

# Differential Genetic Strategies of Burkholderia vietnamiensis and Paraburkholderia kururiensis for Root Colonization of Oryza sativa subsp. japonica and O. sativa subsp. indica , as Revealed by Transposon Mutagenesis Sequencing

Adrian Wallner, Nicolas Busset, Joy Lachat, Ludivine Guigard, Eoghan King, Isabelle Rimbault, Peter Mergaert, Gilles Béna, Lionel Moulin

# ▶ To cite this version:

Adrian Wallner, Nicolas Busset, Joy Lachat, Ludivine Guigard, Eoghan King, et al.. Differential Genetic Strategies of Burkholderia vietnamiensis and Paraburkholderia kururiensis for Root Colonization of Oryza sativa subsp. japonica and O. sativa subsp. indica , as Revealed by Transposon Mutagenesis Sequencing. Applied and Environmental Microbiology, 2022, 88 (14), 10.1128/aem.00642-22 . hal-03747731v1

# HAL Id: hal-03747731 https://hal.inrae.fr/hal-03747731v1

Submitted on 28 Oct 2022 (v1), last revised 18 Jan 2024 (v2)

**HAL** is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



Distributed under a Creative Commons Attribution 4.0 International License

- 1 Title: Differential genetic strategies of Burkholderia vietnamiensis and Paraburkholderia kururiensis for root
- 2 colonization of Oryza sativa ssp. japonica and ssp. indica, as revealed by Tn-seq
- 3 Running title: Tn-seq of burkholderia genes for rice root colonization
- 4 Adrian Wallner<sup>a,\*</sup>, Nicolas Busset<sup>a</sup>, Joy Lachat<sup>b,\*</sup>, Ludivine Guigard<sup>a</sup>, Eoghan King<sup>a</sup>, Isabelle Rimbault<sup>a</sup>,
- 5 Peter Mergaert<sup>b</sup>, Gilles Béna<sup>a</sup>, Lionel Moulin<sup>a,#</sup>
- 6 <sup>a</sup>PHIM Plant Health Institute, Univ Montpellier, IRD, CIRAD, INRAE, Institut Agro, Montpellier, France
- <sup>b</sup>Université Paris-Saclay, CEA, CNRS, Institute for Integrative Biology of the Cell (I2BC), 91198, Gif sur-Yvette, France
- 9 \*Present adresses: AW, Université de Reims Champagne-Ardenne, RIBP EA4707 USC INRAE 1488,
- 10 SFR Condorcet FR CNRS 3417, Reims, France
- JL, Center for Immunology and Infectious Diseases, Cimi-Paris, Inserm, Sorbonne Université, Paris,
   France
- 13
- #Corresponding author: Lionel Moulin, PHIM, 911 Avenue Agropolis 34394 Montpellier cedex 5. Email: <u>lionel.moulin@ird.fr</u>
- 16 Keywords: Rice, *Paraburkholderia kururiensis*, *Burkholderia vietnamiensis*, Root-colonization, Tn-seq,
  17 PGPR.

## 18 Abstract

19 Burkholderia vietnamiensis LMG10929 (Bv) and Paraburkholderia kururiensis M130 (Pk) are bacterial 20 rice growth-promoting models. Besides this common ecological niche, species of the Burkholderia genus 21 are also found as opportunistic human pathogens while Paraburkholderia are mostly environmental and 22 plant-associated species. Here, we compared the genetic strategies used by Bv and Pk to colonize two 23 subspecies of their common host, Oryza sativa ssp. japonica (cv. Nipponbare) and ssp. indica (cv. IR64). 24 We used high-throughput screening of transposon insertional mutant libraries (Tn-seq) to infer which 25 genetic elements have the highest fitness contribution during root surface colonization at 7 days post 26 inoculation. Overall, we detected twice more genes in Bv involved in rice roots colonization compared to 27 Pk, including genes contributing to the tolerance of plant defenses, which suggests a stronger adverse 28 reaction of rice towards Bv compared to Pk. For both strains, the bacterial fitness depends on a higher 29 number of genes when colonizing *indica* rice compared to *japonica*. These divergences in host pressure 30 on bacterial adaptation could be partly linked to the cultivar's differences in nitrogen assimilation. We detected several functions commonly enhancing root colonization in both bacterial strains e.g., Entner-Doudoroff (ED) glycolysis. Less frequently and more strain-specifically, we detected functions limiting root colonization such as biofilm production in Bv and quorum sensing in Pk. The involvement of genes identified through the Tn-seq procedure as contributing to root colonization i.e., ED pathway, c-di-GMP cycling and cobalamin synthesis, was validated by directed mutagenesis and competition with WT strains in rice root colonization assays.

#### 37 Importance

38 Burkholderiaceae are frequent and abundant colonizers of the rice rhizosphere and interesting candidates 39 to investigate for growth promotion. Species of Paraburkholderia have repeatedly been described to 40 stimulate plant growth. However, the closely related Burkholderia genus hosts both beneficial and 41 phytopathogenic species, as well as species able to colonize animal hosts and cause disease in humans. 42 We need to understand to what extent the bacterial strategies used for the different biotic interactions 43 differ depending on the host and if strains with agricultural potential could also pose a threat towards 44 other plant hosts or humans. To start answering these questions, we used here transposon sequencing to 45 identify genetic traits in Burkholderia vietnamiensis and Paraburkholderia kururiensis that contribute to 46 the colonization of two different rice varieties. Our results revealed large differences in the fitness gene 47 sets between the two strains and between the host plants, suggesting a strong specificity in each 48 bacterium-plant interaction.

# 49 Introduction

50 Species of the Burkholderia and closely related Paraburkholderia genera are highly prolific rhizosphere 51 colonizers (1, 2). Their persistence and competitiveness in the rhizosphere environment can be explained 52 by strong secondary metabolite production as well as efficient nitrogen cycling, mineral solubilization and 53 phytohormone biosynthesis (3-5). Furthermore, several Paraburkholderia and at least one Burkholderia 54 species can fix atmospheric nitrogen. Two well studied models, Paraburkholderia kururiensis strain 55 M130 (hereafter called Pk) and Burkholderia vietnamiensis strain LMG10929 (hereafter called Bv) 56 demonstrate strong rice root colonization, endophytic lifestyles and significant plant growth promotion 57 through transfer of fixed nitrogen (6–9). Despite this convergence in their plant beneficial features, both 58 bacteria belong to distinct genetic backgrounds. While Paraburkholderia species are often found in 59 beneficial relationships and symbioses with plants (10-12), Burkholderia members comprise human 60 pathogens and opportunists as well as fungal and plant pathogens (13).

61 Bacterial genes used for plant colonization have been screened in several model bacteria but also on a 62 broader scale using microbiome approaches to reveal plant-associated functions (1). A few studies have 63 compared plant-adapted Burkholderia s.l. (sensu largo, e.g. the former genus that is now regrouping the 64 newly defined Burkholderia, Paraburkholderia, Caballeronia and others) at the genomic level (4, 11), or 65 profiled the transcriptome of bacteria stimulated by root exudates (14, 15). However, there is no record of a comparison between the strategies used by plant-adapted Burkholderia and Paraburkholderia species. 66 67 The host plant genotype's impact on bacterial colonization strategies also remains poorly explored in 68 these bacterial genera. Rice is an interesting model to assess Burkholderia s.l. adaptation to the plant 69 environment as it is hosting plant growth promoting model strains from both Burkholderia and 70 Paraburkholderia genera. It was repeatedly demonstrated that rice genotypes influence the composition 71 of their microbiome at the rhizosphere and rhizoplane levels (16, 17). In particular, a study on 95 different 72 Orvza sativa subsp. indica and subsp. japonica varieties showed significant differences in microbiome 73 composition between both rice subspecies, which was related to their nitrogen use efficiency and the 74 presence of a particular nitrate transporter in indica varieties (18).

Transposon mutagenesis sequencing (Tn-seq), is a high-throughput screening method that combines transposon insertional mutagenesis followed by sequencing of the insertion sites (19). It is a powerful tool that leads to immediate identification of genes of interest improving or reducing the bacteria's fitness in a tested condition. This methodology has been successfully used to unravel important genes and functions in plant-pathogenic or plant-symbiotic interactions (20–22). Commonly between pathogens and mutualists, genes functioning in amino acid and purine metabolism as well as in cell motility were detected to be required for root colonization (21, 23, 24).

82 In the present study, we used Tn-seq, to perform a genome-wide identification of genes involved in rice-83 root colonization in Bv and Pk. In detecting which genes influence the fitness of Bv and Pk we aim at 84 unraveling their commonalities and differences in root colonization strategies. We also analyzed the 85 association strategies of Bv and Pk with the two rice genotypes Oryza sativa subsp. japonica (cv. 86 Nipponbare) and *indica* (cv. IR64) to understand how the host-factor can influence colonization 87 strategies. Overall, we identified a total of 1,404 and 540 genes that influence the fitness of Bv and Pk 88 respectively during rice root colonization. Our results underline the importance of motility, amino acid 89 and vitamin metabolism, stress response as well as biofilms for the efficient association of these bacteria 90 with rice roots.

# 91 Results

92 Quality of Tn-Seq libraries and essential genomes in Pk and Bv

To generate a genome-wide library of insertion mutants for Pk and Bv strains we used a mariner transposon that targets genomic thymine-adenosine (TA) sites (Materials and methods). The genome of 95 Bv contains 95,075 TA sites and we estimated the mutant population at  $1.6 \ge 10^8$  cfu, which represents a 96 1683x coverage of the total TA sites. The saturation level of the *Pk* library was lower although still 97 significant at a 38x coverage given the 4.0  $\ge 10^6$  cfu obtained after transposon mutagenesis for a total of 98 106,136 total TA sites contained in the genome.

99 To further assess the quality of the Tn-seq libraries we determined the essential genomes required by both 100 bacteria for optimal development in a rich liquid growth medium. Both bacteria have similar genome 101 sizes (6,820 and 6,436 genes for By and Pk respectively) and comparable proportions of genes (661 or 9.7% and 638 or 9.9%, respectively) that are required for optimal growth in a rich, liquid medium (Figure 102 103 1A; Supplemental dataset S1). The size of the essential genomes determined in a controlled liquid 104 medium are in the order of magnitude of what has been observed for other prokaryotes, including 105 Burkholderia spp. (26–30). We used the Minimal Gene Set tool implemented in the MicroScope platform 106 (Materials and methods) to extract a core list of the predicted minimal bacterial gene sets from Bv and Pk. 107 On a total of 206 core bacterial genes defined by their conservation among multiple bacterial genomes 108 (25), 151 and 150 were identified as essential by our approach in Bv and Pk, respectively. In 10 and 9 109 cases of genes belonging to the minimal gene set but predicted as non-essential by our approach in Bv and 110 *Pk* respectively, there are duplicates present in the genome.

111 According to their distribution in COG categories, little variance differentiates the essential genomes of 112 Pk and Bv (Figure 1B). As expected in a liquid, rich medium, genes involved in motility, defense and 113 nutrition are largely unessential. Conversely, structural components of the cell and the general replicative 114 machinery are predictably essential (Supplemental dataset S1).

Given the strong saturation level of the TA insertion sites and the coherent essentiality results observed in the control setting, we can safely assume that both mutant libraries allow a reliable analysis of the impact of genes on the bacterial fitness.

#### 118 Colonization of the two rice genotypes by *Pk* and *Bv*

119 Prior to genetic analyses, we assessed the colonization efficiencies of Bv and Pk on Nipponbare 120 (*japonica*) compared to IR64 (*indica*) rice genotypes (Figure 2). Overall, both bacteria display a similar 121 colonization dynamic with an increasing population density during the first week and a decrease in the 122 second week. A host-genotype effect is observed in the colonization phenotype displayed by Pk as 123 significantly different root colonizing populations are observed on IR64 and Nipponbare at 3 dpi and 14 124 dpi (Figure 2). By on the other hand displays a steady colonization pattern between both plant genotypes 125 at all measured time points. These observations confirm that the rice-colonizing populations of Bv and Pk126 can be compared, especially at 7 dpi, which was selected for further analyses. In our following analyses,

we consider that the bacterial adaptations observed result purely from root surface colonization, as the endophytic populations at 7 dpi are inferior by several log scales to the surface colonizing population (6).

#### 129 Identification of Rice colonization genes

130 To assess gene fitness for rice root colonization, and its host-dependent variation in the two model 131 bacteria, we inoculated the Tn-Seq mutant libraries on Nipponbare and IR64 rice genotypes. Seven days 132 after inoculation, we harvested and pooled five root systems per replicate, representing a total of  $1.2 \times 10^7$ 133 colonization events (Figure 2) and a more than 100-fold coverage of the available mutant diversity. We 134 performed a first Tn-seq analysis by pooling the reads of IR64 and Nipponbare isolates together to infer 135 the genes globally required for the association with rice. The read frequencies were compared to a control 136 condition grown in a rich medium with limited growth generations (see Material and Methods) to 137 establish a root fitness score for each gene. In this manner, we identified 1,404 and 540 genes as significantly impacted (enriched or depleted) after root-colonization by Bv and Pk, respectively (Figure 138 **3A**). Colonization-depleted genes will be our major focus as they are positively associated with bacterial 139 140 fitness. Inversely, colonization-enriched genes diminish the bacterial fitness during root colonization. We 141 organized the identified colonization genes according to their general function based on their clusters of 142 orthologous groups (COG) annotation (Figure 3B). Colonization-enriched and -depleted genes share a 143 similar distribution with amino acid metabolism, cell wall/membrane/envelope biogenesis, transcription and cell motility being amongst the most frequent categories, consistent with the expected implication of 144 145 nutrition, motility and morphological adaptations involved in plant colonization.

146 We identified a total of 2,071 core gene families for Bv and Pk, sharing homologues in both strains 147 (Supplemental dataset S2). In Bv and Pk, 192 genes, respectively 24% and 41% of the colonization-148 depleted genes are part of the core-genome and equally depleted in both strains (Figure 4). Bv displays 149 68% more colonization-depleted genes than Pk (Figure 3). Interestingly, a majority of these Bv specific 150 colonization-depleted genes (53%) are part of the core genome (Figure 4). Similarly, a lower, although 151 substantial portion of the genome shared with Bv (37%), is required by Pk specifically for efficient root 152 colonization (Figure 4). Thus, although a large portion of the core genes contribute to root colonization in 153 both strains, many core genes are only required for colonization in one or the other strain, indicating that 154 the impact of these genes on root colonization is dependent on the genome context.

## 155 Rice cultivar dependent colonization specialization

156 Next, we performed a separate analysis of the reads for each rice cultivar and then compared the lists of

- 157 significantly colonization-depleted and -enriched genes to infer cultivar-specificities. For both strains, the
- 158 majority (~65%) of identified genes impacted the bacterial fitness on both IR64 and Nipponbare rice

(Table 1). However, both bacteria displayed a greater number of colonization-depleted genes (41% and
49% higher for *Bv* and *Pk* respectively) on IR64 compared to Nipponbare rice (Table 1).

161 Core genes of Bv and Pk that are colonization-depleted on both cultivars belong primarily to 162 housekeeping categories such as central metabolism, cell cycle control and motility (Figure 5A). The 163 additional genes, depleted during IR64 colonization are chiefly attributed to amino acid metabolism, 164 transcription, cell wall/membrane/envelope biogenesis and energy production in both bacterial strains (Figure 5B & 5C). Interestingly, contrary to the global trend, in Pk the COG categories " replication, 165 166 recombination & repair", "coenzyme transport & metabolism", "carbohydrate metabolism", and 167 "inorganic ion transport" are more strongly impacted on Nipponbare than IR64 (Figure 5C). The same 168 trend is observed in Bv for the COG categories "signal transduction mechanisms" and "coenzyme 169 transport and metabolism" (Figure 5B).

### 170 Tn-seq reveals bacterial functions involved in early rice colonization

171 Many genes significantly impacting the colonization fitness of Bv and Pk are clustered together within 172 operons supporting the validity of our results. In these cases, there is furthermore a strong conservation of 173 either gene enrichment or depletion within the same operon. In several cases explored hereafter, the lack 174 of detection of a complete operon can be explained by the presence of homologues for some of the genes 175 resulting in functional complementation.

#### 176 Functions required for bacterial fitness during rice-root colonization

177 As expected in this kind of colonization assay, mutants affected in motility and chemotaxis functions ( $fl_g$ , 178 flh, fli, mot) were depleted from the root colonizing population (Table 2 & Supplemental dataset S3). 179 For both bacteria, we further detected many genes involved in amino acids (arg, his, ilv, leu, lys, met, pro, 180 ser and trp) and nucleotide synthesis (pur and pyr) that, when mutated, negatively impacted the bacteria's 181 fitness on plants. Genes involved in the synthesis of enzymatic cofactors for amino acid metabolism such 182 as vitamin B1 (thiamin) were also colonization-depleted in both strains. Multiple Tn-seq studies reported 183 that auxotrophy for certain amino acids is disadvantageous for root colonization and can limit plant growth promotion and biocontrol potential (23, 31). 184

Additionally, genes involved in potassium nutrition (kdpA-F) were similarly depleted. Mutants of both Bvand Pk affected in central elements of the Entner-Doudoroff glycolysis (ED) pathway (edd, zwf) suffered a significant fitness decrease on rice. This pathway is involved in the metabolism of gluconate which is not a dominant sugar of rice exudates (32) suggesting an alternative role than sole sugar assimilation for this pathway. The activation of the ED pathway was suggested to be a tolerance strategy towards oxidative stress through the generation of NADPH as essential cofactor for thioredoxins (33, 34). 191 Genes of the type 2 secretion system (T2SS; gspD-M) are the only ones belonging to a macromolecular 192 secretory pathway to be colonization-depleted in both bacteria. While Bv possesses a single T2SS, Pk193 bears two copies (4) out of which only one is colonization-depleted suggesting that these systems are not 194 redundant but are used differently in specific conditions.

195 Several genes involved in DNA maintenance and reparation, i.e., ruvABC, xerCD and recABCD were 196 colonization-depleted in both Bv and Pk while their absence was tolerated in the relatively stress-free 197 control medium (**Supplemental dataset S1**). Thus, rice appears to be inflicting considerable genotoxic 198 stress during the process of colonization. Furthermore, the presence of osmotic stress is exemplified by 199 the depletion of Bv and Pk mutants involved in the synthesis of the osmoprotectant trehalose (otsAB).

200 By seems to suffer additional stress as multiple functions maintaining cell wall and membrane integrity 201 are colonization-depleted. We detected several genes involved in hopanoid synthesis (hpnDEFHJKN), 202 peptidoglycan synthesis (murAI) and maintenance (tolAOR, pal) as well as lytic murein 203 transglycosylation (*mltA*, rlpA and mtgA) to be depleted specifically in Bv mutant populations. This 204 indicates that Bv has an increased requirement to maintain its cellular integrity compared to Pk. 205 Consistent with an increased need in membrane maintenance, the loss of vitamin B7 (biotin) synthesis 206 genes (bioABCDF) had a negative impact on the colonization of Bv. Biotin is a cofactor for many 207 enzymes, especially those involved in fatty acid biosynthesis and amino acid metabolism (35, 36). Roots 208 are known to secrete aromatic phenolic compounds which are toxic to various soil microbes. Accordingly, 209 Bv mutants were depleted during colonization when impacted in genes of the  $\beta$ -ketoadipate pathway 210 (pcaBCDK) which allow to metabolize 4-hydroxybenzoate and protocatechuate. A final sign of the stress 211 By is exposed to during colonization is found in the depletion of mutants for the queuosine synthesis 212 pathway (queACEF, tgt). This hypermodified nucleoside improves translation accuracy, a need that only 213 arose in Bv during rice colonization.

214 While no genes annotated as coding for autotransporter proteins (T5SS) are identified through our 215 approach, the genes tamAB were colonization-depleted in Pk. TamA and TamB can be involved in outer 216 membrane assembly, allowing surface structuration which is essential for adhesion and host-invasion by 217 bacteria in several pathogenic models (37). Genes for vitamin B12 (cobalamine) synthesis are present in 218 both Bv and Pk but were only colonization-depleted in Pk (cobBDHIKLMNQSTUW, cbiB, btur). 219 Intriguingly, several genes of this synthesis pathway were actually colonization-enriched in  $B_{\nu}$ , 220 suggesting an adverse role in colonization for this co-factor involved in a multitude of enzymatic 221 reactions.

222 Functions detrimental for bacterial fitness during rice-root colonization

223 Only two genes were identified to be colonization-enriched in Bv and Pk simultaneously, one of which 224 (AK36 1927/ANSKv1 30041) has an unknown function. The other (*pdeR*) is involved in cyclic di-GMP 225 (c-di-GMP) cycling. In Bv, four other genes having a homologous role in c-di-GMP degradation 226 (phosphodiesterases; Supplemental dataset S3) and two involved in c-di-GMP production (diguanylate 227 cyclases; Supplemental dataset S3) were also colonization-enriched. C-di-GMP levels can play a role in 228 colonization as they were demonstrated to influence biofilm production and motility in Burkholderia species (38). Interestingly, while several genes of the putrescine catabolism (puuBD, speG) were 229 230 colonization-depleted in Bv, the entire cluster responsible for putrescine uptake (potFGHI, puuP) is 231 colonization-enriched. Together, this suggests that putrescine accumulation has a negative impact on root 232 colonization. An accumulation of intracellular putrescine could have an impact on biofilm production as 233 recently suggested (23). Among additional biofilm-related colonization-enriched genes in Bv, we 234 identified a cluster involved in cellulose synthesis (bcsABCE). Another Bv surface element that increases 235 colonization efficiency when mutated is a Tad pilus (tadABCEVZ, rcpAB, flp) suggesting that this 236 attachment or motility mechanism is suitable for an alternative condition than rice colonization. While 237 many genes involved in carbohydrate metabolism in both bacteria were colonization-depleted, Bv mutants were colonization-enriched when impacted in ribose import and catabolism genes (rbsABCK). 238

*Pk* mutants impacted in the ability to sense the bacterial community through quorum sensing (*braR*, *rsaL*) were enriched during colonization as can be expected for organisms that lost the ability to sense population density and reduce growth rates accordingly. Genes belonging to a region enriched in elements of phage origin (ANSKV1\_30083-120) had the same effect when mutated, indicating that the prophage might be induced by the plant colonization process and that its inactivation through mutation benefits the bacteria's fitness.

#### 245 Rice cultivar specific adaptations

246 Genes which are colonization-depleted or -enriched specifically on one cultivar can be more rarely 247 grouped in metabolic pathways and operon structures (Supplemental dataset S4). Still, single genes can 248 have determining impacts on a strain's ecology and metabolism. There are two genes that were 249 colonization-depleted for both Bv and Pk on Nipponbare rice. One is an outer membrane porin of the OmpC family (AK36 1494 / ANSKv1 11218) and the other, amtB, is involved in ammonium uptake 250 251 (Table 3 & Supplemental dataset S4). By mutants for genes involved in phosphate transport (pstAC) and glycerol (ugpABQ) metabolism were depleted on IR64 rice, further suggesting that the metabolic 252 253 requirements of the plant force adaptations on rhizosphere microbes.

Several Bv genes involved in type 6 secretion (*tssGHIJK*) were colonization-depleted on IR64. For Pk, mutants impacted in O-antigen biosynthesis (*rfbABD*) were depleted on IR64. In both cases we can hypothesize that surface elements have significant and diverging repercussions depending on the type of colonized rice-host.

#### 258 Validation of three candidate genes

259 We underlined the importance of various genes involved in the ED pathway, c-di-GMP cycling and 260 cobalamin synthesis among others. To validate their involvement in the colonization of rice roots, we 261 used an insertional mutagenesis approach in Bv and Pk by targeting single copy genes that are key in the 262 respective processes. The 6-phosphogluconate dehydratase Edd is essential for the first step of the ED 263 pathway (39). The metal delivery protein CobW is essential for vitamin B12 synthesis (40). Finally, we 264 chose PdeR, an enzyme with domains predicted to be involved in both c-di-GMP production and 265 degradation. The genomic context of each of these genes, including their fold change between root & 266 rich-medium conditions, is presented in Supplementary Figure S1.

267 Mutants and wild-type strains were used jointly in a competition assay and colonization efficiencies were 268 surveyed at 7 dpi. In Bv, each mutation had the observable impact that had been predicted by the Tn-seq 269 analysis (Figure 6). Disruption of the colonization-depleted gene *edd* reduced *Bv*'s colonization capacity 270 while the deletion of the colonization-enriched gene *pdeR* had the opposite effect. The disruption of *cobW* 271 had been identified in Tn-seq data as having a deleterious impact on Pk but did not significantly alter the 272 root colonization capacity of Bv. The mutagenesis approach confirms that the disruption of cobW has no 273 negative impact on Bv but is required by Pk for efficient root colonization. The only inconsistent 274 observation occurred for the *pdeR* mutant in *Pk*, which was predicted as slightly colonization-enriched by 275 the Tn-seq approach, but its disruption resulted in a colonization deficiency compared to the WT strain.

#### 276 Discussion

277 Understanding which genetic bases are involved in PGPR-cereal interaction is pivotal for a controlled and 278 informed selection of beneficial organisms and a gateway to efficient and directed strain improvement. In 279 rice, the microbiome composition was demonstrated to be significantly modulated by the plant-genotype 280 (16). This suggests a variance in selective pressure forced by the plant onto colonizing microbes. The 281 present Tn-seq analysis demonstrates which genes have the strongest contribution to bacterial fitness in 282 the early steps of rhizosphere colonization of two rice cultivars, Nipponbare and IR64 by two bacteria Pk283 and Bv (Figure 7). At this stage, we cannot exclude that additional genes might be required to colonize 284 more mature plants as the bacteria progress and potentially reach different plant compartments. It is also 285 known that Tn-seq approaches are insensitive to genes whose function is complemented by functional homology either within the same bacteria (gene duplicates, paralogues) or by the surrounding bacterialcommunity, e.g., secretion systems and secreted molecules.

The present study compares insertion mutant libraries that were grown on plant roots for 7 days to libraries grown in a rich liquid medium for a brief period. Thus, caution needs to be taken when interpreting the role of the detected genes in plant interaction. Especially detected nutritional function could be general prerequisites for a sound bacterial development in a poor medium and not specifically involved in a plant colonization task. Still, we find several examples of genes involved in plant interaction

that are supported by independent approaches, as illustrated hereafter.

# 294 Bv and Pk have different strategies for rice root colonization

295 As was previously suggested by microscopy observations and host transcriptomics, Bv and Pk have 296 different approaches to rice colonization (6). The present analysis highlights the respective genetic 297 requirements of By and Pk during their rice colonizing process. Strikingly, a considerable higher number 298 of genes is necessary for a successful colonization by Bv than by Pk, despite similar genome size. This 299 requirement could result from increased plant defenses that Bv is exposed to compared to Pk (6). 300 Supposedly, the pathogenic genetic background and potential of Bv could be responsible for this adverse 301 plant response and necessitate a higher number of genes to act against. Moreover, as Bv is known to colonize multiple eukaryotic hosts, both plants and animals (41–44), we can also hypothesize that Bv302 303 expresses many genes which are not of use for plant colonization, thus a mutation in these genes will 304 increase the bacteria's fitness as the ensuing metabolic cost can be spared. This could explain that plant-305 detrimental genes were found in relatively similar amounts to plant-essential genes in Bv while they are 306 considerably less frequent in Pk (Figure 3A). One striking example is the case of vitamin B12 307 production. This cofactor is among the most complex in bacteria and requires at least 19 enzymatic steps 308 to produce while many enzymes further depend on B12 for their activity (45). Here, the synthesis of B12 309 is beneficial for the fitness of Pk during colonization while it has an adverse effect on Bv. This 310 observation is supported both by the Tn-seq and the insertional mutagenesis approaches. We can 311 hypothesize that the cofactor is involved in enzymatic activities that play a crucial role for colonization in 312 *Pk* but not in *Bv*. As the production of B12 is associated with a high metabolic cost, sparing its expense 313 could explain the fitness advantage displayed by Bv when the B12 synthesis pathway is mutated.

Defects in secretory activities are readily complemented in bacterial communities making the responsible secretion systems opaque for detection by Tn-seq approaches. Indeed, secretion systems are not detected by most studies relying on Tn-seq except when focusing on more sparsely colonized environment such as the apoplast (31). In the present study however, a T2SS is colonization-depleted in Bv and Pk and the type 6 secretion system (T6SS) in Bv during IR64 colonization. While the T6SS can be used for cell adhesion 319 (46), a role beyond complementable secretion remains elusive for the T2SS. Still, as only one T2SS is 320 supposedly involved in colonization it suggests a separate role from the second one present in Pk.

321 The only gene with an identified function that is predicted by the Tn-seq approach to be negatively 322 involved in root colonization in both Bv and Pk is pdeR, Site-directed mutagenesis and a competition 323 assay revealed that its mutation was highly beneficial for the root colonization activity of Bv as expected 324 from the Tn-seq results, but detrimental for Pk, which was inconsistent with the Tn-seq analysis. PdeR is 325 involved in the turnover of the secondary messenger c-di-GMP. In Xanthomonas oryzae, the deletion of 326 pdeR results in a decreased virulence on rice (47). Through its association in two component signal 327 transduction systems, PdeR could have a variety of indirect roles beside c-di-GMP cycling (48). The divergence observed between the Tn-seq and directed mutagenesis for Pk could also be linked to the 328 329 inoculum's concentration which is likely to have been superior in the latter approach and might have 330 triggered a different plant response with the observed detrimental effects on the Pk mutant population 331 (Figure 6).

#### 332 Tn-seq identifies common plant-colonization traits

333 To date, there is no transcriptome data available for Bv that would allow a comparison with our data. 334 However, the transcriptomic response of Pk to rice root macerates has been assessed before (14). Out of 335 the 471 colonization-depleted genes identified by the present approach, 267 are differentially expressed in 336 Pk when stimulated with root macerate (Supplemental dataset S5). Dominantly, these genes are 337 involved in amino acid metabolism, cell motility and cell wall/membrane biogenesis. The prevalence of 338 these functions has further been reported by Tn-seq approaches in other root colonizing models such as 339 Dickeya dadantii and Pseudomonas spp. (21, 23, 24). Our results are congruent with most plant-microbe 340 colonization studies, underlining the importance of genes involved in the production of surface 341 components for cellular attachment to the host (49). Our analysis also presents substantial overlap of 342 colonization-depleted genes with what has been found in the plant growth promoting species Azoarcus 343 olearius and Herbaspirillum seropedicae (50). Notably genes involved in peptidoglycan (ampD) and cell 344 wall (murAI) formation, chemotaxis (cheARW), iron uptake (ferredoxins) and cobalamin synthesis (cbiA, 345 *cobIO*) were detected.

Disruption of the ED pathway was demonstrated to reduce root colonization capacity in *Pseudomonas chlororaphis* with a subsequent loss of ISR stimulation (51). Later works on *Pseudomonas fluorescens* observed that the expression of *edd* was enhanced in the rhizosphere of *Arabidopsis* compared to liquid growing medium (52). Mutants for *edd* in this latter study also failed to stimulate ISR but without the colonization defect observed in *P. chlororaphis*. For pseudomonads, the importance of the ED pathway might be linked to its direct product, pyruvate, that is required for the synthesis of the ISR-eliciting butanediol (51). Other advantages might arise from the ED pathway such as the production of NAD(P)H which is not generated by the EMP glycolysis pathway. This cofactor is used by thioredoxins and could be involved in plant colonization and associated oxidative stress tolerance (33, 34).

355 IR64 rice has more stringent requirements for colonization by *Bv* and *Pk* than Nipponbare rice.

356 For both Bv and Pk, a stronger host-genotype effect on the number of colonization-impacted genes was 357 observed during the association with IR64 compared to Nipponbare rice. For Pk, this genotype-effect is 358 also observed on the root colonization profiles of the two cultivars (Figure 2). One specificity of *indica* 359 rice (IR64) over *japonica* (Nipponbare) is its improved nitrate uptake and assimilation capacity (53), 360 which is linked with the presence of the nitrate transporter NRT1.1B in the former, and proved to impact 361 its microbiota (18). Nipponbare rice preferably imports the alternative nitrogen source ammonium, which 362 can be correlated with the depletion of bacterial mutants impacted in the import of this nutrient (amtB) in the Nipponbare environment in our experimental set-up, as bacteria would compete with rice for the same 363 364 nitrogen source) (Table 3).

365 It was interesting to observe that the T6SS, a major macromolecular system of Bv, was only colonization-366 depleted during inoculation on IR64 roots. The T6SS can be employed by bacteria for competition with 367 other prokaryotes but also interaction with eukaryotes (54). As secretory function should be 368 complemented by the bacterial community, the T6SS of Bv appears to be rather involved in adherence to 369 eukaryotic cells, in a host-dependent manner.

370 Still, most colonization-impacted genes are conserved between rice cultivars and several functions such as 371 motility, amino acid metabolism and biofilm production have been repeatedly described for their role in 372 the general association of bacteria with plants (Figure 7). We have found evidence of an increased stress 373 to which bacteria are exposed in the near vicinity of plants through the depletion of mutants involved in 374 osmoprotection, toxic compound degradation and DNA reparation. The identification of several functions 375 that are part of the core-genome shared by both bacteria but are only colonization-impacted in one strain 376 reinforces the validity of these observations and the different adaptations that both bacteria must undergo 377 during colonization of rice. This is especially true in the present system as Bv and Pk induce different levels of plant defenses (6). 378

An increasing amount of plant microbiome studies rely on Tn-seq to reveal the nature of genes underlying root colonization (20, 21, 23, 24, 31, 50). Tn-seq further offers the benefit over more common methods such as RNAseq, to inform on the genes obstructing colonization (mutants with higher colonization fitness), thus presenting a more complete catalogue of factors driving host-bacteria compatibility. Together with other trending methods such as microfluidic visualization technics and synthetic bacterial communities (49), Tn-seq proves here to be a powerful tool for a better understanding of the genetic bases underlying colonization of different hosts.

## 386 Materials and Methods

#### 387 Bacteria and plant culture conditions

388 The strains used in this study are Burkholderia vietnamiensis LMG 10929 (Bv) and Paraburkholderia 389 kururiensis M130 (Pk), either wild type, or modified by insertion of the pIN29 plasmid conferring 390 chloramphenicol (Cm) resistance and DsRed fluorescence, or spontaneous rifampicin (Rif) and 391 spectinomycin (Spt) resistant strains, or insertional mutants of a non-replicative plasmid in several 392 candidate genes in each strain (Table 4) (6, 55). Spontaneous antibiotic resistant strains for LMG 10929 393 and M130 were obtained by plating 100  $\mu$ L of bacterial liquid culture at OD<sub>600</sub>=1.0 on Luria's low salt LB (LBm; Sigma-Aldrich) with either rifampicin (30 µg.mL<sup>-1</sup>) or spectinomycin (50 µg.mL<sup>-1</sup>). After 48h 394 395 incubation at 28°C, single colonies were selected and streaked on fresh LBm plates with rifampicin or 396 spectinomycin, then grown in broth LBm with antibiotics and stored in 20% glycerol at -80°C. From 397 glycerol stocks, bacterial strains were cultured as follows: bacterial cells conserved at -80°C were plated on LBm agar plates (with antibiotic for mutants) and incubated for 72 h at 28°C. Single colonies were 398 399 used to inoculate 10 mL of LBm broth (with antibiotic for mutants) in 50 mL Falcon tubes and incubated for various amounts of time allowing the different strains to reach an  $OD_{600}=1.0$ . For inoculation 400 purposes, cultures were adjusted to  $5 \times 10^7$  cfu.ml<sup>-1</sup>. 401

Oryza sativa subsp. japonica cv. Nipponbare and Oryza sativa subsp. indica cv. IR64 were cultured as 402 403 described in King et al. (2019). Briefly, seeds were sterilized using successive 70% ethanol and 3.6% 404 sodium hypochlorite treatments and germinated seedlings were transferred onto sterile perlite in an 405 airtight hydroponic system. 5-days old seedlings were inoculated with 1 mL of bacterial solution at 5 x 10<sup>7</sup> cfu.ml<sup>-1</sup> and grown up to 14 days at 28°C (16h light, 8h dark). For competition assays, five plants 406 were inoculated with a mix at  $1.10^7$  cfu.ml<sup>-1</sup> containing 0.5 mL of SptR Bv or Pk strains (previously 407 grown separately in broth LBm + Spt 50 µg.mL<sup>-1</sup> and washed to remove antibiotic) and 0.5 mL of either 408 one of the insertional mutants ( $\Omega cobW$ ,  $\Omega pdeR$  or  $\Omega edd$ ) of the corresponding strain (grown in LBm + 409 Cm 100  $\mu$ g.mL<sup>-1</sup> and washed to remove antibiotic). 410

411 Estimation of root colonizing bacterial population

For the estimation of root colonizing populations, the plants were prepared as described above and inoculated with WT strains at  $1 \times 10^7$  cfu.ml<sup>-1</sup>. Three systems (15 plants) were prepared for each condition and grown for 1, 7 or 14 days (28°C; 16 h light; 8 h dark). At each time point, five plants were harvested 415 for each condition. The entire root systems were sampled and separately transferred to screw cap tubes 416 containing 1 mL of sterile water and a sterile ceramic bead. Roots were weighted and then pulverilzed using a Fastprep-24 (MPbio) at 6 m.s<sup>-1</sup> for 40 seconds. A serial dilution of the resulting solution was 417 418 spotted out on square LBm plates and incubated at 28°C for 48 hours. The number of colony forming units 419 (cfu) were counted and adjusted to the weight of the root systems. The mean colonization values were 420 compared between cultivars and between bacteria genotypes at each time point and for each bacterium 421 separately using a Wilcoxon test. Results were considered significantly different at p < 0.05. For the 422 competition assay between WT and mutant strains, plants were harvested at 7 dpi and processed as 423 described above, except the bacterial dilutions were spotted on LBm plates containing either streptomycin 424 (30 µg.ml<sup>-1</sup>) or Cm (200 µg.ml<sup>-1</sup>) to select respectively the WT or the mutant strains.

#### 425 Tn-seq library preparation

426 A Himar1 mariner transposon carried by the pSAM EC vector in *Escherichia coli* strain MFDpir (56, 57) was introduced into Paraburkholderia kururiensis M130 Rif<sup>R</sup> and Burkholderia vietnamiensis LMG 427 428 10929 Rif<sup>R</sup> through conjugation. Both donor and recipient were grown until  $OD_{600}=1.0$  in liquid low-salt LB (LBm) and the medium of MFDpir was further supplemented with diaminopimelic acid (DAP; 300 429  $\mu$ g.mL<sup>-1</sup>). Cells were spun down and washed once with LBm and concentrated to a final OD<sub>600</sub>=50. Donor 430 and recipient strains were mixed 1:1 and 50 µL suspensions were spotted on square Petri dishes 431 containing LBm and DAP. Growth rates were different for both recipient species and conjugation time 432 433 was adapted accordingly. The mating mix was incubated for 6 h and 24 h at 28°C for LMG 10929 and 434 M130, respectively. The spots where then resuspended in 2 mL LBm per plate. The mating mix was further spread on LBm Petri dishes with rifampicin (30 µg.mL<sup>-1</sup>) and kanamycin (50 µg.mL<sup>-1</sup>) and 435 incubated at 28°C. This positively selects for recipient strains having integrated the transposon and 436 437 negatively selects for the DAP auxotroph E. coli donor. After growth, the colonies were resuspended in 1 438 mL LBm per plate. The library was separated into 1 mL aliquots, and stored in 20% glycerol at -80°C. 439 Additionally, a serial dilution of the mating mix followed by spreading on LBm with rifampicin (30  $\mu$ g.mL<sup>-1</sup>) and kanamycin (50  $\mu$ g.mL<sup>-1</sup>) was used to estimate the abundance of mutants in the library 440 441 through cfu counting.

## 442 Tn-seq experimental setup

Tn-seq libraries were thawed on ice and diluted in sterile water to  $OD_{600}=1$  and then inoculated in 50 mL liquid LBm with rifampicin (30 µg.mL<sup>-1</sup>) and kanamycin (50 µg.mL<sup>-1</sup>) at a final  $OD_{600}=0.1$  and grown at 28°C 150 rpm until  $OD_{600}=1.0$ . One part was conserved for plant inoculation while the other was centrifuged at top speed for 10 minutes and flash frozen for subsequent DNA extraction to serve as control condition. Plants were grown in hydroponic systems as described above. After 5 days of growth, each plant was inoculated with 1 mL Tn-seq library bacterial suspension at 5 x  $10^7$  cfu.mL<sup>-1</sup>. The experimentation was performed in triplicates with each replicate consisting of five plants. At 7 dpi, the roots were harvested and placed in TE buffer.

#### 451 DNA extraction and sequencing methods

The bacterial genomic DNA isolation using CTAB protocol (JGI) was used for the extraction of bacterial DNA from the control and experimental conditions. For the latter, whole roots were used in the first stages of the protocol. Whole roots were immersed in the extraction solution and vortexed for 5 min at each step of the protocol to facilitate bacterial separation. Root grinding was avoided to prevent excessive DNA contamination from the plant material. Roots were removed from the extraction buffer after the lysis steps, before CTAB is added.

458 Ten µg of total DNA were digested with 1 µL of MmeI restriction enzyme (New England Biolabs) in 250 459 µL total volume with 10 µL of S-adenosyl methionine (SAM) 1.5 mM and 25 µL CutSmart buffer during 1 h at 37°C. Then, 1 μL of FastAP Thermosensitive Alkaline Phosphatase 1 U.μL<sup>-1</sup> (ThermoScientific) 460 461 was added and incubated 1h at 37°C. All enzymes were inactivated through a 5 min incubation at 75°C. 462 Digested DNA was column purified using a NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel). 2 µg of purified DNA was ligated to adapters containing specific barcode sequences. Adaptors were 463 464 obtained by annealing the primers 5'-TTCCCTACACGACGCTCTTCCGATCTXXXXXNN-3' and 5'-465 YYYYAGATCGGAAGAGCGTCGTGTAGGGAAAGAGT-3' where NN are random nucleotides for 466 annealing to the dinucleotide overhang generated by MmeI and XXXXX and YYYYY are complementary, barcode-specific sequences. Ligations were performed using 1 µL of 5 mM adapters, 1.5 467 µL of T4 DNA ligase 1 Weiss U.µL<sup>-1</sup> (ThermoScientific) and 2.5 µL 10x ligase buffer in a total volume 468 469 of 25 µL and incubated overnight at 16°C. 1 µL of ligation product was amplified using PCR with a GO 470 Taq DNA (P7 5'polymerase (Promega) and Illumina primers 471 CAAGCAGAAGACGGCATACGAGATAGACCGGGGACTTATCATCCAACCTGT-3'; **P5** 5'-472 AATGATACGGCGACCACCGAGATCTACACTCTTTCCCTACACGACGCTCTTCCGATCT-3'). 22 473 PCR cycles were run (30 s at 92°C; 30 s at 60°C; 1 min at 72°C) with an initial step at 92°C for 2 min and 474 a final step at 72°C for 10 min. PCR products were subjected to a final gel purification using the 475 NucleoSpin Gel and PCR Clean-up kit (Macherey-Nagel).

476 Sequencing was performed at the high-throughput sequencing platform at I2BC (CNRS, Gif-sur-Yvette, 477 France) using Nextseq500 (Illumina) technology and 80 sequencing cycles (single read). Preliminary data 478 analysis (demultiplexing, trimming and mapping) was performed by the sequencing platform. Sequences 479 were trimmed from barcodes and mariner transposon sequences and deposited in European Nucleotide 480 Archive under Bioproject number PRJEB42565.

#### 481 Tn-seq data analysis

Gene essentiality for optimal growth in a liquid LBm broth under agitation was assessed through a hidden Markov model (HMM)-based analysis using the EL-ARTIST pipeline (window size of 7 TA sites; *P* value of 0.03) (58). The pipeline predicts whether a TA site is located in an essential or nonessential region. In some cases, the locus is predicted to be located in a region that contains both domains that are required and dispensable for growth. After the first refinement, the HMM will recalculate the transition probabilities based on the new data and repeat the run until the algorithm reaches convergence (see **Supplemental dataset S6** for TA sites transposon insertion cover in each gene and condition).

For the identification of conditionally essential genetic regions, the sorted aligned sequences were first normalized using TRANSIT 3.1.0 (59) with the Trimmed Total Reads method at default settings and an additional LOESS correction. The normalized datasets were then analyzed using the TnseqDiff function of the R package Tnseq (60). Adjusted p-values were calculated using the Benjamini & Hochberg correction. Genetic loci were considered colonization depleted when their occurrence in the experimental condition was at least 1.5-fold lower than in the control condition at a confidence level of padj<0.05 and inversely for colonization enriched genes.

#### 496 Comparative genomics procedures

497 Core genome compositions were calculated using the Phyloprofile exploration tool implemented in the 498 MicroScope microbial genome annotation and analysis platform (61). Homology constraints were set at 499 minLrap  $\ge 0.8$  and identity  $\ge 50\%$ . We used the COGNiTOR pre-computed COG category classification 500 available on MicroScope to infer COG category affiliations of *Bv* and *Pk* genes.

#### 501 Construction of insertional mutants

502 For the production of the cobW, pdeR or edd insertional mutants of B. vietnamiensis LMG10929 and P. 503 kururiensis M130, a gene fragment of either cobW, pdeR and edd of each strain was amplified using 504 specific primers (Supplementary Table S1). Genes fragments were inserted into the target vector 505 pSHAFT2 (Shastri et al., 2017) carrying a chloramphenicol (Cm) resistance gene using an XbaI/XhoI 506 double digestion. Insert and plasmid were mixed at a 5:1 ratio with regard to their respective molecular 507 size, ligated using a T4 DNA ligase (Promega) and cloned into heat-shock competent *E. coli* DH5a cells. 508 Positive cells were multiplied and their plasmid extracted using the Wizard Minipreps DNA purification 509 system (Promega). Plasmids were electroporated into WT B. vietnamiensis LMG10929 and P. kururiensis 510 M130 strains. Transformed cells were transferred in LBm for 3h at 30°C and then spotted on LBm plates containing Cm (200 µg.ml<sup>-1</sup>) for selection of mutants. Plasmid insertion in each targeted gene was 511

512 checked by PCR. Mutants and WT strains showed identical growth rates in broth LBm or Hoagland 513 medium supplemented with glucose (not shown).

514

# 515 Acknowledgments

The authors acknowledge the IRD itrop HPC (South Green Platform; https://bioinfo.ird.fr http://www.southgreen.fr) at IRD Montpellier for providing HPC resources that have contributed to the research results reported here. We acknowledge the High-throughput sequencing facility of I2BC for its sequencing and bioinformatics expertise. The LABGeM (CEA/Genoscope & CNRS UMR8030), the France Génomique and French Bioinformatics Institute national infrastructures are acknowledged for support within the MicroScope annotation platform.

# 522 Funding

523 The authors gratefully acknowledge financial support from the CGIAR research program (CRP) RICE as

524 well as from the French national research agency (ANR) funding the BURKADAPT project (ANR-19-

525 CE20-0007). AW, JL and LG were supported by PhD fellowships from the French Ministry of Higher

526 Education, Research and Innovation.

# 527 References

- Levy A, Gonzalez IS, Mittelviefhaus M, Clingenpeel S, Paredes SH, Miao J, Wang K, Devescovi G, Lebeis
   S, Jin Z, Mcdonald M, Klein AP, Meghan E. 2018. Genomic features of bacterial adaptation to plants. Nat
   Genet 50:138–150.
- 531 2. Compant S, Nowak J, Coenye T, Clement C, Ait Barka E. 2008. Diversity and occurrence of *Burkholderia*532 spp. in the natural environment. FEMS Microbiol Rev 32:607–626.
- Suárez-moreno ZR, Caballero-mellado J, Coutinho BG. 2012. Common features of environmental and
   potentially beneficial plant-associated *Burkholderia*. Microb Ecol 63:249–266.
- 535 4. Dias GM, de Sousa Pires A, Grilo VS, Castro MR, de Figueiredo Vilela L, Neves BC. 2019. Comparative
   536 genomics of *Paraburkholderia kururiensis* and its potential in bioremediation, biofertilization, and
   537 biocontrol of plant pathogens. Microbiologyopen 8:e00801.
- 538 5. Wallner A, King E, Ngonkeu ELM, Moulin L, Béna G. 2019. Genomic analyses of *Burkholderia* 539 *cenocepacia* reveal multiple species with differential host-adaptation to plants and humans. BMC Genomics
   540 20:803.
- King E, Wallner A, Rimbault I, Barrachina C, Klonowska A, Moulin L, Czernic P. 2019. Monitoring of rice
  transcriptional responses to contrasted colonizing patterns of phytobeneficial Burkholderia s.l. reveals a
  temporal shift in JA systemic response. Front Plant Sci 10:1141.

- Trân Van V, Berge O, Ngô Kê S, Balandreau J, Heulin T. 2000. Repeated beneficial effects of rice
  inoculation with a strain of *Burkholderia vietnamiensis* on early and late yield components in low fertility
  sulphate acid soils of Vietnam. Plant Soil 218/2:273–284.
- Mattos KA, Pádua VLM, Romeiro A, Hallack LF, Neves BC, Ulisses TMU, Barros CF, Todeschini AR,
   Previato JO, Mendonça-Previato L. 2008. Endophytic colonization of rice (*Oryza sativa* L.) by the
   diazotrophic bacterium *Burkholderia kururiensis* and its ability to enhance plant growth. An Acad Bras
   Cienc 80:477–493.
- 9. Baldani V, Oliveira E, Balota E, Baldani J, Kirchhof G, Dobereiner J. 1997. *Burkholderia brasilensis* sp.
  nov., uma nova espécie de bactéria diazotrófica endofítica. An Acad Bras Cienc 69.
- Bournaud C, de Faria SM, dos Santos JMF, Tisseyre P, Silva M, Chaintreuil C, Gross E, James EK, Prin Y,
  Moulin L. 2013. *Burkholderia* species are the most common and preferred nodulating symbionts of the
  Piptadenia Group (Tribe Mimoseae). PLoS One 8:e63478.
- Angus AA, Agapakis CM, Fong S, Yerrapragada S, Estrada-de los Santos P, Yang P, Song N, Kano S,
  Caballero-Mellado J, de Faria SM, Dakora FD, Weinstock G, Hirsch AM. 2014. Plant-associated symbiotic *Burkholderia* species lack hallmark strategies required in mammalian pathogenesis. PLoS One 9:e83779.
- Estrada-de los Santos P, Palmer M, Chávez-Ramírez B, Beukes C, Steenkamp E, Briscoe L, Khan N, Maluk
  M, Lafos M, Humm E, Arrabit M, Crook M, Gross E, Simon M, dos Reis Junior F, Whitman W, Shapiro N,
  Poole P, Hirsch A, Venter S, James E. 2018. Whole genome analyses suggests that *Burkholderia* sensu lato
  contains two additional novel genera (*Mycetohabitans* gen. nov., and *Trinickia* gen. nov.): Implications for
  the evolution of diazotrophy and nodulation in the Burkholderiaceae. Genes (Basel) 9:389.
- 564 13. Eberl L, Vandamme P. 2016. Members of the genus *Burkholderia*: good and bad guys. F1000Research
  565 5:1007.
- Coutinho BG, Licastro D, Mendonça-Previato L, Cámara M, Venturi V. 2015. Plant-Influenced gene
   expression in the rice endophyte *Burkholderia kururiensis* M130. Mol Plant-Microbe Interact 28:10–21.
- Klonowska A, Melkonian R, Miché L, Tisseyre P, Moulin L. 2018. Transcriptomic profiling of
   *Burkholderia phymatum* STM815, *Cupriavidus taiwanensis* LMG19424 and *Rhizobium mesoamericanum* STM3625 in response to *Mimosa pudica* root exudates illuminates the molecular basis of their nodulation
   competitiveness and symbiotic evolutionary history. BMC Genomics 19:1–22.
- Edwards J, Johnson C, Santos-Medellín C, Lurie E, Podishetty NK, Bhatnagar S, Eisen JA, Sundaresan V,
  Jeffery LD. 2015. Structure, variation, and assembly of the root-associated microbiomes of rice. Proc Natl
  Acad Sci U S A 112:E911–E920.
- 575 17. Shenton M, Iwamoto C, Kurata N, Ikeo K. 2016. Effect of wild and cultivated rice genotypes on rhizosphere
  576 bacterial community composition. Rice 9:42.
- 577 18. Zhang J, Liu YX, Zhang N, Hu B, Jin T, Xu H, Qin Y, Yan P, Zhang X, Guo X, Hui J, Cao S, Wang X,
  578 Wang C, Wang H, Qu B, Fan G, Yuan L, Garrido-Oter R, Chu C, Bai Y. 2019. NRT1.1B is associated with
  579 root microbiota composition and nitrogen use in field-grown rice. Nat Biotechnol 37:676–684.
- 580 19. van Opijnen T, Bodi KL, Camilli A. 2009. Tn-seq: High-throughput parallel sequencing for fitness and

- 581 genetic interaction studies in microorganisms. Nat Methods 6:767–772.
- 582 20. Cole BJ, Feltcher ME, Waters RJ, Wetmore KM, Mucyn TS, Ryan EM, Wang G, Ul-Hasan S, McDonald
- 583 M, Yoshikuni Y, Malmstrom RR, Deutschbauer AM, Dangl JL, Visel A. 2017. Genome-wide identification
   584 of bacterial plant colonization genes. PLOS Biol 15:e2002860.
- Royet K, Parisot N, Rodrigue A, Gueguen E, Condemine G. 2019. Identification by Tn-seq of *Dickeya dadantii* genes required for survival in chicory plants. Mol Plant Pathol 20:287–306.
- Wheatley RM, Ford BL, Li L, Aroney STN, Knights HE, Ledermann R, East AK, Ramachandran VK, Poole
  PS. 2020. Lifestyle adaptations of *Rhizobium* from rhizosphere to symbiosis. Proc Natl Acad Sci U S A
  117:23823–23834.
- Liu Z, Beskrovnaya P, Melnyk RA, Hossain SS, Khorasani S, O'Sullivan LR, Wiesmann CL, Bush J,
  Richard JD, Haney CH. 2018. A genome-wide screen identifies genes in rhizosphere-associated *Pseudomonas* required to evade plant defenses. MBio 9:e00433-18.
- Sivakumar R, Ranjani J, Vishnu US, Jayashree S, Lozano GL, Miles J, Broderick NA, Guan C,
  Gunasekaran P, Handelsman J, Rajendhran J. 2019. Evaluation of inseq to identify genes essential for *Pseudomonas aeruginosa* PGPR2 corn root colonization. G3 Genes, Genet 9:651–661.
- 596 25. Gutierrez MG, Yoder-Himes DR, Warawa JM. 2015. Comprehensive identification of virulence factors
   597 required for respiratory melioidosis using Tn-seq mutagenesis. Front Cell Infect Microbiol 5:78.
- Moule MG, Spink N, Willcocks S, Lim J, Guerra-Assunção JA, Cia F, Champion OL, Senior NJ, Atkins
  HS, Clark T, Bancroft GJ, Cuccui J, Wren BW. 2016. Characterization of new virulence factors involved in
  the intracellular growth and survival of *Burkholderia pseudomallei*. Infect Immun 84:701–710.
- Wong Y-C, Abd El Ghany M, Naeem R, Lee K-W, Tan Y-C, Pain A, Nathan S. 2016. Candidate essential
  genes in *Burkholderia cenocepacia* J2315 identified by genome-wide TraDIS. Front Microbiol 7:1288.
- 603 28. Gislason AS, Turner K, Domaratzki M, Cardona ST. 2017. Comparative analysis of the *Burkholderia* 604 *cenocepacia* K56-2 essential genome reveals cell envelope functions that are uniquely required for survival
   605 in species of the genus *Burkholderia*. Microb Genomics 3:e000140.
- 606 29. Grazziotin AL, Vidal NM, Venancio TM. 2015. Uncovering major genomic features of essential genes in
  607 Bacteria and a methanogenic Archaea. FEBS J 282:3395–3411.
- 608 30. Gil R, Silva FJ, Peretó J, Moya A. 2004. Determination of the core of a minimal bacterial gene Set.
  609 Microbiol Mol Biol Rev 68:518–537.
- 610 31. Helmann TC, Deutschbauer AM, Lindow SE. 2019. Genome-wide identification of *Pseudomonas syringae*611 genes required for fitness during colonization of the leaf surface and apoplast. Proc Natl Acad Sci U S A
  612 116:18900–18910.
- Suzuki K, Okazaki K, Tawaraya K, Osaki M, Shinano T. 2009. Gas chromatography–mass spectrometry
  associated global analysis of rice root exudates under aseptical conditions. Soil Sci Plant Nutr 55:505–513.
- 615 33. Chavarría M, Nikel PI, Pérez-Pantoja D, De Lorenzo V. 2013. The Entner-Doudoroff pathway empowers
  616 *Pseudomonas putida* KT2440 with a high tolerance to oxidative stress. Environ Microbiol 15:1772–1785.
- 617 34. Cumming RC, Andon NL, Haynes PA, Park M, Fischer WH, Schubert D. 2004. Protein disulfide bond

- formation in the cytoplasm during oxidative stress. J Biol Chem 279:21749–21758.
- 619 35. Streit WR, Entcheva P. 2003. Biotin in microbes, the genes involved in its biosynthesis, its biochemical role
  620 and perspectives for biotechnological production. Appl Microbiol Biotechnol 61:21-31.
- 621 36. Salaemae W, Booker GW, Polyak SW. 2016. The role of biotin in bacterial physiology and virulence: a
  622 novel antibiotic target for *Mycobacterium tuberculosis*. Microbiol Spectr 4.
- Selkrig J, Belousoff MJ, Headey SJ, Heinz E, Shiota T, Shen HH, Beckham SA, Bamert RS, Phan MD,
  Schembri MA, Wilce MCJ, Scanlon MJ, Strugnell RA, Lithgow T. 2015. Conserved features in TamA
  enable interaction with TamB to drive the activity of the translocation and assembly module. Sci Rep 5:1–
  12.
- Richter AM, Fazli M, Schmid N, Shilling R, Suppiger A, Givskov M, Eberl L, Tolker-Nielsen T. 2019. Key
  players and individualists of Cyclic-di-GMP signaling in *Burkholderia cenocepacia*. Front Microbiol
  10:3286.
- 630 39. Peekhaus N, Conway T. 1998. What's for dinner?: Entner-Doudoroff metabolism in *Escherichia coli*. J
  631 Bacteriol 180:3495–3502.
- Crouzet J, Levy-Schil S, Cameron B, Cauchois L, Rigault S, Rouyez MC, Blanche F, Debussche L, Thibaut
  D. 1991. Nucleotide sequence and genetic analysis of a 13.1-kilobase-pair *Pseudomonas denitrificans* DNA
  fragment containing five cob genes and identification of structural genes encoding Cob(I)alamin
  adenosyltransferase, cobyric acid synthase, and bifunctional cobinamide kinase-cobinamide phosphate
  guanylyltransferase. J Bacteriol 173:6074–6087.
- 637 41. Govindarajan M, Balandreau J, Muthukumarasamy R, Revathi G, Lakshminarasimhan C. 2006. Improved
  638 yield of micropropagated sugarcane following inoculation by endophytic *Burkholderia vietnamiensis*. Plant
  639 Soil 280:239–252.
- Gillis M, Van Van T, Bardin R, Goor M, Hebbar P, Willems A, Segers P, Kersters K, Heulin T, Fernandez
  MP. 1995. Polyphasic taxonomy in the genus *Burkholderia* leading to an emended description of the genus
  and proposition of *Burkholderia vietnamiensis* sp. nov. for N2-fixing isolates from rice in Vietnam. Int. J.
  Syst. Bacteriol., 45:274-289.
- Mahenthiralingam E, Bischof J, Byrne SK, Radomski C, Davies JE, Av-Gay Y, Vandamme P. 2000. DNABased diagnostic approaches for identification of *Burkholderia cepacia* complex, *Burkholderia vietnamiensis*, *Burkholderia multivorans*, *Burkholderia stabilis*, and *Burkholderia cepacia* genomovars I and
  III. J Clin Microbiol 38:3165–3173.
- Tang S-Y, Hara S, Melling L, Goh K-J, Hashidoko Y. 2010. *Burkholderia vietnamiensis* isolated from root
  tissues of nipa palm (Nypa fruticans) in Sarawak, Malaysia, proved to be its major endophytic nitrogenfixing bacterium. Biosci Biotechnol Biochem 74:1972–1975.
- 45. Raux E, Schubert HL, Warren MJ. 2000. Biosynthesis of cobalamin (vitamin B12): a bacterial conundrum.
  652 Cell Mol Life Sci C 2000 5713 57:1880–1893.
- 46. Lertpiriyapong K, Gamazon ER, Feng Y, Park DS, Pang J, Botka G, Graffam ME, Ge Z, Fox JG. 2012.
- 654 *Campylobacter jejuni* type VI secretion system: Roles in adaptation to deoxycholic acid, host cell adherence,

- 655 invasion, and in vivo colonization. PLoS One 7: e42842.
- 47. Yang F, Xue D, Tian F, Hutchins W, Yang CH, He C. 2019. Identification of c-di-GMP signaling
  components in xanthomonas oryzae and their orthologs in xanthomonads involved in regulation of bacterial
  virulence expression. Front Microbiol 10:1402.
- 48. Yang F, Tian F, Sun L, Chen H, Wu M, Yang C-H, He C. 2012. A novel two-omponent system PdeK/PdeR
  regulates c-di-GMP turnover and virulence of *Xanthomonas oryzae* pv. *oryzae*. Mol Plant Microbe
  Interact 25:1361-9.
- Knights HE, Jorrin B, Haskett TL, Poole PS. 2021. Deciphering bacterial mechanisms of root colonization.
  Environ Microbiol Rep. 13:428-444.
- 50. Do Amaral FP, Tuleski TR, Pankievicz VCS, Melnyk RA, Arkin AP, Griffitts J, Tadra-Sfeir MZ, Maltempi
  de Souza E, Deutschbauer A, Monteiro RA, Staceya G. 2020. Diverse bacterial genes modulate plant root
  association by beneficial bacteria. MBio 11:1–15.
- Kim HJ, Nam HS, Anderson AJ, Yang KY, Cho BH, Kim YC. 2007. Mutation in the edd gene encoding the
  6-phosphogluconate dehydratase of Pseudomonas chlororaphis O6 impairs root colonization and is
  correlated with reduced induction of systemic resistance. Lett Appl Microbiol 44:56–61.
- 670 52. Cheng X, Etalo DW, van de Mortel JE, Dekkers E, Nguyen L, Medema MH, Raaijmakers JM. 2017.
  671 Genome-wide analysis of bacterial determinants of plant growth promotion and induced systemic resistance
  672 by Pseudomonas fluorescens. Environ Microbiol 19:4638–4656.
- 673 53. Chanch T, Ohira K. 1982. Comparison of the uptake and assimilation of ammonium and nitrate in indica
  674 and japonica rice plants using the tracer 15N method. Soil Sci Plant Nutr 28:79–90.
- 675 54. Cianfanelli FR, Monlezun L, Coulthurst SJ. 2016. Aim, load, fire: the type VI secretion system, a bacterial
  676 nanoweapon. Trends Microbiol 24:51-62.
- 55. Vergunst AC, Meijer AH, Renshaw SA, O'Callaghan D. 2010. Burkholderia cenocepacia creates an
  intramacrophage replication niche in zebrafish embryos, followed by bacterial dissemination and
  establishment of systemic infection. Infect Immun 78:1495–508.
- 56. Ferrières L, Hémery G, Nham T, Guérout AM, Mazel D, Beloin C, Ghigo JM. 2010. Silent mischief:
  Bacteriophage Mu insertions contaminate products of *Escherichia coli* random mutagenesis performed using
  suicidal transposon delivery plasmids mobilized by broad-host-range RP4 conjugative machinery. J
  Bacteriol 192:6418–6427.
- Wiles TJ, Norton JP, Russell CW, Dalley BK, Fischer KF, Mulvey MA. 2013. Combining quantitative
  genetic footprinting and trait enrichment analysis to identify fitness determinants of a bacterial pathogen.
  PLoS Genet 9:e1003716.
- 58. Pritchard JR, Chao MC, Abel S, Davis BM, Baranowski C, Zhang YJ, Rubin EJ, Waldor MK. 2014.
  ARTIST: High-resolution genome-wide assessment of fitness using transposon-insertion sequencing. PLoS
  Genet 10:e1004782.
- 690 59. DeJesus MA, Ambadipudi C, Baker R, Sassetti C, Ioerger TR. 2015. TRANSIT A software tool for
  691 Himarl TnSeq analysis. PLoS Comput Biol 11.

- 692 60. Zhao L, Anderson MT, Wu W, Harry HL, Bachman MA. 2017. TnseqDiff: Identification of conditionally
  693 essential genes in transposon sequencing studies. BMC Bioinformatics 18:326.
- 694 61. Médigue C, Calteau A, Cruveiller S, Gachet M, Gautreau G, Josso A, Lajus A, Langlois J, Pereira H, Planel
  695 R, Roche D, Rollin J, Rouy Z, Vallenet D. 2019. MicroScope-an integrated resource for community
  696 expertise of gene functions and comparative analysis of microbial genomic and metabolic data. Brief
  697 Bioinform 20:1071-1084.
- 698

# 699 Tables

- Table 1. Distribution of genes significantly impacting bacterial fitness on IR64 and Nipponbare rice.Colonization events on IR64 and Nipponbare rice cultivars were analyzed separately.
- Table 2. Selection of functions and number of corresponding genes impacting rice colonizationefficiencies when mutated. The detailed list of genes is provided in Table S3.
- Table 3. Selection of functions and number of corresponding colonization-depleted genes on IR64 andNipponbare rice.
- Table 4. Bacterial strains and plasmids used in this study.
- 707

#### 708 Figures

Figure 1. Distribution of genes significantly impacting bacterial fitness in a rich liquid medium. (A) Number of genes predicted to be essential, domain-essential (only insertions in part of the gene lead to abundance decrease) or non-essential by the EL-ARTIST pipeline. (B) Principal component analysis of essential and non-essential genes of Bv and Pk in a rich liquid medium. The genes were classified according to their COG categories and the PCA was built in regard to the abundance of these categories.

Figure 2. Rice colonization efficiencies for Bv and Pk. Nipponbare and IR64 rice seedlings were inoculated at 5 days post germination with  $1.10^7$  bacterial cells, total root systems were harvested at different time points and the associated bacterial population was estimated (p<0.05, Wilcoxon test). The boxplots were generated using 5 replicates for each condition. Outliers are represented as black dots.

# Figure 3. Distribution of genes significantly impacting bacterial fitness during rice colonization. Colonization events on IR64 and Nipponbare rice cultivars were pooled for this analysis. (A) Transposon insertions in genes leading to a 1.5-fold or higher decrease in abundance on roots at padj < 0.05 were</p> identified to be colonization- depleted. Those leading to a 1.5-fold or higher abundance were identified to

be colonization-enriched. (B) Distribution of genes significantly depleted or enriched during ricecolonization along COG categories in ratio of total impacted genes.

Figure 4. Distribution of colonization-depleted genes along the core- and specific-genomes of Bv and *Pk*. There are a total of 2072 genes that present significant homologies in the genomes of Bv and Pk and thus considered part of the core-genome (Materials and methods). A fraction of this core-genome is colonization-depleted in both species. Additionally, a substantial proportion of the genes that are found to be colonization depleted in a species or the other also belong to the core-genome.

Figure 5. Distribution of colonization-depleted genes on IR64 and Nipponbare rice cultivars. Distribution along COG categories of: (A) core genes of Bv and Pk that are colonization-depleted on both rice cultivars, (B) genes that are specifically colonization-depleted in Bv and (C) genes that are specifically colonization-depleted in Pk.

Figure 6. Colonization capacity of Bv and Pk insertion mutants in competition assays. Three genes, *cobW*, *edd* and *pdeR*, predicted by the Tn-seq approach to be involved in rice root colonization, were selected for targeted disruption in Bv and Pk. Mutants and WT strains were inoculated simultaneously on Nipponbare rice roots and enumerated at 7 dpi. Significance levels of pairwise comparisons were estimated using a Wilcoxon test (p<0.05). For each mutant, the log2 FC value observed in the Tn-seq approach on Nipponbare is displayed. Positive and negative correlations with the mutagenesis approach are expressed with green and red squares respectively.

Figure 7. Summary diagram of colonization-impacted bacterial functions involved in efficient rice root colonization. Pathways, functions, and genes described throughout this work are synthetized on this integrative representation. Elements specific of Bv are placed in the top part of the schematic bacteria (pink) and Pk specific elements at the bottom (green). Elements that are commonly found in both bacteria are placed at the interface (white). Function and molecular systems which were detected to be colonization-depleted are written in red font whereas colonization-enriched genes are written in blue font.

746

Strain	Colonization impact	Common	IR64	Nipponbare	Total
<b></b>	Enriched	507	83	130	720
B. vietnamiensis LMG10929	Depleted	635	225	134	994
	Total	1142	308	264	1714
P. kururiensis M130	Enriched	48	57	36	141
	Depleted	374	98	50	522
	Total	422	155	86	663

Table 1. Distribution of genes significantly impacting bacterial fitness on IR64 and Nipponbare rice.

	B. vietnamiensis		P. kururiensis	
Function	Depleted	Enriched	Depleted	Enriched
Motility	35	-	35	-
Chemotaxis	4	5	9	1
Cobalamin synthesis (B12)	1	8	15	-
Thiamin synthesis (B1)	4	-	5	-
Biotin synthesis (B7)	5	-	-	-
Potassium uptake	5	-	5	-
Ribose metabolism	-	5	-	-
Entner Doudoroff pathway	1	-	2	-
Aromatic compound metabolism	5	-	-	-
c-di-GMP cycling	1	7	-	1
Putrescine metabolism	3	-	1	-
Putrescine uptake	-	5	-	-
Cellulose synthase	-	5	-	-
Cell wall integrity	5	-	10	-
Peptidoglycan synthesis	3	-	-	-
Lytic transglycolases	4	-	1	-
Tol-Pal system	5	-	3	-
Hopanoid synthesis	7	-	1	-
O-antigen synthesis	2	-	-	-
Osmoprotection	2	-	1	-
DNA repair	9	-	9	-
Flp/Tad pilus	-	12	-	-
Arginine biosynthesis	5	-	4	-
Histidine biosynthesis	9	-	9	-
Isoleucine biosynthesis	6	-	4	-
Leucine biosynthesis	3	-	4	-
Lysine biosynthesis	-	-	1	-
Methionine biosynthesis	3	-	5	-
Proline biosynthesis	2	-	2	-
Serine biosynthesis	2	-	2	-
Tryptophan biosynthesis	6	-	6	-
Purine metabolism	9	1	1	-
Pyrimidine metabolism	5	-	6	-
Queuosine synthesis	6	-	1	-
Quorum sensing	-	-	-	2
T2SS	9	-	8	-
Autotransporter assistance	_	-	2	-
Prophage	-	-	-	22

Table 2. Selection of functions and number of corresponding genes impacting rice colonization efficiencies when mutated.

Table 3. Selection of functions and number of corresponding colonization-depleted genes on IR64 and
Nipponbare rice.

	B. viet	B. vietnamiensis		ruriensis
Function	IR64	Nipponbare	IR64	Nipponbare
T6SS	10	-	-	-
Glycerol metabolism	3	-	-	4
Phosphate import	3	-	-	-
Ammonium transport (AmtB)	-	1	-	1
Oxoprolinase	-	3	-	-
O-antigen biosynthesis	1	-	3	-

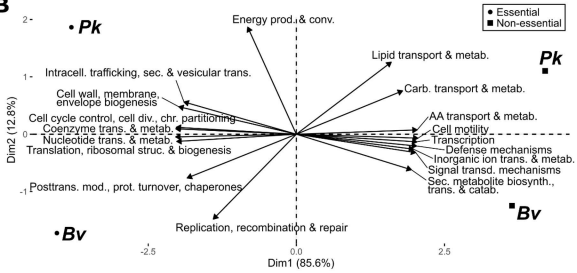
Strains/plasmids	Relevant characteristics and plasmid constructions	Reference/Source
E. coli		
MFDpir	Donor strain containing the pSAM-Ec vector and auxotroph for diaminopimelate, <i>\Delta dapA::(erm-pir)</i>	(Ferrières et al., 2010)
B. vietnamiensis		
LMG10929	Wild-type strain	(Gillis et al., 1995)
Rif <sup>R</sup>	Spontaneous rifampicin resistance clone of LMG10929	This study
Spt <sup>R</sup>	Spontaneous spectinomycin resistance clone of LMG10929	This study
DsRed	LMG1029 + pIN29	(King et al., 2019)
$\Omega cobW$	Insertion mutant of pSHAFT2 in AK36_2246	This study
$\Omega pdeR$	Insertion mutant of pSHAFT2 in AK36 4666	This study
$\Omega$ edd	Insertion mutant of pSHAFT2 in AK36 178	This study
P. kururiensis		
M130	Wild-type strain	(Baldani et al., 1997)
Rif <sup>R</sup>	Spontaneous rifampicin resistance clone of M130	This study
Spt <sup>R</sup>	Spontaneous spectinomycin resistance clone of M130	This study
DsRed	M130 + pIN29	(King et al., 2019)
$\Omega cobW$	Insertion mutant of pSHAFT2 in ANSKv1 10390	This study
$\Omega pdeR$	Insertion mutant of pSHAFT2 in ANSKv1 51226	This study
$\Omega$ edd	Insertion mutant of pSHAFT2 in ANSKv1 70910	This study
Plasmids		-
pIN29	Carries genes conferring DsRed fluorescence and chloramphenicol resistance. DsRed is under control of the highly active pTAC promoter	(Vergunst et al., 2010)
pSAM-Ec	Contains Himar1C9 transposase and a mariner transposon with a kanamycin resistance cassette	(Wiles et al., 2013)
pSHAFT2	Vector used for insertional mutagenesis containing a chloramphenicol resistance gene (non-replicative plasmid)	(Shastri et al., 2017)

Table 4. Bacterial strains and plasmids used in this study.

# Α

ARTIST prediction	B. vietnamiensis LMG10929	P. kururiensis M130
Essential	661	638
Domain-essential	96	97
Non-essential	5845	5638
Unclear	112	77

В



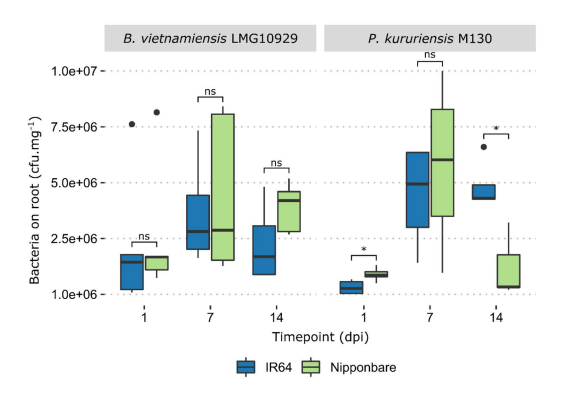
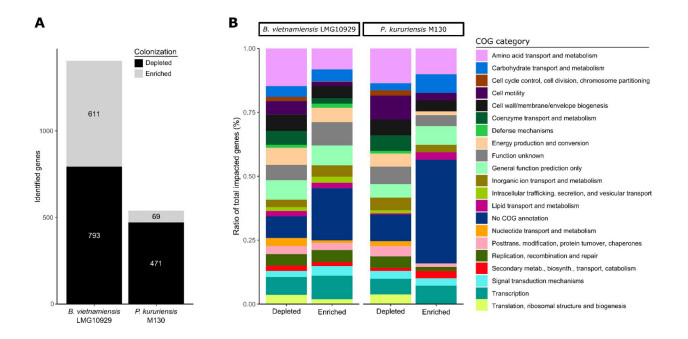
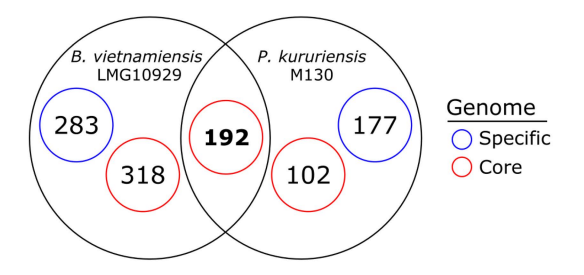


Fig.3





# Fig.5

