

# Comparative genomics of Pseudomonas syringae reveals convergent gene gain and loss associated with specialisation onto cherry (Prunus avium)

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Hulin, M. T., Armitage, A. D., Vicente, J. G., Holub, E. B., Baxter, L., Bates, H. J., Mansfield, J. W., Jackson, R. and Harrison, R. J. (2018) Comparative genomics of Pseudomonas syringae reveals convergent gene gain and loss associated with specialisation onto cherry (Prunus avium). New Phytologist, 219 (2). pp. 672-696. ISSN 1469-8137 doi: 10.1111/nph.15182 Available at https://centaur.reading.ac.uk/76290/

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To link to this article DOI: http://dx.doi.org/10.1111/nph.15182

Publisher: Wiley

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## Comparative genomics of *Pseudomonas syringae* reveals convergent gene gain and loss associated with specialization onto cherry (Prunus avium)

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Received: 9 January 2018 Accepted: 22 March 2018

New Phytologist (2018) 219: 672-696 doi: 10.1111/nph.15182

Key words: avirulence, bacterial canker, comparative genomics, host specialization, prediction, Pseudomonas, toxins, type III effectors.

#### **Summary**

- Genome-wide analyses of the effector- and toxin-encoding genes were used to examine the phylogenetics and evolution of pathogenicity amongst diverse strains of Pseudomonas syringae causing bacterial canker of cherry (Prunus avium), including pathovars P. syringae pv morsprunorum (Psm) races 1 and 2, P. syringae pv syringae (Pss) and P. syringae pv avii.
- Phylogenetic analyses revealed Psm races and P. syringae pv avii clades were distinct and were each monophyletic, whereas cherry-pathogenic strains of Pss were interspersed amongst strains from other host species.
- A maximum likelihood approach was used to predict effectors associated with pathogenicity on cherry. Pss possesses a smaller repertoire of type III effectors but has more toxin biosynthesis clusters than Psm and P. syringae pv avii. Evolution of cherry pathogenicity was correlated with gain of genes such as hopAR1 and hopBB1 through putative phage transfer and horizontal transfer respectively. By contrast, loss of the avrPto/hopAB redundant effector group was observed in cherry-pathogenic clades. Ectopic expression of hopAB and hopC1 triggered the hypersensitive reaction in cherry leaves, confirming computational predictions.
- Cherry canker provides a fascinating example of convergent evolution of pathogenicity that is explained by the mix of effector and toxin repertoires acting on a common host.

#### Introduction

Pseudomonas syringae is a species complex, associated with plants and the water cycle, comprising several divergent clades that frequently recombine (Young, 2010; Parkinson et al., 2011; Berge et al., 2014; Baltrus et al., 2017). It is a globally important pathogen, causing disease on over 180 different host species. P. syringae is responsible for recurring chronic diseases in perennial crops, such as cherry canker (Lamichhane et al., 2014), and also sporadic outbreaks on annual crops, such as bacterial speck of tomato (Solanum lycopersicum) (Şahin, 2001). Individual strains are reported to be specialized and assigned to over 60 hostspecific pathovars; some of these are further divided into races which show host genotype specificity (Joardar et al., 2005). Strains also exist that can infect a variety of crop species, indicating that specialization is not always the norm (Monteil et al., 2013; Bartoli et al., 2015a,b). This complexity makes P. syringae an important model to study the evolution of host specificity (O'Brien et al., 2011; Mansfield et al., 2012).

High-throughput sequencing has become a major tool in bacteriology (Edwards & Holt, 2013). With the increasing number of genomes available, population-level analyses allow complex evolutionary questions to be addressed, such as how disease epidemics emerge and what ecological processes drive the evolution of pathogenicity (Guttman et al., 2014; Monteil et al., 2016). Before genomic methods were available, mutational studies of P. syringae were used to identify functional virulence factors in pathogenesis, such as type III secretion system effector (T3E) repertoires and toxins (Lindgren, 1997; Bender et al., 1999). Some T3Es were also shown to act as avirulence (avr) factors when detected by a corresponding pathogen recognition (R) protein in the host, leading to effectortriggered immunity (ETI) (Jones & Dangl, 2006). ETI is often associated with the hypersensitive response (HR) which is a cell death mechanism important in preventing pathogen spread (Morel & Dangl, 1997). Evidence suggests that P. syringae has evolved a functionally redundant repertoire of effectors, which is postulated to allow pathogen populations to lose/modify expendable avr elicitors, with minimal impact on overall virulence (Arnold & Jackson, 2011). It is proposed that as pathogen lineages specialize, they finetune their effector repertoires to maximize fitness in this niche by ensuring adequate growth and transmission, whilst avoiding detection by the plant immune system. Host range becomes restricted because the pathogen may lose effectors important for disease on other hosts or gain effectors detected by other plant species (SchulzeLefert & Panstruga, 2011). Many genomics studies have therefore focused on linking variation in virulence gene complements with particular diseases (Baltrus *et al.*, 2011, 2012; O'Brien *et al.*, 2012).

Much of the understanding of *P. syringae*–plant interactions has been achieved using herbaceous plant models. Woody pathosystems provide a greater challenge (Lamichhane *et al.*, 2014). Population genomics of *P. syringae* pv *actinidiae*, the causal agent of kiwifruit canker, revealed that three pathogenic clades, with distinct effector sets, have arisen during kiwifruit cultivation (McCann *et al.*, 2013, 2017). A study of the olive pathogen *P. syringae* pv *savastanoi* revealed that the *hopBL* effector family is overrepresented in wood-infecting pathovars (Matas *et al.*, 2014). Genes involved in the metabolism of aromatic compounds, phytohormone production, tolerance to reactive oxygen species and sucrose metabolism have also been associated with virulence on woody tissues (Green *et al.*, 2010; Bartoli *et al.*, 2015; Buonaurio *et al.*, 2015; Nowell *et al.*, 2016).

This study used genomics to examine the evolution of strains that cause bacterial canker on sweet and wild cherry (both Prunus avium). Clades of *P. syringae* that constitute the main causal agents of bacterial canker include P. syringae pv morsprunorum (Psm) race 1 and race 2 and P. syringae pv syringae (Pss) (Bull et al., 2010; Bultreys & Kaluzna, 2010). In addition, P. syringae pv avii causes bacterial canker of wild cherry (Ménard et al., 2003). The cherrypathogenic clades of *P. syringae* are reported to exhibit differences in virulence, host range and lifestyle (Crosse & Garrett, 1966; Scortichini, 2010), making the P. syringae-cherry interaction a good pathosystem to study convergent gain of pathogenicity. The genomic analysis has been coupled with robust pathogenicity testing (Hulin et al., 2018) and functional analysis of potential avirulence (avr) genes. This study provides a proof of concept that genomics-based methods can be used to identify candidate genes involved in disease and will likely become the major tool in disease monitoring, diagnostics and host range prediction.

#### Materials and Methods

#### Bacteria, plants and pathogenicity tests

Methods used for bacterial culture and sources of plants were as described in Hulin *et al.* (2018) and are detailed in Supporting Information Methods S1. Plant species utilized included *P. avium* L. and *Nicotiana tabacum* L. *Pseudomonas* strains are listed in Table 1. *Escherichia coli* was grown on lysogeny broth (LB) agar plates and grown overnight at 37°C. Antibiotic concentrations: kanamycin, 50 μg ml<sup>-1</sup>; gentamycin, 10 μg ml<sup>-1</sup>; spectinomycin, 100 μg ml<sup>-1</sup>, introfurantoin, 100 μg ml<sup>-1</sup>. X-gal was used at a concentration of 80 μg ml<sup>-1</sup>. Tables S1–S3 list the *P. syringae* mutants, plasmids and primers used in this study.

Pathogenicity tests, performed on detached cherry leaves and analysed as in Hulin *et al.* (2018) are described in Methods S1.

### Genome sequencing, assembly and annotation

Bioinformatics commands for analyses performed in this paper are available on Github (https://github.com/harrisonlab/pseud

omonas/). Genome sequencing using Illumina and genome assembly were performed as in Hulin *et al.* (2018). For long-read sequencing, PacBio (Pacific Biosystems, Menlo Park, CA, USA) and MinION (Oxford Nanopore, Oxford, UK) were used. High molecular weight DNA was extracted using a cetyltrimethylammonium bromide method (Feil *et al.*, 2012). For the PacBio sequencing of strains R1-5244, R2-leaf and syr9097, DNA samples were sent to the Earlham Institute (Norwich) for PacBio RSII sequencing.

For MinION sequencing of *Psm* R1-5300, the DNA library was prepared using the RAD001 rapid-prep kit (Oxford Nanopore) and run on the Oxford Nanopore MinION, flowcell vR9.5 followed by basecalling using Metrichor (Oxford Nanopore). MinION reads were extracted from Fast5 files using Poretools (Loman & Quinlan, 2014). The sequencing reads for both long-read technologies were trimmed and assembled using Canu (Berlin *et al.*, 2015), and Circlator was used to circularize contigs (Hunt *et al.*, 2015). The assemblies were polished using error-corrected Illumina reads with Bowtie2, Samtools and Pilon 1.17 (Li *et al.*, 2009; Langmead & Salzberg, 2013; Walker *et al.*, 2014).

Plasmid profiling was performed using an alkaline-lysis method and gel electrophoresis (Moulton *et al.*, 1993; Neale *et al.*, 2013). Genomes were submitted to GenBank and accession numbers are listed in Table 1.

#### Orthology analysis

OrthoMCL (Li *et al.*, 2003) was used to identify orthologous genes. All genomes were re-annotated using RAST (Aziz *et al.*, 2008) to ensure similar annotation quality. For this reason, the Illumina short-read assemblies of the four long-read sequenced genomes (R1-5244, R1-5300, R2-leaf and syr9097) were used in orthology analysis. OrthoMCL was run with default settings and a 50 residue cut-off length. All RAST annotated protein files used in this analysis are available on Github (https://github.com/harrisonlab/pseudomonas/).

## Phylogenetic and genomic analysis of *Pseudomonas* syringae

Nucleotide sequences of single-copy genes present in all genomes were aligned using CLUSTALW (Larkin *et al.*, 2007) and trimmed using GBLOCKS (Castresana, 2000). Gene alignments were concatenated using GENEIOUS (Kearse *et al.*, 2012). RAXML-AVX v.8.1.15 (Stamatakis, 2014) was used in partitioned mode to build the maximum likelihood phylogeny, with a general time reversible (GTR) gamma model of substitution and 100 nonparametric bootstrap replicates. To detect core genes that may have undergone recombination, the program GENECONV (Sawyer, 1989) was used as in Yu *et al.* (2016). Whole-genome alignments were performed using PROGRESSIVEMAUVE (Darling *et al.*, 2010).

#### Virulence and mobility gene identification

All T3E-encoding protein sequences were downloaded from pseudomonas-syringae.org, including the recent classification of

Table 1 List of bacterial strains used in this study, including a range of cherry pathogens and nonpathogens

								Pathogenicity tested	
Strain	Pathovar	Race	PG	Isolation source	Isolator	Prunus cv	Sequenced	on cherry ( <i>Prunus avium</i> )	Accession number
avii5271	avii		1	Prunus avium	Garrett, 1990, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004)	NBAO00000000
R1-5270	morsprunorum	1	3	Prunus avium	Garrett, 1990, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004)	NBAN00000000
R2-7968A	morsprunorum	2	1	Prunus avium	Vicente, 2000, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004)	NBAI00000000
R2-9095	morsprunorum	2	1	Prunus avium	Roberts, 2010, UK	Wild cherry	This work	M. Hulin, pers. obs.	MLED00000000
syr5264	syringae		2	Prunus avium	Garrett, 1990, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004)	NBAQ00000000
syr5275	syringae		2	Prunus avium	Garrett, 1990, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004)	NBAP00000000
syr7928A	syringae		2	Prunus avium	Vicente, 2000, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004)	NBAL00000000
syr8094A	syringae		2	Prunus avium	Vicente, 2001, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004)	NBAK00000000
Ps-7928C	Unknown		2	Prunus avium	Vicente, 2000, UK	Wild cherry	This work	Vicente et al. (2004)	NBAM0000000
Ps-7969	Unknown	1	2	Prunus avium	Vicente, 2000, UK	Wild cherry	This work	Vicente <i>et al.</i> (2004) <b>Hulin <i>et al.</i></b>	NBAJ00000000 CP026557–
R1-5244 R1-5300	morsprunorum morsprunorum		<b>3</b>	Prunus avium Prunus domestica	Crosse, 1960, UK Prunier, UK	<b>Unknown</b> Victoria	This work This work	(2018) Hulin <i>et al.</i>	CP026561 MLEN00000000
R2-leaf	morsprunorum		1	Prunus avium	Hulin, 2014, UK	Napoleon	This work	(2018) Hulin <i>et al.</i>	CP026562-
syr9097	syringae	_	2	Prunus avium	Roberts, 2010,	Unknown	This work	(2018) Hulin <i>et al.</i>	CP026567 CP026568
3912027	syrrigae		_	Trumus avium	UK	Olikilowii	TIII3 WOIK	(2018)	CI 020300
syr2675	syringae		2	Phaseolus vulgaris	1965, Kenya		This work	This work	MLEX00000000
syr2676 syr2682	syringae		2	Phaseolus vulgaris Phaseolus vulgaris	1990, Lesotho		This work This work	nt nt	MLEY00000000 MLFA00000000
syr3023	syringae		2	Syringa vulgaris	1950, Lesotho 1950, UK		This work	This work	MLFD00000000
,	syringae		2	Phaseolus lunatus	1962, Kenya		This work	This work	MLEV00000000
syr100 R1-9326	syringae morsprunorum	1	3	Prunus domestica	Roberts, 2011,	Victoria	Hulin et al.	Hulin et al.	MLEO00000000
R1-9629	morsprunorum	1	3	Prunus domestica	UK Roberts, 2012,	Victoria	(2018) Hulin <i>et al</i> .	(2018) Hulin <i>et al</i> .	MLEP00000000
R1-9646	morsprunorum	1	3	Prunus avium	UK Roberts, 2012,	Stella	(2018) Hulin et al.	(2018) Hulin <i>et al.</i>	MLEE00000000
R1-9657	morsprunorum	1	3	Prunus avium	UK Roberts, 2012,	Kiku-Shidare		(2018) Hulin <i>et al</i> .	MLEF00000000
R2-5255	morsprunorum	2	1	Prunus avium	UK Prunier, UK	Napoleon	(2018) Hulin et al.	(2018) Hulin <i>et al.</i>	MLEC00000000
R2-5260	morsprunorum	2	1	Prunus avium	Garrett, UK	Roundel	(2018) Hulin <i>et al.</i> (2018)	(2018) Hulin <i>et al.</i> (2018)	MLEG0000000
R2-SC214	morsprunorum	2	1	Prunus avium	Roberts, 1983, UK	Wild cherry	Hulin <i>et al.</i> (2018)	Hulin <i>et al.</i> (2018)	MLEI00000000
syr9293	syringae		2	Prunus domestica	Roberts, 2011, UK	Victoria	Hulin <i>et al.</i> (2018)	Hulin <i>et al.</i> (2018)	MLEQ00000000
syr9630	syringae		2	Prunus domestica		Victoria	Hulin <i>et al.</i> (2018)	Hulin <i>et al.</i> (2018)	MLER00000000
syr9644	syringae		2	Prunus avium	Roberts, 2012, UK	Stella	Hulin <i>et al.</i> (2018)	Hulin <i>et al.</i> (2018)	MLEK00000000
syr9654	syringae		2	Prunus domestica	Roberts, 2012, UK	Victoria	Hulin <i>et al.</i> (2018)	Hulin <i>et al.</i> (2018)	MLES00000000
syr9656	syringae		2	Prunus avium	Roberts, 2012, UK	Kiku-Shidare		Hulin <i>et al.</i> (2018)	MLEM00000000
syr9659	syringae		2	Prunus avium	Roberts, 2012, UK	Kiku-Shidare		Hulin <i>et al.</i> (2018)	MLEL00000000



Table 1 (Continued)

								Pathogenicity tested	A accesis :-
Strain	Pathovar	Race	PG	Isolation source	Isolator	Prunus cv	Sequenced	on cherry ( <i>Prunus avium</i> )	Accession number
Ps-9643	Unknown		1	Prunus domestica	Roberts, 2012, UK	Victoria	Hulin <i>et al.</i> (2018)	Hulin <i>et al.</i> (2018)	MLET00000000
avii3846	avii		1	Prunus avium	1991, France	Wild cherry	Nowell <i>et al.</i> (2016)	Ménard <i>et al.</i> (2003)	LIIJ00000000
R1-2341	morsprunorum	1	3	Prunus cerasus	1988, Hungary	Unknown	Nowell <i>et al.</i> (2016)	nt	LIIB00000000
R1-5269	morsprunorum	1	3	Prunus cerasus	Garrett, 1990, UK	Wild cherry	Nowell <i>et al.</i> (2016)	Vicente <i>et al.</i> (2004)	LIHZ00000000
R2-5261	morsprunorum	2	1	Prunus avium	Garrett, UK	Roundel	Nowell <i>et al.</i> (2016)	Vicente <i>et al.</i> (2004)	LIIA00000000
R2-302280	morsprunorum		1	Prunus domestica	USA	Unknown	Baltrus <i>et al.</i> (2011)	Gilbert <i>et al.</i> (2009)*	AEAE00000000
syr2339	syringae		2	Prunus avium	1984, Hungary	Unknown	Nowell <i>et al.</i> (2016)	nt	LIHU00000000
syr7872	syringae		2	Prunus domestica	Lewis, 2000, UK	Opal	Nowell <i>et al.</i> (2016)	Vicente <i>et al.</i> (2004)	LIHS00000000
syr7924	syringae		2	Prunus avium	Vicente, 2000, UK	Wild cherry	Nowell <i>et al.</i> (2016)	Vicente <i>et al.</i> (2004)	LIHR00000000
acer302273	aceris		2	Acer sp.	USA		Baltrus <i>et al.</i> (2011)	nt	AEAO00000000
acti18884	actinidiae		1	Actinidia deliciosa	2010, New Zealand		McCann <i>et al.</i> (2013)	nt	AOKO00000000
acti19073	actinidiae		1	Actinidia deliciosa			McCann <i>et al.</i> (2013)	nt	AOJR00000000
acti212056	actinidiae		1	Actinidia deliciosa	2012, Japan		Sawada <i>et al.</i> (2015)	nt	BBWG00000000
acti302091	actinidiae		1	Actinidia deliciosa	1984, Japan		Baltrus <i>et al.</i> (2011)	nt	AEAL00000000
actiCRAFRU	actinidiae		1	Actinidia deliciosa	2010, Italy		Butler <i>et al.</i> (2013)	nt	ANGD00000000
actiNCPPB3871	actinidiae		1	Actinidia deliciosa	1992, Italy		Marcelletti et al. (2011)	nt	ANGD00000000
aes089323	aesculi		3	Aesculus hippocastanum	India, 1980		Baltrus <i>et al.</i> (2011)	nt	AEAD00000000
aes2250	aesculi		3	Aesculus hippocastanum	2008, UK		Green <i>et al.</i> (2010)	nt	ACXT00000000
aes3681	aesculi		3	Aesculus	1969, India		Green et al.	nt	ACXS00000000
amy3205	amygdali		3	hippocastanum Prunus dulcis	1967, Greece		(2010) Bartoli <i>et al</i> .	nt	JYHB00000000
amyICMP3918	amygdali		3	Prunus dulcis	Panagopoulos, 1967. Greece		(2015a) Thakur et al. (2016)	nt	LJPQ00000000
avelBP631	avellanae		1	Corylus avellana	1976, Greece		O'Brien <i>et al.</i> (2012)	Hulin <i>et al.</i> (2018)	AKBS00000000
avelVe037	avellanae		2	Corylus avellana	1990, Italy		O'Brien <i>et al.</i> (2012)	nt	AKCJ00000000
BRIP34876	Unknown		2	Hordeum vulgare	1971, Australia		Gardiner	nt	AMXK00000000
castCFBP4217	castaneae		3	Castanea crenata	1977, Japan		et al. (2013) Nowell et al.	nt	LIIH00000000
CC1416	Unknown		1	Epilithon	USA		(2016) Baltrus <i>et al</i> .	nt	AVEP00000000
CC1544	Unknown		1	Lake water	France		(2014b) Baltrus <i>et al.</i>	nt	AVEI00000000
CC1559	Unknown		1	Snow	France		(2014b) Baltrus <i>et al.</i>	nt	AVEG00000000
CC94	Unknown		2	Cantaloupe	France		(2014b) Baltrus <i>et al</i> . (2014b)	nt	AVEA00000000

Table 1 (Continued)

Strain	Pathovar	Race	PG	Isolation source	Isolator	Prunus cv	Sequenced	Pathogenicity tested on cherry (Prunus avium)	Accession number
cera6109	cerasicola		3	Prunus yedoensis	1995, Japan		Nowell et al.	nt	LIIG00000000
ceralCMP17524	cerasicola		3	Prunus yedoensis	Japan		(2016) Thakur <i>et al</i> .	nt	LJQA00000000
ciccICMP5710	ciccaronei		3	Ceratonia siliqua	Italy		(2016) Thakur <i>et al.</i> (2016)	nt	LJPY00000000
cunnICMP11894	cunninghamiae		3	Cunninghamia Ianceolata	China		Thakur <i>et al</i> . (2016)	nt	LJQE00000000
daphICMP9757	daphniphylli		3	Daphniphyllum teijsmannii	Japan		Thakur <i>et al.</i> (2016)	nt	LJQF00000000
delphi569	delphinii		1	Delphinium sp.	New Zealand		Thakur <i>et al.</i> (2016)	nt	LJQH00000000
dendro3226	dendropanacis		3	Dendropanax trifidus	1979, Japan		Bartoli <i>et al.</i> (2015a)	nt	JYHG00000000
dendro4219	dendropanacis		3	Dendropanax trifidus	1981, Japan		Bartoli <i>et al.</i> (2015a)	nt	JYHD00000000
dendro9150	dendropanacis		3	Dendropanax trifidus	Japan		Thakur <i>et al.</i> (2016)	nt	LJQG00000000
erio4455	eriobotryae		3	Eriobotrya japonica	USA		Thakur <i>et al.</i> (2016)	nt	LJQ100000000
glyR4	glycinea		3	Glycine max	Cross, 1960, USA		Qi <i>et al.</i> (2011)	nt	AEGH00000000
ICMP19498	Unknown		3	Actinidia deliciosa	2010, New Zealand		Visnovsky et al. (2016)	nt	LKCH00000000
lach301315	lachrymans		3	Cucumis sativus	Japan		Baltrus <i>et al.</i> (2011)	nt	AEAF00000000
lach302278	lachrymans ,		1	Cucumis sativus	USA		Baltrus <i>et al.</i> (2011)	nt	AEAM00000000
lapsaICMP3947	lapsa		2	Zea sp.	Unknown		Thakur <i>et al</i> . (2016)	nt	LJQQ00000000
meli6289	meliae		3	Melia azedarach	Japan		Thakur <i>et al.</i> (2016)	nt	LJQT00000000
morsU7805	morsprunorum		3	Prunus mume	Unknown		Mott <i>et al.</i> (2016)	nt	LGLQ00000000
myriAZ8448	myricae		3	Myrica rubra	Japan		Thakur <i>et al.</i> (2016)	nt	LGLA00000000
neriilCMP16943	savastanoi		3	Olea europea	Spain		Thakur <i>et al</i> . (2016)	nt	LJQW00000000
paniLMG2367	panici ,		2	Panicum miliaceum	Unknown		Liu <i>et al.</i> (2012)	nt	ALAC00000000
papu1754	papulans		2	Malus sylvestris	1973, Canada		Nowell <i>et al</i> . (2016)	nt	JYHI00000000
persNCPPB2254	persicae		1	Prunus persica	1972, France		Zhao <i>et al.</i> (2015)	nt	LAZV00000000
photICMP7840	photiniae 		3	Photinia glabra	Japan		Thakur <i>et al</i> . (2016)	nt	LJQO0000000
pisiPP1	pisi		2	Pisum sativum	Japan		Baltrus <i>et al.</i> (2014a)	nt	AUZR00000000
phas1448a	phaseolicola		3	Phaseolus vulgaris	Teverson, 1965, Ethiopia		Joardar <i>et al.</i> (2005)	Hulin <i>et al.</i> (2018)	CP000058
rhapCFBP4220	rhaphiolepidis		3	Rhaphiolepis umbellata	1980, Japan		Nowell <i>et al.</i> (2016)	nt	LIHV00000000
RMA1	Unknown		1	Aquilegia vulgaris	Jackson, 2012, UK		Hulin <i>et al.</i> (2018)	Hulin <i>et al.</i> (2018)	MLEU00000000
sava3335	savastanoi		3	Olea europea	Stead, France		Rodríguez- Palenzuela	nt	ADMI00000000
sava4352	savastanoi		3	Olea europea	Yugoslavia		et al. (2010) Thakur et al. (2016)	nt	LGKR00000000



Table 1 (Continued)

Strain	Pathovar	Race	PG	Isolation source	Isolator	Prunus cv	Sequenced	Pathogenicity tested on cherry (Prunus avium)	Accession number
savaDAPP-PG722	savastanoi		3	Olea europea	Italy		Moretti <i>et al.</i> (2014)	nt	JOJV00000000
savaPseNe107	savastanoi		3	Olea europea	Balestra, Nepal		Bartoli <i>et al.</i> (2015a)	nt	JYHF00000000
soliICMP16925	solidagae		2	Solidago altissima	Japan		Thakur <i>et al.</i> (2016)	nt	JYHF00000000
syr1212	syringae		2	Pisum sativum	UK		Baltrus <i>et al.</i> (2014a)	This study	AVCR00000000
syr2340	syringae		2	Pyrus sp.	1985, Hungary		Nowell <i>et al.</i> (2016)	nt	LIHT00000000
syr41a	syringae		2	Prunus armeniaca	2011, France		Bartoli <i>et al.</i> (2015a)	nt	JYHJ00000000
syrB301D	syringae		2	Pyrus communis	Garrett, 1959, UK		Ravindran et al. (2015)	nt	CP005969
syrB64	syringae		2	Triticum aestivum	Wilcoxson, USA		Dudnik & Dudler (2014)	nt	ANZF00000000
syrB728a	syringae		2	Phaseolus vulgaris	1987, USA		Feil <i>et al</i> . (2005)	This work	CP000075
syrHS191	syringae		2	Panicum miliaceum	Hayward, Australia, 1969		Ravindran et al. (2015)	nt	CP006256
syrUMAF0158	syringae		2	Mangifera indica	Cazorla, 2010, Spain		Martínez- García <i>et al</i> . (2015)	nt	CP005970
thea3923	theae		1	Camelia sinensis	1974, Japan		Mazzaglia et al. (2012)	nt	AGNN00000000
tomaDC3000	tomato		1	Solanum lycopersicum	1960, UK		Buell <i>et al.</i> (2003)	nt	AE016853
tomaT1	tomato		1	Solanum lycopersicum	1986, Canada		Almeida <i>et al.</i> (2009)	nt	ABSM00000000
UB303	Unknown		2	Lake water	France		Baltrus <i>et al.</i> (2014b)	nt	AVDZ00000000
ulmiICMP3962	ulmi		3	Ulmus sp.	Yugoslavia		Thakur <i>et al.</i> (2016)	nt	LJRQ00000000
USA007	Unknown		1	Stream water	USA		Baltrus <i>et al.</i> (2014b)	nt	AVDY00000000
USA011	Unknown		1	Stream water	USA		Baltrus <i>et al.</i> (2014b)	nt	AVDX00000000

Pathovar designation, phylogroup, isolation information, cherry pathogenicity (reference for when tested; nt, not tested), publication of genome sequence and NCBI accession numbers are listed. Strains in bold were considered pathogenic in cherry. cv, cultivar of sweet cherry or plum. Long-read sequenced genomes are highlighted with shading. Strains are ordered, first with those sequenced in this study, followed by other *Pseudomonas* strains from cherry and plum used in plasmid profiling analysis and previously pathogenicity tested in Hulin *et al.* (2018). Next, further strains isolated from cherry and plum sequenced elsewhere are listed. Finally, the remaining strains were only used in comparative analysis. Note that all 108 genomes were used in initial orthology analysis but only 102 were used in the final phylogeny and comparative genomics.

\*The pathogenic status of MAFF302280 on cherry is debated. This strain is reported to be the pathotype strain of *P. syringae* pv morsprunorum (*Psm*; Sawada *et al.*, 1999), so is assumed to be equivalent to CFBP 2351, NCPPB2995, ICMP5795 and LMG5075. The strain NCPPB2995 was reported to be potentially nonpathogenic (Gardan *et al.*, 1999). Whilst, the 'same' strain LMG5075 tested positive for pathogenicity in a recent publication (Gilbert *et al.*, 2009). There is no definite link showing that MAFF302280 is the same strain as the others listed as it is not linked to them in online databases (http://www.straininfo.net/) or taxonomy-focused publications (Bull *et al.*, 2010). It is assumed to be putatively pathogenic in this study owing to its close relatedness to other *Psm* R2 strains; however, further pathogenicity tests would be required to fully confirm this.

HopF effectors into four alleles (Lo *et al.*, 2016). tBlastn (Altschul *et al.*, 1990) was used to search each genome for homologues with a score of  $\geq$  70% identity and  $\geq$  40% query length to at least one sequence in each effector family. Nucleotide sequences were extracted and manually examined for frameshifts or truncations. Disrupted genes were classed as pseudogenes. A

heatmap of effector presence was generated using heatmap.2 in gplots (Warnes *et al.*, 2016). Interproscan (Quevillon *et al.*, 2005) identified protein domains, and Illustrator for Biological Sequences was used for visualization (Liu *et al.*, 2015). Genomic regions containing effectors were aligned using MAFFT (Katoh *et al.*, 2002).

A similar analysis was performed for phytotoxin and auxin biosynthesis, wood degradation, ice nucleation and plasmid-associated genes. Protein sequences were obtained from NCBI (Table S4) and blasted against the genome sequence as noted earlier. Prophage identification was performed using Phaster (Arndt et al., 2016).

#### Gain and loss analysis

Gain loss mapping engine (GLOOME) was used to plot the gain and loss of genes on the core-genome phylogeny (Cohen *et al.*, 2010). Effector genes were considered present even if predicted to be pseudogenes, as these can still be gained and lost. The optimization level was set to 'very high', a mixture model allowing variable gain and loss distributions was used and the distribution type was GENERAL\_GAMMA\_PLUS\_INV. Highly probable events ( $P \ge 0.80$ ) on the branches leading to cherry-pathogenic strains were extracted.

#### BayesTraits analysis

BAYESTRAITS v.2 was used to correlate T3E gene evolution with pathogenicity (Pagel, 2004). A binary matrix was created of effector family presence and pathogenicity of each strain. The effector matrix was collapsed into effector families, as the different alleles likely perform similar biological functions *in planta* (Cunnac *et al.*, 2011). Putative pseudogenes were considered absent, as they may be nonfunctional. The BAYESTRAITS methodology followed an approach as in Press *et al.* (2013) and is described in detail in Methods S1.

#### Horizontal gene transfer analysis

For each effector family, best-hit nucleotide sequences were aligned using ClustalW (Larkin *et al.*, 2007). RAxML was used to build a phylogenetic tree with a GTR model of evolution and 1000 bootstrap replicates. Incongruence with the core-genome tree was examined visually. To further assess horizontal transfer, a species—gene tree reconciliation method RANGER-DTL v.2 (Bansal *et al.*, 2012) was used, as in Bruns *et al.* (2018). Full details are described in Methods S1.

#### Identification of genomic islands

Genomic islands (GIs) were identified in the PacBio-sequenced cherry pathogenic strains using IslandVIEWER3 (Dhillon *et al.*, 2015). Islands were manually delimited as in McCann *et al.* (2013). Blastn was utilized to determine if these GIs were present in other *P. syringae* genomes. As most GIs exceeded 10 kb and most genome assemblies were highly fragmented, the islands were split into 0.5 kb sections before analysis to prevent false negatives due to contig breaks. An island was concluded as likely to be present if all sections produced hits with a query length > 30%. To validate this approach, the Illumina-sequenced genome assemblies of the PacBio-sequenced strains were searched for their own islands.

#### General DNA manipulations and bacterial transformations

Cloning and other molecular biology techniques, including ectopic expression of potential *avr* genes, were as described in earlier studies (Staskawicz *et al.*, 1984; Arnold *et al.*, 2001; Kvitko & Collmer, 2011). Details are provided in Methods S1.

#### **Results**

#### Genome assembly and sequencing statistics

Eighteen *P. syringae* strains isolated from cherry and plum were phenotyped for pathogenicity and genome sequenced in a previous study (Hulin *et al.*, 2018). To increase this sample, nine strains isolated from wild cherry and five additional non-*Prunus* out-groups were genome sequenced using the Illumina MiSeq. The genomes of eight cherry strains sequenced elsewhere (Baltrus *et al.*, 2011; Nowell *et al.*, 2016) were also downloaded from NCBI.

Information on the origin and pathogenicity of each strain is summarized in Table 1. Twenty-eight were considered pathogenic to cherry, including all *Pss* isolated from cherry and plum. By contrast, three *Psm* race 1 strains from plum (R1-5300, R1-9326 and R1-9629) and one from cherry, strain R1-9657, failed to induce canker on cherry following tree inoculations; and three strains of unknown taxonomy isolated from plum and cherry (Ps-9643, Ps-7928C and Ps-7969) were also non-pathogenic (references in Table 1). The cherry pathogens are referred to as their described pathovar names throughout this study. To simplify figures, cherry pathogens are highlighted and the first few letters of the pathovar name were used. '*Pss*' becomes '*syr*', as otherwise *Pss* could refer to other pathovars beginning with 's' (e.g. *savastanoi*).

All strains included in this study were sequenced using Illumina MiSeq. Three representative strains (R1-5244, R2-leaf and syr9097) were sequenced using PacBio and the nonpathogenic Psm R1 strain R1-5300 was sequenced using Oxford Nanopore, to obtain more complete genomes. Table 2 summarizes the genome assembly statistics for all strains sequenced in this study and Hulin et al. (2018). Illumina genomes assembled into 23-352 contigs, whilst the long-read sequenced genomes assembled into one to six contigs. The number of plasmids in each strain was determined by plasmid profiling (Fig. S1). Psm R1 and R2 strains possessed between two and six plasmids, P. syringae pv avii 5271 possessed six plasmids, whereas, apart from three strains (syr5275, syr7928A, syr9644) with one plasmid each, most cherry-pathogenic strains of Pss did not possess plasmids. The strain syr9097, which was sequenced using PacBio, lacked plasmids. The genomes sequenced with long-read technology all assembled into the correct number of contigs corresponding to chromosome and plasmids, apart from R1-5300. The chromosome of this strain was separated into two contigs (tig0 and tig75), based on a whole-genome alignment with Psm R1 strain R1-5244 (Fig. S2).

The Psm R1 (R1-5244, R1-5300) and Psm R2 (R2-leaf) long-read assemblies revealed putatively complete plasmid contigs

**Table 2** Assembly statistics for all strains sequenced in this study and Hulin et al. (2018)

Assembly	Sequencing	Reference	No. of contigs	Plasmids	Total length	GC%	N50	Coverage	Features
R1-5270	Illumina	This work	185	3	6258 313	58.10	202 152	134	5770
R1-9326	Illumina	Hulin et al. (2018)	268	4	6353 636	57.91	142 021	81	5874
R1-9629	Illumina	Hulin et al. (2018)	216	3	6341 664	57.94	142 021	172	5856
R1-9646	Illumina	Hulin et al. (2018)	171	3	6302 776	58.03	235 429	180	5801
R1-9657	Illumina	Hulin et al. (2018)	191	4	6317 852	57.91	145 272	158	5848
R2-5255	Illumina	Hulin et al. (2018)	206	2	6448 834	58.38	102 760	112	5966
R2-7968A	Illumina	This work	278	6	6498 711	58.42	91 262	134	6016
R2-5260	Illumina	Hulin et al. (2018)	223	3	6495 620	58.41	101 794	458	5995
R2-9095	Illumina	This work	304	2	6418 849	58.48	92 453	100	5887
R2-SC214	Illumina	Hulin et al. (2018)	203	3	6253 818	58.56	108 341	180	5747
avii5271	Illumina	This work	352	6	6243 644	58.56	56 064	127	5809
Ps-9643	Illumina	Hulin et al. (2018)	58	1	5937 102	58.78	243 355	212	5386
syr9293	Illumina	Hulin et al. (2018)	73	0	6135 031	58.84	557 853	196	5302
syr9630	Illumina	Hulin et al. (2018)	57	0	5940 819	59.33	347 701	206	5175
syr9644	Illumina	Hulin et al. (2018)	75	1	6173 193	59.13	251 053	208	5334
syr9654	Illumina	Hulin et al. (2018)	49	0	5941 610	59.37	245 023	147	5148
syr9656	Illumina	Hulin et al. (2018)	39	0	5980 728	59.10	1007 808	205	5184
syr9659	Illumina	Hulin et al. (2018)	51	0	5943 090	59.37	235 830	116	5148
syr5264	Illumina	This work	59	0	6029 896	59.08	380 149	114	5314
syr5275	Illumina	This work	64	1	5994 091	59.30	371 492	145	5207
syr7928A	Illumina	This work	59	1	6129 363	59.26	371 492	141	5338
syr8094A	Illumina	This work	71	0	5942 438	59.33	265 238	106	5184
syr7928C	Illumina	This work	49	1	5994 455	59.17	325 175	124	5318
syr7969	Illumina	This work	92	0	6185 932	59.01	164 374	151	5476
RMA1	Illumina	Hulin et al. (2018)	95	1	6306 889	58.73	187 448	320	5825
syr100	Illumina	This work	23	0	5872 916	59.33	893 822	83	5140
syr2675	Illumina	This work	65	0	5994 384	59.34	227 612	83	5177
syr2676	Illumina	This work	90	1	6158 476	59.30	259 660	78	5387
syr2682	Illumina	This work	185	1	6259 099	59.21	242 212	84	5405
syr3023	Illumina	This work	228	0	6203 212	58.90	456 738	88	5365
R1-5244	PacBio	This work	5	4	6445 963	58.05	6109 228	82	6024
R2-leaf	PacBio	This work	6	5	6576 340	58.41	6242 845	141	6093
syr9097	PacBio	This work	1	0	5929 959	59.30	5929 959	100	5147
R1-5300	MinION	This work	6	4	645 601	57.87	5688 034	100	6449

Cherry pathogens are in bold. Long-read sequenced genomes are highlighted with shading. N50, the weighted median contig size in the assembly; Features, the number of protein encoding and RNA sequences in the RAST annotated genome.

containing plasmid-associated genes (Tables S5–S7). All three strains (R1-5244, R1-5300 and R2-leaf) possessed plasmids with *repA* homologues, indicating they may belong to common plasmid family pPT23A (Zhao *et al.*, 2005). Several plasmids also contained T4SS conjugational machinery (VirB/D), so may be conjugative.

#### Core-genome phylogenetic analysis

To examine the relatedness of strains, an analysis of core genes was carried out using 108 genomes of strains from the well-studied phylogroups 1–3 isolated from both plants and aquatic environments. A maximum likelihood phylogeny based on 1035 concatenated core genes was constructed (Fig. S3). There was low support for certain P2 and P3 clades based on bootstrap analysis. To determine if particular taxa were causing low support, the analysis was systematically repeated for the two phylogroups, with non-cherry strains removed. Support and tree likelihood values were compared (Table S8). Within P3, the removal of *P. syringae* pv *eriobotryae* or *P. syringae* pv *daphniphylli* improved

support, whilst the removal of *P. syringae* pv *syringae* 1212 improved support values in P2 (Figs S4, S5). The global analysis was then repeated with these taxa removed (Figs S6–S9). The final phylogeny (Fig. 1), with the highest mean branch support (92.8%), lacked *P. syringae* pv *eriobotryae*. The phylogeny, built using a 611 888 bp alignment, contained 102 taxa due to the removal of strains found to be identical to others (dendro4219, syr9630, R1-9629, R1-9326 and R1-5269). Most support values exceeded 70%, with good support for branches leading to cherrypathogenic clades.

One explanation for the low support within P2 and P3 was that these clades have undergone core-genome recombination. The program Geneconv (Sawyer, 1989) showed that 140 genes had putatively recombined (127 288 bp total length, 20.8% of the alignment). Table S9 lists the number of recombination events per phylogroup. The most frequent core gene recombination occurred in P3 (73 genes affected), followed by 31 genes in P2 and only 13 in P1.

Cherry pathogens were found in all three phylogroups. The two *Psm* races (R1 in P3, R2 in P1) and *P. syringae* pv *avii* (P1)

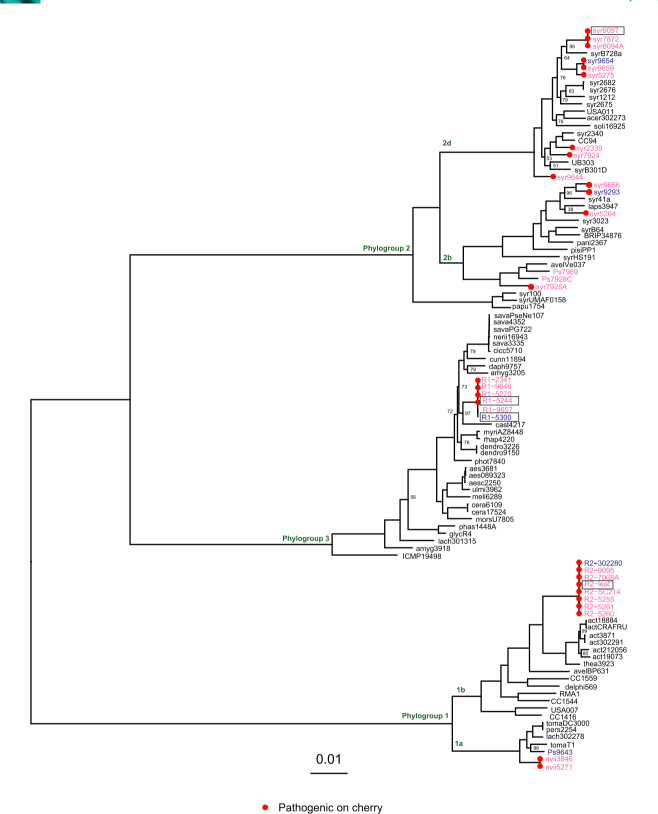


Fig. 1 Core-genome phylogenetic tree. Multi-locus phylogeny based on 1035 genes which represent the core genome of *Pseudomonas syringae*. Strains from cherry and plum are highlighted in pink and blue respectively. Strains pathogenic to cherry (assessed in Hulin *et al.*, 2018; Vicente *et al.*, 2004) are labelled with red circles. Strains with long-read sequenced genomes are in black boxes. Phylogroups are also labelled for reference. Percentage bootstrap support values below 99% are shown for each node. The bar is nucleotide substitutions per site.

formed monophyletic clades. Within *Psm* R1, strains pathogenic to cherry formed a clade distinct from previously classified non-pathogenic strains (Hulin *et al.*, 2018), indicating that there has been divergence in their core genomes. By contrast, *Prunus*-infecting strains of *Pss* were found across P2, interspersed with strains isolated from other plants and aquatic environments. To ensure that genomic comparisons between P2 strains were based on differential pathogenicity, several related non-*Prunus* strains were pathogenicity tested on cherry leaves. *In planta* bacterial populations of non-*Prunus* strains were reduced compared with *Prunus Pss* strains (Fig. S10; Table S10).

#### Search for virulence factors

The *hrp* pathogenicity island All sequenced strains contained the *hrp* pathogenicity island required for conventional Type III secretion. Core effector genes from the adjacent conserved effector locus (Alfano *et al.*, 2000), such as *avrE1*, *hopM1* and *hopAA1*, were present (Fig. S11), However, *hopAA1* was truncated in both *Psm* R1 and R2 due to inversion events. The *hopAA1* gene was truncated in *Psm* R2, whilst in *Psm* R1 both *hopAA1* and *hopM1* were truncated (Fig. S12).

Type III effectors and other virulence genes Genomes were then scanned for known virulence genes and a heatmap of presence, absence and pseudogenization was constructed (Fig. 2). In terms of T3Es, there was variation both between and within the different cherry-pathogenic clades. Notably, *Psm* R1, which contained strains pathogenic and nonpathogenic on cherry, showed clear differentiation in effector repertoires (Table S11). *Psm* R1, R2 and *P. syringae* pv *avii* possessed 24–34 effector genes, whereas *Pss* strains possessed nine to 15. The reduced effector repertoire of *Pss* was representative of P2 strains as previously noted (Baltrus *et al.*, 2011; Dudnik & Dudler, 2014). Table 3 lists the effectors in each long-read genome assembly in order of appearance.

Non-T3 virulence factors were identified. All pathogenic *Psm* R1 strains possessed the coronatine biosynthesis clusters, which were plasmid borne in *Psm* R1-5244. All cherry-pathogenic *Pss* strains possessed at least one biosynthesis gene cluster for the toxins syringomycin, syringolin and syringopeptin, with several strains possessing all three. Strains within clade P2b possessed the biosynthesis genes for mangotoxin. The nonpathogenic cherry P2b strains Ps7928C and Ps7969 lacked all toxin biosynthesis clusters.

A cluster of genes named WHOP (woody hosts and *Pseudomonas*) thought to be involved in aromatic compound (lignin) degradation (Caballo-Ponce *et al.*, 2016) was present in *Psm* R1 and R2, whilst *P. syringae* pv *avii* and most *Pss* strains contained no WHOP homologues. Two cherry P2d strains (syr2339 and syr7924) did, however, possess the catechol *catBCA* cluster. The genomes were also searched for the ice nucleation gene cluster. Members of *Psm* R1, *Pss* and *P. syringae* pv *avii* strains all possessed genes involved in ice nucleation (Fig. 2), whilst *Psm* R2 lacked the complete set of ice nucleation genes.

#### Associating type III effector evolution with host specificity

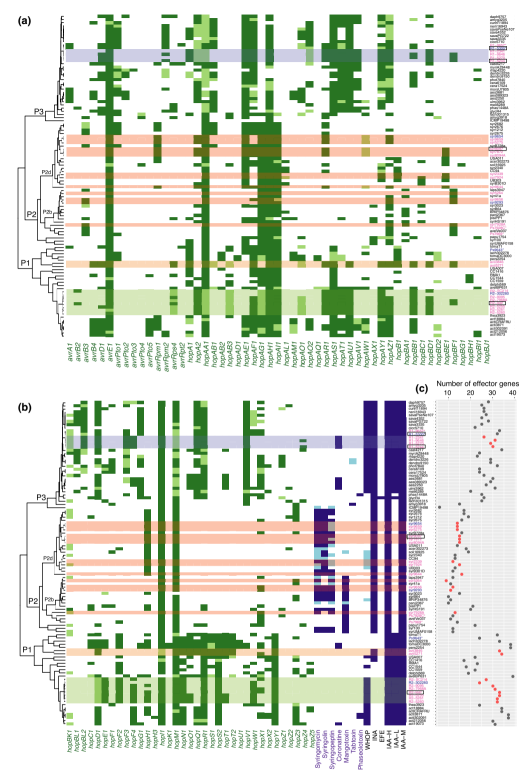
T3E evolution was statistically correlated with cherry pathogenicity, using BAYESTRAITS and GLOOME (Pagel, 2004; Cohen *et al.*, 2010). BAYESTRAITS takes a binary matrix of two traits within a phylogeny and determines if changes in the two characters (effector gene and pathogenicity) have evolved independently or dependently. Fig. 3(a) shows the likelihood ratio of cherry pathogenicity being correlated with each effector family's evolution, with significantly associated effectors highlighted.

BAYESTRAITS analysis using the core-genome phylogeny predicted the evolution of six T3E families was linked to cherry pathogenicity. These were hopBF, hopAB, hopH, hopAR, avrPto and hopBB. To account for any phylogenetic uncertainty, the program was also run 100 times on the full set of 100 bootstrapped trees generated by RAxML. The evolution of T3Es hopBF, hopAR and hopAB was always associated with pathogenicity for all 100 trees in all runs, indicating strong association. However, the T3E genes avrPto, hopBB and hopH were only significantly correlated for 88%, 77% and 62% of trees respectively, averaged across the different runs (Fig. S13). To determine how these genes had been gained or lost across the phylogeny, the program GLOOME was used (Cohen et al., 2010). Fig. 3(b) illustrates the predicted gain and loss of these T3Es on the branches leading to clades pathogenic to cherry. Those putatively associated with pathogenicity (high probability of gain in cherry-pathogenic clades) included hopAR1, hopBB1, hopBF1 and hopH1. The T3Es hopAB1 and avrPto1 were found to be lost from cherry pathogenic Psm R1, whilst the hopAB1 and hopAB3 alleles were pseudogenized in Psm R2 and P. syringae pv avii (Fig. 3b). All effector gain and loss events are presented in Fig. S14 and Table S12. Fig. S15 shows the phylogeny with branch labels used in GLOOME.

GLOOME predicted that key effectors have been gained in multiple clades. The hopAR1 gene has been gained in Psm R1, Psm R2, Pss and P. syringae pv avii. The T3E hopBB1 was present in the majority of strains within Psm R1, R2 and P. syringae pv avii but was absent from Pss strains. It showed high probability of gain on branches leading to both Psm R2 and P. syringae pv avii. However, GLOOME predicted loss in two Psm R1 strains, indicating that the gene may have experienced dynamic evolution in cherry pathogens. The hopBB1 effector is closely related to members of the hopF family and avrRpm2 (Lo et al., 2016). In addition to the significant acquisition of hopBB1 homologues, the hopF family was expanded in cherry pathogens. Pathogenic strains in Psm R1 and R2 all possessed two hopF alleles each (hopF3 and hopF4, and hopF2 and hopF4; see Fig. 2). P. syringae pv avii did not possess any hopF homologues, but had gained *hopBB1*. By contrast, *Pss* strains lacked all *hopF* members.

#### Origins of key effectors in cherry pathogens

To understand the origins of key effectors, gene phylogenies were produced. Incongruence with the core-genome phylogeny indicated that effector sequences had likely experienced horizontal



**Fig. 2** Virulence gene identification. (a) Heatmap of virulence gene presence and absence across *Pseudomonas syringae* (*avrA1-hopBJ1*). The dark green squares indicate presence of a full-length type III effector (T3E) homologue, whereas light green squares indicate that the gene is disrupted or truncated in some way. Strains isolated from cherry and plum are highlighted in pink and blue respectively. Asterisks indicate nonpathogenic to cherry in controlled pathogenicity tests. Strains with long-read sequenced genomes are in black boxes. The cherry-pathogenic clades are illustrated via horizontal shading of cells with *P. syringae* pv *morsprunorum* (*Psm*) R1 in blue, *Psm* R2 in light green, *P. syringae* pv *syringae* (*Pss*) in pink and *P. syringae* pv *avii* in orange. Strains are ordered based on the core-genome phylogenetic tree, which is represented by the dendrogram, with phylogroups labelled (P1–P3). (b) Continuation of (a) for T3Es *hopBK1*–*hopZ5* and additional non-T3 virulence factors, which are coloured in dark blue (all genes full-length and present) and light blue (not all genes present/truncation of genes). (c) The total number of full-length and pseudogenized T3E genes plotted for each strain, with cherry pathogenic strains in red and other strains in grey.

Table 3 Order of effectors in genomes sequenced using PacBio/MinION methods

	Contig	Length	Effector
R1-5244			
Chromosome	tig0	6109 228	hopAZ1, hopA2*, avrE1, hopM1*, hopAA1* <sup>†</sup> , hopZ4, hopAT1, hopQ1, hopD1, hopR1, hopF4, hopBL2, hopAV1, hopAO2*, hopAY1, hopF3, hopAS1, hopI1, hopAE1, hopAF1-2, hopAU1, hopAH1, hopV1, hopAR1, hopBK1*
Plasmid	tig3	168 854	hopAF1-1, hopBF1, avrD1, avrRpm2, hopBD1
Plasmid	tig4	81 536	hopA1
Plasmid	tig5	45 535	
Plasmid	tig6	40 810	_
R1-5300	0-		
Chromosome	tig0	5688 034	hopV1, hopAZ1, avrA1, hopQ1-2, hopA2*, <u>avrE1, hopM1*, hopAA1</u> <sup>†</sup> , <u>hopAB1, hopQ1, hopD1, hopR1,</u> hopAO2*, avrRpm2, avrPto1, hopAS1, <u>hopAT1*, hopBL2</u> *, hopI1, hopAE1, hopAF1-2, hopF3, hopAY1*, hopAU1, hopAH1
Chromosome	tig75	697 453	hopW1, hopBK1*, hopAR1
Plasmid	tig46	52 059	<u> - '                                  </u>
Plasmid	tig65	47 809	hopX1,hopBB1,hopG1
Plasmid	tig84	57 689	hopAO1*
Plasmid	tig113	102 557	avrD1
R2-leaf	<u> </u>		
Chromosome	tig0	6242 845	hopY1, hopAS1, hopAT1, hopH1, hopF4, hopW1, hopR1, hopAG1*, hopAH1-2, hopAl1, hopN1, hopAA1*, hopM1, avrE1 <sup>†</sup> , hopF2, hopE1, hopA2, hopAH1-1, hopAH1-1, hopAB3*, avrRps4, hopS2, hopI1, hopAR1
Plasmid	tig5	102 862	hopAO1*, hopAZ1, hopAY1
Plasmid	tig4	97 840	hopD1*, hopAU1
Plasmid	tig6	69 519	hop AF1-1, hop BF1
Plasmid	tig8	42 783	hopBB1, hopBD1
Plasmid	tig9	20 491	avrB2, hopX1
syr9097	O		<del></del>
Chromosome	tig0	5929 959	$\frac{hopAG1, hopAH1, hopA1I}{hopAH1, hopA2, avrE1, hopM1, hopAA1}^{\dagger}$

Effectors are listed in order of appearance on each assembly contig (labelled as chromosomal or plasmid). Where effectors could be considered as linked (within 10 kb of each other) they are underlined.

gene transfer (HGT) between the pathogenic clades, as their sequences clustered together. There has been possible effector exchange between Psm R1, R2 and P. syringae pv avii. To predict precisely where transfers had occurred on the phylogeny, the program RANGER-DTL was utilized (Bansal et al., 2012). Table 4 reports T3Es that exhibited evidence of HGT between cherry pathogens (gene trees are presented in Figs S16, S17). Full transfer events are listed in Table S13, and Fig. S18 shows the phylogeny with branch labels used in RANGER-DTL. The BAYESTRAITS pathogenicity-correlated T3Es, hopBB and hopBF, both showed evidence of HGT. Fig. 4(a) shows examples of T3Es putatively undergoing HGT between cherry pathogenic clades highlighted in red. Alignments of the flanking regions (Fig. 4b) showed homology between the cherry pathogens and included mobile elements likely involved in recombination events. Putatively transferred effectors were mostly plasmid encoded in the longread genomes (Table 3). In R1-5244, several of these genes were encoded on one plasmid (Contig 3), whilst in R2-leaf they were found on two plasmids (Contig 6 and 8).

The pathogenicity-associated T3E gene *hopAR1* was present in 23/28 cherry pathogens and showed probable gain in pathogenic clades. Phylogenetic analysis of this T3E (Fig. 5a) showed that the sequences for the different cherry pathogenic clades did not

cluster with each other, indicating convergent acquisition. Prophage identification (Table S14) did, however, reveal that this T3E is predicted to have been gained in Psm R1 and R2 within different phage sequences, whilst in Pss it is on a genomic island (Fig. 5b), and so has been acquired via distinct mechanisms. The Psm R1 phage is 51.5 kb, described as intact, and contains both hopAR1 and a truncated version of hopBK1. The Psm R2 phage sequence was 37.1 kb and was described as 'incomplete', indicating it did not have all the components of an active prophage. Further analysis of this region in Psm R2 and P2 strains revealed a shared adjacent tRNA-Thr gene (Fig. S19a,b). Within P2, although cherry Pss strains lacked the phage, several strains isolated from bean (syr2675, syr2676 and syr2682) possessed the hopAR1 gene within a phage homologous to that in Psm R2. The syr2675 hopAR1 sequence was also the most closely related homologue of Psm R2 hopAR1 (Fig. 5a). This evidence suggests that this effector gene may have been transferred via phage between phylogroups.

Many T3Es are mobilized between bacteria on GIs. GIs were identified for the three PacBio-sequenced strains of *Psm* R1, *Psm* R2 and *Pss* (Tables S15–S17). R1-5244 GIs contained the coronatine biosynthesis cluster and six T3Es. In R2-leaf, eight T3E genes were located on GIs, whilst in syr9097 three T3Es were

<sup>\*</sup>Effector gene is disrupted and is labelled as a pseudogene.

<sup>†</sup>Effectors within the conserved effector locus.

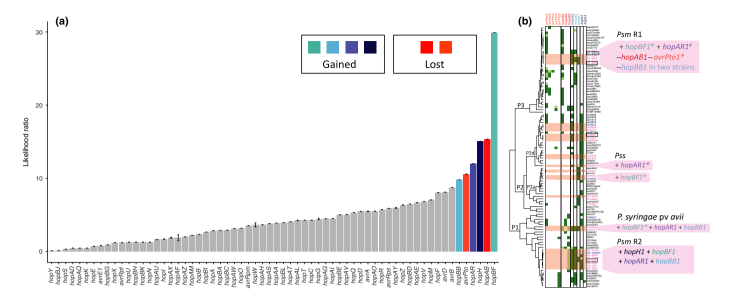


Fig. 3 Association of type III effector evolution with cherry pathogenicity. (a) Barplot showing the likelihood ratio for the correlation of each effector gene family with cherry pathogenicity based on BAYESTRAITS analysis using the core-genome phylogeny. The values are obtained from means of 100 independent runs of the program with error bars showing  $\pm$  SE above and below the mean. Those effectors that were not significantly associated with pathogenicity are coloured in grey. Coloured bars were associated with pathogenicity ( $P \le 0.05$  in > 90% of runs). Those genes that were hypothesized to be gained in cherry pathogens (from gain loss mapping engine ( $G_{LOOME}$ ) analysis) are coloured in shades of blue, whilst where the significant gene was absent in cherry pathogenic clades the bar is coloured in shades of red. (b) Gain and loss of BAYESTRAITS-associated T3Es in cherry-pathogenic clades on the core-genome phylogeny predicted using  $G_{LOOME}$  ( $P \ge 0.8$ ). The phylogeny and heatmap of these effector genes is presented (heatmap as in Fig. 2, effector gene names are colour coded based on the bar colours in (a) and cherry-pathogenic strains are highlighted by pink horizontal shading of columns). Phylogroups (P1–P3) are labelled. Strains with long-read sequenced genomes are in black boxes. \*The probability of this effector being gained/lost was slightly < 0.8, but exceeded 0.65 (see Supporting Information Table S12 for details). \*The hopAR1 gene has been gained on the branch leading to *P. syringae* pv morsprunorum (*Psm*) R1 (including pathogens and nonpathogens).

found on genomic islands. These GIs were then searched for in other *P. syringae* genomes to identify potential sources of transfer, and Fig. S20 shows heatmaps of GI presence. The *Psm* R1 GIs included several found only in pathogenic *Psm* R1 strains, differentiating them from the nonpathogens. These included the coronatine biosynthesis cluster (GI1), *hopF3* (GI6) and *hopAT1* (GI14). Most *Psm* R1 GIs produced hits across *P. syringae*, particularly in P1 and P3. *Psm* R2 GIs were most commonly found in P1. Several were shared with other cherry-pathogenic clades, including those containing *hopAF1* (GI36), *hopAT1* (GI3) and *hopD1* (GI6). Finally, although most islands identified in syr9097 were commonly found across the species complex, those containing T3Es (GI30, GI23 and GI26) appeared to be P2 specific, indicating that cherry-pathogenic strains likely gained these islands from other members of P2.

#### Functional analysis of potential avr genes

To validate predictions from genome analysis, cloning was used to identify avirulence factors active in cherry. The effector genes *avrPto* and *hopAB* were absent from cherry pathogens, and their evolution was theoretically linked to pathogenicity. Several other candidate avirulence effectors were identified that were absent from cherry pathogens but present in close out-groups (Fig. 6). Avirulence-gene identification focused on *Psm* R1, as any T3E variation within this clade may be due to differences in host specificity rather than phylogenetic distance. Potential avirulence

T3E genes included *avrA1*, *avrPto1*, *hopAA1*, *hopAB1*, *hopAO2* and *hopG1*, which had full-length homologues in nonpathogenic *Psm* R1 strains but were absent from or truncated in pathogens. These genes were cloned from the strain R1-5300 (except *hopAO2*, which was cloned from R1-9657).

The effector avrRps4 was also cloned from P. syringae pv avellanae (Psv) BPIC631, a close relative of Psm R2. This effector was absent from most cherry-pathogenic strains (Fig. 6). Several pathogens possessed the full-length gene (R2-leaf, R2-9095 and P. syringae pv avii), but lacked the KRVY domain that functions in planta (Fig. S21) (Sohn et al., 2009). The hopAW1 gene was cloned from Pph1448A as this T3E has undergone two independent mutations in Pss strains, disrupting the beginning of the gene (Fig. S22). Finally, hopC1 was cloned from the Aquilegia vulgaris pathogen RMA1, which is basal to the Psm R2 clade as it is absent from all cherry-pathogenic strains.

Nine effectors were cloned into pBBR1MCS-5 and conjugated into three pathogenic strains (R1-5244, R2-leaf and syr9644). The presence of the plasmids did not affect multiplication *in vitro*. Knock-out strains for the T3SS gene *hrpA* were obtained for R1-5244 and R2-leaf to act as nonpathogenic controls that could not secrete T3Es and failed to cause the HR on tobacco (Fig. S23).

Bacterial multiplication experiments were conducted in cherry leaves. The transconjugants expressing HopAB1 or HopC1 failed to multiply to the same levels as the pathogenic empty vector (EV) controls or produce disease lesions. The ectopic expression

Table 4 List of putative horizontal gene transfer events that have occurred between cherry-infecting clades within Pseudomonas syringae

Effector	Putative transfers	Region	Plasmid location	Predicted in RANGER-DTL
avrD1	R1/R2/P. syringae pv avii	Plasmid	tig3 (R1-5244)	Y
avrRps4*	R2/P. syringae pv avii	Next to cluster of mobile elements	_	Υ
hopAF1	R1/R2/P. syringae pv avii	Plasmid	tig3 (R1-5244), tig6 (R2)	Υ
hopAO1*	R1/R2/P. syringae pv avii	Plasmid	tig5(R2), tig84 (R1-5300)	N
hopAT1	R1/R2	Genomic island	_	N
hopAU1	R2/P. syringae pv avii	Plasmid	tig4 (R2)	Υ
hopAY1	R2/P. syringae pv avii	Plasmid	tig5 (R2)	Υ
hopBB1	R1/R2/P. syringae pv avii	Plasmid	tig8 (R2), tig65 (R1-5300)	Υ
hopBD1	R2/P. syringae pv avii	Plasmid	tig3 (R1-5244), tig8 (R2)	Υ
hopBF1	R1/R2/P. syringae pv avii	Plasmid	tig3 (R1-5244), tig6 (R2)	Υ
hopD1*	R2/P. syringae pv avii	Plasmid	tig4(R2)	N
hopO1	R2/P. syringae pv avii	Next to cluster of mobile elements (next to hopT1)	_	Υ
hopT1	R2/P. syringae pv avii	Next to cluster of mobile elements (next to hopO1)	-	Υ

Where the effector gene is present in the PacBio- or MinION-sequenced strains, its chromosomal or plasmid location is indicated. The type III effector genes *hopO1* and *hopT1* were not present in the PacBio-sequenced strains and therefore it is uncertain if they are on plasmids or chromosomal. \*Effector gene is disrupted in some strains and is labelled as a pseudogene.

of AvrA1, AvrRps4 and HopAW1 also caused significant reductions in growth, but this reduction was not consistently seen across all three pathogenic strains (Fig. 7a). As the addition of the *hopAB1* gene reduced pathogenicity, full-length *hopAB2* and *hopAB3* genes were also cloned from *Psv*BPIC631 and RMA1, and were also found to reduce pathogen multiplication (Fig. 7b).

To investigate the induction of the HR by the HopAB family and HopC1, inoculations were performed at high concentrations  $(2 \times 10^8 \text{ colony-forming units (CFU) ml}^{-1})$  as in Hulin *et al.* (2018). In *Psm* R1 and R2, the addition of these T3Es led to more rapid tissue collapse than observed in EV controls, indicative of HR induction (Fig. 7c,d); HopC1 and HopAB1 were particularly effective. With *Pss*, however, EV transconjugants themselves caused rapid tissue collapse, making it impossible to recognize an induced HR as symptom development was not significantly different.

The *hopAB1* gene is found in a mobile-element-rich c. 40 kb region in the nonpathogenic Psm R1-5300, missing from the pathogen Psm R1-5244 (Fig. 8a). Meanwhile, Psm R2 and P. syringae pv avii possessed putatively pseudogenized hopAB3 alleles (Fig. 8b), and P. syringae pv avii also possessed a truncated hopAB1 gene (Fig. S24). hopAB3 is truncated in Psm R2 due to a 2 bp insertion (GG at position 1404 bp) leading to a premature stop codon, whilst in P. syringae pv avii a 218 bp deletion has disrupted the C-terminus. If expressed, the E3-ubiquitin ligase domain is completely absent from the Psm R2 protein and disrupted in P. syringae pv avii (Fig. 8c). Both HopAB3 alleles were also divergent enough that the Pto-interacting domain (PID) was not identified by Interproscan. To determine if the truncated Psm R2 HopAB3 allele induced any resistance response in cherry leaves, the gene was expressed in Psm R1-5244 and population growth measured. The addition of this gene did not lead to a significant reduction in growth compared with the EV control, unlike other hopAB alleles (Fig. 8d), and the transconjugant was still able to induce disease symptoms 10 d post inoculation (dpi) (Fig. 8e).

Overall, the data supported the conclusion that expressing alleles of *hopAB* and *hopC* reduced bacterial multiplication in cherry

and were consistent with HR induction by *Psm* R1 and R2. However, it should be noted that any growth changes exhibited might have been influenced by aberrant transcription or translation of these effectors in the plant due to expression *in trans*.

#### **Discussion**

#### Core-genome phylogenetics

Phylogenetic analysis confirmed that cherry pathogenicity has evolved multiple times within *P. syringae*. *Psm* R1, R2 and *P. syringae* pv *avii* each formed distinct monophyletic clades, whereas cherry-pathogenic *Pss* strains were distributed across the P2 clade, indicating that cherry pathogenicity has either evolved multiple times within P2 or that this clade is not particularly specialized. To confirm this genomic prediction of pathogenicity, several additional P2 strains isolated from bean, pea and lilac were tested for pathogenicity in cherry. They each produced lower population levels in cherry leaves than cherry pathogens, suggesting that strains isolated from cherry and plum are more pathogenic to their hosts of origin (Fig. S10). Many P2 strains have previously been named *Pss* on the basis of lilac pathogenicity, despite being pathogenic to other plant species (Young, 1991). A new naming system within this phylogroup is desirable.

# Search for candidate effectors involved in cherry pathogenicity

Gains and losses of T3Es were closely associated with pathogenicity. Virulence-associated effectors *hopAR1*, *hopBB1*, *hopH1* and *hopBF1* had been gained in multiple cherry-pathogenic clades. The *hopAR1* effector has been studied in the bean pathogen *P. syringae* pv *phaseolicola* R3 (1302A), as a GI-located *avr* gene (*avrPphB*) whose protein is detected by the corresponding R3 resistance protein *in planta* (Pitman *et al.*, 2005; Neale *et al.*, 2016). HopAR1 also acts as a virulence factor as a cysteine protease which targets receptor-like kinases to interfere with plant

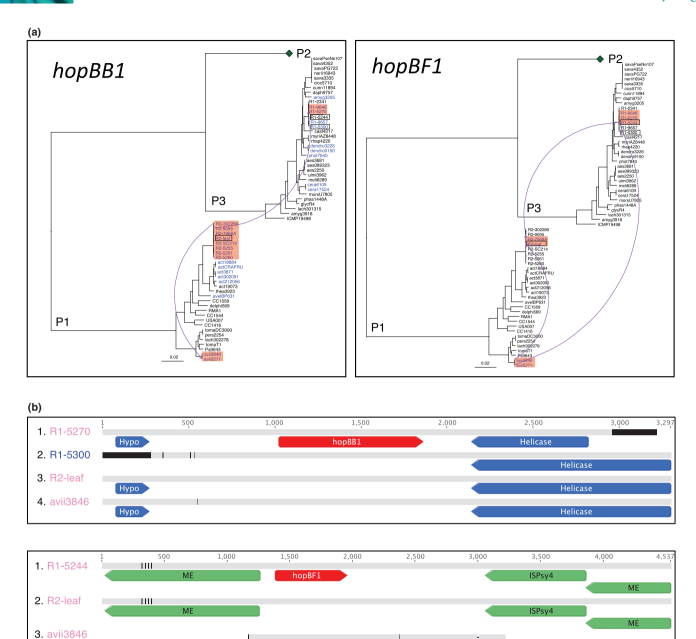
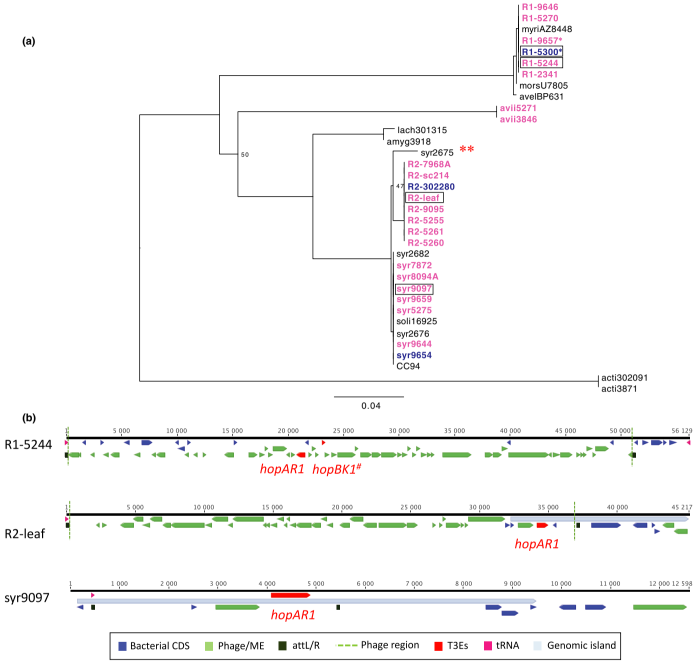


Fig. 4 Horizontal gene transfer has played a key role in the evolution of cherry pathogenicity. (a) Putative horizontal transfer of the hopBB1 and hopBF1 genes between different cherry-pathogenic clades based on the Pseudomonas syringae core-genome phylogeny. The different phylogroups are labelled (P1–P3), with P2 collapsed to concentrate on the other phylogroups. Strains that possess the effector gene are coloured in blue, and those that are cherry-pathogenic are highlighted in red. Strains with long-read sequenced genomes are in black boxes. The transfer events predicted by RANGER-DTL are shown by purple arrows. The bar shows substitutions per site. (b) DNA alignments of genomic regions containing these two effector genes, showing similar flanking regions between cherry pathogens. Alignments are colour coded based on similarity; identical residues are in grey, whereas dissimilar residues appear in black. The effector gene is coloured in red, mobile element genes are in green and other coding sequences are in blue. Cherry-isolated strains are named in pink, whilst the nonpathogenic plum strain R1-5300 is in blue. Gene name abbreviations: Hypo, hypothetical protein gene; ISPsy4, insertion sequence; ME, mobile element.

PAMP-triggered immunity (PTI) responses (Zhang *et al.*, 2010). This effector could play a similar role in PTI suppression in cherry.

HopBB1 and other members of the HopF family were abundant in cherry pathogens. All HopF members share an N-terminus and myristoylation sites for plant cell membrane

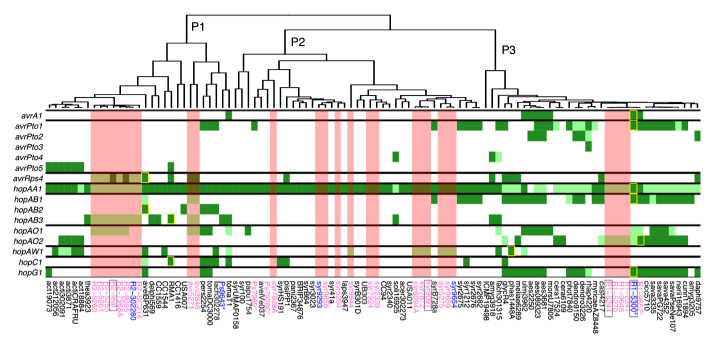
localization (Lo et al., 2016) and interfere with PTI and ETI in model plants (Wang et al., 2010; Wu et al., 2011; Hurley et al., 2014). The presence of multiple hopF homologues in cherry pathogens and specific gain of hopBB1 suggested the importance of their function. In comparison, HopH1 and HopBF1 are understudied. HopH1 is a protease, homologous to the Ralstonia



**Fig. 5** Evolution of *hopAR1* in different clades of *Pseudomonas syringae* containing cherry pathogens. (a) Maximum likelihood phylogenetic tree built using the nucleotide sequences of the *hopAR1* gene. Cherry and plum isolated strains are highlighted in pink and blue respectively; those names followed by single asterisks were nonpathogenic on cherry in controlled pathogenicity tests. Strains with long-read sequenced genomes are in black boxes. Bootstrap supports < 99% are shown. The bar is nucleotide substitutions per site. Double asterisks point to the clustering of *P. syringae* pv *morsprunorum* (*Psm*) R2 sequences with syr2675. (b) Genomic locations of the *hopAR1* gene in the three PacBio-sequenced cherry pathogens. The gene is located within prophage sequences in *Psm* R1 and R2 (see Table S14 for details), whereas in syr9097 it is on a genomic island (GI) adjacent to a transfer RNA (tRNA) gene. Effector genes are coloured in red, other coding sequences in blue, phage genes predicted by Phaster and mobile element genes are in green, tRNA genes in pink and GIs predicted (GI14 in *Psm* R2 and GI23 in *P. syringae* pv *syringae* (*Pss*)) in light blue. Predicted phage *att* sites are in dark green, with sites homologous to R2-leaf in *Pss* 9097 also indicated even though a phage is not predicted here. The ends of predicted prophage sequences are denoted with dashed green lines. #hopBK1 is a pseudogene in this strain. CDS, coding sequence; ME, mobile element; T3E, type III effector.

solanacearum Rip36 protein (Nahar et al., 2014). This T3E gene was found on GI37 in Psm R2-leaf and was within 3 kb of hopF4 (Fig. S25), indicating that these two T3Es may have been gained together. HopBF1 was first discovered in P. syringae pv aptata

and *oryzae* (Baltrus *et al.*, 2011), but its role is undetermined. This study therefore identified candidate T3Es important for cherry pathogenicity that should be the focus of future functional studies.



**Fig. 6** Distribution of putative avirulent type III effector (T3E) genes across the *Pseudomonas syringae* phylogeny. The heatmap shows the presence and absence of T3Es studied in the functional analysis. This was constructed as in Fig. 2. The cherry-pathogenic clades are illustrated via vertical shading of cells in pink. Full-length putative *avr* genes used in cloning work are outlined with yellow boxes.

Phytotoxin biosynthesis gene clusters were also identified. Coronatine is present on a plasmid in pathogenic *Psm* R1 and may be one of the factors that differentiate pathogens from non-pathogens in this clade. Coronatine functions in virulence by downregulating salicylic acid defence signalling (Grant & Jones, 2009). Necrosis-inducing lipodepsipeptide toxins were common in P2. All cherry-pathogenic *Pss* strains possessed at least one biosynthesis cluster. The ability of *Pss* strains to cause necrosis on cherry fruits has been linked to toxins (Scholz-Schroeder *et al.*, 2001). Interestingly, two nonpathogenic P2b cherry strains lacked all phytotoxins, a deficiency that probably contributes to their lack of pathogenicity.

All cherry-pathogenic *Pss* strains had reduced effector repertoires. This observation supports the hypothesis that a phenotypic trade-off exists, with strains retaining few T3Es, whilst relying more on phytotoxins for pathogenicity (Baltrus *et al.*, 2011; Hockett *et al.*, 2014). If this pathogenic strategy has evolved in the P2 clade, it raises the question as to how it affects host specificity and virulence. P2 strains often infect more than one host species (Rezaei & Taghavi, 2014). These strains probably possess fewer ETI-inducing avirulence factors that restrict effector-rich strains to particular hosts, so may be more successful generalists. The reduction in T3E repertoire, however, may be limiting, as strains may be less capable of the long-term disease suppression required at the start of a hemi-biotrophic interaction.

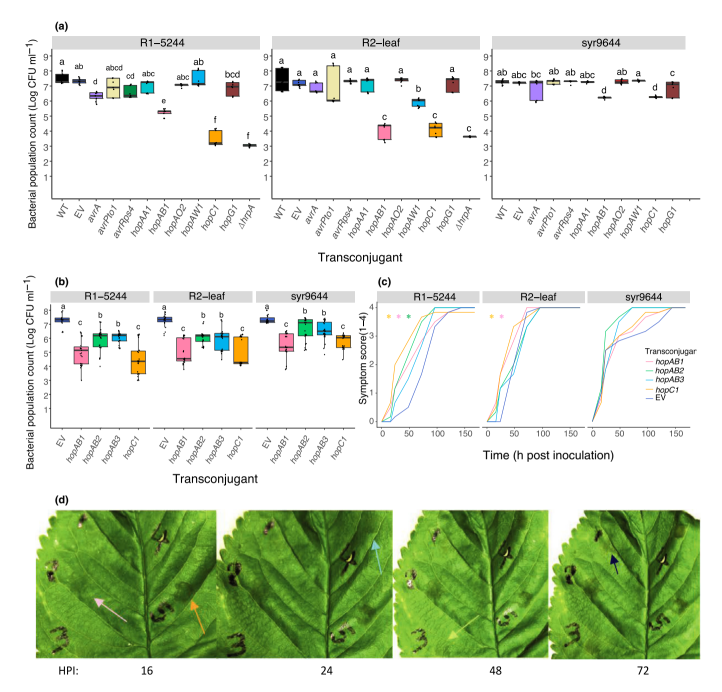
Most cherry-pathogenic clades possessed genes involved in aromatic compound degradation, shown to be important in virulence on olive (Caballo-Ponce *et al.*, 2016), and ice nucleation genes that stimulate frost damage (Lamichhane *et al.*, 2014). The fact that not all cherry-pathogenic clades possessed these genes

suggests they are not essential requirements for bacterial canker; however, they may contribute to niche persistence. For example, Crosse & Garrett (1966) observed that *Psm* R1 survived in cankers for longer than *Pss.* Increased persistence might be linked to genes involved in woody-tissue adaptation.

## Horizontal gene transfer has been important in the acquisition of key effectors

HGT is important for effector shuffling within *P. syringae* (Arnold & Jackson, 2011). Pathogenicity-associated T3Es *hopBB1* and *hopBF1* were plasmid encoded and showed evidence of HGT between the cherry-pathogenic clades in P1 and P3. Plasmid profiling revealed that cherry pathogens in these phylogroups possessed native plasmids, some of which were putatively conjugative, indicating the importance of plasmids in gene exchange. By contrast, most cherry-pathogenic *Pss* strains lacked plasmids.

The T3E *hopAR1* was chromosomal in all long-read sequenced genomes. This gene was found within distinct prophage sequences in *Psm* R1 and R2. To our knowledge this is the first reported example of a plant pathogen T3E located within a prophage sequence. Interestingly, the *Psm* R2 *hopAR1* gene homologue was most similar to *hopAR1* from a P2 bean strain syr2675, which is a close relative of cherry *Pss.* This strain possessed a homologous phage to *Psm* R2, indicating that HGT of this T3E between phylogroups may have been phage mediated. This striking example of convergent acquisition of *hopAR1* in the cherry pathogens, putatively through distinct prophages in *Psm* R1 and R2, and a GI in *Pss* indicates that this T3E may have important roles in virulence. The well-characterized



**Fig. 7** Identification of avirulence factors activating effector-triggered immunity in cherry. (a) Boxplot of an initial 10-d population count analysis of cherry pathogens (R1-5244, R2-leaf and syr9644) transconjugants expressing candidate avirulence genes. The data presented are based on one experiment, with three leaf replicates and three nested technical replicates (n = 9). Boxplots show median and interquartile range (IQR) and whiskers extend to values 1.5× IQR above and below the median. All data points are plotted with circles. Controls included the wild-type strain, a strain containing the empty pBBR1MCS-5 vector and a Δ*hrpA* deletion mutant (for R1-5244 and R2-leaf). A separate ANOVA was performed for each cherry pathogen (R1-5244, R2-leaf and syr9644) and the Tukey-HSD significance groups (P = 0.05; confidence level: 0.95) for each strain are presented above each boxplot. (b) Boxplot of 10-d population counts of cherry pathogens (R1-5244, R2-leaf and syr9644) expressing different HopAB alleles and HopC1. The data presented are based on three independent experiments (n = 27). Tukey-HSD significance groups are presented above each boxplot. (c) Symptom development of R1-5244, R2-leaf, syr9644 transconjugants. Mean symptom score values are presented and represent two independent experiments (n = 6). Symptoms assessed as degree of browning of the infiltration site: 1, limited browning; 2, < 50%; 3, > 50%; 4, 100% of the infiltrated area brown. Analysis was based on area under disease progress curve (AUDPC) values (0-48 h). An ANOVA was performed on AUDPC values, with asterisks indicating significantly different from the empty vector (EV) control. (d) Symptom development over time on a representative leaf inoculated with R1-5244 transconjugants. HPI, hours post inoculation. The order of strains: 1, EV; 2, hopAB1; 3, hopAB2; 4, hopAB3; 5, hopC1. Arrows indicate the first appearance of symptoms associated with each strain and are coloured based on the graph in (c). ANOVA tables for all statistical

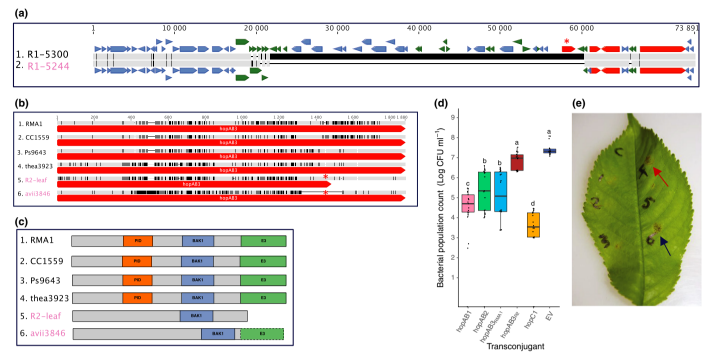


Fig. 8 hopAB alleles have been both lost and truncated in cherry pathogens. (a) Alignment of the DNA region surrounding hopAB1 in Pseudomonas syringae pv morsprunorum (Psm) R1 strains. Grey indicates sequence identity, whereas black indicates divergence. The effector genes are coloured in red, whereas other coding sequences are in blue and mobile element genes are in green. Asterisk indicates the location of hopAB1 in R1-5300, whilst the upstream effectors are hopQ1, hopD1 and hopR1. (b) DNA alignment of the hopAB3 gene of Psm R2 and close out-groups. Asterisks indicate where the hopAB3 gene has been truncated due to a GG insertion at 1404 bp leading to a frameshift in Psm R2, whilst in P. syringae pv avii (avii3846) there is a deletion at the end of the gene. (c) Diagrams showing the location of key domains in the HopAB3 protein including the Pto-interaction domain (PID), BAK1-interacting domain (BAK1) and E3 ubiquitin ligase (E3). The E3 domain is lost completely from the Psm R2 protein, whereas in avii3846 the beginning of this domain is lost. The PID domain was not detected in the cherry pathogen sequences. (d) Boxplot of 10-d population counts of R1-5244 transconjugants expressing three different full-length hopAB alleles, truncated hopAB3<sub>R2-leaf</sub> and hopC1. The boxplots were constructed as in Fig. 7. The data presented are based on two independent experiments (n = 18). Tukey-HSD significance groups (P = 0.05; confidence level: 0.95) are presented above each boxplot (full statistical analysis is in Table S26). (e) Representative image of symptoms 10 d post inoculation (dpi) with the different R1-5244 transconjugants when inoculated at a low concentration ( $2 \times 10^6$  colony-forming units (CFU) ml<sup>-1</sup>) to observe pathogenicity. Arrows point to pathogenic symptoms in the strain expressing hopAB3<sub>R2-leaf</sub> and the empty vector (EV) strain, colour coded as in (d).

*P. syringae* pv *phaseolicola* R3 homologue is not associated with a phage, but has been shown to undergo dynamic evolution on a mobile genomic island *in planta* in resistant bean cultivars (Neale *et al.*, 2016).

Several T3Es in *Psm* R1, R2 and *Pss* were located on GIs. To determine the likely source of GIs in cherry strains, all other *P. syringae* strains were searched for homologous sequences. There was evidence of *Psm* R1 and R2 islands being shared between cherry pathogen clades indicative of HGT events occurring between strains occupying the same ecological niche.

## Functional genomics revealed convergent loss of an *avr* factor

Genes from the *hopAB* and *avrPto* families form a redundant effector group (REG) vital for early PTI suppression in herbaceous species (Jackson *et al.*, 1999; Lin & Martin, 2005; Kvitko *et al.*, 2009). Both effectors also trigger ETI by interacting with the serine-threonine kinase R protein Pto in tomato (Kim *et al.*, 2002).

Across the *P. syringae* complex, the REG was common (Fig. S26), but cherry pathogens all lacked full-length genes. The

hopAB1 gene has been lost from Psm R1, whilst the Psm R2 and P. syringae pv avii predicted HopAB3 proteins lacked the PID and E3-ubiquitin ligase domains through contrasting mutations. P. syringae pv avii also possessed a truncated hopAB1 gene (Fig. S24), lacking the PID domain. The lack of a PID in cherry pathogen HopAB proteins suggested that they could have diverged to avoid a Pto-based recognition system in cherry.

Full-length members of this REG were expressed in cherry pathogens to determine their role *in planta*. The addition of HopAB alleles (HopAB1–3) consistently reduced population growth of pathogenic strains *in planta* and triggered a response consistent with the HR. If this effector does trigger immunity in cherry, there may have been selection pressure for its loss or pseudogenization in cherry pathogens in order to reduce avirulence activity. The truncated version of HopAB3 in R2-leaf was found not to exhibit avirulence activity as its expression did not reduce the growth of R1-5244 *in planta*. Although AvrPto is part of the same REG, its expression had no effect on the ability of cherry pathogens to multiply in leaves. The absence of AvrPto from cherry pathogens is therefore unlikely to be driven by avirulence, but could be due to the lack of HopAB virulence targets *in planta*.

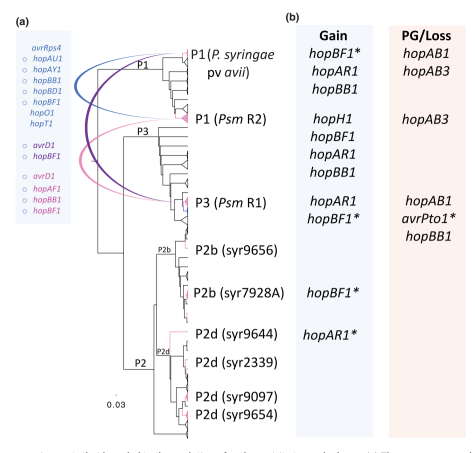


Fig. 9 Model highlighting genomic events that have led to the evolution of pathogenicity towards cherry. (a) The core-genome phylogeny is presented with phylogroups (P1–P3) labelled. Bar shows substitutions per site. For visualization, clades within the phylogenetic tree have been collapsed with clades containing cherry pathogens in pink (*Pseudomonas syringae* pv *morsprunorum* (*Psm*) R1 nonpathogenic strains in blue). Examples of cherry pathogens within each clade of P2 are named. Horizontal gene transfer (HGT) events predicted using RANGER-DTL are shown, with open circles representing those upon plasmids in long-read sequenced strains. (b) The key gains and losses of associated virulence genes in strains pathogenic to cherry are described based on gain loss mapping engine (Gloome) analysis. Asterisks indicate the probability of this effector being gained/lost predicted using Gloome was slightly lower than the threshold of 0.8, but exceeded 0.65 (see Table S12 for details).

As this REG is vital for early disease suppression in model strains, cherry pathogens must rely on other T3Es to fulfil this role.

The variation in *hopAB1* presence in *Psm* R1 is intriguing. *Psm* R1 strains may be pathogenic on both cherry and plum  $(\Delta hopABI)$  or just pathogenic on plum (possessed hopABI) as recorded in Hulin et al. (2018). This suggests that the host proteins in cherry that detect the presence of HopAB are not present/functioning in plum. Future studies may determine how the two host immune responses diverged and could examine hopAB diversity across *Prunus* pathogens. This study focused on bacterial canker of P. avium; however, strains isolated from additional Prunus spp. that cause other diseases were included, such as P. syringae pv cerasicola (bacterial gall of hybrid cherry Prunus × yedoensis; Kamiunten et al., 2000), P. syringae pv morsprunorum FTRSU7805 (canker of apricot), P. syringae pv amygdali (canker of almond) and P. syringae pv persicae (decline and canker of peach) (Table 1). All apart from P. syringae pv amygdali 3205 and P. syringae pv persicae lacked HopAB (Fig. 2), indicating that there may be a conserved resistance mechanism regulating ETI activated by this effector family in *Prunus* species.

#### Linking genomics to the evolution of cherry pathogenicity

Cherry pathogenicity has arisen independently within *P. syringae*, with strains using both shared and distinctive virulence strategies. Cherry-pathogenic clades in P1 and P3 have large effector repertoires. Cherry Pss were found across P2 with reduced T3Es and several phytotoxin gene clusters. Key events in the evolution of cherry pathogenicity (Fig. 9) appear to be the acquisition of virulence-associated effectors, often through HGT. Putatively important T3Es included *hopAR1*, members of the *hopF* family such as hopBB1 and the other T3Es hopBF1 and hopH1. Significantly, the loss/pseudogenization of HopAB effectors has also occurred in multiple clades. Within P2, the different cherry-infecting Pss clades have slight differences in their virulence factor repertoires that may reflect their convergent gain of pathogenicity. Clades differed in T3E content, phytotoxin genes and possession of genes for catechol degradation (Fig. 2), and thus pathogenicity was achieved with variable virulence factor repertoires. This study demonstrates that populations genomics can be used to examine a complex disease of a perennial plant species. A huge dataset was

narrowed down to several candidate host-specificity-associated genes, two of which (*hopAB* and *hopCI*) encode proteins that had putative avirulence functions *in planta*.

#### **Acknowledgements**

We acknowledge the East Malling Trust, University of Reading and BBSRC for funding (BB/P006272/1). Library preparation for PacBio sequencing was performed at the Earlham Institute, whilst for wild cherry strains Illumina MiSeq sequencing, libraries were prepared at the Genomics Facility, University of Warwick. We thank Steve Roberts, Helen Neale, Mateo San José and David Guttman for providing bacterial strains. We thank the East Malling Farm and Glass staff for plant maintenance. The authors declare no conflict of interest.

#### **Author contributions**

M.T.H., J.W.M., R.W.J. and R.J.H. conceived and designed the study as well as writing the manuscript. M.T.H. performed bioinformatics, statistical analysis and laboratory work. J.G.V. isolated some of the strains used in this study, J.G.V. and E.B.H. selected representative strains and prepared DNA for MiSeq sequencing of nine strains and L.B. assembled these nine sequences. H.J.B. performed the MinION library preparation and sequencing. A.D.A. assisted in bioinformatics pipeline development. All authors read and reviewed the final manuscript.

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## **Supporting Information**

Additional Supporting Information may be found online in the Supporting Information tab for this article:

- **Fig. S1** Plasmid profiles of all *Prunus*-infecting sequenced strains and some out-groups for comparison.
- **Fig. S2** Whole-genome alignment of R1-5244 and R1-5300 using ProgressiveMauve.
- Fig. S3 Initial core genome phylogeny of 108 *Pseudomonas syringae* strains.
- **Fig. S4** Core genome phylogenetics of phylogroup 3 with removal of different out-group strains.
- **Fig. S5** Core genome phylogenetics of phylogroup 2 with removal of different out-group strains.

- **Fig. S6** Core genome phylogenetic tree with removal of the outgroup strain *Pseudomonas syringae* pv *daphniphylli* 569.
- **Fig. S7** Core genome phylogenetic tree with removal of the outgroup strains *Pseudomonas syringae* pv *daphniphylli* 569 and *P. syringae* pv *syringae* 1212.
- **Fig. S8** Core genome phylogenetic tree with removal of the outgroup strain *Pseudomonas syringae* pv *eriobotryae* 4455.
- **Fig. S9** Core genome phylogenetic tree with removal of the outgroup strains *Pseudomonas syringae* pv *eriobotryae* 4455 and *P. syringae* pv *syringae* 1212.
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- **Fig. S11** Organisation of the structural T3SS and conserved effector locus (CEL) of the three PacBio-sequenced pathogens (R2-leaf, syr9097 and R1-5244).
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- **Fig. S14** Predicted gain and loss of effector genes occurring on each branch leading to cherry pathogenic clades on the core genome phylogenetic tree.
- **Fig. S15** Core genome phylogenetic tree used in GLOOME gain and loss analysis with the branch labels added.
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- Methods S1 Detailed descriptions of methodology used.
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