

1 The Trk potassium transporter is required for RsmB-mediated activation of virulence in the
2 phytopathogen *Pectobacterium wasabiae*

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15 Running Head: Potassium regulates RsmB-mediated virulence activation

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27 **Abstract**

28 *Pectobacterium wasabiae* (previously known as *Erwinia carotovora*) is an important plant pathogen
29 that regulates the production of plant cell wall-degrading enzymes through both an *N*-acyl-
30 homoserine lactone-based quorum sensing system and the GacS/GacA two component system (also
31 known as ExpS/ExpA). At high cell density, activation of GacS/GacA induces the expression of
32 RsmB, a non-coding RNA which is in turn essential for activation of virulence in this bacterium. A
33 genetic screen to identify regulators of RsmB revealed that mutants defective in components of a
34 putative Trk potassium transporter (*trkH*, *trkA*) had decreased *rsmB* expression. Further analysis of
35 these mutants showed that changes in potassium concentration influenced *rsmB* expression and
36 consequent tissue damage in potato tubers, and that this regulation required an intact Trk system.
37 Regulation of *rsmB* expression by potassium via the Trk system occurred even in the absence of the
38 GacS/GacA system, demonstrating that these systems act independently and are both required for
39 full activation of RsmB and the downstream induction of virulence in potato infection assays.
40 Overall, our results identified potassium as an essential environmental factor regulating the Rsm
41 system, and the consequent induction of virulence, in the plant pathogen *P. wasabiae*.

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43 **Importance**

44 Crop losses from bacterial diseases caused by pectolytic bacteria are a major problem in
45 agriculture. By studying the regulatory pathways involved in controlling expression of plant cell
46 wall-degrading enzymes in *Pectobacterium wasabiae* we showed that the Trk potassium transport
47 system plays an important role in the regulation of these pathways. The data presented further
48 identify potassium as an important environmental factor in the regulation of virulence in this plant
49 pathogen. We showed that a reduction in virulence can be achieved by increasing the extracellular
50 concentration of potassium. Therefore, this work highlights how elucidation of the mechanisms
51 involved in regulating virulence can lead to the identification of environmental factors that can
52 influence the outcome of infection.

53 **Introduction**

54 *Pectobacterium* spp. are Gram-negative, rod-shaped bacteria belonging to the
55 Enterobacteriaceae family. They cause soft-rot disease in several plants, including potatoes, carrots
56 and cabbage. Damage to plant tissues is caused by the action of a mixture of cellulases, proteases,
57 pectate lyases (Pel), pectin lyases and polygalacturonases secreted by these bacteria. The enzymes
58 degrade plant cell wall components, releasing nutrients that fuel bacterial growth. Production of
59 these plant cell wall-degrading enzymes (PCWDEs), is carefully coordinated by a complex
60 multipartite regulatory system that integrates internal and external information to ensure that
61 virulence is only switched on when conditions are optimal (1).

62 In *Pectobacterium wasabiae* [previously *Erwinia carotovora* (2)], production of PCWDEs is
63 regulated mainly through two signal transduction systems. These systems coordinately control
64 expression and activity of the global post-transcriptional regulator, RsmA, which represses the
65 expression of PCWDEs by binding to their mRNAs. Transcription of *rsmA* is regulated by
66 ExpI/ExpR, a typical quorum sensing system that relies on *N*-acyl homoserine lactone (AHL)
67 autoinducer (3–5). The second sensory pathway regulating RsmA, and therefore virulence, is the
68 GacS/GacA two-component system (also known as ExpS/ExpA). The response regulator, GacA, is
69 the major transcriptional activator of RsmB, a non-coding RNA which binds RsmA (6). This
70 binding of RsmA by RsmB inhibits RsmA-mediated repression of PCWDEs, ultimately promoting
71 the expression of these virulence factors (7). Therefore, activation of RsmB expression is essential
72 to induce virulence in *P. wasabiae*. Homologues of the Gac/Rsm system exist in many Gram-
73 negative bacteria, including the Gac/Rsm system in *Pseudomonas* spp. and the BarA/UvrY/Csr in
74 *Escherichia coli*. Though these systems regulate a wide range of important physiological functions
75 in bacteria, including primary and secondary metabolism, biofilm formation, motility and virulence
76 [reviewed in (8, 9)], the chemical identity of the molecule(s) responsible for its activation remains
77 unknown. Accumulation of intermediates of the Krebs Cycle have been shown to stimulate the
78 Gac/Rsm in *Pseudomonas fluorescens* and *Vibrio fischeri* (10, 11), but the physiological conditions

79 that lead to the accumulation of these metabolites with the consequent activation of the Gac/Rsm
80 system are still poorly understood. Additionally, short-chain fatty acids, such as acetate, formate,
81 propionate and butyrate, have been shown to influence the expression of *csrB* (a functional *rsmB*
82 homologue) in *E. coli* or *Salmonella enterica enterica* serovar Typhimurium at low pH, but the
83 stimuli that activate the system at neutral pH have not been identified (12, 13).

84 To identify physiological stimuli involved in the activation of the Gac/Rsm system and
85 understand how this system regulates virulence in the well-characterized *P. wasabiae* strain
86 SCC3193, we performed a genetic screen to identify mutants affected in *rsmB* expression, the main
87 target of the Gac/Rsm system in this bacterium. The screen revealed 5 mutants defective in genes
88 coding for components of a putative homologue of the *E. coli* Trk potassium uptake system (10).
89 Therefore, here we investigated the importance of potassium and the Trk system for the regulation
90 of RsmB. Our results demonstrated that extracellular potassium is a critical environmental factor
91 influencing virulence in *Pectobacterium* spp.

92

93 **Materials and Methods**

94 **Bacterial strains, plasmids and media.** All strains and plasmids used in this study are listed in
95 Table S1. *P. wasabiae* strains are derived from the wild-type (wt) strain SCC3193 (14). Strains
96 were grown at 30°C with aeration in Luria-Bertani (LB) broth, M9 Minimal Media (MM) or
97 Minimal Potassium Media (15) with 0.4% (w/v) Glycerol. When specified, media were
98 supplemented with 0.4% of polygalacturonic acid (PGA; Sigma P3850) to induce the expression of
99 PCWDEs. Where mentioned, KCl was added at various final concentrations. Antibiotics were used
100 at the following concentrations: ampicillin (Amp) 100 mgL⁻¹; streptomycin (Str) 100 mgL⁻¹;
101 kanamycin (Kan) 50 mgL⁻¹; spectinomycin (Spec) 50 mgL⁻¹ and chloramphenicol (Cm) 25 mgL⁻¹.
102 To assess bacterial growth the optical density (OD₆₀₀) was determined by measuring absorbance at
103 600 nm in Bioscreen C Reader System (Multi-plate reader Oy Growth Curves Ab Ltd).

104 **Genetic and molecular techniques.** Primers used in this study are listed in Table S2. The plasmid
105 encoding the *rsmB* promoter fused to green fluorescence protein (GFP) ($P_{rsmB}::gfp$) was constructed
106 using a modified version of the promoterless::*gfp* vector pCMW1 (16). First the chloramphenicol
107 resistance gene (*cm*) was amplified by PCR from pKD3 (17) using P0276-pKD3/4(BamHI) and
108 P0277-pKD3/4(BamHI) primers and introduced into pCMW1, yielding PRSV59. Next a 392 bp
109 fragment containing the promoter region of *rsmB* was amplified by PCR using P0528-rsmB(SphI)
110 and P0529-rsmB(SalI) primers and ligated into PRSV59, yielding pRSV206. Deletion mutants were
111 constructed by chromosomal gene replacement with an antibiotic marker using the λ Red
112 recombinase system (17, 18). The DNA region of *gacA*, including approximately 500 bp upstream
113 and 500 bp downstream of the gene, was amplified by PCR and cloned into pUC18 (19) using SalI
114 and SacI. This construct, containing the *gacA* gene and its flanking regions, was divergently
115 amplified by PCR using primers to introduce an XhoI restriction site in the 5' and 3' regions of the
116 gene. The antibiotic resistance gene (*str*) was amplified from pKNG101 (20) with primers
117 containing the XhoI restriction site. The amplified fragment containing *str*^R was digested with XhoI
118 and introduced into the XhoI digested PCR fragment. The final construct contained the antibiotic
119 resistance marker flanked by the upstream and downstream regions of *gacA*. The 500bp-*str*-500bp
120 fragment was amplified by PCR and approximately 3 ng of DNA were electroporated into a strain
121 expressing the λ Red recombinase system from pKD46, to allow recombination (17). To construct
122 the *trkA*⁺ and *trkH*⁺ complementation plasmids, a 400bp-*trkA*-400bp fragment and a 400bp-*trkH*
123 fragment were amplified from SCC3193 and cloned into pOM18 using SalI and SacI or PstI and
124 XbaI, respectively. pOM18 was constructed by amplifying the multiple cloning site of pUC18 and
125 cloning it into the BglII site created from the divergently amplified pOM1 (21). PCR reactions for
126 cloning purposes were performed using the proof reading Herculase II Polymerase (Agilent).
127 Dream Taq Polymerase (Fermentas) was used for all other PCR reactions. Digestions were done
128 using Fast Digest Enzymes (Fermentas) and ligations performed with T4 DNA ligase (New

129 England Biolabs). All cloning steps were performed in *E. coli* DH5 α . All mutants and constructs
130 were confirmed by sequencing at the Instituto Gulbenkian de Ci \acute{e} ncia sequencing facility.

131 **Isolation of transposon insertion mutants with low *rsmB* expression.** A library of 15,126
132 mutants in SCC3193 was constructed by transposon mutagenesis using the EZ-Tn5TM
133 <R6K γ ori/KAN-2>Tnp TransposomeTM kit, according to the manufacturer's instructions
134 (Epicentre). Mutants were tested in 96-well plates for Pel activity levels using the thiobarbituric
135 acid (TBA) method (22). The pRSV206 plasmid, which contained the P_{*rsmB*}::*gfp* fusion, was
136 introduced into mutants with low Pel activity by electroporation. To measure *rsmB* expression
137 levels, these strains were grown in multi-well plates at 30°C in MM supplemented with Cm and
138 Kan and diluted 1:100 into fresh medium in black multi-well plates. After 24 h of growth, GFP
139 expression was assessed using a multi-label counter (Victor³, PerkinElmer). We identified 29
140 mutants with less than 75% of *rsmB* expression level shown by wt bacteria. The transposon
141 insertion site of these mutants was amplified by a two-step arbitrary PCR using the transposon
142 specific primers, P0058-Kan_SP1 and P0057-Kan_SP2, and the arbitrary primers, P0052-Arb1K,
143 P0053-Arb2k and P0054-Arb6K (23, 24). The insertion site was identified by DNA sequencing
144 coupled with BLAST analysis against the *Pectobacterium* SCC3193 complete genome sequence
145 (NCBI taxid:1166016).

146 **Analysis of expression of P_{*rsmB*}::*gfp*.** Analysis of *rsmB* expression was performed by flow
147 cytometry. Mutant and wt strains of *P. wasabiae* SCC3193 containing the *rsmB* reporter fusion
148 were grown overnight in MM supplemented with Cm and Kan then inoculated into fresh medium in
149 multi-well plates at a starting OD₆₀₀ of 0.02. Aliquots were collected at OD₆₀₀ = 0.3 or 0.4, as
150 indicated in the figures legends. Cells were diluted 1:100 into Phosphate Buffered Saline (PBS) and
151 fluorescence intensity of GFP per cell was assessed immediately in the flow cytometer (LSR
152 Fortessa, BD). Results were analyzed with Flowing Software v2.5.1. A minimum of 5,000 GFP-
153 positive single cells were acquired per sample and analyzed for their *rsmB* expression. *rsmB*

154 expression of the mutants with the $P_{rsmB}::gfp$ fusion is reported as the median GFP expression of the
155 GFP-positive single cells in arbitrary units (a.u.).

156 **Pel activity assay.** Overnight cultures of bacterial strains were diluted to an OD_{600} of 0.02 in fresh
157 LB supplemented with PGA in multi-well plates. Bacteria were sub-cultured to an OD_{600} of 0.4, at
158 which time cell-free supernatants were harvested by centrifugation. Extracellular Pel activity was
159 measured using the previously described TBA colorimetric method (22). Briefly, supernatants were
160 incubated for 3 h with the substrate mixture at 37° C. The reaction was stopped by acidification,
161 TBA (Sigma T5500) was added, and the reaction mixture boiled for 1 h. The pink coloration was
162 measured at 548 nm using a multi-label counter (Victor³, Perkin-Elmer) and normalized to the
163 OD_{600} of the culture.

164 ***P. wasabiae* virulence assay.** Virulence was determined using a modified protocol to assess
165 maceration of potato tubers (25). Potatoes were washed and surface sterilized by soaking for 10 min
166 in 10% bleach followed by 10 min in 70% ethanol. To prepare bacteria for inoculation, overnight
167 cultures were washed twice in PBS, which contained a total of 4.5 mM of potassium, or in PBS
168 with various concentrations of KCl, as indicated. 30 μ L of cells at an OD_{600} of 0.05 were inoculated
169 into previously punctured potatoes and incubated at 28°C with relative humidity above 90% for 48
170 h. After incubation, potatoes were sliced and macerated tissue was collected and weighed. To
171 quantify the inoculum, serial dilutions of this bacterial suspension were performed in PBS, plated
172 and bacterial growth was quantified by the number of colony forming units (cfu) present in 30 μ l.

173 **Statistical analysis.** Data were analyzed using Graphpad Prism6 software and R program version
174 3.0.2. The Mann-Whitney test was performed to determine statistical significance and P-values
175 were adjusted using the Holm-Bonferroni correction for multiple comparisons. An adjusted P-value
176 <0.05 was used as the cut-off for statistical significance. ^{ns}Not significant; *P-value <0.05; **P-
177 value <0.01; ***P-value <0.001.

178

179

180 **Results**

181 **Mutants in the Trk potassium uptake system are impaired in *rsmB* expression and production**
182 **of PCWDEs**

183 To identify regulators of the Gac/Rsm system we generated and screened a library of 15,126
184 transposon mutants in *P. wasabiae* SCC3193 for changes in Pel activity. We obtained 58 mutants
185 with reduced Pel activity compared to wt. *rsmB* expression levels were then tested in these mutants
186 following the introduction of a plasmid-encoded *rsmB* promoter-GFP fusion construct (pRSV206 –
187 *P_{rsmB}::gfp*) into the mutants. Of these, 29 independent mutants showed less than 75% of *rsmB*
188 expression in comparison to wt (Table S3). From those 29 mutants, five were found to have
189 transposon insertions in genes annotated as components of a putative Trk system, which is involved
190 in potassium uptake in *E. coli* (26). As shown in Fig. 1, all five mutants had an approximately two-
191 fold reduction in the expression of the *rsmB::gfp* fusion compared to wt levels. Three of those
192 mutants (RSV108, RVS141 and RSV154) had transposon insertions in the W5S_4400 gene,
193 annotated as the potassium uptake protein, TrkH. The protein encoded by this gene has 89%
194 sequence identity with one of the potassium uptake channel proteins (TrkH) from *E. coli*. This
195 bacterium has two redundant Trk channel proteins, TrkH and TrkG: disruption of both proteins is
196 required for the abolishment of potassium uptake by the Trk system. In contrast, *P. wasabiae*, has
197 only one putative Trk channel protein, similar to most other bacteria containing the Trk transporter.
198 The other two mutants (RSV124 and RSV236) had transposon insertions in the W5S_4132 gene,
199 annotated as the potassium uptake protein, TrkA. The protein encoded by this gene shares 85%
200 sequence identity with the TrkA protein from *E. coli*, a regulator of the Trk potassium uptake
201 system. In *E. coli*, disruption of *trkA* results in a lower rate of potassium uptake by the Trk system
202 (26). To investigate the role of the Trk system in the regulation of RsmB, one mutant with a
203 transposon insertion in the *trkH* gene (RSV141) and another with an insertion in the *trkA* gene
204 (RSV236) were selected for further characterization.

205 We tested the *trkH*::Tn5 and *trkA*::Tn5 mutants for complementation in *trans* with the *trkH*
206 or *trkA* genes, respectively, both for *rsmB* expression and Pel activity (Fig. 2). Due to the growth
207 defect for both *trkH* and *trkA* mutant strains (Fig. S1), cells were collected and analyzed at the same
208 cell density ($OD_{600} = 0.4$). $P_{rsmB}::gfp$ expression was restored to wt levels in *trkH* and *trkA* mutants
209 when the respective genes were expressed under the control of their own promoter, but remained
210 low in mutants carrying the empty vector (Fig. 2A). When tested for their ability to produce
211 PCWDEs, these mutants had a more than two-fold decrease in extracellular Pel activity which was
212 restored to wt levels upon expression of either *trkH* or *trkA* in *trans* (Fig. 2B). These data showed
213 that the reduction in *rsmB* expression, and consequent effect upon the downstream induction of
214 PCWDEs observed, were a consequence of the disruption of *trkH* and *trkA* by the transposon
215 insertion, and suggested that the Trk system could affect regulation of virulence in *P. wasabiae*. We
216 therefore investigated the ability of the selected *trk* mutants to cause tissue damage in potatoes. As
217 shown in Fig. 3, the mutant strains were impaired in virulence, showing an approximately 40%
218 reduction in the mass of macerated potato tuber tissue compared to that in tubers infected with wt
219 bacteria.

220

221 **Extracellular potassium levels influence *rsmB* expression**

222 The requirement for a functional Trk potassium uptake system for full activation of *rsmB*
223 expression indicated that induction of virulence might in fact be regulated by potassium. Therefore,
224 we analyzed *rsmB* expression levels in wt *P. wasabiae* grown in different concentrations of
225 potassium. Bacteria grown in 0.25, 2.5 and 25 mM potassium induced *rsmB* expression by
226 responding positively to increasing concentrations of potassium. However, in cells grown in 250
227 mM potassium, induction was as low as that observed in cells with 0.25 mM potassium (Fig. 4A).

228 To verify that this regulation was dependent upon the Trk system, we analyzed *rsmB*
229 expression in the *trkA* and *trkH* mutants cultured in the same range of potassium concentrations. As
230 neither mutant grew in the lowest concentration tested (0.25 mM, Fig. 4B), it was not possible to

231 determine the level of *rsmB* expression in these mutants under these conditions. Importantly, this
232 lack of growth shows that the Trk system is required for the growth of *P. wasabiae* in low
233 concentrations of potassium, supporting the predicted function of these genes in potassium uptake.
234 Furthermore, as shown in Fig. 4A, the two mutants yielded distinct phenotypes with regards to
235 potassium-dependent regulation of *rsmB* expression, in agreement with the putative functions
236 assigned by similarity to the Trk system in *E. coli*. In the *trkH* mutant, where based on knowledge
237 from *E. coli*, we would expect Trk-dependent potassium uptake to be absent, *rsmB* expression was
238 low when it was grown in the potassium concentrations tested. This result showed that no
239 potassium-dependent regulation of *rsmB* expression was observed upon disruption of the putative
240 Trk potassium channel protein. In contrast, *trkA* mutants retained some ability to induce *rsmB*
241 expression in response to changes in potassium availability, though 100-fold higher concentrations
242 (250 mM) were needed to reach the level of activation seen in the wt cultured with 2.5 mM
243 potassium. This suggests that Trk-dependent potassium uptake is less efficient in *P. wasabiae* *trkA*
244 mutants than in wt, similar to what was reported for *E. coli* (26). Together these phenotypes support
245 the prediction that these genes are part of a functional homologue of the Trk potassium transport
246 system in *P. wasabiae*, and these results show that this system is involved in the potassium
247 mediated regulation of RsmB.

248 Previous studies of the Gac/Rsm system in both pectobacteria and other bacterial species,
249 have established GacA as the key response regulator responsible for activating expression of *rsmB*.
250 To verify whether potassium-dependent regulation of RsmB occurs via this two component system,
251 we measured *rsmB* expression in mutants lacking *gacA* and determined the effect of potassium in
252 the absence of this response regulator. In line with current literature, disruption of *gacA* resulted in
253 reduced expression of *rsmB* compared to that of wt bacteria (Fig. 4A). Nonetheless, *rsmB*
254 expression in the *gacA* mutant could still be induced with intermediate levels of potassium, similar
255 to that observed in wt bacteria. Furthermore, deletion of *gacA* in a *trkA* mutant background (*gacA*
256 *trkA::Tn5* double mutant) resulted in the same potassium-dependent activation of *rsmB* expression

257 observed for the *trkA* single mutant. Therefore, potassium-dependent regulation of RsmB is not
258 mediated by the GacS/GacA signal transduction pathway.

259 Overall, our results demonstrate that *rsmB* expression is regulated by extracellular levels of
260 potassium and that this regulation requires the Trk system, but is independent of the GacS/GacA
261 system.

262

263 **Virulence in *P. wasabiae* is regulated by the extracellular concentration of potassium**

264 Next we tested whether the potassium-dependent effect on *rsmB* expression observed in the
265 *in vitro* studies described above had consequences for virulence *in vivo*, using the maceration of
266 potato tubers. We tested whether supplementation of the inoculum with different concentrations of
267 potassium affected the outcome of infection. In agreement with the *in vitro* results for *rsmB*
268 expression, high concentrations of extracellular potassium (250 mM) had an inhibitory effect on the
269 virulence of wt bacteria: less tissue maceration was observed in potatoes inoculated with cells
270 resuspended in buffer with 250 mM potassium compared to that of potatoes infected with bacteria
271 prepared in buffer supplemented with 4.5 mM potassium or no potassium (squares in Fig. 5). As for
272 the mutants, virulence of the *trkH* mutant was low at all potassium concentrations tested, while that
273 of the *trkA* mutant was also low at low potassium concentrations, though supplementation of high
274 concentrations of potassium (250 mM) resulted in an increase in virulence to near wt levels (Fig. 5).
275 The results for the *trkH* and *trkA* mutant strains are in full agreement with their respective
276 phenotypes for RsmB induction obtained *in vitro* (Fig. 4A).

277 Nonetheless, no increase in virulence of the wt bacteria occurred when potatoes were
278 inoculated with cells prepared in 4.5 mM potassium compared to those prepared without the
279 addition of potassium. However, bacterial inocula were grown in LB prior to infections, a medium
280 which contains approximately 8 mM potassium (15), and because our data showed that
281 concentrations higher than 2.5 mM were sufficient to induce *rsmB* expression in wt bacteria *in vitro*
282 (Fig 4), we reasoned that such induction might sustain *rsmB* expression, and therefore virulence,

283 during infection. To address this possibility, as well as to determine if extracellular potassium at the
284 site of infection could be essential to induce virulence, wt *P. wasabiae* was grown under non-
285 inducing conditions in Minimal Potassium Medium supplemented with low potassium (0.25 mM)
286 and then re-suspended in buffer with different potassium concentrations before inoculation into
287 potatoes. The *trkH* and *trkA* mutant strains were not tested under these conditions due to their lack
288 of growth in low concentrations of potassium (Fig. 4B). As shown in Fig. 6, wt bacteria grown in
289 0.25 mM caused some tissue maceration even when no potassium was added to the inoculum, but,
290 importantly, an increase in virulence was observed with the addition of 4.5 mM potassium (Fig. 6).
291 These results demonstrate that the addition of potassium at the time of infection induced virulence
292 in cells that were grown under non-inducing conditions. Again, high concentrations of potassium
293 (250 mM) inhibited the induction of virulence: the mass of macerated tissue was as low in these
294 conditions as when no potassium was added to the inoculum (Fig. 6).

295

296 Discussion

297 The small RNA, RsmB, has a major role in the signaling network controlling virulence in *P.*
298 *wasabiae* by preventing RsmA-mediated repression of PCWDEs expression. Consequently,
299 activation of *rsmB* transcription is essential for the production of these enzymes, and thus
300 unraveling the signals and mechanisms involved in this regulation is a key step in understanding the
301 environmental factors influencing virulence in *Pectobacterium* spp.

302 In a screen for regulators of RsmB, we isolated five mutants disrupted in genes involved in
303 the putative Trk potassium uptake system (*trkH* and *trkA*). We demonstrated that these mutants
304 presented a reduced *rsmB* expression and that the Trk system is important for production of
305 PCWDEs. These results led us to investigate the role of potassium in expression of *rsmB* and
306 virulence. We showed that intermediate concentrations of potassium (2.5 – 25 mM) were required
307 to induce RsmB, but high potassium concentrations (250 mM) inhibited expression of this
308 regulatory RNA. The conclusion that potassium and Trk participate in RsmB regulation is further

309 supported by the identification of another mutant isolated in our screen, that had the transposon
310 inserted in the third gene of the annotated *sapABCDF* operon (RSV238, Table S3). This operon, in
311 particular SapD, has been implicated in potassium transport via the Trk system in *E. coli*, as
312 mutants in *sapD* present no potassium uptake by the Trk system (27). It is thus possible that a
313 transposon insertion in *sapC* also affects the Trk system.

314 Our analysis of the *trk* mutants showed that in *P. wasabiae* the Trk potassium system
315 functions in a similar way to the *E. coli* Trk system. However, although in *E. coli* the Trk system
316 seems to be important mainly at intermediate concentrations of potassium, in *P. wasabiae* this
317 system appeared to be relevant in a broader range of potassium concentrations. In *E. coli* an
318 inducible high affinity potassium system, the Kdp system, is the major responsible for potassium
319 uptake at concentrations lower than 5mM [reviewed in (28)], but, in *P. wasabiae* the *trk* mutants
320 had a strong growth defect at low potassium concentration (2.5 mM) and did not even grow at 0.25
321 mM. These results showed that the Trk system is important at low potassium concentrations. This,
322 together with the fact that we could not find any *kdp*-like gene in the genome of *P. wasabiae*
323 indicates that *P. wasabiae*, in contrast to *E. coli*, might lack a high affinity potassium transporter
324 and relies solely on Trk at low potassium concentrations.

325 Our results showing that the Trk system regulated *rsmB* expression provided strong support
326 for the hypothesis that extracellular levels of potassium were important for virulence in *P.*
327 *wasabiae*. Accordingly, we observed that extracellular potassium was required to fully induce
328 maceration of plant tissue in wt and that this induction required an intact Trk system. Moreover, we
329 observed that virulence was inhibited at high extracellular concentrations of potassium (250 mM).
330 Importantly, this inhibition of virulence in wt at a concentration of 250 mM potassium is unlikely to
331 be a consequence of growth inhibition since *in vitro*, at this potassium concentration, wt still grows
332 better than the *trkA* mutant. In addition, the *trkA* mutant despite its growth defect, with 250 mM
333 potassium can cause almost as much tissue maceration as the maximal levels observed with the wt
334 (Fig. 4E, Fig. 5 and Table S4). We also determined that supplementation of high concentration of

335 potassium had no effect on the viability of the inoculum applied to the potato tubers in any of
336 strains tested, as shown in Fig. S2.

337 The mechanism by which the extracellular levels of potassium regulate *rsmB* expression via
338 the Trk system is still unclear. As *rsmB* transcription is activated by the GacS/GacA system, we
339 investigated whether the observed effects of potassium were also linked with or dependent upon the
340 function of this two component system. Though disruption of the *gacA* gene decreased the extent to
341 which *rsmB* expression was induced, surprisingly potassium- and Trk-dependent regulation was
342 still observed in a *gacA* mutant. Two additional regulators in *Pectobacterium* spp., KdGR and
343 RsmC, have been shown to repress RsmB expression (7, 29). However, again disruption of these
344 regulators had no effect in potassium-dependent regulation of *rsmB* expression (data not shown).
345 These results provide evidence for additional players in this regulation of RsmB. The identity of
346 such regulators could come from a genetic approach to isolate mutants that no longer respond to
347 changes in extracellular levels of potassium. For example, it is possible that an additional two-
348 component system is involved in such regulation. In fact, some transport systems have been
349 associated with the activation of two-component systems [reviewed in (30)]. In *E. coli*, the
350 DcuS/DcuR two-component system which is activated by C4-carboxylates in the periplasm is also
351 regulated by the DcuB antiporter that uptakes C4-carboxylates. It has been proposed that in the
352 absence of C4-carboxylates, protein-protein interactions between DcuB and DcuS repress the
353 autophosphorylation activity of the DcuS sensor kinase. Upon transport of these compounds, this
354 repression is absent presumably because DcuB releases DcuS which can then be activated by the
355 periplasmic levels of C4-dicarboxylates (31). A similar interaction might be happening between the
356 Trk transport proteins and an unknown two-component system to regulate *rsmB*. Alternatively, it is
357 possible that the observed regulation of *rsmB* is not responding to potassium flux through the Trk
358 system but instead it is sensitive to changes in intracellular potassium concentrations, as it has been
359 shown for the induction of biofilm formation in *Bacillus subtilis* as a response to potassium leakage
360 (32). In this case it was proposed that in the presence of natural compounds that cause potassium

361 leakage, the membrane kinase KinC responds to transient decreases in cytoplasmic potassium
362 concentrations, activating a phosphorylation cascade that results in induction of exopolysaccharides
363 production and biofilm formation.

364 The Trk system is widespread among bacteria and it is the potassium transporter most
365 commonly found in the largest number of species. This transport system is crucial for many of the
366 intracellular functions of potassium, such as maintenance of cell turgor pressure, regulation of
367 intracellular pH, adaptation to osmotic stress and function of many cytoplasmic enzymes that
368 require potassium [reviewed in (28)], but importantly it has also been implicated in regulation of
369 virulence in diverse bacteria. Trk mutants are impaired in virulence in *Vibrio vulnificus*, *Salmonella*
370 *enterica enterica* serovar Typhimurium and *Francisella tularensis*, however the molecular
371 mechanisms involved in such regulation have not been identified (15, 33, 34). As both *V. vulnificus*
372 and *S. enterica* also use homologues of the Rsm system (Csr) to regulate virulence, it is conceivable
373 that Trk regulation of virulence in these organisms might also take place through the regulatory
374 RNAs of the Csr system. Therefore, it is possible that the mechanisms identified here are conserved
375 among the other pathogens that have the Gac/Rsm system. As the Gac/Rsm system has been shown
376 to modulate carbon fluxes (10, 11) we propose that in bacteria that use the Trk transporter to
377 regulate components of the Gac/Rsm pathway cells might benefit from coupling the information
378 obtained from perceiving changes in potassium concentrations with the information on the
379 metabolic state of the cell, to modulate functions that go beyond maintaining the physiological
380 functions of potassium to control activities that contribute to host colonization. For instance, when
381 short-chain fatty acids accumulate during fermentation, bacterial cells typically use potassium
382 transporters to manipulate intracellular potassium levels, to cope with changes in cytoplasmic pH
383 and to control turgor pressure [reviewed in (35)]. It is interesting that the Gac/Rsm homologue
384 system in *E. coli* has been shown to respond under certain conditions to these weak organic acids
385 (12) and thus, it is tempting to speculate that the link between the regulation of potassium transport
386 with the regulation of the Gac/Rsm system might be related to the need to adapt to the presence of

387 short-chain fatty acids. The benefit of regulating virulence in response to changes in potassium
388 concentrations might also be related to the environmental changes typically experienced by
389 pathogens during host invasion. For example, plant pathogens, such as *P. wasabiae*, are typically
390 found in the soil where potassium concentrations are in the 10-100 μM range, contrary to the 100
391 mM concentration found in eukaryotic cells (36). Hence, upon arrival to a wounded host, the local
392 increased potassium concentration might provide a cue for the activation of the production of
393 PCWDEs. The action of these enzymes will further disrupt the host cells with the consequent
394 leakage of intracellular potassium, and the bacteria will have to adapt to the increasing potassium
395 levels which can ultimately reach the levels found inside the eukaryotic plant cells.

396 Crop losses resulting from bacterial diseases remain significant agricultural and economic
397 concerns. Understanding the regulatory networks responsible for the activation of virulence genes
398 will help to define and improve control strategies that target this problem. The results presented
399 here identify potassium as an important environmental factor in the regulation of virulence in the
400 plant pathogen *P. wasabiae* and show that a reduction in wt virulence can be achieved by increasing
401 the extracellular concentration of potassium (Fig. 5 and Fig. 6). Although additional work is
402 required to fully characterize the molecular mechanisms behind the regulation of virulence by
403 potassium via RsmB, our study highlights how potassium levels in the soil could affect the outcome
404 of virulence in the plant pathogen *P. wasabiae*.

405

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507

508 **Legends**

509 **Fig. 1. Trk mutants have low *rsmB* expression.** (A) Expression of $P_{rsmB}::gfp$ promoter fusion from
510 pRSV206 in *P. wasabiae* wt, RSV108 (*trkH*::Tn5), RSV141 (*trkH*::Tn5), RSV154 (*trkH*::Tn5),
511 RSV124 (*trkA*::Tn5) or RSV236 (*trkA*::Tn5) mutant strains was measured by fluorescence flow
512 cytometry of cells grown for 24 h in MM. Error bars represent SD, n = 3.

513

514 **Fig. 2. Complementation of the *trkH* and *trkA* mutants.** (A) Expression of a $P_{rsmB}::gfp$ promoter
515 fusion was measured in wt harboring the control vector (pOM18), the *trkH*::Tn5 mutant carrying
516 either the pOM18 empty vector or the vector expressing *trkH* ($p(trkH^+)$), and the *trkA*::Tn5 mutant
517 carrying either the pOM18 empty vector or the vector expressing *trkA* ($p(trkA^+)$). Fluorescence was
518 measured in cells collected from cultures grown to $OD_{600} = 0.4$ in MM. (B) Pel activity was
519 measured in cell-free supernatants from cultures of the bacterial strains indicated above grown in
520 LB supplemented with PGA at $OD_{600} = 0.4$. Error bars represent SEM, $n = 6$.

521

522 **Fig. 3. Mutants in the Trk system are impaired in virulence.** Virulence of wt (squares),
523 *trkH*::Tn5 (circles) and *trkA*::Tn5 (triangles) was measured by quantification of the mass of potato
524 tuber maceration induced by these bacteria 48 h after inoculation of the potato tubers. Potatoes were
525 inoculated with approximately 3×10^5 cells of the respective strain grown overnight in LB. $n = 7$,
526 **P-value < 0.01. This is a representative experiment from three independent experiments.

527

528 **Fig. 4. Effect of extracellular potassium on *rsmB* expression and growth in *P. wasabiae*.** (A)
529 The expression of the *rsmB* promoter fusion ($P_{rsmB}::gfp$) was measured by flow cytometry of
530 cultures of wt, *trkA*::Tn5 and *trkH*::Tn5, *gacA* and *gacA trkA*::Tn5 grown to $OD_{600} = 0.3$ in Minimal
531 Potassium Media supplemented with a final potassium concentration of 0.25 mM, 2.5 mM, 25 mM
532 or 250 mM. NG stands for no growth. Error bars represent SEM, $n = 6$. (B – D) OD_{600} was
533 measured throughout growth for wt (squares), *trkH*::Tn5 (circles), *trkA*::Tn5 (triangles) in Minimal
534 Potassium Media supplemented with (B) 0.25 mM, (C) 2.5 mM, (D) 25 mM or (E) 250 mM KCl
535 (growth rates in Table S4).

536

537 **Fig. 5. Regulation of virulence by extracellular potassium concentration.** Virulence of wt
538 (squares), *trkH*::Tn5 (circles) and *trkA*::Tn5 (triangles) was measured by quantification of the mass
539 of macerated tissue 48 h after inoculation of potato tubers. Cells cultured overnight in LB

540 (approximately of 8 mM potassium) were harvested and re-suspended in potassium-free PBS (wt
541 only, white squares), or PBS supplemented with a final concentration of either 4.5 mM (grey) or
542 250 mM (black) potassium. Potatoes were inoculated with approximately 3×10^5 cells of the
543 respective strain at the different potassium concentrations. Error bars represent SEM, $n = 6$, **P-
544 value < 0.01 , ^{ns}Not significant. This is a representative experiment from three independent
545 experiments.

546

547 **Fig. 6. Induction of virulence by extracellular potassium.** Virulence of wt bacteria was measured
548 by determining the mass of damaged tissue 48 h after infection of potato tubers. Potatoes were
549 inoculated with approximately 3×10^5 cells of the respective strain grown overnight in Minimal
550 Potassium Media supplemented with a final potassium concentration of 0.25 mM and re-suspended
551 in potassium-free PBS (0 mM, white), in PBS (4.5 mM of potassium, grey) or in PBS supplemented
552 with potassium to a final concentration of 250 mM (black). Error bars represent SEM, $n = 6$, **P-
553 value < 0.01 . This is a representative experiment from two independent experiments.

554











