Flp, a Fis-like protein, contributes to the regulation of type III secretion and virulence processes in the phytopathogen Xanthomonas campestris pv. campestris

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SUMMARY

The ability of the plant pathogen Xanthomonas campestris pv. campestris (Xcc) to cause disease is dependent on its ability to adapt quickly to the host environment during infection. Like most bacterial pathogens, Xcc has evolved complex regulatory networks that ensure expression and regulation of their virulence genes. Here, we describe the identification and characterization of a Fis-like protein (named Flp), which plays an important role in virulence and type III secretion system (T3SS) gene expression in Xcc. Deletion of flp caused reduced virulence and hypersensitive response (HR) induction of Xcc and alterations in stress tolerance. Global transcriptome analyses revealed the Flp had a broad regulatory role and that most T3SS HR and pathogenicity (hrp) genes were down-regulated in the flp mutant. β-glucuronidase activity assays implied that Flp regulates the expression of hrp genes via controlling the expression of hrpX. More assays confirmed that Flp binds to the promoter of hrpX and affected the transcription of hrpX directly. Interestingly, the constitutive expression of hrpX in the flp mutant restored the HR phenotype but not full virulence. Taken together, the findings describe the unrecognized regulatory role of Flp protein that controls hrp gene expression and pathogenesis in Xcc.

Keywords: Fis-like protein, hrpX promoter, type III secretion system, virulence, Xanthomonas.

INTRODUCTION

Xanthomonads are Gram-negative bacteria that are known to cause disease in a range of important crops worldwide. Xanthomonas campestris pv. campestris (Xcc) is one of the best studied of these as it is the causative agent of black rot disease in crucifers, which include many important vegetable brassica crops (Swings and Civero, 1993; Vicente and Holub, 2013). Xcc is also an important model for the study of microbe–plant interactions because of its genetic tractability and cultivability. For these reasons, the study of Xcc has provided a lot of insight into how plant pathogens can adapt to the host environment and cause disease. Studies have revealed many mechanisms that are important for disease and environmental adaptation, including extracellular enzymes (protease, mannanase, etc.), extracellular polysaccharides (EPS), diffusible signal factor (DSF)-dependent cell–cell signalling, and proteins secreted by the type II secretion system (T2SS), type III secretion system (T3SS), type IV secretion system (T4SS) and more recently type VI secretion system (T6SS) (He and Zhang, 2008; Büttner and Bonas, 2010; Ryan et al., 2011, 2015; Notti and Stebbins, 2016; Zhou et al., 2017). In Xcc, one of the best-studied mechanisms that contributes to virulence is the T3SS apparatus. This complex system is encoded by over 20 hypersensitive response and pathogenicity (hrp) genes that when silenced lead to the loss of Xcc’s ability to cause full virulence and hypersensitive response (HR) induction (Alfano and Collmer, 1997; Lindgren, 1997). The activation of hrp genes, as well as some genes that encode secreted effector proteins, is controlled by two main regulators: HrpG (OmpR family regulator) and HrpX (AraC-type transcriptional activator) (Huang et al., 2009; Koebnik et al., 2006; Wengelnik et al., 1996; Wengelnik and Bonas, 1996).

Several other proteins have been shown to control the expression of hrp genes. HpaS, a sensor kinase that putatively constitutes a two-component signal transduction system with HrpG, positively regulates the expression of hrp genes (Li et al., 2014). The zinc uptake regulator (Zur), a key regulator of zinc homeostasis belonging to the Fur family of transcription factors, positively regulates the hrp gene expression via HrpX (Huang et al., 2009). HpaR1 (hrp-associated regulator), a global regulator belonging to the YtrA subfamily of the GntR family, appears to indirectly regulate the expression of hrp genes via HrpG (An et al., 2011). In addition, HpaP, a novel regulatory protein with ATPase and
phosphatase activity, regulates the expression of *hrp* genes by controlling the expression of *hrpX*. However, this is unlikely to be by a direct action (Cui et al., 2018).

Despite these advances, considerable work is still required to understand the regulatory mechanisms associated with control of T3SS and virulence in this important bacterial plant pathogen. In the present study, we report the identification and characterization of a Fis-like protein (Flp), a previously unreported protein regulator of *hrp* and other virulence determinants in *Xcc*. We demonstrate that this protein contributes to virulence, HR induction and a series of other cellular functions in *Xcc*. We present evidence that Flp is likely to regulate the expression of *hrp* genes by controlling the expression of *hrpX* via directly binding to its promoter.

**RESULTS**

**Flp is important for pathogenicity in *Xcc***

In our earlier work aimed at establishing a general view of the factors that contribute to pathogenicity in *Xcc*, we screened a series of mutants which were constructed using a suicide vector (pK18mob) strategy (Windgassen et al., 2000). Virulence assays using the host plant cabbage (*Brassica oleracea*) showed that one of these mutants, 0520nk (Table S1), caused weaker symptoms of disease when compared with the wild-type *Xcc* strain. The 0520nk mutant had a disrupted gene which corresponded to open reading frame (ORF) *XC_0520* in the genome of *Xcc* strain 8004 (accession number CP000050). The *XC_0520* gene encodes a putative DNA-binding protein, with further analysis with the SMART (Simple Modular Architecture Research Tool) program (http://smart.embl-heidelberg.de) has 40% identity with the Fis (factor for inversion stimulation) protein that is found in many other bacteria, in-lberg.de) has 40% identity with the Fis (factor for inversion stimulation) protein that is found in many other bacteria, in-

**Flp influences the regulation of extracellular polysaccharide, motility, stress tolerance and extracellular enzymes production**

To explore if Flp manipulated specific functions that are known to be required for pathogenesis in *Xcc*, we conducted a series of basic phenotypic tests to assess extracellular polysaccharide (EPS) production, extracellular enzymes (cellulase and amylase), cell motility and the adaption to stress and antimicrobials.

The results showed that the *flp*-mutant strain Δflp displayed decreased EPS production (Fig. 2A) and motility (swimming and swarming, tested on 0.28% w/v agar plates and 0.6% w/v agar plates, respectively) (Fig. 2B). However, EPS production and motility of the CΔflp complemented strain were similar to the wild-type under the conditions tested (Fig. 2A,B).

When extracellular enzymes in the *Xcc* wild-type strain, Δflp mutant strain and CΔflp complemented strain were compared, positive differences were seen. The results show that the Δflp mutant exhibits a significant enhancement in cellulase and amylase secretion (*P* < 0.05 by Student’s *t*-test) (Fig. 2C). Moreover, enhancement in the activity of extracellular enzymes could be restored to wild-type levels in the CΔflp complemented strain (Fig. 2C).

The growth characteristics of the *Xcc* strains in liquid medium nutrient-yeast-glycerol (NYG) were also investigated. Results revealed that the *flp*-mutant Δflp displayed small changes in growth properties. The mutant had a reduced growth rate compared to that of the wild-type strain in the early exponential phase (Fig. 2D). However, the growth rate was recovered in the mid-exponential phase and the Δflp mutant grew with a rate exceeded that of the wild-type in the late exponential phase.

**Additional differences were seen when the wild-type strain, the Δflp mutant and the CΔflp complemented strain were assessed for their ability to adapt to environmental stresses (Fig. 3).** For these experiments, kill curve assays were used in which the quantity of living cells on agar plates were supplied with various different concentrations of environmental stresses, including sodium dodecyl sulphate (SDS), *H₂O₂*, NaCl, phenol and heavy metal salt CdCl₂ and CuSO₄ (see Experimental procedures). These tests demonstrated that compared to the wild-type, the Δflp mutant showed reduced survival in the presence of phenol, CdCl₂, CuSO₄ and SDS but not *H₂O₂* and NaCl (Fig. 3). Importantly, in all cases the complementation strain responded similarly to the wild-type (Fig. 3).

Taken together, these findings suggest that Flp regulates positively the EPS production, motility and some stress tolerance but appears to negatively regulate extracellular enzymes in *Xcc* under the conditions tested. Despite these observations, the mechanism of regulation by Flp in these cases remains enigmatic.
Flp has an influence over the expression of genes involved in virulence and various adaptation processes in Xcc

To gain a greater understanding of the regulatory role of Flp in Xcc a set of global gene expression profiles was generated using RNA-Seq. For this, selected Xcc strains were grown to the mid-exponential phase (OD$_{600}$ = 0.6) in XVM2, a medium that mimics more closely the nutrition environment of the plant (Astuá-Monge et al., 2005). Following bacterial RNA extraction, library construction and sequencing, differential gene expression analysis was conducted on the generated data (see Experimental procedures). Of the 4273 annotated genes from the genome of Xcc strain 8004, 279 genes presented differentially expressed (fold changes ≥ 2.0), with 121 genes up-regulated and 158 genes down-regulated according to the transcriptome data (Fig. 4A, Table S2). In order to verify the transcriptome data, 16 genes that showed changes were selected randomly and reverse transcription-polymerase chain reaction (RT-PCR) was performed to examine the relative expression levels of these genes. All selected genes showed expression changes that were comparable with the transcriptome data (Table 1).

Functional clustering analysis, according to the annotation of Xcc strain 8004 genome (Qian et al., 2005), was carried out where the majority of the 279 genes regulated by Flp were assigned to the functional categories ‘pathogenicity and adaption’, ‘cellular process’, ‘central intermediary metabolism’, ‘transport’, ‘energy and carbon metabolism’, ‘cell envelope and cell structure’, ‘regulatory functions’ and ‘signal transduction’. The remaining genes were predicted to encode hypothetical proteins or have not been given a functional category to date (Fig. 4A; Table S2). The most dominant functional categories which genes were assigned to were ‘pathogenicity and adaption’ and ‘cellular processes’ (Fig. 4A). Notably, the deletion of Flp had an impact on genes belonging to the type III secretion system (T3SS). The expression of 16 hrp genes, XC_3003 (hrcC), XC_3004 (hrcI), XC_3006 (hrn), XC_3007 (hrpB5), XC_3009 (hrcH), XC_3010 (hrpB2), XC_3012 (hrcU), XC_3013 (hrcV), XC_3015 (hrcQ), XC_3016 (hrcR), XC_3018 (hpA), XC_3019 (hrpD5), XC_3022 (hpAB), XC_3023 (hrpW), XC_3025 (hrpF) and XC_3076 (hrpX) was decreased in the Flp mutant compared to the wild-type (Fig. 4A, Table S2).

Given that mutation of Flp leads to a significant reduction in virulence, it is feasible that the impact of Flp on the expression of
pathogenicity related genes at the transcriptional level accounts for the phenotypes seen in the flp mutant.

**Flp regulates the expression of T3SS genes by altering the expression of key regulator HrpX**

The gene transcription profile data presented suggest that Flp regulated the expression of the T3SS via hrp gene expression. To confirm this idea, we quantified the expression of several hrp operons (hrpB, hrpC, hrpF, and hrpX) and hrpG, the key regulators of T3SS. This was achieved by using promoterless-gusA transcriptional fusion reporters that we have deployed in previous studies (An et al., 2011; Cui et al., 2018; Huang et al., 2009). Here we constructed a group of reporter plasmids for hrpB, hrpC, hrpF, hrpX and hrpG where the promoter sequence fused in front of the gusA gene so that the activity of gusA (Huang et al., 2009). The reporter plasmids were introduced into the wild-type and Δflp strains, respectively (Table S1). The reporter strains were cultured in XVM2 medium for 8 h, and the activities of β-glucuronidase GUS were determined (see Experimental procedures). The results demonstrate that GUS activities for hrpB, hrpC, hrpF, and hrpX reporters, but not the hrpG reporter, were significantly reduced in the Δflp deletion strain compared to the wild-type (P < 0.05 by Student’s t-test). Growth of the strains was recorded at intervals of 4 h.
Flp is required for stress tolerance in *Xanthomonas campestris pv. campestris* (Xcc). Survival experiments performed by subculturing strains overnight on fresh NYG agar plates supplemented with different concentrations of SDS (A), oxidant H$_2$O$_2$ (B), hyperosmosis NaCl (C), phenol (D) and heavy metal salts CdCl$_2$ (E) and CuSO$_4$ (F). The surviving bacterial colonies on the plates were counted after incubation for 3 days.
Flp is a global regulatory protein that affects the expression of a number of genes, including T3SS genes. (A) Functional categories of differential expressed genes in Δflp mutant. Genome-scale transcriptome profiling of Xanthomonas campestris pv. campestris (Xcc) strains cultured in XVM2 were investigated by RNA sequencing, and 279 genes were found differentially expressed by two-fold or more in Δflp mutant (Table S2). These genes were broadly categorized according to their biological function (He et al., 2007). Notably, due to cross-talk between different metabolism pathways, some genes might be counted into different categories simultaneously. (B) ß-glucuronidase (GUS) activity of hrp gene promoter-gusA reporters in Δflp mutation and wild-type backgrounds. Data shown are mean and standard deviation of triplicate measurements. The experiment was repeated twice, and similar results were obtained. (C) The expression levels of hrp genes in Xcc wild-type strain 8004, Δflp, CΔflp and Δflp/p3X as measured by semi-quantitative RT-PCR.

Flp enhances hrpX expression via binding to its promoter

The findings outlined above raise the question of how the Flp influences hrpX expression. One possible explanation is that Flp binds directly to the promoter of hrpX. To explore the potential interaction between Flp and hrpX promoter we conducted a set of electrophoretic mobility shift assays (EMSA) (see Experimental procedures). The 6 × His-tagged expression construct of Flp was generated and the protein subsequently purified by nickel affinity column chromatography (see Experimental procedures). The purified 6 × His-tagged Flp fusion protein caused a mobility shift of DNA probes spanning the promoter of hrpX [from +70 bp down-stream to –131 bp upstream against the transcription initiation site (TIS) with a pair of 5’ FAM-labelled primers]. The binding of
the hrpX promoter appeared to increase with increasing concentrations of Flp protein (scale from 0 to 1500 nM) to the EMSA assays (Fig. 5A). Furthermore, the shifted bands also could be competed by excess of the unlabelled probes (Fig. 5A). Taken together, the data suggests that under the conditions used Flp binds to the upstream region of hrpX.

Table 1  Confirmation of RNA-Seq gene expression data by semi-quantitative RT-PCR.

| ID     | Gene    | Description                                   | Expression level | Semi RT-PCR WT/Δflp |
|--------|---------|-----------------------------------------------|------------------|---------------------|
| XC_3001| hpa2    | Lysozyme-related protein Hpa2                  | 2.97↓            |                     |
| XC_2324|         | c-di-GMP phosphodiesterase A                   | 4.33↓            |                     |
| XC_3657| copB    | Copper resistance protein B                    | 2.58↓            |                     |
| XC_3129| pmrC    | Inner membrane protein                         | 3.59↓            |                     |
| XC_3597| hns     | DNA-binding protein                             | 3.03↓            |                     |
| XC_3437| lptD    | LPS-assembly protein                            | 2.29↓            |                     |
| XC_2004| avrXccC | Avirulence protein                              | 8.31↓            |                     |
| XC_3694| ompW    | Outer membrane protein                          | 2.59↓            |                     |
| XC_2827| hpaR    | MarR family transcriptional regulator          | 2.25↓            |                     |
| XC_0158|         | Xylosidase/arabinosidase                        | 2.51↓            |                     |
| XC_2659| gcd     | Quinoprotein glucose dehydrogenase             | 7.31↓            |                     |
| XC_1273| trkA    | Voltage-gated potassium channel                 | 2.36↓            |                     |
| XC_1314| lptC    | Lipopolysaccharide export system protein LptC   | 2.51↓            |                     |
| XC_3652| fabB    | β-ketoacyl-[ACP] synthase                       | 2.05↑            |                     |
| XC_0783| celS    | Cellulase S                                    | 2.51↑            |                     |
| 16S rRNA|         | Internal reference                             |                  |                     |

Note: 16 genes of the transcriptome data were chosen to validate the integral accuracy via semi-quantitative RT-PCR. RNA, extracted from the cultures of Xanthomonas campestris pv. campestris wild-type (8004) and Δflp, respectively, was reverted into cDNA, and then cDNA was used as the template in semi-quantitative PCR. In this study, false discovery rate (FDR) ≤ 0.05 and absolute value of log₂ fold change ≥ 1 were used as the cut-off values. The acquired results were accordant to the transcriptome data;↑, up-regulated; ↓, down-regulated.

The fact that Flp binds to the promoter of hrpX suggests that Flp may directly regulate the expression of the hrpX gene. To validate this, we performed an in vitro transcription assay. Template DNA fragments of 311 bp, extending from −131 to +179 relative to the TIS of the hrpX promoter, were incubated with RNA polymerase holoenzyme from Escherichia coli with increasing
Flp regulates HrpX by interacting with the promoter sequence of hrpX. (A) Electrophoretic mobility shift assays (EMSA) to examine the interaction between the hrpX promoter sequence and Flp protein. In this experiment, DNA is acquired from purified PCR products of the hrpX promoter sequence with 5′-FAM labelled primers. The DNA is then incubated with purified Flp protein (protein final density 0, 0.25, 0.5, 0.75, 1 and 1.5 μM) and the hrpX promoter DNA is 0.10 μmol). A competition assay was also conducted with unlabelled DNA. When the concentration of unlabelled DNA increased from 10 to 30 μM, the Flp–DNA complexity decreased, indicating the unlabelled DNA of hrpX brings competition to the 5′-FAM-labelled hrpX in forming DNA–protein complex with Flp. F, free DNA strips; R, retarded DNA strips. (B) In vitro transcription experiment using Flp protein (final density arranged from 0, 0.1, 0.20 and 0.5 μM) in the transcription system. A template DNA fragment containing the clp promoter (Liu et al., 2019) and a 121-bp hrpX fragment extending from +1 to +121 relative to the TIS were used as controls.

Fig. 5 Flp regulates HrpX by interacting with the promoter sequence of hrpX. (A) Electrophoretic mobility shift assays (EMSA) to examine the interaction between the hrpX promoter sequence and Flp protein. In this experiment, DNA is acquired from purified PCR products of the hrpX promoter sequence with 5′-FAM labelled primers. The DNA is then incubated with purified Flp protein (protein final density 0, 0.25, 0.5, 0.75, 1 and 1.5 μM) and the hrpX promoter DNA is 0.10 μmol). A competition assay was also conducted with unlabelled DNA. When the concentration of unlabelled DNA increased from 10 to 30 μM, the Flp–DNA complexity decreased, indicating the unlabelled DNA of hrpX brings competition to the 5′-FAM-labelled hrpX in forming DNA–protein complex with Flp. F, free DNA strips; R, retarded DNA strips. (B) In vitro transcription experiment using Flp protein (final density arranged from 0, 0.1, 0.20 and 0.5 μM) in the transcription system. A template DNA fragment containing the clp promoter (Liu et al., 2019) and a 121-bp hrpX fragment extending from +1 to +121 relative to the TIS were used as controls.

Flp regulates T3SS in planta by altering the expression of hrpX, which has an impact on the induction of HR and plant defence

Given that mutation of Flp leads to a reduction in the expression level of hrp genes, the influence of flp on the induction of HR was evaluated using an infiltration assay. Bacterial suspensions of the wild-type, Δflp, CΔflp and a hrpX mutant (negative control) were inoculated into the leaves of non-host pepper (Capsicum annuum cv. ECW-10R) through the use of a pressurized syringe (see Experimental procedures). Following 8 h inoculation, the Δflp strain resembled the negative control by showing no visible HR symptoms (Fig. 7A). Conversely, the wild-type and CΔflp strains induced comparable HR symptoms (Fig. 7A). It was not until after 16 h post-inoculation that Δflp strain appeared to induce any visual HR symptoms. These results suggest that the Δflp mutant had a delayed and weakened HR compared to the wild-type.

To provide a quantitative assessment of HR induction an electrolyte leakage assay was used (see Experimental procedures). Here, leaf tissues within the infiltration areas were collected at three time points (8, 16 and 24 h) post-inoculation of the Xcc strains. The results show that wild-type and CΔflp induced similar levels of electrolyte leakage, whereas Δflp and the hrpX mutant induced much lower levels at 8 and 16 h, compared to the wild-type (Fig. 7A). However, at 24 h post-inoculation, it appeared that Δflp generated a very similar to the wild-type and CΔflp strains but the hrpX mutant still retained lower levels (Fig. 7A).

In order to confirm Flp influence on T3SS in planta was due to modulation of hrpX expression, we tested an flp mutant that constitutively expressed hrpX in several HR induction assays. To achieve this, the entire ORF and Shine–Dalgarno (SD) sequence of hrpX gene was introduced into the pLARF3 plasmid (under the control of the lacZ promoter) to generate the p3X construct (Huang et al., 2009). This construct was introduced into the Δflp mutant to obtain the Δflp/p3X strain. Importantly, the Δflp/p3X strain constitutively expressing hrpX retained the ability to induce HR (Fig. 7A).

Callose deposition is required for disease resistance against many pathogens, including Xcc. It has been shown previously that Xcc induces defence responses in host plant Arabidopsis in a T3SS-dependent manner (Rong et al., 2010). Therefore, to examine potential differences in defence response we monitored callose deposition in the wild-type, Δflp, CΔflp, Δflp/p3X and hrpX mutant strains. As before, we inoculated wild-type, Δflp, CΔflp, Δflp/p3X and hrpX strains into cabbage leaves using the
infiltration method (see Experimental procedures). Subsequently, leaves were collected at 20, 40 and 60 h post-infiltration and callose disposition was examined (see Experimental procedures). Like HR induction the wild-type, Δflp and Δflp/p3X showed similar callose disposition across the leaves tested (Fig. 7B). The Δflp showed reduced levels of callose disposition compared to the wild-type but the level, which was greater than what the hrpX produced, was restored to the wild-type by the constitutive expression of hrpX (Fig. 7B).

To determine if growth in planta contributes to the HR induction and plant defence phenotypes observed, we determined the growth of strains. We examined the growth of wild-type, Δflp, Δflp/p3X and hrpX strains following inoculation into cabbage by leaf-cutting methods and recorded the growth variation within 10 days post-inoculation (see Experimental procedures). The results show that the Δflp and Δflp/p3X have slower growth in planta compared to wild-type and CΔflp strains (Fig. 7C). Notably, a comparison between Δflp and Δflp/p3X growth profiles showed that the latter was slower initially in colonization, which might be due to the deregulation of HrpX in this strain. Conversely, the hrpX mutant grew at a similar growth rate to the wild-type and complementary strain CΔflp strains, indicating that HrpX is not initially required for host colonization.

These results together indicate that a constitutive expression of hrpX could restore T3SS, HR and host plant defence induction ability of the Δflp strain, which is consistent with the idea that Flp regulates T3SS by modulating the expression of hrpX in planta. Furthermore, Flp appears to be important for plant colonization.

**Flp influences previously uncharacterized virulence factors independently of HrpX regulation in Xcc**

The data presented above demonstrate that Flp regulates hrpX by directly interacting with the promoter and therefore has an impact on T3SS. However, results from the transcriptome profile and hrpX complementation of the Δflp strain suggest that Flp plays a greater role in virulence than just through regulation by HrpX/T3SS.

To confirm this idea we compared the transcriptome profile of the Δflp strain (generated in this study) and the hrpX mutant strain (generated in Jiang et al., 2018), which identified a set of target genes that appeared to be regulated by Flp and not HrpX. To confirm this we examined the expression of XC_1273, XC_2659, XC_2827, XC_3129, XC_3694 and XC_4002 by RT-PCR by extracting RNA from wild-type, Δflp, CΔflp and Δflp/p3X cultured in XVM2, respectively. The RT-PCR results confirmed that expression of these selected genes was consistent with the data from the transcriptome analyses and appeared to be under the regulation of Flp (Fig. 8A). Furthermore, the expression of these genes was restored to wild-type levels by in trans expression of flp in the Δflp mutant background (CΔflp) (Fig. 8A). Interestingly, the expression levels of genes (XC_1273, XC_2659, XC_3694, XC_4002) were not restored to wild-type levels in the Δflp mutant background expressing hrpX (Fig. 8A). This suggests that genes are under the control of Flp and not HrpX.

The role of the Flp-regulated genes in virulence to Chinese cabbage was tested by the use of a panel of insertional mutants (Table S1). The virulence of each mutant was tested by measurement of the lesion length after bacteria that were introduced into the vascular system of Chinese cabbage by leaf clippings. Mutation of three genes (XC_2827, XC_3129, XC_3694) gave a significant reduction in virulence (P < 0.05...
by Student’s t-test) (Fig. 8B). Nevertheless, the remaining genes (XC_1273, XC_2659, XC_4002) showed no difference to the wild-type strain (Fig. 8B). Despite a significant impact on virulence, none of the genes led to a complete loss of virulence. Many of these genes had not been previously associated with Xcc virulence.
This subset of new virulence factors for Xcc includes outer membrane protein (XC_3694) and inner membrane protein (XC_3129), both under the control of Flp. Interestingly, many of these proteins have homologues in other plant-associated bacteria, including other Xanthomonas species and Stenotrophomonas.

**DISCUSSION**

Fis protein was initially identified as a factor for inversion stimulation of the homologous Hin and Gin site-specific DNA recombinases of E. coli (Kahmann et al., 1985). Subsequently, its diverse roles have been described, including roles in regulating bacterial virulence factors and optimizing bacterial adaptation to various environments. Fis is an abundant bacterial nucleoid-associated protein that influences DNA topology by directly binding and bending DNA (Dillon and Dorman, 2001). It has been suggested that Fis alteration of DNA can occur in multiple tandem sites in a non-random fashion (Schneider et al., 2001; Kahramanoglu et al., 2011). Fis has been shown to display a preference for binding to regions upstream of ORFs which can influence gene expression (Kahramanoglu et al., 2011). In addition, Fis can also directly activate and repress transcription at promoters by interacting with RNA polymerase (Browning et al., 2010). Despite extensive studies showing that Fis serves as a global transcription factor that activates a diverse range of virulence functions, including quorum sensing, capsule production, adhesion and type III secretion in many mammalian pathogens, e.g. pathogenic E. coli (Falconi et al., 2001; Goldberg et al., 2001), Shigella flexneri (Falconi et al., 2001), Salmonella enterica serovar Typhimurium (Kelly et al., 2004) and Yersinia pseudotuberculosis (Green et al., 2016), few works have been carried out in plant bacterial pathogens. Additionally, no such protein has been identified or characterized in Xcc or other bacteria from the Xanthomonas genus. In this study, we identified the ORF XC_0520 from Xcc that encodes a small protein with 40% amino acid sequence identity to the Fis protein characterized in other bacterial species. This Fis-like protein (or Flp) is identical in all three sequenced Xcc strains (8004, ATCC33913 and B100). Here, the deletion of Flp in Xcc caused a series of changes in virulence and HR-related phenotypes. Although virulence factor regulation by Fis has been seen in other bacterial plant pathogens such as Dickeya dadantii (Ouafa et al., 2012) and Dickeya zeae (Lv et al., 2018), no role has been attributed to its regulation of HR, suggesting a different role in Xcc.

Our transcriptome analysis revealed that expression of T3SS genes, including master regulator HrpX, was down-regulated in a flp mutant. We therefore investigated the regulatory effect of Flp on hrp genes. Using GUS-based reporter plasmids, electrophoretic mobility shift and in vitro transcription assays, we demonstrated that Flp regulates the expression of hrp genes and T3SS by controlling the expression of hrrX directly. In Xanthomonas spp., the hrp genes are highly conserved and comprise more than 20 genes. The expression of hrp genes is mainly controlled by two key regulators, HrpG and HrpX. HrpG and HrpX form a regulatory cascade: HrpG regulates the expression of hrrX and HrpX then activates the expression of other hrp genes as well as some effector genes (Huang et al., 2009; Koebnik et al., 2006; Wengelnik et al., 1996; Wengelnik and Bonas, 1996). Besides HrpG and HrpX, several regulators have been identified as being involved in the regulation of the expression of hrp genes. However, these regulators and their regulatory mechanisms are distinct in different Xanthomonas species, e.g. a histone-like nucleoid-structuring (H-NS) protein XrvC and a LysR-type transcriptional regulator GamR directly control the transcription of both hrrG and hrrX in Xanthomonas oryzae pv. oryzae (Liu et al., 2016; Rashid et al., 2016), a T3SS component HrcT positively regulates the expression of hrrX via binding to its promoter in X. oryzae pv. oryzicola (Liu et al., 2014), and the zinc uptake regulator Zur and a GntR family transcriptional regulator HpaR1 indirectly control the expression of hrp genes via HrpX and HrpG, respectively, in Xcc (An et al., 2011; Huang et al., 2009). Here our experimental evidences suggest a novel regulatory pathway control T3SS in Xcc. However, our analysis cannot eliminate the possibility that Flp may interact with multiple ORFs to control the transcription of hrp genes. Furthermore, given that Flp functions as a global regulator, illustrated by our global transcriptional analysis, it is possible that Flp directly/indirectly regulates hrp expression via other avenues.

Our transcriptome analysis also demonstrated that Flp affects the expression of a series of other genes, such as flagellar and pilus biosynthesis, nutrition transport, stress response, amino acid and cofactor biosynthesis, galactose and starch metabolism. These genes are unlikely to be under the control of HrpX that contribute to virulence and other phenotypes such as motility and cellular stress response. This is supported by the fact that expression of HrpX in the Flp mutant cannot rescue virulence in planta (Fig. 1). Importantly, the transcriptome data, as well as RT-PCR, mutation and pathogenicity experiments, unveiled a range of previously uncharacterized proteins, e.g. XC_3694 (putative outer membrane protein) and XC_3129 (putative inner membrane protein), which are required for full virulence in Xcc. These virulence factors appear to be regulated by a Flp-dependent and no HrpX-dependent mechanism. However, whether Flp have directly regulatory effects on these proteins remains unknown. Given many of them have homologues in other plant pathogens, the roles of the novel virulence factors in the pathogenicity of Xcc merit further investigation.

Previous studies in various bacteria have shown that Fis plays a pleiotropic role in bacterial virulence (Duprey et al., 2014). Our data further illustrate this in Xcc as Flp has a role in regulating HrpX and T3SS but also in a range of previously uncharacterized virulence factors that appear to be regulated independently.
of HrpX. Additionally, we demonstrated that Flp is required for swimming and swarming motility, stress tolerance, and extracellular polysaccharide and extracellular enzymes (cellulase and amylase) production. Further studies are needed to examine the role of Flp in the regulation of these phenotypes but this suggests several other questions need to be addressed: What are the environmental cues that activate the expression and activity of Flp? What genes are directly regulated by Flp? Does Flp have a conserved binding site? Does Flp regulate a different set of genes during plant colonization? Does Flp modulate the activity of HrpG directly? How does Flp affect gene expression of the new genes identified as Xcc virulence?

Our work demonstrates the complexity of the signalling pathways involved in the regulation of virulence in Xcc and describes Flp, a Fis-like protein, an extensive regulator that controls hrp gene expression, HR induction and pathogenesis. The characterization of virulence protein regulators such as Flp is required to develop strategies for disease control in plant pathogens such as Xcc.

**EXPERIMENTAL PROCEDURES**

**Bacterial strains, plasmids and growth conditions**

The bacterial strains and plasmids used in this work are listed in Table S1. *Escherichia coli* strains were cultured in Luria Bertani medium (Miller, 1972) at 37 °C. Xcc strains were cultured in NYG medium (Daniels et al., 1984), NY medium (NYG medium but without glycerol), and the mimic medium XVM2 (Wengelnik and Bonas, 1996) at 28 °C and 200 rpm. Antibiotics were used according to the concentrations as required: kanamycin at 25 µg/mL, rifampicin at 50 µg/mL, ampicillin at 100 µg/mL, spectinomycin at 50 µg/mL and Tet at 5 µg/mL for Xcc strains and 15 µg/mL for *E. coli* strains.

**Nucleic acid manipulations**

The nucleic acid manipulations followed the procedures described by Sambrook et al. (1989). Conjugation between the Xcc and *E. coli* strains was performed as described by Turner et al. (1985). The restriction endonucleases, T4 DNA ligase and *Pfu* polymerase were provided by Promega (Shanghai, China). The total RNAs were extracted from the cultures of the Xcc strains with a total-RNA extraction kit (Invitrogen, Waltham, MA, USA) and cDNA generated using a cDNA synthesis kit (Invitrogen). These kits were used with reference to the manufacturer’s instructions. For semi-quantitative RT-PCR, the obtained cDNA was diluted and used as a template with selected primers for target genes (Table S3).

**Construction of mutant strains**

In order to construct the in-frame deleted mutant of *flp* (XC_0520), 333 bp (EcoRI and BamHI) of the upstream and 339 bp (BamHI and HindIII) of the downstream sequences of the *flp* gene were amplified by PCR using the relevant primers (Table S3). After being digested by restriction enzymes, these DNA fragments were fused with the suicide plasmid pK18mobsacB (Schäfer et al., 1994) and transformed into *E. coli* DH5α. The acquired recombinant plasmid was introduced into Xcc with the help of plasmid pRK2073 (Table S1). The original ORF of XC_0520 will be deleted from genomes through allelic exchange and homologous recombination.

To complement the *flp* deletion mutant, an 297-bp DNA fragment of the *flp* gene coding sequence was PCR-amplified from Xcc strain 8004 and inserted into the plLCFR3 vector at the BamHI/HindIII restriction site, creating the plasmid pLCfrp (Table S1). This plasmid was introduced into the mutant by triparental mating.

**Pathogenicity tests, HR assays, leakage assays and in-plant growth curve**

The virulence of the Xcc strain to the host cabbage plant (*Brassica oleracea ‘Jingfeng No. 1’*) was tested by the leaf-clipping method (Wang et al., 2007). Cabbage seedlings were planted and grown in the greenhouse for 30 days and the leaves were used for inoculation. Xcc strains, collected from overnight culture, were washed and adjusted to the same final density (OD_{600} = 0.6, approximately 1 × 10^8 CFU/mL). The bacterial resuspension was then diluted to 1 × 10^6 CFU/mL. The lesion and symptoms were measured 10 days post-inoculation (Dow et al., 2003).

HR was tested on pepper leaves (*Capsicum annuum cv. ECW-10R*) as previously described (Li et al., 2014). Briefly, bacteria suspensions (1 × 10^8 CFU/mL) were infiltrated into the abaxial side of the pepper leaves. These inoculated plants were kept in the greenhouse at 28 °C to observe the HR symptoms and gauge conductivity. For conductivity measurements, samples (leaf discs of 0.4 cm^2) were collected using a hole-puncher. These acquired samples were soaked in 10 mL ultrapure water with shaking at 200 rpm. The leaf discs were then removed and the conductivity of water was measured with a DDS-307A conductivity meter.

Bacterial in planta growth was estimated as previously described (Li et al., 2014). Briefly, Xcc strains were inoculated onto cabbage leaves using leaf clippings. At intervals of 24 h, four clipped leaves from every group of inoculated plants were collected and homogenized. Homogenate was serially diluted using NYG medium and 100 µL dilution was used for spread plate counting (NYG medium). The amounts of bacteria were calculated after 2 days of incubation.

**Calllose deposition assay**

Calllose deposition assay was performed on cabbage leaves as recently described by Hamdoun et al. (2018). Thirty-day-old cabbage leaves were inoculated with Xcc strains by infiltration (detailed above). At intervals of 20 h, the leaves were collected and individually soaked in 70% ethanol for 2 h, then 50% ethanol for...
2 h and finally sterile distilled water for 2 h. When the chlorophyll was completely leached, the leaves were stained in 0.01% aniline blue. Callose deposition was observed using an Olympus BH-2 epifluorescent microscope.

**GUS activity assays**

GUS activity in bacterial strains was measured as described by Henderson et al. (1985). Wild-type and mutant strains harbouring reporter plasmids were cultured in XVM2 media at 28 °C for 8 h. Bacteria cells were collected by centrifugation and resuspended in 375 µL of 1 mM p-nitrophenyl-β-d-glucuronide extraction buffer (50 mM sodium dihydrogen phosphate, 0.1% Triton X-100 and 10 mM β-mercaptoethanol, pH 7.2) and incubated at 37 °C for 10 min, and then terminated by 200 µL of 2.5 M 2-amino-2-methyl-1,3-propanediol. Enzyme assays were carried out in triplicate from at least three independent cultures.

**Protein manipulation**

In order to obtain purified Flp protein, the flp gene was cloned and introduced into the expression vector pQE30 [harbouring a lac promoter and a ribosome binding site (RBS) in front of the multiple cloning site (MCS)]. The Flp protein was expressed and purified with the methods modified from An et al. (2011). To remove imidazole, 6 × His-tagged Flp was dialysed against 200 volumes of Tris-HCl buffer [10 mM Tris-HCl (pH 8.0) and 1 mM dithiothreitol (DTT)] at 4 °C.

**Electrophoretic mobility shift assays**

The method that was deployed for EMSA assays was modified from that previously described by Su et al. (2016). Here fragments of the gene encoding HrpX were PCR amplified with the primers described in Table S3. These DNA fragments were labelled with FAM at the 5′ terminal. The Flp protein and selected DNA fragments were mixed with the binding buffer and incubated for 20 min at 28 °C. The reaction samples were loaded onto a 6% polyacrylamide-Tris-glycine-EDTA gel. Electrophoresis was performed in TGE buffer (pH 7.6) and visualized using an autoradiograph.

**In vitro transcription assays**

In vitro transcription assays were performed as previously described (Su et al., 2016). Promoter sequence fragments (311-bp) of hprX were PCR amplified with PCR (see Table S3). 6 × His-tagged Flp protein and DNA fragments were incubated for 30 min at room temperature in transcription buffer. Then, a NTP mixture (250 µM each of ATP, CTP and GTP; 250 µM biotin-16-UTP) and 0.5 U of *E. coli* RNA polymerase holoenzyme (New England BioLabs, Ipswich, MA, USA) were added to initiate transcription. After incubation at 28 °C for 30 min, the reactions were terminated and the transcription products were analysed by electrophoresis. The transcripts obtained were visualized using a phosphor imager screen (GE AI600).

**Stress tolerance assay**

The minimal inhibitory concentration (MIC) method (Su et al., 2016) was employed to test the sensitivity of the Xcc strains to several environmental stresses, including sodium dodecyl sulphate (SDS), hydrogen peroxide (H₂O₂), hyperosmosis (NaCl), phenol and heavy metal salts (CdCl₂, CuSO₄) stress. Briefly, Xcc strains were cultured overnight and diluted to an OD₆₀₀ of 0.1, then 100 µL of the diluted culture was plated on NYG plates supplemented with different concentrations of each reagent, respectively. The surviving colonies on the plates were counted after 3 days of incubation at 28 °C.

**Exopolysaccharide and extracellular enzymes assays**

Exopolysaccharide (EPS) and extracellular enzymes assays were performed as previously described (Su et al., 2016; Tang et al., 1991). To estimate EPS production, Xcc strains were inoculated into 100 mL NY liquid medium containing glucose (2% w/v) at 28 °C, 200 rpm for 5 days. EPS was precipitated from the culture supernatant with ethanol and dried at 55 °C using an oven and weighed. For quantitative estimation of the activity of the extracellular enzymes endoglucanase (cellulase) and amylase, Xcc strains were cultured in NYG medium for 12 h. For endoglucanase, 10 µL of enzyme-containing extracts was added to 200 µL of indicator buffer containing 1% (wt/vol) carboxymethylcellulose (CMC, Sangon, Shanghai, China) as the substrate. The reactions were carried out for 30 min at 28 °C. The released reducing sugars were measured as g-glucose equivalents, as described by Miller (1959). One unit (U) of the endoglucanase activity was defined as the amount of enzyme releasing 1 µmol of reducing sugar per minute. Amylase activity quantification was conducted in the same way as for the endoglucanase measurement, except that the substrate 1% (wt/vol) CMC was replaced by 1% (wt/vol) starch solution.

**Transcriptome analysis of the Flp mutant**

Transcriptome analysis were performed as previously described (Cui et al., 2018). Briefly, RNA was prepared from cell culture of OD₆₀₀ = 0.6. Contaminating genomic DNA was removed using RNase-free DNase I. After the quantity determination and quality assessment, total RNA was sent to Novogene (Beijing, China) for library construction and strand-specific RNA sequencing. Sequencing libraries were generated using a NEBNext Ultra™ Directional RNA Library Prep Kit for Illumina (New England BioLabs), and sequenced on an Illumina (CA, USA) HiSeq 2000 platform. Clean reads were mapped to the reference genome and the RPKM (reads per kilobase per million mapped reads) method was used to calculate the gene expression.
expression levels. False discovery rate FDR ≤ 0.05 and |log2FC| (log2 of the fold changes) ≥1 were considered for differentially expressed genes.

**ChIP assay**

ChIP assay was performed as previously described with minor modifications (Liu et al., 2019). In brief, a strain producing an Flp protein fused with 3 × Flag-tag (3 × Flag::Flp) at the N-terminus of Flp was first constructed. To do this, a DNA fragment encoding Flp fused with 3 × Flag peptide was obtained using PCR with the primer set F-Flag-F/R (Table S3) and cloned into the BamHI/HindIII sites of the vector pLAFR3. The acquired recombinant plasmid pFlp-Flag was introduced into Xcc flp deletion strain Δflp, resulting in strain Δflp/ pFlp-Flag. Xcc wild-type strain 8004 containing the empty vector pLAFR3 (8004/pLAFR3) was used as a negative control. Δflp strains were grown in XVM2 medium for 12 h and cross-linked using formaldehyde. Bacterial cells were collected and then lysed by sonication. For each ChIP sample, 50 μL of anti-Flag (agarose conjugated) was added to the bacterial lysates and incubated overnight. Unbound DNA fragments were washed and the bound DNA fragments and proteins were eluted by 0.25 M glycine (pH 2.5).

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**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of this article at the publisher’s web site:

Fig. S1 The homology of Flp and its position in evolution. (A) Sequence alignments between Flp and Fis proteins in *Yersinia pseudotuberculosis*, *Dickeya zeae* and *E. coli*. The sequences of these proteins were acquired from NCBI website and the alignment was proceeded with the software NTI Vector. (B) The position of Flp in evolutionary tree. A series of Fis family proteins were acquired from NCBI, and the alignment was proceeded with the software NTI Vector. A series of Fis family proteins were acquired from NCBI, and the alignment was proceeded with the software NTI Vector. (C) The position of Flp in evolutionary tree. A series of Fis family proteins were acquired from NCBI, and the alignment was proceeded with the software NTI Vector.

Table S1 Bacterial strains and plasmids used in this work. Note: *Rif*, Kan*, Tet* and Spc* indicate resistance to rifampicin, kanamycin, tetracycline and spectinomycin, respectively.

Table S2 Genes expressed by the Δflp mutant strain when grown in XVM2.

Table S3 Primers used in this study.