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Relationship between pathogenicity, race and vegetative compatibility grouping among Algerian populations of *Fusarium oxysporum* f. sp. *pisi* causing pea wilt

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Abstract

Fusarium oxysporum f. sp. *pisi* (FOP) is a significant and destructive pathogen of field pea in Algeria. In the present study, 50 isolates of *F. oxysporum* f. sp. *pisi*, the causal agent of pea (*Pisum sativum*) wilt, collected from different parts of western Algeria and representing four races of the pathogen, were analyzed for virulence. The wilt incidence ranged from 6.66 to 88.33% on a highly susceptible cultivar (Little Marvel). Twenty-one isolates belonging to four races of FOP and one nonpathogenic *F. oxysporum* (FO) isolate were analyzed for vegetative compatibility in order to reveal the genetic structure of the population and to check the reliability of the method for the identification of physiological races of FOP. Obtained results showed that the FOP isolates could be classified into four main vegetative compatibility groups (VCGs) that corresponded to races 1, 2A, 2B and 5. The race 6 isolate fell into the race 1 VCG. To our knowledge, this is the first such study in Algeria of its kind.

Key words: *Fusarium oxysporum* f. sp. *pisi*, *nit* mutants, pathogenicity, pea wilt, physiologic races, vegetative compatibility group

Introduction

Fusarium oxysporum f. sp. *pisi* (FOP), the causal organism of pea wilt, occurs as a number of races identified by tests on differential hosts of pea. So far, four races: 1, 2, 5 and 6, have been distinguished. Races 1, 2 and 6 have been reported in Europe, while all four races are found in the USA (Haglund and Kraft 2001) and in Algeria (Merzoug *et al.* 2014). The pathogenicity test to determine the race of an isolate is time-consuming and greatly affected by environmental conditions. In the absence of a sexual stage another way of grouping *Fusarium* isolates was proposed, based on their ability to form heterokaryons by anastomosis. Vegetative compatibility groups (VCGs) in fungi are a genetic-trait controlled by *vic* or *het* loci and identical alleles at each loci must be present in two compatible hyphae before anastomosis takes place (Leslie 1993). Vegetatively compatible isolates of a fungal species are placed

in the same VCG. Isolates within the same VCG often share common biological, physiological, and pathological characteristics. VCGs are identified by using nitrate nonutilizing (*nit*) auxotrophic mutants that show thin but expansive growth on minimal medium with nitrate as sole nitrogen source. Isolates are considered vegetatively compatible when they complement *nit* mutants anastomose and produce wild type growth (Puhalla 1985). That exchange of genetic material would be limited to compatible isolates within a VCG and so each VCG represents a genetically isolated population. Vegetative compatibility has been used to classify isolates of *F. oxysporum* belonging to distinct formal specials (Klein *et al.* 2005), races within a special form (Elena and Pappas 2006) and special forms and/or races within or between different geographical origins (Di Primo *et al.* 2002; Pasquali *et al.* 2005). VCG has

been used to distinguish between nonpathogenic and pathogenic populations on the same host species (Lori *et al.* 2004). In general, pathogenic isolates of *F. oxysporum* in the same VCG are assumed to be associated with the same clonal lineage, even if they are geographically isolated (Leslie 1993). However, isolates belonging to different VCGs can also belong to the same clonal lineage (Baayen *et al.* 2000). Several formae speciales of *F. oxysporum* have been characterized by VCG analysis (Lori *et al.* 2004; Abo *et al.* 2005), as well as by random amplified polymorphic DNA (RAPD) analysis (Alves-Santos *et al.* 2007; Bayraktar *et al.* 2008; Baysal *et al.* 2010) and several studies have combined both markers (Vakalounakis *et al.* 2004; Nagarajan *et al.* 2006). This technique is appropriate for developing countries with inadequate facilities for molecular work and can rapidly determine genetic groups of many fungal pathogens and their relation to pathogenicity (Cumagun *et al.* 2008). The objective of this work was to determine VCGs in a collection of 21 isolates of different pathogenic races *F. oxysporum* f. sp. *pisi* from different geographical origins in four different agro-climatic zones in western Algeria (coastal plains, interior plains, the high plateaus, and the Sahara) and compatibility between pathogenic and nonpathogenic *F. oxysporum* isolates obtained from asymptomatic pea rhizospheres.

Materials and Methods

Fungal isolates

The study was conducted with 50 isolates representing the four races of FOP collected from different regions of western Algeria (Table 1). Pathogenicity tests were performed on the standard pea differential cultivars. The inoculum production and inoculation method have been previously described (Merzoug *et al.* 2014). Twenty-one isolates of different pathogenic races *F. oxysporum* f. sp. *pisi* and one nonpathogenic isolate (np) were selected for vegetative compatibility tests. The choice of this population was based on pathogenicity, races and areas. The isolates were maintained as single-spore colonies in tubes with potato dextrose agar (PDA), at 4°C. The isolates selected for the tests are listed in Table 2.

Vegetative compatibility tests

Culture media

The media used in the vegetative compatibility study were PDA with 1.5% chlorate (PDC), salt minimal medium with 1.5% chlorate (MMC), minimal medium (MM) with nitrate, chlorate, nitrite and hypoxanthine added according to Puhalla (1985) and Correll *et al.* (1987).

Determination of VCGs

Selection, characterization and storage of *nit* mutants

Vegetative compatibility groups were determined through the complementation of nitrate nonutilizing (*nit*) mutants as a visual indicator of heterokaryon formation (Puhalla 1985). Generation, isolation and characterization of *nit* mutants were carried out according to the methodology of Correll *et al.* (1987). The *nit* mutants were selected by Puhalla's method (Puhalla 1985). For this, 2 mm PDA blocks with fungal mycelium were transferred to 9 cm plates that contained PDC or MMC (Puhalla 1985), and were incubated at 25°C for 1 to 3 weeks. Fast growing sectors were transferred to minimal medium (MM) and those which grew as fine and expanding colonies, without aerial mycelium, were considered to be *nit* mutants. In cases where we were not able to select mutants under the conditions described above, we introduced some modifications, in order to increase the chlorate concentration in MMC or PDC up to 5%, or two cultures consecutive on MMC (1.5%) chlorate. Changes were inspired by the methodology described by Nogales Moncada *et al.* (2009).

Phenotype identification

Nit mutants were grown on basal medium amended with different nitrogen sources: sodium nitrate, sodium nitrite and hypoxanthine. Petri dish cultures were incubated under the previously described conditions and examined after 5 days for mutant identification. Based on an ability to grow on the different nitrogen sources, three types of mutants were easily identified. *Nit1* mutants were mutated in a structural locus of nitrate reductase. Those mutants did not grow with nitrate as the only nitrogen source, but they did grow in the presence of nitrite or hypoxanthine. *Nit3* mutants were mutated in a locus which was probably involved in the regulation of both nitrate reductase and nitrite reductase. Consequently, these mutants did not grow with nitrate or nitrite as the only nitrogen source, but they did grow in the presence of hypoxanthine. Lastly, *nitM* mutants were mutated in one of the loci which were involved in the synthesis of the molybdenum co-factor, which is necessary for both the reduction of nitrate and the hydroxylation of hypoxanthine. These mutants cannot grow on nitrate or hypoxanthine as the only nitrogen source, but they will grow on nitrite (Correll *et al.* 1987).

Complementary tests

Vegetative compatibility was determined on the basis of the formation of heterokaryons in the contact area between the two colonies, as shown by the growth of aerial mycelium similar to that of the wild colony.

Table 1. Place of origin, races and wilt incidence of *Fusarium oxysporum* f. sp. *pisi* isolates

No.	Geographic origin	Departments	Isolates codes	Pathogenicity	
				wilt incidence [%]	races
1	Coastal plains	Ain Temouchent	Fop A1	58.33 b	1
2		Ain Temouchent	Fop A2	56.66 b	1
3		Ain Temouchent	Fop A16	25.66 d	6
4		Ain Temouchent	Fop A18	21.00 d	2
5		Ain Temouchent	Fop A19	26.66 d	1
6		Ain Temouchent	Fop A21	36.33 c	5
7		Ain Temouchent	Fop A22	41.33 c	6
8		Mostaganem	Fop M42	36.66 c	5
9		Mostaganem	Fop M43	38.00 c	2
10		Mostaganem	Fop M44	45.00 c	1
11		Mostaganem	Fop M45	40.33 c	1
12	Interior plains	Tlemcen	Fop TL24	25.66 d	6
13		Tlemcen	Fop TL27	56.33 b	1
14		Tlemcen	Fop TL28	60.00 b	6
15		Tlemcen	Fop TL29	71.66 b	1
16		Tlemcen	Fop TL31	69.66 b	1
17		Tlemcen	Fop TL32	69.00 b	1
18		Chlef	Fop C20	78.00 b	1
19		Chlef	Fop C21	63.33 b	1
20		Relizane	Fop R17	60.00 b	1
21		Relizane	Fop R26	70.00 b	2
22		Relizane	Fop R27	46.66 c	1
23		Relizane	Fop R28	43.33 c	1
24		Relizane	Fop R29	83.33 a	1
25		Relizane	Fop R31	66.66 b	1
26		Relizane	Fop R32	63.33 b	6
27		Mascara	Fop Ma6	60.66 b	2
28		Mascara	Fop Ma9	46.66 c	2
29		Mascara	Fop Ma13	60.33b	1
30		Mascara	Fop Ma15	60.66 b	2
31	Mascara	Fop Ma19	65.00 b	1	
32	Sidi Belabes	Fop Sb1	86.66 a	1	
33	Sidi Belabes	Fop Sb3	60.00 b	2	
34	Sidi Belabes	Fop Sb4	62.33 b	1	
35	Sidi Belabes	Fop Sb11	67.00 b	1	
36	Sidi Belabes	Fop Sb14	66.66 b	1	
37	High plateaus	Tiaret	Fop T46	35.00 c	1
38		Tiaret	Fop T47	40.00 c	2
39		Tiaret	Fop T48	23.33 d	1
40		Tiaret	Fop T49	50.00 c	6
41		Tiaret	Fop T50	44.33 c	1
42		Tiaret	Fop T51	39.00 c	1
43		Tiaret	Fop T52	40.00 c	1
44		Saida	Fop S56	88.33 a	1
45		Saida	Fop S58	34.33 c	1
46	Sahara	Adrar	Fop Ad60	6.66 e	np
47		Adrar	Fop Ad67	88.33a	2
48		Adrar	Fop Ad62	60.00 b	2
49		Adrar	Fop Ad63	68.33 b	1
50		Adrar	Fop Ad66	48.33 c	1

np – nonpathogenic *F. oxysporum*

The values with a common letter do not differ significantly at 5% level using the Newman-Keuls test

Table 2. Number of each *nit* mutant type selected on characterization media in relation to geographic origin

Geographic origin	Department	Isolates	Race	<i>nit1</i>	<i>nit3</i>	<i>nitM</i>	Total
Coastal plains	Ain Temouchent	A2	1	10 (100%)*	0 (0%)	0 (0%)	10
	Ain Temouchent	A21	5	10 (41.66%)	8 (33.33%)	6 (25%)	24
	Ain Temouchent	A22	6	8 (66.66%)	0 (0%)	4 (33.33%)	12
	Mostaganem	M42	5	9 (69.23%)	0 (0%)	4 (30.76%)	13
	Mostaganem	M44	1	11 (45.83%)	1 (4.16%)	12 (50%)	24
Interior plains	Tlemcen	TI 24	6	7 (58.33%)	0 (0%)	5 (41.66%)	12
	Tlemcen	TI27	1	13 (86.66%)	0 (0%)	2 (13.33%)	15
	Tlemcen	TI28	6	16 (88.8%)	0 (0%)	2 (11.11%)	18
	Chlef	C20	1	6 (60%)	2 (20%)	2 (20%)	10
	Cheliff	C21	1	6 (54.5%)	1 (9%)	4 (36.3%)	11
	Relizane	R26	2	11 (100%)	0 (0%)	0 (0%)	11
	Relizane	R28	1	2 (20%)	2 (20%)	6 (60%)	10
	Mascara	Ma13	1	14 (63.6%)	0 (0%)	8 (36.36%)	22
	Mascara	Ma15	2	7 (35%)	7 (35%)	6 (30%)	20
	Sidi Belabes	Sb1	1	8 (88.8%)	1 (11.1%)	0 (0%)	9
High plateaus	Sidi Belabes	Sb11	1	12 (46.15%)	0 (0%)	14 (53.84%)	26
	Tiaret	T49	6	5 (21.73%)	16 (69.56%)	2 (8.69%)	23
	Tiaret	T51	1	2 (9.09%)	4 (18.18%)	16 (72.72%)	22
Sahara	Saida	S58	1	5 (33.33%)	5 (33.33%)	5 (33.33%)	15
	Adrar	Ad62	2	6 (60%)	0 (0%)	4 (40%)	10
	Adrar	Ad63	1	10 (83.33%)	0 (0%)	2 (16.66%)	12
	Adrar	Ad60	np	2 (14.28%)	10 (71.42%)	2 (14.28%)	14
Total			22	180 (52.47%)	57 (16.41%)	106 (30.9%)	343

*percentage of isolates in that group; np – nonpathogenic *F. oxysporum*

A portion of the *nitM* colony was placed in the middle of the culture dish containing MM, and one portion each of *nit1* and *nit3* mycelium from other strains was placed at an equal distance from the *nitM* colony. Complementation between *nit1* and *nit3* mutants occurs less frequently than complementation between one of these mutants with a *nitM* mutant. Each *nitM* was paired with a *nit1* and a *nit3* from other strains in all possible combinations. All pairings were replicated twice. The complementation test was considered: negative – when there was no prototrophic growth in the mycelial line of contact; weak – when interaction became evident by the appearance of a thin, sometimes noncontinuous, zone of prototrophic growth with very little aerial mycelium; strong – when a dense line of prototrophic growth with abundant aerial mycelium was obtained. Weak and strong reactions were taken as evidence of compatibility. Training the heterokaryon, visible by the appearance of a thick, dense aerial mycelium, was noted after 5, 10, 15, 25 days and up to 4 weeks.

Statistical analysis

The results were subjected to statistical analysis. All the collected data were submitted to ANOVA analysis using Statistica software v. 5.5 (Statsoft, Ed'99) and Biostat 2009 and the significance of differences among treatments was recorded at $p < 0.05$. Multiple comparisons of the means were conducted according to the Newman-Keuls test at $p < 0.05$.

Results

Results of percentage wilt incidence of 50 isolates of *F. oxysporum* f. sp. *pisi* collected from different parts of western Algeria and representing four races of the pathogen ranged from 6.66 to 88.33% on the highly susceptible cultivar (Little Marvel) (Table 1). A partial account of this work has previously been published (Merzoug *et al.* 2014).

Isolation, characterization and storage of *nit* mutants

The medium PDC was selected, that proved most favorable to the growth of mutants sectors. Isolation of chlorate-resistant mutants and phenotypic diversity of *nit* mutants were obtained from all FOP isolates, but there was considerable variation in their recovery rate. Whenever possible, at least two different *nit* mutants, preferably a *nit1* and a *nitM*, were selected and stored as indicated in Materials and Methods.

Vegetative compatibility tests

The results of the characterization of 343 chlorate-resistant mutants (Table 2), indicate that in 180 (52.47%) of them are *nit1*, 106 (30.9%) *nitM* and 57 (16.41%) *nit3*. The overall distribution of three types of *nit* mutant according to the races, shows that race 1 forms the greatest number of mutants with a total of 186 (54.22%), which represents more than half of all the mutants followed by race 6 with 65 (18.95%). The highest number of mutants *nit1* was obtained from race 1 with 99 (55%). Race 2 had the lowest number of mutants *nit3* with 7 (12.28%) (Table 3).

Development of the pairs of mutants

When possible, a *nit1* and a *nitM* mutant from each isolate were selected and paired in all possible combinations. The positive complementarity between different *nit* mutants resulted in the growth of aerial mycelium in the confrontation zone, resulting from the anastomosis hyphae matched mutant colonies. In this study, the line of fusion of the hyphae varied in its density and appeared 10–15 days after transfer of cuttings on the MM medium. It consisted of a continuous growth line of thick and abundant mycelium, or as a weak aerial mycelium line dispersed and a more or less dense or a less defined line separated with a mycelium +/- flush that appeared after 15 days up to a month (Fig. 1). The results of various comparisons between *nit* mutants complementary to a same isolate were self-compatible for all isolates tested except for two isolates of race 1 (Sb1 and A2) and an isolate of race 2 (R26) which was self-incompatible knowing that for the three isolates only *nit1* was obtained. The nonpathogenic isolate was self-compatible.

Vegetative compatibility group (VCG)

Analysis of results of the complementation test listed in Table 4 shows:

1. Race 2 isolates were classified into 2 subgroups, the Ma15 isolate was self-compatible but incompatible with all other isolates of race 2. Isolate R26 was

self-incompatible but it was compatible with isolate Ad62. These 2 isolates had distant geographical origins (internal plains and Sahara) while Ma15 and R26 which were incompatible had the same geographical origin.

2. Race 5 isolates were classified in the same VCG. They were compatible with each other and showed a slow complementation that occurred after a month of confrontation with isolates of race 1 (Sb1, Ad63, S58, C24 and A2). The only positive complementation was observed between M44 and M42, which were from the same region (Mostaganem).
3. Race 1 isolates were classified in the same VCG. All isolates were self-compatible with the exception of Sb1 and A2 isolates which were self-incompatible and formed only *nit1* that produce a negative or inconclusive complementation. Compatibility relationships within the race 1 VCG are not simple. The reaction of compatibility between isolates was variable. For example, the Sb1 isolate was only compatible with the TL27 and had a weak reaction with R28, while TL27 was compatible with all isolates of race 1 with the exception of S58. Ma13 was compatible with TL27 but not compatible with Sb1.
4. Race 6 isolates were classified in the same VCG as race 1. Race 6 isolates were all self-compatible. Complementation was positive or partial between most isolates of race 6.
5. The nonpathogenic isolate (np) was self-compatible. It was compatible with race 1 and race 6 isolates but was incompatible with the R2 races and race 5 isolates.

There was no compatibility between the VCGs that comprised races 2A and 2B and any other VCGs. However, there was a weak interaction between some race 1 and race 5 isolates, although these interactions only developed after at least 4 weeks incubation (Table 5).

The results obtained show that the FOP isolates from western Algeria can be classified into four main VCGs that corresponded to races 1, 2A, 2B and 5. The race 6 isolate, fell into the race 1 VCG, and although compatibility was not complete with any race 1 isolate tested. In addition, after several weeks of incubation there was also weak compatibility to some race 5 isolates. According to the procedure for determining the VCG described by Correll (1991) and the systematic numbering proposed by Katan and Katan (1999). The VCGs FOP code number is 0070-0073.

Discussion

In this study, virulence and vegetative compatibility were analyzed for FOP isolates representing four races prevalent in the west of Algeria. The range of wilt

Table 3. Distribution of numbers and percentages mutants (*nit*) according to the different physiological races of *Fusarium oxysporum* f. sp. *pisi*

<i>Nit</i> mutant	R1	R2	R5	R6	np	Total
<i>nit1</i>	99 (55%)*	24 (13.3%)	19 (10.5%)	36 (20%)	2 (1.1%)	180
<i>nit3</i>	16 (28%)	7 (12.28%)	8 (14.03%)	16 (28%)	10 (17.5%)	57
<i>nitM</i>	71 (66.98%)	10 (9.43%)	10 (9.43%)	13 (12.26%)	2 (1.88%)	106
Total	186 (54.22%)	41 (11.95%)	37 (10.78%)	65 (18.95%)	14 (4.08%)	343

np – nonpathogenic *F. oxysporum*; *percentage of isolates in that group

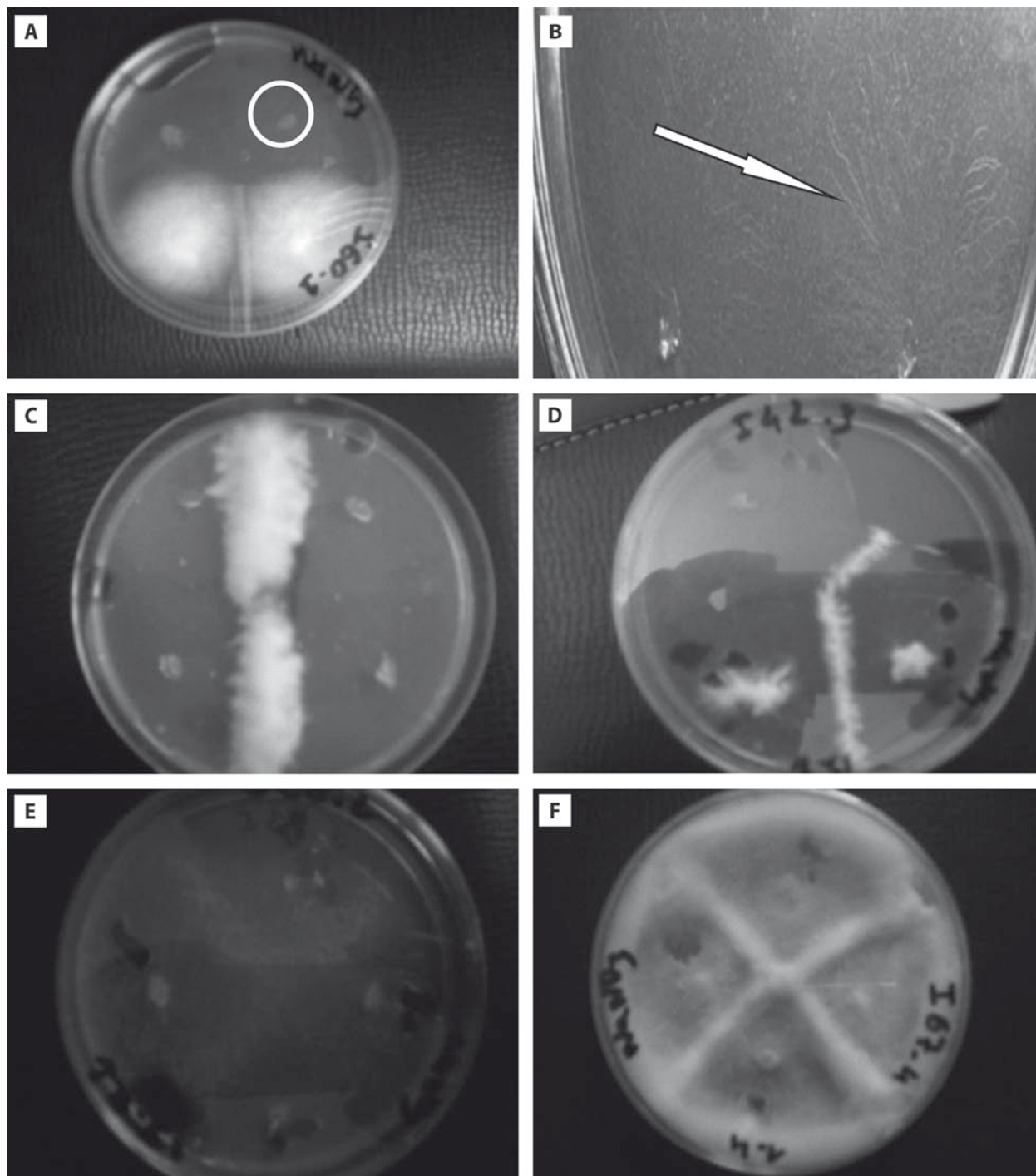


Fig. 1. Vegetative compatibility tests. A–B – growth of *nit* mutants on minimal medium (MM). The elliptical areas – the chlorate-resistant sectors; the arrow – development of thin mycelium. Pairing of *nit* mutants subcultured on MM; C – continuous growth line of a thick and abundant aerial mycelium; D – weak aerial mycelium line dispersed; E – less defined line separates with a mycelium +/- flush, that appears after a month; F – self-compatibility (np isolate Ad67)

Table 4. Vegetative compatibility relationships between race isolates of *Fusarium oxysporum* f. sp. *pisi*

Race	R2			R5		R6				R1															
	isolates	Ma15	Ad62	R26	M42	A21	T49	Tl28	Tl24	A22	Sb1	Sb11	Tl27	Ad63	S58	Ma13	C20	A2	Tl51	M44	R28	C21	Ad60		
R2	Ma15	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	
	Ad62		+	+	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	
	R26			-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	
R5	M42				+	+	-	-	-	-	-	-	-	+/-	+/-	-	-	+/-	-	+	-	-	-	-	
	A21					+	-	-	-	-	-	-	-	+/-	-	-	+/-	+/-	-	-	-	-	-	-	
R6	T49						+	+	+/-	+/-	-	-	+	+/-	+/-	+	-	+	-	+	-	+	-	+	
	Tl28									+	+/-	+	-	+	+/-	-	+/-	-	+	-	-	+	-	+/-	
	Tl24												+	+/-	-	+	-	+/-	-	-	-	-	-	+	
	A22																+	-	+	+	-	-	-	-	
R1	Sb1									-	-	+	-	-	-	-	-	-	-	-	+	-	-	+/-	
	Sb11										+	-	-	-	+	+/-	-	-	-	-	+	-	-	+	
	Tl27											+	+/-	-	+	+	+	+	+	+	+	+	+	+	
	Ad63												+	+/-	-	+	-	+	+	+	+	+	+	+	
	S58													+	-	+	-	+	+	+	+	-	-	+	
	Ma13															+	-	-	+/-	+	-	-	-	+/-	
	C20																+	-	+/-	+	-	+	-	+	
	A2																	-	+/-	+	+/-	-	-	+/-	
	Tl51																			+	-	+	-	-	+/-
	M44																				+	+	+	+	+/-
	R28																					+	-	-	+
C21																						+	+	+/-	
np	Ad60																							+	

"+" – compatible interaction; "-" – incompatible interaction; "+/-" – partially compatible interaction; np – nonpathogenic *F. oxysporum*

Table 5. Correlation between vegetative compatibility groups (VCG) and races

Race	R1	R2A	R2B	R5	R6	np
R1	+	-	-	+/-	+	+
R2A		+	-	-	-	-
R2B			+/-	-	-	-
R5				+	-	-
R6					+	+
NP						+

"+" – compatible interaction; "-" – incompatible interaction; "+/-" – partially compatible interaction; np – nonpathogenic *F. oxysporum*

incidence (6.33–88.33%) caused by the isolates showed variability in the virulence of the pathogen even on a single susceptible variety of pea. In the present work, the three types of mutant's nit whose highest percentage is the *nit1* followed by *nitM* and the lower the *nit3* without these differences may be related to a specific characteristic of isolates (geographical origin, race or virulence). According to Bowden and Leslie (1992) and Puhalla (1985), the frequency of *nit1* mutants is higher than the frequency of the other types of nit mutants. The percentage of *nit3* is always lower (Belabid

and Fortas 2002). The study of the organization of 21 isolates in the formae speciales *pisi* from different geographical origins (western Algeria) and belonging to the four races of FOP showed the existence of four VCGs, reflecting its genetic diversity. Such diversity has been reported for many formae speciales, such as *F. oxysporum* f. sp. *radicis-lycopersici* (Katan and Katan 1999), *F. oxysporum* f. sp. *ciceris* (Nogales Moncada *et al.* 2009) and *F. oxysporum* f. sp. *batatas* (Rodriguez-Molina *et al.* 2013). Comparing the VCG distribution and race distribution defined by means of differential hosts, showed that isolates which belong to different races are included in the same VCG (Saabale and Dubey 2014). This is the case of the isolates of race 1 and 6 *F. oxysporum* f. sp. *pisi* which are in the same VCG, whereas isolates of race 2 are included in two other VCGs. This situation is found in other formae speciales for which the VCG-race relationships are complex. Indeed, it is known that more than one race may appear in the same VCG and the same race may contain more than one VCG (Correll 1991). A similar degree of complexity has been noted in the *F. oxysporum* f. sp. *cubense* (Somrith *et al.* 2011), many of which belong to the same race and multiple VCGs correspond to a single race. It is the same for *F. oxysporum* f. sp.

lycopersici (Katan and Katan 1999) and *F. oxysporum* f. sp. *melonis* (Elena and Pappas 2006).

The results of the weak or partial vegetative compatibility observed between isolates of races 1 and 5 suggest a higher degree of similarity with isolates of race 1 than of races 2A and 2B. According to Whitehead *et al.* (1992), it seems likely that races 5 and 6 derived from race 1. The greatest degree of intra-racial variation seen among race 1 isolates may suggest that this is the oldest race. The differences between races 1 and 5 compared to the similarity between races 1 and 6 would indicate that race 6 may have evolved much more recently than race 5. Race 2 is very different from races 5, 6 and 1 and may have a distinct evolutionary origin. Races 2A and 2B are clearly very closely related, but the total lack of vegetative compatibility between them demonstrates that they are genetically distinct populations (Whitehead *et al.* 1992). It is possible that 2A and 2B are two separate races and might be distinguished by pathogenicity testing if the range of differential lines was extended; new line differential has been demonstrated recently for a sharper characterization race 2 *Fusarium oxysporum* f. sp. *phaseoli* (Alves-Santos *et al.* 2002). Generally, in various formae speciales all isolates of the same VCG are compatible with each other. In addition, some combinations between the race 1 isolates produced weak or partial reactions. These differences in the intensity of crop interactions have been reported in *F. oxysporum* and could reflect the result of the action of alleles at different loci affecting vegetative compatibility (Nogales Moncada *et al.* 2009). Moreover, some isolates in the VCG were incompatible; that is, two isolates might be vegetatively incompatible even though each is compatible with a 3rd isolate. Concerning vegetative compatibility in *Verticillium dahliae*, Rataj-Guranowska *et al.* (2016) observed bridges between American testers VCG 2 and VCG 4, also between American and Dutch testers and between subgroup testers VCG 2A and 2B. The occurrence of these bridging strains is well known in *F. oxysporum* (Vakalounakis and Fragkiadakis 2004). Another difference between FOP and other formae speciales is demonstrated by the anomalies found in the pairings of race 1 isolates. The presence of two subgroups is not explained by geographical factors since isolates from both Europe and North America are present in the two subgroups (Whitehead *et al.* 1992). The weak complementation found between some race 1 and race 5 isolates even though they are in separate VCGs is not unique. Somrith (2011) reported that bridging isolates were capable of forming heterokaryons with isolates in both of two closely related VCGs in *F. oxysporum* f. sp. *cubense*. According to the analysis of Whitehead *et al.* (1992) it may be possible for weak complementation to occur in *Fusarium oxysporum* f. sp. *pisi* (FOP) even when two isolates differ slightly at

one or more loci. Such differences could come about, for example, if a series of missense alleles of a particular het gene were present in the population. Some combinations might lead immediately to compatibility whereas others either would not or only do so after a long time, depending on the nature and position of the amino acid change (Nogales Moncada *et al.* 2009). This could explain both the variation found in race 1 and the weak compatibility between some isolates of race 1 and race 5. Studies undertaken by different molecular techniques have made it possible to determine that races 1, 5 and 6 are closely linked and that race 2 is distinct (Okubara *et al.* 2005; Sharma *et al.* 2006).

In this study, the complementation between pathogenic isolates FOP and nonpathogenic isolate FO was tested. The nonpathogenic isolate was vegetatively compatible with some isolates of races 1 and 6. It was grouped in the same VCG as races 1 and 6, which includes pathogenic and nonpathogenic isolates. According to a study conducted by Nogales Moncada *et al.* (2009) on pathogenic and nonpathogenic isolates, most of the non-pathogenic isolates were compatible with a yellowing-type low virulent *F. oxysporum* f. sp. *ciceris* isolate, thus suggesting the possibility of the existence of transition isolates between pathogenic and nonpathogenic populations. Vegetative compatibility testing supported that *F. oxysporum* f. sp. *betae* is polyphyletic and that pathogenic isolates cannot be differentiated from nonpathogenic *F. oxysporum* using vegetative compatibility (Webb *et al.* 2013). Little is known, however, about the ecological significance and the population dynamics of the non-pathogenic strains of *F. oxysporum*, particularly at the subspecies level. Differentiating strains among the nonpathogenic *F. oxysporum* isolates is very important since they could provide a means of identifying and characterizing the various subpopulations of *F. oxysporum*. No specific relationship was observed between pathogenicity, VCGs and geographic origin of the isolates in this study. These results are in agreement with results from previous studies with other *Fusarium* spp. (Mohammadi and Mofrad 2009; Mohammadi *et al.* 2012) and on various formae speciales of *F. oxysporum* (Belabid and Fortas 2002).

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