The filamentous phage XacF1 causes loss of virulence in Xanthomonas axonopodis pv. citri, the causative agent of citrus canker disease

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INTRODUCTION

Xanthomonas axonopodis pv. citri, Xac (syn. Xanthomonas campestris pv. citri), is the causative agent of Asiatic citrus canker disease (ACC), one of the most serious citrus plant diseases in the world (Civerolo, 1984; Graham et al., 2004). Under natural conditions, the bacterium can invade all aboveground parts of plants, entering through natural openings and wounds (Brunings and Gabriel, 2003; Vojnov et al., 2010). A characteristic symptoms include raised corky lesions surrounded by a water or oil-soaked margin on leaves, stems, and fruits, including defoliation, twigs dieback, general tree decline, blemished fruit, and premature fruit drop in severely infected trees (Graham et al., 2004). Management of ACC relies on an integrated approach that includes: (1) replacement of susceptible citrus species with resistant ones; (2) production of disease-free nursery stock; (3) reduction of pathogen spread by establishing windbreaks and fences around groves; (4) preventative copper sprays; and (5) application of insecticide to control Asian leafminer. Because of the limited effectiveness of the current integrated management strategies, citrus canker disease continues to be an economically serious problem for field-grown crops worldwide (Balogh et al., 2010). Hence, alternative control methods are necessary.

Bacteriophages have been recently evaluated for controlling a number of phytopathogenic bacteria and are now commercially available for some diseases (Balogh et al., 2010). The use of phages for disease control is a fast expanding area of plant protection, with great potential to replace existing chemical control measures. In this study, filamentous phage XacF1, which can infect Xanthomonas axonopodis pv. citri (Xac) strains, was isolated and characterized. Electron microscopy showed that XacF1 is a member of the family Inoviridae and is about 600 nm long. The genome of XacF1 is 7325 nucleotides in size, containing 13 predicted open reading frames (ORFs), some of which showed significant homology to Ff-like phage proteins such as ORF1 (pII), ORF2 (pV), ORF6 (pIII), and ORF8 (pVI). XacF1 showed a relatively wide host range, infecting seven out of 11 strains tested in this study. Frequently, XacF1 was found to be integrated into the genome of Xac strains. This integration occurred at the host dif site (attB) and was mediated by the host XerC/D recombination system. The attP sequence was identical to that of Xanthomonas phage Cf1c. Interestingly, infection by XacF1 phage caused several physiological changes to the bacterial host cells, including lower levels of extracellular polysaccharide production, reduced motility, slower growth rate, and a dramatic reduction in virulence. In particular, the reduction in virulence suggested possible utilization of XacF1 as a biological control agent against citrus canker disease.

Keywords: filamentous phage, loss of virulence, citrus canker, biocontrol
In the current study, we isolated and characterized a novel filamentous phage and showed that changes occurred at a cellular level in *X. axonopodis pv. citri* strains following infection. This filamentous phage might be a unique biological agent for use against bacterial citrus canker disease.

**MATERIALS AND METHODS**

**BACTERIAL STRAINS AND GROWTH CONDITIONS**

Ministry of Agriculture, Forestry, and Fisheries (MAFF) strains of *X. axonopodis pv. citri*, Xac (Table 1) were obtained from the National Institute of Agrobiological Sciences, Japan. Strain KC33 was obtained from the National Institute of Fruit Tree Science, the National Agriculture and Food Research Organization, Japan. All strains were stored at −80°C in 0.8% nutrient broth (NB) (BBL, Becton Dickinson and Co., Cockeysville, MD, USA) supplemented with 30% (v/v) glycerol. The strains were grown on nutrient agar (NA) medium (Difco, BBLBD, Cockeysville, MD, USA) supplemented with 30% (v/v) glycerol. The strains were grown on nutrient agar (NA) medium (Difco, BBLBD, Cockeysville, MD, USA) at 28°C. For preparation of bacterial suspension, Xac strains were cultured for 24 h at 28°C with shaking at 220 rpm in NB medium.

For time course experiments, phage-infected and uninfected cells were grown overnight in 5 mL of NB media. Then, 0.5 mL of the cell suspensions (10⁸ cfu/mL) were transferred into 100-mL flasks containing 30 mL of NB medium. Cultures were grown at 28°C with agitation at 200 rpm, and OD₆₀₀ nm measurements were taken every 3 h over the course of 48 h using a spectrophotometer. Three replicates were included at each time point. The experiments were repeated twice (Li and Wang, 2011).

**BACTERIOPHAGE ISOLATION, PURIFICATION, AND CHARACTERIZATION**

The presence of filamentous phages in collected soil samples from cropping fields in Japan was detected by the spot test and plaque-forming assay technique (Yamada et al., 2007). Approximately 10 g of soil was placed in a sterile 50 mL conical centrifuge tube that then was filled to the top with tap water, and allowed to stand for 20 min with periodic inversions. The tubes were then centrifuged at 15,000 × g for 20 min and the supernatant was passed through a membrane filter (0.45-μm pore size) (Millipore Corp., Bedford, MA, USA). One-hundred-microliter aliquots of the soil filtrate were subjected to spot test and plaque-forming assay with strains of Xac (Table 1) as host on NB plates containing 1.5% (w/v) agar. Phages were propagated and purified from single-plaque isolates. An overnight culture of bacteria grown in NB medium (1 mL) was diluted 100-fold with 100 mL of fresh NB medium in a 500 mL flask. To collect a sufficient amount of phage particles, a total of 2 L of bacterial culture was grown. When the cultures reached an OD₆₀₀ of 0.2, bacteriophage was added at a multiplicity of infection (moi) of 0.001–1.0. After further growth for 12–24 h, the cells were removed by centrifugation in a Hitachi Himac CR21E centrifuge with an R12A2 rotor at 8000 × g for 15 min at 4°C. The supernatant was passed through a 0.45-μm-pore membrane filter followed by precipitation of the phage particles in the presence of 0.5 M NaCl and 5% (v/v) polyethylene glycol 6000 (Kanto Chemical Co., Tokyo, Japan). The pellet was collected by centrifugation in a Hitachi Himac CR21E centrifuge with an RPR20-2 rotor at 15,000 × g for 30 min at 4°C, and was dissolved in SM buffer [50 mM Tris/HCl at pH 7.5, 100 mM NaCl, 10 mM MgSO₄ and 0.01% gelatin (w/v)]. Phages were stored at 4°C in complete darkness. Phage titers were determined by serial dilution and subsequent plaque-forming assays (Yamada et al., 2007). The purified phage [10¹³ pfu/mL] was stained with sodium phosphotungstate prior to observation in a Hitachi H600A electron microscope, according to the methods of Dykstra (1993)].

**PHAGE SUSCEPTIBILITY AND ADSORPTION ASSAYS**

The phage susceptibility assays were based on a standard agar overlay method with dilution series of phage preparations (Yamada et al., 2007; Ahmad et al., 2014). Small turbid plaques, typical of Ff-phages, always appeared at reasonable frequencies depending on input phage titers (usually 300–600 pfu/plate), if the bacterial strain was sensitive to the phage. No spontaneous phages (induced prophages) appeared from either strain tested under usual plaque assay conditions. In the phage adsorption assay, exponentially growing cells (OD₆₀₀ 0.1) of the test strain were mixed with XacF1 phage at moi of 0.1, and the mixture was incubated for 0 min (no adsorption) and 30 min at 28°C to allow binding of the phage to the cell surface. Following centrifugation at 15,000 × g for 5 min at 4°C in a Sakuma SS-1500 microcentrifuge (Sakuma Seisakusho, Tokyo, Japan), the phage titer in the supernatant was determined by a standard plaque assay with the indicator strain (MAFF301080). *Escherichia coli* JM109 was used as a negative control.

**DNA ISOLATION AND MANIPULATION**

Standard molecular biological techniques for DNA isolation, digestion with restriction enzymes and other nucleases, and construction of recombinant DNAs were followed, according to Sambrook and Russell (2001). Phage DNA was isolated...
from the purified phage particles by phenol extraction. In some cases, extrachromosomal DNA was isolated from phage-infected Xac cells by the minipreparation method (Ausubel et al., 1995). Replicative-form (RF) DNA for sequencing was isolated from host bacterial cells infected with XacF1 phage, treated with S1 nuclease, and then shotgun-sequenced by Hokkaido System Science Co. (Sapporo, Japan) using a Roche GS Junior Sequence System. The draft assembly of the obtained sequences was assembled using GS De novo Assembler v2.6. The analyzed sequences corresponded to 156 times the final genome size of XacF1 (7325 bp). Computer-aided analysis of the nucleotide sequence data was performed using DNASTAR v3.6 (Hitachi Software Engineering Co., Tokyo, Japan). Potential ORFs larger than 80 bp were identified using the online program ORF Finder (http://www.ncbi.nlm.nih.gov/gorf/gorf.html) and the DNASTAR program. Sequence alignment was performed using the ClustalW (Larkin et al., 2007) program. To assign possible functions to the ORFs, DDBJ/EMBL/GenBank databases were searched using the FASTA, FASTX, BLASTN, and BLASTX programs (Altschul et al., 1997).

**DETERMINATION OF attL AND attR SEQUENCES IN Xac MAFF673010**

Chromosomal DNA was extracted from Xac MAFF673010 after infection with XacF1 and subjected to PCR to amplify fragments containing left and right attachment sites (attL and attR). The attL was amplified using a 29-base forward primer, 5′-TGC GAT CGA GCA GCT TCC CAG TTG GCG AT-3′ (primer P1) and a 30-base reverse primer, 5′-TTC GAT GGT CAC GGT GCC TGT AGT AGA GGC-3′ (primer P2), while attR was amplified using a 30-base forward primer, 5′-ATA ATT TGC TTG ACA CC GTG TGC AGT AGT CTT-3′ (primer P3) and a 28-base reverse primer, 5′-CCT TGA CCG TCA GGG ACT GCA TCA GCC TCT-3′ (primer P4). The primer sequences were based on the dif (attB) region sequence of Xanthomonas citri subsp. citri Aw12879 (DDBJ accession no. CP003778.1). The PCR products were purified from an agarose gel and subjected to sequencing.

**SOUTHERN HYBRIDIZATION**

Genomic DNA from bacterial cells was prepared by the minipreparation method according to Ausubel et al. (1995). Following digestion with restriction enzyme Hincll, DNA fragments were separated by agarose gel electrophoresis, blotted onto a nylon membrane (Piodyne; Pall Gelman Laboratory, Closter, NJ, USA), hybridized with a probe (the entire XacF1 DNA digested by EcoRI), labeled with fluorescein (Gene Images Random Prime labeling kit; Amersham Biosciences, Uppsala, Sweden), and detected with a Gene Images CDP-Star detection module (Amersham Biosciences). Hybridization was performed in buffer containing 5× SSC (0.75 M NaCl, 0.075 M sodium citrate), 0.1% (w/v) sodium dodecyl sulfate (SDS), 5% liquid block, and 5% (w/v) dextran sulfate for 16 h at 65°C. The filter was washed at 60°C in 1× SSC and 0.1% (w/v) SDS for 15 min and then in 0.5× SSC and 0.1% (w/v) SDS for 15 min with agitation, according to the manufacturer’s protocol. The hybridization signals were detected by exposing X-ray film (RX-U; Fuji Film, Tokyo, Japan) to the filter.

**EPS ASSAY**

EPS in bacterial culture supernatants was determined quantitatively as described previously (Guo et al., 2010). Briefly, bacterial strains were grown in NB supplemented with 2% (w/v) D-glucose for 24 h at 28°C with shaking at 200 rpm. A 10-ml portion of the culture was collected, and the cells were removed by centrifugation (5000 × g for 20 min). The supernatant was mixed with three volumes of 99% ethanol and the mixture was kept at 4°C for 30 min. To determine the dry weights of EPS, the precipitated EPS was collected by centrifugation and dried at 55°C overnight prior to measurement. Three replicates were used for each strain and the test was repeated three times.

**MOTILITY ASSAY**

Swimming and swarming motilities were examined on NB containing 0.3% (w/v) and 0.7% (w/v) agar (Difco, Franklin Lakes, NJ, USA), respectively. Overnight cultures of bacteria grown in NB were centrifuged at 8000 × g for 2 min at 4°C, washed twice with ddH2O, and resuspended in ddH2O (OD600 = 1.0). Two microliters of the suspension were spotted onto NA plates (diameter, 90 mm; containing 20 mL of NA) and incubated at 28°C. The migration zones were measured, and used to evaluate the motility of Xac cells (Li and Wang, 2011; Addy et al., 2012). For twitching motility assays, overnight bacterial culture in NB were centrifuged at 8000 × g for 2 min at 4°C, washed twice with ddH2O, resuspended in ddH2O (OD600 = 1.0), and spotted on minimal medium (MM) plates (Addy et al., 2012). Plates were incubated at 28°C, and the morphology of the colony edge was observed under a light microscope (100× magnification).

**PATHOGENICITY ASSAY**

After careful washing with tap water, immature fully expanded lemon leaves were sterilized by soaking for 2 min in sodium hypochlorite, followed by rinsing in sterilized water. Leaves were placed on the surface of filter paper with abaxial surfaces facing upwards. Lemon leaves were inoculated with bacterial suspension of Xac phage-uninfected and phage-infected strains (108 cfu/mL in sterile water) using an infiltration or a needle prick method. The infiltration method was conducted by pushing a needleless syringe containing the bacterial suspension against the surface of a citrus leaf supported by a finger on the opposite side of the leaf. The treated areas were immediately marked following inoculation (Chen et al., 2012). Needle-prick inoculation was performed by pricking the leaves, and droplets (10 μL) of bacterial suspensions were applied to each inoculation site. In both methods, the inoculated leaves were covered with a plastic bag for 48 h to facilitate the infection. Leaves were incubated in a growth chamber at 28°C with a photoperiod of 12 h light and 12 h dark for 4 weeks (Vernière et al., 1998; Li and Wang, 2011; Malamud et al., 2012).

**PHAGE STABILITY TEST**

We used Xac strain MAFF301080 because it was free from a XacF1 sequence in the genome. After infection with XacF1, a single colony was isolated and confirmed for its production of phage particles, the presence of XacF1 DNA in the cells by miniprep, and no integration of XacF1 DNA in the chromosome by PCR with a primer set of chromosomal sequences, 5′-ACT CGC TTT
GCA TGA AAT TCG CTA GCG AT-3’ (forward) and 5’-TTC GAT GGT CAC GGT GCC TGT AGT AGA GGC (reverse). After cultivation in NB at 28°C for several generations, random colonies spread on NA plates were picked and subjected to plaque assay, miniprep for XacF1 DNA, and PCR to detect lysogeny with the same primers as above.

Nucleotide sequence accession number

The sequence of the XacF1 genome has been deposited in the DDBJ under accession no. AB910602.

RESULTS

ISOLATION, MORPHOLOGY, AND HOST RANGE OF XacF1

A total of 20 phages were isolated from soil samples collected from citrus fields in Japan using a plaque assay on Xac strains (see Experimental Procedures), one of which formed small and turbid plaques (designated XacF1). A single plaque of this phage was picked for propagation, purification, and further experiments. Electron micrographs using highly purified phage particles (10^{13} pfu/mL) showed that XacF1 virions have typical filamentous phage features, with a long fibrous shape approximately 600 nm in length (Figure 1A). To determine the host range of the phage, Xac strains infecting different citrus species were tested for phage susceptibility (Table 1). The host range of the XacF1 phage was relatively wide, infecting 7 out of 11 Xac strains tested in this study (Table 1).

NUCLEOTIDE SEQUENCE AND GENOMIC ORGANIZATION OF XacF1

The genomic DNA of XacF1 was obtained as a replicative form of the XacF1 genomic DNA. Lanes: 1, undigested XacF1 DNA (RF); 2, digested with S1 nuclease; 3, cII; 4, EcoRI digestion produced a single band corresponding to approximately 7.3 kb on an agarose gel (Figure 1B, lane 4). The genomic DNA isolated from phage particles was completely digested by S1 nuclease treatment (data not shown), suggesting that the XacF1 genome is a circular single stranded DNA, like those of all other filamentous phages.

To determine the entire nucleotide sequence of XacF1, DNA was shotgun-sequenced. The results showed that the complete genome was 7325 nucleotides long, with a G+C content of 57.8%, which was significantly lower than that of the host genome (i.e., 64.7% for strain 306, accession no. NC_003919). There were 13 putative open reading frames (ORFs), of which 11 were located on the same strand and two were on the opposite strand (Table 2 and Figure 2). When databases were searched for sequences homologous to the XacF1 DNA sequence using BLAST and BLASTX programs, nine ORFs showed high similarity to ORFs previously reported for other filamentous phages, especially to ORFs of X. campestris pv. citri phage Cf1c (Kuo et al., 1991) (accession no. NC_001396), X. campestris pv. vesicatoria Cf1 phage (YP_364205.1), and X. campestris pv. campestris phi-Lf phage (X70328) (Table 2). XacF1 ORFs could be arranged in a similar modular structure to that of previously characterized filamentous phages of the Ff group (Model and Russel, 1988; Marvin, 1998), as shown in Figure 2. Within the putative replication module (Figure 2), we identified ORF1 and ORF2. The peptide encoded by ORF1 was homologous to filamentous phage phl replication initiation protein II (98% amino acid sequence identity) (Table 2). This gene encodes the pII protein, which is necessary for rolling-circle replication of phage genomes (Model and Russel, 1988). The deduced amino acid sequence encoded by ORF2 was homologous to peptides that mapped at the same position as the ssDNA binding protein (gV gene) of Ff phages, and its size was similar to that of this binding protein (Figure 2 and Table 2). Within the putative structural module of XacF1, we predicted five ORFs. ORF3 showed similarity to a hypothetical Xanthomonas protein (Table 2), with 32% amino acid sequence identity to a transmembrane motif (WP_005416529), supporting the hypothesis that ORF3 belongs to the module of structural genes (Figure 2). Moreover, ORF4, ORF5, and ORF7 (Figure 2 and Table 2) were the same size and in the same position as genes encoding the coat proteins of Ff phages. Another possible ORF included in this module was ORF6 (with similarity to coat protein Cf1c phage cp3, Kuo et al., 1991), which was similar in both size and location to gIII of the Ff phage. gIII encodes pIII, a minor coat protein that recognizes and interacts with receptors and coreceptors on the host cells (Armstrong et al., 1981; Lubkowski et al., 1999; Heilpern and Waldor, 2003) (Figure 2 and Table 2). It also showed 28% amino acid sequence identity to phage adsorption protein of Xanthomonas citri subsp. citri (YP_007649573). Therefore, ORF6 could be a homolog of gIII in XacF1. In the third putative module of XacF1, the assembly module, we found that ORF8 showed the highest homology to the cp4 protein of Cf1c phage (Figure 2 and Table 2), and to the zot protein of Xanthomonas vesicatoria (WP_005997731), with 59% amino acid sequence identity. Also, based on its size and position, it seems that ORF8 is a homolog of pl. XacF1 does not encode a pl homolog, hence like many filamentous phages it must use a host encoded pl homolog, outer membrane protein of the secretin family. Interestingly, we found that ORF12 might encode a regulator gene similar to those found in several filamentous phages, because amino acids encoded by this ORF exhibited similarity
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Table 2 | Predicted ORFs found in the XacF1 genome.

| Coding sequence | Strand | Position 5′-3′ | GC content (%) | Length of protein (aa) | Molecular mass (Kda) | Amino acid sequence identity/similarity to best homologs (no. of amino acid identical; % identity) | E-value Accession no. |
|-----------------|--------|---------------|----------------|-----------------------|---------------------|--------------------------------------------------------------------------------------------------|----------------------|
| ORF1            | +      | 1–1080        | 57.7           | 360                   | 40.5                | Filamentous phage phiL replication initiation protein II (340; 98%)                                | 0.0 0.005637352      |
| ORF2            | +      | 1077–1373     | 57.3           | 99                    | 9.1                 | V protein Xanthomonas phage Cf1c (73; 99)                                                        | 2e-46 0.005637352    |
| ORF3            | +      | 1405–1605     | 51.4           | 67                    | 7.2                 | Hypothetical protein-Xanthomonas (65; 98)                                                         | 2e-40 0.013787278    |
| ORF4            | +      | 1611–1868     | 60.5           | 62                    | 8.4                 | B coat protein-Xanthomonas phage Cf1c (62; 100)                                                  | 6e-33 0.005637352    |
| ORF5            | +      | 1928–1995     | 55             | 23                    | 5.9                 | No significant similarity                                                                        | –                    |
| ORF6            | +      | 1996–3474     | 55.8           | 493                   | 51.7                | A coat protein-Xanthomonas phage Cf1c (383; 96)                                                  | 0.0 0.005637352      |
| ORF7            | +      | 3474–3791     | 54.2           | 106                   | 11.5                | Hypothetical protein-Xanthomonas campestris (103; 98)                                           | 5e-67 0.013787278    |
| ORF8            | +      | 3788–4954     | 59.0           | 389                   | 42.8                | Hypothetical protein Cf1cp4- Xanthomonas phage Cf1c (388; 100)                                   | 0.0 0.005637352      |
| ORF9            | +      | 4954–5601     | 59.4           | 216                   | 23.5                | Hypothetical protein Cf1cp5- Xanthomonas phage Cf1c (214; 99)                                    | 4e-148 0.013787278   |
| ORF10           | +      | 5617–6009     | 56.6           | 131                   | 14.4                | Hypothetical protein Cf1cp6- Xanthomonas phage Cf1c (130; 100)                                   | 4e-88 0.013787278    |
| ORF11           | −      | 6047–6430     | 59.7           | 128                   | 14.4                | Hypothetical protein Cf1cp7- Xanthomonas phage Cf1c (127; 100)                                   | 1e-86 0.013787278    |
| ORF12           | −      | 6427–8885     | 57.6           | 153                   | 16.4                | - Filamentous phage Cf1 protein-Xanthomonas campestris pv. vesicatoria str. 85-10 (146; 90)       | 3e-88 0.013787278    |
|                 |        |               |                |                       |                     | - 18.2K protein-Xanthomonas phage Cf1c (164; 99)                                                  | 5e-113 0.013787278   |
| ORF13           | +      | 7015–7203     | 54.8           | 63                    | 6.8                 | Hypothetical protein- Xanthomonas axonopodis (59, 95)                                            | 6e-32 0.013787278    |

FIGURE 2 | Genomic organization of bacteriophage XacF1. Linear genomic maps of E. coli phage M13 (A), XacF1 (B), and Cf1c (C) are compared. Arrows oriented in the direction of transcription represent ORFs or genes. The functional modules for replication (R), structure (S), and assembly-secretion (A-S) are indicated according to the M13 model (Marvin, 1998). Map for Cf1c was drawn according to the genomic sequence (accession no. NC_001396, Kuoy et al., 1991). ORF sizes (in amino acids) are in parentheses. IG (intergenic region), and attP are also shown.

to several putative transcriptional regulators and DNA-binding helix-turn-helix proteins of phages (e.g., Cp8 of X. campestris pv. citri phage Cf1c (99% amino acid identity) (Shieh et al., 1991); phage repressor of Vibrio parahaemolyticus V-223/04, exhibiting 45% amino acid identity, EVU16279, E-value = 0.71). ORFs 9, 10, 11, and 13 had homology to hypothetical proteins of phages and bacteria, but did not appear to belong to any of the previously described modules.

XacF1 USES HOST XerCD RECOMBINASES TO INTEGRATE INTO THE XANTHOMONAS GENOME

Homology searches of the DDBJ/EMBL/GenBank databases for the XacF1 sequence revealed similar sequences in the genomes of some Xanthomonas species. This result suggested possible integration of this kind of phage into the host genome. To test this possibility, we performed genomic Southern blot analysis of 11 strains of Xac using a XacF1 DNA probe. The results, shown in Figure 3A, indicated that eight of the 11 strains contained hybridizing bands and, among them, seven strains showed similar hybridization patterns with variations in signal intensity. Therefore, XacF1 likely has a lysogenic cycle and integrates frequently into the host genome. Regarding the integration mechanism of XacF1, we could not find any genes or ORFs that encode a phage integrase in the genome (Table 2). In several cases, involvement of the host recombination system by XerC/D in integration of filamentous phages into host genomes has been established, including Vibrio cholerae phage CTXφ (Huber and Waldor, 2002; Das et al., 2011). In CTXφ integration, the dif site of the host genome (attB) forms a recombination complex with dif-like sequences on the phage genome (attP) (Val et al., 2005). We therefore looked for a possible dif-like sequence for attP on the XacF1 sequence and found a 15-bp dif core sequence of
5′-TAT ACA TTA TGC GAA (XacF1 positions 6504–6518). This sequence showed a high degree of homology to attP sequences of phages Cf1c (accession no. NC_001396) (Kuo et al., 1991), Cf16-v1 (M23621), φLt (X70328) phages of _X. campestris_. The attP core sequences are shown in red. Changed nucleotides are shown in blue. (C) Alignment of att sequences in the XacF1 integration system. The attL and attR sequences were determined for a lysogenic strain newly established with Xac MAFF673010 infected with XacF1. The attB sequence (dif) of _X. axonopodis pv. citri_ 306 was obtained from the genome database (accession no. AE008923.1, Jalan et al., 2011). Common 15 bases are shown in red and chromosomal sequences are in blue. (D) Putative XerC, XerD binding sites of the XacF1 system are compared with those of _Vibrio cholerae_ dif.

**FIGURE 3** Site specific integration of XacF1 DNA into the chromosome of Xac. (A) Southern blot hybridization analysis showing integration of the XacF1 phage into Xac host chromosomes. Genomic DNA of _X. axonopodis pv. citri_ strains was digested with HinII and hybridized with a probe (the entire XacF1 phage genomic DNA digested with EcoRI). (B) The attP sequence indicated for XacF1 is compared with attP sequences (putative in some cases) of filamentous phages, including Cf1c (accession no. NC_001396), Cf16-v1 (M23621), and Lt (X70328) phages of _X. campestris_. The dif core sequences are shown in red. Changed nucleotides are shown in blue. (C) Alignment of att sequences in the XacF1 integration system. The attL and attR sequences were determined for a lysogenic strain newly established with Xac MAFF673010 infected with XacF1. The attB sequence (dif) of _X. axonopodis pv. citri_ 306 was obtained from the genome database (accession no. AE008923.1, Jalan et al., 2011). Common 15 bases are shown in red and chromosomal sequences are in blue. (D) Putative XerC, XerD binding sites of the XacF1 system are compared with those of _Vibrio cholerae_ dif.

**FIGURE 4** (A) Effects of XacF1 infection on the growth of _X. axonopodis pv. citri_ (MAFF301080). (B) Effects of XacF1 infection on EPS production. The pellet of uninfected Xac MAFF301080 cells (right) was yellow, while the pellet of XacF1-infected cells (left) was white, indicating a defect in xanthan production.

**EFFECTS OF XacF1 INFECTION ON THE GROWTH RATE OF _X. AXONOPODIS PV. CITRI_**

Unlike other bacterial viruses, the Ff phages do not kill their hosts, but establish a persistent coexistence in which new virions are continually released (Model and Russel, 1988). Because of this non-lytic mode of viral replication, it is possible to grow high-titer cultures of the virus. Similarly, infection by XacF1 did not cause lysis of host cells, but established a persistent association between the host and phage, releasing phage particles from the growing host cells. Although cells infected with XacF1 could continue to grow and divide indefinitely, the process caused the infected cells to grow at a significantly lower rate than uninfected cells (Figure 4A).

**EFFECT OF XacF1 INFECTION ON HOST EPS PRODUCTION**

EPS production was compared between uninfected and XacF1-infected cells of strain MAFF301080. The XacF1-infected cells used in this experiment were confirmed to be free from prophage by plaque assay of the culture supernatant, Southern hybridization (Figure 3A), and PCR. The amount of EPS produced by the infected cells was significantly lower than that of the wild-type cells. We observed that following centrifugation, the culture pellets of the infected cells turned white, reflecting a low production of xanthan, which is the major component of EPS and is responsible for the yellow color of _Xanthomonas_ culture in the media (Figure 4B). Our prediction was confirmed by an EPS quantitation assay, which showed that the XacF1-infected cells had significantly lower EPS production (0.6 mg/10^10 cfu) than uninfected cells (3.35 mg/10^10 cfu).

**EFFECT OF XacF1 INFECTION ON HOST MOTILITY**

Swimming, swarming, and twitching motilities of uninfected and XacF1-infected cells of strain MAFF301080 were compared. A significant reduction in swimming and swarming motility was observed in XacF1-infected cells (Figures 5A–D). When visualized with a microscope, the colony margin of uninfected cells had a highly irregular shape, indicating profuse twitching motility, whereas the colony edge of XacF1-infected cells was smooth (Figure 5E), suggesting a decrease or loss of twitching motility. Because twitching motility is the surface movement associated with type IV pili (Marques et al., 2002; Meng et al., 2005),
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**FIGURE 5** | Impact of XacF1 infection on the motility of Xac MAFF301080 cells. Two microliters of bacterial solution [10^8 colony forming units (CFU)/mL] were inoculated in the swimming assay [0.3% (w/v) agar] (A,B), swarming assay [0.7% (w/v) agar] (C,D), and twitching motility assay (minimal agar medium) (E). The movement of bacterial cells was photographed 5 and 8 days post-inoculation (dpi) on the swimming and swarming plates, respectively. Twitching motility of bacteria was observed under a microscope 5 dpi on the twitching plates.

XacF1 infection may have affected the type IV pilus structures and/or functions of the host cells. We examined whether cell surface structural components were affected by XacF1 infection. Cell surface structure proteins were prepared by passing bacterial cells through a hypodermic needle, separated by SDS-PAGE, and compared between XacF1-infected and uninfected cells. XacF1-infected cells had considerably decreased levels of PilA, the major component of type IV pili, and decreased levels of FilC, flagellin (Supplemental Figure S1).

**EFFECTS OF XacF1 INFECTION ON VIRULENCE OF X. AXONOPODIS PV. CITRI**

Wild-type cells of strain MAFF301080 caused infection symptoms 4 days post-infection, and formed clear cankers 1 week post-inoculation (**Figure 6A**). Starting from 2 weeks post-infection, the lesion became brown in color and its center became raised and spongy or corky, typical canker symptoms (Graham et al., 2004) that reflected the aggressive virulence of this strain. In contrast, the symptoms of XacF1-infected MAFF301080 cells were relatively weak, and no mature canker symptoms were observed up to 4 weeks post-infection, except for marginal lesions formed around the pricking site (**Figure 6A**). To be more precise, we measured lesion size (**Figure 6B**), which showed that in uninfected cells, the lesions were large with a smooth center, spongy raised top, and their distribution around the infected area reached more than 6.5 mm in width 4 weeks post-infection. In contrast, the lesions formed by XacF1-infected cells remained weak and dry, and they did not expand more than 1 mm in width. Another inoculation method, in which we infiltrated the bacterial suspension into the lemon leaves, showed that XacF1-uninfected MAFF301080 cells incurred water soaking at the inoculation site 3 days post-infection, and then an erumpent tissue reaction was obvious 1 week after inoculation. The erumpent tissue expanded to an aggressive canker area on both sides of the leaf, and then
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**Figure 6** | Lesions on detached lemon leaves inoculated with cells of Xac MAFF301080. (A) Canker symptoms that had developed on leaves 1, 2, 3, and 4 weeks post-infection by the needle-pricking method. Leaves were inoculated with uninfected cells (upper panels) or XacF1-infected cells (lower panels). Leaf areas shown by a square were examined by photomicroscopy and the microscopic images are shown under each corresponding leaf. Characteristic canker lesions occurred with uninfected cells, while no obvious cankers developed on XacF1-infected cells. (B) Comparison of the size of lesions formed on lemon leaves. (C) Lesions formed on lemon leaves by infiltration of bacterial cells. Uninfected MAFF301080 cells were applied to two areas of the leaf (left half of the abaxial side), and XacF1-infected cells were similarly applied to the right side (upper panels). Lesions on the axial side are also shown in lower panels. Lesions on both lower and upper surfaces of leaves inoculated with the uninfected cells showed severe symptoms, expanding with time. No lesions formed on either surface of the leaves infected with XacF1-infected cells.

The lesions became dark and decayed with a yellow halo at the inoculation site 4 weeks post-infection (Figure 6C). However, in XacF1-infected cells, a slight water-soaking area on the leaf surface was only visible 2 weeks after inoculation, and weak canker symptoms could be seen 4 weeks post-infection. In all cases, leaves inoculated with ddH2O showed no canker symptoms.

**Discussion**

In this study, we isolated and characterized a filamentous phage, named XacF1, that infects X. axonopodis pv. citri strains. The isolated phage had a relatively wide range of host bacterial strains. Of particular interest, this study showed that along with the phage infection, the infected cells had decreased ability to form citrus cankers and a loss of virulence. We demonstrated that the canker symptoms of XacF1-infected lemon leaves were dramatically mitigated up to 4 weeks post-infection using both pricking and infiltration methods of inoculation (Figures 6A–C). The significant reduction in EPS (xanthan) production caused by XacF1 phage could be one of the reasons for such a dramatic decrease in canker formation. Virulence of numerous phytopathogenic bacteria, particularly various Xanthomonas species, is correlated with their ability to produce EPS (Dolph et al., 1988; Bellemann and Geider, 1992; Kao et al., 1992; Chou et al., 1997; Katzen et al., 1998; Dharmapuri and Sonti, 1999; Yu et al., 1999; Kemp et al., 2004). The multiple functions of EPS in virulence include protection of bacteria from toxic plant compounds, reduction...
of bacterial contact with plant cells to minimize host defense responses, promotion of bacterial multiplication by prolonging water soaking of tissues, and supporting invasion or systemic colonization of bacterial cells (Denny, 1999). Another possible role of EPS is to confer epiphytic fitness. It was previously suggested that EPS functions during both epiphytic and pathogenic phases of infection in *X. campestris pv. campestris* (Poplawsky and Chun, 1998; Rigano et al., 2007). As significant differences in virulence were observed between wild-type and xanthan-deficient mutant strains of other *Xanthomonas* species, Dunger et al. (2007) proposed that in citrus canker, xanthan supports epiphytic survival in citrus canker, but is not required for colonizing nearby tissue. Without xanthan the bacteria were unable to retain water and could not withstand abiotic stress and, thus, could not survive on the leaf surface. Therefore, xanthan works in two ways: to enhance bacterial virulence and to block the host defense. The drastic reduction in host EPS production caused by XacF1 infection may explain why the XacF1-infected cells showed dramatically decreased virulence.

Another major finding is the significant reduction in the swimming, swarming, and twitching motilities of Xac cells following infection by XacF1. Bacteria use a variety of motility mechanisms to colonize host tissues. These mechanisms include flagella-dependent swimming and swarming for movement in liquid surfaces, and flagella-independent twitching, gliding, and sliding for movement on solid surfaces (O’Toole and Kolter, 1998; Mattick, 2002; Harshey, 2003). Recent reports propose that bacterial adhesion and motility are required in the initial stages of Xac biofilm formation, whereas lipopolysaccharide and EPS play important roles in the establishment of mature biofilms (Li and Wang, 2011). The reduction in motility of XacF1-infected cells may be because filamentous phages such as XacF1 assemble on the host cell membrane and protrude from the cell surface, and so the nature of the host cell surface may change drastically during phage production (Addy et al., 2012). As shown in Supplemental Figure S1, XacF1-infected cells had considerably decreased levels of PilA, the major component of type IV pili.

Frequent protrusion of XacF1 particles from the infected cell surface may somehow compete with the formation of type four pili (Tfp). As reported by Kang et al. (2002), Tfp is responsible for twitching motility and adherence to multiple surfaces and is required for virulence. Interestingly, ORF 9 of the XacF1 phage (Table 2) showed significant homology to a TraX family protein (H8P1E6, E-value = 1e-70) and a putative F pilin acetylation protein (Q3B7T0, E-value = 4e-70), involved in pilus modification. Therefore, the loss of virulence in the XacF1-infected cells seems to be, at least partly, caused by the reduction or modification of Tfp formation and decrease in swimming, swarming, and twitching motilities.

Several works have described the use of phages for control of bacterial citrus canker caused by *X. campestris pv. citri* (Balogh et al., 2010). In those cases, the bacteriophages used for foliar plant diseases interacted with the target bacteria on the leaf surface, the phylloplane. The phylloplane is a constantly changing environment; there are changes in temperature, sunlight irradiation, leaf moisture, relative humidity, osmotic pressure, pH, microbial flora, and, in the case of agricultural plants, chemical compounds (Jones et al., 2012). These factors may be harmful to bacteriophages to varying extents. Sunlight irradiation, especially in the UVA and -B range, is mainly responsible for eliminating bacteriophages within hours of application (Jones et al., 2012). To avoid quick inactivation of XacF1, we propose the application of XacF1-infected cells instead of XacF1 phage alone. The XacF1-infected cells can grow and continue to produce infectious phage, so the XacF1 phage may serve as an efficient long-lasting tool to control citrus canker by decreasing the virulence of the pathogen. Concerning the stability of XacF1-infected cells, we observed relatively high stability of “a free phage state” in the infected cells. After several bacterial generations, 100% cells contained XacF1 and more than 70% of them were at the state of producing free phages without integration into the host chromosome (confirmed by PCR) (data not shown). Even if once prophage states were established, we observed frequent spontaneous excision and production of phage particles.

Another possible way to use XacF1 for biological control may be given as a phage cocktail with other lytic phages, such as Cp1 and Cp2, originating from Japan, which can infect more than 97% of Xac strains and was recently characterized by Ahmad et al. (2014).

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**SUPPLEMENTARY MATERIAL**
The Supplementary Material for this article can be found online at: http://www.frontiersin.org/journal/10.3389/fmicb.2014.00321/abstract

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