1	Inhibition of glycine cleavage system by pyridoxine 5'-phosphate causes synthetic
2	lethality in glyA yggS and serA yggS in Escherichia coli
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#### 15 Abstract

16 The YggS/Yb1036c/PLPBP family includes conserved pyridoxal 5'-phosphate (PLP)-17 binding proteins that plays a critical role in the homeostasis of vitamin  $B_6$  and amino acids. Disruption of members of this family causes pleiotropic effects in many organisms by 18 19 unknown mechanisms. In *Escherichia coli*, conditional lethality of the *yggS* and *glyA* 20 (encoding serine hydroxymethyltransferase) has been described, but the mechanism of 21 lethality was not determined. Strains lacking vggS and serA (3-phosphoglycerate 22 dehydrogenase) were conditionally lethality in the M9-glucose medium supplemented 23 with Gly. Analyses of vitamin B<sub>6</sub> pools found the high-levels of pyridoxine 5'-phosphate 24 (PNP) in the two yggS mutants. Growth defects of the double mutants could be eliminated 25 by overexpressing PNP/PMP oxidase (PdxH) to decrease the PNP levels. Further, a serA 26 pdxH strain, which accumulates PNP in the presence of yggS, exhibited similar 27 phenotype to serA yggS mutant. Together these data suggested the inhibition of the glycine cleavage (GCV) system caused the synthetic lethality. Biochemical assays 28 29 confirmed that PNP disrupts the GCV system by competing with PLP in GcvP protein. Our data are consistent with a model in which PNP-dependent inhibition of the GCV 30 system causes the conditional lethality observed in the *glyA* yggS or *serA* yggS mutants. 31 32

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#### 35 Introduction

36 Pyridoxal 5'-phosphate (PLP) is the biologically active form of vitamin B<sub>6</sub>, and is used as 37 a cofactor in various kinds of enzymes in most organisms. PLP-dependent 38 enzymes catalyze more than 140 distinct reaction types, including transamination, 39 decarboxylation, racemization, and elimination or replacement of chemical groups at the  $C_{\beta}$  or  $C_{\gamma}$ , and are involved in sugar, lipid, and amino acid metabolism (Percudani et al., 40 2003; Eliot et al., 2004; Toney, 2011). In addition to the coenzyme function, PLP may 41 42 function as an antioxidant (Bilski et al., 2000) and modulator of steroid hormone (Tully 43 et al., 1994) or P2X purinoceptor7 (Lambrecht et al., 2002; Thériault et al., 2014). Most 44 of the bacteria utilize PLP-dependent transcriptional factors to modulate metabolisms of 45 vitamin B<sub>6</sub> and/or amino acid (Belitsky, 2004; Belitsky, 2014; Takenaka et al., 2015; 46 Tramonti et al., 2018).

47 Two different pathways, deoxyxylulose 5-phosphate (DXP)-dependent or DXP-48 independent pathways, are known for PLP biosynthesis (Laber et al., 1999; Fitzpatrick et 49 al., 2007; di Salvo et al., 2011). The DXP-dependent pathway contains multiple reactions 50 and is found in a few members of the  $\gamma$ -proteobacteria and *Escherichia coli*. This pathway 51 utilizes 4-phospho-hydroxythreonine and DXP to synthesize pyridoxine 5'-phosphate (PNP) as a precursor of PLP. The DXP-independent pathway is found in archaea, fungi, 52 most eubacteria, and plants, and PLP is formed directly from glutamine, ribose 5-53 54 phosphate/ribulose 5-phosphate, and glyceraldehyde 3-phosphate/dihydroxyacetone 55 phosphate by the action of PLP synthase (Burns et al., 2005; Raschle et al., 2005; di Salvo et al., 2011; Fitzpatrick et al., 2007). Mammals do not possess the PLP biosynthetic 56 pathway and thus obtain B<sub>6</sub> vitamers from their diet. In all organisms, B<sub>6</sub> vitamers are 57 converted to PLP by a salvage pathway that involves pyridox(am)ine 5'-phosphate 58 59 oxidase (EC 1.4.3.5), phosphatases, and/or pyridoxal/pyridoxine kinase (EC 2.7.1.35) 60 (Fitzpatrick et al., 2007; di Salvo et al., 2011). In mammals, PLP-dependent enzymes play essential roles in the metabolism of neurotransmitters including  $\gamma$ -aminobutyric acid, 61

dopamine, epinephrine, serotonin, serine, and histamine. Therefore, a deficiency of PLP
is implicated in neurological disorders. Excess PLP is thought to be toxic to mammals,
probably due to the non-specific binding of reactive aldehyde group at C4' to amines
and/or thiols.

In man, disruption of B<sub>6</sub> homeostasis, including interfering with B<sub>6</sub> synthesis, recycling 66 67 (PNPO deficiency), and transportation (tissue non-specific alkaline phosphatase 68 deficiency), and inactivation of PLP by accumulation of a metabolite that reacts with PLP 69 (ALDH7A1 or ALDH4A1 deficiency), causes vitamin B<sub>6</sub>-dependent epilepsy, which is 70 treatable with high-dose of B<sub>6</sub> vitamers such as pyridoxine (PN) and PLP (Farrant et al., 71 2001; Narisawa et al. 2001; Mills et al., 2005; Mills et al., 2006; Wilson et al., 2019). 72 Darin et al. recently found that a mutation in PROSC, proline synthetase co-transcribed 73 homolog protein (renamed as PLPBP) causes vitamin B<sub>6</sub>-dependent epilepsy (Darin et al., 74 2016). The PLPBP-deficiency exhibits complex phenotypes including altered levels of 75 vitamin B<sub>6</sub> vitamers and imbalances of neurotransmitters with evidence of reduced 76 activity of PLP-dependent enzymes (Plecko et al., 2017; Shiraku et al., 2018; Johnstone 77 et al., 2019).

78 PLPBP is a member of a highly conserved PLP-binding protein family (COG0325, 79 referred to herein as YggS/Ybl036c/PLPBP) and the members of this protein family are 80 present in three domains of life. The YggS/Ybl036c/PLPBP family exhibits typical TIM-81 barrel structure and the PLP is bound to a Lys residue through Schiff-base linkage (Eswaramoorthy et al., 2003; Ito et al., 2013; Tremiño et al., 2017). The PLP-binding is 82 83 essential for the function of this protein family (Ito et al., 2013; Tremiño et al., 2018). In 84 human cells, the PLPBP localizes in mitochondria and cytosol (Johnstone et al. 2019). 85 Although this protein family shows structural similarity to bacterial alanine racemase and eukaryotic ornithine decarboxylase (Eswaramoorthy et al., 2003; Tremiño et al., 2017), 86 representatives of fold-type III PLP-dependent enzyme, no enzymatic activity has been 87 detected with the purified proteins (Ito et al., 2013). Previous studies showed that in E. 88

89 coli mutation of this protein family (yggS) causes pleiotropic phenotypes, including 90 altered flux in the Thr-Ile-Val metabolic pathways (Ito et al., 2013; Ito et al., 2016; Ito et 91 al., 2019), increased sensitivity to PN, accumulation of pyridoxine 5'-phosphate (PNP) 92 in the cells (Prunetti et al., 2016), and synthetic lethality with glyA (serine 93 hydroxymethyltransferase) (Nichols et al., 2011; Côté et al., 2016; Prunetti et al., 2016). 94 A recent study showed that the two phenotypes, PN sensitivity and the aberration of amino 95 acid metabolism in Thr-Ile-Val pathway, are caused by high-levels of PNP via an 96 unidentified mechanism (Ito et al., 2019). In cyanobacteria, disruption of the 97 cyanobacterial member of this protein family (PipY) increases sensitivity to antibiotics 98 targeting essential PLP-dependent enzymes and causes synthetic lethality with putative 99 cysteine synthase CysK (Labella et al., 2017). Studies with the yeast strain lacking 100 Ybl036c suggest a role of this protein in mitochondrial metabolism (Johnstone et al., 2019). The same authors demonstrated that the *plpbp*<sup>-/-</sup> zebrafish larvae exhibit seizure 101 phenotype and impaired biosynthesis and/or homeostasis of PLP-dependent 102 103 neurotransmitters (Johnstone et al., 2019).

104 Although the mutation of the YggS/Ybl036c/PLPBP family causes significant effects on 105 the metabolisms of vitamin B<sub>6</sub> and amino acid, limited information is available for the 106 molecular mechanisms. In this study, we undertook a detailed study about the conditional 107 lethality of glvA and yggS observed in E. coli. GlyA catalyzes the PLP-dependent 108 conversion of Ser to Gly while transferring the hydroxymethyl group to tetrahydrofolate 109 (THF) to generate 5, 10-methylenetetrahydrofolate (5, 10-mTHF). The GlyA reaction provides most of Gly and 5, 10-mTHF in E. coli and the latter is required for Met, 110 111 thymidine, and purine biosynthesis (Blaklet, 1955; Blakley 1969; Schirch et al., 1985) 112 (Fig. 1). The reason why the yggS mutation induces lethality under the glyA-background remained to be elucidated. 113

Here we show that the *yggS* mutation under the *serA* (3-phosphoglycerate dehydrogenase)-background also causes conditional lethality in the M9-glucose medium

- 116 supplemented with Gly. Further studies demonstrated that high-levels of PNP, which is
- 117 induced by *yggS* mutation, disrupts glycine cleavage (GCV) system and causes lethality
- 118 under the *glyA* or *serA*-background. The PNP-dependent inhibition of the GCV system
- 119 may provide a plausible reason for the diverse phenotypes observed in *E. coli*.
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#### 121 **Results and Discussion**

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#### 123 **Poor growth of** *glyA yggS* double mutant and its alleviation by inosine

124 Null alleles of *yggS* and *glvA* (encoding serine hydroxymethyl transferase; EC 2.1.2.1) 125 in *E. coli* are synthetically lethal on LB medium, although both the single mutant grow 126 well on the same medium (Nichols et al., 2011; Côté et al., 2016; Prunetti et al., 2016). 127 We constructed a glyA yggS double mutant using LB medium supplemented with 128 methionine and thymidine. These two compounds partially alleviate the growth defect of 129 the double mutant on LB (Prunetti et al., 2016). In our hands, the double mutant grew in 130 LB medium, although it showed severe growth defect (Fig. 2A). A previous study suggested that this LB sensitivity phenotype is due to amino acids such as Thr and Ala 131 132 (Prunetti et al., 2016). The apparent inconsistency could be due to the different nutrients 133 composition of the media that affect the growth of the glyA yggS mutant. Consistent with 134 previous results, the glyA yggS mutant had a significant growth defect in an M9-glucose 135 medium supplemented with 0.4% casamino acid (M9-Casa medium) when compared to 136 the glyA mutant (Fig. 2B). Plasmid-borne expression of yggS completely restored the growth of the double mutant to the level of the glyA mutant, confirming the growth defect 137 was due to the yggS mutation (Fig. 2B). The addition of Ser or Gly (5 mM) to the M9-138 139 Casa medium did not significantly affect the growth of the glyA yggS mutant (data not 140 shown). These data suggested that neither Ser toxicity nor Gly limitation caused the 141 growth defect in the *glyA* yggS double mutant.

In some conditions, a *yggS* mutant overproduces Val, which is toxic to *E. coli* K-12 (Ito et al., 2019). To ask if increased Val could be responsible for inhibition of growth of the *glyA yggS* mutant, the amino acid pool of the double mutant grown in the M9-Casa medium was analyzed. A significantly altered amino acid pool was found in the double mutant as compared to that of the *glyA* mutant strain (Fig. 2C). The *glyA yggS* double mutant accumulated Met, 2-AB, Val, Ile, and Leu compared to the *glyA* mutant (Fig. 2C). 148 Ile alleviates Val toxicity in *E. coli* K-12 strain (Leavitt et al. 1962; Lawther et al., 1981), 149 but providing Ile (0.1 - 1 mM) to the M9-Casa medium did not improve the growth of 150 the double mutant (data not shown). These results indicated that Val toxicity is not the 151 cause of the growth defect in the *glyA yggS* mutant, which accumulated 2.5-times more 152 Val than the *glyA* mutant.

153 Supplementation with a variety of metabolites found that guanosine or inosine (0.2 mM) 154 eliminated the growth defect of the glyA yggS mutant in the M9-Casa medium (Fig. 3A, 155 3C). Adenosine was also effective for restoring the growth of the glvA yggS mutant (Fig. 156 3A). Supplementation of either of Gly, Met, thymidine, cytidine, or pantothenate (0.2 157 mM), or the combination of the two did not significantly support the growth of the double 158 mutant under the condition examined (Fig. 3A, B, C). Inosine is probably phosphorylated 159 to inosine-5'-phosphate (IMP) by a salvage pathway and used as a purine source in E. coli 160 (Xi et al., 2000). The growth characteristics suggest that yggS mutation under the glvA-161 background influences the purine biosynthesis. Because the purine biosynthesis requires 162 one-carbon unit metabolism, it was also possible that the one-carbon unit metabolism is 163 affected in the *yggS* mutant.

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### Growth of a *serA yggS* double mutant suggests compromised glycine cleavage (GCV) system

167 In a wild-type strain, GlyA and/or GCV system play role in the production of 5, 10mTHF (Fig. 1), and a glyA mutation, in combination with a lesion in any of the genes of 168 169 the GCV complex (gcvT, gcvH, gcvP, or lpd) is synthetically lethal (Côté et al., 2016). In 170 the glyA mutant, the GCV system is responsible for the production of 5,10-mTHF 171 (Kikuchi et al., 2008) (Fig. 1). The possibility that the GCV system was inhibited in a 172 yggS mutant was tested in the serA-background. The serA gene encodes 3phosphoglycerate dehydrogenase, and a serA mutant requires either Ser or Gly for growth. 173 174 When Ser is provided, both Gly and 5,10-mTHF are generated by GlyA. In the presence

of Gly, the GCV system is responsible for the production of 5, 10-mTHF, which is used for the synthesis of Ser by GlyA (Fig. 1). A *serA* mutant in which the GCV system is compromised cannot utilize Gly as a Ser source (Plamann et al. 1983). Therefore, we can assess the function of the GCV system *in vivo* using a *serA* mutant and providing Gly and/or Ser supplementation (see Fig. 1).

180 Growth of serA and serA yggS mutants was indistinguishable in the M9-glucose medium 181 supplemented with 2 mM Ser (M9-Ser medium), (Fig. 4A). When grown in the M9-182 glucose medium supplemented with 2 mM Gly (M9-Gly medium), the growth of the serA 183 *yggS* double mutant was significantly compromised compared to the *serA* mutant (Fig. 184 4B). The addition of Ser (1 mM) to the M9-Gly medium completely restored the growth 185 of the serA yggS double mutant, indicating that Ser limitation caused the growth defect 186 (data not shown). The *yggS* mutation did not affect the incorporation of Gly, since a *yggS* 187 single mutant was able to use Gly as the sole nitrogen source (data not shown). These 188 data supported the hypothesis that the GCV system is compromised in the yggS mutant.

189 To obtain further insight into the yggS mutation on the cellular metabolism, intracellular 190 amino acid compositions in the two strains were investigated. The levels of many amino 191 acids were altered significantly when grown in the M9-Gly medium (Fig. 4C, 4D). 192 Concentrations of 2-AB, GABA, ophthalmic acid (OA), Ile, Met, Gly, Thr, His, and Glu 193 were significantly elevated in the serA yggS double mutant, whereas those of Ala, Asp, 194 and Leu were slightly decreased (Fig. 4D). Many of the amino acids changed significantly 195 were the same ones previously reported (Ito et al. 2013; Ito et al., 2016; Ito et al., 2019), 196 suggested the perturbation of Thr/Ile/Val metabolic pathways in the serA yggS mutant.

197 GCV activities in the cell-free extract of *serA* and *serA yggS* strains grown in the M9-198 Gly medium were measured (Table 1). Unexpectedly, the GCV activity in the cell-free 199 extract of *serA yggS* strain was approximately 3-times higher than in the *serA* mutant (1.2 200  $\pm 0.1$ ,  $0.4 \pm 0.1$  nmol/min/mg, respectively). Similar experiments were performed with 201 the cell-free extract of *serA* and *serA yggS* strains grown in the M9-Ser medium. The GCV activity for the *serA yggS* strain was approximately 6-times higher than that for the *serA* mutant  $(0.31 \pm 0.02 \text{ nmol/min/mg}, 0.05 \pm 0.01 \text{ nmol/min/mg}, respectively})$ . These data eliminated the simple possibility that *yggS* mutation lowered the expression of some of the GCV components to cause the growth defect of the *serA yggS* mutant. The reason for the increased GCV activity in the *serA yggS* mutant is unknown. GlyA activities in these two strains on M9-Gly were not significantly different (*serA*; 0.22 ± 0.01 nmol/min/mg protein, *serA yggS*; 0.20 ± 0.01 nmol/min/mg protein).

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### 210 Perturbation of B<sub>6</sub> vitamer pool correlates with growth defect of *serA yggS* mutants

211 A caveat of *in vitro* enzyme assays is that the cell breakage can eliminate allosteric, or 212 competitive inhibition of enzymes due to dilution of relevant metabolites. Significantly, 213 the YggS/Yb1036c/PLPBP family has been implicated in vitamin B<sub>6</sub> homeostasis (Darin 214 et al., 2016; Prunetti et al., 2016; Johnstone et al., 2019; Ito et al., 2019), and the P-protein 215 of GCV (GcvP) is a PLP-dependent enzyme (Kikuchi et al., 2008). We found that 216 exogenous PN (> 1  $\mu$ M) significantly inhibited the growth of serA yggS double mutant in 217 the M9-Gly medium, but not in the presence of Ser (Fig. 5A). The growth of serA strain 218 was not affected by exogenous PN on either M9-Ser or M9-Gly media (data not shown). 219 It was possible that altered B<sub>6</sub> availability affected the GCV activity in vivo.

220 The concentration of total B<sub>6</sub> vitamers in serA and serA yggS mutants grown with Ser 221 or Gly were analyzed (Table 2). When grown in the M9-Gly medium the serA yggS 222 mutant accumulated approximately 7-times more PNP ( $30 \pm 5.8 \mu$ M) than the serA mutant  $(6.9 \pm 0.2 \,\mu\text{M})$  in the cells. The elevated PNP was eliminated by plasmid-borne expression 223 224 of YggS (by pUS) in the serA yggS mutant. High-levels of PNP was also detected when 225 the serA yggS mutant was grown in the M9-Ser medium (serA:  $5.9 \pm 0.4 \mu$ M, serAyggS: 226  $34.4 \pm 0.7 \mu$ M). We also quantified the levels of B<sub>6</sub> vitamers in the glyA and glyA yggS 227 double mutant. The glyA yggS double mutant also exhibited elevated levels of PNP in the 228 cells (Table 2). We have shown that exogenous PN significantly increases the intracellular

229 concentration of PNP in the yggS mutant (Ito et al. 2019). The PNP content in the *serA* 230 yggS mutant grown in the M9-Ser medium supplemented with 1  $\mu$ M PN was 25% higher 231 than when grown in the absence of PN (Fig. 5B). The GCV system is a multi-enzyme 232 complex composed of P-protein (GcvP, glycine dehydrogenase), T-protein (GcvT, 233 aminomethyltransferase), L-protein (Lpd, lipoamide dehydrogenase), and H-protein 234 (GcvH, lipoprotein), and the GcvP protein requires PLP for the function. These data led 235 to hypothesize that PNP inhibits the GCV system *in vivo*.

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#### 237 PNP inhibits the GCV system in vivo

238 The occurrence of the PNP-dependent inhibition of the GCV system was investigated 239 by eliminating the accumulation of PNP with PNP/PMP oxidase (PdxH). The 240 overexpression of the *pdxH* significantly restored the growth defect of the *serA yggS* 241 mutant (Fig. 6A and 6B). We found that the overexpression of gcvT-gcvH-gcvP operon also supported the growth of serA yggS double mutants in the M9-Gly medium. Further, 242 243 the expression of only gcvP alleviated the growth defect of serA yggS mutants (Fig. 6C and 6D). These data supported the idea that the high-levels of PNP disrupt the function 244245 of the GCV system by inhibiting the GcvP function in vivo.

246 It was formally possible that another factor(s) affected by yggS mutation inhibits the 247 GCV system. To eliminate this possibility, we constructed a serA pdxH double mutant. 248 Mutation of *pdxH* prevents further catabolism of PNP and induces accumulation of PNP under the  $yggS^+$  background (Table 2). If the growth defect of the serA yggS strain was 249 due to the accumulation of PNP, the serA pdxH double mutant was predicted to elicit a 250 251 similar phenotype to the serA yggS mutant. When grown in the M9-Ser medium supplemented with 10  $\mu$ M PL, the serA and serA pdxH (yggS<sup>+</sup>) strains had 252 253 indistinguishable growth rates (Fig. 6E). In contrast, when grown in the M9-Gly medium 254 supplemented with 10 µM PL, the serA pdxH double mutant did not grow (Fig. 6F). These phenotypic data are consistent with the idea of the PNP-dependent- and YggS-255

256 independent inhibition of the GCV system.

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#### 258 **PNP inhibits the GCV system** *in vitro*

259 The data shown in Table 2 represent the total intracellular concentration of B<sub>6</sub> vitamers 260 including that which are bound to enzymes and do not reflect the concentrations of free 261 PLP or PNP available in the cells. We thus estimated the intracellular concentrations of free B<sub>6</sub> vitamers in the serA and serA yggS mutants. We quantified the B<sub>6</sub> levels presented 262 263 in the protein-free fraction, which was obtained by passing through the cell-free 264 supernatant using a centrifugal filter device with the molecular wight cut-off of 10 kDa. 265 This experiment found that most of the intracellular PLP (~70%) present as protein-266 binding form, whereas most of the PNP and PMP ( $\sim$ 70%) exist as free-form (Fig. 7). This 267 difference probably reflects the fact that PLP forms Schiff-base in proteins. These data 268 indicate that the concentration of free PNP (estimated to be approximately 21  $\mu$ M) is 269 comparable to that of free PLP in the serA yggS mutant (Fig. 7).

270 In the GCV system, GcvP protein catalyzes the decarboxylation of Gly in a PLPdependent manner. We hypothesized that PNP competes with PNP in GcvP protein and 271 272 inhibits the GCV system. To test this possibility, GCV activity was measured in crude 273 extracts with added PLP (5  $\mu$ M) and varying concentrations of PNP from 0 - 250  $\mu$ M. 274 PNP reduced the GCV activity in a dose-dependent manner (Fig. 8A). In the presence of 275 50 µM PNP, activity was decreased by 50% at 250 µM PNP by 73%. The half-maximal inhibitory concentration (IC<sub>50</sub>) under the condition was calculated as 59  $\pm$  2  $\mu$ M. We 276 277 performed similar assays in the presence of 50 µM PLP. The experiments found that the 278 GCV activity was inhibited by 24% at 50  $\mu$ M PNP and 38% at 250  $\mu$ M PNP (IC<sub>50</sub> > 279 400 µM) (Fig. 8B). These data demonstrated that PNP inhibits the GCV system by 280 competing with PLP for binding to the enzyme. Considering the estimated free PNP level  $(21 \,\mu\text{M})$  and the IC<sub>50</sub> value (59  $\mu\text{M}$ ), we speculated that this inhibition may be more 281 282 pronounced in vivo. Little GCV activity was detected when assays were performed in the

absence of added PLP, suggesting that the PLP was not strongly bound to the GCVcomplex.

285 The ability of PNP to inhibit GlyA activity was also tested. Unlike the GCV system, 286 GlyA activity was not inhibited by PNP (Fig. 8C). The slight activation observed in the 287 presence of 250 µM PNP may be due to the contamination of PLP. The PNP stock used 288 in this experiment had contaminating (~2%) PLP. Some other PLP-dependent enzymes 289 including threonine synthase (ThrC), threonine dehydratase (IlvA), and branched-chain 290 transaminase (IlvE) are also PNP insensitive (Ito et al., 2019). As shown in Table 2, the 291 intracellular PNP concentrations in the *serA* yggS strain were not significantly affected by 292 the extracellular amino acid (M9+Ser: 30 µM and M9+Gly: 34 µM). In contrast, the 293 amino acid pool of serA yggS strain was significantly perturbed only in the M9+Gly 294 medium (Fig. 4C and 4D). The GCV reaction is essential for the growth in the M9+Gly 295 medium but dispensable in the M9+Ser medium. Therefore, we can estimate that the GCV system is the most PNP-sensitive target in E. coli under the condition examined. Wild-296 297 type E. coli cells produce most of their one-carbon units required by the GlyA reaction, and the GCV system plays a minor role (Meedel et al., 1974). This fact may explain the 298 299 milder phenotypes of the yggS mutants observed under the  $glyA^+$  background.

300 GCV system is highly conserved in diverse organisms ranging from bacteria to humans, 301 where it plays an important role in the degradation of Gly as well as one-carbon unit 302 metabolism. GcvP of E. coli, a PLP-dependent component of the GCV system, exhibits 303 approximately 50% sequence identity to the corresponding protein of human and yeast 304 protein. We found that the GCV system of Bacillus subtilis, whose glycine decarboxylase 305 (P-protein, subunit 1 and subunit 2) shows more than 40% identity to that of E. coli GcvP, 306 is weakly inhibited by PNP (Fig.5D). Although PNP is not known to accumulate in B. 307 subtilis, this result suggested the PNP-sensitive feature of the GCV system. The PLPBP-308 deficient human cells (fibroblasts and HEK293 cells) accumulate a considerable amount 309 of PNP (Johnstone et al., 2019). The PLPBP-deficient strain of Saccharomyces cerevisiae also accumulates high-levels of PNP in the cells (Vu et al, unpublished). It is interesting

311 to examine the occurrence of PNP-dependent inhibition of the GCV system and/or other

312 PLP-dependent enzymes in the PLPBP-deficient organisms.

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#### 314 Working model for diverse phenotypes observed in the *yggS* mutant

The *yggS* mutant accumulates some metabolites in the Thr/Ile/Val metabolic pathway (Val, ketoisovalerate (KIV), Ile, 2-AB, and OA) and exhibits decreased level of coenzyme A (Ito et al., 2013; Ito et al. 2016; Ito et al., 2018). Some of these changes correlate with increased levels of PNP, but the mechanism was unknown (Ito et al., 2019). The present findings may explain the mechanism of diverse phenotypes observed in the *yggS* mutant as described below and shown in Fig. 8.

- 321 The *yggS* mutation induces PNP accumulation by an unidentified mechanism. The high-322 levels of PNP inhibit the GCV system and decrease cellular 5,10-mTHF levels. The 323 limitation of 5,10-mTHF would lower the activity of 3-methyl-2-oxobutanoate 324 hydroxymethyltransferase (PanB), which catalyzes the conversion of the 5,10-mTHF and KIV to form 2-dehydropantoate and THF (Teller et al., 1976; Powers et al., 1976). It can 325 326 decrease the production of the CoA precursor 2-dehydropantoate and increase the KIV 327 (and its transamination product Val) in the cells. Val activates threonine dehydratase 328 (IIvA) and inhibits acetohydroxy acid synthases (AHAS I/III) to produce more 2-329 ketobutyrate (Leavitt et al., 1962; Eisenstein, 1991). 2-ketobutyrate is the precursor of Ile, 330 2-AB, and OA (Fig. 7).
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#### 332 Conclusion remarks

We demonstrated that high-levels of PNP induced by *yggS* mutation cause the conditional lethality in the *glyA- or serA-*background by inhibiting the GCV system. The PNP-dependent inhibition of the GCV system can provide a plausible reason for the diverse phenotypes observed in *E. coli*. Further investigations focusing on the metabolism

- of PNP will shed light on the molecular function of the YggS/Ybl036c/PLPBP family and
- 338 the link between the perturbation of  $B_6$  pools and diverse metabolic pathways.
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#### 340 **Experimental procedures**

#### 341 *Bacterial strains and culture conditions.*

342 The strains used in this study are listed in Table 3. When added to the medium, ampicillin 343 (Amp), chloramphenicol (Cm) and kanamycin (Km) were used at concentrations of 100 344 µg/ml, 30 µg/ml, and 50 µg/ml, respectively. Single-gene deletion mutants of E. coli 345 (Keio collection) and folD-expression plasmid (pCA24N-folD, ASKA clone) were 346 obtained from NBRP (Baba et al., 2006; Kitagawa et al., 2005). E. coli strains were grown 347 in an LB medium or M9-glucose medium as previously described (Ito et al. 2013; Ito et 348 al. 2019). Ser, Gly, or Met were added at a final concentration of 2 mM. Casamino acid 349 or nucleotide was supplemented at a final concentration of 0.4% or 0.2 mM, respectively. Unless otherwise noted, E. coli strains were grown at 30°C. Cells growth was recorded 350 351 by the OD-Monitor C&T apparatus (Taitec Co., Ltd., Koshigaya, Japan) using glass test tubes (16.5 mm in diameter by 165 mm in height), or the ELx808 (Biotec, Winooski, VT, 352 353 USA) using 96-wells plate.

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#### 355 *Molecular genetics and sequence analysis.*

356 The deletion of the yggS gene was performed using the bacteriophage  $\lambda$ -Red recombinase 357 system described by Datsenko and Wanner (Datsenko et al., 2000). The glyA yggS double mutant was constructed as follows. An *E. coli glyA* mutant (Keio collection, JW2535-KC) 358 359 harboring pCP20 (Cherepanov et al., 1995) was streaked on an LB plate and grown at 42°C, forming glyA<sup>-Km</sup> strain. A PCR product was generated with primers yggS-H1 and 360 361 yggS-H2 (Ito et al., 2009) using Tks Gflex DNA polymerase (TaKaRa) and pKD13 as a 362 template. The PCR product was purified from agarose-gel and electroporated into the glyA<sup>-Km</sup> harboring pKD46. Resultant transformants appeared on an LB plate containing 1 363

364 mM Met, 0.1 mM thymidine, and 30 µg/ml kanamycin were screened by PCR for the 365 appropriate insertion of the kanamycin-resistant gene with the primers yggS-200up and 366 yggS-300dwn (Ito et al., 2009). Construction of the serA yggS double mutant and serA 367 pdxH double mutant was performed in a similar way using the JW2880-KC (Keio 368 collection) as parental strains, respectively. Primers pair of pdxH-H1 (5'-ATGTCTGATAACGACGAATTGCAGCAAATCGCGCATCTGCGCCGTGAATGTGT 369 370 AGGCTGGAGCTGCTTCG-3') and pdxH-H2 (5'-371 TCAGGGTGCAAGACGATCAATCTTCCACGCATCATTTTCACGCTGGTCATATG 372 AATATCCTCCTTAG-3') were used for the construction of *pdxH*-deficient strain. 373 Insertion of kanamycin-resistant cassette at the pdxH locus was confirmed with the 374 primers pdxH-100up (5'-CGCATCGTCTTGAATAACTGTCAG-3') and pdxH-100dwn 375 (5'-CACCTTTGCCGGTACACGACTTTTC-3'). The gcv operon (gcvT-gcvH-gcvP) or 376 amplified with gcvT-fw *gcvP* gene was the primers (5'-377 GCTAACAGGAGGAATTAACCATGGCACAACAGACTCCTTTGTACG-3') or gcvP-378 fw (5'-GGCTAACAGGAGGAATTAACCATGACACAGACGTTAAGCCAGCTTG-3') 379 and gcvP-rv (5'-380 GATGAGTTTTTGTTCTACGTCGCCGAAGCGCCTTTAGAAAATAG-3') and cloned 381 into pBAD/MycHisC plasmid (Invitrogen) using In-Fusion HD Cloning Kit (Takara) 382 according to the manufacture's instruction.

383

#### 384 GCV assay

GCV activity was assayed as described previously with some modifications (Meedel et al., 1974; Nagarajan et al., 1997). *E. coli* cells (*serA* or *serA yggS* strains) were grown in an M9-glucose medium supplemented with 1 mM Ser or 1 mM Gly. To examine the effect of PNP to GCV system, a cell-free extract of *glyA*-deficient strain grown in M9-glucose medium supplemented with 1 mM Gly was used. The *gcvP*-deficient *E. coli* was used as a negative control. The cells pellet (500 mg) was sonicated in 5 ml of phosphate buffer 391 (20 mM potassium phosphate buffer (pH 7.4), 10% glycerol) and centrifuged (20,000 g, 392 30 min, 4°C). The protein concentrations in the cell-free extract were determined using 393 the BioRad Bradford assay (BioRad). The reaction was initiated by mixing 125 µl of the 394 cell-free extract and 125 µl of a reaction mixture containing 40 mM potassium buffer (pH 395 7.4), 4 mM dithiothreitol (DTT), 2 mM tetrahydrofolate (THF), 5 µM or 50 µM PLP, 0 to 250  $\mu$ M PNP, 4 mM nicotinamide adenine dinucleotide (NAD<sup>+</sup>), and 1.2 nmol [2-<sup>14</sup>C] 396 397 Gly (53.0 mCi / mmol). After 2 hours of incubation at 30°C, 250 µl of a solution (a 398 mixture of 150 µl of 1 M sodium acetate (pH 4.5) and 100 µl of 0.1 M formaldehyde) and 399 then 150 µl of 0.4 M dimedon in 50% ethanol was added to the reaction mixture. After 5 400 min incubation at 65°C, 2.5 ml of toluene was added and vortexed vigorously for 1 minute. 401 After centrifugation, the radioactivity of the toluene layer (2 ml) was measured by liquid 402 scintillation counting using AccuFLEX LSC-7200 (Hitachi Aloka Medical Co., Ltd.). 403 PNP was synthesized by reduction of PLP with NaBH<sub>4</sub>, and purified as described 404 previously (Argoudelis et al. 1986).

405

#### 406 GlyA assay

407 GlyA activity in the cell-free extract of serA or serA yggS double mutant was assayed 408 according to the protocol described previously with some modifications (Schirch et al., 409 1968). Cell-free extracts of serA or serA yggS double mutant were prepared as described 410 above. Reaction was initiated by adding 10 µl of the cell-free extract to the 890 µl of reaction mixture containing 50 mM Hepes-NaOH (pH 7.5), 0.4 mM THF, 0.2 mM 411 412 NADP<sup>+</sup>, 5 mM Ser, 5 µM PLP, and 0.1% 2-mercaptoethanol. After 20 min incubation at 413 37°C, 100 µl of 0.1 M potassium carbonate buffer (pH 9.5) was added to terminate the 414 reaction. Then 20 µl of the purified His-tagged E. coli 5, 10-methylene tetrahydrofolate 415 dehydrogenase (FolD, 5 mg/ml) was added, and the absorbance change at 340 nm was 416 recorded. The recombinant FolD was prepared from an E. coli AG1 strain harboring 417 pCA24N-folD by His-tag affinity chromatography using a conventional method.

418

#### 419 Amino acid and B<sub>6</sub> vitamer analysis

420 The glyA or glyA yggS double mutants were grown in an M9 medium supplemented with 421 0.4% casamino acid. The serA or serA yggS double mutants were grown in an M9 medium 422 containing 1 mM Ser or 1 mM Gly. The log-phase cells ( $OD_{600}$  of 0.4 - 0.6) were collected 423 by centrifugation. For the quantification of total B<sub>6</sub> content, the cell pellet was 424 resuspended in 10 volumes (v/w) of 0.8 M HClO<sub>4</sub> (100 µl of the HClO<sub>4</sub> solution for 10 425 mg E. coli wet cells). The suspension was vortexed and incubated for 30 min at 4°C. 5 426 volumes (v/w) of 0.8 M K<sub>2</sub>CO<sub>3</sub> solution (50 µl for 10 mg E. coli cells) was added. Debris 427 was removed by centrifugation (20,000  $\times$  g for 20 min at 4°C). For the analyses of "free 428 B<sub>6</sub>" (B<sub>6</sub> vitamer that is not tightly bound to large-molecule), the cells pellet was sonicated 429 in 5 volumes of 50 mM Hepes-NaOH buffer (pH 7.5). The cell-free supernatant was 430 obtained by centrifugation (20,000 g, 30 min). Protein concentration in the soluble 431 fraction was determined with the BioRad protein assay kit (BioRad) using bovine serum 432 albumin as standard. The protein-free solution was prepared by passing the cell-free 433 supernatant through the Amicon-ultra centrifugal membranes with 10 kDa molecular 434 weight cutoff. These solutions (180 µl) were deproteinized with HClO<sub>4</sub> (8 M, 20 µl) and 435 neutralized with 100 µl of the 0.8 M K<sub>2</sub>CO<sub>3</sub>. After centrifugation, samples were diluted 436 and used for the HPLC analyses as previously described (Ito et al., 2018; Ito et al., 2019). 437 Previous results reported that 1 mg of wet cell weight corresponds to 0.23 mg of dry cell 438 weight, and the amount of cytoplasmic water in cells grown in the M9-glucose is 2.0 µl/mg dry cell weight (Glazyrina et al. 2010, Cayley et al. 1991). We thus estimated that 439 440 1 pmol/mg wet cells of PLP correspond to 2.17 µM of intracellular PLP.

441

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630	٠	
631		

### 632 Table 1 Activity of GCV or GlyA in the cell-free extract of *serA* and *serA yggS*633 strains

The *serA* or *serA yggS* strains harboring pU0 plasmid were grown in the M9-Ser or M9-Gly medium. Ampicillin was added for plasmid maintenance. The *E. coli* strains were collected at log-phase and disrupted by sonication. The enzyme activities were analyzed as described in Experimental Procedures. The data represent the averages and standard deviations from triplicate experiments.

- 639
- 640

	GCV activity		GlyA activity	
	(5, 10-mTHF production (pmol)/mg protein)		(5, 10-mTHF production (pmol)/mg protein/min)	
	+ Ser	+ Gly	+ Ser	+ Gly
serA	11 ± 1	$69 \pm 4$	$170 \pm 6$	$220\pm12$
serA yggS	$66 \pm 1$	$215\pm19$	$192 \pm 8$	$202\pm15$

641

#### 643 Table 2 Total vitamin B<sub>6</sub> levels in the *E. coli* strains

The *serA* or *serA yggS* strains harboring pU0 plasmid or complementary pUS plasmid (expresses YggS protein) were grown in the M9-Ser or M9-Gly medium. The *serA pdxH* strain was grown in the M9-Ser medium in the presence of 10  $\mu$ M PL. The *glyA* and *glyA yggS* strains were grown in the M9-Casa medium. Amp (100  $\mu$ g/ml) was added for plasmid maintenance. The *E. coli* strains were collected at log-phase and the total B<sub>6</sub> pools were analyzed as described in Experimental Procedures. The data represent the averages and standard deviations from triplicate experiments.

651

	PLP conc. (µM)	PNP conc. (µM)	PMP conc. (µM)
(M9 + Gly)			
serA	$57 \pm 4$	$7\pm0.2$	$146\pm14$
serA yggS	$49\pm10$	$32\pm5.8$	$121\pm24$
serA yggS/yggS <sup>+</sup>	$62 \pm 2$	$9\pm0.6$	$147\pm14$
(M9 + Ser)			
serA	$68 \pm 5$	$7\pm0.4$	$145 \pm 3$
serA yggS	$78 \pm 4$	$38 \pm 1$	$132\pm 6$
serA yggS/yggS <sup>+</sup>	$69 \pm 3$	$6\pm0.3$	$138 \pm 6$
serA pdxH	$55.7 \pm 3.6$	$159\pm36$	$44 \pm 10$
(M9 + Casa)			
glyA	$83 \pm 7$	N.D. <sup>a</sup>	$84 \pm 2$
glyA yggS	98 ± 11	$28 \pm 1$	$127 \pm 6$

<sup>a</sup>, N.D.: Not Detected

Strains glyA E. coli BW25113 glyA::Km (JW2535-KC) Keio collection  $glyA^{-Km}$ E. coli BW25113 glyA This study E. coli BW25113 glyA yggS::Km glyA yggS This study E. coli BW25113 serA::Km (JW2880-KC) Keio collection serA ser ser ser gcv рU pU pB.

Table 3 E. coli strains and plasmids used in this study 654

serA <sup>-Km</sup>	E. coli BW25113 serA	This study	
serA yggS	E. coli BW25113 serA yggS::Km	This study	
serA pdxH	<i>E. coli</i> BW25113 <i>serA pdxH</i> ::Km	This study	
gcvP	<i>E. coli</i> BW25113 <i>gcvP</i> ::Km (JW2871-KC)	Keio collection	
Plasmids			
pU0	pUC19 containing a partial sequence of <i>yggS</i>	Ito et al., 2009	
pUS	pUC19 expressing yggS	Ito et al., 2009	
pBAD24	pBAD24 empty vector	Laboratory collection	
pBAD24-pdxH	pBAD24 containing <i>pdxH</i> from <i>S. enterica</i>	Vu et al. to be	
		published	
pBAD-gcvTHP	pBAD-MycHisC containing gcvT-gcvH-gcvP	This study	
pBAD-gcvP	pBAD-MycHisC containing gcvP	This study	
pCA24N-folD	pCA24N containing <i>folD</i> (JW0518-AM)	ASKA clone	
pKD13	A template plasmid for gene disruption. The	Datsenko et al. 2000	
	Km <sup>r</sup> gene is flanked by FRT sites.		
pKD46	Lambda Red recombinase expression plasmid	Datsenko et al. 2000	
pCP20	Yeast Flp recombinase expression plasmid with	Nagarajan et al. 1997	
	temperature-sensitive replication.		

#### 657 Figure legends

#### 658 Figure 1 Reaction of GlyA and GCV system

- GlyA is PLP-dependent enzyme and catalyzes conversion of Ser to Gly, while transferring the hydroxymethyl group to tetrahydrofolate (THF), and generates 5, 10-methyltetrahydrofolate (5,10-mTHF). GCV system cleaves Gly to  $CO_2$ , ammonia and provides 5,10-mTHF. GlyA and GCV reactions require PLP. In the wild-type *E. coli*, GlyA can provide most of the 5,10-mTHF (Meedele et al., 1974). In the absence of *glyA*, GCV system provides 5,10-mTHF for one-carbon biosynthesis.
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### Figure 2 Effect of *yggS* mutation under *glyA* background on the growths and intracellular amino acid pool

- 668 (A) Growth of the glyA and glyA yggS mutants in the LB medium. (B) Growth of the glyA 669 strain and glyA yggS mutants both harboring pU0 plasmid (pUC19 containing partial 670 sequence of yggS) and glyA yggS mutant harboring pUS plasmid (yggS expression vector) 671 (Ito et al. 2009) in the M9-Casa medium. Cells growth was recorded by the ELx808. (C) 672 Intracellular amino acid pool of glyA or glyA yggS mutants grown in the M9-Casa medium. Amino acid pools were analyzed as described in Experimental procedure. Disruption of 673 yggS under glyA background affects Ile/Val and Met metabolisms. (\*p < 0.05, \*\*p < 0.01, 674 \*\*\*p <0.001, \*P < 0.05, \*\*P < 0.01, student's t-test) 675
- 676

#### Fig. 3 Effect of nucleotide and/or amino acid on the growth of glyA yggS

Growths of the glyA (circle) and glyA yggS double mutant (square) in the M9-Casa medium in the presence of nucleotide and/or amino acid. The concentration of nucleotide or amino acid (Met or Gly) was 0.2 mM or 2 mM, respectively. Guanosine (A), adenosine (B), or inosine (C) supported the growth of glyA yggS double mutant. Adenine inhibited the growth of the glyA (A). Other nucleotides and amino acids did not significantly affect the growth of glyA strain (data not shown). The data represent the averages and standard 684 685 deviations from triplicate experiments. Cells growth was monitored by the ELx808.

- Fig. 4 Effect of *yggS* mutation under *serA* background on the growths and
   intracellular amino acid pool
- 688 (A, B) Growths of the serA and serA yggS double mutant in the (A) M9-Ser or (B) M9-Gly medium. When grown in the M9-Ser medium, the two strains exhibited almost 689 690 identical growth. When grown in the M9-Gly medium, the serA vggS double mutant 691 showed poor growth. Cells growth was recorded by the ELx808. (C, D) Differences of 692 amino acid pools of serA and serA yggS grown in the (C) M9-Ser or (D) M9-Gly medium. 693 When grown in the M9-Gly medium, the amino acid pool of serA yggS strain was 694 significantly different from that of serA strain. Experiments were performed in triplicate, 695 and data are represented as the fold-change. (\*p < 0.05, \*\*p < 0.01, \*\*\*p < 0.001, \*P < 0.00696 0.05, \*\*P < 0.01, student's t-test)
- 697

## Fig. 5 Effect of PN on the growths and concentrations of total B<sub>6</sub> vitamers of serA *yggS* strain

(A) Growths of the *serA yggS* double mutant in the M9-Ser (black) or M9-Gly medium (red) in the presence (open symbol) or absence of PN (1  $\mu$ M) (closed symbol). When grown in the M9-Gly medium, the growth of *serA yggS* double mutant was further inhibited by PN. Cells growth was recorded by the OD-Monitor C&T apparatus. (B) Concentrations of B<sub>6</sub> vitamers in *serA yggS* mutant grown in the absence or presence of 1  $\mu$ M of PN. Exogenous PN significantly increased the intracellular content of PNP.

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#### 707 Fig. 6 PNP inhibits GCV system in vivo

(A, B, C, D) Growths of *serA* and *serA yggS* double mutant harboring pBAD empty vector

709 (pBAD), pBAD-pdxH, pBAD-gcvTHP ( $gcvTHP^+$ ), or pBAD-gcvP ( $gcvP^+$ ) plasmid

710  $(pdxH^+)$  were compared in the M9+Ser or M9+Gly medium. Expression of gcvT-gcvH-

711 gcvP (gcvTHP<sup>+</sup>), gcvP, or pdxH was induced by 0.2% or 0.02% arabinose, respectively. 712 The expression of *pdxH*, *gcvT-gcvH-gcvP*, or *gcvP* significantly improved the growth of 713 serA yggS double mutant in the M9-Gly medium. Note that growth of serA mutant strain 714 was not significantly affected by the overexpression of gcvP or pdxH. (E, F) Growth of 715 serA and serA pdxH double mutant in a (E) M9-Ser + 10  $\mu$ M PL or (F) M9-Gly + 10  $\mu$ M 716 PL medium. The serA pdxH double mutant exhibited lethality in the M9-Gly + 10  $\mu$ M PL 717 medium. Cells growth was recorded by the OD-Monitor C&T apparatus (panels C, D) or 718 the ELx808 (panels A, B, E, F) using 96-wells plate.

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#### Fig. 7 Estimation of free B<sub>6</sub> levels in the *serA yggS* mutant

The *serA* or *serA yggS* double mutant was cultivated in the M9+Gly medium. The cells were disrupted and centrifuged. The resultant cell-free fraction was passed through the centrifugal filter device (10 kDa-cut off) and obtained the protein-free fraction. The B<sub>6</sub> levels in the cell-free fraction (total B<sub>6</sub>) or the protein-free-fraction were determined as described in the Experimental procedure. In the *serA yggS* mutant, most of the PNP was presented as free-form and the concentration was almost identical to the free PLP concentration.

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#### 730 Fig. 8 PNP inhibits GCV system in vitro

GCV activity was measured in the presence of (A) 5  $\mu$ M or (B) 50  $\mu$ M of added PLP, and various concentrations of PNP (0, 5, 50, or 250  $\mu$ M). Cell-free extract of *glyA* strain grown in the M9-Gly medium was used for the analyses. No GCV activity was detected in the absence of added PLP. (C) Effect of PNP on GlyA activity was also assayed using a purified GlyA in the presence of 5  $\mu$ M PLP and various concentration of PNP. (D) Effect of PNP on GCV activity of *B. subtilis* was also assayed using cell-free extract in the presence of 5  $\mu$ M PLP and various concentration of PNP. Experiments were performed in triplicate, and data represent the averages and standard deviations of the means.

# Fig. 9 Connection of PNP, GCV system, and phenotypes observed in the *yggS*deficient *E. coli*.

Deletion of *yggS* in *E. coli* induces accumulation of PNP by unidentified mechanism.
High-levels of PNP compete with PLP and inhibit GCV system. Disruption of GCV
system can decrease 5,10-mTHF supply, which may decrease flux into pantothenate
production and increase flux for Val production. Val stimulates threonine dehydratase
(IlvA) to produce more 2-ketobutyrate (2-KB) as a precursor for 2-aminobutyrate (2-AB),
Ile, and ophthalmic acid (OA).

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