

Variations in type III effector repertoires, pathological phenotypes and host range of *Xanthomonas citri* pv. *citri* pathotypes

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SUMMARY

The mechanisms determining the host range of *Xanthomonas* are still undeciphered, despite much interest in their potential roles in the evolution and emergence of plant pathogenic bacteria. *Xanthomonas citri* pv. *citri* (*Xci*) is an interesting model of host specialization because of its pathogenic variants: pathotype A strains infect a wide range of Rutaceous species, whereas pathotype A^{*}/A^W strains have a host range restricted to Mexican lime (*Citrus aurantifolia*) and alemow (*Citrus macrophylla*). Based on a collection of 55 strains representative of *Xci* worldwide diversity assessed by amplified fragment length polymorphism (AFLP), we investigated the distribution of type III effectors (T3Es) in relation to host range. We examined the presence of 66 T3Es from xanthomonads in *Xci* and identified a repertoire of 28 effectors, 26 of which were shared by all *Xci* strains, whereas two (*xopAG* and *xopC1*) were present only in some A^{*}/A^W strains. We found that *xopAG* (= *avrGf1*) was present in all A^W strains, but also in three A^{*} strains genetically distant from A^W, and that all *xopAG*-containing strains induced the hypersensitive response (HR) on grapefruit and sweet orange. The analysis of *xopAD* and *xopAG* suggested horizontal transfer between *X. citri* pv. *bilvae*, another citrus pathogen, and some *Xci* strains. A strains were genetically less diverse, induced identical phenotypic responses and possessed indistinguishable T3E repertoires. Conversely, A^{*}/A^W strains exhibited a wider genetic diversity in which clades correlated with geographical origin and T3E repertoire, but not with pathogenicity, according to T3E deletion experiments. Our data outline the importance of taking into account the heterogeneity of *Xci* A^{*}/A^W strains when analysing the mechanisms of host specialization.

INTRODUCTION

Several factors may contribute to the emergence of new diseases: the introduction of an existing pathogen into new geographical areas, environmental changes that favour the spread of a pathogen or the broadening of the pathogen host range through rapid evolution to overcome host defences and an increase in fitness (Anderson *et al.*, 2004). The investigation of the mechanisms underlying the host range broadening or restriction of a pathogen might aid in the understanding of emergences. Asiatic canker caused by *Xanthomonas citri* pv. *citri* (*Xci*) is the most common and most aggressive form of citrus canker. This invasive disease occurs in more than 30 countries throughout the world, and is currently emerging in Africa (Balestra *et al.*, 2008; Derso *et al.*, 2009; Leduc *et al.*, 2011; Traore *et al.*, 2008). It affects most commercial varieties of citrus, limiting citrus production worldwide (Gottwald *et al.*, 2002). In South America, it has been reported to supplant the milder, geographically and host-restricted South American canker (*Xanthomonas citri* pv. *aurantifolii*), referred to as canker B and C, which has a much lower incidence on citrus (Schubert *et al.*, 2001). *Xci* enters host plant tissues through stomata or wounds, and multiplies in the mesophyll. This pathogen induces erumpent, callus-like lesions with a water-soaked margin on leaves, fruit and stem tissue. Severe attacks result in extensive defoliation, premature fruit drop and twig dieback. Strains of *Xci* have been classified into three pathogenic variants (i.e. pathotypes) as they differ in host range but not in symptomatology (Vernière *et al.*, 1998). Strains of pathotype A are present worldwide and have the widest host range (nearly all *Citrus* species and some related genera), although with differences in susceptibility among species or cultivars. In contrast, strains of pathotypes A^{*} and A^W have a narrow host range, mainly restricted to Mexican lime (*Citrus aurantifolia*) and alemow (*Citrus macrophylla*) in natural conditions (Sun *et al.*, 2004; Vernière *et al.*, 1998). Strains of pathotype A^W have been distinguished from strains of pathotype A^{*} on the basis of hypersensitive response (HR)-like symptoms on grapefruit (*Citrus paradisi*), sweet orange

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(*Citrus sinensis*), sour orange (*Citrus aurantium*), citron (*Citrus medica*), Orlando tangelo (*Citrus reticulata* × *Citrus paradisi*) and trifoliolate orange (*Poncirus trifoliata*) (Sun *et al.*, 2004). A previous description of the diversity of a collection of *Xci* by amplified fragment length polymorphism (AFLP) analysis, insertion sequence ligation-mediated polymerase chain reaction (IS-LM-PCR) typing and multi-locus variable-number tandem-repeat analysis (MLVA) showed a congruent structure of *Xci* separating pathotypes A and A*/A^W (Bui-Thi-Ngoc *et al.*, 2009). Furthermore, this study emphasized that the *Xci* population probably had a predominant clonal structure and that A* strains showed a higher genetic diversity than A strains. This result raised interesting questions about the phylogenetic relationship and evolutionary histories of *Xci* pathotypes A, A* and A^W.

Today, much remains to be learned about the molecular determinants involved in pathogenicity and host range variations reported among *Xci* pathotypes. The pathogenicity of most *Xanthomonas* depends on the conserved syringe-like type III secretion system (T3SS, encoded by the chromosomal *hrp*—HR and pathogenicity—gene cluster) capable of traversing the host cell wall and membrane to inject effector proteins, called type III effectors (T3Es), into the host cell cytoplasm (Alfano and Collmer, 1997). In compatible interactions, T3Es promote bacterial growth and virulence. In incompatible interactions, they trigger specific defences in resistant plants, and induce HR (Gurlebeck *et al.*, 2006; White *et al.*, 2009). They constitute prime targets for mutation and selection to evade the host immune system.

Recent work has speculated the likely role of T3Es as a group of factors or 'repertoire' shaping the host ranges of individual *Pseudomonas syringae* strains (Sarkar *et al.*, 2006) and many xanthomonads (Hajri *et al.*, 2009, 2012a, 2012b). Documented examples have shown the contribution of one or a few effectors in the virulence of a specific strain on a specific host or cultivar (Arnold and Jackson, 2011). In the *Xanthomonas* genus, effectors named *Xanthomonas* outer proteins (Xops) have been described according to studies based on functional assays, combined with sequence and structural similarity searches of genomic data. They are classified into 66 families (White *et al.*, 2009) (*Xanthomonas* resource website: <http://www.xanthomonas.org/>; R. Koebnik, IRD, Montpellier, France, personal communication). In *Xci*, a functional T3SS is necessary for infection, the formation of canker symptoms on host plants and HR induction in nonhost plants (Dunger *et al.*, 2005). Twenty-six effector families are represented in the genome of *Xci* IAPAR 306 (Da Silva *et al.*, 2002). Furthermore, the effector *avrGf1* (=xopAG) was subsequently identified in A^W strains (Rybak *et al.*, 2009), enlarging the *Xci* repertoire to 27 T3Es. XopAG has been shown to be responsible for the induction of HR-like reactions on grapefruit, putatively contributing to the specific interaction of the pathogen. The deletion of *xopAG* partially restored canker development on host species differentiating A and A^W strains, yet canker symptoms were less severe. This effector is also found in

Xanthomonas campestris pv. *campestris* sequenced genomes (strains ATCC33913, B100, 8004) (Da Silva *et al.*, 2002; Qian *et al.*, 2005; Vorhoelter *et al.*, 2008), in *Xanthomonas vesicatoria* ATCC35937 (Potnis *et al.*, 2011), but not in the *Xanthomonas euvesicatoria* sequenced strain (85-10) (Thieme *et al.*, 2005), which is phylogenetically closer to *Xci* IAPAR 306. The involvement in pathogenicity of 13 of the 26 Xops found in the genome of the model strain *Xci* IAPAR 306 was assessed by site-directed mutagenesis on each individual effector (Figueiredo *et al.*, 2011). This study also included other candidate coding sequences revealed by the presence of a conserved promoter element, known in *Xanthomonas* as plant-inducible promoter (PIP) boxes (Koebnik *et al.*, 2006; Wengelnik and Bonas, 1996). Apart from *hrp* genes and the mutation of the key canker-inducing protein PthA4 belonging to the transcription activator-like (TAL) effector family (Swarup *et al.*, 1991, 1992), none of the tested mutants reduced the ability to cause citrus canker disease on grapefruit and Mexican lime leaves. An understanding of the role of these effectors in host range variations among *Xci* pathotypes requires further investigation.

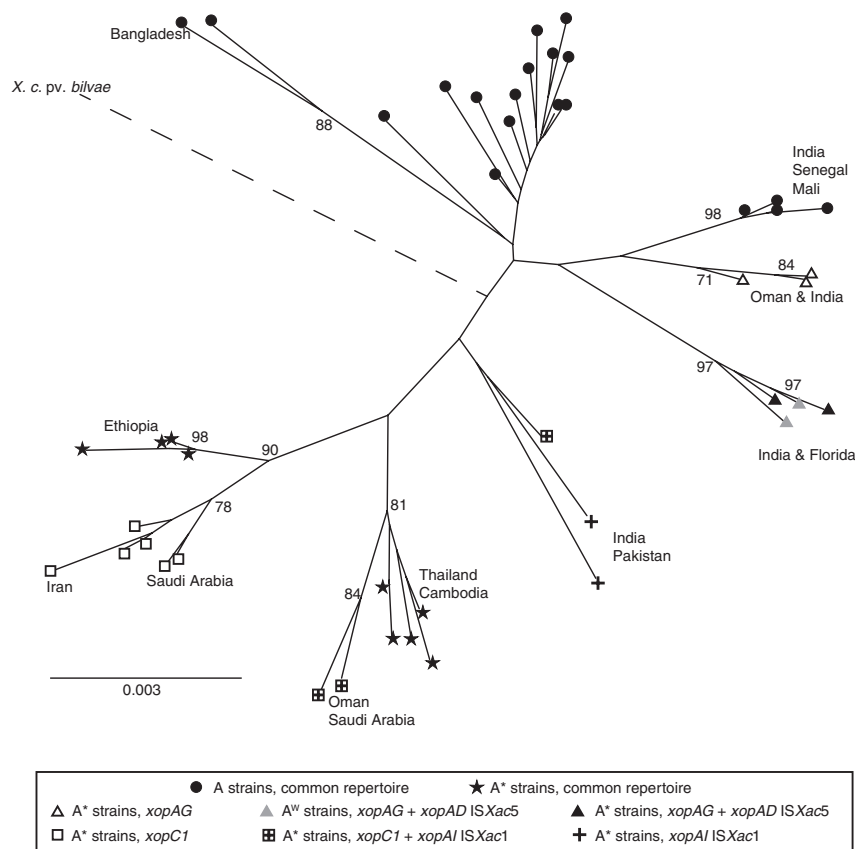
The aim of this work was to investigate the role of T3Es in defining the host specialization of *Xci* pathotypes. For this purpose, we worked on a comprehensive worldwide collection of *Xci* strains chosen to represent the known genetic diversity of the pathovar. We characterized the pathogenicity of the strains based on several tests to better assess their aggressiveness and host range, and we identified their T3E repertoires. We based this screen on the potential Xop candidates described in the *Xanthomonas* genus, and conducted PCR and Southern blot hybridization. In the case of pathotype-specific candidate effectors, site-directed mutagenesis was conducted to evaluate the role of single T3Es in the plant–bacterium interaction. We also investigated how the different T3E repertoires correlate with the genetic diversity of *Xci*.

RESULTS

AFLP analysis reveals the *Xci* global diversity and geographical structure of A*/A^W strain clusters

A collection of 55 *Xci* strains was built to represent *Xci* genetic diversity based on a previous analysis and including recently isolated strains (Bangladesh) and strains corresponding to recent outbreaks (Mali and Senegal). A total of 169 AFLP fragments was recorded, 105 of which were polymorphic. A total of 49 haplotypes was identified within the strain collection. Graphical representations, a neighbour-joining (NJ) tree (Fig. 1) and metric multidimensional scaling (MDS) plots (Fig. S1, see Supporting Information) suggested a larger polymorphism for A*/A^W strains. This higher diversity was supported by Nei's diversity indices: HE = 0.10 for A*/A^W strains vs. HE = 0.06 for A strains. A majority of A*/A^W clusters were supported by high bootstrap values (≥80%). Some of these clusters contained strains originating from a single

Fig. 1 Amplified fragment length polymorphism (AFLP) neighbour-joining tree. Tree constructed using evolutionary genome divergences among 55 *Xanthomonas citri* pv. *citri* strains based on 169 AFLP markers. Bootstrap values under 70% are not indicated. The branch containing *X. citri* pv. *bilvae* should be twofold longer and is represented by a broken line. Symbols indicate the type III effector (T3E) contents of each strain, either as the common repertoire with additional genes or the common repertoire with genes containing an insertion. Geographical origins are indicated when they are consistent with branching.



country (e.g. Ethiopia), whereas others contained strains from several countries (e.g. Oman and India). Most A strains clustered irrespective of their geographical origin, except in rare cases (e.g. Bangladesh). A cluster of the three A* strains JF90-8*, LG100* and LG116* from Oman and India (bootstrap of 71%) was found close to a cluster of A strains containing LE116-1, LE117-1, LH1-3, LH37-1 and NCPPB 3562 from Mali, Senegal and India (supported by a bootstrap of 98%). Strains LG115* and NCPPB 3608* from India and X2002-1035^W and X2003-3218^W from Florida formed a subclade supported by a maximal bootstrap value (97%), and were closer to A strains than to the other A* strains, based on the MDS plot and NJ tree (Figs S1 and 1).

All *Xci* share a common repertoire of 26 T3Es and only the host-restricted A* and A^W strains contain a variable repertoire

Repertoires of T3Es were defined in terms of the absence or presence of portions of the corresponding sequence after PCR amplification with two pairs of specific primers (Table S4, see Supporting Information). We were able to define a common repertoire of 26 candidate T3Es, present in all tested *Xci* strains, and a variable set of two candidate T3Es: *xopC1* and *xopAG* (Table 1). *xopC1* was present in only 13 A* strains. *xopAG* was present in the

two A^W tested strains and in five A* strains (Fig. 1 and Table 1). Among the 26 common T3Es, the two T3Es *xopAD* and *xopAI* were detected by PCR with larger sizes than expected (2 kb and 1.2 kb, respectively) in some A* and A^W strains, suggesting the presence of ISs (see below). The pathogen *X. citri* pv. *bilvae* strain NCPPB 3213, responsible for spot lesions on Rutaceae, shared 23 effectors with *Xci* and contained two additional effectors: *xopJ5* and *xopB*. Southern blots were performed to validate the presence of T3Es that were detected by PCR in only a few *Xci* strains or that were not detected by PCR and previously reported in canker-forming strains. We looked for the presence of two T3Es detected in a few *Xci* strains by PCR: *xopC1* and *xopAG* (false-negative results putatively induced by primer mismatches). We also looked for the presence of four T3Es [*xopB*, *xopE4*, *xopAF* and *xopJ5* (=avr*XccB*)] reported in South American citrus canker-forming strains *Xanthomonas citri* pv. *aurantifolii* type B ICPB 11122 (*XauB*) and type C ICPB 10535 (*XauC*) (Moreira *et al.*, 2010), and *xopF1* previously reported in other repertoire studies, including strains of *Xci* (Hajri *et al.*, 2009). The two effectors *xopC1* and *xopAG* were only detected from strains with positive results in PCR, and no hybridization was detected in *Xci* strains for the five markers *xopB*, *xopE4*, *xopAF*, *xopJ5* and *xopF1*. Overall, strains sharing the same variable repertoire cluster in the same AFLP clades were supported by high bootstrap values (>70%) (Fig. 1).

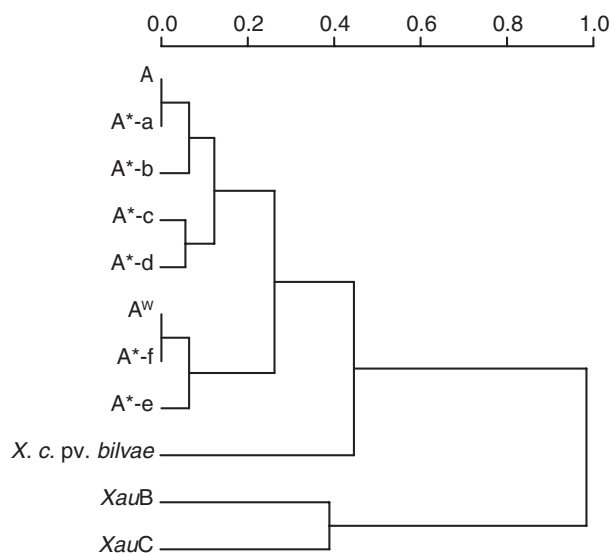


Fig. 2 Relationships between strains according to their variable repertoire. Dendrogram built by Ward's method (Maechler *et al.*, 2012) based on the variable repertoires of *Xanthomonas citri* pv. *citri* (*Xci*), *X. citri* pv. *bilvae* strain NCPPB 3213, and *X. citri* pv. *aurantifolia* strains ICPB 11122 (*XauB*) and ICPB 10535 (*XauC*). All strains contain the common repertoire with the following variations for A* strains: (a) common repertoire; (b) *xopAI*Δ*ISXac1*; (c) *xopC1*; (d) *xopAI*Δ*ISXac1* and *xopC1*; (e) *xopAG*; (f) *xopAG* and *xopAD*Δ*ISXac5*. A^w strains contain the common repertoire with *xopAD*Δ*ISXac5* and *xopAG*.

Hierarchical clustering based on T3E content

Hierarchical clustering by Ward's method was performed using a matrix of T3E presence/absence built after homology searches of the T3E candidate list on 27 *Xci*, *X. citri* pv. *bilvae* strain NCPPB 3213 draft genome sequences and genomes of *XauB* and *XauC*. This clustering showed that the T3E repertoire does not distinguish pathotypes, but rather reflects strain phylogenetic divergence, the *X. citri* pv. *bilvae* strain NCPPB 3213 being closer to the *Xci* strains than the two tested *X. citri* pv. *aurantifolia* strains (Fig. 2).

Sequencing analysis of two variable T3Es of *Xci* reveals potential horizontal gene transfer (HGT) events with other *Xanthomonas* strains

Sequencing analyses of the two variable T3Es and their flanking regions were performed to better understand their distribution and to assess whether differences in sequence could explain some phenotypic variability among strains sharing the same variable repertoire.

Sequence analysis of *xopC1*

The *xopC1* gene was sequenced from four A* strains belonging to three different AFLP clusters: CFBP 2911* from Pakistan, JF90-2* from Oman and both JS584* from Iran and JK2-10* from Saudi Arabia belonging to the same cluster as they displayed distinct phenotypes on sweet orange and grapefruit (see below). Our

analysis showed that *xopC1* DNA coding sequences were completely identical between *Xci* A* strains and had 98.8% homology with *xopC1* from *X. euvesicatoria* strains 85-10 and 75-3. Analysis of the *xopC1* upstream flanking region (1460 bp) and downstream region (750 bp) for all four strains revealed that they were highly homologous (98.8%–99.0%) to the corresponding regions bordering *xopC1* in *X. euvesicatoria* 85-10. *xopC1* had a G + C content of 48%, contrasting with the genomic average of 65.2%, and was flanked by the insertion sequences IS1595 and IS1478. The presence of IS1595 had not been documented in *Xci* previously.

Sequence analysis of *xopAG*

The *xopAG* gene was sequenced from four *Xci* strains found in two AFLP clades and of various geographical origins: X2003-3218^w from Florida and NCPPB 3608* from India, which were found in the same subclade, and strains JF90-8* and LG116* from Oman and India, respectively. The gene *xopAG* was also sequenced from *X. citri* pv. *bilvae* strain NCPPB 3213. *xopAG* from A^w strains differed by one nonsynonymous single nucleotide polymorphism (SNP) from *xopAG* found in A* strains NCPPB 3608*, JF90-8* and LG116*, and had a G + C content of 56.9%, which was again lower than the average value of the rest of the genome (65.2%). Orthologues of *xopAG* were found in *X. campestris* pv. *campestris* B100 (80% homology), *XauC* (55% homology) and interrupted by IS1479 in *XauB* (55% homology) and *X. citri* pv. *bilvae* NCPPB 3213 (99% homology), which induced a strong HR on grapefruit and sweet orange. Based on unpublished draft genome sequence data, BLAST analyses revealed the presence of the right-end inverted repeat of ISXac4, 1240 bp downstream of *xopAG* (data not shown). Both G + C content and the presence of IS elements suggested that both *xopC1* and *xopAG* might have been acquired through HGT.

Sequencing analysis of the larger amplicons of *xopAD* and *xopAI* reveals the presence of IS elements

The larger amplicon obtained for *xopAI* was sequenced in strain JF90-2*, and sequence analysis revealed that it was interrupted at position 714 after the start codon (total length, 906 bp) by ISXac1, a member of the IS4 family. Similarly, the larger amplicons obtained by PCR for *xopAD* were sequenced from strains NCPPB 3608* and X2003-3218^w, representative of the two pathotypes, and sequence analysis revealed that *xopAD* was interrupted at position 7989 (total length, 8652 bp) by ISXac5 related to ISRs019, which belongs to the IS21 family. ISXac5 and ISRs019 had a similar size and shared 72.5% DNA homology, similar inverted repeats and displayed two putative peptides which were 67% and 74% identical, respectively. Both ISs were bordered by direct repeats of 6 bp. Comparison of the *xopAD* sequence (without considering the IS element) showed that *xopAD* from strains NCPPB 3608* and X2003-3218^w shared 99.2% DNA homology with the corresponding *xopAD* allele from *X. citri* pv. *bilvae* strain NCPPB 3213. Draft genome sequence data indicated

that, in strains possessing *xopAG* (including JF90-8*, NCPPB 3608* and X2003-3218^w), the sequence of *xopAD* showed less homology with *xopAD* orthologues in the other A* strains (95.9%) and in A strains (97.6%) (data not shown) than with *X. citri* pv. *bilvae* strain NCPPB 3213.

Phenotypic characterization on detached leaf assay reveals that all A strains produce an identical phenotypic response, whereas A*/A^w strains show heterogeneous responses depending on the host and the strains

Phenotypic characterization was conducted to assess the role of T3Es in the observed responses during the interaction with various hosts. A disease scale was used to assess strain pathogenicity each day post-inoculation (dpi) (Fig. S2, see Supporting Information), and a disease development index (DDi) was calculated to assess the global response of each tested strain, as values of the area under the disease progression curve. As expected, *X. citri* pv. *bilvae* strain NCPPB 3213 did not cause canker-like symptoms, but produced extensive water-soaked lesions on Mexican and Tahiti limes and alemow, and HR-like responses on Pineapple sweet orange and Marsh grapefruit.

All tested *Xci* strains showed similar typical erumpent, callus-like lesions at 14 dpi on the nondiscriminant host species (Mexican lime, Tahiti lime and alemow) with minor differences in disease development and time of appearance of the epidermis rupture (data not shown). As expected, inoculation of two pathotype-discriminating host species, Pineapple sweet orange and Marsh grapefruit, showed heterogeneous responses depending on the strains. On both host species, A strains produced typical canker-like symptoms at 14 dpi, whereas A* strains elicited a

range of reactions without any correlation with the T3E content, from no symptoms to erumpent symptoms for the three A* strains from Iran (JS581*, JS582* and JS584*). The majority of A* strains induced defence-like responses, which consisted of necrotic tissues on grapefruit and/or sweet orange with variability in days of appearance from 3 to 21 dpi. Representation of the disease progression curves combined with the appearance of HR for a subset of strains representative of the various pathogenic profiles observed in this assay showed no HR-like responses for A strains on these host species (Fig. S3, see Supporting Information). Strains JS581*, JS582* and JS584*, which are further described below, showed the same disease progression curve, and only strain JS584* is represented. Based on DDi on sweet orange and grapefruit, three clusters of strains were identified. Cluster I contained all 17 A strains which showed a homogeneous response to inoculations, cluster II contained the three Iranian A* strains with intermediate phenotypes (JS581* and JS582* showed exactly the same phenotypic response and thus are not distinguishable on Fig. 3), and cluster III contained all other A* and A^w strains with no typical canker lesions on discriminant host species. Overall, sweet orange provided a better discriminative ability than grapefruit, with Iranian A* strains being distinguished from A strains by a delayed development of canker-like lesions. Strains of cluster II were further inoculated, together with three A strains, on Temple mandarin, Fairchild mandarin SRA30, Meyer lemon and Ortanique tangor (mandarin *C. reticulata* × sweet orange). Inoculation on Meyer lemon and the two mandarin cultivars did not allow the cluster II strains to be distinguished more clearly than on sweet orange (data not shown). Interestingly, all cluster II strains induced defence-like responses on Ortanique tangor without canker-like lesions. A strains produced canker-like lesions, but the observed reactions suggested a moderate aggressiveness on this host

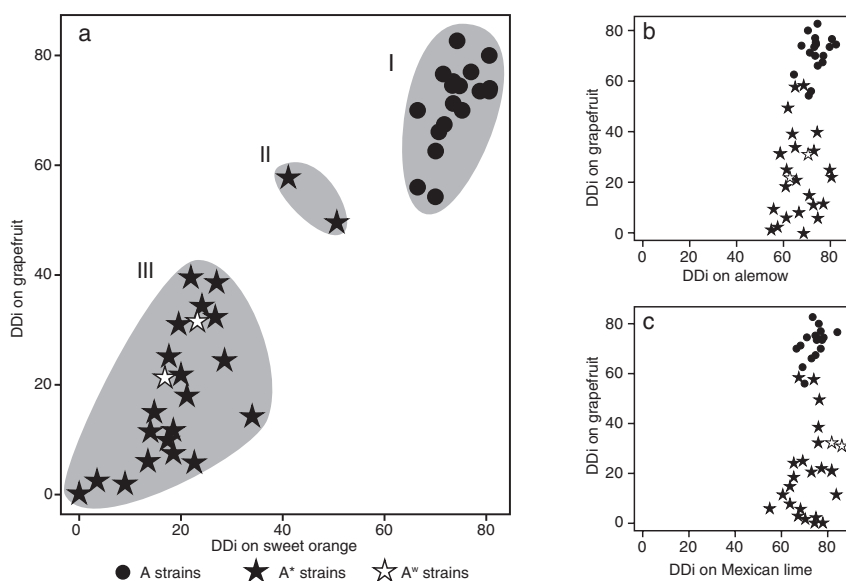


Fig. 3 Disease development indices (DDi) on four host species. Forty-seven strains were inoculated on detached leaves of grapefruit and sweet orange (a), grapefruit and alemow (b), and grapefruit and Mexican lime (c). Clusters I, II and III represent subsamples according to Tukey's tests.

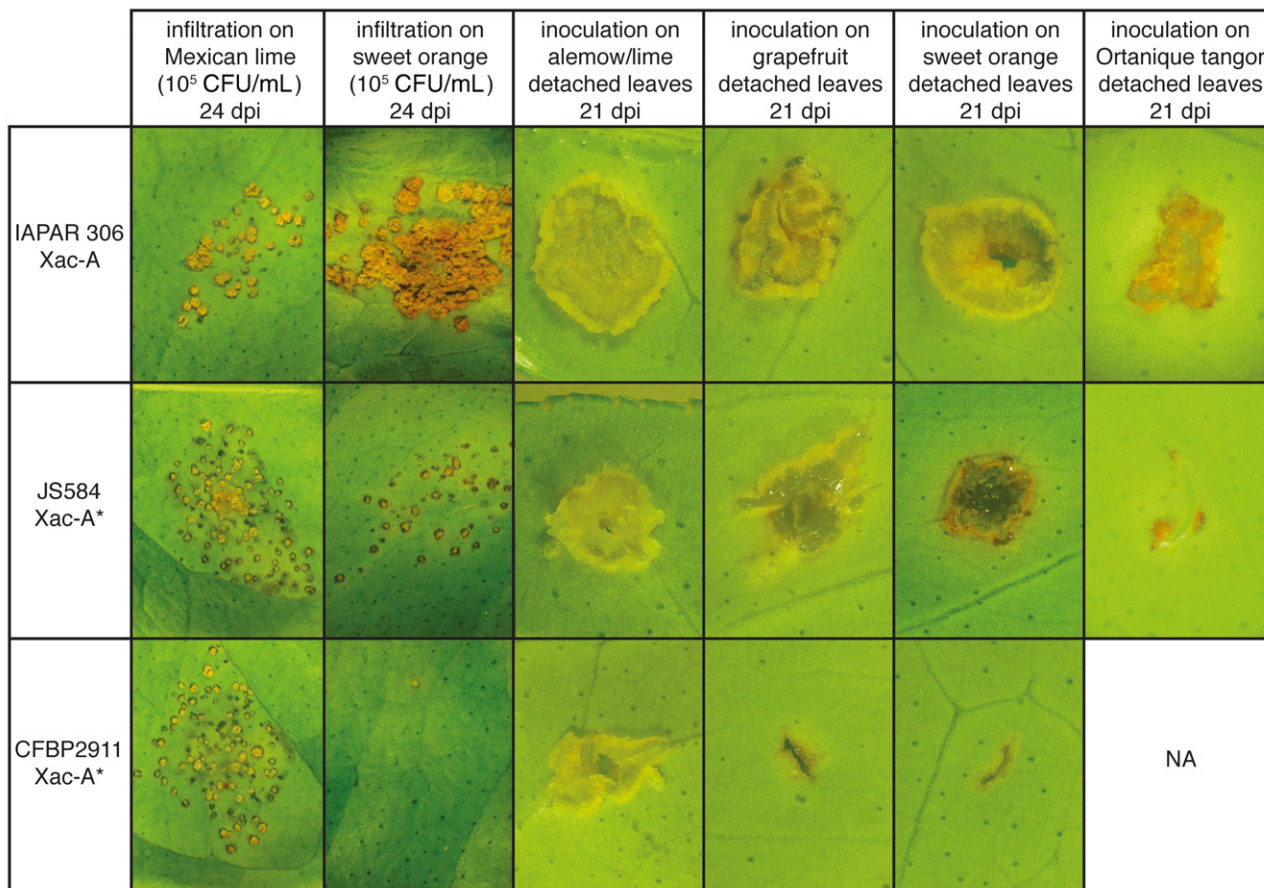


Fig. 4 Symptomatology of A and A* strains. Symptoms observed after infiltration of attached leaves (first two columns) and after wound inoculation of detached leaves (last four columns) with A strain IAPAR 306, and A* strains JS584* and CFBP 2911*. The width of each photograph is approximately 15 mm for the first two columns and 1.5 mm for the last four columns. CFU, colony-forming units.

species (Figs 4 and S2). Overall, the *Xci* phenotypic response was variable for A* strains, but did not correlate with their T3E content.

Comparison of bacterial population *in planta* confirms A* heterogeneous response on sweet orange

In order to better characterize the atypical interaction on detached leaf assay of A* strains from Iran, which correspond to cluster II (Fig. 3), we compared *in planta* bacterial multiplication of strain JS584* (selected as a reference for cluster II) with A strain IAPAR 306 and A* strain CFBP 2911* (selected as references for clusters I and III, respectively). As shown in Fig. 5, on Mexican lime at 24 dpi, all three strains reached population densities of 2.5×10^7 to 1.3×10^8 colony-forming units (CFU)/cm², which is typical of canker symptoms. On sweet orange at 8 dpi, strains JS584* and IAPAR 306 showed nonsignificantly different mean population sizes, whereas strain CFBP 2911* showed a much smaller population size (Fig. 5). At 24 dpi, only strain IAPAR 306 reached large population sizes (6.3×10^8 CFU/cm²), whereas strains CFBP 2911*

and JS584* showed significantly smaller population sizes (according to Tukey's tests), thus confirming previous results (Fig. 5) (Bui-Thi-Ngoc *et al.*, 2010). Visual comparison at 24 dpi of the symptoms caused by all three strains showed overall differences in symptomatology that illustrated the differences observed in terms of bacterial population sizes (Fig. 4), and confirmed that strain JS584* is much less aggressive than A strains on sweet orange. Strain JS584* induced small canker-like symptoms on sweet orange, which is unexpected for A* strains, compared with the symptomless strain CFBP 2911*. However, JS584* symptoms were attenuated compared with the typical canker symptoms induced by the A strain IAPAR 306, and this is consistent with population sizes recovered *in planta*.

All A* and A^w strains containing *xopAG* induce HR-like responses on sweet orange and grapefruit

Considering that A^w strains were first described as causing HR-like reaction on various hosts, and that such responses depend on the

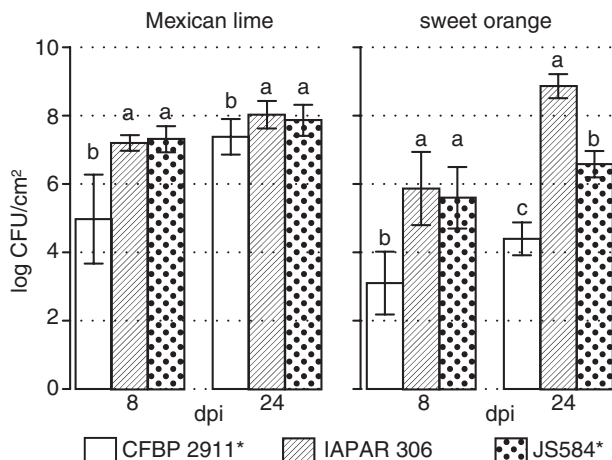


Fig. 5 Bacterial multiplication in Mexican lime and sweet orange. Means of log colony-forming units (CFU) from 20 leaf discs (two experiments) after inoculation at 10^5 CFU/mL. Error bars indicate \pm standard deviation (SD). Distinct letters indicate significant differences between strains at a given time on a given host according to Tukey's test.

T3E *xopAG* (Rybak *et al.*, 2009), all *xopAG*-containing strains were submitted to infiltration to look for HR or symptom development, and compared with a subset of 17 strains representative of *Xci* genetic diversity and repertoire variability. After inoculation with bacterial suspensions (1×10^8 CFU/mL), typical water-soaked lesions that evolved as canker symptoms at 4–6 dpi were observed on Mexican lime, irrespective of the pathotype. On sweet orange, mandarin and grapefruit, HR-like symptoms appeared at 3–6 dpi for strains X2003-3218^W and X2002-1035^W, and for strains NCBBP 3608*, LG100*, LG115* and LG116* (from India) and JF90-8* (from Oman), which all carry *xopAG*. Interestingly, none of the strains that did not contain *xopAG* produced HR-like symptoms on sweet orange, mandarin and grapefruit (Fig. S4, see Supporting Information, and data not shown). On these hosts, A strains induced typical canker symptoms, whereas A* strains exhibited variable responses: chlorosis, blister-like lesions with water soaking in the infiltrated area or canker-like symptoms, depending on the strain and host species. *Xanthomonas citri* pv. *bilvae* strain NCPPB 3213 induced water-soaking lesions on Mexican lime and HR-like symptoms at 3 dpi on mandarin, sweet orange and grapefruit. This strain contains *xopAG*, but also possesses other T3Es absent from *Xci* strains, and we cannot conclude that *xopAG* is responsible for the HR-like symptoms observed for this pathovar.

Deletion of *xopC1* or *xopAD* does not alter *Xci* pathogenicity

Xci is characterized by a variable repertoire of two T3Es: *xopAG*, whose role in HR-like symptoms has already been shown (Rybak

et al., 2009), and *xopC1*. We also showed that both *xopAD* and *xopAI* are interrupted by IS elements in several strains. The role of *xopAI* has already been assessed by Figueiredo *et al.* (2011). They showed that the pathogenicity of A strain IAPAR 306 was not altered when *xopAI* was deleted. Therefore, we focused our study on *xopC1* and *xopAD*. *xopC1* was deleted in strains JS584*^R and CFBP 2911*^R, as these two strains showed different phenotypic responses on sweet orange and pomelo. *xopAD* is interrupted in several A* and A^W strains by IS*Xac5*, and so may not be functional. To assess the role of the *xopAD* full-length protein in A strains, strain IAPAR 306 was deleted for *xopAD*. Mutants JS584*^R Δ *xopC1*, CFBP 2911*^R Δ *xopC1* and IAPAR 306^R Δ *xopAD* showed similar symptoms to the rifampicin-derivative strains when inoculated on detached leaves, as well as when infiltrated at a concentration of 1×10^5 CFU/mL (Fig. S5, see Supporting Information). *In planta* population sizes between a given mutant and its wild-type-resistant derivative were not significantly different at 8 and 24 dpi, according to Tukey's tests. Population size differences did not exceed 0.48 log CFU/cm² at 8 dpi and 0.30 log CFU/cm² at 24 dpi (Table S2, see Supporting Information).

DISCUSSION

Xci pathotype A* shows a wider genetic diversity than pathotype A and a broad extent of reaction on several *Citrus* species

Most studies aimed at understanding *Xci* virulence mechanisms and host range specialization have focused on one strain (Dunger *et al.*, 2012; Figueiredo *et al.*, 2011; Rybak *et al.*, 2009; Yan and Wang, 2012) or a set of strains (Al-Saadi *et al.*, 2007). The choice of a particular strain often relies on the availability of genomic data, or of well-characterized phenotypes obtained by random mutagenesis. Therefore, only one of few pathological profiles can be tested simultaneously. In both cases, the representativeness of a particular strain or set of strains remains arbitrary and the genetic diversity of *Xci* was not taken into account when studying determinants of its host range. In this work, we wanted to characterize the role of T3Es in *Xci* host range specialization, and we rationalized our choice of strains to represent the worldwide genetic and phenotypic diversity of the bacterium, based on previous studies (Bui-Thi-Ngoc *et al.*, 2009) and including strains from recent outbreaks (Balestra *et al.*, 2008; Derso *et al.*, 2009; Leduc *et al.*, 2011; Traore *et al.*, 2008). We first analysed the genetic relatedness between *Xci* pathotypes using AFLP markers, which have the advantage of covering the genome and being theoretically neutral. This analysis indicated that strains from *Xci* pathotypes are differentiated into several clusters which do not correlate with *Xci* pathotypes: all A* and A^W strains which contain *xopAG* cluster closer to A strains than to A* strains (Figs 1 and S1). Pathotype A* strains are genetically more diverse than pathotype A strains. In addition, pathogenicity tests revealed

that this difference in terms of homogeneity/heterogeneity is correlated with their aggressiveness and virulence. Our results clearly revealed that A strains are homogeneous in terms of host, types of symptom and aggressiveness, whereas the pathogenicity of A* strains is highly diverse on sweet orange, grapefruit and mandarin (Figs 3, 4 and S3). The two tested A^W strains exhibited one of the several reactions observed for A* strains and, consistent with previous data (Bui-Thi-Ngoc *et al.*, 2010), A^W and most A* strains were not distinguished on the basis of their host range. Within pathotype A*, no correlation was found between pathogenicity profiles and genetic structure. We observed that A* Iranian strains of our collection induced canker-like symptoms when inoculated on detached or attached leaves of grapefruit and sweet orange, although the size of the canker symptoms was often more moderate than those induced by A strains. Our results highlighted the difficulty of assigning strains to a given pathotype on the sole basis of detached leaf assay on grapefruit and sweet orange, and suggested that Ortanique tangor may be a useful species for pathotype identification based on pathogenicity test data. The genetic relatedness between A* strains did not correlate with differences in symptomatology, as the latter A* strains from Iran (corresponding to cluster II in Fig. 3), which had a high virulence on sweet orange and grapefruit, belonged to the same clade as strains from Saudi Arabia, which induced no symptoms on detached leaf assay or when infiltrated (Figs 1 and 4). The classification of Iranian strains within pathotype A* was confirmed by the cumulative information obtained by *in planta* population measurements on sweet orange (two log units lower than those obtained for strain IAPAR 306 at 24 dpi), response to inoculations on several host species and genotyping data (this study and data not shown).

The *Xci* variable repertoire does not explain host specialization between pathotypes

Our T3E gene distribution analysis showed that *Xci* possesses a common repertoire of 26 known T3Es which corresponds to the average size of *Xanthomonas* repertoires described so far (Hajri *et al.*, 2012a, b; White *et al.*, 2009). *Xci* repertoire variations do not explain host specialization between *Xci* pathotypes. Several A* strains shared the same repertoire as A strains, but belonged to distant genetic clusters and induced different symptoms (Fig. 1). Only A* strains exhibited variability in known T3Es (*xopC1* and *xopAG*), which showed a restricted distribution among *Xanthomonas* sequenced strains (*Xanthomonas* resource website: <http://www.xanthomonas.org/>; R. Koebnik, personal communication). However, the distribution of T3Es among A* strains was not correlated with symptomatology variation within this pathotype. Deletion of *xopC1* did not seem to alter pathogenicity, regardless of the inoculation technique used (Fig. S5). The secretion of *xopC1* *in planta* was shown for *X. euvesicatoria*, but no function modification could be evidenced after deletion (Noel *et al.*, 2003). The

close relatedness between the *xopC1* orthologues and the presence of identical IS elements in the flanking regions suggests that this T3E belongs to a 'mobile module' that has been inserted in genomes by HGT mediated by mobile elements. The region in which it is inserted in *X. euvesicatoria* contains multiple coding sequences associated with mobile elements (Noel *et al.*, 2003) and is homologous with two nonadjacent regions of the genome of *Xci* strain IAPAR 306 (one on the chromosome, one on plasmid pXac64), in which the mobile module containing *xopC1* is absent (Fig. S6, see Supporting Information). We can therefore speculate that this region is favourable for the insertion of mobile elements, which could explain why *xopAI* (if it is located in the same area) is interrupted by an IS in several A* strains. However, deletion of *xopC1* (this study, Fig. S5) and *xopAI* (Figueiredo *et al.*, 2011) did not show any selective advantage related to the plasticity of this region for colonization on the tested citrus species. Similarly, the effector *xopAD* is interrupted by the ISXac5 related to *ISRso19* in several A^W strains, but the deletion of this effector (this study, Fig. S5) did not modify its host range.

The *Xci* A*/A^W T3E repertoire is closer to that of *Xci* A than to that of South American canker-forming strains despite homologies in host range

The variability of the response of A* strains on two pathotype-discriminating host species, Pineapple sweet orange and Marsh grapefruit, can be compared with the symptomatology and host ranges characterizing South American canker-causing strains *XauB* and *XauC*. A* strains from Iran (corresponding to cluster II in Fig. 3) share many similarities in symptomatology with strains causing canker B, which was described as primarily restricted to lemon (*C. limon*), but also mildly pathogenic to sweet orange and grapefruit, and was characterized by an *in planta* multiplication ability lower than that of A strains (Moreira *et al.*, 2010). Similarly, A*/A^W strains that contain *xopAG* exhibit the same pattern of response as strains causing canker C, which is restricted to Mexican lime and induces HR-like reactions on most other species (Moreira *et al.*, 2010). All strains of our collection shared a common T3E repertoire identical to the repertoire of the sequenced *Xci* A strain IAPAR 306. Although some T3Es could be missed in our PCR screen, because of sequence divergence between *Xci* and strains from which primers were designed, homology searches of the effector candidate list on 28 *Xci* draft genome sequences confirmed that T3E repertoires determined in this study correspond to the T3E deduced from the genomic sequences (data not shown). Despite the fact that host range-restricted *Xci* A* strains and *X. citri* pv. *aurantifolii* display similar host ranges, searches against the whole genome shotgun database revealed several differences between citrus canker-forming strains in their non-TAL effector repertoires. In addition, the 17.5-repeat TAL-effector PthA4 from *Xci* IAPAR 306 and its homologues

PthB in *XauB* and PthC in *XauC* have been reported to be functionally interchangeable to induce canker symptoms when transformed in strains A, A*, A^w, B and C, without incidence on the host range (Al-Saadi *et al.*, 2007). Therefore, our results suggest that the T3E repertoire may reflect phylogenetic divergence rather than pathogenicity features, such as host range (Fig. 2).

A*/A^w strains that contain *xopAG* are closely related to A strains, and therefore *xopAG* may have contributed to their host range restriction

The difficulty in demonstrating virulence roles for individual effectors has been documented previously and is generally imputed to functional redundancy (Kvitko *et al.*, 2009). The role of single T3Es has often been characterized in nonhost interactions, where they are recognized by a corresponding resistance gene, resulting in local cell death (HR). Here, we investigated the role of the *Xci* variable T3E repertoire, but were unable to identify a single or combination of effectors to explain *Xci* host range specialization. In agreement with previous work showing that XopAG (AvrGf1) is responsible for HR-like symptoms caused by *Xci* A^w strains (Rybak *et al.*, 2009), we found that all A* and A^w strains carrying *xopAG* induced necrotic symptoms (HR-like) on grapefruit and sweet orange within a few days. Mutants of A^w strains lacking *xopAG* produced small pustules, typical of reactions induced by some A* strains, but to a less severe extent than the disease reactions induced by A strains in grapefruit (Rybak *et al.*, 2009), thus indicating a possible role of another determinant of pathogenicity in host range specialization of A^w strains. All strains inducing similar phenotypic responses contained *xopAG* orthologues that were 99.9% homologous at the protein level between strains. Both groups of strains carrying *xopAG* were more closely related to A strains than to other A* strains (Fig. 1). Strains carrying *xopAG* might have a common ancestor in the Indian region and might have participated in the evolution of *Xci* local host range specializations driving to the current state: two clades of A*/A^w strains with a narrow host range share homologies in symptomatology because of the effector *xopAG*, and both are closely related to A strains.

The variable T3E repertoire of *Xci* is related to phylogenetic placement and the geographical origins of *Xci* pathotypes

Interestingly, *xopC1* was found only in strains originating from western Asia, but genetically distant. This finding is consistent with the hypothesis of HGT between *Xci* strains. *xopAG* was first described in A^w strains from Florida, but an Indian origin of the Florida A^w strains is very likely (Schubert *et al.*, 2001). The *Xci* variable T3E repertoire, consisting of *xopC1* and *xopAG*, but also DNA rearrangements within *xopAI* and *xopAD*, can therefore be linked to the geographical origin of the strains and their intrap-

athovar phylogenetic position, rather than to differences in host range. Such results suggest that these *Xci* AFLP clades might have distinct evolutionary histories.

Comparison of *xopAG* sequences among xanthomonads pathogenic to citrus showed that *xopAG*_{*Xci*} was more closely related to *xopAG* from *X. citri* pv. *bilvae*, which also induces HR on grapefruit and sweet orange, than to *xopAG* from *XauB* and *XauC* (99% vs. 55% DNA homology). *xopAD* was present in all tested *Xci* strains but, in strains that also possessed *xopAG* (which were the only ones inducing HR on grapefruit and sweet orange), *xopAD* was more homologous to the corresponding orthologue found in *X. citri* pv. *bilvae* than to the orthologue present in *Xci* strains that did not contain *xopAG*. *Xanthomonas citri* pv. *bilvae* has been reported solely from India, where it shares Mexican lime (Chakravarti and Chaudhary, 1988) and, to a lesser extent, *Aegle marmelos* and *Feronia elephantum* (Khan and Hingorani, 1970) as host species with *Xci*. On the basis of these data, we hypothesize that several T3Es may have been horizontally transferred between *X. citri* pv. *bilvae* and some *Xci* strains in the Indian subcontinent and that *xopAG*-containing strains may have subsequently spread to Oman. This result is consistent with the very close genetic relatedness between the *xopAG*-containing strains JF90-8* from Oman and LG116* from India.

The monophyletic pathovar *Xci* is an interesting model for the study of the contribution of the T3E repertoire to the intrapathovar structure and host range variations reported among pathotypes. We showed that, at the pathotype level, the T3E repertoire did not explain *Xci* host range specialization. Previous studies have shown that, in other *Xanthomonas* species, including *X. axonopodis*, *X. arboricola* and *X. oryzae*, the composition of the T3E repertoires and the host specificity of pathovars are correlated (Hajri *et al.*, 2009). A possible limitation is that we worked on Xops previously identified in the genus *Xanthomonas*. Although these effectors were identified on the basis of avirulence activity, co-regulation with the T3SS or *in silico* prediction based on complete genome sequences (White *et al.*, 2009), there are probably other unsuspected T3Es to be discovered. Determinants of pathogenicity, other than those associated with the T3SS, could be involved in the specialization of the host range in *Xci*. However, the fact that *Xci* pathotypes A and A* induce canker symptoms on both limes and alemow suggests that common mechanisms allow them to escape both innate and specific host immunity, and to colonize citrus species to produce canker symptoms. The presence and/or functionality of a few or even single T3Es can be a founder event to colonize a new plant host (Schulze-Lefert and Panstruga, 2011). Therefore, the hypothesis of the role of T3Es in the definition of the host range of *Xci* pathotypes remains strong. Screening was made for the presence of T3Es, without considering whether or not a gene was functional. Alternatively, sequence allelic variation of a given T3E may also be involved in host specificity. Minute variations in these genes can have important implications for host range

and pathogenicity, as illustrated in *X. campestris* pv. *vesicatoria*, in which point mutations in *avrBs2* resulted in the breakdown of *Bs2*-based field resistance in pepper (Gassmann *et al.*, 2000), and for *P. syringae*, in which allelic variants of the T3E HopZ1 were shown to be differentially recognized by *Nicotiana benthamiana* and soybean (Zhou *et al.*, 2009), and where a key amino-acid polymorphism in *AvrPto* led the pathogen to overcome tomato resistance (Kunkeaw *et al.*, 2010). Host range changes have also been observed via inactivation of the T3E *avrBs2* by a 5-bp insertion in *X. euvesicatoria* (Swords *et al.*, 1996). We are now working on *Xci* draft genome sequences to assess the role of *xop* in the definition of the pathotypes by exploring their allelic variation. Access to *Xci* draft genome sequences might lead to the identification of new T3Es potentially involved in *Xci* pathogenicity. Together with the description of the phenotypic responses of *Xci* pathotypes presented here, the study of T3E sequences, combined with a better knowledge of housekeeping genes, will help us to better understand the mechanism underlying the interaction of the bacterium with its host, what determines the host range specialization and to provide clues about the evolution of the pathogenicity of *Xci* pathotypes. Our results also illustrated the dynamic nature of the host range of a pathogen (Schulze-Lefert and Panstruga, 2011). Working on a large collection of strains revealed the complexity of *Xci* pathotype A*, for which no correlation was found between the reported genetic diversity and the observed phenotypic responses. *Xci* host specialization may involve several evolutionary mechanisms affecting the capacity of a strain to have a narrow or a wide host range. It is therefore tempting to speculate that the diversity of *Xci* A* phenotypes described here corresponds to several steps in this dynamic process. Under this hypothesis, a better characterization of the clades which encompass A* Iranian strains with unexpected aggressiveness on sweet orange and grapefruit and A* strains causing no symptoms on both hosts should be of great interest, and might help us to understand the contribution of mutations or loss/gain of virulence genes in *Xci* host range specialization. Similarly, the close relatedness of some A*/A^W strains to A strains highlights that *Xci* variability should not be ignored when describing the mechanisms involved in pathogen evolution and host specialization.

EXPERIMENTAL PROCEDURES

Bacterial strain culture and DNA extraction conditions

The bacterial strains used in this study are listed in Tables S1 and S3 (see Supporting Information). All 55 *Xci* strains were isolated from citrus canker lesions on various *Citrus* species and collected from 21 different countries. These strains are a subset of the collection used previously (Bui-Thi-Ngoc *et al.*, 2009) and were selected in order to maximize the diversity in terms of host, year and country of isolation. *Xanthomonas citri* pv. *bilvae* strain NCPPB 3213, which is responsible for spot lesions on Rutaceae, was included in the study for comparative purposes. Positive and negative

controls for T3E screening consisted of *Xanthomonas* published genomes or draft genome sequences. *Xanthomonas* strains were stored and cultivated on YPGA (yeast extract 7 g/L, glucose 7 g/L, peptone 7 g/L, agar 18 g/L, pH 7.2), as described previously (Bui-Thi-Ngoc *et al.*, 2010). Suspensions were used for DNA extraction with the Wizard® Genomic DNA Purification Kit (Promega, Charbonnières, France). Rifampicin derivatives from strains IAPAR 306, JS584* and CFBP 2911* were selected on plating of YPGA supplemented with 1000 µg/mL rifampicin. For each strain, two isolates showing resistance to rifampicin were inoculated on detached leaves of Pineapple sweet orange, Mexican lime, Tahiti lime and Marsh grapefruit, and confirmed to be as pathogenic as the corresponding wild-type strains. One of each was selected for further experiments and referred to as IAPAR 306^R, JS584*^R and CFBP 2911*^R, respectively. Prior to triparental conjugation, *Xanthomonas* strains were grown at 28 °C on MOKA-rich medium (yeast extract 4 g/L, casamino acids 8 g/L, K₂HPO₄ 2 g/L, MgSO₄·7H₂O 0.3 g/L, agar 18 g/L, pH 7.2; Blanvillain *et al.*, 2007). *Escherichia coli* cells were cultivated on Luria–Bertani medium at 37 °C. Antibiotics were used at the following concentrations: for *Xci*: rifampicin, 100 µg/mL; kanamycin, 50 µg/mL; tetracycline, 10 µg/mL; chloramphenicol, 10 µg/mL; for *E. coli*: ampicillin, 50 µg/mL; chloramphenicol, 100 µg/mL; kanamycin, 25 µg/mL; tetracycline, 10 µg/mL; spectinomycin, 40 µg/mL.

Pathogenicity tests

Detached leaf assays

Inoculations of the 47 strains (Table S1), submitted to pathogenicity tests on detached leaves, were performed as described previously (Vernière *et al.*, 1998) on five citrus species: Pineapple sweet orange (*C. sinensis*), grapefruit (*C. paradisi*), Mexican lime (*C. aurantifolia*), alemow (*C. macrophylla*) and Tahiti lime (*C. latifolia*). Leaves were inoculated with bacterial suspension containing 1×10^6 CFU/mL, and were observed for the development of symptoms at 5, 7, 10, 14 and 21 dpi. Each strain was inoculated on three leaves. The experiment was repeated at least twice, which made a total of at least 48 replicates on six different leaves for each strain–host combination. A subset of six strains was subsequently inoculated onto Mandarin temple (*C. temple*), Mandarin Fairchild SRA30 (*C. reticulata*), Meyer lemon (*C. meyeri*) and Ortanique tangor (*C. reticulata* × *C. sinensis*) to further describe the atypical pathogenicity of Iranian strains and to compare them with the typical A strain responses and with another A* strain (Table S1).

Disease intensity was evaluated visually, and scored using a scale of 0–5 as follows: 0, no symptoms; 0.5, water-soaked margin surrounding the wound sites; 1, more extensive water soaking (diameter \geq 1 mm); 2, small pustules with no visible rupture of the epidermis or isolated pustules with rupture of the epidermis; 3, numerous pustules with visible rupture of the epidermis; 4, small canker (observation of callus-like material); 5, wide canker formation (at least twice the diameter of the pinprick). Detailed illustrations of these symptoms are presented in Fig. S2. Although no typical HR could be visualized using the detached leaf assay, defence-like reactions that appeared over time were also recorded (Fig. S1). To assess the global response of each tested strain, a DDi was calculated as the area under the disease progression curve (see the 'Data analysis' section below for further details). DDis on grapefruit, combined with the DDis of each of three other species (sweet orange, alemow or Mexican lime), were then plotted to give an overview of *Xci* pathogenicity using this mode of inoculation.

Attached leaf assays

All plants for leaf infiltration assays were pruned to produce uniformly aged shoots. Twenty-four strains were submitted to infiltration to look for HR or symptom development (Table S1). Leaves of Mexican lime, Pineapple sweet orange, Marsh grapefruit and Page mandarin were inoculated by infiltration of bacterial suspensions containing 1×10^8 and 1×10^5 CFU/mL, as described previously (Ah-You *et al.*, 2007). Symptom development was evaluated at 4, 6, 8, 10 and 14 dpi, and HR was scored at 4 and 6 dpi.

In planta bacterial population sizes were determined from infiltrated leaves (1×10^5 CFU/mL) of Mexican lime and Washington Navel sweet orange, as described previously (Bui-Thi-Ngoc *et al.*, 2010) in two experiments: (1) the first aimed to characterize the atypical interaction of cluster II A* strains from Iran and strain JS584* (selected as a reference for cluster II) in comparison with A strain IAPAR 306 and A* strain CFBP 2911*; (2) mutant strains and their corresponding rifampicin-resistant derivatives were compared. Each experiment was performed twice. Inoculated leaf fragments were sampled at 8 and 24 dpi. *Xanthomonas* enumerations were performed on YPGA-KC semi-selective medium (YPGA complemented with 20 mg/L of kasugamycin and 40 mg/L of cephalexin; Pruvost *et al.*, 2005) according to the method described earlier (Ah-You *et al.*, 2007; experiment 1), or on YPGA-KC and YPGA supplemented with 100 mg/L rifampicin (YPGAR; experiment 2).

T3E screening

We screened for the presence of 66 *Xanthomonas* T3Es (Xop) described on the *Xanthomonas* resource website (<http://www.xanthomonas.org/>; R. Koebnik, personal communication). The screen was performed using two pairs of primers for each Xop (Table S4). Screening of the 26 Xops found in *Xci* strain IAPAR 306 was performed using pairs of primers targeting different regions of the T3E genes. One of the two pairs of primers was designed to amplify at least 80% of the open reading frame (ORF), except for *xopAD*, which is too long (8652 bp) to be amplified in a single reaction, for which we amplified two fragments of 1660 and 608 bp. Screening of *xopAG* (*avrGf1*) was performed using primers designed on the sequence of *avrGf1* (accession number GI: 82571048, LOCUS DQ275469). In order to screen the 40 remaining effectors absent in *Xci* IAPAR 306, primers were designed in conserved regions based on the alignments of Xop orthologues from *Xanthomonas* available sequences to avoid false-negative results caused by point mutations or length differences between orthologues. Positive and negative controls are listed in Table S1. PCR amplification was verified *in silico* (<http://insilico.ehu.es/PCR/Amplify.php>) on the available *Xanthomonas* sequenced genomes. All PCR runs were performed with a GeneAmp PCR system 9700 thermocycler (Applied Biosystems, Saint Aubin, France). PCRs were performed in 20- μ L reaction mixtures containing $1 \times$ Gotaq@ green buffer (Promega), 1.5 mM MgCl₂, 200 μ M of each deoxynucleoside triphosphate (dNTP), 0.3 μ M of each primer, 2 ng of template genomic DNA and 0.8 U of GoTaq@ Polymerase. The amplification programme consisted of 35 cycles of denaturation at 95 °C for 45 s, annealing at 55 °C for 45 s and extension at 72 °C for 0.5 to 2 min, depending on the length of the PCR product (1 kb/min). Positive and negative controls were systematically included in the PCR amplification. PCR amplicons of the expected size were considered as positive responses for the presence of each marker. Amplified PCR products showing an unexpected size were sequenced to confirm identity.

DNA sequencing

All PCR products to be sequenced were sent to Beckman Coulter Genomics (Stansted, Essex, UK). They consisted of Xop ORF and flanking regions for mutagenesis, and fragments from the screen with unexpected size. Sequence alignments and sequence analyses were performed using Geneious software (v5.5.6; Biomatters Ltd, Auckland, New Zealand). Sequences were submitted to the nucleotide BLAST program available online (<http://www.ncbi.nlm.nih.gov/blast/Blast.cgi>; accessed July 2012) when necessary.

Southern blot hybridization

DNAs (10 μ g) from the 56 strains of the collection were digested with *Bam*HI (New England Biolabs, Evry, France), according to the supplier's instructions. DNA fragment separation and blotting were performed as described previously (Gagnevin *et al.*, 1997). Hybridization probes were obtained by PCR with primers used for the screen, under the same PCR conditions (Table S4). Probe labelling, hybridization and detection were performed with the ECL kit (Amersham ECL Direct Nucleic Acid Labelling and Detection Systems, GE Healthcare Limited, Amersham, Buckinghamshire, UK), according to the manufacturer's instructions, under moderate stringency. Blots were then exposed to Hyperfilm™ ECL for several minutes. Southern blot was preferred to dot blot to prevent false-positive results.

Construction of *Xci* mutants

Deletion mutant strains were constructed using the *sacB* system (Schäfer *et al.*, 1994). Briefly, 400–600-bp upstream and downstream regions of the full-length targeted gene were amplified by PCR on the DNA of the targeted strain as template. PCR for plasmid construction was performed with PHusion® High Fidelity DNA polymerase (Finnzymes, Waltham, MA, USA) following the manufacturer's instructions. Both amplicons were then cloned into GoldenGate-compatible nonreplicative suicide plasmid pK18::*sacB* (L. Noël, CNRS, Castanet-Tolosan, France, unpublished data; Schäfer *et al.*, 1994). GoldenGate is a cloning method based on the use of type II restriction enzymes (*Bsa*I in our study), which cuts outside of the enzyme recognition sites (Engler *et al.*, 2008). On triparental conjugation using pRK2073 as a helper (Leong *et al.*, 1982), primary transformants were selected on MOKA medium supplemented with kanamycin. Then, Kan^R colonies were plated onto MOKA medium containing 10% sucrose to select for second recombination. Colonies found to be sensitive to kanamycin and resistant to 10% sucrose were assayed by PCR for deletion of the targeted gene. Primers used for deletion are listed in Table S5.

AFLP analysis

The experiments were performed in 96-well plates in a GeneAmp PCR system 9700 thermocycler (Applied Biosystems), as described previously (Ah-You *et al.*, 2007; Bui-Thi-Ngoc *et al.*, 2010). Four AFLP conditions, each involving the two restriction enzymes *Msp*I and *Sac*I and two selective nucleotides, were used; two independent DNA extractions were used for all strains, and strain IAPAR 306 was used as a control in each experiment. The amplified fragments were scored as present (1) or absent (0) to create binary matrices. Only bands with an intensity above a preset level (500 relative fluorescence units) were scored.

Data analysis

Data were analysed using R software (version R 2.15.0; R Development Core Team, Vienna, Austria). Evolutionary genome divergence (EGD) values were calculated on the basis of Dice similarity indices (Mougel *et al.*, 2002) and were used as distances to construct a weighted NJ tree with the 'ape' package version 3.0–2 in R (Paradis, 2006). The robustness of the tree was assessed by bootstrap (1000 resamplings). MDS was used to represent distances between strains based on the Dice dissimilarity matrix. MDS transforms a distance matrix (which cannot be analysed by eigendecomposition) into a cross-product matrix, and then solves the eigenvector problem to find the coordinates of individuals, so that distortions to the distance matrix are minimized. As in principal component analysis, individuals are projected into n dimensions (Abdi, 2007). MDS was performed using the cmd-scale function. Populations *in planta* were analysed by analysis of variance (ANOVA). Whenever F values were significant ($P = 0.05$), mean population sizes were compared using Tukey's contrasts at level 0.05. The area under the disease progress curve (AUDPC) was calculated using the audpc function from the package agricolae. Hierarchical agglomerative clustering was performed on a simple matching dissimilarity matrix based on a matrix of T3E presence/absence built after homology searches of the T3E candidate list on 28 *Xci* draft genome sequences and genomes of *XauB* (accession number ACPX00000000) and *XauC* (accession number ACPY00000000). We used Ward's method implemented in 'cluster' package version 1.14.2 in R (Maechler *et al.*, 2012).

ACKNOWLEDGEMENTS

We thank C. Boyer and F. Maillot for their technical expertise, and F. Chiroleu and P. Lefeuvre for advice on data analysis. The European Union (FEDER), Conseil Régional de La Réunion and Centre de Coopération Internationale en Recherche Agronomique pour le Développement provided financial support. Financial support for LDN was provided by the LABEX TULIP (ANR-10-LABX-41) and an Agence Nationale de la Recherche–Jeunes Chercheurs grant (Xopaque ANR-10-JCJC-1703-01).

GenBank accession numbers for original sequences in this work are as follows: ISXac5 from *Xci* strain NCPPB 3608: JX566665; *xopAG* from *Xci* strain NCPPB 3608: JX566667; *xopC1* from *Xci* strain JK2-10: JX566666; *xopAG* from *X. citri* pv. *bilvae* strain NCPPB 3213: JX556152.

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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 Amplified fragment length polymorphism (AFLP)-based metric multidimensional scaling (MDS).

Fig. S2 Disease scale used in this study.

Fig. S3 Disease severity evolution on detached leaves.

Fig. S4 Hypersensitive response (HR) symptoms induced by A^w and A* strains containing *xopAG*.

Fig. S5 Symptomatology of A and A* mutants.

Fig. S6 *xopC1* regions.

Table S1 Bacterial strains used in this study.

Table S2 *In planta* populations of mutants and corresponding rifampicin-derivative strains.

Table S3 Bacterial strains and plasmids used for mutagenesis experiments.

Table S4 Primers for polymerase chain reaction (PCR) amplifications of *Xanthomonas* type III effector (T3E) genes.

Table S5 Polymerase chain reaction (PCR) primers for PCR amplification for deletion of *Xanthomonas* type III effector (T3E) genes.