Differential expression of Spiroplasma citri surface protein genes in the plant and insect hosts

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Abstract

Background: Spiroplasma citri is a cell wall-less, plant pathogenic bacteria that colonizes two distinct hosts, the leafhopper vector and the host plant. Given the absence of a cell wall, surface proteins including lipoproteins and transmembrane polypeptides are expected to play key roles in spiroplasma/host interactions. Important functions in spiroplasma/insect interactions have been shown for a few surface proteins such as the major lipoprotein spiralin, the transmembrane S. citri adhesion-related proteins (ScARPs) and the sugar transporter subunit Sc76. S. citri efficient transmission from the insect to the plant is expected to rely on its ability to adapt to the different environments and more specifically to regulate the expression of genes encoding surface-exposed proteins.

Results: Genes encoding S. citri lipoproteins and ScARPs were investigated for their expression level in axenic medium, in the leafhopper vector Circulifer haematoceps and in the host plant (periwinkle Catharanthus roseus) either insect-infected or graft-inoculated. The vast majority of the lipoprotein genes tested (25/28) differentially responded to the various host environments. Considering their relative expression levels in the different environments, the possible involvement of the targeted genes in spiroplasma host adaptation was discussed. In addition, two S. citri strains differing notably in their ability to express adhesin ScARP2b and pyruvate dehydrogenase E1 component differed in their capacity to multiply in the two hosts, the plant and the leafhopper vector.

Conclusions: This study provided us with a list of genes differentially expressed in the different hosts, leading to the identification of factors that are thought to be involved in the process of S. citri host adaptation. The identification of such factors is a key step for further understanding of S. citri pathogenesis. Moreover the present work highlights the high capacity of S. citri in tightly regulating the expression level of a large set of surface protein genes, despite the small size of its genome.

Keywords: Spiroplasma, Spiroplasma citri, Lipoproteins, Adhesins, Gene expression

Background

Spiroplasma citri is the etiological agent of stubborn disease of citrus in the Mediterranean area and California [1] as well as horseradish brittle root disease in the United States [2]. It has a complex life cycle that involves multiplication in the insect vector and in the host plant, indicating that S. citri has the ability to adapt to two very different hosts. S. citri is transmitted from plant to plant in a persistent propagative manner by phloem sap-feeding insects of the order Hemiptera. Once ingested from the phloem vessels of an infected plant by the leafhopper vectors, Circulifer tenellus or Circulifer haematoceps, S. citri invades the entire insect. The circulative route of S. citri through its leafhopper vector is well established: spiroplasmas cross the insect gut wall, move into the hemolymph where they multiply, circulate, and invade most of the insect organs including the salivary glands, and are released in the main salivary...
duct leading to the stylet’s salivary canal. They are then introduced into the plant phloem along with salivary secretions during feeding [3–5]. In the host plant, *S. citri* multiplies in the phloem sieve elements and triggers severe symptoms. In the experimental host plant (periwinkle *Catharanthus roseus*), *S. citri* induces leaf yellowing, wilting and stunting [1]. The molecular mechanisms underlying the interaction between *S. citri* and its host plant remain largely unknown. Nonetheless, a perfect correlation between the ability of *S. citri* to use fructose and its ability to induce symptoms in the host plant was demonstrated [6].

In bacterial pathogens, many lipoproteins have been shown to play a key role in virulence-associated functions such as adhesion, invasion and colonization [7, 8]. In *S. citri*, surface proteins are suspected to recognize the insect gut and/or salivary glands epithelium, possibly participating in both adsorption and endocytotic events mediated by receptor ligand interactions [4]. The *S. citri* GII-3 genome (1,820 kbp) [9] encodes 645 membrane proteins including 68 putative lipoproteins, as predicted by the presence of a consensus lipobox in the first 28–30 amino-acids [10] and 577 transmembrane proteins [11]. The major lipoprotein at the cell surface of *S. citri* is the protein named spiralin [12], which is required for efficient transmission of *S. citri* by its leafhopper vector [13] and, which was further shown to act *in vitro* and *in vivo* as a lectin able to bind glycoproteins of the vector insect [14, 15]. In addition, the surface lipoprotein Sc76 homolog to a solute-binding protein of an ABC transporter was also found to be involved as disruption of the gene dramatically reduced *S. citri* ability to be transmitted by *C. haematoceps* [16]. The *S. citri* GII-3 genome is also characterized by an abundant extrachromosomal DNA content, including seven plasmids, pSciA and pSci1 to pSci6, present as 10 to 14 copies per cell [17]. Plasmid genes also encode proteins associated with spiroplasma transmission. Recently, the role of the 8 surface adhesion-related proteins (ScARPs) encoded by plasmids pSci 1 to pSci 5 [17] has been studied. As compared to the wild-type strain GII-3, the *S. citri* mutant G6/6 [18] and the non-insect-transmissible, strain 44 [19] both lacking pSci1 to 5, were affected in their ability to adhere and enter into the leafhopper cells [20, 21]. The role of ScARPs in adhesion and entry in leafhopper cells was clearly demonstrated for the ScARP3d, which possesses the whole set of domains found in ScARPs as well as the largest repeated domain [21]. Only a few membrane proteins have been investigated for their implication in transmission of *S. citri* by its vector insect, and the putative role of surface-exposed proteins in plant disease has not been studied. Moreover, despite surface proteins have been shown to be involved in transmission of *S. citri* by the leafhopper vector, very few is known about their regulation. Only one study dealt with gene regulation in *S. citri* [22]. In this work, the genes encoding the glucose and trehalose permeases were shown to be up-regulated in the presence of the respective sugars, reflecting the capacity of *S. citri* to adapt to environments with distinct carbohydrate contents [22].

Understanding the changes in membrane protein gene expression in response to different environmental conditions (plant and insect) is an important step in unraveling the possible functions of these genes and the transcriptional regulation mechanisms in *S. citri*. To investigate the molecular adaptation of *S. citri* in its different environments we compared the expression level of 28 putative lipoprotein genes including *spiralin* and *sc76* in *S. citri*-infected leafhoppers with those in *S. citri*-infected periwinkle plants. Expression profile of the *S. citri* ScARP genes was also assessed in the plant and in the leafhopper host. In addition, considering that spiroplasmas persisting in plants for a long period of time may lose the expression of genes necessary for insect host adaptation or may overexpress genes involved in plant long-term adaptation, insect-infected plants were compared to graft-inoculated plants for *S. citri* lipoprotein gene expression.

**Methods**

**Spiroplasmas, plants and leafhoppers**

The *S. citri* wild-type strain GII-3 (GII-3 wt) was first isolated from its leafhopper vector, *C. haematoceps*, in Morocco [23]. To avoid loss of transmissibility due to extensive in vitro passaging, the working strain was periodically subcloned and selected clones were submitted to experimental transmission to periwinkle plants via injection to its leafhopper vector *C. haematoceps* to confirm transmissibility and pathogenicity. After isolation from symptomatic plants, batches of *S. citri* cultures with low passage numbers (less than 5p) were stored at -25 °C until use.

*S. citri* G-GIP (this study) was isolated from *S. citri* GII-3 wt, graft-infected periwinkles (GIP) five months after grafting. In this strain the *scarp2b* mRNA transcript was not detected in the first passage culture (strain G-GIP1), but was readily detected after ten passages in the culture medium (strain G-GIP10). Spiroplasmas were grown at 32 °C in SP4 medium [24].

Intra-abdominal microinjection of *S. citri* into *C. haematoceps* leaffoppers and transmission to periwinkle host plant (*Catharanthus roseus*) were previously described [6, 25]. Leafhoppers were injected with low passage (7 to 10p) cultures of *S. citri* GII-3 and caged on healthy stock plants (*Matthiola incana*) for 2 weeks before being randomly divided in 2 groups. The first group was used directly for insect DNA extraction; the second
group was transferred onto young periwinkle host plant (six-to-eight-leaf stage, 10 insects per plant) for a period of 3 weeks (transmission period). Plants with symptoms were designated in our study as ‘leafhopper-infected periwinkles’. (LIP).

*S. citri* GII-3 was also maintained into periwinkle plants by successive graft inoculations without any insect transmission. In this case, inoculum sources were symptomatic branches of 1 year old plants originally infected with *S. citri* GII-3 via leafhopper transmission.

**DNA isolation and spiroplasma quantification by quantitative PCR**

Total DNA from pure culture of *S. citri* was isolated by using the Wizard genomic DNA purification kit (Promega, Madison, Wis. USA). Five hundred milligrams of midribs collected on infected periwinkles (GIP or LIP) were ground in a plastic bag in a Homex 6 homogenizer (Bioreba AG, CH-4153 Reinach BL1, Switzerland). Total DNA was extracted using the CTAB (cetyl trimethyl ammonium bromide) method according to Murray and Thompson [26]. DNA from leafhoppers was also purified by the CTAB method. DNA preparations were kept at -20 ° C.

For quantitative real time PCR, the LightCycler® 480 SYBR Green I Master Mix (04887352001, Roche) was used. The SYBR Green reaction was performed in a 30 µl reaction mixture containing 1 X master mix, 0.15 mM of each primer, and 1 µg of total DNA preparation. The LightCycler® 480 System (Roche Diagnostics GmbH Mannheim, Germany) was used with the following program for DNA amplification: 95 °C for 15 min, 40 cycles each at 95 °C for 30 s, 67 °C for 30 s, 72 °C for 30 s, and a final extension at 72 °C for 10 min. Primers used in quantitative amplification of *S. citri* DNA were designed from the spiralin gene [EMBL:Q2YHQ8]; the sequences of forward primer SQ1 and reverse primer SQ2 were 5′ ACAACGAAGGTACATCATTAACAAC 3′ and 5′ TTTGCTGGAGTAATTTGAACATAAAC 3′, respectively, and led to an amplicon of 80 bp.

For absolute quantification, plasmid pES3’ [27] containing the spiralin gene was used to construct the calibration curve and calculate the PCR reaction efficiencies. Knowing the number of plasmid molecules in 5 µl, tenfold serial dilutions of the plasmid DNA were prepared and used to generate the standard curve. To determine the theoretical sensitivity and the reliability of the qPCR, three repetitions of the assay were performed.

**RNA extraction from infected and uninfected hosts and cDNA synthesis**

Total RNA from spiroplasma-infected plants, infected leafhoppers, and from spiroplasma cells in culture were isolated by using Trizol Reagent according to manufacturer’s guidelines (Invitrogen CA, USA). For periwinkle plants, 5 to 10 leaf midribs were ground in a mortar by freezing with liquid nitrogen and homogenized in 1 mL of Trizol reagent. Fresh leafhoppers (‘10) were ground directly in the Trizol Reagent. Total RNA from frozen *S. citri* cell pellets harvested by centrifugation during the exponential growth phase was extracted following the same procedure as above for the different hosts.

Subsequently, all RNA samples were treated with RNase-free RQ1 DNase (Promega, Madison, WI, USA) for 1.5 h at 37 °C to remove residual DNA, ethanol-precipitated, and finally dissolved in water following the protocol described by the supplier. DNase I treated RNA samples were tested in conventional PCR with primers SQ1-SQ2 without the RT step to confirm the absence of significant amounts of contaminating genomic DNA.

For each sample (spiroplasmas, healthy and infected hosts) 1 µg of DNA-free RNA was used for cDNA synthesis using Superscript Reverse Transcriptase III and random or specific primers according to manufacturer's guidelines (Invitrogen, CA, USA).

**Quantitative real time reverse transcription PCR (RT-PCR)**

Quantitative real time RT-PCR assays were performed on cDNA templates using the SYBR green chemistry detection system according to the manufacturer's instructions (Roche Diagnostics GmbH Mannheim, Germany). In order to validate changes in transcript levels, identification of reference genes whose expression is independent of the environmental conditions is required. The genes selected to be tested as possible reference transcripts are listed in Table 1. Quantitative real time RT-PCR was performed using 5 µl of template in 1 X Light Cycler 480 SYBR Green Master Mix and 0.15 µM of each primer in a total volume of 25 µl. Primers used in the qRT-PCR assays for evaluating the expression of *S. citri* lipoproteins in leafhoppers and plants were listed in Table 2.

The relative quantification method (ΔΔCₜ) [28] was used to evaluate quantitative variation between the hosts (plant or leafhopper) and the culture arbitrary designated as “calibrator” in our study. For the calibrator sample, the average Ct value of the reference gene was subtracted from the Ct value of the target gene under investigation to give ΔCₜ_calibration.

The ΔCₜ values for plants or insects were calculated using the same procedure. The ΔΔCₜ value calculated for each host was obtained by subtracting the respective ΔCₜ of the target gene in the calibrator sample from those of the target gene in the host. The data were analyzed using the following equation [28].
Experiments were carried out on three independent biological replicates, each consisting of three replicate reactions. A significant change in $\Delta\Delta C_T$ value in host versus axenic medium was considered if $\Delta\Delta C_T$ value was superior to 1. To identify genes for which expression was significantly different in various hosts, statistical analyses were performed using Student’s t test ($P < 0.05$).

**Bi-dimensional gel electrophoresis and nLC-MS/MS analysis**

Spiroplasma proteins from 150 mL culture were prepared as previously described [29]. Proteins (300 $\mu$g) solubilized in 1–5 mL of a rehydration solution containing 7 M urea, 2 M thiourea, 4 % (w/v) CHAPS, 2 % Triton X-100, 10 mM DTT and 2 % (v/v) Ampholine pH 3–10 were submitted to 2-D gel electrophoresis as described before [29]. Gels intended for LC-MS/MS analysis were stained using Coomassie brilliant blue [30]. For the nLC-MS/MS analysis, the gel spot (1 × 1 mm) present in the 2D-gel obtained for *S. citri* GII-3 wt expressing the *scarp2b* gene was excised then treated with destaining solution consisting of 25 mM ammonium bicarbonate and 50 % ACN. The gel piece was rinsed twice in ultrapure water and shrunk in ACN for 10 min. After ACN removal, the gel piece was dried at room temperature, covered with trypsin solution (10 ng/μL in 40 mM NH₄HCO₃ and 10 % ACN), rehydrated at 4 °C for 10 min, and finally incubated overnight at 37 °C. The gel piece was then incubated for 15 min in 40 mM NH₄HCO₃ and 10 % ACN at room temperature with rotary shaking. The supernatant was collected, and an H₂O/ACN/HCOOH (47.5:47.5:5) extraction solution was added onto the gel piece for 15 min. The extraction step was repeated twice. Supernatants were pooled and concentrated in a vacuum centrifuge to a final volume of 25 μL. Digests were finally acidified by addition of 2.4 μL of formic acid (5 %, v/v) and stored at -20 °C.

The peptide mixture was analyzed on a Ultimate 3000 nanoLC system (Dionex, Amsterdam, The Netherlands) coupled to an Electrospray LTQ-Orbitrap XL mass spectrometer (Thermo Fisher Scientific, San Jose, CA). The conditions of peptide separation and data acquisition were identical to those described in [31]. Data were searched by SEQUEST through Proteome Discoverer 1.4 (Thermo Fisher Scientific Inc.) against a custom made *S. citri* GII-3 database.

### Table 1

| Name | GenBank accession | Symbol | Function | Primer sequences (5′ to 3′) | Amplitcon length |
|------|------------------|--------|----------|-------------------------------|-----------------|
| Putative chromosomal replication initiator protein dnaA | SPIC101B_001 | dnaA | Replication | ATGAGTAATCCAGGAGTTAGTCTTTG CCACCGAACTCTG | 116 |
| DNA gyrase subunit b protein | SPIC101B_003 | gyrB | Topoisomerase | GGGAGATCTGCGTGAGGAGTTCTTT AATACCTGCTCCTAATGCG | 167 |
| DNA gyrase subunit a protein | SPIC101B_004 | gyrA | Topoisomerase | TTTGCCCAAAGGAAAGTAGCTCAGT AATGTCATGACAAATTC | 195 |
| DNA-directed ma polymerase beta chain protein | SPIC101B_073 | rpoB | Transcription | TGGTCCATTAGTGCGCTAAGCATCTCTTC AATACAAATCGAGTTT | 179 |
| Hypothetical chromosome replication initiation and membrane attachment protein | SPIC103_040 | dnaB | Replication | AATTACAAATTCGGCAAATGTGCTTTGT CTTTCTATTATTAAC | 131 |
| 50s ribosomal protein L3 | SPIC103_102 | rplC | Ribosomal protein L3 | AAGCCTGAGCATTGGGAGACGCTCAAA CAACTACAAGTGG | 252 |
| Spiralin lipoprotein | SPIC104_139 | spi | Lipoprotein | ACAACGAAAGGTTACATTTAACAACCTTG ATCGAAGTTAATGGAACATTAAC | 80 |
| Pyruvate kinase protein | SPIC104_141 | pyK | Glycolysis | GGGAGATTAAAAAACAATTTTCTGCAC AATCAAAATGTC | 171 |
| Fibril protein | SPIC112_006 | fib | Cytoskeleton Structure | TAAGCATTGACAGAGATACAAACTGCCC ATATTAAATCAACATTTTC | 246 |
| Cell shape-determining protein mreB1 | SPIC113_009 | mreB1 | Cell morphogenesis | AGGAACACAACGACTTGCAGTACCC CATATTGAGACAC | 125 |
| 16S rRNA | ND | 16S | 30S ribosome subunit RNA component | CAAATTCCTGGAGCTAACTGCGTAGAC ACTAGGATATCAT | 204 |
| Name         | Primers sequences (5’ → 3’)                          | Gene product                              | Amplicon length (bp) | Annealing temperature (°C) | Efficiency (%) |
|--------------|------------------------------------------------------|--------------------------------------------|----------------------|----------------------------|----------------|
| pSci4_02     | GCGAATGACTCCATCAAGTCTCGT and TGGTTCTTTACTGTTGAGTGG  | Hypothetical lipoprotein                   | 221                  | 52                         | 99.9           |
| pSci4_06     | ATCAGTAAACATGCTCAGG and TATCAAGCCAGTTGCCTGCTC       | Hypothetical lipoprotein                   | 334                  | 52                         | 91.0           |
| pSci6_18     | AGTTGTCGCTCGTCTTAG and GATCTGCTCGTAC                | Truncated adhesion-related protein         | 173                  | 60                         | 93.9           |
| SPICI01A_047 | GATGTAGAATTCGCCAAG and TCGATTCGCCTCGTC              | Hypothetical lipoprotein                   | 563                  | 52                         | 102.8          |
| SPICI02_046  | TGCAACAAACAAAGTCAACG and TAGCAAGAACCGTATTCCATG      | Hypothetical lipoprotein                   | 288                  | 60                         | 95.9           |
| SPICI03_030  | AGTAAACATCCACACCTTATTG and ATCCGTTGCTACACAC        | Hypothetical lipoprotein                   | 219                  | 60                         | 103.0          |
| SPICI03_098  | GTTACAGGGAGGGCCGATG and TGGCAAGAACTGCTGATTG         | Hypothetical lipoprotein                   | 573                  | 60                         | 100.2          |
| SPICI03_180  | TTGGGAAAAAGGAGGTTGGAATG and CTGGTCAACCGG           | Hypothetical lipoprotein                   | 659                  | 60                         | 100.0          |
| SPICI03_317  | GAAATAGTGTAAATGATGTTAG and GCAGTTAACATTACCAAGAC    | Hypothetical lipoprotein                   | 184                  | 52                         | 108.4          |
| SPICI04_017  | CACCAACTTCAATACATATCC and ATTTACAGTACGACTCAG       | Hypothetical lipoprotein                   | 86                   | 60                         | 99.7           |
| SPICI04_108  | ACTCTGCCGCTCTATTACCTCAG and CCTGTCGTAACCAAGC       | Hypothetical lipoprotein                   | 157                  | 60                         | 100.5          |
| SPICI04_139  | ACAACGAAGTCACTATTACCAAC and ATTTACAGTACGACTCAG     | Hypothetical lipoprotein                   | 80                   | 60                         | 100.8          |
| SPICI05_014  | GCCGTAACACTCTTTTGACTC and ATTTAGTCGCAATTACCAAG     | Hypothetical lipoprotein                   | 138                  | 60                         | 96.1           |
| SPICI06_025  | CTAATACACAAACAGCCGCC and CTTTATCAATGAGTACGGT       | Hypothetical lipoprotein                   | 161                  | 60                         | 103.1          |
| SPICI07_030  | CTTCCGCATACACTACTAAG and ATCTACTAACAGTACG          | Hypothetical lipoprotein                   | 160                  | 52                         | 108.8          |
| SPICI09_027  | TGCCCGCCTAAATCTTTTG and TGATTTATGATGATGATCAG       | Hypothetical lipoprotein                   | 153                  | 60                         | 108.8          |
| SPICI11_054  | CTCCTCGGCTGCATC and CAAGCTTGTGCTAAGTACG           | Hypothetical lipoprotein                   | 505                  | 60                         | 97.8           |
| SPICI11_055  | GGTAGCAAGAAATGATGACG and CCTGCCTGAAGTACG           | Hypothetical lipoprotein                   | 206                  | 60                         | 96.1           |
| SPICI11_003  | GTCAATCAAGATAGTTGCCAG and GTGCAATTAAAAGAGTA         | SC76                                      | 157                  | 52                         | 104.6          |
| SPICI12_020  | TGCAATCGGCAGTATGATGTC and CTGAATTACATACAG          | Hypothetical lipoprotein                   | 200                  | 60                         | 96.3           |
| SPICI12_021  | TATGCTTGGTATGATGTC and CTCAATTACGTCACAC           | Hypothetical lipoprotein                   | 536                  | 60                         | 98.2           |
| SPICI12_028  | ACGTATTAACACATTTTGAGT and TCCAGACAGGAGCCTCTT       | Hypothetical lipoprotein                   | 126                  | 60                         | 95.6           |
| SPICI13_014  | AACAATTAAACCACCAAGAAG and CACATAGACAAATTGCTTG      | Hypothetical lipoprotein                   | 228                  | 60                         | 98.4           |
| SPICI16_011  | GTCAGTACCCAGTTTTAGGC and AGCAAGAAGGAAACACGC        | Hypothetical lipoprotein                   | 535                  | 60                         | 90.6           |
| SPICI20_004  | GAATTATGAGGAGGAGAC and AAGTTAAGTGAATCCTCTCC        | Hypothetical lipoprotein                   | 191                  | 60                         | 93.3           |
database. Spectra from peptides higher than 5000 Da or lower than 350 Da were rejected. The search parameters were as follows: mass accuracy of the monoisotopic peptide precursor was set to 10 ppm and peptide fragments tolerance was set at 0.6 Da. Only b- and y-ions were considered for mass calculation. Oxidation of methionines (+16 Da) and deamidation of asparagine and glutamine (+1 Da) were considered as variable modifications and two missed trypsin cleavages were allowed. Only high-confidence matches corresponding to false positive rate of 1 % at peptide level were considered.

Results and discussion
Quantification of spiroplasmas in periwinkle plants and insects
From the standard curve constructed with serial dilutions of the plasmid pES3′, real-time PCR assay was used to accurately quantify the spiroplasmas cells in plant and insect extracts used in this study. One ng of pES3 contains 10⁸ molecules of plasmid, each containing one copy of the spiralin gene. Because this gene is present in a single copy on the spiroplasma chromosome, 1 ng of pES3′ corresponds to 10⁸ spiroplasmas. The number of spiroplasmas in 1 μg of DNA extracted either from periwinkle plants infected by insects (LIP) or grafted periwinkle plants (GIP) fresh midribs or from infected insects was similar and corresponded to 1.5 ± 0.5 × 10⁵, 1.5 ± 0.7 × 10⁵, and 1.8 ± 0.6 × 10⁵, respectively. The average number of spiroplasmas in 1 g of fresh midribs from LIP reached 2.4 ± 0.8 × 10⁷; from GIP the average value was 3.3 ± 1.7 × 10⁷. The number of spiroplasmas in leafhoppers was equivalent to 6.4 ± 2.4 × 10⁶ spiroplasma cells per insect.

Selection of reference genes for transcript quantification in plants and leafhoppers
The overall cycle threshold values C<sub>t</sub> for the 11 genes selected as candidates for normalization of transcript level determination (Table 1) in plants and leafhoppers were distributed from 15 to 30. Control reactions without S. citri template, performed with cDNAs from healthy plants and insects, remained below the threshold for genes 16S, spi, fib, pyk, rplC. Weak unspecific signals detected with rpoB, gyrA and B, dnaA and B, and mreB in the no template controls excluded these genes from the study.

For four candidate genes fib, pyk, rplC, and spiralin, transcript level in the 2 hosts was analyzed by absolute quantification. For each gene, a standard curve prepared with known concentrations of the same gene previously cloned in a plasmid gave regression lines with an average slope value of -3.542 and an average error value of 0.04. All PCRs displayed an efficiency ranging from 96 to 99 %. Such efficiency is considered acceptable and this relatively high-efficiency value results in a better sensitivity at low target concentrations.

As shown on Fig. 1, expression of spiralin and rplC varied according to the hosts and were expressed at higher levels in leafhoppers than in plants. Also spiralin transcripts were 10 times more abundant in both hosts than the other gene expression products. Thus these 2 genes cannot be used as internal controls. Comparison of the transcript levels of fib and pyk in infected leafhoppers and plants (Fig. 1) showed that each of these genes was equally expressed in both environments. Transcript

| Table 2 List of primers used to study lipoprotein genes expression in Spiroplasma citri GII-3 (Continued) |
|---------------------------------------------------------------|
| SPIC120_057 | TTGATGAACTGCCTTCTATTG and CTTG TGCATTATTGAAAAC | Hypothetical lipoprotein | 360 | 60 | 95.5 |
| SPIC120_065 | GTCAGGCAGCTACTCC and GCTGA GCCAGACCTGGAAC | Hypothetical lipoprotein | 562 | 60 | 100.3 |
| SPIC120_066 | TTAGGCTATGATAGTG GG and ATAC CTGGTGGTGGTC | Hypothetical lipoprotein | 445 | 60 | 91.8 |

Fig. 1 Absolute quantification of 4 candidate reference genes (spiralin, fib, pyk and rplC) in leafhopper-infected plants (LIP) and in insects. Grey bars indicate the number of transcripts detected in LIP and the black bars the transcript level in insects. Bars correspond to the standard deviation obtained with three independent replicates.
levels of fib and pyk in leafhoppers and plants were then compared to those in the culture medium. The number of transcripts per spiroplasma was not significantly different in leafhoppers, in Lip and in the culture medium (0.1 ± 0.02 for fib and 0.07 ± 0.001 for pyk). Considering that fib and pyk were equally transcribed in the three environments these genes could be used as internal controls. Given that the fibril protein, but not Pyk, is specific to spiroplasmas [32], fib was used as the reference gene in further experiments.

Lipoprotein genes expression profiles in the different environments

To investigate the molecular adaptation of S. citri in plants and leafhoppers, we examined the expression of spiroplasma lipoprotein genes. Among the 68 genes predicted to encode lipoproteins, genes with redundant sequences (of viral origin), pseudogenes (with the exception of pSci6_18), as well as genes, for which no satisfactory amplification primers could be identified were removed from the study. Among the 28 selected genes, 3 were carried by plasmids pSci4 and pSci6, and 25 others, including genes spiralin (SPICI04_139) and sc76 (SPICI11_003) were carried by the chromosome (Table 2). Evaluation of mRNA expression profiles in both environments was conducted in leafhoppers, and in the 2 types of periwinkle plants Lip and Gip (see section Methods). The relative gene expression level was calculated as described in Methods where the transcript level in SP4 medium of the target and reference (fib) genes were chosen as calibrators. All PCRs displayed an efficiency ranging from 90.6 to 108.8 %.

The calculated -ΔΔCt values for the 28 tested genes are shown in Fig. 2 and in Additional file 1: Table S1. A positive −ΔΔCt value indicates an up-regulation of the gene’s expression whereas a negative −ΔΔCt indicates down-regulation as normalized to the fib reference gene.

Lipoprotein gene expression profiling in plant LIP and leafhopper hosts

Lipoprotein gene expression profiles in infected insects versus spiroplasmas cultured in SP4 (black bars, Fig. 2). During cultivation in SP4 medium all 28 S. citri lipoprotein genes were expressed. Expression of fifteen of them significantly changed once the spiroplasmas were introduced in insects, indicating the strong transcriptional response of S. citri to environmental changes. Among these genes, 12 (including SPICI11_003 (sc76) and SPICI12_021 (oppA)) were up-regulated, while 3 were down-regulated in insects. These were operationally defined as “insect-up-regulated” and “insect-down-regulated”, respectively.

Lipoprotein gene expression profiles in spiroplasmas from leafhopper inoculated plants (LIP) versus those in spiroplasmas grown in culture (grey bars, Fig. 2). The pattern of lipoprotein gene expression was found to be clearly different in LIP. In the host plant, 8 lipoprotein genes were up-regulated and 7 genes (including spiralin) were down-regulated.

Lipoprotein gene expression profiles in spiroplasmas from leafhopper inoculated plants (LIP) versus those in spiroplasmas from leafhopper bodies (grey bars versus black bars, Fig. 2). Eight genes were up-regulated in both plant and insect hosts, whereas 2 were significantly down-regulated in both hosts. Fifteen genes (for which a \( |\Delta\Delta C_T| > 1 \) was obtained for at least one host compared to SP4) were differentially expressed in insects and in plants (indicated by an asterisk on Fig. 2). For most of them (12/15), the transcript level was higher in insects than in plants. Four genes (SPICI04_108, spiralin, SPICI06_025 (prophage element), SPICI12_020) were down-regulated in plants but not in the leafhopper. Genes pSci4_02, SPICI12_021, SPICI12_028, SPICI16_011 were overexpressed in insects but neither up- nor down-regulated in LIP. Expression of SPICI20_065 transcripts was repressed only in insects (no activation or repression in infected LIP) compared to SP4. Six genes followed the same type of regulation in both hosts but showed significantly different transcript levels in LIP and in insects (SPICI01A_047, more strongly up-regulated in plants; SPICI02_046, SPICI03_098, SPICI10_054 more strongly up-regulated in insects; SPICI13_014, more strongly down-regulated in insects; SPICI10_055 more strongly down-regulated in plants).

To summarize, during in vitro cultivation of S. citri in SP4 medium, the 28 lipoprotein genes tested were expressed. Once the spiroplasmas were introduced into the leafhopper host, 12 of the lipoprotein genes were up-regulated. Among these genes that are up-regulated, and thus putatively involved in adaptation of the bacteria to its insect host, the majority encode hypothetical lipoproteins with no assigned function (Table 1). However, two of them (SPICI11_003 (sc76) and SPICI12_021 (oppA)) are noticeable, as they encode proteins sharing sequence identity with substrate binding units of two distinct ABC transporters. The superfamily of ABC transporters plays an important role in the export of proteins and polysaccharides and in the import of sugars, inorganic ions, and oligopeptides [33]. SPICI11_003 (sc76) was found to be up-regulated in both insects and plants, suggesting that this gene may be required in both hosts. Sc76 is a solute binding protein of a sugar ABC transporter, for which the sugar specificity has not yet been identified [16]. Sc76 could play a role in S. citri GII-3 growth, as the S. citri mutant G76 having a truncated sc76 gene multiplies to low titers in plants and leafhopper salivary glands compared to GII-3 wt [16]. In plants sucrose is the most abundant sugar and the spiroplasmas preferred
sugar is fructose [34], while the main sugar in the insect hemolymph and salivary glands is trehalose [35, 36].

The fact that, in plants and insects, the transcription of sc76 was up-regulated at a similar level suggests that expression of sc76 might be regulated by a sugar present in both hosts. Even if glucose, which is abundant in SP4 medium can be easily excluded, it remains to be investigated which sugar might be transported through the Sc76-containing ABC transport system.

Interestingly, S. citri pathogenicity was severely impaired in fructose operon mutants [6], and transcription of the genes coding for the phosphoenolpyruvate transferase systems (PTS) responsible for glucose and fructose import into the spiroplasma cell were stimulated by the respective sugar [22, 37]. The present work provides further evidence of the crucial role of sugar metabolism in spiroplasma pathogenicity, and suggests that the sugar transported via Sc76 could participate in S. citri’s adaptive capacity to multiply in distinct hosts.

The lipoprotein SPICI12_021 shares identity with the substrate binding unit OppA of the oligopeptide permease. Over the past 10 years OppA has been characterized as a multifunctional lipoprotein in mollicutes. As for an example, OppA was not only involved in oligopeptide import into the cytoplasm, but also in cytadherence to and invasion of epithelial surfaces of the human urogenital tract by M. hominis [38]. In M. pneumoniae a lipoprotein gene (mpn456) having with homology to a gene encoding a predicted oligopeptide ABC transport system was also up-regulated in response to adhesion to a human cell line [39]. More generally, in several bacteria other than mollicutes, such as Mycobacterium tuberculosis, solute binding proteins of ABC transporters are lipoproteins that play a role in bacterial growth and contribute to virulence [8]. As in the human mycoplasmas M. hominis and M. pneumoniae, the protein SPICI12_021 could play a key role in the spiroplasmas’ interactions with leafhopper cells, which are crucial steps for transmission of S. citri by its vector insect.

The spiralin gene was shown to be strikingly down-regulated in plant (LIP) whilst it was abundantly expressed in insects and in culture. Spiralin is the most abundant lipoprotein of S. citri membrane, and covers the entire spiroplasma cell surface [12, 40]. This lipoprotein was designated as a lectin interacting with insect glycoproteins [14, 15] and was required for adhesion and entry of S. citri into insect cells [15]. Thus it could be hypothesized that, during transmission of S. citri to plant hosts, over-expression of spiralin within the leafhopper vector would occur to enable adhesion and internalization of spiroplasmas into midgut and salivary glands cells.
These results revealed infection regulatory programs common to both hosts as well as genes submitted to insect- or plant-specific regulation, indicating a fine-tuned regulation of several lipoprotein genes depending on the *S. citri* environment, despite the reduced genome size of this bacterium [9]. Guell et al. [41] have analyzed large transcriptomic data sets obtained with *M. pneumoniae* cultivated under a broad range of conditions and submitted to diverse stresses. Their study highlighted the unanticipated, high transcriptome complexity in mollicutes. Considering that *M. pneumoniae* possesses one of the smallest genomes among mollicutes, it is plausible that a high level of transcriptional regulation also occurs in other mollicutes upon environmental changes. Nevertheless, most transcriptional variations that occur in mollicutes upon environmental changes have been recorded *in vitro*, and there are only a few *in vivo* studies [39, 42, 43]. Unlike mycoplasmas, *S. citri* invades two very different hosts and our data demonstrate differential expression of genes encoding membrane-anchored proteins in plants and in insects. This study provided us with a list of lipoprotein genes putatively involved in *S. citri* adaptation to its hosts and possibly underlying virulence and/or host specialization. Genes pSci4_02, SPICI10_054, SPICI12_021, SPICI12_028, and SPICI16_011, which, like spiralin, sc76, and oppA, are overexpressed in the leafhopper, and SPICI04_108, SPICI06_025 (prophage element), and SPICI12_020, which are down-regulated in plants are therefore good candidates for being involved in the adaptation of *S. citri* to its insect host. On the contrary, genes that are strongly up-regulated in LIP (such as SPICI01A_047) or are less repressed in plants than in insects (SPICI13_014, SPICI20_065) are expected to encode proteins involved in adaptation to the host plant.

**Comparison of lipoprotein gene expression in LIP and GIP plants**

To analyze the putative adaptive response of *S. citri* during long-term plant infection, lipoprotein genes’ expression in leafhopper-infected periwinkles LIP (recent infection) was compared to those in graft-infected periwinkles GIP, in which *S. citri* GII-3 wt was inoculated through grafting 5 months earlier (old infection). In both cases (graft- and insect-inoculation) the infected plants share similar symptoms suggesting they share similar physiological responses to *S. citri* infection.

Six spiroplasma lipoprotein genes were up-regulated in GIP and LIP infected hosts (Fig. 3 and Additional file 1: Table S1). Among them, SPICI01A_047 was less expressed in insects. The up-regulation of this gene could be involved in the protection of *S. citri* from plant defence or in the successful colonization of host plant by the spiroplasma. Three genes (pSci6_18, SPICI04_108, SPICI04_139 (spi)) were down-regulated in both GIP and LIP hosts and to a similar extent, suggesting that these genes are likely not to contribute to the adaptation of *S. citri* to the plant host.

For one gene SPICI 05_014 overexpression level was significant in GIP but not in LIP, in which the transcript level was similar to that measured in the leafhopper and in the SP4 culture medium. Instead, the plasmid gene pSci4_06, and the chromosomal genes SPICI04_017, SPICI20_066 were transcribed at a very low level in GIP, while their expression did not significantly differ in LIP and in leafhoppers. The changes of expression of these genes in GIP compared to LIP could be due either to the age of the host plant, to the duration of infection, or to the mode of inoculation (graft vs insect-infection), three variables differing in LIP and GIP infected-plant models. Due to differences between old and young plants in the efflux of essential micronutrients such as sugar and amino acids in the phloem sap, changes in gene expression between old-grafted (GIP) and young (LIP) periwinkle plants could reflect an adaptation to the old plant environment. Variation of gene expression in GIP compared to LIP could also be observed for genes having a role during the early stage of infection but not for spiroplasma persistence, when *S. citri* is well adapted to the host plant (GIP). Finally, in the case of GIP, the lack of an interim period of habitation in insects might be responsible for down-regulation of spiroplasma genes that are non-vital for survival in plants but are involved in adaptation to insects. Indeed the lack of exposure to the selective pressure exerted in the insect host is likely to alter the expression of such genes.

**Transcriptional analysis of scars in *S. citri***

Eight *S. citri* adhesion-related proteins (ScARPs) are encoded by plasmids pSci1 to 5 whose presence has been associated with the ability of *S. citri* to be transmitted by its leafhopper vector [20, 21]. To determine whether *scarp* genes, similarly to several lipoprotein genes, were regulated by environmental conditions, their expression levels in SP4, in infected insects, GIP and LIP were compared. To avoid misinterpretation due to variable copy number of the different plasmids, the *scarp* genes *scarp5a* and *scarp2b*, carried by the same plasmid (pSci5) [17], were chosen for comparison.

In *S. citri* GII-3 wt culture the levels of *scarp 2b* and *5a* transcripts per µg of RNA were similar, respectively 10 ± 1 × 10^6 and 15 ± 1.25 × 10^6. In infected leafhoppers, 12 days after injection the number of spiroplasma cells was 6 × 10^6 per insect and the *scarp 2b* and *5a* transcript amounts were the same (35 ± 0.4 × 10^5 for 1 µg of total infected leafhopper RNA). Plants infected by *S. citri* through leafhopper transmission (LIP) developed symptoms within 3 weeks and spiroplasmas reached a titer of
$10^6 \text{ to } 10^7$ per g of fresh midribs. Scarp2b and scarp5a were equally transcribed to $10 \pm 1 \times 10^4$ transcripts per μg of total infected plant RNA. These results indicated that expression of scarp 2b and 5a were similar in the culture medium, the infected leafhoppers and in LIP (Fig. 4, protocol A). On the contrary, scarp2b transcript was not detected in GIP (Fig. 4, protocol B), whereas expression level of scarp5a was identical to that detected in infected LIP. PCR amplifications and sequencing the scarp2b and 5a coding sequences and the non-coding regions upstream of scarp genes revealed that in both LIP and GIP the sequences were 100% identical to those of GII-3 wt (data not shown), indicating that no sequence deletions had occurred in these regions.

There are no existing mechanisms that explain how a given gene such as scarp2b can be down regulated in old grafted plants. To further investigate the influence of a long period of multiplication in plants (GIP) on S. citri adaptation, spiroplasmas were isolated from GIP, subcultured for 10 passages, and injected into leafhoppers (Fig. 4, protocol B). The expression levels of scarp2b were compared in the different environmental conditions explained in Fig. 4 (protocol B vs A).

In the first passage of the S. citri culture obtained from GIP, the scarp2b mRNA transcript was not detected. However after further passaging in the SP4 culture medium, the scarp2b transcript was detected and reached a $10^7$ transcripts/μg of RNA at the 10th passage. Interestingly, in the leafhoppers injected with this 10 passages culture as well as in those injected with the early passage culture from GIP (which we named G-GIP1), expression level of scarp2b was similar to that obtained with leafhoppers injected with S. citri GII-3 wt. Unexpectedly, in periwinkle plants infected by these insects, expression of scarp2b was not detected whereas that of scarp5a was equivalent to that obtained in plants infected by GII-3 wt. Furthermore, in a periwinkle plant graft inoculated with a shoot coming from the newly infected plant (protocol B), scarp2b transcript was still undetectable. The fact that the scarp2b transcript was detected in leafhopper-infected periwinkles (LIP) (Fig. 4, protocol A) but not in plants infected or graft-inoculated by the S. citri originally isolated from GIP (Fig. 4, all symptomatic plants obtained through protocol B) strongly suggested a difference between the GIP-isolated spiroplasmas and S. citri GII-3 wt (used in protocol A). Therefore we investigated whether these results could be explained by a phenotypic (on/off expression of scarp2b) heterogeneity in the bacterial population present in GIP.
**Spiroplasma mixture in grafted periwinkles**

To challenge the hypothesis that GIP may contain a mixed bacterial population, spiroplasmas from GIP were directly plated on solid SP4 medium. After 10 days, spiroplasmas from 18 colonies were cultivated in liquid medium (only one passage) and submitted to qRT-PCR for detecting RNA transcription of *scarp2b*. Out of 18 spiroplasma cultures tested, 5 expressed the *scarp2b* RNA at levels similar to those obtained for GII-3 wt (10⁷/μg RNA) while the other 13 did not. Given that *scarp2b* transcripts could not be detected in GIP, such a high proportion of *scarp2b*-expressing colonies was unexpected. Following the hypothesis of a mixed population in GIP, *scarp2b*-expressing spiroplasmas could overtake the population that does not express *scarp2b* in the SP4

**Fig. 4** Schematic representation of the protocols used for *scarp* expression studies in the different hosts and expression of scarps under these conditions. Positive and negative detection (see Methods for details) of *scarp2b* and *scarp5a* transcripts in the different hosts are noted ‘−’ or ‘+’, respectively. Protocol A (left column): A culture of *S. citri* in axenic medium was injected in insects, which then were fed on young periwinkles that became symptomatic within 3 weeks (leafhopper-infected plants LIP). *S. citri* extracted from LIP were then cultivated in axenic medium. After one passage, *scarp2b* transcripts were undetectable. After 10 passages, *scarp2b* transcripts could be detected, and spiroplasmas were microinjected to insects. The insects were fed on young periwinkles. Symptomatic periwinkles were used to graft a new batch of periwinkle plants, which developed symptoms (grafted plants second generation).

Protocol A

| Transcripts (presence/absence) | Protocol B |
|-------------------------------|------------|
| *scarp2b* | Grafted Plant (GIP) |
| *scarp5a* | Spiroplasma isolation |

Protocol B

| *S. citri* GII-3 wt culture | S. citri culture (1st passage) (G-GIP1) |
|----------------------------|-------------------------------------|
| + | + |

| Insects | S. citri culture (10th passage) (G-GIP10) |
|---------|----------------------------------------|
| + | + |

| Transmission | Transmission |
|-------------|-------------|

| Symptomatic Plant (LIP) | Symptomatic Plant |
|------------------------|-------------------|
| + | + |

| Spiroplasma isolation | Graft transmission |
|-----------------------|--------------------|

| *S. citri* culture | Symptomatic grafted plant 2nd generation |
|-------------------|------------------------------------------|
| + | + |

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culture medium. Following this assumption the proportion of \textit{scarp2b}-expressing colonies would provide an overestimation of \textit{scarp2b}-expressing cells in GIP. This assumption is consistent with the finding that the \textit{scarp2b} transcript level continuously increased during passaging in the culture medium.

One spiroplasma clone that expressed \textit{scarp2b} and one that did not were separately injected into leafhoppers to investigate transmission to periwinkle plants. Ten days after transmission, all plants developed symptoms. In these plants, however, the \textit{scarp2b} transcript was detected or not according to the initial spiroplasma inoculum. Thus, the phenotypic differences, in particular regarding \textit{scarp2b} expression, between the \textit{S. citri} clones are stable in plants and do not modify the pathogenicity to plants. These results confirmed the hypothesis that two phenotypes of \textit{S. citri} co-existed in the original GIP. In these plants, the number of spiroplasma cells expressing \textit{scarp2b} may fall below a detectable level whereas spiroplasmas lacking \textit{scarp2b} expression efficiently multiplied. Other environments such as SP4 medium and leafhoppers probably constitute a better environment for propagation of \textit{scarp2b} expressing spiroplasmas. The nature of the environmental selective pressure encountered by spiroplasmas differs in plants and in insects. This change may be responsible for the differential multiplication of the two phenotypic variants depending on the host. Taken together, our data suggest that this \textit{scarp} gene may not be essential in plants and in insects (the \textit{scarp2b} non-expressing strain was transmissible), and argue in favor of the functional redundancy of \textit{scarp} genes.

Protein extracts from \textit{S. citri} GII-3 wt and \textit{S. citri} G-GIP1, which did not express \textit{scarp2b}, were separated by 2D-electrophoresis to determine whether lack of \textit{scarp2b} expression was associated with major changes in soluble protein expression profiles (Fig. 5). Two-dimensional electrophoresis patterns of the two strains were very similar for relative intensities of the protein spots, suggesting that these strains did not strikingly differ in their soluble protein expression profiles. However, one spot was clearly present in GII-3 wt strain and absent in G-Scarp-2b". Due to the lack of their expression in spiroplasmas well-adapted to plants, the proteins present in this spot were considered as good candidates for being involved in adaptation of \textit{S. citri} to the insect. \textit{LC-MS/MS} analysis of trypsinized peptides from the gel spot identified that 24 peptide fragments were derived from protein SPICI03_175, the alpha subunit of pyruvate dehydrogenase E1 subunit (Sequest score, 583; sequence coverage, 67 %), and that 16 peptide fragments were derived from protein SPICI01B_002, the beta chain of DNA polymerase III (Sequest score, 99; sequence coverage, 46 %). In \textit{M. pneumoniae}, pyruvate dehydrogenase E1 beta component can be surface translocated and binds host fibronectin [44]. Thus in this mycoplasma, pyruvate dehydrogenase E1 component functions in adherence in addition to its biosynthetic activity. Other cases of multifunctional proteins, such as the ABC transporter subunit OppA in \textit{M. hominis} [38] or the phosphoglycerate kinase in \textit{S. citri} [45, 46], involved in both cell metabolism and adhesion have been described in mollicutes. In \textit{S. citri} GII-3 wt, a role of dehydrogenase E1 subunit in

![Fig. 5 Bi-dimensional gel electrophoresis of total extract proteins of \textit{S. citri} GII-3 wt and \textit{S. citri} strain deficient in \textit{scarp2b} expression \textit{S. citri} G-GIP1. Circled spot was further analyzed by LC-MS/MS (see text for details)]
adhesion of spiroplasma to insect cells in addition to its role in pyruvate metabolism cannot be ruled out. In this case however, the presence of this subunit would not be essential for efficient transmission of *S. citri* by its vector.

Conclusions

Host specialization by bacterial pathogens requires a repertoire of virulence factors and dynamic regulation of gene expression. The present study provided us with a snapshot of the spiroplasma's response to its hosts and offered the opportunity to identify protein candidates required for maintenance and/or virulence in the different hosts for further functional studies. Consistently with the idea that pathogenic bacteria adapt to various host environments by varying synthesis of surface components, several *S. citri* lipoprotein genes were shown to be regulated in *C. haematocceps* and in periwinkles. Spiroplasma ability to regulate gene expression in both hosts is probably at least partially responsible for its capacity to multiply in plant as well as in its leafhopper vector and for its remarkable abilities to survive in a wide range of leafhoppers and plants [1]. In addition, most lipoprotein genes tested were up-regulated in insects compared to plants. It seems that insects are more favorable than plants for the lipoprotein gene expression of *S. citri*. This implies that these lipoprotein genes could be involved in adhesion and/or in invasion of insect cells during the transmission process. In addition to lipoprotein genes, gene encoding the E1 component of pyruvate dehydrogenase represents a good candidate for being involved in spiroplasma adaptation to its vector.

Finally, despite the small size of the *S. citri* chromosome, the regulation at transcription gene level in *S. citri* likely plays a significant role in its adaptive capacities to its hosts and could constitute an efficient mean for shaping the spiroplasma surface in response to the environmental conditions. Transcriptional regulation upon interaction with the host environment seems to be more developed in *S. citri* than in *M. pneumoniae* [39], *M. gallisepticum* [42] or in *M. hyopneumoniae* [43]. It has been suggested that the close adaptation to specific mucosal environments, such as the human lung epithelium for *M. pneumoniae*, was associated to restricted regulating capacities at the gene level [47]. Following this assumption, the important transcriptional regulating capacities in *S. citri* compared to these three mycoplasmas may be associated to the versatile environment (two distinct hosts) encountered by the spiroplasma.

Availability of data and materials

The data sets supporting the results of this article are included within the article and its additional file.
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