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Use of Comparative Genomics-Based Markers for Discrimination of Host Specificity in *Fusarium oxysporum*

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ABSTRACT The polyphyletic nature of many formae speciales of Fusarium oxysporum prevents molecular identification of newly encountered strains based on conserved, vertically inherited genes. Alternative molecular detection methods that could replace labor- and time-intensive disease assays are therefore highly desired. Effectors are functional elements in the pathogen-host interaction and have been found to show very limited sequence diversity between strains of the same forma specialis, which makes them potential markers for host-specific pathogenicity. We therefore compared candidate effector genes extracted from 60 existing and 22 newly generated genome assemblies, specifically targeting strains affecting cucurbit plant species. Based on these candidate effector genes, a total of 18 PCR primer pairs were designed to discriminate between each of the seven Cucurbitaceaeaffecting formae speciales. When tested on a collection of strains encompassing different clonal lineages of these formae speciales, nonpathogenic strains, and strains of other formae speciales, they allowed clear recognition of the host range of each evaluated strain. Within Fusarium oxysporum f. sp. melonis more genetic variability exists than anticipated, resulting in three F. oxysporum f. sp. melonis marker patterns that partially overlapped with the cucurbit-infecting Fusarium oxysporum f. sp. cucumerinum, Fusarium oxysporum f. sp. niveum, Fusarium oxysporum f. sp. momordicae, and/or Fusarium oxysporum f. sp. lagenariae. For F. oxysporum f. sp. niveum, a multiplex TaqMan assay was evaluated and was shown to allow quantitative and specific detection of template DNA quantities as low as 2.5 pg. These results provide readyto-use marker sequences for the mentioned F. oxysporum pathogens. Additionally, the method can be applied to find markers distinguishing other host-specific forms of F. oxysporum.

IMPORTANCE Pathogenic strains of *Fusarium oxysporum* are differentiated into *formae speciales* based on their host range, which is normally restricted to only one or a few plant species. However, horizontal gene transfer between strains in the species complex has resulted in a polyphyletic origin of host specificity in many of these *formae speciales*. This hinders accurate and rapid pathogen detection through molecular methods. In our research, we compared the genomes of 88 strains of *F. oxysporum* with each other, specifically targeting virulence-related genes that are typically highly similar within each *forma specialis*. Using this approach, we identified marker sequences that allow the discrimination of *F. oxysporum* strains affecting various cucurbit plant species through different PCR-based methods.

KEYWORDS cucurbits, genome analysis, host range, molecular markers, pathogen detection, pathogenic fungi

A ccurate and rapid pathogen detection is necessary to take appropriate action against plant diseases. *Fusarium oxysporum* is a soilborne fungus that includes both nonpathogenic and plant-pathogenic strains. Pathogenic strains of *F. oxysporum*

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	Abbreviation in strain		
Forma specialis	designations	Host	Reference(s)
F. oxysporum f. sp. cucumerinum	Focuc	Cucumber (Cucumis sativus)	66
F. oxysporum f. sp. melonis	FomIn	Muskmelon (Cucumis melo)	67
F. oxysporum f. sp. niveum	Foniv	Watermelon (Citrullus lanatus)	68
F. oxysporum f. sp. radicis-cucumerinum	Forcu	Multiple cucurbits (including cucumber, melon, watermelon, and gourd)	69
F. oxysporum f. sp. momordicae	Fomom	Bitter gourd (Momordica charantia)	70
F. oxysporum f. sp. lagenariae	Folag	Calabash gourd (Lagenaria spp.)	32, 71
F. oxysporum f. sp. luffae	Foluf	Sponge gourd (Luffa cylindrica)	7

TABLE 1 Formae	speciales of F.	oxvsporum	affecting	members o	f the	Cucurbitaceae f	familv
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cause vascular wilt and cortical rot disease in a wide variety of agricultural crop species. They are classified into host-specific forms (*formae speciales*) and are often further subdivided into races based on their capacity to infect different cultivars of a plant species (1–3).

Fusarium wilt and root rot in cucurbits are among the most prominent and destructive diseases affecting this plant family (4–6). In total, seven cucurbit-infecting *formae speciales* have been described: *Fusarium oxysporum* f. sp. *cucumerinum, Fusarium oxysporum* f. sp. *niveum, Fusarium oxysporum* f. sp. *radicis-cucumerinum, Fusarium oxysporum* f. sp. *lagenariae, Fusarium oxysporum* f. sp. *momordicae*, and *Fusarium oxysporum* f. sp. *luffae* (Table 1). The last three are mostly restricted to Southeast Asia (7), while the *formae speciales* affecting cucumber, melon, and watermelon are globally distributed and more important from an economic standpoint (4, 7, 8).

Currently, there are no effective curative treatments for *Fusarium* disease (9). Use of resistant varieties or rootstocks is the only practical measure for controlling the disease in the field (10–12). In greenhouses, soil sterilization by fumigation with methyl bromide can be performed (10, 13). Most efforts are directed toward prevention of the disease. Routine methods that provide reliable subspecific identification, sensitive detection, and accurate quantification of *F. oxysporum* are of high importance (14) and could prevent unnecessary efforts to suppress harmless fungal populations (15). Development of these types of markers has thus far been complicated by the polyphyletic nature of most *formae speciales* of *F. oxysporum* (14).

As many *F. oxysporum* strains have been found to be nonpathogenic, endophytic, or even applicable as biocontrol strains (16–18), discrimination between pathogenic and abundantly present nonvirulent strains is very important (19). Discrimination of *F. oxysporum formae speciales* and races is routinely done through labor- and time-intensive disease assays (20–22). Molecular detection methods are therefore highly desired.

Formae speciales are often of polyphyletic origin (23), and pathogenic strains may share a higher level of sequence similarity of conserved genes with strains that are nonpathogenic or pathogenic toward another host (24, 25). Diagnostics based on genes like that encoding translation elongation factor 1-alpha ($EF1\alpha$) or the ribosomal intergenic spacer (IGS) are therefore only useful to discriminate between fungal species (26, 27). In several cases they have been suggested for subspecies discrimination, but these often prove to be unreliable for this purpose (8, 27, 28).

Several molecular markers for the cucurbit-infecting *F. oxysporum* f. sp. *cucumerinum*, *F. oxysporum* f. sp. *radicis-cucumerinum*, *F. oxysporum* f. sp. *niveum*, and *F. oxysporum* f. sp. *luffae* have been developed. These are all based on random amplified polymorphic DNA (RAPD) fragments, resulting in sequence-characterized amplified region (SCAR) markers. SCAR markers are suboptimal for *forma specialis* discrimination because they are based on genomic regions that are not necessarily required for virulence. Furthermore, as they can be localized anywhere on the genome, there are often little to no sequence data available in public databases for comparison with other sequences. The robustness of the markers can be verified only by screening against a large collection of strains (14).

Interestingly, closer inspection of previously developed *forma specialis*-distinguishing SCAR markers showed that the selected sequences were often (part of) a transposable element, such as *Fot1* (*Fusarium oxysporum* f. sp. *albedinis*, *Fusarium oxysporum* f. sp. *chrysanthemi*, and *Fusarium oxysporum* f. sp. *dianthi*) and *Folyt1* (*F. oxysporum* f. sp. *radicis-cucumerinum*) and *Impala* (*Fusarium oxysporum* f. sp. *ciceris* and *F. oxysporum* f. sp. *dianthi*) (29), or pathogenicity-associated genes like *FTF1* (*Fusarium oxysporum* f. sp. *phaseoli*) (30). A race 1-specific *Fusarium oxysporum* f. sp. *lactucae* marker was developed by amplifying and cloning regions between long terminal repeats of retrotransposons in the genome (31). For *F. oxysporum* f. sp. *lagenariae*, *F. oxysporum* f. sp. *momordicae*, and *F. oxysporum* f. sp. *melonis*, only DNA fingerprinting results have been described thus far (32).

It was recently shown that host specificity is associated with the suite of effector genes present in the genomes of F. oxysporum strains (33). Both presence-absence polymorphisms and the sequence type of individual effector genes turned out to be predictive for a strain's host range. These genes therefore form the most solid base for discrimination of formae speciales within the F. oxysporum species complex (FOSC) (14, 20, 25). Indeed, use of virulence genes to identify fungal plant pathogens has proven successful in the past for other Fusarium species (34, 35). Within the FOSC, this approach has been applied to distinguish Fusarium oxysporum f. sp. cubense tropical race 4 by targeting a candidate effector gene (36). Additionally, Fusarium oxysporum f. sp. lycoperici and F. oxysporum f. sp. cubense can be discriminated from other formae speciales through the use of PCR primers designed to detect specific secreted in xylem (SIX) effector gene sequences (15, 25, 37). At the time of these studies, however, no (or limited) comparative genomics analyses could be performed due to the lack of available genome sequences. All SIX genes have homologs in other host-pathogenic forms of F. oxysporum (e.g., SIX1, SIX5, and SIX6 [33, 38-40]). For these, marker specificity could not be evaluated beforehand and cross-reaction with nontarget formae speciales was found (25).

Since it is not yet viable to sequence every individual strain encountered, we decided to design effector candidate-based markers. In this way, we aimed to be able to distinguish cucurbit-affecting *formae speciales* from (i) each other, (ii) other *formae speciales*, and (iii) nonpathogenic strains. Therefore, we used whole-genome sequences of a number of representative cucurbit-infecting *F. oxysporum* strains as a starting point and identified putative effector genes suitable as markers. An advantage of using molecular markers over whole-genome sequencing is that they can also be applied to infected soil or plant tissue samples; the fungus does not need to be isolated and cultured (28). Techniques such as TaqMan real-time PCR even allow for a quantitative evaluation of pathogen abundance, e.g., on DNA isolated from soil (41).

The genetic bases for host specificity of FOSC strains toward plants belonging to the Cucurbitaceae family are similar (33), making these *formae speciales* relatively difficult to separate. This means that this is a good test case for host specificity discrimination, and the results presented here can be exemplary for application to other plant species where disease caused by *F. oxysporum* is a pressing problem.

RESULTS

Several cucurbit-infecting formae speciales have a polyphyletic origin. In order to be able to select forma specialis-wide marker sequences, it is necessary to collect the genetic variety for that forma specialis as completely as possible. We made use of 66 previously published genome sequences and added *de novo* genome assemblies generated from Illumina paired-end read data of 22 new strains (see Data S2 in the supplemental material).

In order to assess the genetic diversity of the *formae speciales* under investigation, we generated a phylogenetic tree based on over 400 core genomic gene sequences from each of their genomes (Fig. 1). This showed that *F. oxysporum* f. sp. *cucumerinum*, *F. oxysporum* f. sp. *melonis*, *F. oxysporum* f. sp. *niveum*, and *F. oxysporum* f. sp. *lagenariae* occupied multiple clades in the tree (5, 3, 3, and 3, respectively), indicating that they



FIG 1 *F. oxysporum* f. sp. *cucumerinum*, *F. oxysporum* f. sp. *melonis*, *F. oxysporum* f. sp. *niveum*, and *F. oxysporum* f. sp. *lagenariae* are of polyphyletic origin. A total of 441 conserved core genes from all genomes were extracted, aligned, and concatenated into

belong to different clonal lines. In our set of strains, we have 6 of 7 described *F. oxysporum* f. sp. *cucumerinum* vegetative compatibility groups (VCGs) (6, 42), 3 of 8 *F. oxysporum* f. sp. *melonis* VCGs (43), all 3 *F. oxysporum* f. sp. *niveum* VCGs (43), and both *F. oxysporum* f. sp. *radicis-cucumerinum* VCGs (43). For *F. oxysporum* f. sp. *lagenariae* (3 VCGs described [44]), *F. oxysporum* f. sp. *momordicae* (4 VCGs described [44]), and *F. oxysporum* f. sp. *luffae* (unknown number of VCGs), no VCG information was available for our strains, although they group into three, one, and one clade(s), respectively (Fig. 1).

Candidate effector gene phylogenies display clear grouping of host specificity. Unlike conserved core genes, virulence-related genes tend to be identical across members belonging to the same polyphyletic *forma specialis* of *F. oxysporum* (15, 33). For this reason, they have predictive value for a strain's host range. *Forma specialis* markers are essentially the smallest possible set of effector genes that is shared by all strains of a *forma specialis* and absent or different in sequence (at least as a set) in all other strains (33).

We extracted the sequences of the candidate effector genes from the work of van Dam et al. (33) from all assemblies and generated a multiple-sequence alignment (MSA; see Data S1 in the supplemental material) and phylogenetic tree for each of them (three examples in Fig. 2 and continued in Data S2). A custom python script identified those genes in which all members of a *forma specialis* grouped together in a separate clade. From the genes displaying such grouping, the genes that facilitated the best discrimination were selected based on manual inspection of the MSA to come to a final selection of marker sequences per *forma specialis* (Table 2).

Some of the selected genes show multiple *forma specialis*-specific clades; therefore, multiple markers targeting different *forma specialis* could be designed on these genes. An example is candidate effector 99, a hypothetical protein-encoding gene that is used as a marker for *F. oxysporum* f. sp. *niveum*, *F. oxysporum* f. sp. *lagenariae*, *F. oxysporum* f. sp. *cucumerinum*, and *F. oxysporum* f. sp. *luffae* (Fig. 2B). *F. oxysporum* f. sp. *melonis* strain FomIn010 possesses a copy identical to both candidate effector 99 homologs present in the *F. oxysporum* f. sp. *niveum* strains as well as a copy identical to the *F. oxysporum* f. sp. *lagenariae* gene sequence. To still be able to distinguish these *formae speciales* from one another, it is therefore of importance to use multiple markers for each *forma specialis*.

Discrimination of cucurbit-infecting formae speciales by PCR. PCR primers were designed specifically on polymorphic regions of the selected DNA sequences (Table 2; see also Data S1), aiming to generate a PCR product sized above 120 and below 700 nucleotides (nt) for quick and reliable application. The Fusarium extracellular matrix 1 gene (FEM1) (45) was taken along as a positive control. To verify the applicability of the markers, PCRs were executed for each of the primer pairs on a subset of the strains that were used for marker design, i.e., of which the host range has been confirmed and the genome had been sequenced. This included strains belonging to the cucurbit-infecting formae speciales, several other formae speciales (Fusarium oxysporum f. sp. vasinfectum, Arabidopsis infecting, F. oxysporum f. sp. lycopersici, Fusarium oxysporum f. sp. radicislycopersici, Fusarium oxysporum f. sp. nicotianae, Fusarium oxysporum f. sp. melongenae, Physalis infecting, F. oxysporum f. sp. cubense, Fusarium oxysporum f. sp. pisi, Fusarium oxysporum f. sp. tulipae, and Fusarium oxysporum f. sp. gladioli) and two nonpathogenic F. oxysporum strains: Fo47 (16) and MN14 (33). The strains were selected based on their differential phylogenetic distribution in Fig. 1 as well as the presence and absence of selected marker sequences in their genome assembly.

FIG 1 Legend (Continued)

a multiple-sequence alignment. Phylogeny was inferred with 100 bootstrap iterations. All strains fall within the three main clades of the FOSC. Focuc, *F. oxysporum* f. sp. *cucumerinum*; Fomln, *F. oxysporum* f. sp. *melonis*; Foniv, *F. oxysporum* f. sp. *niveum*; Forcu, *F. oxysporum* f. sp. *radicis-cucumerinum*; Folag, *F. oxysporum* f. sp. *lagenariae*; Foluf, *F. oxysporum* f. sp. *luffae*; Fomom, *F. oxysporum* f. sp. *momordicae*. For abbreviations of other *formae speciales*, see Data S1.



0.09

FIG 2 Phylogenetic trees of three genes selected as markers for *F. oxysporum* f. sp. *cucumerinum*: 66 (A), 99 (B), and 21 (C). Separation of a clade that includes all strains belonging to a *forma specialis* indicates sequence similarity within and sequence dissimilarity between *formae speciales*. Colored areas in the tree reflect the target *forma specialis* of the marker. Hypothetical protein-encoding genes 99 and 21 are used as markers for multiple *formae speciales*.

All except one of the *forma specialis*-specific PCR markers behaved like expected (Table 3), showing PCR products only in the expected combinations of genomic DNA and marker primers. One false-positive PCR product was found, in the combination of *F. oxysporum* f. sp. *pisi* HDV247 and marker 130 (*F. oxysporum* f. sp. *momordicae*). In the genome assembly of HDV247, this gene was found to be present with 97% sequence similarity, although the downstream region of this gene provided sufficient sequence diversity for primer design (Data S1).

Marker 94, targeting all cucurbit-affecting *formae speciales*, gave a band of the correct size for all cucurbit-affecting isolates tested, except FomIn010. Furthermore, strain FomIn010 displayed an atypical *F. oxysporum* f. sp. *melonis* marker pattern, as it yielded PCR products that were not seen in the other *F. oxysporum* f. sp. *melonis* isolates for *F. oxysporum* f. sp. *niveum* markers 99 and 100 and *F. oxysporum* f. sp. *lagenariae* marker 99. This pattern, designated pattern B in Table 3, was not unexpected, since presence of identical sequences for these three markers as well as absence of the gene encoding hypothetical protein 94 had been observed in the genome assembly of FomIn010 (sequence similarity of marker 99 shown in Fig. 2B). The presence of identical effector candidate sequences across *formae speciales* affecting similar plant species was

	TABLE 2 Selected	marker	aenes	and	their	respective	target	formae	speciales
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			Gene tree
Gene ID ^a	Target gene	Target forma specialis ^b	illustration
Positive control	FEM1	Positive control	
94	HPEG ^c	All cucurbit-infecting formae speciales	Data S2A
13	SIX13	radicis-cucumerinum	Data S2B
70	HPEG	radicis-cucumerinum	Data S2C
66	HPEG	cucumerinum	Fig. 2A
99	HPEG	cucumerinum	Fig. 2B
21	Fom effector 7	cucumerinum	Fig. 2C
1	SIX1	melonis	Data S2D
20	Fom effector 6	melonis	Data S2E
18	Fom effector 3	melonis + niveum	Data S2G
99	HPEG	niveum	Fig. 2B
100	HPEG	niveum	Data S2H
21	Fom effector 7	niveum	Fig. 2C
	(pseudogenized)		
98	HPEG	momordicae	Data S2I
130	HPEG	momordicae	Data S2J
1	SIX1	lagenariae + momordicae	Data S2D
71	HPEG	lagenariae	Data S2F
99	HPEG	lagenariae	Fig. 2B
99	HPEG	luffae	Fig. 2B

^aID, identifier.

^bAll formae speciales listed are F. oxysporum formae speciales; to save space, a shortened form of the

organism name is used.

 $^c\mbox{HPEG},$ hypothetical protein-encoding gene.

not surprising since they share part of their genetic toolset allowing for pathogenic colonization of these plants (33). It does, however, make marker selection more challenging. While screening for specific differentiation of, for instance, *F. oxysporum* f. sp. *melonis* and *F. oxysporum* f. sp. *niveum*, it is therefore important to check multiple markers.

Evaluating *forma specialis* **classification using markers.** After testing of the markers on sequenced strains to verify that they worked as anticipated, an extended set of strains originating from around the world (strain information in Data S1) was screened. Most strains were isolated from *Fusarium*-affected cucurbit plants and were described as one of the pathogenic forms listed in Table 1. The aim was to either confirm or reject their reported host specificity with our markers. A number of strains isolated from noncultivated soil samples was also taken along. The expectation was that these nonspecialized strains do not possess many effector genes and therefore would test negative for all of the 18 markers.

As can be seen from Table 4, marker analysis confirmed the reported *forma specialis* of most strains that were tested. However, some strains behaved differently than expected. For example, PCR products were identified for FomIn017, FomIn021, FomIn024, and FomIn026 for *cucumerinum* markers 66 and 21, as well as *F. oxysporum* f. sp. *momordicae/F. oxysporum* f. sp. *lagenariae* marker 1. Intriguingly, none of the *F. oxysporum* f. sp. *melonis* markers tested positive in these strains (marker pattern C in Table 4). Additionally, a third *F. oxysporum* f. sp. *melonis* pattern was observed with strain FomIn023 that was nearly identical to the pattern of FomIn010 (pattern B in Table 3). Finally, *F. oxysporum* f. sp. *melonis* marker 1 cross-reacted with Foniv041 genomic DNA, showing that this marker is not 100% specific for *F. oxysporum* f. sp. *melonis*.

Cucurbit marker 94 did not test positive for four individual *F. oxysporum* f. sp. *melonis* strains and one *F. oxysporum* f. sp. *momordicae* strain. However, while it does not detect all cucurbit-infecting strains, it did not result in false positives (Tables 3 and 4), meaning that it can still be used in addition to the other *forma specialis*-specific markers.

Several strains showed a marker pattern typically observed for another *forma specialis*, indicating that their reported host specificities might not be accurate. Strains Focuc014 and Focuc040, reported as *F. oxysporum* f. sp. *cucumerinum*, clearly showed

TABLE 3 PCR markers allowing discrimination of cucurbit-affecting formae speciales of F. oxysporuma

Strain	forma specialis	FEM (+)	94 cucurbits	13 Forcu	70 Forcu	66 Focuc	99 Focuc	21 Focuc	1 Fomln	20 Fomln	18 Fomln + Foniv	99 Foniv	100 Foniv	21 Foniv	98 Fomom	130 Fomom	1 Fomom + Folag	71 Folag	99 Folag	99 Foluf	Marker profile
Forcu016	radicis-cucumerinum	+	+	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum
Forcu031	radicis-cucumerinum	+	+	+		-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum
Focuc001	cucumerinum	+	+	-	-	+	+	+	-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum
Focuc013	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum
Focuc015	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum
Focuc018	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum
Focuc035	cucumerinum	+	+	-	-	+	+	+	-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum
Fomln001	melonis	+	+	-	-	-	-	-	+			-	-	-	-	-	-	-	-	-	melonis "A"
Fomln006	melonis	+	+	-	-	-	-	-	+			-	-	-	-	-	-	-	-	-	melonis "A"
Fomln010	melonis	+	-	-	-	-	-	-	+	+	+	+	+	-	-	-	-	-	+	-	melonis "B"
Foniv002	niveum	+	+	-	-	-	-	-	-	-	+			+	-	-	-	-	-	-	niveum
Foniv010	niveum	+	+	-	-	-	-	-	-	-	+			+	-	-	-	-	-	-	niveum
Foniv015	niveum	+	+	-	-	-	-	-	-	-	+			+	-	-	-	-	-	-	niveum
Foniv020	niveum	+	+	-	-	-	-	-	-	-	+	+	+	+	-	-	-	-	-	-	niveum
Fomom001	momordicae	+	+	-	-	-	-	-	-	-	-	-	-	-			+	-	-	-	momordicae
Fomom004	momordicae	+	+	-	-	-	-	-	-	-	-	-	-	-	+	+	+	-	-	-	momordicae
Folag001	lagenariae	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-			+	-	lagenariae
Folag004	lagenariae	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-	+	+	+	-	lagenariae
Foluf001	luffae	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-		luffae
Foluf002	luffae	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	+	luffae
Fovas 25433	vasinfectum	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Focon5176	conglutinans	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Folyc 4287	lycopersici	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Folyc MN25	lycopersici	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Folyc069	lycopersici	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Folyc072	lycopersici	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Forly CL57	radicis-lycopersici	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Fonic001	nicotianae	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Fomel001	melongenae	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Fophy KOD886	physali	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Focub II5	cubense	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Fopis HDV247	pisi	+	-	-	-	-	-	-	-	-	-	-	-	-	-	±	-	-	-	-	-
Fotul Tu67	tulipae	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Fogla G76	gladioli	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
Fo47	non-pathogenic	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
FoMN14	non-pathogenic	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
H2O	_	I -	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	_

"Symbols: +, positive test result; -, negative test result; ±, weak positive test result (very faint PCR product of the expected size present).

a positive result for both *F. oxysporum* f. sp. *radicis-cucumerinum* markers and an absence of all three *F. oxysporum* f. sp. *cucumerinum* markers, suggesting that they are in fact *F. oxysporum* f. sp. *radicis-cucumerinum* strains. Another interesting candidate was strain 14150, reportedly an isolate belonging to "*F. oxysporum* f. sp. *cucurbita-cearum*," a *forma specialis* proposed to encompass all *formae speciales* affecting cucurbits (46). This strain also showed the marker pattern typically observed for *F. oxysporum* f. sp. *radicis-cucumerinum*. Four strains (one reported as *F. oxysporum* f. sp. *cucumerinum* and three as *F. oxysporum* f. sp. *niveum*) displayed an absence of all 18 markers tested, while another strain (reported as *F. oxysporum* f. sp. *cucumerinum* tested positive only for *F. oxysporum* f. sp. *cucumerinum* marker 21, suggesting that they are not capable of infecting any of the cucurbit plants. As expected, each of the environmental strains tested negative for all of the markers.

Disease assays confirm marker predictions. The strains of which the reported forma specialis did not match the marker pattern were tested in a bioassay on susceptible cucurbit varieties to evaluate their actual host range (Table 4, rightmost column; Data S3). Strains Focuc014, Focuc040, and 14150 caused severe crown rot symptoms in both cucumber and melon, meaning that they are in fact *F. oxysporum*

TABLE 4 PCR testing of the markers on a set of 48 worldwide isolates for verification of their reported formae speciales

		(+)]	icurbits	orcu	orcu	ocuc	ocuc	ocuc	nln	nlm	omln + Foniv	oniv	oniv	niv	mom	Tomom	nom + Folag	olag	olag	oluf		
Strain	Reported f sp	EM	4 cu	3 F(0 Fe	6 F	9 F(1 Fe	Foi	0 Fc	8 F	9 F(100	1 Fe	8 F	30 I	Foi	1 Fe	9 F(9 F(Marker profile ^a	Bio-assay result ^b
Focuc014	cucumerinum	+	6	+	+	-	<u>_</u>	6	-	1	-	<u> </u>	-	1	<u> </u>	-	-	-	-	10	radicis-cucumerinum	radicis-cucumerinum
Focuc040	cucumerinum	+	+	+		-	_	-	-	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum	radicis-cucumerinum
Forcu005	radicis-cucumerinum	+	+	+		-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum	not tested
Forcu017	radicis-cucumerinum	+	+	+		-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum	not tested
Forcu020	radicis-cucumerinum	+	+	+		-	-	-	-	-	-	-	-	- 1	-	-	-	-	-	-	radicis-cucumerinum	not tested
Forcu028	radicis-cucumerinum	+	+	+		-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum	not tested
Forcu029	radicis-cucumerinum	+	+	+		-	-	-	- 1	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum	not tested
14150	cucurbitacearum	+	+	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	radicis-cucumerinum	radicis-cucumerinum
Focuc009	cucumerinum	+	+	-	-	+			- 1	-	-	-	-	-	-	-	-	-	-	-	cucumerinum	not tested
Focuc010	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum	not tested
Focuc016	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum	not tested
Focuc017	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum	not tested
Focuc026	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum	not tested
Focuc027	cucumerinum	+	+	-	-	+			- 1	-	-	-	-	-	-	-	-	-	-	-	cucumerinum	not tested
Focuc036	cucumerinum	+	+	-	-	+			-	-	-	-	-	-	-	-	-	-	-	-	cucumerinum	not tested
Focuc038	cucumerinum	+	+	-	-	+	+	+	-	-	-	÷.	-	-	-	-	-	-	-	-	cucumerinum	not tested
FomIn018	melonis	+	+	-	-	-	-	-				-	-	-	-	-	-	-	-	-	melonis "A"	not tested
FomIn019	melonis	+	+	-	-	-	-	-				-	-	-	-	-	-	-	-	-	melonis "A"	not tested
Fomin020	melonis	+	+	-	-	-	-	-				-	-	-	-	-	-	-	-	-	melonis "A"	not tested
Fomin027	melonis	+	+	-	-	-	-	-				-	-	-	-	-	-	-	-	-	melonis "A"	not tested
Fomin023	melonis	Ť	1	-	-	-	-	-				-	-	-	-	-	-	-	-	-	melonis A	meionis
Fomin002	melonis	- T +	-	-	-	-	-	-	T	- -		-	-	-	-	-	-	-	-	-	melonis A melonis "A"	non-pathogenic
FomIn023	melonis	+	1			-	-	-		+		+	+	Û.		-	-	-	+	Ê.	melonis "R"	melonis/niveum
FomIn024	melonis	+	+		2	+		+		-	<u>+</u>	-	-		2		+	1	-		melonis "C"	melonis
FomIn017	melonis	+	+		-				-	-	-		-	_	-	-		-	-	-	melonis "C"	melonis
Fomln021	melonis	+	+	-	-		-		-	-	-	-	-	-	-	-		-	-	-	melonis "C"	melonis
Fomln026	melonis	+	+	-	-		-		-	-	-	-	-	-	-	-		-	-	-	melonis "C"	melonis
Foniv011	niveum	+	+	-	-	-	-	-	-	-	+	+	+	+	-	-	-	-	-	-	niveum	not tested
Foniv017	niveum	+	+	-	-	-	-	-	-	-	+				-	-	-	-	-	-	niveum	not tested
Foniv018	niveum	+	+	-	-	-	-	-	-	-	+				-	-	-	-	-	-	niveum	not tested
Foniv033	niveum	+	+	-	-	-	-	-	-	-	+				-	-	-	-	-	-	niveum	not tested
Foniv039	niveum	+	+	-	-	-	-	-	-	-	+				-	-	-	-	-	-	niveum	not tested
Foniv040	niveum	+	+	-	-	-	-	-	- 1	-	+				-	-	-	-	-	-	niveum	not tested
Foniv041	niveum	+	+	-	-	-	-	-	±	-	+	+	+	+	-	-	-	-	-	-	niveum	not tested
Fomom002	momordicae	+	+	-	-	-	- 1	-	-	-	-	-	-	- 1	+		+	-	-	-	momordicae	not tested
Fomom003	momordicae	+	-	-	-	-	-	-	-	-	-	-	-	-	+	+	+	-	-	-	momordicae	not tested
Folag003	lagenariae	+	+	-		-	-	-	-	-	-	-	-	-	-	-				-	lagenariae	not tested
Folag006	lagenariae	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-				-	lagenariae	not tested
Folag007	lagenariae	+	+	-	-	-	-	-	-	-	-	-	-	-	-	-				-	lagenariae	not tested
Folag008	lagenariae	+	+	-	-	-	-1	-	-	-	-	-	-	-	-	-	+	+	+	-	lagenariae	not tested
Focuc022	cucumerinum	+	-	-	-	-	-	+	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	non-pathogenic
Focuc028	cucumerinum		-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	non-pathogenic
Foriv034	ninoum		1	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	non-pathogenic
Foniv034	niveum	+			-	-	-	-		-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	non-pathogenic
Foniv038	niveum	+	10			-	-		-		-		-					-	-	-	non-pathogenic	non-pathogenic
RBG1687	environmental	+	-	-	-	-	-	-	-	-	-	_	-	-	-	-	-	-	_	-	non-pathogenic	not tested
RBG1693	environmental	+		-	-	-	-	_	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
RBG5713	environmental	+		-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
RBG5786	environmental	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
RBG5789	environmental	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
RBG5791	environmental	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
RBG5798	environmental	+	1.	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
RBG5820	environmental	+	-	-	-	-	-	-	-	-	-	-	-	- 1	-	-	-	-	-	-	non-pathogenic	not tested
RBG5824	environmental	+	-	-	-	-	- 1	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
RBG5827	environmental	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	non-pathogenic	not tested
HaO		1																				

^a"Non-pathogenic" means not pathogenic toward any of the seven hosts listed in Table 1.

b"Non-pathogenic" means no symptom development in susceptible plant hosts of the originally reported forma specialis.

f. sp. radicis-cucumerinum strains, as predicted by our PCR analysis. The strains that were predicted to be nonpathogenic based on their marker patterns indeed did not cause symptom development when tested on susceptible cucumber (Focuc028 and Focuc039) or watermelon (Foniv034, Foniv035, and Foniv038) plants. Strains FomIn017, FomIn021, FomIn024, and FomIn026 (profile C) as well as FomIn023 (profile B) were all

able to cause disease in susceptible musk melon plants, even though their marker pattern was different from the most common profile in our set of isolates (profile A [Tables 3 and 4]). Fomln023, which tested positive for two of the three *F. oxysporum* f. sp. *niveum* markers, was also tested on susceptible watermelon plants. This strain was found to also be capable of causing disease in these plants, whereas Fomln010, with an almost identical marker pattern, was not (33). Fomln002 did not cause symptoms in susceptible melon plants, showing that possessing effector gene sequences alone is not always sufficient for pathogenicity and false positives may show up.

The fact that the bioassay data confirmed the suspected *forma specialis* predicted by the reported markers indicates that they provide a robust tool for identifying whether an isolate indeed belongs to the suspected *forma specialis* or not. PCR cross-reaction between *F. oxysporum* f. sp. *cucumerinum*, *F. oxysporum* f. sp. *melonis*, and *F. oxysporum* f. sp. *niveum* markers and the cross-pathogenicity of strain FomIn023 suggest a shared evolutionary origin of the *formae speciales* affecting cucumber, melon, and watermelon.

Specific detection of F. oxysporum f. sp. niveum using a TaqMan assay. TaqMan real-time PCR has added benefits over traditional PCR: samples can easily be multiplexed, the fluorescent probe provides additional sequence specificity, and the technique allows for quantification of a target DNA sequence, for example, on DNA isolated from soil or diseased plant tissue. A TagMan experiment was conducted using two of the marker genes in this study, F. oxysporum f. sp. niveum markers 21 and 100. These markers showed good specificity and displayed no cross-reaction with nontarget strains in the PCRs (Tables 3 and 4). TaqMan-specific primers and probes were designed in such a way that 116- and 138-bp F. oxysporum f. sp. niveum-specific amplicons were formed, respectively. As a fluorescent dye, hexachlorofluorescein (HEX; $\lambda_{emission} = 556$ nm), was used for marker 21 and 6-carboxyfluorescein (FAM; $\lambda_{emission} = 518$ nm) was used for marker 100. As an internal control for sample/DNA quality that would allow for normalization of the tested markers during multiplexing experiments, a set of primers and a probe with a different fluorescent dye (6-carboxytetramethylrhodamine [TAMRA]; $\lambda_{\text{emission}} = 580$ nm) was designed on a region of EF1 α conserved in all F. oxysporum strains. To test the efficiency of the primers and probe sets, a dilution series of F. oxysporum genomic DNA was made and used as the template in a TaqMan assay.

A linear relationship was found between Foniv002 genomic DNA concentration and real-time quantification cycles (Data S4; $R^2_{marker21} = 0.999$; $R^2_{marker100} = 0.998$; $R^2_{EF1\alpha} = 0.999$). The pathogen could be detected at template concentrations as low as ~ 2.5 pg (Data S4).

The TaqMan assay was performed on isolates for which marker genes 21 and 100 were identified in the genome assembly (Fig. 2C; see also Data S2H). Each sequence type was included, with the addition of strains of *Fusarium proliferatum* and a *Fusarium* sp. that were identified to have candidate effector 100 in a recent study (47). *F. oxysporum* f. sp. *lycopersici* 4287, *F. oxysporum* f. sp. *cubense* II5, and biocontrol strain Fo47 were included as negative controls, since these do not have either of the marker genes. No cross-reactions were found, except in the case of FomIn010, which possesses a gene sequence identical to marker 100 in *F. oxysporum* f. sp. *niveum* isolates (Fig. 3). These results show the applicability of the TaqMan assay for specific detection of *F. oxysporum* f. sp. *niveum* DNA in very small quantities.

DISCUSSION

In the current study, we tried to make use of comparative genomics to design robust markers based on candidate effector genes. Effectors are functional elements in the pathogen-host interaction and have been found to show very limited sequence diversity between members of the same *forma specialis* (48, 49). This means that they form ideal targets for marker design (20). Effector gene sequences are often different between *formae speciales*, although several cases of identical gene sequences have been found in a previous study in our lab (33). For example, the *SIX6* and *SIX11* homologs present in some isolates belonging to *F. oxysporum* f. sp. *niveum*, *F. oxysporum* f. sp. *melonis*, and *F. oxysporum* f. sp. *radicis-cucumerinum* have 100% nucleotide



FIG 3 TaqMan primer-probe combinations show amplification of *F. oxysporum* f. sp. *niveum* DNA (Foniv002) when markers 21 (A) and 100 (B) are tested in duplex with *EF1* α . No amplification of these markers was detected in any of the non-*F. oxysporum* f. sp. *niveum* strains, with the exception of FomIn010, which has an identical gene sequence for hypothetical protein-encoding gene 100. High threshold cycle (C_7) values (\geq 35 cycles) under the detection threshold in the water control of marker 21 are probably caused by primer-dimer formation. Four technical replicates were used per sample, each represented by a bar.

identity. These sequences can therefore not be used for differentiation of these *formae speciales*. They do, however, give insight into the evolutionary history of pathogenicity of *F. oxysporum* toward cucurbits; the presence of sequences that are completely identical between relatively distantly related strains implies recent horizontal transfer of genetic material.

The benefit of using comparative genomics for marker design is that the specificity of the designed markers can directly be evaluated in other genome assemblies (as opposed to RAPD-derived marker sequences). Within the FOSC, one study has reported the use of comparative genomics for *forma specialis* marker development. This resulted in markers based on unique (random) sequences distinguishing *F. oxysporum* f. sp. *conglutinans* from 19 other *formae speciales* of *F. oxysporum* (50).

Our goal was to differentiate between *formae speciales* affecting the Cucurbitaceae family. The respective hosts are highly similar to each other, and incidental cross-pathogenicity between these *formae speciales* has been described (51–53). We designed a set of 18 primer pairs aiming to discriminate seven cucurbit-infecting *formae speciales* from each other as well as from other host-specific forms and nonpathogenic strains of *F. oxysporum*. We found that for *F. oxysporum* f. sp. *cucumerinum*, *F. oxysporum* f. sp. *lagenariae*,

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F. oxysporum f. sp. *momordicae*, and *F. oxysporum* f. sp. *luffae*, the marker sets allowed clear recognition of the host range of each evaluated strain. Marker 94, designed on a gene encoding a hypothetical protein present in all cucurbit-infecting *formae speciales*, was positive for all target strains, with the exception of several *F. oxysporum* f. sp. *melonis* strains and one *F. oxysporum* f. sp. *momordicae* strain. This gene was not identified in the genome sequence of FomIn010.

Within F. oxysporum f. sp. melonis strains, more genetic variability exists than what had been taken into account as a starting point used for marker design (the 10 F. oxysporum f. sp. melonis strains with a sequenced genome). Indeed, F. oxysporum f. sp. melonis has been described as a highly heterogeneous forma specialis, encompassing at least eight VCGs (33, 43, 54). Several F. oxysporum f. sp. melonis strains showed overlap in their effector gene contents with cucurbit-infecting F. oxysporum f. sp. cucumerinum, F. oxysporum f. sp. niveum, F. oxysporum f. sp. momordicae, and/or F. oxysporum f. sp. lagenariae (Table 4). So far, no SCAR or other marker sequences have been reported for F. oxysporum f. sp. melonis, possibly due to its heterogeneous nature. Two marker patterns were observed that were different from the marker patterns found in the majority of our F. oxysporum f. sp. melonis strains. Two strains (FomIn010 and FomIn023) tested positive for F. oxysporum f. sp. melonis as well as F. oxysporum f. sp. niveum markers (pattern B). Interestingly, FomIn023 was capable of causing severe wilting symptoms both in melon and in watermelon, while FomIn010 was not (33). This raises the question of whether the separation of these formae speciales is justified, similar to the question of whether strains pathogenic toward both cucumber and melon should be regarded as F. oxysporum f. sp. cucumerinum or F. oxysporum f. sp. melonis. Cafri et al. (53) decided in their study that since the F. oxysporum f. sp. cucumerinum strains they tested were more aggressive toward cucumber than melon and no cross-pathogenicity was found the other way around, these formae speciales should indeed remain distinct. In the case of strain FomIn023 in the current study, disease severities were comparable between watermelon and melon plants, indicating that this strain is a "bridging" forma specialis, and its marker gene pattern reflects this. Isolates bridging multiple host species are not commonly described in the literature, although most isolates admittedly are not tested against a large variety of plant species to confirm host specificity. It would be interesting to compare the genomes of strains with a wider host range with those that are highly specific to one plant species, which may have implications for the current nomenclature system within the FOSC.

We recently demonstrated that clustering isolates based on patterns of presence or absence of candidate effector genes divided F. oxysporum f. sp. cucumerinum into two groups, separated from each other by F. oxysporum f. sp. melonis and F. oxysporum f. sp. niveum strains (33). The cucurbit-infecting isolates formed a supercluster from other formae speciales, indicating that they share a significant number of effector genes between them. Not much is known regarding the evolution of host specificity of F. oxysporum toward cucurbits, but FomIn023 might contain accessory genetic material originating from both an F. oxysporum f. sp. niveum and an F. oxysporum f. sp. melonis strain. Likewise, strains FomIn017, FomIn021, FomIn024, and FomIn026 tested positive for two F. oxysporum f. sp. cucumerinum markers and the F. oxysporum f. sp. melonis markers used all tested negative (pattern C). This indicates that these F. oxysporum f. sp. melonis and F. oxysporum f. sp. cucumerinum strains also share accessory genetic material. However, FomIn017 is, like most F. oxysporum f. sp. melonis isolates, highly specific to melon plants. It would be interesting to further investigate these strains, for example, through long-read sequencing of their genomes and analysis of their pathogenicity chromosome(s) compared to those of other F. oxysporum f. sp. cucumerinum, F. oxysporum f. sp. melonis, and F. oxysporum f. sp. niveum strains. This could shed light on how pathogenicity toward cucurbits has evolved in the FOSC.

Horizontal gene and chromosome transfer has been described as an important contributor to genetic diversity and the generation of new (pathogenic) clonal lines in fungi (55, 56). The different effector sequences and presence/absence patterns between and even within some cucurbit-infecting *formae speciales* suggest that it is

possible that multiple horizontal transfers of accessory genome material have taken place in the evolutionary trajectory, resulting in pathogenicity of *F. oxysporum* toward cucurbits. This is in contrast to the case with *F. oxysporum* f. sp. *lycopersici*: the four clonal lines that were tested in the work of van Dam et al. all have nearly identical sets of effectors and effector gene sequences (25, 33).

Minor cross-reaction (a much lighter band) was found with one of the markers (*F. oxysporum* f. sp. *momordicae* marker 130) with an unrelated *forma specialis*. *F. oxysporum* f. sp. *pisi* HDV247 (as well as *F. oxysporum* f. sp. *raphani* PHW815) indeed possesses this gene, although its downstream flank on which the reverse primer (FP7336) was designed was deemed to be sufficiently different from the copy in *F. oxysporum* f. sp. *momordicae*; only the four 5' nucleotides matched between these two sequences. The forward primer contained only a single nucleotide polymorphism (SNP), meaning that it probably binds in the nontarget sequence of *F. oxysporum* f. sp. *pisi*, too. A similar observation was made for *F. oxysporum* f. sp. *melonis* marker 1 and Foniv041 genomic DNA. For this isolate, however, no genome sequence is available. Through quantitative PCR techniques such as TaqMan, (cross-)reactions with a significantly smaller amount of product can probably be distinguished from genuine positives.

As a proof of concept, a TaqMan test was developed for two of the markers. The TaqMan real-time PCR technique has several advantages over traditional PCR. Since it makes use of a sequence-specific fluorescently tagged probe in addition to the primer sequences, marker specificity is potentially higher. Additionally, the technique allows for quantification of the targeted DNA sequence (and thus of the pathogen in soil or infected plant tissue). Quantification of pathogenic *F. oxysporum* propagules in soil, seeds, or plant tissues may aid in deciding if and when to take action. Also, it is possible to test multiple markers by multiplexing, using several different fluorescent dyes at once (57–59). The markers that were tested in duplex for *F. oxysporum* f. sp. *niveum* behaved like expected: no amplification was identified in other strains (except FomIn010 with marker 100), even those that do possess the target gene. The technique allows for identification of sequences slightly different from the target sequence; the cycle number of a single copy marker with SNPs would be distinguishably higher than that of a positive-control single-copy gene like *EF1* α .

These findings illustrate the hurdles that can be experienced in the process of designing *forma specialis*-specific markers based on candidate effector genes, specifically if the *formae speciales* infect members of the same plant family and possibly arose through a shared and recent evolutionary history. Nonetheless, the combination of marker sequences described here can be used with relatively high fidelity to discriminate the seven cucurbit-affecting *formae speciales*, particularly when multiple markers are tested simultaneously in the analysis. It is possible—perhaps even likely—that more diversity exists among the seven *formae speciales* targeted in this study, since for several of the *formae speciales* not all VCGs were sampled for genome sequencing due to unavailability of these strains. This means that the markers might require revision in the future. The availability of more whole-genome sequences like the ones generated in this study will allow easier marker design and comparison in the future.

MATERIALS AND METHODS

Fungal strains and accession numbers. An overview of the strains that were used in this study and their respective genome assembly or nucleotide sequence accession numbers can be found in Table 5.

Whole-genome sequencing and *de novo* **assembly.** *F. oxysporum* genomic DNA was isolated through phenol-chloroform extraction from freeze-dried mycelium that was harvested from 5-day-old NO₃ medium (0.17% yeast nitrogen base, 3% sucrose, 100 mM KNO₃) cultures as described in detail in reference 33. Library preparation of insert size 550 bp and Illumina HiSeq 2500 paired-end sequencing were performed at Keygene N.V. (Wageningen, Netherlands).

Sequencing reads were trimmed for quality and to remove adapter sequences with FastqMcf v1.04.676 (https://github.com/ExpressionAnalysis/ea-utils/blob/wiki/FastqMcf.md; quality threshold = 20). *De novo* assemblies were generated using CLC-workbench 8.0. Default settings were used, except "minimum contig length = 500."

For generating a core phylogeny, homologs of 15,956 Fol4287 core genes (including introns) were searched in all genomes using BLASTN with default parameters. We selected all sequences that overlapped >70% with the query sequence and with more than 80% identity to the query. We then

TABLE 5 Overview of fungal strains used in this study and their NCBI genome accession numbers

	Original	Forma specialis or				Source or	GenBank assembly
Strain	designation	description	VCG	Race	Origin of strain	reference ^a	accession no.
Folyc 4287	NRRI 34936	lycopersici	0030	2	Snain	Broad Institute	GCA 000149955 2
Focon 5176	F05176	On Brassica	0050	2	Australia	Broad Institute	GCA_000222805.1
Folyc MN25	NRRI 54003	lycopersici	0033	з		Broad Institute	GCA_000259975.2
Fonis HDV247	NIRRI 37622	nisi	0055	5	05/(Broad Institute	GCA 000260075 2
Forly CL 57	NRRI 26381	radicis-lycopersici	0094		USA (Florida)	Broad Institute	GCA_000260155.3
Fovas 25433	NRRI 25433	vasinfectum	0004		China	Broad Institute	GCA_000260175.2
Focub II-5	NRRI 54006	cubense	01213	TR4	Indonesia	Broad Institute	GCA_000260195.2
Focon PHW808	NRRI 54008	conclutinans	0101	2	maonesia	Broad Institute	GCA_000260215.2
Foran PHW815	NRRI 54005	raphani	0102	2		Broad Institute	GCA_000260235.2
FomIn001	NRRI 26406	melonis	0136	1	Mexico	Broad Institute	GCA_000260495.2
Fo47	NRRI 54002	Nonpathogen biocontrol	0150	·	France	Broad Institute	GCA_000271705.2
Forli FOSC 3-a	NRRI 32931	Clinical isolate from human			USA (Massachusetts)	Broad Institute	GCA_000271745.2
l och i obe b u		blood			05/1 (Massachasetts)	broad institute	00/20002/17/15/2
Focub N2	N2	cubense		1	China	72	GCA 000350345 1
Focub R2	R2	cubense		4	China	72	GCA 000350365 1
Focuc013	000 <i>4</i> _1	cucumerinum	0186	7	China	72 22	MARI0100000
Focuc015	9906-3	cucumerinum	0184		China	22	MABK01000000
Focuc021	ATCC 16416	cucumerinum	0180		LISA (Florida)	22	MABL 01000000
Focuc018	$\Lambda f_{11}=50(R)$	cucumerinum	0180		Croto, Grooco	22	MABL0100000
Focuc030		cucumerinum	0180		Israel	22	MABN01000000
Focuc035	NETH 11170	cucumerinum	0100		Nothorlands	22	MABO01000000
Focuc037	Tf_212	cucumerinum	0185		lanan	22	MARD01000000
Forcu016	22	radicis-cucumarinum	0105		Canada	22	MABOO200000
Forcu024	Δfu 11(Λ)	radicis cucumorinum	0200		Croto Grooco	22	MARDQ02000000
Forcu024	AIU-II(A) AK_2	radicis-cucumerinum	0200		Crete, Greece	22	MARS0100000
Focuc011	0003-1	cucumarinum	0201		China	22	MABT01000000
Forde011	5905-1 Eom 0122	malonis	0124	1	Spain	72	MAKV0100000
Forminious Forme001	For 1	riteionis	0134	1	Japan	75 D I	MAKT01000000
Focución Fomin004	FOC-1 Fom 0122	malonis	0105	0	Spain	D.L. 72	MAL 40100000
Fomin004	Fom 0122	malonis	0124	2	Spain	75	MALA01000000
Fomini000	F0111 0124	malonis	0125	2	Jerzol	75	MALC0100000
Fomin009		melonis	0155	2 1	Israel	73	MALC01000000
FomIn012	ML 2	melonis	0134	0	131001	73	MALE01000000
Fomin012	Fom26	melonis	0134	1		73	MALE01000000
FomIn013	101120	melonis	0134	י כ	Spain	73	MAL C0100000
Folyc004	IDO1530/B1	lycoparsici	0030	2 1	Nothorlands	73	MALH01000000
Folyc007	101020/01	lycopersici	0030	י כ	France	74 74	MAL 101000000
Folyc014	1511-3	lycopersici	0030	1	LISA (Louisiana)	74	MAL 101000000
Folyc026	RRIP 14844 (M1943)	lycopersici	0030	2	Australia	74	MALK01000000
Folyc018		lycopersici	0030	2	LISA (Louisiana)	74	MALL0100000
Folyc016	BEOL-51	lycopersici	0031	1	USA (Louisiana)	74	MAL M01000000
Folyc029	5307	lycopersici	0030	2	USA (Elorida)	74	MALN0100000
Folyc038	CA92/95	lycopersici	0030	3	USA (California)	25	MAL 001000000
Folyc069	DF0-23	lycopersici	0035	2	USA (California)	75	MAL P01000000
Folyc072	DF0-38	lycopersici	0031	2	USA (California)	75	MAL 001000000
Folyc073	DF0-40	lycopersici	0030	2	USA (California)	75	MAL R01000000
Folyc074	DF0-41	lycopersici	0030	3	USA (California)	75	MAL S01000000
Folyc075	DF0-62	lycopersici	0031	2	USA (California)	75	MALT01000000
FoMN14	MN-14	Nonpathogen from tomato	0051	2	USA (California)	76	MALLI01000000
		nlant			obri (camorna)	70	
Foniv002	CBS 418 90	niveum			Israel	22	MALX0100000
Foniv005	TY_471_1	niveum	0080	0	IISA (Toyac)	51	MAL X01000000
Foniv010	F-016-1	niveum	0000	1	USA (Maryland)	51	MAL 701000000
Foniv013	F_014_2	niveum	0002	2	USA (Maryland)	51	MAMA0100000
Foniv015	F-063-1	niveum	0002	2	USA (Maryland)	51	MAMR01000000
Foniv010	TX-X1D	niveum	0002	2	USA (Tevec)	51	MAMC0100000
Foniv020	F_000_1	niveum	0082	∠ 2	USA (Texas)	51	MAMD0100000
Foniv020	MD-7F622	niveum	0005	∠ ג	USA (Maryland)	51	MAME0100000
Foniv037	NRRI 38530	niveum		2	Israel	77	MAME0100000
Folyc002	WCS862/F241	lyconersici	0020	2	Netherlands	74	MAMG0100000
FomIn011	WCJ002/ L241	melonis	0050	0	Israel	73	MAMH0100000
Fogla G14	G14	aladioli	03/1	U	Netherlands	78	NICM01000000
Fogla G2	62	aladioli	0340		France	78	NICI 0100000
. 3914 02		3.301011	01-00		unice		

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TABLE 5 (Continued)

	Original	Forma specialis or				Source or	GenBank assembly
Strain	designation	description	VCG	Race	Origin of strain	reference ^a	accession no.
Fogla G76	G76	gladioli	0343		Italy	78	NJCK0100000
Folag001	01-03008	lagenariae			Japan	32	NJCJ0100000
Folag002	03-05118	lagenariae			Japan	32	NJCI0100000
Folag004	Lag:3-1 (JCM9293)	lagenariae			Japan	32	NJCH01000000
Folag005	Lag:1-1	lagenariae			Japan	32	NJCG01000000
Folil Fol39	Fol39	lilii			Netherlands	79; J.V.D.	NJCF01000000
Foluf001	Fol-114	luffae			Taiwan	25	NJCE0100000
Foluf002	Fol-167	luffae			Taiwan	25	NJCD01000000
Fomel001	J-71	melonaenae				IPO	NJCC01000000
Fomom001	NRRI 26413	momordicae			Taiwan	ARS	NICB0100000
Fomom004	90NF2-1 (ICM9292)	momordicae			lanan	32	NICA0100000
Fonar Na5	Na5	narcissi	2		Netherlands	79. IV D	NICV01000000
Fonic001	FON-1	nicotianae	2		USA (Connecticut)	80	NIR701000000
Fonic002	10012	nicotianae	0272		USA (Connecticut)	80 80	NJRV0100000
Fonic005	TU915	nicotianae	0373		USA (Marth Carolina)	80	
	FL-ROD	nicolianae	0576		USA (North Carolina)	80 80	
FONICUT2	Ft-1512	nicotianae				80	NJC00100000
Fophy KOD886	KOD886	physali			USA (California)	К.О.	NJBW0100000
Fophy KOD887	KOD887	physali			USA (California)	К.О.	NJBV01000000
Fophy KOD888	KOD888	physali			USA (California)	К.О.	NJBU01000000
Fo Tu58	Tu58	Nonpathogenic, from			Netherlands	79; J.V.D.	NJBT01000000
Fatul Turcz	T. (7	symptomatic tulip)			N ath a vlavada		
	14150	tunpae			Nethenlands	79; J.V.D.	
14150	14150	(redesignated			Netherlands	NAKI	Not available
F. a	0020	radicis-cucumerinum)	0107		China	22	
FOCUCUU9	0020	cucumennum	0187		China	22	Not available
FOCUCUIU	9901-2	cucumerinum	0186		China	22	Not available
Focuc014	9906-2	cucumerinum (redesignated radicis-cucumerinum)	0184		China	22	Not available
Focuc016	9909-2	cucumerinum	0185		China	22	Not available
Focuc017	9909-3	cucumerinum	0186		China	22	Not available
Focuc022	ATCC 36330	cucumerinum (redesignated a nonpathogen)	0180		Israel	22	Not available
Focuc026	ATCC 201950	cucumerinum	0180		USA (Florida)	22	Not available
Focuc027	Cu:4-1 Koma 4	cucumerinum	0181		lapan	22	Not available
Focuc028	Cu:5-0 Koma 5	<i>cucumerinum</i> (redesignated	0183		Japan	22	Not available
Focuc036	NRRI 26437	cucumerinum			USA (South Carolina)	ARS	Not available
Focuc038	NRRI 26744	cucumerinum (redesignated			lanan	ARS	Not available
1 ocucoso		a nonpathogen)			Jupun	7113	Not available
Focuc039	NRRL38591	cucumerinum			New Zealand	ARS	Not available
Focuc040	07-08969	cucumerinum (redesignated			Netherlands	NAKT	Not available
		radicis-cucumerinum)					
Folag003	07-27503	lagenariae			Japan	32	Not available
Folag006	Lag:7-1	lagenariae			Kumamoto, Japan	32	Not available
Folag007	No. 87	lagenariae			Tochigi, Japan	32	Not available
Folag008	No. 134	lagenariae			Tochigi, Japan	32	Not available
FomIn002	CBS 420 90	melonis			Israel	22	Not available
FomIn002	CBS 423.90	melonis			Israel	22	Not available
FomIn017	NIDDI 22519	melonis				Q1	Not available
Fomin017		molonis			CIA Erança	01	Not available
FOIIIIIIUIO	NRRL22519	meionis			FIGILE	01	Not available
FominU19	NRRL22520	meionis			USA	81	Not available
FomIn020	NRRL22521	melonis			Belgium	81	Not available
FomIn021	NRRL26172	melonis			China	ARS	Not available
FomIn023	NKRL26174	melonis			China	ARS	Not available
FomIn024	NRRL26745	melonis			Japan	ARS	Not available
FomIn025	NRRL26746	melonis			Japan	ARS	Not available
FomIn026	NRRL38516	melonis			New Zealand	ARS	Not available
FomIn027	NRRL38524	melonis			New Zealand	ARS	Not available
Fomom002	NRRL26748	momordicae			Japan	82	Not available
Fomom003	90NF1-2	momordicae			Kagoshima, Japan	32	Not available
Foniv011	F-086-1	niveum	0082	1	USA (Maryland)	83	Not available
Foniv017	F-097-2	niveum	0082		USA (Delaware)	51	Not available

(Continued on next page)

TABLE 5 (Continued)

Strain	Original designation	Forma specialis or description	VCG	Race	Origin of strain	Source or reference ^a	GenBank assembly accession no.
Foniv018	F-100-2	niveum	0082	2	USA (Delaware)	51	Not available
Foniv033	NRRL26747	niveum			Japan	ARS	Not available
Foniv034	NRRL36275	niveum (redesignated a nonpathogen)				ARS	Not available
Foniv035	NRRL38278	niveum (redesignated a nonpathogen)			USA	ARS	Not available
Foniv036	NRRL38503	niveum			New Zealand	ARS	Not available
Foniv038	NRRL38552	niveum (redesignated a nonpathogen)			Israel	ARS	Not available
Foniv039	LB	niveum		0		NAKT	Not available
Foniv040	IPO 30288	niveum		1		IPO	Not available
Foniv041	CBS 419.90	niveum			Israel	22	Not available
Forcu005	14	radicis-cucumerinum			Canada	22	Not available
Forcu017	34	radicis-cucumerinum			Canada	22	Not available
Forcu020	38	radicis-cucumerinum			France	22	Not available
Forcu028	Afu-58	radicis-cucumerinum	0260		Crete, Greece	22	Not available
Forcu029	Afu-68(A)	radicis-cucumerinum	0260		Crete, Greece	22	Not available
RBG1687	RBG1687	Nonpathogen, from			Australia	M.L.	Not available
		Wollemia nobilis seedling leaves					
RBG1693	RBG1693	From flannel flower roots			Australia	M.L.	Not available
RBG5713	RBG5713	Nonpathogen, from soil			Australia	49	Not available
RBG5786	RBG5786	Nonpathogen, from soil			Australia	49	Not available
RBG5789	RBG5789	Nonpathogen, from soil			Australia	49	Not available
RBG5791	RBG5791	Nonpathogen, from soil			Australia	49	Not available
RBG5798	RBG5798	Nonpathogen, from soil			Australia	49	Not available
RBG5820	RBG5820	Nonpathogen, from soil			Australia	49	Not available
RBG5824	RBG5824	Nonpathogen, from soil			Australia	49	Not available
RBG5827	RBG5827	Nonpathogen, from soil			Australia	49	Not available
F. proliferatum Fol3	Fol3	From a Lilium sp.			Netherlands	47	NJCT01000000
<i>Fusarium</i> sp. strain Na10	Na10	From a Narcissus sp.			Netherlands	47	NJCS01000000

^aAbbreviations and initials: NAKT, NAKtuinbouw, Netherlands Inspection Service for Horticulture, Roelofarendsveen, Netherlands; B.L., Bart Lievens, Scientia Terrae Research Institute, Belgium; J.V.D., Joop van Doorn, PPO Research Centre, Lisse, Netherlands; M.L., Matthew Laurence, Plant Disease Diagnostic Unit of the Royal Botanic Gardens and Domain Trust, Sydney, Australia; ARS, Agricultural Research Service, USDA, USA; IPO, Plant Research International (formerly Instituut voor Planteziektenkundig Onderzoek), Wageningen, Netherlands; K.O., Kerry O'Donnell, USDA ARS, Peoria, IL.

selected query genes for which we found only a single hit in each genome, leaving us with 440 genes. We used ClustalO (60) to construct a multiple-sequence alignment for each gene and a custom python script to concatenate these alignments. This alignment was subsequently trimmed using trimAl-strictplus. We used PhyML v20120412 (61) with 100 bootstraps to infer phylogeny and ETE v3.0.0b35 (62) to visualize the tree.

Marker discovery and primer design. A custom python script was written to extract the sequence (plus 150 bp up- and downstream) of candidate effector genes from each of the genome assemblies using BLASTN with default parameters. MUSCLE v3.8.31 (63) was used to generate alignments of each gene, and phylogeny was inferred using PhyML v20120412 with 100 bootstraps. Another python script was used to traverse the tree in ETE v3 to identify instances where all isolates belonging to a *forma specialis* were clustering together in a separate clade, indicating sequence similarity that could potentially be used for primer design. Highlighting, drawing, and rendering of the gene trees were done using ETE v3. Visual inspection of each of the gene trees allowed for the selection of a final set of marker genes per *forma specialis*. Scripts are available upon request.

Primers were designed manually based on the sequence alignment per gene (see Data S1 in the supplemental material). In cases where only a few SNPs were identified to separate host specificity of isolates, we aimed to target the mismatching nucleotides toward the 3' end of the primer, as described in reference 64.

DNA isolation. Genomic DNA isolation for testing of markers was performed using 10- to 20-day-old mycelium scraped off a peptone-dextrose agar (PDA) plate as starting material. The tissue was disrupted by shaking it in a TissueLyser (Qiagen) for 2 min at 30 Hz in the presence of 400 μ l of Tris-EDTA (TE), 300 μ l of phenol-chloroform (1:1), and glass beads. The aqueous phase was transferred to a fresh tube, and an equal volume of chloroform was added. The DNA in the aqueous phase was transferred to a fresh tube and diluted 10× with sterile Milli-Q water prior to use in PCR. DNA quantity was estimated for the TaqMan standard curve using a Qubit fluorometer (Thermo Fisher Scientific).

PCR. PCR was executed using Sphaero-Q Super *Taq* (Gorinchem, Netherlands) in 20- μ l reaction volumes which included the following components (final concentration): 1× Sphaero-Q Super *Taq*

					T _{ann}	Product
Gene ID	Target gene	Target forma specialis	Primer name	Primer sequence (5'-3')	(°C)	size (nt)
+	FEM1	Positive control	fp157	ATGAAGTACACTCTCGCTACC		
			fp158	GGTGAAAGTGAAAGAGTCACC	54	274
94	HPEG	All cucurbit infecting	fp7304	GCCTCATTGAAGTTTCAACA		
		5	fp7321	TGGTAAAGGACACGACCATT	54	346
13	SIX13	radicis-cucumerinum	fp7305	TTGCCCAAAATGGCATGTTT		
			fp7322	CATTGACACTGTAAGTGGG	56	328
70	HPEG	radicis-cucumerinum	fp7306	TACAACCTCTCTCTTTCCTT		
			fp7323	GCTGAATTCTAGCAGAGAAT	54	454
66	HPEG	cucumerinum	fp7307	CCGTTATGGCCAGAGATC		105
			fp7324	CCAACAAACAGAGCAAAACTAA	54	425
99	HPEG	cucumerinum	fp7308	CTACCAATCTCTCCTGAGTG		
			fp7325	GTCGATTGCAGTGCTAGTCT	54	445
21	Fom effector 7	cucumerinum	fp7309	CAGTCTAACCCTGTCTCATT		
			fp7326	CGCCAATAGATAGTGATGGA	54	381
1	SIX1	melonis	fp7310	CCTCTCAGTCCTTGGGTCT		
			fp7327	ACTCGCTTCAGCTTACCGA	54	397
20	Fom effector 6	melonis	fp7406	TGAAAGTCTTGGCGGGTGT		
			fp7328	TCCTCTCCATCCTCATCAGT	56	305
18	Fom effector 3	melonis + niveum	fp7312	TTAGTGCAGCTTTTCTCCTC		
			fp7329	AGTGGTTAGTCAAGTGGTAA	54	299
99	HPEG	niveum	fp7313	TGCCGGGCTAGTTAATATAGT	- 4	100
			fp7330	ACCATTTTTCTGTTGGGGTTG	54	406
100	HPEG	niveum	fp7314	ATTTTGCTAGCTTCAGCAGTT		
			fp7331	ATCCTGAACGGTGACTAGAG	54	482
21	Fom effector 7	niveum	fp7315	CGCTCGCTATAATTCAAACG	- 4	120
			fp7332	GGAGGAGCACTACAACTAAT	54	139
71	HPEG	lagenariae	fp7407	TAGTCCAATCTGCCTCAGCAA	54	270
			fp7410	GGAAGTGAGCATTCTTCCGTA	54	270
99	HPEG	lagenariae	fp7408	TCGTATCTCTCAGTAGTATGG	- 4	267
			fp7411	AATGGATACCTTATAAGGGCT	54	367
1	SIX1	lagenariae + momordicae	fp7409	TTGGGATTGCGGCTTATGCT	5.4	462
			fp7412	AAAGTGGTACACTCCGTGC	56	463
98	HPEG	momordicae	fp7318	AGGTGCAGCGTTTTTAGGT	60	160
			fp7335	GAGGGCTGGTTGAGAACTA	60	469
130	HPEG	momordicae	fp7319	TCTACGCTTCGAGGATGGTA	54	260
			fp7336	TCGTTTAGACGACTACAACC	50	308
99	HPEG	luffae	fp7320	TACTCTCCTAGAGTCAGTCT	54	606
			fp7337	CACGCCATCATCCTTTATTC	54	606

TARI F	6	Primers	and	annealing	temperatures	used	in	this	study
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buffer, 0.25 U of Sphaero-Q Super *Taq*, 5 pmol of each primer, deoxynucleoside triphosphates (dNTPs; 0.2 mM each), and 1 μ l of template DNA. The following PCR program was used: 2 min at 94°C; 35 cycles of 30 s at 94°C, 30 s at annealing temperature (T_{ann}), and 40 s at 72°C; 5 min at 72°C; and a pause at 16°C. The PCR primer sequences and corresponding annealing temperatures are listed in Table 6. *Fusarium FEM1* primers were used as a positive control and sterile Milli-Q was used as a negative control for each of the primer combinations instead of template DNA.

TaqMan real-time PCR. TaqMan reverse transcription-PCRs (RT-PCRs) were performed on a Quant-Studio 3 system (Thermo Fisher Scientific). Primers and probes were designed using Primer3web v4.0.0 (http://primer3.ut.ee/), and their sequences can be found in Table 7. A total volume of 10 μ l of the reaction mixture included the following components (final concentrations): 1× Sphaero-Q Super *Taq*

IABLE / Primers and probes used for the Tagivian experiment
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Gene ID	Target gene	Target forma specialis	Primer		Product
			name	Sequence (5'–3')	size (nt)
21	Fom effector 7	niveum	fp7589	CCGGTACCCCAGCTTTATGT	
			fp7590	CAGCAACGTTCTGAAAGCGT	116
			probe_3	HEX-TGCAGGTTGGCAGGCCCCTG-BHQ1	
100	HPEG	niveum	fp7591	CACCAACAACTATGCGGCAC	
			fp7592	GCAATTGACCCAGCTGCAAT	138
			probe_4	FAM-AGTCGCCGGCCACCACATTGA-BHQ1	
EF1α	Elongation factor 1 α	All strains	fp7710	CGCTGAGCTCGGTAAGGG	
	-		fp7711	CCAGAGAGCAATATCGATGGTGA	97
			probe_7	TAMRA-ACGCCTGGGTTCTTGACAAGCTCA-BHQ2	

buffer, 0.25 U of Sphaero-Q Super *Taq* (Gorinchem, Netherlands), 3 pmol of each primer, 1 pmol of each probe, dNTPs (0.2 mM each), $0.1 \times$ ROX reference dye (Thermo Fisher Scientific), and 1 μ l of template DNA. Four simultaneous amplifications were performed for each sample to confirm the reproducibility of the results. A negative-control sample consisted of sterile Milli-Q substituted for the DNA template. The PCR program was set as follows: 2 min at 94°C and 40 cycles of 30 s at 94°C, 48 s at 60°C, and 12 s at 60°C (data collection).

Disease assays. Pathogenicity tests were performed using the root dip method (65). In short, conidia were isolated from 5-day-old cultures in NO₃ medium (0.17% yeast nitrogen base, 3% sucrose, 100 mM KNO₃) by filtering through Miracloth (Merck; pore size of 22 to 25 μ m). Spores were centrifuged, resuspended in sterile Milli-Q water, counted, and brought to a final concentration of 10⁷/ml. When the first true leaves were emerging (after ~10 days), 5 to 8 seedlings per treatment were uprooted, inoculated, individually potted, and kept at 20°C (*F. oxysporum* f. sp. *radicis-cucumerinum*) or 25°C (all other *formae speciales*) in the greenhouse. The following plant cultivars were used: *Cucumis sativus* cv. Paraiso, *Cucumis melo* cv. Cha-T, and *Citrullus lanatus* cv. Black Diamond. Two weeks after inoculation, disease was scored using a disease index from 0 to 4 as described in detail by van Dam et al. (33).

Accession number(s). Whole-genome shotgun projects for the newly sequenced strains of *F. oxysporum* have been deposited at GenBank under BioProject no. PRJNA389501. Raw sequence data have been deposited into the Sequence Read Archive under accession number SRP109253. All publicly available genome sequences that were used were obtained from GenBank. Their NCBI accession numbers can be found in Table 5.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at https://doi.org/10.1128/AEM .01868-17.

SUPPLEMENTAL FILE 1, PDF file, 4.4 MB.

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