

ORIGINAL ARTICLE

Positive selection is the main driving force for evolution of citrus canker-causing *Xanthomonas*Yunzeng Zhang^{1,5}, Neha Jalan^{1,5}, Xiaofeng Zhou¹, Erica Goss², Jeffrey B Jones², João C Setubal³, Xiaoling Deng⁴ and Nian Wang¹¹Citrus Research and Education Center, Department of Microbiology and Cell Science, University of Florida, Lake Alfred, FL, USA; ²Department of Plant Pathology, University of Florida, Gainesville, FL, USA;³Departamento de Bioquímica, Instituto de Química, Universidade de São Paulo, São Paulo, Brazil and⁴Department of Plant Pathology, South China Agricultural University, Guangzhou, Guangdong, China

Understanding the evolutionary history and potential of bacterial pathogens is critical to prevent the emergence of new infectious bacterial diseases. *Xanthomonas axonopodis* subsp. *citri* (Xac) (synonym *X. citri* subsp. *citri*), which causes citrus canker, is one of the hardest-fought plant bacterial pathogens in US history. Here, we sequenced 21 Xac strains (14 XacA, 3 XacA* and 4 XacA^W) with different host ranges from North America and Asia and conducted comparative genomic and evolutionary analyses. Our analyses suggest that acquisition of beneficial genes and loss of detrimental genes most likely allowed XacA to infect a broader range of hosts as compared with XacA^W and XacA*. Recombination was found to have occurred frequently on the relative ancient branches, but rarely on the young branches of the clonal genealogy. The ratio of recombination/mutation ρ/θ was 0.0790 ± 0.0005 , implying that the Xac population was clonal in structure. Positive selection has affected 14% (395 out of 2822) of core genes of the citrus canker-causing *Xanthomonas*. The genes affected are enriched in ‘carbohydrate transport and metabolism’ and ‘DNA replication, recombination and repair’ genes ($P < 0.05$). Many genes related to virulence, especially genes involved in the type III secretion system and effectors, are affected by positive selection, further highlighting the contribution of positive selection to the evolution of citrus canker-causing *Xanthomonas*. Our results suggest that both metabolism and virulence genes provide advantages to endow XacA with higher virulence and a wider host range. Our analysis advances our understanding of the genomic basis of specialization by positive selection in bacterial evolution.

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Introduction

Approximately 160 emerging infectious bacterial diseases have been discovered in the past 70 years (Jones *et al.*, 2008). The recent increase in emerging bacterial diseases is partly because of agricultural practices and climate change, among other factors (Wolfe *et al.*, 2007; Jones *et al.*, 2008). The agroecosystem has played a critical role in the recent emergence and spread of pathogenic bacteria in plants. Prevention of emergence of new bacterial diseases is one essential task of the scientific community. Emergence of plant bacterial diseases depends largely on the life history traits and evolutionary potential of the corresponding bacteria.

Among plant pathogenic bacteria, those of the genus *Xanthomonas* in the gamma division of Proteobacteria can cause disease in virtually all plant species (Leyns *et al.*, 1984; Chan and Goodwin, 1999). Individual members of *Xanthomonas* are highly host specific (Hayward, 1993). Evolutionary analysis of *X. axonopodis*, which is a complex species composed of multiple pathovars (host-specific subspecies), identified two main diversification steps based on sequence analysis of selected loci including seven housekeeping genes and several virulence-associated genes (Mhedbi-Hajri *et al.*, 2013). The first diversification led to a clustering of generalist pathogens over the past 25 000 years with no apparent connection to host or geography. The second step led to specialized pathotypes grouped according to their host range and their symptomatology over the past two centuries. Eventually, secondary contacts likely occurred between host-specialized bacteria that enabled genetic exchange of virulence-associated genes. For pathogens, acquisition of virulence genes is not the end, but instead, the beginning of the process of host

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specialization (Rohmer *et al.*, 2011). During the specialization process, these pathogens experience new niches and strong directional selection from hosts, and eventually the more adapted lineages will emerge and sweep through evolution, including accumulating beneficial mutations, acquiring beneficial genes, losing antivirulence genes and so on (Lieberman *et al.*, 2011; Rohmer *et al.*, 2011). The process of host specialization of *Xanthomonas* has not been well understood. In this study, we investigated the evolutionary history and potential of *X. axonopodis* pv. *citri* (Xac) (synonym *X. citri* subsp. *citri*) by genome-wide analysis.

Xac is the causal agent of citrus canker that infects most commercial citrus cultivars and causes distinctive raised necrotic lesions on leaves, stems and fruit. Severe infections can cause defoliation, blemished fruit, premature fruit drop, twig dieback and general tree decline (Gottwald *et al.*, 2002). Citrus canker is distributed worldwide and found in most citrus-producing countries. Importantly, it is endemic in the top three citrus-producing countries: Brazil, United States and China. Owing to its importance, citrus canker is one of the hardest-fought plant bacterial diseases in the US history. Citrus originated in Southeast Asia, that is, North-east India, Burma and the Yunnan province of China (Scora, 1975; Gmitter and Hu, 1990). Xac is also suggested to have originated in the same area (Gottwald, 2000). Xac was introduced into the United States in at least three different events in 1910, 1986 and 1995. These introductions were followed by eradication programs, in which billions of dollars were spent (Gottwald and Irey, 2007).

Xac consists of multiple pathotypes, that is, A, A* and A^w. XacA is widespread worldwide and can infect many citrus species, hybrids between citrus species and the citrus relative trifoliolate orange *Poncirus trifoliolate* as well as some other plants in the Rutaceae family (Graham *et al.*, 2004), whereas XacA^w and XacA* have a restricted host range of Mexican lime (*Citrus aurantifolia*) and alemow (*Citrus macrophylla*) (Sun *et al.*, 2004). However, the genetic basis of the host range and virulence difference between them has not been well determined (Jalan *et al.*, 2013a).

In addition to Xac, *X. fuscans* subsp. *aurantifolii* (Xau) (previously named as *X. axonopodis* pv. *aurantifolii*) also causes citrus canker disease. Xau causes canker B and canker C on restricted hosts and these strains have only been found in South America (Graham *et al.*, 2004). Given that Xau B and C have evolved to be citrus canker-causing pathogens from a separate lineage from Xac strains (Mhedbi-Hajri *et al.*, 2013), analysis of how these *Xanthomonas* pathogens have evolved to specialize on citrus and associated relatives to cause citrus canker disease via convergent evolution may reveal common mechanisms underlying this disease.

The evolution of Xac has only been examined using multilocus sequence analysis or genotyping

(Ngoc *et al.*, 2009; Mhedbi-Hajri *et al.*, 2013). To investigate the evolutionary history and potential of Xac, we sequenced 21 representative Xac strains from North America and Asia. We included strains from the XacA* and XacA^w pathotypes to identify potential mechanisms underlying the host range difference within Xac. The importance and relative contribution of recombination and mutation in evolutionary history of Xac was analyzed. Furthermore, the importance of positive selection in the process of host specialization for the citrus canker-causing *Xanthomonas* was discussed.

Materials and methods

Strains

The strains for sequencing were chosen to represent Xac that are genetically and geographically diverse. We selected 21 strains of Xac from North America (United States) and Asia, with a time span of over two decades for genome sequencing (Supplementary Table S1). The complete genomes of strain A306 (da Silva *et al.*, 2002) and A^w12879 (Jalan *et al.*, 2013b) were also used in this analysis.

Sequencing, assembly and annotation

Quantity and quality of the DNA were measured using Agilent 2100 BioAnalyzer (Agilent Technologies, Inc., Santa Clara, CA, USA). Whole-genome sequencing of all strains was performed using 1/4 of a lane of paired-end 100-cycle sequencing using an Illumina genome analyzer Ix (Illumina, Hayward, CA, USA) at the Interdisciplinary Center for Biotechnology Research, University of Florida. In addition, for XacA*270 paired-end 454 pyrosequencing was performed using Roche 454 GS-FLX plus system (454 Life Sciences, Branford, CT, USA) at the Interdisciplinary Center for Biotechnology Research. A *de novo* BamHI optical map of the genome of XacA*270 was generated by OpGen Technologies (Madison, WI, USA). The draft genome of XacA*270 was closed using the method described in our previous study (Jalan *et al.*, 2011).

All reference-based assemblies were performed on CLC Genomics Workbench 6.0 (CLC bio, Cambridge, MA, USA), using default parameters, a length fraction of 0.9 and similarity of 0.9 for reference assemblies to XacA306, XacA^w12879 and XacA*270. Assembly of all strains was annotated using the 'isolated genome gene calling' pipeline from the Integrated Microbial Genomes-Expert Review (IMG/ER; Markowitz *et al.*, 2009).

Comparative genomic analysis

The ANIm values between genomes were calculated using the NUCmer algorithm v3.1 integrated in Jspecies v1.2.1 (Richter and Rossello-Mora, 2009). The pan-genome and core genome were calculated by using OMCL and bidirectional best hit (BDBH)

methods implemented in *get_homologues* package (Contreras-Moreira and Vinuesa, 2013) with parameters: *e*-value: $1e-5$, identity: 60% and coverage: 75%. For pan-genome size calculation, the power law model was employed. In short, according to the power law model, the new genes $N(n)$ found when the n th genome is added to the pan-genome can be described as $N(n) = A \times n^\alpha$, and the pan-genome is 'open' if $\alpha > 0$, is 'logarithmical' if $\alpha = 0$ or is 'closed' if $\alpha < 0$, while A is a constant (Donati *et al.*, 2010). The core genes were aligned using MUSCLE (Edgar, 2004) and concatenated using Gblocks0.91 (Castresana, 2000). The best nucleotide substitution model was obtained by MODELTEST analysis (Posada and Crandall, 1998), and the subsequent maximum likelihood phylogenetic tree was constructed by using PAUP4b10 (Swofford, 1998).

To calculate the ratio of recombination rate to mutation rate (ρ/θ) and the relative contribution of recombination and mutation (r/m), a whole-genome sequence alignment of the selected strains was created using progressiveMauve (Darling *et al.*, 2010) and the sequences of the first nine longest alignments (3.06 Mb, ~58% of the whole genomes) for the 23 *Xac* strains or three longest alignments (1.77 Mb, ~33% of each genome) for the 25 strains (23 *Xac* plus *XauB* and *XauC*) were concatenated using Gblocks0.91 (Castresana, 2000), and then calculated by using ClonalFrame v1.2 (Didelot and Falush, 2007). Four independent runs of ClonalFrame were performed, each consisting of 40 000 iterations, and the first half was discarded as MCMC burn-in. The convergence and mixing properties revealed by manual comparison between the four runs suggested that the obtained parameters were confident. The presence or absence of intragenic recombination was assessed by the single break-point method (Pond *et al.*, 2006) ($P < 0.05$). Fast Unconstrained Bayesian Approximation (FUBAR) (Murrell *et al.*, 2013) implemented in HYPHY2.2 package was used to evaluate the contribution of positive selection on the citrus canker-causing *Xanthomonas*. For each gene cluster, 400 Grid points were used, and five independent runs of FUBAR were performed, each consisting of 2 000 000 iterations. The first half was discarded as MCMC burn-in.

Effector prediction was conducted as previously described (Bart *et al.*, 2012). In short, a data set of all known effectors from animal and plant pathogens was obtained and *tblastn* (Blast+ 2.2.28) was used to identify potential effectors in *Xac* strains with at least 45% amino acid homology and 80% coverage length.

LPS extraction and host response analysis

Lipopolysaccharide (LPS) extractions were performed as previously described with modifications (Marolda *et al.*, 2006). Briefly, bacterial cells from 20 ml culture ($OD_{600} = 1$) were centrifuged and resuspended in 500 μ l distilled water. LPSs were

extracted with hot phenol (70 °C) three times and aqueous phases were subjected to dialysis against distilled water at 4 °C (Pur-A-Lyzer Midi 1000 Dialysis kit, cutoff = 1 kDa, Sigma-Aldrich, St Louis, MO, USA). Prepared LPSs (~50 μ l per leaf) were infiltrated into plant leaves using a needleless syringe. Leaf samples were collected at 0 (right after treatment), 6 and 24 h post inoculation. Time point 0 was used as the control. The expression levels of host defense-related genes were performed using a QuantiTect SYBR green RT-PCR kit (Qiagen, Valencia, CA, USA) on a 7500 fast real-time PCR system (Applied Biosystems, Foster City, CA, USA). Citrus *GAPDH* gene was used as an internal control. The relative fold change was calculated as previously described (Livak and Schmittgen, 2001).

Data access

The finished genomes of 14 *XacA* and 4 *XacA^w* strains and draft genomes of 3 *XacA^{*}* strains studied in this project have been deposited in GenBank under the project accession number PRJNA255042 (Supplementary Table S1).

Results and discussion

Genomic features of citrus canker-causing *Xac* strains

We sequenced 21 representative *Xac* strains (14 *XacA* strains, 3 *XacA^{*}* strains and 4 *XacA^w* strains) encompassing three countries (United States, China and Saudi Arabia), and isolated over a period of more than two decades (Supplementary Table S1). The reads from *XacA* and *XacA^w* strains were reference assembled against *XacA306* and *XacA^w12879* genome (da Silva *et al.*, 2002; Jalan *et al.*, 2013b), respectively. As no *XacA^{*}* genome was publically available, *De Novo* assembly was conducted for the genome of *XacA^{*}270* (Supplementary Table S2) using previously described methodology (Jalan *et al.*, 2011), and the two remaining *XacA^{*}* strains were reference assembled based on the *XacA^{*}270* genome. The comparison of our reads and assemblies (Supplementary Table S3) with other published assemblies for *Xanthomonas* (Bart *et al.*, 2012) indicated that our assemblies are of high quality. Assembly of all strains was annotated using the IMG/ER annotation pipeline and submitted to GenBank, the results for which are presented in Supplementary Table S1.

The ANIm values of the 23 *Xac* strains (21 strains sequenced in this study and *XacA306* and *XacA^w12879* sequenced previously; da Silva *et al.*, 2002; Jalan *et al.*, 2013b) were 99.65–99.94% (Supplementary Table S4), demonstrating highly conserved genomic backgrounds among these strains, despite differences in host ranges. To reveal the mechanisms underlying differential host ranges of the three pathotypes A, A* and A^w, we calculated the pan-genome of the 23 *Xac* strains using the

OMCL method. The core genome comprised 3912 orthologous clusters, whereas the pan-genome contained 5147 orthologous clusters. Interestingly, hierarchical clustering of the accessory orthologous clusters (that is, the clusters not harbored by all of the 23 strains) of these Xac strains produced a distribution in which the XacA^w and XacA* strains were grouped together and formed a separate clade from the XacA strains (Figure 1), consistent with the phylogenetic tree constructed based on the Xac/Xau core genomes (Figure 2 and Supplementary Figure S1). Given that XacA strains have wide host ranges whereas the host range of XacA^w and XacA* strains is restricted, we infer that the clade-specific genes probably contributed to the host range difference between the XacA group and XacA^w and XacA* groups. Through comparative genome analysis, 89 XacA- and 121 XacA^w- and XacA*-specific genes were identified (Supplementary Data Set 1).

The gene *avrGf1* (*xopAG*), which is known as the type III effector restricting the host range of XacA^w strains (Rybak *et al.*, 2009), was absent in the three XacA* strains. A previous study also reported that some XacA* strains harbor the *avrGf1* gene whereas others do not (Escalon *et al.*, 2013). Furthermore, the XacA^w Δ *avrGf1* mutant obtained the ability to infect more citrus species and cause canker on hosts other than limes, but the virulence was not as strong as XacA strains (Rybak *et al.*, 2009), suggesting that other genomic feature(s) might contribute to the host range and virulence difference between XacA and XacA^w. Among the type III effector genes identified in the 23 Xac genomes, only one effector, *xopAF*, was found to be common and specific to XacA^w and XacA* (Supplementary Table S5); however, mutation of *xopAF* in XacA^w12879 resulted in lower virulence and no change of host range (Jalan *et al.*,

2013a), suggesting that *xopAF* is not related to the host range determination. Two other effectors, *xopC1* and *xopL*, were found to differentiate XacA from A^w or A*; *xopC1* was specifically harbored by the three XacA* strains whereas all the 23 strains harbored *xopL*, but it was frameshift mutated in XacA* strains (Supplementary Table S5). However, *XopC1* and *XopL* do not contribute to virulence or host range determination (Dunger *et al.*, 2012; Escalon *et al.*, 2013). Thus, there should be other mechanisms other than the known type III effectors underlying the difference in host range and virulence between XacA and XacA^w and XacA* strains. Notably, compared with the specific genes belonging to XacA^w and XacA*, XacA-specific genes were significantly enriched in the COG category 'metabolism' (Fisher's exact test, $P = 0.007$; Table 1). Therefore, we reasoned that XacA-specific genes, especially the 'metabolism' genes, provide advantages to endow XacA with higher virulence and a wider host range (Rohmer *et al.*, 2011).

We identified a gene region involved in LPS biosynthesis (J151_03781-J151_03787, equal to XAC3596-XAC3601, totally 7 genes, *nlxA* is recently determined in the intergenic region of XAC3597 and XAC3598; Yan *et al.*, 2012) in XacA that was highly variable from that of XacA^w and XacA*, but the upstream and downstream genes were conserved (Figure 3a). Three of the genes in this region, *nlxA*, *wzt* and *wzm*, are 'metabolism' genes. This variable region in XacA, XacA^w and XacA* was all identified as acquired genomic content via horizontal transfer as predicted by both Alien_Hunter and Island-Viewer programs (Vernikos and Parkhill, 2006; Langille and Brinkman, 2009), suggesting XacA acquired this region from different source(s) as compared with XacA^w and XacA*. Some of the

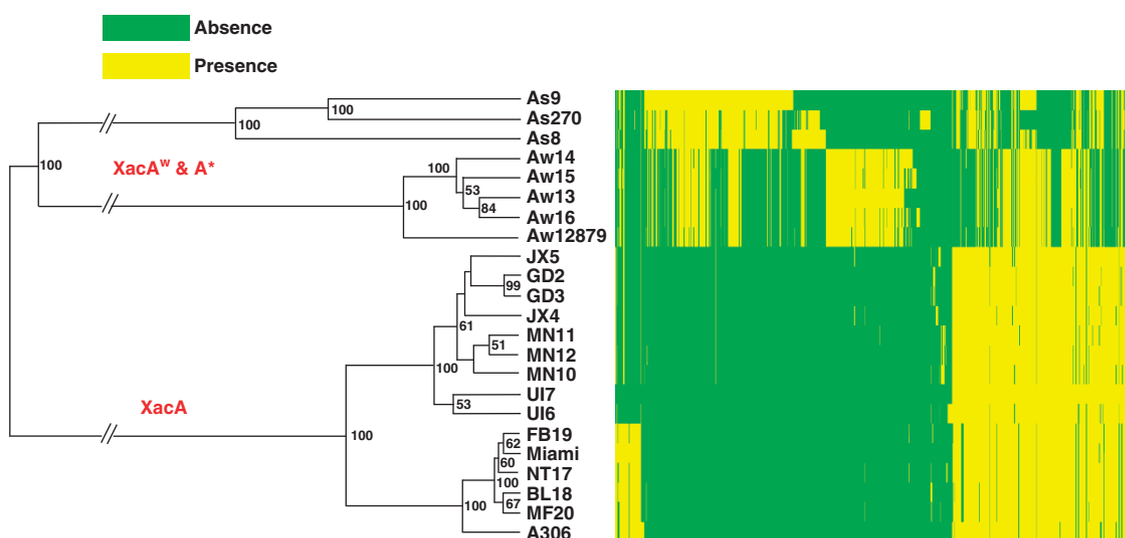


Figure 1 Hierarchical clustering of 23 Xac strains based on heat map of 1235 accessory orthologous clusters. Presence and absence of the homolog for each cluster are indicated in yellow and green, respectively. A total of 100 bootstrap replicates were made, and bootstrap values of >50% were indicated at each node. The UPGMA tree was generated by using DendroUPGMA (<http://genomes.urv.es/UPGMA/>) with Jaccard coefficient.

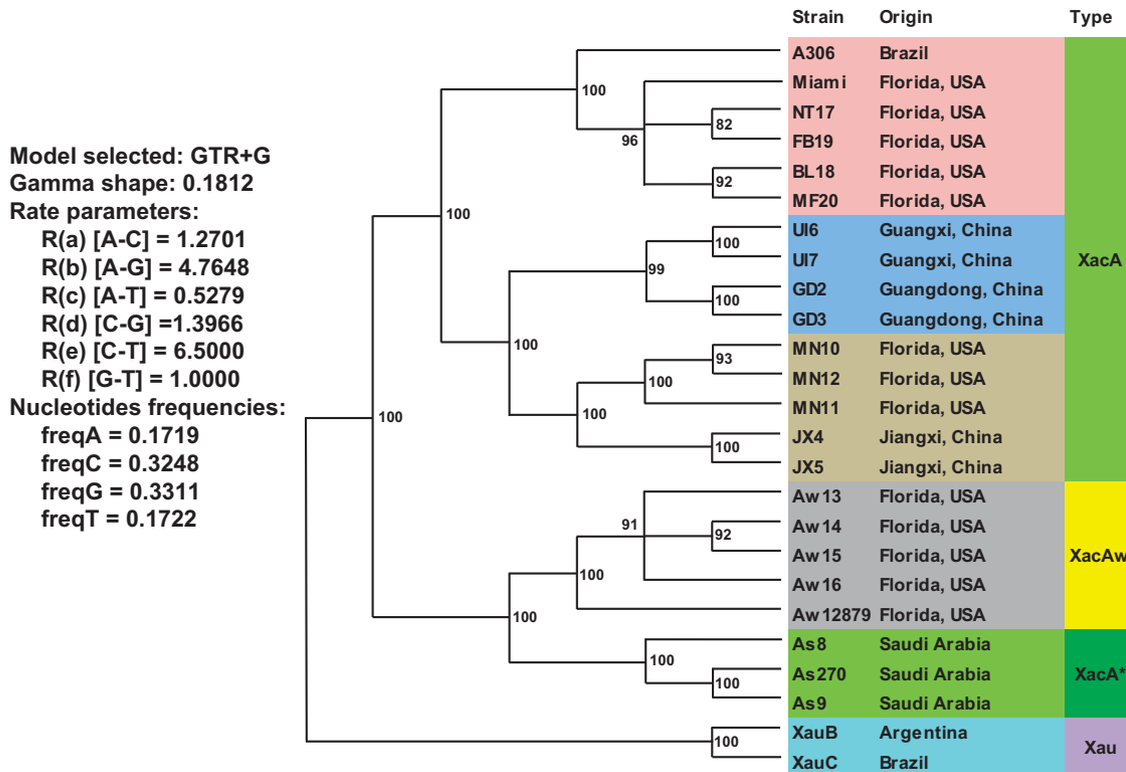


Figure 2 Maximum likelihood (ML) tree reconstructed based upon the concatenated sequences of 2822 core genes of the 25 citrus canker-causing *Xanthomonas* strains. A total of 100 bootstrap replicates were made, and bootstrap values are indicated at the branch points. The parameters on the left were calculated using Modeltest 3.7 and were used by PAUP4.0b10 to construct the ML tree.

genes in this region, such as *wxacO*, *rbcC*, *nlxA* and *wzt*, have been demonstrated to be involved in O-antigen synthesis and transport and be associated with biofilm formation on the host and contribute to virulence in XacA (Casabuono *et al.*, 2011; Li and Wang, 2011; Petrocelli *et al.*, 2012; Yan *et al.*, 2012). Among the three components, lipid A, core and O-antigen that are composed of the entire Xac LPS, the O-antigen region is probably the most important for host basal response during citrus-pathogen interaction (Casabuono *et al.*, 2011). O-antigen is highly variable with regard to its composition, length and the branching of carbohydrate subunits, whereas the core and lipid A are conserved among different bacterial species (Nicaise *et al.*, 2009). LPS analysis by SDS-polyacrylamide gel electrophoresis clearly demonstrated that the O-antigen region of XacA group was different from that of XacA^w (Figure 3b). We hypothesized that the O-antigen differences between XacA and XacA^w or A* group may induce different defence responses in different host plants. As shown in Figure 3c, LPS isolated from XacA and XacA^w acted as pathogen-associated molecular pattern and induced the gene expression of pathogen-associated molecular pattern-triggered immunity markers *GST1* and *WRKY22* (Asai *et al.*, 2002), salicylic acid metabolism *PAL1* (Greenberg *et al.*, 2009) and pathogen-associated molecular pattern-triggered immunity signaling kinase *MKK4* (Zhao *et al.*, 2014) in both sweet orange and

Mexican lime leaves. Defence-related gene expression triggered by XacA LPS is significantly lower than that triggered by XacA^w LPS in sweet orange. This is consistent with the fact that XacA^w is not pathogenic on sweet orange whereas XacA causes disease on sweet orange. LPS of XacA or XacA^w induced the gene expression of *GST1*, *WRKY22*, *PAL1* and *MKK4* in Mexican lime leaves despite both strains are pathogenic on Mexican lime. The defence-related gene expression triggered by XacA LPS is slightly higher than or at similar level as that triggered by XacA^w LPS in Mexican lime. This indicates that both XacA and XacA^w could suppress the pathogen-associated molecular pattern-triggered immunity induced by LPS in Mexican lime, probably using type III effectors (Jones and Dangl, 2006).

In addition, several genes that have been shown to contribute to biofilm formation, including XAC1469, XAC3285, XACb0003-0004 and XACb0050, were also found to be specific to XacA strains (Laia *et al.*, 2009; Li and Wang, 2011; Malamud *et al.*, 2013). It remains to be determined how the specific genes other than known type III effectors for XacA, A* and A^w contribute to the host range of Xac.

Evolution of the core and accessory genomes of Xac

We hypothesize that XacA evolved to be a separate group from XacA^w and XacA*. Our evidence is that

Table 1 COG distribution of Xac-specific genes and positive selection-affected genes of the citrus canker-causing *Xanthomonas*

Individual functional categories	<i>XacA</i> group-specific genes ^a			Core genome of 25 citrus canker-causing strains			
	<i>XacA</i>	<i>A^w</i> and <i>A[*]</i>	P-value ^b	Positive selection ^c	Core genome	P-value ^b	
B: Chromatin structure and dynamics	0	8	0	15	0.5	0	0
J: Translation, ribosomal structure and biogenesis	1	1	1	21	139	0.61	
K: Transcription	5	7	7	19	169	0.56	
L: Replication, recombination and repair	2	7	7	22	103	0.047	
D: Cell-cycle control, cell division, chromosome partitioning	2	13	0	23	0.46	3	29
V: Defense mechanisms	2	3	3	8	41	0.36	
T: Signal transduction mechanisms	1	7	7	22	182	0.74	
U: Intracellular trafficking, secretion and vesicular transport	0	1	1	8	90	0.33	
N: Cell motility	0	3	3	7	80	0.32	
O: Posttranslational modification, protein turnover, chaperones	4	3	3	22	129	0.3	
M: Cell wall/membrane/envelope biogenesis	4	6	6	22	187	0.65	
P: Inorganic ion transport and metabolism	2	16	0	7	0.007	20	163
F: Nucleotide transport and metabolism	0	0	0	5	61	0.33	
G: Carbohydrate transport and metabolism	4	3	3	38	183	0.017	
E: Amino-acid transport and metabolism	1	1	1	33	209	0.35	
H: Coenzyme transport and metabolism	3	2	2	17	126	1	
I: Lipid transport and metabolism	1	1	1	13	120	0.58	
C: Energy production and conversion	2	0	0	23	160	0.724	
Q: Secondary metabolites biosynthesis, transport and catabolism	3	0	0	9	57	0.56	
R: General function prediction only	12	15	11	14	0.31	39	316
S: Function unknown	3	3	3	23	238	0.13	
No COG	42	70	70	0.16	53	326	0.279

^aFor the Xac-specific genes, the individual Clusters of Orthologous Groups (COG) were combined into four categories as described by Popa *et al.* (2011). Gray colored, 'Information storage and processing'; light green colored, 'cellular processes and signaling'; light blue colored, 'metabolism'; light purple colored, 'unknown'. The COG annotation was assigned to XacA-specific genes based on XacA306 annotation (da Silva *et al.*, 2002). The COG annotation was assigned to XacA^w- and A^{*}-specific genes by blasting protein sequences against COG v1.0 database (<http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi>).

^bFisher's exact test.

^cThe COG distribution of the 395 genes under positive selection.

the XacA strains formed a separate clade from the XacA^w and XacA^{*} strains as revealed by the phylogenetic tree reconstructed based on the core genome (Figure 2 and Supplementary Figure S1), the phyletic distribution of accessory orthologous clusters (Figure 1) of the strains analyzed and the different origin of the LPS genomic island of the two groups. This finding is consistent with the previous report based on MLVA (multilocus variable number of tandem repeat Analysis) genotyping method (Pruvost *et al.*, 2014). The slight but significant difference of genome size (5.24 ± 0.02 Mb (mean \pm s.d., same for following ones) for XacA and 5.36 ± 0.05 Mb for A^w and A^{*} ($P = 0.0003$)), CDS number (4443 ± 22 for XacA and 4591 ± 39 for A^w and A^{*} ($P = 3.7E - 6$)) and percentage of GC content (64.754 ± 0.001 for XacA and 64.656 ± 0.001 ($P = 7.5E - 6$ for A^w and A^{*})) of XacA strains and XacA^w and XacA^{*} also suggested the divergence of the two groups (Supplementary Figure S2). During their divergent evolutionary history, the acquisition of beneficial genes and loss of genes that restrict the host range of the pathogens (for example, *avrGf1*) may have provided XacA advantages to become more adapted to diverse citrus hosts than XacA^w and XacA^{*} (Rohmer *et al.*, 2011; Merhej *et al.*, 2013). Noteworthy, the phyletic distribution of accessory genes was globally similar to the phylogeny of core

genomes, suggesting that the gain and loss of accessory genes for each strain was also under selection and might be driven by similar evolution mechanism as core genomes. In line with the positive selection of accessory genes, compared with XacA^w and XacA^{*}, the more adapted XacA strains were found to harbor much fewer 'No COG' genes as well as genes involved in signal transduction (COG category 'T') (Table 1). This phenomenon probably results from loss of useless and/or superfluous genes (for example, those involved in process such as signal transduction) for the pathogens after the pathogens have specialized on certain hosts (Merhej *et al.*, 2013). Furthermore, our data suggest that the pan-genome of the 23 Xac strains is closed ($A = 155.01 \pm 35.65$ and $\alpha = -1.69 \pm 0.09$). The depletion curves of core genome and accumulation curves of pan-genome shows that these 23 strains comprehensively sample the pan-genome of Xac (Supplementary Figure S3). These results suggest infrequent import of new genes from outside species to Xac and there was loss of genes (for example, those involved in process such as signal transduction) in the recent evolution of Xac during host specialization (Merhej *et al.*, 2013). One explanation for gene loss may be intensified citrus culture, allowing Xac to have a complete infection cycle on citrus without the need for an alternative host or

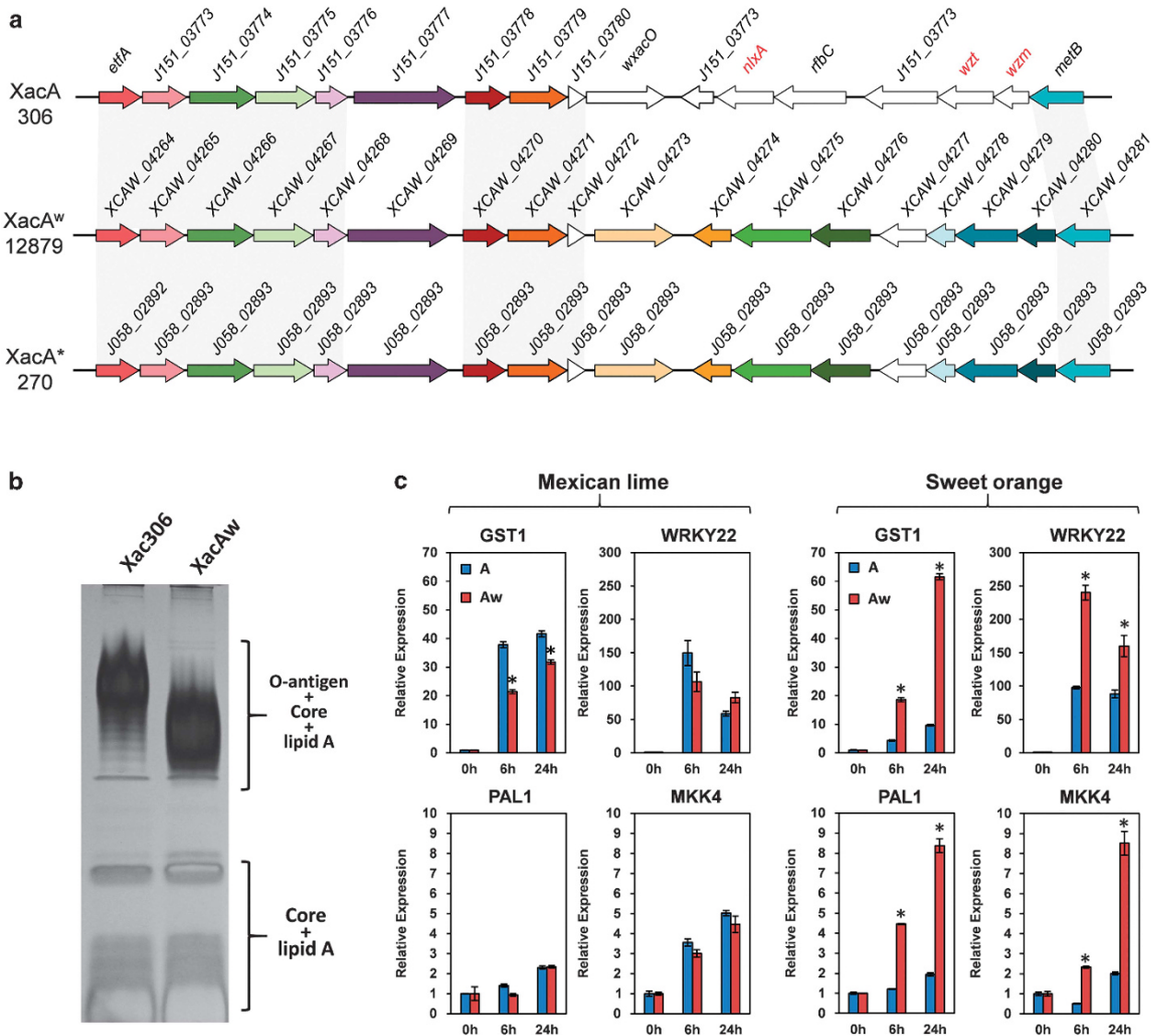


Figure 3 (a) Comparison of the LPS gene clusters of *X. axonopodis* subsp. *citri* str. A306 (XacA306), XacA^w12879 and XacA^{*}270. Conserved and homologous genes (>50% identity) are colored. The *nlxA*, *wzt* and *wzm* are 'metabolism' genes. (b) SDS–polyacrylamide gel electrophoresis (SDS–PAGE) analysis of LPSs extracted from XacA306 and XacA^w12879. LPSs were isolated using hot phenol method, subjected to analysis on polyacrylamide gel (12% in the resolving gel and 5% in the stacking gel) and visualized by silver staining. The different regions of LPSs are shown as indicated. (c) Expression analysis of four pathogen-associated molecular pattern-triggered immunity (PTI)-related genes using quantitative reverse-transcription PCR. Relative expressions (calculated using the formula $2^{-\Delta\Delta CT}$) were monitored from young orange leaves inoculated with XacA306 and XacA^w12879 LPSs. Leaves were sampled at 0, 6 and 24 h post inoculation. Bars are means \pm s.e. ($n = 3$). The experiment was repeated once with similar results. Student's *t*-test was conducted. The asterisks indicate a statistically significant difference ($*P < 0.01$ vs A strain).

survival in nonhost environments (Gottwald *et al.*, 2002). The constant citrus hosts provide a relatively stable environment and nutrition source, and environmental threats and antagonists would be relatively rare.

To evaluate the frequency and importance of recombination and mutation in the evolution of Xac, the ratio of recombination rate to mutation rate (ρ/θ) and the relative contribution of recombination and mutation (r/m) were calculated on the basis of ungapped sequences of the longest 9 aligned blocks of the 23 genomes, generated by progressive MAUVE (Darling *et al.*, 2010) (3.06 Mb, ~58% of

the whole genomes), using ClonalFrame (Didelot and Falush, 2007). The clonal genealogy calculated by ClonalFrame (Supplementary Figure S4) was consistent with the phylogenetic tree based on the core genome (Figure 2). The r/m value was 1.061 ± 0.006 , suggesting that recombination contributed as much as mutation to the observed diversity of Xac as recombination can introduce a segment of foreign DNA into the genome. However, recombination was found to have only occurred frequently on the relative ancient branches representing the ancestors of the XacA and XacA^{w/A*} lineages, but rarely on the young branches

Table 2 Shimodaira–Hasegawa (SH) analysis of gene phylogeny of individual sequences of 10 common type III effectors shared by all 25 citrus canker-causing *Xanthomonas* strains in comparison with the reference phylogeny based on core genome

Gene ^a	– ln L ^b	Diff – ln L ^c	P-value ^d	Positive selection ^e
<i>xopA_XAC0416</i>	4 740 381	91 517.66	0.000*	Y
<i>avrBs2_XAC0076</i>	4 663 871	15 008.52	0.000*	N
<i>xopE3_XAC3224</i>	4 768 447	119 583.9	0.000*	N
<i>hpaA_XAC0400</i>	4 768 447	119 583.9	0.000*	Y
<i>xopX_XAC0543</i>	4 667 601	18 738.44	0.000*	Y
<i>xopQ_XAC4333</i>	4 768 447	119 583.9	0.000*	N
<i>xopV_XAC0601</i>	4 659 360	10 496.79	0.000*	Y
<i>xopK_XAC3085</i>	4 659 796	10 933.24	0.000*	N
<i>xopAI_XAC3230</i>	4 663 627	14 764.36	0.000*	N
<i>xopE1_XAC0286</i>	4 768 447	119 583.9	0.000*	N
Core genome	4 648 863	(Best)		–

^aGene name and corresponding gene loci in XacA306.

^bln L, negative log-likelihood values correspond to those for the constrained topology.

^cScore differences between the unconstrained and constrained trees.

^dSH test. * $P < 0.05$.

^eUnder positive selection, ‘Y’ denotes yes, ‘N’ denotes no and ‘–’ denotes not detected.

(Supplementary Figure S4). The ratio ρ/θ is 0.0790 ± 0.0005 , implying that the Xac population was effectively clonal in structure (Fraser *et al.*, 2007). Although other genotyping methods have also suggested a clonal population structure for Xac (Ngoc *et al.*, 2009; Vernière *et al.*, 2014), these results imply that the Xac pathotypes have been clonal and recombination deficient across a majority of the core genome.

Phylogenetic analysis of citrus canker-causing *Xanthomonas*

To uncover the mechanism underlying infection and disease development of citrus canker-causing pathogens, we conducted a comparative genomic analysis of 23 Xac strains as well as two other sequenced citrus canker-causing *Xanthomonas*, XauB and XauC (Moreira *et al.*, 2010). Core orthologous gene clusters were computed for 25 total genomes using the BDBH method, and 2822 gene clusters were identified, accounting for 61.0–74.2% of the total protein coding sequences of each genome. A maximum likelihood phylogenetic tree was reconstructed based on concatenated sequences of these 2822 clusters generated by Gblocks0.91b (Castresana, 2000) using PAUP4b10 (Swofford, 1998) with the best model suggested by ModelTest 3.7 (Posada and Crandall, 1998). The high bootstrap values suggested that the topology structure of this tree was highly confident (Figure 2). The neighbor-joining tree reconstructed using MEGA6 (Tamura *et al.*, 2013) (Supplementary Figure S1) also gave a consistent topology. The clade formed by XauB and XauC was separate from the clade of XacA, XacA* and XacA^w strains, demonstrating the variability of genome content between these two groups. Based on the phylogenetic tree, XacA and variants in Florida were grouped into three separate clades that most likely correspond to three independent introductions that took place in Florida (Gottwald *et al.*,

2002). The XacA^w strains in Florida may have been introduced from India (Schubert *et al.*, 2001). Previous studies demonstrated that XacA^w strains collected from Florida were closely related to some XacA* strains isolated from India (Ngoc *et al.*, 2009). The XacA strains from central Florida and Miami grouped together with the Brazilian strain XacA306 and formed a separate branch from other isolates from Florida and China, suggesting that the central Florida, Miami and Brazilian strains share closer ancestry than other strains. The Manatee (MN) strains of Florida are closely related to strains from China, indicating that they share closer ancestry than other strains.

Positive selection analysis of the core genome of citrus canker-causing *Xanthomonas*

When XauB and XauC were taken into account, the citrus canker-causing population was still effectively clonal (the ρ/θ was 0.0545 ± 0.0006). Furthermore, when tested on gene level, only one gene encoding an aminoacyl-tRNA synthetase, a gene family known to have already undergone complex horizontal gene transfer events in its evolutionary history (O’Donoghue and Luthey-Schulten, 2003), was found to have undergone intragenic recombination ($P < 0.05$) among the 2822 core genes. These results emphasized the rare occurrence of recombination with relative high occurrence of mutation across the core genome of the citrus canker-causing *Xanthomonas*.

Positive selection is an important driving force for pathogens in adaptation process (Lieberman *et al.*, 2011; Trivedi and Wang, 2013). Knowledge of the genes affected by positive selection across citrus canker-causing species can show which genes may be involved in citrus canker development and host specificity. To evaluate the contribution of positive selection to the citrus canker-causing *Xanthomonas* population, FUBAR (Murrell *et al.*, 2013) method

implemented in HYPHY2.2 package (Pond *et al.*, 2005) was employed. This uses a site-specific model of dN/dS. Out of 2822 core genes, 395 genes (14%) were identified as containing codons under positive selection (posterior probability >0.90) (Table 1). These genes were found to be enriched in the 'carbohydrate transport and metabolism' (Fisher's exact test, $P=0.017$; Table 1), further demonstrating the specialization of the pathogens on citrus, and how citrus exerts selection on the pathogens. The 'DNA replication, recombination and repair' (3R) genes were also enriched in the gene set under positive selection (Fisher's exact test, $P=0.047$; Table 1). The 3R genes are crucial for stable maintenance and propagation of DNA for organisms. Interestingly, an important gene involved in DNA repair, *mutY* (XAC2553), the defects of which can cause excess mutations but not inter-genomic recombination (Huang *et al.*, 2006), was found under positive selection. In addition to the two enriched categories, multiple genes known to contribute to virulence were detected. For example, *gumC* and *gumB* were found to be under positive selection. The gum genes, responsible for biosynthesis of xanthan gum, are crucial for *Xac* to form biofilms and survive on the surface of citrus tissues to initiate infection (Facincani *et al.*, 2014). *Xanthomonas* have a diverse complement of T3SEs that contribute to overall pathogenicity and host range variation (Ryan *et al.*, 2011). Among all the effectors identified in the strains analyzed as previously described (Bart *et al.*, 2012) (Supplementary Table S5), 10 common effectors that were shared by all 25 strains were identified in the 2822 core genes. Although none of the 10 effectors was found to have been affected by intragenic recombination as suggested by single breakpoint method result, the maximum likelihood gene trees of all the 10 effectors were found to be incongruent with the maximum likelihood reference tree based on 2822 core genes revealed by Shimodaira-Hasegawa test ($P<0.001$; Table 2) (Shimodaira and Hasegawa, 1999). Four of them, including *xopA* (*hpa1*), *xopV*, *xopX* and *hpaA*, were found to be affected by positive selection. Interestingly, the four effector genes contained more codons under positive selection (*xopA* (2 codons), *xopV* (4), *xopX* (10) and *hpaA* (7)) than average (1.59 codons per gene). T3SEs play important roles in the stepwise arms race that has been taking place between pathogens and their plant hosts during their co-evolutionary history as described by the so-called 'zig-zag' model (Jones and Dangl, 2006), and positive selection of effector genes is an important driving force for host adaptation (Dong *et al.*, 2014). These results suggest that these effectors underwent horizontal gene transfer or intergenic recombination during their evolutionary history, but they had been strongly affected by positive selection during the arms race after their acquirement. In addition, T3SS is the essential machinery responsible for delivering

effectors from the bacterial cytosol directly into the interior of host cells (Ghosh, 2004), and its necessary role in virulence and host adaptation makes the involved genes a hot spot of selection (McCann and Guttman, 2008). This study revealed that six T3SS genes, *hpaB*, *hrpD5*, *hrcQ*, *hpaP*, *hrpB7* and *hpa2*, have been under positive selection, *hrpB7* was found to contain four codons under positive selection and the remaining ones contained one codon affected by positive selection. Positive selection was found for the T3SS and effector genes that are important for virulence and the capacity to overcome the host immune system. They could thus be good candidates for further experimental studies.

Taken together, these results indicate that the evolution of citrus canker-causing *Xanthomonas* is characterized by selection on new mutations that enhance virulence.

Conclusions

We sequenced and conducted genomic comparison and evolutionary analyses of 21 representative *Xac* strains. We found that the recent evolution of *XacA* is characterized by loss of genes and selection on new mutations rather than recombination. Early in its divergence from the narrow host range *Xac* pathotypes, *XacA* likely acquired beneficial genes through recombination, but the *XacA* genome has diversified through clonal expansion as it has colonized citrus worldwide. Mutation occurred much more frequently than recombination for *Xac*, and also for all the citrus canker-causing *Xanthomonas*, suggesting selection, rather than recombination, has been likely important in host range and virulence. Positive selection was observed for 'carbohydrate transport and metabolism' genes, 'DNA replication, recombination and repair' genes and genes involved in T3SS and T3SEs, likely providing a fitness advantage to drive adaptation of these pathogens to their citrus hosts.

Given the closed nature of the pan-genome and the clonal structure of the *Xac* population, the future evolutionary potential of this citrus canker pathogen may be limited. However, rare horizontal transfer events as well as selection on beneficial mutations in the core or accessory genome will likely continue to challenge management of this pathogen.

Conflict of Interest

The authors declare no conflict of interest.

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