

Identification of virulence factors and type III effectors of phylotype I, Indian *Ralstonia solanacearum* strains Rs-09-161 and Rs-10-244

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Abstract. *Ralstonia solanacearum* is a well-known phytopathogen causing bacterial wilt in a large number of agriculturally important crops. The pathogenicity of *R. solanacearum* is expressed due to the presence of various virulence factors and effector proteins. In this study, various virulence factors and type III effector proteins of *R. solanacearum* that are present in the strains Rs-09-161 and Rs-10-244 were identified through bioinformatics approach and compared with other reference strains. *R. solanacearum* strains, Rs-09-161 and Rs-10-244 belong to the phylotype I, biovar3, and are the only sequenced strains from India infecting solanaceous vegetables. Similarity matrix obtained by comparing the sequences of virulence genes of Rs-09-161 and Rs-10-244 with other reference strains indicated that Rs-09-161 and Rs-10-244 share more than 99% similarity between them and are closely related to GMI1000. The virulence factors in *R. solanacearum* appear to be highly conserved in the *R. solanacearum* species complex. Rs-09-161 has 72 type III effectors whereas Rs-10-244 has 77. Comparison of the complete genes of type III effectors of Rs-09-161, Rs-10-244 has four unique effectors. Phylogenetic trees of RipA, RipG, RipH and RipS effector sequences resulted in the grouping of the isolates based on their phylotypes. Group 1 consists of strains that belong to phylotype I including Rs-09-161 and Rs-10-244. Phylotype III strain CMR15 forms a group closely associated with phylotype I. The strains belonging to phylotypes II and IV have separated to form two different groups.

Keywords. bacterial wilt; type III effector; virulence factor; phylogenetic analysis; Ralstonia solanacearum.

Introduction

Ralstonia solanacearum is a phytopathogen, which causes bacterial wilt disease in many crop plants and is responsible for huge losses in agriculturally important crops (Genin and Denny 2012). It has been ranked second in the list of top 10 of the most studied bacterial plant pathogens (Mansfield *et al.* 2012). Due to the extensive diversity that exists among the strains, the organism is now referred as *R. solanacearum* species complex (RSSC) and is divided into four phylotypes (Fegan and Prior 2005). These phylotypes represent their geographical origin: phylotype I (Asia), phylotype II (America), phylotype III (Africa) and phylotype IV (Indonesia). The phylotype IV also includes *R. syzygii* and the banana blood disease bacterium (BDB) (Genin and Denny 2012).

R. solanacearum finds its way into the plant through wounds in the roots and initiates wilting by impairing transport of water in the xylem that ultimately leads to the death of the infected plant (Genin and Denny 2012). Exopolysaccharide (EPS) produced by the bacterium is the primary virulence factor and impairs water transport within its susceptible host (Schell 2000). In addition to EPS, the type II secretory system (T2SS), chemotaxis, swimming, twitching motility and type III secretory system (TTSS) also contribute towards the virulence of the bacterium (Saile *et al.* 1997). The T2SS secretes various plant cell wall degrading enzymes

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(PCWDE) that include cellulolytic and pectinolytic enzymes which promote the colonization of the bacterium in the plant tissue. Chemotaxis and twitching motility also act as important virulence factors and aid in locating and attaching the bacterium to the host roots (Alvarez *et al.* 2010; Genin and Denny 2012).

The TTSS is an essential pathogenicity determinant and is encoded by the hypersensitive response and pathogenicity (hrp) regulon in R. solanacearum (Boucher et al. 1987). The hrp regulon is so named because it induces a hypersensitive response (HR) in nonhost or resistant plants and pathogenicity in the susceptible plants (Hueck 1998). A defect in the T3SS results in the loss of the ability to induce both; a hypersensitive response and pathogenicity in plants (Alfano and Collmer 2004). Type III-dependent protein secretion was first identified in the animal pathogen Yersinia enterocolitica (Heesemann et al. 1984) and later was found to be present in a variety of gram-negative phytopathogenic bacteria (Hueck 1998). The T3SS enables a bacterium to translocate pathogenicity proteins called as 'type III effectors (T3E)' into the cytosol of eukaryotic host cells. The effectors act as toxins and target the host immune system. This translocation is brought about by hrp dependant filamentous structure called as hrp pili (Van Gijsegem et al. 2000, 2002). In situ immunogold labelling experiments suggest that the hrp pili acts as a needle to provide a protein transport channel for transport of effector proteins into the host cytosol. All the hrp mutants that lack the hrp pilus protein (Hrp Y) cannot secrete hrp substrate proteins like hairpins and effectors (Van Gijsegem et al. 2002).

The expression of *R. solanacearum* T3SS is induced in the presence of poor nutritional conditions which mimics that of intracellular spaces in plants (Genin *et al.* 1992). The T3SS is an emerging area of study among many molecular biologists with plant and animal pathogens like *Pseudomonas, Xanthomonas, Salmonella* etc. Many T3Es are validated and many are under process in *R. solanacearum* through translocation studies using reporter-based systems like the Cya reporter and HA reporter systems (Cunnac *et al.* 2004; Mukaihara and Tamura 2009; Mukaihara *et al.* 2010; Sole *et al.* 2012).

In India, *R. solanacearum* has been isolated from various agriculturally important crops like eggplant, chilli, ginger, tomato, potato, capsicum etc. (Kumar *et al.* 2004; Chandrashekara *et al.* 2012; Ramesh and Phadke 2012). Even though the bacterial wilt is a severe issue and affects various crops in India, there are only two strains (Rs-09-161 and Rs-10-244) infecting solanaceous vegetables sequenced from India. These strains belong to race 1, biovar 3, phylotype I and based on endoglucanase (*egl*) gene sequence analysis the isolates belong to two different representative subgroups (Ramesh *et al.* 2014a,b). Both the strains are highly pathogenic on tomato and eggplant, cause 100% wilt within 15 days after inoculation (R. Ramesh, G. Achari, S. Gaitonde and T. Asolkar Bacterial wilt in solanaceous vegetables, *unpublished data*).

In RSSC, 16 strains are sequenced from different phylotypes (Salanoubat *et al.* 2002; Gabriel *et al.* 2006; Remenant *et al.* 2010, 2011; Xu *et al.* 2011; Ramesh *et al.* 2014a). Data on T3E of phylotype I strains is majorly contributed by studies on GMI1000 and no information on Indian strains is available. This study aims to identify and analyse various virulence factors and T3Es of *R. solanacearum* strains Rs-09-161 and Rs-10-244 using bioinformatics approach.

Materials and methods

R. solanacearum strains

R. solanacearum strains Rs-09-161 and Rs-10-244 were selected to analyse the virulence factors and T3Es in this study. These are the whole genome sequenced strains and belong to phylotype I from India (Ramesh *et al.* 2014a) and are being maintained in the culture collection of Plant Pathology Lab, ICAR-CCARI, Goa. Annotation of various virulence genes of *R. solanacearum* Rs-09-161 and Rs-10-244 was carried out using Eugene-P, with GMI1000 (phylotype I) as standard reference strain. The general features of all *R. solanacearum* strains used in this study are provided in the table 1.

Analysis of virulence factors

Various virulence genes involved in the colonization and wilting of the host were identified in the strains Rs-09-161 and Rs-10-244 based on the annotation data. These include the genes coding for EPS (epsA, epsB, epsC, epsD, epsE, epsF, epsP and epsR), PCWDE (PehA, PehB, PehC, Pme, Egl and CbhA), chemotaxis (CheA and CheW), swimming motility (FliC and FlgM) and twitching motility (PilA and PilP). The coding sequences of these virulence genes were compared with representative strains of R. solanacearum from different phylotypes, namely GMI 1000 (phylotype I), CFBP2957 (phylotype IIA), Po82 (phylotype IIB), CMR15 (phylotype PIII) and Psi07 (phylotype IV). The nucleotide sequences for virulence factors of Rs-09-161 and Rs-10-244 were retrieved from annotated files and of the reference strains were extracted from NCBI database (http://www.ncbi.nlm.nih.gov). Virulence sequences were submitted to GenBank and accession numbers were obtained (for details see table 1 in electronic supplementary material at http://www.ias.ac.in/jgenet/). The sequences were aligned pairwise using Clustal W (Thompson et al. 1994) and the evolutionary similarity matrix was constructed using MEGA ver. 6 software with p-distance method and bootstrap value of 1000 (Tamura et al. 2013).

Strain	Phylotype	Geographical origin	Isolated from	Genome size (Mb)	Total no. of T3Es	Hypothetical effectors	Accession	Remarks/purpose in this study
Rs-09-161	-	India	Egenlant	5.65	72	6	PR.INA217471	Test strain
Rs-10-244	(India	Chili	5.66	<u></u>		PRJNA236788	Test strain
GM11000		French Guyana	Tomato	5.811	71	1	PRJNA13	Virulence factors, T3E
RS1000		Japan	Tomato	NA	65	0		T3E
Y45		China	Tobacco	5.712	50	ŝ	PRJNA182081	T3E
FQY_4		China	Bacterial Wilt Nursery	5.805	52	7	PRJNA182081	T3E
CFBP2957	IIA	French West Indies	Tomato	5.683	72	5	PRJEA50685	Virulence factors, T3E
IPO1609	IIB	Netherlands	Potato	5.313	09	5	PRJNA32087	T3E
UW551		Kenya	Geranium	5.895	58	ŝ	PRJNA15601	T3E
Molk2		Philippines	Banana	5.961	76	8	PRJNA32085	T3E
Po82		Mexico	Potato	5.43	75	4	PRJNA66837	Virulence factors, T3E
CMR15	III	Cameroon	Tomato	5.593	68	5	PRJEA50681	Virulence factors, T3E
Psi07	IV	Indonesia	Tomato	5.606	72	7	PRJEA50683	Virulence factors, T3E
BDB R229		Indonesia	Banana	5.159	57	4	PRJNA53877	T3E
R. syzygii R24		Indonesia	Clove	5.424	48	2	PRJNA53879	T3E
Information was	compiled bas	sed on the published lite	Information was compiled based on the published literature and NCBI database.					

Table 1. R. solanacearum strains used in this study

Identification of T3E

The preliminary identification of T3E genes was carried out by screening the presence of *hrpII* box element (TTCGn16TTCG) in the region 500-bp upstream of the start codon using PatScan where only one mismatch was allowed. The presence of T3SS dependent export pattern in the T3E genes was detected by analysis of 50 amino acid N-terminal domain. The T3E was considered positive for N-terminal domain if it fulfilled at least two of the three criteria mentioned below: (i) serine + proline content should be greater than 30% (ii) leucine content should be lesser than 10% (iii) acidic residues should be absent within the first 12 amino acids.

Prediction of the start codon of the gene was carried out by the multiple sequence alignment of the region located downstream of the *hrpII* box element. The more distal 5' initiator codon conserved among different strain sequences was considered as the start codon. The predicted T3E genes were also analysed for frame-shift mutations and pseudogenes. T3E genes that had open reading frames disrupted by the insertion of IS element, altered structure (< 50%) of the gene or evidence that the T3E gene product is not translocated by the T3SS were considered as pseudogenes (Peeters *et al.* 2013). The identification of candidate T3Es in the genomes of Rs-09-161 and Rs-10-244 was carried out using 'Scan Your Genome' (Peeters *et al.* 2013).

Analysis of the T3Es

The phylogenetic analysis based on the gene families of T3Es of *R. solanacearum* was studied. The gene families analysed include RipA (*AWR* family), RipG (*GALA* family), RipH (*HLK* family) and RipS (*SKWP* family). The coding sequences of effectors belonging to each gene family were arranged in concatenated manner and compared with other strains. Reference strains used to study the phylogenetic relation of T3E are indicated in table 1. Phylogenetic analysis was performed in MEGA ver. 6.0 (Tamura *et al.* 2013) by using neighbour-joining (NJ) and the algorithm of Jukes and Cantor (1969) with 1000 bootstrap resamplings.

Results and discussion

With the availability of genomic data through whole genome sequencing, it has become interesting to study *R. solanacearum* at the genomic level. We therefore have studied virulence associated genes of Indian isolates Rs-09-161 and Rs-10-244 and compared them with the isolates available globally. This involves identifying the virulence factors and T3Es present in Indian strains and analysing the coding sequences of these genes for generating similarity matrix and phylogenetic trees. The basic architecture of the strains Rs-09-161 and Rs-10-244 and the sizes of the two

replicons; the chromosome and the megaplasmid are similar to that of the previously sequenced strains (Salanoubat *et al.* 2002; Remenant *et al.* 2010). The details are provided in table 2 of electronic supplementary material.

Analysis of virulence factors

The virulence genes in the test strains were identified using GMI1000 (phylotype I) as standard reference strain. The details of virulence genes of *R. solanacearum* strain Rs-09-161 and Rs-10-244, their location in the genome and the probable function are provided in table 1 in electronic supplementary material. The nomenclature of virulence genes is with the prefix RALSO161 and RALSO244 for Rs-09-161 and Rs-10-244, respectively. A similarity matrix obtained by comparing the sequences of virulence genes of Rs-09-161 and Rs-10-244 with other reference strains is given in table 3 in electronic supplementary material. The sequences of Rs-09-161 and Rs-10-244 share more than 99% similarity between them and are closely related to GMI1000.

The main virulence factor of *R. solanacearum*, the EPS is secreted by seven genes, namely *epsA*, *epsB*, *epsC*, *epsD*, *epsF*, *epsP* and *epsR*. The gene *epsD* is absent in phylotype I strains and all the EPS contributing genes are located on the megaplasmid. The sequences of CMR15 (phylotype III) and Psi07 (phylotype IV) share more than 90% similarity with Rs-09-161 and Rs-10-244. Both Rs-09-161 and Rs-10-244, have the presence of all six PCWDE genes. The sequences of PCWDE genes of Rs-09-161 and Rs-10-244 share 99% similarity with that of GMI1000. *PehB* is the only gene present on the chromosome and this observation is consistent with the reports of other *R. solanacearum* strains. This shows that along with the major housekeeping genes, some of the essential virulence associated genes are also present on the chromosome (Genin and Denny 2012).

Motility associated genes in R. solanacearum help the bacterium to locate and invade the host root for colonization (Meng et al. 2011). The genes identified for chemotaxis (*CheA* and *CheW*) and swimming motility (*FliC* and *FlgM*) are located on the megaplasmid whereas twitching motility (*PilA* and *PilP*) are located on the chromosome. The mutants of swimming motility are highly reduced in the degree to cause virulence on tomato plants under natural conditions (Tans-Kersten et al. 2001) and of chemotaxis are completely nonchemotactic (Yao and Allen 2006). The swimming motility and chemotaxis associated genes FliC, *PilP* and *CheW* are found to be highly conserved among all phylotypes and share more than 95% similarity (table 3 in electronic supplementary material). These are probably the regions which are not evolving or are conserved across the phylotypes. Twitching motility is a trait associated with the type IV pili and plays an important role in autoaggregation and biofilm formation (Kang et al.

2002). The *pilP* gene of Rs-10-244 shares 100% similarity with GMI1000 whereas Rs-09-161 is 99% similar. The *PilA* gene shares 89% similarity within the two strains and 91% with GMI1000. *PilA* exhibits diversity in the sequence among the other phylotypes of RSSC and is probably the region which undergoes evolution and thus can be used for designing primers for the strain-wise differentiation. *PilA* has been used to study the genetic diversity in soil bacterium *Myxococcus xanthus* strains and has shown highest polymorphism in comparison to that of other genes used (Vos and Velicer 2006). Further, we observed that there are no major differences between Indian strains and reference strain in the virulence gene sequences except *pilA* gene.

Sequences of virulence associated genes in strains Rs-09-161 and Rs-10-244 are found to be more close to phylotypes III and IV strains. Similar results were also observed by Ramesh *et al.* (2014b), when *egl* and *hrp* gene sequences from phylotype I strains were analysed.

Analysis of T3Es

We have identified 72 T3Es in Rs-09-161 and 77 T3Es in Rs-10-244 (including one multiple copy T3E) based on the identification criteria (table 2). The identified T3Es are assigned the names with prefix Rip (Ralstonia injected protein) as per the newly proposed nomenclature by Peeters et al. (2013) and the locus tag of the T3Es is represented by the prefix 161_ and 244_ for Rs-09-161 and Rs-10-244, respectively. The T3E gene RipTPS is present in multiple copies in both the strains. Rs-09-161 has the presence of three candidate effectors (Rs_T3E_Hyp6, Rs_T3E_Hyp7 and Rs_T3E_Hyp15) and three pseudogenes (RipF1, RipAX1 and Rs_T3E_Hyp8). Rs-10-244 has the presence of one candidate effectors (Rs_T3E_Hyp7) and four pseudogenes (RipO1, RipAX2, Rs T3E Hyp8 and Rs_T3E_Hyp15). Pseudogenes are nonfunctional genes and its presence can be attributed to the fact that either these genes are been mutated due to the presence of transposable elements within the gene, leading to its disruption or due to errors in sequencing. Comparison of the functional T3Es genes of R. solanacearum strains Rs-09-161, Rs-10-244 and GMI1000 revealed 60 common T3Es within the three strains. Rs-09-161 has two unique T3Es (Rs T3E Hyp6 and Rs T3E Hyp15) and shares 63 common effectors with GMI1000 and 66 T3Es with Rs-10-244. Rs-10-244 bears four unique T3Es (RipC2, RipE2, RipP3 and RipBB) and shares 66 common effectors with GMI1000 (figure 1). Majority of the R. solanacearum strains have an average of 70–75 T3E, which is much larger than many other bacterial plant pathogens like *P. syringe* and Xanthomonas sp., where it is in the range of 30-40 (Zumaquero et al. 2010; Hajri et al. 2011). Hence, it is presumed that an ancestor of R. solanacearum probably possessed a large number of effectors since the majority

Reference LT* Locus tag Rs-09-161	Reference LT*	Locus tag Rs-09-161	GC (%)	Length	% Blast	Locus tag Rs-10-244	GC (%)	Length	% Blast
	D6-7120	00206 121	26.92	LL 1 C	00	00136 146	CC 03	1000	00
Din A1	RSC2139 D S=0000	101_20090	06.00 2.07	01/10 2270	66 00	244_30120 244_00730	70.00 74 01	5204 2204	66 00
Dip A2	DShould	06040 ⁻¹⁰¹	2.07	0/02	00	244-00/20	71.06	2104	00
CENTRA Din AA	D Spuce D	101_40//0	11.90	20/0	66 00	09/04-101	2 17	3000	00
Rind5	RSP1024		60.51	5020	00	244_00210	60 43	3776	00
Ring	RSc0245	161_72300	10.00	1584	6	004070777 0740 18450	69 I	1479	66
RinC1	R Sn1730	161_022201	66.87	7837	00	244 11640	07:1 66 41	2745	66
RinCo	RCFRP mn20032		0.00	7007 U		244 24680	59.83	0407	<u> 8</u>
RinD	RCI DI LIIPZUUJZ	161 35070	63.61	1037	00		63 71	1037	10
RinFl	RSr3369	161_31680	67.84	1778	10	244 47820	68 38	1778	00
RinE2	RCFBP mp10565	Nil	0	0	<u></u>	244 24320	57.64	606	88
RipF1	RSp1555	Pseudogene	0	0	0	244 08360	64.41	2181	66
RipF2	RCFBP_mp30453	Nil	0	0	0	Nil	0	0	0
RipG1	RSp0914	$161_{-}41320$	65.7	1986	66	244_08450	65.71	1998	66
RipG2	RSp0672	$161_{-}39180$	69.35	3129	98	244_06280	69.41	3129	66
RipG3	RSp0028	161_{-33870}	68.28	1797	98	$244_{-}00270$	68.06	1794	66
RipG4	RSc1800	161_17570	71.34	1389	66	$244_{-}33040$	71.34	1389	66
RipG5	RSc1801		69.88	1617	66	244_33050	70.19	1617	66
RipG6	RSc1356	161_13550	68.85	1875	88	244_29420	68.75	1866	88
RipG7	RSc1357	161_13560	64.97	1836	80	244_29430	66.89	1935	98
RipG8	CMR15v4_10224	Nil	0	0	0	Nil	0	0	0
RipH1	RSc1386	161_13830	69.3	2304	66	244_29700	69.32	2298	66
RipH2	RSp0215	161_{-35090}	67.76	2274	98	244_01730	68.28	2229	98
RipH3	RSp0160	161_{-34560}	67.54	2160	96	244_{-01190}	67.68	2160	99
RipH4	RPSI07_mp0161	Nil	0	0	0 0	NI	0	0	0 0
RipI	RSc0041	161_00420	65.5 27	1206	96 97	244_16460	64.73	1293	66 00
KıpJ	RSc2132	161_{-41350}	5/.41	21/2	8/	244_36060	/8.80	1296	66 0
KIPK Dint	KCFBF_mp10024	INI 161 34000	0 70 03	0	0 00	1NII 244 01 530	n D	0 17 0	0
RIPL	K2pU195 D So 1475	161_34900	69.03 71 5	41/0	86	244_01250	00.00 71 5	41/5	66
DinN	D Co 1136	101-10100	(1.7 66.1	CC/1 2011	66 00	010201442	C.17	2071 2071	66 00
RinO1	R Sh0373	101_43400 161_36140	00.1 61 2	1530	99 80	Denidogene	00.00 67 36	1423	00
RinO2	RALSY mn30159	liN	0.10	0	ς C	I octubent Nil	0.20	0	
RipP1	RSp0826	Nil	0	0	0	244 24210	55.46	1107	100
RipP2	RSp0868	$161_{-}08570$	59.71	1467	66	$244_{-}24510$	59.71	1467	66
RipP3	$RSY45_{33690}$	Nil	0	0	0	244_46550	61.41	1161	66
RipQ	RSp1277	$161_{-}44830$	68.85	1557	66	$244_{-}12020$	69.04	1557	66
RipR	RSp1281	161_{-44870}	70.6	5229	66	$244_{-}12060$	70.66	5229	66
RipS1	RSc3401		69.04	7056	98	Nil	0	0	0
RipS2	RSp1374	$161_{-}45440$	70.18	7539	66	$244_{-}12410$	66.69	7458	66
RipS3	RSp0930	161_41420	66.81	6876	99 9	244_08550	66.85	6876	99
RipS4	RSc1839	161_17940	70.78	7725	66 90	244_33400	70.78	7725	66 8
csqix	KSP0296		CC./0	/10/	66 00	244_02630	C./0	/10/	66 90
KipS6	RSc2130	161_20590	66.62	2424	<u>ور</u>	244_36050	66.35	2547	<u>ور</u>
/sdix	1071dm ^{-/} 01CM	INI	n		0	IINI	n	0	

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(contd)
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Table

and armit	(course)								
	Reference LT*	Locus tag Rs-09-161	GC (%)	Length	% Blast	Locus tag Rs-10-244	GC (%)	Length	% Blast
RipS8	RPSI07_1850	$161_{-}17980$	68.77	9303	92	244_3343	68.89	9294	92
RipT	RSc3212	Nil	0	0	0	$244_{-}46320$	56.62	996	66
RipU	RSp1212	161_{-44210}	62.57	879	0	244_11370	61.84	891	100
RipV1	RSc1349	161_{13470}	67.95	2013	100	244_29350	68 0	2013	66 °
Kip V2	C6210/-1893	11N 111 26130	0	0	05	11NI 244 41770	0	01140	0 00
Kipw DioV	C//72CX	161_20420	10.00	006	56 20	0// 14-741//0	90.50 20 23	1140	66 00
NipA DinV	1/000CX	161_41090	00.70 672	066 0020	00	244_0020U	20.10	CZU1	06
Rin7	R Sn1031	161_02290	27/0 22/0	4110 4110	00	01001-442	00.00	4113 4113	700
Rinda	RSPIN7 7747	161_06050	60 56	804	C 8	06660-442	60 4 67 4	202	L 8
RinAB	RSn0876	161 41080	65 14	525	66 60	244 08250	64 76	525	66
RinAC	RSp0875	161 - 41070	63.59	2958	67	244 08240	62.92	2643	66
RipAD	RSp1601	161 - 47560	64.11	978	98	244 14660	64.21	978	66
RipAE	RSc0321	161_{-03080}	66.4	1932	66	244_{-19150}	67.02	1932	66
RipAF1	RSp0822	$0161_{-}4053$	1020	846	95	244_07700	68.78	1041	67
RipAF2	RALSY_20037	Nil	0	0	0	Nil	0	0	0
RipAG	RSc0824	$161_{-}08850$	59.59	297	98	$244_{-}24200$	60.6	297	100
RipAH	RSc0895	Nil	0	0	0	244_24720	62.88	291	100
RipAI	RSp0838	161_40690	70.75	612	66 ĭ	244_07860	71.07	612	66 0
RipAJ	RSc2101	161_{-20300}	69.15	1005	99	244_35750	69.05	1005	96
RipAK	RSc2359	NI	οï	0 0	0 00	244_38200	65.72	2430	99
RipAL	RPS107_mp0618	161_39680	70	1350	66 80	244_06790	70	1350	66 8
KipAM	KSc3272	161_30700	/1.82	465 2005	66 90	244_46830	/1.18	465	66 00
KipAN	KSp0845	161_40/60	6/.96 28.07	4323	66	244_0/940	67.82	4323	66 90
	KSPU8/9		08.90	14/9 2002	76	244_08280	10.60	14/9	86.00
RipAP Din AO	CMR15V4_10224 D G 0005	161_44240 161_41170	61.32 60 6	2803	93 00	244_11400	67.49 60.5	1242	93 00
RinAR	RSn1236	161_411/0	02.0	2112	96 96	244_00540 244_11600	09.5 60 63	2112 1818	66 80
RindS	RSn1384	095540 161 45560	71 86	2802	00	244_11500 244_12510	CD.CD 27 17	2802	00
RinAT	RSn1388	161 45600	70.65	1755	66	244 12550	70.59	1755	66
RipAU	RSp1460	161 - 46300	67.19	939	66	244 13250	67.3	939	66
RipAV	RSp0732	161_{-39740}	68.3	2538	66	244_{-06850}	68.7	2538	66
RipAW	RSp1475	$161_{-}46450$	66.29	1347	66	$244_{-}13400$	66.44	1347	97
RipAX1	RSc3290	Pseudogene	0	0	0	244_47020	58.68	759	100
RipAX2	RSp0572	$161_{-}38220$	57.99	657	66	Pseudogene	0	0	0
RipAY	RSp1022	$161_{-}42340$	64.48	1236	97	$244_{-}09440$	64.56	1236	67
RipAZI	RSp1582	161_{-47400}	61.03	834	66	$244_{-1}4480$	60.91	834	66
RipAZ2	RALSY_20407	Nil	0	0	0 0	Nil	0	0	0 0
KipBA	KSc0227	$161_{-0.2140}$	54.71	594 2	99	244_182/0	55.21	594	98
KipBB	KPSI0/_mp05/3		0 0	0 0	0 0	244_05100	61.43	1320	0 0
RipbC Damp	BATEV 20184								
Ringe	R \$1000 R 1010	Nil				Nil Nil			
DieDE	D DCIUZ Jecz	THAT				III I			
RipBC	COOL_VICIN	INI Nël				INI Nël			
Dadra	CO/DOMINICAL	IINT				IINT	0		

	Reference LT*	Locus tag Rs-09-161	GC (%)	Length	% Blast	Locus tag Rs-10-244	GC (%)	Length	% Blast
RipBH	RPSI07 mp30113	Nil	0	0	0	Nil	0	0	0
RipBI	RCFB mp30113	Nil	0	0	0	Nil	0	0	0
RipTAL	RSc1815	161_17720	66.04	3726	66	244_{33180}	66.05	3738	66
RipTPS	RSp0731	161_{-39730}	68.68	1785	66	244_06840	68.57	1785	66
			¢	¢	c	244_10270	¢	¢	c
RS_T3E_Hyp1	$RSPsi07_0331$	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp2	$RSPsi07_1883$	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp3	RSPsi07_mp0834	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp4	RSPsi07_mp1047	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp5	RSPsi07_mp1559	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp6	CMR15v4_30001	161_{-32360}	55.45	1311	88	Nil	0	0	0
RS_T3E_Hyp7	RSMK06225	161_{32180}	55.1	519	0	$244_{-}48280$	55.1	519	0
RS_T3E_Hyp8	RSMK02655	Pseudogene	0	0	0	Pseudogene	0	0	0
RS_T3E_Hyp9	RRSL_01783	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp10	RSMK02638	Nil	0	0	0	Nil	0	0	0
RS T3E Hyp11	RSMK01187	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp12	RSMK03335	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp13	RSPO_m01098	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp14	$BDB mp_{40006}$	Nil	0	0	0	Nil	0	0	0
RS_T3E_Hyp15	$RSPsi07_1860$	161_{-35100}	52.73	2799	94	Pseudogene	0	0	0
RS_T3E_Hyp16	RSc3174	Nil	0	0	0	Nil	0	0	0

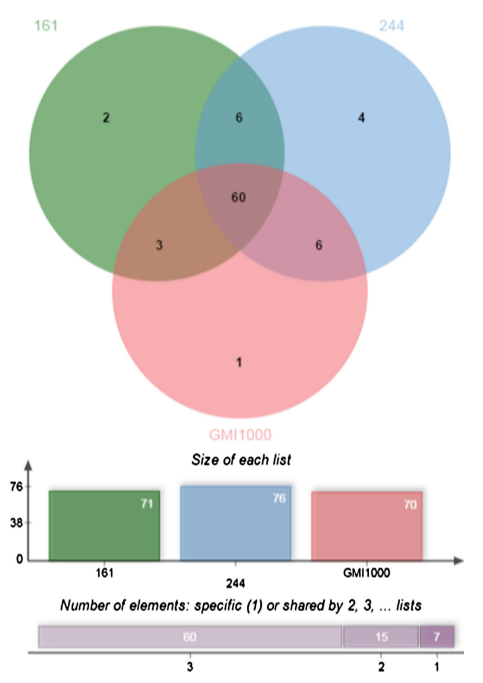


Figure 1. Venn diagram constructed using T3Es complete genes of *R. solanacearum* strain Rs-09-161(green), Rs-10-244 (blue) and GMI1000 (pink). The merged regions display the T3Es shared between the strains. In Rs-09-161 and Rs-10-244, one effector is present in multiple copies which is not indicated in the bar graph.

of the strains possess a high number of effectors. The only exception to this is BDB, which has less number of effectors (Genin and Denny 2012).

A majority of *R. solanacearum* T3Es have been validated through translocation studies (Cunnac *et al.* 2004; Mukaihara *et al.* 2004; Mukaihara and Tamura 2009; Mukaihara *et al.* 2010; Sole *et al.* 2012). Many of the effectors share homology with those of other bacterial plant pathogens like *P. syringae, Xanthomonas* sp. and Acidovorax sp. Few of these effectors are present as effector families and have three to eight effectors. These include RipA (AWR family), RipG (GALA family), RipH (HLK family) and the RipS (SKWP family) (Poueymiro and Genin 2009; Mukaihara et al. 2010; Remigi et al. 2011; Sole et al. 2012; Peeters et al. 2013). These effectors possess certain inherent characters about the sequence such as specific internal repeats within them, which characterizes them to constitute a family. The T3SS secretes T3Es in a highly specialized manner in the eukaryotic hosts and is used by many plant and animal pathogenic bacteria, as a tool to colonize in their respective hosts. The various motifs or domains present on the T3Es interact with the host cells and benefits the bacteria in colonization. The T3Es are secreted into the cytosol of the host through the *hrp* pili. Unique T3E of Indian strains (Rs-09-161 and Rs-10-244) were analysed for functional/conserved motifs through homology search in NCBI revealed the presence of various motifs. The T3E RipBB present in Rs-10-244 exhibits presence of ankyrin repeats which mediate protein-protein interactions in diverse families of proteins. RipC2 shares homology with haloacid dehalogenase (HAD)-like hydrolases and RipP3 which is also known as PopP3 displays YopJ serine/threonine acetyltransferase activity. The T3E Rs T3E Hyp15 present in Rs-09161 displays presence of serine/threonine protein kinase domain within it.

RipA (*AWR* family) effectors include five effectors (RipA1 to RipA5), and both the strains, Rs-09-161 and Rs-10-244 have all the RipA effectors present in them. Among the RipA effectors, RipA1 is present only in phylotype I strains, whereas RipA2 along with RipA4 and RipA5 is present in all the phylotypes of *R. solanacearum* isolates studied till date. RipA5 is also present in multiple copies in some of the phylotype II strain (Molk2, IPO1609, UW551 and Po82). The RipA effectors consist of a conserved region containing the alanine–tryptophan–arginine tryad and can be virulent or avirulent depending on the host with which *R. solanacearum* interacts and RipA2 was found to be a major contributor to the virulence among the AWR family (Sole *et al.* 2012).

RipG (*GALA* family) possesses eight T3Es (RipG1– RipG8) and seven (RipG1–RipG7) are present in both Rs-09-161 and Rs-10-244. The RipG has the presence of leucine rich repeats (LRR) and F-box domain with them. The F-box protein forms a component of E3ubiquitin ligase complex and is found in eukaryotes. This complex plays an important role in ubiquitination of proteins which leads to the degradation or modification of the activity of the targeted protein (Hua and Vierstra 2011). RipG8 is present only in CMR15 (phylotype III). More isolates from phylotype III needs to be studied to identify if the RipG is specific to phylotype III strains.

RipH (*HLK* family) consists of four effectors (RipH1– RipH4); Rs-09-161 and Rs-10-244 has the presence of RipH1, RipH2 and RipH3 with an average size of \sim 600 amino acids. RipH4 is found to be present only among phylotype IV strains. The RipH (*HLK* family) is named so because of the presence of histidine–leucine–lysine triad in a conserved C-terminal region. Phylogenetic analysis of the RipH effectors indicates an ancestral strain of *R. solanacearum* most likely had only three RipH effectors and the fourth one has evolved later independently (Chen *et al.* 2014).

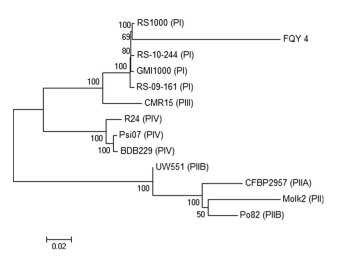


Figure 2. Phylogenetic tree constructed using NJ method based on the coding sequences of effectors of RipA (*AWR* family) of *R. solanacearum* isolates. The isolates are represented with their names followed by the phylotypes in parenthesis. The tree was generated by MEGA-6 (Tamura *et al.* 2013) software using the NJ and the algorithm of Jukes and Cantor (1969) with 1000 bootstrap resamplings. Numbers at each branch indicate bootstrap value. The scale indicates the genetic distance.

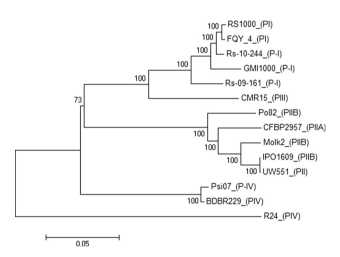


Figure 3. Phylogenetic tree constructed using NJ method based on the coding sequences of effectors of RipG (*GALA* family) of *R. solanacearum* isolates. The isolates are represented with their names followed by the phylotypes in parenthesis. The tree was generated by MEGA-6 (Tamura *et al.* 2013) software using the NJ and the algorithm of Jukes and Cantor (1969) with 1000 bootstrap resamplings. Numbers at each branch indicate bootstrap value. The scale indicates the genetic distance.

RipS (*SKWP* family) has eight effectors (RipS1–RipS8); RipS7 is absent in Rs-09-161 whereas RipS1 and RipS7 are absent in Rs-10-244. RipS7 is absent in all phylotype I strains studied till date and is present in all phylotype IV strains. RipS1 and RipS6 is absent in all phylotype IV isolates. Phylotype II lacks RipS6 and RipS8. The structure of RipS (*SKWP* family) effectors is found to be related to heat/armadillo repeat domain. The RipS proteins exert

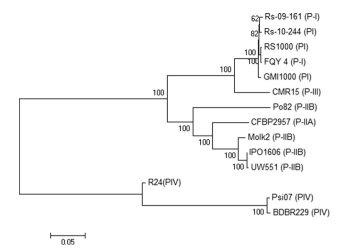


Figure 4. Phylogenetic tree constructed using NJ method based on the coding sequences of effectors of RipH (*HLK* family) of *R. solanacearum* isolates. The isolates are represented with their names followed by the phylotypes in parenthesis. The tree was generated by MEGA-6 (Tamura *et al.* 2013) software using the NJ and the algorithm of Jukes and Cantor (1969) with 1000 bootstrap resamplings. Numbers at each branch indicate bootstrap value. The scale indicates the genetic distance.

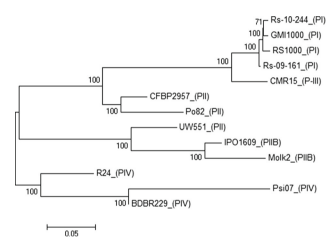


Figure 5. Phylogenetic tree constructed using NJ method based on the coding sequences of effectors of RipS (*SKWP* family) of *R. solanacearum* isolates. The isolates are represented with their names followed by the phylotypes in parenthesis. The tree was generated by MEGA-6 (Tamura *et al.* 2013) software using the NJ and the algorithm of Jukes and Cantor (1969) with 1000 bootstrap resamplings. Numbers at each branch indicate bootstrap value. The scale indicates the genetic distance.

their virulence on their host plant by interaction through the SKWP domain (Mukaihara and Tamura 2009).

Phylogenetic analysis of T3Es

The phylogenetic trees constructed for RipA, RipG, RipH and RipS effectors are depicted in figures 2–5. Effector gene sequences appear to be conserved and thus have revealed grouping of the isolates based on their phylotypes. Group 1 consists of strains that belong to phylotype I including Rs-09-161 and Rs-10-244. Phylotype III strain CMR15 forms a group closely associated with phylotype I. The strains belonging to phylotype II (IIA and IIB) and phylotype IV have separated to form two different groups. This grouping is consistent with the four gene families studied here. Similar results are also observed by Remenant *et al.* (2011) and Peeters *et al.* (2013), where isolates from phylotype I and III have clustered together. It is likely that the isolates from phylotype I and phylotype I and phylotype III did not undergo much evolution and hence form a major group (Remenant *et al.* 2011).

In conclusion, in this study, analysis of virulence genes and T3E genes of *R. solanacearum* strains Rs-09-161 and Rs-10-244 indicated that a majority of the virulence associated genes are present in both the strains. It was observed that all the virulence genes of Rs-09-161 and Rs-10-244 are highly conserved and share high level of similarity except for *pilA* gene, which shares a minimum of 72% similarity. Seventy-two T3E genes were identified in *R. solanacearum* strain Rs-09-161 and 77 in Rs-10-244. Phylogenetic analysis of T3E genes of RipA, RipG, RipH and RipS revealed close association between phylotype I and phylotype III strain of *R. solanacearum*.

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References

- Alfano J. R. and Collmer A. 2004 Type III secretion system effector proteins: double agents in bacterial disease and plant defense. *Annu. Rev. Phytopathol.* 42, 385–414.
- Alvarez B., Biosca E. and Lopez M. 2010 On the life of *Ralsto-nia solanacearum*, a destructive bacterial plant pathogen. In *Current research, technology and education topics in applied microbiology and microbial biotechnology.* (ed. A. Mendez Vilas), vol. 1, pp. 267–279. Formatex, Badajoz.
- Boucher C. A., Van Gijsegem F., Barberies P. A., Arlat M. and Zischek C. 1987 *Pseudomanas solanacearum* genes controlling both pathogenicity on tomato and hypersensitivity on tobacco are clustered. *J. Bacteriol.* 169, 5626–5633.
- Chandrashekara K. N., Kumar M. P. and Saroja S. 2012 Aggressiveness of *Ralstonia solanacearum* isolates on tomato. J. Exp. Sci. 3, 05-09.
- Chen L., Shirota M., Zhang Y., Kiba A., Hikichi Y. and Ohnishi, K. 2014 Involvement of HLK effectors in *Ralstonia* solanacearum disease development in tomato. J. Gen. Plant Pathol. 80, 79–84.
- Cunnac S., Occhialini A., Barberis P., Boucher C. and Genin S. 2004 Inventory and functional analysis of the large Hrp regulon in *Ralstonia solanacearum*: identification of novel effector

proteins translocated to plant host cells through the type III secretion system. *Mol. Microbiol.* **53**, 115–128.

- Fegan M. and Prior P. 2005. How complex is the *Ralstonia* solanacearum species complex? In *bacterial wilt disease and* the *Ralstonia solanacearum species complex* (ed. C. Allen, P. Prior and A. C. Hayward), pp. 449–461. APS Press, St Paul.
- Gabriel D. W., Allen C., Schell M., Denny T. P., Greenberg J. T., Duan Y. P. *et al.* 2006 Identification of open reading frames unique to a select agent: *Ralstonia solanacearum* race 3 biovar 2. *Mol. Plant Microbe Interact.* **19**, 69–79.
- Genin S., Gough C. L., Zischek C. and Boucher C. A. 1992 Evidence that the hrpB gene encodes a positive regulator of pathogenicity genes from *Pseudomonas solanacearum*. *Mol. Microbiol.* 6, 3065–3076.
- Genin S. and Denny T. P. 2012 Pathogenomics of the *Ralstonia* solanacearum species complex. Annu. Rev. Phytopathol. **50**, 67– 89.
- Hajri A., Pothier J. F., Fischer-Le Saux M., Bonneau S., Poussier S. et al. 2011 Type three effector genes distribution and sequence analysis provides new insights into pathogenicity of plant pathogenic Xanthomonas arboricola. Appl. Environ. Microbiol. 78, 371–384.
- Heesemann J., Algermissen B. and Laufs R. 1984 Genetically manipulated virulence of *Yersinia enterocolitica*. *Infect. Immun.* 46, 105–110.
- Hua Z. and Vierstra R. D. 2011 The cullin-RING ubiquitinprotein ligases. Annu. Rev. Plant Biol. 62, 299–334.
- Hueck C. J. 1998 Type III protein secretion systems in bacterial pathogens of animals and plants. *Microbiol. Mol. Biol. Rev.* 62, 379–433.
- Jukes T. H. and Cantor C. R. 1969 Evolution of protein molecules. In*Mammalian protein metabolism* (ed. H. N. Munro), pp. 21–132. Academic Press, New York.
- Kang Y., Liu H., Genin S., Schell M. A. and Denny T. P. 2002 *Ral-stonia solanacearum* requires type 4 pili to adhere to multiple surfaces and for natural transformation and virulence. *Mol. Microbiol.* 46, 427–437.
- Kumar A., Sarma Y. R. and Anandaraj M. 2004 Evaluation of genetic diversity of *Ralstonia solanacearum* causing bacterial wilt of ginger using REP-PCR and PCR-RFLP. *Curr. Sci.* 87, 1555–1561.
- Mansfield J., Genin S., Magori S., Citovsky V., Sriariyanum M., Ronald P. et al. 2012 Top 10 plant pathogenic bacteria in molecular plant pathology. *Mol. Plant Pathol.* 13, 614–629.
- Meng F., Yao J. and Allen C. 2011 A MotN mutant of *Ralsto-nia solanacearum* is hypermotile and has reduced virulence. J. Bacteriol. **193**, 2477–2486.
- Mukaihara T., Tamura N., Murata Y. and Iwabuchi M. 2004 Genetic screening of Hrp type III-related pathogenicity genes controlled by the HrpB transcriptional activator in *Ralstonia solanacearum. Mol. Microbiol.* **54**, 863–875.
- Mukaihara T. and Tamura N. 2009 Identification of novel *Ralstonia solanacearum* type III effector proteins through translocation analysis of hrpB-regulated gene products. *Microbiology* 155, 2235–2244.
- Mukaihara T., Tamura N. and Iwabuchi M. 2010 Genome-Wide identification of a large repertoire of *Ralstonia solanacearum* type III effector proteins by a new functional screen *Mol. Plant Microbe Interact.* 23, 251–262.
- Peeters N., Carrere S., Anisimova M., Plener L., Cazale A. C. and Genin S. 2013 Repertoire, unified nomenclature and evolution of the type III effector gene set in the *Ralstonia solanacearum* species complex. *BMC Genomics* 14, 859–877.
- Poueymiro M. and Genin S. 2009 Secreted proteins from *Ralsto-nia solanacearum*: a hundred tricks to kill a plant. *Curr. Opin. Microbiol.* 12, 44–52.

- Ramesh R. and Phadke G. S. 2012 Rhizosphere and endophytic bacteria for the suppression of eggplant wilt caused by *Ralstonia solanacearum*. Crop Prot. 37, 35–41.
- Ramesh R., Gaitonde S., Achari G., Asolkar T., Singh N. P., Carrere S. *et al.* 2014a Genome sequencing of *Ralstonia solanacearum* Biovar 3, phylotype I, strains Rs-09-161 and Rs-10-244, ssolated from eggplant and chili in India. *Genome Announc.* 2, 1–2.
- Ramesh R., Achari G. A. and Gaitonde S. 2014b Genetic diversity of *Ralstonia solanacearum* infecting solanaceous vegetables from India reveals the existence of unknown or newer sequevars of Phylotype I strains. *Eur. J. Plant Pathol.* 140, 543– 562.
- Remenant B., Coupat-Goutaland B., Guidot A., Cellier G., Wicker E., Allen, C. *et al.* 2010 Genomes of three tomato pathogens within the *Ralstonia solanacearum* species complex reveal significant evolutionary divergence. *BMC Genomics.* 11, 379–395.
- Remenant B., de Cambiaire J-C., Cellier G., Jacobs J. M., Mangenot S., Barbe V. *et al.* 2011 Ralstonia syzygii, the blood disease bacterium and some Asian *R. Solanacearum* strains form a single genomic species despite divergent lifestyles. *PLoS One* 6, e24356.
- Remigi P., Anisimova M., Guidot A., Genin S. and Peeter N. 2011 Functional diversification of the GALA type III effector family contributes to *Ralstonia solanacearum* adaptation on different plant hosts. *New Phytol.* **192**, 976–987.
- Saile E., McGarvey J., Schell M. and Denny T. 1997 Role of extracellular polysaccharide and endoglucanase in root invasion and colonization of tomato plants by *Ralstonia solanacearum*. *Phytopathology* 87, 1264–1271.
- Salanoubat M., Genin S., Artiguenave F., Gouzy J., Mangenot S., Arlat M. *et al.* 2002 Genome sequence of the plant pathogen *Ralstonia solanacearum. Nature* **415**, 497–502.
- Schell M. A. 2000 Control of virulence and pathogenicity genes of *Ralstonia solanacearum* by an elaborate sensory network. *Annu. Rev. Phytopathol.* 38, 263–292.
- Sole M., Popa C., Mith O., Sohn K. H., Jones J. D., Deslandes L. et al. 2012 The awr gene family encodes a novel class of *Ralstonia solanacearum* type III effectors displaying virulence and avirulence activities. *Mol. Plant Microbe Interact.* 25, 941– 953.
- Tamura K., Stecher G., Peterson D., Filipski A. and Kumar S. 2013 MEGA6: molecular evolutionary genetics analysis version 6.0. *Mol. Biol. Evol.* **30**, 2725–2729.
- Tans-Kersten J., Huang H. and Allen C. 2001 Ralstonia solanacearum needs motility for invasive virulence on tomato. J. Bacteriol. 183, 3597–3605.
- Thompson J. D., Higgins D. G. and Gibson T. J. 1994 CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. *Nucleic Acids Res.* 22, 4673–4680.
- Van Gijsegem F., Vasse J., Camus J.-C., Marenda M. and Boucher C. 2000 *Ralstonia solanacearum* produces Hrp-dependent pili that are required for PopA secretion but not for attachment of bacteria to plant cells. *Mol. Microbiol.* **36**, 249–260.
- Van Gijsegem F., Vasse J., Rycke R. D., Castello P. and Boucher C. 2002 Genetic dissection of the *Ralstonia solanacearum* hrp gene cluster reveals that the HrpV and HrpX proteins are required for Hrp pilus assembly. *Mol. Microbiol.* 44, 935–946.
- Vos M. and Velicer, G. J. 2006 Genetic population structure of the soil bacterium *Myxococcus xanthus* at the centimeter scale. *Appl. Environ. Microbiol.* **72**, 3615–3625.
- Xu J., Zheng H. J., Liu L., Pan Z. C., Prior P., Tang B. et al. 2011 Complete genome sequence of the plant pathogen Ralstonia solanacearum strain Po82. J. Bacteriol. 193, 4261–4262.

- Yao J. and Allen C. 2006 Chemotaxis is required for virulence and competitive fitness of the bacterial wilt pathogen *Ralstonia solanacearum*. J. Bacteriol. **188**, 3697–3708.
- Zumaquero A., Macho A. P., Rufian J. S. and Beuzon C. R. 2010 Analysis of the role of the type III effector inventory of *Pseudomonas syringae* pv. phaseolicola 1448a in interaction with the plant. *J. Bacteriol.* **192**, 4474–4488.

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