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1	Novel 3,6-Dihydroxypicolinic Acid Decarboxylase Mediated Picolinic Acid
2	Catabolism in Alcaligenes faecalis JQ135
3	
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#### 22 ABSTRACT

Alcaligenes faecalis strain JO135 utilizes picolinic acid (PA) as sole carbon and 23 nitrogen source for growth. In this study, we screened a 6-hydroxypicolinic acid 24 25 (6HPA) degradation-deficient mutant through random transposon mutagenesis. The mutant hydroxylated 6HPA into an intermediate, identified as 3,6-dihydroxypicolinic 26 acid (3,6DHPA) with no further degradation. A novel decarboxylase PicC was 27 identified that was found to be responsible for the decarboxylation of 3,6DHPA to 28 2,5-dihydroxypyridine. Although, PicC belonged to amidohydrolase\_2 family, it 29 30 shows low similarity (<45%) when compared to other reported amidohydrolase\_2 family decarboxylases. Moreover, PicC was found to form a monophyletic group in 31 the phylogenetic tree constructed using PicC and related proteins. Further, the genetic 32 33 deletion and complementation results demonstrated that picC was essential for PA degradation. The PicC was  $Zn^{2+}$ -dependent non-oxidative decarboxylase that can 34 specifically 35 catalyze the irreversible decarboxylation of 3.6DHPA to 36 2,5-dihydroxypyridine. The  $K_{\rm m}$  and  $k_{\rm cat}$  towards 3,6DHPA were observed to be 13.44  $\mu$ M and 4.77 s<sup>-1</sup>, respectively. Site-directed mutagenesis showed that His163 and 37 His216 were essential for PicC activity. 38

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#### 41 **IMPORTANCE**

Picolinic acid is a natural toxic pyridine derived from L-tryptophan metabolism 42 and some aromatic compounds in mammalian and microbial cells. Microorganisms 43 44 can degrade and utilize picolinic acid for their growth, and thus, a microbial degradation pathway of picolinic acid has been proposed. Picolinic acid is converted 45 into 6-hydroxypicolinic acid, 3,6-dihydroxypicolinic acid, and 2,5-dihydroxypyridine 46 in turn. However, there was no physiological and genetic validation for this pathway. 47 This study demonstrated that 3,6DHPA was an intermediate in PA catabolism process 48 49 and further identified and characterized a novel amidohydrolase\_2 family decarboxylase PicC. It was also shown that PicC could catalyze the decarboxylation 50 process of 3,6-dihydroxypicolinic acid into 2,5-dihydroxypyridine. This study 51 52 provides a basis for understanding PA degradation pathway and the underlying 53 molecular mechanism.

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#### 58 **INTRODUCTION**

Decarboxylation is a fundamental process in nature (1, 2). A variety of organic 59 compounds, including carbohydrates, fatty acids, aromatic compounds, and 60 61 environmental xenobiotics, are involved in decarboxylation. The decarboxylase family can be subdivided into two groups based on the cofactor involved (1). Some 62 enzymes require organic cofactors, such as flavin or NAD(P)<sup>+</sup>; while others utilize 63 inorganic cofactors, such as  $Zn^{2+}$  or  $Mn^{2+}$ . Recently, the amidohydrolase\_2 family 64 decarboxylases that use inorganic ions as cofactors, have been gaining more 65 attentions (3-6). The enzymes in this family are usually involved in the catabolism of 66 important natural compounds such as  $\alpha$ -amino- $\beta$ -carboxymuconate- $\varepsilon$ -semialdehyde 67 (ACMSD) (7, 8).  $\gamma$ -resorcylate (9), 2,3-dihydroxybenzoate 68 (10).2,5-dihydroxybenzoate (11), 3,4-dihydroxybenzoate (2), 4-hydroxybenzoate (12), 69 5-carboxyvanillate (13), vanillate (14), and 2-hydroxy-1-naphthoate (3). However, 70 71 most of these studied compounds are benzene ring derivatives, while no 72 decarboxylase has been studied that is involved in the catabolism of pyridine 73 derivatives.

Picolinic acid (PA) is a typical C2-carboxylated pyridine derivate that is widely 74 produce from physiological metabolism in mammalian and microbial cells (15). PA is 75 76 a natural dead-end metabolite of L-tryptophan produced via kynurenine pathway in humans and other mammals (16-18). Moreover, it can be produced in other biological 77 processes such as the microbial degradation of 2-aminophenol, catechol, and 78 nitrobenzene (19-21). PA was found to be toxic and it inhibited the growth of normal 79 80 rat kidney cells and T cell proliferation, thus, enhancing seizure activity in mice, and inducing cell death via apoptosis (22-25). PA cannot be metabolized by humans, thus 81 gets excreted through urine or sweat (26). However, PA can be degraded by 82

83 microorganisms in the natural environment (15). Numerous PA-degrading bacterial strains have been isolated including Achromobacter (27), Aerococcus (28), 84 Alcaligenes (29), Arthrobacter (30), Bacillus (31), Burkholderia (32), or Streptomyces 85 (33). The metabolic pathway of PA in microorganisms has been partially elucidated in 86 previous studies (15, 28, 32) (Fig. 1). In other studies, the crude enzyme facilitating 87 the conversion of PA to 6HPA has been preliminarily purified in Arthrobacter 88 89 picolinophilus DSM 20665 and an unidentified gram-negative bacterium (designated as UGN strain) (30, 34). Nevertheless, the functional genes or enzymes involved in 90 91 PA degradation has not been cloned or characterized yet.

92 In our previous work, we demonstrated that Alcaligenes faecalis strain JQ135 utilizes PA as sole carbon, nitrogen, and an energy source and 6-hydroxypicolinic 93 94 acid (6HPA) was the first intermediate of PA.(35). Further studies showed that maiA 95 gene was essential for PA catabolism (36). In the present research, we reported the fully characterized intermediate compound, 3,6-dihydroxypicolinic acid (3,6DHPA) 96 97 (Fig. 1). Further, a novel non-oxidative 3, 6-dihydroxypicolinic acid decarboxylase gene (*picC*) was cloned from A. *faecalis* strain JQ135, and the respective product was 98 characterized. 99

100

#### 102 **RESULTS**

#### 103 Transposon mutant and identification of the intermediate 3,6DHPA

A library of A. faecalis JQ135 mutants incapable of 6HPA utilization was 104 constructed by random transposon mutagenesis. More than 30 mutants that could not 105 grow on 6HPA were selected from approximately 10 000 clones and their ability to 106 convert 6HPA was examined. The concentration of 6HPA was 1 mM and the 107 inoculum of mutant was set at a final  $OD_{600}$  of 2.0. HPLC results showed that one 108 mutant (designated as Mut-H4) could convert 6HPA into a new intermediate with no 109 110 further degradation (Fig. 2). After liquid chromatography/time of flight-mass spectrometry (LC/TOF-MS) analysis, it was found that the molecular ion peak 111 112  $([M+H]^+)$  of this new intermediate was 156.0295 (Ion Formula, C<sub>6</sub>H<sub>6</sub>NO<sub>4</sub><sup>+</sup>, calculated 113 molecular weight 156.0297 with -3.2 ppm error), indicating that one oxygen atom was added to 6HPA (C<sub>6</sub>H<sub>5</sub>NO<sub>3</sub>). According to previously predicted PA degradation 114 pathway, the intermediate is most likely to be 3,6DHPA (15, 31, 34). In the present 115 study, 3,6DHPA was chemically synthesized and characterized by UV-visible, 116 LC/TOF-MS, <sup>1</sup>H NMR and <sup>13</sup>C NMR spectroscopies (Fig. S1 and S2) and HPLC 117 analysis showed that the retention time of the new intermediate was identical to that 118 of the synthetic sample of 3,6DHPA (Fig. 2). Thus, this intermediate compound was 119 120 identified as 3.6DHPA.

121

#### 122 Screening of the 3,6DHPA decarboxylase gene

123 The transposon insertion site of mutant Mut-H4 was identified using the genomic 124 walking method (37). The insertion site of the transposon was located in gene 125 *AFA\_15145* (genome position 3298929). Gene *AFA\_15145* was a 972 bp length ORF 126 starting with GTG. AFA\_15145 exhibited the highest sequence similarities to several

127	non-oxidative decarboxylases such as $\gamma$ -resorcylate decarboxylase ( $\gamma$ -RSD, 45%
128	identity) (9), 2,3-dihydroxybenzoate decarboxylase (2,3DHBD, 36% identity) (10),
129	5-carboxyvanillate decarboxylase (5CVD, 27% identity) (13), and hydroxynaphthoate
130	decarboxylase (HndA, 22% identity) (3) (Fig. 3). All these decarboxylases belong to
131	the amidohydrolase_2 family proteins (COG2159) that contain a triosephosphate
132	isomerase (TIM)-barrel fold. Based on the phenotype of the mutant Mut-H4 and
133	bioinformatics analysis, it was predicted that the AFA_15145 (designated as picC)
134	encoded the 3,6DHPA decarboxylase.

# 136 **Function identification of** *picC* **gene in PA degradation in** *A. faecalis* **JQ135**

137 To confirm whether *picC* is involved in PA degradation, a *picC*-deleted mutant 138 JQ135 $\Delta$ *picC* was constructed. The mutant JQ135 $\Delta$ *picC* lost the ability to grow on PA, 139 6HPA, or 3,6DHPA. The complementation strain, JQ135 $\Delta$ *picC*/pBBR-*picC* 140 completely restored the phenotype of growth on PA, 6HPA, and 3,6DHPA. These 141 results showed that *picC* gene was essential for the degradation of PA in *A. faecalis* 142 JQ135.

143

## 144 The *picC* encodes 3,6DHPA decarboxylase

The recombinant PicC was overexpressed in *E. coli* BL21(DE3) cells containing the plasmid pET-*picC*. SDS/PAGE analysis showed the presence of an intense band, consistent with the  $6 \times$  His-tagged PicC (37 kDa) (Fig. 4). The degradation of 3,6DHPA by purified PicC was monitored spectrophotometrically. The maximum absorption was shifted from 340 nm (3,6DHPA) to 320 nm (2,5DHP). LC/TOF-MS analysis suggested that the molecular ion peak of the product was 112.0400 (M+H<sup>+</sup>), which was identical to that of 2,5DHP (36, 38). Further, the HPLC analysis showed that the retention time of the product was identical to that of the authentic sample of
2,5DHP. The 3,6DHPA was degraded completely with the formation of equal molar of
2,5DHP. Moreover, the PicC did not catalyze the reverse carboxylation of 2,5DHP in
a reaction mixture containing NaHCO<sub>3</sub>.

156

### 157 Biochemical properties of PicC

The recombinant PicC was highly active at pH 7.0 and 40°C (Fig. S3). The  $K_{\rm m}$ and  $k_{\rm cat}$  values for 3,6DHPA were found to be 13.44 µM and 4.77 s<sup>-1</sup>, respectively (Table 1). The enzyme was unstable at room temperature and could retain only 50% of initial activity when incubated at 30°C for 24 h. In addition, PicC could not convert the structural analogues of 3,6DHPA including 3-hydroxy-picolinic acid, gentisic acid, 2,3-dihydroxybenzoic acid, and 2,6-dihydroxybenzoic acid. This can be attributed to the substrate specificity of PicC towards 3,6DHPA.

The effects of various metal ions and inhibitors, on decarboxylase activity are 165 presented in Table 2. PicC activity was not affected by metal ions such as Ca<sup>2+</sup>, Cd<sup>2+</sup>. 166 Co<sup>2+</sup>, Fe<sup>3+</sup>, Mg<sup>2+</sup>, Mn<sup>2+</sup>, and Zn<sup>2+</sup> but was strongly inhibited by Ag<sup>+</sup>, Co<sup>2+</sup>, and Hg<sup>2+</sup> 167 ions. In addition, several inhibitors such as EDTA, 8-hydroxy-quinoline-5-sulfonic 168 acid (8-HQSA, a zinc metal specific inhibitor), phenylmethylsulfonyl fluoride (PMSF, 169 serine and cysteine specific inhibitor), and sodium iodoacetate (cysteine specific 170 171 inhibitor) showed relatively low effects on PicC activity. However, diethylpyrocarbonate (DEPC), a histidine residue modifier, strongly inhibited PicC 172 activity, indicating the presence of active-site histidine residues. 173

Further, absorption spectroscopy analysis revealed the presence of  $Zn^{2+}$  at 0.85±0.1 mol per mol of protein that is similar to several other non-oxidative decarboxylases of the amidohydrolase\_2 superfamily (1, 3). The moderate effects,

exhibited by additional  $Zn^{2+}$  or EDTA in the reaction system, indicated the presence of  $Zn^{2+}$  in the center of PicC.

179

# 180 Site-directed mutagenesis

In order to assess their roles in the function of PicC, seven histidine residues 181 (H12, H135, H163, H172, H177, H194, and H216) were substituted with Ala residue 182 through site-directed mutagenesis (Fig. 3B). The seven PicC mutants obtained were 183 expressed and purified and their activities were determined (Fig. S4; Table 1). 184 PicC<sup>H135A</sup> showed a slight increase in decarboxylase activity, while PicC<sup>H172A</sup> and 185  $PicC^{H194A}$  showed a slight reduction in decarboxylase activities (10%~50%) and 186 PicC<sup>H12A</sup> and PicC<sup>H177A</sup> strongly reduced the decarboxylase activities (>90%). 187 Moreover, the mutant proteins PicC<sup>H163A</sup> and PicC<sup>H216A</sup> completely lost their 188 decarboxylase activities. 189

#### 191 **DISCUSSION**

PA is a natural and toxic mono-carboxylated pyridine derivative. The studies on 192 the microbial degradation mechanism of PA began 50 years ago (28). A partial 193 194 catabolic pathway of PA has been proposed (Fig. 1): PA was dehydrogenated to 6HPA, and then gets converted into 3,6DHPA via hydroxylation leading to the 195 decarboxylation of 3,6DHPA into 2,5DHP. The intermediate, 6HPA has been 196 197 substantially identified in most strains including Aerococcus sp. (28), Alcaligenes faecalis DSM 6269 (29), Arthrobacter picolinophilus DSM 20665 (30, 39), 198 199 Burkholderia sp. ZD1 (32), Streptomyces sp. Z2 (33), and the UGN strain (34). Further, another intermediate compound, 2,5DHP has been detected in the media 200 during PA degradation in few strains (32, 34). However, the intermediate 3,6DHPA, a 201 202 key link between 6HPA and 2,5DHP, was hardly detectable. This could be most attributed to its immediate degradation before excreting out of the cells. Previously, 203 3,6DHPA has only been theoretically proposed in Bacillus sp. (31) and the UGN 204 205 strain (34). In this study, to the best of our knowledge, we demonstrated the chemical properties (UV-visible, LC/TOF-MS, <sup>1</sup>H NMR and <sup>13</sup>C NMR spectroscopies) of 206 3,6DHPA for the first time (Fig. S2) and detected it in the media using the transposon 207 mutant strain, thus confirming that 3,6DHPA is a catabolic intermediate of PA. 208

Some previous studies have attempted to identify the genes and enzymes involved in PA degradation, such as the PA dehydrogenase in *Arthrobacter* (39) and 2,5DHP dioxygenase in UGN strain (34). However, their amino acid sequences and respective coding genes remained unknown, with no biochemical, physiological or genetic evidence to explain the decarboxylation of 3,6DHPA to 2,5DHP. In this context, we cloned a decarboxylase gene *picC* through random transposon mutagenesis, and ascertained that PicC was responsible for 3,6DHPA decarboxylation

216 to form 2,5DHP. We found that PicC shared homology with the amidohydrolase 2 family proteins and contained the conserved triosephosphate isomerase (TIM)-barrel 217 fold of amidohydrolase\_2 family (Fig. 3). It has been previously reported that the 218 219 amidohydrolase\_2 family protein (PF04909) catalyzes the decarboxylation reaction (C-C bond) of several benzene derivatives, whereas the amidohydrolase\_1 family 220 protein (PF01979) catalyzes the hydrolytic reactions (C-N, C-Cl, or C-P bond) (4). 221 The ACSMD was the first member of amidohydrolase\_2 family to be reported (8) 222 followed by other members, including y-RSD (9), 2,3DHBD (10), 5CVD (13), and 223 224 HndA (3). A phylogenetic tree of PicC and related proteins showed that PicC was clustered with amidohydrolase 2 but not amidohydrolase 1 family proteins (Fig. 3). 225 However, the identities between PicC and reported decarboxylases were low (less than 226 227 45%), and PicC formed a separate branch in the phylogenetic tree (Fig. 3). In addition, PicC was found to be specific toward its substrate 3,6DHPA. Thus, it can be 228 concluded that PicC could be a novel amidohydrolase\_2 family decarboxylase. 229

230 The amidohydrolase 2 family proteins contain a few conserved amino acid residues, which are usually the active sites. In ACSMD, the His177, His228, and 231 D294 were the  $Zn^{2+}$ -binding sites (4). These three residues have been found in all 232 reported amidohydrolase\_2 family proteins including PicC (His163, His216, and 233 D283) (Fig. 3B). In addition, the results of site-directed mutagenesis of PicC 234 235 confirmed that H163 and H216 also played essential roles in PicC-mediated catalysis. Another  $Zn^{2+}$ -binding motif 'HxH' has been found in the N-terminal of ACSMD (4) 236 and HndA (3), whereas this motif was replaced by 'EEH' in  $\gamma$ -RSD (9) or 'EEA' in 237 5CVD (13) (Fig. 3B). In PicC, the corresponding motif has been found to be similar 238 to that of  $\gamma$ -RSD. After substituting His12 by Ala, the resultant enzyme PicC<sup>H12A</sup> still 239 exhibited 10% activity, suggesting a variation in the third residue of this motif. In 240

addition, site-directed mutagenesis results demonstrated that several other histidine
residues His172, His177, and His194 were important for PicC activity.

In conclusion, this study revealed that 3,6DHPA was a catabolic intermediate in PA degradation by bacteria. The 3,6DHPA decarboxylase (PicC) was identified and characterized. To the best of our knowledge, PicC is also the first non-oxidative decarboxylase belonging to the amidohydrolase\_2 family that catalyzes the irreversible decarboxylation of pyridine derivative. This study will expand our understanding of the bacterial degradation mechanisms of pyridine derivatives.

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#### 251 MATERIALS AND METHODS

#### 252 Chemicals

PA, 6HPA, and 2,5DHP were purchased from J&K Scientific Ltd. (Shanghai,
China). EDTA, 8-HQSA, PMSF, sodium iodoacetate, DEPC, and other reagents of
analytical grade were purchased from Sangon Biotech Co., Ltd. (Shanghai, China).
3,6DHPA was chemically synthesized (detailed in the supplemental materials). The
structure of 3,6DHPA was confirmed by UV-visible, LC/TOF-MS, and NMR
spectroscopy (Fig. S1 and S2).

259

#### 260 Strains, plasmids, and primers

All bacterial strains and plasmids used in this study are listed in Table 3. 261 Alcaligenes faecalis JQ135 (CCTCC M 2015812) is the wild-type PA-degrading 262 strain (35). E. coli DH5a was used as the host for the construction of plasmids. E. coli 263 BL21(DE3) was used to over-express the proteins. Bacteria were cultivated in LB 264 265 medium at 37°C (E. coli) or 30°C (Alcaligenes and their derivatives). Antibiotics were added at the following concentrations (as required): chloramphenicol (Cm), 34 mg/L; 266 gentamicin (Gm), 50 mg/L; kanamycin (Km), 50 mg/L; and streptomycin (Str), 50 267 mg/L. Primer synthesis and the sequencing of PCR products or plasmids were 268 performed by Genscript Biotech (Nanjing, China) (40). The primers used in this study 269 270 are listed in Table 4.

271

### 272 Transposon mutagenesis, mutant screening, and gene cloning

A transposon mutant library of *A. faecalis* JQ135 was constructed using the transposon-based plasmid pSC123 (kanamycin-resistance gene) as described previously (35). In this study, the PA was replaced by its intermediate 6HPA. Mutants that could not utilize 6HPA as the sole carbon source were selected. The flanking sequences of the transposon in the mutants were amplified using the DNA walking method (37). The amplified PCR products were sequenced and analyzed. The insertion sites were confirmed by comparison with the genome sequence of *A. faecalis* JQ135.

281

#### 282 Gene knockout and genetic complementation of A. faecalis JQ135

The genes or DNA fragments from A. faecalis JQ135 were were amplified by 283 284 PCR using corresponding primers (Table 4). The fusion of DNA fragments and cut plasmids was carried out using the ClonExpress MultiS One Step Cloning Kit 285 (Vazyme Biotech Co.,Ltd, Nanjing, China). Gene deletion mutant of the picC in A. 286 faecalis JQ135 was constructed using a two-step homogenetic recombination method 287 with the suicide plasmid pJQ200SK (41). Two homologous recombination-directing 288 289 sequences were amplified using primers, kopicC-UF/kopicC-UR and 290 kopicC-DF/kopicC-DR, respectively. Two PCR fragments were subsequently ligated into SacI/PstI-digested pJQ200SK generating pJQ- $\Delta picC$ . The pJQ- $\Delta picC$  plasmid 291 was then introduced into A. faecalis JQ135 cells. The single-crossover mutants were 292 screened on a LB plate containing Str and Gm. The gentamicin-resistant strains were 293 then subjected to repeated cultivation in LB medium containing 10% sucrose with no 294 295 gentamicin. The double-crossover mutants that lost their plasmid backbone and were sensitive to gentamicin, were selected on LB Str plates. Deletion of the *picC* gene was 296 confirmed by PCR. This procedure resulted in the construction of the deletion mutant 297 298 strain JQ135 $\Delta picC$ .

299 Knockout mutants were complemented as follows. The intact picC gene was 300 amplified using the primers picC-F and picC-R, and then ligated with the

301 XhoI/HindIII-digested pBBR1-MCS5, generating pBBR-*picC*. The pBBR-*picC* was 302 then transferred into the mutant strain JQ135 $\Delta picC$  to generate the complemented 303 strain JQ135 $\Delta picC$ /pBBR-*picC*.

304

#### 305 Expression and Purification of the His-tagged PicC and its mutations

For the over-expression of *picC* gene in *E. coli* BL21(DE3), the complete ORF 306 without the stop codon (genome position 3298274-3299242) were amplified using 307 genomic DNA of strain JO135 and inserted into the NdeI/XhoI-digested plasmid 308 309 pET29a(+), resulting in the plasmid pET-PicC. E. coli BL21(DE3) cells (containing pET-PicC) were initiated by the addition of 0.3 mM IPTG when the optical density of 310 the culture (OD<sub>600</sub>) reached 0.5-0.8 and was incubated for an additional 12 h at 16°C.. 311 Cells were harvested by centrifugation at 4 °C, sonicated, and then centrifuged again 312 to remove cell debris. The supernatant was used for recombinant protein purification 313 using Ni-NTA agarose column (Sangon, Shanghai, China). The purified  $6 \times$ 314 315 His-tagged protein were then analyzed using 12.5% SDS-PAGE. The protein 316 concentrations were determined using the Bradford method (42).

For site-directed mutagenesis of PicC, the *picC* fragments were amplified from plasmid pET-PicC through overlap PCR using the primers carrying point mutations (Table 4). Amplified fragments were fused into plasmid pET29a(+), resulting in pET-PicC<sup>H12A</sup>, pET-PicC<sup>H135A</sup>, pET-PicC<sup>H163A</sup>, pET-PicC<sup>H172A</sup>, pET-PicC<sup>H177A</sup>, pET-PicC<sup>H194A</sup>, and pET-PicC<sup>H216A</sup>. Resultant constructs were confirmed by sequencing. The expression and purification of the mutations were performed as described in the section above.

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#### 326 Enzymatic assays of 3,6DHPA decarboxylase

For the decarboxylase activity, enzyme reaction mixture contained 50 mM PBS 327 (pH 7.0), 0.3 mM 3,6DHPA, and 5 µg purified PicC (in 1 mL) and incubated at 40°C. 328 329 The enzymatic activities were determined spectrophotometrically by the disappearance of 3,6DHPA at 360 nm ( $\epsilon$ =4.4 cm<sup>-1</sup> mM<sup>-1</sup>). To determine the effect of one condition, 330 other conditions were kept at fixed concentration of the standard reaction. The 331 optimum pH of the PicC protein was determined using various buffers such as 50 mM 332 citric acid-sodium citrate (pH 4 to 6), 50 mM KH<sub>2</sub>PO<sub>4</sub>-K<sub>2</sub>HPO<sub>4</sub> (pH 6 to 8), and 50 mM 333 334 glycine-NaOH (pH 8.0 to 9.8) at 40°C. The optimum temperature of the PicC protein was determined to be in between 10°C to 50°C in PBS (pH 7.0). Purified PicC was 335 pre-incubated with various metal ions and inhibitors at 4°C for 30 min to study their 336 effects on the enzyme. The activity was expressed as a percentage of the activity 337 obtained in the absence of the added compound. To determine the kinetic constants for 338 3,6DHPA, a range of 3,6DHPA concentrations (2 to 150 µM) were used. The values 339 340 were calculated through non-linear regression fitting to the Michaelis-Menten equation. One unit of the activity was defined as the amount of enzyme that catalyzed 1 µmol of 341 3,6DHPA in 1 min. 342

The measurement of carboxylase activity of PicC was similar with a previous study (43). The reaction mixture contained 50 mM PBS (pH 7.0), 0.3 mM 2,5DHP, 5.0 mM NaHCO<sub>3</sub> and 5  $\mu$ g purified PicC in 1 mL mixture at 40 °C.

346

#### 347 Analytical methods

The UV-VIS spectra was observed by a UV2450 spectrophotometer (Shimadzu). The determination of PA and 6HPA, 3,6DHPA, and 2,5DHP concentrations were performed by HPLC analysis on a Shimadzu AD20 system equipped with a Phecda

351	C18 reversed phase column (250 mm $\times$ 4.60 mm, 5 $\mu m$ ). The concentrations of the
352	compounds were calculated using standard samples. The mobile phase was consisted
353	of methanol : water : formic acid (12.5:87.5:0.2, v/v/v) at a flow rate of 0.6 mL/min,
354	at 30 °C. LC/TOF-MS analysis was performed in a TripleTOF 5600 (AB SCIEX)
355	mass spectrometer, as described previously (44). The $Zn^{2+}$ concentration in the PicC
356	protein was analyzed using inductively coupled plasma optical emission spectrometry
357	(ICP-OES) according to a previous study (45).
<b></b>	

# 359 Nucleotide sequence accession numbers

The PicC and the complete genome sequence of *A. faecalis* JQ135 have been deposited in the GenBank database under accession numbers <u>ARS01287</u> and <u>CP021641</u>, respectively.

363

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372

#### 373 **Conflict of interest**

- The authors declare no conflict of interest.
- 375

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#### 498 Figure legends

501

- 499 Fig. 1 Proposed PA degradation pathway in *A. faecalis* JQ135. The 3,6DHPA and
- 500 2,5-DHP are shown in blue color. TCA, tricarboxylic acid cycle.
- 502 Mut-H4. (A and C) the authentic sample of 6HPA and 3,6DHPA, respectively. (B) The

Fig. 2 HPLC and LC/TOF-MS profiles of the conversion of 6HPA by mutant

sol indi in (in and c) the addicate sample of offirm and s, obtinn, respectively. (b) the

503 conversion of 6HPA into 3,6DHPA by mutant Mut-H4. The detection wavelength was

set at 310 nm. (D) LC/TOF-MS spectra of 3,6DHPA produced in panel B.

Fig. 3 Amino acid sequence analysis of PicC. (A) Phylogenetic analysis of PicC 505 506 and related decarboxylases. Each item was arranged in the following order: protein name, accession number, and strain. AmiH 1 and 2 are amidohydrolase 1 and 2 507 family, respectively. The phylogenetic tree was constructed using the neighbor-joining 508 509 method (with a bootstrap of 1000) with software MEGA 6.0. The bar represents amino acid substitutions per site. (B) Multiple sequence alignment of PicC and seven 510 decarboxylases. The predicted N-terminal motifs for  $Zn^{2+}$  binding are denoted by blue 511 box. The other three  $Zn^{2+}$ -binding sites are denoted by purple diamonds. The seven 512 histidine residues for site-directed mutagenesis are denoted by blue triangles. 513

Fig. 4 Characterization of PicC. (A) SDS-PAGE of purified PicC. Lane M, protein marker. Lane 1, purified PicC. (B) Spectrophotometric changes during transformation of 3,6DHPA by purified PicC. The reaction was initiated by adding 3,6DHPA. Spectra were recorded every 1 min. The arrow denotes the biotransformation of 3,6DHPA into 2,5DHP. (C) HPLC analysis of the transformation of 3,6DHPA into 2,5DHP by PicC. The detection wavelength was 310 nm.

#### 521 Tables

522

#### 523

# 524 **Table 1 Kinetic constants of wild-type PicC and mutants.**

525

Enzyme	K <sub>m</sub>	$V_{\max}$	$k_{\rm cat}$	$k_{\rm cat}/K_{ m m}$
	(µM)	$(\mu mol min^{-1} mg^{-1})$	$(s^{-1})$	$(s^{-1} mM^{-1})$
PicC	$13.44 \pm 2.91$	$7.73 \pm 0.05$	4.77	354.54
PicC <sup>H12A</sup>	$53.06 \pm 4.05$	$2.83 \pm 0.09$	1.75	32.94
PicC <sup>H135A</sup>	9.38±0.68	$6.14 \pm 0.11$	3.78	403.40
PicC <sup>H163A</sup>	ND	ND	ND	ND
PicC <sup>H172A</sup>	$15.80 \pm 1.21$	$3.33 \pm 0.08$	2.06	130.14
PicC <sup>H177A</sup>	$110.81 \pm 14.70$	$2.18 \pm 0.01$	1.35	12.14
PicC <sup>H194A</sup>	$11.38 \pm 0.64$	$5.15 \pm 0.07$	3.18	279.05
PicC <sup>H216A</sup>	ND	ND	ND	ND

526 ND: not detected

527

# 530 **Table 2 Effect of metal ions and inhibitors on PicC activity.**

531

ions/inhibitor	Relative activity (%)*	ions/inhibitor	Relative activity (%)
BLANK	100	AgNO <sub>3</sub>	4.63±0.53
CaCl <sub>2</sub>	$90.68 \pm 1.49$	CdCl <sub>2</sub>	99.98±2.43
CoCl <sub>2</sub>	84.47±2.21	CuCl <sub>2</sub>	23.89±0.67
FeCl <sub>3</sub>	$97.59 \pm 1.44$	$HgCl_2$	$2.36 \pm 0.40$
MgSO <sub>4</sub>	87.59±1.95	MnSO <sub>4</sub>	91.30±1.13
ZnSO <sub>4</sub>	$97.11 \pm 1.84$	DEPC	$4.05 \pm 0.19$
EDTA	$60.94 \pm 1.64$	8-HQSA	$55.21 \pm 0.91$
PMSF	$97.27 \pm 0.32$	Sodium iodoacetate	81.68±0.99

\* Metal ions were tested at 0.1 mM. DEPC: 0.1%. PMSF: 1%. Sodium iodoacetate: 0.025 mM.

533 8-HQSA: 100 μg/mL. EDTA: 100 μg/mL.

534

# 537 Table 3 Strains and plasmids used in this study.

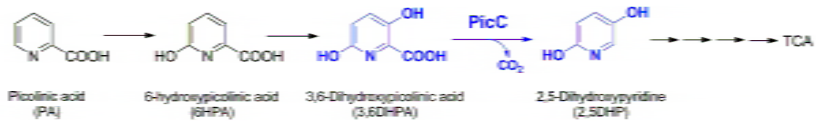
Strains or plasmids	Description	Source
Alcaligenes faecalis strains		
JQ135	Str <sup>r</sup> ; PA-degrading bacterium. Gram-negative. Wild type.	CCTCC M 2015812
Mut-H4	Str <sup>r</sup> , Km <sup>r</sup> ; picC (AFA_15145) mutant of A. faecalis JQ135 inserted by transposon	This study
$JQ135\Delta picC$	Str <sup>1</sup> ; <i>picC</i> -deletion mutant of JQ135	This study
JQ135∆picC/pBBR-picC	Str <sup><math>r</math></sup> , Gm <sup><math>r</math></sup> ; JQ135 $\Delta picC$ complemention with pBBR- <i>picC</i>	This study
E. coli strains		
DH5a	F <sup>-</sup> recA1 endA1 thi-1 hsrdR17 supE44 relA1 deoRΔ(lacZYA-argF) U169 φ80lacZ ΔM15	TaKaRa
BL21(DE3)	$F^{-}$ ompT hsdS( $rB^{-}mB^{-}$ ) gal dcm lacY1(DE3)	TaKaRa
$SM10_{\lambda pir}$	Donor strain for biparental mating	Lab stock
Plasmids		
pET29a(+)	Km <sup>r</sup> , expression plasmid	Novagen
pSC123	Cm <sup>r</sup> , Km <sup>r</sup> ; suicide plasmid, mariner transposon	Lab stock
pJQ200SK	Gm <sup>r</sup> , mob <sup>+</sup> , <i>orip</i> 15A, <i>lacZa<sup>+</sup></i> , <i>sacB</i> ; suicide plasmid	Lab stock
pBBR1MCS-5	Gm <sup>r</sup> ; broad-host-range cloning plasmid	Lab stock
pJQ-∆ <i>picC</i>	$Gm^r$ ; <i>picC</i> gene deletion plasmid	This study
pBBR-picC	Gm <sup>r</sup> ; the fragment containing the <i>picC</i> gene inserted into XhoI/HindIII-digested pBBR1MCS-5	This study
pET-PicC	Km <sup>r</sup> ; <i>Nde</i> I-XhoI fragment containing <i>picC</i> gene inserted into pET29a(+)	This study
pET-PicC <sup>H12A</sup>	Km <sup>r</sup> ; <i>Nde</i> I-XhoI fragment containing <i>picC<sup>H12A</sup></i> gene inserted into pET29a(+)	This study
pET-PicC <sup>H135A</sup>	Km <sup>r</sup> ; <i>Nde</i> I-XhoI fragment containing <i>picC</i> <sup>H135A</sup> gene inserted into pET29a(+)	This study
pET-PicC <sup>H163A</sup>	Km <sup>r</sup> ; <i>Nde</i> I-XhoI fragment containing <i>picC</i> <sup>H163A</sup> gene inserted into pET29a(+)	This study
pET-PicC <sup>H172A</sup>	Km <sup>r</sup> ; <i>Nde</i> I- <i>Xho</i> I fragment containing <i>picC</i> <sup>H172A</sup> gene inserted into pET29a(+)	This study
pET-PicC <sup>H177A</sup>	Km <sup>r</sup> ; <i>Nde</i> I- <i>Xho</i> I fragment containing <i>picC</i> <sup>H177A</sup> gene inserted into pET29a(+)	This study
pET-PicC <sup>H194A</sup>	Km <sup>r</sup> ; <i>Nde</i> I- <i>Xho</i> I fragment containing <i>picC</i> <sup>H194A</sup> gene inserted into pET29a(+)	This study
pET-PicC <sup>H216A</sup>	$\text{Km}^{r}$ ; <i>NdeI-XhoI</i> fragment containing <i>picC</i> <sup>4216A</sup> gene inserted into pET29a(+)	This study

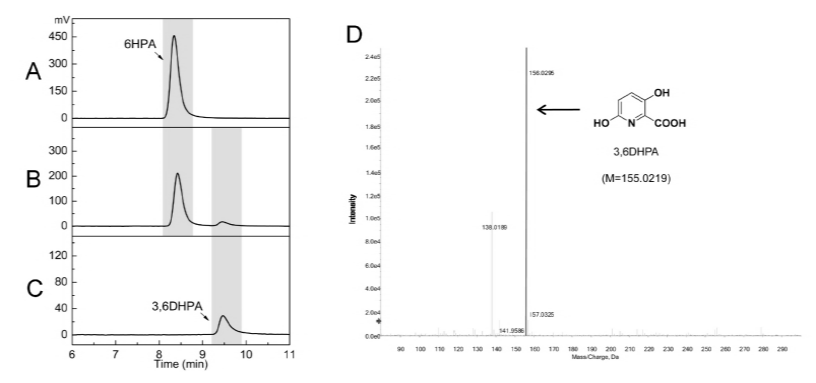
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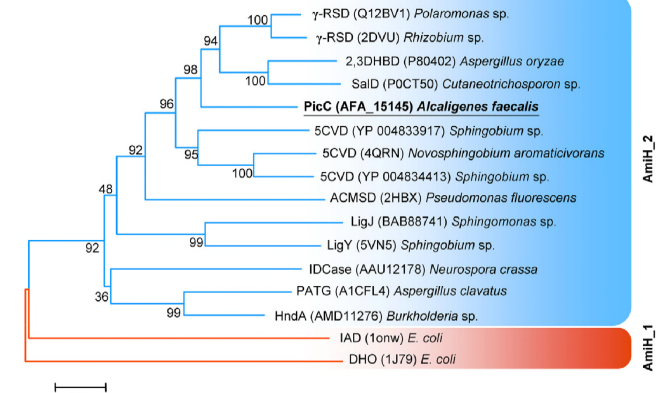
# **Table 4 Primers used in this study.**

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Primers	Sequence(5'-3')	Description
kopicC-UF	AGCTTGATATCGAATTCCTGCAGTTCTTCTTGCTTTAGCTCGGC	To construct plasmid pJQ-Δ <i>picC</i>
kopicC-UR	TCTAGAACTAGTGGATCCAGTCTGCGACAGCACGACGTAC	
kopicC-DF	GGATCCACTAGTTCTAGAGTGTTGTTCTCGGTGGACTACC	
ko <i>picC-</i> DR	AGGGAACAAAAGCTGGAGCTCAGAAGGCGCGCATATCCTCAG	
picC-F	CAGGAATTCGATATCAAGCTTAGTGGCTTTCAGCCTGTTGCC	To construct plasmid pBBR-picC
picC-R	GGTACCGGGCCCCCCTCGAGGTAATGCTGACGAATGCCATTGG	
expPicC-F	CTTTAAGAAGGAGATATACATATGAAACGTATTAAAAAAATAGC	To construct plasmid pET-PicC
expPicC-R	GTGGTGGTGGTGGTGGTGCTCGAGGCTCCGGTCCAGCTTGAACAG	
Mut-H12A-1	CTTTAAGAAGGAGATATACATATGAAACGTATTAAAAAAATAGCACTGGAGG	To construct plasmid pET-PicC <sup>H12A</sup>
Mut-H12A-2	AAAAAAATAGCACTGGAGGAGGCATTCAACGCCGTTGG	
Mut-H12A-3	TCAGTGGTGGTGGTGGTGGTGCTCGAGGCTCCGGTCCAGCTTGAACAG	
Mut-H135A-1	GCTTTGGTCAACGGTGCTACGCATGGTGTGTAC	To construct plasmid pET-PicC <sup>H135A</sup>
Mut-H135A-2	GTACACACCATGCGTAGCACCGTTGACCAAAGC	
Mut-H163A-1	GTGCCGTTCTATCTGGCTCCCTTTGATGCTTACG	To construct plasmid pET-PicC <sup>H163A</sup>
Mut-H163A-2	CGTAAGCATCAAAGGGAGCCAGATAGAACGGCAC	
Mut-H172A-1	GCTTACGAAATGCCAGCTGCTTACACAGGCCAC	To construct plasmid pET-PicC <sup>H172A</sup>
Mut-H172A-2	GTGGCCTGTGTAAGCAGCTGGCATTTCGTAAGC	
Mut-H177A-1	CACGCTTACACAGGCGCTCCGGAGCTGGTTGGG	To construct plasmid pET-PicC <sup>H177A</sup>
Mut-H177A-2	CCCAACCAGCTCCGGAGCGCCTGTGTAAGCGTG	
Mut-H194A-1	GTAGAAACCGGCACCGCAGCGCTGCGCATGTTG	To construct plasmid pET-PicC <sup>H194A</sup>
Mut-H194A-2	CAACATGCGCAGCGCTGCGGTGCCGGTTTCTAC	
Mut-H216A-1	AAGCTGGTGCTGGGTGCAATGGGTGAAGGCCTG	To construct plasmid pET-PicC <sup>H216A</sup>
Mut-H216A-2	CAGGCCTTCACCCATTGCACCCAGCACCAGCTT	







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PicC	1	EEH(/A) MKRIK <mark>KI</mark> AI <mark>EEHF</mark> NAVGFEDYSKAFVKHIDSADARELMAR.LHDFDAQ <mark>RL</mark> EV <mark>MD</mark>
gammaRSD 5CVD 2,3DHBD SalD PATG HndA ACMSD	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	MKRIKKIALEEH NAVGFEDYSKAFVKH
PicC	54	RAGIEYVVLSQTGFGGVQVEKDVSVAI.ERARQNNDFLAQQIAHHPDRLGGFATLP.MQDPAAAAQELTRAVQDLGLKG
gammaRSD	49	EHGIETMILSLNAPAVQAIADSTRAN.ETARRANDFLAEQVAKQPTRFRGFAALP.MQDPELAARELERCVKELGFVG
5CVD	79	ETGIDVAILSMTSPGGQVFEADEAKA.LVSEANDVLKAACERYPTRYYGMISIV.PQDPAWSVAEIRRGKEELGFRG
2,3DHBD	51	KHGVGYQILSYTAPGVQDIWDPVEAQ.ALAVEINDYIAEQVRVNPDRFGAFATLS.MHNPKEAADELRRCVEKYGFKG
SalD	53	KHGIGYTIYSIYSPGPQGWTERAECE.EYARECNDYISGEIANHKDRMGAFAALS.MHDPKQASEELTRCVKELGFIG
PATG	45	VLKVQTTILSVTAPGPGIETDPGKAA.ALARLCNEEAAAIRDAHPLQYGFFASVPSLFDTAAVLAEIEHAFTNLHADG
HndA	48	SLEIETGVLSLTAPG.VQGWNGSARRDMARRINEYVAGLVAQWPARFGNFATLP.LPDVDGTLAEIGHAFDALNADG
ACMSD	69	AQGVDVQVTCATPVMFGYTWEANKAA.QWAERMNDFALEFAAHNPQRIKVLAQVP.LQDLDLACKEASRAVA.AGHLG
PicC gammaRSD 5CVD 2,3DHBD SalD PATG HndA ACMSD	130 125 154 127 129 122 123 144	ALVNGHTHGVYYDGREYDAFWETVQKLDVPFYLHPFDAYEMPHAYTGHPELVGATWGWGVETGTHALRM ALVNGFSQDNRSA.VPLYYDMAQYWPFWETVQALDVPFYLHPRNPLP.SDARIYDGHAWLLGPTWAFGQETAVHALRL VMVNSHTKGQYLDEPQFDPILRACAEQDLPLYIHPQSPPDGMIAGMVEAG.LDGAIFGFGVETGYHLLRL ALVNDTQRAGPDGDDMIFYDNADWDIFWQTCTELDVPFYMHPRNPTGTIYEKLWADRKWLVGPPLSFAHGVSLHVLGM ALVNDVQHAGPEGETHIFYDQPEWDIFWQTCVULDVPFYLHPEPPFGSYLRNQYEGRKYLIGPPVSFANGVSLHVLGM VTLYTRYGAGHSYLGDERFRPIWAELSKRRAVVFIHPTHAVDTQLINSWMPQPMFDYPHETGRTAMDL VVLSNYGGKDLDDATLEAFLTHCANEDIPILVHPWDMMGGQRMKKWMLPWLVAMPAETQLAILSL
PicC	199	LFGGVFDRCPE.VKLVLGHMGEGLPFORWRYDSRFAVYPHGVTLKRKPSEYIGSNILITTSGVCSAPTLMG
gammaRSD	201	MGSGLFDKYPA.LKIILGHMGEGLPYSMWRIDHRNAWIKTTPKYPAKRKIVDYFNENFYLTTSGNFRTOTLID
5CVD	223	LTTGVFDRYPN.LQVVVGHGGEAIPNWLFRVDYMHKAGVRSQR.YERLKPLQHDMFHYMRNNVLVTTSGMASEPTIKL
2,3DHBD	205	VTNGVFDRHPK.LQIIMGHLGEHVPFDMWRINHWFEDRKKLLGLAETCKKTIRDYFAENIWITTSGHFSTTTLNF
SalD	207	IVNGVFDRFPK.LKVILGHLGEHIPGDFWRIEHWFEHCSRPLAKSRGDVFAEKPLLHYFRNNIWLTTSGNFSTETLKF
PATG	190	LTRGVIRDYPG.CKIILSHAGGTLPYLIHRAATMLPFMPRNLGMSREEIVEAARTFYFDTAISANPVTLKA
HndA	187	VLNGVIARYPN.VRIILSHAGGFLPYAAYRFAELAPGVRHDVPD.RDGVLDLLRRFYFDTAISA.PSALPS
ACMSD	210	LLSGAFERIPKSLKICFGHGGGSFAFLLGRVDNAWRHRDIVREDCPRPPSEYVDR.FFVDSAVFNPGALEL
PicC	269	AIGEMGAEAVLFSVDYPYESTELAADFIEAAPMDDKTREL.VCYGNAARLFKLDRS
gammaRSD	273	AILEIGADRILFSTDWPFENIDHAADWFENTSISEADRKK.IGWGNAQNLFKLNR
5CVD	299	CMEQLGEDRVMYAMDYPYEYVADEVRVHDNLAIPFAQKKK.LMQTNAERVFKL
2,3DHBD	279	CMAEVGSDRILFSIDYPFETFSDACEWFDNAELNGTDRLK.IGRENAKKLFKLDSYKDSSA
SalD	284	CVEHVGAERILFSVDSPYEHIDVGCGWYDDNAKAIMEAVGGEKAYKDIGRDNAKKLFKLGKFYDSEA
PATG	260	LLEFAKPGHVLFGSDFFNAPRGAITHFTSFLEGYDNMSEETRRLVEREAALELFPRLRGQSTRACL
HndA	255	LVAFAQPDRVLYGSDFFYAPASVGTSFTAALDAYPAPDADRHAAIHRTNALPLFPRLAALAR
ACMSD	280	LVSVMGEDRVMLGSDFFPLGEQKIGGLVLSSNLGESAKDKISGNASKFFNINV

