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Phytoplasma phoenicium'**

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1 *Asymmetrasca decedens* (Cicadellidae, Typhlocybinae), a natural vector of
2 ‘*Candidatus Phytoplasma phoenicium*’

3

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17

18

19 **Abstract**

20 ‘*Candidatus Phytoplasma phoenicium*’ is associated with a lethal disease of almond, peach and
21 nectarine named almond witches’-broom disease (AlmWB). The disease spread rapidly in Lebanon
22 from coastal areas to elevations exceeding 1200 m, killing over 150,000 trees in a span of two
23 decades. The mode of spread suggested the involvement of efficient vector(s) and *Asymmetrasca*
24 *decedens* (Hemiptera, Cicadellidae) was suspected as it is the most abundant leafhopper species
25 present in Lebanese stone fruit orchards. Living *A. decedens* specimens were collected from fields
26 heavily infested by AlmWB and used in transmission trials on healthy peach almond hybrid GF-677
27 and peach GF-305 seedlings with an inoculation-access period of 30 days. PCR analysis supported
28 by sequencing showed that *A. decedens* is a carrier of the phytoplasma, and that the phytoplasma
29 was detected in insect salivary glands and in some inoculated GF-677 and GF-305 seedlings. One
30 year post-inoculation, ‘*Ca. P. phoenicium*’ was detected in newly emergent leaves of inoculated
31 seedlings. However, the characteristic symptoms of witches’-broom were not observed. PCR
32 amplified fragments from phytoplasma-positive seedlings and from *A. decedens* samples showed
33 99.9% nucleotide identity in their 16S RNA region and phylogenetic analysis using a neighbour
34 joining tree confirmed that the phytoplasmas detected in both insects and inoculated seedlings
35 belonged to 16SrIX-B (D). The present manuscript is the first known report for a leafhopper vector
36 of ‘*Ca. P. phoenicium*’ and shows that the incubation period of the disease in plants may be longer
37 than 1 year. The importance of phytosanitary control measures, the adoption of a national strategy
38 and regional cooperation in order to contain the further spread of the disease are discussed.

39

40 **Keywords**

41 *Asymmetrasca decedens*; ‘*Ca. Phytoplasma phoenicium*’; leafhopper; phytoplasma transmission;
42 phytoplasma vector; salivary glands.

43 **Introduction**

44 In the 1990s, a devastating disease on almond trees appeared in Lebanon, characterised by
45 proliferation, small yellowish leaves, bushy growth, dieback and appearance of witches'-broom on
46 the stems. Infected trees either did not produce any fruits, or produced a limited number of
47 deformed fruits, resulting in practically 100% marketable yield loss. The disease was named
48 almond witches'-broom (AlmWB), it spread rapidly and killed about 100,000 trees over a period of
49 10 years (Abou-Jawdah *et al.*, 2002). The disease was associated with '*Candidatus* Phytoplasma
50 phoenicium' strains belonging to the pigeon pea witches'- broom (PPWB) group (16SrIX) (Abou-
51 Jawdah *et al.*, 2002; Verdin *et al.*, 2003), subgroup 16SrIX-B (also called 16SrIX-D) and its genetic
52 variants (subgroups 16SrIX-F and -G) (Molino Lova *et al.*, 2011). More recent surveys identified
53 over 40,000 new almond, peach and nectarine trees infected with AlmWB (Molino Lova *et al.*,
54 2011). The disease epidemic spreads rapidly from coastal areas to high mountainous areas (>1200
55 m), encompassing several ecological niches. Furthermore, AlmWB was found to infect properly
56 managed orchards, abandoned orchards and isolated wild trees. These observations suggested the
57 presence of efficient aerial vectors.

58 Phytoplasmas are bacteria devoid of cell walls that are capable of growing in specific insect vectors
59 as well as in the phloem tissue of host plants (Lee *et al.*, 2000). Phytoplasmas are mainly
60 transmitted by phloem-feeding insects which belong to the families Cicadellidae, Cixiidae,
61 Psyllidae, Cercopidae, Delphacidae, Derbidae, Meenoplidae and Flatidae in the order Hemiptera
62 (Weintraub & Beanland, 2006). Of these families, only some species can act as vectors because of
63 vector-pathogen-host specificity (Bosco *et al.*, 2009). The most common vectors of phytoplasmas
64 appear to be leafhoppers (Cicadellidae), planthoppers (Cixiidae) and psyllids (Psyllidae) (Weintraub
65 & Gross, 2013).

66 Field surveys were conducted in AlmWB-infested almond orchards located in South and North
67 Lebanon. *Asymmetrasca decedens* (Hemiptera, Cicadellidae, Typhlocybinae) was the most
68 abundant hemipteran species representing over 82% of total leafhoppers caught in sticky yellow

69 traps and in malaise traps (Dakhil *et al.*, 2011). *Asymmetrasca decedens* is a polyphagous species
70 which may feed on a wide variety of economic crops such as peach, almond, citrus, grapevine,
71 beans, beet, cotton, lucerne and potatoes (Jacas *et al.*, 1997). PCR tests showed that *A. decedens*
72 along with eight other leafhopper species carried 16SrIX phytoplasma and may represent potential
73 vectors (Dakhil *et al.*, 2011). However, phytoplasmas may be acquired by insects but may not be
74 transmitted during feeding (Marzachì *et al.*, 2004). Phytoplasmas are transmitted in a persistent
75 propagative manner (Marzachì *et al.*, 2004). For an insect carrier to become a vector, an intimate
76 association with the phytoplasma is required (Suzuki *et al.*, 2006). The phytoplasma must be able to
77 multiply in the vector, circulate in the hemolymph, accumulate in the salivary glands and be
78 secreted with the saliva upon feeding on plant phloem cells (Hogenhout *et al.*, 2008). Such a cycle
79 may take several days to several months. For example, in the case of *Cacopsylla pruni*, the vector of
80 ‘*Ca. P. prunorum*’ (agent of European stone fruit yellows, ESFY), most transmissions occur only
81 after an effective latency of 8 months (Thébaud *et al.*, 2009).

82 Only appropriate transmission tests can provide definitive evidence of the role of an insect as a
83 vector, while the detection of a phytoplasma in an insect is just considered as a preliminary step.
84 Moreover, controlled transmission tests are not always straightforward. Many vectors do not
85 survive easily in captivity, and various life stages may vary in the efficiency of transmission.
86 Symptom development on the inoculated plants and incubation period may also span from 1 week
87 to more than 24 months (Hogenhout *et al.*, 2008). In the case of ESFY, it may take 4–5 months and
88 some hosts may remain symptomless (Carraro *et al.*, 1998). Hence, molecular techniques may play
89 an important role in phytoplasma detection in asymptomatic and susceptible hosts during the
90 incubation or latent period (Mehle *et al.*, 2010). The major objective of this work was to investigate
91 the capacity of *A. decedens* to transmit AlmWB phytoplasma.

92

93 **Materials and methods**

94 *Plant material*

95 Certified tissue culture seedlings of two stone fruit rootstocks were imported from Italy, peach
96 almond hybrid ‘GF-677’ rootstock (*Prunus persica*×*Prunus dulcis* (Mill.) D.A. Webb.) and peach
97 seedling ‘GF-305’. The seedlings (30–35cm in length) were transplanted into 25 cm diameter pots
98 containing a mixture of potting soil, sand and perlite (2:1:1) and maintained in insect-proof cages,
99 within an insect-proof net house.

100

101 *Leafhoppers collection and transmission trials*

102 Insects were collected in two stone fruit orchards infested with AlmWB, an almond orchard in
103 Feghal, North Lebanon, and a nectarine orchard in Kfarkela, South Lebanon. A hand-held
104 mechanical aspirator (D-Vac Vacuum Insect Net-Model 122, Rincon-Vitova Insectaries, Ventura,
105 CA, USA) was used to collect insects from AlmWB-infected trees. *Asymmetrasca decedens*
106 leafhoppers were sorted out by mouth aspirator and transported to a cold room where they were
107 counted and dispensed into falcon tubes. Transmission trials were initiated the day of insect
108 collection. Collected insects were released either into 10 small insect-proof cages containing a
109 single seedling each or into 4 large cages containing 6 seedlings per cage. Twenty-five leafhoppers
110 were used for each seedling in individual cages and 150 leafhoppers were released into each of the
111 bigger cages. The leafhoppers were allowed an inoculation access feeding on GF-677 and GF-305
112 seedlings for 30 days. Afterwards, the insects were sprayed with insecticides at 5-day intervals
113 (spinosad and acetamiprid, in alternation). A total of 34 seedlings (15 GF-305 and 19 GF-677) were
114 inoculated in these tests. Two types of controls were used, six healthy seedlings maintained in an
115 insect-proof cage and six healthy seedlings maintained in another cage but subjected to feeding by a
116 total of 150 leafhoppers collected from a nectarine orchard in Wata Al Jawz, an AlmWB-free region.
117 Observations on symptom development were recorded at weekly intervals. Leaf samples were
118 collected periodically (1, 2, 3 and 12 months post-inoculation [mpi]) from treatments and control
119 seedlings and tested by polymerase chain reaction (PCR) for the presence of ‘*Ca. P. phoenicium*’.
120 Five batches of samples each containing five leafhoppers were collected from each of the

121 AlmWBinfested almond and nectarine orchards or from AlmWBfree regions to be tested by PCR,
122 as well as one batch that was taken from each cage during the transmission studies. Moreover, the
123 salivary glands of nine selected *A. decedens* specimens were dissected out of the insects and
124 analysed by PCR in order to assess presence of ‘*Ca. P. phoenicium*’ and to get further confirmation
125 of the insect vectorship capability. Briefly, leafhopper heads were removed from the rest of the
126 insect body, and the salivary glands from subsets of three insects were dissected and pooled into a
127 microfuge tube (1.5 mL) containing 25 μ L Sodium Chloride-Tris-EDTA (STE) Buffer. DNA in the
128 samples was extracted and used for phytoplasma detection by PCR.

129

130 **Molecular diagnosis**

131 *Total nucleic acid extraction*

132 For plant samples, the total nucleic acids (TNA) were extracted from 100mg of leaf midribs
133 following the CTAB protocol as described previously (Abou-Jawdah *et al.*, 2002). Samples from
134 leafhoppers collected from AlmWB-infested orchards or from AlmWB-free orchards were also
135 tested by PCR. Groups of five *A. decedens* insects were put in a 1.5mL Eppendorf tube and the
136 TNAs were extracted according to the procedure described by Marzachi *et al.* (1998). The final
137 TNA precipitate was suspended in 50 μ L of sterile water. TNA extracts were analysed in a 1%
138 agarose gel electrophoresis to determine their quality. Total DNA were quantified using a
139 NanoDrop 2000c (NanoDrop Technologies, Wilmington, DE, USA) and stored at -20°C .

140

141 *Phytoplasma detection by polymerase chain reaction*

142 The semi-specific primer pair, ALW-F2/ALW-R2, which amplifies a DNA fragment of 390 bp
143 from 16SrIX phytoplasmas, was used in PCR assays as described previously (Abou-Jawdah *et al.*,
144 2003). Each amplification reaction was performed in 20 μ L reaction mixture containing 2 μ L of
145 template DNA (20–50 ng), 10 μ L of REDTaq® ReadyMix™ PCR Reaction Mix (Sigma-Aldrich,
146 St Louis, MO, USA), 0.25 μ M of each primer and 7 μ L of sterile water. Amplifications were done

147 with a Bio-Rad Thermal Cycler 1000 (Bio-Rad Laboratories, Hercules, CA, USA). Positive
148 leafhoppers and inoculated plant samples detected by previous direct PCR, were retested by nested
149 PCR using forward primer P1 (Deng & Hiruki, 1991) and reverse primer P7 (Smart *et al.*, 1996)
150 followed by R16F2n/R16R2 (F2n/R2) to confirm phytoplasma attribution to '*Ca. P. phoenicium*'
151 following nucleotide sequence analyses (Lee *et al.*, 1998; Abou-Jawdah *et al.*, 2003). The F2n/R2
152 amplicons were purified with the Illustra™ GFX PCR DNA and Gel Band Purification kit (GE
153 Healthcare, Waukesha, UK) and cloned using the pGEM-T Easy Vector System II (Promega,
154 Madison, WI, USA). Sequencing of the cloned PCR products was performed at Macrogen Inc.
155 (Seoul, South Korea) in both forward and reverse directions. The nucleotide sequence data were
156 assembled by employing the Contig Assembling programme of the sequence analysis software
157 BIOEDIT, version 7.0.0 (<http://www.mbio.ncsu.edu/Bioedit/bioedit.html>).

158

159 *16S rRNA gene analysis*

160 F2n/R2 fragments amplified from insect bodies, salivary glands and inoculated seedlings were
161 sequenced and a representative sequence from each host was submitted to GenBank. Sequences
162 were compared with the Gen- Bank database using the algorithm BLASTN ([http://www.](http://www.ncbi.nlm.nih.gov/BLAST/)
163 [ncbi.nlm.nih.gov/BLAST/](http://www.ncbi.nlm.nih.gov/BLAST/)) in order to determine the best sequence identity hit and to establish the
164 species affiliation of phytoplasmas detected in *A. decedens* specimens and in plants used in
165 transmission trials. Multiple alignment using Geneious R6 (v6.0.5, Biomatters, Auckland, New
166 Zealand) was performed for sequences obtained from the insect vectors and the inoculated seedlings
167 to ascertain that the same phytoplasma species occur in both hosts.

168 The alignments were exported to the MEGA 6 software (Tamura *et al.*, 2013) for distance and
169 phylogenetic analyses. Neighbour-Joining (NJ) (Saitou & Nei, 1987) tree was constructed using 500
170 replicates for bootstrap analysis (Felsenstein, 1985) and *Acholeplasma laidlawii* was used as an out
171 group.

172

173 **Results**

174 *Symptom development*

175 The transmission trials using leafhoppers were initiated on May 2012, and symptoms were
176 monitored at weekly intervals. Symptoms started to develop on 16 inoculated seedlings within 25
177 days post-inoculation (dpi). By 30 dpi, 4 out of 15 GF-305 seedlings and 12 out of 19 GF-677
178 seedlings had developed symptoms. The observed symptoms were not typical of AlmWB
179 phytoplasma; they consisted mainly of downward leaf curling or rolling and proliferation of new
180 growth at the leaf axils. The curled leaves were smaller than normal leaves but were not chlorotic;
181 moreover, many growing tips were burned. Similar symptoms were observed on some of the control
182 plants which were inoculated with leafhoppers originating from an area free of AlmWB
183 phytoplasma.

184 In August, 3 mpi, a new flush of growth appeared which looked normal. During winter, the leaves
185 dropped and in early March the new growth had vigorous growth that was similar to that emerging
186 from healthy controls.

187

188 *Molecular diagnosis*

189 At 1 and 2 mpi, PCR tests showed that out of the 34 inoculated seedlings, 16 gave positive results
190 using the AlmWB semi-specific primer pair, ALW-F2/R2 (Table 1). When the new summer flush
191 appeared in August (about 3 mpi), the new growth looked normal. Young leaf samples were
192 collected and the PCR results showed that only three samples of the GF-305 seedlings were positive
193 out of the four that were positive at 2 mpi, while with the GF-677 seedlings, only 8 seedlings tested
194 positive out of the 12 seedlings that were previously positive. Both populations of *A. decedens*
195 collected from almond or nectarine orchards were able to transmit the phytoplasma (Table 2). The
196 direct PCR results were confirmed by nested PCR, sequencing and BLAST analyses. During winter,
197 all the leaves dropped. In the following spring season, new growth emerged which appeared normal.
198 PCR tests were repeated and all the seedlings, whose summer flush tested positive, were also

199 positive with the new spring growth (Tables 1 and 2, Fig. 1). Therefore out of a total of 34
200 inoculated seedlings only 11 seedlings got infected as revealed by PCR tests about 1 year post-
201 inoculation; however, none developed AlmWB-associated symptoms.

202 Leafhoppers collected from Wata Al Jawz, an AlmWBfree area, gave negative PCR results using
203 the semispecific primer pair, ALW-F2/R2. The 14 composite samples of *A. decedens* leafhoppers
204 used in the inoculation tests gave positive results (Fig. 2). When the salivary glands of three
205 representative *A. decedens* samples, each composed of pooling salivary glands of three insects
206 collected from the AlmWB-infested orchard also tested positive for Alm WB (Fig. 2).

207 Sequences of the F2n/R2 amplified products from four samples, one sample each from the insect
208 body (GenBank accession number: KF359551), the salivary glands (KF488577), the inoculated GF-
209 677 seedlings (KF500029) and from GF-305 (KF500030) were deposited in GenBank. BLAST
210 analysis showed 99.9% identity with '*Ca. P. phoenicium*'. Results obtained by the NJ tree showed
211 that phytoplasmas present in the insects and in the inoculated seedlings were all similar and were
212 members of the species '*Ca. P. phoenicium*', subgroup 16SrIX-B (D) (Fig. 3).

213

214 **Discussion**

215 '*Candidatus Phytoplasma phoenicium*' is associated with a devastating and lethal disease of almond,
216 peach and nectarine that has so far only been reported in Lebanon and Iran (Abou-Jawdah *et al.*,
217 2002; Verdin *et al.*, 2003). '*Candidatus Phytoplasma phoenicium*' has all the characteristics of a
218 severe quarantine pathogen. It is associated with a lethal disease of three major stone fruit crops;
219 cannot be controlled by classical control measures, has the potential to occupy different ecological
220 niches, and its unaided transmission across natural barriers seems limited because it has been
221 reported in only two countries. The rapid spread of AlmWB in Lebanon suggests the presence of
222 one or more efficient vectors. Previous surveys carried out in Lebanese orchards showed that
223 several leafhopper species are carriers and may represent potential vectors of '*Ca. P. phoenicium*'

224 (Dakhil *et al.*, 2011). In this study, transmission trials were performed with *A. decedens*, the most
225 dominant leafhopper detected in stone fruit orchards, to investigate its vectoring activity.

226 The initial symptoms, observed 1 mpi, were not attributed to phytoplasma infection. They were
227 most likely correlated with leafhopper feeding, because leafhoppers feed mainly on leaves and
228 cause a symptom known as the ‘hopperburn’ (Backus *et al.*, 2005). In eastern Spain, a high
229 infestation of *A. decedens* in almond orchards induced stunted shoots with small curled leaves that
230 were only observed on young flush. The damage was mainly destructive to nursery seedlings,
231 young nonbearing trees and over-grafted plants (Jacas *et al.*, 1997).

232 Transmission trials performed in this study showed that 11 out of 34 inoculated stone fruit seedlings
233 got infected with ‘*Ca. P. phoenicium*’, as evidenced by PCR detection in emergent tissue 3 and 12
234 mpi. PCR data were confirmed by BLASTN and iPhyClassifier analyses of nucleotide sequences,
235 highlighting that the same phytoplasma ‘*Ca. P. phoenicium*’, subgroup 16SrIX-B (D) was detected
236 in the leafhoppers used in transmission trials and in the inoculated seedlings. The detection of ‘*Ca.*
237 *P. phoenicium*’ in leafhoppers and in inoculated certified seedlings provides strong evidence for the
238 role of *A. decedens* as a vector of ‘*Ca. P. phoenicium*’. Interestingly, both populations of *A.*
239 *decedens* collected from almond or nectarine orchards located in two different regions were able to
240 transmit the phytoplasma. A possible explanation for getting negative PCR results, 3 and 12 months
241 after inoculation, from five seedlings which showed positive results, 1 and 2 mpi, is that the vectors
242 successfully inoculated the leaf tissues but the phytoplasma failed to induce systemic infection and
243 thus was not detected in the new growth that emerged later on.

244 Two important features resulting from transmission experiments should be discussed. (1) The
245 number of insects used in transmission trials and (2) The long incubation period. First, the high
246 number of insects used per seedling led to transient phytotoxicity symptoms. In future tests, to
247 reduce phytotoxicity symptoms, a lower number of leafhoppers should be used per plant and one
248 seedling per cage may be preferable. However, it is worth mentioning that *A. decedens* was the
249 most abundant species in a surveyed almond orchard and 544, 2760 and 3901 insects were collected

250 on six yellow sticky traps during the months of March, April and May 2002, respectively (Dakhil *et*
251 *al.*, 2011). These results were confirmed in a recent survey with a slight difference in timing,
252 whereby 3 800, 11 700 and 7 200, were trapped in May, June and July 2012 (H. Abdul-Nour,
253 personal communication). This experiment was conducted in insect-proof cages under greenhouse
254 conditions, and a large number of leafhoppers died within 2 weeks of transfer to the insect-proof
255 cages, suggesting that the survival potential of *A. decedens* under the experimental conditions was
256 limited. Moreover, in an effort to study the transmission characteristics, mainly the latency period in
257 *A. decedens*, several attempts failed to rear this leafhopper in insect-proof cages.

258 Even though several leafhopper species belonging to the Cicadellidae family and sub-families were
259 reported to transmit phytoplasmas, only one report mentions *A. decedens* as a potential phytoplasma
260 vector based on transmission trials (Pastore *et al.*, 2004). Moreover, most leafhoppers in the
261 subfamily Typhlocybinae are reported to be mesophyll feeders (Nault & Rodriguez, 1985). This
262 characteristic reduces their potential to act as phytoplasma vectors. However, *A. decedens* and its
263 close relative, *Empoasca decipiens*, the two predominant species in stone fruit orchards in Lebanon,
264 were found to be carriers of AlmWB phytoplasma (Abou-Jawdah *et al.*, 2011; Dakhil *et al.*, 2011).
265 In Italy, these two species were also found to be positive for ESFY in PCR assays. Preliminary
266 trials showed that *Empoasca decedens* (a synonym to *A. decedens*) may transmit ESFY from
267 *Prunus armeniaca* L. to *P. armeniaca* (Pastore *et al.*, 2004), however, more recent trials failed to
268 confirm it (Pastore *et al.*, 2001). In Cuba, 67 *Empoasca* spp. samples were examined by PCR and
269 63 were found carrying 'Ca. P. aurantifolia' (Arocha *et al.*, 2006).

270 Second, phytoplasma symptoms can start to appear on plants as soon as 7 days after the insect has
271 introduced the phytoplasma, but this is not always the case because the symptoms may also take 6
272 to over 24 months to develop depending on both the phytoplasma and the plant host species
273 (Hogenhout *et al.*, 2008). Even in grafting experiments, symptoms may take a long time to appear,
274 for example, it took around 18 months for the ESFY symptoms to appear on patch grafted 3-year-old
275 plum and peach seedlings (Pastore *et al.*, 2001). Flavescence dor´ee of grapevine is symptomless in

276 some cultivars, and it also has a long (up to three years) latent period before symptoms can be seen
277 (Belli *et al.*, 2010). These data may be explained by the fact that phytoplasmas live inside plants as
278 symbiont but they can become pathogens in later stages when suitable conditions occur such as
279 special weather conditions or changes in the production practices (Mehle *et al.*, 2010). The long
280 incubation period poses a problem in early visual disease detection, and may have played a role in
281 the spread of the AlmWB disease to distantly isolated regions in Lebanon, through the production
282 of AlmWB-infected asymptomatic seedlings. This observation necessitates stricter phytosanitary
283 control measures on stone fruit nurseries and mother stock plants. For this reason, specific AlmWB
284 detection methods based on PCR and qPCR are being developed to survey accurately the plant
285 materials within the stone fruit nurseries.

286 The rapid spread of the disease over distantly located regions, and the detection of AlmWB
287 phytoplasma in eight other leafhopper species (Dakhil *et al.*, 2011) may indirectly represent a
288 hypothesis that other potential vectors for AlmWB phytoplasma may be present. Effectively, for
289 many phytoplasma diseases more than one vector was reported. For example, '*Ca. Phytoplasma*
290 *solani*' (16SrXII-A), agent of the bois noir (BN) disease of grapevine (Quaglino *et al.*, 2013), is
291 transmitted by Cixiidae; *Hyalesthes obsoletus* is the major reported vector, but recently *Reptalus*
292 *panzeri* was reported also as a natural vector of the disease, and several other vectors are suspected
293 (Cvrković *et al.*, 2013). The other potential vectors of AlmWB phytoplasma may not be common
294 pests of stone fruits, but may infest stone fruits only during part of their life cycles or occasionally
295 when their natural hosts become limited. For example, even though the vector of 'bois noir' (BN)
296 *Hyalesthes obsoletus* cannot live on grapevines, it feeds on different crops and has been proved to
297 accidentally transmit the phytoplasma from weeds to grapevine (Maixner, 1994; Weintraub *et al.*,
298 2009). Therefore, the preferred host(s) for some suspected phytoplasma vectors may be weeds or
299 other plants (Maniyar *et al.*, 2013). In view of the concurrent results that Cixiidae may play a role in
300 '*Ca. P. phoenicium*' transmission (R. Tedeschi, personal communication) from 'wild' or alternative

301 hosts to stone fruits, it seems that *A. decedens* plays a major role in spreading the disease within or
302 to nearby stone fruit orchards.

303 In conclusion, the detection of ‘*Ca. P. phoenicium*’ in the salivary glands of *A. decedens* along with
304 the transmission trial results confirm that this leafhopper is a vector of ‘*Ca. P. phoenicium*’, the
305 suspected causal agent of AlmWB. This constitutes the first report of *A. decedens* as a vector of
306 AlmWB disease and another experimental proof that it may act as a phytoplasma vector in stone
307 fruits (Pastore *et al.*, 2004). Further research is needed on the modality of transmission (efficiency
308 of different life stages, latency period), and the possibility of the occurrence of other potential
309 vectors. Therefore, further studies must be conducted on the epidemiology of the disease including
310 its alternative hosts and their relative importance in disease spread. In addition to vector control,
311 screening for resistant germplasms may also represent a possible option to perform, although all the
312 almond varieties present in Lebanon are susceptible. In view of the importance and severity of
313 AlmWB disease, regional and international cooperation should be established in order to develop an
314 integrated pest management approach to contain the disease, prevent its further spread and to reduce
315 its negative impact on the stone fruit industry.

316

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421 **Table 1** PCR detection of ‘*Ca. P. phoenicium*’ with ALW-F2/ALW-R2 primers in stone fruit
 422 seedlings 1, 2, 3 and 12 mpi using *A. decedens* as a vector

Variety	Seedling code	1 mpi	2 mpi	3 mpi ^a	12 mpi ^a
GF-305	AF3	+	+	+	+
	AF7	+	+	+	+
	AK4	+	+	+	+
	AK5	+	+	-	-
	AK10, 11, 12, 13, 14, 15	-	-	-	-
	AF8, 9, 10, 11, 12	-	-	-	-
	AF1	+	+	-	-
	AF2	+	+	+	+
	AF4	+	+	-	-
	AF5	+	+	+	+
GF-677	AF6	+	+	-	-
	AK1	+	+	+	+
	AK2	+	+	+	+
	AK3	+	+	-	-
	AK6	+	+	+	+
	AK7	+	+	+	+
	AK8	+	+	+	+
	AK9	+	+	+	+
	AK16, 17, 18	-	-	-	-
	AF13, 14, 15, 16	-	-	-	-
Total		16/34	16/34	11/34	11/34

423 ^aResults for 3 and 12 mpi are for leaf samples collected from new growths that were not subjected
 424 to direct leafhopper feeding.

425

426 **Table 2** Transmission of ‘*Ca. P. phoenicium*’ by *A. decedens* to seedlings

427 of two stone fruit rootstocks (GF-305 and GF-677)

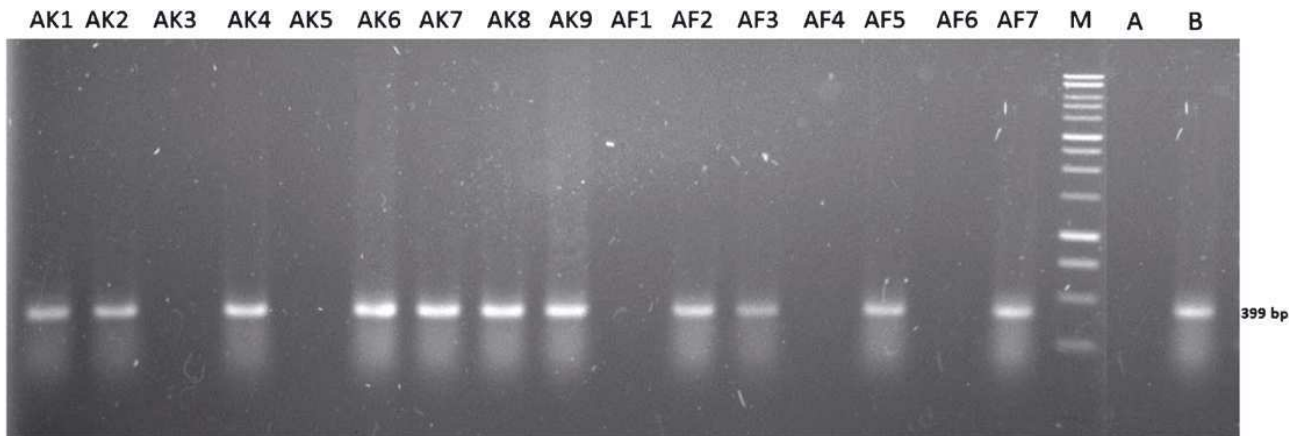
	No. of PCR positive/total tested	
	GF-305	GF-677
Single seedling/cage	0/1 ^a	1/1 ^a
	1/1 ^b	1/1 ^b
	0/1 ^a	0/1 ^a
		1/1 ^a
		1/1 ^a
		1/1 ^b
Multiple seedlings/cage	2/6 ^a	2/6 ^a
	0/6 ^b	1/6 ^b
		0/1 ^b
Total	3/15	8/19

428

429 The leafhoppers were collected from AlmWB-infested almond or nectarine orchards and the
 430 inoculations were conducted using either single or multiple seedling(s) per cage.

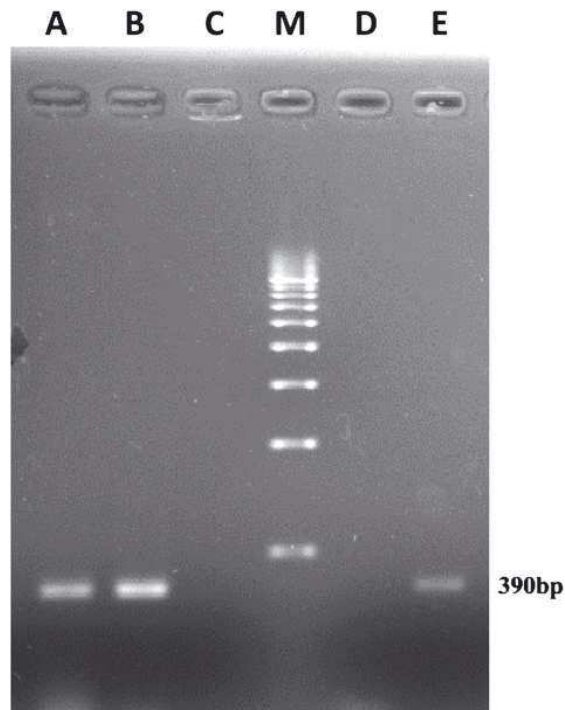
431 ^a*Asymmetrasca decedens* collected from nectarine orchard, Kfarkela region, South Lebanon.

432 ^b*Asymmetrasca decedens* collected from almond orchard, Feghal region, North Lebanon.



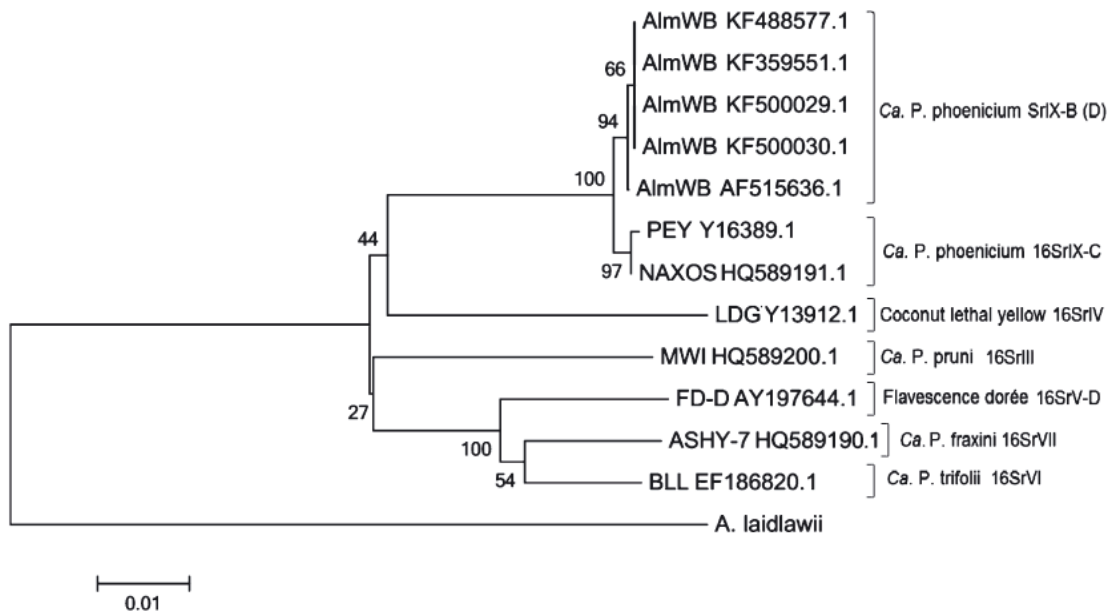
433

434 Figure 1 Agarose gel electrophoresis of PCR products using the semi-specific primer pair
 435 ALWF2/ALWR2 amplifying an amplicon of about 390 bp of the 16S-ITS23S region. DNA samples
 436 were extracted at 12 mpi, from 16 seedlings inoculated with *A. decedens* carrying '*Ca. P.*
 437 *phoenicium*. A: Healthy seedling, B: '*Ca. P. phoenicium*' positive control, M: 1 Kbp ladder.



439

440 Figure 2 Agarose gel electrophoresis of PCR products using the semispecific primer
441 pairALWF2/ALWR2 amplifying an amplicon of about 390 bp of the 16S-ITS23S region. DNA
442 samples were extracted from A: body and B: salivary glands of *A. decedens* collected from
443 AlmWB-infested orchard, C: *A. decedens* collected from healthy orchard, D: healthy control, E: '*Ca.*
444 *P. phoenicium*' positive control, M: 1 Kbp ladder.



445

446 **Figure 3** Neighbour-joining tree of R16F2n/R16R2 amplified fragment of the 16S rRNA gene.

447 Numbers at the nodes indicate bootstrap values; bars, substitutions per nucleotide position; 16S

448 rRNA GenBank sequence accession number is indicated following the strain acronym; 16S rRNA

449 group and subgroup are indicated following the phytoplasma strain; *A. laidlawii* (NR074448.1) was

450 used as an outgroup. AlmWB sequences are from insect body (KF359551) and the salivary glands

451 (KF488577) of *A. decedens*, from the inoculated seedlings of GF-677 (KF500029) and GF-305

452 (KF500030). The AlmWB reference strain (AF515636) was also used as well as seven more

453 reference phytoplasmas closely related to AlmWB.