

1 **Phosphogluconolactonase as the linchpin of an efficient pentose phosphate pathway**

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15 **ABBREVIATIONS:**

16 6PGL : 6-Phosphogluconolactone

17 δ -6PGL : δ -6-phosphogluconolactone

18 GNT: Gluconate

19 6PGNT: 6-phosphogluconate

20 oxPPP: oxidative branch of the pentose-phosphate pathway

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23

24 **ABSTRACT:**

25 The metabolic networks of microorganisms are remarkably robust to genetic and environmental
26 perturbations. This robustness stems from redundancies such as gene duplications, isoenzymes,
27 alternative metabolic pathways, and also from non-enzymatic reactions. In the oxidative branch of the
28 pentose-phosphate pathway (oxPPP), 6-phosphogluconolactone hydrolysis into 6-phosphogluconate
29 is catalysed by 6-phosphogluconolactonase (Pgl) but in the absence of the latter, the oxPPP flux is
30 thought to be maintained by spontaneous hydrolysis. However, in Δpgl *Escherichia coli*, an
31 extracellular pathway can also contribute to pentose-phosphate synthesis. This raises question as to
32 whether the non-enzymatic reaction can compensate for the absence of 6-phosphogluconolactonase
33 and, ultimately, on the role of 6-phosphogluconolactonase in central metabolism. Our results indicate
34 that in the absence of Pgl, this bypass pathway accounts for the entire flux into the oxPPP, suggesting
35 that non-enzymatic hydrolysis does not compensate for the absence of Pgl and demonstrating that Pgl
36 is critical for an efficiently functioning oxPPP.

37

38 **INTRODUCTION**

39 Metabolic networks are a set of interconnected chemical reactions, most of which are catalysed by
40 enzymes. The interplay between chemical reactions provides alternative routes for adaptation to
41 genetic or environmental perturbations; the network organisation of metabolic systems thus
42 underpins the metabolic robustness and adaptability of cells. The central metabolism of *Escherichia*
43 *coli* is a model of metabolic robustness since very few of the associated genes are indispensable for
44 growth on glucose minimal medium^{1,2}. Furthermore, *E. coli* knock-out mutants lacking key central
45 metabolic enzymes have similar growth phenotypes^{3–7}. This robustness is in part the result of
46 metabolic flux rerouting^{6,7} but stems also from local compensatory mechanisms based on redundancy,
47 such as the presence of isozymes and alternative pathways^{2–5,7}.

48 Compensatory mechanisms involving non-enzymatic reactions are harder to identify because they
49 cannot be studied by metabolic reconstruction using comparative genomics⁸. These mechanisms are
50 based on specific or non-specific chemical reactions that occur either exclusively non-enzymatically
51 within the metabolic network, or in parallel to existing enzyme functions⁹. The second step in the
52 oxidative branch of the pentose-phosphate pathway (oxPPP), the spontaneous hydrolysis of δ-6-
53 phosphogluconolactone (δ-6PGL) into 6-phosphogluconate (6PGNT) is an archetypal example of a
54 reaction that can occur enzymatically, catalysed by 6-phosphogluconolactonase (Pgl, EC 3.1.1.31), and
55 non-enzymatically, without Pgl activity. However, Pgl's main role, rather than maintaining the flux
56 through the PPP, is thought to be preventing the formation of unwanted side-products by covalent
57 modification with highly reactive 6PGLs (δ (1-5) and γ (1-4))^{10–12}. In pioneering studies, Kupor and
58 Fraenkel^{13,14} found that absence of the *pgl* gene indeed slowed the growth of *E. coli* on glucose, but
59 more importantly perhaps, they also discovered an alternative pathway bypassing Pgl, involving
60 dephosphorylation and secretion of gluconolactone, spontaneous abiotic hydrolysis in the medium,
61 and re-import of gluconate and phosphorylation¹⁴. This “Pgl bypass” can therefore provide the same
62 anabolic precursors (NADPH and pentose-phosphates) as the canonical oxPPP.

63 In *E. coli*, one fifth of glucose uptake is directed towards the oxPPP to provide the required anabolic
64 precursors⁶, highlighting the importance of this pathway. In the absence of Pgl, it is thought that this
65 flux is maintained by rapid non-enzymatic hydrolysis of δ-6PGL rather than by the Pgl bypass^{7,11,14,15}.
66 However, 6PGLs have a non-negligible lifetime¹¹, raising the questions whether spontaneous δ-6PGL
67 hydrolysis is fast enough to maintain a high flux through the oxPPP in the absence of Pgl and how the
68 bypass pathway might contribute.

69 In this study, we used a metabolic flux approach to investigate *E. coli* oxPPP function in the absence of
70 Pgl to (i) reinvestigate the extracellular bypass pathway identified by Kupor and Fraenkel using state-
71 of-the-art methods, (ii) quantify its contribution to oxPPP flux, and (iii) elucidate the metabolic function
72 of Pgl. In the absence of Pgl, we found that virtually all carbon flux through the oxPPP was channelled
73 through the Pgl bypass, and that the contribution of non-enzymatic δ-6PGL hydrolysis was negligible.
74 This in turn suggests that the catalytic function of Pgl is required to maintain high flux through the
75 canonical oxPPP, and that the metabolic role of this enzyme is more substantial than simple house-
76 cleaning duties.

77

78 **RESULTS**

79 **Slow growth in *E. coli* Δ pgl is accompanied by gluconate accumulation in the culture medium**

80 We first sought to determine the growth phenotype of an *E. coli* K-12 strain lacking *pgl*, grown on
81 minimal medium with glucose (15 mM) as sole carbon source (Table 1 and supplementary 1 figure 1).
82 In agreement with Kupor and Fraenkel's¹⁴ findings, the Δ pgl strain grew much slower than the WT. The
83 rate of acetate production was markedly increased in the Δ pgl strain, resulting in a higher acetate yield
84 (0.54 ± 0.03 mol/mol) than in the WT strain (0.31 ± 0.02 mol/mol), as observed previously⁷. In addition
85 to acetate and other metabolites previously detected by 1 H NMR¹⁶, significant amounts of gluconate
86 were also detected, but only in the culture medium of the Δ pgl strain (Table 1 and supplementary 1
87 figure 2). Gluconate accumulation represented approximately 2% of the total carbon flux entering the
88 central metabolic pathways of the Δ pgl strain (Table 1). Note that no 6PGL was detected in the culture
89 supernatant of the Δ pgl strain (Supplementary 1 figure 2). Complementation of the Δ pgl strain with
90 the WT gene restored the growth rate determined for the WT *E. coli*, without any gluconate
91 accumulation (Table 1), demonstrating that reduced growth and gluconate excretion are related to the
92 absence of Pgl.

93 **Gluconate is formed by abiotic degradation of gluconolactone**

94 In the extracellular Pgl bypass proposed by Kupor and Fraenkel¹⁴, gluconolactones are hydrolysed into
95 gluconate abiotically in the culture medium. The obvious explanation for the presence of gluconate in
96 the extracellular medium is therefore that gluconolactones are fully converted into gluconate within
97 the time taken to sample the culture medium and store the samples. To prevent abiotic hydrolysis, the
98 culture medium was sampled by filtration, placed immediately on ice and then analysed by NMR, with
99 the first spectrum recorded approximately 5 min after sample withdrawal. Both isomeric forms of
100 gluconolactone (δ and γ) were detected in equal amounts whereas the gluconate concentration was
101 below the detection limit (Supplementary 1 figure 3). 6PGL was not detected either. Importantly, the
102 total amount of gluconolactones was similar to the amount of gluconate detected in the corresponding

103 sample obtained with the initial procedure, indicating that rapidly reducing the sample temperature
104 prevents the spontaneous degradation of gluconolactones. To confirm this effect, we measured the
105 rate constants of gluconolactone spontaneous non-enzymatic hydrolysis in fresh minimal medium
106 containing commercial gluconolactone using real-time $^1\text{H-NMR}$ (Figure 1). The degradation constants
107 obtained by fitting the gluconolactone concentration–time curves assuming a first-order process¹⁷
108 were $0.36 \pm 0.01 \text{ h}^{-1}$ at 7°C and $2.58 \pm 0.02 \text{ h}^{-1}$ at 37°C . At 7°C therefore, the amount of gluconolactone
109 hydrolysed during 5 min of sample preparation is negligible. Using the same approach, the hydrolysis
110 rate of δ -6PGL was found to be $3.97 \pm 0.04 \text{ h}^{-1}$ at 37°C , about 6-7 times higher than at 5°C ¹¹
111 (Supplementary 1 figure 4) and the same order of magnitude as measured for gluconolactone.

112 The detection of gluconate in the culture medium in the above experiments is therefore an
113 experimental artefact arising from the hydrolysis of gluconolactones during the sampling process. The
114 Δpgl strain produces gluconolactone (not gluconate) during growth on glucose.

115 **Glucose-derived gluconate is metabolized by canonical gluconate metabolism**

116 Under growth conditions at 37°C , gluconolactones in the culture medium spontaneously hydrolyse to
117 gluconate. However, gluconate remained below the limit of detection when the culture medium was
118 sampled by filtration and rapidly cooled. We therefore hypothesized that the cells consumed gluconate
119 faster than the rate of production by spontaneous hydrolysis. To confirm this, we blocked gluconate
120 utilization by knocking out the two gluconokinases, GntK and IdnK, known to activate gluconate after
121 uptake^{18,19}. As reported previously²⁰, the $\Delta\text{gntK}\Delta\text{idnK}$ double mutant did not grow on D-gluconate as
122 sole carbon source. As expected therefore, gluconate accumulation was observed when the
123 $\Delta\text{pgl}\Delta\text{gntK}\Delta\text{idnK}$ triple mutant was grown on glucose, with an excretion rate of 1.13 ± 0.04
124 mmol.(g_{CDW}·h)⁻¹. These results are consistent with the Pgl bypass topology described by Kupor and
125 Fraenkel¹⁴ (Figure 2).

126 An unforeseen consequence of gluconokinase deletion was that the $\Delta\text{pgl}\Delta\text{gntK}\Delta\text{idnK}$ triple mutant
127 had a much slower growth rate than the Δpgl strain ($0.29 \pm 0.01 \text{ h}^{-1}$ vs. $0.43 \pm 0.01 \text{ h}^{-1}$). This highlights

128 the importance of metabolic activity in the bypass pathway to sustain growth. To investigate this
129 further, we quantified the metabolic flux through the extracellular bypass using ^{13}C metabolic flux
130 analysis. Since the bypass pathway and the non-enzymatic hydrolysis of δ -6PGL operate in parallel
131 without carbon scrambling, their relative contributions to the oxPPP flux cannot be determined using
132 stationary ^{13}C labelled experiments alone. We therefore investigated the contributions of both routes
133 in the Δpgl strain using stationary and non-stationary metabolic flux analyses.

134 **Deletion of *pgl* barely modifies the contribution of the oxidative branch of the PPP to glucose
135 metabolism**

136 We first quantified the distribution of glucose-6-phosphate between the Embden-Meyerhof-Parnas
137 (EMP) pathway and the oxPPP in the Δpgl strain and the WT strain (as control) grown on minimal M9
138 medium supplemented with [$1-^{13}\text{C}$]-glucose as sole carbon source. Metabolic fluxes were inferred from
139 quantitative measurements of ^{13}C incorporation into alanine *via* pyruvate (Figure 3). In keeping with
140 previous results^{6,21}, $17 \pm 1\%$ of glucose was channelled through the oxPPP in the WT strain. In the Δpgl /
141 strain, the oxPPP accounted for $15 \pm 1\%$ of glucose uptake, a similar contribution as in the WT strain.
142 This suggests that the absence of Pgl does not lead to a rewiring of central metabolic fluxes, in contrast
143 to what has been observed in the absence of glucose-6-phosphate dehydrogenase, the first enzyme in
144 the oxPPP^{6,7}. In the Δpgl strain, the metabolic flux through the oxPPP must therefore be maintained by
145 the Pgl bypass and/or by non-enzymatic intracellular hydrolysis of δ -6PGL.

146 **In the absence of Pgl, oxPPP flux passes exclusively through the Pgl bypass**

147 To quantify the contributions of the extracellular bypass and of intracellular non-enzymatic δ -6PGL
148 hydrolysis to oxPPP flux, we performed a non-stationary carbon labelling experiment wherein ^{13}C
149 labelled glucose was added at the mid-exponential phase to *E. coli* Δpgl growing on unlabelled glucose
150 (Figure 4a and 4c). We used [$2-^{13}\text{C}$]-glucose because the NMR signals corresponding to [$2-^{13}\text{C}$]-glucose
151 and [$2-^{13}\text{C}$]-gluconolactones formed from [$2-^{13}\text{C}$]-glucose, are well resolved in 1D, ^1H NMR spectra,
152 allowing accurate quantification of the total concentrations of unlabelled and labelled gluconolactone

153 (Figure 4b) over time, sampling the culture medium every 30 min after ^{13}C labelled glucose was added.

154 Metabolic fluxes were calculated using a differential equation model of the metabolic network shown

155 in Figure 4d. The flux through the oxPPP and the constant rate of gluconolactone hydrolysis were set

156 to their experimentally determined values (Figure 1d and 3c). The metabolic fluxes were then obtained

157 by fitting them to the time-course concentrations of biomass, (unlabelled and labelled) glucose and

158 (total and ^{13}C -labelled) gluconolactone. Sensitivity analysis confirmed that the system was identifiable

159 based on the available data, meaning that all fluxes were determined with high precision (relative

160 standard deviation below 5 % in each of the three independent biological replicates).

161 The estimated glucose uptake rates and the growth rates were consistent with data obtained from

162 unlabelled experiments (Table 1). The intracellular branch accounts for a negligible proportion of

163 oxPPP flux ($< 1 \pm 1\%$ of glucose uptake), which is fully channelled through the extracellular bypass (99

164 $\pm 1\%$). The bypass carries a flux of $1.07 \pm 0.08 \text{ mmol} \cdot (\text{g}_{\text{CDW}} \cdot \text{h})^{-1}$ which is similar to the flux through the

165 canonical oxPPP in the WT strain and much higher than the gluconolactone accumulation flux in

166 unlabelled experiments ($0.17 \text{ mmol} \cdot (\text{g}_{\text{CDW}} \cdot \text{h})^{-1}$, Table 1). Overall, these results demonstrate the

167 absence of intracellular flux through the canonical oxPPP under these circumstances, and reveal the

168 key contribution of the extracellular Pgl bypass to R5P biosynthesis in the Δpgl strain. These results

169 also explain the severe growth restriction observed when the Pgl bypass is blocked by the deletion of

170 gluconokinases.

171 DISCUSSION

172 To our knowledge this work is the first time the metabolic activity of the Pgl bypass has been

173 quantified, despite its discovery in *E. coli* more than fifty years ago¹⁴. Our NMR results confirm the

174 bypass topology proposed by Kupor and Fraenkel, although the 6PGL dephosphorylation and (P-

175)gluconolactone secretion steps remain to be elucidated. Metabolite phosphatases are generally

176 cytosolic, although some have been shown to be periplasmic, notably those involved in nucleotide

177 salvage²². If 6PGL is secreted in its phosphorylated form, it should be found in the culture medium

178 based on the spontaneous hydrolysis rate estimated in this work. Since 6PGL was not detected, this
179 suggests it is dephosphorylated intracellularly and that it is the resulting gluconolactone that is then
180 secreted. Nevertheless, further studies are required to identify the genes involved in
181 dephosphorylation and secretion of gluconolactones and to unequivocally resolve the first two steps
182 of the Pgl bypass.

183 Because of this bypass, absence of Pgl leads to substantial gluconolactone excretion – about 17% of
184 the total carbon flux entering the cell. This process, which resembles directed overflow mechanism²³,
185 prevents 6PGL intracellular accumulation, avoiding covalent modifications of proteins²⁴ and other
186 nucleophiles¹⁰ in the presence of these highly reactive electrophiles. Gluconolactone excretion may
187 therefore be a cleansing mechanism that avoids the formation of toxic metabolites, the function
188 generally ascribed to Pgl¹². The Pgl bypass may therefore have arisen fortuitously because the unstable
189 gluconolactones excreted into the environment spontaneously hydrolyse into gluconate, one of *E.*
190 *coli*'s preferred carbon sources²⁵.

191 The Pgl bypass produces pentose-phosphates and NAPDH from G6P with the same stoichiometry as
192 the canonical oxPPP. However, this extracellular metabolic route has a significant energetic cost for
193 the cell. First, one extra mole of ATP is required for the phosphorylation of gluconate by gluconokinases
194 (GntK and IdnK). Second, gluconate import is mediated by proton symport systems driven by the
195 proton motive force (with a stoichiometry of one proton per gluconate^{26,27}), thereby consuming
196 protons that could otherwise be used by ATP synthases for ATP production. Assuming a maximal
197 H⁺/ATP ratio of 4²⁸, the total energetic cost of the Pgl bypass is 1.25 ATP equivalents per molecule of
198 glucose entering the oxPPP, corresponding to an extra ATP demand of 1.33 mmol·[g_{CDW}·h]⁻¹. In other
199 words, roughly 20% of the energy used by the cell for glucose uptake and subsequent phosphorylation
200 by the PTS system is wasted in the Pgl bypass. Furthermore, because gluconolactone hydrolysis occurs
201 non-enzymatically outside the cell, it escapes direct metabolic control and the gluconolactone and
202 gluconate are potentially accessible to other cells competing for carbon sources. An illustrative

203 example of this kind of cross-feeding is the recovery of normal growth in $\Delta zwf\Delta pgi$ *E. coli* in the
204 presence of Δpgl mutants, likely because the gluconolactone or gluconate released by the latter feed
205 the oxPPP of the $\Delta zwf\Delta pgi$ cells¹⁵. The downsides of this extracellular bypass are therefore that it is
206 more energetically expensive, partially beyond metabolic regulation and subject to hijack by
207 competing cells.

208 Two oxPPP topologies must therefore be considered in metabolic flux analyses depending on whether
209 *Pgl* is present (the canonical oxPPP) or absent (with *Pgl* bypass). The latter configuration is generally
210 ignored however^{7,29}, particularly in analyses of *E. coli* BL21^{30,31}, which is widely used in biotechnological
211 applications, notably for heterologous protein production³². *E. coli* BL21 and derived strains lack the
212 *pgl* gene and there is evidence of *Pgl* bypass activity in this strain³³.

213 Our results show that metabolic activity in the *Pgl* bypass oxPPP is similar to that of the canonical oxPPP
214 in the WT strain. As observed previously though^{7,14,15}, the Δpgl mutant grows about 30% slower than
215 the WT. The growth rate is reduced even further if gluconate utilisation is also blocked (down to about
216 50% of the WT's growth rate), highlighting the metabolic contribution of the *Pgl* bypass. Finally, of the
217 three enzymes involved in the oxPPP (*Pgl*, G6P dehydrogenase, and 6PGNT dehydrogenase, the latter
218 two being encoded by *zwf* and *gnd* respectively), *Pgl* is the only one whose absence is associated with
219 a noticeable growth defect. In particular, the silent growth phenotype of the Δzwf mutant
220 demonstrates that flux in the oxPPP is not absolutely required for bacterial growth on glucose. Deletion
221 of *zwf* leads to a global reorganization of metabolic fluxes through *E. coli*'s central metabolism to
222 compensate for the absence of flux in the oxPPP while providing the required precursors and energy
223 for anabolism^{6,34,35}. In contrast, *pgl* deletion leads to a local rearrangement of carbon fluxes that fails
224 to support optimal growth. Furthermore, in spite of the secretion mechanism described in this study,
225 absence of *Pgl* must nevertheless lead to intracellular accumulation of δ -6PGL, *Pgl*'s substrate, and γ -
226 6PGL, which forms faster by isomerization of δ -6PGL than δ -6PGL spontaneously hydrolyses to
227 6PGNT¹¹. Indeed, δ -6PGL accumulation has been observed in *E. coli* BL21 growing on glucose³³. The

228 accumulation of these reactive compounds may be detrimental to cells. The growth defect of the Δpgl
229 mutant can therefore be explained by the energy cost of the Pgl bypass and/or intracellular
230 accumulation of toxic intermediates.

231 Pgl is often described as predominantly a house-cleaning enzyme, preventing the formation of
232 undesirable by-products^{9,12}. According to this view, oxPPP flux should mainly be maintained by non-
233 enzymatic hydrolysis of 6PGL into 6PGNT^{7,12,14}. Our results shown on the contrary that the contribution
234 of non-enzymatic δ -6PGL hydrolysis to the canonical oxPPP in the absence of Pgl is negligible, a
235 situation (absence of Pgl) that should promote non-enzymatic hydrolysis of δ -6PGL, since the
236 intracellular concentration of the latter is increased, as mentioned above³³. We can therefore assume
237 that non-enzymatic hydrolysis of δ -6PGL is also negligible when Pgl is present (*i.e.* in the WT strain),
238 meaning that the Pgl's catalytic activity is crucial for 6PG formation. Our results therefore suggest that
239 Pgl's catalytic role is crucial for the efficient functioning of the central metabolic network. While Pgl
240 does also help prevent intracellular accumulation of 6PGLs, our results suggest that this role is
241 mediated by an additional process involving the excretion of these compounds from the cell.

242 In summary, our comprehensive functional analysis of a knock-out mutant in *E. coli*'s central
243 metabolism confirms the existence of a long described compensatory mechanisms and reveals
244 fundamental gaps in our current understanding of an enzyme operating in parallel to a non-enzymatic
245 reaction. Because the reaction catalysed by Pgl can occur spontaneously, the role of Pgl in metabolism
246 has remained unclear. Our results indicate that Pgl's catalysis of 6PGL hydrolysis into 6PGNT is crucial
247 to maintain the required flux through the oxPPP in an energy-efficient manner.

248

249 **METHODS**

250 **Strains and media.**

251 *Escherichia coli* K-12 BW25113 was selected as the experimental model (wild-type strain) for this study.

252 *E. coli* BW25113 Δpgl was constructed from a Δpgl strain in the Keio collection³⁶, with kanamycin

253 cassette removal with a pCP20 plasmid encoding FLP recombinase³⁷. The deletion mutant $\Delta pgl\Delta gnt$

254 was obtained via a one-step disruption protocol³⁸, and the triple deletion mutant $\Delta pgl\Delta gnt\Delta idnK$

255 similarly from the double $\Delta pgl\Delta gnt$ mutant. To confirm the mutations, polymerase chain reaction (PCR)

256 was used to amplify fragments containing the modified sequences. The lengths of the amplified

257 fragments were tested by agarose gel electrophoresis and compared with those of the previous

258 mutant strain. The complemented strain ($\Delta pgl::pgl$) was obtained by transformation of the Δpgl strain

259 by a pZA23::*pgl* plasmid carrying the *pgl* gene under the control of the pLac promoter. The *pgl* gene

260 was amplified from the *E. coli* BW25113 WT strain by PCR and inserted in pZA23 plasmid using the In-

261 Fusion® HD Cloning Plus CE Kit (Takara).

262 All *E. coli* strains were grown in M9-based minimal synthetic medium as described in⁶, complemented

263 with 15 mM glucose. Cultures (50 mL) were performed in triplicate in baffled shake flasks at 37 °C and

264 200 rpm. Growth was monitored by measuring the optical density at 600 nm (OD₆₀₀) using a Genesys

265 6 spectrophotometer (Thermo, USA), and a correlation factor of 0.37 (g_{CDW}·L⁻¹)·OD₆₀₀⁻¹ was used to

266 calculate biomass concentration.

267 **NMR analysis**

268 NMR spectra were recorded on a Bruker Avance III HD 800 MHz spectrometer equipped with a 5-mm

269 quadrupole-resonance QCI-P (H/P-C/N/D) cryogenically cooled probe head. D₂O (10 vol.%) was added

270 to the samples for field/frequency locking and 1 mM TSP-d4 (dissolved in D₂O) was added as an internal

271 standard for frequency calibration and concentration measurements. Spectra were recorded and

272 processed using Bruker TopSpin 3.2. 1D ¹H NMR spectra were acquired using the zgpr30 sequence at

273 280 K with 32 or 64 scans, 64k points, an acquisition time of 2 s, and an recycle delay of 8 s.

274 **Extracellular metabolite concentration measurements and calculation of extracellular fluxes.**

275 Metabolite concentrations (glucose, acetate, gluconate (GNT)) were quantified by 1D¹H NMR from the
276 supernatant obtained by centrifugation (12,000 g, 3 min) of culture broth. The 1D ¹H NMR data were
277 acquired as described above (64 scans). For samples containing gluconolactone (GL), the metabolites
278 (including glucose, gluconate and gluconolactone) were quantified immediately after filtration (0.2
279 µm, polyethersulfone membrane) of 500 µL of culture. The filtrate was kept on ice and rapidly analysed
280 by NMR as described above (32 scans).

281 Extracellular fluxes (*i.e.* glucose uptake, GNT accumulation and growth rates) were determined from
282 the time course concentrations of biomass, substrates, and products using PhysioFit
283 v2.0.4³⁹(<https://github.com/MetaSys-LISBP/PhysioFit>).

284 **Stationary ¹³C- labelling experiments**

285 After preculture on LB, strains were grown in 50 ml M9 minimal synthetic medium complemented with
286 15 mM [1-¹³C] glucose in baffled shake flasks at 37 °C and 200 rpm. At OD 1.2, cells were harvested by
287 centrifugation (5 min at 10,000 g) and the pellet was resuspended in 1.250 ml of Milli-Q H₂O. The
288 suspension (250 µl) was mixed with 250 µl of HCl 12N and hydrolysed at 110 °C for 18 h. HCl was
289 evaporated using a vacuum concentrator, the pellet was washed twice with Milli-Q H₂O and
290 resuspended in 100 µl H₂O. This sample was diluted 100 times for analysis. The carbon isotopologue
291 distribution of alanine was measured in three independent biological replicates, as detailed in⁴⁰.

292 **¹³C-Pulse experiments**

293 *E. coli* BW25113 *Δpgl* was grown in 50 ml M9 minimal synthetic medium complemented with 15 mM
294 glucose in a baffled shake flask at 37 °C and 200 rpm. At OD 1.2, [2-¹³C] glucose was added to the
295 culture medium to obtain approximately 50 % of ¹³C-labelled glucose. Growth was monitored using
296 OD₆₀₀ measurements as described above. Extracellular compounds were quantified directly by NMR as
297 described above.

298 **Kinetics of phosphogluconolactone and gluconolactone spontaneous hydrolysis**

299 *Real time monitoring of gluconolactone (GL) degradation.* δ -Gluconolactone (δ -GL) degradation was
300 monitored by ^1H NMR using a pseudo-2D pulse program (noesyphpr) at 310K. A 3 mM δ -GL (Sigma-
301 Aldrich) solution was prepared in M9-based minimal synthetic medium. Acquisitions were started after
302 rapid homogenisation and temperature stabilisation, and spectra were recorded every 65 s with 8
303 scans each (acquisition time, 3 s; recycle delay, 5 s) for a total of 120 time points. δ -GL degradation at
304 280 K was monitored using the same procedure and a freshly prepared δ -GL solution.

305 *Real time monitoring of 6-phosphogluconolactone (6PGL) degradation.* 6PGL was produced
306 enzymatically from glucose-6-phosphate (G6P) using commercial *L. mesenteroides* NADP $^+$ dependent
307 G6P-dehydrogenase (G6PDH) expressed in *E. coli* (Sigma-Aldrich). The reaction mix consisted of 5 mM
308 G6P, 5 mM NADP $^+$, 12 mM MgCl $_2$, 100 mM phosphate buffer (at pH = 7.2) and 1 mM TSP-d4 6 . ^1H and
309 ^{31}P 1D NMR spectra of the mix were recorded using the zgpr30 sequence with 16 scans and the zg
310 sequence with 64 scans, respectively. One enzyme unit of G6PDH was then added to the NMR tube
311 and the production and hydrolysis of 6PGL were monitored at 310 K using dual reception (^1H and ^{31}P)
312 pseudo-2D spectra (2DDR zggpw5) 41 . Spectra were recorded every 115 s with 64 scans each for a total
313 of 64 time points (^1H acquisition time, 0.7 s; recycle delay; ^{31}P acquisition time 0.6 s, recycle delay, 1
314 s). The concentrations of G6P, 6PGL and 6PGNT were determined from the rows of the pseudo-2D ^{31}P
315 spectra.

316 *Calculation of degradation constants.* The degradation constants of GL and 6PGL were determined by
317 fitting their time-course concentrations assuming a first-order process using COPASI 42 (v4.27), as
318 detailed in 17 .

319 **Relative contributions of glycolysis and the oxPPP**

320 The contributions of the oxPPP and Emben-Meyerhof-Parnas (EMP) pathway to glucose metabolism
321 were estimated by quantifying alanine isotopologues. [1- ^{13}C] glucose metabolised through the EMP
322 pathway forms unlabelled and [1- ^{13}C] pyruvate in equal proportions, while C $_1$ -decarboxylation of

323 glucose through the oxPPP only produces unlabelled pyruvate. The contributions of the oxPPP and the
324 EMP pathway were thus estimated from the fraction of the M1 isotopologue of alanine, using the
325 following algebraic equations²¹:

326
$$Glycolysis = 2 \times Ala_{M1}$$

327
$$PPP = 2 \times (0.5 - Ala_{M1})$$

328

329 **Dynamic ¹³C-flux model of glucose metabolism to quantify the contribution of the Pgl bypass**

330 To quantify the contributions of the canonical oxPPP and of the Pgl bypass to ribose-5-phosphate
331 biosynthesis, we constructed a dynamic ¹³C-flux model of glucose metabolism following the formalism
332 detailed in⁴³. The model contains 21 reactions, 25 species, and 2 compartments (the environment and
333 the cell), and represents five processes: i) growth, ii) glucose uptake and phosphorylation into G6P,
334 utilisation of G6P through iii) the EMP pathway and iv) the intracellular branch of the oxPPP, and v) the
335 extracellular Pgl bypass (Figure 4).

336 The differential equations, which balance the concentrations of extracellular compounds (biomass,
337 glucose, gluconolactone and gluconate) and intracellular compounds (G6P, glyceraldehyde-3-
338 phosphate, 6PGL, 6PGNT and pyruvate), were completed with isotopic equations for parameter
339 estimation. As detailed in ^{43,44}, we considered all reactions (except biomass synthesis) separately for
340 unlabelled and labelled reactants. Fluxes were assumed to be constant over time, in line with the
341 metabolic steady-state assumption, except for the gluconolactone hydrolysis flux which was modelled
342 using the mass action law to represent first-order degradation kinetics¹⁷, in keeping with the abiotic
343 hydrolysis of gluconolactone into gluconate (see Results).

344 The final model has 8 free parameters in total (Supplementary 2). The contribution of the oxPPP to
345 glucose metabolism (i.e. the sum of the flux through the intra- and extracellular branches of the oxPPP)
346 and the gluconolactone (abiotic) hydrolysis rate were fixed at the experimental values determined in
347 this study, as detailed in the Results section. The remaining parameters (p) were estimated by fitting

348 to the experimentally determined concentration dynamics of biomass and unlabelled and labelled
349 glucose and gluconolactone, by minimising the objective function f defined as the weighted sum of
350 squared errors:

$$351 \quad f(p) = \sum_i \left(\frac{x_i - y_i(p)}{\sigma_i} \right)^2$$

352 where x_i is the experimental value of data point i , with an experimental standard deviation σ_i , and $y_i(p)$
353 is the corresponding simulated value. The objective function f was minimised using the particle swarm
354 optimisation algorithm (2,000 iterations with a swarm size of 50). The experimental and fitted data of
355 one biological replicate are shown in Figure 4, and the data for all replicates are provided in the
356 Supplementary 2.

357 The model was constructed and analysed using COPASI⁴² (v4.27) and is provided in SBML and COPASI
358 formats in the Supplementary 2 and at https://github.com/MetaSys-LISBP/GL_GNT_bypass. The
359 model has also been deposited in the Biomodels database (<https://www.ebi.ac.uk/biomodels>)⁴⁵ with
360 the identifier MODEL2310250001 to ensure reproducibility and reusability.

361 **Supplementary Material**

362 Supplementary material 1 : Figure 1, Figure 2, figure 3 et Figure 4

363 Supplementary material 2: Model documentation

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471

472

473 **TABLES**

474 **Table 1: Growth parameters of the different *E. coli* K12 BW25113 strains grown in minimal media**
475 **with 15mM glucose.**

476 The growth of the *E. coli* BW25113 Δpgl strain was compared to the WT strain's and that of the
477 complemented strain, $\Delta pgl::pZA23pgl$, obtained by transformation of the Δpgl strain with pZA23::pgl
478 plasmid and of the $\Delta pgl::pZA23$ strain with an empty plasmid. μ_{\max} , specific growth rate; q_{Glc} , glucose
479 uptake rate; q_{Ac} , net acetate production rate; q_{Gnt} , gluconate accumulation rate. Results are the mean
480 \pm SD of three biologically independent samples.

Parameters	WT	Δpgl	$\Delta pgl::pZA23pgl$	$\Delta pgl::pZA23$
μ_{\max} (h ⁻¹)	0.61 \pm 0.01	0.43 \pm 0.01	0.57 \pm 0.01	0.41 \pm 0.01
q_{Glc} (mmol · [g _{CDW} · h] ⁻¹)	7.77 \pm 0.17	8.52 \pm 0.39	10.11 \pm 0.83	7.60 \pm 0.08
q_{Ac} (mmol · [g _{CDW} · h] ⁻¹)	2.37 \pm 0.15	4.56 \pm 0.16	7.60 \pm 0.31	3.95 \pm 0.16
q_{Gnt} (mmol · [g _{CDW} · h] ⁻¹)	0	0.17 \pm 0.01	0	0.14 \pm 0.02

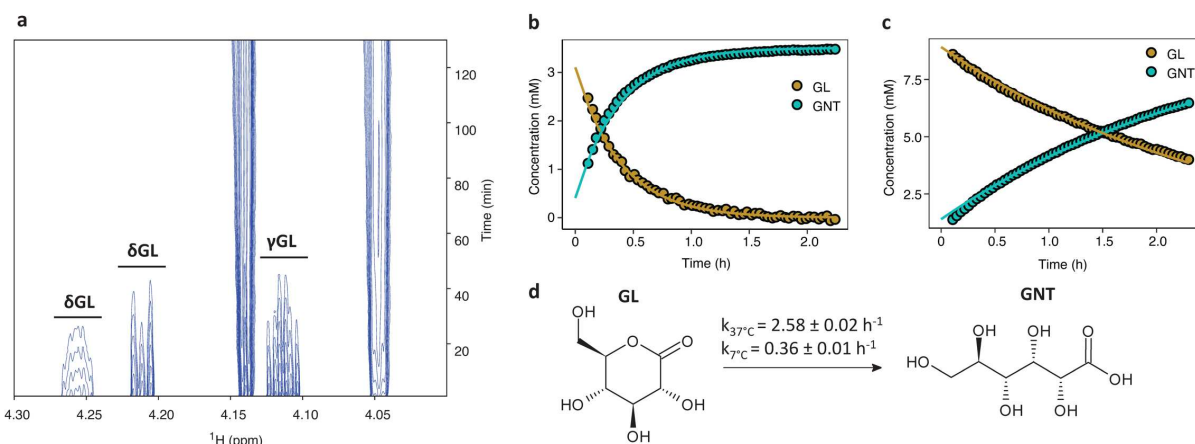
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485 **FIGURES (6 max. in the main text)**



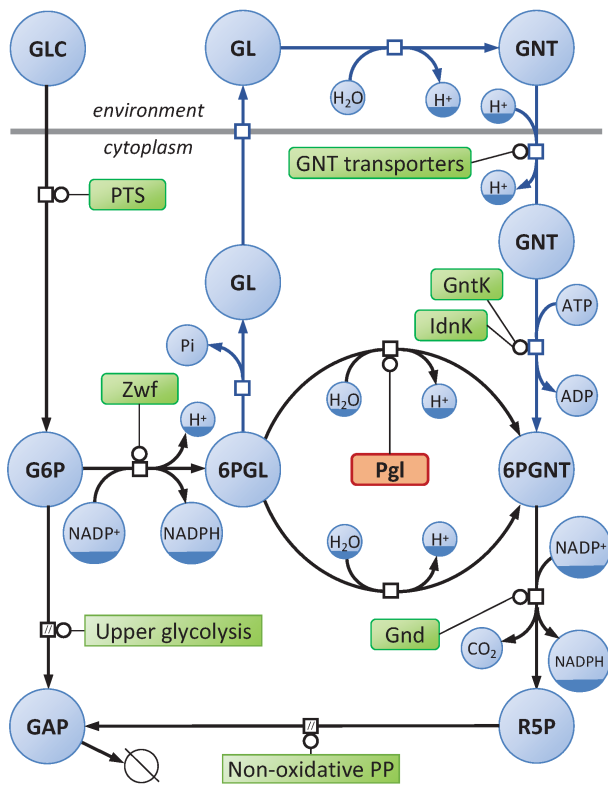
486

487 **Figure 1. Gluconolactone spontaneous hydrolysis.** a. Pseudo-2D ¹H spectra at 37°C, pH7.2 in M9
488 synthetic minimal media of spontaneous gluconolactone hydrolysis. T0 corresponds to the beginning
489 of NMR acquisition (= 6 min 35 s after gluconolactone addition). b,c. GL and GNT concentrations
490 extracted from pseudo-2D spectra (dots) and fitted by COPASI model (lines) at 37 °C (b) and 7 °C (c). d.
491 Reaction scheme for the spontaneous hydrolysis of gluconolactone (GL) into gluconic acid (GNT) with
492 the degradation constant obtained at 37 °C and 7 °C.

493

494

495 **Figure 2:**

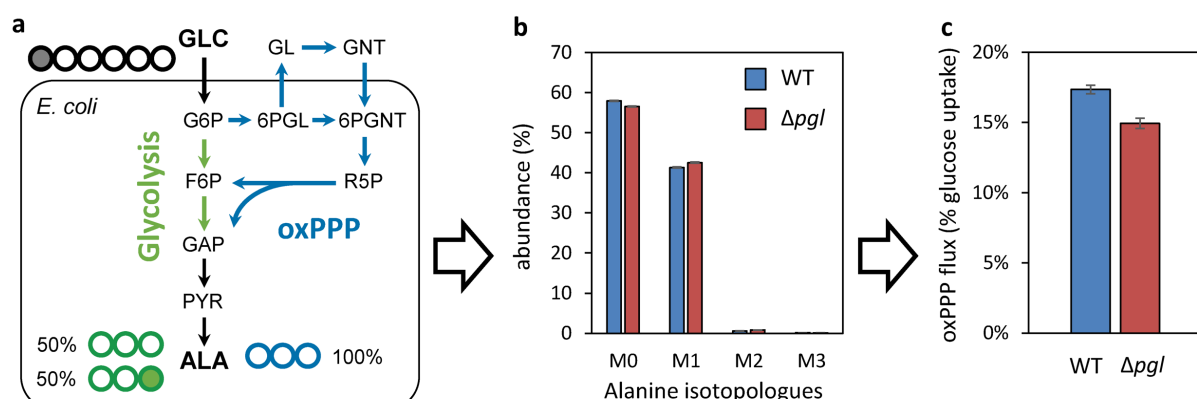


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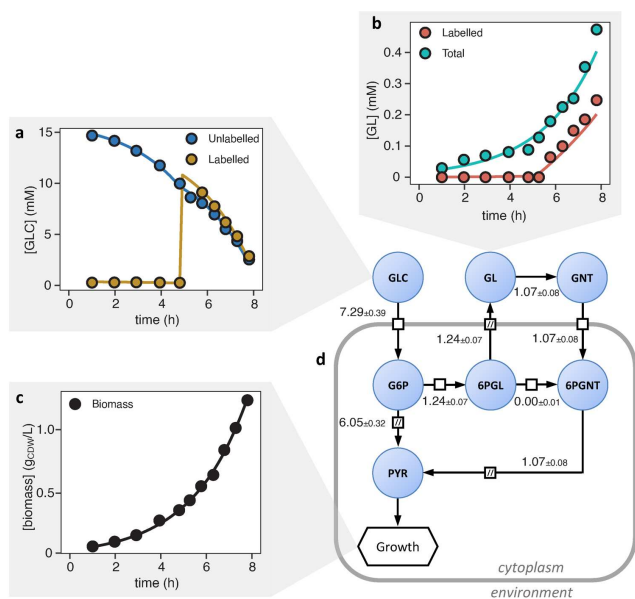
497 **Figure 2. Schematic representation of the Pgl bypass and of the oxidative branch of the pentose-
498 phosphate (PP) pathway in *E. coli*, in Systems Biology Graphical Notation format⁴⁶
499 (<http://sbgn.org.io>). Circles represent metabolites and rounded rectangles represent enzymes. GLC,
500 glucose; G6P, glucose-6-phosphate; GAP, glyceraldehyde-3-phosphate; 6PGL, 6-
501 phosphogluconolactone; GL, gluconolactone; GNT, gluconate; 6PGNT, 6-phosphogluconate; R5P,
502 ribulose-5-phosphate; PTS, Phosphoenolpyruvate:glucose phosphotransferase system; Zwf, glucose-
503 6-phosphate dehydrogenase; GntK, gluconate kinase; IdnK, thermosensitive gluconate kinase; Gnd, 6-
504 phosphogluconate.**

505

506 **Figure 3**



516 **Figure 4 :**



517

518 **Figure 4. Fluxes in the oxPPP of *E. coli* BW25113 Δ pgl quantified by non-stationary ^{13}C -metabolic flux
519 analysis.**

520 **a.** Concentrations of labelled and unlabelled glucose. $[2-^{13}\text{C}]$ -glucose was added at mid-exponential
521 growth (approx. 5 h). **b.** Gluconolactone (labelled and total) concentrations. **c.** Growth of the Δ pgl
522 strain as a function of time. **d.** Metabolic network of the model used to calculate fluxes involved in the
523 Pgl bypass, with flux results (obtained from three independent biological replicates).

524

525