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Characterization of 'Candidatus Phytoplasma solani' associated with a maize leaf reddening disease in Turkey

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1	Multigene characterization of ' <i>Candidatus</i> Phytoplasma solani' associated
2	with a maize leaf reddening disease in Turkey
3	
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11	
12	Abstract
13	In recent years, empty or poor grain bearing corn plants with leaf reddening symptoms have
14	been observed in some commercial maize production areas in Adana province of Turkey. The
15	disease has an ongoing increasing trend and cause economic losses. To verify the possible
16	presence of phytoplasmas, leaf samples were tested with P1/Tint and R16F2n/R2 phytoplasma
17	specific primer pairs after DNA extraction. The 16S ribosomal gene sequence analysis followed
18	by sequence in silico enzyme digestions and phylogenetic analyses, allowed the identification
19	of 'Candidatus Phytoplasma solani', subgroup of 16SrXII-A related-strain in all symptomatic
20	samples. The positive samples were subjected to multigene analyses for the molecular
21	characterization of <i>tuf</i> and <i>vmp1</i> genes. <i>In silico</i> restriction fragment length polymorphism and
22	sequence analyses allowed the inclusion of all the positive samples into tuf-type b and V15

tested samples (n=12). This is the first molecular characterization of a 'Ca. P. solani' strain infecting maize plants in Turkey. The identified pathogen is listed in the EPPO A2 list.

molecular types. The sequence analysis indicated the presence of a single genotype in all the

27 Key words: Zea mays L., phytoplasma, 16SrXII-A subgroup, PCR, tuf, vmp1

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# 30 1 | INTRODUCTION

31 Maize (Zea mays L.) is a widely cultivated annual species in the Poaceae family followed by wheat and rice throughout the world. Maize is generally cultivated in humid 32 regions in most parts of the world where 90% of its production is used for human and livestock 33 feeding. It is grown in almost all regions of Turkey and has a great economic value to the 34 national economy with a production of 6,500,000 tons. Adana province significantly 35 contributes to the Turkish economy with the 12.6% of national maize production (819.978 tons) 36 (Anonymous, 2020). Diseases associated with phytoplasmas in maize have been reported in 37 some European countries including Serbia, Romania, Bulgaria, and more recently north Italy, 38 39 Hungary, Bosnia and Herzegovina, Russia and India (Jović et al., 2009; Calari et al., 2010; Acs et al., 2011; Kovačević et al., 2014; Grigoryevna, 2015; Rao et al., 2017). Maize redness, also 40 known as corn reddening, is a severe disease associated with the presence of 'Ca. P. solani' or 41 42 'Ca.P. asteris' that belongs to the 16SrXII-A and 16SrI-B ribosomal subgroups, respectively (Quaglino et al., 2013; Rao et al., 2017). Maize redness disease is transmitted in Serbia by the 43 cixiid planthoppers, Reptalus panzeri (Löw) and Hyalesthes obsoletus Signoret (Hemiptera: 44 Cixiidae) (Jović et al., 2007; 2009; Mori et al., 2013). Duduk and Bertaccini (2006) reported 45 the first presence of phytoplasmas in corn with reddening symptoms in Serbia, then, maize 46 redness disease associated with the same phytoplasma was reported in Hungary and Bulgaria 47 (Acs et al., 2011; Genov et al., 2014). 'Ca. P. solani' has a wide range of diverse host plants 48 and many of them are of economic importance. Several studies have focused on its molecular 49 identification and multilocus typing, moreover its genetic diversity was evaluated by different 50

protein coding-genes and molecular approaches (Cimerman et al., 2009; Fialová et al., 2009; 51 Quaglino et al., 2019). Based on tuf gene, 'Ca. P. solani' consists of two main strains named 52 tuf-type a and b; while the tuf-type a strain is having an epidemic cycle that involves the stinging 53 nettle (Urtica dioica), the tuf-type b is primarily associated with field bindweed (Convolvulus 54 55 arvensis) (Langer and Maixner, 2004). The gene encoding variable membrane protein 1 (*vmp1*) has been employed for molecular characterization of immunodominant proteins linked to 56 phytoplasma-host interactions (Cimerman et al., 2009); it is more variable and has shown its 57 usefulness for 'Ca. P. solani' epidemiological studies (Murolo et al., 2010; Aryan et al., 2014; 58 Conigliaro et al., 2020). 59

In Turkey, phytoplasma symptoms in maize plants have recently been detected and studied due to their increasing incidence in fields (Çağlar *et al.*, 2016, 2019). In the last two years, symptoms such as redness on leaf veins, empty and poor seed set in corn cobs were commonly observed in maize fields between April and May. This study was, therefore, designed to determine if these symptoms on maize plants are associated with '*Ca*. P. solani' presence, and to verify the possible presence of genetic diversity based on *tuf* and *vmp1* genes in the detected phytoplasma strain(s).

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# 68 2 | MATERIALS AND METHODS

## 69 2.1 | Sample collection

Leaf samples were collected from commercial maize fields expressing reddening symptoms in Adana province, Turkey and tested to verify possible phytoplasma presence and identity. Samples were collected in an area of two hectares, during surveys between April-May 2020. A total of 12 plants exhibiting the symptoms of reddening disease in the midrib of leaf and stem carrying empty corn cob with poor seed set were sampled (three per field) from four neighboring fields (Figure 1). One asymptomatic plant sample from each field was alsocollected as a negative control.

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#### 78 2.2 | Nucleic acid extraction

DNA was extracted to determine the possible phytoplasma occurrence from midribs of 79 fresh leaves following the method of Ahrens and Seemüller (1992) with some minor 80 modifications. In brief, 1 g midrib was homogenized in 4 ml of CTAB buffer pH 8.0 (2% w/v 81 cetyl trimethyl ammonium bromide, 1.4 mol L<sup>-1</sup> NaCl, 20 mmol L<sup>-1</sup> EDTA, 100 mmol L<sup>-1</sup> Tris 82 HCl, 0.2% 2-\beta-mercaptoethanol, 2% polyvinylpyrrolidone-40) followed by an incubation at 83 65°C for 30 min. After elimination of plant tissue residues by centrifugation at 13,000 g for 10 84 min, the nucleic acids in suspension were clarified through chloroform-isoamyl alcohol (24:1), 85 and precipitated with isopropanol at -20°C overnight. The precipitated nucleic acids were then 86 collected by centrifugation at 13,000 g for 15 min, washed with 70% ethanol, dried at room 87 temperature and finally dissolved in 50 µL of sterile double distilled water (SDDW). Before 88 using in polymerase chain reaction (PCR) assays, nucleic acids were quantified and diluted with 89 SDDW to a final concentration of 20 ng/ $\mu$ L. 90

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## 92 2.3 | Detection and molecular characterization of phytoplasma DNA

The phytoplasma detection was carried out through nested PCR assays using the primer pair P1 (Deng and Hiruki, 1991) / Tint (Smart *et al.*, 1996) amplifying a fragment of 1,600 bp from the 16S rRNA gene and the beginning of the 16S-23S spacer region, followed by R16F2n/R16R2 primers (Gundersen and Lee, 1996), which amplify fragments of 1,250 bp in the 16S rRNA gene. All the samples amplified with R16F2n/R16R2 primer pair were used for amplification of *vmp1* and *tuf* genes using fTuf1/rTuf1 and StolH10F1/R1 primer pairs

followed by nested-PCR with fTufAY/rTufAY and TYPH10F/R primer pairs, respectively 99 (Schneider et al., 1997; Cimerman et al., 2009; Fialová et al., 2009). A DNA concentration of 100 40 ng was used as template in the final volume of 50 µL containing 1 µL dNTPs (10 mmol L<sup>-1</sup> 101 mM), 1 µL of each primer (10 pmol), 5 µL of 10X Dream Taq green buffer, 0.25 µL Dream 102 Taq DNA polymerase (5 U/µL) (Thermo Scientific, USA) and 40.75 µL SDDW. PCRs using 103 P1/Tint, fTuf1/rTuf1 and H10F1/R1 were carried out after initial denaturation for 3 min at 95°C 104 followed by 35 cycles of 1 min at 95°C, 1 min at 60°C (for P1/Tint) and 55°C (for fTuf1/rTuf1 105 and H10F1/R1) and 2 min at 72°C followed by an extension step of 10 min at 72°C in Mini 106 Amp Plus Thermal Cycler (Applied Biosystems by Thermo Fisher Scientific). Nested-PCR 107 assays using R16F2n/R16R2, fTufAY/rTufAY and TYPH10F1/R1 were performed using 1 µL 108 of a 1: 50 dilution in SDDW of the PCR product. All the parameters were identical in direct 109 and nested PCR except for the annealing temperatures, which were 55-, 50- and 53°C for the 110 primers R16F2n/R16R2, fTufAY/rTufAY and TYPH10F1/R1, respectively. Samples without 111 DNA template were added as negative controls in all PCR reactions. Total nucleic acid of the 112 BGWL-Tu-M1 ('Ca. P. cynodontis', GenBank accession number HE599389) strain (Çağlar et 113 114 al., 2013) was employed as positive control. A strain of 'Ca. P. solani' (Güldür et al., 2018) was used as positive control in the PCR studies with *tuf* and *vmp1* genes. Seven microlitres of 115 amplicons were electrophoresed in 1% of 1X TAE agarose gel, stained with ethidium bromide 116 and photographed under a UV transilluminator. 117

118 The nested-PCR products obtained using R16F2n/R16R2 and fTufAY/rTufAY primers 119 from symptomatic maize plants were analyzed by RFLP. Enzymatic digestions were carried 120 out in a final volume of 30  $\mu$ L including 2  $\mu$ L Fast Digest Green Buffer, 10  $\mu$ L amplicons and 121 1 U restriction enzyme *Hpa*II (Thermo Scientific, Lithuania), according to the manufacturer's 122 instruction at 37°C for 20 min. The digested products were separated in 1.5% of 1X TAE 123 agarose gel, stained with ethidium bromide and photographed under a UV transilluminator. 124

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# 2.4 | Sequencing and phylogenetic relationships

The R16F2n/R2 and TYPH10F1/R1 amplicons from four fields were purified using 126 ExoSAP-IT® for PCR product clean-up (Thermo Fischer) according to the manufacturer's 127 instructions. The purified products were quantified in a nanodrop spectrophotometer and 128 129 sequenced in both directions with the same primers used for amplification in an automated sequencing (Macrogen, The Netherlands and Medsantek, Turkey). Sequence editing and 130 assembly were carried out using MEGA 7 (Kumar et al., 2016). The sequences obtained were 131 compared with those of phytoplasmas using the Basic Local Alignment Search Tool (BLASTn) 132 system of the National Center for Biotechnology Information (https://www.ncbi.nlm.nih.gov/). 133 134 Partial sequences from the 16S rRNA gene corresponding to the R16F2n/R2 fragments were subjected to in silico RFLP analyses with the iPhyClassifier tool (Zhao et al., 2009). In addition 135 to 16S rDNA, the RFLP patterns of *vmp1* gene amplicons was studied to verify the 136 phytoplasma identification accuracy. The nucleotide sequences were virtually digested with 137 RsaI endonuclease via pDRAW32 software version 1.1.146 (AcaClone software, 138 http://www.acaclone.com). 139

140 Phylogenetic trees were generated using MEGA 7 with the neighbor-joining method and 1,000 replicates for the bootstrap analysis to estimate the stability of the inferred subclades. 141 142 The 16S rRNA gene sequence from Acholeplasma laidlawii (GenBank accession number M23932) was used as an outgroup for the 16S rDNA tree root. Twenty-one nucleotide 143 sequences shown in Table 1 were downloaded from the GenBank database and used together 144 with the nucleotide sequences generated in this study to reconstruct the phylogenetic 145 relationships according to *vmp1* gene. 146

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**3 | RESULTS** 148

The symptomatic maize plants, mainly located at the edge of the surveyed fields, 149 exhibited severe reddening symptoms in leaves and midribs, and showed atrophied 150 inflorescence and empty corn cob with poor quality of seeds (Figure 1). The use of P1/Tint 151 primer pair enabled to amplify DNA bands (about 1.6 kb) from the positive control and from 152 153 all symptomatic leaf tissues. However, no bands were observed from the asymptomatic plants negative control (data not shown). The 16S rDNA partial sequences of samples Tr-AcM 1, -2, 154 -3 and -4 showed 100% identity with each other and 99.75% with that of 'Ca. P. solani' 155 (GenBank accession number AF248959) and were submitted to GenBank under the accession 156 numbers MW344059, MW344060, MW344061, MW344062, respectively. The identification 157 was made with iPhyClassifier based on in silico RFLP patterns (similarity coefficient 1.00) and 158 this confirmed that the detected phytoplasma belong to the 16SrXII-A subgroup (Figure 2). 159 This identification was further supported by the clustering analysis with other 'Candidatus 160 Phytoplasma' species and 'Ca. P. solani' strains deposited to GenBank from Turkey and 161 neighboring countries (Figure 3). 162

For the *tuf* and *vmp1* genes, amplification of approximately 1.4 kb and 0.9 kb, was 163 obtained from the nested-PCR assays with TYPH10F1/R1 and fTufAY/rTufAY primer pairs, 164 respectively (data not shown). Restriction patterns of PCR products from the *tuf* gene with 165 HpaII were identical to each other and confirmed that all maize strains belong to tuf-type-b 166 (Figure 4). The primer pair TYPH10F1/R1 also amplified the expected length fragment in all 167 the maize samples and positive control but not from the negative controls. The *vmp1* amplicons 168 of Tr-AcM 1, -2, -3 and -4 samples were sequenced in both directions, and the sequences were 169 deposited in GenBank under the accession numbers MW984610, MW984611, MW984612, 170 MW984613, respectively. The unrooted phylogenetic tree (Figure 5) constructed using the 171 *vmp1* sequences listed in Table 1 and those obtained from corn from Turkey showed that the 172 nucleotide sequences of maize strains clustered according to the typology of the RsaI/RFLP 173

(Figure 6). *Vmp1* gene amplicons of the four studied strains were virtually digested using the
pDraw32 software that showed only one *RsaI* profile, identical to the one of the '*Ca*. P. solani'
strains enclosed in the lineage V15 (Figure 6).

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### 178 4 DISCUSSION

179 In this study, the presence of 'Ca. P. solani' in commercial maize production areas in Southern Anatolia including Adana province, was reported for the first time in Turkey. While the 180 phytoplasmas in the 16SrXII group are divided in numerous subgroups distributed in 181 agricultural crops worldwide, only subgroup 16SrXII-A was detected, indicating the presence 182 of a homogeneous phytoplasma population. A prerequisite to manage newly occurring diseases 183 184 is the detection and proper identification of the associated pathogen. The identification of this phytoplasma strain in corn is, therefore, a first step for the management of the reddening of 185 186 maize disease via cultural, chemical and breeding approaches.

The molecular characterization of 'Ca. P. solani' in maize by using the tuf and vmp1 genes 187 confirmed the finding and the phytoplasma identity and represents a new epidemiological 188 information about 'Ca. P. solani' in Turkey. Based on the tuf gene characterization, all the 189 190 symptomatic maize samples showed the presence of only tuf-type b which agrees with reports from Bosnia and Herzegovina (Kovačević et al. 2014; Delić et al., 2016). Tuf-type b is 191 primarily associated with field bindweed (Convolvulus arvensis L.) (Langer and Maixner, 192 2004) that is its dominant reservoir plant species throughout most of the Europe and Turkey 193 (Ember et al., 2011; Ozturk et al., 2017). With the abundance of these plants, 'Ca. P. solani' 194 has a strong potential to spread out to maize and other crop plants when the insect vector(s) are 195 present. Typing *vmp1* gene with *RsaI* digestion and sequence analysis indicated that all maize 196 strains in the present study belonged to the same molecular type named V15. It was previously 197 identified in the 22% of 56 Hyalestes obsoletus in Spanish vineyards (Sabaté et al., 2014), in 198

diseased grapevine and Convolvulus arvensis (Murolo et al., 2010; Sabaté et al., 2014; 199 Quaglino et al., 2016; Jamshidi et al., 2019), and in tomato (Contaldo et al., 2021) in other 200 Mediterranean Countries. In the present study, all symptomatic maize samples resulted infected 201 by this phytoplasma strain. Identification of the same genetic variant of 'Ca. P. solani' from 202 203 different fields in the present study indicated that the pathogen population is strongly homogeneous, and the pathogens have a probable common origin. A similar case with severe 204 corn reddening symptoms associated with a similar strain of phytoplasma was reported in 205 Bulgaria (Genov et al., 2014) considering that close geographical areas could create potential 206 danger for insect vector mobility. Limited studies have reported the presence of H. obsoletus 207 and Dictyophara europaea, insect vectors of 16SrXII-A phytoplasmas, in vineyards of Turkey. 208 However, the phytoplasma presence in those insect species were negatively reported previously 209 (Bayram et al., 2014). Moreover, Laodelphax striatellus (Fallen, 1826), a possible vector of 210 16SrI, 16SrXII-A and 16SrIII group phytoplasmas (Prota et al., 2007), was previously detected 211 in Turkish maize fields (Sertkaya et al., 2010; Bayram et al., 2014). Çukurova region is a large, 212 flat and fertile area suitable for agricultural crop production where almost all agricultural 213 214 products except tea and hazelnuts could be grown. Since this region is suitable for polycultivation due to the suitable climatic conditions and long vegetation period, two or more 215 harvests can be possible for different crops in one season. As a result of this, any occurrence of 216 new disease or emerging pathogen has great potential in terms of spread mainly due to insect 217 218 vectors. It is known that phytoplasmas are spread primarily by insects families such as Cicadellidae (leafhoppers), Fulgoridae (planthoppers), and Psyllidae (jumping plant lice), 219 220 which feed in the phloem of plants, and, in case of infected plants, they may transmit them to 221 the next plant on which they feed. Thus, the host range of phytoplasmas strongly depends on the insect vector (Weintraub and Beanland, 2006). In the reported case, it is suggested that insect 222 vectors might have played a significant role for the spread of phytoplasmas to maize plants. 223

Therefore, the cultivation of maize varieties resistant to the diseases associated with 224 phytoplasmas and the control of insect vectors may be of great significance to manage this 225 emerging disease situation. Further studies are, however, necessary to identify the potential 226 insect vectors, the origin of the disease and its geographical distribution to develop a sustainable 227 disease management. With the global increase in temperature, xerothermic insect vectors might 228 migrate towards northern areas, and the cultivation of various host plant species in the same 229 area would probably increase the possibility of phytoplasma dissemination (Maixner and 230 Langer, 2006; Krishnareddy, 2013; Mamay and Şimşek, 2017). The phytoplasma-associated 231 disease emergence in new geographic areas might create epidemics where susceptible host plant 232 species are cultivated. Therefore, the main strategy for the control of the phytoplasma diseases 233 is to monitor their presence and consider the biological activities of the insect vectors. 234

235

## 236 CONFLICT OF INTEREST

All the authors declare that there is no conflict of interest in this study.

238

### 239 AUTHOR CONTRIBUTION

BKÇ was responsible for establishing and designing the surveys and experiments. BKÇ conducted the Nested-PCR experiments based on 16S rDNA, BKÇ, EŞ and MD conducted the experiments based on *Tuf* and *VMP1*. AB conducted phylogenetic studies. BKÇ, EŞ, AB and MD wrote the manuscript and analysed the data. All authors have read and approved the manuscript.

245

#### 246 DATA AVAILABILITY STATEMENT

The data that support the findings of this study are available from the corresponding authorupon reasonable request.

249

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Figure 1. Disease symptoms associated with the presence of '*Ca*. P. solani' in maize. Top: maize reddening symptoms with red streak on vein of leaf and stem; bottom: empty corn cob and poor seed set formation from symptomatic maize plants.

386

**Figure 2.** Virtual RFLP patterns derived from *in silico* digestions of Tr-AcM-1, Tr-AcM-2, Tr-AcM-3 and Tr-AcM-4 strain sequences obtained from R16F2n/16R2 products (about 1.2 kb) and phytoplasma 16S rDNA gene fragments from strains enclosed in the 16SrXII subgroups with four key enzymes (*AluI*, *MseI*, *Bst*UI and *HhaI*). Red boxes show the identity in the profiles of the strains (*AluI* and *MseI*), green boxes show the difference of the strains using *Bst*UI and *HhaI*. MW: Marker,  $\Phi$ X174 DNA profile digested with *HaeIII* with the following fragments in base pairs from top to bottom: 1,353; 1,078; 872; 603; 310; 281; 271; 234; 194; 118; 72.

394

**Figure 3.** Phylogenetic tree constructed by neighbor-joining method inferred from the '*Candidatus* Phytoplasma' species nucleotide sequences of the 16S rDNA gene. The tree shows the relationships among the phytoplasma strains obtained from maize plants from Turkey (red squares) and some '*Candidatus* Phytoplasma' species available in GenBank. The tree was constructed using MEGA 7 software. GenBank accession number and ribosomal subgroups are reported on the right of the strain names. *A. laidlawii* was employed as outgroup to root the tree.

402

Figure 4. RFLP analysis of the fTufAY/rTufAY amplicons of the elongation factor Tu (tuf)
gene with *Hpa*II. Only '*Ca*. P. solani' tuf-type b was detected. M, marker 1 kb DNA ladder; +,
Con: positive control from 16SrXII-A subgroup; UdC, undigested amplified control.

406

Figure 5. Unrooted phylogenetic tree inferred from the '*Ca*. P. solani' strain nucleotide sequences of the *vmp1* gene. Minimum evolution analysis was carried out using the neighbor-joining method and bootstrap replicated 1,000 times. The full names of the phytoplasma strains included in the phylogenetic analysis are given in the Table 1. The GenBank accession number of each sequence is given in parenthesis; gene sequences obtained in the present study are indicated with red squares. Square brackets show the grouping obtained by digestion of the amplicons with the endonuclease *Rsa*I.

413

Figure 6. Virtual RFLP patterns of *vmp1* gene sequence amplified with the primer pair TYPH10F/R of
strains Tr-AcM1, -2, -3 and -4 with the P7 strain from periwinkle from Lebano as control (GenBank

416	accession number AM992100). M: marker, $\Phi$ X174 DNA profile digested with <i>Hae</i> III with the
417	following fragments in base pairs from top to bottom: 1,353; 1,078; 872; 603; 310; 281; 271;
418	234; 194; 118; 72.

419

420 Table 1. '*Ca.* P. solani' *vmp1* gene sequences available in GenBank used in the phylogenetic analysis.
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