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New insights in phytoplasma-vector interaction: acquisition and inoculation of Flavescence dorée phytoplasma by *Scaphoideus titanus* adults in a short window of time

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14 **Running title:** Phytoplasma acquisition and transmission by *Scaphoideus titanus* adults

16 Abstract:

17 The leafhopper Scaphoideus titanus is able to transmit 16SrV phytoplasmas agents of grapevine's 18 Flavescence dorée (FD) within 30-45 days, following an Acquisition Access Period (AAP) of a few 19 days feeding on infected plants as a nymph, a Latency Period (LP) of 3-5 weeks becoming 20 meanwhile an adult, and an Inoculation Access Period (IAP) of a few days on healthy plants. 21 However, several aspects of FD epidemiology suggest how the whole transmission process may 22 take less time, and may be start directly with adults of the insect vector. Transmission experiments 23 have been set up under lab condition. Phytoplasma-free S. titanus adults were placed on broad bean (BB) plants (Vicia faba) infected by FD-C (16SrV-C) phytoplasmas for an AAP=7 days. 24 Afterwards, they were immediately moved onto healthy BB for IAP, which were changed every 7 25 days, obtaining three timings of inoculation: IAP 1, IAP 2 and IAP 3, lasting 7, 14 and 21 days from 26 27 the end of AAP, respectively. DNA was extracted from plants and insects, and PCR were performed 28 to identify FD phytoplasmas. Insects were dissected and fluorescent in situ hybridization was made 29 to detect the presence of phytoplasmas in midguts and salivary glands. The rate of infection in 30 insects ranged 46-68% without significant differences among IAPs. Inoculation in plants succeeded 31 in all IAPs, at a rate of 16-23% (no significant differences). Phytoplasma load was significantly 32 higher in IAP3 than IAP 1-2 for both plants and insects. Phytoplasmas were identified both in 33 midgut and salivary glands of *S. titanus* at all IAP times. The possible implications of these results

34 in the epidemiology of Flavescence dorée are discussed.

36 Key words: Flavescence dorée, Scaphoideus titanus, Vicia faba, acquisition by adults, latency

for per period

- 37 access period, transmission process

39 Introduction

Flavescence dorée (FD) is a serious disease of grapevine caused by 16SrV phytoplasmas (subgroups C and D) (Arnaud et al., 2007) transmitted by Scaphoideus titanus Ball (Chuche & Thiery, 2014), and to a lesser extent by Dictyophara europaea (L.) (Filippin et al., 2009) and Orientus ishidae (Matsumura) (Lessio et al., 2016). However, unlike these two last ones, S. titanus is able to accomplish its whole life cycle only on grapevine, both Vitis vinifera L. and American grapevines' and/or wild rootstocks (Maixner et al., 1993; Vidano, 1966; Lessio et al., 2007; Chuche & Thiery, 2014). S. titanus is a univoltine species, and overwinters as egg, laid under the bark of 2-yr old wood, although sometimes 1-yr old wood can be used as well (Bagnoli et al., 2011; Lessio & Alma, 2013). During its postembryonic development, it undergoes five nymphal instars: N1, N2, N3, N4 and N5. The time elapsing between molts depends mostly on temperature (Falzoi et al., 2014). Usually, eggs start hatching in the beginning of May. Hatching dynamic depends on the winter temperatures eggs are exposed to: the colder is the winter, the shorter will be the hatching period (Chuche & Thiery, 2009).

Transmission of phytoplasmas by insect vectors follows a typical sequence of events: Acquisition Access Period (AAP), Latency Access Period (LAP), or Latency Period (LP), an Inoculation Access Period (IAP) (Alma et al., 2015). The transmission is therefore considered as "persistent-propagative". According to the state-of art, S. titanus nymphs (N3, N4 and N5) can acquire phytoplasmas by feeding on infected grapevines (both European and American), and during their LAP they become adults able to move from plant to plant and to transmit phytoplasmas to healthy plants (Chuche & Thiery, 2014; Lessio et al., 2007). In S. titanus, firstly, the LAP was thought lasting about 28-35 days, with a minimum of 7 days for both AAP and IAP starting from N3 nymphs (Schvester et al., 1961; 1969). Afterwards, Caudwell et al. (1970) demonstrated a possible inoculation to broad bean (BB) (Vicia faba L.) and grapevine after 21 days of AAP + LAP and 7 days of IAP (total days: 28).

Acquisition efficiency of phytoplasmas by S. titanus depends on a number of factors. The source of inoculum plays an important role: highly susceptible vine varieties are a better source than tolerant ones (Bressan et al., 2005b; Galetto et al., 2016); as well, a higher phytoplasma concentration increases acquisition efficiency (Galetto et al., 2014; 2016), which on the other hand is lower in the case of recovered grapevines (Roggia et al., 2014). Moreover, acquisition is more efficient in late summer because of a higher phytoplasma load in grapevines (Galetto et al., 2014). However, in late summer there are no nymphs left, but only adults. If AAP is performed just by nymphs, this aspect would be biologically unrealistic, whereas it might be possible that adults acquire phytoplasmas too.

At present, the whole literature about *S. titanus* and transmission of FD phytoplasma (FDP)
involves the nymphs for AAP, and adults for IAP, after a LAP of 28- 35 days. It is therefore
necessary to ascertain if the whole transmission period could be completed within adults' lifespan.
The aim of this research was to investigate if *S. titanus* adults are able to complete the transmission

process starting from AAP and finishing with IAP. Moreover, since adults have never been used for
AAP before, we reviewed the LAP in order to state if the whole transmission process could be
completed within a shorter period.

81 Materials and methods

83 <u>Insect source</u>

S. titanus was reared under lab conditions in order to obtain healthy individuals. Grapevine canes containing eggs (two-year old wood, or older) were collected during winter in 2014 and 2015 in several Piedmontese grapevine growing areas, in particular Asti Province and Canavese district, where S. titanus was detected in great number during the previous summer, by means of yellow sticky traps. Canes were cut in pieces of approximately 20 cm length, placed in plastic bags, periodically sprinkled to avoid desiccation of eggs, and preserved into a cool chamber (+4°C) before use. The twigs were placed into BugDorm® insect rearing cages (47.5 x 47.5 x 138 cm) made of mesh and polyethylene at springtime, along with potted plants of healthy BB. Broad beans were used to rear S. titanus through its whole life cycle to avoid any phytoplamas contamination of the insects before the transmission experiments, as 16SrV phytoplasmas are thought not being transmitted through seeds (Duduk and Bertaccini, 2009). The cages were kept either outdoors, under a shelter against rain and sunlight, from May 15th to September 1st, or into a greenhouse from March 15th to May 15th and from September 1st to November 15th, and periodically sprinkled. Insects were reared up to the adult stage, and then used for transmission experiments. Adults from such rearing were also directly collected and dissected for fluorescence in situ hybridization (FISH) analysis, as a negative control.

102 Transmission experiments

Transmission experiments were performed in a climatic chamber (T=25°C, RH=75%). Acquisition
(AAP) of phytoplasmas by *S. titanus* adults was made on BBs previously infected with FD-C
phytoplasmas by means of *Euscelidius variegatus* (Kirschbaum) (Caudwell *et al.*, 1972; Salar *et*

al., 2013). Plants used for AAP were 30-40 cm high, and had 8-10 leaves; before use, phytoplasma infection was checked by qPCR (see below). AAP was performed by placing 1-2 infected plants into BugDorm® insect rearing cages (47.5 x 47.5 cm) along with 20-30 newly emerged adults of S. titanus for 7 days. The procedure was repeated several times until enough live infected adults to be used in IAPs were obtained. After AAP, insects were divided into batches of 5 individuals, and each batch was placed onto a healthy BB seedling (height 5 cm, 2 leaves), inside a Plexiglas cylinder (h=20 cm; diameter=12 cm) with the top covered by a fine mesh. Every 7 days, live insects were moved onto another plant up to three times to perform a total of three IAPs, named IAP1-3 (Fig. 1). Cylinders were checked daily, and dead insects were removed and preserved into a freezer ($-20^{\circ}C$) for molecular analyses to test for FDP (see later). The same set up was repeated for FISH analysis: at the end of each IAP five live leafhoppers from single batches (one for each of the IAPs) were collected and dissected to isolate midgut and salivary glands. Hence, a total of 28 batches with 140 specimens (25 batches with 125 specimens for qPCR, and 3 batches with 15 insects for FISH, Fig 1) started IAP 1, however the number of batches progressively decreased due to insect mortality or collection in IAPs 2 and 3. Inoculated plants were treated with an insecticide (Dichlorvos, 0.5 g/L) and kept in a climatic chamber, inside an insect-proof cage to avoid contamination, for three weeks from the beginning of phytoplasma inoculation before molecular analyses.

DNA extraction and quantitative Real Time PCR analysis

Total DNA extraction was performed from whole, single insects used in transmission experiments and from inoculated BB. Nucleic acids extraction from S. titanus was carried out by following a procedure previously described for leafhoppers (Marzachi et al., 1998). Plant DNA was extracted from leaf tissue previously grounded with liquid nitrogen in a sterile mortar, according to the DNeasy Plant Mini Kit protocol instructions (Qiagen, Milan, Italy). Quantitative real-time PCR (qPCR) was carried out to measure the presence and concentration of phytoplasma genome units in insect and plant samples. A Chromo4 real-time detector (Bio-Rad, Milan, Italy) was used with PrecisionPlusTM -SY Mastermix (Primerdesign, Chandler's Ford, UK). Reactions targeting the 16S rRNA gene of group 16SrV phytoplasmas were carried out on all samples by using the fAY/rEY primer pair (Marcone et al., 1996; Marzachi et al., 2001), with the conditions described by Galetto et al. (2005). To calculate the average FDP Genome Units (GU) / sample, 16S rRNA gene copy numbers were divided by two, because this gene is estimated to be in double copy in the genome of

phytoplasmas (Schneider & Seemuller, 1994). Additionally, qPCR targeting the insect's 18S rRNA was performed on insect DNA gene to normalize the absolute phytoplasma density. Primers MqFw / MqRv were used according to Marzachì & Bosco (2005). Hence, normalized phytoplasma GU were calculated per pg of insect 18Sr RNA gene. On the other hand, qPCR results from BBs were expressed as FDp GU per 100 mg of leaf.

Standard curves were constructed by using dilutions of PCR-amplified 16S rRNA gene of FDP cloned with the pGEM T-easy Vector Cloning Kit (Promega, Milan, Italy). The detection limit was calculated as the highest dilution of cloned amplicons used for standard curves which was successfully amplified, and corresponded to 1.0×10^{0} FDP GU.

- Fluorescent in situ hybridization

FISH analyses were performed on S. titanus midguts and salivary glands. All of the experiments were carried out using a fluorescent probe specifically targeting group 16SrV phytoplasmas, along with Mollicutes-specific and eubacterial probes. Specifically, the 16SrV phytoplasma-specific probe ph1298, labelled with Cy5 (indodicarbocyanine, absorption/emission at 650/670nm), the Mollicutes probe MCP52, labelled with fluorescein isothiocyanate (FITC, absorption/ emission at 494/520 nm), and the eubacterial probe Eub338, labelled with Texas Red (absorption and emission at 595 nm and 620 nm, respectively) were used as described by Gonella et al. (2011) and Lessio et al. (2016).

In order to perform whole mount FISH, salivary glands were dissected in a sterile saline solutions, then fixed for 2 min at 4°C in 4% paraformaldehyde and washed in PBS. All hybridization experiment steps were performed following Gonella et al. (2011). After hybridization, the samples were mounted in antifading medium and then observed in a laser scanning confocal microscope SP2- AOBS (Leica).

Statistical analysis

> Statistical analyses were performed by mean of SPSS Statistics 24® (IBM Corp. Released 2016, Armonk, NY). A generalised linear model (GLM) was run for analysing infection rates (binomial: positive/negative) and quantitative PCR (qPCR) data, concerning both S. titanus adults and plants, as dependent variables, whereas the inoculation access period (IAP) was the categorical variable in both cases, counting three levels (IAP 1, IAP 2, and IAP 3). Concerning qPCR, negative values were excluded from GLM analyses. A binomial distribution with a Logit link function was used for

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infection rates, whereas a normal distribution and a logarithm link function was chosen for qPCR
data. When IAP effects were significant, Helmert contrasts were performed to identify differences
in infection rates and qPCR data (for both plants and insects), as follows: IAP1 vs. mean (IAP2,
IAP3) and IAP2 vs. IAP3.

180 **Results**

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182 FDP transmission by adult *S. titanus*

184 The acquisition of FDP by adult leafhoppers was performed on a total of eight experimentally 185 infected BBs. The phytoplasma concentration in those plants was stable, ranging from 1.46×10^4 to 186 1.60×10^5 FDP GU / sample, with an average density of 6.64×10^4 FDP GU / sample.

Quantitative PCR analysis on insects and BBs showed that transmission of FDP did actually occur, 187 188 and started from the first week after AAP (IAP1). The percentage of FDP-infected S. titanus adults 189 and their corresponding BBs obtained at the end of each IAP are presented in Table 1, along with 190 the average concentration of phytoplasma cells found in positive samples. FISH experiments 191 confirmed the results of qPCR analyses, as at the end of all IAPs, 16SrV phytoplasmas were found 192 not only in the midgut of leafhoppers (Fig. 2 A-C), but also in salivary glands (Fig. 2 D-F). 193 Phytoplasma- specific hybridization was not observed in insects directly collected form the mass 194 rearing, where the eubacterial signal was only detected (Fig. 2 G-I).

195 Considering S. titanus individuals, the infection rate reached 70% of the total, whereas infected BBs 196 were up to 25%. For both insects and plants, infection rates peaked at the end of IAP2, while the 197 lowest percentages of positive samples were recorded at the end of IAP1 as expected; however, according to binomial GLM analysis, no significant differences between IAP levels were found (S. 198 *titanus*: $\chi^2 = 5.359$; df = 2; P = 0.07; BBs: $\chi^2 = 0.634$; df = 2; P = 0.73). Symptoms (such as 199 yellowing and curling, especially of apical leaves) were observed on 9 BBs out of 13 which resulted 200 201 FD-positive after molecular analyses (69%). Symptomatic plants were evenly distributed among 202 IAP times (2 plants in IAP 1 and 3, 3 plants in IAP 2). On the other hand, none of the 49 FD-203 negative plants have shown any symptom.

The average concentration of phytoplasma cells ranged from 1.30×10^2 to 1.40×10^3 FDP GU / sample for leafhoppers and from 5.74×10^0 to 5.76×10^2 FDP GU / sample for inoculated plants, and in both cases IAP factor was significant (GLM, *S. titanus*: $\chi^2 = 16.60$, df = 2; P < 0.001; BBs: $\chi^2 =$ 29.76, df = 2; P < 0.001). Helmert tests showed significant differences between IAP levels, except from IAP 1 vs. mean (IAP 2, IAP 3) in broad beans (Table 2).

Discussion The results of the present study highlighted that, under laboratory conditions, S. titanus is able to acquire FD-C phytoplasma from infected BB as an adult too, and not only at the nymphal stage, as previously reported (Caudwell et al., 1970; Bressan et al., 2005b; Galetto et al., 2014). Indeed, phytoplasmas were successfully acquired after seven days of AAP, blooming to detectable loads after only seven days of LAP. Additionally, the phytoplasma load in insects increased over a period of 28 days (7 days AAP + 21 days LAP), indicating that the pathogen multiplies inside the leafhopper's body, in agreement with a persistent-propagative transmission model (Alma et al., 2015). However, overall phytoplasma concentrations were lower compared to those reported about other vectors, such as E. variegatus and Macrosteles quadripunctulatus (Kirschbaum), in similar experiments (Rashidi et al., 2014; Bosco et al., 2007). Furthermore, the mortality of S. titanus adults, which may be related to experimental conditions, since BB is not the favorite plant host for S. titanus (Chuche et al., 2016), suggests limited chance for adults to reach high levels of phytoplasma load, at least in laboratory. Under natural conditions, and especially on grapevine, adults' lifespan is longer, and acquisition of FDP from infected vines may therefore be more likely. This is in agreement with previous observations: S. titanus adults were thought to live up to 40 days (Vidano, 1964). Yet, recent (unpublished) results show how lifespan of adults is much longer under semi-natural conditions. If these data were confirmed, adults would be able to complete their transmission cycle (AAP+LAP+IAP) acquiring and transmitting phytoplasmas to vines, especially in the last part of the season, when no insecticides are used in vineyards. This is particularly threatening because of adults incoming from wild grapevine nearby (Lessio et al., 2014): a similar pattern in FD transmission implies that adults could acquire phytoplasmas directly on infected grapevines in a vineyard, even if coming from outside. Yet, another issue which needs further investigation is the influence of FDP on the fitness of S. titanus when AAP occurs at the adult stage. It has been demonstrated that FDP have a negative influence on lifespan and fecundity of S. titanus when AAP occurs at the nymphal stage (Bressan et al., 2005a). If confirmed for adults too, this could explain the low survival obtained in our experimental conditions. In addition, our results underlined that phytoplasma inoculation may occur after a shorter LAP than previously believed (Schvester et al., 1969; Caudwell et al., 1970), following an AAP performed

by adults. In effect, transmission to BBs was detected as soon as seven days after the end of AAP, with no significant differences in transmission efficiency with respect to longer LAPs, in spite of the higher phytoplasma titer recorded at the last IAP. Since similar percentages of infected BBs

were found at the end of IAPs 1-3, it can be assumed that the presence of positive leafhoppers at the end of IAP1, although bearing a low titer, is not merely due to transient ingested phytoplasma cells, but more likely to effective colonization of insects, which rapidly become infective. The transmission of a phloem-restricted plant pathogen by adult vectors has been previously demonstrated only for *Diaphorina citri* Kuwayama, vectoring 'Candidatus Liberibacter asiaticus'. However, diverging results were obtained by different studies: while Pelz-Stelinski et al. (2010) and Wu et al. (2016) reported successful adult transmission, even though with lower efficiency than in nymphs, Inoue et al. (2009) found no adults transmitting the pathogen. Moreover, Pelz-Stelinski et al. (2010) demonstrated that a one day long IAP is enough for D. citri to transmit liberibacter, but with an efficiency lower than 10%. Our results show that S. titanus adults are much more efficient in transmitting FDP, especially with short IAP, suggesting that different insect-pathogen dynamics occur for liberibacters and phytoplasmas, causing divergent transmission pathways.

Temperature may have played a role concerning multiplication of FDP both in insects and in plants. It has been (partially) demonstrated that, under lab conditions, multiplication of FDP in S. titanus was faster at low temperatures and CO_2 concentrations, whereas this trend was inverted in plants (Galetto et al., 2011). Our experiments took place at 25 °C, but under field conditions temperatures change along with the day and the season. It is possible therefore that the multiplication of phytoplasmas occur faster in the last part of summer, increasing the threat caused by long-living adults.

Although successful transmission was observed, the average phytoplasma concentration in infected BB samples was always lower if compared to densities reported about phytoplasmas in wooden plants (Raddadi et al., 2011; Galetto et al., 2016; Jawhari et al., 2015), possibly because of the herbaceous nature of BB. Moreover, higher FDP loads were recorded in BBs experimentally infected by E. variegatus (Salar et al., 2013); low phytoplasma concentration found in inoculated plants may be due to the limited number of infected leafhoppers hosted by each plant during our experiments and the reduced pathogen load in inoculating insects.

Even though it must be pointed out that our results were obtained with BB, where S. titanus was reported to acquire the phytoplasma more rapidly than from grapevine in laboratory conditions (Chuche & Thiery, 2014), this work suggests a potential role of S. titanus adults in acquiring FDP from infected grapevines in vineyards, and a consequent inoculation to healthy plants within insects' lifespan. This evidence opens to concerning scenarios for viticulture, explaining at least partially the epidemic development of FD which is observed in some cases, in spite of continuous control. Indeed, insecticide sprays are generally limited with respect to the occurrence of adults in the vineyard, which may be prolonged until autumn (Lessio et al., 2011) due to global increase of

temperatures, therefore the total contribution of adults to FDP transmission can be relevant.
Moreover, additional insecticide treatments in late summer, targeting adults, may be difficult to
apply due to food safety issues. In addition, the presence of hotbeds with overgrown grapevine's
rootstocks is a serious threat for vineyards due to incoming adults during late summer (Lessio *et al.*, 2014; Lessio *et al.*, 2015; Pavan *et al.*, 2012).

The results of this work, by showing that *S. titanus* is able to acquire and transmit FDP at the adult stage with reduced LAP in laboratory, represent the necessary scientific background for a deeper study of interactions involving adult leafhoppers, pathogens and host plants in different agricultural models, including the grapevine-FD pathosystem. Further researches are needed to shed light on such relationships, in order to improve the control of phytoplasma diseases.

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427 Table 1. Results of inoculation trials of FDP to broad beans (BB) by means of *S. titanus* adults.
428 Data of qPCR analyses on leafhoppers ("*S. titanus*" columns) and inoculated plants ("BB" columns)
429 are indicated.

 Table 2. Results of Helmert contrasts between levels of the categorical variable "Inoculation
Access Period (IAP)", for the dependent variable "quantitative PCR (qPCR)" concerning both *S. titanus* (ST) and broad beans (BB).

Figure 1. Experimental design of FDP transmission trials. Adult *S. titanus* [25 batches with 125 insects for qPCR analysis (A) and three batches with 15 insects for FISH analysis (B)] were caged for phytoplasma acquisition at day 0, and then from day 7 they were maintained on BB seedlings in groups of five. Every seven days live specimens were moved onto a new plant to perform up to three IAPs: 14 days (IAP1), 21 days (IAP2), and 28 days (IAP3) after the beginning of AAP, respectively.

Figure 2. FISH on S. titanus midgut and salivary glands. Exemplificative micrographs showing results from FISH of: midgut of a specimen at the end of IAP2 (A-C), salivary glands of an adult at the end of IAP3 (D-F), and midgut of a mass-reared individual (negative control) (G-I). A, D and G show interferential contrast micrographs of the organs. Hybridizations with Mollicutes probe (green signal in B) and eubacterial probe (green signal in E and H) are presented; moreover, hybridization with 16SrV phytoplasma probe is shown in C (cyan signal), F (red signal), and I. Bars = 75 μ m. The phytoplasmas were effectively detected in the organs of leafhoppers after transmission experiments, while they were absent in the negative control.

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	S. titanus				BB			
Rank	Ν	Positive samples	Infection rate ¹ \pm SE	$Titer^2 \pm SE$	N	Positive samples	Infection rate $^{1} \pm SE$	Titer ± SE
IAP1	65	30	0.46 ± 0.06	$1.30 \times 10^2 \pm 3.99 \times 10^1$	25	4	0.16 ± 0.07	$6.57 \times 10^{1} \pm 5.43 \times 10^{1}$
IAP2	38	26	0.68 ± 0.08	$5.05{\times}10^2{\pm}1.39{\times}10^2$	24	6	$0.25.\pm0.09$	$5.74{\times}10^{0}{\pm}1.33{\times}10^{0}$
IAP3	22	14	0.64 ± 0.10	$1.40{\times}10^3{\pm}6.42{\times}10^2$	13	3	0.23 ± 0.12	$5.76{\times}10^2{\pm}1.98{\times}10^2$

¹ Rate of 16SrV phytoplasma-positive individuals related to the total tested samples. ² 16SrV phytoplasma GU per sample (single insect or 100 mg of plant tissue). Values below the detection limit (1.00×10⁰ GU /sample) were considered negative (cut-off value). SE: Standard Error.

	IAP 1 vs. mean (IAP 2, IAP 3)			IAP 2 vs. IAP 3		
	χ ²	df	P	χ^2	df	P
BB qPCR	2.078	1	0.00	6.736 29.597	1	0.0



Experimental design of FDP transmission trials. Adult *S. titanus* [25 batches with 125 insects for qPCR analysis (A) and three batches with 15 insects for FISH analysis (B)] were caged for phytoplasma acquisition at day 0, and then from day 7 they were maintained on BB seedlings in groups of five. Every seven days live specimens were moved onto a new plant to perform up to three IAPs: 14 days (IAP1), 21 days (IAP2), and 28 days (IAP3) after the beginning of AAP, respectively.

209x180mm (300 x 300 DPI)



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59 60



FISH on *S. titanus* midgut and salivary glands. Exemplificative micrographs showing results from FISH of: midgut of a specimen at the end of IAP2 (A-C), salivary glands of an adult at the end of IAP3 (D-F), and midgut of a mass-reared individual (negative control) (G-I). A, D and G show interferential contrast micrographs of the organs. Hybridizations with Mollicutes probe (green signal in B) and eubacterial probe (green signal in E and H) are presented; moreover, hybridization with 16SrV phytoplasma probe is shown in C (cyan signal), F (red signal), and I. Bars = 75µm. The phytoplasmas were effectively detected in the organs of leafhoppers after transmission experiments, while they were absent in the negative control.

180x181mm (300 x 300 DPI)