


Identification of *Xanthomonas* species associated with bacterial leaf spot of tomato, capsicum and chilli crops in eastern Australia

R. Roach  · R. Mann · C. G. Gambley · R. G. Shivas · B. Rodoni

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Abstract Several species of *Xanthomonas* cause bacterial leaf spot, a disease that affects solanaceous crops worldwide. The diversity of 64 Australian isolates of *Xanthomonas* spp. associated with bacterial leaf spot in tomato, capsicum and chilli crops in eastern Australia was determined using multi-locus sequence analysis of *atpD*, *dnaK*, *efp* and *gyrB* genes, species-specific PCR assays and biochemical analyses. At least five species of *Xanthomonas* associated with bacterial leaf spot were identified in Australian tomato, capsicum and chilli crops and their pathogenicity assessed. Phylogenetic and biochemical analyses identified *X. euvesicatoria*,

X. perforans and *X. vesicatoria* as the most frequently recovered pathogenic species. Non-pathogenic and weakly pathogenic species were also identified. The suitability of the identification methods used and the implications of the detection of these species will be discussed.

Keywords amylolytic · pectolytic · *Solanaceae* · disease management

Introduction

Bacterial leaf spot (BLS) is a disease of solanaceous crops that occurs worldwide, especially in warm and humid climates (Jones et al. 2014). Several species of *Xanthomonas* are reported to cause BLS of tomato (*Solanum lycopersicum*), capsicum and chilli (*Capsicum annuum*) (Potnis et al. 2015). The symptoms of BLS are small, brown, angular, water-soaked lesions on leaves, stems and fruit, and result in defoliation and direct fruit damage. Severe infection may result in extensive damage to crops with significant yield losses (Pemezný et al. 2003). Species reported to cause BLS all produce similar symptoms on their hosts, making precise diagnosis difficult from visual symptoms alone. The impact of pathological convergence and importance of phylogenetic testing in the case of BLS are further highlighted by Hajri et al. (2009). The causal bacteria are spread by wind and water, and may survive in crop residues, weeds and volunteer plants (Jones et al. 1986). A link between field disease and seed contamination has

GenBank accession numbers: KY658725–KY658788, KY658789–KY658852, KY658853–KY658916, KY658917–KY658975, KY364014–KY364018.

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R. Roach · R. G. Shivas
Department of Agriculture and Fisheries, Ecosciences Precinct,
Brisbane, QLD, Australia

R. Roach (✉) · R. Mann · B. Rodoni
Department of Environment and Primary Industries, La Trobe
University, Melbourne, VIC, Australia
e-mail: roach.r@students.latrobe.edu.au

C. G. Gambley
Department of Agriculture and Fisheries, Applethorpe Research
Facility, Stanthorpe, QLD, Australia

been apparent for almost a century (Higgins 1922; Shekhawat and Chakravarti 1979). Control of BLS is achieved primarily through the use of resistant lines, antibiotic application and copper sprays. Reliance on a limited range of chemicals has seen copper and antibiotic resistance develop rapidly in *Xanthomonas* populations (Griffin et al. 2017; Martin et al. 2004; Minsavage et al. 1990).

Four species of *Xanthomonas* reported to cause BLS are *X. euvesicatoria* (Jones et al. 2006), *X. gardneri* (ex Susic 1957; Jones et al. 2006), *X. perforans* (Jones et al. 2006), and *X. vesicatoria* (ex Doidge 1920; Vauterin et al. 1995). These species were once classified in the *X. campestris* pv. *vesicatoria* species complex (Young et al. 1978) and the most recent Australian report of BLS refers to *X. campestris* pv. *vesicatoria* (Martin et al. 2004). A recent study by Constantin et al. (2016) proposed merging *X. perforans* and *X. euvesicatoria* as they and others have noted synonymy in these species (Potnis et al. 2015). The identification and detection of these species is typically achieved through the use of various molecular techniques (eg. PCR, MLSA, NGS) as the taxonomy has moved beyond the point where traditional biochemical techniques effectively distinguish these groups. *Xanthomonas* spp. that cause BLS in tomato and capsicum crops are reported from several countries (EPPO 2013; Timilsina et al. 2015). Strains of *Xanthomonas arboricola* (Vauterin et al. 1995) have been shown to cause BLS on capsicum in Korea and has also been associated with disease outbreaks on tomato in Tanzania (Mbega et al. 2012; Myung et al. 2010). A *Xanthomonas* sp. reported on tomato in Tanzania was genetically distinct from the four commonly reported species based on a phylogeny of *fyuA* sequence (Mbega et al. 2012). *Xanthomonas campestris* pv. *raphani* has also been noted to cause BLS on tomato (Punina et al. 2009). Non-pathogenic species of *Xanthomonas* have also been recorded, potentially complicating the detection of pathogens (Vauterin et al. 1996). The distribution and prevalence of BLS-causing *Xanthomonas* species in Australia is relatively unknown (EPPO 2013).

BLS was first reported in Australia in 1944 (Anonymous 1944) and continues to impact crop production. The 2014/15 production value of Australian capsicum and chilli industries were valued at \$144.7 M and \$9.6 M respectively. The production

value of Australian tomato was \$548 M, with the majority of all three crops grown in Queensland (HIA 2016). Though outbreaks of BLS occur in most commercial growing regions in Australia, little has been done to investigate the genetic diversity and distribution of the causal *Xanthomonas* species. The development of effective management strategies, primarily the selection of resistant plant material, relies upon the accurate identification of pathogens and an understanding of their diversity and pathogenicity. This study describes the identification and diversity of pathogenic and non-pathogenic *Xanthomonas* spp. associated with BLS in Australia based on biochemical and molecular analysis.

Materials and methods

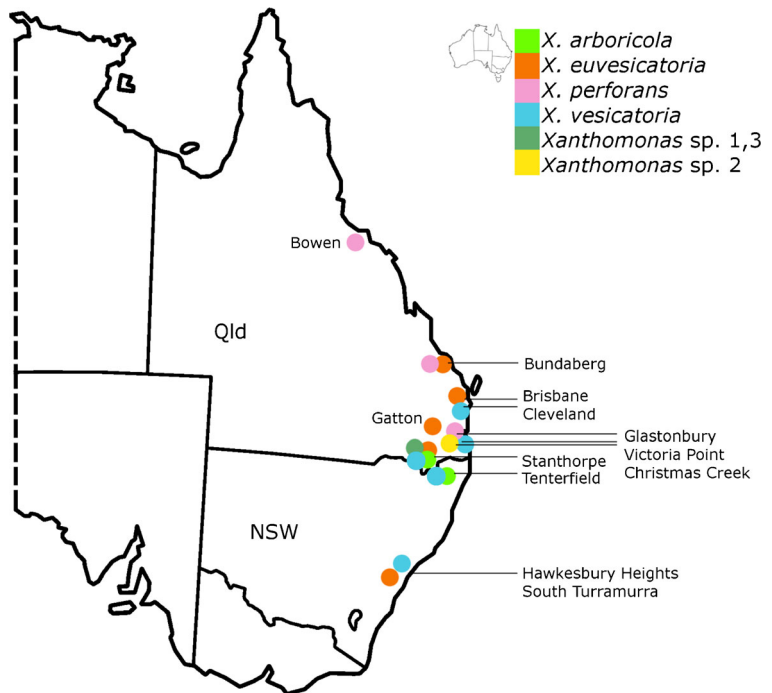
Specimen collection

Capsicum, chilli and tomato crops were surveyed for BLS symptoms during the 2015 growing season in Queensland (Qld) and New South Wales (NSW). Symptomatic tissue was collected from sites around eastern Australia (Fig. 1). Lesions were examined for bacterial streaming and initial isolations were made on Nutrient Agar (NA, Astral Scientific Pty Ltd) (Schaad 2001). *Xanthomonas*-like bacteria were single colony purified and stored at -80°C in 2 ml Microbank™ vials (Pro-Lab Diagnostics Inc.) and deposited in the culture collection of the Queensland Plant Pathology Herbarium (BRIP). Type strains for *X. euvesicatoria* (NCPPB 2968), *X. gardneri* (NCPPB 881), *X. perforans* (NCPPB 4321) and *X. vesicatoria* (NCPPB 422) were imported under permit from the National Collection of Plant Pathogenic Bacteria (NCPPB). A further seven unidentified isolates of *Xanthomonas* from solanaceous hosts were sourced from the BRIP culture collection. All of the bacterial isolates examined in this study are shown in Table 1, with additional reference sequence from GenBank shown in Table 3 (supplementary data).

Biochemical analysis and pathogenicity

Biochemical tests were conducted on 48 h-old cultures taken from storage and grown on NA at room temperature. Oxidase, catalase and potassium hydroxide (KOH) tests and media preparation were done as described by Schaad (2001). Cultures were streaked on to

Fig. 1 *Xanthomonas* species distribution across the surveyed areas of eastern Australia. Colours represent species determined by phylogenetic clades generated using multi-locus sequence analysis (Fig. 3)



starch, pectin and yeast extract-dextrose-calcium carbonate (YDC) media, and the morphology recorded after 48–72 h (Schaad 2001). Biochemical data was displayed as a cluster analysis (Fig. 2) by assigning each result a binary value (1 = positive, 0 = negative). The cluster and dendrogram were created in R (R core team 2016).

To confirm pathogenicity on the host of isolation, bacteria were grown on nutrient agar overnight (Schaad 2001) at room temperature and then suspended in sterile distilled water. The concentration of the inoculum solution was measured using a BioDrop DUO (BioDrop) spectrophotometer and adjusted to $OD_{600} = 0.2$ (1×10^8 cfu/ml) with sterile distilled water. Leaf panels of *Capsicum annuum* var. Jupiter or *Solanum lycopersicum* var. Grosse Lisse at the 2–4 true leaf stage were infiltrated with 300 μ l of inoculum using a 1 ml syringe. Sterile distilled water was used as a control. Water-soaked lesions were deemed a susceptible reaction and tan papery lesions a hypersensitive response. Isolates representing each phylogenetic clade (62409–62412, 62414–62418, 62432, 39016, 38864, 62555, 63565, 62428) were then prepared as above and spray inoculated to confirm symptomatic pathogenicity. Isolates from non-crop hosts BRIP 39016 and 38864

were inoculated on tomato by infiltration and spray as described above.

Gene amplification and sequencing

DNA template was prepared by suspending a single pure colony in 100 μ l milliQ water, incubated at 95 °C for 7 min and then used for species-specific and MLSA associated PCRs. PCR protocols developed to differentiate *X. euvesicatoria*, *X. gardneri*, *X. perforans* and *X. vesicatoria* were used for all isolates including type strains which were used as positive controls (Koenraadt et al. 2009). PCRs were performed as described in Koenraadt et al. (2009) using 2.5 U of MangoTaq™ (Bioline) and 1 μ l of template. Primers and their annealing temperatures are listed in Table 2.

MLSA-PCR primers targeting the chaperone protein *dnaK* (*dnaK*), elongation factor P (*efp*), ATP synthase subunit beta (*atpD*) and DNA gyrase subunit B (*gyrB*) genes (Table 2) were used as described (Boudon et al. 2005; Ah-You et al. 2009; Ngoc et al. 2010; Hamza et al. 2012). Annealing temperatures were optimised for the MLSA primers targeting all four genes. PCR products were purified and sequenced by Macrogen Inc. (South Korea) in both directions using an Applied Biosystems 3730xl DNA Analyser.

Table 1 Collection data, GenBank accession numbers and species-specific PCR results for isolates collected and type isolates used in this study

Organism ^c	Accession no.	Location	Host	Collection year	Species specific PCR result ^b
<i>X. arboricola</i>	BRIP 62410	Stanthorpe	Tomato	2015	0
<i>X. arboricola</i>	BRIP 62412	Stanthorpe	Tomato	2015	0
<i>X. arboricola</i>	BRIP 62414	Stanthorpe	Tomato	2015	0
<i>X. arboricola</i>	BRIP 62416	Stanthorpe	Tomato	2015	0
<i>X. arboricola</i>	BRIP 62432	Tenterfield	Tomato	2015	0
<i>X. euvesicatoria</i>	NCPBP 2968 ^T	USA	Chilli (<i>Capsicum frutescens</i>)	1977	Xe
<i>X. euvesicatoria</i>	BRIP 38855	Bundaberg	Capsicum	1981	Xe
<i>X. euvesicatoria</i>	BRIP 63464	Bundaberg	Capsicum	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62439	Bundaberg	Chilli ^a	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62440	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62441	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62442	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62443	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62555	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62656	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62757	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62858	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62959	Bundaberg	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 38997	Bundaberg	Chilli	1986	Xe
<i>X. euvesicatoria</i>	BRIP 62454	Bundaberg	Chilli (<i>C. chinense</i>)	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62390	Gatton	Capsicum	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62391	Gatton	Capsicum	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62392	Gatton	Capsicum	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62393	Gatton	Capsicum	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62394	Gatton	Capsicum	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62395	Gatton	Capsicum	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62396	Gatton	Chilli	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62399	Gatton	Chilli	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62400	Gatton	Chilli	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62401	Gatton	Chilli	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62402	Gatton	Chilli	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62403	Gatton	Chilli	2014	Xe
<i>X. euvesicatoria</i>	BRIP 62434	Gatton	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62435	Gatton	Chilli	2015	Xe
<i>X. euvesicatoria</i>	BRIP 39000	Glastonbury	Chilli	1986	Xe
<i>X. euvesicatoria</i>	BRIP 62438	Hawkesbury Heights	Capsicum	2015	Xe
<i>X. euvesicatoria</i>	BRIP 62425	Stanthorpe	Capsicum	2015	Xe
<i>X. gardneri</i>	NCPBP 881 ^T	Yugoslavia	Tomato	1961	Xg
<i>X. perforans</i>	NCPBP 4321 ^T	USA	Tomato	2004	Xp
<i>X. perforans</i>	BRIP 62383	Bowen	Tomato	2012	0
<i>X. perforans</i>	BRIP 62384	Bowen	Tomato	2012	0
<i>X. perforans</i>	BRIP 62385	Bowen	Tomato	2012	0
<i>X. perforans</i>	BRIP 62387	Bowen	Tomato	2012	0

Table 1 (continued)

Organism ^c	Accession no.	Location	Host	Collection year	Species specific PCR result ^b
<i>X. perforans</i>	BRIP 62398	Bowen	Tomato	2014	0
<i>X. perforans</i>	BRIP 62404	Bowen	Tomato	2014	0
<i>X. perforans</i>	BRIP 62397	Brisbane	Tomato	2014	0
<i>X. perforans</i>	BRIP 62386	Bundaberg	Tomato	2012	0
<i>X. perforans</i>	BRIP 62405	Bundaberg	Tomato	2014	0
<i>X. perforans</i>	BRIP 63262	Bundaberg	Tomato	2015	0
<i>X. perforans</i>	BRIP 63565	Bundaberg	Tomato	2015	0
<i>X. perforans</i>	BRIP 63666	Bundaberg	Tomato	2015	0
<i>X. perforans</i>	BRIP 62389	South Turrumurra	Tomato	2013	0
<i>X. vesicatoria</i>	NCPPB 422 ^T	NZ	Tomato	1957	Xv
<i>X. vesicatoria</i>	BRIP 38864	Bowen	<i>S. peruvianum</i>	1982	Xv
<i>X. vesicatoria</i>	BRIP 39109	Cleveland	Tomato	1976	Xv
<i>X. vesicatoria</i>	BRIP 62388	South Turrumurra	Tomato	2012	Xv
<i>X. vesicatoria</i>	BRIP 62423	Stanthorpe	Tomato	2015	Xv
<i>X. vesicatoria</i>	BRIP 62429	Tenterfield	Tomato	2015	Xv
<i>X. vesicatoria</i>	BRIP 62430	Tenterfield	Tomato	2015	Xv
<i>X. vesicatoria</i>	BRIP 62428	Tenterfield	Tomato	2015	Xv
<i>X. vesicatoria</i>	BRIP 38861	Victoria Point	Tomato	1981	Xv
<i>X. vesicatoria</i>	BRIP 62413	Stanthorpe	Tomato	2015	0
<i>Xanthomonas</i> sp.	BRIP 39016	Christmas Creek	<i>N. physalodes</i>	1973	0
<i>Xanthomonas</i> sp.	BRIP 62409	Stanthorpe	Tomato	2015	0
<i>Xanthomonas</i> sp.	BRIP 62411	Stanthorpe	Tomato	2015	0
<i>Xanthomonas</i> sp.	BRIP 62415	Stanthorpe	Tomato	2015	0
<i>Xanthomonas</i> sp.	BRIP 62417	Stanthorpe	Tomato	2015	0
<i>Xanthomonas</i> sp.	BRIP 62418	Stanthorpe	Tomato	2015	0

^T indicates type strains from the National Collection of Plant Pathogenic Bacteria (NCPPB), Fera, UK

^a *C. annuum* variety unless otherwise stated

^b positive reaction in the species-specific PCR for *X. euvesicatoria* (Xe), *X. gardneri* (Xg), *X. perforans* (Xp) or *X. vesicatoria* (Xv), negative in all four PCRs indicated by 0

^c species designation based on MLSA (Fig. 3)

Sequence analysis

Geneious (v. 9.1.2) was used to process and analyse sequence data (Kearse et al. 2012). Sequence reads were manually trimmed by quality scores and compared to the non-redundant database in GenBank using the BLASTn algorithm (Madden 2003). Sequences generated in this study were deposited in GenBank (Table 4 supplementary data). Trimmed sequences were aligned with reference sequences of *Xanthomonas* spp. *atpD*, *dnaK*, *efp*, and *gyrB* genes (supplementary Table 3) available on GenBank (Benson et al. 2005) using clustalW (Larkin et al. 2007). Sequences from Hamza et al. (2010, 2012) and the top BLASTn hit for each gene of isolates

in unresolved clades were included in the MLSA analysis. The outgroup taxa were *Stenotrophomonas maltophilia*, *X. albilineans* and *X. campestris* pv. *raphani*. Other sequences of *X. arboricola* from capsicum and tomato were not included in this analysis due to lack of available sequence data. Bayesian inference (BI) was used to construct a phylogeny of concatenated gene sequences of *atpD*, *dnaK*, *efp*, and *gyrB*. Phylogenetic trees were produced for individual genes and concatenated gene sequences using MrBayes ver. 3.2.6 (Huelsenbeck and Ronquist 2001) for Bayesian estimation with a GTR gamma model chosen with jmodeltest2 (Darriba et al. 2012; Guindon and Gascuel 2003) and 1,100,000 chain with a burn-in of 100,000 sampled

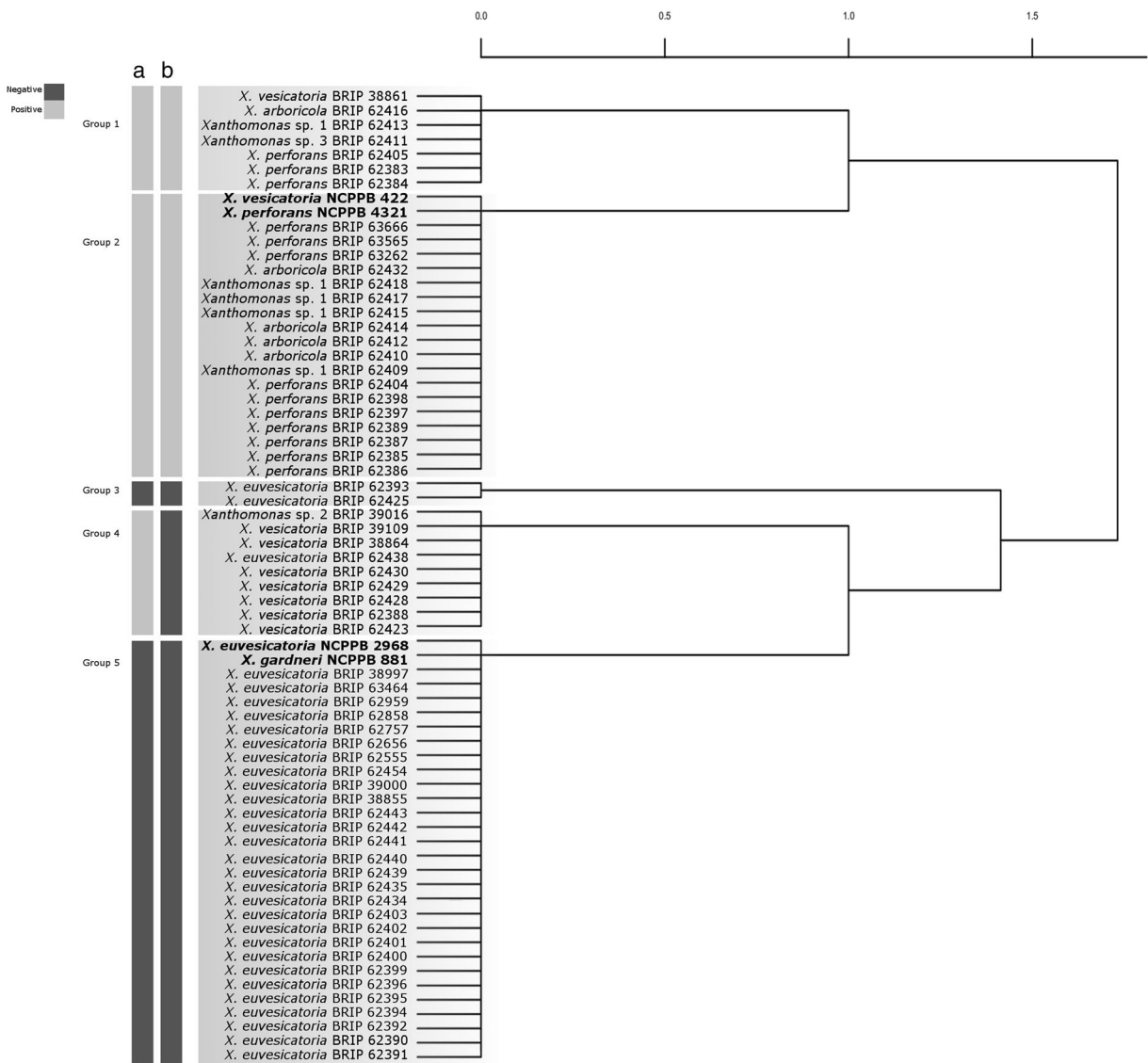


Fig. 2 Hierarchical cluster analysis of biochemical traits from the collected and type isolates. Names are designated as per the clades in Fig. 3. **a**: starch reaction **b**: pectin reaction

every 200 trees. Branch support is reported as posterior probability (PP). A maximum likelihood RaxML (version 7.2.8) tree was also generated for the concatenated gene sequences using a GTR gamma model with 1000 bootstrap replicates (Stamatakis 2014). Images were annotated using FigTree ver. 1.4.2 (Rambaut 2016) and GIMP ver. 2.8.14 (The GIMP team 2014).

The Species Delimitation plugin for Geneious was used to measure relatedness in the generated phylogenies (Masters et al. 2011). Clades and individual sequences were selected to determine which clades were most closely related and supplement the bootstrap values generated in the phylogeny. Single gene and

concatenated data sets were checked for recombination in *atpD* with RDP (Martin et al. 2015). *AtpD* sequence was included in this MLSA to better distinguish the *X. euvesicatoria* and *X. perforans* groups.

Results

Specimen collection

Fifty-seven *Xanthomonas*-like bacteria isolated from symptomatic tomato, capsicum and chilli crops plants from Qld and NSW from 2012 to 2015 and an additional

Table 2 PCR primer details for *Xanthomonas* species-specific and MLSA assays

Target	Name	Sequence	Ta	Product size
<i>X. vesicatoria</i>	Bs-XvF ^a	CCA TGT GCC GTT GAA ATA CTT G	64 °C	138 bp
	Bs-XvR ^a	ACA AGA GAT GTT GCT ATG ATT TGC		
<i>X. euvesicatoria</i>	Bs-XeF ^a	CAT GAA GAA CTC GGC GTA TCG	64 °C	173 bp
	Bs-XeR ^a	GTC GGA CAT AGT GGA ATA CTT G		
<i>X. perforans</i>	Bs-XpF ^a	GTC GTG TTG ATG GAG CGT TC	64 °C	197 bp
	Bs-XpR ^a	GTG CGA GTC AAT TAT CAG AAT GTG G		
<i>X. gardneri</i>	Bs-XgF ^a	TCA GTG CTT AGT TCC TCA TTG TC	64 °C	154 bp
	Bs-XgR ^a	TGA CCG ATA AAG ACT GCG AAA G		
<i>Xanthomonas dnaK</i> gene	dnaK-F ^c	TGG GCA AGA TCA TTG GTA TT	67 °C	765 bp
	dnaK-R ^c	ACC TTC GGC ATA CGG GTC TG		
<i>Xanthomonas atpD</i> gene	atpD-F ^c	GGG CAA GAT CGT TCA GAT	66 °C	768 bp
	atpD-R ^c	GCT CTT GGT CGA GGT GAT		
<i>Xanthomonas gyrB</i> gene	gyrB-F ^b	GCC GAG GTG ATC CTC ACC GT	69 °C	774 bp
	gyrB-R ^b	GGC CGA GCC ACC TGC CGA GT		
<i>Xanthomonas efp</i> gene	efp-F ^c	GTG AAG AAC GGC ATG AAG A	65 °C	387 bp
	efp-R ^c	TCG TCC TGG TTG ACG AAC		

^aKoenraadt et al. 2009^bAh-You et al. 2009^cBoudon et al. 2005

seven isolates from the BRIP collection dating back to 1973, were included in this study (Table 1). Of these 64 isolates, 10 isolates were from capsicum, 21 from chilli, 31 from tomato, and one from each of *Nicandra physalodes* (apple-of-Peru) and *Solanum peruvianum* (wild tomato). Additional data from GenBank was included in the MLSA as reference sequence (Table 3 supplementary data).

Biochemical results and pathogenicity

All isolates gave susceptible reactions by infiltration on their host of isolation apart from BRIP 62409, 62411, 62415, 62417 and 62418. These isolates are recorded as non-pathogenic on tomato. Isolates BRIP 62410, 62412, 62414, 62416 and 62432 displayed pathogenic reactions when infiltrated but did not display typical lesions when spray inoculated. These isolates are described as weakly pathogenic. All other spray inoculated isolates produced typical BLS symptoms, including isolates from non-crop hosts, BRIP 39016 and 38864.

The biochemical description for members of the *Xanthomonas* genus is gram-negative, oxidase-negative, catalase positive rods producing yellow colonies on YDC (Schaad 2001). All 68 bacteria (64 Australian and four type isolates) tested KOH positive (therefore gram-negative), produced yellow colonies on YDC and were negative for the presence of an oxidase enzyme. Sixty-one isolates (including the four type isolates) were strongly or weakly positive for catalase while nine isolates were negative. Variable biochemical test results (starch, pectin and catalase) are described below (Fig. 2).

The biochemical profile of *X. euvesicatoria* (NCPBP 2968) and *X. gardneri* (NCPBP 881), and 28 Australian isolates from capsicum and chilli were catalase-positive, starch-negative and pectin-negative. The type strains of *X. perforans* (NCPBP 4321) and *X. vesicatoria* (NCPBP 422), together with 18 Australian isolates from tomato were catalase-positive, starch-positive and pectin-positive. The remaining Australian isolates had different biochemical profiles than those of the four type strains. Seven of these isolates from tomato were starch-positive, pectin-positive and catalase-negative. One isolate from *S. peruvianum*, one from *N. physalodes*, one from capsicum and six isolates from tomato (nine total) were catalase-positive, starch-positive and pectin-negative. The remaining two isolates from capsicum were

catalase-negative, starch-negative and pectin-negative (Fig. 2).

The cluster analysis of the above biochemical profiles (Fig. 2) identified five groups among the 68 isolates based largely on the variable traits of starch and pectin degradation (Fig. 2). Group 1 of the dendrogram contained 7 isolates, all of which were starch-positive and pectin-positive. Group 2 clustered most closely to group 1 and contained 20 isolates, including the type strains of *X. vesicatoria* and *X. perforans*, which were also starch-positive and pectin-positive. Group 3 contained 2 starch and pectin-negative isolates that were also catalase-negative. Group 4 contained nine starch-positive and pectin-negative isolates. Group 5 contained 30 starch-negative and pectin-negative isolates, including the type strains of *X. euvesicatoria* and *X. gardneri*. Isolates in groups 1, 2 and 4 were from tomato, with the exception that group 4 also contained isolates from *N. physalodes* and *S. peruvianum*. Groups 2 and 5 contained isolates from capsicum and chilli.

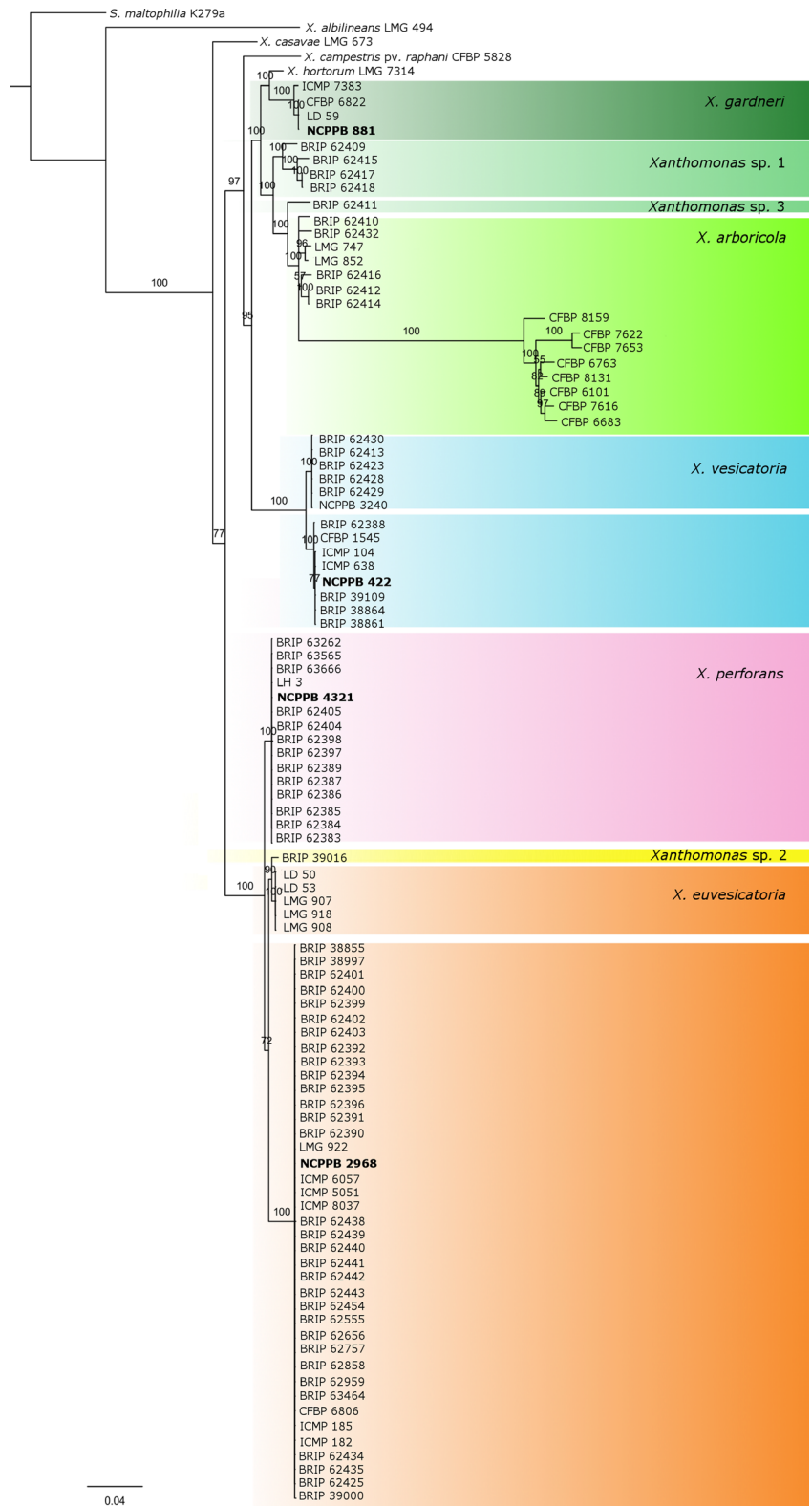
Species specific PCR

Type strains of *X. euvesicatoria*, *X. gardneri*, *X. perforans* and *X. vesicatoria* were positive for their respective specific PCR tests and negative for the remaining three PCRs (Table 1). Thirty-one isolates, all derived from capsicum and chilli, tested positive for *X. euvesicatoria*. Eight isolates tested positive using the *X. vesicatoria*-specific PCR, seven of which were isolated from tomato and one from *S. peruvianum*. The remaining 25 pathogenic and non-pathogenic isolates (24 from tomato and one from *N. physalodes*) were negative for all 4 species-specific PCR assays. Assays for *X. euvesicatoria* and *X. vesicatoria* generated amplicons, while assays for *X. perforans* and *X. gardneri* did not react with any Australian isolates.

MLSA

Bayesian inference analysis resolved all 64 Australian isolates collected in this study (Table 1) and additional Genbank sequence (Table 3 supplementary data) in eight strongly supported clades (Fig. 3). Phylogenetic analysis of the *dnaK*, *efp* and *gyrB* genes (Fig. 4 supplementary data) gave the same general topology as the tree including *atpD* and grouped isolates in the same clades, as did the RaxML phylogeny (Fig. 5 supplementary data). Each individual gene phylogeny

Fig. 3 Phylogenetic Bayesian inference tree of concatenated gene sequences (in the order of *gyrB*, *efp*, *dnaK*, *atpD*) with Genbank reference isolates and posterior probability (percent) branch support values. Type strains are indicated in bold



failed to distinguish at least one clade from the other clades. *atpD* sequence was included in this MLSA as its absence did not significantly alter the tree topology and its inclusion better differentiated *X. euvesicatoria* and *X. perforans* clades. The recombination events detected in *atpD* by Hamza et al. (2012) were not detected by RDP in the single gene or concatenated data sets of these isolates. The species delimitation results supported the relatedness seen in the above phylogenies (supplementary Table 5).

The largest clade (Fig. 3) contained 31 Australian isolates as well as the type strain of *X. euvesicatoria* (NCPBP 2968) from chilli in the USA. This clade also contained seven GenBank reference sequences of *X. euvesicatoria* (supplementary Table 3) from capsicum (Brazil, Cook Islands, New Zealand and USA) and chilli (USA). All Australian isolates in this clade were sourced from capsicum and chilli at Bundaberg and Gatton, Qld and Glastonbury and Hawkesbury Heights, NSW. Five *X. euvesicatoria* reference isolates from Hamza et al. (2010) grouped outside the largest *X. euvesicatoria* clade, closest to but separate from the *N. physalodes* (BRIP 39016) isolate. The clade containing the type strain of *X. perforans* (NCPBP 4321) from tomato (USA) also included 13 Australian isolates and a GenBank reference of *X. perforans* (LH 3). These Australian isolates were recovered from tomato at Bowen, Bundaberg and Brisbane (QLD) and South Turrumurra (NSW). Two closely related but distinct clades (100% PP) contained GenBank reference sequence of *X. vesicatoria* and Australian isolates. The first of these clades contained GenBank references of *X. vesicatoria* (NCPBP 3240) from tomato in Spain and five Australian isolates from tomato at Stanthorpe (QLD) and Tenterfield (NSW). The second clade contained the type strain of *X. vesicatoria* (NCPBP 422) from tomato (New Zealand), three GenBank references of *X. vesicatoria* from tomato (France, New Zealand and the USA), three Australian isolates from tomato (Cleveland, Bowen and Victoria Point, QLD) and one from *S. peruvianum* (Bowen). The clade containing the type strain of *X. gardneri* (NCPBP 881) from tomato (Yugoslavia) also included three GenBank references of *X. gardneri* from Canada, Argentina and New Zealand (supplementary Table 3). None of the Australian isolates were in this clade. A currently uncharacterised clade (*Xanthomonas* sp.) sister to the *X. gardneri* clade (100% PP) contained four Australian isolates from tomato in Stanthorpe, QLD.

The type strain of *X. arboricola* (LMG 747) from walnut (New Zealand) was resolved in a clade that contained a Genbank accession of *X. arboricola* pv. *pruni* from plum (New Zealand), five Australian isolates from tomato (Stanthorpe and Tenterfield, QLD) and a sub-clade (100% PP) of eight GenBank accessions of *X. arboricola*. One isolate (BRIP 62411, designated *Xanthomonas* sp. 3) from tomato at Stanthorpe (QLD) was resolved outside the *X. arboricola* clade and the *Xanthomonas* sp. clade (Fig 3).

Discussion

Xanthomonas arboricola, *X. euvesicatoria*, *X. perforans*, *X. vesicatoria* and at least one undescribed *Xanthomonas* spp. were identified amongst isolates associated with BLS on capsicum, chilli and tomato crops in eastern Australia on the basis of MLSA, species-specific PCR, pathogenicity and biochemical analysis. Of these, only *X. vesicatoria* and *X. euvesicatoria* were previously reported to cause BLS in Australia (EPPO 2013). This study represents the first report of *X. perforans* causing bacterial leaf spot on tomato in Australia. *Xanthomonas gardneri* was not detected amongst the Australian isolates, and thus may represent a biosecurity threat for Australia. The rapid global spread of *X. gardneri* is highlighted previously (Timilsina et al. 2015), and its absence here further emphasises the importance of seed testing and updated diagnostic protocols as highlighted in other studies (Potnis et al. 2015). The suitability of standard identification techniques and the relationships of Australian *Xanthomonas* isolates is discussed below.

The methods used to identify the *Xanthomonas* species associated with BLS in this study highlight a need for updated diagnostic protocols for this disease. The cluster analysis displayed biochemical traits as an unreliable indicator of species and pathogenicity (Fig. 2). The limited reliability of biochemical tests has been observed in other diagnostic and diversity studies, and our results confirm that biochemical data is not sufficient for distinguishing Australian BLS-associated isolates (Berge et al. 2014). Considering further molecular characterisation was necessary, species-specific PCR and MLSA were evaluated. The species-specific PCR assays developed to detect four common *Xanthomonas* species causing BLS (Jones et al. 2004; Koenraad et al. 2009) are endorsed as a diagnostic tool by EPPO

(EPPO 2013). However, these assays provided only partial detection of the Australian BLS-causing species, failing to detect *X. perforans* isolates and one *X. vesicatoria* isolate. The assays were developed with isolates from various geographic regions and subsequently validated with Brazilian isolates, though did not include any Australian isolates (Araujo et al. 2012). MLSA of four genes resolved Australian BLS-causing isolates in species clades consistent with the literature (Hamza et al. 2010) and distinguished them from non-pathogenic strains. The presence of *Xanthomonas* spp. that were not pathogenic on their host of isolation (*Xanthomonas* spp., Fig. 3) suggests potential for confounding diagnostics. Based on the findings of this study we suggest that updated diagnostic assays for Australian BLS isolates is needed.

The identification of *X. perforans* from tomato in QLD and NSW represents a new record for Australia. *Xanthomonas perforans* showed little genetic diversity amongst isolates from Australia and GenBank accessions of *X. perforans*, including the type strain, which is consistent with overseas observations of this species (Timilsina et al. 2015). The biochemical profile for starch and pectin utilisation of the Australian *X. perforans* isolate matched that of the *X. perforans* type strain (NCPFB 4321). Despite this apparent genetic and phenotypic similarity, these isolates did not react with the species-specific PCR that targets *X. perforans* (Koenraad et al. 2009), possibly due to variation in the primer binding region (eg. indels, mismatch). Additional diagnostic assays will need to be validated for use with Australian BLS-causing *X. perforans* and other *Xanthomonas* spp. The similarity of *X. perforans* and *X. euvesicatoria* has been noted previously (Timilsina et al. 2015). Merging these species has been proposed, however we refer to *X. perforans* in this study to distinguish this clade from our *X. euvesicatoria* clade. The polyphyletic results of other studies and the highly related sequences of this study indicate whole genome phylogenies with diverse sample groups would further resolve the taxonomy.

Isolates of *X. vesicatoria* from tomato identified in this study separated into two strongly supported subclades that each contained at least one isolate identified as *X. vesicatoria* by Hamza et al. (2012). The *X. vesicatoria*-specific PCR detected the Australian isolates from both subclades. These distinct subclades of *X. vesicatoria* may represent different introductions of

the same species from different origins or possibly the existence of two distinct species in Australia. Geographical clade separation of isolates of *X. vesicatoria* from New Zealand and South America has been observed (Timilsina et al. 2015). The biochemical profiles of Australian *X. vesicatoria* isolates differed from the type strain in pectin utilisation, although all were starch-positive. This gives further support to the observation that biochemical traits are not a reliable discriminating test for these species.

Isolates of *X. euvesicatoria* from Australian capsicum and chilli crops were genetically similar to isolates from overseas based on phylogenetic analysis of the *atpD*, *efp*, *dnaK*, and *gyrB* genes (Ah-You et al. 2009; Hamza et al. 2010, 2012). Australian isolates of *X. euvesicatoria* were positive in the *X. euvesicatoria* species-specific PCR, supporting the grouping of these isolates in the MLSA phylogeny. The *X. euvesicatoria* isolates were generally starch and pectin negative, apart from one isolate (BRIP 62438) that was starch positive. Starch utilisation has historically provided a point of variation for the differentiation of populations of BLS since the early description of *X. campestris* pv. *vesicatoria* (Doidge 1921; Gardner and Kendrick 1921). Variable starch utilisation has been noted amongst isolates of *X. euvesicatoria* (Bouzar et al. 1996), suggesting some variation in this trait is not unusual and supporting the findings above of unreliable biochemical tests. An isolate from *N. physalodes* (BRIP 39016; *Xanthomonas* sp. 2) collected at Christmas Creek (QLD) in 1973 was negative in all of the species-specific PCRs and grouped most closely to the *X. euvesicatoria* clades (Fig. 3) and particularly with reference isolates from Hamza (2010) that formed a small *X. euvesicatoria* clade. Based on these results, it is likely that BRIP 39016 belongs to *X. euvesicatoria*, though genomic data is need to clarify this.

The inclusion of multiple *X. arboricola* pathovars in the MLSA revealed a highly diverse clade. Five isolates of *X. arboricola* were most closely related to *X. arboricola* pv. *pruni* and the type strain of *X. arboricola* pv. *juglandis* by phylogenetic analysis of the *atpD*, *efp*, *dnaK*, and *gyrB* genes. One isolate, BRIP 62411, resolved outside of the *X. arboricola* clade in the MLSA analysis and currently remains unclassified (Fig. 3). *Xanthomonas arboricola* pv. *juglandis* is starch-positive (Scortichini et al. 2001), as were the five Australian isolates, however *Xanthomonas arboricola* pv. *pruni* is recorded as being starch-negative,

indicating this trait is not uniform among *X. arboricola* (EPPO 2006). The five Australian isolates were also negative in all species-specific PCRs. The MLSA in this study indicates that additional genetic data may provide more accurate resolution of this species. High genetic diversity has been observed in *X. arboricola* previously (Fischer-Le Saux et al. 2015), supporting the levels of diversity reported in this study and indicating further review of the *X. arboricola* species may be necessary. The weakly pathogenic nature of these isolates suggests they may be opportunistic, and *X. arboricola* isolates of uncertain pathogenicity have been previously observed (Fischer-Le Saux et al. 2015). BLS-causing isolates of *X. arboricola* have also been reported in Korea and Tanzania (Myung et al. 2010; Mbega et al. 2012). These were not included in this MLSA due to a lack of sequence data.

Three species are responsible for BLS outbreaks in Australia. Further investigation into the genetics behind the pathogenicity of these species will provide more detailed knowledge of pathogen population dynamics and spread. Instances of race shifts and mutations in populations of *Xanthomonas* associated with BLS have been recorded overseas (Dahlbeck and Stall 1979; Kousik and Ritchie 1996; Ma et al. 2011), demonstrating that knowledge of population distribution is critical to crop protection. An understanding of the genetic diversity of BLS associated *Xanthomonas* species in Australia will aid in the development of resistant host lines and diagnostic assays, while also furthering the global understanding of bacterial disease detection and management. Australia's capability for detecting exotic or newly evolved strains will also be improved, especially given the absence of *X. gardneri*, as the observations made in this and subsequent studies will be applicable to exotic *Xanthomonas* species.

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Compliance with ethical standards

Conflict of interest The authors state they have no conflict of interest

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