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Development of a sensitive TaqMan qPCR assay for detection and quantification of venturia inaequalis in apple leaves and fruit and in air samples

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Development and validation of a highly sensitive real-time PCR TaqMan® assay for specific
 detection and quantification of *Venturia inaequalis* in apple leaves and fruit and in air samples

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11 Abstract

A TaqMan real-time PCR assay, based on the translation elongation factor 1-α gene, was developed 12 13 for the quantification of Venturia inaequalis in leaves and fruits of Malus x domestica and in spore trap samples. The designed primers and probe amplified a specific 86 bp fragment for V. inaequalis. 14 15 The specificity of the assay was tested using 35 strains of V. inaequalis and 20 different fungal species, including common pathogens of apple and other species of Venturia. The limit of detection 16 17 was 20 fg, which is lower than a single genome of V. inaequalis. The selectivity of the assay was tested using DNA from three cultivars of Malus x domestica and no influence on pathogen 18 amplification was found. The assay was also validated for repeatability and reproducibility. With this 19 assay, it was possible to detect and quantify V. inaequalis in four cultivars ('Ambrosia', 'Fiorina', 20 'Golden Delicious' and 'Mondial Gala') in both symptomatic and asymptomatic leaves, and in 21 symptomatic 'Golden Delicious' apple fruit, stored for 2 months. Furthermore, the assay was 22 successfully tested on air samples coming from apple orchards. The quantification of the molecular 23 assay, when compared with the estimated number of V. inaequalis cells using an optical microscope, 24 showed a correlation coefficient of 0.8186. The developed technique could be used to detect V. 25 26 inaequalis in asymptomatic samples and could be a promising tool for timely application of fungicides in orchards and to improve the efficacy of disease management. 27

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29 Keywords: apple scab, *Malus* x *domestica*, real-time PCR, TaqMan, *Venturia inaequalis*.

30

31 1. Introduction

Apple scab is a worldwide disease affecting apple cultivation (*Malus x domestica*, Bork H.) and it is caused by the ascomycete *Venturia inaequalis* (Cooke) G. Winter (Sutton et al., 2014). The disease is particularly severe in temperate climate regions characterized by humid and cool springs (Bowen

- et al., 2011) and, if not appropriately managed, can cause huge economic losses (MacHardy, 1996).
 Disease control requires an integrated strategy, based on prophylaxis practice, application of
 fungicides, and increasing use of resistant cultivars (MacHardy et al., 2001).
- V. inaequalis has a hemibiotrophic life cycle. Ascospores, which are released from pseudothecia and 38 form during winter in the fallen leaves, are the sexual reproductive structures of the pathogen, and 39 cause primary infections during the growing season (spring - early summer). If the weather conditions 40 are favorable, asexual conidia are released and cause secondary infections during the growing season 41 (Carrisse et al., 2000, Bowen et al., 2011). The entire life cycle of V. inaequalis is strongly influenced 42 43 by the humidity and temperature conditions (MacHardy, 1996). In order to reduce scab infections, fungicide application programs are scheduled based on the local weather conditions, on disease 44 45 prediction models and on the level of infection in spy plants (Mills and Laplante 1951; Gadoury and MacHardy 1986; Meitz-Hopkins et al., 2014; Carisse et al., 2000). The evaluation of the level of 46 47 infection is based on a visual scoring, which is an operator-dependent method used to attribute a disease severity index (Bock et al., 2010; Gusberti et al., 2012). 48
- The detection and quantification of pathogens, even in asymptomatic tissues, currently rely on molecular techniques. Different studies have been reported for specific amplification of *V. inaequalis* using conventional PCR (Schnabel et al. 1999; Stehmann et al. 2001, Koh et al., 2013). Daniëls et al.
- 52 (2012) developed a qPCR assay using the housekeeping genes ATP-binding cassette transporter 2
- 53 (ABC2) and the elongation factor (EF1), while Gusberti et al. (2012) developed a qPCR assay, based
- on primers and probe designed on the internal transcribed spacer (ITS) to detect *V. inaequalis*.
- Spore traps were used in combination with qPCR in order to quantify the air-borne inoculum of
 different plant pathogens (Carisse et al., 2009; Klosterman et al., 2014; Huang et al., 2016), including *V. inaequalis* (Meitz-Hopkins et al., 2014).
- To date, no studies have been carried out using the TaqMan technology combined with spore traps. 58 The aim of the current work was to develop and validate a specific TagMan quantitative PCR (qPCR) 59 60 assay that could be used to detect and quantify V. inaequalis on different Malus x domestica cultivars, both in the leaves and in the fruits. V. inaequalis isolates from different fields and growing seasons 61 62 were used to evaluate the assay specificity. Moreover, different plant materials (symptomatic and asymptomatic) and artificially inoculated leaves were tested. The assay was validated for specificity 63 64 on 20 different species, including Venturia asperata, which has recently been reported as a pathogen on scab-resistant varieties of apple having the Rvi6 gene (Caffier et al., 2012; Turan et al., 2019). 65 66 Furthermore, the assay was tested on spore trap samples in order to detect and quantify the airborne 67 inoculum of V. inaequalis.
- 68

69 **2. Materials and methods**

70

71 Fungal strains

72 Strains of V. inaequalis were isolated from leaves of apple tree (Malus x domestica) 'Ambrosia', 73 'Golden Delicious' and 'Mondial Gala', cultivated in northern Italy (Piedmont) during 2015 and 2016. Thirty-five isolates were selected for this study. Other apple pathogenic or commonly present 74 in orchard fungal species were isolated from apple leaves. All the isolates were identified through the 75 amplification of the ribosomal DNA internal transcribed spacer (ITS) region, following the protocol 76 77 of White et al. (1990). Reference strains (V. inaequalis CBS 815.69, V. asperata IRHS 2345, V. pirina CBS 120.825, V. nashicola CBS 794.84, V. cerasi CBS 444.54 and Fusicladium carpophilum CBS 78 79 497.62) and strains from the Agroinnova collection were used to verify the assay specificity. The 80 strains used in this study are summarized in Table 1. The stains were maintained as monoconidial 81 cultures in tubes of malt extract agar (MEA) (Sigma Aldrich, Germany) and stored at 4° C.

82

83 Fungal DNA extraction and quantification

Strains of *Venturia* spp. were grown on MEA for 30 days at 20±1 °C in the dark, while the other fungal species were grown for 10 days at 25±1 °C. The fungal DNA was extracted from approximately 200 mg of fresh-weight mycelium using an Omega E.Z.N.A. Fungal DNA Mini Kit (VWR, USA), according to the manufacturer's instructions. The DNA quality and the concentration were measured using a Nanodrop 2000 Spectrophotomer (Thermo Scientific, Wilmington, DE, USA).

90

Sequence analysis of the translation elongation factor gene and design of primers and TaqMan probe

The translation elongation factor 1 alpha (*EF1-\alpha*) sequences were amplified, for the 35 V. *inaequalis* 93 94 strains, using EF1 (CGAGAAGTTCGAGAAGGT) and EF2 (CCAATGACGGTGACATAG) primers. PCR was carried out in a total volume of 25 µL containing 2.5 µL of Buffer 10 X, 0.5 µL 95 96 of MgCl₂, 0.75 µL of dNTPs (10 mM), 1 µL of each primer (10 mM), 0.2 µL of Taq DNA polymerase (Qiagen, Germany) and 20 ng of template DNA. The thermal cycling program was performed 97 according to Gladieux et al. (2010). After agarose gel electrophoresis, the PCR products were purified 98 using a QIAquick[®] PCR purification Kit (Qiagen), and sequenced in both directions by Macrogen, 99 Inc. (The Netherlands). The consensus sequences were obtained by assembling forward and reverse 100 sequences, using DNA Baser (Heracle Biosoft, Romania). The consensus sequences obtained for the 101 102 V. inaequalis strains were compared with those deposited in GenBank and a multi alignment was

performed using the CLUSTALW algorithm, through Molecular Evolutionary Genetics Analysis
 (MEGA6) software, version 6.0. The alignment was used to design the primers and probe used in this
 study.

Six sets of primers and two probes were designed using Primer ExpressTM software 3.0 (Applied 106 Biosystem, Foster City, USA) and are listed in Table 2. The TaqMan probes were labelled at the 5'-107 end with the 6-carboxyfluorescein dye (6-FAM) reporter and Black Hole Quencher (BHQ1) or the 108 non-fluorescent quencher minor groove binder (NF-MGB) at the 3'-end. OligoCalc tool 109 (http://biotools.nubic.northwestern.edu/OligoCalc.html) was used to verify the presence of hairpins 110 and potential secondary structures, while *in silico* specificity was verified using the BLASTN tool of 111 the National Centre of Biotechnology Information (http://blast.ncbi.nlm.nih.gov/Blast.cgi). 112 Invitrogen (Carlsbad, USA) and Metabion (Steinkirchen, Germany) synthesized primers and probes, 113 respectively. 114

115

116 Conventional end-point PCR and qPCR optimization and amplification

117 The primer pair sets were first assessed by means of conventional end-point PCR using DNA from V. inaequalis strains and other selected species. The PCR endpoints were performed in a 25 µl 118 119 reaction, composed of 2 µL of Buffer 10x, 0.8 µL of MgCl₂, 1 µL of dNTPs (10 mM), 1 µL of each primer (10 mM), 0.2 µL of Taq Platinum Pfx DNA polymerase (Invitrogen, USA) and 20 ng of DNA. 120 The PCR thermal cycler conditions were 3 min at 95°C followed by 30 cycles of 95°C for 45 s, 54°C 121 for 45 s, 72°C for 1 min and a final extension of 5 min. The PCR products were run on 1% agarose 122 gel in a TBE buffer, and visualized under UV transilluminator using the Quantity One software 123 (BioRad Labs, Hercules, USA). After performing end-point PCR, the primer pairs that gave the best 124 results were selected and used in qPCR with SYBR Green in order to compare them with the 125 specificity and sensitivity of the TaqMan assay. Real-time reactions were performed using a 126 StepOnePlus qPCR system (Applied Biosystems) with 96 well-plates (Optical reaction plate, Applied 127 Biosystems) sealed with MicroAmp optical adhesive film (Applied Biosystems). 128

SYBR Green reactions were carried out using $10 \,\mu\text{L}$ of Power SYBR Green Mastermix 10x (Applied Biosystems), $1 \,\mu\text{l}$ of each primer (3 μ M) and $1 \,\mu\text{l}$ of template DNA. Amplification conditions were of $95 \,^{\circ}\text{C}$ for $10 \,\text{min}$, followed by 40 cycles of 57 $^{\circ}\text{C}$ for $1 \,\text{min}$ and $95 \,^{\circ}\text{C}$ for $15 \,\text{s}$. The melting curves were acquired after each run at the following conditions: $95 \,^{\circ}\text{C}$ for $15 \,\text{s}$, $60 \,^{\circ}\text{C}$ for $15 \,\text{s}$ and $95 \,^{\circ}\text{C}$ for 15 s. Sterile water was used as a negative control. In order to determine the sensitivity of the SYBR Green assay, a standard curve was obtained with *V. inaequalis* DNA using 1b14 strain 8-fold serially diluted (from 20 ng to 2 fg).

The TaqMan assay was performed using TaqMan Universal Mastermix 2X (Applied Biosystems). Each 96-well plate was loaded with a negative control, standard DNA and a positive control in triplicate. Different primer concentrations (from 3 μ M to 0.3 μ M) and different temperatures were initially tested at the annealing stage (57 to 60 °C). Reactions were carried out in a final volume of 20 μ l, with 1 μ l of DNA, 0.4 μ l of each primer (3 μ M) and 0.2 μ l of probe (5 μ M). Sterile water was used as a negative control. Amplification conditions were 95 °C for 10 min, 40 cycles of 57 °C for 1 min and 95 °C for 15 s. The newly developed TaqMan assay was compared, in terms of sensitivity

- and specificity, with the TaqMan assay previously published by Gusberti et al., (2012).
- 144 The Ct values generated by qPCR were compared with the standard curve to quantify the DNA of *V*.

inaequalis in target samples. The approximate number of cells was calculated by dividing the DNA quantity by the weight of the genome of *V. inaequalis* (0.0000597 ng; Deng et al., 2017), thereby

obtaining the number of target cells (*V. inaequalis*) per µL of reaction.

148

149 Specificity, sensitivity, selectivity, repeatability and reproducibility of the TaqMan qPCR

The TaqMan assay was validated according to the international EPPO standard PM 7/98 (EPPO 2014). In order to evaluate the specificity of the assay for *V. inaequalis*, 20 different species, including other pathogenic *Venturia* species that affect different hosts and *V. asperata*, which has recently been reported in Italy on apples, were used (Table 1). The presence of aspecific amplification signals was evaluated after 40 cycles in different experiments.

In order to determine the sensitivity of the TaqMan assay, a standard calibration curve was obtained 155 with V. inaequalis DNA using the 1b14 strain 8-fold serially diluted (from 20 ng to 2 fg) in sterile 156 157 deionized water. Furthermore, in order to verify the influence of the host DNA on V. inaequalis amplification, the pathogen DNA was 8-fold diluted in Malus x domestica 'Ambrosia', 'Mondial 158 Gala' and 'Golden Delicious' DNA. The standard curve reaction was carried out in triplicate and used 159 as an internal control in order to quantify the target DNA in different samples. Repeatability was 160 161 checked by running three independent assays of the test. Two different operators tested the reproducibility of the assay in two different laboratories and on different days. 162

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164 Detection of *V. inaequalis* in naturally infected *Malus* x *domestica* leaves and in apple fruit

During the 2015-2017 seasons, leaves from *Malus* x *domesica* plants showing symptoms of apple scab were harvested and stored for DNA extraction. From 'Ambrosia', 'Golden Delicious' and 'Mondial Gala' leaves, respectively 11, 11 and 5 samples were collected. Furthermore, four apples that had been stored for 2 months and which showed apple scab symptoms were selected for DNA extraction. DNA from the infected plants/fruit was extracted from approximately 100 mg of freshweight leaves or fruit. The samples were previously freeze-dried, ground with liquid nitrogen and
then extracted with E.Z.N.A. Plant DNA kit (VWR, USA), following the manufacturer's instructions.

172

173 Detection of *V. inaequalis* in asymptomatic *Malus* x *domestica* leaves

A total of 30 samples of asymptomatic *Malus* x *domestica* 'Ambrosia', 'Fiorina' and 'Golden Delicious' leaves were analyzed in May 2018 to assess the presence of *V. inaequalis* using the TaqMan assay. The leaves were divided into two equal parts using sterile blades and two different extractions were performed (100 mg each). The DNA extraction was performed as previously described.

179

180 Detection of *V. inaequalis* in artificially inoculated *Malus* x *domestica* leaves

V. inaequalis conidia were obtained from a single spore strain (1b14), according to Parker et al. 181 182 (2005). Leaves were obtained from Malus x domestica 'Fiorina' plants. The absence of V. inaequalis in the plant material was confirmed using a binocular microscope (Nikon Eclipse 55i, Tokyo, Japan). 183 184 Leaf disks (1 cm diameter) were collected for the assay, disinfected with a 10% solution of sodium hypochlorite, washed by immersion in sterile deionized water and air-dried. The inoculation of the 185 leaf disks was performed on the adaxial leaf surface with an initial concentration of 10² conidia/mL 186 of V. inaequalis, which was subsequently serially diluted (1:2, 1:3, 1:5, 1:10, 1:15, 1:20, 1:50 and 187 1:100). Control leaves were prepared in a similar way with sterile deionized water. Two replicates 188 were tested for each conidial concentration. DNA was extracted from artificially inoculated leaves, 189 following previously described procedures. 190

191

192 Detection of *V. inaequalis* from spore trap samples

A volumetric spore sampler (Burkard Manufacturing Co. Ltd., Rickmansworth, Hertfordshire, UK), 193 placed in an apple orchard in Manta (Cuneo, Italy; 44.609217; 7.502627), was used to detect the 194 195 airborne conidia of V. inaequalis. The tapes were collected at 24 h intervals for 14 days. Each daily tape was first visualized under an optical microscope to count the V. inaequalis cells at $40\times$ 196 magnification. The same segments were subsequently placed in 50 mL tubes and stored at 4 °C until 197 processing. An aliquot of 25 mL of a polyethylene glycol (PEG) alkaline buffer (50 g/L PEG average 198 Mn 4600; 20 mM KOH; pH 13.5) was added to each tube and vortexed for 20 minutes for the DNA 199 extraction. After incubation of 1 h at 65 °C, the samples were vortexed for 20 min and centrifuged 200 for 30 min at 8,000 g. The supernatant was collected and the DNA was extracted using an E.Z.N.A. 201 Fungal DNA Mini kit, according to the manufacturer's protocol, by adjusting the buffer volumes. 202

203 Repeatability of the assay on air samples was evaluated by performing 6 independent reactions using204 the same set of samples.

205

Data analysis. StepOne[™] software was used to automatically generate the baseline range and the
 qPCR standard curves, as well as to determine the Ct values. Student's t-test was used to analyze the
 reproducibility of the assay.

- 209
- 210 **3. Results**
- 211

212 TaqMan assay optimization

213 The EF1- α gene was selected because of the presence of a conserved region and the deletion of 31 nucleotides in the V. inaequalis species, able to differentiate from the other Venturia species, 214 215 including V. asperata. The alignment of partial EF1- α gene sequences from different V. inaequalis strains and 5 other *Venturia* species was used to design the primers and probes reported in Table 2. 216 217 The F1/R11 primer pair, which gave the best results with conventional end-point PCR, was selected for the real-time assays. The optimization of the TaqMan qPCR assay was tested using different 218 219 primer and probe concentrations and considering different cycling conditions. The best conditions were found for a primer concentration of 3 µM at 57 °C for 1:00 for the annealing stage. The Ven1 220 probe and F1/R11 primer pair were selected for the TaqMan assay as they showed the most suitable 221 amplification. 222

223

224 Specificity, sensitivity, selectivity, repeatability and reproducibility of the TaqMan qPCR

The TaqMan qPCR, with the designed primer pair and probe, was able to amplify the 86 bp fragment of the *EF1-a* gene in different experiments from 35 *V. inaequalis* strains (ST1). No amplification signal was detected after 40 cycles for *V. asperata*, *V. pirina*, *V. carpophila*, *V. cerasi*, *V. naschicola* or for the other tested species.

The DNA of V. inaequalis, serially diluted from 20 ng to 0.2 fg in sterile distilled water, was used to 229 230 build a standard curve in order to evaluate the limit of detection (LOD). The pathogen was quantifiable from 20 ng to 20 fg (Fig. 1), and a LOD threshold cycle (Ct), ranging between 36 and 231 37, was obtained. The LOD of 20 fg is lower than a single genome of V. inaequalis, (0.0000597 ng, 232 Deng et al., 2017). The mean value of the regression slope was -3.28, and the mean relative efficiency 233 was between 99% and 110%. No influence was observed on the selectivity of the TaqMan assay when 234 V. inaequalis DNA, serially diluted in Malus x domestica 'Ambrosia', 'Golden Delicious' and 235 236 'Mondial Gala' DNA, was used (Fig. 2). The amplifications showed similar PCR efficiencies and a reliable correlation between the Ct values and the amount of measured *V. inaequalis* DNA (Fig. 2).
No statistical differences (p>0.05) were found for the results of the student's t-test, which was used
to analyze the repeatability and reproducibility of the assay.

240

241 Specificity and sensitivity comparison of the SYBR Green and TaqMan assays

The SYBR Green assay by using the selected F1/R11 primer pair provided positive results for the target DNA from different cultivars, and no amplification was observed for the other *Venturia* species after 40 cycles. However, aspecific signals were detected for some species used as negative controls, i.e. *Cladosporium cladosporioides*, *Colletotrichum fioriniae*, *Fusarium equiseti*, *Penicillium expansum* and *Phoma* sp., after 34 to 36 cycles. The sensitivity of the assay with SYBR Green revealed a 10-fold higher LOD (200 fg) than the TaqMan assay designed on the *EF1-a* gene.

The TaqMan assay previously developed by Gusberti et al. (2012) gave a cross-reaction for the strain
MALT1 *Alternaria* sp. isolated from apple leaves, while no other amplification was obtained for the

other tested species. Sensitivity, assessed using the method of Gusberti et al. (2012) on the V. *inaequalis* 1b14 strain, showed a LOD of 100 fg (ST2).

252

253 Detection of *V. inaequalis* in naturally infected *Malus* x *domestica* leaves and fruit

The TaqMan assay was used to quantify *V. inaequalis* in naturally infected leaves of three *Malus* x *domestica* cultivars and apple fruit. No influence of the host DNA was detected, and all the analyzed samples resulted positive for the target amplification (Fig. 3A and ST3). The average number of cells/ μ L was 5.26 x 10⁴ for the 'Ambrosia' samples, 4.28 x 10⁴ for the 'Mondial Gala' samples and 3.76 x 10⁴ for the 'Golden Delicious' samples. The assay was able to detect 2.83 *V. inaequalis* cells/ μ L in the naturally infected leaves (mean Ct 34.35) and 182 cells/ μ L in the naturally infected fruit (mean Ct 28.38) (Fig. 3A and ST3).

261

262 Detection of *V. inaequalis* in asymptomatic *Malus* x *domestica* leaves

The TaqMan assay was able to detect *V. inaequalis* in both the asymptomatic resistant and susceptible cultivars (Fig. 3B and ST4). The lowest concentration was 3.21 cells/ μ L and it was found in the As7 sample (resistant 'Fiorina'), while the highest concentration was found in the susceptible 'Ambrosia' samples, with 4.50 x 10³ cells/ μ L. The mean *V. inaequalis* concentration was 1.90 x 10² cells/ μ L for 'Fiorina', while 3.46 x 10² cells/ μ L were found for 'Golden Delicious' and 1.91 x 10³ cells/ μ L for 'Ambrosia'.

269

270 Detection of *V. inaequalis* in artificially inoculated *Malus* x *domestica* leaves

In order to test the developed TaqMan assay on environmental-like samples, *Malus* x *domestica* leaves were artificially inoculated with a *V. inaequalis* conidial suspension. The results obtained for the quantification of *V. inaequalis* are reported in Figure 4. All the samples amplified with a clear amplification signal, with Ct values ranging from 28, for the samples inoculated with highest concentration, to 36 for the lowest. No amplification was obtained for the negative controls. The amplification showed linearity in the serial dilutions, and the assay allowed us to quantify from 1.65 x 10² cells/µL, for the initial sample, to 1.21 cells/µL, for the 100-fold diluted sample.

278

279 Detection of *V. inaequalis* from spore trap samples

The TaqMan assay was used to detect and quantify the presence of airborne inoculum of V. inaequalis 280 in the spore trap samples. The estimated conidial concentrations, based on microscope counts, were 281 higher than the estimated concentrations calculated from the amount of V. inaequalis DNA detected 282 283 by the TaqMan assay. Only one of the 14 analyzed samples showed no amplification (Table 3). The estimated mean number of cells/ μ L obtained from the microscope count was 1.01 x 10⁴, while it was 284 5.26×10^3 for the TagMan assay. The correlation coefficient between the two assays was positive. 285 with an R^2 value of 0.8186. The repeatability of the assay was variable, with at least one positive 286 287 amplification per sample over six reactions (Table 3).

288

289 4. Discussion

The molecular techniques currently applied for the detection of plant pathogens are often used to specifically identify and quantify fungal species in crops and food commodities (Postollec et al., 2001; Capote et al., 2012; Aslam et al., 2017). In this study, a highly sensitive TaqMan real-time assay has been developed for the specific detection and quantification of *V. inaequalis*, and it has successfully been used with symptomatic and asymptomatic leaves, fruit and air samples.

Conventional molecular methods, based on PCR, showed specificity when used for the detection of 295 296 Venturia species (Schnabel et al., 1999;, Koh et al., 2013), but there is a lack of effective quantitative results (Suarez et al., 2005). Real-time PCR gives more specific and sensitive results than 297 298 conventional PCR and allows pathogens to be quantified (Selma et al., 2008; Mirmajlessi et al., 2015; 299 Baskarathevan et al., 2016; Kuzdraliński et al., 2017). Real-time PCR was used in previous studies to detect and quantify V. inaequalis in different types of samples. Both SYBR Green I and the 300 TaqMan assay were utilized. The first technique exploited a fluorescent measurement for DNA 301 amplification through double strand DNA binding dyes (SYBR Green I®), although there were some 302 limits pertaining to aspecific amplifications and difficulties in the interpretation of the results after 303 the melting curve (Martinez et al., 2011). These issues could be solved by using the second technique, 304

i.e. TaqMan chemistry, which includes specific fluorescent probes into the PCR (Amaral Carneiro et
 al., 2017).

- In preliminary studies, different target sequences were explored for the development of the TaqMan real-time assay, following multiple sequence alignments (data not shown). The used sequences were obtained from strains isolated in Italy and from sequences available in the GenBank public database. Due to the high intraspecific variability, the ITS region and the beta-tubulin gene were discarded (data not shown), whereas the *EF1-a* gene proved a highly conserved species-specific region for *V*. *inaequalis* and it was therefore selected to design primers and probes.
- 313 The developed assay was compared using SYBR Green I and with the TaqMan assay, previously 314 published (Gusberti et al., 2012). The primers designed on the $EF1-\alpha$ gene cross-reacted with various 315 tested species when used with SYBR Green I. Cross-reaction was also observed in the work of Meitz-Hopkins et al. (2014), where two qPCR assays, based on the CYP51A1 gene and the ITS region, were 316 317 developed. The study of Daniëls et al. (2012), using a qPCR assay based on the ITS region, also showed cross-reaction. The main fungal species that showed cross-amplification with the 318 319 aforementioned molecular tools were Alternaria sp. and V. asperata, V. nascicola, V. pirina, V, cerasi 320 and V. carpophila. Our data demonstrated that the use of SYBR Green allowed to reach a sensitivity 321 of 200 fg, which in turn results in a 5 times lower than the detection limit obtained by Meitz-Hopkins 322 et al. (2014), but 2 times higher than that reported by Daniëls et al. (2012).
- When the specificity was tested using the TaqMan probe technology, no cross-reaction was detected for the other *Venturia* species, including *V. asperata*, or for other pathogens. On the contrary, when the specificity was tested with the TaqMan probe developed by Gusberti et al. (2012), a positive amplification was obtained for a strain of *Alternaria* sp. from apple leaves. A blast search of the primers and probes used in the study of Gusberti et al. (2012) gave 100% homology and 100% coverage with *A. solani* (accession number CPO22033.1) and 100% homology and 72% coverage with *A. alternata* (accession number XM_01852723.1).
- Because of the high intraspecific variability that exists within the species *V. inaequalis* (Tenzer and Gessler, 1999; Tenzer et al., 1999; Ebrahimi et al., 2016), the specificity of the assay was confirmed on DNA extracted from a large number of pure cultures of *V. inaequalis* strains, isolated from different cultivars over different years, and positive amplifications were observed for all the strains. The assay showed a low variation in the Ct values obtained in independent experiments and resulted
- to be highly reproducible.
- When the sensitivity of the assays was compared, the TaqMan assay resulted in a lower detection limit than the SYBR Green assay. The advantage of using the TaqMan assay has been reported in a qPCR comparative study by Soltany-Rezaee-Rad et al. (2015). A qPCR with SYBR Green could

inhibit the Taq DNA polymerase, thereby reducing the sensitivity of the assay (Kermekchiev et al.,

2009). The LOD of our TaqMan assay (20 fg) proved to be more sensitive than that reported by
Gusberti et al. (2012) for which the LOD was 100 fg. The LOD of the TaqMan assay (20 fg) is similar

to those reported for other species i.e. *Botrytis cinerea* (Suarez et al., 2005), *Fusarium solani* (Bernal-

343 Martinez et al., 2012), Aspergillus fumigatus (Fernandez-Molina et al., 2014), Lichtheimia

344 *corymbifera* (Springer et al., 2016), *Fusarium fujikuroi* (Amaral Carneiro et al., 2017) and *Fusarium*

345 *culmorum* (Bilska et al., 2018).

The TaqMan assay was also tested to quantify the target DNA in the presence of the plant material. The sensitivity was not affected by the presence of the DNA of *Malus* x *domestica* from different cultivars, and the assay allowed to detect and quantify *V. inaequalis* in inoculated samples with significant linearity. Positive amplifications were also obtained from leaf samples, collected both from susceptible and resistant cultivars, that were, respectively, symptomatic or asymptomatic. In addition, the assay proved to be useful for the detection of the pathogen in fruit samples.

When the assay was performed on DNA obtained from the spore trap, 13 samples out of 14 resulted 352 353 positive to V. inaequalis, even with a low conidial concentration. In our study, the overall concentration estimated with a microscope count resulted higher than that estimated with the TaqMan 354 355 assay, but with a linear relationship (\mathbb{R}^2 : 0.8186), as previously reported by Carisse et al. (2009), pertaining to the quantification of Botrytis squamosa. However, the repeatability of the assay was 356 variable for air samples, as inhibition of PCR may occur when DNA is extracted from air samples, 357 thus false-negative plate readings and a reduced amplification efficiency may be obtained. The 358 reasons for these false negatives could be linked to competition in the amplification due to: high 359 amounts of non-target DNA, inoculum density, co-extraction of contaminants (PCR inhibitors) or 360 unequal distribution of conidia on tapes (McDevitt et al., 2007; Rogers et al., 2009; Williams et al. 361 2001; Bilodeau et al 2012; Klosterman et al., 2014; Dung et al., 2015). 362

In conclusion, the development of a highly sensitive species-specific assay is important to detect 363 pathogen at low concentrations, even during latent infection or in asymptomatic samples. In our study, 364 the use of a TaqMan real-time assay increased the sensitivity of the molecular tool and led to the 365 366 advantage of being able to detect less than a single target cell. The assay proved to be specific and highly sensitive for the detection of V. inaequalis, both in symptomatic and asymptomatic apple 367 leaves. The developed qPCR could be used in apple scab risk management systems to quantify the 368 inoculum in the field and to plan phytosanitary treatments. Furthermore, the technique proved to be 369 370 sensitive with air samples, though assessed on a limited number of samples.

This newly developed TaqMan assay could be a useful tool, combined with a weather-based data model, to plan phytosanitary treatments in order to control apple scab. The here developed TaqMan assay could also be used, together with nanobiosensor technology, to quantify airborne inoculum in
the field, without time consuming DNA extraction and processing, and could therefore be used to
implement a decision support system for apple scab management.

376

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530 Tables

Table 1 – Strain ID, host, source of isolation and origin of the fungal species used in this study to
 develop the TaqMan real-time PCR assay.

Species	Strain ID	Host	Source	Origin
Venturia inaequalis	1b1	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	1b5	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	1b10	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	1b7	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	1b6	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	116_c2	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	216_b4	Malus x domestica 'Golden Delicious'	Leaf	Italy
Venturia inaequalis	3b6	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	3b2	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	3b5	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	516_6	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	516_2	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	516_3	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	3aa	Malus x domestica 'Golden Delicious'	Leaf	Italy
Venturia inaequalis	3a1	Malus x domestica 'Golden Delicious'	Leaf	Italy
Venturia inaequalis	3ac	Malus x domestica 'Golden Delicious'	Leaf	Italy
Venturia inaequalis	416_a12	Malus x domestica 'Mondial Gala'	Leaf	Italy
Venturia inaequalis	416_a5	Malus x domestica 'Mondial Gala'	Leaf	Italy
Venturia inaequalis	416_a9	Malus x domestica 'Mondial Gala'	Leaf	Italy
Venturia inaequalis	516_h	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	516_4	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	516_5	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	3af	Malus x domestica 'Golden Delicious'	Leaf	Italy
Venturia inaequalis	416_b2	Malus x domestica 'Golden Delicious'	Leaf	Italy
Venturia inaequalis	416_a1	Malus x domestica 'Mondial Gala'	Leaf	Italy
Venturia inaequalis	516_10	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	1b2	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	416_a13	Malus x domestica 'Mondial Gala'	Leaf	Italy
Venturia inaequalis	1b9	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	3b3	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	3b4	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	3b10	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	1b14	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	416_a3	Malus x domestica 'Mondial Gala'	Leaf	Italy
Venturia inaequalis	1b13	Malus x domestica 'Ambrosia'	Leaf	Italy
Venturia inaequalis	CBS 815.69	Malus sylvestris	Fruit	The Netherlands
Venturia cerasi	CBS 444.54	Prunus cerasus	Unknown	Germany
Venturia pirina	CBS120.825	Pyrus communis	Unknown	Brazil

Species	Strain ID	Host	Source	Origin
Venturia nashicola	CBS 794.84	Pyrus serotina var Culta	Unknown	Japan
Venturia asperata	IRHS 2345	Malus x domestica	Fruit	France
Venturia carpophila	CBS 497.62	Prunus mirabelle	Unknown	Switzerland
Monilia fructicola	MON1	Malus x domestica	Leaf	Italy
Monilia laxa	LAXA3	Prunus persica	Fruit	Italy
Botryosphaeria dothidea	BOTRYO1	Malus x domestica	Leaf	Italy
Sclerotinia sclerotiorum	SCLA2	Unknown	Unknown	Italy
Colletotrichum fioriniae	COLLRIV	Malus x domestica	Fruit	Italy
Epicoccum nigrum	EPI2	Malus x domestica	Leaf	Italy
Stemphylium sp.	STEM	Pyrus communis	Fruit	Italy
Cladosporium cladosporioides	CLAD1	Malus x domestica	Leaf	Italy
Alternaria mali	CBS 106.24	Malus sylvestris	Unknown	The USA
Alternaria sp.	MALT1	Malus x domestica	Leaf	Italy
Botrytis cinerea	BOT1	Malus x domestica	Leaf	Italy
Phoma sp.	PH5	Malus x domestica	Leaf	Italy
Fusarium equiseti	3FEQS	Unknown	Unknown	Italy
Penicillium expansum	PEX16	Malus x domestica	Leaf	Italy
Ramularia mali	RM2	Malus x domestica	Fruit	Italy

535	Table 2. Primer pairs and probes, designed on $EF1-\alpha$ gene used for the detection of V. inaequalis with
536	the TaqMan real-time assay.

Primers and Probes A	Amplicon (bp)	Sequence (5'-3')
F 3.2/R11	191 bp	ACCCGGATTTCATTTCGAAACT GCAATCGTTAGCATCGTCATAGTG
F 4.4/R11	154 bp	TTTTGCACTGTGGCAGCCC GCAATCGTTAGCATCGTCATAGTG
F 1/R11	86 bp	CACTTCCCCGCTATTCACGT GCAATCGTTAGCATCGTCATAGTG
F2.3/R6	127 bp	TTGCCCCTCCAAAATTACAGTG GGCGGCTTCCTATTGCAATC
F1/R6	100 bp	CACTTCCCCGCTATTCACGT GGCGGCTTCCTATTGCAATC
F4.3/R6	171 bp	AAATTTTGCACTGTGGCAGC GGCGGCTTCCTATTGCAATC
F 2.2/R6	127 bp	TTGCCCCTCCAAAATTACAGT GGCGGCTTCCTATTGCAATC
Vina	-	[FAM] - AGCCCAACTTTCTCCGGTCC - [NF-MGB]
Ven1	-	[FAM] - CTCAAGGCAGCCCAACTTTCTCCGGT - [BHQ1]

Table 3 – Number of *V. inaequalis* cells estimated using the optical microscope compared with the number of cells estimated with the TaqMan real-time assay for the analyses of spore trap samples, and repeatability of the assay, expressed as the number of positive amplifications in six different reactions.

Sample	Estimated number of cells (optical microscope)	Estimated number of cells (TaqMan assay)	Number of positive results
ST1	2.73 x 10 ⁴	5.49 x 10 ³	6/6
ST2	5.21 x 10 ⁴	6.88 x 10 ³	6/6
ST3	1.31 x 10 ⁴	$1.12 \text{ x } 10^4$	5/6
ST4	$7.94 \text{ x } 10^3$	$4.42 \text{ x } 10^3$	6/6
ST9	$1.30 \ge 10^3$	2.72×10^3	3/6
ST10	$2.05 \text{ x } 10^3$	$4.80 \ge 10^3$	3/6
ST10/2	$4.67 \ge 10^2$	*	*
ST11	8.92 x 10 ²	$1.66 \ge 10^3$	2/6
ST12	8.51 x 10 ³	9.34 x 10 ³	1/6
ST13	$1.37 \ge 10^3$	2.58×10^3	4/6
ST17	2.83 x 10 ³	5.14 x 10 ³	2/6
ST21	$1.09 \text{ x } 10^4$	7.93 x 10 ³	2/6
ST27	$1.36 \ge 10^3$	3.70×10^3	2/6
ST28	2.21 x 10 ³	2.52×10^3	2/6

542

543 *the extraction failed for the ST10/2 sample.

544 **Captions**

- Figure 1. Standard curve obtained with genomic DNA of the *V. inaequalis* 1b14 strain showing the correlation between the *V. inaequalis* DNA amount and the Ct values (the standard deviation values range from 0.04 to 0.22, thus the standard deviation bars are too small to display on the graph).
- 548

Figure 2. Standard curve obtained with genomic DNA of the *V. inaequalis* 1b14 strain diluted in *Malus x domestica* DNA 'Ambrosia', 'Golden Delicious' and Mondial Gala. The *V. inaequalis* DNA amount is plotted against the Ct values (the standard deviation values range from 0.03 to 0.53, thus the standard deviation bars are too small to display on the graph).

553

Figure 3. Detection of *V. inaequalis* with the TaqMan real-time PCR on: **A.** naturally infected leaves of *Malus x domestica* 'Ambrosia', 'Golden Delicious' and 'Mondial Gala'; **B.** asymptomatic leaves of *Malus x domestica* 'Ambrosia', 'Golden Delicious' and 'Fiorina'. The values are expressed as the mean \pm SD (n=9).

558

Figure 4. Detection of *V. inaequalis* with the TaqMan real-time PCR on *Malus* x *domestica* 'Ambrosia' leaf disks artificially inoculated at different concentrations. The values are expressed as

561 the mean \pm SD (n=9).

562 Supplementary Tables

Supplementary Table 1 - Results of the q-PCR amplification (Ct values ± standard deviation) with

the target DNA (\cong 20 ng) of different *V. inaequalis* strains used to evaluate the specificity of the TaqMan real-time assay.

Sample ID	Species	Cultivar	Ct Mean ±SD*
1b1	V. inaequalis	Ambrosia	$18.57 \pm 0.04 $
1b5	V. inaequalis	Ambrosia	$18.23 \pm 0.24 $
1b10	V. inaequalis	Ambrosia	$17.42 \pm 0.04 $
1b7	V. inaequalis	Ambrosia	$17.13 \pm 0.23 $
1b6	V. inaequalis	Ambrosia	17.58 ± 0.19
116_c2	V. inaequalis	Ambrosia	18.12 ± 0.30
216_b4	V. inaequalis	Ambrosia	17.32 ± 0.20
3b6	V. inaequalis	Ambrosia	17.14 ± 0.20
3b2	V. inaequalis	Ambrosia	$18.22 \pm 0.11 $
3b5	V. inaequalis	Ambrosia	17.20 ± 0.13
516_6	V. inaequalis	Ambrosia	$18.35 \pm 0.30 $
516_2	V. inaequalis	Ambrosia	17.41 ± 0.13
516_3	V. inaequalis	Ambrosia	17.81 ± 0.17
3aa	V. inaequalis	Golden Delicious	$17.49 \pm 0.22 $
3a1	V. inaequalis	Golden Delicious	$17.79 \pm 0.03 $
3ac	V. inaequalis	Golden Delicious	17.52 ± 0.12
416_a12	V. inaequalis	Mondial Gala	18.49 ± 0.10
416_a5	V. inaequalis	Mondial Gala	17.92 ± 0.31
416_a9	V. inaequalis	Mondial Gala	$18.29 \pm 0.30 $
516_h	V. inaequalis	Ambrosia	17.44 ± 0.37
516_4	V. inaequalis	Ambrosia	18.14 ± 0.37
516_5	V. inaequalis	Ambrosia	18.09 ± 0.22
3af	V. inaequalis	Golden Delicious	17.75 ± 0.31
416_b2	V. inaequalis	Golden Delicious	17.76 ± 0.36
416_a1	V. inaequalis	Mondial Gala	17.76 ± 0.39
516_10	V. inaequalis	Ambrosia	$18.45 \pm 0.32 $
1b2	V. inaequalis	Ambrosia	$17.66 \pm 0.23 $
416_a13	V. inaequalis	Mondial Gala	17.47 ±0.24
1b9	V. inaequalis	Ambrosia	17.94 ± 0.11
3b3	V. inaequalis	Ambrosia	17.11 ± 0.24
3b4	V. inaequalis	Ambrosia	17.68 ± 0.11
3b10	V. inaequalis	Ambrosia	17.13 ± 0.25
1b14	V. inaequalis	Ambrosia	18.00 ± 0.39
416_a3	V. inaequalis	Mondial Gala	17.85 ± 0.28
1b13	V. inaequalis	Ambrosia	17.34 ± 0.07

^{*} The values are expressed as the mean \pm SD (n=9)

Supplementary Table 2 – Standard curve obtained for the quantification of the *V. inaequalis* 1b14 strain using the TaqMan real-time PCR assay developed by Gusberti et al. (2012). Ct mean values \pm standard deviation.

V. inaequalis DNA concentration (ng)	Ct values ± SD
10	20.13 ± 0.11
1	23.24 ± 0.12
0.1	26.74 ± 0.10
0.01	30.10 ± 0.04
0.001	33.69 ± 0.71
0.0001	36.21 ± 0.86

572	Supplementary Table 3 – Ct mean values \pm standard deviation and the estimated number of V.
573	inaequalis cells/ μ L \pm standard deviation for the symptomatic Malus x domestica 'Ambrosia',
574	'Mondial Gala' and 'Golden Delicious' leaves obtained using the TaqMan real-time PCR assay.

Sample ID	Cultivar	Ct mean ± SD*	V. inaequalis cells/ μ L ± SD*
1b	Ambrosia	21.93 ± 0.10	$1.67 \text{ x } 1 \pm 0^4 \pm 1137.31$
3b	Ambrosia	25.15 ± 0.20	$1.59 \text{ x } 10^3 \pm 209.73$
Amba17	Ambrosia	19.89 ± 0.27	$8.63 \ x \ 10^4 \pm 15945.86$
316s2	Ambrosia	18.86 ± 0.18	$1.25 \ge 10^5 \pm 15918.36$
216	Ambrosia	25.38 ± 0.54	$1.91 \text{ x } 10^3 \pm 777.22$
516	Ambrosia	22.35 ± 0.64	$2.19 \text{ x } 10^4 \pm 8523.70$
Amb1	Ambrosia	23.89 ± 0.34	$5.59 \ge 10^3 \pm 1262.95$
Amb4	Ambrosia	25.29 ± 0.90	$1.53 \text{ x } 10^3 \pm 829.27$
Col1	Ambrosia	18.71 ± 0.29	$1.36 \ge 10^5 \pm 26719.38$
Col2	Ambrosia	20.37 ± 0.38	$5.90 \ge 10^4 \pm 1673.62$
Col3	Ambrosia	19.12 ± 0.48	$1.23 \ x \ 10^5 \pm 40605.49$
416a1	Gala	27.96 ± 0.12	$3.20 \text{ x } 10^3 \pm 476.83$
416a2	Gala	18.12 ± 0.05	$2.48 \ge 10^5 \pm 8191.52$
GalaA	Gala	20.76 ± 0.79	$4.31 \ge 10^4 \pm 17722.88$
Gala1	Gala	25.81 ± 0.03	$9.33 \ge 10^2 \pm 261.42$
Gala2	Gala	24.85 ± 0.24	$1.82 \text{ x } 10^3 \pm 311.25$
Gala3	Gala	24.98 ± 0.18	$2.01 \text{ x } 10^3 \pm 399.74$
Gala4	Gala	27.79 ± 0.49	$2.38 \ge 10^2 \pm 73.01$
Gold11	Golden Delicious	21.04 ± 0.39	$3.16 \ge 10^4 \pm 8540.33$
116	Golden Delicious	18.32 ± 0.24	$2.07 \text{ x } 10^5 \pm 34735.35$
1a	Golden Delicious	34.35 ± 0.37	$0.28 \ x \ 10^1 \pm 0.67$
416b2	Golden Delicious	22.58 ± 0.17	$1.13 \text{ x } 10^4 \pm 1400.69$
416b3	Golden Delicious	23.53 ± 0.23	$5.85 \text{ x } 10^3 \pm 977.49$
Gold22	Golden Delicious	21.26 ± 0.35	$3.35 \ge 10^4 \pm 8582.51$
3a	Golden Delicious	24.25 ± 0.23	$3.46 \ge 10^3 \pm 582.48$
Gold3a	Golden Delicious	20.17 ± 0.31	$6.19 \text{ x } 10^4 \pm 13818.83$
GoldA	Golden Delicious	20.29 ± 0.14	$5.79 \text{ x } 10^4 \pm 5549.25$
Gold4	Golden Delicious	31.65 ± 0.65	$3.30 \ge 10^1 \pm 337.26$
Gold1	Golden Delicious	26.12 ± 0.43	$1.05 \text{ x } 10^3 \pm 337.26$
Mti2	Golden Delicious	36.36 ± 0.84	$7.58 \ge 10^2 \pm 359.66$
MtiA	Golden Delicious	26.11 ± 0.44	$3.21 \text{ x } 10^4 \pm 9293.85$
MtiB	Golden Delicious	21.51 ± 0.06	$7.04 \text{ x } 10^3 \pm 294.32$
MtiC	Golden Delicious	28.38 ± 0.07	$1.82 \text{ x } 10^2 \pm 0.39$

* Values are expressed as the mean \pm SD (n=9).

Sample ID	Cultivar	Ct mean ± SD*	V. inaequalis cells/µL ±SD*
As1	Fiorina	27.21 ± 0.42	$4.23 \times 10^2 \pm 56.27$
As2	Fiorina	28.86 ± 0.11	$1.30 \ge 10^2 \pm 10.25$
As3	Fiorina	27.68 ± 0.12	$2.98 \times 10^2 \pm 26.45$
As4	Ambrosia	25.56 ± 0.07	$1.31 \times 10^3 \pm 6.18$
As5	Ambrosia	23.81 ± 0.21	$4.50 \ge 10^3 \pm 175.73$
As6	Fiorina	26.20 ± 0.18	$8.40 \text{ x } 10^2 \pm 10.92$
As7	Fiorina	34.48 ± 0.07	$0.03 \text{ x } 10^2 \pm 0.22$
As8	Fiorina	33.18 ± 0.12	$0.06 \text{ x } 10^2 \pm 0.55$
As9	Fiorina	34.94 ± 0.13	$0.02 \text{ x } 10^2 \pm 0.29$
As10	Ambrosia	24.32 ± 0.16	$3.14 \times 10^3 \pm 157.47$
As11	Ambrosia	24.15 ± 0.16	$3.53 \times 10^3 \pm 7.29$
As12	Ambrosia	24.16 ± 0.15	$3.51 \times 10^3 \pm 15.75$
As13	Fiorina	33.54 ± 0.95	$0.06 \ge 10^2 \pm 1.12$
As14	Fiorina	28.89 ± 0.18	$1.28 \times 10^2 \pm 15.92$
As15	Golden Delicious	28.11 ± 0.17	$2.21 \text{ x } 10^2 \pm 25.84$
As16	Golden Delicious	29.87 ± 0.51	$6.65 \text{ x } 10^1 \pm 22.08$
As17	Fiorina	29.91 ± 0.33	$6.31 \ge 10^1 \pm 13.57$
As18	Ambrosia	28.91 ± 0.05	$1.26 \ge 10^2 \pm 33.85$
As19	Ambrosia	27.57 ± 0.14	$3.21 \times 10^2 \pm 30.04$
As20	Ambrosia	25.90 ± 0.20	$1.03 \times 10^3 \pm 43.97$
As21	Golden Delicious	26.72 ± 0.13	$5.82 \times 10^2 \pm 39.72$
As22	Golden Delicious	27.85 ± 0.07	$2.64 \times 10^2 \pm 13.54$
As23	Golden Delicious	26.17 ± 0.25	$8.61 \times 10^2 \pm 15.32$
As24	Golden Delicious	27.64 ± 0.07	$3.04 \text{ x } 10^2 \pm 15.71$
As25	Golden Delicious	26.60 ± 0.19	$6.34 \text{ x } 10^2 \pm 89.10$
As26	Ambrosia	25.55 ± 0.67	$1.43 \text{ x } 10^3 \pm 6.01$
As27	Golden Delicious	27.31 ± 0.19	$3.86 \times 10^2 \pm 51.03$
As28	Golden Delicious	28.00 ± 0.18	$2.39 \ x \ 10^2 \ \pm 29.63$
As29	Golden Delicious	30.81 ± 0.30	$3.37 \times 10^1 \pm 6.58$
As30	Golden Delicious	29.47 ± 0.21	$8.50 \ge 10^1 \pm 12.54$

Supplementary Table 4 – Ct mean values ± standard deviation and the estimated number of V.
 inaequalis cells/µL ± standard deviation for the asymptomatic Malus x domestica 'Fiorina',
 'Ambrosia' and 'Golden Delicious' leaves obtained using the TaqMan real-time PCR assay.

580 *	^c Values a	re expressed	as the mean	±SD (n=9).
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