Phytopathologia Mediterranea

The international journal of the Mediterranean Phytopathological Union Firenze University Press www.fupress.com/pm



Citation: Ayad D., Aribi D., Hamon B., Kedad A., Simoneau P., Bouznad Z. (2019) Distribution of large-spored *Alternaria* species associated with early blight of potato and tomato in Algeria. *Phytopathologia Mediterranea* 58(1): 139-149. doi: 10.13128/Phytopathol_Mediterr-23988

Accepted: November 19, 2018

Published: May 15, 2019

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Data Availability Statement: All relevant data are within the paper and its Supporting Information files.

Competing Interests: The Author(s) declare(s) no conflict of interest.

Editor: Vladimiro Guarnaccia, University of Torino, Italy.

Research Papers

Distribution of large-spored *Alternaria* species associated with early blight of potato and tomato in Algeria

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Abstract. Potato and tomato are important crops in Algerian agriculture, and both are threatened by abiotic and biotic stresses, and early blight is a major disease affecting both crops. Surveys carried out from 2012 to 2015 in 12 major growing regions for these crops yielded a total of 247 *Alternaria* isolates having morphological and cultural characteristics of sections *Alternaria* and *Porri*. Since early blight symptoms and morphological characteristics of the isolates did not allow sharp distinction between the different large-spored species of *Alternaria*, the isolates in section *Porri*, often considered primary causes of the diseases, were selected for molecular characterization by diagnostic PCR using specific primers. This allowed species identification of 147 *Alternaria* isolates as *A. solani*, *A. protenta*, *A. grandis* or *A. linariae*. These species were present on potato and tomato crops at varying frequencies, depending on the hosts and on bioclimatic locations. Pathogenicity tests for the four species, on detached leaflets and whole seedlings, showed that all were pathogenic to potato and tomato, with varying virulence. These results suggest that parasitic specialization of these *Alternaria* species on solanaceous plants should be reconsidered.

Keywords. Solanaceous crops, parasitic specialization, prevalence.

INTRODUCTION

Potato (Solanum tuberosum L.) and tomato (S. lycopersicum L.) are important crops grown in Algeria. They are distributed differently in the various bioclimatic zones of the country, and are grown with different cultural practices. Potato crops occupy significant area in several regions of Algeria, from north (Ain Defla) to south (El Oued), and from east (Skikda, Guelma) to west (Mostaganem, Mascara). Tomato, however, is mainly cultivated in the northern part of the country, less extensively and mostly in plastic houses, as in the Biskra region. Both crops are often cultivated side by side or follow one another in rotations for several years, in many coastal regions of the country. Intensification of these two crops without compliance to good agricultural practices led to the development of important diseases caused by fungi and oomycetes, including early blight and late blight.

The first reference mentioning A. solani Sorauer (= Macrosporium solani Ellis & G. Martin) as a pathogen causing potato leaf blight was that of Galloway (1891) in Australia. Chester (1892) then reported this pathogen in the United States of America, other solanaceous cultivated plants (Van der Waals et al., 2001; Odilbekov, 2015). A few years later, A. solani and A. alternata (Fr.) Keissl. were identified on potato leaves but A. alternata was considered first as saprophytic (Jones and Grout, 1897). It was not until 1984 that A. alternata was recognized as pathogenic on several solanaceous plants (Droby et al. 1984). The name "early blight" for the disease caused by A. solani was coined by Jones (1893) to distinguish this from late blight. The symptoms of the early blight appear on the leaves as dark, elongated or circular lesions each with concentric rings surrounded by a yellow halo.

On the basis of cultural techniques and morphological characteristics of the conidia (size and beak length), Simmons (2000, 2007) described several new Alternaria species on Solanaceae, that may be responsible for early blight on potato and tomato. Thus, new species were described, which are morphologically very similar to A. solani. These include A. grandis E.G. Simmons on S. tuberosum, A. tomatophila E.G. Simmons on S. lycopersicum, A. cretica E.G. Simmons & Vakal. On S. lycopersicum var. esculentum, and A. subcylindrica E.G. Simmons & R.G. Roberts on S. lycopersicum var. cerasiforme.

More recently, molecular and genomics techniques applied to Alternaria allowed redefinition of various species with large conidia commonly isolated from Solanaceae, and considered as the primary causes of early blight. Woudenberg et al. (2014), by synonymizing A. linariae (Neerg.) E.G. Simmons with A. cretica E.G. Simmons & Vakal., A. cucumericola E.G. Simmons & C.F. Hill, A. subcylindrica, A. tabasco E.G. Simmons & R.G. Roberts, and A. tomatophila, have expanded the host range of A. linariae to Solanaceae, Cucurbitaceae, and Scrophulariaceae. Alternaria linariae, a common pathogen of tomato, was also reported on potato during surveys carried out in Algeria (Ayad et al., 2018). Alternaria Grandis, which was confused for a long time with A. solani, has been regularly reported in many countries on potato (Lourenço et al., 2009; Cardoso, 2014; Bessadat et al., 2016; Landschoot et al., 2017). Alternaria grandis was also recently isolated from tomato (Bessadat et al., 2017) in the northwest of Algeria. Alternaria protenta, which is closely related to A. solani and whose host range was extended to Asteraceae, Euphorbiaceae, Gramineae and *Solanaceae* by Woudenberg *et al.* (2014), was recently detected on potato in Belgium (Landschoot *et al.*, 2017) and in Algeria (Ayad *et al.*, 2017).

The aims of the present study were, firstly, to identify at the species level the large-spored *Alternaria* isolates obtained from surveys carried out during three cropping years on different potato and tomato fields across the main bioclimatic areas of Algeria. Since early blight symptoms and the morphological characteristics of the isolates do not allow distinction between the different large-spored *Alternaria* spp., specific identification of the isolates was based on PCR techniques. The second objective was to assess the aggressiveness of isolates on their respective potato and tomato hosts, and to clarify the parasitic specialization of replace by large-spored *Alternaria* species by artificial cross inoculations *in vitro* and under conditions similar to those occurring in the field.

MATERIALS AND METHODS

Surveys and sampling

Surveys were carried in 60 locations across 12 regions in Algeria, with different cropping systems (fields and greenhouses). Three to four plots were sampled in each region during three successive cropping years: 2012-2013, 2013-2014 and 2014-2015. The majority of the localities were selected from the main potato- and tomato-producing regions represented by different bioclimatic areas of Algeria: Mediterranean climate (Algiers, Tipaza, Guelma and Skikda), semiarid climate (Mostaganem, Chlef, Mascara, Ain Defla and Bouira), and arid climate (Laghouat, Biskra and El Oued) (Figure 1). Sampling was carried out from potato and tomato plant organs showing the typical symptoms of early blight: 164 isolates were collected from potato leaves essentially and 83 from leaves, stems and fruits of tomato.

Isolation, purification and conservation of isolates

Isolations were performed according to the method of Van der Waals *et al.* (2004). Small tissue pieces (3 to 4 mm²) were cut from individual lesion edges, disinfected in 1% active NaOCl for 3 min, washed twice in sterile distilled water, and then dried with absorbent sterile paper. Four pieces of tissues were plated on individual Petri plates containing Potato Dextrose Agar (PDA), and these were incubated at 22°C under continuous light for 1 to 2 weeks. The purification of the *Alternaria* spp. isolates as single conidium cultures was car-



Figure 1. Geographic localities in the different bioclimatic areas: 1 = Mediterranean climate, 2 = semi-arid climate, 3 = arid climate

ried out between the 3rd and the 5th day of incubation. Plates containing the single conidium cultures were then placed under near-UV (12h dark and 12h light) to induce sporulation and allow further morphological characterization.

Single conidium cultures for short-term use were grown in Petri dishes containing a PDA for 12 d then stored at 4°C. For medium term use, 12-d-old PDA cultures in test tubes were stored at 4°C. For long term use, mycelium/agar plugs (8 mm diam.) were stored at 80°C in cryotubes containing sterile glycerol (30% w/w).

Identification of the large-spored Alternaria species using molecular markers

On the basis of the cultural and the morphological characteristics, 155 isolates with large conidia (out of a total 247 isolates) were selected for molecular characterization by PCR using specific primers, for analysis of genomic DNA by PCR/RFLP, using double enzymatic digestion of a portion of the calmodulin gene and sequencing the calmodulin and the RPB2 genes. Isolates were grown on PDA medium at room temperature for 10 d. Total genomic DNA from fungal samples was extracted according to the method described by Goodwin and Lee (1993). The DNA extracts of the 155 isolates were amplified using two sets of specific primer pairs to differentiate between A. linariae and A. solani and related species (A. grandis and A. protenta) (Gannibal et al., 2014). The primer pair OasF7 and OasR6 amplified a 164 bp fragment from the Alt a1 gene of A. solani, A. grandis and A. protenta, and the primer pair OatF4 and OatR2 amplified a 438 pb fragment from the calmodulin encoding gene of A. linareae. The PCR conditions were as described by Gannibal et al. (2014). To differentiate between A. grandis and A. solani sensu lato (i.e. including A. protenta), a portion of the calmodulin gene was amplified from the isolates that each gave a positive signal with the primer pair OasF7 - OasR6 using the primer pair CALDF1 / CALDR1 (Lawrence et al., 2013). The resulting PCR products were double digested with the restriction enzymes RsaI and HaeII. The digestion products were then separated by electrophoresis in 2% agarose gels and visualized under UV light after staining with ethidium bromide. Predicted restriction patterns mainly differed by the size of the larger fragment, i.e. 420 bp for A. solani sensu lato and 292 bp for A. grandis (Figure 2). Identification of the Alternaria species was validated by sequencing the fragment of the calmodulin locus amplified as described above from 42 isolates. To confirm the presence of A. protenta, a portion of the RPB2 locus was amplified from the isolates identified as A. solani based on their calmodulin sequence, using the primer pair RPB2-5F2 (Sung et al., 2007) and RPB2-7cR (Liu et al., 1999), and the resulting PCR product was sequenced. Sanger sequencing was performed by GATC Biotech. Sequence analyses were carried out using the Phylogeny.fr web service (Dereeper et al., 2008). Multiple sequence alignments were generated with MUSCLE and curated using the Gblocks algorithm. Maximum likelihood (ML) analyses were performed with PhyML. The robustness of the ML topologies was evaluated using the Shimodaira-Hasegawa (SH)-like test for branches.

Pathogenicity tests of large-spored Alternaria species on their respective hosts

To confirm pathogenicity of isolates, inoculations onto detached leaflets and whole plants were performed on susceptible varieties of tomato (cv. Marmande) and



Figure 2. Typical restriction patterns obtained by double digestion of PCR product corresponding to a portion of the calmodulin gene from *A. solani* and *A. grandis.* Size of the larger fragment, 420 bp for *A. solani sensu lato* (S) and 292 bp for *A. grandis* (G)

potato (cv. Spunta). These isolates were selected for their ability to produce conidia. Conidium suspensions were prepared from 15-d-old cultures. The final concentration of the suspensions was adjusted to 10^4 conidia mL⁻¹. The inoculations were performed by depositing drops (20 µL each) on detached leaflets or by spraying suspensions onto whole plants. Disease severity was evaluated at 15 d –post-inoculation, using a visual rating scale from 1 to 9 expressing the extension of necrosis CIP (International Potato Center) (2008).

RESULTS

Characterization of the large-spored Alternaria species according to early blight symptoms on potato and tomato

Isolation from plant samples showing typical early blight symptoms yielded 247 isolates (164 from potato and 83 from tomato). On the basis of the cultural and morphological characteristics of the colonies and conidia, the isolates were divided in three groups. The first (Group A: 92 isolates) included all the isolates with abundant conidium production and small catenulate conidia and identified as species belonging to section Alternaria. The second (Group B: 92 isolates) contained all isolates producing large solitary beaked conidia, related to species belonging to the section Porri. The third (Group C: 63 isolates) consisted of large-spored isolates that only formed sterile mycelium after culture purification. In relation to the host plants, 89% of the isolates belonging to the section Porri originated from potato, and 73% of the non-sporulating isolates originated from tomato. No correlation was found between the presence of yellow halos on symptomatic leaves and the production of a diffusible pigment on growth medium by the isolates from the diseased samples. Similarly, the various forms of symptoms generated colonies producing either large or small conidia. Thus, there was no relationships between necrosis size (small, medium or large necrotic lesions), the types of the colonies in culture medium (sporulating or non-sporulating) and the conidium types (large solitary or small catenulate conidia).

Molecular characterization and identification of largespored Alternaria species

Given the very similar morphological characteristics of the conidia, and the difficulty to distinguish the different *Alternaria* species within the section *Porri*, molecular markers and DNA-sequencing were used for their identification. The small-spored isolates (Group A) were not included in this examination. The three molecular approaches allowed clear species identification for 147 of 155 isolates from Groups B and C, while eight isolates could not be identified due to problems in DNA extraction and amplification at the selected loci (Table 1). Preliminary tests were carried out for the identification of the 147 isolates using two pairs of specific primers that differentiate A. solani, A. protenta and A. grandis (OasF7/OasR6) from A. linariae (OATF4/ OATR2). All the tested isolates generated positive signals with one of the two primer pairs, confirming the presence of these species on potato and tomato crops in Algeria. In this analysis, 37 isolates including seven from potato and 30 from tomato, were identified as A. linariae by amplification of the 438 pb from the gene encoding for calmodulin (Table 1). A PCR product corresponding to a fragment of the Alt a1 gene (164 bp) was amplified from the remaining 110 isolates. These isolates, of which 86 were isolated from potato and 24 from tomato, could be assigned to either A. solani or to phylogenetically related-species, i.e. A. grandis and A. protenta. Further identification was conducted in order to differentiate between A. solani and A. grandis isolates using PCR/RFLP, by amplifying the calmodulin locus followed by restriction enzyme digestion (HaeII and RsaI). Based on the characteristic electrophoretic profiles of the restricted PCR products, A. solani and A. protenta were separated from A. grandis. Thus, of the 110 isolates that gave positive signals with the OasF7/ OasR6 primer pair, 93 were A. solani, 12 were A. grandis and five showed abnormal enzymatic restriction digestion patterns that may be due to partial digestion with at least one of the restriction enzymes. These five isolates, as well as 37 other representative isolates obtained from the two host plants collected in nine of the twelve surveyed Algerian regions and representing the three identified species (A. linariae, A. grandis and A. solani), were selected for species confirmation by DNA sequencing which was firstly performed at the calmodulin locus. Phylogenetic analyses of sequences derived from the calmodulin locus of the 42 isolates (Figure 3) separated the isolates into two clades: the first (A) contained the isolates previously identified by PCR with specific primers as A. linariae, and the second (B) included two sub clades (B1 and B2) regrouping isolates previously identified as A. grandis (sub clade B1) and A. solani (sub clade B2). DNA sequencing at the RPB2 locus then carried out on the isolates of the sub clade B2 gave the phylogenetic tree shown in Figure 4, in which among the 17 isolates of A. solani, three were identified as A. protenta, two of which were isolated from the potato and one from tomato.

Table	1.	Identification	of lar	ge-spored	of A	Alternaria	spp	isolates.

Isolate	Species ^a	Host	Geographic origin	PCR OATF4/ OATR2	PCR OAsF7/OAsR6	PCR RFLP Profile ^b	CAL	Rpb2
DA001	A. linariae	Potato	Tipaza	+	-	ND	MH243795	_
DA002	A. linariae	Potato	Tipaza	+	-	ND	MH243769	_
DA003	A. linariae	Potato	Tipaza	+	-	ND	MH243793	_
DA005	A. solani s.l	Potato	Alger	-	+	S	_	_
DA006	A. linariae	Potato	Alger	+	-	ND	MH243789	_
DA007	A. linariae	Potato	Alger	+	-	ND	MH243794	_
DA008	A. solani	Potato	Alger	-	+	S	MH243805	MH243818
DA009	A. grandis	Potato	Alger	-	+	?	MH243790	_
DA010	A. solani s.l	Potato	Alger	-	+	S	_	_
DA011	A. solani	Potato	Alger	-	+	S	MH243806	MH243820
DA012	A. solani s.l	Potato	Alger	-	+	S		
DA013	A. solani s.l	Potato	Alger	-	+	S	_	_
DA014	A. solani s.l	Potato	Alger	-	+	S	MH243796	_
DA015	A. solani s.l	Potato	Bouira	-	+	S		—
DA016	A. solani s.l	Potato	Bouira	-	+	S	_	—
DA017	A, solani s.l	Potato	Bouira	-	+	S	-	_
DA018	A. solani s.l	Potato	Bouira	-	+	S	-	_
DA019	A. solani	Potato	Bouira	-	+	S		– MH243822
DA020	A, solani s l	Potato	Bouira	-	+	S	11112 10000	
DA021	A solani s l	Potato	Bouira	_	+	S	_	-
DA022	A solani s l	Potato	Bouira	_	+	s	-	-
DA025	A solani	Potato	Bouira	_	+	s	– MH243786	– MH243823
DA025	A solani s l	Potato	Bouira	_		s	11112 137 00	101112 15025
DA028	A solani s l	Potato	Bouira		-	S	-	-
DA020	A. solani s.l	Potato	Bouira	-	-	S	-	-
DA029	A. solani s.l	Dotato	Bouira	-	+	5	-	-
DA030	A. solani s.l	Potato	Bouira	-	+	5	-	-
DA031	A. soluni s.i	Potato	Doulla	-	+	S	-	-
DA032	A. solani s.l	Potato	Bouira	-	+	S	-	-
DA035	A. soluni s.l	Potato	Bouira	-	+	5	-	-
DA034	A. solani s.l	Polato	Bouira	-	+	5	-	_
DA035	A. solani s.l	Potato	Mascara	-	+	8	-	-
DA036	A. solani s.l	Potato	Mascara	-	+	3	-	_
DA037	A. linariae	Iomato	Mostaganem	+	-	ND	-	-
DA038	A. grandis	Potato	Alger	-	+	G	MH243770	-
DA039	A. linariae	Potato	Alger	+	-	ND	_	-
DA040	A. linariae	Tomato	Mascara	+	-	ND	MH243792	_
DA041	A. linariae	Tomato	Mostaganem	+	-	ND	-	_
DA042	A. linariae	Tomato	Mostaganem	+	-	ND	-	_
DA043	A. solani s.l	Potato	Mostaganem	-	+	S	-	_
DA045	A. solani s.l	Potato	Mascara	-	+	S	-	_
DA046	A. solani s.l	Potato	Mostaganem	-	+	S	-	-
DA047	A. grandis	Potato	Mostaganem	-	+	G	MH243771	_
DA048	A. solani s.l	Potato	Mostaganem	-	+	S	_	_
DA049	A. solani s.l	Potato	Chlef	-	+	S	_	_
DA050	A. solani	Potato	Chlef	-	+	S	MH243772	MH243809
DA051	A. grandis	Potato	Chlef	-	+	G	_	_
DA052	A. grandis	Potato	Chlef	-	+	G	MH243773	_

(Continued)

Table 1. (Continued).	•
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Isolate	Speciesª	Host	Geographic origin	PCR OATF4/ OATR2	PCR OAsF7/OAsR6	PCR RFLP Profile ^b	CAL	Rpb2
DA053	A. solani s.l	Potato	Chlef	-	+	S	_	_
DA054	A. grandis	Potato	Mostaganem	-	+	G	MH243774	_
DA055	A. grandis	Potato	Mostaganem	-	+	G	_	_
DA056	A. solani s.l	Potato	Mostaganem	-	+	S	_	_
DA057	A. solani	Potato	Mascara	-	+	?	MH243791	MH243819
DA058	A. solani s.l	Potato	Mascara	-	+	S	_	_
DA059	A. solani s.l	Potato	Mostaganem	-	+	S	_	_
DA060	A. grandis	Potato	Mostaganem	-	+	G	MH243775	_
DA061	A. grandis	Potato	Mostaganem	-	+	G	_	_
DA062	A. protenta	Tomato	Mostaganem	-	+	S	MH243797	MH243810
DA063	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA064	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA065	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA066	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA067	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA068	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA069	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA070	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA071	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA072	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA073	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA074	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA075	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA076	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA077	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA078	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA079	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA080	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA081	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA082	A. protenta	Potato	El Oued	-	+	?	KX870505	KX870507
DA083	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA084	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA085	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA086	A. protenta	Potato	El Oued	-	+	S	KX870506	KX870508
DA087	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA088	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA089	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA090	A. solani	Potato	El Oued	-	+	S	MH243776	MH243811
DA091	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA092	A. solani s.l	Potato	El Oued	-	+	S	_	_
DA093	A. solani s.l	Potato	Alger	-	+	S	_	_
DA094	A. solani	Potato	Alger	-	+	S	MH243798	MH243812
DA096	A. solani s.l	Potato	Alger	-	+	S	_	_
DA097	A. solani s.l	Potato	Alger	-	+	S	_	_
DA098	A. solani s.l	Potato	Alger	-	+	S	_	_
DA099	A. grandis	Tomato	Alger	-	+	G	MH243777	_
DA100	A. linariae	Tomato	Alger	+	-	ND	MH243807	-

(Continued)

Table 1	I. (Cont	inued).
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Isolate	Speciesª	Host	Geographic origin	PCR OATF4/ OATR2	PCR OAsF7/OAsR6	PCR RFLP Profile ^b	CAL	Rpb2
DA101	A. linariae	Tomato	Alger	+	-	ND	MH243778	_
DA102	A. linariae	Tomato	Alger	+	-	ND	_	_
DA103	A. linariae	Tomato	Alger	+	-	ND	_	_
DA104	A. linariae	Tomato	Alger	+	-	ND	_	_
DA107	A. linariae	Tomato	Alger	+	-	ND	_	_
DA108	A. solani s.l	Potato	Guelma	-	+	S	_	_
DA109	A. linariae	Tomato	Tipaza	+	-	ND	MH243779	_
DA110	A. linariae	Tomato	Tipaza	+	-	ND	_	_
DA111	A.linariae	Tomato	Tipaza	+	-	ND	MH243780	_
DA112	A. linariae	Potato	Skikda	+	-	ND	_	_
DA113	A. solani	Tomato	Guelma	-	+	S	MH243799	MH243821
DA114	A. solani	Potato	Guelma	-	+	?	MH243788	MH243813
DA115	A. solani s.l	Potato	Bouira	-	+	S	_	_
DA116	A. solani s.l	Potato	Bouira	-	+	S	_	_
DA117	A. linariae	Tomato	Biskra	+	-	ND	_	_
DA118	A. linariae	Tomato	Biskra	+	-	ND	_	_
DA119	A. linariae	Tomato	Biskra	+	-	ND	MH243800	_
DA120	A. linariae	Tomato	Biskra	+	-	ND	_	_
DA121	A. linariae	Tomato	Biskra	+	-	ND		
DA122	A. linariae	Tomato	Biskra	+	-	ND	_	—
DA123	A. solani s.l	Tomato	Biskra	-	+	S	_	—
DA124	A. linariae	Tomato	Biskra	+	-	ND	—	—
DA125	A. linariae	Tomato	Biskra	+	-	ND	—	—
DA126	A. linariae	Tomato	Biskra	+	-	ND	-	—
DA127	A, linariae	Tomato	Biskra	+	-	ND	_	-
DA128	A, solani s.l	Tomato	Biskra	-	+	S	_	-
DA129	A. solani s.l	Tomato	Biskra	-	+	S	-	-
DA130	A solani s l	Tomato	Biskra	_	+	S	-	-
DA131	A solani s l	Tomato	Biskra	_	+	s	-	-
DA132	A solani s l	Tomato	Biskra	_	+	S	_	_
DA133	A solani s l	Tomato	Biskra	_		S	_	_
DA134	A linariae	Tomato	Biskra		-	ND	-	_
DA135	A solani sl	Tomato	Biskra	I	-	S	-	_
DA135	A. soluni s.i	Tomato	Diskra	-	Ŧ	S	- MU242901	- MU242814
DA130	A. solani cl	Tomato	Diskia	-	+	S	WII1243001	M11243014
DA137	A. solani s.l	Tomato	Diskra	-	+	5	-	_
DA138	A. solani s.l	Tomato	Diskra	-	+	S	-	_
DA139	A. solani s.l	Tomato	Diskra	-	+	S	-	- MU242015
DA140	A. solani	Tomato	Diskra	-	+	5	MH243802	MH243815
DA141	A. solani	Tomato	Biskra	-	+	8	MH243803	MH243816
DA142	A. solani	Iomato	Biskra	-	+	8	MH243804	MH243817
DAI43	A. solani s.l	Tomato	Biskra	-	+	S	-	-
DA144	A. linariae	Tomato	Biskra	+	-	ND	-	_
DA145	A. solani s.l	Tomato	Biskra	-	+	S	_	-
DA146	A. solani s.l	Potato	Tipaza	-	+	S	_	-
DA147	A. linariae	Tomato	Tipaza	+	-	ND	MH243781	_
DA148	A. linariae	Tomato	Mostaganem	+	-	ND	_	_
DA149	A .linariae	Tomato	Mostaganem	+	-	ND	_	

(Continued)

Isolate	Speciesª	Host	Geographic origin	PCR OATF4/ OATR2	PCR OAsF7/OAsR6	PCR RFLP Profile ^b	CAL	Rpb2
DA150	A. linariae	Tomato	Mostaganem	+	-	ND	_	_
DA152	A. grandis	Tomato	Alger	-	+	G	MH243782	_
DA153	A. linariae	Tomato	Alger	+	-	ND	MH243783	_
DA154	A .grandis	Tomato	Alger	-	+	?	MH243787	_
DA155	A. grandis	Tomato	Alger	-	+	G	MH243784	_
DA156	A. grandis	Tomato	Alger	-	+	G	MH243785	-

Table 1. (Continued).

^a Species names in bold characters were supported by sequence data; *A. solani s.l* stands for *A. solani sensu lato* (i.e. including *A. protenta*). ^b PCR-RFLP profiles: S = typical of *S. solani*; G = typical of *A. grandis*; ND = not determined; ? = atypical restriction profile



Figure 3. Phylogenetic tree reconstructed by the maximum likelihood method from the alignment of calmodulin sequences of 42 isolates belonging to the *Alternaria* section *Porri*. Bootstrap support values greater than 80 % are indicated by arrows. The calmodulin sequences of the following strains were included as references: *A. linariae* CBS 109156 (GenBank number JQ646257), *A. protenta* CBS 116696 (GenBank number JQ646236), *A. solani* CBS 109157 (GenBank number KJ397981), *A.grandis* CBS 109158 (GenBank number JQ646249). The calmodulin sequence from *A. alternata* EGS 34016 (Genbank number JQ646208) was used to root the tree.

Distribution and prevalence of the large-spored Alternaria species in the bioclimatic regions of Algeria

The surveys carried out through the different Algerian potato and tomato growing regions revealed the



Figure 4. Phylogenetic tree reconstructed by the maximum likelihood method from the alignment of RPB2 sequences of 17 isolates belonging to cluster B2 in the calmodulin phylogeny. Bootstrap support values greater than 80 % are indicated by arrows. The RPB2 sequences of the following strains were included as references: *A. linariae* CBS 105.41 (GenBank number KJ718353), *A. solani* CBS 106.21 (GenBank number KJ718410) *A. protenta* CBS 347.79 (GenBank number KJ718392). The RPB2 sequence from *A. alternata* EGS 34016 (Genbank number JQ646490) was used to root the tree.

presence of *A. grandis, A. linariae* and the species complex *A. solani-A. protenta* at variable levels. The frequencies of these species varied according to the host plants (potato or tomato) and to the geographical locations (Table 2). *Alternaria solani sensu lato* (i.e. including *A. protenta*) was the most common species in Algeria, representing 65% of the large-spored isolates, followed by *A. linariae* (25%) and *A. grandis* with low isolation frequency (10%). Isolation frequencies of these species varied with the sampled crop, although they were all found on both potato and tomato. *Alternaria solani sensu lato* (80%) and *A. grandis* (64%) were more prevalent on potato than on tomato. Conversely, *A. linariae*

Table 2. Frequencies of the four large-spored Alternaria speciesaccording to the host plants (potato and tomato) and to the geo-graphical locations

Origin	A. solani- A protenta	A. grandis	A. linariae	
Host plant origin				
Potato	$80.2^{a} - 82.8^{b}$	64.3-9.6	18.9–7.5	
Tomato	19.8-35.2	35.7-9.2	81.1-55.6	
Geographic origin area)	(bioclimatic			
Mediterranean	16.7-39.0	50.0-17.0	48.6-44.0	
Semi-arid	34.3-70.2	50.0-14.9	18.9–14.9	
Arid	49.0-79.7	0-0	32.4-20.3	

^a Percentage of the total number of isolates of the same species according to the host plant or the geographic origin

^b Percentage of the total number of isolates from the same host or from the same geographic origin

was more frequently isolated from tomato than from potato. Locations of sample collection could be divided into three temperature and precipitation zones (Mediterranean, semi-arid or arid). In Mediterranean and arid zones, isolates were almost equally distributed between potato and tomato. Alternaria linariae was over-represented in the Mediterranean zone (44% of the collected isolates) compared to the arid zone where it represented only 20% of the collected isolates. The reverse situation was observed for A. solani sensu lato. Alternaria grandis was not isolated from samples collected from the arid zone, although it represented 15% of isolates from the semi-arid zone and 17% of those from the Mediterranean zone. Only small-spored Alternaria isolates were obtained from samples in two locations, i.e. Ain Defla and Laghouat.

Pathogenicity and parasitic specialization of Alternaria species on potato and tomato

Fourteen isolates representing the four large-spored *Alternaria* species were used in the pathogenicity tests. These were: *A. solani* (isolates DA008, DA114, DA140 and DA141), *A. grandis* (isolates DA009, DA060, DA099 and DA152), *A. protenta* (DA062, DA082 and DA086) and *A. linariae* (DA002, DA007 and DA153). The symptoms (necroses surrounded by yellow halos) obtained on detached leaflets in Petri dish assays were confirmed by high degrees of aggressiveness obtained on whole seedling plants of potato and tomato (Figure 5). All the tested isolates produced symptoms on both plant species irrespective of their host plant of origin (Figure 6). The



Figure 5. Symptoms on detached leaflets (a, d) and whole tomato (c, e) or potato (b, f) plants inoculated by *A. linariae* isolate DA002 (a, b and c) or by *A. grandis* isolate DA009 (d, e and f)



Figure 6. Disease severity evaluated using a visual rating scale (0-100%) at 15 days post-inoculation of tomato (black bars) and potato (grey bars) leaves with isolates representing the four large-spored *Alternaria* species. Letters (P or T) under the isolates identities refer to the host of origin (Potato or Tomato).

degrees of aggressiveness for each host/pathogen combination varied for the isolates, however. On average, *A. solani* isolates were more aggressive on both tomato and potato, while *A. protenta* isolates were less aggressive with a marked preference for their original host plants.

DISCUSSION

Early blight has long been attributed to large-spored *Alternaria* species, i.e. *A. solani* and *A. grandis* on potato and *A. solani* and *A. linariae* on tomato. Small-spored species related to *A. alternata* are considered to be responsible for the brown spot disease (Nolte, 2008, Tymon *et al.*, 2016). The surveys carried out in Alge-

ria between 2012 and 2015 have shown the presence of several Alternaria species producing large conidia in samples from both hosts with typical early blight symptoms, as well as in samples with symptoms that resemble brown spot. On the other hand, many Algerian isolates of the section Alternaria collected by Bessadat et al. (2017) have caused typical early blight symptoms after artificial inoculations on tomato, and were therefore considered pathogenic but with low aggressiveness. This complicated the attribution of a given symptom to a given Alternaria species. Moreover, morphological similarities between species within the Alternaria section Porri precludes reliable identification at the species level based only on these criteria. In the present work, molecular tools were used to confirm the identity of large-spored isolates originating from tomato and potato leaves. The diagnostic species-specific primers developed by Gannibal et al. (2014) were successfully used to quickly characterized 147 isolates from a collection the 155 largespored isolates, and to distinguish A. linariae from isolates that could correspond to either A. solani, A. grandis or A. protenta. No amplification signal was obtained for eight isolates either due to poor quality of the extracted DNA or because these isolates corresponded to another species within section Porri. Although never previously observed in Algeria, it has recently been reported that A. macrospora may be responsible for leaf spot of tomato in China (Zhang et al., 2017). To discriminate between A. grandis, on one hand, and A. solani or A. protenta, on the other, we took advantage of sequence polymorphism within the calmodulin gene to develop a PCR-RFLP assay. Reliability of these diagnostic methods was checked by sequencing the calmodulin locus, and taken together they allowed us to accurately type more than 90% of the large-spored isolates without the need for nucleotide sequencing. Applying these tools to our isolates, we observed that A. solani and A. linariae predominate on their respective potato or tomato hosts in most of the surveyed areas. In agreement with recent observations (Bessadat et al., 2017, Ayad et al., 2018), the presence of A. linariae and A. grandis in Algeria, traditionally considered to be restricted, respectively, to either tomato or potato, was also confirmed on these two plant hosts. Alternaria linariae was found on potato and A. grandis on tomato in the coastal regions of Algiers and Tipaza, where these two crops often exist side by side or follow each other in the same plot during the seasons. Alternaria grandis has not been isolated from the Saharan regions that are characterized by warm and arid periods during each year. Analyses of nucleotidic polymorphism at the RPB2 locus has shown that A. protenta was recorded for the first time on tomato in the coastal region of Mostaganem (northwest of Algeria), where the climate is very different from that of the South. This species has already been reported on potato in Belgium (Landschoot *et al.*, 2017), but also in a particular region of the south of Algeria (El Oued), where the temperatures are high and the potato crops are grown under overhead irrigation (Ayad *et al.*, 2017).

Cross inoculations carried out with isolates representing the four identified large-spored Alternaria species (A. solani, A. grandis, A. linariae and A. protenta) on potato and tomato, showed that aggressiveness toward the two host plants varied according to the isolates but not with respect to particular fungal species. Similar experiments carried out on detached leaves with A. linariae and A. grandis (Rodrigues et al., 2010; Cardoso, 2014; Gannibal et al., 2014) have previously shown that these two species were able to cause disease on both plants with more severe symptoms on their respective traditional hosts. This suggests that the parasitic specialization of large-spored Alternaria species on solanaceous crops should be reconsidered.

ACKNOWLEDGEMENTS

The authors thank the UMR 1345 IRHS, Université d'Angers-INRA-Agrocampus Ouest (France) for hosting D. Ayad, and the Ministry of Higher Education and Scientific Research of Algeria for awarding her scholarship.

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