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2	Full Title:
3	Genome wide identification of genes required for bacterial plant infection by Tn-seq
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5	Short title: Dickeya dadantii virulence genes in chicory
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15 Abstract

Soft rot enterobacteria (Dickeva and Pectobacterium) are major pathogens that provoke 16 17 diseases on plants of agricultural importance such as potato and ornamentals. Long term 18 studies to identify virulence factors of these bacteria focused mostly on plant cell wall 19 degrading enzymes secreted by the type II Out secretion system and the regulation of their expression. To identify new virulence factors we performed a Tn-seq genome-wide screen of 20 21 a transposon mutant library during chicory infection followed by high-throughput sequencing. This allowed the detection of mutants with reduced but also increased fitness in 22 23 the plant. Virulence factors identified differed from those previously known since diffusible 24 ones (secreted enzymes, siderophores or metabolites) were not detected by this screen. In addition to genes encoding proteins of unknown function that could be new virulence factors, 25 26 others could be assigned to known biological functions. The central role of the FlhDC 27 regulatory cascade in the control of virulence was highlighted with the identification of new members of this pathway. Scarcity of the plant in certain amino acids and nucleic acids 28 29 required presence of the corresponding biosynthetic genes in the bacteria. Their products 30 could be targets for the development of antibacterial compounds. Among the genes required for full development in chicory we also identified six genes involved in the glycosylation of 31 the flagellin FliC, a modification which in some other plant pathogenic bacteria contributes to 32 33 virulence. 34 35

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40 Introduction

Dickeva are broad-host range phytopathogenic bacteria belonging to the 41 42 Pectobacteriaceae family [1] that provoke the soft rot disease on many plant species. They 43 are the cause of important losses on economically important crops such as potato, chicory and ornamentals. Identification and studies on the virulence factors of these bacteria have been 44 45 performed mostly on the model strain D. dadantii 3937 and focused mainly on three 46 domains/aspects, known to be important for disease development: plant cell wall degrading enzymes, the type III secretion system and iron metabolism [2]. Secretion of plant cell wall 47 48 degrading enzymes has long ago been identified as the bacteria main virulence factor. Many 49 studies focused on the identification and characterization of these secreted enzymes, mostly pectinases [3], of the regulators controlling their production (kdgR, pecS, pecT, hns, gacA), 50 51 [4-8] of the genes whose expression is coregulated with that of the secreted enzyme genes [9, 52 10], and of the mechanism of their secretion by the Out type II secretion system [11]. Although of a lesser importance for *Dickeya* virulence, the same type of approach has been 53 54 used to identify Hrp type III secretion system regulators and effectors [12] [13] [14]. Competition for iron within the plant is strong. D. dadantii acquires this metal through 55 production of two siderophores, chrysobactin and achromobactin [15] [16] [17]. More 56 recently, omics approaches have also been used to identify genes induced during plant 57 infection [18] [19] [20]. These studies now give a view on a complex network of factors 58 59 required for *D. dadantii* virulence [2, 21]. However, these approaches may have missed some important factors not targeted by these analyses. More global screens need to be performed to 60 identify these factors. Libraries of transposon-induced mutants were tested on plants to find 61 62 mutants showing reduced virulence with Pectobacterium carotovorum and atrosepticum, two other soft rot enterobacteria [22, 23]. These studies identified auxotrophs, mutants defective 63 in production or secretion of exoenzymes and in motility. Other mutants with a more 64

65 complex phenotype were not characterized at this time. Moreover, the number of tested mutants was limited by the necessity to test individually each mutant on plant. This type of 66 67 work has never been performed on *Dickeya* strains. To have a more complete view of the genes required for the virulence of Dickeya, we used a high-throughput sequencing of a 68 saturated transposon library (Tn-seq) to screen tens of thousands random insertion mutants of 69 D. dadantii in laboratory medium and during infection of chicory. Tn-seq has been 70 71 extensively used to uncover essential genes required for mouse colonization by human pathogens Vibrio cholerae [24], Pseudomonas aeruginosa [25] and Streptococcus 72 73 pneumoniae [26]. Thus, Tn-seq is a very powerful method to identify genes required for 74 bacterial growth in their host. However, this technology has not yet been employed to unveil genes required for the multiplication of a phytopathogen in a plant host. By applying this 75 76 technique to screen a *D. dadantii* mutant library in chicory, we identified metabolic pathways and bacterial genes required for growth *in planta*. Among them, we found a cluster of genes 77 required for flagellin glycosylation, a modification known to be important for several plant 78 79 pathogenic bacteria virulence.

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- 82 Results and discussion
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84 Characterization of *D. dadantii* 3937 *Himar1* transposon library

Many tools are available to perform Tn-seq [27]. In order to perform a Tn-seq experiment with *D. dadantii* 3937, we used a *Himar9* mariner transposon derivative carrying MmeI restriction sites in the inverted repeats (IR) and a kanamycin resistance cassette between the IRs [28]. We realized a biparental mating between *E. coli* and *D. dadantii* on M63 agar medium without carbon source and amino acids. We obtained approximately 300 000

90 colonies that were pooled. Subsequent DNA sequencing (see below) showed the presence of 91 transposon insertions in amino acid biosynthesis pathways, demonstrating that mating on 92 M63 minimal medium does not prevent the obtention of auxotroph mutants. To identify 93 essential genes, mutants were grown in LB medium during several generations. Two DNA libraries were prepared from two cultures and subjected to high-throughput sequencing. The 94 95 mariner transposon inserts into TA dinucleotides. The TPP software [29] was used to 96 determine the number of reads at each TA site for each biological replicate. D. dadantii genome has 171 791 TA sites that can be targeted by the Himar9 transposase. Pairs of 97 98 biological replicates were compared. 37 794 and 48 101 unique insertions in TAs were 99 detected in each sample, which corresponds to 22 and 28% density of insertion respectively (Table 1). The average number of reads per TA is 88 and 75, respectively. The results were 100 101 reproducible with a Pearson correlation coefficient of 72% (Fig. S1) The location of the 102 unique insertions showed an even distribution around the chromosome (Fig. 1A). For each 103 gene, we calculated a log₂FC corresponding to a ratio between the measured number of reads 104 and the expected number of reads. The density plot (Fig. 1B) indicates that essential and non-105 essential genes are easily distinguishable, confirming the good quality of our Tn-seq libraries. 106 Then, gene essentiality of the Tn-seq input libraries was determined by using the TRANSIT 107 software [29]. We decided to use the Hidden Markov Model (HMM) which predicts 108 essentiality and non-essentiality for individual insertion sites because it has been shown to 109 give good prediction in datasets with density as low as 20% [29]. The HMM analysis led to 110 the identification of 665 genes essential for growth in LB (ES), representing 14% of the genes of D. dadantii 3937, a number in the range of those found for this type of analysis with 111 112 bacteria. 552 genes were categorized as Growth Defect genes (GD, i.e. mutations in these 113 genes lead to loss of fitness), 125 as growth advantage genes (GA, i.e mutations in these 114 genes lead to gain of fitness) and 3320 as non-essential genes (NE) (table S2).

115	TABLE 1	Tn-Seq	analysis	of Dickeya	dadantii 3937

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Mutant pool	Total no. of reads	No. of reads containing Tn end		No. of mapped read to unique TA sites	t No. of mapped reads to unique TA sites after LOESS correction	Density (%) ^b	Mean read count ^c
LB #1	23,152,186	22,647,343	18,748,028	13,166,770 (70 %)	12,904,900 (69 %)	28 %	75
LB #2	30,105,412	27,963,154	18,748,028	15,535,291 (83 %)	15,195,582 (81 %)	22 %	88
Chicory #1	18,925,029	18,748,028	18,748,028	17,535,146 (94 %)	14,906,888 (79 %)	24 %	87
5	27,607,717	26,555,297	18,748,028	17,477,706 (93 %)	16,955,724 (90 %)	23 %	99

^a The number of reads containing the sequence of a Tn end were normalized for each sample

118 according to the number of reads for the sample Chicory #1

^b Dickeya dadantii 3937 genome has 171,791 TA sites. The density is the % of TAs for which mapped

120 reads has been assigned by the TPP software.

121 ^c the mean value of mapped reads per TA.

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Genes necessary for chicory leaf maceration. We used chicory leaf infection as a model to 123 identify D. dadantii genes required for growth in plant tissues. Biological duplicates were 124 125 performed to insure the reproducibility of the results. Each chicory was inoculated with 10^7 bacteria from the mutant pool and after 2 days more than 10¹⁰ bacteria were collected from 126 the rotten tissue. Sequencing transposon insertion sites in these bacteria followed by the TPP 127 128 analysis indicated a density of unique insertion in TAs comparable to that of the input datasets (23-24%). Surprisingly, the results were more highly reproducible than in LB with a 129 very high Pearson correlation coefficient of 98% (Fig. S1). 130

In order to identify genes of D. dadantii conferring loss or gain of fitness in planta, we 131 performed the RESAMPLING analysis of the TRANSIT software. The RESAMPLING 132 133 method is a variation of the classical permutation test in statistics that sums the reads at all TA sites for each gene in each condition. It then calculates the difference of the sum of read-134 counts between the input (LB) and output (chicory) datasets. The advantage of this statistical 135 136 method is to attribute for each gene an adjusted p-value (q-value). Genes with a significant difference between total read-counts in LB and chicory achieve a q-value ≤ 0.05 . The method 137 also calculates a log₂ fold-change (log₂FC) for each gene based on the ratio of the sum of 138

139 read counts in the output datasets (chicory) versus the sum of read counts in the input (LB) 140 datasets [29]. Applied to our Tn-seq datasets and selecting only genes achieving a q-value \leq 141 0.05, we identified 122 genes out of 4666 required for fitness in planta, as shown with the 142 volcano plot of RESAMPLING results comparing replicates grown in LB versus in planta (Fig. S2). From this 122 genes, we applied an additional cutoff by removing 20 genes with a 143 144 mean read count in LB <5 (less than 5 reads in average / TA). These genes were categorized 145 as ES or GD in LB. We also removed from the analysis 6 genes with a log₂FC comprised between -2 and 2. By applying all these criteria, we retained only 96 genes for a further 146 147 analysis (Fig. 2). 92 of them were identified as GD genes in the chicory ($\log_2 FC \leq 2$), the 4 148 left as GA genes in the chicory (log₂FC \geq 2). Some of these genes, in bold in Fig 2, were already known to play a role in D. dadantii virulence, confirming the validity of the Tn-seq 149 150 approach. Using the Kyoto Encyclopedia of Genes and Genomes (KEGG) [30], we 151 discovered that certain metabolic pathways and biological functions are very important for 152 growth in chicory (Table S3). We highlight some of them in the next sections of the article.

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154 Analysis of the genes of *D. dadantii* required for plant colonization.

155 (i) Metabolism. Chicory appears as an environment in which amino acids, nucleic acids and some vitamins (pyridoxal) are scarce. Of the 92 genes identified as GD genes in planta, 8 are 156 157 involved in purine and 7 in pyrimidine metabolisms (table S3). In the purine metabolism 158 pathway, the inosine monophosphate (IMP) biosynthesis pathway that produces IMP from L-159 glutamine and 5-phosphoribosyl diphosphate is particularly important for D. dadantii in planta since 5 out of the 10 genes of this pathway are significantly GD genes in planta (Fig. 160 161 3). IMP is the precursor of adenine and guanine. Next, IMP can be converted in xanthosine 162 5'-phosphate (XMP) by the IMP dehydrogenase GuaB. guaB gene is also a GD gene in 163 planta, with a strong log₂FC of -10.06 (Fig. 3). In the pyrimidine synthesis, the uridine monophosphate (UMP) biosynthesis pathway that converts L-glutamine to UMP, a precursor
of uracyl, is very important *in planta* since *carAB*, *pyrB*, *pyrC* and *pyrE*, involved in this
enzymatic pathway, are all required for growth *in planta* (Fig. 3). This pyrimidine
biosynthesis pathway is specific to bacteria. It is noteworthy that in the human pathogen *S*. *pneumoniae*, mutants of this pathway have a fitness defect in the nasopharynx of infected
mice [26]. Hence, it looks that the pyrimidine biosynthesis pathway is particularly important
for multiplication of some bacterial species in the host.

171 Mutants in genes involved in the synthesis of sulfur-containing amino acids (cvsIJO, metB), 172 lysine (lysA) and leucine (leuABC) are disadvantaged in chicory (Fig. 2 and Fig. 4A). These 173 amino acids are known to be present in low concentration in plant tissues. Other amino acids 174 seem to be present in quantity sufficient for growth of *D. dadantii* auxotrophs. Low level of 175 certain amino acids probably induces the stringent response in the bacteria. Reduced growth 176 in the plant of the *relA* mutant, unable to synthesize the alarmone ppGpp, supports this 177 hypothesis. Glucose is the main sugar present in plant tissue, present as a circulating sugar or 178 a cellulose degradation product. Mutants in the PTS glucose transport system genes *ptsI* and 179 ptsG have a reduced growth in bacteria (Fig. 2) showing its importance as a carbon source in 180 planta.

Degradation of cell wall pectin by a battery of extracellular enzymes is the main determinant of *Dickeya* pathogenicity. Mutants unable to produce or to secrete these enzymes by the type II Out secretion system were not disfavored in chicory since these mutants could use for their growth the pectin degradation compounds produced by enzymes secreted by other bacteria. The redundancy of oligogalacturonate specific porins (KdgM and KdgN) and inner membrane transporters (TogT and TogMNABC) allow entry of these compounds into the bacteria even in a mutant in one of these transport systems. However, *kduI* mutants, blocked

in the intracellular part of the pectin degradation pathway, have a limited growth *in planta*,confirming the importance of the pectin degradation pathway in the disease progression.

190 (ii) Stress resistance. Plant is an hostile environment for the bacteria that has to cope with 191 antimicrobial peptides, ROS, toxic compounds and acidic pH [31]. We observed that the pump AcrABTolC, that can efflux a wide range of compounds [32], is important for survival 192 193 in chicory (Fig. S3). Stress can lead to the accumulation of phospholipids in the outer 194 membrane. This accumulation makes the bacteria more sensitive to small toxic molecules [33]. Such a phospholipid accumulation probably occurs when the bacteria infect chicory 195 196 since *mlaC* and *mlaF* mutants, which are unable to prevent phospholipid accumulation in the 197 outer membrane, have a reduced growth in plant. Production of exopolysaccharides (EPS) was shown to protect the bacteria during the first steps of infection [9]. We observed that 198 199 rffG and wzx mutants unable to synthesize EPS have a growth defect in chicory. A set of 200 genes required to repair or degrade altered proteins (*clpA*, *degO*, *trxB*) are also important for survival in planta. No gene directly involved in detoxification of ROS was detected in our 201 202 analysis. However, ROS can create DNA damage. The two helicases involved in DNA repair, 203 UvrD and HelD, give growth advantage in plant. Osmoregulated periplasmic glycans (OPG) 204 are polymers of glucose found in the periplasm of α , β and γ proteobacteria. Their exact role 205 is unknown but their absence leads to avirulence in certain bacteria such as D. dadantii [34]. 206 This absence induces a membrane stress that is sensed and transduced by the Rcs envelope 207 stress response system. This system controls the expression of many genes, including those 208 involved in motility, and those encoding plant cell wall degrading enzymes through the RsmA-RsmB system [35-37]. Thus, mutants defective in OPG synthesis are expected to have 209 210 a reduced virulence. Indeed, in our experiment, mutants in the two genes involved in OPG 211 synthesis, *opgG* and *opgH* were non competitive in chicory (Fig. 2).

212 (iii) Iron uptake. D. dadantii produces two types of siderophores, achromobactin and 213 chrysobactin, that are required for the development of maceration symptoms in the iron 214 limited environment of plant hosts [38]. Once iron loaded, the siderophores are imported into 215 the bacteria. Import through the outer membrane requires a specific outer membrane channel 216 and the energy transducing complex formed by TonB ExbB and ExbD. While the absence of 217 synthesis of one of the siderophores can be compensated by the presence of siderophore 218 secreted by other bacteria in the growth medium, mutants of the TonB complex are totally unable to acquire iron and thus are unable to grow in the plant. In accordance, tonB was 219 220 essential in chicory while the genes coding for siderophore synthesis or secretion were not. 221 Similarly a mutant devoid of the iron-loaded chrysobactin transport gene (fct) is non-222 competitive.

223 (iv) Regulation. Mutants in several genes controlling virulence factor production have a 224 growth defect in the plant. The master regulator FlhDC acts as a regulator of both flagella and 225 virulence factor synthesis in many bacteria such as Yersinia ruckeri, Edwardsiella tarda and 226 Ralstonia solanacearum [39-41]. In D. dadantii FlhDC has recently been shown to control, in 227 addition to flagellar motility, type III secretion system and virulence factor synthesis through 228 several pathways [42]. We observed that flhC gives a growth advantage in chicory. In addition, we uncovered that some genes regulating *flhDC* in other bacteria regulate *D*. 229 230 dadantii virulence, probably by controlling *flhDC* expression. *rsmC* is a poorly characterized 231 gene in *D. dadantii* but that has been studied in *Pectobacterium carotovorum*. It negatively 232 controls motility and extracellular enzyme production through modulating transcriptional activity of FlhCD [43]. HdfR is a poorly characterized LysR family regulator that controls the 233 234 std fimbrial operon in S. enterica and FlhDC expression in E. coli [44]. rsmC mutants were 235 overrepresented in the chicory (Fig. 4B), indicating a gain of virulence for these mutants.

hdfR conferred fitness benefits during growth in chicory and could also act in *D. dadantii* as
activator of *flhDC* expression.

238 The GGDEF proteins are c-di-GMP synthase. Their gene are often located next to their 239 cognate EAL diguanylate phosphodiesterase gene. *ecpC (vhiH)* encodes an EAL protein that was shown to activate virulence factor production in D. dadantii [45]. gcpA, which is located 240 next to ecpC encodes a GGDEF protein. However, a gcpA mutant could not be constructed 241 242 and analysed in this previous study. We observed that gcpA mutants (Dda 03858) were 243 overrepresented in chicory (Fig. 2). This increased virulence is a phenotype opposite to that 244 described for the *ecpC* mutants, indicating that overproduction of c-di-GMP could reduce D. 245 dadantii virulence.

Of the eighteen regulators of the LacI family present in *D. dadantii*, four of them were found to be involved in plant infection [46]. One of those, LfcR, which has been found important for infection of chicory, Saintpaulia and Arabidopsis, was identified as important for chicory infection in our experiment. LfcR is a repressor of adjacent genes [46]. Surprisingly none of these genes appeared to play a role for chicory infection suggesting that other targets of LfcR probably remain to be discovered.

Finally, it is noteworthy to mention that the *ackA* and *pta* genes are GD *in planta*. These genes constitute the reversible Pta-AckA pathway. The steady-state concentration of acetylphosphate (acetyl-P), a signaling molecule in bacteria, depends upon the rate of its formation catalyzed by Pta and of its degradation catalyzed by AckA [47]. The GD phenotype of *D*. *dadantii ackA* and *pta* mutants during infection suggests that acetyl-P might play a crucial signaling role in the adaptation of *D*. *dadantii* to the plant tissue.

(v) Motility. Motility is an essential virulence factor of *D. dadantii* required to move on the
surface of the leaf, enter the wounds and propagate into the plant tissue [48-50]. Accordingly,
all the genes required for flagella synthesis, the flagella motor and genes regulating their

synthesis (*flhC*, *flhD*, *fliA*) (see above) are necessary for fitness during chicory infection (Fig. 4C and 5A). All the genes responsible for the transduction of the chemotaxis signal (*cheA*, *B*, *R*, *W*, *X*, *Y* and *Z*) also confer a benefit in planta (Fig. 2). No methyl-accepting chemoreceptor gene mutant was found. Like other environmental bacteria, *D. dadantii* encodes many such proteins (47). They probably present some redundancy in the recognized signal which prevented their detection by our screen.

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268 D. dadantii flagellin is modified by glycosylation

269 A group of six genes located between *fliA* and *fliC* retained our interest since insertions in 270 these genes led to a growth defect in chicory (Fig. 5A). Dda3937 03424 encodes an O-linked N-acetylglucosamine transferase and Dda3937 03419 encode a protein with a nucleotide 271 272 diphospho sugar transferase predicted activity. The other ones could be involved in the 273 modification of sugars (predicted function of Dda3937 03423: nucleotide sugar 274 transaminase, Dda3937 03422: carbamoyl phosphate synthase, Dda3937 03421: 275 oxidoreductase; Dda3937 03420: methyltransferase). Their location let suppose that this 276 group of genes could be involved in flagellin glycosylation. Analysis by SDS-PAGE of FliC 277 produced by the wild type, and the A4277 and A3422 strains (Dda3937 03424 and Dda3937 03419 mutants, respectively) showed that in the two latter strains the molecular 278 279 weight of the protein diminished (Fig. 5B). The molecular weight determined by mass 280 spectroscopy was 28911 Da for FliCA4277, 31034 Da for FliCA3422 and 32258 Da for the 281 WT Flic. Thus, in the wild type strain Flic is modified by the products of the genes Dda 03424 to Dda 03419, probably by multiple glycosylation with a disaccharide. Absence 282 283 of modification did not modify D. dadantii motility (data not shown). The flagellin of the 284 plant pathogens Pseudomonas syringae pv tabaci and Burkholderia cenocepacia are also glycosylated and absence of this modification lowered the ability of these bacteria to cause 285

disease on tobacco and arabidopsis, respectively [51, 52]. Accordingly, in *D. dadantii*, FliC
modification appears important for multiplication of the bacteria in plant.

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289 Additional genes could be involved in virulence

290 Several genes have a $\log_2 FC >4$ or <-4 but do not satisfy the statistical permutation test adjusted by the false discovery rate method (q-value) (table S4). However, most of them 291 292 belong to the categories described above and could be required for growth in planta. Among those with a log₂FC< -4 can be found genes involved in amino acid and nucleic acid synthesis 293 294 (cysH, ilvC, pyrF, pyrD, purC, thrC, metA, cysK, lysC), flagella and motility (flgJ, fliO, flgC, 295 fliS, flgG, flgA, flgL, cheW, fliN, fliP, fliK, fliG, fliL), pectin and glucose metabolism (kduD, pgi), EPS synthesis (gmd), flagella glycosylation (vioA) and regulation (zur, ecpC and the 296 297 general RNA chaperone *hfq*).

298 Among the genes with a $\log_2 FC > 4$, three regulators can be noticed: *pecS*, *pecT* and Dda3937 00840. pecS and pecT are known regulators of D. dadantii controlling the 299 300 expression of many factors involved in virulence (ref, ref). Thus, their mutation could confer an increased fitness of the bacteria in chicory. D. dadantii possesses a functional expl-expR 301 quorum sensing system which does not seem to control plant virulence factor production 302 303 (Nasser). However, several LuxR family regulator genes which are not associated with a *luxI* 304 gene are present in the genome of the bacteria. Mutants of one of them (Dda3937 00840) are 305 overrepresented in the chicory. Its product is probably a repressor of genes conferring an 306 increased fitness in planta.

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309 Validation of the Tn-seq results.

310 To validate the Tn-seq results, we performed coinoculation experiments in chicory leaves 311 with the wild type strain and various mutants in genes conferring a growth advantage, a 312 growth defect or no fitness benefit in a 1/1 ratio. We calculated a competitivity index (CI) by 313 counting the number of each type of bacteria in the rotten tissue after 24 h. We found a 314 correlation between the competitivity index of mutants with the log_2FC of the corresponding 315 genes with a Pearson coefficient of 0.50 (Fig. 6), indicating that Tn-seq is a reliable technique 316 to identify genes involved in plant colonization and virulence. Additional validations were 317 performed by inoculation of uncharacterized leucine and cytosine auxotroph mutants in 318 chicory leaves which confirmed that these mutants are unable to grow in plant and are thus 319 avirulent (Fig S4).

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322 Conclusion

This Tn-seq experiment highlights new factors required for *D. dadantii* successful rotting of 323 324 chicory. Many genes known to be important for pathogenesis were not found in this screen 325 because their products are secreted and can be shared with other strains in the community. This includes all the proteins secreted by the type II secretion system and small molecules 326 327 such as siderophores and butanediol. Other categories of genes were not found: for example, no genes involved in response to acidic or oxidative stresses were identified. Chicory has 328 329 been described as an inadequate model to study the response of *D. dadantii* to oxidative stress 330 (touati, expert). Similarly, the type III *hrp* genes were not identified in our study. The Hrp system is not necessary for *D. dadantii* virulence and in our experimental conditions (high 331 332 inoculum on isolated chicory leaves) the necrotrophic capacities of *D. dadantii* (production of 333 plant cell wall degrading enzymes) is probably sufficient to provoke the disease. Studies on other host plants will allow to determine all the full virulence repertoire of the bacteria. 334

Our results give also some information on the metabolic status of the plant. The amount of nucleic acids and of the amino acids cysteine, leucine, methionine, threonine and isoleucine is too low in chicory to allow the multiplication of mutant bacteria. Some enzymatic steps involved in their synthesis are specific to bacteria and fungi. Thus, they could be good target for the development of specific inhibitors [53] to fight *D. dadantii*.

While Tn-seq has been used to study genes required for the infection of animals, there has been no genome-wide study of factors necessary for a plant pathogen to develop and provoke disease on a plant. Besides the genes of known function described in the Result section, this study allowed the identification of several genes of unknown function required for chicory rotting. Repetition of this experiment with other strains or on other plants will tell if these genes encode strain or host specific virulence factors.

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347 Methods

348 Bacterial strains and growth conditions. Bacterial strains, phages, plasmids and 349 oligonucleotides used in this study are described in Table S1. D. dadantii and E. coli cells 350 were grown at 30 and 37°C respectively in LB medium or M63 minimal medium supplemented with a carbon source (2 g/L). When required antibiotics were added at the 351 352 following concentration: ampicillin, 100 µg/L, kanamycin and chloramphenicol, 25 µg/L. 353 Media were solidified with 1.5 g/L agar. Transduction with phage PhiEC2 was performed according to [54]. The motility of each mutant was compared with that of the wild-type strain 354 355 on semisolid (0.4%) LB agar plates as previously described [55].

356 Construction of the transposon library

Five mL of an overnight culture of *D. dadantii* strain A350 and of *E. coli* MFDpir/pSamEC
were mixed and centrifuged 2 min at 6000 g. The bacteria were resuspended in 1 mL of M63

medium and spread onto a 0.45 µm cellulose acetate filter placed on a M63 medium agar 359 plate. After 8h, bacteria were resuspended in 1 mL M63 medium. An aliquot was diluted and 360 361 spread onto LB agar + kanamycin plates to estimate the efficiency of mutagenesis. The other part was inoculated in 100 mL of LB medium + kanamycin and grown for 24 h at 30°C. To 362 confirm that the bacteria that grew were D. dadantii strains with a transposon but without 363 plasmid pSamEC, we checked that all the grown bacteria were kan^R, amp^S and 364 365 diaminopimelate (DAP) prototrophs (MFDpir is DAP⁻). The bacteria were frozen in 40% 366 glycerol at -80°C and represent a library of about 300 000 mutants.

367 DNA preparation for high-throughput sequencing

368 An aliquot of the mutant library was grown overnight in LB medium + kanamycin. To identify essential genes in LB, the culture was diluted 100-fold in LB and grown for 6 h. To 369 infect chicory, the overnight culture was centrifuged and resuspended at $OD_{600} = 1$ in M63 370 371 medium. Chicories cut in half were inoculated with 10 µL of this bacterial suspension and incubated at 30°C with maximum moist. After 60 h, the rotten tissue was collected and 372 373 filtered through a cheese cloth. The bacteria were collected by centrifugation and washed 374 twice in M63 medium. DNA was extracted from 1.5 mL aliquots of bacterial suspension adjusted to OD₆₀₀1.5 with the Promega Wizard Genomic DNA purification kit. Next steps of 375 376 the DNA preparation methods were adapted from [25]. All DNA gel-extraction were 377 performed onto a blue-light transilluminator of DNA stained with gel-green (Biotium) to avoid DNA mutation and double-stranded breaks. 50 µg of DNA samples were digested with 378 50 U MmeI in a total volume of 1.2 mL for one hour at 37°C according to manufacturer's 379 380 instructions, then heat-inactivated for 20 minutes at 80°C, purified (QIAquick, PCR 381 purification kit Qiagen) and concentrated using a vacuum concentrator to a final volume of 382 25 µL. Digested DNA samples were run on a 1% agarose gel, the 1.0–1.5 kb band containing the transposon and adjacent DNA was cut out and DNA was extracted from the gel according 383

to manufacturer's instructions (Qiaquik Gel Extraction Kit, Qiagen). This allowed recovery 384 of all the fragments containing genomic DNA adjacent to transposons (1201 bp of 385 386 transposable element with 32-34 bp of genomic DNA). A pair of single-stranded 387 complementary oligonucleotides containing an unique 5-nt barcode sequence (LIB AdaptT and LIB AdaptB) was mixed and heated to 100°C, then slowly cooled down in a water bath 388 to obtain double-stranded adaptors with two-nucleotide overhangs. 1 µg DNA of each sample 389 390 was ligated to the barcoded adaptors (0.44 mM) with 2000 U T4 DNA ligase in a final 391 volume of 50 μ L at 16°C overnight. Five identical PCR reactions from the ligation product 392 were performed to amplify the transposon adjacent DNA. One reaction contained 100 ng of 393 DNA, 1 unit of Q5 DNA polymerase (Biolabs), 1X Q5 Buffer, 0.2 mM dNTPs, 0.4 µM of the forward primer (LIB PCR 5, which anneals to the P7 Illumina sequence of the transposon) 394 395 and the reverse primer (LIB PCR 3, which anneals to the P5 adaptor). Only 18 cycles were 396 performed to keep a proportional amplification of the DNA. Samples were concentrated using a vacuum concentrator to a final volume of 25 µL. Amplified DNA was run on a 1.8% 397 398 agarose gel and the 125 bp band was cut-out and gel extracted (OIAquick, PCR purification 399 kit Qiagen). DNA was finally dialysed (MF-Millipore[™] Membrane Filters) for 4 hours. Ouality control of the Tn-seq DNA libraries (size of the fragments and concentration) and 400 401 High-throughput sequencing on HiSeq 2500 (Illumina) was performed by MGX (CNRS 402 sequencing service, Montpellier). 6 DNA libraries were multiplexed on one flow-cell. After 403 demultiplexing, the total number of reads was comprised between 18 and 31 millions (Table 404 1).

405

406 **Bioinformatics analysis:**

407 Raw reads from the fastQ files were first filtered using cutadapt v1.11 [56] and only reads
408 containing the *mariner* inverted left repeat (ACAGGTTGGATGATAAGTCCCCGGTCTT)

409 were trimmed and considered bona fide transposon-disrupted genes. Trimmed reads were 410 then analyzed using a modified version of the TPP script available from the TRANSIT 411 software v2.0.2 [29]. The mapping step was modified to select only reads mapping uniquely 412 and without mismatch in the D. dadantii 3937 genome (Genbank CP002038.1). Then, the 413 counting step was modified to accurately count the reads mapping to each TA site in the 414 reference genome according to the Tn-seq protocol used in this study. Read counts per 415 insertion were normalized using the LOESS method as described in [57]. We next used the 416 TRANSIT software (version 2.0) to compare the Tn-seq datasets. 417

418 Strain construction. To construct the A4277 strain, gene Dda3937_03424 was 419 amplified with the oligonucleotides 19732+ and 19732-. The resulting fragment was 420 inserted into the pGEM-T plasmid (Promega). A *uidA*-kan^R cassette [58]was inserted 421 into the unique AgeI site of the fragment. The construct was recombined into the *D*. 422 *dadantii* chromosome according to [59]. Recombination was checked by PCR.

423

424 Protein techniques. Flagella were prepared from overnight LB grown cells.
425 Bacteria were pelleted, resuspended in 1/10 volume of water and passed 20 fold through
426 a needle on a syringe. Cells and cells debris were removed by centrifugation 5 min at 20
427 000 x g [55]. Proteins were analyzed by SDS-polyacrylamide gel electrophoresis (SDS428 PAGE).

429

430 **Coinoculation experiments**. To determine the competitivity index, the wild type strain 431 and the mutant to test, marked with an antibiotic resistance gene, were grown overnight in 432 M63 + glycerol medium. Bacteria were washed in M63 medium and the OD_{600} was adjusted 433 to 1.0. Bacteria were mixed to a 1:1 ratio and diluted 10-fold. 10 uL of the mixture were

434	inoculated into chicory leaves. The wound was covered with mineral oil and the leaves were
435	incubated at 30 °C at high humidity. After 24 h the rotten tissue was collected, homogenized,
436	diluted in M63 and spread onto LB and LB + antibiotic plates. After 48 h at 30°C, colonies
437	were counted. The competitivity index is the ratio (number of mutant bacteria/number of WT
438	bacteria) in the rotten tissue / (number of mutant bacteria/number of WT bacteria) in the
439	inoculum.
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441	References
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445	Legend of figures
446	
447	Fig. 1. Quality control of the Tn-seq <i>D. dadantii</i> 3937 libraries.
447 448	Fig. 1. Quality control of the Tn-seq <i>D. dadantii</i> 3937 libraries.(A) Frequency and distribution of transposon sequence reads across the entire <i>D. dadantii</i>
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448 449 450 451 452	(A) Frequency and distribution of transposon sequence reads across the entire <i>D. dadantii</i> 3937 genome. The localization of transposon insertions shows no bias throughout the genome of <i>D. dadantii</i> 3937. B) Density plot of \log_2 FC (measured reads/expected reads per gene).
448 449 450 451	 (A) Frequency and distribution of transposon sequence reads across the entire <i>D. dadantii</i> 3937 genome. The localization of transposon insertions shows no bias throughout the genome of <i>D. dadantii</i> 3937. B) Density plot of log₂FC (measured reads/expected reads per gene). Fig 2. Genes identified by Tn-seq exhibiting a fitness change in chicory compared to LB.
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448 449 450 451 452 453 454 455 456	 (A) Frequency and distribution of transposon sequence reads across the entire <i>D. dadantii</i> 3937 genome. The localization of transposon insertions shows no bias throughout the genome of <i>D. dadantii</i> 3937. B) Density plot of log₂FC (measured reads/expected reads per gene). Fig 2. Genes identified by Tn-seq exhibiting a fitness change in chicory compared to LB. Data were obtained with the TRANSIT software. From left to right : (a) classification of the gene with the HMM method performed in the LB dataset. (b) Number of TA sites left after removing 10% of 5' and 3' extremities. (c) Mean read number per gene in the LB or chicory dataset. (d) Ratio of read number between chicory and LB expressed in log₂. (e) p-values

459 Fig 3. Scheme of the purine and pyrimidine biosynthesis pathways in *D. dadantii* that

460 produce XMP (purine metabolism) and UMP (pyrimidine metabolism) from L-

- 461 glutamine.
- 462 In red are indicated the growth defect genes in chicory that pass the permutation test (q-value
- 463 ≤ 0.05). The log₂FC of read numbers between chicory and LB for each gene is indicated in
- 464 bracket. Some genes do not pass the permutation test (in black) but have a strong negative
- 465 log₂FC. PRPP: 5-phosphoribosyl-1-pyrophosphate ; GAR: 5'-phosphoribosyl-1-glycinamide
- 466 ; FGAM: 5'-phosphoribosyl-*N*-formylglycinamide ; AIR: 5'-phosphoribosyl-5-
- 467 aminoimidazole ; CAIR: 5'-phosphoribosyl-5-aminoimidazole carboxylic acid ; SAICAR: 5'-
- 468 phosphoribosyl-4-(*N*-succino-carboxamide)-5-aminoimidazole ; AICAR: 5-aminoimidazole-
- 469 4-carboxamide ribonucleotide ; IMP: inosine monophosphate ; XMP: xanthine
- 470 monophosphate ; UMP: uridine monophosphate.
- 471

472 Fig 4. Examples of essential and important genes revealed by Tn-seq.

473 Number of reads at each transposon location in the sample grown in LB or in chicory. Data

474 are averaged from biological replicates and normalized as described in the method section.

475 Three regions of the genome representative of Tn-seq results are shown, with the predicted

476 genes represented at the bottom of each panel. Peaks represent read number at TA sites.

- 477 Black arrows represent genes that pass the permutation test (q-value ≤ 0.05). (A) Essentiality
- 478 of leucine biosynthetic genes in chicory. (B) Insertions in the 5' region of *rsmC* generate

growth advantage for the bacteria in chicory. (C) Importance of genes involved in motility for

480 growth in chicory.

481 Fig 5. Modification of FliC revealed by Tn-seq analysis and SDS-PAGE. (A) Importance

482 of 6 genes located between *fliA* and *fliC* for growth in chicory. Log₂FC are indicated in

483 bracket. Dda3937_03425 and Dda3937_03426 are duplicated transposase genes that have

484	been removed from the analysis. Black arrow: GD in chicory (q-value ≤ 0.05); white arrow:
485	genes that do not pass the permutation test (q-value > 0.05). (B) Analysis by SDS-PAGE of
486	FliC produced by the wild type (lane 1), the A3422 (lane 2) and A4277 (lane 3) strains.
487	
488	Fig 6. Correlation between the competitivity index (IC) of several mutant strains with
489	the log ₂ FC of the corresponding genes. IC values were determined in chicory leaves as
490	described in Methods. The IC values for <i>dltB</i> , <i>anrB</i> , <i>phoS</i> and <i>phoH</i> are from [60]. Each
491	value is the mean of 5 experiments.
492	
493	Fig S1. Biological reproducibility of the Tn-seq results.
494	Pairs of Tn-seq assay results are compared, with the total number of reads per gene plotted.
495	Analysis of DNA samples corresponding to two independent cultures of the mutant pool
496	grown (A) in LB medium (correlation coefficient $R = 0.72$) and (B) in chicory (correlation
497	coefficient $R = 0.98$). Values represent average numbers of reads per gene from the pairs of
498	biological replicates.
499	
500	Fig S2. Volcano plot of resampling results comparing replicates grown in chicory versus
501	in LB. Significant hits have $q < 0.05$ or $-\log_{10} q > 1.3$. Growth defect (GD) or growth
502	advantage (GA) genes are indicated by a red frame.
503	Fig S3. acrAB are essential in chicory. Number of Tn-seq reads at each insertion site in the
504	acrA acrB region in samples grown in LB or in chicory. Data are averaged from biological
505	replicates and normalized as described in Methods. <i>dnaX</i> which encodes both the tau and

gamma subunits of DNA polymerase is represented by a grey arrow. *dnaX* is essential gene in

507 LB. *acrAB* represented by grey arrows are GD in chicory (q-value ≤ 0.05).

508 Fig S4. Pathogenicity tests on chicory with leucine (A5969) and cytosine (A5968)

509 mutants of D. dadantii.

510 Each leaf was inoculated with 10^6 bacteria. Length of rotten tissue was observed after 24h.

511

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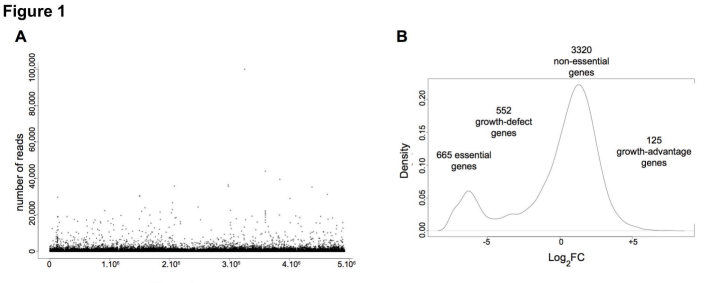
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TA coordinates

rsmC

gcpA

mltD

mrcA

Dda3937_03858

Dda3937 03971

Dda3937_00363

global regulatory protein RsmC

penicillin-binding protein 1A (PBP1A)

outer membrane-bound lytic murein transglycosylase D

hypothetical protein

State Mean reads Locus Gene Function No. of TAs^b Chicory log₂FC^d in LB LB q-value Dda3937 00335 glpD glycerol-3-phosphate dehydrogenase -12.56 Dda3937_03379 -11.91 0.00 purL phosphoribosylformyl-glycineamide synthetase NE 73 378 0 Glucans biosynthesis protein G precursor Dda3937 03564 opgG GA 40 1976 1 -11 41 0.00 Dda3937 00244 purH phosphoribosylaminoimidazolecarboxamide formyltransferase/IMP cyclohydrolase NE 37 145 0 -11.25 0.00 Dda3937_00432 hflK 28 339 FtsH protease regulator GD 0 -11.12 0.03 Dda3937 02515 purM phosphoribosylaminoimidazole synthetase NE 344 0 -10.57 0.00 Dda3937_02627 129 4-hydroxythreonine-4-phosphate dehydrogenase NE 26 0 -10.06 0.00 Dda3937 00004 guaB IMP dehydrogenase NE 33 151 0 -997 0.00 Dda3937 03563 opgH Glucans biosynthesis glucosyltransferase H GA 62 1409 2 -9.79 0.00 17 Dda3937 01284 159 -9.68 0.00 nvrB aspartate carbamovltransferase NE 0 Dda3937_03924 rffG dTDP-glucose 4,6-dehydratase NE 23 317 -9.38 0.02 1 Dda3937 01389 carB carbamoyl-phosphate synthase large subunit NE 48 249 0 -9.23 0.00 Dda3937 03299 MexE family multidrug efflux RND transporter periplasmic adaptor subunit 34 196 0 -9.03 0.00 acrA NE Dda3937_03300 Dda3937_03258 acrR multidrug efflux system protein 89 422 1 -8 90 0.00 NE. 14 **DVr**E orotate phosphoribosyltransferase NE 175 0 -8.81 0.00 33 Dda3937 02336 nlpI lipoprotein GD 27 0 -8.69 0.00 Dda3937 02506 nlpB (bamC) outer membrane protein assembly factor BamC NE 20 47 0 -8.69 0.00 Dda3937 04018 phosphate acetyltransferase GD 36 579 2 -8.59 0.02 pta pyrC Dda3937 03554 dihydro-orotase NE 25 343 1 -8.44 0.00 Dda3937 04573 InxM acyl (myristate) transferase NE 33 63 0 -8.31 0.00 Nitrogen regulation protein NR(I), Two-component system Dda3937 01116 glnG NE 26 39 0 -8.22 0.00 Dda3937 02099 purF amidophosphoribosyltransferase NE 32 107 0 -8.19 0.00 29 Dda3937 04019 ackA acetate kinase A and propionate kinase 2 NE 45 0 -8.16 0.00 Dda3937_02189 yejM Membrane-anchored periplasmic protein, alkaline phosphatase superfamily GA 34 4160 15 -8.08 0.00 Dda3937_01390 carA carbamoyl-phosphate synthase small subunit NE 21 69 0 -8.05 0.00 Phosphoenolpyruvate-protein phosphotransferase of PTS system 45 Dda3937 01426 ntsI NE 0 -785 0.00 Dda3937 00161 cvs0 3'(2'),5'-bisphosphate nucleotidase NE 16 44 0 -7.81 0.02 Dda3937 00210 sulfite reductase beta subunit NE 40 252 -7.65 0.00 cysl Dda3937_04075 -7.51 lysR LysR family transcriptional regulator NE 13 2385 13 0.00 Dda3937 02526 conserved protein NE 18 50 0 -7.50 0.00 Dda3937 03888 metB Cystathionine gamma-synthase NE 118 1 -7.34 0.01 Dda3937 00195 relA 55 -7.12 0.00 (p)ppGpp synthetase I/GTP pyrophosphokinase NE 256 2 Dda3937_02532 Fructose repressor FruR, LacI family NE 15 399 3 -7.04 0.00 lfcR Dda3937_02226 4 fliF Flagellar M-ring protein fliF NE 46 476 -7.02 0.00 Dda3937 02206 flgE Flagellar hook protein flgE NE 50 597 5 -7.00 0.00 Dda3937 04507 phosphogluconate dehydrogenase (NADP(+)-dependent, decarboxylating) GD 36 0 -6.91 0.00 gnd Dda3937 00697 deoO Protease NE 28 80 -6.87 0.01 Dda3937 03631 thioredoxin-disulfide reductase 25 GD 0 -6.85 0.03 trxB 16 Dda3937 00361 yrfF (igaA) intracellular growth attenuator protein GD 38 0 -6.78 0.03 22 Dda3937 00588 cysB Transcriptional dual regulator, O-acetyl-L-serine-binding protein NE 29 90 -6.75 0.00 Dda3937_03783 carboxy-terminal protease for penicillin-binding protein 3 NE 46 243 -6.71 0.00 2 Dda3937 00433 hflX predicted GTPase GD 16 0 -6 69 0.04 33 Dda3937 03427 flagellar filament structural protein (flagellin) NE -6.61 0.03 fliC 96 1 Dda3937_02223 fliI Flagellum-specific ATP synthase fliI NE 42 236 3 -6.56 0.00 29 Dda3937 04419 hdfR DNA-binding transcriptional regulator NE 117 -6.34 0.00 Dda3937_00209 41 -6.25 cvsJ sulfite reductase alpha subunit NE 180 2 0.00 flgH Dda3937 02209 Flagellar L-ring protein flgH NE 23 586 8 -6.22 0.01 Dda3937 02246 fahF beta-ketoacyl-[acyl-carrier-protein] synthase II GD 41 10 0 -6.15 0.00 Dda3937_00301 uvrD ATP-dependent DNA helicase UvrD/PcrA NE 42 29 0 -6.11 0.00 Dda3937 02212 Flagellar hook-associated protein flgK NE 63 116 2 -6.07 0.00 flgK Dda3937 04046 purU Formyltetrahydrofolate deformylase NE 28 51 1 -5.84 0.00 flhA Dda3937 03965 redicted flagellar export pore protein NF 49 106 -5.80 0.00 2 Dda3937 02205 flgD Flagellar basal-body rod modification protein flgD NE 227 4 -5 73 0.01 Dda3937 01352 21 leuC 3-isopropylmalate dehydratase large subunit NE 139 3 -5.73 0.01 Dda3937_02784 20 9 -5.66 flhC Flagellar transcriptional activator flhC NE 477 0.01 Dda3937_02782 motB Flagellar motor rotation protein motB NE 40 109 2 -5.55 0.01 0.00 Dda3937 02210 flgI Flagellar P-ring protein flgI NE 26 163 4 -5.49 Flagellar protein fliJ Dda3937 02222 fliJ NE 14 182 4 -5.44 0.03 Dda3937 02219 fliM Flagellar motor switch protein fliM NE 143 3 -5.40 0.00 Dda3937 02774 32 -5.31 flhB Flagellar biosynthesis protein flhB NE 186 5 0.00 Dda3937 02777 cheB Chemotaxis response regulator protein-glutamate methylesterase CheB NE 31 282 8 -5.14 0.00 Dda3937 02783 motA Flagellar motor rotation protein motA NE 24 39 -5.06 0.00 Dda3937_00565 tonB TonB protein NE. 14 106 3 -5.00 0.05 Dda3937_00427 fructose-bisphosphatase GA 33 805 27 -4 92 0.01 fbp Dda3937_02781 50 -4.89 cheA Chemotaxis protein CheA NE 5 0.00 Dda3937_03422 13 Carbamoyl-phosphate synthase small subunit NE 43 379 -4.85 0.02 Dda3937 02577 diaminopimelate decarboxylase NE 23 332 0 -4.79 0.00 lysA Flagellar basal-body rod protein flgF Dda3937 02207 flgF NE 21 35 -4.76 0.00 Dda3937_02230 fliD Flagellar hook-associated protein fliD NE 47 93 4 75 0.00 3 Dda3937 04301 leuA 2-isopropylmalate synthas NE 36 35 1 -4.69 0.02 Dda3937_02778 cheR Chemotaxis protein methyltransferase CheR NE 30 462 18 -4.67 0.05 Dda3937 02228 fliT Flagellar biosynthesis protein fliT GD 0 -4.63 0.05 16 8 Dda3937_04404 285 -4.63 leuB 3-isopropylmalate dehydrogenas NE 16 12 0.05 Dda3937 02214 fliR Flagellar biosynthesis protein fliR NE 33 268 11 -4 56 0.00 Dda3937 03727 kdul 4-deoxy-L-threo-5-hexosulose-uronate ketol-isomerase NE 26 70 3 -4.54 0.03 Dda3937 03267 O-antigen, teichoic acid lipoteichoic acids export membrane protein ES 107 89 4 -4.33 0.05 Dda3937_00415 16 -4.27 0.02 epd D-erythrose 4-phosphate dehydrogenase NE 26 316 Dda3937 02337 polynucleotide phosphorylase/polyadenylase GD 50 0 -3.97 0.00 pnp Dda3937 01683 purK N5-carboxyaminoimidazole ribonucleotide synthase NE 16 90 0 -3.49 0.01 Dda3937_00689 vrbF (mlaF. predicted toluene transporter subunit GA 9 1254 114 -3.47 0.01 Dda3937 02829 26 helD DNA helicase IV NE 99 9 -3 46 0.01 Dda3937 02252 PTS system glucose-specific IICB component NE 37 81 8 -3.38 0.03 ptsG Dda3937_00726 tolC NE 34 -3.35 0.00 transport channel 184 0 44 Dda3937 02363 clpA ATP-dependent Clp protease ATP-binding subunit NE 64 -3.02 0.03 Dda3937_02470 magnesium and cobalt ions transport 13 159 -2.90 0.02 corC NE 21 Dda3937 00692 vrbC (mlaC) predicted ABC-type organic solvent transporter GA 23 740 106 -2.81 0.01 Dda3937 02045 envC murein hydrolase activator NE 17 71 12 -2.59 0.00 Dda3937 01807 nuoM NADH-quinone oxidoreductase subunit M 29 -2.47 0.03 NE 57 10 Dda3937 03668 sufB Fe-S cluster assembly protein NE 32 116 21 -2.44 0.00 -2.33 Dda3937 02080 trkG Potassium uptake protein NE 36 65 13 0.05 Dda3937_03042 ferrichrysobactin outer membrane receptor NE. 80 244 51 -2.25 0.01 fct Dda3937 01287 argi Ornithine carbamoyltransferase NF 24 279 59 2 22 0.03 Dda3937 02456 221,705

NE

GA 55

NE 46

NE

116

3728

276

85

10.90

5.30

2.47

140.136 5.23

10,885

468

0.028

0.00

0.00

0.021

HMM

RESAMPLING

