

Hybridizations between formae speciales of Venturia inaequalis pave the way for a new biocontrol strategy to manage fungal plant pathogens

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13 Abstract

Hybridization and adaptation to new hosts are important mechanisms of fungal disease 14 15 emergence. Evaluating the risk of emergence of hybrids with enhanced virulence is then key 16 to develop sustainable crop disease management. We evaluated this risk in Venturia inaequalis, the fungus responsible for the common and serious scab disease on Rosaceae 17 hosts, including apple, pyracantha and loquat. Field isolates from these three hosts and 18 19 progenies obtained from five crosses between formae speciales isolates collected from 20 pyracantha (f.sp. pyracantha) and apple (f.sp. pomi) were tested for their pathogenicity on the three hosts. We confirmed a strict host specificity between isolates from apple and 21 pyracantha, and showed that most isolates were able to cause disease on loquat. None of the 22 23 251 progeny obtained from five crosses between V. inaequalis f.sp. pyracantha and V. inaequalis f. sp. pomi could infect apple. If confirmed on more crosses, the inability of the 24 25 hybrids to infect apple could lead to a novel biocontrol strategy based on a sexual hijacking of V. inaequalis f.sp. pomi by a massive introduction of V. inaequalis f.sp. pyracantha in apple 26 orchards. This strategy, analogous to the sterile insect approach, could lead to the collapse of 27 28 the population size of *V. inaequalis* and dramatically reduce the use of chemicals in orchards. 29

Keywords : Host specificity, Biological control, Crossing experiments, Scab, Apple, Firethorn,
 Eriobotrya, Sexual hijacking

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Plant-pathogenic fungi show variation in the width of their host range, from generalist pathogens that cause disease on multiple host species to specialized pathogens that infect a single host species. Fungal species strictly specialized to different hosts are referred to as *formae speciales*. Reproductive barriers between lineages – including *formae speciales* – Page **3** of **17**

specialized to different hosts may be incomplete, which may lead to hybridization (Le Gac and 37 Giraud 2008; Depotter et al. 2016). If hybrids are less fit than their parents on the same host, 38 39 they should ultimately disappear, but still may act as a "genetic bridge" allowing transfer of 40 pathogenicity traits from one species to another by backcrossing (Brasier 2001; Feurtey and Stukenbrock 2018; Stukenbrock et al. 2012). Hybridization can also result in extreme 41 phenotypes, a phenomenon referred to as transgressive segregation (Rieseberg et al. 1999). 42 43 Hybrid individuals can display increased virulence, as observed for hybrids between anther 44 smut pathogens Microbotryum lychnidis-dioicae and M. silenes-dioicae (Gibson et al. 2014), or display an expanded or new host range such as in the triticale pathogen Blumeria graminis 45 f.sp. triticale that was formed by hybridization between the wheat and rye pathogens B. 46 graminis f.sp. tritici and B. graminis f.sp. secalis (Menardo et al. 2016). As hybridization of 47 different formae speciales can be a driving force in the emergence of new diseases with 48 49 expanded host ranges we investigated the potential for this phenomenon to occur in the fungal pathogen Venturia inaequalis. 50

V. inaequalis is responsible for scab disease on apple (Malus x domestica), the most cultivated 51 52 fruit tree in temperate areas worldwide. Apple scab reduces both the quality and quantity of fruit. The control of the disease is mainly based on the use of fungicides. For a sustainable and 53 risk-free environmental control of apple scab, there is a need to develop the use of apple 54 55 resistant cultivars together with sanitation (Didelot et al. 2016), and to develop new control 56 methods, especially for biological control which lacks efficiency in orchards (Carisse and Rolland 2004; Kohl et al. 2015). V. inaequalis also causes scab disease on other Rosaceae 57 58 hosts, including pyracantha and loquat (Jones and Aldwinckle 1991; Sivanesan 1977). Loquat (Eriobotrya japonica Lindl.) is grown for fruit production in regions with subtropical climate, 59 60 while pyracantha (*Pyracantha* spp) is grown as an ornamental plant sometimes planted in

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hedges in an area extending from Southwest Europe to Southeast Asia. A strict host specificity 61 has been observed for isolates of V. inaequalis from pyracantha, which are able to cause 62 63 disease on pyracantha but not on apple (*i.e. V. inaequalis* f.sp. pyracantha), and isolates from apple, which are able to cause disease on apple but not on pyracantha (*i.e. V. inaequalis* f.sp. 64 pomi) (Le Cam et al. 2002). It was suggested that these two formae speciales are in "a process 65 of isolation due to host specialization" (Le Cam et al. 2002). The status of isolates from loquat 66 is less clear. Raabe and Gardner (1972) considered isolates from pyracantha and from loguat 67 68 as belonging to the same *forma specialis* because they were able to infect both pyracantha and loquat. Sanchez-Torres et al. (2009) showed that four isolates of V. inaequalis sampled on 69 apple were not able to cause disease on loquat. Gladieux et al. (2010) used six nuclear loci to 70 71 show that V. inaequalis populations from loquat and those from pyracantha have more 72 recently diverged than V. inaequalis populations from apple and those from pyracantha. 73 Recently, using whole genome sequencing data, Le Cam et al. (2019) confirmed that 74 populations of V. inaequalis isolated from apple, pyracantha and loquat represent distinct genetic lineages. 75

The aim of this study was to investigate the host specificity of *V. inaequalis* isolates from apple, pyracantha and loquat, and to test the hypothesis that crosses between *V. inaequalis formae speciales* represent a disease emergence risk by generating hybrids pathogenic on both hosts, or potentially with higher virulence on existing hosts.

80

A strict host specificity was observed on apple and pyracantha, while most *V. inaequalis* isolates were able to cause disease on loquat.

Thirty-two monoconidial isolates were recovered from scabbed leaves of apple (11 isolates, named *Pomi*), pyracantha (11 isolates, named *Pyr*) and loquat (10 isolates, named *Loq*), Page 5 of 17

sampled from different locations in nine, seven and seven different countries, respectively 85 (Table S1). These isolates were checked as being non-clonal using eight Simple Sequence 86 87 Repeat markers (SSRs) developed on V. inaequalis (Table S2; Guérin et al. 2004; Guérin et al. 2007; Tenzer et al. 1999). These isolates were inoculated individually on two apple cultivars 88 (Gala and Top Red), four pyracantha cultivars (Kasan, Orange Charmer, Red Column and Soleil 89 d'or), and two loquat cultivars (Peluche and Alfonso). These cultivars were selected for their 90 susceptibility to scab (Bertrand et al. 1992; Bus et al. 2011; Sanchez-Torres et al. 2009). Plants 91 92 were grown in a glasshouse in a scab-free environment, and transferred into a growth chamber for the tests of pathogenicity. Actively growing plants with expanding leaves which 93 are highly susceptible to V. inaequalis were used in six different experiments, because all 94 isolates could not be inoculated simultaneously across all hosts in the growth chamber. In 95 order to avoid cross-contamination during inoculation, plants were placed in polycarbonate 96 97 compartments and each compartment was inoculated with a single isolate. Each experiment included isolates of each origin and at least one cultivar of each host, except experiment 1 98 99 where no pyracantha was present and experiment 6 where no loquat was present. Each 100 isolate was grown on a cellophane sheet placed on malt-agar medium at 17°C to obtain spores 101 (Caffier et al. 2014). Spores were suspended in water by shaking the cellophane sheet. After 102 filtration through medical gauze, the spore suspension was adjusted to a final concentration of 2.5 x 10⁵ spores ml⁻¹ and was inoculated using a manual sprayer on three plants of each 103 104 cultivar. After inoculation, plants were kept in darkness under a plastic sheet to maintain leaf wetness. Climatic conditions in the growth chamber were set to 95% of relative humidity at 105 17°C for 48 hours. Afterwards, the plastic sheet was removed and the relative humidity was 106 107 reduced to 80% at day and 90% at night, with 12 hours of light per day. In order to evaluate

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the ability of each isolate to cause disease, the presence/absence of scab symptoms on each
plant was scored at 14, 21 and 28 days post inoculation.

110 Not all Pomi and Pyr isolates caused symptoms on their respective hosts, while all Log isolates 111 could infect both loquat cultivars (Figure 1). This result suggests the existence of resistance genes in the apple cultivar Top Red and in all four pyracantha cultivars. No Pyr isolate was able 112 to cause disease on the apple cultivars, and no Pomi isolate was able to cause disease on any 113 114 of the four pyracantha cultivars (Figure 1), which agrees with previous results and confirms 115 the status of formae speciales V. inaequalis f.sp. pomi and V. inaequalis f.sp. pyracantha (Le Cam et al. 2002). All Pyr isolates were able to cause disease on loguat, which agrees with data 116 from Raabe and Gardner (1972). However, the reciprocal was not true, as only one of the 10 117 Loq isolates was able to cause disease on pyracantha. Nine out of 11 Pomi isolates were able 118 119 to cause disease on the loquat cultivars Peluche and/or Alfonso. No Log isolate was pathogenic 120 on apple except isolate 2305 that was previously described as resulting from a one-off shift from apple to loquat, on the basis of its polymorphism at six nuclear loci (Gladieux et al. 2010). 121 To further investigate the pathogenicity of isolates on loquat and compare their 122 123 aggressiveness in relation to their host of origin, five isolates from each host were inoculated 124 on both loquat cultivars (three plants per cultivar) in the same experiment. Loquat plants were randomized within three blocks to have in each block one plant replicate for each isolate and 125 126 each cultivar. We performed a quantitative scoring through visual evaluation of the 127 percentage scabbed leaf area 21 days post inoculation, from no disease to 100% leaf area scabbed. The effect of the host of origin on the percentage scabbed leaf area was tested in R 128 129 4.0.3 (R Core Team 2020) using a linear mixed effect model with the LMER package (Kuznetsova et al. 2017). The 'isolate' factor was treated as a random factor nested within the 130 'host of origin' factor. The data were Log transformed prior to statistical analysis. This 131

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experiment showed that the *Pomi* isolates were significantly less aggressive on the loquat cultivars Peluche and Alfonso than the *Pyr* and *Loq* isolates (Figure 2). There was no significant difference in aggressiveness on loquat between Pyr and Loq isolates. This result agrees with the scenario proposed by Gladieux et al. (2010), where populations from loquat and pyracantha have diverged more recently than populations from apple and pyracantha.

137

Most progenies from a cross between f.sp. *pomi* and f.sp. *pyracantha* can infect pyracantha,
but none can infect apple.

As *V. inaequalis* f.sp. *pyracantha* and *V. inaequalis* f.sp. *pomi* isolates can be crossed *in vitro* (Le Cam et al. 2002), we tested the hypothesis that hybrids may have increased virulence on one of the hosts (apple or pyracantha) or that they may have an extended host range (able to cause disease on both apple and pyracantha).

144 We performed five different crosses, three crosses between one Pyr isolate (1669, 1381 and 1383) and one Pomi isolate (EU-B04 and 2556) and two crosses between one Pyr isolate (1387 145 and 2299) and a mixture of five Pomi isolates (104 + 2416 + 2429 + 2444 + 2557, Table S1, 146 Table 1), on sterile foliar disks of apple cultivar Gala, as described in Le Cam et al. (2002). The 147 use of a mixture of isolates enabled us to diversify the origin of the *Pomi* parent. Mature 148 pseudothecia were recovered from the leaf disks and crushed individually to obtain a 149 150 suspension of ascospores that was spread on malt (10g per liter) agar. After 24 to 48 hours of 151 incubation at 17°C, germinated ascospores were picked up individually with a needle under a stereomicroscope and transferred to new medium. To ensure the hybrid status of each 152 progeny, we performed a multilocus genotyping using eight SSRs (Table S2; Guérin et al. 2004; 153 154 Guérin et al. 2007; Tenzer et al. 1999; D. Sargent, unpublished data). Two hundred and fiftyPage 8 of 17

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one progeny, checked as being hybrids by SSRs, and the parental isolates were grown oncellophane sheets.

Spore suspensions from 37 progeny of the cross between 1669 and EU-BO4 isolates and from 157 both parents were adjusted to a final concentration of 2.5 x 10⁵ spores ml⁻¹ and inoculated in 158 a single experiment on three plants of pyracantha cultivar Kasan as described above. 159 160 Pyracantha plants were randomized within three blocks to have in each block one host 161 replicate for each isolate. At 15, 20, 23 and 27 days post inoculation, disease severity was 162 measured visually as the percentage of sporulating leaf area on the most scabbed leaf on each plant. These 37 progeny produced a large range of disease levels on the pyracantha cultivar 163 Kasan (Figure 3). No progeny was more aggressive than the Pyr parent 1669. Only one progeny 164 isolate was unable to cause disease on Kasan, similarly to the parental Pomi isolate. All other 165 166 progenies gave a disease level that was intermediate between the parental Pomi isolate EU-167 B04 (0% leaf area scabbed) and the parental Pyr isolate 1669 (90 % leaf area scabbed), suggesting a complex genetic determinism for pathogenicity on this host. This result differed 168 from previous studies in other pathosystems, for which a few genes explained host specificity. 169 170 For example, two to four genes may be responsible for the pathogenicity of the two sibling 171 species Ceratocystis manginecan and Ceratocystis fimbriata on Acacia mangum and Ipomoea batatas (Fourie et al. 2018; Fourie et al. 2019). Similarly, one to two genes have been shown 172 173 to be responsible for host specificity of Magnaporthe oryzae on foxtail millet in a cross 174 between an isolate pathogenic on wheat and an isolate pathogenic on foxtail millet (Murakami et al. 2003), and three genes have been shown to be responsible for host specificity of M. 175 176 oryzae on wheat in a cross between an isolate pathogenic on rice and an isolate pathogenic on wheat (Tosa et al. 2006). The loss of a single locus, AVR-Co39, was involved in the host shift 177 of *M. oryzae* from Setaria millet to rice (Couch et al. 2005). In the same way a single gene was 178

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shown to be involved in the host specificity of *Blumeria graminis* f.sp. *secalis* on wheat (Tosa1994).

181 The 251 progeny obtained from the five different crosses were inoculated on apple, either 182 four seedlings obtained from seeds collected from the apple cultivar Gala or six apple plants grafted on the rootstock MM106 (three Gala and three Golden Delicious) at a concentration 183 of 2 to 4 x 10⁵ spores ml⁻¹. The viability of each inoculum was checked by counting germinated 184 185 and non-germinated spores over a total of 100 spores for each isolate 24 hours after the 186 deposition of the suspension on malt agar incubated in darkness at 17°C. Only progeny with a concentration higher than 8 x 10⁴ viable spores ml⁻¹ were kept for analysis. Nine experiments 187 were needed to test all progeny. In each experiment, two parental isolates (a Pomi isolate and 188 a Pyr isolate) were used as control to check that the conditions were suitable for scab 189 190 infection. None of the 251 progeny were pathogenic on apple, *i.e.* all progeny behaved as the 191 Pyr parental isolates 1669, 1381, 1383, 1387 and 2299 on apple, whereas the Pomi isolate used as a control gave at least 45% and 80% leaf area scabbed (mean of the most infected leaf 192 of three plants for each cultivar), on Gala and Golden Delicious plants, respectively. This result 193 194 is consistent with the study of Tosa et al. (2006) on *M. oryzae*, which showed that none of the 195 94 progeny obtained from a cross between an isolate pathogenic on rice and an isolate pathogenic on wheat was virulent on rice. Comparative genomic analysis of these progeny 196 197 and parental isolates are needed to understand pathogenicity mechanisms and to identify 198 putative genes involved in specificity of V. inaequalis on apple and pyracantha.

199

200 Towards a new strategy of biocontrol against apple scab.

201 While hybridization between divergent populations or phytopathogenic species is often 202 depicted as a threat for agriculture because it can increase the host range of the pathogen or

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enhance its virulence, we showed that hybridization between *formae speciales* of *V. inaequalis*can be detrimental to fungal pathogens. If confirmed on more crosses, our finding that hybrids
are non-pathogenic on apple paves the way for a new biocontrol method to collapse pathogen
population size in apple orchards. Indeed, we propose to exploit this finding for biological
control based on a sexual hijacking strategy, similar to the sterile insect technique proposed
by Vanderplank (1944) to control the tsetse fly, which is largely developed now on many insect
species (Klassen and Curtis 2005).

As sexual reproduction is the main way for V. inaequalis f.sp. pomi to survive in winter in 210 temperate areas and as heterothallic mating is induced on dead leaves after leaf fall 211 212 (MacHardy 1996), a massive introduction in autumn of Pyr isolates representing the two mating types by spraying of spores on the senescent leaves before or just after leaf fall should 213 214 lead to the production of hybrid ascospores unable to cause disease on apple in spring. This strategy is expected to be efficient to reduce primary inoculum throughout most regions of 215 apple production, except maybe in regions with mild winter where V. inaequalis can survive 216 217 as asexual spores within buds (Holb et al. 2005; Passey et al. 2017). To maximize impact on 218 apple scab, this strategy may be associated with other methods, like sanitary measures and 219 use of resistant cultivars (Didelot et al. 2016).

Because the proposed biocontrol strategy is based on an organism which is widely distributed, that has not undergone genetic modification, and whose effect is highly specific, we do not expect any toxicity to humans or the environment, with the possible exception of pyracantha shrubs located in the immediate vicinity of orchards. The efficiency of this sexual hijacking strategy remains to be evaluated experimentally in orchards, and so does the risk of emergence of hybrids pathogenic on apple trees or other cultivated Rosaceae hosts. Since this Page **11** of **17**

- strategy could be implemented with other fungi having *formae speciales* and a sexual stage,
- 227 our work opens a new area of research in the biocontrol of plant pathogenic fungi.

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228 Patent for this novel strategy of biocontrol

- 229 Patent application PCT/FR2020/052580 « Méthode de bio-contrôle pour lutter contre la
- propagation des champignons et oomycètes phytopathogènes », jointly filed on 21/12/2020
- 231 by INRAE, Université d'Angers, Institut National Supérieur des Sciences Agronomiques,
- 232 Agroalimentaires, Horticoles et Paysage.

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349 Table 1. Description of the crosses between *Venturia inaequalis* f.sp. *pomi* (*Pomi*) and *V. inaequalis* f.sp.

350 *pyracantha* (*Pyr*) and pathogenicity of 251 progeny on apple

Pyr isolate	Pomi isolate	Number of progeny non pathogenic on apple / tested on apple
1669	EU-B04	76 / 76
1381	2556	44 / 44
1383	2556	44 / 44
1387	mixture of isolates (104, 2416, 2429, 2444, 2557)	49 / 49
2299	mixture of isolates (104, 2416, 2429, 2444, 2557)	38 / 38

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353 Legends of the figures

Figure 1. Pathogenicity of 32 isolates of *Venturia inaequalis* isolated from apple (*Pomi*), pyracantha (*Pyr*) or loquat (*Loq*) on two cultivars of apple, four cultivars of pyracantha and two cultivars of loquat. Each isolate was inoculated under controlled conditions on three plants of each cultivar. In red: presence of scab symptoms (with light red indicating less than 5% of the leaf area with scab symptoms), in blue: absence of scab symptoms, in white: not tested.

359 Figure 2. Boxplots of the percentage leaf area scabbed on two cultivars of loquat Alfonso and Peluche 21 days 360 after inoculation with isolates of Venturia inaequalis isolated from three hosts: apple (Pomi in dark grey), 361 pyracantha (Pyr in middle grey) and loquat (Loq in light grey). For each origin, five isolates were tested (Pomi: 362 190, 1395, 2315, 2367, EU-NL19; Pyr: 1669, 2267, 2269, 2299, 2309; Log: 2258, 2264, 2300, 2303, 2304). Each 363 isolate was inoculated under controlled conditions on three plants of each cultivar. The box represents the lower 364 and upper quartiles. The thick horizontal line represents the median. The whiskers represent the highest and 365 lowest values falling within 1.5 times the interquartile range. The black dots represent outliers. Pomi isolates are 366 significantly less aggressive than Pyr and Loq isolates (P = 0.0095 and P = 0.0164, respectively, linear mixed effect 367 model).

368 Figure 3. Percentage leaf area scabbed on the pyracantha cultivar Kasan after inoculation with 37 progeny of

the cross between the Venturia inaequalis f.sp. pomi isolate EU-B04 (Pomi isolate) and the V. inaequalis f.sp.

370 *pyracantha* isolate 1669 (*Pyr* isolate). The pathogenicity of the parents EU-B04 and 1669 are presented on the

371 left and right sides, respectively. Each isolate was inoculated under controlled conditions on three plants of

372 cultivar Kasan, and scoring was performed at four dates: 15, 20, 23 and 27 days post inoculation.

373

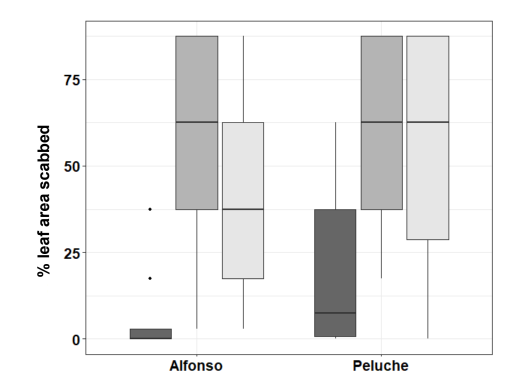
374 Titles of the e-Xtras

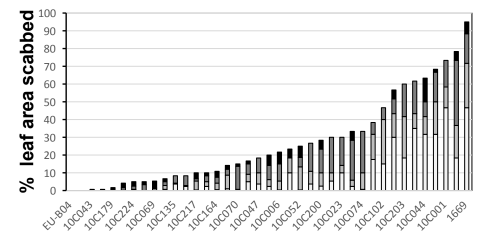
Table S1. Description of the origin of the *Venturia inaequalis* isolates used in the present study and indication of
their mating type

377 Table S2. Simple Sequence Repeats used to check the clone status of each sampled isolate of *Venturia inaequalis*

and the hybrid status of each progeny isolate

		Inoculated host Apple Pyracantha Loquat							
		Ар	pie	P	yrac	ann	a	a Loquat	
Is olate origin	ls olate name	Gala	Top Red	Kasan	Orange charmer	R ed column	Soleil d'or	Peluche	Alfonso
	190								
	301								
	912								
	1170								
	1395								
Pomi	2311								
	2315								
	2367								
	EU-B04								
	EU-D42a								
	EU-NL19								
	186								
	1381								
	1383								
	1387								
	1400								
Pyr	1669								
	2267								
	2269								
	2299								
	2307								
	2507								
	1389								
	2258								
	2260								
	2262								
Loq	2263								
Loq	2264								
	2300								
	2303								
	2304								
	2305								





🗆 mean 15dai 🗖 mean 20dai 🗖 mean 23 dai 🗖 mean 27 dai

1 Table S1. Description of the origin of the *Venturia inaequalis* isolates used in the present

2 study and indication of their mating type

Host of origin	Isolate	Sampling year	Sampling country	Sampling cultivar	Mating type
Apple	104	1978	France	Golden Delicious	a ²
(Pomi)	190	1987	France	Golden Delicious	a¹
	301	1988	Germany	81/19-53	α ²
	912	1992	Spain	unknown	α1
	1170	1995	The Netherlands	Discovery	a¹
	1395	1995	Colombia	Golden Dorsett	α1
	2311	2006	Algeria	Gala	α ¹
	2315	2006	USA	Gala	α ¹
	2367	2005	China	Fuji	a¹
	2416	2006	France	J25	a¹
	2429	2006	France	J25	a¹
	2444	2007	France	Gala	a¹
	2556	2006	France	J108	α ¹
	2557	2009	France	E063	a¹
	EU-B04	1998	Belgium	Golden Delicious Reinhardt	α ²
	EU-D42a	1998	Germany	Prima	a¹
	EU-NL19	1998	The Netherlands	Golden Delicious	α1
Pyracantha	186	1987	Ireland	Red Column	a ²
(Pyr)	1381	1998	France	unknown	a¹
	1383	1998	France	unknown	a¹
	1387	1998	France	unknown	α ¹
	1400	1995	Germany	unknown	a¹
	1669	2001	USA	unknown	a ²
	2267	2003	United Kingdom	Thornless	a¹
	2269	2003	Sweden	unknown	a ²
	2299	2004	France	unknown	α ¹
	2307	2004	Chile	unknown	a¹
	2507	2004	Chile	unknown	α ²
Loquat	1389	1999	France	unknown	a¹
(Loq)	2258	2006	Marocco	unknown	a¹
,	2260	2006	Portugal	unknown	α1
	2262	2004	France	unknown	a¹
	2263	2006	Spain	Algerie	α ²
	2264	2006	Spain	unknown	a¹
	2300	2006	Iran	unknown	a¹
	2303	2004	New Zealand	unknown	a¹
	2304	2004	New Zealand	unknown	a ¹
	2305	2003	Chile	unknown	a ¹

³

4 ¹ Mating types defined with primer sequences F: CCCCTCTGACTCTGAACAGC and R: TGTCGAAATCGTCACTCTGC for Mat a,

5 and primer sequences F: CACCTCTTTCCAGCAGAAGG and R: CGATCTGCAGGAACTTGTCA for Mat α

6 ² Mating types defined by genome sequencing (Le Cam et al., 2019)

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8 Table S2. Simple Sequence Repeats used to check the clone status of each sampled isolate of

9 Venturia inaequalis and the hybrid status of each progeny isolate

Primer name	Primer sequences (5' - 3')	Primer locus	Reference	Clone status	Hybric status
M1	F: [HEX]TCGAGATCCTCAAACTTCCTT R: TTTTAACTGTGCGGCCTG	1tc1a	Tenzer et al, 1999	х	Х
M2	F: [FAM]CGATTGGGGATATGAAGACTT R: TTAGTAATCAAATCGCACCCA	1tc1b	Tenzer et al, 1999	x	x
M4	F: [FAM]AGCGCTAGGTCGTGAAATC R: TTTCTGAAGTGTGTGGGACAT	1aac3b	Tenzer et al, 1999	х	
M15	F: [TAMRA]GCACCTGCTCTGTCTATCTC R: AAGGTTCAGGCACTGGAG	Vica9/152	Guerin et al., 2004	х	
M20	F: [TAMRA]TGTCAGCCACGCTAGAAG R: CACCGGACGAATCATGC	Vicacg8/42	Guerin et al., 2004	х	Х
M42	F: [TAMRA]CCAGACCTCCTTATTCAC R: TAACTCCTGAAGACGGCATG		Guerin et al., 2007	х	Х
M43	F: [HEX]GCCTGGTTGTGGATCTGTC R: ATCCTGCTACATCGACCTTC	Viga7/116	Guerin et al., 2004	х	Х
M51	F: [HEX]TCGCGCATCACTATCTACAC R: AGACAGGAATGTGGTGGAAG	Vica9/X	Guerin et al., 2004	х	
283036	F: [6- FAM]CAAGCTGAAAGGGCAAAGAG R: ATATGGGGCATTGGGAAACT		Sargent, unpublished		х
276191	F: [6- FAM]TGAGGGGAGAGATTTTGGTG R: ATGGTGGGGGCTTGACTAATG		Sargent, unpublished		x
139795541	F: [HEX]GTGGTTATGTTGTGGGAGTGG R: GTACTCTCTCGGCCTAAACTCG		Sargent, unpublished		x

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