The *gpsX* gene encoding a glycosyltransferase is important for polysaccharide production and required for full virulence in *Xanthomonas citri* subsp. *citri*

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**Abstract**

**Background:** The Gram-negative bacterium *Xanthomonas citri* subsp. *citri* (*Xac*) causes citrus canker, one of the most destructive diseases of citrus worldwide. In our previous work, a transposon mutant of *Xac* strain 306 with an insertion in the XAC3110 locus was isolated in a screening that aimed at identifying genes related to biofilm formation. The XAC3110 locus was named as *bdp24* for biofilm-defective phenotype and the mutant was observed to be affected in extracellular polysaccharide (EPS) and lipopolysaccharide (LPS) biosynthesis and cell motility. In this study, we further characterized the *bdp24* (XAC3110) gene (designated as *gpsX*) using genetic complementation assays and expanded the knowledge about the function of the *gpsX* gene in *Xac* pathogenesis by investigating the roles of *gpsX* in EPS and LPS production, cell motility, biofilm formation on host leaves, stress tolerance, growth *in planta*, and host virulence of the citrus canker bacterium.

**Results:** The *gpsX* gene encodes a putative glycosyltransferase, which is highly conserved in the sequenced strains of *Xanthomonas*. Mutation of *gpsX* resulted in a significant reduction of the amount of EPS and loss of two LPS bands visualized on sodium dodecylsulphate–polyacrylamide gels. Biofilm assays revealed that the *gpsX* mutation affected biofilm formation by *Xac* on abiotic and biotic surfaces. The *gpsX* mutant showed delayed bacterial growth and caused reduced development of disease symptoms in susceptible citrus leaves. The *gpsX* mutant was more sensitive than the wild-type strain to various stresses, including the H\(_2\)O\(_2\) oxidative stress. The mutant also showed attenuated ability in cell motility but not in flagellar formation. Quantitative reverse transcription-PCR assays indicated that mutation of *gpsX* did not affect the expression of virulence genes such as *pthA* in *Xac* strain 306. The affected phenotypes of the *gpsX* mutant could be complemented to wild-type levels by the intact *gpsX* gene.

**Conclusions:** Taken together, our data confirm that the *gpsX* gene is involved in EPS and LPS synthesis and biofilm formation in *Xac* and suggest that the *gpsX* gene contributes to the adaptation of *Xac* to the host microenvironments at early stage of infection and thus is required for full virulence on host plants.

**Background**

Citrus canker, caused by the Gram-negative plant pathogenic bacterium *Xanthomonas citri* subsp. *citri* (*Xac*) (syn. *Xanthomonas axonopodis* pv. *citri*) [1,2], is one of the most important diseases of citrus crop worldwide [3]. Citrus canker is widely distributed in wet subtropical citrus growing areas and affects most commercial citrus varieties [3,4]. The canker symptom is characterized by raised necrotic lesions on leaves, stems and fruit of infected trees; and in severe cases, defoliation, twig die-back, general tree decline, blemished fruit and premature fruit drop can occur [3,4]. Wind-blown rain is the primary short- to medium-distance spread mechanism for citrus canker and long-distance dissemination is usually caused by transportation of infected citrus fruits and plant materials [5]. The decrease of yield and less value or entirely unmarketable of infected fruit are responsible...
for serious economic losses [3]. Moreover, this disease has a significant impact on commerce due to restrictions to national and international fruit trade from canker-affected areas [3]. Economic losses are also resulting from costly eradication programs and heave use of chemical treatments such as copper-based bactericides for prevention from and control of citrus canker disease [6].

Several members of the genus *Xanthomonas*, including *Xac*, have been extensively studied as model organisms to dissect the mechanisms of bacterial-plant interactions, including the molecular basis of pathogenesis and plant disease resistance [7-9]. *Xac* is considered to be a hemibiotrophic pathogen because it is able to obtain nutrients from living host cells, multiply in the apoplast (intercellular spaces) and then infect neighbouring tissues, after invading citrus host directly through natural openings, such as stomata, and through wounds [4]. The apoplast is a nutrient-limited environment that is guarded by plant defenses [10]. *Xac*, like many other plant pathogenic bacteria, has evolved several strategies to adapt to and successfully colonize this *in planta* niche by overcoming the plant defense and creating a favourable environment for bacterial growth, which include, among others, the type III secretion system (TTSS) and its effectors, cell wall degrading enzymes, and bacterial polysaccharides [8]. Bacterial polysaccharides of plant pathogenic bacteria, including extracellular polysaccharides (EPS), lipopolysaccharides (LPS) and capsular polysaccharides (CPS), have been shown to play a role in a number of different diseases. They collectively or individually contribute to the bacterial growth and survival *in planta*, and also are involved in the bacterium-plant interaction [8].

Progress has been made in elucidating the biosynthesis of bacterial polysaccharides over the decades [11]. The biosynthesis of bacterial polysaccharides occurs in successive steps. Firstly, nucleotide sugars are produced, which provide specifically activated monosaccharides as precursors for the subsequent synthesis steps. Secondly, monosaccharide moieties from the nucleotide sugar precursors are sequentially transferred by highly specific glycosyltransferases (GTs) to sugar or nonsugar acceptors, resulting in the formation of saccharide repeating units. Finally, the repeating units are polymerized and the polymer is exported from the cell. Bacterial GTs have been reported to be involved not only in the biosynthesis of EPS, LPS, CPS, peptidoglycans, and glycolipids, but also in protein and lipid glycosylation, showing enormous diversity of biological functions and substrates [12-14].

Much effort has been made to identify genes that encode GTs, their enzymatic functions, and the structures of these enzymes. Currently, there are more than 94 GT families in the Carbohydrate-Active EnZymes (CAZy) database (http://www.cazy.org) based on amino acid sequence similarities [15,16]. Two main three-dimensional folds, named GT-A and GT-B, have been observed for structures of nucleotide sugar-dependent GTs [12,13]. There is high sequence variability, although the relatively low structural variety and it is not yet possible to reliably predict the precise function of a given GT.

Mutations in GTs encoding genes have profound biological effects in a variety of bacteria. For example, mutation in *spxA* of *Bacillus subtilis* resulted in an altered spore coat [17]. In *Escherichia coli* strain 2787 (O126: H27), an *aah*-deletion mutant was attenuated in adherence to host cells [18], and the *pgaC* mutant of *E. coli* K-12 was impaired in surface binding, intercellular adhesion, and biofilm formation [19]. Mutation of *orfN* in *Pseudomonas aeruginosa* PAK affected the flagellin glycosylation [20]. In *X. campestris* pv. *campestris* strain 8004, mutation of *xagB* (*XC_3555*) led to decreased EPS production, abolished biofilm formation and attenuated bacterial resistance to oxidative stress [21], and the *XC_3814* mutant was significantly reduced both in EPS production and virulence on host plants [22]; while the *rfbC* mutation in *Xac* strain 306 resulted in altered O-antigen of LPS, reduced biofilm formation and attenuated bacterial resistance to environmental stresses [23].

In our previous work, an EZ-Tn5 transposon mutant of *Xac* strain 306 with an insertion in the *XAC3110* locus was isolated in a screening that aimed at identifying genes involved in biofilm formation. The *XAC3110* locus was named as *bdp24* for biofilm-defective phenotype and the mutant was observed to be affected in EPS and LPS biosynthesis, cell motility and biofilm formation on abiotic surfaces [24]. Due to the nature of our previous study in genome-wide identification of biofilm related genes, we focused on big picture rather than individual genes. It is necessary to further characterize the novel genes identified in our previous study and provide conclusive genetic evidence in complementation. In this study, we further characterized the *bdp24* (*XAC3110*) gene (renamed as *gpsX*) that encodes a putative glycosyltransferase using genetic complementation assays. The data obtained confirmed that the novel gene *gpsX* plays a role in EPS and LPS biosynthesis, cell motility, biofilm formation on abiotic surfaces and host leaves, stress tolerance, growth *in planta*, and host virulence of the citrus canker bacterium. These findings suggest that the *gpsX* gene contributes to the adaptation of *Xac* to the host microenvironments at early stage of infection and thus is required for full virulence on host plants.

**Results**

The *gpsX* gene encodes a glycosyltransferase involved in polysaccharide biosynthesis in *X. citri* subsp. *citri*

The *XAC3110* locus was identified as a biofilm formation-related gene of *bdp24* that may be involved in EPS and LPS biosynthesis, following screening a transposon
insertion mutant library of Xac strain 306 in our earlier work [24]. The XAC3110 open reading frame (ORF) is 2028 bp in length and located in the genome sequence at position 3655217-3657244 (Figure 1). XAC3110 consists of a single transcriptional unit, whereas the adjacent upstream and downstream genes were transcribed separately from this ORF in reverse orientation [25]. XAC3110 was annotated as a 675 aa glycosyltransferase [7]. The predicted pI and molecular weight (MW) of the putative enzyme are 6.67 and 73.9 kD (http://web.expasy.org/compute_pi/), respectively. The predicted protein contained a glycosyltransferase family 2 domain (PF00535, 2.00e-28) (residues 50-216) at the N-terminal and a UDP-Glycosyltransferase/glycogen phosphorylase superfamily domain (SCOP:d1f6da_, 2.00e-12) (residues 340-660) at the C-terminal (Figure 1). In addition, PSI-BLAST analysis revealed that the XAC3110 belongs to glycosyltransferase family II (GT-2) in the Pfam Protein Family Database [26]. The predicted XAC3110 protein processes several conserved catalytic residues of glycosyltransferases including the DXD motif (D\textsuperscript{234}TD\textsuperscript{236}) for the divalent metal ion binding in glycosyltransferases with a common GT-A structural fold [27,28] (Figure 2). Database search revealed that XAC3110 are highly conserved in other sequenced Xanthomonas species, including X. oryzae, X. campestris, X. fuscans, X. perforans, X. vesicatoria, X. gardneri, and X. albilineans, with over 70% amino acid identity (Table 1). All these homologues are putative glycosyltransferases with unknown functions. Their homologues with about 35-70% identity are also present in Acetobacter aceti, Clostridium spp., Xylella fastidiosa, Chlorobium phaeobacteroides, Saccharopolyspora erythraea, Thiorhodococcus drewsii, Rhodospirillum centenum, Stenotrophomonas spp., and Burkholderia spp.; among which, several are putative GT-2 proteins (data not shown). These findings strongly suggested that XAC3110 may be a member of GT-2. Collectively, given the role in polysaccharide production (see below), the bdp24 (XAC3110) gene was renamed as gpsX (glycosyltransferase for polysaccharide synthesis in X. citri subsp. citri).

To exclude the possibility of multiple EZ-Tn5 insertions in the genome of the gpsX mutant 223 G4 (gpsX-), and the complemented strain C223G4 (gpsX+), was assayed for EPS and LPS production. The results showed that the total EPS production of the gpsX mutant in NB containing 2% glucose at 24 hours post inoculation could be restored to the wild-type level by the plasmid pJU3110, but not by the empty vector pUFR053 (Figure 3A). Both the mutant strains 223 G4 (gpsX-) and 223G4V (gpsX-) produced significantly less

Figure 1 Schematic diagram of the gpsX (XAC3110) gene in the genome of X. citri subsp. citri strain 306. Open arrows indicate length, location and orientation of the ORFs. The triangle indicates the EZ-Tn5 insertion site in mutant 223 G4 (gpsX-). Gene colour represents operon membership. The middle element shows the 2299 bp PCR fragment cloned into the plasmid pUFR053 for complementation of the gpsX mutant 223 G4 (gpsX-). The lower element shows domain structure analyses of the putative GpsX protein. The domain structure prediction was performed using the SMART program program http://smart.embl-heidelberg.de/. Domain symbols: Glycos_transf_2: glycosyltransferase family 2 domain, SCOP:d1f6da_: UDP-Glycosyltransferase/glycogen phosphorylase superfamily.
EPS than the wild-type strain 306. The complemented strain C223G4 (gpsX+) had a similar level of EPS production to the wild-type strain 306. Sodium dodecylsulphate-polyacrylamide gel electrophoresis (SDS-PAGE) analysis showed that LPS of the gpsX mutant was different from that of the wild-type strain 306 (Figure 3B). Two bands corresponding to the O-antigen containing LPS were completely lost in the gpsX mutant, compared to wild-type strain 306. The LPS pattern of the complemented gpsX mutant was similar with that of the wild-type strain 306. The empty vector pUFR053 did not complement LPS biosynthesis in the gpsX mutant (Figure 3B). These findings indicated that the transposon insertion mutation in gpsX could be complemented by the wild type ORF in trans and, the gpsX locus is involved in polysaccharides biosynthesis in X. citri subsp. citri.

To further confirm the role of gpsX in polysaccharides biosynthesis, the EPS production of the mutants grown in XVM2 liquid medium supplemented with 2% of various carbohydrates was quantitatively estimated. As summarized in Table 3, the gpsX mutant produced about 30-50% less EPS than the wild-type strain 306 when cultured in fructose-, galactose-, glucose-, maltose-, mannose-, or sucrose-containing medium; while the EPS yield of the complemented mutant strains showed no significant difference from that of the wild-type. In contrast, no significant difference in capsule staining was observed between the gpsX mutant strain

Figure 2 Sequence alignment of N-terminal residues of GpsX including the Glycosyltransferase family 2 domain and its glycosyltransferase homologues. The motifs predicted to be involved in the catalytic activity of GpsX are highlighted in gray background and indicated by the symbols (*). Abbreviations are as follows: GpsX, X. citri subsp. citri glycosyltransferase (NCBI Accession No. NP_643419); Xpe_GT, X. perforans glycosyltransferase (ZP_08188792); Xoo_GT, X. oryzae pv. oryzae glycosyltransferase (YP_200377); Xoc_GT, X. oryzae pv. oryzicola glycosyltransferase (ZP_02244158); Xcamv_GT, X. campestris pv. vasculorum glycosyltransferase (ZP_06483586); Xau_GT, X. fuscans subsp. aurantiifoli glycosyltransferase (ZP_06732262); Xve_GT, X. vesicatoria glycosyltransferase (ZP_08176519); Xcv_GT, X. campestris pv. vesicatoria glycosyltransferase (YP_364973); Xga_GT, X. gardneri glycosyltransferase (ZP_08185487); Xcc_GT, X. campestris pv. campestris glycosyltransferase (YP_242265); Xan_GT, X. albilineans glycosyltransferase (AELO8167); Xan_GT, X. albilineans glycosyltransferase (YP_003376724).
we were interested in determining whether the 

oryzae pv. 

Kmr Tra+, conjugation helper plasmid [32]

C223G4 (223G4V (223G4 (pUFR053 IncW Mob+mob(P) lacZ+ Par+, Cm’, Gm’, shuttle vector

puJ3110 2,299-bp KpnI- HindIII fragment containing wild-type gpsX cloned in puJFR053; Cm’, Gm’

Table 1 GpsX/XAC3110 homologues in Xanthomonas spp

| Strains | Gene/locus_tag | Putative product | Size (aa) | Domain structure | Identity (%) |
|---------|----------------|------------------|-----------|-----------------|-------------|
| Xac 306 | gpsX/XAC3110    | glycosyltransferase | 675       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 94         |
| Xpe 91-118 | XPE_2818     | glycosyltransferase | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 97         |
| Xoo KACC10331 | XOO_1639     | glycosyltransferase | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 94         |
| Xoo MAFF311018 | XOO_1639     | glycosyltransferase | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 94         |
| Xoo PXO99A | PXO_01594     | glycosyltransferase | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 94         |
| Xoc BL5256 | Xonyp_01010016275 | glycosyltransferase | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 94         |
| Xcv NCPPB702 | XcampvH_01010002613 | glycosyltransferase | 698       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 93         |
| Xau ICBP10535 | XAUC_30140    | glycosyltransferase | 694       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 93         |
| Xau ICBP11122 | XAUB_29140    | glycosyltransferase | 694       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 93         |
| Xve ATCC35937 | XVE_0383      | glycosyltransferase | 701       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 93         |
| Xcv 85-10 | XCV3242       | glycosyltransferase | 694       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 92         |
| Xga ATCC19865 | XGA_4540       | glycosyltransferase | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 92         |
| Xcc 8004 | XC_1175       | glycosyltransferase | 675       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 90         |
| Xcc ATCC33931 | XCC2933       | glycosyltransferase | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 89         |
| Xcc B100 | xccb100_1219  | hypothetical protein | 700       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 89         |
| Xcr 756C | XCR_3304      | glycosyltransferase | 698       | Glycos_transf_2 (1); SCOP:1f6da_(1) | 89         |

a Xac 306: X. axonopodis pv. citri strain 306 (GenBank accession number: AE008923); Xpe 91-118: X. perforans 91-118 (AEQW00000000); Xoo KACC10331: X. oryzae pv. oryzae KACC10331 (AE0013583); Xoo MAFF311018: X. oryzae pv. oryzae MAFF311018 (AP008229); Xoo PXO99A: X. oryzae pv. oryzae PXO99A (CP000967); Xoc BL5256: X. oryzae pv. oryzicola BL5256 (AAQW00000000); Xcv NCVPB702: X. campestris pv. vasculosum NCVPB702 (ACHS00000000); Xau ICBP10535: X. fuscans subsp. aurantifoli XCB11122 (ACP00000000); Xve ATCC35937: X. sibiricus ATCC35937 (AEQV00000000); Xcv 85-10: X. campestris pv. vesicatoria 85-10 (AM039952); Xga ATCC19865: X. gardneri ATCC19865 (AEQX00000000); Xcc 8004: X. campestris pv. campestris 8004 (CP000059); Xcc ATCC33931: X. campestris pv. campestris ATCC 33931 (AE008892); Xcc B100: X. campestris pv. campestris B100 (AM920689); Xcc 756C: X. campestris pv. raphani 756C (CP002789); Xan GPE PC73: X. albilineans GPE PC73 (FP565176). Dominant Domains were predicted by the SMART program http://smart.embl-heidelberg.de/. Domain symbol: Glycos_transf_2, glycosyltransferase family 2 domain; Glycos_transf_1, glycosyltransferase family 1 domain. The total number of the domains was indicated in the bracket. *According to a BLASTP search.

and the wild-type strain 306 in capsule assays (data not shown).

GpsX was required for full virulence and growth of X. citri subsp. citri in host plants

Since both EPS and LPS have been demonstrated to contribute to host virulence of X. citri subsp. citri [23,34,35], we were interested in determining whether the gpsX gene is associated with pathogenicity of the canker bacterium. The virulence of the gpsX mutant was assessed on the host plant grapefruit using two inoculation methods: pressure infiltration and spray. Upon infiltration at a concentration 10^8 colony-forming units (cfu/ml), no disease symptom was observed on the gpsX mutant or wild type strain inoculated leaves at 7 days post inoculation (dpi); however, the gpsX mutant induced less canker lesions than the wild-type strain 306 at 14 dpi (Figure 4A). When infiltrated at a higher concentration (10^8 cfu/ml),

Table 2 Bacterial strains and plasmids

| Strains and plasmids | Characteristics | Reference or source |
|----------------------|-----------------|---------------------|
| Strains              |                 |                     |
| E. coliDH5α          | F: recA1 endA1 hsdR17 supE44 thi-1 gyrA96 relA1 Δ (argE-lacZYA)169 o80 lacZ Δ M15 | [29] |
| HB101                | F: supE44, hsdS20(rB- mB-) recA1 ara-14, proA2, lacY1, galK2, rpsL20, xyl-5, mtl-1, leuB6, thi | [30] |
| X. citri subsp. citri|                 |                     |
| 306                  | Syn. X. axonopodis pv. citri strain 306; wild type, Rf | [31] |
| 223G4 (gpsX+)        | gpsX (XAC3110): EZ-Tn5 derivative of strain 306, Km’, Rf | [24] |
| 223G4V (gpsX+)       | 223G4 (gpsX+) containing puJFR053, Cm’, Gm’, Km’, Rf | This study |
| C223G4 (gpsX+)       | 223G4 (gpsX+) containing puJ3110, Cm’, Gm’, Km’, Rf | This study |
| Plasmids             |                 |                     |
| puRQ013              | ColE1 Km’ Tra+, conjugation helper plasmid | [32] |
| puJFR053             | IncW Mob* mob(P) lacZ+ Par+, Cm’, Gm’, shuttle vector | [33] |
| puJ3110              | 2,299-bp KpnI- HindIII fragment containing wild-type gpsX cloned in puJFR053; Cm’, Gm’ | This study |

*aP*, Cm’, Gm’, Km’, and Rf indicate resistance to ampicillin, chloromycetin, gentamicin, kanamycin and rifamycin, respectively.
the gpsX mutant induced significantly less lesions than wild type at 7 dpi, but caused similar disease symptoms as wild type at 14 dpi. In both cases, the complemented mutant strain with the intact gpsX cloned in pUFR053 showed no difference from the wild-type strain (Figure 4A). Plant inoculation by spray, a method that mimics the natural infection, showed that the gpsX mutant was reduced in virulence on grapefruit compared with the wild-type strain 306. After 21 days post inoculation the number of canker lesions on leaves infected with the gpsX mutant was significantly less than that inoculated with wild type strain. Symptom induction by the gpsX mutant could be restored to the wild-type level by complementary plasmid pJU3110, but not by the empty vector (Figure 4B).

Although there were no differences between the wild type and the gpsX mutant strains in the ability to grow in XVM2 medium (data not shown), the growth of gpsX mutant 223 G4 (gpsX-) was significantly reduced in planta compared to the growth of the wild-type strain. After inoculation by infiltration at 10^5 cfu/ml, the bacterial population of the gpsX mutant moderately reduced in planta, and between 24 and 48 h, the bacterial population began to increase; whereas the bacterial population of the wild type strain 306 continued to increase after inoculation (Figure 5A). The bacterial population

### Table 3 EPS production in X.citri subsp. citri strains

| Strain     | EPS yield (g/L) |
|------------|-----------------|
|            | Fructose | Galactose | Glucose | Maltose | Mannose | Sucrose | Xylose |
| 306        | 1.73 ± 0.23 a | 1.08 ± 0.24 a | 1.83 ± 0.17 a | 1.22 ± 0.11 a | 1.54 ± 0.27 a | 1.54 ± 0.21 a | 1.15 ± 0.17 a |
| 223G4 (gpsX-) | 0.83 ± 0.14 b | 0.64 ± 0.11 b | 1.22 ± 0.25 b | 0.75 ± 0.19 b | 0.94 ± 0.12 b | 0.68 ± 0.11 b | 1.15 ± 0.17 a |
| C223G4 (gpsX+) | 1.91 ± 0.36 a | 1.22 ± 0.25 a | 1.96 ± 0.34 a | 1.14 ± 0.16 a | 1.45 ± 0.19 a | 1.76 ± 0.31 a | 1.53 ± 0.25 a |

*Strains were cultured in XVM2 liquid medium supplemented with various carbon sources. Data presented are means and standard errors of three replicates from one representative experiment and similar results were obtained in two other independent experiments. Different letters in each data column indicate significant differences at *P* < 0.05 (Student’s t-test).
of the \textit{gpsX} mutant recovered from the infected leaves was approximately 10 to 100-fold lower than that of the wild-type strain at each of the test points (Figure 5A). Similar differences in growth of the wild type and mutant strains were observed following infiltration at $10^8$ cfu/ml (Figure 5B). The bacterial population of the complemented strain was similar to that of the wild-type at each test point (Figure 5A and 5B). The populations of the \textit{gpsX} mutant compared to wild type strain were also analyzed after inoculation via spraying and significant differences were observed. As shown in Figure 5C, after inoculation, the population of the wild type strain remained approximately constant until 4 dpi, whereas the population of the \textit{gpsX} mutant declined significantly. At 4 dpi, the population size of the \textit{gpsX} mutant was nearly 100 times lower than for the wild type strain. From that point forward, the population sizes of the \textit{gpsX} mutant began to increase slowly, whereas growth of the wild type strain continued after inoculation, so that, at 14 dpi, the difference in population size was one to two orders of magnitude. The affected growth of the \textit{gpsX} mutant was restored to wild type levels by complementation with the cloned \textit{gpsX} gene (Figure 5C). Overall, these findings suggest that \textit{gpsX} is required for \textit{X. citri} subsp. \textit{citri} to proliferate well and to achieve full virulence in host plants.

**Figure 4** \textit{GpsX} is important for host virulence of \textit{X. citri} subsp. \textit{citri}. (A) Suspensions of each strain [approximately $10^5$ and $10^8$ cfu/ml, respectively] were inoculated into the intercellular spaces of fully expanded, immature grapefruit (\textit{C. paradise} cv. Duncan) leaves by pressure infiltration with a needleless syringe. A representative leaf from four replicates was photographed at 7 and 14 dpi, respectively. W: wild-type strain 306; M: \textit{gpsX} mutant 223 G4 (\textit{gpsX}-); MV: \textit{gpsX} mutant 223G4V (\textit{gpsX}-) with empty vector pUFR053; CM: complemented \textit{gpsX} mutant C223G4 (\textit{gpsX}+). (B) Bacterial cell suspensions (approximately $10^8$ cfu/ml) of wild-type strain 306 and its derivatives were inoculated onto fully expanded, immature grapefruit by spray. Images are representative of five independent replicates at 21 dpi.

**Mutation in \textit{gpsX} affected biofilm formation of \textit{X. citri} subsp. \textit{citri} on abiotic surfaces and host leaves**

Biofilm has been well characterized as a virulence trait in many plant pathogenic bacteria [36]. Our earlier study indicated that \textit{gpsX} is related to biofilm formation [24]. In order to confirm the role of \textit{gpsX} in biofilm formation in \textit{X. citri} subsp. \textit{citri}, biofilm formation of the \textit{gpsX} mutant was examined on three different kinds of surfaces: polystyrene, glass and host leaves. The \textit{gpsX} mutant 223 G4 (\textit{gpsX}-) exhibited a significant reduction in biofilm formation both on polystyrene surface and in glass tubes compared to that of the wild-type, where the level of biofilm...
formation were approximately 30% and 40% of the wild-type level, respectively; and the complemented C223G4 (gpsX+) strains were restored to levels similar to that of the wild-type strain (Figure 6A and 6B). Similar to the observations on polystyrene surface and in glass tubes, the gpsX mutant showed declined biofilm formation on citrus leaf surfaces and, the complemented C223G4 (gpsX+) strains were restored the wild-type levels (Figure 6C), suggesting that the gpsX gene is involved in biofilm formation of _X. citri_ subsp._citri_.

Mutation of _gpsX_ caused impairment in cell motility but no impact on flagellar formation
Several studies have indicated that both EPS and LPS are associated with the flagella-independent motility in a couple of bacteria including _X. citri_ subsp._citri_ [21,24,37]. To verify whether a mutation in _gpsX_ has any effect on the cell motility of _X. citri_ subsp._citri_, the _gpsX_ mutant was evaluated for the mobile ability on swimming and swarming plates, respectively. The results showed that a significant reduction (_P_ < 0.05, student’s t-test) both in swimming and swarming motility was observed in the _gpsX_ mutant C223 G4 (gpsX-), compared with the wild-type strain (Figure 7). On the tested plates, the diameter of the growth zones resulting from migration away from the inoculation points on the agar surface were about 2.5 cm (swimming plates) and 2.0 cm (swarming plates) for the _gpsX_ mutant, and 4.2 cm (swimming plates) and 3.0 cm (swarming plates) for the wild type. The diameter of the complemented strain and the wild-type strain were not significantly different, indicating that the mobility of the mutant could be restored to wild-type levels by _gpsX_ in _trans_. Flagellum visualization assays using transmission electron microscope (TEM) showed that both the wild-type and the _gpsX_ mutant strains formed a polar flagellum at the cell surface (data not shown), suggesting that mutation of _gpsX_ has no impact on flagellar formation in _Xac_. These results indicated that the _gpsX_ is implicated in bacterial motility in _X. citri_ subsp._citri_.

GpsX contributed to stress tolerance of _X. citri_ subsp._citri_
The decrease in bacterial population _in planta_ of the _gpsX_ mutant immediately after inoculation (Figure 5A, B...
Biofilm formation by *X. citri* subsp. *citri* strain 306 and its derivatives. Biofilm formation in polystyrene 96-well plates (A), glass tubes (B) and on citrus abaxial leaf surfaces (C) was visualized using crystal violet staining. Biofilm formations in glass tubes were quantified by measuring the optical density at 590 nm after dissolution in ethanol-acetone (80:20, v/v). WT: wild-type strain 306; M: *gpsX* mutant 223 G4 (*gpsX*<sup>-</sup>); MV: *gpsX* mutant 223G4V (*gpsX*<sup>-</sup>) with empty vector pUFR053; CM: complemented *gpsX* mutant C223G4 (*gpsX*<sup>+</sup>); CK-: XVM2 medium without inoculation of bacteria. All experiments were performed in quadruplicate and repeated three times with similar results, and only one representative result is presented. Means ± standard deviations are shown.

Motilities of *X. citri* subsp. *citri* strains. Cells were inoculated onto NA plates supplemented with 0.25% or 0.60% agar from bacterial cultures at exponential stage and photographed after 3 days (0.25% agar plate) and 7 days (0.60% agar plate) of incubation at room temperature (22-23°C). WT: wild-type strain 306; M: *gpsX* mutant 223 G4 (*gpsX*<sup>-</sup>); MV: *gpsX* mutant 223G4V (*gpsX*<sup>-</sup>) with empty vector pUFR053; CM: complemented *gpsX* mutant C223G4 (*gpsX*<sup>+</sup>). The *gpsX* mutant 223 G4 (*gpsX*<sup>-</sup>) is reduced in cell motility, which could be restored to the wild-type level by providing a plasmid bearing the *gpsX* gene [i.e., the complemented strain C223G4 (*gpsX*<sup>+</sup>)].
and 5C) suggested that the gpsX gene might play a role in the adaptation of X. citri subsp. citri to the conditions of the host microenvironments. To test this hypothesis, the survival of the gpsX mutant was investigated under various stresses that would be likely experienced at the early stage of infection when the bacteria has to attach to the leaf surface and later when the bacteria has to survive inside the host plant, including UV radiation, heat shock, saline stress, osmotic challenge, desiccation stress, SDS exposure, and the H$_2$O$_2$ oxidative stress. These assays revealed that the gpsX mutant 22G4 (gpsX-) was more sensitive than the wild-type strain to UV radiation, heat shock, desiccation stress, SDS exposure, and H$_2$O$_2$ (Table 4). After 20 min of exposure to UV radiation, there were greater numbers of surviving cells of the wild-type strain than that of the gpsX mutant. Following 15 min of exposure of bacteria to heat (50°C), viable cells of the gpsX mutant declined more rapidly than the wild-type. When exposed to air and dried for 60 min, the gpsX mutant showed significantly decreased survival compared with the wild-type strain. After treatment with SDS (0.1%) for 10 min, the survival rate of the gpsX mutant was significantly lower than that of the wild-type strain. The gpsX mutant also showed higher sensitivity than the wild-type strain to hydrogen peroxide (exposure to 0.03% H$_2$O$_2$ for 20 min). The levels of stress tolerance of the complemented strain were similar to those of the wild type (Table 4), indicating that the affected stress tolerance of the gpsX mutant could be restored by gpsX in trans. There were no differences between the gpsX mutant and wild type strain in survival under saline stress or osmotic challenge.

**Mutation of gpsX has no impact on expression of virulence-related genes**

Reduced virulence could result from down-regulation of key virulence genes. In order to test whether mutation of the gpsX gene affected the expression of virulence-related genes, quantitative reverse transcription-PCR (QRT-PCR) assays were performed to monitor the expression profiles of six genes which were selected based on the alternated mutant phenotypes mentioned above. For total RNA preparation, the gpsX mutant and wild type strains were cultured to exponential phase in XVM2 medium that has been reported to mimic the environment of plant intercellular spaces [38]. The six target genes included one EPS biosynthesis gene (gumB), one LPS synthesis gene (rfbC), one catalase gene (katE), one TTSS component gene (hrcV), one TTSS regulator genes hrpX, and one TTSS effector gene (pthA). The results showed that none of the six genes was significantly differently expressed in the mutant 22G4 (gpsX-) compared with wild-type strains when grown in XVM2 medium (Table 5), based on a student’s t-test ($P < 0.05$). Specifically, the primer set used for pthA is present in pthA4 and its homologues pthA1, pthA2, and pthA3, but not in any other genes. Thus we refer it as pthA rather than differentiating them. The qRT-PCR result based on this primer should detect the expression of pthA4, pthA1, pthA2, and pthA3. It is very likely that pthA4, pthA1, pthA2, and pthA3 have similar gene expression pattern due to the same promoter sequences. The sequences are 100% identical in the 213 bp upstream of pthA4, pthA1, pthA2, and pthA3 including the predicted promoter region (data not shown). Consequently, the qRT-PCR result will represent the relative fold change in gene expression for pthA4, pthA1, pthA2, and/or pthA3 since it is relative fold change and not absolute expression value.

**Discussion**

In this work we have extended the characterization of the XAC3110 gene locus, previously identified and named bdp24 for involvement in Xac biofilm formation [24]. We conclude from several independent lines of evidence that this gene is required for EPS and LPS biosynthesis, and consequently required for biofilm formation and full virulence of Xac on host plants. For this reason, we have

| Strains | Survival rate (%)$^*$ |
|---------|----------------------|
| 306     | 3.2 ± 1.2a 0.04 ± 0.02a 2.7 ± 0.7a 10.1 ± 3.1a 1.6 ± 0.5a 4.9 ± 2.3a 6.1 ± 2.4a |
| 223G4 (gpsX) | 0.9 ± 0.3b 0.004 ± 0.003b 0.4 ± 0.1b 0.05 ± 0.02 b 0.05 ± 0.02b 3.8 ± 1.4a 3.9 ± 2.2a |
| 223G4V (gpsX) | 1.1 ± 0.5b 0.005 ± 0.003b 0.7 ± 0.2b 0.08 ± 0.03 b 0.12 ± 0.04b 4.1 ± 1.7a 5.5 ± 1.7 a |
| C223G4G (gpsX+) | 4.2 ± 1.6a 0.05 ± 0.03a 3.5 ± 1.3a 8.2 ± 2.5a 2.2 ± 0.4a 5.5 ± 2.4a 7.4 ± 2.8a |

$^*$Bacterial cell viability was estimated by plating on NA agar before (T0) and after (T1) the treatment. Percentage survival was calculated as the ratio of viable cell counts at T1 to that at T0. The treatments were applied as follows: for UV radiation, overnight bacterial culture was exposed to short-wave UV irradiation (254 nm) for 20 min; for heat-shock, bacterial culture was incubated at 50°C for 15 min; for desiccation stress, bacterial culture was placed on glass coverslips and air dried in a laminar flow apparatus for 60 min and then resuspended in 0.85% NaCl and plated; for SDS exposure, bacterial culture was treated with 0.1% SDS for 20 min; for sensitivity to hydrogen peroxide, bacterial culture was exposed to 0.03% H$_2$O$_2$ for 20 min; for osmotic stress, bacterial culture was treated with 40% D-sorbitol for 40 min; for saline stress, bacterial culture was treated with 1.0 M NaCl for 20 min. Bacterial cells were serially diluted with NB medium and colony-forming units (cfu) were counted after being cultured on NA plates at 28°C for 48 h. Each test, plated in triplicate, was repeated three times with similar results. Data shown are means and standard errors of three replicates from one representative experiment. Different letters in each data column indicate significant differences at $P < 0.05$ (Student’s t-test).
changed the name of this gene to gpsX, for glycosyltransferase for polysaccharide synthesis in Xac, to reflect the apparent multiple function of the gene product.

Several lines of evidence indicate that the gpsX locus is involved in polysaccharide biosynthesis. First, GpsX contains a glycosyltransferase family 2 domain and shares the conserved catalytic residues of glycosyltransferases (Figure 1 and 2). Second, the general role of glycosyltransferases in polysaccharide biosynthesis [12,13]. Third, similar genes associated with polysaccharide biosynthesis have been identified in other bacterial pathogens (see below). Homologues of GpsX widely occur in the genomes of related phytopathogenic bacteria of Xanthomonas (Table 1). The biochemical characteristics and physiological roles of these homologous proteins remain unknown.

Some glycosyltransferase genes have already been identified in Xanthomonas spp. For example, as mentioned previously, the rfbC gene encodes a glycosyltransferase, which serves as a truncated O-antigen biosynthesis protein involved in EPS production in X. citri subsp. citri [23]. Both the ORFs XC_3814 and XC_3555 (xagB) in X. campestris pv. campestris are implicated in EPS production, but not EPS production [21,22]. In addition, the gumDMHK1 genes encode different glycosyltransferases that direct the biosynthesis of the xanthan EPS in X. campestris pv. campestris [39,40]. However, these enzymes in Xanthomonas are mono-functional, i.e., involved either in EPS or LPS production. Our data showed that the gpsX gene is involved in both EPS and LPS production (Figure 3). The low similarities between GpsX and these proteins (data not shown) may suggest the differences in substrates and products.

Table 5 Comparison of virulence gene expression in the wild type and the gpsX mutant cultured in XVM2 medium by QRT-PCR

| Gene       | locus_tag | Function of protein product | ΔΔC \text{p}\_t ± SD\_p | Fold change ± SD | P \text{p} |
|------------|-----------|----------------------------|--------------------------|-----------------|------|
| gumB       | XCA2385   | EPS xanthan biosynthesis    | 0.2665 ± 0.1912          | 0.8314 ± 0.1102 | 0.0524 |
| rfbC       | XAC3598   | LPS O-antigen biosynthesis  | -0.2018 ± 0.1467         | 0.8695 ± 0.0841 | 0.0621 |
| katE       | XAC1211   | Monofunctional catalase     | 0.0758 ± 0.1346          | 0.9485 ± 0.0871 | 0.4407 |
| pthA       | NS \_p   | TTSS effector               | -0.1703 ± 0.2407         | 1.1253 ± 0.1845 | 0.3128 |
| hrcX       | XAC1266   | TTSS regulator              | 0.2578 ± 0.1638          | 0.8364 ± 0.0997 | 0.2442 |
| hrcV       | XAC0405   | TTSS component              | 0.1828 ± 0.1348          | 0.8811 ± 0.0832 | 0.1119 |

\( ^{\text{a}} \) Both 16S rRNA and gyrA genes were used as endogenous controls in the QRT-PCR experiments and similar results were obtained when the data were normalized against the two genes respectively. Only the data obtained with 16S rRNA gene as control were shown.

\( ^{\text{b}} \) The ΔΔC\_p \_t was determined using four biological repeats. The experiment was repeated two times with similar results. Data from one experiment are shown.

\( ^{\text{c}} \) The expression change (mutant/wild type) in mutant 223 G4 (gpsX) was calculated using \( ^{\text{c}} \)ΔΔC\_p \_t.

\( ^{\text{d}} \) P value, analyzed by Student’s t-test. Values are significantly different when \( ^{\text{d}} \)P < 0.05.

\( ^{\text{e}} \) No specific locus_tag. This represents the gene expression of pthA1, pthA2, pthA3 and pthA4.

Bacterial polysaccharides are usually synthesized from intracellular nucleotide sugar precursors and, most bacterial polysaccharides contain polymerized saccharide repeating units, the assembly of which involves glycosyltransferases that sequentially link monosaccharide moieties from nucleotide sugars to the growing sugar chain (saccharide acceptors) [11]. Different classes of bacterial polysaccharides can be distinguished on basis of their biosynthesis mechanisms and the precursors required. However, it is worth mentioning that, in some instances, mutation of single genes simultaneously affected biosynthesis of different polysaccharides, similar with the observation in this work. For example, in X. campestris pv. citrumelo, the role of gpsX, a homologue of waaF (rfaF) which codes for a heptosyltransferase for LPS synthesis in E. coli, affected biosynthesis of LPS and EPS [41]. In addition, mutants in xanA and xanB, involved in UDP-Glucose and GDP-Mannose biosynthesis in X. campestris pv. campestris, respectively, showed a decrease in EPS production and an altered LPS [42]. Mutants in galE, encoding a UDP-galactose epimerase in Erwinia amylovora, were deficient in EPS production and produced a LPS with an altered side chain structure [43]. The dual effect of certain genes on EPS and LPS may be due to the shared pathways for EPS and LPS synthesis in these bacteria. As discovered in Salmonella, the same precursor molecule, UDP-glucose, is used for LPS O-antigen polysaccharide and capsular polysaccharide [44]. The major EPS produced by xanthomonads, xanthan, composed of polymerized pentasaccharide repeating units, consisting of glucose, mannose and glucuronic acid [39]. Most recently, glucose and mannose were found to be components of LPS in X. citri. Given the altered O-antigen containing LPS profile of the gpsX mutant and its decreased level of EPS production, it was likely that the gpsX-encoded glycosyltransferase was involved in the formation of saccharide repeating units that might be found in X. citri subsp. citri EPS and LPS, by transferring the glucose and/or mannose monosaccharide moiety from certain nucleotide sugar precursors to corresponding acceptors.
However, biochemical evidence for this proposed function of GpsX is needed.

Interestingly, the gpsX gene is located outside of the LPS gene cluster even though it is involved in the O-antigen biosynthesis. The LPS cluster is responsible for synthesis of O-antigen polysaccharide. The locus presents a size of 19.9 kb and contains 17 ORF [7,46]. The LPS cluster contains three glycosyltransferases, i.e. XAC3598 (RfbC), ORF5, and XAC3595. RfbC was annotated as a 614-amino-acid truncated O-antigen biosynthesis protein containing two separate glycosyltransferase family 2 (GT2) domains. The involvement of rfbC in O-antigen biosynthesis has been confirmed in our previous study [23]. The orf5 has been annotated to encode a putative glycosyltransferase [46], whereas XAC3595 shows significant homology to the glycosyltransferase A (GtrA) family [46]. It remains to be determined how GpsX and other glycosyltransferases are involved in O-antigen biosynthesis in Xac.

The attenuation in virulence and growth in planta of the gpsX mutant both in epiphytic (Spray) and wound (pressure infiltration) inoculations (Figure 4 and 5) may result, at least partially if not completely, from the reduction in EPS production (Figure 3A) and the alteration of LPS profile (Figure 3B), and consequently impaired cell motility (Figure 7) and biofilm formation (Figure 6), rather than from an effect on the virulence genes (Table 5). EPS has been shown to act as an important virulence factor that contributes to epiphytic survival and/or bacterial in planta growth and disease symptom formation in several Xanthomonas spp. including X. campestris pv. campestris, X. oryzae pv. oryzae, and X. citri subsp. citri [8]. EPS can suppress plant basal defense responses by chelating divalent calcium that are required for the activation of plant defense responses [47,48]. It also contributes to biofilm formation [21,24,34,49], which promotes bacterial resistance to environment stresses [23,36]. LPS has also been shown to be an important virulence factor in various plant pathogenic bacteria including several Xanthomonas spp. [8], Erwinia amylovora [50] and Pseudomonas syringae [51]. It can serve as a physical barrier protecting bacteria from plant defense responses [51]. It may also contribute to biofilm formation [23,24]. In addition, both EPS and LPS are related to cell motility in a couple of bacteria including X. citri subsp. citri [21,24,37]. In certain phytopathogenic bacteria, e.g., E. amylovora, P. syringae, andRalstonia solanacearum, motility has been suggested to contribute to bacterial virulence in the early stages such as invasion and colonization [52-54]. X. citri subsp. citri is an intercellular space-colonizing pathogen that invades host plants via stomata or wounds, and multiplies in the apoplasts [4]. Before entering the host, the pathogen persists as epiphytes on the plant surface and has to confront environment stresses. Once entering the host, the pathogen needs to tolerate preformed defense molecules to establish a successful infection. Therefore, the alteration of LPS of the gpsX mutant and the reduction in its ability to produce EPS and to form biofilm formation may attenuate its adaptation to the conditions of the host microenvironment, which result in its growth deficiency, and consequently reduced virulence. Consistent with this, it has been demonstrated that both EPS and LPS biosyntheses are required for growth and survival on leaf surfaces and full virulence in X. citri subsp. citri [23,34]. Finally, gpsX may aid bacterial survival at early stage of infection when the bacterium attaches to the leaf surface and later survives inside the plant tissue. Consistent with the hypothesis, the gpsX mutant was attenuated in resistance against various stresses including oxidative stress (Table 4), which is one of the early plant defense responses triggered by bacterial infections [55].

In summary, in this work we expanded the knowledge about the function of the novel glycosyltransferase encoding gene gpsX from X. citri subsp. citri. Based on its apparently unique function in polysaccharide synthesis and the widely conserved occurrence in sequenced strains of Xanthomonas, this enzyme may represent a novel virulence-related factor of phytopathogenic Xanthomonas including X. citri subsp. citri. Additional study of this gene and its protein product should yield new insights into the biochemistry and physiological roles of bacterial glycosyltransferase of the citrus canker bacterium X. citri subsp. citri.

Conclusions

In this report we characterized the novel gpsX gene in X. citri subsp. citri. We demonstrated that the gpsX mutant is affected in EPS and LPS production, cell motility, biofilm formation, stress tolerance, growth in planta, and virulence on host plants and that the genetic complementation with the wild type gpsX gene, fully restored the affected phenotypes of the gpsX mutant to wild-type levels. In conclusion, the gpsX gene is important for polysaccharide synthesis and biofilm formation and thus, plays an important role in the adaptation of X. citri subsp. citri to the host microenvironments at early stage of infection and required for full virulence on host plants.

Methods

Bacterial strains, plasmids and growth conditions

The bacterial strains and plasmids used in this study are listed in Table 2. E. coli strains were grown in Luria-Bertani (LB) medium at 37°C. Xac wild type strain306 (rifamycin resistant) and the EZ-Tn5 insertion mutant strain 223 G4 (gpsX-) have been described previously [24]. Xac strains were grown in nutrient broth/agar (NB/NA) or XVM2 medium [38] at 28°C. Antibiotics were added at the following concentrations when required: ampicillin (Am) 50 μg/ml; chloramphenicol
(Cm), 35 μg/ml; gentamycin (Gm), 5 μg/ml; Kanamycin (Km), 50 μg/ml; and rifampycin (Rf), 50 μg/ml.

**DNA manipulations**

Bacterial genomic DNA and plasmid DNA were extracted using a Wizard genomic DNA purification kit and a Wizard miniprep DNA purification system following manufacturer’s instructions (Promega, Madison, WI, USA). The concentration and purity of DNA were determined using a Nanodrop ND-1000 spectrophotometer (NanoDrop Technologies, Wilmington, DE, USA). The standard methods [56] were used for restriction digestion, DNA ligation, agarose gel electrophoresis and DNA transformation of *E. coli*. The restriction endonucleases, T4 DNA ligase and Pfu polymerase were provided by Promega (Promega Corporation, Madison, WI).

**Complementation of the gpsX mutant**

For complementation of the *gpsX* mutant 223 G4 (*gpsX-*), a 2,299-bp DNA fragment containing the intact open reading frame (ORF) of *gpsX* and 230 bp upstream of the 5’ end to 21 bp downstream of the 3’ end of the ORF, was amplified from the genomic DNA of *Xac* strain 306 using the primers C10-F (5’- tcgagttcggtgtcctc-3’) and C10-R (5’- tgcagttcccacgcct-3’), respectively containing *Kpn*I and *Hind*III restriction enzyme sites (underlined). The PCR product was digested with *Kpn*I and *Hind*III and cloned into the complementary vector pUFR053 [33] to construct the recombinant plasmid pJU3110 (Table 2). The recombinant plasmid was transferred into the *gpsX* mutant 223 G4 (*gpsX-*) by triparental conjugation as described previously [57], resulting in strain C233G4 (*gpsX+*) (Table 2).

**Quantitative determination of EPS production**

To estimate EPS production, strains were cultured in 100 ml NB or XVM2 liquid medium containing 2% (wt/vol) various sugars (fructose, galactose, glucose, maltose, mannose, sucrose, and xylose) at 28°C with shaking at 200 rpm for 24 hours (in NB) or 48 hours (in XVM2). EPS was precipitated from the culture supernatant at different time point post inoculation with ethanol, dried, and weighed as described elsewhere [35].

**Lipopolysaccharides (LPS) analysis**

Bacterial strains were cultured at 28°C in NB or XVM2 liquid medium with shaking (200 rpm). Five-milliliter samples of cultures at the exponential stage were collected and the LPS samples were isolated as described previously [23]. LPS was separated by SDS-PAGE and visualized using silver staining following the manufacturer’s instructions (Bio-Rad Laboratories, Inc., Hercules, CA). Standard LPS from *Salmonella enterica* serovar Typhimurium was obtained from Sigma. The test was performed three times independently.

**Capsule assays**

Bacterial capsules were stained using a capsule-staining kit (Eng Scientific Inc., Clifton, NJ, USA) following the manufacturer’s instructions. The samples were photographed using a light microscope (Leica DMLB2; Leica Microsystems GmbH, Wetzlar, Germany) with a digital camera. The experiment was repeated three times.

**Biofilm formation assays**

Biofilm formation on polystyrene and glass surfaces were examined as described previously [23] with modifications. The average of four replicates was used for quantitative measurement. Assays of biofilm formation on leaf surfaces were conducted as described previously [58] with modifications. Briefly, 20 μl of each bacterial suspension (10^8 cfu/ml) was incubated on the abaxial surface of citrus leaves, and the leaves were kept at 28°C in a humidified chamber. After 24 h of incubation, biofilm formation on the leaf surface was visualized using crystal violet staining. The experiments were repeated three times.

**Motility assays**

To test cell motility, 2 μL of bacterial cultures at the exponential stage in NB (OD600 of 0.8) was spotted onto NA plates (diameter, 150 mm; each containing 50 mL of NA) containing 0.25% (wt/vol) agar (Difco, Franklin Lakes, NJ) for swimming motility testing or 0.6% (wt/vol) agar for swarming motility testing. Plates were incubated at room temperature for 7 days. The diameters of the areas occupied by the strains were measured, and the values were used to indicate the motility of *Xac* strains. The experiment was repeated three times with three replicates each time.

**Electron microscopy**

For flagella visualization, cells grown on NA plates were harvested at 48 hours post inoculation (hpi) and suspended in 0.85% NaCl. One drop of cell suspension was placed onto a 400-mesh Formvar carbon-coated grid. Excess water was removed by blotting onto Whatman filter paper no. 1 (Whatman Inc, Piscataway, NJ, USA). One drop of 1% uranyl acetate solution was then added, and excess solution was removed. The grids were left at room temperature for 30 min. Samples were viewed with a Philips FEI Morgagni 268 transmission electron microscope (FEI Company, Eindhoven, Netherlands) operating at 80 kV.

**Stress tolerance assays**

The assays were performed as described previously with modifications [23]. Bacterial culture at early exponential
stage (OD$_{600nm}$ = 0.1) in NB were used to test survival under stresses: UV radiation, heat shock, saline stress, osmotic challenge, desiccation stress, SDS stress and oxidative stress. In each stress treatment, cell viability was determined by plate-counting of cfu. The survival rate was defined as the percentage of viable cell counts from the culture with stress treatment compared with those from the non-treated culture. The stress treatments were applied as follows: for UV radiation, the cells were exposed to short-wave UV radiation (254 nm in a biological safety cabinet) at a distance of 60 cm for 20 min; for heat-shock stress, the culture was transferred to 50°C for 15 min; for sodium stress, NaCl (pH 7.5) was added to the bacterial culture at a final concentration of 1.0 M, and survival was estimated after 20 min, respectively; for osmotic challenge, D-sorbitol (pH 7.0) was added to the bacterial culture at a final concentration of 40%, and survival was estimated after 40 min; for desiccation stress, the bacterial culture was placed on glass coverslips (18 mm × 18 mm), air dried in a laminar flow apparatus for 60 min and then resuspended in 0.85% NaCl and plated; for SDS stress, SDS (pH 7.5) was added to the bacterial culture at a final concentration of 0.1%, and survival was estimated after 10 min; for oxidative stress, H$_2$O$_2$ was added to the bacterial culture at a final concentration of 0.03%, and survival was estimated after 20 min. Each stress test was repeated three times with three replicates each time. Student’s $t$-test was used to test the significance of the differences.

**Pathogenicity assays**

Pathogenicity assays were performed as described previously [59]. Briefly, fully expanded, immature leaves of young (about 10-week-old) grapefruit (*Citrus paradise* cv. Duncan grapefruit) were prepared in a quarantine foundation. This work has been supported by Citrus Research and Education Foundation.

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**Authors’ contributions**

JL and NW conceived and designed the experiments, performed the experiments, analyzed the data and wrote the paper. All authors read and approved the final manuscript.

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