The plant-pathogenic mollicute *Spiroplasma citri* inhabits the phloem sieve tubes of infected plants. It is transmitted naturally in a propagative manner from plant to plant by sap-feeding leafhopper vectors: two *Circulifer* spp. (23,35,37) and three *Scaphytopius* spp. (23,35,37). *S. citri* has been cultured since 1970 from sweet orange trees affected with stubborn disease (43). It infects many plant species other than citrus, including Madagascar periwinkle (*Catharanthus roseus*), in which it induces stunting, leaf yellow, leaf curling, and wilting (9).

The circulative route of *S. citri* through an experimental leafhopper vector, *Euuscelis plebejus* (33), and a natural vector, *Circulifer tenellus* (29), is well established. In order to be transmitted to plants, two physical barriers within the leafhopper, the gut and salivary glands walls, must be crossed by *S. citri*. During insect feeding on infected plants, spiroplasmas move from the food canal through the foregut and accumulate in the midgut and hindgut. Then, they move across the basal lamina, circulate in the hemocoel where they multiply, enter the salivary glands, and are released in the salivary canal. Once they are present in the salivary canal, they can flow with salivary secretions through the salvia duct into the host plant during feeding. Electron microscopic observations of *S. citri* in membrane-bound cytoplasmic vesicles at the basal plasmalemma of gut epithelial cells suggest that *S. citri* invades the gut and also the salivary glands of its vector *C. tenellus* by endocytosis (15,25). Recent findings concerning the movement of *S. kunkellii* through the gut epithelial cell layer of its insect vector *Dalbulus maidis* are in agreement with this process (38). Earlier studies reporting that spiroplasmas traverse the epithelial cell layer of gut between cells in a process called diacytosis (1,28) were not confirmed by these results.

In the case of human and animal mycoplasmas, successful colonization of the host cells required adhesion as the first step. This event is mediated by mycoplasma surface proteins and, among these proteins, adhesins play an important role in invasion and pathogenicity (4,41).

Little is known about the biology of recognition and invasion of insect gut and salivary glands by *S. citri*. The passage of *S. citri* through both physical barriers probably is mediated by receptor-ligand interactions (25), although no receptors on the surface of insect’s organs were identified. However, in *S. citri*, one adhesion-related protein, P89, was found to be directly involved in the spiroplasma–insect cell interaction (56).

During the last two decades, much attention has been focused on the possibility that cellular recognition is mediated by carbohydrates and lectins. Because the basal lamina of insect organs is highly glycosylated (2,14), it is reasonable to postulate that carbohydrate moieties might be likely targets for *S. citri* interaction.

To further investigate the nature of the spiroplasma–insect cell interaction and to characterize the nature of the specific molecules involved in this interaction, we developed a protein-blotting protein-overlay assay (Far-western). Our results suggest that *S. citri* undergoes a specific interaction with *C. haematoceps* glycoproteins during insect invasion. Various experimental approaches were used to identify *S. citri* proteins that bind to insect carbohydrates. One *S. citri* protein, the spiralin, identified as a lectin, binds to two insect glycoproteins with apparent masses of 50 and 60 kDa.
MATERIALS AND METHODS

Preparation of *S. citri* proteins for sodium dodecyl sulfate-polyacrylamide gel electrophoresis and protein overlay assay. *S. citri* GII-3, originally isolated from its leafhopper vector *C. haematoceps* captured in Morocco (51), was cultivated in SP4 medium (48) at 32°C. *S. citri* cells from exponential growth phase were harvested by centrifugation at 20,000 × g for 20 min at 4°C and washed three times in washing buffer (HEPES, 8 mM; sucrose, 280 mM; pH 7.4). Protein concentration was determined by the Bradford procedure (8) using the a protein assay kit (Bio-Rad Laboratories, Hercules, CA) with ovalbumin as standard. For overlay assays, cells from a 50-ml culture were re-suspended in 200 µl of phosphate-buffered saline (PBS), pH 7.4 (2 mM KH₂PO₄, 8 mM Na₂HPO₄, 0.14 M NaCl, 2 mM KCl) containing EDTA-free protease inhibitors cocktail (one tablet per 10 ml) (Roche, Meylan Cedex, France). Then, cells were lysed by submitting them to five rounds of freezing at −20°C and thawing on ice. Protein concentrations were usually near 5 mg/ml. Aliquots of the suspension were frozen at −20°C until used.

Membrane proteins of *S. citri* were extracted with Triton X-114 according to a previously described method (7). *S. citri* polyclonal and monoclonal antibodies. Polyclonal serum was from a rabbit that had been immunized with *S. citri* strain GII-3. Immunoglobulins were purified as described before (44). Hybridomas producing monoclonal antibodies were obtained as described by Foissac et al. (17). Monoclonal antibody 12G9 reacted with the spiralin of different *S. citri* strains (17), including strain GII-3 (unpublished data).

**Sodium dodecyl sulfate polyacrylamide gel electrophoresis and blotting.** *S. citri* membrane proteins in sample buffer (26) were loaded (50 µg/well) on a 12.5% polyacrylamide gel and subjected to electrophoresis as previously described (34). After electrophoresis, proteins were transferred to nitrocellulose membrane for 60 min at 10 V (47) using a semi-dry transfer apparatus (Bio-Rad). For renaturation of proteins and blocking the unoccupied spaces, membrane was incubated overnight at 4°C in PBS, pH 7.4, containing 6% nonfat dry milk (Sigma-Aldrich, St. Louis). After blocking, membrane was used immediately for Western or negative Far-western experiments. A healthy colony of *C. haematoceps* was reared in an insect-proof cage at 30°C on stock plants as described previously (16). Adult leafhoppers were ground in a potter-Elvehjem-grinder (VWR International, Fontenay-sous-Bois Cedex, France) with sample buffer (26) (40 insects per 500 µl). Then the mixture was boiled for 15 min and centrifuged for 1 min at 200 × g. Proteins were not further purified to avoid inadvertently removing proteins of interest. Aliquots of supernatant (25 µl) were subjected to sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) (12.5%), and proteins were visualized by staining with Coomassie brilliant blue. For a preparative SDS-PAGE (12.5%), the suspension (500 µl) was applied to a 14-cm-wide slot under denaturing conditions as described by Laemmli (26). After electrophoresis, proteins were transferred to nitrocellulose membranes as above. The saturated membranes were used immediately for overlay assay with *S. citri* proteins.

**Leafhopper two-dimensional PAGE.** Forty *C. haematoceps* adults were ground in liquid nitrogen and the powder was dispersed in ice-cold acetone containing 10% trichloroacetic acid (TCA) and 0.07% β-mercaptoethanol. After 1 h at −20°C, the suspension was centrifuged at 14,000 × g for 10 min and the pellet was washed twice in the same solution without TCA. After the last centrifugation, the pellet was resuspended in 20 µl of rehydration buffer (Bio-Rad). Approximately 1 mg of protein was obtained.

Imobilized pH gradient strips (IPGs), pH 3 to 10 (Bio-Rad), 17 cm long, were incubated for 13 h in rehydration buffer (300 µl) containing 200 µg of insect proteins. Isoelectric focusing (IEF) was carried out using a Protean IEF cell (Bio-Rad). After IEF, the IPGs were incubated in two equilibration buffers for 10 min each according to the manufacturer’s instructions.

The proteins in the equilibrated IPGs were separated on 12.5% SDS-PAGE using a Protean xi cell. Electrophoresis was carried out at 50 V at room temperature until the dye front ran off the gel. Proteins were visualized by Coomassie brilliant blue or transferred onto nitrocellulose membrane and membrane was blocked as described above. After blocking, membrane was used immediately for Far-western experiments.

**Far-western experiments.** The Far-western protocol was used as described (21,49) with some modifications. After blocking, blots of total insect proteins or insect glycoproteins separated by SDS-PAGE were cut into 7-mm strips. Each strip was incubated with shaking (60 oscillations/min) overnight at 4°C in 2 ml of PBS buffer with 0.1% of Tween 20 (PBS-Tween) containing 10 µg of whole cell lysate of *S. citri* proteins. A blot with proteins separated by two-dimensional (2D) electrophoresis was incubated in the same conditions, in 20 ml of PBS-Tween containing 100 µg of *S. citri* proteins. Control experiments were conducted omitting spiroplasma proteins. After three 15-min washings in PBS-Tween, blots were incubated for 1 h at room temperature in PBS buffer with purified polyclonal immunoglobulin Gs (IgGs) against total *S. citri* proteins at a final concentration of 5 µg/ml. After three washings, blots were incubated at room temperature for 1 h with peroxidase-conjugated goat anti-rabbit IgGs (Sigma-Aldrich). Blots were rinsed three times in PBS-Tween and antigen-antibody complexes were detected with SuperSignal West Pico chemiluminescent substrate according to the manufacturer (Pierce, Rockford, IL). Then, blots were exposed for ≈10 min onto X-ray film. For sugar competition experiments, proteins of *S. citri* in PBS solution with different amounts (0.1, 0.5, or 1 M) of D(+)-mannose, methyl-α-mannopyranoside, or D(+)-glucose were added to insect protein blots.

**Identification of *S. citri* proteins involved in interaction with insect proteins.** Negative Far-western experiments were carried out to identify the proteins of *S. citri* implicated in the interaction with insect proteins. Insect protein overlay assays were conducted in a manner similar to that described above for the *S. citri* protein overlay assay.

Blots of spiroplasma membrane proteins, after blocking, were incubated with shaking, overnight at 4°C with a clarified insect suspension (two insects per milliliter of PBS) at 2 ml/strip. Immunodetection of *S. citri* proteins was performed with polyclonal IgGs (5 µg/ml) or monoclonal antibodies (5 µg/ml) against spiralin in the same conditions described above for the Far-western. The proteins of *S. citri* implicated in the interaction with insect proteins were masked and not recognized by antibodies. Detection of the antibody-antigen complexes were performed as above. Sugar competition experiments were carried out with 2 ml of PBS containing proteins from five insects and methyl-α-mannopyranoside (0.1, 0.2, 0.5, or 1 M). Control blots with no insect protein overlay were directly incubated in *S. citri* polyclonal IgGs or monoclonal antibodies against the spiralin.

**Glycoproteins detection.** Saturated blots of *S. citri* and insect proteins were incubated overnight in PBS containing peroxidase-labeled Concanavalin A at 50 µg/ml. After washing three times in PBS-0.05% Tween 20, the interaction was visualized with SuperSignal West Pico chemiluminescent substrate as above.

**Purification of insect glycoproteins by lectin affinity chromatography.** The HiTrap Lectin Test Kit (Amersham Biosciences Europe, Orsay Cedex, France), consisting of four glycoprotein binding columns (HiTrap concanavalin A [Con A], HiTrap Lens culinaris (lentil) agglutinin [LCA], HiTrap wheat germ agglutinin [WGA], HiTrap peanut agglutinin [PNA]), was used for purification of insect glycoproteins. The lectins used, their abbreviations, their major sugar specificities, and the elution buffers are listed in Table 1.
Four hundred healthy C. haematoceps adults were ground in liquid nitrogen and the powder was resuspended in 1 ml of solubilization buffer (SB) (2% Chaps in PBS, pH 7.0). The insoluble materials were removed by centrifugation at 4,000 × g at room temperature for 2 min. Aliquots of the supernatant (200 µl) were diluted with 800 µl of binding buffer specific for the lectin column used, as recommended by the manufacturer. Total volume (1 ml) was directly loaded on lectin columns for glycoproteins purification.

After elution with recommended elution buffers, glycoproteins were precipitated in ice-cold acetone, centrifuged at 20,000 × g for 30 min at 4°C, air dried, and resuspended in PBS buffer. Protein concentrations were estimated by the Bradford method. Puriﬁed glycoproteins were separated on SDS-PAGE (10 µg/well) and silver stained. For electroblotting onto nitrocellulose membrane, the amount of glycoproteins was 50 µg/well. Before use in Far-western experiments, the membrane was blocked as above.

**Lectins detection in S. citri.** Blots of S. citri proteins blocked with 6% dry milk, as above, were incubated for 2 h at room temperature in 2 ml of PBS containing different amounts of ovalbumin (5, 10, and 15 µg). The control was made by omitting ovalbumin. After washing, blots were incubated with peroxidase-labeled Con A (50 µg/ml) in PBS buffer for 1 h. Then, the blots were rinsed in PBS-Tween and detection of the lectin-glycoprotein complex was performed with SuperSignal West Pico chemiluminescent substrate as described above.

**Spiralin purification.** A pellet from 1 liter of S. citri culture was washed and resuspended in 5 ml of Hapes-sucrose buffer. Spiroplasma cells were disrupted by sonication (rate: 50% pulses/s, 50 watts, 0°C, 1 min of sonication and 2 min on ice alternatively, five times) (Vibracell Sonicator; sonic & Materials, Inc., Danbury, CT). The mixture was clarified by centrifugation at 20,000 × g for 30 min at 4°C. The pellet was suspended in 5 ml of 0.1 M citrate buffer, pH 5.6, with 10 mM of phenylmethylsulfonyl fluoride (PMSF). The suspension first was fractionated on an anionic column (Mono S HR 5/5; Amersham Biosciences). The sample composed of the proteins which were not absorbed on the anionic column (pI > 5.6) was diluted with phosphate buffer, pH 7.5, and subjected to separation on a cationic column (Mono Q HR 5/5) because of the spiralin pI, which is 8.3. Proteins absorbed on the column (pI > 7.5) were eluted with a 20-ml NaCl linear gradient from 0 to 1 M in phosphate buffer (pH 7.5). The eluted proteins were monitored by absorbance at 280 nm and some fractions were analyzed on SDS-PAGE. Western blot analyses were performed on selected fractions containing a protein of 24 kDa, using monoclonal antibodies against spiralin.

**RESULTS**

**Affinity of S. citri proteins for leafhopper proteins.** Insect proteins separated by SDS-PAGE in one dimension were stained with Coomassie brilliant blue (Fig. 1A) or blotted onto nitrocellulose and probed with S. citri proteins (5, 10, or 15 µg). Seven significant bindings were revealed by polyclonal IgGs against total S. citri proteins (Fig. 1B, lanes 2, 3, and 4). In a control experiment in which S. citri proteins were omitted from the overlay assay, no reactions with polyclonal IgGs were found (Fig. 1B, lane 1). To ascertain the binding of spiroplasma proteins to one immobilized leafhopper protein, the gel overlay assay was carried out on protein blots issued from 2D-SDS-PAGE (Fig. 2A). Spiroplasma proteins were found to bind to a set of insect proteins having apparent molecular masses of 60, 50, 35, 30, 25, 18, and 16 kDa (Fig. 2B). In order to determine whether the binding of S. citri proteins to insect proteins occurred in a lectin-mediated manner, Far-western experiments were performed in the presence of increasing amounts of D(+)-mannose, methyl-α-mannopyranoside, and D(+) glucose as competitors (Fig. 3A to C). In all six replications of this assay, the presence of sugars interfered in the binding of S. citri proteins with insect proteins. A difference in the intensity of the signals was observed near insect proteins of 35, 30, 25, 18, and 16 kDa. Furthermore, in the presence of 0.1 M of different sugars (Fig. 3A to C, lane 1), faint signals (reduced bindings) were detected near insect proteins of 50 and 60 kDa. With increasing amounts of sugar (0.5 and 1 M), these bindings were completely inhibited (Fig. 3A to C, lanes 2 and 3). These sugarsensitive bindings suggested that the interactions between S. citri and insect proteins of 50 and 60 kDa were lectin-glycoprotein.

**Fig. 1.** Far-western: binding of Spiroplasma citri proteins to Circulifer haematoceps proteins separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). A. Whole-body proteins of one leafhopper were analyzed by SDS-PAGE (12.5%), stained with Coomassie brilliant blue. B. Proteins of 40 insects (800 µg) were separated on a preparative polyacrylamide gel and electrophoretically transferred onto nitrocellulose membrane. Saturated membrane was cut into 20 strips (width, 7 mm). The strips were probed with different amounts of whole S. citri cell lysate proteins (lane 1, 0 µg; lane 2, 5 µg; lane 3, 10 µg; lane 4, 15 µg). Anti-S. citri polyclonal immunoglobulin Gs (IgGs) were used to detect bound spiroplasma proteins. Peroxidase-conjugated goat anti-rabbit IgGs were used as secondary antibodies and detection was performed with chemiluminescent substrate. Arrows on the right indicate the seven significant bindings between proteins of S. citri and insect proteins having apparent molecular masses of 60, 50, 35, 30, 25, 18, and 16 kDa. M = molecular mass marker; relative molecular masses are indicated on the left.

<table>
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<tr>
<th>Lectins</th>
<th>Specificity</th>
<th>Elution buffer</th>
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<tbody>
<tr>
<td>Concanavalin A (Con A)</td>
<td>Branched mannose, carbohydrates with terminal mannose or glucose (αMan&gt;αGlc&gt;αGlcNAc)</td>
<td>1 M methyl-α-mannopyranoside, 20 mM Tris-HCl, pH 7.4</td>
</tr>
<tr>
<td>Lens culinaris agglutinin (LCA)</td>
<td>Branched mannose with fucose linked α(1,6) to the N-acetylglucosamine, (αMan&gt;αGlc&gt;αGlcNAc) Core of N-linked oligosaccharides, [GlcNacβ1,4GlcNAcβ1,3GalNAc&gt;βGlcNAc]</td>
<td>1 M methyl-α-mannopyranoside, 20 mM Tris-HCl, pH 7.4</td>
</tr>
<tr>
<td>Wheat germ agglutinin (WGA)</td>
<td>Terminal β-galactose, (Galβ1,3GalNAc&gt;α and βGal)</td>
<td>0.5 M N-acetylglucosamine, 20 mM Tris-HCl, 0.5 M NaCl, pH 7.4</td>
</tr>
<tr>
<td>Peanut agglutinin (PNA)</td>
<td>0.5 M NaCl, pH 7.4</td>
<td>1 M D-galactose, 10 mM sodium phosphate, 0.15 M NaCl, 0.5 mM MgCl₂, 0.5 mM CaCl₂, pH 7.4</td>
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Detection and purification of glycoproteins. The classical methods used for detection of glycoproteins directly in gels, such as selective carbohydrate silver staining, periodic acid Schiff staining, and lectin (Con A) blotting, have not allowed the detection of glycoproteins in S. citri. Glycoproteins were detected by these classical methods only in insects (data not shown).

Purification of insect glycoproteins were performed on four lectin columns. With WGA, Con A, and LCA columns, glycoproteins were in the elution fractions. With the PNA column, no glycoproteins were obtained in the elution buffer. Using the same columns, glycoproteins also were not obtained from S. citri. These results are consistent with those obtained with the classical methods.

Affinity of S. citri proteins for insect glycoproteins. To determine whether spiroplasma proteins display affinity for insect glycoproteins, Far-Western assays were performed using whole S. citri cell lysate proteins as the overlay (10 µg). With insect glycoproteins purified from WGA, Con A, and LCA columns (Fig. 4A to C, lane 1), two signals were obtained only for glycoproteins of 50 and 60 kDa (Fig. 4A to C, lane 2). These interactions between spiroplasma proteins and insect glycoproteins explain the lack of binding observed with sugars in the same molecular mass region (Fig. 3A to C).

Negative Far-western and competition with methyl-α-mannopyranoside. To identify the proteins of S. citri involved in the interaction with the insect proteins, Far-Western experiments were carried out on spiroplasma membrane protein blots with insect proteins as the overlay. Membrane proteins of S. citri involved in bindings were masked and not detected by S. citri antibodies. Antibodies against whole S. citri cells recognized many proteins among the membrane proteins of S. citri (Fig. 5A, lane 1). Previous results have shown that the protein which is the major antigen recognized by the polyclonal antibodies is the 24-kDa spiralin (55). The comparison between the pattern obtained in the protein overlay assay revealed by the polyclonal IgGs (Fig. 5A, lane 2) and Western blot (Fig. 5A, lane 1) showed that one or several proteins near 24 kDa were masked by insect proteins and were not detected by the antibodies. These masked proteins appeared when insect proteins with increasing amounts of methyl-α-mannopyranoside sugar were used as the overlay (Fig. 5A, lanes 3, 4, 5, and 6). Similar experiments performed with the monoclonal antibodies against spiralin demonstrated that one of these proteins that bound to insect proteins was spiralin (Fig. 5B, lanes 3, 4, 5, and 6). The sugar-sensitive interaction observed between spiralin and insect proteins suggested that the spiralin might be a lectin.

Lectin detection in S. citri. A blot of S. citri proteins was incubated in PBS buffer, pH 7.4, containing ovalbumin as a glycoprotein, followed by binding to peroxidase-labelled Con A. With increasing amount of ovalbumin, one protein of 24 kDa appeared as the only lectin in the S. citri proteins pattern (Fig. 6, lanes 2, 3, and 4). No protein was detected when experiment was carried out without ovalbumin (Fig. 6, lane 1).

Affinity between spiralin and insect glycoproteins. Aliquots of eluted fractions from the cationic column were analyzed on SDS-PAGE (Fig. 7A, lanes 2, 3, 4, 5, and 6) and one aliquot was stained with Coomassie brilliant blue or Schiff staining, and lectin (Con A) blotting, have not allowed the detection of glycoproteins in S. citri.
analyzed in a Western blot with monoclonal antibodies against spiralin. A single band was observed in lane 6 (Fig. 7A) corresponding to a protein of 24 kDa. This protein was identified as spiralin by monoclonal antibodies (Fig. 7B, lane 2). Blobs of purified insect glycoproteins (Fig. 7C, lane 1) were overlaid with 10 µg of S. citri whole-cell proteins (lane 2) or with 1 µg of spiralin (lane 3) and probed with monoclonal antibodies against spiralin. Significant bindings between the two insect glycoproteins of 50 and 60 kDa and the purified spiralin were observed (Fig. 7C, lane 3). Same bindings were also observed in lane 2. The spiralin did not bind any glycoproteins from insect vector except those of 50 and 60 kDa.

**DISCUSSION**

As a prelude to understanding the molecular mechanisms underlying *S. citri* transmission, we have used a protein overlay assay (Far-western) to investigate binding of spiroplasma to proteins of its vector *C. haematoceps*. Far-western assays commonly are used in plant virus research to investigate the relationship of the viruses with their vectors (5,45,50). Similar assays were used to study human and animal mycoplasma interactions with host eukaryotic cells (24,27,57,58). Glycolipid host receptors for *Mycoplasma hyopneumoniae* have been identified (58) by using a thin-layer chromatography overlay assay.

In our work, using a Far-western assay, *S. citri* proteins bound to seven insect proteins. Because of the sugar-dependent nature of the observed interactions, suggesting lectin-glycoprotein bindings, we have looked for glycoproteins and lectins in both partners.

The nondetection of glycoproteins in total *S. citri* proteins was in good agreement with previous reports, in which the authors failed to detect glycoproteins on *S. citri* protein blots after staining of polyacrylamide gel with periodic acid Schiff reagent (22,55). However, binding of some lectins to insect proteins was observed before (22) and a protein that specifically binds Con A was isolated from the membrane of *S. citri* (46). Unfortunately, the preparation was only partially pure, being contaminated by a small amount of another protein of the membrane and also by an exogenous polypeptide tentatively identified as a Concanavalin A subunit (53).

With the different techniques used in this study, glycoproteins were detected only in the insect. With the PNA lectin column, which has an affinity for N-acetyl galactosamine, the common core structure of O-glycans, we did not detect any glycoproteins.

![Fig. 4. Binding experiments between purified insect glycoproteins and whole *Spiroplasm citri* cell lysate proteins. Glycoproteins purified from *A*, wheat germ agglutinin, *B*, concanavalin A, and *C*, *Lens culinaris* agglutinin columns were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and silver stained (strip 1: 10 µg/well) or electrotransferred (strip 2: 50 µg/well). After blocking, all strips were incubated with 10 µg of whole *S. citri* cell lysate proteins. Bindings between *S. citri* proteins and insect glycoproteins were detected as in Figure 1. Relative molecular mass markers are indicated on the left.](image)

![Fig. 5. Negative Far-western experiments: binding of insect proteins to *Spiroplasm citri* membrane proteins separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). *S. citri* membrane proteins (Triton X-114 insoluble fraction) were loaded on SDS-PAGE (50 µg/well) and blotted onto nitrocellulose. Western blot analysis was carried out with *A*, polyclonal immunoglobulin Gs (IgGs) against *S. citri* proteins strip 1) or *B*, monoclonal antibodies against spiralin (strip 1) followed by immunological detection as in Figure 1. Other strips were incubated with insect proteins (40 µg) in the presence of increasing amounts of methyl-α-mannopyranoside (strip 2, no sugar; strip 3, 0.1 M; strip 4, 0.2 M; strip 5, 0.5 M; strip 6, 1 M). *A*, Polyclonal IgGs and *B*, monoclonal antibodies were used as the primary antibodies. Immunodetection procedure was performed as described in Figure 1. Arrows on the left of A1 and B1 indicate the molecular mass of spiralin. Note that spiroplasma proteins bound to insect proteins were masked and not recognized by polyclonal IgGs (*A*, strips 2, 3, and 4) or monoclonal antibodies (*B*, strips 2, 3, and 4).](image)

![Fig. 6. Lectin detection in *Spiroplasm citri*. Blobs of *S. citri* proteins were incubated with increasing amounts of ovalbumin as a glycoprotein (blot 1, 0 µg; blot 2, 5 µg; blot 3, 10 µg; blot 4, 15 µg) and probed with peroxidase-labeled concanavalin A (50 µg/ml). Detection of peroxidase was performed with chemiluminescent substrate. Arrow on the right side indicates the lectin detected. Arrow on left side of the *S. citri* proteins blot (W) probed with polyclonal immunoglobulin Gs indicates spiralin (spi).](image)
in *C. haematoceps*, probably due to the low amount of O-glycans in the insect. This is consistent with previous studies which have shown that insect cells have a reduced capability for O-glycosylation of foreign polypeptides (2,19,30). Numerous glycoproteins were purified from WGA, LCA, and Con A columns, suggesting that the glycoproteins of the whole insect are predominantly N-glycans with some possible O-glycans. From each purified glycoprotein suspension, only two glycoproteins of 50 and 60 kDa bound to *S. citri* proteins. According to the specificity of WGA, LCA, and Con A, these insect glycoproteins were N-glycoproteins that displayed a high content of mannose with fucose-linked α(1.6) to N-acetylgalactosamine.

These results indicate that bindings could arise if surface lectins expressed by the bacteria were able to interact with sugar groups on the insect glycoproteins, which could be receptors for spiroplasmas. It is well known that various forms of carbohydrates covering the eukaryotic cell surface are used by microbial pathogens as attachment sites. Host cell surface sialoglycoproteins and glycolipids have been reported to be receptors for the attachment of human and animal mycoplasmas (31,39,40,58). Studies on the adherence of *S. citri* to a monolayer of *C. tenellus*-cultured cells have demonstrated that carbohydrates have no effect on spiroplasma–insect attachment, suggesting that glycoconjugates are not involved in the *S. citri* adherence to insect cells (56). In our study, competition with carbohydrates reduced the binding of spiroplasma proteins to insect glycoproteins of 50 and 60 kDa. It is important to note that the concentration of sugar used (1 M) is higher than that used by Yu et al. (56). For clover phyllody phloemplasma, a gene encoding a putative O-sialoglycoprotein endopeptidase was found. As reported by the authors, this enzyme may act on host cell glycoproteins to enhance the attachment of the phloemplasma to insect vector gut tissues (12).

In our analysis, among purified insect glycoproteins, sialo- glycoproteins could be present, because WGA lectin selectively binds terminal N-acetylglucosamine and sialic acid residues (54). The ability of hemipteran insect cells to produce sialylated-N-glycans was not investigated. However, it is known that, in some insects, such as members of *Lepidoptera*, cells contain small amounts of sialic acid and sialylation of glycoproteins is a highly specialized function that probably occurs rarely (20,32). If the glycosylation pathways in leafhopper cells are similar to those of *Lepidoptera* spp., these findings strongly suggest that *S. citri* receptors in insect are N-glycosylated proteins with some possible sialoglycoproteins.

Negative Far-western experiments showed that a 24-kDa protein of *S. citri* binds to insect glycoproteins. Competition with increasing amounts of sugar allowed detection of this protein as a lectin. Using monoclonal antibodies against spiralin, we demonstrated that this protein is the spiralin. Because the search for lectin among all *S. citri* proteins revealed only one lectin of ≈24 kDa, we postulate that this unique lectin and the spiralin are the same protein. In addition, two insect glycoproteins of 50 and 60 kDa displayed strong affinity to purified spiralin. Spiralin is one of the most thoroughly characterized *S. citri* membrane proteins (6,11,17). A model in which spiralin could form a protein carpet covering the entire spiroplasma surface was proposed (10) and its requirement for efficient transmission of *S. citri* by its leafhopper was described (13).

Recently, ultrastructural studies on phytopathogenic spiroplasmas indicated that helical spiroplasma cells possess a tip structure at the end of the helices (3). Various proteins on the tip structure may participate in the spiroplasma orientation and adherence to insect cell surface proteins, as is the case for adhesion proteins found on the tip structure of *M. pneumoniae* (42). Spiralin covering the entire surface of the spiroplasma (10) could not play a role in vivo in the adherence mediated via the terminal tip structure. It could be involved in the interaction between the pleiomorphic forms of spiroplasma and the membranous vesicles within the cytoplasm of the leafhopper midgut or salivary glands epithelial cells observed by Waydande et al. (52).

When taken together, the results of the current study suggest that, in vitro, the interaction of *S. citri* with two leafhopper glycoproteins with apparent masses of 50 and 60 kDa is mediated in a lectin manner. The most likely *S. citri* candidate for this molecular interaction is the spiralin. These findings were based on our binding studies that might have excluded certain interactions present in vivo which are dependant on environmental aspects.

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**Fig. 7.** Far-western experiment: binding of purified spiralin to insect glycoproteins purified from wheat germ agglutinin (WGA) lectin column. A, Whole *Spiroplasm citri* cell lysate proteins (50 µg) (lane 1) and aliquots (10 µl) of five fractions eluted from the cationic column (lane 2 to 6) were analyzed on sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) (12.5%) and silver stained. B, Total *S. citri* proteins (lane 1) and aliquots of fraction 6 eluted from the cationic column were transferred onto nitrocellulose membrane. Saturated blots were probed with monoclonal antibodies against spiralin followed by immunological detection as described in Figure 1. C, Glycoproteins purified from WGA column were separated on SDS-PAGE (50 µg/well) silver stained (lane 1) or blotted onto membrane. Saturated blots were incubated with 10 µg of whole *S. citri* cell lysate proteins (lane 2) or with 1 µg of purified spiralin (lane 3). Then, blots were probed with monoclonal antibodies against spiralin. Peroxidase-conjugated goat anti-rabbit immunoglobulin Gs were used as secondary antibodies and detection was done with chemiluminescent substrate.

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LITERATURE CITED


