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1 Detection and molecular characterization of a 16SrI-F phytoplasma in

potato showing purple top disease in Ecuador

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- 4 Samanta Paltrinieri¹, Johanna Buitrón Bustamante², Assunta Bertaccini¹, Carmen Castillo
- 5 Carrillo²
- 6 ¹Alma Mater Studiorum- University of Bologna, DipSA, Plant Pathology, viale G. Fanin,
- 7 42, 40127 Bologna, Italy
- 8 ²Estación Experimental Santa Catalina, Instituto Nacional de Investigaciones
- 9 Agropecuarias (INIAP), Departamento Nacional de Biotecnología y Departamento
- 10 Nacional de Protección Vegetal. Panamericana Sur km 1, Quito, Ecuador
- 11 Corresponding author: Carmen Castillo Carrillo, email address:
- 12 carmen.castillo@iniap.gob.ec

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Abstract

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- Potato purple top (PPT) disease has been reported worldwide, including Canada, United
- 17 States, Mexico, Russia and Pakistan. During surveys carried out in a recent epidemic in
- 18 Ecuador of potato purple top disease phytoplasmas were identified in symptomatic potatoes
- as well as in some tissue cultured potato shoots by PCR/RFLP analyses and by sequencing
- and phylogenetic clustering. The majority of the samples resulted positive to phytoplasmas
- 21 enclosed in the ribosomal subgroup 16SrI-F, for the first time detected in potato and in the
- 22 American continent. Further studies will allow verification of the transmission and further
- presence in Ecuador of this phytoplasma associated with a purple top disease of potatoes.

25 Keywords

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27 Phytoplasmas, Purple top disease, Molecular detection, Sequencing, Potato

Introduction

Potato purple top (PPT) disease has been reported worldwide, including Canada, United States, Mexico, Russia and Pakistan causing significant economic losses (Nasir 2007, Lee et al. 2004, Santos-Cervantes et al. 2010. Several phytoplasmas from 16Sr groups and subgroups have been reported as agents associated with PPT (Lee et al. 2004, Santos-Cervantes et al. 2010, Caicedo et al. 2015). In 2013, in the northern potato production region of Ecuador, symptoms of PPT appeared in the fields on scattered plants. Two years later the disease was more than 80% in the Superchola variety fields, with losses reaching up 50% (INIAP-2014, E. Reina, personal communication). Characteristic symptomatology of this disease such as yellow and purple coloration of the upper leaflets, apical leaf curling, axillary buds, aerial tubers and early senescence starts to appear after flowering (Fig. 1). In 2015, 'Candidatus Phytoplasma aurantifolia' (16SrII group) was reported associated with PPT in Ecuador (Caicedo et al. 2015). Considering that the disease was present in several localities, surveys were carried out from asymptomatic and symptomatic potato plants collected in the northern provinces of

17	Carchi and Pichincha-in the highlands in 2015 and 2016 to further confirm phytoplasma

48 presence, identity and association with the disease.

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Materials and Methods

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Symptomatic (Fig. 1) and asymptomatic potato tissues were-collected from different localities and from diverse portions of the plant (Table 1) and stored at -80°Cs-Micropropagated material was also tested-DNA was extracted using a CTAB method described by Ferreira and Grattapaglia (1998). The universal phytoplasma primer pair P1/P7 (Deng and Hiruki 1991; Schneider et al. 1995) was used to amplify a 1.8 kb fragment of the ribosomal operon consisting of the 16S rRNA gene, the 16S-23S spacer region and a portion of the 5' region of 23S rRNA gene. A 1: 30 dilution of the PCR product was used as template for nested PCR with the primer pair R16(I)F1/R1 which amplifies an internal DNA fragment of 1,100 bp in the 16S rRNA gene (Lee et al. 1994). After the initial denaturation step of 94°C for 1 min, PCR was performed for 35 cycles in an automated thermal cycler (Biometra, Germany) at 94°C for 1 min, 55°C for 2 min (50°C for nested PCR) and 72°C for 3 min, followed by a final extension step at 72°C for 7 min. Phytoplasma positive controls included DNA from phytoplasma strains maintained in periwinkle [Catharanthus roseus (L.) G. Don.] (Bertaccini, 2014): primula yellows from Germany (PRIVA, 16SrI-L); aster yellows from USA (AY, 16SrI-B); Achillea yellows from Italy

(ACH, 16SrI-C) and aster yellows from apricot from Spain (A-AY, 16SrI-F). Samples devoid of DNA template were added as negative controls for the direct and nested PCR reactions. PCR mixtures (25 µl) containing 20-60 ng of total DNA, 0.5 µM of each primer, 0.025 U of Taq DNA polymerase (Sigma Aldrich), 1X PCR buffer containing 1.5 mM MgCl₂ and 0.2 mM dNTPs. PCR products (6 μl) were electrophoresed in 1% agarose gel, stained with ethidium bromide and visualized with a UV transilluminator. RFLP analyses of amplified fragments were performed using 100-200 ng DNA per sample that was digested separately with the restriction enzymes Tru1I, RsaI and TaqI (Fermentas, Vilnius, Lithuania and New England Biolabs) according to the manufacturer's instructions and the digested amplicon. The restriction products were then separated by electrophoresis through a 6.7% polyacrylamide gel-stained and visualized as described above. Direct sequencing was carried out in both directions from selected positive samples with primers R16(I)F1/R1 after purification with a QIAquick PCR Purification Kit (QIAGEN, Valencia, CA). The sequences obtained were assembled by MEGA6 (Tamura et al. 2013) for phylogenetic analysis. The evolutionary history was inferred using the Maximum Likelihood method based on the Tamura-Nei substitution model (Tamura and Nei 1993). Initial tree(s) for the heuristic search were obtained automatically by applying Neighbor-Joining and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. Evolutionary analyses were conducted in MEGA6. Further PCR product analyses were carried out to verify the presence of 'Candidatus Liberibacter solanacearum' (Lso) using primers OA2 and OI2c (Liefting et al.

2008,; 2009) according to the methodology described by Crosslin et al. (2011) and

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employing the *Taq* DNA polymerase recombinant (Termo Fischer Scientific). PCR products were visualized using agarose gel electrophoresis. Positive controls were obtained from Dr Joseph Munyaneza's laboratory at USDA-ARS in WA, USA, which showed clear bands corresponding to the expected 1168-bp products (Liefting et al. 2009), with negative controls as described above. Further verification was performed with the same primers and under the same cycling conditions using 0.16 μl (1.25 units) Promega GoTaq G2 DNA polymerase (Promega) and using primers Lso TX 16/23 F/R under reported conditions which showed the expected 385 bp product in the control samples (Ravindran et al. 2011).

Results

The direct amplification of phytoplasma DNA did not result in any detectable positive reaction from any of the potato samples, while only the positive control DNAs -resulted in the expected 1,800 bp amplicons. However, the use of nested PCR with R16(I)F1/R1 primers resulted in the expected 1,100 bp amplicons in the majority of the samples tested from symptomatic field plants and from micropropaged shoot. In some cases the nested PCR results were in contrast to the symptomology. In particular, symptomatic tissue from one aerial tuber, one stolon and one micropropagated shoot gave a negative result while asymptomatic tissue from a sprout, petiol and one micropropagated shoot

115	gave a positive result (Table 1). The negative
116	controls were always negative in both direct and nested PCR (data not shown).
117	RFLP and sequence analysis verified that the Ecuadorian phytoplasmas from potato
118	samples were similar to A-AY control sample from Spain (Lee et al. 1998). The RFLF
119	profile with restriction enzymes Tru1I, RsaI and KpnI was
120	identical to the A-AY control (Fig. 2). Likewise, DNA
121	sequences-from samples 7, 12 and 72 (GenBank under accession numbers
122	MG272306, MG272307, MG272308, respectively).
123	clustered with the A-AY strain (GenBank accession number
124	X68338) (Fig. 3) confirming the RFLP results.—These results sugest that the
125	phytoplasma identified in the potato samples in the aster yellows
126	cluster, and in subgroup 16SrI-F.
127	No bands were obtained in the testing for 'Ca. L. solanacearum' using all above
128	reported primers and conditions from the tested samples, except for the positive controls
129	(data not shown).
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132	Discussion

The epidemiology of PPT in Ecuadorian fields corresponds

Diseases caused by other pathogens like Rhizoctonia, which

to a complex of new emergent pathogens such as phytoplasmas located in the plant phloem,

spread by infected seed and insect-vectors and mainly detected through molecular analyses,-

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produces sclerotia on tubers, rot and blight in the stems, and aerial tubers near the base of the plant (main factor that misleads farmers to identify the causal agent of purple top in potato plants in Ecuador), do no present the same symptoms in the field as phytoplasma. At the high point of the purple top outbreak in Carchi province in March of 2015, symptoms were visualized in more than 80% of the fields of the potato production area (AGROCALIDAD 2015). The rapid spread of the symptomatic plants does not correspond to the epidemiology of other pathogens such as Rhizoctonia;. Although, Rhizoctonia was, in fact, isolated from some PPT symptomatic plants along with the viruses PVY, PVX and PRLV, and Fusarium cylindrocarpun, Pectobacterium sp. and Pseudomonas viridiflava (Castillo et al. 2017), the epidemiological behaviors and the symptoms correspond to infections associated with phytoplasmas presence. The transmission of PPT suggests more similarity to_-pathogens transmitted by insect vectors and/or spread by infected propagation materials than-to a spreading of a soil fungus like Rhizoctonia. A previous study detected the presence of 'Candidatus Phytoplasma aurantifolia' (16SrII group) (Caicedo et al. 2015). Our study allows to verify the presence of another phytoplasma enclosed in 16SrI-F subgroup. Phytoplasmas enclosed in this subgroup has only been reported in Spain (strain ACLR-AY = A-AY) and in Germany (strain CVB) (Lee et al. 1998; Bertaccini, 2014) in apricot and periwinkle respectively. Our study is the first report of the 16SrI-F subgroup -in potatoes. Past studies have shown that similar symptomatology has resulted in the detection of both

'Candidatus Phytoplasma' and 'Candidatus Liberibacter solanacearum' in potatoes

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(Gutiérrez et al. 2013) and in carrots (Satta et al. 2017). ¬

Nevertheless the results of the present survey

indicate that 'Ca. L. solanacearum' is not currently present in Ecuador. More research is needed to verify if this phytoplasma is inducing the symptomatology observed in the field although strong association was observed in the present survey. A pathogen complex might be involved in producing purple top symptoms, and therefore further research on other possible prokaryotes together with presence of the insect vector or vectors that may transmit these pathogens in the field is necessary, along with a broader sampling in time and space.

Acknowledgments

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Table 1. Nested PCR detection of phytoplasma in potato from

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ID	Plant part	Dumple to-	Duovinas	Altitude	Primer
ID		Purple top	Province	(m asl)	16SrI
3	Petiol	Symptomatic	Carchi	3070	positive
7	Sprouts	Symptomatic	Carchi	2991	positive
30	Aerial tuber	Symptomatic	Carchi	3070	positive
39	Underground tuber	Symptomatic	Carchi	3070	positive
59	Aerial tuber	Symptomatic	Carchi	3070	positive
60	Aerial tuber	Symptomatic	Carchi	3070	positive
62	Aerial tuber	Symptomatic	Carchi	3070	negative
64	Underground tuber	Symptomatic	Carchi	2991	positive
67	Underground tuber	Symptomatic	Carchi	2991	positive
72	Sprouts ^a	Asymptomatic	Carchi	2991	positive
86	Aereal tuber	Symptomatic	Carchi	3070	positive
12	In vitro plant ^a	Asymptomatic	Laboratory	3050	positive
127	In vitro plant	Asymptomatic	Laboratory	3050	negative
80	Aereal tuber	Symptomatic	Pichincha	3050	positive
88	Stolon	Symptomatic	Pichincha	3050	negative
103	Flower petiol	Symptomatic	Pichincha	3050	positive
105	Petiol	Symptomatic	Pichincha	3050	positive
114	Petiol	Asymptomatic	Pichincha	2945	positive
117	Petiol	Symptomatic	Pichincha	2945	positive

246 | a Sequenced samples 250



Figure 1. (a) Symptomatic plant in the field (initial, red arrow); (b) curly yellow-purple leaf coloration and (c) aerial tubers located in the upper part of the plant.

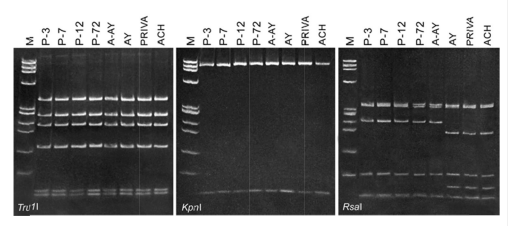


Figure. 2. Polyacrylamide gels 6.7% showing the restriction fragment length polymorphism patterns of R16(I)F1/R1 amplicons from phytoplasma strains from potatoes (P-3, P-7. P12, P72) and controls (A-AY, 16SrI-F; AY, 16SrI-B; PRIVA, 16SrI-L, and ACH, 16SrI-C), the restriction enzymes employed are indicated at the bottom of each gel. M, marker phiX174 DNA digested with *Hae*III length from top to bottom fragments in bp: 1,353; 1,078; 872; 603; 310; 281; 271; 234; 194; 118 and 72.

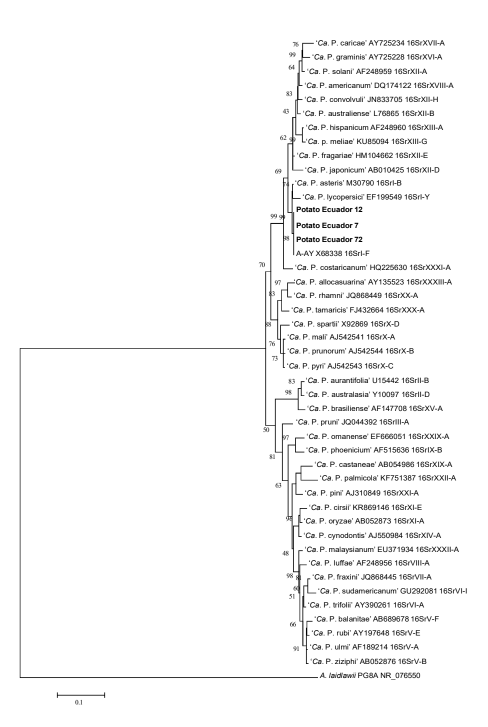


Figure. 3. Molecular Phylogenetic analysis using Maximum Likelihood method. http://purl.org/phylo/treebase/phylows/study/TB2:S21871 The tree with the highest log likelihood (-6080.8085) is shown. Bootstrap values above 40 are shown. Forty-six closely related nucleotide sequences from the 'Candidatus Phytoplasma' species and aster yellows (16Srl) The sequences of phytoplasmas from potatoes from Ecuador obtained in this work are in bold. On the right of the branch the GenBank accession number of the strains employed and the ribosomal group/subgroup. Acholeplasma laidlawii was used as out group to root the tree.