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Detection and identification of a 16SrIII-J subgroup phytoplasma associated with faba bean in Peru

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Published Version:

Torres Suarez G., Gamarra Gamarra D., Villar C.M., Llacza Munive S.L., Satta E., Carrasco Lozano E.C., et al. (2021). Detection and identification of a 16SrIII-J subgroup phytoplasma associated with faba bean in Peru. JOURNAL OF PHYTOPATHOLOGY, 169(4), 203-208 [10.1111/jph.12977].

Availability:

This version is available at: https://hdl.handle.net/11585/846401 since: 2022-01-19

Published:

DOI: http://doi.org/10.1111/jph.12977

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1	Detection and identification of a 16SrIII-J subgroup phytoplasma associated
2	with faba bean in Peru
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19	KEYWORDS: Vicia faba, 16SrIII phytoplasma, dwarfism, abnormal leaf sprouts,
20	Peru.
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25 Abstract

Faba bean samples with symptoms of yellowing, dwarfism, shoot proliferation, 26 internode shortening, leaf sprouts and lack of pod and seed production were 27 collected from Huancayo and Chupaca provinces, Junin-Peru, and analyzed to 28 verify phytoplasma presence and identity. After total nucleic acid extraction, the 29 amplification with universal phytoplasma primers, using nested polymerase chain 30 reactions, on the 16S ribosomal gene followed by restriction fragment length 31 polymorphism analysis and sequence analysis allowed the classification of the 32 33 detected phytoplasma in the subgroup 16SrIII-J. The alignment of 30 16S ribosomal gene sequences from 15 faba bean symptomatic samples from single 34 plants allowed verifying the consistent presence of 5 single nucleotide 35 polymorphisms that are however not modifying the phytoplasma classification. The 36 phytoplasma identity was also corroborated by the amplification and the restriction 37 fragment length polymorphism analyses carried out on the ribosomal protein gene 38 amplicons obtained with primers specific for the phytoplasmas enclosed in the 39 16SrIII group. This is the first description in Peru of a disease associated with 40 phytoplasmas in faba beans. 41

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43 2 | INTRODUCTION

Vicia faba L., known as faba bean, is a leguminous species originated in the
Mediterranean region and grown worldwide for its high content of carbohydrate,
protein, vitamins, minerals (Zn, Fe, Mg, Ca) and omega-3 (Etemadi et al. 2019). In
2017, Peru had a cropping area of 65,985 ha, with an average yield of 11,478
kg/ha and total seed production of 75,736 tons (FAOSTAT 2019). Faba bean is

worldwide infected by various fungi, bacteria, chromista, viruses, and nematodes. 49 The main diseases reported are chocolate spot (caused by Botrytis fabae), leaf 50 blight (Xanthomonas campestris), bacterial spot (Pseudomonas syringae), root rot 51 (Fusarium solani), powdery mildew (Erysiphe cichoracearum), and infections by 52 viral complexes (Etemadi et al. 2019). During the last decades V. faba was also 53 reported to be infected by phytoplasmas (Castro and Romero 2004; Al-Saleh and 54 Amer 2014; Salehi et al. 2016; Omar 2017), that are bacteria discovered in 1967 55 56 (Doi et al. 1967) lacking the cell wall and having a diameter of 200 to 800 nm. Their 57 genomes are of small size and possess variable numbers of extrachromosomal fragments (Nishigawa et al. 2003). Phytoplasmas are hosted in the plant phloem 58 and in the hemolymph of their insect vectors that are mainly those having siphon-59 like mouth systems, as leafhoppers and psyllids (Weintraub and Beanland 2006). 60 They infect hundreds of plant species in which different symptoms can be 61 different phytoplasmas 62 observed; moreover, could be associated with indistinguishable symptomatology in diverse plant host species (Bertaccini 2007, 63 Namba 2019). These phytoplasma-associated symptoms include leaf yellowing, 64 growth reduction, virescence, and witches' broom, shortening of internodes, small 65 leaves, phyllody, floral malformation, and plant death (Salehi et al. 2016; Kumari et 66 al. 2019). 67

Since their discovery it was difficult to detect and identify the phytoplasmas in symptomatic plants since they are present in a reduced concentration in the infected tissues where are unevenly distributed (Berges et al. 2000). Phytoplasmas belong to the class *Mollicutes* provisionally enclosed in the '*Candidatus* Phytoplasma' genus (IRPCM 2004). PCR/RFLP analyses on their 16S ribosomal

gene allowed the phytoplasma classification in ribosomal groups and subgroups
that is integrating the provisional classification for further differentiation among the
detected strains (Bertaccini and Lee 2018).

In recent years in the faba bean fields located in the Junin region, symptoms related to phytoplasma presence have been observed with 8% incidence, however, to date the possible presence and identity of phytoplasmas was not studied. Therefore, the objective of this work was to identify the phytoplasma associated with the disease of the faba bean in the provinces of Huancayo and Chupaca in the Junin region of Peru.

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83 2 | MATERIALS AND METHODS

84 **2.1**| Sample collection and DNA extraction

Faba bean samples with symptoms of yellowing, dwarfism, shoot proliferation, 85 internode shortening, leaf sprouts and lack of pod and seed production (Figure 1a-86 b) were collected from Huancayo (15 samples) and Chupaca provinces (16 87 samples), Junin-Peru, and analyzed to verify phytoplasma presence and identity. 88 The 31 samples each collected from each from a separate plant, were processed 89 and dehydrated with silica gel, then leaf tissues were sprayed with liquid nitrogen, 90 and 25 mg were weighed for the extraction of DNA with innuPREP Plant DNA Kit 91 (Analytik Jena, Germany) according to the instructions of the manufacturer. For the 92 93 negative control, DNAs of four symptomless plants were also extracted following the same procedure. As positive control, the DNA of a strain from Euscelidius 94 variegatus (16SrIII-B, API) maintained in the micropropagated collection 95 (Bertaccini, 2014) was used. The quality of the obtained DNA was verified by 96

97 means of a 1% agarose gel and quantified with a spectrophotometer NanoDrop™
98 (ThermoScientific™).

99 2.2| PCR detection and RFLP analysis

Phytoplasma detection was achieved using PCR assays with P1/P7 (Deng and 100 Hiruki 1991; Schneider et al. 1995), R16mF2/mR1 primers followed in nested PCR 101 reaction by R16F2n/R2 (Gundersen and Lee 1996) and fU5/rU3 (Lorenz et al. 102 1995) primers. The total nucleic acid used was about 20-40 ng for the PCR 103 reactions and amplicons, diluted in sterile distilled water 1:30, were used as 104 105 template for the nested PCR reactions. Cycling conditions reported (Schaff et al. 1992) were employed in all the PCR reactions. The amplicon visualization was 106 obtained by electrophoresis in 1% agarose gel followed by ethidium bromide (10 107 ng/µl) staining. The R16F2n/R2 and fU5/rU3 amplicons were analyzed by RFLP 108 using Tru1 and Hpall (Fermentas, Vilnius, Lithuania) restriction enzymes and 109 compared with the corresponding profile of the phytoplasma strain API in 6.7% 110 polyacrylamide gels stained and visualized as for the agarose gels reported above. 111 112 Further molecular characterization was achieved in nested PCR on the ribosomal protein (rp) operon with primer pairs rpL2F3/rp(I)R1A and rp(III)F1/rp(III)R1, 113 specifically amplifying about 800 bp in the rp coding gene of the phytoplasmas 114 enclosed in ribosomal group 16SrIII, under the reported conditions (Martini et al. 115 2007). The following phytoplasma strains enclosed in the ribosomal group 16SrIII 116 117 and maintained in periwinkle (Catharanthus roseus) as a collection of micropropagated shoots (Bertaccini 2014) were used as controls: X disease 118 phytoplasma (16SrIII-A, CX), a strain from *Euscelidius variegatus* (16SrIII-B, API), 119 Vaccinium witches' broom (16SrIII-F, VAC), spirea stunt (16SrIII-E, SPI), and 120

poinsettia branching (16SrIII-H, JRI). The amplicons were analyzed by RFLP with
the restriction enzyme *Tru1*I, observed after electrophoresis and visualized as for
the agarose gels reported above.

124 **2.3**| Sequencing and phylogenetic analysis of 16S rDNA

The extraction from agar of R16mF2/R16mR1 and fU5/rU3 amplified fragments of 125 approximately 1,400 and 876 bp respectively, was performed with the PCR Clean-126 Up System Kit (Promega, USA). Amplicons were then sent for both directions 127 sequencing using the primer pairs employed in the amplification at Macrogen 128 129 (Seoul, Korea). The sequence alignment and manual revision were done with the BLASTn program, followed by the comparison with phytoplasma sequences in the 130 GenBank. The consensus sequence obtained was then subjected to virtual RFLP 131 analysis with the *i*PhyClassifier using all the 17 available restriction enzymes with 132 the 16SrIII subgroup strains (Zhao et al. 2009). Phylogenetic analysis with the 133 Minimum Evolution system (Rzhetsky and Nei 1992) was also carried out using 134 MEGA 6 (Tamura et al. 2013) with the 43 described 'Ca. Phytoplasma' species and 135 with a number of phytoplasma strains enclosing those classified in 16SrIII-J 136 subgroup or reported from Peru and other South American agricultural areas. 137

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140 3| **RESULTS**

From the 31 symptomatic samples collected from individual plants no or weak amplification was obtained with P1/P7 primers while two samples were amplified with primers R16mF2/R16mR1. By using primers amplifying shorter fragments in nested PCR the detection of phytoplasma expected length bands was obtained in

all the samples from symptomatic plants; in particular primers fU5/rU3 (amplicons 145 of approximately 850 bp) provided 100% of amplification from all the samples from 146 symptomatic plants and primers R16F2n/R2 (amplicons of approximately 1,200 bp) 147 provided 80% of amplification (data not shown). The samples from asymptomatic 148 plants and the sterile water used as negative controls did not produce bands in any 149 PCR reaction. From the fU5/rU3 amplified fragments 10 samples were selected for 150 direct sequencing, the remaining fragments were subjected to RFLP analysis with 151 152 Tru1 and Hpall (Figure 2a). The profiles obtained resulted identical to those of the 153 strain from Euscelidius variegatus (16SrIII-B, API) (data not shown). The two 1,400 bp amplicons obtained with R16mF2/R16mR1 were sequenced in both directions 154 and the consensus sequence of 1,320 bp (4 times coverage for each nucleotide in 155 each position) was deposited in the GenBank under the accession number 156 MT358307; this strain is referred here as faba bean yellows and witches' broom 157 (FB-YWB) phytoplasma. The R16F2n/R2 sequence (1,240 bp) analyzed in silico 158 RFLP by the *i*PhyClassifier (Figures 2b, c and d) was having a 1.00 as similarity 159 160 coefficient to the phytoplasma strains enclosed in the subgroup 16SrIII-J. The phylogeny allowed clustering the FB-YWB sequence with phytoplasma strains 161 enclosed in this ribosomal subgroup detected in other geographic areas of South 162 America (Figures 3a and b). The alignment of 30 16Sr sequences amplified with 163 fU5/rU3 primers (about 800 bp) from the faba bean phytoplasma strains detected 164 165 in Peru showed the consistent presence of three SNPs (*i.e.* present in more than three sequences) at positions 384 (A/G), 442 (C/T), 831 (C/A) and of three 166 insertions T at 492, C at 827, G at 1015 counted 3' to 5' in the deposited 167 consensus sequence (data not shown). The specific amplification of the 168

phytoplasmas with rp 16SrIII group-specific primers followed by RFLP analyses
with *Tru1*I (Figure 4), corroborated the close genetic relationship of this
phytoplasma with those classified in the 16SrIII subgroup.

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173 4 DISCUSSION

The detection of 16SrIII-J in symptomatic faba bean samples is the first 174 identification of phytoplasmas enclosed in this ribosomal subgroup in Peru; they 175 176 were reported first in Brazil (Montano et al. 2000) and later infecting several other 177 crop species in South American states such as Argentina (Galdeano et al. 2013; Fernández et al. 2020), Brazil (Munhoz et al. 2019), and Chile (Quiroga et al., 178 2020). The identification of variability at the 16Sr level is not unusual in 179 phytoplasmas classified in the 16SrIII group where the frequent presence of 180 interoperon heterogeneity is reported in several strains (Davis et al. 2013). 181

The first studies of phytoplasmas in Peru date back to the late 1970s (Nault 182 et al. 1979). Based on serological and microscopic techniques, the presence of 183 phytoplasmas, spiroplasmas, and viruses in corn crops was verified. After that, a 184 survey of phytoplasmas in different regions and crops allowed identifying 185 phytoplasmas enclosed in ribosomal group 16Srl in diverse plants species, of 186 group 16SrII in potato (Solanum tuberosum), while phytoplasmas in the 16SrIII 187 group were detected in tomato (Solanum lycopersicum) and dandelion (Taraxacum 188 189 sp.) (Hodgetts et al. 2009). More recently, a phytoplasma classified in the 16SrXV-A was detected in grapevine (Wei et al. 2017). 190

191 Phytoplasmas have not been identified in faba bean in Peru previously, 192 however, a phytoplasma enclosed in the 16SrIII group was reported in Spain

(Castro and Romero 2004), while in Saudi Arabia (Al-Saleh and Amer 2014, Omar, 193 2017) and Iran (Salehi et al. 2016) 16SrII phytoplasma strains were identified in 194 symptomatic plants. The identified phytoplasma is largely reported in South 195 American countries it is therefore necessary to continue the studies to verify the 196 presence of insect vectors following the recent demonstration of Paratanus 197 exitiousus and Bergallia valdiviana in Chile (Quiroga et al. 2019). Furthermore, 198 assays to verify a possible seed transmission, reported for other phytoplasmas in 199 200 tomato, corn (Zea mays) and carrot (Daucus carota) (Calari et al. 2011; Satta et al. 201 2020) should be planned.

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203 ACKNOWLEDGEMENTS

This research was financed by the Vice-Rector's Office for Research (CANON Projects, resolution No. 1565-R-2017) of the Universidad Nacional del Centro del Peru. We are also thankful to the members of the research team of the Center for Plant Molecular Biology Research. Research data are not shared.

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Figure 1 *Vicia faba* plants with symptoms of yellowing, dwarfism, leaf distortion, and abnormal leaf sprouts near to healthy plants (a) and in a close up (b) in a production field in the province of Chupaca, Peru.



Figure 2 a) Polyacrylamide 6.7% gels showing the restriction profiles of 16S 359 ribosomal amplicons of faba bean phytoplasma strains from Peru: from left 360 digestion with Tru1 of R16F2n/R16R2 amplicons followed by digestion of the 361 same samples with the same enzyme of fU5/rU3 amplicons; after the second 362 marker (P) the same amplicons digested with Hpall. Samples 1, 2, and 4, broad 363 bean samples from the diverse localities. b), c) and d) In silico RFLP patterns of 364 the R16F2n/R2 sequence of the FB-YWB strain (GenBank accession number 365 MT358307) in comparison with the 16SrIII available phytoplasma subgroups using 366 the *i*PhyClassifier. Ribosomal subgroups are listed at the top; lane with no name is 367 strain FB-YWB. Tru1I and MseI are isoschizomer enzymes. P and MW, marker 368 ΦX174 digested with HaeIII and having the fragment sizes from top to bottom of 369 370 1,353; 1,078; 872; 603; 310; 281; 271; 234; 194; 118 and 72 base pairs, respectively. 371

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Figure 3 a) and b) Phylogenetic trees showing relationships at the 16S ribosomal 373 RNA gene level of the faba bean yellows and witches' broom phytoplasma strain 374 from Peru (FB-YWB) (in red) compared with the classified 'Ca. Phytoplasma' 375 species and with some phytoplasmas classified in the group 16SrIII (reported 376 ribosomal subgroups are listed on the right in blue) using the Minimum Evolution 377 method. The replicate tree percentage indicating the clustering of the associated 378 taxa are next to the branches (Felsenstein 1985). The Tamura-Nei method (1993) 379 was used to calculate the evolutionary distances that are expressed in base 380 substitutions per site, only values above 50 are reported. The out group employed 381 382 is the Acholeplasma laidlawii. The numbers of the GenBank of the sequences employed are on the right for the 'Ca. Phytoplasma' species and on the left for the 383 strains in the 16SrIII group. 384



387 Figure 4 Restriction fragment length polymorphism patterns of ribosomal protein 388 amplicons from faba bean yellows and witches' broom phytoplasma from Peru (Fb1, Fb2, and Fb4) and other phytoplasma strains classified in the ribosomal 389 group 16SrIII after polyacrylamide 6.7% gel electrophoresis. The amplification was 390 carried out in nested PCR with rpIIIF1/rpIIIR1 primers; the enzyme used is listed at 391 392 the bottom. Lanes 1, X disease (16SrIII-A, CX), 2, strain from Euscelidius variegatus (16SrIII-B, API), 3, Vaccinium witches' broom (16SrIII-F, VAC), 4, spirea 393 stunt (16SrIII-E, SPI), 5, poinsettia branching (16SrIII-H, JRI). P, marker ФX174 394 digested with HaellI and having the fragment sizes from top to bottom of 1,353; 395 1,078; 872; 603; 310; 281; 271; 234; 194; 118 and 72 base pairs. 396