### **CONTINUING PROJECT REPORT**

Project Title: Development of New Biocontrol Strains from Washington Native Trees

PI: Professor Sharon Doty Organization: University of Washington Telephone: (206) 616-6255 Email: sldoty@uw.edu Address: Winkenwerder Hall, Box 352100 City/State/Zip: Seattle, WA 98195-2100

#### **Cooperators**:

Dr. Achour Amiri (WSU-Wenatchee), a.amiri@wsu.edu Dr. Tianna DuPont (WSU-Wenatchee), tianna.dupont@wsu.edu Dr. Patricia Okubara (WSU-Pullman), patricia.okubara@usda.gov

Total Project Request: Year 1: \$46,229

WTFRC Budget: None

Budget 1

Organization Name: University of Washington Contract Administrator: Carol Rhodes, Director, Office of Sponsored Programs Telephone: (206) 543-4043 Email address: osp@uw.edu

Item	2020	2021
Salaries	\$28,632	
Benefits	\$8,775	
Wages		
Benefits		
Equipment		
Supplies	\$5,000	
Travel	\$822	
Miscellaneous	\$3,000	
Plot Fees		
Total	\$46,229 (Total Year 1)	0

## **OBJECTIVES**

- 1) Testing for biocontrol of *Erwinia amylovora* (causal agent of fire blight) using native plant microbiota from Washington State
  - a. Assay development
  - b. Testing of our fully characterized biocontrol strains in the assays optimized in Objective 1a
  - c. Screening for new microbial strains with activity against *E. amylovora*.
  - d. Genomic sequencing of selected strains
- 2) Testing for biocontrol of pre- and post-harvest apple fruit pathogens using native plant microbiota from Washington State
  - a. Testing our fully characterized biocontrol strains
  - b. Screening for new microbial strains with activity against the pre- and post-harvest apple pathogens
  - c. Genomic characterization of selected strains.

# SIGNIFICANT FINDINGS

- Through this project, we isolated over a hundred new endophyte strains from the Wenatchee, Entiat, Yakima, and Methow areas
- 14 strains showed inhibition of *Penicillium expansum*. Since the strains grew in the presence of this fungus known to produce the antimicrobial compound, patulin, they may have the capacity to degrade it
- 27 strains inhibited *Botrytis cinerea*
- 21 strains inhibited *Neofabraea perennans*
- 38 strains inhibited *Phacidiopycnis washingtonensis*
- 40 strains inhibited *Erwinia amylovora*
- Several of the strains appeared to inhibit the pathogenic fungi through production of volatile compounds
- The current status of this project (continuing through a No Cost Extension) is the preparation of genomic DNA and full genomic sequencing of the top-performing candidate strains

### **METHODS**

**Isolation of new endophyte strains from natural areas near to the fruit tree growing areas.** Doty obtained the required permits and sampled a variety of native plants in natural sites in the Wenatchee, Entiat, Yakima, and Methow areas. Microbial endophyte strains (bacteria and yeast from within plant tissues) were isolated through maceration in bacterial media and streak purification.

In vitro assay for inhibition of the post-harvest decay pathogens, *Penicillium expansum*, *Botrytis cinerea*, *Neofabraea perennans*, and *Phacidiopycnis washingtonensis*. The fungal samples were obtained from the Amiri Lab at the WSU Tree Fruit Research and Extension Center in Wenatchee. Using a modification of the dual plate assay we had used previously (Kandel et al. 2017), we pipetted 10 µl fungal spore/suspended hyphae preparations to the center of agar plates containing medium appropriate for fungal growth, PDA (potato dextrose agar). The fungi were allowed to grow at room temperature until robust fungal growth was evident in the center of the plate. Endophyte isolates, which had been grown on rich media (MGL), were then spotted around the edge of the plate with up to eight isolates per plate (**Figure 1**). Due to the slower growth of *Neofabraea perennans* only up to four isolates were spotted per plate and half the distance from the hyphae edge (**Figure 2A**). Growth of the fungus was monitored and inhibition was scored when the fungal growth reached the

perimeters of the plates, except *Neofabraea perennans* which was scored based on hyphae growth disruption (**Figure 2B**).

*Erwinia amylovora in vitro* inhibition assay. Three *Erwinia amylovora*. isolates were provided by Dr. Tianna DuPont, however after initial testing indicated the three strains displayed identical inhibition patterns, assays were carried out on a mixture of the three strains provided. 100  $\mu$ l of *Erwinia amylovora* with an optical density of 0.01 at 600nm were spread onto rich medium appropriate for *Erwinia* (NYDA). Endophyte isolates were grown on MGL and then spotted onto these plates of *Erwinia amylovora*. Clear zones on the *Erwinia* lawns were scored as inhibitory activity (Figure 3).

# **RESULTS AND DISCUSSION**

A total of 38 strains inhibited the growth of *Phacidiopycnis washingtonensis*, 21 strains inhibited *Neofabraea perennans*, 27 strains inhibited *Botrytis cinerea*, and 14 strains inhibited *Penicillium expansum*. Many of the strains strongly inhibited the growth of *Erwinia amylovora*, with a total of 40 inhibitory strains. (**Table 1**).

**Figure 1.** Inhibition of *Botrytis cinerea* by some of the endophyte strains. A) Example plate showing no inhibition of the fungus. B) Strong inhibition of the fungus by sample # 88. C) Apparent inhibition by volatiles produced by some of the strains, as indicated by the overall reduced fungal growth and the bubbling appearance of some of the samples.



**Figure 2.** Inhibition of *Neofabraea perennans*. A) Example of four spotted endophyte isolates. B) Close up of disrupted hyphae growth near sample #4, with the leading edge of fungal growth becoming filiform as opposed to the smooth edge seen near sample #1.



**Figure 3.** Inhibition of *Erwinia amylovora*. The pathogen was inhibited by several of the endophyte strains as indicated by clearing zones on the lawn of *Erwinia* growth. A. Overall screening results. B. Close up of one of the assay plates showing the strong inhibition of *Erwinia* growth.



**Table 1.** Apple biocontrol project screening results. Endophyte strains with any activity against each of the pathogens. Pw *Phacidiopycnis washingtonensis*, Np *Neofabraea perennans*, Bc *Botrytis cinerea*, Pe *Penicillium expansum*, and Ea *Erwinia amylovora*. Strain names preceded by a number indicate the site from which they were isolated: 1 Wenatchee area, 2 Entiat/Okanagan area, 3 Yakima River area near Ellensburg, and 4 Methow area.

Pw	Np	Bc	Pe	Ea
1SS-L-D	1SS-L-D	1SS-L-D	1SS-L-H	1 SS-L-C
ISS-L-E	1SS-L-F	4_2_2	1Cv-L-C	4RDLD
1SS-L-F	1SS-L-J	WP 40	WP 40	2RDLC
1SS-L-H	4_5_3	WP 41	WP 41	2RDLD
1SS-L-I	4_4_2	WP 42	WP 42	3Pop12L1
1SS-L-J	WP 40	AFE 4A	AFE 4A	3YPLB
1Cv-L-C	WP 41	AFE 21B	WPB	2ALE2
1 SS-S-A	WP 42	AFE 5	AFE 3	2PtLE
4_2_2	AFE 4A	1 SS-A	2PtLD	2OPSB
4_5_3	AFE 21B	1 SS-B	3YPLB	3RS1
4_3_2	1 SS-S-B	1 Cv-S-A	4ASD	3RS3
4_4_2	WW7B	AFE 8	4RDLI	3Pop12L4
WP 40	AFE5	WPB	4RDLJ	3YPS2
WP 41	AFE9	4RLD	4RDLG	3YPS3
WP 42	AFE14	4RDLD		2 OPSB
PTD1	2PtLD	3ThS2		2PtLC2
AFE 4A	2SASA	20PSA		2PTLF1
AFE 21B	2ALE2	2PtLD		2SASD

4RDLD	2RDSA	2SASA	2RDSB
3ThS2	2PtLE	2RDLC	2RDLA
2PtLD	2OPSB	2RDLD	2ALA1A2
2SASA		4SBLB-	2ALB
2RDLC		3WL2	4RLA
2RDLD		3WL3	4RLE
3WL2		3Pop12L1	4RFA
3WL3		3YPLB	4RFB
3Pop12L1		3YPLD	4RSC
3YPLB			4ASA
3YPLD			4ALB
3RS1			4ALC
4ASD			4RDLA
3RS3			4RDLE
3ThS1			4RDLF
3Pop12S3			4HNLA
3Pop12L3			4HNLB
3Pop12L4			4SBLA
3YPS2			3RF1
3YPS3			3RL2
			3ThS3
			3ThL1

**Table 2.** Endophyte isolates chosen for sequencing after rRNA preliminary identification.

3YPLB	Pseudomonas
1SS-L-D	Erwinia
3Pop12L1	Pseudomonas
1SS-L-F	Schwanniomyces
3ThS2	Pseudomonas
3WL2V	Acinetobacter
3WL3V	Acinetobacter
3YPLD	Pseudomonas
3RS1	Enterobacter/Pantoea
4ASD	Sphingomonas
3YPS3	Pseudomonas
4RDLJ	Sphingomonas
4RDLG	Rhizobium
1SS-L-J	Rhodotorula
2PtLD	Serratia

**Genomic DNA Sequencing.** Fifteen strains were chosen for full genomic sequencing based on the number of pathogens towards which the strain was inhibitory, the strength of the inhibitory activity, and uniqueness of the strain compared to the other top-performing strains (**Table 2**). We now have an agreement with GeneWiz for the sequencing of these strains and can begin preparing the genomic DNA. Sequence data is expected by late March, prior to the start of our proposed Phase 2 project.

## KEYWORDS, ABSTRACT AND EXECUTIVE SUMMARY

Keywords: Fire blight; Erwinia amylovora, post-harvest decay; Penicillium expansum, Botrytis cinerea, Neofabraea perennans, Phacidiopycnis washingtonensis

New microbial endophyte strains were isolated from native plants in natural areas near apple tree growing areas of Wenatchee, Entiat/Okanagan, Yakima, and Methow. A total of 119 strains (15 previously characterized and 104 new isolates) were screened using *in vitro* assays for inhibition of the post-harvest decay pathogens, *Penicillium expansum, Botrytis cinerea, Neofabraea perennans,* and *Phacidiopycnis washingtonensis*, as well as the causal agent of fire blight, *Erwinia amylovora.* Two to three dozen inhibitory strains for each pathogen were identified. Preliminary rRNA sequence characterization was used to screen for isolates related to potential human pathogens and a subset of the isolates with the strongest or broadest activities was selected for full genomic sequencing.