# 1 Comparison of real-time PCR and droplet digital PCR for the

# 2 detection of *Xylella fastidiosa* in plants

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- 13 **Declarations of interest**: The authors have no competing interests to declare.

### 14 Abstract

15 *Xylella fastidiosa* (*Xf*) is a quarantine plant pathogen bacterium originating from the Americas 16 and that has emerged in Europe in 2013. Xf can be detected directly on plant macerate using molecular methods such as real-time PCR, which is a sensitive technique. However, some 17 18 plants may contain components that can act as PCR reaction inhibitors, which can lead to false 19 negative results or an underestimation of the bacterial concentration present in the analyzed 20 plant sample. Droplet digital PCR (ddPCR) is an innovative tool based on the partitioning of 21 the PCR reagents and the DNA sample into thousands of droplets, allowing the quantification 22 of the absolute number of target DNA molecules present in a reaction mixture, or an increase 23 of the detection sensitivity. In this study, a real-time PCR protocol, already used for Xf detection 24 in the framework of official surveys in the European Union, was transferred and optimized for Xf detection using ddPCR. This new assay was evaluated and compared to the initial real-time 25 26 PCR on five plant matrices artificially inoculated and on naturally infected plants. In our conditions, this new ddPCR enabled the detection of Xf on all artificially inoculated plant 27 28 macerates with a similar limit of detection, or a slight benefit for Quercus ilex. Moreover, 29 ddPCR improved diagnostic sensitivity as it enabled detection of Xf in samples of Polygala 30 myrtifolia or Q. ilex that were categorized as negative or close to the limit of detection using 31 the real-time PCR. Here, we report for the first time a ddPCR assay for the detection of the 32 bacterium Xf.

33 Keywords: *Xylella fastidiosa*, ddPCR, molecular diagnostics, quarantine plant pathogenic
34 bacteria.

### 35 **1** Introduction

*Xylella fastidiosa* (*Xf*) is a plant pathogenic bacterium known worldwide. Located in the xylem 36 vessels of plants, its natural way of transmission is sap-feeding insect vectors (Almeida and 37 38 Nunney, 2015). To date, five different subspecies (subsp.) have been described: subsp. 39 fastidiosa, subsp. morus, subsp. multiplex, subsp. pauca, and subsp. sandyi (Nunney et al., 40 2014; Schaad et al., 2004; Schuenzel et al., 2005). In Europe, Xf was first detected in Italy, in 41 the Apulia area in 2013, where the subsp. pauca was identified on olive trees (Saponari et al., 42 2013). Then, in France in 2015 (Corsica and in French Riviera region), the subsp. *multiplex* was 43 reported firstly on *Polygala myrtifolia* and then on a large range of ornamental or wild plants 44 (Denancé et al., 2017). More recently in 2016, the subsp. fastidiosa, multiplex and pauca were

identified in the Balearic Islands (Spain), on olive trees, grapevines and sweet cherry trees; and 45 in 2017, the presence of the subsp. *multiplex* was also identified in Spain near Alicante, on 46 47 almond trees (Landa, 2017). Currently, 563 plant species distributed in 82 botanical families are reported to be hosts of Xf, and this list includes plants of major socio-economic interest such 48 49 as olive trees, citrus or grapevine (EFSA, 2018). Since 2017, Xf has been classified in Annex 50 I/A2 of Council Directive 2000/29/EC revised in 2017, and in the A2 list of the EPPO as a 51 quarantine pathogen present on the EU territory and requiring mandatory control (C/2017/4883, 52 2017; EPPO, 2018a).

53 As isolation and cultivation of Xf is fastidious, detection and identification tests are applied 54 directly on plant extracts (Denancé et al., 2017). Nowadays, different molecular tools targeting specific DNA regions are available to detect the bacterium at the species level or to specifically 55 56 detect one of the subspecies. Conventional PCRs such as Minsavage et al. (2014) have been 57 developed, but they are less sensitive than Real-Time PCR (Baldi and La Porta, 2017). Among 58 the real-time PCR techniques developed, the method designed by Harper et al. (2010) was 59 identified as one of the most suitable methods for Xf detection. It allow to detect all the Xf 60 subspecies, its limit of detection determined on different plant species is low, it is sensitive, and no cross-reactions with other bacterial species have been reported (Francis et al., 2006; Harper 61 62 et al., 2010; Li et al., 2013; Modesti et al., 2017; Ouyang et al., 2013).

63 Even though real-time PCR is very sensitive in most cases, low bacterial contamination levels 64 of plants and the presence of PCR inhibitors can lead to false negative results, and the underestimation of positive samples for some plant species (Modesti et al., 2017; Schrader et 65 66 al., 2012). These PCR inhibitors include polyphenols, polysaccharides, pectin and xylan 67 (Harper et al., 2010; Schrader et al., 2012; Wei et al., 2008). Studies have revealed that the 68 improvement of DNA extraction methods, or the addition of bovine serum albumin (BSA) 69 during the PCR assay, may reduce the impact of PCR inhibitors (Harper et al., 2010; Schrader 70 et al., 2012). For some plants, such as Nerium oleander, Prunus dulcis and Vitis vinifera, a 71 tenfold dilution prevented PCR inhibition and led to successful detection of Xf (Francis et al., 72 2006; Minsavage et al., 1994; Modesti et al., 2017). However, DNA dilution cannot be applied 73 to every sample, due to low Xf concentrations in some infected plants. It can therefore be rather 74 difficult to find a universal method, because of the wide range of Xf host plants. Moreover, even 75 though real-time PCR produces quantitative data when using a calibration curve, the results are 76 often only interpreted qualitatively for Xf detection (Cruaud et al., 2018; Modesti et al., 2017).

77 Digital PCR was set up in 1999 by Vogelstein and Kinzler and named later by Morley in 2014. 78 By compartmentalizing the PCR reaction into thousands of droplets, ddPCR offers the promises 79 of absolute quantification without the need for calibration (Hindson et al., 2011; Huggett et al., 2013; Morley, 2014; Voegel and Nelson, 2018; Vogelstein and Kinzler, 1999). At first, ddPCR 80 81 was designed to identify rare mutations in a small number of cells (Vogelstein and Kinzler, 82 1999). Already used for medical purposes (Bharuthram et al., 2014; Cao et al., 2015; Hindson 83 et al., 2011; Nixon et al., 2014; Ramírez et al., 2019), this method was transferred as a detection 84 and quantification tool to other fields, such as environmental sciences (Doi et al., 2015; Hoshino 85 and Inagaki, 2012), food safety control (Bian et al., 2015; Wang et al., 2018), GMO detection (Košir et al., 2017; Morisset et al., 2013) or the agronomic field (Dreo et al., 2014; Maheshwari 86 87 et al., 2017; Rački et al., 2014; Voegel and Nelson, 2018; Zhao et al., 2016). The transfer of 88 real-time PCR assays to ddPCR assays has already provided successful results for the detection 89 and the quantification of plant pathogenic bacteria (Dreo et al., 2014; Maheshwari et al., 2017; 90 Zhao et al., 2016). For example, ddPCR was more efficient than real-time PCR to detect low 91 concentrations of *Ralstonia solanacearum* in potatoes (Dreo et al., 2014). It also increased the 92 detection threshold of other pathogens such as Xanthomonas citri subsp. citri in citrus, Pepper 93 mild mottle virus in plants, soil and water, or of GMOs in maize seed powder (Morisset et al., 94 2013; Rački et al., 2014; Zhao et al., 2016). ddPCR was reported to be up to 1,000 fold more 95 sensitive than conventional PCR developed to detect adenovirus in live attenuated vaccines 96 (Dong et al., 2018). It allows the detection and quantification of pathogen abundance, such as 97 Agrobacterium vitis in grapevines, for which previous methods lacked sensitivity (Voegel and 98 Nelson, 2018). An additional advantage of ddPCR is its tolerance to PCR inhibitors present in 99 plants, soil, water or food (Cao et al., 2015; Maheshwari et al., 2017; Morisset et al., 2013; 100 Rački et al., 2014; Zhao et al., 2016). ddPCR presents many advantages that could make it an 101 alternative for Xf detection.

102 The aim of this study was to transfer the real-time PCR developed by Harper et al. (2010) into 103 a ddPCR assay, in order to improve the detection of Xf at low concentrations in plant matrices rich in PCR inhibitors. ddPCR was compared to real-time PCR using five artificially inoculated 104 105 plant matrices and naturally infected plants sampled in France. The plant species used as 106 matrices were selected for their level of PCR inhibitors reported by the Plant Health Laboratory 107 (PHL) of the French Agency for Food, Environmental and Occupational Health & Safety 108 (ANSES) following the analysis of thousands of different plant samples collected since 2015 109 in the context of the national Xf survey in France (personal communication, Bruno Legendre).

Polygala myrtifolia was selected as a matrix containing a low concentration of inhibitors, Lavandula angustifolia, Olea europaea, Quercus ilex and Rosmarinus officinalis were selected as matrices containing high concentrations of inhibitors. Experimental assays were set up following the standard PM7/98 and the digital MIQE guidelines. The following performance criteria were evaluated: analytical sensitivity, repeatability, and diagnostic specificity (EPPO, 2014; Huggett *et al.*, 2013).

116 2 Materials and methods

### 117 2.1 Bacterial strains

118 Bacterial strains Xf subsp. multiplex CFBP 8416, isolated in France (Corsica) in 2015 from 119 symptomatic P. myrtifolia (Denancé et al., 2017) and Xf subsp. fastidiosa CFBP 7970, isolated 120 in the United States (Florida) from Vitis vinifera were cultivated on modified PWG media at 121  $28^{\circ}$ C for two weeks (EPPO, 2018b). Bacterial suspensions of pure culture of Xf were prepared 122 in sterile demineralized water and suspensions titer was estimated by immunofluorescence (IF) (EPPO, 2009, 2018b). The antiserum, used to count Xf, was especially produced in 123 collaboration with the UR1268 BIA - Team Allergy of the French National Institute for 124 125 Agricultural Research (INRA) of Angers-Nantes. It resulted from the inoculation of rabbits 126 with nine strains of Xf chosen to be as diverse as possible in terms of subspecies, geographical 127 location, and host plant species. The initial concentration of the CFBP 8416 bacterial suspension was estimated by IF at 1.84x10<sup>9</sup> bacteria/mL (b/mL). This suspension was used to 128 129 spike all the artificially inoculated samples in this study. The bacterial suspension of the strain CFBP 7970 was calibrated at  $1 \times 10^7$  b/mL and used as a positive control for the PCR reactions. 130

### 131 2.2 Plant materials

Healthy plant materials used for spiking assays were collected in 2018. *L. angustifolia*, *O. europaea*, *Q. ilex*, *R. officinalis* were sourced from Maine-et-Loire, a French department known to be *Xf* free. *P. myrtifolia* was produced in a nursery in Brittany (*Xf* free) and had a European phytosanitary passport certifying its healthy status. Moreover, no symptoms were recorded on these five plants. In this study, the healthy status of the five matrices was first checked using the real-time PCR assay Harper *et al.* (2010).

Naturally infected samples of *Calicotome* sp. (one sample), *L. angustifolia* (four samples), *P. myrtifolia* (13 samples), and *Q. ilex* (4 samples) were collected in the context of the national

survey, between 2016 and 2018 in Corsica and in the PACA region of France. These 22 samples
were already found to have a positive status or to be at the limit of detection by the PHL, using
the real-time PCR developed by Harper *et al.* (2010).

## 143 2.3 Plant spiking

The artificially inoculated plant samples were prepared by mixing 1 g of healthy plant petiole in 4.5 mL of sterile distilled water and spiked with 0.5 mL of a known concentration of bacterial suspension. Each matrix was spiked in order to reach a range dilution of  $1x10^5$  b/mL;  $5x10^4$ b/mL;  $1x10^4$  b/mL;  $5x10^3$  b/mL;  $1x10^3$  b/mL;  $5x10^2$  b/mL; and  $1x10^2$  b/mL. The negative template control (NTC) was obtained by mixing 1 g of healthy plant petiole with 5 mL of sterile distilled water.

### 150 2.4 DNA extraction

The bacterial strain suspension used as a positive control for all the PCRs was inactivated by thermal lysis. A volume of 1 mL of bacterial suspension was heated at 100°C for 5 min and then frozen at -20°C for at least 15 min.

The macerates of all the spiked plants and naturally infected samples were crushed, prior to DNA extraction. DNA extractions and purifications were carried out using the QuickPick<sup>TM</sup> SML Plant DNA Kit (Bio-Nobile, Turku, Finland). Extraction, washing, and elution of the DNA were automated using KingFisher<sup>TM</sup> mL (Thermo Scientific). DNA extracts were kept at 4°C for a week, or stored at -20°C for a longer period.

### 159 2.5 Real-time PCR Harper et al., 2010

160 The real-time PCR assays were performed on the thermal cycler CFX96 real-time System C1000 Touch (Bio-Rad), using 96-well plates (Hard-Shell® 96-Well PCR Plates, #hsp9601, 161 Bio-Rad). The following thermal cycling program used was: 50°C for 2 min, 94°C for 10 min, 162 163 then 40 cycles of two step of 94°C for 10 s and 62°C for 40 s. The reaction mix was prepared 164 in a final volume of 20 µL containing: 1x TaqMan Fast Universal Master Mix (Applied Biosystems), 300 nM of each Xf forward and reverse primers (XF-F and XF-R, respectively), 165 166 100 nM of 6'FAM/BHQ-1 labeled probe (XF-P), 300 µg/µL of BSA, and 2 µL of DNA sample. For the artificially contaminated plant material, each sample was amplified in triplicate and on 167 168 three independent PCR runs to obtain nine Ct values per sample. For the naturally infected plant 169 material, each sample was amplified in duplicate on the same PCR run.

The data acquisitions and data analyses were performed using Bio-Rad CFX Manager, v 3.0.
The determination of Ct values was done using the regression mode. A Ct higher than 38 was
considered to be a negative result, according to the cut-off indicated by (Harper *et al.*, 2010).
For all the following analyses, the limit of detection was fixed at 100%, meaning the lowest
concentration at which all replicates gave a positive signal.

### 175 **2.6** Optimization and evaluation of the ddPCR assay

176 Two thermal gradients were tested to determine the optimal hybridization temperature ranging 177 from 54.6 to 64.6°C, and from 57 to  $62^{\circ}$ C. Thermal gradients were applied on samples of P. *myrtifolia*, spiked with suspensions of Xf ranging from  $1 \times 10^5$  b/mL to  $1 \times 10^3$  b/mL. BSA was 178 179 tested at the concentration determined by Harper *et al.* (2010) (300  $\mu$ g/ $\mu$ L) on a sample of *P*. *myrtifolia* spiked with Xf at a concentration of  $1 \times 10^5$  b/mL. In order to optimize the assay for 180 181 the five matrices spiked at  $1 \times 10^5$  b/mL, four different DNA volumes added to the reaction mix 182 were tested:  $2 \mu L$ ;  $4 \mu L$ ;  $6 \mu L$  and  $8 \mu L$ . Using the optimized protocol, tenth dilutions of L. 183 angustifolia and R. officinalis spiked with  $1 \times 10^3$  b/mL were tested. All the experiments 184 conducted to optimize the ddPCR assay were amplified in triplicate.

### 185 2.7 Optimized ddPCR assay

186 The optimized ddPCR reaction mix conditions retained were a final reaction volume of 20 µL containing: 1x ddPCR<sup>TM</sup> Supermix for Probes (No dUTP) (Bio-Rad), 900 nM Xf forward and 187 reverse primers (XF-F and XF-R), 250 nM 6'FAM/BHQ-1 labeled probe (XF-P), and 8 µL of 188 DNA sample. Droplets were generated with the QX200<sup>TM</sup> Droplet Digital<sup>TM</sup> System (Bio-Rad) 189 in a cartridge containing 20 µL of the reaction mix and 70 µL of Droplet Generation Oil for 190 191 Probes (ddPCR<sup>TM</sup> 96-Well Plates #12001925, Bio-Rad). The entire emulsion volume was 192 transferred from the cartridge to a 96-well PCR plate (Bio-Rad) and the PCRs were performed 193 on the thermal cycler CFX96 real-time System C1000 Touch (Bio-Rad). Optimal 194 thermocycling conditions retained were: DNA polymerase activation of 95°C for 10 min, then 195 40 cycles of two-steps of 94°C for 30 s for denaturation and 59°C for 60 s for hybridization and 196 elongation, followed by a final step at 98°C for 10 min for droplet stabilization. According to 197 Bio-Rad recommendations, a temperature ramp of 2°C/s was fixed on all PCR steps and the 198 lead was heated at 105°C. After amplification, the PCR plate was directly transferred to the 199 droplet reader QX200<sup>TM</sup> Droplet Digital<sup>TM</sup> System (Bio-Rad). QuantaSoft 1.7.4.0917 software 200 was used for data acquisition and data analysis. The entire concentration range of spiked 201 matrices was first amplified in triplicate on the same run. Then, for each matrix, samples with 202 concentrations equal to and below the limit of detection identified by real-time PCR were 203 amplified in triplicate on two other independent runs, in order to ultimately obtain nine results 204 for these samples. Finally, the amplifications of the naturally infected samples were performed 205 in one replicate in one run.

## 206 2.8 ddPCR analysis

Data were analyzed directly with QuantaSoft<sup>TM</sup> Analysis Pro software. For each plate and each 207 matrix, a threshold was manually set up just above the amplitude value of the cloud 208 209 corresponding to the negative droplets, also considered as the background, and according to the 210 results of the corresponding NTC (Lievens et al., 2016). This threshold enabled the 211 differentiation of droplets by categorizing them as positive (high level of fluorescence) or 212 negative (low and constant level of fluorescence). The PCR reactions with fewer than 10,000 213 droplets generated were excluded from the analysis, and a result was considered positive if at 214 least two positive droplets were detected. The software provided the results in target copies by 215 reaction using the following formula:

$$216 \qquad C = -\ln\left(1 - \frac{P}{P+N}\right) * \frac{1}{V}$$

Where C corresponded to the concentration in target DNA copies/well (cp/well), P the positive droplet number, N the negative droplet number, and V the mean volume in  $\mu$ L of one droplet. According to Bio-Rad, V is equal to  $0.85 \times 10^{-3} \mu$ L. Primers and probe targeted a part of the *rimM* gene, which is present in a single copy in the *Xf* genome. Therefore, the result can be directly converted into cp/ $\mu$ L in the initial samples, by multiplying it with the total volume of reaction mix (20  $\mu$ L), and then dividing it by the volume of DNA sample added to the reaction mix (8  $\mu$ L) at the beginning of the assay.

A bias, meaning the under or over-estimation of the quantification estimated by ddPCR, in comparison with the expected concentration, was calculated with the following formula:

$$226 \qquad Bias = \frac{detected DNA amount-assumed DNA provided}{assumed DNA provided} * 100$$

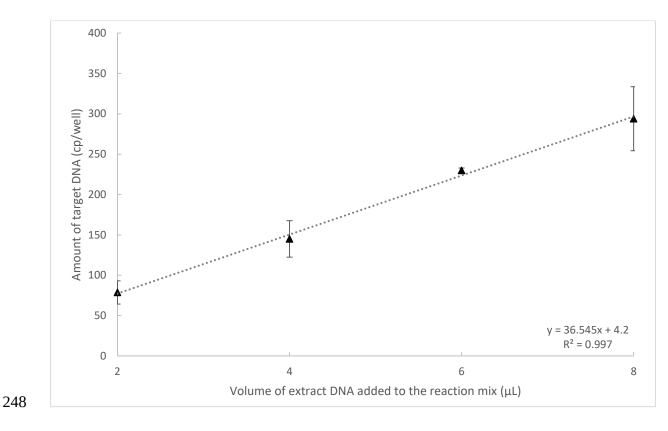
#### 227 **3 Results**

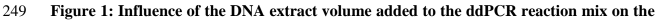
## 228 **3.1** Optimization of the ddPCR assay

229 The ddPCR appeared to be efficient for Xf detection at all tested temperatures. The first thermal 230 gradient tested allowed us to identify 58.5°C as the most suitable hybridization temperature, for 231 which positive droplets showed the highest fluorescence amplitude and were well distinguished 232 from the negative droplets. The second thermal gradient confirmed this preliminary result and 233 made it possible to fix the optimum hybridization temperature for ddPCR at 59°C. At this 234 temperature, positive droplets presented the highest fluorescence amplitude, the less "rain" (i.e. 235 droplets ranging between the positive and negative ones), and better separation from negative 236 droplets (data not shown).

Addition of BSA to the reaction mix did not increase the number of droplets amplified, the number of target DNA detected, nor the amplitude of the fluorescence signal. However, as it increased the standard deviation between replicates, no BSA was added for the optimized ddPCR protocol retained in this study (data not shown).

Like for *O. europaea* shown in Figure 1, *Xf* detection was successful for all the five matrices, regardless of the volume of DNA extract added to the PCR mix (Supplemental Data 1). Increasing the DNA volume added to the mix increased the number of DNA copies detected, without complete inhibition of the reaction. As the aim of the ddPCR assay in this study was also to improve the limit of detection of *Xf* in low-level contaminated samples, the final volume of DNA chosen was 8  $\mu$ L. This corresponded to the highest volume of DNA that could be added to ddPCR reaction mix in this study.





amount of DNA target detected for *O. europaea*.

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A tenth dilution of the DNA of *L. angustifolia* and *R. officinalis* spiked with  $1x10^3$  b/mL, corresponding to the first concentration under the limit of detection obtained with ddPCR, was tested. For *L. angustifolia*, none of the six replicates provided positive droplets. In only one of the six replicates tested for *R. officinalis*, two positive droplets were detected. Dilution of the DNA extract of *L. angustifolia* and *R. officinalis* did not improve the limit of detection or was not reproducible enough in this study.

## 258 3.2 Xf detection by ddPCR in spiked plant samples

259 The healthy status of the five plants was validated before spiking, by applying real-time PCR 260 Harper et al. (2010). Indeed, no Ct value was obtained for all the five plant matrices (Table 1). 261 As expected, in these assays, no cross-reaction was found, as no positive droplets were found for any of the NTCs, for the plant matrices. With the exception of O. europaea (curve could not 262 263 be drawn since only two concentrations gave some results), all the ddPCR matrix standard 264 curves showed high linearity and amplification efficiency, as the correlation coefficient (R<sup>2</sup>) 265 was higher than 0.96. This indicated the successful outcomes and good performances of all the 266 assays (Table 2).

267 Table 1: Mean Ct values obtained with real-time PCR for the bacterial suspension and the five spiked plant matrices.

Dilution range (b/mL)	Bacterial suspensi	on	L. angustifolia	O. europaea	P. myrtifolia	a	Q. ilex		R. officinal	lis
1x10 <sup>5</sup>	$32.18 \pm 0.38^{a}$ (9)	9) <sup>b</sup>	$30.76 \pm 0.18$ (9)	$36.02 \pm 0.43$ (9)	$30.38 \pm 0.17$	(9)	$31.85\pm0.10$	(9)	$32.24\pm0.17$	(9)
5x10 <sup>4</sup>	$33.14 \pm 0.45$ (	9)	$31.24 \pm 0.11$ (9)	$36.59 \pm 0.99$ (9)	$31.41\pm0.18$	(9)	$32.00\pm0.15$	(9)	$32.53\pm0.14$	(9)
$1x10^{4}$	$35.46 \pm 0.64$ (	9)	$33.68 \pm 0.20$ (9)	$37.32 \pm 0.05$ (3)	$33.85\pm0.34$	(9)	$34.90\pm0.50$	(9)	$34.83 \pm 0.66$	(9)
5x10 <sup>3</sup>	$35.55 \pm 0.21$ (	9)	$34.93 \pm 0.20$ (9)	$37.86 \pm 0.49$ (2)	$34.59 \pm 0.28$	(9)	$35.31\pm0.55$	(9)	$36.09 \pm 1.05$	(9)
1x10 <sup>3</sup>	$37.42 \pm 0.37$ (	6)	$37.45 \pm 0.58$ (9)	$37.78 \pm 0.53$ (2)	$38.00\pm0.56$	(9)	$37.62\pm0.86$	(7)	$38.05\pm0.50$	(6)
5x10 <sup>2</sup>	38.81 ± 0.12 (	5)	$37.44 \pm 0.00$ (1)	$36.61 \pm 0.00$ (1)	$37.62\pm0.18$	(5)	$37.76\pm0.29$	(4)	$38.25\pm0.13$	(4)
$1x10^{2}$	38.00 ± 0.19 (	6)	$38.46 \pm 1.01$ (2)	nd (0)	$38.29 \pm 0.13$	(4)	$36.75\pm0.19$	(6)	nd	(0)
0	nd <sup>c</sup> (	(0)	nd (0)	nd (0)	nd	(0)	nd	(0)	nd	(0)

268 <sup>a</sup>: Average Ct ± Standard Deviation (SD)

269 <sup>b</sup>: Number of positive replicates on nine replicates analyzed

270 <sup>c</sup>: Not detected

## 271 Table 2: Curve information for the *Xf* bacterial suspension and the five *Xf* spiked plant matrices.

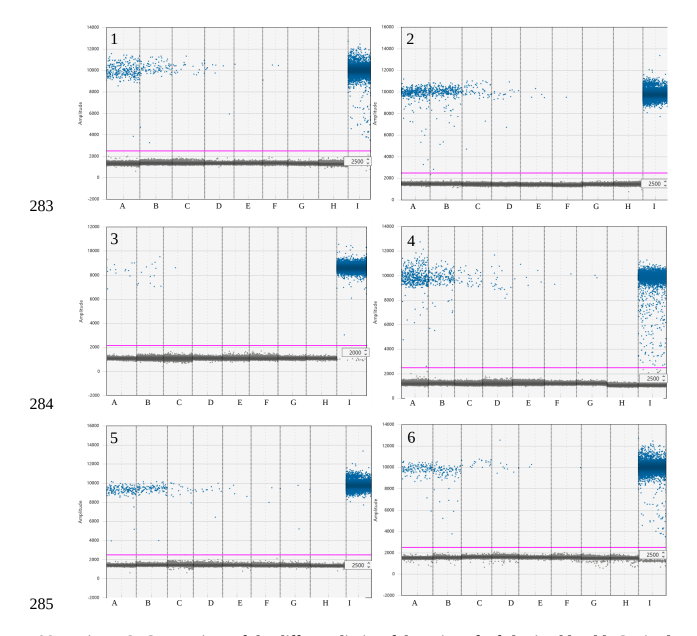
	Curve equation	R <sup>2</sup>
ddPCR		
Bacterial suspension	y = 0.4092x + 1136.7	0.99
L. angustifolia	$y = 85x + 2\ 007.35$	0.96
O. europaea	N/A <sup>a</sup>	
P. myrtifolia	y = 0.90x - 781.02	1.00
Q. ilex	$y = 0.36x + 1\ 187.91$	0.99
R. officinalis	y = 0.37x + 1363.74	0.98
REAL-time PCR		
Bacterial suspension	$y = 2.06 \times 10^{16} e^{-8,08 \times -01 \times 10^{16}} e^{-8,08 \times 10^{16}} e^{-8,0$	0.97
L. angustifolia	$y = 6.40 \times 10^{13} e^{-6,66E-01x}$	0.99
O. europaea	$y = 1.18 \times 10^{33} e^{-1.79 x}$	0.98
P. myrtifolia	$y = 1.10 \times 10^{13} e^{-6,13E-01x}$	0.99
Q. ilex	$y = 1.74 \times 10^{15} e^{-7,48E-01x}$	0.98
R. officinalis	$y = 2.32 \times 10^{15} e^{-7,48E-01x}$	0.99

 $a^{-1}$  As there were results for only two concentrations (10<sup>5</sup> b/mL and 5.10<sup>4</sup> b/mL), no curve could be drawn.

273 The results obtained for the five matrices and the bacterial suspension showed clear distinctions 274 between positive (blue) and negative (grey) droplets (Figure 2). The background (negative 275 droplets) had a similar fluorescence amplitude between samples of the same matrix, and 276 between the five matrices (mean amplitude of fluorescence of negatives droplets ranged from 277 1,070 to 1,606). The threshold was manually set at an amplitude of fluorescence between 2,000 278 and 3,000 for each ddPCR run. Compared to the positive control, which is a lysed suspension 279 of pure culture of Xf, very less rain was observed on the spiked sampled plots, showing high 280 efficiency of the PCR reactions.

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### 282 The figure n°2 should be print in color.



286 Figure 2: Comparison of the different limits of detection of Xf obtained by ddPCR in the bacterial suspension and spiked plant matrices. Pink line: threshold separating negative from 287 288 positive droplets. Blue dots: positive droplets with amplification. Grey dots: negative droplets 289 with no amplification. 1: Bacterial suspension. 2: Lavandula sp. 3: O. europaea. 4: P. 290 myrtifolia. 5: Q. ilex. 6: R. officinalis. Wells A to G bacterial suspension range of Xf, A: 1x10<sup>5</sup> b/mL; B: 5x10<sup>4</sup> b/mL; C: 1x10<sup>4</sup> b/mL; D: 5x10<sup>3</sup> b/mL; E: 1x10<sup>3</sup> b/mL; F: 5x10<sup>2</sup> b/mL and G: 291 292 1x10<sup>2</sup> b/mL. Well H: NTC specific to each matrix. Well I: positive control (lysis suspension of  $1 \times 10^7 \text{ b/mL of } X_f$ ). 293

- ddPCR enabled the detection of *Xf* in all the matrices, but at different concentrations (Table 3).
- 295 The limit of detection was fixed at  $5 \times 10^4$  b/mL, i.e,  $2.5 \times 10^5$  b/g of plant for *O. europaea*, at
- 296  $5x10^3$  b/mL, i.e,  $2.5x10^4$  b/g of plant for *L. angustifolia* and *R. officinalis*, and at  $1x10^3$  b/mL,
- 297 i.e.,  $5 \times 10^3$  b/g of plant for *Q*. *ilex*, *P*. *myrtifolia* and the bacterial suspension. The bias between
- the detected amount of DNA and the presumed amount of DNA provided was calculated for
- 299 each matrix at the limit of detection. Compared to the quantity of DNA expected, DNA
- 300 quantifications of Xf were overestimated by 6.54% and 23.96% in Q. ilex and in the bacterial
- 301 suspension, respectively. In L. angustifolia, R. officinalis, P. myrtifolia, and O. europaea, the
- 302 DNA quantifications of Xf were underestimated by 3.08%, 24.36%, 32.03% and 95.60%,
- 303 respectively.

Dilution range (b/mL)	Bacterial susp	pension	L. angustif	olia	O. europe	aea	P. myrtife	olia	Q. ilex	£	R. officin	alis
1x10 <sup>5</sup>	$\begin{array}{r} 4.02 x 10^{4a} \\ \pm \ 3.63 x 10^{3} \end{array}$	(3/3) <sup>b</sup>	$\begin{array}{c} 8.07 x 10^{4} \\ \pm \ 3.70 x 10^{3} \end{array}$	(3/3)	$\begin{array}{c} 3.38 x 10^3 \\ \pm \ 5.67 x 10^2 \end{array}$	(9/9)	$\begin{array}{c} 9.08 x 10^4 \\ \pm  4.95 x 10^3 \end{array}$	(3/3)	$\begin{array}{c} 3.58 x 10^4 \\ \pm \ 2.96 x 10^3 \end{array}$	(3/3)	$\begin{array}{c} 3.62 x 10^4 \\ \pm \ 3.71 x 10^3 \end{array}$	(3/3)
5x10 <sup>4</sup>	$\begin{array}{c} 2.53 x 10^4 \\ \pm \ 2.20 x 10^3 \end{array}$	(3/3)	$\begin{array}{c} 5.81 x 10^4 \\ \pm \ 2.02 x 10^3 \end{array}$	(3/3)	$\begin{array}{c} 2.20 x 10^3 \\ \pm \ 5.45 x 10^2 \end{array}$	(9/9)	$\begin{array}{c} 4.03 x 10^{4} \\ \pm \ 4.71 x 10^{3} \end{array}$	(3/3)	$\begin{array}{c} 2.30 x 10^4 \\ \pm \ 9.92 x 10^2 \end{array}$	(3/3)	$\begin{array}{c} 2.38 x 10^4 \\ \pm \ 3.20 x 10^3 \end{array}$	(3/3
1x10 <sup>4</sup>	$\begin{array}{c} 5.73 x 10^{3} \\ \pm \ 1.14 x 10^{3} \end{array}$	(3/3)	$\begin{array}{c} 1.17 x 10^{4} \\ \pm \ 2.19 x 10^{3} \end{array}$	(3/3)	$\begin{array}{c} 5.28 x 10^2 \\ \pm \ 1.90 x 10^2 \end{array}$	(4/9)	$\begin{array}{c} 6.17 x 10^{3} \\ \pm \ 1.39 x 10^{3} \end{array}$	(3/3)	$\begin{array}{c} 4.95 x 10^{3} \\ \pm \ 4.04 x 10^{2} \end{array}$	(3/3)	$\begin{array}{c} 5.49 x 10^{3} \\ \pm \ 2.98 x 10^{2} \end{array}$	(3/3
5x10 <sup>3</sup>	$\begin{array}{c} 2.84 x 10^{3} \\ \pm \ 8.78 x 10^{2} \end{array}$	(9/9)	$\begin{array}{c} 4.85 x 10^{3} \\ \pm \ 1.21 x 10^{3} \end{array}$	(3/3)	$\begin{array}{c} 4.25 x 10^2 \\ \pm \ 0.00 \end{array}$	(1/9)	$\begin{array}{c} 5.14 x 10^{3} \\ \pm \ 1.48 x 10^{3} \end{array}$	(3/3)	$\begin{array}{c} 2.81 x 10^{3} \\ \pm \ 7.90 x 10^{2} \end{array}$	(9/9)	$\begin{array}{c} 3.78 x 10^3 \\ \pm \ 2.16 x 10^3 \end{array}$	(9/9
1x10 <sup>3</sup>	$\begin{array}{c} 8.54 x 10^2 \\ \pm \ 2.57 x 10^2 \end{array}$	(9/9)	$\begin{array}{c} 5.66 x 10^2 \\ \pm \ 1.63 x 10^2 \end{array}$	(6/9)	No Call	(0/9)	$\begin{array}{c} 6.80 x 10^2 \\ \pm \ 2.68 x 10^2 \end{array}$	(9/9)	$\begin{array}{c} 1.07 x 10^{3} \\ \pm \ 5.58 x 10^{2} \end{array}$	(9/9)	$\begin{array}{c} 9.04 x 10^2 \\ \pm \ 1.61 x 10^2 \end{array}$	(7/9
5x10 <sup>2</sup>	$\begin{array}{c} 6.20 x 10^2 \\ \pm \ 1.19 x 10^2 \end{array}$	(8/9)	No Call	(0/9)	No Call	(0/9)	$\begin{array}{c} 5.15 x 10^2 \\ \pm \ 1.43 x 10^2 \end{array}$	(4/9)	$\begin{array}{c} 7.17 x 10^2 \\ \pm \ 2.59 x 10^2 \end{array}$	(7/9)	$\begin{array}{c} 6.18 x 10^2 \\ \pm \ 4.56 x 10^2 \end{array}$	(5/9
1x10 <sup>2</sup>	$\begin{array}{c} 5.84 x 10^2 \\ \pm \ 2.06 x 10^2 \end{array}$	(6/9)	$\begin{array}{c} 3.73 \text{x} 10^2 \\ \pm \ 0.00 \end{array}$	(1/9)	No Call	(0/9)	$\begin{array}{c} 1.77 \mathrm{x} 10^{2} \\ \pm \ 0.00 \end{array}$	(1/9)	$\begin{array}{c} 6.23 x 10^2 \\ \pm \ 1.50 x 10^2 \end{array}$	(7/9)	$\begin{array}{c} 2.01 x 10^2 \\ \pm \ 2.92 x 10^1 \end{array}$	(2/9
0	No Call	(0/9)	No Call	(0/9)	No Call	(0/9)	No Call	(0/9)	No Call	(0/9)	No Call	(0/9

304Table 3: Mean concentrations estimated in copies/mL (cp/mL) obtained with ddPCR for the bacterial suspension and the five spiked305plant matrices.

306  $\overline{a}$ : Average Ct ± SD

307 <sup>b</sup>: Number of positive replicates/number of replicates analyzed

## 308 3.3 Real-time PCR vs ddPCR for Xf detection in spiked samples

309 As for ddPCR, all the real-time PCR matrix standard curves showed high linearity and 310 amplification efficiency, as the correlation coefficient ( $R^2$ ) was greater than 0.97. This indicated 311 the successful outcomes and good performances of all the assays (Table 2).

312 Real-time PCR Harper et al. (2010) was used as a reference method in this study. For all the 313 assays, no mean Ct values exceeding 38 were recorded at a concentration equal to or higher 314 than the limit of detection, meaning that the limit of detection and positive results were 315 consistent. Moreover, for O. europaea the EPPO PM 7/24 protocol mentions a limit of detection for samples artificially contaminated with Xf subsp. multiplex of 100% at  $1 \times 10^5$  b/mL. In this 316 317 study, the limit of detection of  $5 \times 10^4$  b/mL was close to that presented in the PM 7/24 (EPPO, 2018b). The limit of detection of Xf in P. myrtifolia is known to be  $1 \times 10^3$ , which is the same as 318 319 the value we found (Legendre B., personal communication). The limits of detection of Xf subsp. 320 *multiplex* for the other matrices could not be compared, as there are no available data.

321 Real-time PCR and ddPCR technology provided equivalent limits of detection for Xf in the 322 following matrices: O. europaea, P. myrtifolia and R. officinalis (Table 1, Table 3). The ddPCR 323 technology presented a slightly higher limit of detection of 0.5 log for L. angustifolia. However, 324 a decrease in the limit of detection for Xf of 0.5 log for Q. ilex and the bacterial suspension 325 wereobserved. In the conditions of DNA extraction used for this study, and according to the volume of DNA added to the real-time PCR assay, the theoretical limit of detection should be 326  $1 \times 10^2$  b/mL for the five plant matrices. These results revealed that L. angustifolia and P. 327 328 myrtifolia may contain fewer real-time PCR inhibitors than Q. ilex and R. officinalis. Moreover, the limit of detection of the bacterial suspension was  $5 \times 10^3$  b/mL, meaning that the QuickPick 329 330 extraction kit may not be 100% efficient to extract the DNA of bacteria in pure culture.

331

## 332 3.4 Xf detection in naturally infected samples: real-time PCR vs ddPCR

A total of 22 samples from infected areas were tested using real-time PCR and ddPCR. Of these, 20 had a mean Ct value below 38 (ranging from 23.30 to 37.00) (Table 4). However, two samples, P13 and Q04, had a Ct value above 38 (38.65 and 39, respectively), and were considered negative.

# 337 Table 4: Mean Ct values and concentrations estimated in cp/mL of naturally infected

## 338 samples obtained after real-time PCR and ddPCR.

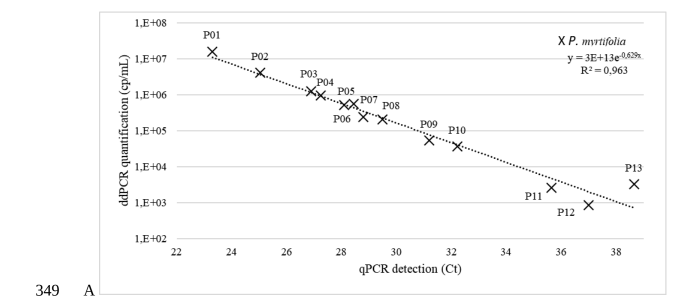
Matrices	Sample name	real-time PCR Ct means	ddPCR concentration (cp/mL)			
Calicotome sp	C01	36.41 (1/2) <sup>a</sup>	NA (1/12618) <sup>b</sup>			
L. angustifolia	L01	26.08 (2/2)	1.59E+06 (6 614/15 814)			
L. angustifolia	L02	26.69 (2/2)	8.82E+05 (3 581/13 826)			
L. angustifolia	L03	31.39 (2/2)	3.65E+04 (196/ 15 876)			
L. angustifolia	L04	31.78 (2/2)	4.19E+04 (227/16 056)			
P. myrtifolia	P01	23.30 (2/2)	1.61E+07 (17 615/ 17 690)			
P. myrtifolia	P02	25.04 (2/2)	4.12E+06 (11 714/ 15 550)			
P. myrtifolia	P03	26.90 (2/2)	1.28E+06 (5 896/16 720)			
P. myrtifolia	P04	27.25 (2/2)	9.52E+05 (4 395/15 894)			
P. myrtifolia	P05	28.10 (2/2)	5.24E+05 (2 895/17 733)			
P. myrtifolia	P06	28.44 (2/2)	5.70E+05 (2 985/16 936)			
P. myrtifolia	P07	28.80 (2/2)	2.39E+05 (1 446/18 565)			
P. myrtifolia	P08	29.48 (2/2)	2.11E+05 (1 272/18 376)			
P. myrtifolia	P09	31.20 (2/2)	5.38E+04 (296/16 332)			
P. myrtifolia	P10	32.21 (2/2)	3.66E+04 (216/17 459)			
P. myrtifolia	P11	35.64 (2/2)	2.63E+03 (16/17 898)			
P. myrtifolia	P12	37.00 (2/2)	8,56E+02 (3/10 309)			
P. myrtifolia	P13	38.65 (2/2)	3,25E+03 (12/10 848)			
Q. ilex	Q01	32.29 (2/2)	2.13E+04 (113/15 673)			
Q. ilex	Q02	34,85 (2/2)	1,02E+04 (39/11 269)			
Q. ilex	Q03	35,80 (2/2)	3,21E+03 (13/11 925)			
$\tilde{Q}$ . ilex	Q04	39.00 (1/2)	4,41E+02 (2/13 340)			

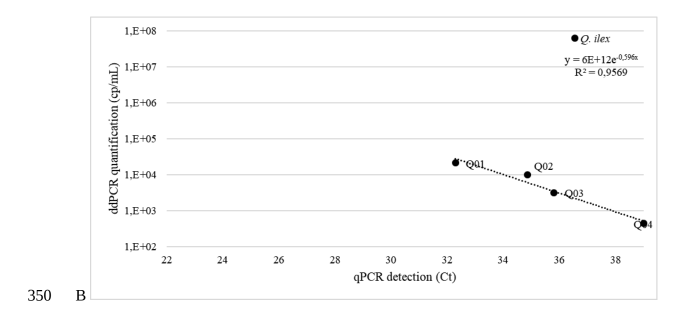
339

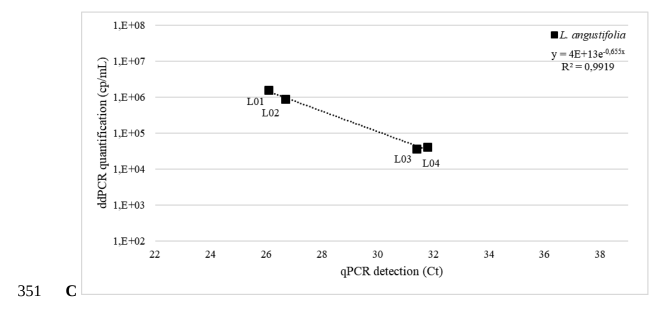
<sup>a</sup> Number of replicates positive for *Xf* detection / total number of analyzed replicates

<sup>b</sup> Number of positive droplets / total number of droplets (i.e. positives and negatives).

- 341 The 22 naturally infected samples were then analyzed by ddPCR. The presence of Xf was
- detected in 21 of them, including samples P13 and Q04, with at least two droplets, and a total
- 343 concentration ranging from  $4.41 \times 10^2$  cp/mL to  $1.61 \times 10^7$  cp/mL, confirming the ability of
- 344 ddPCR to detect Xf in naturally infected samples. As only one positive droplet was detected for
- 345 sample C01, this sample was considered negative by ddPCR. With the exception of samples
- L04, P06 and P13, the decrease in the Ct value was correlated with an increase in the quantity
- of DNA detected by ddPCR (Table 4). For each matrix the results obtained by ddPCR and real-
- 348 time PCR were compared and were highly correlated (Figure 3).







21

- 352 Figure 3: Correlation between the Ct values obtained by real-time PCR and the amount
- 353 of target DNA quantified by ddPCR (log cp/mL) for the naturally infected samples
- analyzed. A: samples of *P. myrtifolia*; B: samples of *Q. ilex*; C: samples of *L. angustifolia*.

### 355 **4 Discussion**

It has been observed that tenfold dilutions of the extracted DNA could reduce the effects of real-time PCR inhibitors (Francis *et al.*, 2006; Minsavage *et al.*, 1994; Modesti *et al.*, 2017). In this study, diluting the DNA extract of *L. angustifolia* and *R. officinalis* did not reduce the limit of detection using ddPCR, or the obtained results were not sufficiently reproducible. This approach does not seem to be useful and appropriate for *Xf* detection in these two matrices, using ddPCR. Nevertheless, more tests should be carried out to support this assumption.

- 362 Compared to real-time PCR, ddPCR can be considered as a controversial method. Some studies 363 have revealed that ddPCR was useful to improve pathogen detection sensitivity and to decrease 364 the impact of PCR inhibitors on PCR efficiency (Arvia et al., 2017; Bharuthram et al., 2014; 365 Dong et al., 2018; Rački et al., 2014; Zhao et al., 2016). In other cases, ddPCR was 10 and 100 fold less sensitive than real-time PCR in detecting cytomegalovirus and Leishmaniasis parasite 366 367 DNA, respectively (Hayden et al., 2013; Ramírez et al., 2019). Dreo et al., reported that ddPCR 368 benefits were dependent of the pathosystem studied (Dreo et al., 2014). The detection of 369 Erwinia amylovora showed similar levels using real-time PCR and ddPCR, while the detection 370 of R. solanacearum in low-level infected samples was improved by ddPCR (Dreo et al., 2014). 371 In our study, the two methods showed the same limit of detection for O. europaea, P. myrtifolia 372 and R. officinalis. Real-time PCR allowed better detection of 0.5 log for L. angustifolia, and 373 ddPCR allowed better detection of 0.5 log for Q. ilex and bacterial suspension.
- 374 ddPCR was also compared with real-time PCR on 22 naturally infected samples. Using real-375 time PCR, two samples P13 and Q04 had a Ct value higher than 38, and were thus considered 376 negative, i.e. not infected by Xf. As these two samples were frozen at -20°C for one year, it is 377 possible that the DNA was altered, explaining the higher Ct values obtained in this study. The 378 other 20 samples, considered positive, had a Ct value lower than 38. Using ddPCR, Xf was 379 detected in 21 samples, as more than two positive droplets were obtained. ddPCR did not reveal 380 the presence of Xf in sample C01 of Calicotome sp., unlike real-time PCR that detected Xf in 381 only one of the duplicates tested. These results could highlight the presence of PCR inhibitors 382 in this matrix. Moreover, ddPCR technology enabled the detection of Xf in both samples P13 383 and Q04, considered in this study as not infected by Xf using real-time PCR. As shown by Dreo 384 et al. (2014) for the detection of R. solanacearum, ddPCR technology could offer a real 385 advantage for the detection of pathogenic bacteria, and can be applied to the detection of Xf in

contaminated plants with low concentrations of target DNA (Dreo *et al.*, 2014). It also
successfully confirmed the positive samples identified using real-time PCR.

#### 388 **5** Conclusion

389 In this work, we proposed the first suitable ddPCR assay for the detection of Xf in plants. We 390 easily transferred a well-known routinely used real-time PCR technique for Xf detection in 391 ddPCR. Here, we reported all the set up steps leading to the optimal protocol and its comparison 392 with the current routine method. The results demonstrated the usefulness of ddPCR technology 393 as an alternative method for Xf detection in plants. However, the ddPCR assay is more time-394 consuming than real-time PCR and does not seem to be suitable for routine analysis. This 395 technology requires more steps than real-time PCR. Furthermore, the reaction mix has to handle 396 with care to ensure the generation of the appropriate number of generated droplets. 397 Nevertheless, as only two droplets are needed to confirm a sample as positive with ddPCR, this 398 method could confirm the status of samples found to be negative by real-time PCR due to high 399 Ct values, and could improve Xf detection in low-level infected samples. ddPCR should be 400 tested on insects to see whether this technology would still be efficient, and whether it offers a 401 benefit for *Xf* detection in this matrix.

402

### 6 Acknowledgements

We thank Marie-Agnès Jacques, Philippe Reignault, Pascal Gentit and Mathieu Rolland forfruitful discussions and critical reading of the manuscript.

### 405 **7 Funding**

406 This work was supported by ANSES and the Inra-SPE department. Enora Dupas was co-funded407 by ANSES and the Inra-SPE department.

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