Complete Genome Sequences of Six Copper-Resistant *Xanthomonas citri* pv. *citri* Strains Causing Asiatic Citrus Canker, Obtained Using Long-Read Technology

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**ABSTRACT** The gammaproteobacterium *Xanthomonas citri* pv. *citri* causes Asiatic citrus canker. Pathotype A strains have a broad host range, which includes most commercial citrus species, and they cause important economic losses worldwide. Control often relies on frequent copper sprays. We present here the complete genomes of six *X. citri* pv. *citri* copper-resistant strains.

Asiatic citrus canker, caused by *Xanthomonas citri* pv. *citri*, causes important economic losses in most tropical and subtropical areas. Its development impacts crop yields and also limits exports because of the quarantine status of the pathogen in some countries. The pathogen is currently classified into three pathotypes. Whereas pathotypes A* and A have a restricted host range and geographical distribution, pathotype A has a wider host range, which has resulted in major geographical expansion of the pathogen (1). In the early stages of emergence, eradication is feasible through tree removal, burning, and quarantine restrictions, whereas integrated pest management (IPM) is conducted after the disease has spread in an area. IPM typically includes repetitive copper applications. Copper-resistant (Cu*) strains were first described in 1994 in Argentina (2), and 20 years later in the French islands of Réunion (3) and Martinique (4). To investigate the genetic basis of this resistance, six Cu* strains were sequenced using long-read PacBio RSII technology with one single-molecule real-time (SMRT) cell per strain. Three strains originated from Réunion (LH201, LH276, and LJ207-7), two from Argentina (LM199 and LM180/H11005 A44), and one from Martinique (LL074-4). Minisatellite typing revealed that all Cu* strains were assigned to genetic lineage 1 within pathotype A (1).

The *de novo* genome assembly was conducted using the SMRT Analysis HGAP version 2.3 protocol. Circularization of the contigs was attempted using a combination of the Minimus assembler (5) and the SMRT Analysis resequencing version 1 protocol. The protein-coding sequences of the plasmid that confers Cu* to LH201 were predicted using the MaGe genome annotation platform. In order to improve the assembly of transcription activator-like genes, additional assembly steps were performed in a manner similar to that as previously described (6).

The chromosomes of the six strains were fully reconstructed, with sizes ranging from 5,148,274 to 5,198,476 bp. Depending on the strains, two to four additional contigs were circularized into plasmids, with sizes ranging from 28,949 to 249,697 bp. All six strains hosted a plasmid carrying Cu* genetic determinants. These plasmids were highly conserved among strains (despite remote geographical origins), apart from LM199.
The six closed genome sequences have been deposited in GenBank. The accession numbers are summarized in Table 1.

Whereas the other five strains hosted the copLAB gene system previously described for Xanthomonas citri pv. citri (7), LM199 encoded copABCD, which has already been described on the chromosome of X. arboricola pv. juglandis (8). Additional putative resistance gene clusters were found on the CuR plasmid: cusAB/smmD, czcABCD, and arsBHCR. However, LM199 lacked czcABCD and arsBHCR. MICs were evaluated as previously described (9): zinc chloride, 32 mg/L; copper sulfate, 128 to 256 mg/L; sodium arsenite, 8 to 16 mg/L (LM199), 64 mg/L (LH201), 128 mg/L (LL074-4), >128 mg/L (LH276, LM180, and LJ207-7); cadmium sulfate, 6.4 to 12.8 mg/L; and cobalt chloride 16 to 32 mg/L. Because strain LM199 (which did not contain the putative czcABCD region) and other sequenced strains (containing czcABCD) shared similar MIC values for cadmium sulfate, zinc chloride, and cobalt chloride, the function of this efflux pump remains unclear. These sequences highlight the potential of microbial adaptation through horizontal gene transfer and the mosaic structure of the genetic elements carrying adaptive genes.

Accession number(s). The six closed genome sequences have been deposited in GenBank. The genome accession numbers are summarized in Table 1.

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REFERENCES


**TABLE 1** Characteristics of six Xanthomonas citri pv. citri strains

<table>
<thead>
<tr>
<th>Strain</th>
<th>Other number(s)</th>
<th>Accession no.</th>
<th>Copper resistance location</th>
<th>Place of isolation</th>
<th>Yr of isolation</th>
<th>Host</th>
</tr>
</thead>
<tbody>
<tr>
<td>LM199</td>
<td>X. citri pv. citri 15-4632 (INTA)</td>
<td>MSQW000000000</td>
<td>Plasmid</td>
<td>Argentina</td>
<td>2015</td>
<td>Orange</td>
</tr>
<tr>
<td>LM180</td>
<td>X. citri pv. citri 03-1638 (INTA); A44</td>
<td>MSQW000000000</td>
<td>Plasmid</td>
<td>Argentina</td>
<td>2003</td>
<td>Grapefruit</td>
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<td>LL074-4</td>
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<td>CP018847 to CP018849</td>
<td>Plasmid</td>
<td>Martinique</td>
<td>2014</td>
<td>Grapefruit</td>
</tr>
<tr>
<td>LJ207-7</td>
<td></td>
<td>CP018850 to CP018853</td>
<td>Plasmid</td>
<td>Réunion</td>
<td>2012</td>
<td>Kaffir lime</td>
</tr>
<tr>
<td>LH276</td>
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<td>Réunion</td>
<td>2010</td>
<td>Kaffir lime</td>
</tr>
<tr>
<td>LH201</td>
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<td>CP018858 to CP018860</td>
<td>Plasmid</td>
<td>Réunion</td>
<td>2010</td>
<td>Kaffir lime</td>
</tr>
</tbody>
</table>

aINTA, Instituto Nacional de Tecnologia Agropecuaria.

bAs designated by Behlau et al. (7).