

## Renewal Progress Report for CDFA Agreement Number 12-0224-SA

### Elucidating process of cell-cell communication in *Xylella fastidiosa* to achieve Pierce's disease control by pathogen confusion

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#### Introduction:

Our work has shown that *Xylella fastidiosa* (*Xf*) uses DSF perception as a key trigger to change its behavior within plants. Under most conditions DSF levels in plants are low since cells are found in relatively small clusters, and hence they do not express adhesins that would hinder their movement through the plant (but which are required for vector acquisition) but actively express extracellular enzymes and retractile pili needed for movement through the plant. Disease control can be conferred by elevating DSF levels in grape to "trick" the pathogen into transitioning into the non-mobile form that is normally found only in highly colonized vessels. While we have demonstrated the principles of disease control by so-called "pathogen confusion", more work is needed to understand how best to alter DSF levels in plants to achieve even higher levels of disease control. Until now we have suffered from a lack of sensitive methods to detect DSF levels in plants (the *Xanthomonas*-based bioassay we have used previously is relatively insensitive to the chemically distinct forms of DSF produced by *Xf*). That is, while we showed that DSF-producing endophytes, direct application of DSF, and transgenic plants producing DSF all conferred some resistance to disease, we had no way to know why they were not more resistant, nor what would be needed to improve control measures since we could not measure the direct effect of our efforts to increase DSF levels in plants. However, in the last 2 years we have developed several new sensitive biosensors that enable us to measure *Xf* DSF both in culture and within plants. We have recently found that *Xf* apparently produces more than one molecule that can act as a DSF signal molecule and that the molecules made by *Xf* are dependent on its growth environment. We thus need to ascertain which form is most active, whether the various forms all have the same effect on regulating traits in *Xf*, and what are their fates when applied to plants in various ways. We also know that related molecules can interfere with the signaling process and thus plant susceptibility may be influenced by the suite of compounds normally present in their xylem sap. Thus the overall goals of the proposed work

is to use these new biosensors for DSF to examine how DSF levels can best be altered by the various methods we have previously identified. As disease control should be directly proportional to both the concentration of and dispersal of DSF within plants we will quantitatively explore the effectiveness of different strategies to elevate DSF levels throughout plants.

We also have made the discovery that XF produces abundant extracellular membranous vesicles which are shed from the cell. Importantly, the content of outer membrane proteins including the adhesion XadA are controlled by DSF accumulation in cultures of XF, and even more importantly, the shedding of these vesicles from the cell is apparently suppressed by the accumulation of DSF. We therefore are testing the model that DSF signaling in XF involves two very different processes both of which lead to a rapid, cell density dependent change in its adhesiveness. When cells of XF are found in relatively low numbers within the xylem vessel they have accumulated little DSF, and because of this they do not produce large amounts of the cell surface adhesins including XadA, HxfA, and others, yet such cells shed large numbers of vesicles. However, when cell density, and thus DSF concentrations increase membrane vesicles are not shed by the cell, and the higher concentration of afimbral adhesins would be retained on the surface of the XF cells rather than being fed into the environment. Such a process would tend to maximize the adhesiveness of XF when DSF levels increased. This increased adhesiveness is apparently needed for their acquisition by insect vectors but would be expected to suppress their ability to move in the plant. In contrast, the shedding of vesicles would tend to prevent access from adhering to surfaces because most cell surface adhesins would no longer be attached to the cell. In this project, we are testing the role of the membranous vesicles, since preliminary data revealed that they may interfere with the adherence of XF to surfaces. That is, by shedding adhesive vesicles which themselves adhere to the surface of plants, access may prevent its own adherence to such surfaces because they are now coded would such vesicles.

### **Specific objectives:**

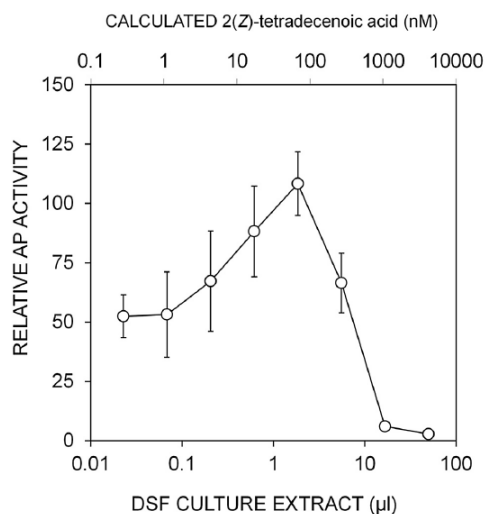
- 1) Identify additional DSF molecules made by *Xf* that contribute to cell-cell signaling and determine their movement and stability when applied to plants in various ways to improve disease control.
- 2) Determine the contribution of membrane vesicles shed by *Xf* in the absence of DSF to its virulence and the utility of measurement of vesicular presence within plants as a sensitive means to assess the success of strategies of disease control by pathogen confusion.

### **Description of activities conducted to accomplish each objective and summary of accomplishments:**

Objective 1: Finding new DSF species. We have optimized methods to use *Xf* itself to detect DSF. Among the several genes that we know to be most strongly regulated by DSF are genes such as *hxfA* and *HxfB* which are involved in cell-surface adhesion. Previous attempts to establish *gfp* or *inaZ*-based transcriptional fusions in *Xf* failed, presumably due to its incapability to express foreign genes properly. We now have successfully used the endogenous *phoA* gene (encoding alkaline phosphatase) as a bioreporter of gene expression in *Xf*. The PhoA-based biosensor in which *phoA* is driven by the *hxfA* promoter is quite responsive to exogenous DSF from extracts of *Xf* cultures as well as C14-cis itself. This biosensor, which we will refer to as *Xf:phoA* has been used extensively as it provides direct measures of the responsiveness of *Xf* to different molecules.

Assay of *Xf* extracts by *Xf* DSF-specific biosensors provide evidence of more than one *Xf* DSF molecule. The use of a *Xanthomonas*-based GCF-*gfp* biosensor and *Xf:pho* reveals that *Xf* produces at least 3 different compounds with DSF signaling activity. *Xf* appears to be more responsive to enoic acids of longer chain length than those produced by *Xcc*. While the

Xcc biosensor pKLN55 was maximally responsive to enoic acids of 12 Carbon in length (the size of each of the three DSF species made by Xcc) the GCF-gfp biosensor was maximally responsive to enoic acids of 13 and 14 Carbons in length. It thus would be expected that the various DSF molecules made by Xf would tend to be longer than those made by Xcc. We have recently published data showing that 2-Z-tetradecenoic acid (hereafter called C14-cis) is produced by Xf. We also have obtained circumstantial evidence that it also produces C12-cis. The relative activity of C12-cis in Xf is apparently somewhat less than that of C14-cis that we have more extensively studied. Importantly, an unidentified DSF species, presumably another enoic acid, is apparently much more active than either C12-cis or C14-cis. The amount of DSF, and apparently also the composition of species comprising the DSF signaling molecules are apparently strongly influenced by the environment in which Xf grows. A very high yield of DSF signaling molecules as detected with GCF-gfp was obtained in Xf cultures grown in PD3 medium compared to PWG: the signaling activity of cells Xf:phoA grown in PD3 as detected by the GCF-gfp biosensor was MUCH higher than that that in PWG media. Chemical analysis of these extracts revealed the presence of only very small amounts of either C14-cis or C12-cis, indicating that the very high signaling activity is due to a new compound (Figure 1). That is, induction of alkaline phosphatase activity in the Xf:phoA bioreporter was detected in culture extracts that contained as little as about 2 nM C14-cis, indicating that another molecular species must account for the induction of the biosensor since at least 500nM C14-cis is needed to induce this biosensor.



**Figure 1.** Alkaline phosphatase activity exhibited by Xf:phoA incubated in PD3 broth containing increasing amounts of DSF containing culture extract of PD three plate cultures of a wild type X.fastidiosa strain. The concentration of C14-cis (XfDSF1) in the extract determined by HPLC is shown by the upper X axis.

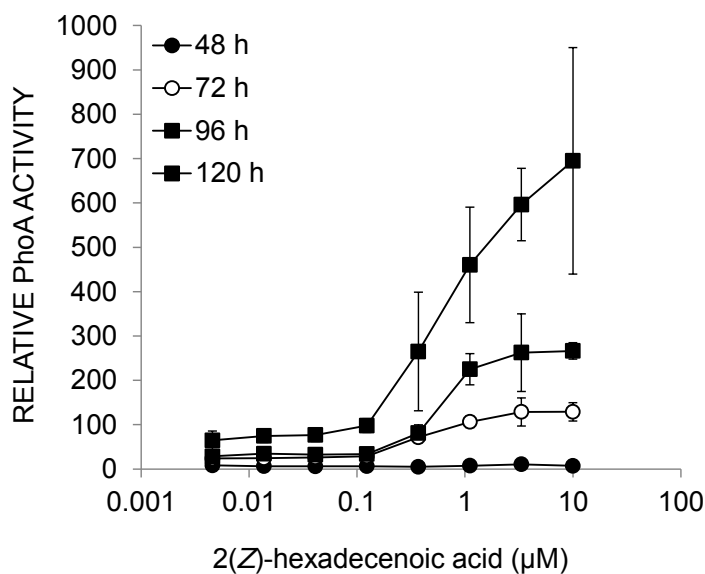
A similarly strong effect of the growth environment of *Xanthomonas* on the species of DSF molecules made has also been observed, and the DSF species originally described in Xcc is uncommon in some culture media. Thus our prior work on identification of DSF species in Xf, which has routinely relied on cultures in PWG may have inadvertently prevented us from observing a highly active DSF species present in other culture media, and presumably in plants. Chemical separation of DSF-containing chemical species from extracts of Xf cultures on the basis of polarity has also revealed the presence of fractions with DSF signaling activity as

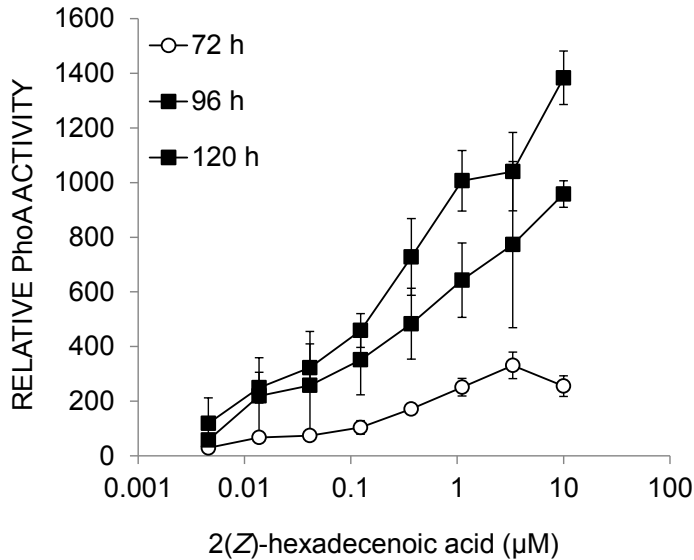
detected by both GCF-gfp and *Xf:phoA* that do not correspond to those containing either C14-cis or C12-cis. Our analysis of the material collected by HPLC from these cultures using electro-spray MS revealed it to be an unsaturated C16 fatty acid which we presume to have a structure similar to that of C14-cis. We therefore chemically synthesized this presumptive derivative which we will call C16-cis or *Xf* DSF2 (Figure 2).



**Figure 2.** Structure of C16-cis (*Xf* DSF2)

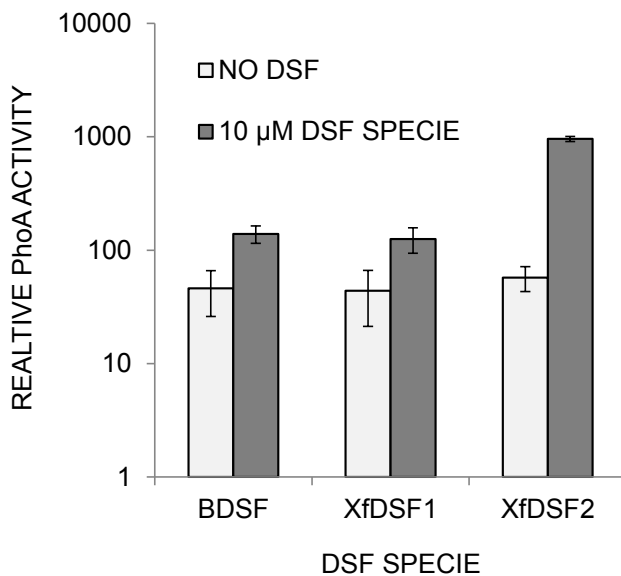
The biological activity of the synthetic C16-cis was tested by the addition of this material at various concentrations to a *rpfF\** mutant strain of *Xf* harboring the *hxfA:phoA* reporter gene fusion and grown in PD3 medium. The *rpfF\** mutant is unable to synthesize DSF due to two mutations introduced into the catalytic site of the DSF synthase, yet this mutant is still able to respond to exogenous DSF. Importantly, this *Xf:phoA* biosensor exhibited very high alkaline phosphatase activity upon the addition of as little as 100 nM C16-cis in all experiments (Figure 3).





**Figure 3.** Alkaline phosphatase activity exhibited by the *Xf:phoA* DSF biosensor when grown in PD3 containing various amounts of C16-cis shown on the abscissa when measured at various times after inoculation in two different experiments.

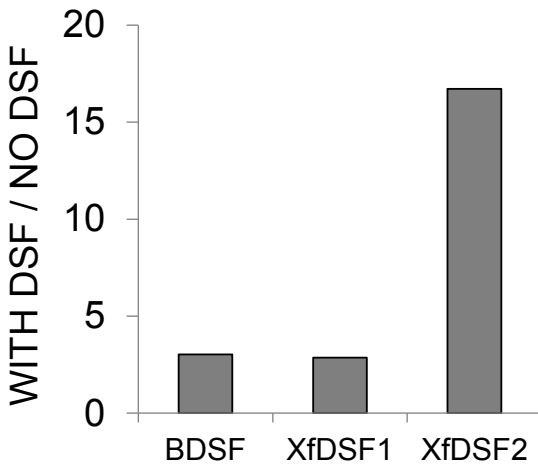
*Xf* appears to be MUCH more responsive to C16-cis than to other DSF species such as C14-cis (*Xf*DSF1) or C12-cis (BDSF). The biological activity of the synthetic C16-cis (*Xf*DSF2) was compared with that of C14-cis (*Xf*DSF1) or C12-cis (BDSF) by the addition of this material at various concentrations to a *rpfF\** mutant strain of *Xf* harboring the *hxfA:phoA* reporter gene fusion and grown in PD3 medium. The alkaline phosphatase activity of this bioreporter was more than 10 times higher in cultures grown with *Xf*DSF2 than with either *Xf*DSF1 or BDSF (Figure 4). This is a very exciting results instead appears that molecules such as *Xf*DSF2 are much more active than our previously discovered *Xf*DSF1.



**Figure 4.** Alkaline phosphatase activity exhibited by the *Xf:phoA* DSF biosensor when grown in PD3 containing 10  $\mu\text{M}$  of BDSF, *Xf*DSF1, or *Xf*DSF2 as shown on the abscissa. Light bars indicate the relatively low amounts of alkaline phosphatase activity exhibited in cultures lacking

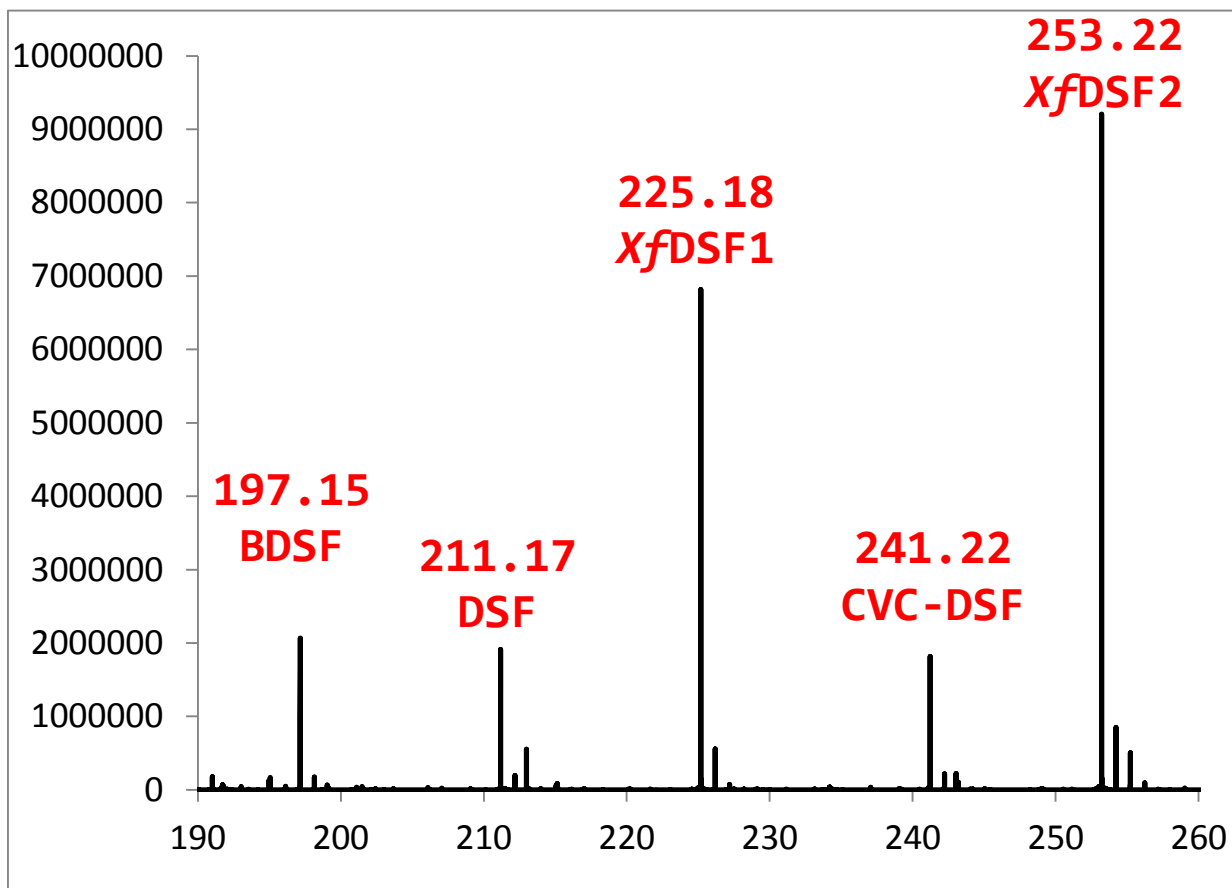
any added DSF species while the dark bars reflect the alkaline phosphatase activity in the presence of the indicated DSF species.

Not only was *Xf* responsive to very low concentrations of added *Xf*DSF2, but more importantly, the expression of DSF-responsive genes such as *hxfA* was as much as 10 fold greater than when cells were exposed to the same concentration of either BDSF or *Xf*DSF1 (Figure 5). Thus not only does *Xf* appear to make different DSF species when grown in different environments, but our finding of extraordinarily high biological activity of *Xf*DSF2 relative to that of *Xf*DSF1 which we had previously studied extensively, suggests that *Xf*DSF2 might have great utility in altering the behavior of *Xf* when applied directly to plants. We are currently preparing plants for topical application of *Xf*DSF2 to evaluate its efficacy in the control of Pierce's disease.



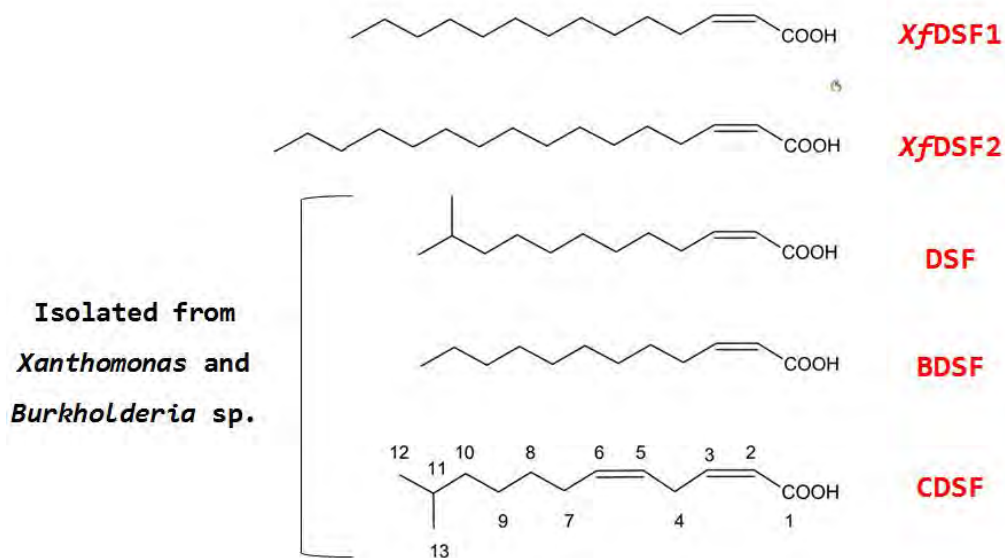
**Figure 5:** Ratio of alkaline phosphatase activity exhibited by an *Xf*-DSF biosensor when grown in PD3 containing 10  $\mu$ M of BDSF, *Xf*DSF1, or *Xf*DSF2s compared to alkaline phosphatase activity in medium without added DSF.

To better understand the chemical species of DSF produced by *Xf* in various conditions in culture and when in infected grape plants, we examined ethyl acetate extracts of chemical species obtained from both culture and *in planta* conditions using electrospray ionization mass spectrometry (ESI-MS). It was possible to identify all of the potential DSF species which we observed in *Xf* as well as those observed by others in other bacterial species such as *Xanthomonas* species based on their highly precise and distinctive ratio of mass to charge (Figure 6). The structure of these different molecular species as shown in Figure 7.



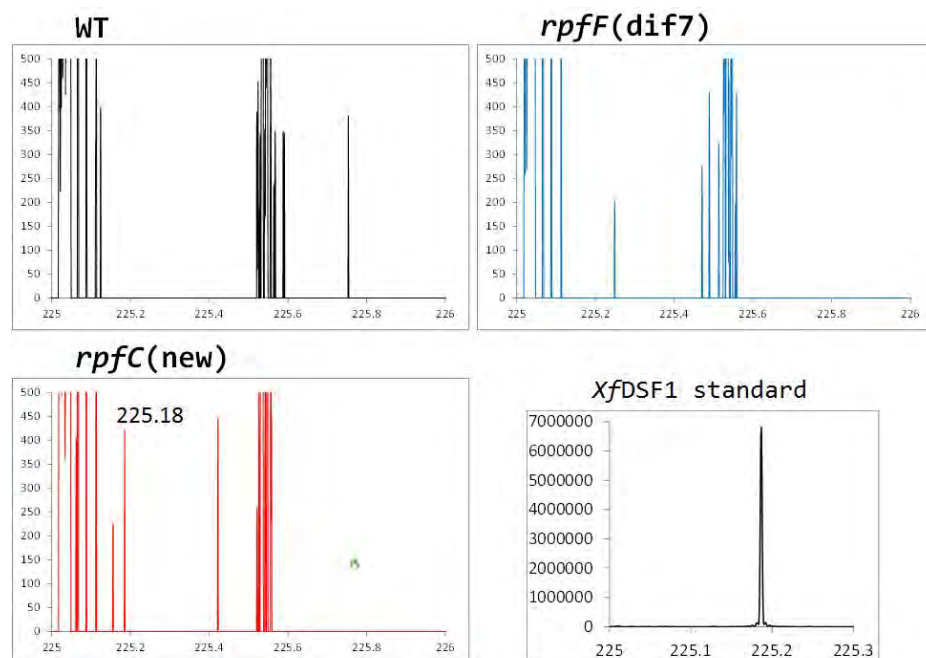
**Figure 6.** M/Z for ionized DSF molecular species determined by ESI-MS

### Known DSFs produced by bacteria

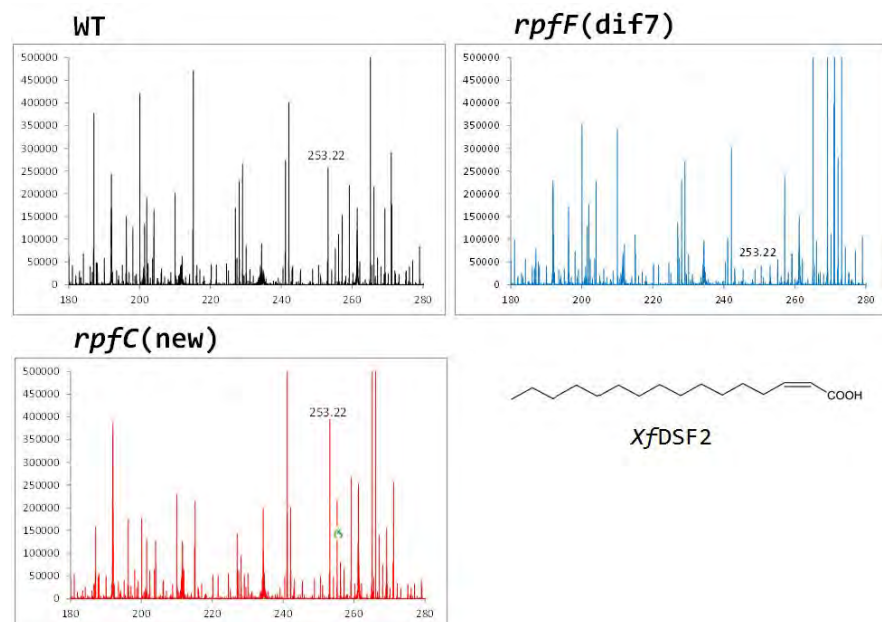


**Figure 7.** chemical species of DSF produced by various bacteria including XfDSF1 and XfDSF2 produced by *Xylella fastidiosa*.

Using ESI-MS we could detect both *Xf*DSF1 and *Xf*DSF2 in culture supernatants of *Xf* grown in PD3 media (Figures 8 and 9). Given that *Xf*DSF2 had never been detected in PWG medium, these results confirm that the production of DSF by *Xf* is rather plastic and somewhat dependent on the growth environment of cells.



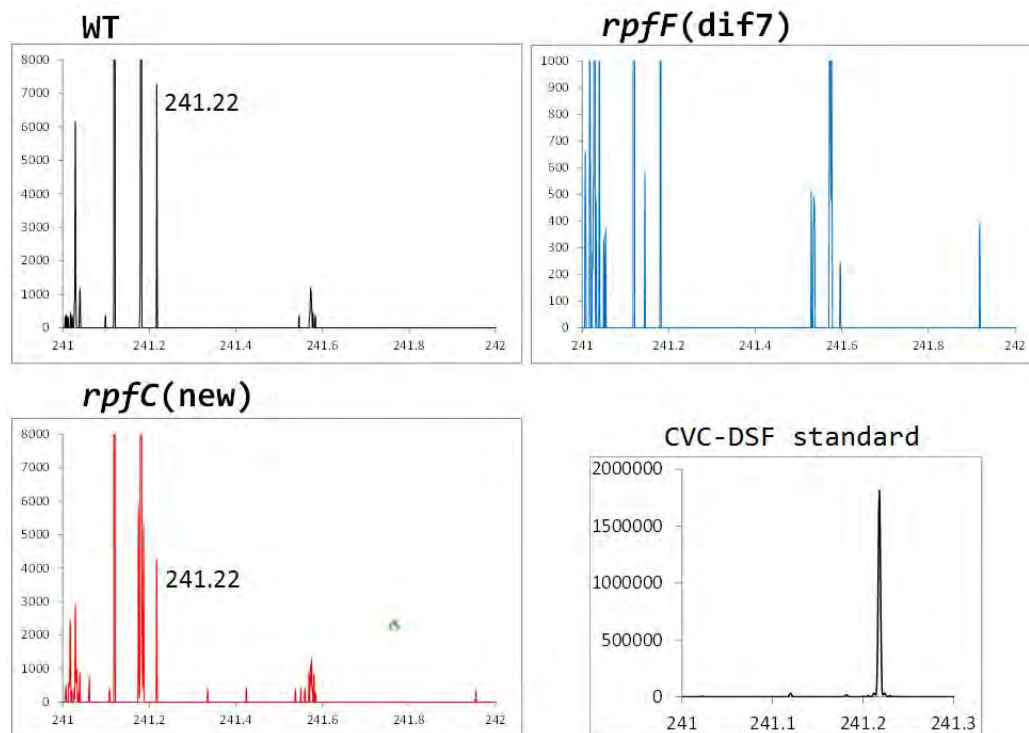
**Figure 8.** Abundance of ions with the M/Z ratios shown on the abscissa or culture extracts of a wild type strain of *Xylella fastidiosa* (upper left panel), of an RpfF mutant unable to produce DSF (upper right panel), of an RpfC mutant which overproduces DSF (lower left panel), as well as a standard of *Xf*DSF1 (lower right panel). Note that *Xf*DSF1 has a M/Z ratio of 225.18.





**Figure 9.** Abundance of ions with the M/Z ratios shown on the abscissa or culture extracts of a wild type strain of *Xylella fastidiosa* (upper left panel), of an RpfF mutant unable to produce DSF (upper right panel), of an RpfC mutant which overproduces DSF (lower left panel). Note that XfDSF2 has a M/Z ratio of 253.22.

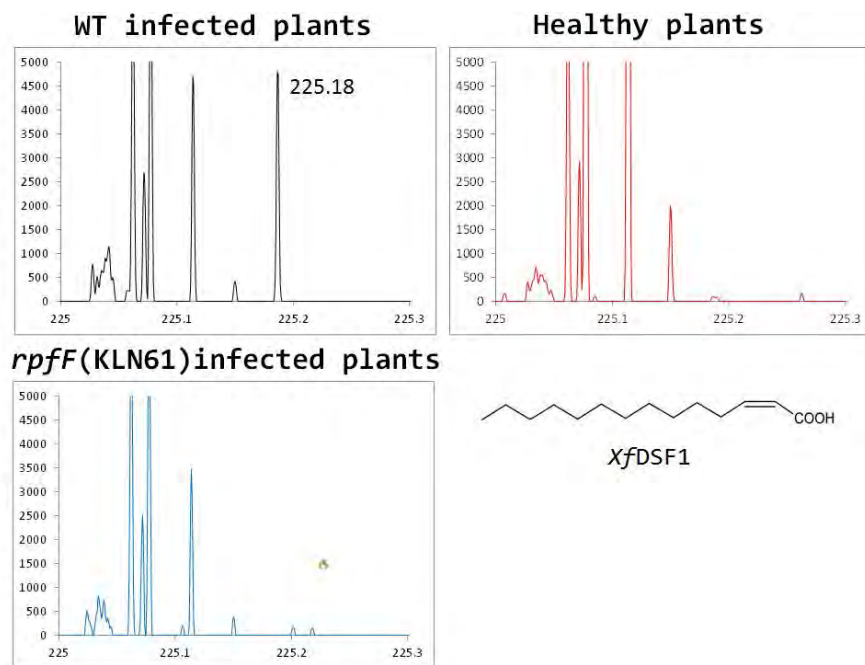
Surprisingly, we also detected small amounts of a molecule which had previously been described as the DSF from CVC strains of *Xylella fastidiosa* (which we term CVCDSF) (Figure 10). While our identification of this molecule is tentative, it is noteworthy that it was found only in wild type and a RpfC mutant of *Xf* but not in an RpfF mutant (Figure 10).



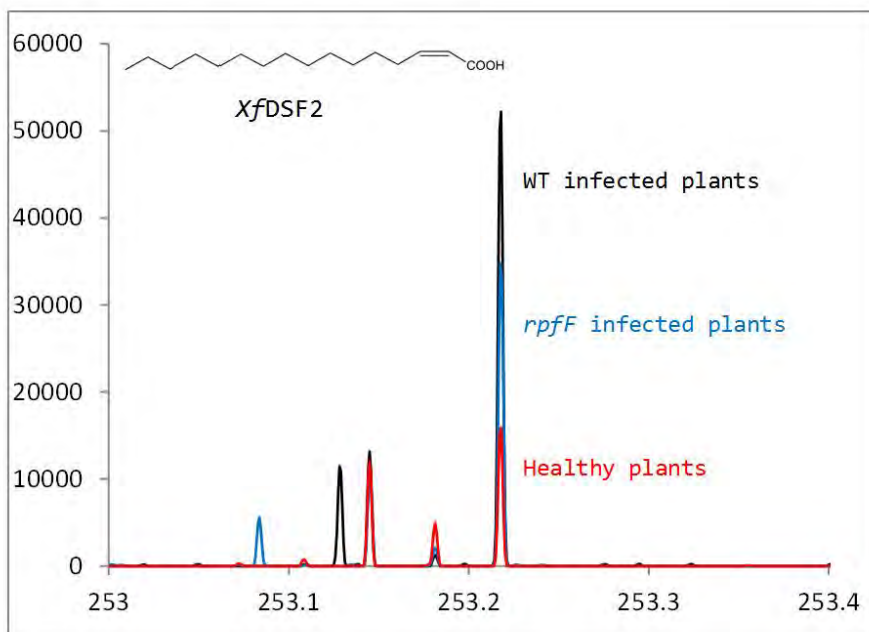
**Figure 10.** Abundance of ions with the M/Z ratios shown on the abscissa or culture extracts of a wild type strain of *Xylella fastidiosa* (upper left panel), of an RpfF mutant unable to produce DSF (upper right panel), of an RpfC mutant which overproduces DSF (lower left panel), as well as a standard of XfDSF1 (lower right panel). Note that CVCDSF1 has a M/Z ratio of 241.22.

To address the DSF species that *Xf* would produce under natural conditions such as within the xylem of grape plants, we extracted DSF-like molecules from plants infected with a wild type strain as well as from an RpfF mutant as well as from healthy plants and subjected these chemicals to identification of separation by ESI-MS. We could easily detect C14-cis (XfDSF1) in plants infected with the wild type strain of *Xf* but this DSF species was not present in either plants infected with the RpfF mutant of *Xf* or in healthy plants (Figure 11). We also could detect elevated concentrations of a molecule presume to be XfDSF2 in plants infected with the wild type except strain (Figure 12). While smaller amounts of this molecule having a M/Z ratio of 253.22 were found in healthy plants and plants infected with the RpfF mutant of *Xf* (Figure 12), it seems likely that these lesser amounts in these negative control plants was associated with 16:1 unsaturated fatty acids found in plant lipids, and not with XfDSF2 itself. Further

characterization of the DSF species found in plants infected with the wild type strain of *Xf* are proceeding to identify other DSF species that are produced under these natural conditions. We find it exciting however that *Xf*DSF2 can be detected in infected plants, suggesting that this highly active DSF species is biologically relevant and may contribute to much of the cell density dependent behavior of *Xf*. For this reason, we are continuing studies to alter the abundance of *Xf*DSF2 in plants by topical applications, injections, and soil drenches.



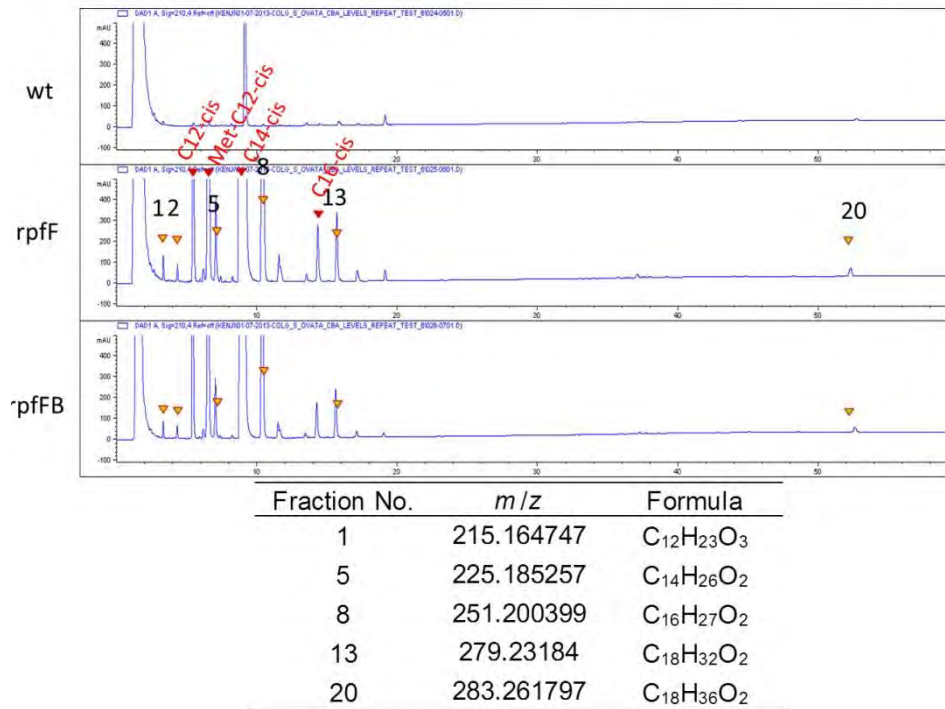
**Figure 11.** Abundance of ions with the M/Z ratios shown on the abscissa in extracts of plants infected with a wild type strain of *Xylella fastidiosa* (upper left panel), infected with a RpfF mutant unable to produce DSF (lower left panel), or from a healthy plant (upper right panel). Note that *Xf*DSF1 has a M/Z ratio of 225.18.



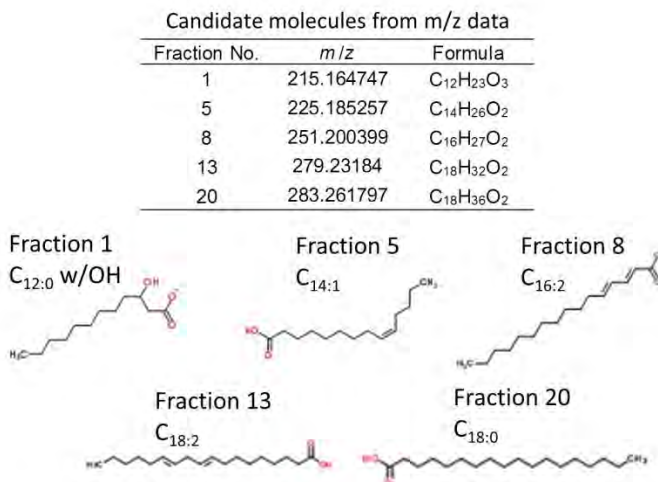
**Figure 12.** Abundance of ions with the M/Z ratios shown on the abscissa made from extracts of plants infected with a wild type strain of *Xylella fastidiosa* (black), from plants infected with an RpfF mutant of XF (blue), or from healthy plants (red). Note the much higher concentration of the ion with a M/Z ratio of 253.22 corresponding to XfDSF2.

To better understand the variety of different DSF species that might be produced by Xf, we chemically fractionated cultures of *Erwinia herbicola* harboring RpfF, the DSF synthase from Xf. *Erwinia herbicola* was chosen as a surrogate host for the DSF synthase because much larger numbers of cells of the species can be produced than that of Xf itself. Ethyl acetate extracts of culture supernatants were separated using HPLC. Several chemical entities were identified in extracts of *Erwinia herbicola* containing RpfF or containing both RpfF and RpfB from Xf but not in control cultures of *Erwinia herbicola* lacking these genes from Xf (figure 13). To obtain putative chemical structures for these chemical species that were present only in the presence of the DSF synthase we physically collected fractions of the separated material and subjected to ESI-MS to obtain high-resolution M/Z ratios for these materials (Figure 13). While a precise molecular formula for these molecules could be obtained this process, it is not possible to obtain an unambiguous structure, given that there are often several solutions that would match the particular formula weight. However, given the constraints that the DSF species are very likely to be fatty acids, we have proposed tentative structures for these novel DSF-like species (Figure 14). The location of the double bond in these molecules however cannot be placed unambiguously based on their mass alone, and further work is underway to obtain sufficient amounts of these molecules to enable them to be subjected to NMR and other techniques that would localize the unsaturation.

ESI-MS analyses on unknown peaks derived from transgenic *Erwinia* strains



**Figure 13.** Optical absorption patterns (lambda = 220 nm) for chemical species separated on a methanol: water gradient by HPLC. Note peaks that are found in extracts of *Erwinia herbicola* harboring RpfF from XF (center panel), or harboring both RpfF and RpfB (bottom panel) but which are not found in extracts of *Erwinia herbicola* lacking these heterologous genes from XF (top panel). The M/Z ratios of the numbered peaks and the molecular formula for molecules in each peak are noted below.

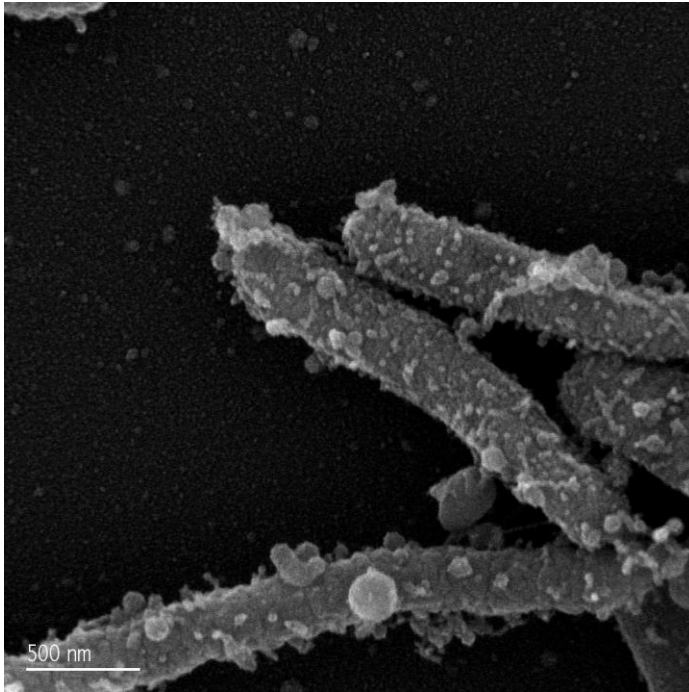


These candidates probably do NOT have any aromatic moieties in their structure, respectively because of NO absorbance >250nm.

**Figure 14.** Tentative structures for novel DSF species produced in *Erwinia herbicola* harboring RpfF from *Xylella fastidiosa*.

## Objective 2: Role of extracellular vesicles

Our continuing work reveals that *Xf* is a very prolific producer of extracellular vesicles. For a large numbers of vesicles (>400/cell) can be associated both with the surface of the bacterial cell, as well as a high portion that are shed by the cells to the extracellular environment (Figure 15). The vesicles are generally quite small, ranging in size from about 0.01 to 0.1  $\mu\text{m}$  in diameter.

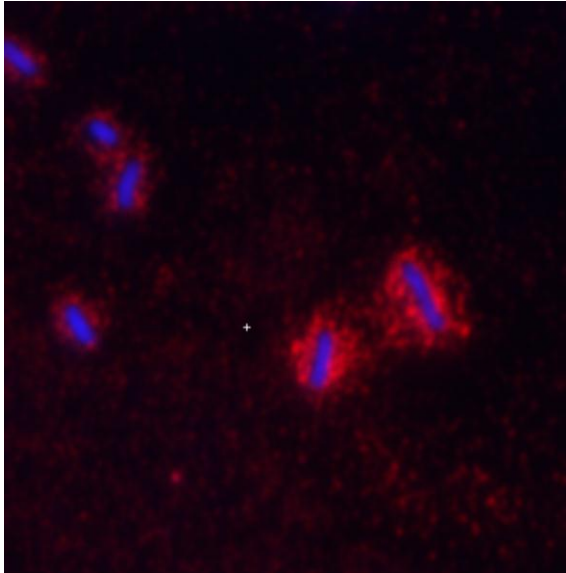


**Figure 15.** Visualization of membranous vesicles forming on the surface of cells of a wild type strain of *Xylella fastidiosa*. Not shown to the method of preparation are those vesicles that have been shed from cells.

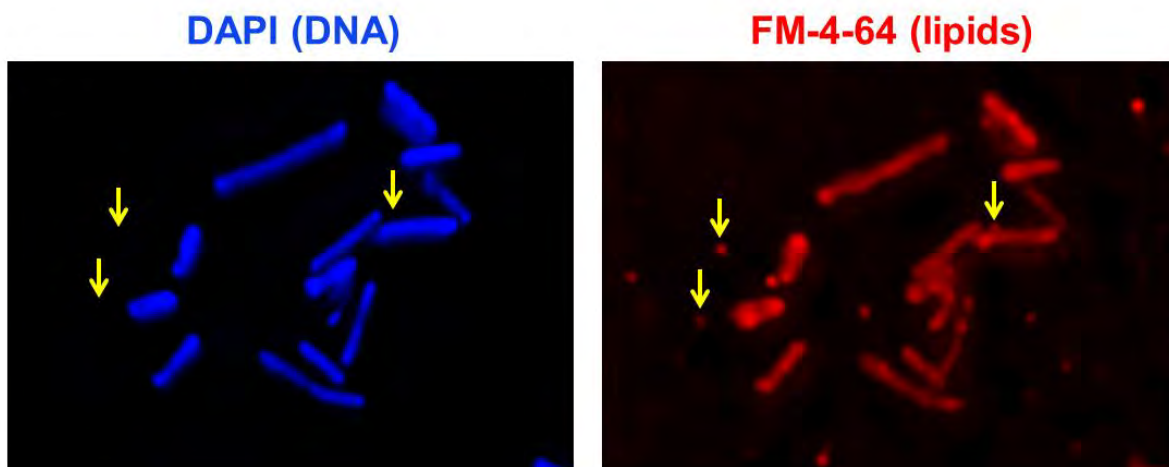
Our ability to quantify membrane vesicles and to determine those factors which control their production and release from cells has been facilitated by our finding that a major outer membrane adhesins XadA is a significant component in these membranous vesicles. Since we have obtained antibodies specific to XadA, it is possible to visualize membrane vesicles fusing light microscopy (Figure 16). Using anti-XadA antibodies with a red fluorescent tag it is clear that the surface of *Xf* cells harbors a large constellation of membranous vesicles which surround the cell has somewhat of a “cloud”. In addition to those vesicles which are relatively closely associated with the cell (although apparently not physically linked), are vesicles that can be found at further distances away from cells (Figure 16). These distantly located vesicles are clearly not simply “mini-cells” of *Xf* since they do not contain DNA as determined with a DNA-binding dyes such as DAPI (Figure 17). Quantification of vesicles by both deconvolution fluorescence microscopy as well as flow cytometry reveals that the higher number of vesicles (normalized for the number of *Xf* cells) are present in RpfF mutants, suggesting strongly that DSF accumulation suppresses the release of such vesicles.

To utilize the outer membrane protein XadA as a molecular marker for the presence of extracellular vesicles it was necessary to demonstrate that it is not secreted as a free protein into the extracellular environment, and is always found associated with membranous vesicles.

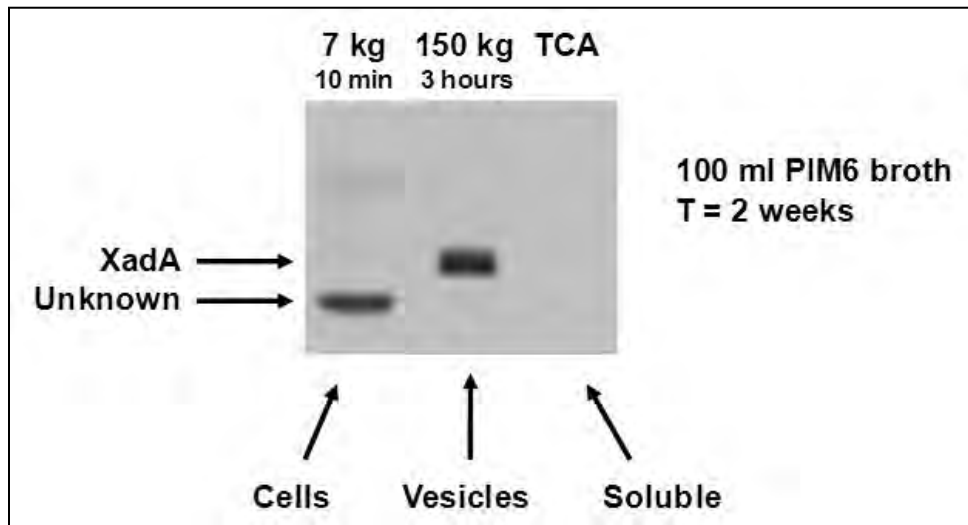
To demonstrate this we isolated total proteins from vigorously washed cells of an RpfF mutant of *Xf*, from a pallet of extracellular material could be recovered after high-speed centrifugation (150,000 x g), as well as free protein that was not pelleted after high-speed centrifugation. While small amounts of XadA were associated with the attack cells of *Xf*, large amounts of XadA were found in membranous vesicles that could be obtained after high-speed centrifugation, and importantly, no XadA was free and solution of cultures of *Xf* (Figure 18). These results clearly show that XadA is abundantly released in the form of membranous vesicles from cells of *Xf*, and since it is always associated with membranes, it makes an excellent marker for membranous vesicles.



**Figure 16.** Visualization of membranous vesicles produced by cells of *Xylella fastidiosa* whose DNA is stained with DAPI and appearing blue in this image. Vesicles are read in this image due to their binding to fluorescently labeled anti-XadA antibodies.

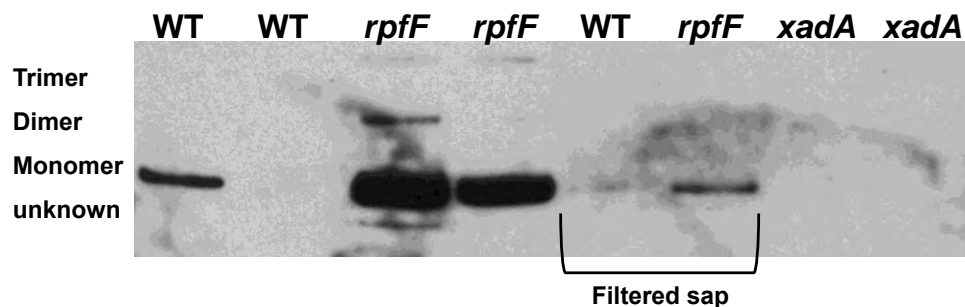


**Figure 17.** Visualization of cells of *Xylella fastidiosa* stained with the DNA-binding stain DAPI (left panel) as well as with the lipid binding stain FM-4-64 (right panel). Note the location of small circular red objects indicating presence of membranous material distal to the location of adjacent *Xylella fastidiosa* cells.



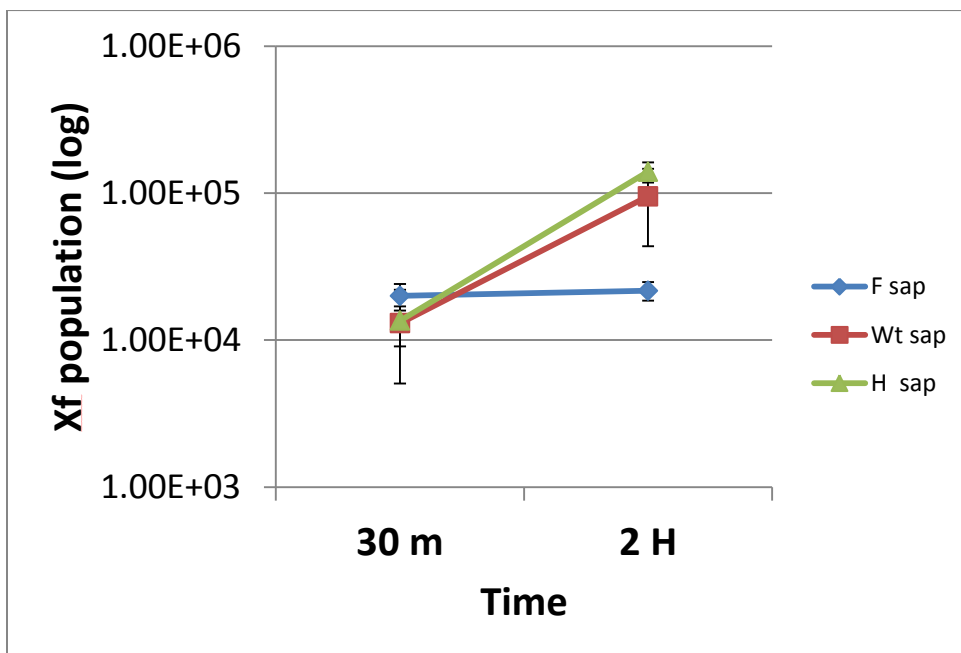
**Figure 18.** Quantification of the outer membrane protein XadA by Western blot analysis using anti-XadA antibodies from washed cells of *Xylella fastidiosa* (left lane) as well as from material recovered from high-speed centrifugation of culture supernatants (center lane), or in proteins precipitated from culture supernatants after high-speed centrifugation by TCA (right lane).

Strong evidence has been obtained that *Xf* releases factors into the xylem fluids of plants that it is colonizing that prevents their adherence to surfaces. XadA that was not associated with intact cells of *Xf* (and hence associated with membranous vesicles), was readily detected in the xylem sap of plants infected with both the wild type strain as well as an RpfF mutant of *Xf* (Figure 19). Substantially more vesicles were apparently present in the sample plants infected with the RpfF mutant, consistent with our finding that such a mutant produces more membranous vesicles in culture.



**Figure 19.** Abundance of XadA, indicative of the presence of membranous vesicles, as determined by Western blot analysis in xylem sap of Thompson seedless grapes infected with either a wild type or an RpfF mutant of *Xf*. Total XadA in sap which had not been filtered to remove intact cells are shown in the leftmost four lanes, while that in sap that had been filtered to remove intact cells (retaining only membranous vesicles) are shown in lanes 5 and 6. Note the high abundance of XadA in filtered xylem sap, especially from plants infected with the RpfF mutant.

Xylem sap containing membranous vesicles was shown to reduce the adherence of various bacteria to surfaces. Xylem fluid was collected by pressure bomb from healthy Thompson seedless grape, as well as from plants infected with a wild type strain of *Xf* or with an RpfF mutant of *Xf*. To test the differential adherence of wild type cells of *Xf* to surfaces such as insects in the presence of these different sample collections, we immersed small sections of hind wings of glassy winged sharpshooter in each of these xylem sap samples to which we also added a wild type strain of *Xf* (final concentration  $10^8$  cells/ml). Cell suspensions in these various sap samples were allowed to incubate with the wings for various times, the wing segments were then removed from the cell suspension in the sap, washed briefly to remove non-adhering cells, and the wing then macerated to release the adhering cells which were then enumerated by dilution plating. While relatively small numbers of cells had attached to wings suspended either of the sap suspensions after a short incubation period (30 min.), a much larger number of cells had attached after a two-hour incubation period. Importantly, many more cells that had been suspended in samples from healthy plants attached than from cells suspended in sap from plants infected with the wild type *Xf* strain, particularly from sap infected with an RpfF mutant of *Xf* (Figure 20). Over 20-fold more bacterial cells were attached when suspended in sap from the healthy plant compared to that of sap from the plant infected with the RpfF mutant of *Xf* (Figure 20). It is thus clear that the xylem sap environment of plants infected with an RpfF mutant of *Xf* is much less conducive to the adherence of *Xf* to surfaces than that of healthy plants. In fact, the cells sap environment of plants infected with the wild type strain of *Xf* is also somewhat less conducive to adherence.



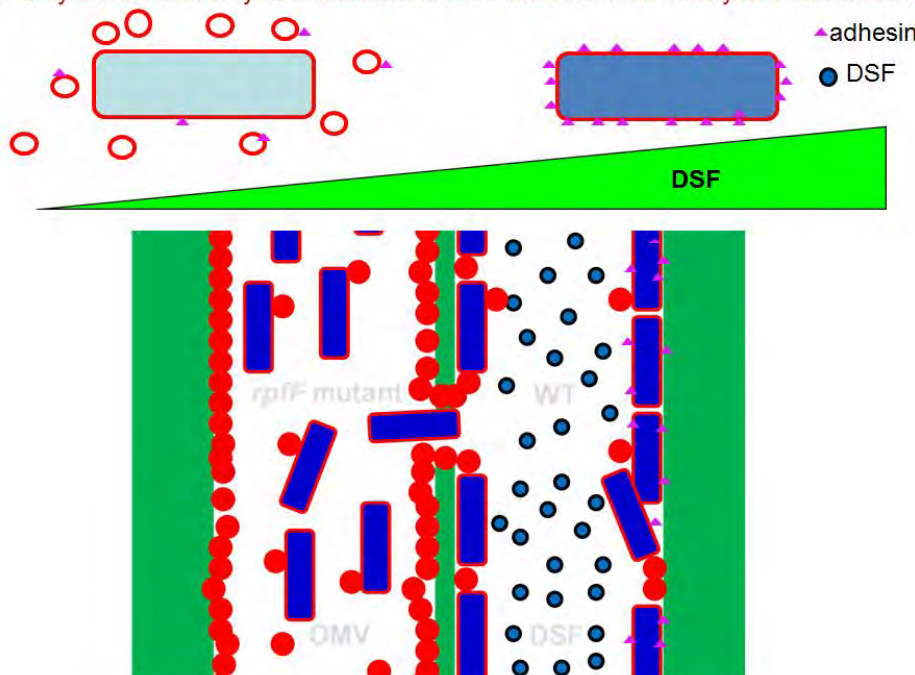
**Figure 20.** The number of cells of a wild type strain of *Xylella fastidiosa* that had adhered to wings of the glassy winged sharpshooter suspended in xylem sap from plants infected with an RpfF mutant (blue), or a wild type strain of *Xylella fastidiosa* (red), or from healthy plants (green) after incubation for either 30 min. or two hours.

Strong circumstantial evidence suggests that the release of extracellular membranous vesicles by the RpfF mutant is responsible for the suppression of adherence of *Xf* to surfaces. Since the RpfF mutant of *Xf* does not accumulate DSF, which in turn suppresses the release of



extracellular vesicles, a higher concentration of extracellular vesicles would be expected in plants infected with the RpfF mutant. At least some extracellular vesicles would also be expected in the sap of plants infected with the wild type strain as well. A higher concentration of extracellular vesicles, as estimated by the abundance of XadA (which we can use as a marker protein for these membranous vesicles), is found in plants infected with the RpfF mutant of *Xf* compared to that of the wild type strain (Figure 21). These results further support our model of a “Teflon mechanism” of virulence of *Xf* whereby it releases adhesive vesicles, especially one found at relatively low cell densities where DSF would not have accumulated. These vesicles would be expected to attached to surfaces such as that of the walls of the xylem vessels. By so attaching, these vesicles would prevent the attachment of *Xf* cells themselves to such surfaces. Only upon reaching relatively high cell concentrations in a particular vessel would DSF concentrations increase to a level that would suppressed the release of the membranous vesicles, thereby retaining adhesive molecules on the surface of *Xf* cells themselves, thus allowing the bacterial cells themselves to attached to surfaces, such as that of insect vectors (Figure 21).

Two-way transition to vary adhesiveness of *X. fastidiosa* modulated by DSF accumulation



**Figure 17.** “Teflon Model” of DSF regulated virulence involving DSF-mediated control of shedding of extracellular vesicles in *Xylella fastidiosa*.

#### Reports and presentations:

1. Beaulieu, E., M. Ionescu, S. Chatterjee, K. Yokota, D. Trauner, and S.E. Lindow. 2012. Characterization of a diffusible signaling factor from *Xylella fastidiosa*. *mBio* 4(1): doi:10.1128/mBio.00539-12
2. Wang, N., Li, J.-L., and Lindow, S.E. 2012. RpfF-dependent regulon of *Xylella fastidiosa*. *Phytopathology* 102:1045-1053.

3. Almeida, R.P.P., Killiny, N., Newman, K.L., Chatterjee, S., Ionescu, M., and Lindow, S.E. 2012. Contribution of *rpfB* to cell-to-cell signal synthesis, virulence, and vector transmission of *Xylella fastidiosa*. MPMI 25:453-462.

Presentation at the University of Florida, Department of plant pathology entitled “the complex lifestyles of *Xylella fastidiosa* coordinated by cell- cell signaling: achieving disease control by pathogen confusion” presented on February 14, 2013.

Presentation at the headquarters of the American society for microbiology for “microbes after hours” webcast entitled “the complex lifestyles of *Xylella fastidiosa* coordinated by cell-cell signaling: achieving disease control by pathogen confusion” presented on January 28, 2013.

#### **Research relevance statement:**

We are very excited about results to date that show that several means of elevating DSF levels in plants have provided disease control via a strategy of “pathogen confusion”. Given the limitations in standard methods of disease control, we are optimistic that DSF interference represents a promising strategy for PD control. Control of Pierce’s disease by direct application of DSF is a very attractive disease control strategy since it could be quickly implemented and would utilize commonly used agricultural equipment and methods and would not require the use of transgenic technologies. Our earlier work had shown that C14-cis, a component of *Xf* DSF, conferred some reduction of disease after topical application, but less than might have been expected compared to application of crude DSF-containing extracts of *Xf*. Our recent studies using biosensors more responsive to the DSF molecules made by *Xf* reveal that at least 2 additional molecules related to C14-cis are biologically active in *Xf*, at least one of which may be much more active than C14-cis. We have now found several additional DSF species that are synthesizing them in testing their biological activity. We will determine which of these molecules are most biologically active, whether they all have similar effects on gene expression in *Xf*, and which are most abundant within plants infected with *Xf*. Our new sensitive biosensors will be used to document the absorption and translocation of these molecules by grape after application in various ways. This should enable us to greatly increase disease control by direct application of the most appropriate molecule. (Obj. 1). Our recent results indicate that *Xf* changes the nature of the xylem sap during its colonization of plants to restrict its adhesion to plants via its shedding of outer membrane vesicles and that DSF accumulation restricts such shedding. Such vesicles represent important virulence factors for the pathogen, presumably by preventing *Xf* adhesion to pit membranes because the vesicles themselves will adhere to and coat plant targets. The presence of such vesicles would also be easily assessed indicators of the cell-cell signaling status of *Xf* in the plant. Our work to date has shown that these vesicles are both very abundant and have very strong effects on the behavior of *Xf* within plants. We thus will continue to further explore the role of vesicles as virulence factors and as tools to determine the effects of strategies of pathogen confusion by elevating DSF in plants.

#### **Layperson summary:**

*X. fastidiosa* produces an unsaturated fatty acid signal molecule called DSF. Accumulation of DSF in *Xf* cells, which presumably normally occurs as cells become numerous within xylem vessels, causes a change in many genes in the pathogen, but the overall effect is to suppress its virulence in plants by increasing its adhesiveness to plant surfaces and also suppressing the production of enzymes and genes needed for active movement through the plant. We have investigated DSF-mediated cell-cell signaling in *X.fastidiosa* with the aim of developing cell-cell signaling disruption (pathogen confusion) as a means of controlling Pierce’s disease. Elevating

DSF levels in plants or artificially reduces its movement in the plant. In this study we have investigated the variety of different fatty acid molecules that can serve as cell-cell signaling agents in *Xf*. Several new DSF species have been found including a 16 carbon unsaturated fatty acid appears to be far more active than the 14 carbon unsaturated fatty acid that we have previously investigated. The release of extracellular membranous vesicles by *Xf* is responsible for the suppression of its adherence to surfaces. These vesicles attach to surfaces such as that of the walls of the xylem vessels. By so attaching, these vesicles prevent the attachment of *Xf* cells themselves to such surfaces. Only upon reaching relatively high cell concentrations in a particular vessel would DSF concentrations increase to a level that would suppress the release of the membranous vesicles, thereby retaining adhesive molecules on the surface of *Xf* cells themselves, thus allowing the bacterial cells themselves to attach to surfaces, such as that of insect vectors.

**Status of funds:**

While this project was approved as of July 1, 2012, because of delays in the sponsored projects office at the University of California, Berkeley, this contract was not formally completed until February, 2013. Since some funding remained in closely related project 10-0276 for which a no cost extension was granted through June 30, 2013, much of the work presented here was conducted with carry-forward funding that project. For that reason, as of March, 2013, much of the originally budgeted funding remains, and is anticipated that a no cost extension will be requested at the end of the currently scheduled budget period.

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**Summary and status of intellectual property associated with the project:**

A patent application (12/422,825) entitled "biological control of pathogenicity of microbes that use alpha, beta unsaturated fatty acid signal molecules" had been submitted March 13, 2009. While many of the claims had been rejected earlier, the University of California patent office has filed on March 13, 2012 a motion requesting reconsideration of the application with clarification of, and justification for, claims related to the production of transgenic plants transformed with the *rpfF* gene from *Xylella fastidiosa*. This petition was approved in June, 2012 and patent US 8,247,648 B2 was issued on August 21, 2012.