



## 2-Tridecanone impacts surface-associated bacterial behaviours and hinders plant-bacteria interactions

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1 **2-Tridecanone impacts surface-associated bacterial behaviours and hinders plant-**  
2 **bacteria interactions**

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22 Running title: 2-tridecanone: a new chemical cue for bacteria

23 **Originality-Significance Statement:** Our work assigns new roles to the volatile  
24 methylketone 2-tridecanone (2-TDC), which was known previously as a natural  
25 insecticide. We demonstrate that 2-TDC is an airborne cue that affects surface motility  
26 and biofilm formation in phylogenetically distant bacteria, including plant and animal  
27 pathogens, without any effect on growth. Moreover, we found that 2-TDC protects  
28 plants from bacterial infections. These observations open new perspectives not only for  
29 fundamental research, but also for the development of biotechnological solutions to  
30 prevent/control bacterial infections, minimizing the risk of generating antimicrobial  
31 resistance.

32

### 33 **Summary**

34 Surface motility and biofilm formation are behaviours which enable bacteria to infect  
35 their hosts and are controlled by different chemical signals. In the plant symbiotic  
36 alpha-proteobacterium *Sinorhizobium meliloti*, the lack of long-chain fatty acyl-  
37 coenzyme A synthetase activity (FadD) leads to increased surface motility, defects in  
38 biofilm development, and impaired root colonization. In this study, analyses of lipid  
39 extracts and volatiles revealed that a *fadD* mutant accumulates 2-tridecanone (2-TDC), a  
40 methylketone known as a natural insecticide. Application of pure 2-TDC to the wild-  
41 type strain phenocopies the free-living and symbiotic behaviours of the *fadD* mutant.  
42 Structural features of the methylketone determine its ability to promote *S. meliloti*  
43 surface translocation, which is mainly mediated by a flagella-independent motility.  
44 Transcriptomic analyses showed that 2-TDC induces differential expression of iron  
45 uptake, redox, and stress-related genes. Interestingly, this methylketone also influences  
46 surface motility and impairs biofilm formation in plant and animal pathogenic bacteria.  
47 Moreover, 2-TDC not only hampers alfalfa nodulation but also the development of

48 tomato bacterial speck disease. This work assigns a new role to 2-TDC as an  
49 infochemical that affects important bacterial traits and hampers plant-bacteria  
50 interactions by interfering with microbial colonization of plant tissues.

51

## 52 **Introduction**

53 The behaviour of bacteria on surfaces is crucial for host-microbe interactions. For  
54 successful host colonization, the ability to spread over and colonize preferred niches as  
55 well as to establish a long-lasting community, are important attributes. Different  
56 mechanisms are used by bacteria to translocate over surfaces (Harshey, 2003; Jarrell  
57 and McBride, 2008). Swarming, twitching and gliding are active bacterial motilities  
58 mediated by flagella, type IV pili and focal adhesion complexes, respectively. Bacteria  
59 can also spread on surfaces using sliding, a passive translocation mechanism in which  
60 surfactants and other compounds that diminish friction between cells and surfaces play  
61 a crucial role (Hölscher and Kovács, 2017). Amongst the different types of surface  
62 translocation, swarming motility has been the most extensively studied because it is a  
63 trait closely connected with virulence in pathogenic bacteria (Verstraeten *et al.*, 2008).  
64 Swarming requires flagellar action as does the well-known swimming motility. The  
65 latter is the movement of individual bacteria in liquid environments. Unlike swimming,  
66 swarming is a coordinated multicellular migration of bacteria on top of solid surfaces  
67 (Kearns, 2010). Several studies have reported an intimate link between swarming and  
68 the formation of surface-attached communities of bacteria known as biofilms (O'Toole  
69 *et al.*, 2000). Both processes are multicellular surface-associated behaviours which  
70 confer resistance to different stressors. In addition, they are often controlled by the same  
71 regulatory pathways (Verstraeten *et al.*, 2008; Prüß, 2017), and influenced by the same  
72 chemical cues, which include quorum sensing signals and volatile compounds.

73 Quorum sensing (QS) is a cell-to-cell communication system that allows bacteria to  
74 collectively modify behaviours by regulating gene expression in response to the  
75 population density (Papenfort and Bassler, 2016; Whiteley *et al.*, 2017). It depends on  
76 the synthesis and detection of different extracellular chemical signals. *N*-  
77 acylhomoserine lactones (AHL), *cis*-2-unsaturated fatty acids of the Diffusible Signal  
78 Factor (DSF) family or  $\alpha$ -hydroxyketones are some of the QS signals recognized to  
79 control motility and biofilm formation (Papenfort and Bassler, 2016). More recently,  
80 several volatile compounds emitted by bacteria have also been found to impact motility  
81 and biofilm formation with different effects depending on the bacterial species or the  
82 volatile concentration (Audrain *et al.*, 2015).

83 Swarming and biofilm formation occurs also in beneficial microorganisms such as  
84 rhizobia. These soil bacteria can induce the formation of nitrogen-fixing nodules in  
85 legume plant roots following a complex molecular interchange (Gibson *et al.*, 2008;  
86 Oldroyd and Downie, 2008). Data about genetic determinants and environmental cues  
87 that control surface motility and biofilm formation in rhizobia, as well as the role that  
88 these two traits play in the plant-microbe interaction, are scarce. Swarming motility has  
89 been reported for the alfalfa symbiont *Sinorhizobium meliloti* strains GR4 and Rm1021,  
90 although the latter can also translocate over surfaces using a type of flagellum-  
91 independent motility, which requires the amphiphilic siderophore rhizobactin 1021  
92 (Nogales *et al.*, 2010; Nogales *et al.*, 2012; Bernabéu-Roda *et al.*, 2015). Recent studies  
93 performed with these rhizobial strains revealed the existence of mechanisms which  
94 regulate swarming and biofilm formation inversely to promote efficient plant root  
95 colonization (Amaya-Gómez *et al.*, 2015). One of these mechanisms involves the fatty  
96 acid catabolism-related enzyme FadD. In *S. meliloti*, *fadD* inactivation promotes surface  
97 motility, obstructs normal biofilm development, and impairs colonization and

98 nodulation efficiency on alfalfa roots (Soto *et al.*, 2002; Nogales *et al.*, 2010; Amaya-  
99 Gómez *et al.*, 2015; Bernabéu-Roda *et al.*, 2015). FadD is an acyl-coenzyme A (CoA)  
100 synthetase which converts long-chain fatty acids into acyl-CoAs which can enter  $\beta$ -  
101 oxidation (Rock, 2008). Recently, FadD has been associated with the turnover of QS  
102 signals of the DSF family (Zhou *et al.*, 2015; Zhou *et al.*, 2017).

103 The phenotypes exhibited by *S. meliloti fadD* mutants led us to hypothesize that  
104 lipidic compounds differentially produced in the wild-type and mutant strains, may act  
105 as cues that control bacterial behaviours on surfaces and influence the interaction with  
106 the plant host. To identify such a compound, in a previous work, the lipidic composition  
107 of *S. meliloti* wild-type and *fadD* mutant strains was investigated. It was determined that  
108 the *fadD* mutant accumulated significant amounts of free fatty acids in the stationary  
109 phase of growth (Pech-Canul *et al.*, 2011). However, none of the fatty acids analysed  
110 was able to induce surface motility of the wild-type strain. In the present study, we  
111 found that the *fadD* mutant accumulates 2-tridecanone (2-TDC), and that  
112 overproduction of this compound might be responsible for the phenotypes exhibited by  
113 the mutant strain. 2-TDC is a volatile methylketone (MK) known as a natural  
114 insecticide that is produced in prodigious amounts by specialized trichomes of wild  
115 tomato plants (*Solanum habrochaites* ssp *glabratum*), which makes these plants  
116 naturally resistant to a wide spectrum of arthropod pests (Williams *et al.*, 1980; Fridman  
117 *et al.*, 2005). Production of 2-TDC has also been reported for several bacterial species  
118 (Forney and Markovetz, 1971; Elgaali *et al.*, 2002; Blom *et al.*, 2011; Lemfack *et al.*,  
119 2014; Raza *et al.*, 2016), but not for rhizobia. More importantly, the impact of this  
120 methylketone on bacterial behaviour and on the establishment of plant-bacteria  
121 interactions has not been hitherto described. Here, we provide evidence for a new role  
122 of 2-TDC that affects bacterial lifestyles.

123

124 **Results**125 *A Sinorhizobium meliloti fadD mutant accumulates 2-tridecanone*

126 To identify the putative FadD-related compound responsible for the pleiotropic  
127 phenotype of a *S. meliloti fadD* mutant, lipidic extracts obtained from cells and spent  
128 culture media of *S. meliloti* wild-type GR4 and the *fadD* mutant QS77 were analysed by  
129 Gas Chromatography/Mass Spectrometry (GC/MS). Besides the already reported  
130 accumulation of free fatty acids (Pech-Canul *et al.*, 2011), the methylketone (MK) 2-  
131 tridecanone (2-TDC) and the aldehyde dodecanal, were detected in the spent media  
132 from the *fadD* mutant (Supporting Information Table S1 and Fig. S1 and S2). Peaks  
133 corresponding to dodecanal or 2-TDC were not detected in the spent media obtained  
134 from the wild-type strain or from the *fadD* mutant QS77 complemented with the *fadD*-  
135 harbouring plasmid pBBRD4. Dodecanal and 2-TDC were present in similar amounts  
136 and each represented about 3% of the total lipidic compounds detected in the spent  
137 media from the *fadD* mutant (Supporting Information Table S1).

138 Since dodecanal and 2-TDC are known as volatile compounds, we decided to also  
139 analyse by Solid Phase MicroExtraction (SPME)-GC/MS the compounds emitted into  
140 the headspace by the wild-type and *fadD* mutant strains grown on MM (1% agar).  
141 Whereas dodecanal was absent in these analyses, peaks corresponding to three  
142 methylketones (2-TDC, 2-pentadecanone, and 2-heptadecanone) were identified in both  
143 strains, albeit at very low levels and close to the detection limit in the case of the wild-  
144 type. We specifically focused on 2-TDC since this was also detected in the analyses of  
145 lipidic extracts from liquid cultures. As shown in Fig. 1, the peak area corresponding to  
146 2-TDC in the *fadD* mutant was 7.6-fold larger than that of the wild-type strain  
147 (49021±7532 vs 6411±1793 counts). These results indicate that *S. meliloti* produces

148 volatile 2-TDC upon growth on semisolid surfaces and that the absence of FadD activity  
149 increases production of the MK, both in liquid and surface-grown cultures.

150

### 151 *2-TDC promotes surface migration and reduces biofilm formation in S. meliloti*

152 In order to investigate whether accumulation of 2-TDC in the *fadD* mutant could be  
153 responsible for the phenotypes exhibited by this strain, the behaviour of *S. meliloti* wild-  
154 type strains in response to the application of 2-TDC was analysed. Firstly, possible  
155 effects of the MK on bacterial growth were tested. It was found that *S. meliloti* cannot  
156 use 2-TDC as the sole carbon source (Supporting Information Fig. S3). Moreover,  
157 bacterial growth rates in liquid MM remained unaffected in the presence of different  
158 concentrations of 2-TDC up to 500  $\mu\text{M}$  (Supporting Information Fig. S4A). In contrast  
159 to the lack of swarming-inducing activity observed for several fatty acids (Pech-Canul  
160 *et al.*, 2011), addition of 2-TDC into semisolid MM at 5  $\mu\text{g/ml}$  (25  $\mu\text{M}$ ) or 10  $\mu\text{g/ml}$   
161 (50  $\mu\text{M}$ ), which correspond to total amounts of 0.5 or 1  $\mu\text{mol}$  in the medium,  
162 respectively, induced surface translocation in the wild-type strain GR4. Interestingly,  
163 the same effect was observed in response to airborne 2-TDC generated from solutions  
164 applied onto the lid of the Petri dish. The stimulatory effect was more pronounced using  
165 less permissive conditions for surface migration MM (1% agar) (Bernabéu-Roda *et al.*,  
166 2015). Airborne 2-TDC generated from solutions containing from 0.2 to 10  $\mu\text{mol}$  of the  
167 MK was able to stimulate surface motility with similar dose-dependent effects on GR4  
168 and Rm1021 (Fig. 2A), two *S. meliloti* strains which exhibit different modes of surface  
169 translocation (Bernabéu-Roda *et al.*, 2015). In contrast, swimming motility remained  
170 unaffected in the presence of 2-TDC (Supporting Information Fig. S4B), indicating that  
171 2-TDC acts by specifically activating surface translocation in *S. meliloti*.



172 Next, we investigated whether 50  $\mu\text{M}$  2-TDC could also impact the ability of *S.*  
173 *meliloti* to form surface-attached communities. The defect shown by *S. meliloti fadD*  
174 mutants on biofilm development on glass surfaces has been shown previously (Amaya-  
175 Gómez *et al.*, 2015) and was exhibited after long periods of incubation. Due to the  
176 volatile nature of 2-TDC, its effect was analysed at earlier stages of biofilm formation  
177 (3 days). Under control conditions and as previously reported (Amaya-Gómez *et al.*,  
178 2015), GR4 and Rm1021 developed flat, unstructured biofilms that covered most of the  
179 glass surface. In the presence of 2-TDC, biofilm formation in both *S. meliloti* strains  
180 was significantly impaired and significant reductions in surface area colonization and  
181 biofilm mass were observed (Figure 2B and Supporting Information Fig. S5A). These  
182 data show that 2-TDC affects surface-associated behaviours in *S. meliloti*, without  
183 affecting bacterial growth.

184

#### 185 *2-TDC impairs nodulation of alfalfa plants*

186 The effects caused by exogenous application of 2-TDC on surface translocation and  
187 biofilm formation in *S. meliloti* wild-type strains suggested the possibility that the  
188 phenotypes exhibited by the *fadD* mutant were triggered by 2-TDC accumulation. To  
189 investigate whether 2-TDC also accounts for the symbiotic deficiency demonstrated by  
190 this mutant, the nodulation kinetics of alfalfa plants inoculated with the *S. meliloti* wild-  
191 type strain GR4 were determined in the absence or presence of 2-TDC added to the  
192 plant mineral solution at the time of inoculation. Compared with control conditions, the  
193 application of 2-TDC led to an important delay (2-3 days) in the appearance of the first  
194 nodules and to a significant reduction in the number of nodules formed per plant, an  
195 effect that was dose-dependent (Fig. 3A).

196 Impaired nodulation of alfalfa roots was also detected when plants were treated with  
197 2-TDC days before inoculation but also when the MK was applied 2 days after rhizobial  
198 inoculation (Fig. 3B and Supporting Information Fig. S6A-C). However, no significant  
199 effect was observed when 2-TDC was applied 4 days after rhizobial inoculation (Fig.  
200 3B and Supporting Information Fig. S6D) indicating that 2-TDC adversely interferes in  
201 the early stages of the *Rhizobium*-legume interaction. Nodulation inhibition caused by  
202 2-TDC cannot be explained by either a direct bacteriostatic or bactericidal effect  
203 (Supporting Information Fig. S4A) or impaired nodulation (*nod*) gene expression  
204 (Supporting Information Table S2). Nevertheless, 2-TDC was found to negatively  
205 interfere with the bacterial ability to efficiently colonize plant roots (Fig. 3C). The  
206 difference in colonization ability was especially remarkable 24 h postinoculation when  
207 the presence of 2-TDC reduced by 100-fold the number of bacteria that were associated  
208 to roots compared to control plants. This effect, which is in line with the inhibition of  
209 biofilm formation caused by 2-TDC, might account for the interference exerted by the  
210 MK on the *Rhizobium*-legume symbiosis.

211

212 *Structural features of 2-TDC are important to promote surface motility in S. meliloti*

213 To evaluate whether the effect caused by 2-TDC on *S. meliloti* surface translocation is a  
214 general property of aliphatic ketones, nine different compounds, which varied from 2-  
215 TDC either in the acyl chain length or in the position of the carbonyl group, were  
216 applied at three different amounts and tested in surface motility assays with strain GR4  
217 (Fig. 4A). In contrast to 2-TDC, treatments with 3-tridecanone (3-TDC), 4-tridecanone  
218 (4-TDC), or 7-tridecanone (7-TDC) did not promote surface translocation, indicating  
219 that the position of the carbonyl group is crucial for biological activity. Our data  
220 indicate that the acyl chain length is another structural attribute that influences

221 biological activity of MKs. With the exception of 2-heptanone (2-Hp), which did not  
222 affect surface motility at any of the amounts tested, the different MKs used in our assay  
223 exhibited certain motility-promoting activity. In general, biological activity decreased  
224 with MKs of shorter acyl chain length. For a given MK, the activity increased when  
225 raising the concentration at which it was applied. A remarkable exception to this rule  
226 was 2-pentadecanone (2-PDC), whose activity at the lowest amount tested (0.2  $\mu\text{mol}$ )  
227 was even higher than that obtained with 1  $\mu\text{mol}$  2-TDC (Fig. 4A). Significant promotion  
228 of GR4 surface motility ( $24.9 \pm 3$  and  $8.8 \pm 0.6$  mm) was still observed in response to  
229 40 nmol and 20 nmol 2-PDC, respectively, amounts at which 2-TDC did not show any  
230 effect.

231 Interestingly, a competitive interaction was observed between 2-TDC and 7-TDC.  
232 GR4 surface translocation promoted by 2-TDC decreased proportionally as the amount  
233 of 7-TDC applied to the assay increased, being completely abolished by the highest  
234 amount of 7-TDC used (Fig. 4B). Altogether, these data suggest that 2-TDC triggers  
235 surface motility in *S. meliloti* upon specific recognition.

236

237 *Exogenous application of 2-TDC promotes surface migration in S. meliloti by*  
238 *stimulating flagella-dependent and independent mechanisms*

239 We sought to investigate the mechanism by which 2-TDC specifically promotes surface  
240 migration in *S. meliloti*. It is known that inactivation of the *fadD* gene promotes surface  
241 motility in *S. meliloti* GR4 and Rm1021 by stimulating a flagella-independent  
242 mechanism exhibited by the surface spreading displayed by the flagella-less  
243 GR4fadDflaAB and 1021fadDflaAB strains (Bernabéu-Roda *et al.*, 2015; Fig. 5). We  
244 found that application of volatile 2-TDC promoted surface motility not only in GR4 and  
245 Rm1021 but also in the flagella-less derivative strains GR4flaAB and 1021flaAB (Fig.

246 5A and 5B, respectively). Therefore, like FadD loss-of-function, application of volatile  
247 2-TDC triggers a flagella-independent surface translocation in *S. meliloti*. Addition of 2-  
248 TDC had a greater influence on promoting surface motility in GR4 and Rm1021 than  
249 inactivation of the *fadD* gene in the corresponding genetic backgrounds, suggesting a  
250 concentration-dependent effect of the MK.

251 To the best of our knowledge, the only flagellum-independent surface migration  
252 described in *S. meliloti* requires production of rhizobactin 1021 (Rhb1021), a lipid-  
253 containing siderophore with surfactant properties (Nogales *et al.*, 2010, 2012;  
254 Bernabéu-Roda *et al.*, 2015). Rhb1021 is produced by *S. meliloti* strain Rm1021 in  
255 response to low iron concentrations but is not produced by strain GR4 (Nogales *et al.*,  
256 2010), which explains the different surface motility behaviour of the two strains on  
257 semisolid MM. The defect in surface translocation exhibited by a siderophore-defective  
258 mutant (1021rhbD) is restored when the *fadD* gene is inactivated and also with  
259 application of volatile 2-TDC (Fig. 5B). These results demonstrate that the surface  
260 motility triggered by 2-TDC is not mediated by Rhb1021 siderophore.

261 Data shown in Fig. 5 indicate that application of 2-TDC induces mainly a flagella-  
262 independent surface translocation in *S. meliloti*. Nevertheless, the increased surface  
263 migration exhibited by wild-type strains compared to their corresponding non-  
264 flagellated strains in response to 2-TDC suggests that flagella might contribute to the  
265 enhanced motility triggered by 2-TDC. In a previous study, we found that expression of  
266 the flagellin *flaA* gene was higher in the *fadD* mutant than in the wild-type strain GR4  
267 under swarming-inducing conditions (Soto *et al.*, 2002). Here, we investigated whether  
268 2-TDC could have a similar effect on *flaA* gene expression by analysing the  $\beta$ -  
269 galactosidase activity of GR4 cells harbouring a *flaA::lacZ* transcriptional fusion and  
270 grown in MM (1% agar) in the absence or presence of volatile 2-TDC. In parallel,

271 expression of the kanamycin promoter was also tested in cells carrying a *km::lacZ*  
272 transcriptional fusion, in order to detect unspecific changes in gene expression caused  
273 by 2-TDC treatment. No differences in the expression of the *lacZ* gene driven by the  
274 kanamycin promoter were detected between cells grown in the presence ( $559 \pm 60$   
275 Miller Units) and in the absence ( $432 \pm 75$  Miller Units) of 2-TDC. In contrast,  
276 expression of the *flaA::lacZ* fusion was 2.6-fold higher in cells exposed to volatile 2-  
277 TDC than that exhibited by cells grown under control conditions ( $29085 \pm 3296$  versus  
278  $11169 \pm 2176$  Miller Units; ANOVA test  $P \leq 0.05$ ). These results indicate that, like FadD  
279 loss-of-function, 2-TDC promotes surface translocation in *S. meliloti* by stimulating  
280 both flagella-dependent and independent mechanisms.

281

#### 282 *Genome wide analysis of 2-TDC-regulated genes in S. meliloti*

283 To gain insights into the molecular mechanisms responsible for the effects caused by 2-  
284 TDC in *S. meliloti*, the transcriptomes of Rm1021 cells grown in broth and semisolid  
285 MM in the absence or presence of 2-TDC were compared. While no differential gene  
286 expression was detected in response to 2-TDC treatment in *S. meliloti* cells grown in  
287 liquid media, 88 genes altered their transcription levels when cells were grown on the  
288 surface of semisolid media exposed to volatile 2-TDC (49 genes up-regulated, 39 genes  
289 down-regulated; Supporting Information Table S3). Previously, we verified that volatile  
290 2-TDC treatment did not cause any effect on growth.

291 The up-regulation of up to 9 genes coding for proteins with putative functions in  
292 redox reactions was remarkable. Indeed, the two most up-regulated genes in response to  
293 2-TDC code for oxidoreductases: *smb20343* coding for a putative isoquinoline 1-  
294 oxidoreductase, and *azoR* (*smc01329*) which codes for a predicted FMN-dependent

295 NADH-azoreductase, an enzyme involved in resistance to thiol-specific stress in  
296 *Escherichia coli* (Liu *et al.*, 2009). In addition to oxidoreduction-related genes, the  
297 presence of 2-TDC increased the expression of 6 genes involved in the synthesis of the  
298 amino acids glutamate, methionine and cysteine, and 5 genes coding for transcriptional  
299 regulators that might control the expression of drug resistance genes. Interestingly, 56%  
300 of the genes down-regulated in response to 2-TDC treatment (22 genes) were related to  
301 iron uptake and metabolism, including siderophore Rhb1021 biosynthesis genes. These  
302 microarray data, which were validated by RT-qPCR (Supporting Information Table S4),  
303 suggest that volatile 2-TDC reduces iron uptake while inducing stress-related responses.

304

305 *Surface-associated behaviours in plant and animal pathogenic bacteria are also*  
306 *influenced by 2-TDC*

307 Different airborne chemical signals have been implicated in interspecific bacterial  
308 signalling, regulating important microbial behaviours such as motility, biofilm  
309 formation, drug resistance or virulence (Kim *et al.*, 2013; Audrain *et al.*, 2015; Schmidt  
310 *et al.*, 2015). To test if 2-TDC could also influence the behaviour of bacteria other than  
311 *S. meliloti*, the surface motility phenotypes of several plant and animal pathogenic  
312 bacteria were assessed in the presence of the MK. As shown in Fig. 6A, volatile 2-TDC  
313 increased surface motility in the plant pathogens *Pseudomonas syringae* pathovars  
314 *syringae* (Pss) and *tomato* (Pst). In contrast, 2-TDC inhibited and reduced swarming  
315 motility in *Salmonella enterica* serovar Typhimurium and *Pseudomonas aeruginosa*,  
316 respectively (Fig. 6B). The effect of swarming reduction in *P. aeruginosa* was not  
317 caused by alterations in rhamnolipid production (Supporting Information Fig. S7).  
318 Furthermore, twitching motility was also significantly inhibited by increasing

319 concentrations of 2-TDC in *P. aeruginosa* (Fig. 6C), suggesting that type IV pili may be  
320 affected by this molecule. However, no changes in *pilA* gene expression were observed  
321 in response to 2-TDC (Supporting Information Fig. S8). As in *S. meliloti*, no effect of 2-  
322 TDC on bacterial growth rate or swimming motility was detected for any of the  
323 pathogenic bacteria used in this study.

324 We also examined the influence of 2-TDC on biofilm formation in *P. syringae* and  
325 *P. aeruginosa*. The presence of 2-TDC in the culture medium significantly decreased  
326 the ability of the bean and soybean pathogen Pss BT111 to develop biofilm firmly  
327 attached to the glass surface (Fig. 7A). Likewise, static biofilms developed on glass  
328 surfaces by *P. aeruginosa* were significantly inhibited by 2-TDC (Fig. 7B and  
329 Supporting Information Fig. S5B). Interestingly, 2-TDC did not have an apparent  
330 impact on cyclic dimeric guanosine monophosphate (c-di-GMP) or exopolysaccharide  
331 levels when *P. aeruginosa* was grown on LB or NB agar plates using a transcriptional  
332 reporter fusion to *cdrA* (regulated by c-di-GMP), or translational fusions to *pelA* and  
333 *pslA* (exopolysaccharide biosynthetic genes) (Supporting Information Fig. S9). This  
334 suggests that the negative effect of 2-TDC on *P. aeruginosa* biofilms is not mediated  
335 via these determinants of biofilm formation. In contrast, a significant decrease in  
336 expression of the quorum sensing *rhII* and *pqsA* genes was detected in response to 2-  
337 TDC (Fig. 7C), which could explain the negative effect of the MK on *P. aeruginosa*  
338 biofilms since it has been observed that these two QS systems stimulate biofilm  
339 formation (Duan and Surette, 2007; Guo *et al.*, 2014) and are required for biofilm  
340 maturation (Rampioni *et al.*, 2010).

341

342 *Application of 2-TDC to tomato plants prevents infection by the phytopathogen P.*  
343 *syringae pv. tomato*

344 Considering the above mentioned results, the hypothesis that 2-TDC could interfere not  
345 only with the *Rhizobium*-legume symbiosis but also with the establishment of  
346 pathogenic interactions was worth testing. With this aim, we analysed the effect of 2-  
347 TDC treatment on the pathogenicity of *P. syringae pv. tomato* DC3000 (Pst DC3000)  
348 on tomato plants. Pst DC3000 is the causative agent for bacterial speck on tomato and it  
349 is used as a relevant model in plant pathology. Interestingly, we found that the  
350 development of disease symptoms (brown necrotic spots in leaves) (Fig. 8A) as well as  
351 growth of Pst DC3000 populations within leaf tissues (Fig. 8B) were significantly  
352 reduced in tomato plants which were sprayed with 2-TDC at the time of bacterial  
353 inoculation. The protective effect was more notable in plants which were sprayed with  
354 the highest concentration of 2-TDC. Therefore, the presence of 2-TDC during early  
355 stages of plant-bacteria interactions protects plants from bacterial infections.

356

## 357 **Discussion**

358 The aim of this study was to identify putative lipidic compounds responsible for the  
359 pleiotropic phenotype exhibited by a *fadD* mutant of *S. meliloti*. The results show that  
360 the mutant overproduces 2-tridecanone (2-TDC), and that the accumulation of this  
361 methylketone (MK) might account for the increased surface motility, decreased biofilm  
362 formation and impaired alfalfa root colonization shown by the *fadD* strain. This is the  
363 first time that 2-TDC has been shown to be produced by rhizobia. Likewise, to the best  
364 of our knowledge, this is the first report that relates FadD inactivation with levels of 2-  
365 TDC production in bacteria. More importantly, we demonstrate for the first time that 2-  
366 TDC acts as a chemical cue that affects bacterial behaviours and the establishment of



367 plant-microbe interactions. Therefore, our study assigns new biological roles to 2-TDC,  
368 which previously was known only as a strong pesticide responsible for the natural  
369 resistance against insect herbivory exhibited by 2-TDC-hyperproducing wild tomato  
370 plants (Williams *et al.*, 1980).

371 Production of 2-TDC was initially detected exclusively in the spent media from the  
372 *S. meliloti fadD* mutant cultures. Later, we found that the wild-type strain also produces  
373 volatile 2-TDC during growth on the surface of MM (1% agar), although the amount  
374 emitted was only 13% of that of the *fadD* mutant. These results demonstrate that *S.*  
375 *meliloti* can produce 2-TDC and indicate that growth conditions and especially FadD  
376 activity somehow modulate 2-TDC synthesis in this bacterium. Why the *fadD* mutant  
377 accumulates 2-TDC remains unclear because the biochemical pathways implicated in  
378 the synthesis of MKs are largely unknown. In tomato plants, 2-TDC is derived from 3-  
379 ketomyristoyl-ACP, an intermediate formed during fatty acid biosynthesis (Yu *et al.*,  
380 2010), but information on how bacteria produce MKs is scarce (Forney and Markovetz,  
381 1971; Patel *et al.*, 1980). We hypothesize that, in a *fadD* mutant, 2-TDC may derive  
382 from substrates and enzymatic reactions similar to those described for tomato  
383 (Supporting Information Fig. S10). We are currently investigating the pathway for 2-  
384 TDC formation in *S. meliloti* and how a deletion in *fadD* leads to a significant increase  
385 in 2-TDC production.

386 The concentrations of 2-TDC used in our assays are higher than those detected by  
387 SPME-GC/MS in the *fadD* mutant. However, it is very likely that only a fraction of the  
388 total amount of 2-TDC synthesized by this strain was detected due to limited volatility  
389 of the MK and partition into the cell membranes. For the same reasons, not all of  
390 exogenously applied 2-TDC might reach its target in the cell to trigger biological  
391 effects. Nevertheless, our study shows that the application of volatile 2-TDC promotes

392 surface translocation in several strains of *S. meliloti*, phenocopying the effect caused by  
393 a *fadD* mutation (Fig. 5). Likewise, without affecting growth, the presence of 2-TDC  
394 impairs biofilm formation and alfalfa root colonization by *S. meliloti* wild-type strains,  
395 resembling the effect of *fadD* inactivation. These results suggest that the accumulation  
396 of 2-TDC is responsible for the altered behaviour in the *fadD* mutant. Notably, the  
397 effects caused by 2-TDC are not exclusive to *S. meliloti* as application of the MK  
398 impacts surface motility and biofilm formation in phylogenetically distant bacteria,  
399 including plant and animal pathogens. Whereas the application of 2-TDC stimulates  
400 bacterial surface motility in plant-interacting bacteria (rhizobia and *P. syringae*  
401 pathovars), the presence of the volatile MK hampers surface translocation in the two  
402 animal pathogenic bacteria tested (*P. aeruginosa* and *S. enterica*). Importantly, the  
403 application of 2-TDC impaired biofilm formation in all the different bacteria tested,  
404 without affecting bacterial growth. These properties could be harnessed as a potential  
405 synergistic treatment against biofilm-forming pathogenic bacteria.

406 The mechanisms by which bacteria perceive and respond to volatiles are mostly  
407 unknown (Kim *et al.*, 2013; Audrain *et al.*, 2015; Schmidt *et al.*, 2015). In this study,  
408 different approaches have been used to unveil how 2-TDC impacts bacterial surface  
409 behaviours. We found that promotion of surface translocation is not a general property  
410 of aliphatic ketones since structural features of the molecule determine the compound's  
411 bioactivity, which suggests specific recognition. The bacterial response to 2-TDC  
412 cannot be explained by an effect on key motility mechanisms, since swimming motility  
413 remains unaffected. This result contrasts with the effect observed for two volatiles  
414 produced by *Bacillus subtilis*, glyoxylic acid and 2,3-butanedione, which were found to  
415 reduce both swimming and swarming motilities in different bacteria (Kim *et al.*, 2013).  
416 In *S. meliloti*, the increased surface motility triggered by 2-TDC is mainly the result of

417 promoting a flagellum-independent migration. The nature of this motility, which cannot  
418 be explained by 2-TDC acting as a surfactant (Kotowska and Isidorov, 2011), and that  
419 is not mediated by the surfactant activity of the Rh1021 siderophore, warrants further  
420 investigation.

421 The phenotypes altered by 2-TDC are often under quorum sensing (QS) regulation  
422 (Verstraeten *et al.*, 2008). In *S. meliloti*, we could not detect changes in the expression  
423 level of the QS genes *sinI* and *wgeA* (Charoenpanich *et al.*, 2013) in response to the MK  
424 (Supporting Information Fig. S11). However, in *P. aeruginosa*, swarming and biofilm  
425 inhibition caused by 2-TDC could be explained by the decreased transcription of the QS  
426 genes *rhlI* and *pqsA*, as these two QS systems stimulate biofilm formation and are  
427 required for biofilm maturation (Diggle *et al.*, 2003; Duan and Surette, 2007; Guo *et al.*,  
428 2014; Rampioni *et al.*, 2010). The negative impact of 2-TDC on twitching may also  
429 explain the reduction on biofilm formation since this type of motility has been  
430 associated with the early stages of biofilm formation (Oliveira *et al.*, 2016). Under the  
431 conditions tested, no changes in transcription of the type IV major pilin gene *pilA* could  
432 be detected. Nevertheless, decreased *pqsA* expression could influence transcriptional  
433 activity of the cognate regulator PqsR, which is required for twitching motility (Guo *et*  
434 *al.*, 2014). Considering the results obtained in *S. meliloti* and *P. aeruginosa*, and that  
435 surface translocation and biofilm formation are controlled by different mechanisms  
436 depending on the bacterial species, studies performed in different model bacteria are  
437 required in order to identify common and/or specific targets of 2-TDC.

438 The transcriptomic analyses performed in *S. meliloti* revealed that 2-TDC alters gene  
439 expression in cells grown in semisolid but not in liquid media. This could indicate that  
440 cell responses to 2-TDC might require the physiological conditions conferred upon  
441 growth on surfaces (Nogales *et al.*, 2010). Differential expression of redox and iron

442 uptake-related genes were detected in response to volatile 2-TDC. Curiously, in the  
443 cotton bollworm *Helicoverpa armigera*, 2-TDC increases the expression of a transferrin  
444 gene coding for a putative iron-binding protein and of several genes coding for heme-  
445 containing P450 enzymes which catalyse different oxidative reactions. Whereas  
446 increased expression of P450 enzymes was related to the retardant effect on insect  
447 development, transferrin was associated with moth tolerance to 2-TDC by controlling  
448 iron and H<sub>2</sub>O<sub>2</sub> levels (Zhang *et al.*, 2015; Zhang *et al.*, 2016). Future research will be  
449 focused on investigating whether in bacteria a link also exists between iron homeostasis,  
450 oxidative stress and 2-TDC. The control of iron levels is not a novel mechanism  
451 underlying the effect of bacterial volatiles. The inorganic volatile H<sub>2</sub>S renders bacteria  
452 highly resistant to oxidative stress through a mechanism involving H<sub>2</sub>S-mediated  
453 sequestration of free iron, thus avoiding damaging Fenton reactions (Shatalin *et al.*,  
454 2011; Mironov *et al.*, 2017).

455 The present study not only describes for the first time the impact of 2-TDC on  
456 bacterial behaviours that are relevant for niche colonization and persistence, but also its  
457 effect on the establishment of plant-microbe interactions. The presence of 2-TDC  
458 hampers the bacterial ability to efficiently colonize plant tissues. This effect, which  
459 correlates with the reduced biofilm formation on abiotic surfaces observed with 2-TDC,  
460 might account for the negative effect that the MK exerts on the development of plant-  
461 bacteria interactions. However, whether 2-TDC exerts an effect directly on the plant  
462 cannot be ruled out since it is known that some bacterial volatiles can elicit plant  
463 responses, including induced systemic resistance (Ryu *et al.*, 2004; Audrain *et al.*, 2015;  
464 Chung *et al.*, 2016). In any case, we demonstrate that in addition to the protective role  
465 against insect pests, 2-TDC can also protect plants from detrimental bacteria. As a  
466 natural, non-growth limiting chemical that targets bacterial traits which influence the

467 virulence of pathogenic bacteria, 2-TDC opens perspectives for the development of  
468 biotechnological solutions to control bacterial infections, minimizing the risk of  
469 generating antimicrobial resistance. The identification of genes involved in 2-TDC  
470 metabolic pathways, as well as deciphering specific mechanisms through which 2-TDC  
471 impacts bacterial behaviours and plant-bacteria interactions will undoubtedly contribute  
472 to the development of new approaches for environmental-friendly agricultural practice.

473

#### 474 **Experimental procedures**

##### 475 *Bacterial strains, plasmids, and growth conditions*

476 Bacterial strains and plasmids used in this work are listed in Supporting Information  
477 Table S5. *S. meliloti* strains were grown in either complex tryptone yeast (TY) medium,  
478 in Bromfield medium (BM), or in Robertsen minimal medium (MM) as described  
479 previously (Nogales *et al.*, 2012). *E. coli*, *S. enterica* and *Pseudomonas* spp. strains  
480 were routinely grown in Lysogeny Broth (LB) medium. Nutrient Broth (NB) medium  
481 (Rashid and Kornberg, 2000) containing 0.8% Nutrient broth No.2 (Oxoid) and 0.5% D  
482 glucose was also used for *P. aeruginosa*. When required, antibiotics were added to final  
483 concentrations ( $\mu\text{g/ml}$ ) of: 200 ampicillin, 50 kanamycin, 10 tetracycline for *E. coli*; 200  
484 kanamycin, 10 tetracycline and 75 hygromycin for *S. meliloti*; 10 rifampicin for *P.*  
485 *syringae* pv. *tomato* DC3000; 10 gentamicin for *P. aeruginosa*. *S. meliloti* and *P.*  
486 *syringae* were routinely grown at 30°C. *E. coli*, *S. enterica* and *P. aeruginosa* strains  
487 were grown at 37°C.

488

##### 489 *Analyses of lipid extracts by Gas Chromatography/Mass Spectrometry (GC/MS)*

490 One litre-cultures of *S. meliloti* strains were grown for 26 h ( $\text{OD}_{620} = 1.2$ ) at 30°C in  
491 Sherwood minimal medium as described previously (Pech-Canul *et al.*, 2011). Lipids

492 from cell pellets were extracted following the method of Bligh and Dyer (Bligh and  
493 Dyer, 1959), and spent media were extracted twice with 400 ml of acidified ethyl  
494 acetate (0.01 % acetic acid). Each of the extracts was dried by rotary evaporation at  
495 45°C and dissolved in 1 ml of 1:1 methanol:chloroform. Each extract (1 ml) was  
496 fractionated on a 5 g silica gel (Merck) column (1.5 cm diameter) by eluting with ethyl  
497 acetate:hexanes:acetic acid (60:40:5). Twenty fractions of 2 ml were collected and  
498 aliquots of each fraction were analysed by thin layer chromatography (TLC) using as  
499 mobile phase ethyl acetate:hexanes:acetic acid (60:40:5). TLC plates were revealed with  
500 iodine vapours and fractions showing a similar pattern were pooled, dried with a stream  
501 of nitrogen, and dissolved in 600 µl of methanol:chloroform (1:1). Extractions were  
502 carried out from two biological repetitions. One µl of the extract was used for analysis  
503 by GC (GC 6890 Agilent Technologies, Santa Clara, CA, U.S.A.) coupled to MS  
504 detection (Quadrupole MSD HP 5973, Agilent Technologies, Santa Clara, CA).  
505 Samples were analysed on a 5% phenyl, 95% methylpolysiloxane capillary column  
506 (HP-5MS, Agilent Technologies, Santa Clara, CA, 25 m x 0.20 mm x 0.33 µm film  
507 thickness). The oven temperature was set to 40°C for 2 min followed by a temperature  
508 gradient of 40°C to 269°C at 10°C min<sup>-1</sup> and held at 269°C for 20 min. Helium was used  
509 as the carrier gas, with a flow rate of 40 cm s<sup>-1</sup> at 100°C. The molecules were ionized at  
510 70 eV by electron impact and the NIST database version 1.7a used for identification.

511

#### 512 *Analyses of volatiles by Solid Phase MicroExtraction (SPME)-GC/MS*

513 Ten µl of washed, 10-fold-concentrated of *S. meliloti* cultures grown in TY broth to the  
514 late exponential phase were inoculated onto the surface of 3 ml of MM (1% agar) in 10  
515 ml glass vials equipped with silicone septa and incubated at 30°C for two days. For  
516 control samples 10 µl of liquid MM were applied onto the medium. Volatiles were

517 collected from the headspace of the vials by SPME (Zhang and Pawliszyn, 1993). For  
518 that, a Divinylbenzene/Carboxen/Polydimethylsiloxane (DVB/CAR/PDMS) coated  
519 StableFlex SPME fiber of 50/30 $\mu\text{m}$  (Supelco 57298-U), was conditioned for 1 h at  
520 250°C in a stream of helium and then introduced into the headspace of the vial and  
521 incubated at 30°C for 60 min. Then the fiber was inserted in the injector of GC/MS  
522 (Varian 450GC 240MS, Ion Trap) and desorbed at 280°C (6 min). Samples were  
523 separated in a DB5MS-UI column (30 m, 0.25-mm inside diameter, 0.25  $\mu\text{m}$ ). The  
524 program used for oven temperature was 40°C (3 min), 140°C (15°C min<sup>-1</sup>) and 300°C  
525 (15°C min<sup>-1</sup>), and held for a further 5 min. Helium was used as the carrier gas, with a  
526 flow rate of 1.2 ml min<sup>-1</sup>. The mass spectrometer was run at 220°C in the positive  
527 electron ionization mode at 70 eV and 220°C with scans performed from 40 to 600  $m/z$ ,  
528 which allow detection of characteristic ions of medium and long-chain aldehydes and  
529 MKs. The mass spectra of detected peaks were compared with the NIST database. The  
530 production of 2-TDC was confirmed by comparison with a pure 2-TDC standard  
531 prepared and analysed using SPME fibers in the same manner as the biological samples.

532

### 533 *Methylketone (MK) treatments*

534 MK solutions were prepared in ethanol. When the treatment was applied in the culture  
535 medium, a 1000-fold concentrated solution was diluted accordingly. Control treatment  
536 contained 0.1% ethanol. For treatments as a volatile, 20  $\mu\text{l}$  of either solutions containing  
537 the desired amount of the MK (e.g. 20  $\mu\text{l}$  of 50 mM 2-TDC contain 1  $\mu\text{mol}$  of the MK)  
538 or ethanol (control) were applied onto the lid of the plate just before incubation. Plates  
539 were sealed with parafilm and incubated face-down. Furthermore, treatments with  
540 different MKs were kept in independent compartments to avoid possible interferences  
541 between them.

542

543 *Motility assays*

544 Swimming motility was assayed on BM plates with 0.3% Bacto agar as reported  
545 previously (Nogales *et al.*, 2012). Surface motility assays for *S. meliloti* and *P. syringae*  
546 were carried out according to Soto *et al.* (2002) in MM containing 1% Noble Agar  
547 (Difco, BD). Swarming assays for *S. enterica* were carried out according to Harshey and  
548 Matsuyama (1994) on LB medium containing 0.4% glucose and 0.6% Bacto Agar  
549 (Difco). Swarming assays for *P. aeruginosa* were performed by inoculating 3  $\mu$ l of an  
550 overnight culture on the surface of NB medium containing 0.5 % Bacto Agar (Difco).  
551 Twitching motility in *P. aeruginosa* was assessed by stab inoculation of overnight  
552 cultures right to the bottom of plates containing 10 ml of LB (1% agar). After 48 h of  
553 incubation at 37°C, the agar was eliminated and plates were stained with 1% crystal  
554 violet.

555

556 *Biofilm formation assays*

557 Biofilms of *S. meliloti* and *P. aeruginosa* strains expressing the gene coding for the  
558 green fluorescence protein (GFP) were established under static growth conditions on  
559 chambered cover glass slides (Lab-Tek® II Chamber Slide™ no. 70379-82, Electron  
560 Microscopy Sciences, Hatfield, PA, USA). Late exponential phase cultures in MM  
561 broth of *S. meliloti* strains harbouring the *gfp*-expressing vector pHC60 were washed  
562 and diluted to an OD<sub>600nm</sub> of 0.2 in the same culture medium containing either 50  $\mu$ M 2-  
563 TDC or 0.1% ethanol (control). In the case of *P. aeruginosa*, an overnight culture of the  
564 GFP producing strain PADP44 was diluted to an OD<sub>600nm</sub> of 0.05 in NB broth  
565 containing 100  $\mu$ M 2-TDC or 0.1% ethanol (control). Five hundred microliters of the  
566 corresponding bacterial suspensions were inoculated into each chamber and incubated at



567 30°C for three days (*S. meliloti*) or at 37°C overnight (*P. aeruginosa*). Prior to  
568 visualization, supernatants were carefully removed by pipetting to avoid biofilm  
569 detachment. Biofilms were observed using a confocal laser inverted microscope Zeiss  
570 LSM 700 (Carl Zeiss, Oberkochen, Germany). To quantify the mass of biofilms  
571 established on glass surfaces, total fluorescence emitted by each biofilm was determined  
572 using ImageJ (Abràmoff *et al.*, 2004). At least three replicates were analysed to obtain a  
573 mean value. To assess biofilm formation of *P. syringae* pv *syringae*, strain BT111 was  
574 grown in LB broth with 50 µM 2-TDC or 0.1% ethanol (control) for 24 h at 30°C. After  
575 the incubation period, liquid from the tubes was removed and bacterial biofilm stained  
576 with 0.1% crystal violet during 20 minutes.

577

#### 578 *Transcriptomics*

579 Total RNA was isolated from cells grown either in MM broth or on the surface of  
580 semisolid MM in the presence or absence of 2-TDC. 2-TDC (50 µM) or ethanol (0.1%)  
581 were added to broth cultures at mid-exponential phase ( $OD_{600nm} = 0.5$ ), and cells were  
582 collected after 1 and 4 h of treatment. RNA was also extracted from cells grown for 7 h  
583 on the surface of semisolid MM as described previously (Nogales *et al.*, 2010) in the  
584 presence of volatile 2-TDC (1 µmol) or ethanol (control). RNA extraction, cDNA  
585 synthesis from 10 µg of total RNA, Cy3- and Cy5-labelling, hybridization to Sm14kOLI  
586 microarrays, image acquisition and data analysis were performed as previously reported  
587 (Calatrava-Morales *et al.*, 2017). Normalization and t-statistics were carried out using  
588 the EMMA 2.8.2 microarray data analysis software (Dondrup *et al.*, 2009). Data were  
589 collected and analysed from three independent biological replicates. Genes were  
590 regarded as differentially expressed if they showed a *P* value  $\leq 0.05$ , average signal to  
591 noise value  $\geq 7$  and  $\log_2$  experiment/control ratio value  $\geq 1$  or  $\leq -1$ . Raw data resulting

592 from the microarray experiments and detailed protocols have been deposited in the  
593 ArrayExpress database with accession number E-MTAB-3893.

594

#### 595 *Reverse transcription quantitative real-time PCR (RT-qPCR)*

596 One  $\mu\text{g}$  of RNA was reversely transcribed using Superscript II reverse transcriptase  
597 (Invitrogen) and random hexamers (Roche) as primers. qPCR was performed on an  
598 iCycler iQ5 (Bio-Rad, Hercules, CA, USA) according to Calatrava-Morales *et al.*  
599 (2017). The primer sequences for qPCR are listed in Supporting Information Table S6.  
600 Data normalization with 16S rRNA was performed with the critical threshold ( $\Delta\Delta\text{CT}$ )  
601 method (Pfaffl, 2001).

602

#### 603 *Measurement of $\beta$ -galactosidase activity*

604  $\beta$ -galactosidase activity of *S. meliloti* cells containing *lacZ* fusions and grown in either  
605 liquid or semisolid MM, was assayed as described previously (Soto *et al.*, 2002).

606

#### 607 *Construction of a *pilA-lux* transcriptional fusion*

608 To construct a *pilA-lux* transcriptional fusion, a  $\sim 0.3$  kb fragment generated from  
609 PAO1-L chromosomal DNA by PCR with primers PPpilAFw (5'-  
610 TATAAGCTTCTTTTCGTCGAGTAGATTGG-3') and PPpilARv (5'-  
611 ATAGAATTCGTTGATTATGTATAGGCCTA-3'), was cloned into a *Hind*III and  
612 *Eco*RI-cut miniCTX-*lux*(Gm<sup>R</sup>) plasmid. The plasmid obtained was mobilized from *E.*  
613 *coli* S17-1  $\lambda$ pir and mini-CTX elements inserted in the chromosome of PAO1-L wild-  
614 type by mating.

615

#### 616 *Analysis of bioluminescence activity*

617 To study the effect of 2-TDC on the expression of QS regulators and type IV pili, *P.*  
618 *aeruginosa* PAO1 containing the corresponding transcriptional reporter fusion was  
619 grown overnight, and cultures were spotted onto NB-agar plates in the presence and  
620 absence of 2  $\mu\text{mol}$  of 2-TDC inoculated on the plate lid (20  $\mu\text{L}$ ). After an incubation  
621 time of 8 h or 16 h at 37  $^{\circ}\text{C}$ , cells were harvested from the plate, dissolved in phosphate  
622 buffer saline (PBS), and analysed for bioluminescence output activity over growth using  
623 a TECAN Genios Pro spectrophotometer.

624

#### 625 *Rhamnolipid determination by TLC*

626 Three  $\mu\text{l}$  from overnight *P. aeruginosa* cultures grown in LB were diluted in water and  
627 then spread on swarming plates with the help of glass beads. The next day, the whole  
628 agar content was recovered and mixed with 20 ml of 1 %  $\text{KHCO}_3$ , pH=9, and the  
629 mixture was incubated 1 day at 4 $^{\circ}\text{C}$ . The agar was then removed by centrifugation and  
630 filtration with 0.2  $\mu\text{m}$  pore filters (Sartorius), and the liquid phase was extracted 3 times  
631 with equal volumes of ethyl acetate.

632 Normal phase TLC was used to study rhamnolipid production, based on a  
633 previously published method (Wittgens *et al.*, 2011). The mobile phase consisted of  
634 chloroform:methanol:glacial acetic acid in a 65:15:2 ratio. Once the samples were run,  
635 mono- and di-rhamnolipids were put in contact with a detection agent comprised of 0.15  
636 g orcinol monohydrate, 8.2 ml 60 %  $\text{H}_2\text{SO}_4$  and 42 ml water, and subsequently  
637 incubated at 110  $^{\circ}\text{C}$  for 10 min. The areas corresponding to rhamnolipids on the TLC  
638 plates were measured using GNU Image Manipulation Program (GIMP) version 2.8.10.

639

#### 640 *Nodulation and colonization assays*

641 Alfalfa (*Medicago sativa* L. cv. Aragón) seeds were surface-sterilized, germinated and  
642 the seedlings were grown in hydroponic cultures under axenic conditions in glass tubes  
643 as described previously (Olivares *et al.*, 1980). For nodulation assays, ten day-old plants  
644 (1 plant/tube) were inoculated with 1 ml of a rhizobial suspension containing  $5 \times 10^6$   
645 cells (Bernabéu-Roda *et al.*, 2015). After inoculation, the number of nodules per plant  
646 was recorded daily. For colonization assays, one-week-old alfalfa plants (5 plants in  
647 each tube) were inoculated with 1 ml of a rhizobial suspension as described above. At  
648 defined times, 15 roots from each treatment were collected and processed to determine  
649 colony-forming units (CFU) as described previously (Amaya-Gómez *et al.*, 2015).

650

#### 651 *Tomato infection assays*

652 *Solanum lycopersicum* “Moneymaker” seedlings were grown in a growth chamber with  
653 70% relative humidity, 16/8 h light/dark photoperiod and 24/20°C day/night  
654 temperature. *P. syringae* pv. *tomato* DC3000, grown on LB plates containing rifampicin  
655 for 48 h at 28°C, was resuspended in 10 mM MgCl<sub>2</sub> and the OD<sub>600</sub> was adjusted to 0.3-  
656 0.4 to obtain a bacterial suspension of 10<sup>8</sup> CFU/ml. Fifty µM 2-TDC, 500 µM 2-TDC,  
657 or 0.1% ethanol as control, were added to the bacterial suspension. Four- to five-week-  
658 old plants were inoculated with each of the three different bacterial suspensions (4  
659 plants/treatment) using an airbrush. The evolution of symptoms and sampling were  
660 performed 3 h after inoculation (time 0) and several days (3, 6 and 10) after inoculation.  
661 To determine the number of bacteria in infected leaves, five 10 mm diameter-disks (3.9  
662 cm<sup>2</sup>) were obtained per leaf with a cork borer, homogenized by mechanical disruption  
663 into 1 ml of 10 mM MgCl<sub>2</sub> and the CFU determined by plating serial dilutions. At least  
664 4 different homogenized samples obtained from 2 leaves from 2 plants were analysed  
665 per treatment.

666

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681

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839 303.
- 840

841 **Figure legends**

842 **Fig. 1.** Identification of volatile 2-TDC in *S. meliloti* by SPME-GC/MS.

843 A. Representative GC/MS chromatograms in the selected ion monitoring (SIM) mode  
844 for ion 58 of SPME trapped volatiles emitted into the headspace by *S. meliloti fadD* and  
845 wild-type strains grown on MM (1% agar), or a pure 2-TDC standard applied onto the  
846 same medium after 48 h of incubation at 30°C. The scale at the Y axis is in arbitrary  
847 units. The mean peak area  $\pm$  standard error (given in counts) corresponding to 2-TDC  
848 produced by the two *S. meliloti* strains and calculated from at least four biological  
849 measurements, is indicated with A.

850 B. Mass spectrum of the compound identified in *S. meliloti fadD* or wild-type strains  
851 and the 2-TDC standard. The arrow points to the molecular ion ( $m/z$  198).

852

853 **Fig. 2.** 2-Tridecanone promotes surface motility and impairs biofilm formation in *S.*  
854 *meliloti*.

855 A. Surface motility assays on MM (1% agar) with *S. meliloti* wild-type GR4 and  
856 Rm1021 strains in the presence of volatile 2-TDC. Twenty  $\mu$ l of either ethanol (control)  
857 or solutions containing the desired amount ( $\mu$ mol) of 2-TDC were applied to the lid of  
858 the plates just before incubation. Representative pictures of the motilities exhibited after  
859 48 h of incubation are shown. Values below each picture represent average surface  
860 migration (given in millimetres). Means and standard errors were obtained from at least  
861 3 replicates in three independent experiments (n=9). Different letters indicate significant  
862 differences according to an analysis-of-variance (ANOVA) test ( $P \leq 0.05$ ). \* Indicates  
863 that the treatment was applied as a volatile.

864 B. Three-dimensional reconstructions of overlapped confocal image stacks obtained  
865 from 3-day old biofilms developed by GFP-marked GR4 and Rm1021 after growth in  
866 MM in the presence of 0.1% ethanol (control) or 50  $\mu\text{M}$  2-TDC under static conditions.

867

868 **Fig. 3.** 2-TDC impairs nodulation of alfalfa plants.

869 A. Nodulation kinetics of alfalfa plants inoculated with *S. meliloti* GR4 in the absence  
870 or presence of 2-TDC. 2-TDC was added to final concentrations of 5  $\mu\text{M}$  or 25  $\mu\text{M}$  to  
871 the mineral solution at the moment of rhizobial inoculation. Data represent the average  
872 number of nodules per plant. The bars represent standard errors from twenty  
873 independent replicates. Asterisks indicate significant differences compared to the  
874 control according to an ANOVA test ( $P \leq 0.05$ ). One representative experiment out of at  
875 least three repetitions is shown.

876 B. Percentage of nodules formed by alfalfa plants 10 days after inoculation with *S.*  
877 *meliloti* GR4 and treated with 25  $\mu\text{M}$  2-TDC at 4 days before inoculation (4dbi), 2 days  
878 before inoculation (2dbi), at the same time as inoculation (0), 2 days postinoculation  
879 (2dpi), and 4 days postinoculation (4dpi). Data represent the average percentage  
880 calculated relative to the corresponding mean of the control treatment at the same  
881 moment as 2-TDC application. Error bars indicate the standard error. Values followed  
882 by the same letter do not differ significantly according to an ANOVA test ( $P \leq 0.05$ ).  
883 Additional data can be found in Fig. S6.

884 C. Efficient alfalfa root colonization by *S. meliloti* is hindered by the presence of 2-  
885 TDC. Colony forming units (CFU) recovered from alfalfa roots were determined 2, 24,  
886 48 and 72 h postinoculation with *S. meliloti* GR4. Data are expressed per gram of root  
887 fresh weight (RFW). Error bars indicate the standard error from the mean. According to  
888 ANOVA test ( $P \leq 0.05$ ), values followed by the same letter do not differ significantly.

889

890 **Fig. 4.** Structural features of aliphatic ketones determine their biological activity.

891 A. Surface motility of *S. meliloti* GR4 48 h after application of different ketones in  
892 volatile form. Twenty  $\mu\text{l}$  of either ethanol (control) or concentrated solutions containing  
893 the desired amount of each ketone were applied to the lid of the plates just before  
894 incubation. Data represent the means and standard errors from at least nine  
895 measurements obtained in three independent experiments. Different letters indicate  
896 significant differences compared with control and 2-TDC treatments according to an  
897 ANOVA test ( $P \leq 0.05$ ). 2-TDC, 2-tridecanone; 3-TDC, 3-tridecanone; 4-TDC, 4-  
898 tridecanone; 7-TDC, 7-tridecanone; 2-Hp, 2-heptanone; 2-DC, 2-decanone; 2-UDC, 2-  
899 undecanone; 2-DDC, 2-dodecanone; 2-PDC, 2-pentadecanone; 2-HxDC, 2-  
900 hexadecanone.

901 B. Suppression of 2-TDC-induced surface motility by different amounts of 7-TDC. Data  
902 are shown as the proportion of the surface motility exhibited by GR4 when only 2-TDC  
903 (1  $\mu\text{mol}$ ) was applied.

904

905 **Fig. 5.** Effects on surface translocation caused by the *fadD* mutation or volatile 2-TDC  
906 in flagella-less and Rhb1021-defective *S. meliloti* strains.

907 Surface motility assays on MM (1% agar) with GR4 derivative strains (A) and Rm1021  
908 derivative strains (B). The genotype of each strain is indicated. Twenty  $\mu\text{l}$  of either  
909 ethanol (control) or a solution containing 1  $\mu\text{mol}$  2-TDC were applied to the lid of the  
910 plates just before incubation. Representative pictures of the motilities exhibited by each  
911 strain are shown. Values represent average surface migration (given in millimetres)  
912 obtained after 48 h of incubation at 30°C. Means and standard errors were obtained

913 from at least 3 replicates in three independent experiments (n=9). \* Indicates that the  
914 treatment was applied as a volatile.

915

916 **Fig. 6.** 2-TDC impacts surface motility of plant and animal pathogenic bacteria.

917 Swarming of (A) the plant pathogens *Pseudomonas syringae* pv. *syringae* (Pss) BT111  
918 and *P. syringae* pv. *tomato* (Pst) DC3000 and (B) the animal pathogens *Salmonella*  
919 *enterica* serovar Typhimurium (St) LT2 and *P. aeruginosa* PAO1-L in response to  
920 volatile 2-TDC. Twenty  $\mu$ l of ethanol (control) or a solution containing 1  $\mu$ mol 2-TDC  
921 were applied to the lid of the plates just before incubation. \* Indicates that the treatment  
922 was applied as a volatile.

923 C. Twitching motility of *P. aeruginosa* PAO1-L assayed in the presence of volatile 2-  
924 TDC generated from solutions containing different amounts of the MK and applied to  
925 the lid of the plates. The zone of twitching is indicated with arrows. Error bars represent  
926 one standard deviation from three independent replicates. A one-way ANOVA was used  
927 to assess statistical significance; \*\*\*\*, p value<0.0001.

928

929 **Fig. 7.** Biofilm formation in plant and animal pathogenic bacteria is hampered by 2-  
930 TDC.

931 A. Effect of 2-TDC (50  $\mu$ M) on the crystal violet-stained biofilm formed by Pss BT111  
932 on glass tubes after 24 h of growth in LB broth.

933 B. Three-dimensional reconstructions of overlapped confocal image stacks obtained  
934 from biofilms developed by *gfp*-expressing *P. aeruginosa* PAO1-L after growth in NB  
935 in the presence of ethanol (control) or 2-TDC (100  $\mu$ M) under static conditions.  
936 Representative images of the biofilms developed after incubation overnight at 37°C are  
937 shown.



938 C. Impact of 2-TDC on the expression of *P. aeruginosa* quorum sensing regulatory  
939 genes. Strains harbouring transcriptional fusions were spotted onto NB plates alongside  
940 the corresponding promoterless construct, in the absence or presence of volatile 2-TDC  
941 (2  $\mu$ mol applied onto the plate lid). Plates were incubated for 8 h or 16 h at 37°C. \*  
942 Indicates that the treatment was applied as a volatile. Relative luminescent units (RLU)  
943 were divided by the OD<sub>600</sub>. Results are presented standardised by the RLU/OD and by  
944 the background RLU/OD of the promoterless pminiCTX-lux PADP228. Error bars  
945 represent one standard deviation of the means. Multiple t-tests, with the Holm-Sidak  
946 correction (cut off value=0.05) were performed; a, p-value<0.0001.

947

948 **Fig. 8.** Tomato plants treated with 2-TDC are more resistant to the development of  
949 bacterial speck disease.

950 A. Bacterial speck symptoms developed on tomato leaves 10 days after spray-  
951 inoculation with a suspension of *P. syringae* pv. tomato DC3000 (10<sup>8</sup> CFU/ml) in the  
952 absence or presence of 2-TDC (50 or 500  $\mu$ M). The different treatments were applied at  
953 the time of inoculation with the pathogen. Control plants were treated with an  
954 equivalent amount of ethanol (0.1%).

955 B. Time course of *P. syringae* pv. tomato DC3000 growth in leaves of tomato plants.  
956 CFUs were determined 3 h (day 0) and 3, 6, and 10 days postinoculation. Data represent  
957 the average and standard errors obtained from at least three different measurements.  
958 Asterisk indicates significant differences compared with control treatment according to  
959 an ANOVA test ( $P \leq 0.05$ ).

960

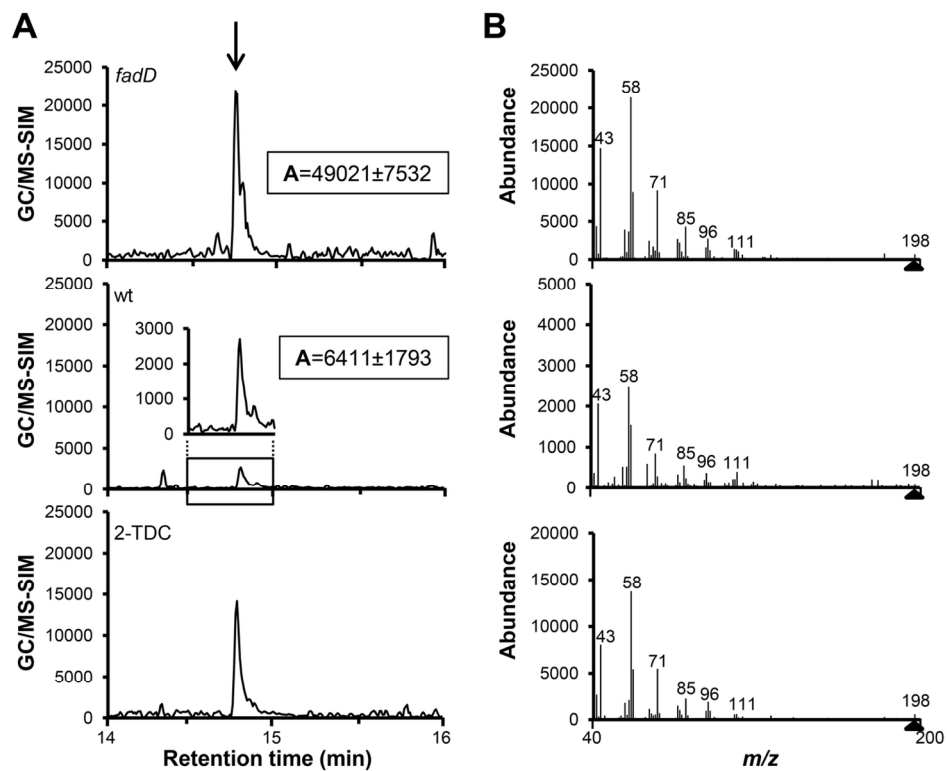


Fig. 1. Identification of volatile 2-TDC in *S. meliloti* by SPME-GC/MS.

A. Representative GC/MS chromatograms in the selected ion monitoring (SIM) mode for ion 58 of SPME trapped volatiles emitted into the headspace by *S. meliloti fadD* and wild-type strains grown on MM (1% agar), or a pure 2-TDC standard applied onto the same medium after 48 h of incubation at 30°C. The scale at the Y axis is in arbitrary units. The mean peak area  $\pm$  standard error (given in counts) corresponding to 2-TDC produced by the two *S. meliloti* strains and calculated from at least four biological measurements, is indicated with A.

B. Mass spectrum of the compound identified in *S. meliloti fadD* or wild-type strains and the 2-TDC standard. The arrow points to the molecular ion ( $m/z$  198).

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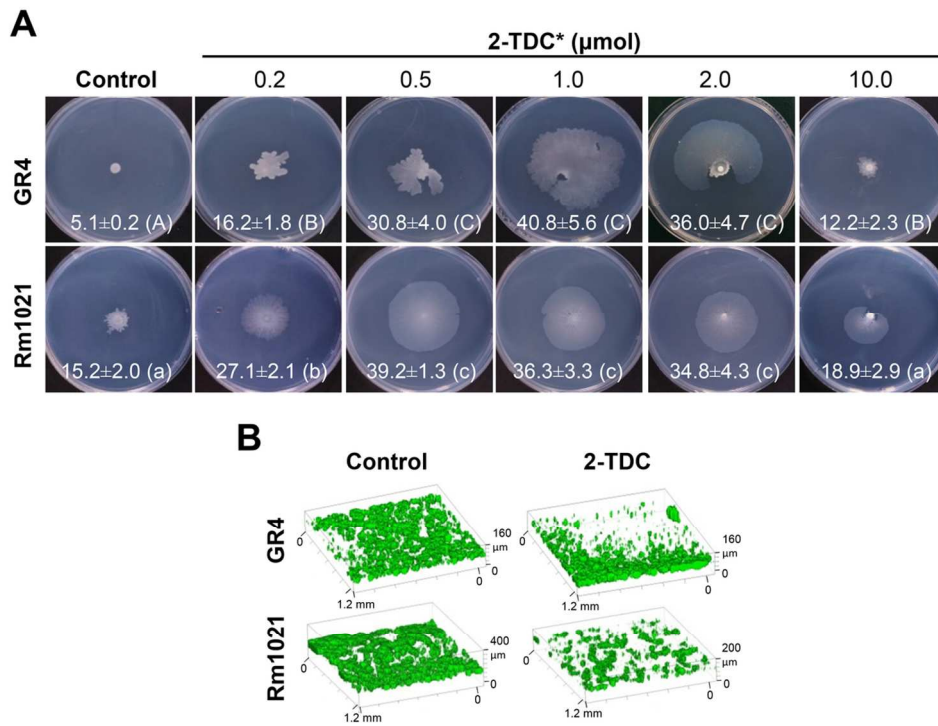


Fig. 2. 2-Tridecanone promotes surface motility and impairs biofilm formation in *S. meliloti*.

A. Surface motility assays on MM (1% agar) with *S. meliloti* wild-type GR4 and Rm1021 strains in the presence of volatile 2-TDC. Twenty  $\mu\text{l}$  of either ethanol (control) or solutions containing the desired amount ( $\mu\text{mol}$ ) of 2-TDC were applied to the lid of the plates just before incubation. Representative pictures of the motilities exhibited after 48 h of incubation are shown. Values below each picture represent average surface migration (given in millimetres). Means and standard errors were obtained from at least 3 replicates in three independent experiments ( $n=9$ ). Different letters indicate significant differences according to an analysis-of-variance (ANOVA) test ( $P\leq 0.05$ ). \* Indicates that the treatment was applied as a volatile.

B. Three-dimensional reconstructions of overlapped confocal image stacks obtained from 3-day old biofilms developed by GFP-marked GR4 and Rm1021 after growth in MM in the presence of 0.1% ethanol (control) or 50  $\mu\text{M}$  2-TDC under static conditions.

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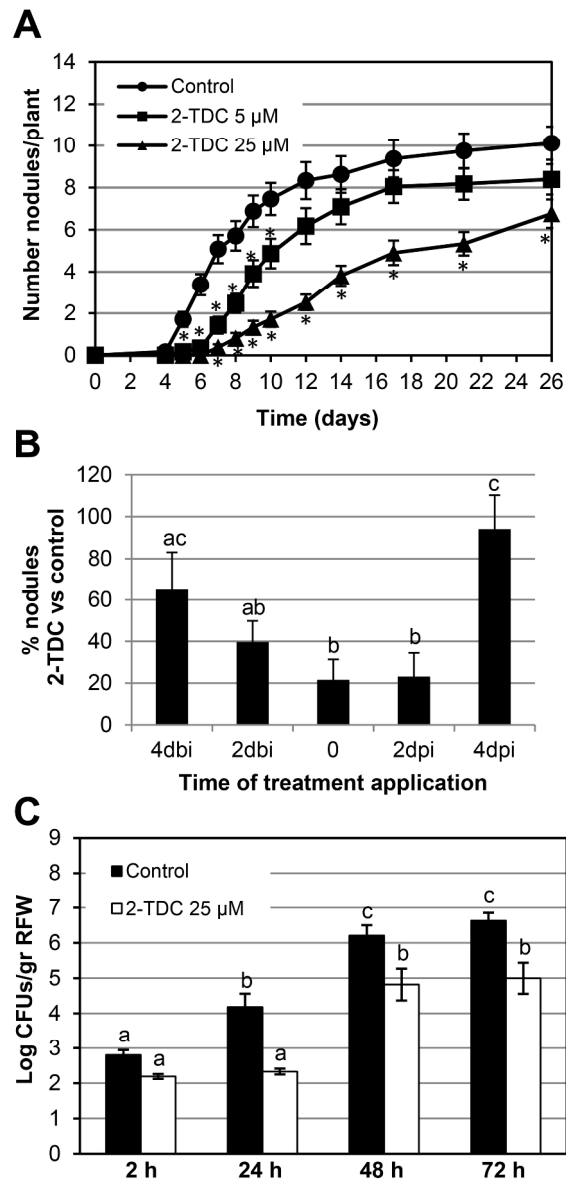


Fig. 3. 2-TDC impairs nodulation of alfalfa plants.

A. Nodulation kinetics of alfalfa plants inoculated with *S. meliloti* GR4 in the absence or presence of 2-TDC. 2-TDC was added to final concentrations of 5  $\mu$ M or 25  $\mu$ M to the mineral solution at the moment of rhizobial inoculation. Data represent the average number of nodules per plant. The bars represent standard errors from twenty independent replicates. Asterisks indicate significant differences compared to the control according to an ANOVA test ( $P \leq 0.05$ ). One representative experiment out of at least three repetitions is shown.

B. Percentage of nodules formed by alfalfa plants 10 days after inoculation with *S. meliloti* GR4 and treated with 25  $\mu$ M 2-TDC at 4 days before inoculation (4dbi), 2 days before inoculation (2dbi), at the same time as inoculation (0), 2 days postinoculation (2dpi), and 4 days postinoculation (4dpi). Data represent the average percentage calculated relative to the corresponding mean of the control treatment at the same moment as 2-TDC application. Error bars indicate the standard error. Values followed by the same letter do not differ significantly according to an ANOVA test ( $P \leq 0.05$ ). Additional data can be found in Fig. S6.

C. Efficient alfalfa root colonization by *S. meliloti* is hindered by the presence of 2-TDC. Colony forming units (CFU) recovered from alfalfa roots were determined 2, 24, 48 and 72 h postinoculation with *S. meliloti* GR4. Data are expressed per gram of root fresh weight (RFW). Error bars indicate the standard error from the mean. According to ANOVA test ( $P \leq 0.05$ ), values followed by the same letter do not differ significantly.

172x327mm (300 x 300 DPI)

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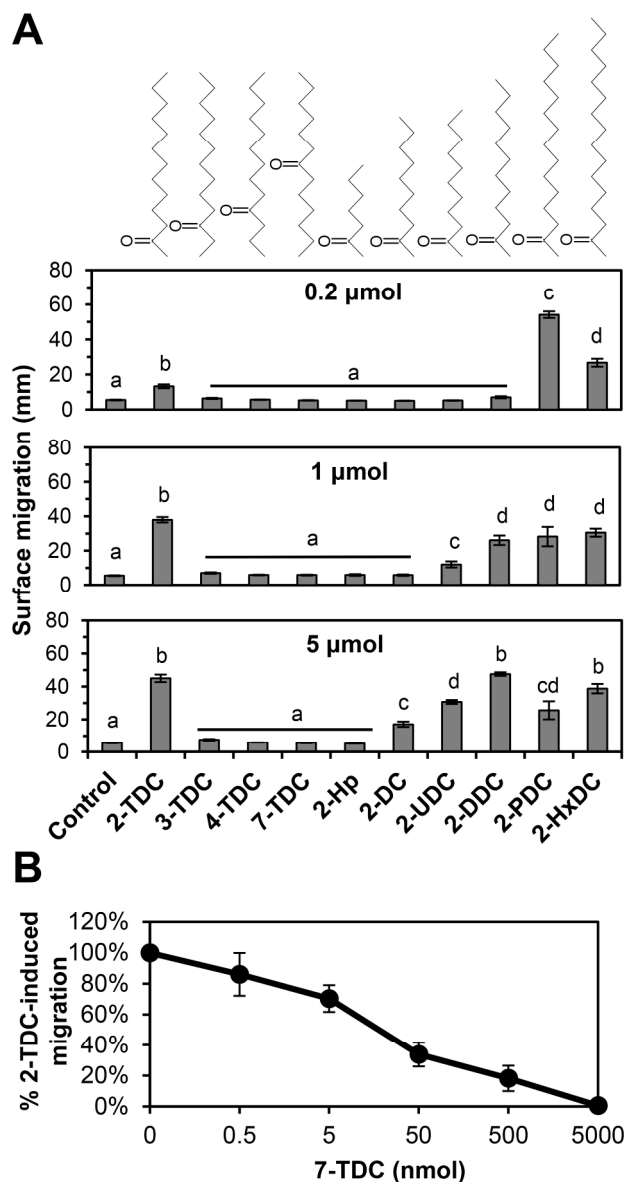


Fig. 4. Structural features of aliphatic ketones determine their biological activity.

A. Surface motility of *S. meliloti* GR4 48 h after application of different ketones in volatile form. Twenty  $\mu\text{l}$  of either ethanol (control) or concentrated solutions containing the desired amount of each ketone were applied to the lid of the plates just before incubation. Data represent the means and standard errors from at least nine measurements obtained in three independent experiments. Different letters indicate significant differences compared with control and 2-TDC treatments according to an ANOVA test ( $P \leq 0.05$ ). 2-TDC, 2-tridecanone; 3-TDC, 3-tridecanone; 4-TDC, 4-tridecanone; 7-TDC, 7-tridecanone; 2-Hp, 2-heptanone; 2-DC, 2-decanone; 2-UDC, 2-undecanone; 2-DDC, 2-dodecanone; 2-PDC, 2-pentadecanone; 2-HxDC, 2-hexadecanone.

B. Suppression of 2-TDC-induced surface motility by different amounts of 7-TDC. Data are shown as the proportion of the surface motility exhibited by GR4 when only 2-TDC (1  $\mu\text{mol}$ ) was applied.

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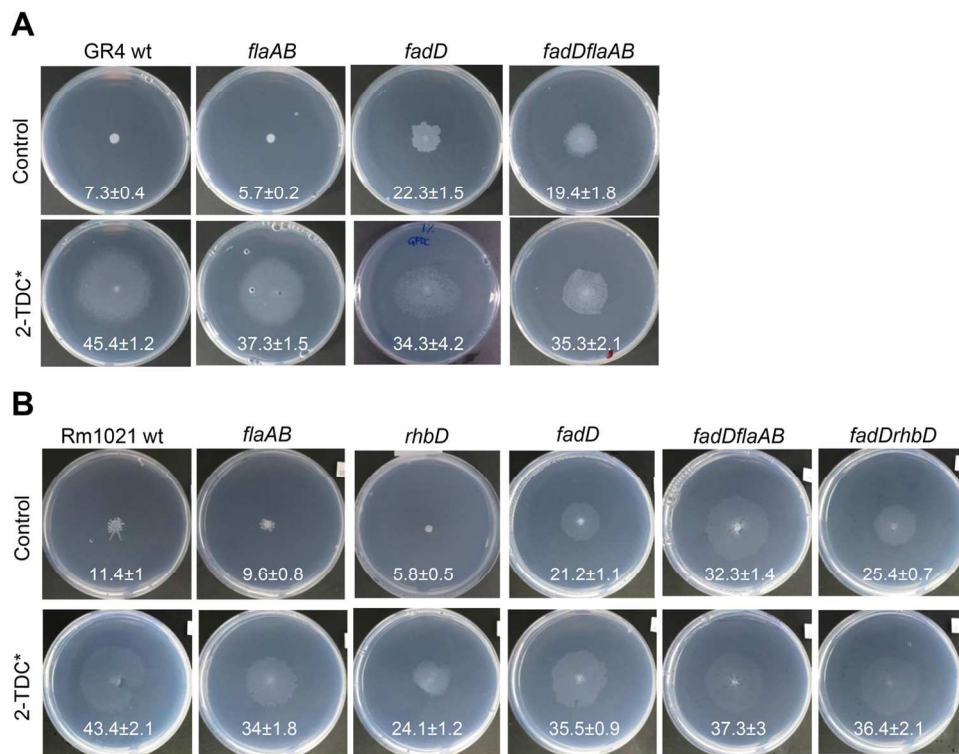


Fig. 5. Effects on surface translocation caused by the *fadD* mutation or volatile 2-TDC in flagella-less and Rhb1021-defective *S. meliloti* strains. Surface motility assays on MM (1% agar) with GR4 derivative strains (A) and Rm1021 derivative strains (B). The genotype of each strain is indicated. Twenty  $\mu$ l of either ethanol (control) or a solution containing 1  $\mu$ mol 2-TDC were applied to the lid of the plates just before incubation. Representative pictures of the motilities exhibited by each strain are shown. Values represent average surface migration (given in millimetres) obtained after 48 h of incubation at 30°C. Means and standard errors were obtained from at least 3 replicates in three independent experiments (n=9). \* Indicates that the treatment was applied as a volatile.

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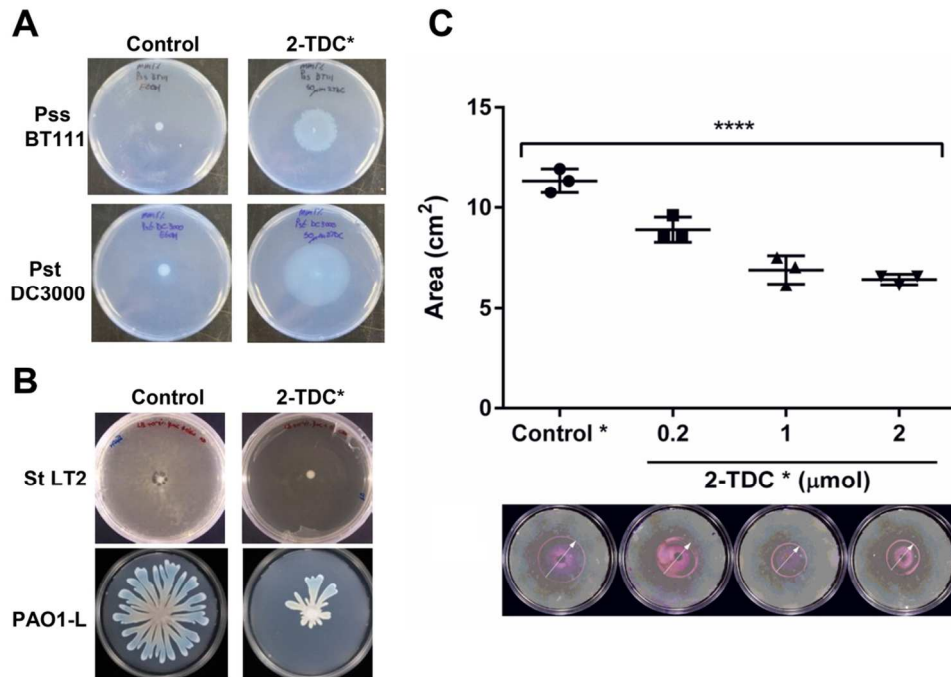


Fig. 6. 2-TDC impacts surface motility of plant and animal pathogenic bacteria. Swarming of (A) the plant pathogens *Pseudomonas syringae* pv. *syringae* (Pss) BT111 and *P. syringae* pv. *tomato* (Pst) DC3000 and (B) the animal pathogens *Salmonella enterica* serovar Typhimurium (St) LT2 and *P. aeruginosa* PAO1-L in response to volatile 2-TDC. Twenty  $\mu$ l of ethanol (control) or a solution containing 1  $\mu$ mol 2-TDC were applied to the lid of the plates just before incubation. \* Indicates that the treatment was applied as a volatile.

C. Twitching motility of *P. aeruginosa* PAO1-L assayed in the presence of volatile 2-TDC generated from solutions containing different amounts of the MK and applied to the lid of the plates. The zone of twitching is indicated with arrows. Error bars represent one standard deviation from three independent replicates. A one-way ANOVA was used to assess statistical significance; \*\*\*\*,  $p$  value < 0.0001.

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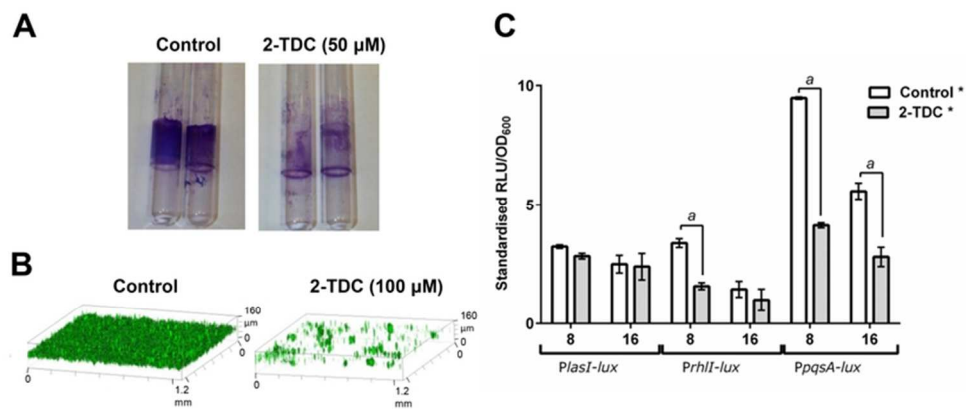


Fig. 7. Biofilm formation in plant and animal pathogenic bacteria is hampered by 2-TDC.

- A. Effect of 2-TDC (50  $\mu\text{M}$ ) on the crystal violet-stained biofilm formed by Pss BT111 on glass tubes after 24 h of growth in LB broth.
- B. Three-dimensional reconstructions of overlapped confocal image stacks obtained from biofilms developed by *gfp*-expressing *P. aeruginosa* PAO1-L after growth in NB in the presence of ethanol (control) or 2-TDC (100  $\mu\text{M}$ ) under static conditions. Representative images of the biofilms developed after incubation overnight at 37°C are shown.
- C. Impact of 2-TDC on the expression of *P. aeruginosa* quorum sensing regulatory genes. Strains harbouring transcriptional fusions were spotted onto NB plates alongside the corresponding promoterless construct, in the absence or presence of volatile 2-TDC (2  $\mu\text{mol}$  applied onto the plate lid). Plates were incubated for 8 h or 16 h at 37°C. \* Indicates that the treatment was applied as a volatile. Relative luminescent units (RLU) were divided by the OD<sub>600</sub>. Results are presented standardised by the RLU/OD and by the background RLU/OD of the promoterless *pminiCTX-lux PADP228*. Error bars represent one standard deviation of the means. Multiple t-tests, with the Holm-Sidak correction (cut off value=0.05) were performed; a,  $p\text{-value} < 0.0001$ .

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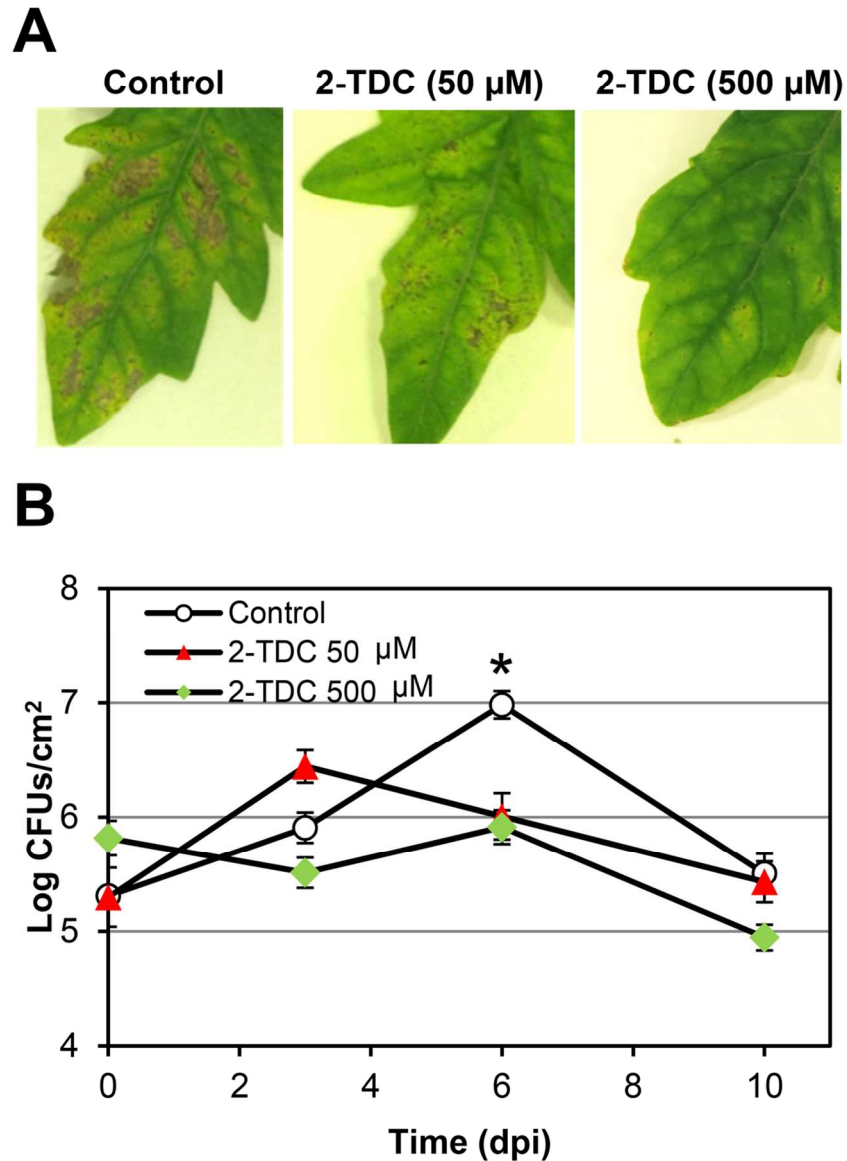


Fig. 8. Tomato plants treated with 2-TDC are more resistant to the development of bacterial speck disease. A. Bacterial speck symptoms developed on tomato leaves 10 days after spray-inoculation with a suspension of *P. syringae* pv. tomato DC3000 (108 CFU/ml) in the absence or presence of 2-TDC (50 or 500  $\mu\text{M}$ ). The different treatments were applied at the time of inoculation with the pathogen. Control plants were treated with an equivalent amount of ethanol (0.1%).

B. Time course of *P. syringae* pv. tomato DC3000 growth in leaves of tomato plants. CFUs were determined 3 h (day 0) and 3, 6, and 10 days postinoculation. Data represent the average and standard errors obtained from at least three different measurements. Asterisk indicates significant differences compared with control treatment according to an ANOVA test ( $P \leq 0.05$ ).

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