         |
| hypothetical protein          |                   | B9Q16_23835 | Hypothetical protein CDS | 192         |
| hypothetical protein          |                   | B9Q16_23830 | Hypothetical protein CDS | 978         |
| surE                          | 5′-nucleotidase SurE | B9Q16_23825 | 5′/3′ nucleotidase SurE | 777         |
| styA                          | transcriptional regulator StyA | B9Q16_23820 | Hypothetical protein CDS | 426         |
| acrF                          | Multidrug export protein AcrF | B9Q16_23815 | ACR family transporter CDS | 3,084       |
| bepF                          | Efflux pump periplasmic linker BepF | B9Q16_23810 | Efflux transporter periplasmic adapter subunit CDS | 1,146       |
| baeS                          | Signal transduction histidine-protein kinase BaeS | B9Q16_23805 | Two-component sensor histidine kinase CDS | 1,170       |
| phoP                          | Alkaline phosphatase synthesis transcriptional regulatory protein PhoP | B9Q16_23800 | DNA-binding response regulator CDS | 714         |
| mtpA                          | MltA-interacting protein MtpA | B9Q16_23795 | MltA-interacting MtpA family protein CDS | 783         |

(Continued)
TABLE 4 | Continued

| Gene     | Prokka annotation | NCBI locus | NCBI PGAP annotation | Length (bp) |
|----------|-------------------|------------|-----------------------|-------------|
| **ONION VIRULENCE REGION C (OVR-C)** |                  |            |                       |             |
| yjaB     | Uncharacterized N-acetyltransferase YjaB | B9Q16_23705 | Acetyltransferase      | 444         |
| puuR     | DNA-binding transcriptional repressor PuuR | B9Q16_23700 | Transcriptional regulator CDS | 585         |

**ONION VIRULENCE REGION D (OVR-D)**

| Gene     | Prokka annotation | NCBI locus | NCBI PGAP annotation | Length (bp) |
|----------|-------------------|------------|-----------------------|-------------|
| Major Facilitator Superfamily |             | B9Q16_23450 | Hypothetical protein CDS | 1,251        |
| Hypothetical protein |             | B9Q16_23445 | Hypothetical protein CDS | 903         |
| Cupin superfamily protein |             | B9Q16_23440 | Hypothetical protein CDS | 930         |
| Hypothetical protein |             | B9Q16_23435 | Hypothetical protein CDS | 1,170        |
| Hypothetical protein |             | B9Q16_23435 | Hypothetical protein CDS | 1,170        |
| speB     | Agmatinase       | B9Q16_23430 | Hypothetical protein CDS | 1,110        |
| eamA     | EamA-like transporter family | B9Q16_23425 | Hypothetical protein CDS | 880         |
| Hypothetical protein |             | B9Q16_23420 | Hypothetical protein CDS | 618         |
| sn-glycerol-3-phosphate dehydrogenase subunit A | | B9Q16_23415 | Hypothetical protein CDS | 1,134        |

Genes identified in Georgia pathogenic strains, and absent in non-pathogenic strains of Pantoea ananatis.
1 Predicted Transmembrane protein (TMHMM v 2.0).
2 Tat/ Sec Secreted target signal present (Signal P 4.1).

The OVR-A locus begins with a gene for a Cre-like phage recombinase, which may indicate integration of the 31-kb OVR-A locus as a mobilized genomic island. A gene closely following Cre is predicted to be a TetR family transcriptional regulator. Closely following this regulator are sugar transport proteins including a sugar ABC transporter permease and ribulose-phosphate-3-epimerase. The OVR-A contains three genes associated with sulfur, redox and oxidative stress, such as a thiol reductase and alkylhydroperoxidase. Several efflux pumps and transporters are present including an ABC transporter and PTS galactitol transporter subunit IIA-C. Amino acid metabolism-transport and plant cell wall degrading enzymes (CWDE) including a pectate lyase, and rhamnogalacturonoase B are also present in the OVR-A region (Table 4, Figure 5).

OVR-B consists of 16 CDS. Three proteins are predicted to have DNA-binding regulatory functions such as the AraC family transcriptional regulator. A ureidoglycylate lyase is associated with amino acid degradation. Two outer membrane transport proteins are present including an ACR family transporter. Two protein kinases are present including a predicted two-component sensor histidine kinase and a phosphatidylethanolamine-binding protein with a predicted secretion target signal. In addition, three proteins are present related to transposable DNA elements including an IS110 family transposase. An MlaA-interacting family protein lytic transglycosylases, which is related to protein binding membrane, is also present (Table 4).

OVR-C contains a transcriptional repressor with homology to PuuR, and an acetyl-transferase YjaB (Table 4). OVR-D contains nine predicted CDS. The PGAP annotation pipeline characterized all as hypothetical proteins. Prokka annotation predicted two transport associated proteins including a Major Facilitator Superfamily CDS (Table 4). In addition, two proteins involved in carbon and amino acid metabolism, including agmatinase are also present.

Several of the CDS in the regions specific to onion pathogenic strains have predicted transmembrane or Tat-secreted target signal. CDS of interest with Tat-secreted target signals include: CWDE pectate lyase, and rhamnogalacturonoase in the OVR-A and two protein signal regulators in OVR-B. Most proteins predicted to be transport-related contained transmembrane domains (Table 4).

**DISCUSSION**

*Phenotypic Variation amongst P. ananatis Strains*

A set of fifty *P. ananatis* strains from Georgia was assembled to maximize diversity of host and geographic sources of isolation as well as diversity of previously determined onion pathogenicity and ice nucleation phenotypes. We observed extensive phenotypic variation among a sub-set of 33 strains tested for pathogenicity and aggressiveness on multiple *Allium* hosts. In addition to the 15 primary onion and onion seed strains, 10 of 25 PANS strains collected from weed and thrips species were re-confirmed to be pathogenic on onion. However, aggressiveness of strains on *Allium* sp. varied considerably. Among the 33 strains, we observed similar levels of variation in pathogenic potential and aggressiveness on shallot and leek although the strains were generally less aggressive on these hosts than on onion. Pathogenetic potential and aggressiveness of a strain on onion or leek or shallot did not generally correspond with its potential to be pathogenic on the other *Allium* hosts. In addition, there were multiple examples of onion non-pathogenic strains that were pathogenic on leek or shallot and vice versa such as PANS 99-23. This could indicate some degree of host specialization among *P. ananatis* strains tested. Unlike other *Allium* hosts, the chive cv. Dolores was infected by only a single *P. ananatis* strain with mild symptoms. This may highlight the
potential of *Allium schoenoprasum* to serve as a source of genetic resistance against *P. ananatis*.

We used MLSA and rep-PCR based genetic diversity analysis to determine whether phenotypic diversity correlated with underlying phylogeny. Contrary to the high phenotypic variation, both analyses indicated little genetic diversity amongst the fifty *P. ananatis* strains. By MLSA all strains strain were found to be >99% identical. The majority of branches in the phylogeny had poor bootstrap value support and were composed of a mixture of PNA and PANS strains. We observed no underlying correlation between the isolate phylogeny and pathogenic phenotypes or isolation sources using either method of diversity analysis.

Motivated by the high degree of *P. ananatis* phenotypic variation that is unexplained by the low levels of core genome diversity, we sought to conduct a pan-genome analysis to determine whether fine scale variations in gene presence-absence might correlate with phenotypic variation. A sub-set of 10 strains from the original fifty strains were selected for whole genome Illumina shotgun sequencing with the goal of maximizing diversity of phenotypes and sources of isolation. Additionally, the Georgia prototype onion pathogenic strain PNA 97-1R was sequenced using PacBio to create a high quality near-closed genome assembly. Among the 10 strains selected for WGS, six were able to cause foliar necrosis in onion blades and five of those also were able to clear red onion scales (Figure 2). The phenotypic variation seen in red-onion scale assays may potentially be a useful tool for identifying isolates with onion pathogenic potential as it is easy to replicate and rapid with minimal space requirements. It would be beneficial to utilize more strains to determine if correlations between red onion scale clearing and onion foliar blight potential occur.

**Pan-Genome Analysis of Sequenced Isolates**

Dendrograms generated from pan-genomic gene presence-absence analysis clearly indicate a clustering of onion pathogenic strains from non-pathogenic strains. This suggests an underlying genetic difference between the two groups that was not detected using MLSA or rep-PCR approaches. The majority of the gene differences between onion pathogenic and non-pathogenic strains resides in four contiguous gene clusters the largest being 31 kb that we termed as the Onion Virulence Region (OVR-A). RAST BlastP further illustrated this point by identifying genes present in onion pathogenic strains, but absent in non-pathogenic strains when compared with the pathogenic prototype strain PNA 97-1R. The level of core genome homology amongst all the strains is notable. There is a high degree of CDS similarity. Most gene variation is the result of phage, transposase, and integrase proteins. In addition, homologs to the LuxI, type VI, V, I s, and Tat secretion systems previously noted as candidate virulence factors were identified in both onion pathogenic and non-pathogenic strains. Phytotoxin synthetic clusters were not identified (Weller-Stuart et al., 2017).

The *de novo* genome assembly of the Georgia prototype strain PNA 97-1R resulted in three contigs, of 4.9 Mbp, 295 kbp, and 180 kbp (CP020943-CP020945). The 4.9 Mbp contig corresponds to the *Pantoea* circular chromosome. The 295 kbp contig was identified as the Large Pantoea Plasmid LPP-1 based on previously described genes and MAUVE alignment. The genomes of the sequenced strains contain LPP-1 with a high level of conservation (Supplementary Figure 3). The third 180 kbp PNA 97-1R contig encompasses the four OVR loci which correlate with onion pathogenicity based on pan genomic analysis. This contig aligned with single contigs of varying sizes (74–167 kbp) from onion pathogenic WGS strains. The 180 kbp contig shares complete homology in 6 MAUVE alignment blocks with all genes in a 60 kb contig (CAEI100108.1) of a single sequenced strain B1-9 of *P. ananatis* in the NCBI database (NZ_CAEI00000000.1). Strain B1-9 is an isolate from green onion described by researchers in South Korea associated with growth promotion of pepper (Kim et al., 2012). It is intriguing that a growth promoting strain from South Korea share all genes associated with onion virulence among our Georgia strains. We know that the pathogenicity of *P. ananatis* strains can vary widely between different species of *Allium* (Table 1). It would be interesting to determine if B1-9 is pathogenic on onion and on other *Allium* sp. In addition to sharing homology with this contig from B1-9, the 180 kbp contig shares some genes with those found on the pEcWSU1 plasmid of the onion pathogen, *Enterobacter cloacae* associated with onion bulb-rot (Schoeder et al., 2009). Highest homology is shared between a histidine kinase protein in OVR-B, and the PTS galactitol transporter system A-C in the OVR-A loci. Interestingly, plasmid partitioning genes and repA plasmid replication genes were present on the WGS contigs that contained the OVR-A loci in each of the onion pathogenic strains indicating that these contigs may, in fact, represent an accessory plasmid.

*Pantoea ananatis* is a species that is frequently associated with diverse eukaryotic hosts. In the onion-center rot system, it is associated epiphytically with weed species, and pathogenically and endophytically with onions, and persists temporarily in the thrips vector commonly associated with center-rot disease outbreaks (Gitaitis et al., 2002; Walcott et al., 2002; Dutta et al., 2014). Our strains were isolated from various sources such as thrips, weeds, seeds, and symptomatic onions. Despite these strains inhabiting different niches, there existed a high degree of genetic similarity among them. Furthermore, a high average nucleotide identity was observed among Georgia isolates and other strains isolated in different regions around the world (Table 3). This suggests that the core genome is highly conserved among *P. ananatis* strains (Supplementary Figure 3). The core genome of 3,750 CDS identified through ROARY pan-genomic analysis is consistent with the core genome identified through previous pan-genomic analysis methods (De Maayer et al., 2014; Sheibani-Tezerji et al., 2015). The accessory genome, where most differences were identified, revealed the Onion Virulence regions (OVR-AD), as well as differences among poorly characterized hypothetical coding sequences, phages, transposases, integrases, and mobile genetic elements. The presence of the *Cre* recombinase in the OVR-A loci could be indicative of a possible horizontal gene transfer mechanism.
Genetic Content of the OVR-A Loci

The OVR-A loci has not been previously described and may offer insights into the ability of *P. ananatis* to cause bacterial disease in onions. This region was present in all Georgia strains that caused the foliar symptoms on onion (Figure 2). Except for strain PNA 200-3, this region was present in all *P. ananatis* strains that caused scale clearing in the red onion scale assay. The region begins with a Cre recombinase followed by amino acid transporters, metabolic enzymes, transcriptional regulators and CWDEs (Table 4).

The presence of redox-related/sulfur reducing enzymes could play a role in withstanding host sulfur containing antimicrobials, or interrupting ROS signaling that is important during plant immunity (Torres et al., 2006). In addition, amino acid transporters may play a role in the absorption of host nutrients. The specific localization of *P. ananatis* in host tissue has not been well characterized and may offer insights into nutrient acquisition for this bacterium (Fatima and Senthil-Kumar, 2015).

Sugar transporters and major super facilitator proteins have been shown to play an important role in the pathogenicity of bacterial pathogens such as *Dickeya dadantii* where they can play a role in the anti-microbial resistance, and chemotaxis (De Maayer et al., 2011).

Plant cell wall degrading enzymes such as *pelE* and rhamnogalacturonanase, have been found to contribute to pathogenicity in a number of other bacterial pathogens, notably *Dickeya dadantii* (Hassan et al., 2013). A protein in OVR-D was computationally annotated as a member of the *EamA* transport family shares homology with the *PecM*, a transporter in *D. dadantii*, which plays a major role in regulating virulence factor synthesis such as, *PelE* and other CWDEs (Praillet et al., 1996). However, as *P. ananatis* lacks a T2SS it is unclear by what mechanism these proteins would be secreted out of the cell. The protein kinase and kinase inhibitor found in OVR-B are intriguing as they both have predicted secretion signals (putative kinase inhibitor, two-component sensor histidine kinase) (Table 4). They could play a role in signal transduction in the bacteria or perhaps could be involved in disrupting host immunity by some as-of-yet uncharacterized host cell delivery mechanism. Transporter and transmembrane proteins located in onion virulence correlated regions could be involved in secretion of proteins across the outer membrane, or surface interactions between the bacteria and host cells.

The localization of OVR A-D to single contigs from the genome constructs offers some supportive evidence that the region is plasmid localized. Symbiotic and pathogenic plasmids are important for pathogenicity in a number of phytopathogenic bacterial species. In plant pathogens mega plasmids have been shown to contribute to virulence in bacteria such as *Pseudomonas syringae pv. phaselicola* (Jackson et al., 1999). The contigs constructs containing OVR loci contain aA plasmid partitioning proteins. In our future work work we will seek to confirm whether the OVR loci are, in fact, plasmid-borne.

Few pan-genomic analyses of *P. ananatis* strains have been conducted. The first pan-genomic study of *P. ananatis* utilized NCBI draft and complete genomes of strains isolated from various hosts. It is difficult to infer the genomic diversity of strains isolated from a single geographic location or host-pathosystem from such, studies, as hosts, isolation sources, and locations varied dramatically (De Maayer et al., 2014).

Currently, only one study elucidated the genomic variation among *P. ananatis* strains, isolated from the same host, maize seed. The researchers observed a high degree of genetic similarity among *P. ananatis* strains with diverse phenotypes; epiphytic, plant growth-promoting, and pathogenic (Sheibani-Tezerji et al., 2015). Despite high levels of genetic similarity, minute differences in genes encoding protein secretion systems, putative effectors, and transpose/integrase/phage related genes were noted. For example, the pathogenic *P. ananatis* genome contained genes relating to cell cycle control, cell division, chromosome partitioning, and amino acid transport that differentiated it from the growth promoting, and endophytic strain. This study found subtle variations between the pathogenic and non-pathogenic strains. Other protein coding sequences that differed between the strains included transposase related proteins, chemotaxis proteins, and T6SS loci (Sheibani-Tezerji et al., 2015). Conversely, our results identified a distinct subset of genes in contiguous regions that correlated with onion pathogenicity.

The presence of genes within the OVR-A loci associated with virulence and pathogenicity in other onion pathogens such as *Enterobacter cloacae* illustrates a need to further characterize genes in this region and other onion virulence correlated regions. Future work will include the deletion of the OVR-A loci from PNA 97-1R to determine what role this region plays in onion pathogenicity.

Strain PNA 200-3 has the OVR-A region but lacks the scale clearing phenotype. The negative-scale clearing phenotype may be explained by SNP missense mutations in genes encoding CWDEs of PNA 200-3s OVR-A or could indicate variation in expression of genes associated with this loci. The OVR-A loci may simply represent a vertically inherited locus or plasmid common to Georgia onion pathogenic strains through a common clonal origin of onion pathogenic strains in Georgia, and possible relation to strains in South Korea. Thus, while the OVR-A loci is common among Georgia onion pathogenic strains, it may not in itself, carry pathogenicity determinants that are critical to infection. Many other currently sequenced strains have been reported as pathogenic on onion, while apparently lacking the OVR loci (Morohoshi et al., 2007; Shyntum et al., 2015).

CONCLUSIONS

Overall, our study illustrates the importance of WGS in differentiating closely related strains. MLSA and rep-PCR analysis indicated little diversity among the *P. ananatis* strains in Georgia, although the strains were phenotypically diverse for *Allium* infection phenotypes. The WGS revealed that although these strains share similar core genomes, the universal plasmid LPP-1, and previously identified candidate virulence factors, differences in the mobile genetic elements, as well as the OVR-A loci differentiated the sequenced strains into two groups that shared common scale-clearing and foliar pathogenicity
phenotypes. Thus, the differences between pathogenic and non-pathogenic strains in *P. ananatis* may not be as subtle as previously suggested (Sheibani-Tezerji et al., 2015). In any case, the contribution of the OVR loci to *P. ananatis* onion virulence remains to be determined. Subtle differences in loci gene expression and host defense reactions may further differentiate host range and aggressiveness of *P. ananatis* strains. The transcriptional analysis of mutants with and without these loci may provide insights into the nature of this regions contributions to virulence. Future studies will involve the use of mutational analysis to assess the role of the OVR-A loci in onion pathogenicity and determine if this genetic locus is chromosomal or plasmid-borne.

**AUTHOR CONTRIBUTIONS**

SPS and SDS contributed equally to this work. BK and BD are co-senior on this work. SPS, SDS, RG, BK, and BD conceived and designed the experiments. SDS performed MLSA, rep-PCR and phenotypic testing. SPS conducted experiments and analysis related to whole genome sequencing and Pan-genome analysis. SPS, SDS, BK, and BD wrote and edited the manuscript.

**FUNDING**

We would like to acknowledge the Vidalia Onion Committee for partial funding to BD as well as NIFA HATCH (Project: GEO00791) and the State of Georgia for funding to BK.

**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: [https://www.frontiersin.org/articles/10.3389/fmicb.2018.00184/full#supplementary-material](https://www.frontiersin.org/articles/10.3389/fmicb.2018.00184/full#supplementary-material)

**Supplementary Figure 1** | Dendrogram derived by cluster analysis of similarities between *Pantoea ananatis* strains based on repetitive extragenic palindromic polymerase chain reaction (REP-PCR) DNA genomic profiles. Analysis was performed using Dice’s (1945) coefficient in BioNumerics software package (Applied Math, Kortrijk, Belgium).

**Supplementary Figure 2** | Large Pantoea Plasmid LPP-1 alignment of proto-type strain PNA 97-1 with AJ13355 PE9320 (Accession CP020943; NC_017533.1). Alignment generated with Geneious MAUVE plugin.

**Supplementary Figure 3** | Venn diagram of genes shared among the pathogenic isolate PNA 97-1R; non-pathogenic isolates PANS 99-36, and genomic type strain LMG 20103 (Accession CP001875.2). The number of coding sequences are derived from the gene_presence_absense.csv output of the ROARY pan-genome analysis using default parameters (Supplementary File 2).

**Supplementary Figure 4** | wgMLST cladogram of Pantoea strains generated through PGAdo-builder platform. Outgroups *P. agglomerans* P10C and *P. stewartii* subsp. indologens LMG 2632 (Accession NZ_LIME0000000.1 and JPK00000000.1).

**Supplementary Figure 5** | Phaster predicted phage regions in PNA 97-1R. Green indicated intact phage regions whereas red indicates incomplete phage regions.

**Supplementary Figure 6** | BRIG analysis of *P. ananatis* strains and the OVR A-D loci to the reference strain PNA 97-1R. BRIG is a locally run analysis that generates figures based on blastn comparisons of query genomes to a reference strain.

**Supplementary Table 1** | Sequencing and assembly statistics for Illumina draft-genomes of *Pantoea ananatis* strains.

**Supplementary Table 2** | RAST functional categories for predicted protein coding sequences of *Pantoea ananatis* strains.

**Supplementary Table 3** | Primers and conditions used for amplification and sequencing.

**Supplementary Table 4** | IslandPath DIMOB genomic islands identified in WGS *P. ananatis* strains.

**Supplementary File 1** | ROARY 10 strain WGS Analysis. The presence of each CDS in the respective genome is listed below each labeled strain column. If a cell is blank the respective strain does not have the gene. Logic equations were used to highlight cells, resulting in genes present in pathogenic strains and absent in non-pathogenic strains.

**Supplementary File 2** | ROARY 3 strain WGS Analysis. The presence of each CDS in the respective genome is listed below each labeled strain column. If a cell is blank the respective strain does not have the gene. Summation equations were used to calculate the numeric values for the Venn Diagram in Supplementary Figure 3.

**REFERENCES**

Abe, K., Watabe, S., Emori, Y., Watanabe, M., and Arai, S. (1989). An ice nucleation active gene of erwinia-ananas-sequence similarity to those of pseudomonas species and regions required for ice nucleation activity. *FEBS Lett.* 258, 297–300. doi: 10.1016/0014-5793(89)81678-3

Alíkhán, N. F., Petty, N. K., Ben Zakour, N. L., and Beatson, S. A. (2011). BLAST Ring Image Generator (BRIG): simple prokaryote genome comparisons. *BMC Genomics* 12:402. doi: 10.1186/1471-2164-12-402

Bertelli, C., Laird, M. R., Williams, K. P., Lau, B. Y., Hoad, G., Winsor, G. L., et al. (2017). IslandViewer 4: expanded prediction of genomic islands for larger-scale datasets. *Nucleic Acids Res.* 45, W30–W35. doi: 10.1093/nar/gkw343

Carr, E. A., Bonasera, J. M., Zaid, A. M., Lorbeer, J. W., and Beer, S. V. (2010). First report of bulb disease of onion caused by *Pantoea ananatis* in New York. *Plant Dis.* 94, 916–916. doi: 10.1094/PDIS-94-7-0916B

Carr, E. A., Zaid, A. M., Bonasera, J. M., Lorbeer, J. W., and Beer, S. V. (2013). Infection of onion leaves by *Pantoea ananatis* leads to bulb infection. *Plant Dis.* 97, 1524–1528. doi: 10.1094/PDIS-06-12-0597-RE

Coplın, D. L., Frederick, R. D., Majerczak, D. R., and Tuttle, L. D. (1992). Characterization of a gene-cluster that specifies pathogenicity in erwinia-stewartii. *Mol. Plant Microbe Interact.* 5, 81–88. doi: 10.1094/MPMI-5-081

Coutinho, T. A., Preisig, O., Mergaert, J., Cnockaert, M. C., Riedel, K. H., Swings, J., et al. (2002). Bacterial blight and dieback of Eucalyptus species, hybrids, and clones in South Africa. *Plant Dis.* 86, 20–25. doi: 10.1094/PDIS.2002.86.1.20

Coutinho, T. A., and Venter, S. N. (2009). *Pantoea ananatis*: an unconventional plant pathogen. *Mol. Plant Pathol.* 10, 325–335. doi: 10.1111/j.1364-3703.2009.00542.x

Darling, A. C. E., Mau, B., Blattner, F. R., and Perna, N. T. (2004). Mauve: multiple alignment of conserved genomic sequence with rearrangements. *Genome Res.* 14, 1339–1403. doi: 10.1101/gr.2289704

De Baere, T., Verhelst, R., Labit, C., Verschraegen, G., Wauters, G., Claeyss, G., et al. (2004). Bacteremic infection with *Pantoea ananatis*. *J. Clin. Microbiol.* 42, 4393–4395. doi: 10.1128/JCM.42.9.4393-4395.2004

Delétot, E., Decré, D., Courant, S., Passet, V., Audo, J., Grimont, P., et al. (2009). Phylogeny and identification of *Pantoea* species and typing of *Pantoea* agglomerans strains by multilocus genome sequencing. *J. Clin. Microbiol.* 47, 300–310. doi: 10.1128/JCM.01916-08

De Maeyer, P., Chan, W. Y., Blom, J., Venter, S. N., Duffy, B., Smits, T. H. M., et al. (2012). The large universal Pantoea plasmid LPP-1 plays a major role in biological and ecological diversification. *BMC Genomics* 13:625. doi: 10.1186/1471-2164-13-625
De Maayer, P., Chan, W. Y., Rubagotti, E., Venter, S. N., Toth, I. K., Birch, P. R. J., et al. (2014). Analysis of the Pantoea ananatis pan-genome reveals factors underlying its ability to colonize and interact with plant, insect and vertebrate hosts. BMC Genomics 15:404. doi: 10.1186/1471-2164-15-404

De Maayer, P., Venter, S. N., Kamber, T., Duffy, B., Coutinho, T. A., and Smits, T. H. M. (2011). Comparative genomics of the type VI secretion systems of Pantoea and Erwinia species reveals the presence of putative effector islands that may be translocated by the VgrG and Hcp proteins. BMC Genomics 12:576. doi: 10.1186/1471-2164-12-576

Dutta, B., Barman, A. K., Srinivasan, R., Avci, U., Ullman, D. E., Langston, D. B., et al. (2014). Transmission of Pantoea Ananatis and P-agglomerans, causal agents of center rot of onion (Allium cepa), by Onion Thrips (Thrips tabaci) through feces. Phytopathology 104, 812-819. doi: 10.1094/PHYTO-07-13-0199-R

Fatima, U., and Senthil-Kumar, M. (2015). Plant and pathogen nutrient acquisition strategies. Front. Plant Sci. 6:750. doi: 10.3389/fpls.2015.00750

Gaititis, R. D., and Gay, J. D. (1997). First report of a leaf blight, seed stalk rot, and bulb decay of onion by Pantoea ananatis in Georgia. Plant Dis. 81, 1096–1096. doi: 10.1094/PD.I.S.1997.81.1096C

Gaititis, R., Walcott, R., Culpepper, S., Sanders, H., Zolobowska, L., and Langston, D. (2002). Recovery of Pantoea ananatis, causal agent of center rot of onion, from weeds and crops in Georgia, USA. Crop Prot. 21, 983–989. doi: 10.1016/S0965-2191(02)00078-9

Hassan, S., Shevchik, V. E., Robert, X., and Hugouvieux-Cotte-Pattat, N. (2013). PelN is a new pectate lyase of Dickeya dadae with unusual characteristics. J. Bacteriol. 195, 2197–2206. doi: 10.1128/JB.02118-12

Jackson, R. W., Athanassopoulos, E., Tsiamis, G., Mansfield, J. W., Sesma, A., Hassan, S., Sheibani-Tezerji, R., Naveed, M., Jehl, M. A., Sessitsch, A., Rattei, T., and Hugouvieux-Cotte Pattat, N. (2013). Pantoea ananatis utilizes a Type VI secretion system for pathogenesis and bacterial competition. Mol. Plant Microbe Interact. 28, 420–431. doi: 10.1094/MPMI-07-14-0219-R

Tabei, H., Kvitko, N. A., and Dutta, A. (2007). Natural infestation of onion seed by Pantoea ananatis. Plant Dis. 91, 323–323. doi: 10.1094/PDIS-93-7-0323A

Zhai, Ban, Y., Wu, J. Y., Yang, J. H., Sun, S. X., Xiao, J. F., and Yu, J. (2012). PGAAP, pan-genomes analysis pipeline. Bioinformatics 28, 416–418. doi: 10.1093/bioinformatics/bts655

Conflict of Interest Statement: 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 © 2018 Stice, Stumpf, Gaititis, Kritko and Dutta. 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 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.