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1	Potential and Limitations of Klebsiella pneumoniae as a Microbial Cell Factory Utilizing
2	Glycerol as the Carbon Source
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18 Abstract

Klebsiella pneumoniae is a Gram-negative facultative anaerobe that metabolizes 19 glycerol efficiently under both aerobic and anaerobic conditions. This microbe is considered 20 21 an outstanding biocatalyst for transforming glycerol into a variety of value-added products. Crude glycerol is a cheap carbon source and can be converted by K. pneumoniae into useful 22 compounds such as lactic acid, 3-hydroxypropionic acid, ethanol, 1,3-propanediol, 2,3-23 butanediol, and succinic acid. This review summarizes glycerol metabolism in K. 24 pneumoniae and its potential as a microbial cell factory for the production of commercially 25 26 important acids and alcohols. Although many challenges remain, K. pneumoniae is a promising workhorse when glycerol is used as the carbon source. 27 Key words: Klebsiella pneumoniae; Glycerol; 3-Hydroxypropionic acid; 1,3-Propanediol; 28

29 2,3-Butanediol; Lactic acid; Ethanol

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30 **1. Introduction**

Crude glycerol is produced in large quantities as a byproduct of the biodiesel industry. 31 Global biodiesel production was 22.7 million tons in 2012 and is forecasted to reach 36.9 32 33 million tons by 2020. As a result, worldwide production of glycerol climbed from 1 million 34 tons in 2000 to 3 million tons in 2011 and is expected to reach 6 million tons by 2025 (Ciriminna et al., 2014; OECD, 2011; Katryniok et al., 2013; Sun et al. 2017). A significant 35 36 amount of glycerol is also generated during bioethanol fermentation by yeast. The liquid fraction of the stillage obtained from ethanol separation contains up to 2% glycerol. Another 37 38 source of glycerol is industrial waste generated from vegetable oils and animal fats. For example, the concentration of glycerol in waste streams from the oleochemical industry is 39 55%-90% (da Silva et al., 2009; Yazdani and Gonzalez, 2007). The increased availability of 40 41 glycerol has caused a substantial reduction in its cost. According to a recent report, the price 42 of crude glycerol is \$0.24/kg and that of pure USP grade glycerol is \$0.9/kg (Kong et al., 2016). Extensive research has been conducted to investigate ways to utilize this surplus crude 43 glycerol. The annual number of research articles addressing the use of glycerol has increased 44 to >7000, doubling in number from the year 2000 to 2007. Numerous chemical conversions 45 to valuable products have been reported (Ciriminna et al., 2014). 46 Glycerol can be used for the microbial production of 1,3-propanediol (PDO), ethanol, 47 48 2,3-butanediol (BDO), butanol, lactic acid, 3-hydroxypropionic acid (3-HP), and succinic

49 acid (Fig. 1) (Clomberg and Gonzalez, 2013; Mattam et al., 2013; Yazdani and Gonzalez,

50 2007). Because of its reduced nature, glycerol generates twice the number of reducing

51 equivalents per carbon than do traditional carbohydrates (glucose, xylose, and sucrose) when

- 52 converted into the glycolytic intermediates phosphoenolpyruvate (PEP) or pyruvate
- 53 (Dharmadi et al., 2006). As a consequence, glycerol gives higher yields of reduced
- 54 metabolites (e.g., succinate, ethanol, PDO, and butanol) than does glucose (Yazdani and

55 Gonzalez, 2007). Table 1 compares the maximum theoretical yield of various metabolites, ATP, and redox balances between glycerol and glucose. 56

Despite its great potential as a carbon source, glycerol is not efficiently metabolized 57 by many microorganisms under anaerobic or oxygen-limited conditions. The ability of K. 58 pneumoniae to assimilate glycerol under oxygen-limited conditions is outstanding and has 59 been studied extensively. Several review articles addressing this issue have also been 60 61 published (Clomberg and Gonzalez, 2013; Mattam et al., 2013; Yazdani and Gonzalez, 2007), but the biotechnological potential and physiological aspects of this strain have not been fully 62 63 elucidated. This review explores and discusses glycerol metabolism in K. pneumoniae and the production of value-added chemicals from glycerol. The potential and challenges of using K. 64 pneumoniae as a microbial cell factory are also discussed. 65

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2. Glycerol metabolism in K. pneumoniae

Many microorganisms can metabolize glycerol in the presence of external electron 67 acceptors (respiratory metabolism), but relatively few under non-respiratory conditions 68 69 (without exogenous electron acceptors). Among the latter, genera including *Citrobacter*, Enterobacter, Bacillus, Propionibacterium, Anaerobiospirillum, Klebsiella, Clostridium, and 70 71 Lactobacillus are well documented (da Silva et al., 2009; Yazdani and Gonzalez, 2007). Some Lactobacillus species including L. reuteri, L. brevis, and L. bucheneri convert glycerol 72 73 to PDO but only in the presence of a major carbon source for growth. Glycerol does not 74 support cell growth, limiting its biotechnological applications. In contrast, *Clostridium sp.* grow well on glycerol as the sole carbon source. However, the use of *Clostridium* is limited 75 by its slow growth and requirement of strict anaerobic conditions. The lack of a convenient 76 77 genetic tool box for large-scale metabolic engineering is another serious drawback of Clostridium (Branduardi et al., 2014; Lütke-Eversloh and Bahl, 2011). 78

From a biotechnological standpoint, Klebsiella sp., especially K. pneumoniae, have 79 many advantages over the aforementioned microbes. They grow on glycerol as the sole 80 carbon source under both aerobic and anaerobic conditions (Arasu et al., 2011; Kumar et al., 81 82 2012). Their growth is fast, typically growing at a rate >0.9/h, even on glycerol minimal medium (Arasu et al., 2011). They naturally produce coenzyme B₁₂, an essential cofactor for 83 the synthesis of the commercially important chemicals PDO and 3-HP (Huang et al., 2013a; 84 85 Oh et al., 2012a). Furthermore, their evolutionary and biochemical proximity to E. coli allows for the application of most of gene manipulation methods developed for E. coli without much 86 87 modification (Celińska, 2012; Kumar et al., 2013a).

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2.1 Overview of glycerol metabolism in K. pneumoniae

In *K. pneumoniae*, glycerol is metabolized oxidatively and/or reductively (Ashok et
al., 2011, Kumar et al., 2013a). The oxidative pathway provides energy and carbon
constituents for the synthesis of cell biomass, while the reductive pathway reduces glycerol
and regenerates NAD⁺, enabling the oxidative assimilation of glycerol under anaerobic
conditions.

Glycerol oxidation proceeds in either a respiratory or a fermentative manner. 94 Respiration requires an exogenous electron acceptor such as oxygen, nitrate, or fumarate 95 (Ashok et al., 2013a,b) and is carried out by enzymes encoded by the glp regulon. During 96 97 respiration, glycerol is first phosphorylated by glycerol kinase (glpK) to yield sn-glycerol-3-98 phosphate using ATP or phosphoenolpyruvate (PEP) as a phosphate donor. Next, snglycerol-3-phosphate is oxidized to dihydroxyacetone phosphate (DHAP) with electron 99 100 transfer to a quinone molecule, which is linked to the electron transport chain (ETC). This 101 transformation is catalyzed by glycerol-3-phosphate dehydrogenase, which is expressed under both aerobic (glpD) and anaerobic (glpABC) conditions. In contrast, fermentation of 102 103 glycerol is conducted without a supply of external electron acceptors, and the reactions are

104 catalyzed by enzymes encoded by the *dha* regulon. Here, glycerol is oxidized to

105 dihydroxyacetone (DHA) by glycerol dehydrogenase (mainly encoded by *dhaD*) using NAD⁺

106 as an electron acceptor. DHA then is phosphorylated by dihydroxyacetone kinase, an enzyme

107 encoded by the ATP-dependent *dhaK1/dhaK* and/or PEP-dependent

108 *dhaK2/dhaKLM/dhaK123*. DHAP obtained from respiration and/or fermentation is channeled

109 into the glycolytic pathway (Fig. 2). A variety of organic acids and alcohols, including acetic

110 acid, lactic acid, succinic acid, ethanol, BDO, and formic acid, are produced by the

downstream glycolytic pathway (Ashok et al., 2011; Kumar et al., 2013a).

The anaerobic growth capability of *K. pneumoniae* using glycerol as a carbon source is attributed to its well-established reductive metabolism. In the reductive pathway, glycerol first undergoes a difficult rearrangement reaction catalyzed by glycerol dehydratase (GDHt) to yield 3-hydroxypropioaldehyde (3-HPA). Catalysis by GDHt requires coenzyme B₁₂, the *de novo* synthesis of which is limited to only few genera of microorganisms. The 3-HPA is

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subsequently reduced to PDO by at least two 1,3-propanediol oxidoreductases (1,3-PDORs),

118 including NADH-dependent DhaT (PDOR) and NADPH-dependent hypothetical

119 oxidoreductase (HOR), along with the regeneration of $NAD(P)^+$ (Fig. 3). *E. coli* and other

120 enterobacter sp. have the oxidative (respiratory) pathway of glycerol assimilation. However,

121 these organisms lack the reductive pathway and cannot grow on glycerol under anaerobic

122 conditions (Dharmadi et al., 2006; Kumar et al., 2013a).

123 2.2 Genes and enzymes involved in fermentative metabolism of glycerol

124 The *dha* regulon, induced by the DHA molecule and expressed in the absence of 125 exogenous electron acceptors, encodes numerous genes needed for both the oxidative

126 (fermentative) and reductive pathways (Celińska, 2012; Forage and Foster, 1982) (Fig. 4A).

127 The *dha* regulon of *K*. *pneumoniae* contains the following genes arranged in the order as

128 appears: Kpk_0615/dhaK, dhaM, dhaL, dhaK, dhaD, dhaR, orfW, CdAT, orfX (dhaG), dhaT,

orfY, dhaB, dhaC, dhaE, orfZ (dhaF), and glpF. Of these, dhaB, dhaC, dhaE, orfX, orfZ, and 129 *dhaT* are relatively well documented for their functions in the reductive metabolism of 130 glycerol. The *orfW* and *orfY* genes, although commonly found in the *dha* regulons of many 131 132 organisms, are not as well described. The proteins encoded by the *dha* regulon of *K*. pneumoniae have high similarity (80%–95%) with those of Citrobacter species but not with 133 those of *Clostridium* species (30%–80%). Surprisingly, the sequence similarity of *dha* 134 135 proteins among *Clostridium* species is very low in some cases, even lower than their similarity to the corresponding proteins in K. pneumoniae or C. freundii (Celińska, 2012; Sun 136 137 et al., 2003; Wei et al., 2014).

138 The complete *dha* regulon is found in only a few dozen species belonging to five

139 different taxonomic groups, including four bacterial species [Actinobacteria, Firmicutes,

140 Fusobacteria, and Proteobacteria (Gammaproteobacteria and Deltaproteobacteria)] and one

141 archaea species, *Halalkalicoccus jeotgali* (Martins-Pinheiro et al., 2016). Interestingly, an

142 incomplete *dha* regulon is present in more than 100 prokaryotes, suggesting that the enzymes

143 of the regulon have other functions not restricted to the anaerobic growth on glycerol. A well-

144 documented example is the one found in *Lactobacillus* species. *Lactobacillus* cannot grow on

145 glycerol as the sole carbon source because it lacks the enzymes for oxidative assimilation of

146 glycerol. However, the enzymes encoded in the incomplete *dha* regulon enable the

147 production of PDO from glycerol, which accelerates cell growth in *Lactobacillus* grown on

148 glucose (Årsköld et al., 2008; Martins-Pinheiro et al., 2016).

In the following sections, the major enzymes involved in fermentative glycerolmetabolism and their genes are described in detail.

151 2.2.1 Glycerol dehydrogenase

152 Encoded by the *dhaD* and/or *gldA* gene, glycerol dehydrogenase is an oxidoreductase
153 that converts glycerol to DHA. The enzyme extracts electrons from glycerol and transfers

them to NAD⁺. DhaD plays a major role during the anaerobic growth of *K. pneumoniae* on
glycerol. Quantification of mRNA expression indicates that *dhaD* is highly induced when
glycerol is used as carbon source. GldA expression is lower than that of DhaD and is thought
to serve as a back up to DhaD during anaerobic growth on glycerol. However, DhaD and
GldA exhibit a high level of similarity in amino acid sequence and have the same substratebinding site (Wang et al., 2014).

160 Recently, DhaD was reported to catalyze the interconversion of acetoin and BDO for NADH disposal and recovery (Wang et al., 2014). The promiscuity of DhaD is responsible 161 162 for BDO production in the $\Delta budC$ mutant of K. pneumoniae. The contribution by DhaD to or its participation in BDO production is also supported by the observed increase in BDO 163 synthesis and increased concentration of NADH in the presence of the *dhaD* gene. NADH is 164 165 generated by the oxidation of glycerol and consumed by the production of BDO from acetoin. 166 The dual physiological functions of DhaD allow K. pneumoniae to control the intracellular redox level. In response to the intracellular level of NADH, DhaD is able to switch its role 167 from NADH production (glycerol oxidation) to NADH consumption (BDO production) 168 (Wang et al., 2014). The diversion of glycerol flux from organic acids toward alcohols such 169 170 as BDO also helps the cells to counteract intracellular acidification (Petrov and Petrova, 2009). Thus, this promiscuity confers different physiological roles to the enzyme, which 171 include regulation of the intracellular NADH/NAD⁺ ratio, prevention of acidification, and 172 173 storage of carbon and energy (Wang et al., 2014).

174 **2.2.2 Dihydroxyacetone kinases**

Dihydroxyacetone kinase (DHAK) phosphorylates DHA to DHAP, an intermediate of
the glycolytic pathway. Sun et al. (2003) analyzed the genome of *K. pneumoniae* MGH
78578 and found, in addition to ATP-dependent DhaK I, a PEP-dependent DhaK II. DhaK I
is encoded by the *dhaK* (*Kpk* 0615) gene, while DhaK II is encoded by three genes, *dhaK*,

179 *dhaL* and *dhaM*, which correspond to the medium, small, and large subunits of the enzyme, respectively (Celińska, 2012; Sun et al., 2003; Wei et al., 2014). The names dhaKLM and 180 181 *dhaK123* are synonyms used without distinction (Martins-Pinheiro et al., 2016; Sun et al., 182 2003; Wei et al., 2014). DhaK enzymes are found in many microorganisms. PEP-dependent DhaK II is present in E. coli, L. lactis, and C. butyricum, whereas ATP-dependent DhaK I is 183 present in Citrobacter freundii (Garcia-Alles et al., 2004; Raynaud et al., 2011). The PEP-184 dependent DhaK II of E. coli, encoded by dhaKLM, is homologous to dhaK123 (dhaKLM) of 185 K. pneumoniae (Gutknecht et al., 2001). In a recent study, Wei et al. (2014) further elucidated 186 187 the physiological roles of DhaK I and DhaK II of the *dha* regulon of *K. pneumoniae*. They found that disruption of ATP-dependent DhaK I had no significant effect on glycerol uptake, 188 indicating that PEP-dependent DhaK II is the major contributor to the conversion of DHA to 189 190 DHAP. As in E. coli, the subunits of DhaK II regulate expression of the dha regulon; dha 191 regulon expression was suppressed by the disruption of *dhaK1* (*dhaK*) and *dhaK2* (*dhaL*). However, in contrast to E. coli, mutation of dhaK3 (dhaM) in K. pneumonia did not 192

upregulate the *dha* regulon but rather decreased cell growth and glycerol uptake.

194 2.2.3 Glycerol dehydratase and reactivating factor

Glycerol dehydratase (GDHt) is the first enzyme in the reductive pathway and 195 196 catalyzes the dehydration of glycerol to 3-HPA. The same reaction can be carried out by diol dehydratase (DDHt), although the latter has a higher activity on 1,2-propanediol than on 197 glycerol. GDHt requires coenzyme B₁₂ or S-adenosyl methionine (SAM) as a cofactor. K. 198 pneumoniae has B₁₂-dependent GDHt only, while strict anaerobes such as clostridia sp. have 199 200 SAM-dependent GDHt. The three subunits of GDHt are encoded by the genes dhaB1 (~1670 bp), dhaB2 (~590 bp) and dhaB3 (~430 bp), for which the gene products are the α (large, 201 60.7 kDa), β (medium, 21.3 kDa), and γ (small, 16.1 kDa) subunits, respectively (Xu et al., 202 2009a). GDHt of K. pneumoniae is known to be present as a dimer of heterotrimers, $(\alpha\beta\gamma)_2$. 203

The subunits of GDHt have high homology among *K. pneumoniae* strains; for example, *K. pneumoniae* XJPD-Li and U30909 strains showed >99% identity.

Coenzyme B_{12} is present in the reaction center and plays an essential role in the 206 207 catalysis of GDHt. The catalytic cycle often results in the mechanism-based inactivation of coenzyme B₁₂ due to the irreversible breakage of the chemical bond between cobalt (Co) and 208 a carbon of the adenosyl moiety. Oxygen accelerates breakage of the Co-C bond. Oxygen is 209 210 also known to inactivate GDHt, even in the absence of the substrate glycerol. The gene products of orfZ/dhaG (small subunit) and orfX/dhaF (large subunit) act as reactivating 211 212 factors of GDHt (GdrAB) (Celińska, 2012; Sun et al., 2003). They reactivate GDHt by catalyzing the exchange of damaged for intact coenzyme B_{12} in the presence of ATP and 213 Mg²⁺/Mn²⁺ (Celińska, 2012; Kumar et al., 2013a; Maervoet et al., 2011; Sun et al., 2003; Wei 214 215 et al., 2014). The reactivase is a molecular chaperone that functions as a heterotetramer 216 containing two elongated α subunits (63 kDa) and two globular β subunits (14 kDa). Structurally, the α subunit resembles both GroEL and Hsp70 chaperones, while the β subunit 217 resembles that of the β subunit of glycerol dehydratase, except that it lacks some of the amino 218 acids responsible for coenzyme B_{12} binding (Liao et al., 2003). 219 In some organisms, the GDHt enzyme varies in terms of the number of subunits and 220 their copies (Fig. 4B) (Liu et al., 2010). For example, in Mesorhizobium loti and 221 222 Mesorhizobium opportunistum, the large and medium subunits (dhaB1 and dhaB2) are fused together and encoded by a single gene. The gene fusion may arise by frameshift mutations. 223 224 The active site of the fused enzyme differs slightly from that of unfused enzyme. Moreover, these microorganisms lack the genes encoding the reactivation factor, indicating that the 225 226 reactivation process may not exist or is carried out by a different route. Mycobacterium smegmatis has two subunits (fused dhaB1 + dhaB2 and dhaB3) that are similar to those of M. 227 *loti* and *M. opportunistum*; however, the reactivating factor differs in that it has one (large) 228

229 subunit. No small subunit has been identified in *M. smegmatis*. Fusobacterium *Ilyobacter polytropus* is unique in that it has genes coding for both B₁₂-independent (SAM-dependent) 230 and B₁₂-dependent GDHt. The genes for B₁₂-independent GDHt are present in the genomic 231 232 DNA, while those encoding B₁₂-dependent GDHt along with its reactivation factor are present in two copies, one in the chromosome and the other in a plasmid. Surprisingly, even 233 with this gene redundancy, no significant production of 3-HPA and/or PDO has been reported 234 235 in *I. polytropus* (Stieb and Schink, 1984). The B₁₂-independent proteins showed no homology with B₁₂-dependent GDHt but significant similarity with the pyruvate formate lyases (PFL) 236

and PFL-activating enzymes and their homologues (Raynaud et al., 2003).

238 2.2.4 1,3-Propanediol oxidoreductase and hypothetical oxidoreductase

The 1,3-propanediol oxidoreductase (PDOR), responsible for the second step of the 239 240 reductive pathway of glycerol, catalyzes the conversion of 3-HPA to PDO with the oxidation 241 of NAD(P)H. 3-HPA is a toxic intermediate, and PDOR protects the cell from deleterious damage by reducing 3-HPA to PDO (Celińska, 2012). In K. pneumoniae, PDOR is encoded 242 by the *dhaT* gene and is strictly NADH dependent. Active PDOR is a homo-octamer with a 243 monomeric molecular mass of 43 kDa. Despite its importance, the kinetic properties of DhaT 244 were elucidated only recently (Lama et al., 2015). DhaT has high specificity for 3-HPA 245 among various aldehydes. Under physiological conditions at pH 7.0 and 37°C, the catalytic 246 247 efficiency (k_{cat}/K_m) of the forward reaction (3-HPA to PDO) was found to be 77.2/s. mM, 248 which is 57-fold higher than that of the reverse reaction (1.3/s. mM; PDO to 3-HPA). The $K_{\rm m}$ (mM) values for 3-HPA and NADH (forward reaction) are 0.77 and 0.03, respectively, while 249 those for PDO and NAD⁺ (backward reaction) are 7.4 and 0.23, respectively. Furthermore, 250 251 the maximum activity of DhaT for the reverse reaction was obtained at pH 9.0 and 55°C. These results strongly suggest that, under physiological conditions, DhaT catalyzes the 252 reduction of 3-HPA to PDO rather than the oxidation of PDO to 3-HPA. Nevertheless, the 253

reverse reaction (PDO→3-HPA) becomes significant as the PDO concentration increases.
This factor is considered a serious drawback of using DhaT for PDO production from
glycerol.

K. pneumoniae has several oxidoreductases in addition to PDOR that can reduce 3-257 HPA to PDO. One such enzyme, designated as hypothetical oxidoreductase (HOR), has been 258 studied extensively and shares 89% identity with YqhD of E. coli (Zhu et al., 2009; Zhuge et 259 al., 2010). Expression of yqhD in wild-type K. pneumoniae is not high, with an mRNA level 260 ~20-fold lower than that of *dhaT*. However, deletion of *dhaT* increases the transcription of 261 262 *yqhD* by more than 10-fold (Ko et al., 2015). This observation suggests that in addition to its wide-ranging activity on short-chain aldehydes, YqhD is a detoxification enzyme for many 263 aldehydes (Jarboe, 2011). YqhD exhibits negligible activity on PDO (backward reaction), 264 265 which is an important advantage over DhaT for use in the production of PDO from glycerol 266 (Celińska, 2012; Chen et al., 2011; Zheng et al., 2006). However, YqhD is kinetically inferior to DhaT because of its low catalytic activity (k_{cat}/K_m) on 3-HPA and the requirement of 267 NADPH as a cofactor. The k_{cat}/K_m value of YqhD on 3-HPA (pH 7.0 and 37°C) is estimated 268 to be 2.1/s. mM, which is 36-fold lower than that of DhaT (unpublished data). The use of 269 270 NADPH as a cofactor prohibits this enzyme from being directly coupled to the reactions catalyzed by DhaD and other enzymes in glycolysis and the TCA cycle, by which NADH is 271 mainly produced. Furthermore, excessive use of NADPH for PDO production can 272 273 significantly reduce the cellular NADPH level and this can disturb many anabolic reactions requiring NADPH (Wang et al., 2003; Jarboe, 2011). NADH and NADPH differ in a 274 phosphate group only, and it has been suggested that electrostatic interaction of this 275 276 phosphate is the major factor distinguishing NADH from NADPH for enzymes using these cofactors. Comparison of the amino acid sequences of DhaT and HOR from K. pneumoniae 277 278 and YqhD from E. coli has revealed that DhaT contains aspartic acid (Asp) at position 41,

while HOR and YqhD have valine (Val) at this position. The steric hindrance and
electrostatic repulsion between Asp in the active site and the phosphate group of NADPH
impede their binding. In contrast, the hydrophobic amino acid Val poses no such obstruction;
thus, YqhD uses NADPH as a cofactor (Ma et al., 2010).

283 Despite their difference in cofactor preference, DhaT and YqhD are similar (44%) in 284 amino acid sequences. *K. pneumoniae* has five enzymes that have at least 40% similarity in 285 amino acid sequence with DhaT and YqhD: putative alcohol dehydrogenase (PaldH), L-1,2-286 propanediol oxidoreductase (FucO), ethanolamine-utilization enzyme (EutG), propanediol-287 utilization propanol dehydrogenase (PduQ), and bifunctional alcohol dehydrogenase (AdhE) 288 (Ko et al., 2015). The roles of these enzymes in PDO production have been investigated.

289 2.2.5 Regulatory protein

290 The *dhaR* gene product is an important transcription factor that is responsible for 291 expression of the *dhaT* and *dhaB* genes (Sun et al., 2003). This gene is found in K. pneumoniae, C. freundii, and Clostridium botulinum but not in every organism containing a 292 293 complete *dha* regulon (Martins-Pinheiro et al., 2016). The presence of DhaR is thought to allow the *dha* operon to function as a separate regulatory system, independent of fumarate 294 295 nitrate reduction (FNR). Under anaerobic conditions, FNR acquires DNA-binding properties and induces the expression of many anaerobiosis-related genes. DhaR contains domains for 296 GAF (52-199), PAS (203-267), σ^{54} factor interaction, and histidine HTH-8. The PAS senses 297 internal levels of energy charge, light, oxygen, and redox potential. The HTH-8 domain has a 298 helix-turn-helix conformation and acts as a DNA binding structure. The σ^{54} factor interaction 299 domain interacts with the σ^{54} factor of RNA polymerase and activates RNA transcription 300 from the σ^{54} promoters (Buck et al., 2000; Taylor and Zhulin, 1999). Thus, the DhaR protein 301 triggers the transcription of *dha* genes in response to intracellular levels of several important 302 physiological parameters (Celińska, 2012; Sun et al., 2003). Zheng et al. (2006) investigated 303

the effect of overexpression of the putative regulatory gene dhaR in the dha regulon on 304 glycerol metabolism in K. pneumoniae. dhaR overexpression increased PDOR activity up to 305 6.7-fold over that of the control wild-type strain, confirming the role of DhaR as a positive 306 regulator of the *dhaT* gene. However, *dhaR* overexpression did not increase the activity of 307 glycerol dehydratase (DhaB). Furthermore, despite the significant increase in PDOR activity, 308 PDO production was unexpectedly low compared with that of the wild type (440 vs. 656.23 309 310 mM, respectively). More studies are required to determine the precise role of DhaR and its regulatory function in the *dha* regulon. 311

312 DhaR in E. coli (70% identity to that of K. pneumoniae) has also been studied. This protein stimulated transcription of the *dhaKLM* operon from a σ^{70} promoter. Interestingly, in 313 E. coli, DhaL was a positive regulator of the dhaKLM operon while phosphorylated DhaM 314 and DhaK were negative regulators of the same operon. DhaK contains the DHA-binding site. 315 DhaL uses ADP as a cofactor for the double displacement of phosphate from DhaM to DHA. 316 DHA is the oxidation product of glycerol metabolism; binding of DHA to DhaK reduces the 317 affinity of DhaK for DhaR. In the presence of DHA, DhaL::ATP is dephosphorylated, 318 leading to the displacement of DhaK by DhaL::ADP, which stimulates DhaR activity. In the 319 absence of DHA, DhaL::ADP is phosphorylated back to DhaL::ATP by the phototransferase 320 system (PTS), thereby inhibiting the binding of the phosphorylated complex (DhaL::ATP) to 321 DhaR. This phosphorylation is mediated by DhaM, providing a phospho-histidine relay 322 323 between the PTS and DhaL::ADP. This double-check mechanism of binding and turnover increases the selectivity such that the binding of nonphosphorylated compounds does not 324 induce the *dha* operon (Bächler et al., 2005). 325

326 **2.3 Genes and enzymes involved in respiratory assimilation of glycerol**

The *glp* regulon comprises the genes encoding proteins required for the conversion of glycerol to DHAP via sn-glycerol-3-phosphate. The five operons that constitute the *glp*

regulon are *glpFK*, *glpD* (anaerobic glycerol-3-phosphate dehydrogenase), *glp ACB* (aerobic 329 glycerol-3-phosphate dehydrogenase), *glpTQ* (glycerol-phosphate 330 permease/glycerophosphodiesterase), and *glpEGR*. The *glpF* gene encodes a cytoplasmic 331 332 membrane protein that facilitates the diffusion of glycerol into the cell. The *glpE* gene encodes an acidic, cytoplasmic protein of 108 amino acids with a molecular weight of 333 12,082. The glpG gene encodes a basic, cytoplasmic membrane-associated protein of 276 334 335 amino acids with a molecular weight of 31,278 (Lin, 1976; Yang and Larson, 1998; Zeng et al., 1996). The biochemical functions of GlpE and GlpG are unknown. The *glpR* gene 336 337 encodes the GlpR repressor protein. These five operons are located at three different positions on the chromosome. Transcription of these operons is subject to multiple controls, including 338 catabolite repression mediated by cAMP-CRP and respiratory control mediated by the FNR 339 340 and ArcA/ArcB systems. Moreover, each of the operons is negatively controlled by a 341 repressor specific for the regulon (Lin, 1987; Zeng et al., 1996).

342 2.3.1 GlpR and other regulatory proteins

The GlpR repressor protein, predicted to contain 252 amino acids with a molecular 343 weight of 28,048, belongs to the DeoR bacterial regulatory factor super family. Although not 344 studied extensively, GlpR of K. pneumonia is expected to have the same biochemical 345 properties and regulatory functions as that of E. coli. GlpR negatively controls all the glp 346 347 operons by binding to operators that overlap or are close to the *glp* promoters. In case of *E*. 348 *coli*, glycerol-3-phosphate acts as an inducer and has a high affinity for GlpR ($K_d = 31 \mu M$) (Lin, 1987; Weissenborn et al., 1992; Zeng et al., 1996). When complexed with glycerol-3-349 350 phosphate, GlpR cannot bind to the operator sites, and repression is relieved. Glycerol-3-351 phosphate is a product of GlpK and a substrate of GlpD/GlpABC; thus, the absence of GlpD causes a substantial increase in the level of glycerol-3-phosphate, resulting in induction of the 352 353 glp regulon (Fig. 2). The GlpR protein exhibits differential binding affinity for the operators

354 of corresponding operons, as follows: glpFK > glpD > glpTQ > glpACB. The glpFK promoter appears to be the strongest among the *glp* promoters. Therefore, the maximum strength of the 355 *glpFK* promoter combined with the high affinity of GlpK for glycerol ($K_m = 10 \mu M$) should 356 357 result in sufficient accumulation of the inducer, glycerol-3-phosphate, in the presence of glycerol to relieve repression by GlpR. The relatively lower strength of the *glpD* promoter 358 along with the low affinity of the GlpD enzyme for glycerol-3-phosphate ($K_m = 1 \text{ mM}$) 359 prevents excessive degradation of glycerol-3-phosphate that is needed for phospholipid 360 synthesis when the exogenous source of glycerol-3-phosphate becomes limiting, until after 361 362 accumulation of the inducer (Lin, 1987; Weissenborn et al., 1992). Recently, Jung et al. (2014) investigated the role of GlpR in the aerobic production of 3-HP from glycerol in E. 363 coli. They found that elimination of this regulatory factor caused increased assimilation of 364 365 glycerol and higher production of 3-HP and suggested that the deletion of glpR led to the 366 upregulation of genes involved in glycerol transport and assimilation. K. pneumoniae has several other regulatory genes in addition to *glpR*: the global regulatory genes *hdeB*, *hdeD*, 367 and *yfdX*, with unknown functions and *kvgS* and *kvgA*, encoding two proteins comprising a 368 two-component signal transduction system (Sun et al., 2003). 369

370

2.3.2 Glycerol uptake/transport facilitator and other components

The glycerol facilitator GlpF is an integral membrane protein that forms aqueous 371 372 pores. The GlpF pores selectively allow passive transport of glycerol and other molecules 373 such as DHA and urea across the cytoplasmic membrane, but not water or ions (Heller et al., 1980; Reizer et al., 1993). The GlpF protein contains six membrane-spanning helices that are 374 unique among prokaryotic transport proteins. These transmembrane helices are arranged in 375 376 two bundles. The GlpF protein may function as a homodimer with the two six-member domains arranged in the membrane to form a channel (Weissenborn et al., 1992). K. 377 pneumoniae has two glycerol transporter genes, one next to orfZ/dhaF in the fermentative 378

379 dha regulon (see Fig. 4A) and the other in the respiratory glpFK operon. Both glpF genes have a high similarity (85 %) to each other. When one but not both of the glpF genes is 380 disrupted, the glpF deletion mutants of K. pneumoniae can consume glycerol and convert it to 381 382 PDO at a high rate, similar to the wild type (unpublished data; Supplementary Fig. S1). This observation indicates that both *glpF* genes encode active glycerol transporter proteins. 383 However, their presence is not essential for glycerol utilization in K. pneumoniae because the 384 385 double-deletion mutant for both facilitators can still uptake glycerol and convert it to PDO. In E. coli, only one glpF gene is present in the glpFK operon; the GlpF protein shows a high 386 387 similarity (80.9%) to that (encoded from *glpFK*) of *K. pneumoniae* (Sun et al., 2003). The importance of GlpF in E. coli also seems to be limited only at low levels of glycerol (<8 388 mM), as glycerol itself may pass through the membrane by diffusion at high concentrations 389 390 (Richey and Lin, 1972).

2.3.3 Simultaneous operation of fermentative and respiratory pathways

The presence of the *dha* and *glp* regulons affords *K*. *pneumoniae* metabolic flexibility. 392 393 The *dha* regulon enables fermentative utilization of glycerol, while the *glp* regulon enables respiratory metabolism of glycerol. During fermentative metabolism of glycerol, redox 394 395 constraints force the production of PDO from glycerol, with a low ratio of carbon conversion yield to cell biomass. Moreover, ATP is formed through substrate-level phosphorylation, and 396 397 NADH oxidation occurs through the formation of reduced metabolites. In the presence of 398 oxygen, the redox constraints forcing the production of PDO are relieved; therefore, the carbon loss required for regeneration of NAD⁺ is greatly reduced. As a consequence, glycerol 399 utilization, biomass synthesis, and even the formation of many metabolites increase (Durnin 400 401 et al., 2008). Interestingly, the fermentative route (dha regulon) is in operation under a wide range of oxygen levels, from microaerobic to highly aerobic conditions, as evidenced by 402 403 PDO formation. For example, a high titer PDO (102.7 g/L) was produced under the condition

of 2.0 vvm aeration and even at 3.5 vvm, although the production was a little lowered in the
latter case (Oh et al., 2012a). These results suggest that the fermentative and respiratory
pathways might have evolved to complement each other for optimal control of cell growth
under a wide range of aeration conditions except the strictly anaerobic one.

408

2.4 Carbon catabolite repression

Carbon catabolite repression (CCR) refers to the suppression of the use of less-409 preferred carbon sources, when a more preferred carbon, most prominently glucose, is 410 present. Cyclic AMP, cAMP receptor protein (CRP), and the enzyme EIIA^{Glc}, an 411 412 intermediate in the phosphorylation cascade of the PTS, are key players in the CCR of enterobacteria. The EIIA^{Glc}, a cytosolic protein, exists in phosphorylated and 413 414 unphosphorylated forms. The phosphorylated form stimulates membrane-bound adenylate 415 cyclase, which catalyzes the formation of cAMP; the cAMP-CRP complex then induces expression of the catabolic genes of less-preferred carbon sources. The unphosphorylated 416 form of EIIA^{Glc} is also responsible for inhibiting the transport of less-preferred carbon 417 418 sources by inhibitory binding to permeases, causing so-called inducer exclusion (Eppler and Boos, 1999; Eppler et al., 2002). Glycerol assimilation in K. pneumoniae is greatly inhibited 419 by CCR in the presence of glucose. This repression is related to not only low cAMP and 420 cAMP-CRP but also inhibition of GlpK (which functions mainly under aerobic conditions) 421 by fructose 1,6-bisphosphate, a metabolite produced from glucose. The unphosphorylated 422 EIIA^{Glc} is also known to allosterically inhibit GlpK (Holtman et al., 2001). Anaerobic 423 glycerol metabolism mediated by the *dha* operons is also suppressed by the presence of 424 glucose. Enzyme-level inhibition similar to that of the aerobic GlpK has not been reported, 425 426 but we have observed that, in K. pneumoniae, transcription of the dha operons is greatly reduced in the presence of glucose (Suman et al., 2017). 427

428 Interestingly, glycerol exerts catabolite repression on the assimilation of maltose in E. *coli*. To exert catabolite repression, glycerol must be phosphorylated to glycerol-3-phosphate 429 (Hogema et al., 1998; Eppler et al. 2002). However, unlike other carbon sources, further 430 431 metabolism of glycerol is not required. According to Eppler and Boos (1999), glycerol-3phosphate reduces expression of MalT, a positive activator of all mal genes. The key players 432 in this repression are adenylate cyclase, EIIA^{Glc}, and CRP. The growth on maltose of the 433 mutants lacking EIIA^{Glc} or containing truncated adenylate cyclase was no longer repressed by 434 glycerol, and CRP-independent transcription of *malT* was also not influenced by glycerol 435 either. In addition, stimulation of adenylate cyclase by phosphorylated EIIA^{Glc} was controlled 436 by glycerol-3-phosphate. The mutation in adenylate cyclase relieved the repression caused by 437 glycerol-3-phosphate (Eppler et al., 2002). Although not investigated, we assume that the 438 439 glycerol repression on maltose catabolism also hold true in K. pneumoniae.

Glycerol also exerts catabolite repression on the assimilation of citrate in *K*. *pneumoniae*. Citrate can be used as the sole carbon and energy source during anaerobic
growth of *K. pneumoniae*, but the presence of glycerol suppresses the expression of the
citrate fermentation genes. A reduced concentration of the cAMP–CRP complex has been
postulated to be the cause of this repression (Meyer et al., 2001). Further research is needed
to test the hypothesis.

3. *K. pneumoniae* as a microbial cell factory for the production of commodity chemicals
During glycerol fermentation, *K. pneumoniae* produces many metabolites, including
PDO, BDO, and lactic acid. This section describes the biotechnological production of bulk
chemicals from glycerol in *K. pneumoniae*. 3-HP is not a natural product during the growth
of *K. pneumoniae* on glycerol. However, its production is possible using a minor
modification of the PDO synthetic pathway and is included here. The biochemical pathways

used for the synthesis of these chemicals are explained, and the metabolic engineering used toimprove their production is discussed.

454 **3.1 1,3-Propanediol**

PDO, also known as trimethylene glycol, is a promising platform chemical that has
two hydroxyl groups. With terephthalic acid, it can be co-polymerized to the novel copolymer polytrimethylene terephthalate (PTT). PDO also has applications in the food and
cosmetic industries. Currently, commercial PDO is mainly produced by recombinant *E. coli*(Maervoet et al., 2011; Saxena et al., 2009). The global demand for PDO was 60.2 kilotons in
2012 and is predicted to reach approximately 150 kilotons by 2019. The market for PDO is
growing rapidly, from \$157 million in 2012 with projections up to \$560 million in 2019

462 (MarketsANDMarkets, 2012; Lee et al., 2015).

463 K. pneumoniae is one of the best native producers of PDO from glycerol. GDHt and PDOR are two important enzymes for the conversion of glycerol to PDO (Fig. 3). 464 Physiologically, the synthesis of PDO from glycerol is essential for the anaerobic growth of 465 K. pneumoniae, as described previously. The production of PDO requires two cofactors, 466 coenzyme B₁₂ and NAD(P)H. K. pneumoniae has a de novo pathway for the biosynthesis of 467 coenzyme B₁₂ that comprises more than 20 genes. NAD(P)H is generated through the 468 oxidative metabolism of glycerol, during the production of cell biomass and/or oxidized 469 470 (pyruvate and acetate) or partially reduced (BDO, lactic acid) metabolites. The production 471 yield of PDO from glycerol varies depending on the cost of carbon for NAD(P)H production; the lower the cost, the higher the yield of PDO from glycerol. NADPH can also be the 472 electron donor for the conversion of 3-HPA to PDO when YqhD is present. However, the 473 474 contribution of NADPH to PDO production in native K. pneumoniae seems to be limited because the PP pathway, which is the main source of NADPH, is not active when glycerol is 475

476	the sole carbon source. Furthermore, K. pneumoniae has very low transhydrogenase activity
477	for the conversion of NADH to NADPH (Zhang and Xiu, 2009; Chen et al., 2011).
478	Several metabolic engineering strategies have been employed to improve PDO
479	production by K. pneumonia (Table 2). Amplification of the reductive pathway by
480	overexpressing DhaB, PDOR, or both has been attempted by many research groups. As
481	indicated previously, 3-HPA is highly toxic, and its accumulation is detrimental to cell
482	viability and toxic to PDO production. To reduce 3-HPA accumulation, overexpression of
483	dhaT has been attempted. In one study, 3-HPA accumulation was reduced by dhaT
484	overexpression, but no improvement in PDO production resulted (Hao et al., 2008).
485	Overexpression of both DhaT and DhaD increased PDO production (56.3%) in batch culture
486	but no increase in fed-batch culture (Chen et al., 2009). With another K. pneumoniae strain,
487	DSM 2026, the same experiments were repeated but also with no increase in PDO production
488	(Zheng et al., 2006). These results suggest that the reductive pathway in K. pneumoniae is
489	evolutionarily well established (perhaps in concert with the oxidative pathway) and that PDO
490	production is not limited by expression of the <i>dhaB</i> and <i>dhaT</i> genes of the reductive pathway.
491	However, in one study where the <i>E. coli yqhD</i> gene was highly overexpressed to give ~10-
492	fold higher PDOR activity (110 IU/mg total protein), a significant increase in PDO titer (25%;
493	67.6 g/L) and yield (from 0.53 to 0.62 mol/mol), along with less 3-HPA accumulation, was
494	achieved (Zhu et al., 2009). This observation suggests that in the recombinant strain, NADPH
495	can play an important role as a cofactor and that PDO production can be increased by
496	modifying the reductive pathway. It is also probable that the effect of DhaB and PDOR on
497	PDO production is highly dependent on the strain and/or culture conditions adopted.
498	Lactic acid, BDO, ethanol, and acetic acid are the major byproducts of PDO synthesis
499	in <i>K. pneumoniae</i> . Inactivation of the metabolic pathways that give rise to these products has
500	resulted in increased PDO synthesis. In particular, deletion of the <i>ldhA</i> gene encoding lactate

501 dehydrogenase has been highly beneficial. For example, by disrupting the *ldhA* gene, Xu et al. (2009b) significantly increased PDO production without lactic acid production. They 502 observed increases in the PDO titer (95.4–102.1 g/L), conversion yield (0.48–0.52 mol/mol), 503 504 and productivity (1.98–2.13 g/L h; Table 2). Similarly, Oh et al. (2012a) achieved the high PDO titer of 102.7 g/L by deleting ldhA in their own K. pneumoniae isolate. When lactic acid 505 production was eliminated, more ethanol and succinic acid were generated from the pyruvate 506 507 node. Reduction of these compounds has also been attempted by deleting *adhE* (encoding alcohol dehydrogenase) and *frdA* (encoding fumarate reductase). Although the production of 508 509 ethanol and succinate was substantially reduced in the triple mutant ($\Delta ldhA\Delta adhE\Delta frdA$), PDO production was only marginally increased compared with that of the single $\Delta ldhA$ 510 mutant. The failure of the triple mutant ($\Delta ldhA\Delta adhE\Delta frdA$) to increase PDO production was 511 512 mainly due to an increase in BDO production. Disruption of the BDO pathway encoded by the *bud* operon (*budA*, *budB*, and *budC*) has also been studied (Fig. 5). Although partially 513 effective, individual inactivation of each of the three genes did not successfully decrease 514 BDO production (Oh et al., 2012a; Wu et al., 2013). Deletion of the entire bud operon 515 completely eliminated BDO synthesis (Kumar et al., 2016), but seriously hampered cell 516 growth and glycerol consumption. Consequently, no increase in PDO production was resulted. 517 Wu et al. (2013) attempted to decrease BDO production while increasing PDO production by 518 519 inserting the *fdh* gene from *Candida boidinii* (NADH-forming formate dehydrogenase) into 520 the budC locus. This protocol increased the PDO titer (62.3-72.2 g/L) and yield (0.47-0.57 mol/mol) and decreased the production of BDO by 52.2% and formic acid by 73.4%. 521 Acetic acid is one of the most toxic metabolites, accumulating in large quantities 522 523 during glycerol fermentation by K. pneumonaie (Celińska, 2012). The toxic effects of acetate is realized even at the low concentration of 20 mM. Physiologically, acetate is a preferred 524 metabolite for many Enterobacter sp. because its production is accompanied by ATP 525

generation. In a typical bioreactor run for PDO production by K. pneumoniae, the 526 accumulation of acetic acid to >300 mM by the end is not uncommon, which often 527 completely stops glycerol fermentation. Acetate is produced by Pta-Ack and PoxB when 528 529 pyruvate formation is faster than its consumption, which is so-called 'overflow metabolism' (Fig. 6) (De Mey et al., 2007). In K. pneumoniae, the contribution of the poxB gene is not 530 significant and its deletion does not much affect acetate production. In comparison, deletion 531 532 of the *pta-ack* genes greatly reduces cell growth and results in high accumulation of pyruvate and pyruvate-based metabolites (Ko et al., 2017). Extensive studies to reduce or eliminate 533 534 acetate production have been conducted in E. coli. Compared to poxB, ackA and/or pta were much more significant in reducing acetate production. However, deletion of the latter genes 535 was not beneficial because the glycolytic flux and cell yield were seriously reduced and the 536 537 accumulation of other byproducts such as pyruvate, lactate, and formate was greatly increased (Causey et al., 2004; De Mey et al., 2007). Several indirect approaches to 538 decreasing pyruvate/acetate accumulation were investigated: stimulation of the anaplerotic 539 pathway by overexpression of pyruvate carboxylase, PEP carboxylase, or PEP carboxykinase 540 (Abdel-Hamid et al., 2001; Causey et al., 2004; Chao and Liao, 1993; Sanchez et al., 2005); 541 overexpression of acetyl-CoA synthetase (ACS); increase of the TCA cycle throughput by 542 deleting the transcriptional repressor arcA gene; and stimulation of the glyoxylate shunt by 543 544 deleting the transcriptional repressor *iclR* gene (De Mey et al., 2007; Jeong et al., 2004; Lin et 545 al., 2006). Moreover, the use of a mutant pyruvate dehydrogenase which is less sensitive to inhibition by NADH has also been examined (Kim et al., 2008). In E. coli, these approaches 546 proved to be effective to varying extents under properly selected culture conditions. However, 547 548 they have not yet been fully investigated in K. pneumoniae.

To eliminate byproduct formation, reduction of glycerol flux through the oxidativepathway at the glycerol node has also been attempted. The deletion of glycerol

551 dehydrogenase (dhaD) and/or dihydroxyacetone kinase (dhaK) resulted in decrease of the oxidative flux and increase of the conversion yield of glycerol to PDO. In addition, 552 production of the byproducts lactate, ethanol, and succinate but not acetate decreased 553 554 significantly, even without blocking the pathways leading to these byproducts (Seo et al. 2009; Horng et al. 2010). However, cell growth and glycerol consumption rate were also 555 reduced. Interestingly, in these mutant strains, deletion of *dhaT* has almost no effect on PDO 556 production. In the absence of *dhaT*, expression of an HOR, highly homologous to YqhD, was 557 upregulated (see section 2.2.4). 558

559 The theoretical maximum yields of PDO on glycerol are 0.875 and 0.844 mol/mol under aerobic and anaerobic conditions, respectively (Celińska, 2012). However, 560 experimentally, the yields are reported between 0.35 and 0.65 mol/mol (Saxena et al., 2009; 561 562 Sun et al., 2008). The low yield is mainly attributed to the accumulation of unnecessary byproducts and inefficient metabolism for generating reducing equivalents (NADH) and ATP. 563 If acetyl-CoA is fully oxidized through the TCA cycle (without byproducts formation) and 564 the activity of the electron transport chain (ETC) oxidizing NADH is properly controlled, a 565 significant increase in PDO yield is expected. Attaining this goal requires extensive pathway 566 engineering of cellular metabolism. In bioreactor operation, the aeration rate should be 567 optimized. Oxygen regulates gene expression and the enzymatic reactions of glycerol 568 metabolism and determines the diversion of DHA to the PP pathway. Regeneration of NADH 569 570 by the TCA cycle and its oxidation by ETC are also controlled by oxygen levels.

571 **3.2 3-Hydroxypropionic acid**

As with PDO, 3-HP is an important platform chemical and has been selected by the US Department of Energy (DOE) as one of the top 12 chemicals obtainable from biomass. 3-HP is a C3 bifunctional molecule and a structural isomer of lactic acid (Werpy and Petersen, 2004). 3-HP can be converted into a variety of useful chemicals such as acrylic acid and acryl

576	amide (Pina et al., 2011; Kumar et al., 2013a). 3-HP can be synthesized by chemical routes
577	from acrylic acid, 3-propiolactone, 3-hydroxypropionitrile, allyl alcohol, vinyl acetate, and
578	PDO. However, none of the chemical processes is commercially feasible at present due to the
579	high cost of the starting materials, toxicity of intermediates, and/or the environmental
580	incompatibility of the processes (Jiang et al., 2009; Pina et al., 2011; Kumar et al., 2013a).
581	Many microorganisms, including both prokaryotes and eukaryotes, can naturally synthesize
582	3-HP as either an intermediate or end product through a range of metabolic pathways.
583	However, the production of 3-HP by these native microorganisms is too low to be
584	commercially meaningful (Jiang et al., 2009; Kumar et al., 2013a).
585	In studies pursuing commercial production of 3-HP, glycerol is first converted to 3-
586	HPA by GDHt. 3-HPA then is converted to 3-HP via the coenzyme A (CoA)-dependent or
587	CoA-independent pathway (Fig. 3). In the CoA-independent pathway, 3-HPA is oxidized to
588	3-HP by aldehyde dehydrogenase (ALDH); in the CoA-dependent pathway, 3-HPA is
589	converted to 3-HP via 3-HP-CoA and 3-HP-phosphate. In K. pneumoniae, both the CoA-
590	dependent and CoA-independent pathways are present. However, the titer of 3-HP produced
591	by the CoA-dependent pathway is low as <5.0 g/L (Luo et al., 2011 and 2012). High
592	production of 3-HP requires the disruption of oxidoreductases (e.g., DhaT, YqhD) and the
593	overexpression of a proper aldehyde dehydrogenase (ALDH). Ashok et al. (2013a) developed
594	a recombinant <i>K. pneumoniae</i> in which <i>dhaT</i> and <i>yqhD</i> were deleted, and the homologous
595	<i>puuC</i> gene encoding the NAD ⁺ -dependent γ -glutamyl- γ -aminobutyraldehyde dehydrogenase
596	was overexpressed. The recombinant K. pneumoniae produced 3.8 g/L 3-HP in 12 h of flask
597	culture, but only under appropriate aeration (i.e., microaerobic) conditions. Under anaerobic
598	conditions, PDO (instead of 3-HP) was obtained as the main product, even though two
599	oxidoreductases were disrupted and <i>puuC</i> was highly overexpressed. In contrast, highly
600	aerobic conditions produced a high cell mass without much accumulation of either 3-HP or

601	1,3-PDO. In a glycerol-fed batch bioreactor experiment under a constant dissolved oxygen
602	(DO) concentration of 5% (considered "proper" microaerobic condition), the recombinant K .
603	<i>pneumoniae</i> $\Delta dhaT \Delta yqhD$ overexpressing both PuuC and DhaB produced >28 g/L 3-HP in
604	48 h, with a yield of >40%. In contrast, the same strain produced only 10.5 g/L 3-HP when
605	cultivated under the higher 10% DO conditions. Several important conclusions regarding 3-
606	HP production by K. pneumoniae were made, as follows. First, in addition to DhaT and
607	YqhD, K. pneumoniae has more unidentified oxidoreductases that can produce PDO from 3-
608	HPA. Second, to produce 3-HP rather than PDO, proper aeration to regenerate NAD^+ is
609	essential. Third, excessively high aeration decreases the expression of the Dha regulon and
610	synthesis of coenzyme B_{12} , an essential cofactor for GDHt catalysis.
611	To eliminate PDO production even under limited aeration conditions, Ko et al.
612	attempted to identify and disrupt other potential PDORs from K. pneumoniae (Ko et al.,
613	2015). A mutant strain devoid of <i>dhaT</i> , <i>yqhD</i> , <i>ahpF</i> , and <i>adhE</i> genes was developed;
614	however, the mutant neither eliminated 1,3-PDO production nor increased 3-HP production.
615	Again, this result suggests that K. pneumoniae has more unidentified oxidoreductases, and
616	thus, the complete elimination of 1,3-PDO production during 3-HP production is highly
617	challenging. To address the important problem, of NAD^+ regeneration and coenzyme B_{12}
618	production, Ashok et al. (2013b) studied anaerobic respiration using nitrate as an external
619	electron acceptor. They attempted to regenerate NAD^+ from NADH by nitrate reduction
620	while maintain anaerobic conditions to synthesize and stabilize the oxygen-sensitive
621	coenzyme B_{12} . Disruption of the <i>glpK</i> gene (encoding for glycerol kinase) was also necessary
622	because with $glpK$ intact, the rate of anaerobic respiration was too fast, and most of the
623	glycerol was metabolized through GlpK to produce biomass, as in the case of highly aerobic
624	cultivation. The anaerobic glycerol fermentation in the presence of nitrate was successful:
625	1.74 g/L 3-HP was obtained from a 12-h flask culture and 22 g/L 3-HP was produced in a 48-

626 h fed-batch bioreactor culture (Ashok et al., 2013b) (Table 3). However, disadvantages were noted, including the requirement for large amounts of nitrate and the toxic effect of nitrite 627 generated as a reduced intermediate of nitrate. In a recent study, Li et al. (2016) optimized a 628 629 promoter for ALDH expression, culture conditions, and metabolic flux to achieve high-level production of 3-HP in their K. pneumoniae isolate. One recombinant strain, K. pneumoniae 630 (pTAC-puuC) expressing puuC under the IPTG-inducible tac promoter, produced 73.4 g/L 3-631 632 HP in a bioreactor with a molar yield of glycerol of 0.52 and productivity of 1.53 g/L h. Further, elimination of the *ldh1*, *ldh2*, and *pta* genes elevated the titer and molar yield to 83.8 633 634 g/L 3-HP and 0.54 mol/mol, respectively, with decreased cell growth and productivity. This titer of 3-HP is the highest produced by K. pneumoniae to date. 635

As another approach to addressing the problem associated with NAD⁺ regeneration 636 637 and coenzyme B₁₂ synthesis in 3-HP production, co-production of 3-HP and PDO has been investigated. The simultaneous production of 3-HP and PDO can eliminate the dependency 638 on oxidative metabolism of glycerol (and electron transport chain), as the cofactor required 639 for 3-HP production is regenerated by PDO production or vice versa. Furthermore, the 640 problems associated with the expression of genes for the assimilation of vitamin B_{12} and 641 glycerol can be alleviated substantially because co-production can be performed under 642 anaerobic or microaerobic conditions (Kumar et al., 2012, 2013b). Ashok et al. (2011) 643 developed a recombinant strain of K. pneumoniae DSMZ by overexpressing ALDH and 644 645 deleting *dhaT*. With *dhaT* intact, PDOR activity was too high compared with that of ALDH, even with overexpression of the latter enzyme from a multicopy plasmid. The recombinant K. 646 pneumoniae DSMZ accumulated 16.0 g/L 3-HP and 16.8 g/L PDO in 24 h, and the 647 648 cumulative yield of these two metabolites on glycerol was 51% (Table 3). Huang et al. (2012) studied the same co-production using their K. pneumoniae isolate overexpressing ALDH and 649 obtained 24.4 3-HP and 49.9 g/L PDO, respectively, after 24 h under anaerobic conditions. 650

651 The cumulative molar yield of the two metabolites reached 0.61 (0.18 for 3-HP and 0.43 for PDO). In a following study, they showed that the maximum titer of 48.9 g/L 3-HP (along 652 with 25.3 g/L PDO) was achieved at 1.5 vvm, while the highest concentration of 38.4 g/L 653 654 PDO (along with 16.6 g/L 3-HP) was obtained at 0.1 vvm (Huang et al., 2013a). During 3-HP production from glycerol by K. pneumoniae, the accumulation of such 655 byproducts as lactic acid, ethanol, acetic acid, formic acid, and 2,3-BDO is a serious 656 657 consideration, as in the case of 1,3-PDO production (Ashok et al., 2013a,b; Li et al., 2016). Deletion of *ldhA* reduces lactic acid formation and increases the 3-HP yield. However, 658 659 deletion of other genes such as *adhE*, *pfl*, *budBAC*, and *pta-ack* had little desired effect and seriously hampered cell growth and glycerol assimilation (unpublished data). Even with the 660 co-production of 3-HP and 1,3-PDO, where oxidative glycerol assimilation was greatly 661 662 reduced, accumulation of these byproducts was a serious drawback (Ko et al., 2017). To prevent the formation of lactic acid along with other byproducts, Kumar et al. (2013b) 663 employed resting cells of recombinant K. pneumoniae J2B overexpressing ALDH and devoid 664 of lactate dehydrogenase (*ldhA*). For this strain, final titers of 3-HP and PDO of 22.7 and 23.5 665 g/L, respectively, were obtained without lactic acid accumulation. The cumulative product 666 yield increased to 0.77. 667 In addition to K. pneumoniae, E. coli has been extensively studied for 3-HP 668

669 production from glycerol. Several successful results have been reported by Samsung Ltd. Co.

670 Korea. One study (Chu et al., 2015) used an engineered strain (*E. coli* W3110 ΔackA-pta

671 ΔyqhD_dhaB_mutant gabD4) harboring an active ALDH mutant (designated as GabD4) from

672 *Cupriavidus necator*, producing 71.9 g/L 3-HP with a productivity of 1.8 g/L h. However,

673 unlike *K. pneumoniae*, *E. coli* does not synthesize vitamin B₁₂ naturally, and it was necessary

to add the coenzyme externally. Furthermore, a significant amount of glucose for cell growth

675 was required because for *E. coli*, glycerol is a less-preferred carbon source than glucose.

676 Consequently, the researchers at Samsung initially grew cells to a high density (40-300 OD_{600}) on glucose and then produced 3-HP by feeding glycerol (Chu et al., 2015; Kim et al., 677 2014a). In comparison, K. pneumoniae can produce a high concentration of 3-HP growing 678 purely on glycerol at a much lower cell concentration (OD₆₀₀, <20) (Huang et al., 2013a,b; Li 679 et al., 2016). Together, the use of glucose for cell proliferation, exogenous addition of 680 expensive cofactor vitamin B_{12} , and low cumulative yield of 3-HP (on glucose plus glycerol) 681 682 increase the production cost. Thus, K. pneumoniae can be considered a better biocatalyst than E. coli. However, for the commercial production of 3-HP, many other factors are important, 683 684 including biosafety, process stability, and downstream processing. Further studies to assess the potential of these strains as hosts for 3-HP production are needed. 685

686 **3.3 2,3-Butanediol**

BDO has many applications in the pharmaceutical, biomedical, and other chemical industries for the production of printing inks, perfumes, fumigants, spandex, moistening and softening agents, and plasticizers (e.g., cellulose nitrate, polyvinyl chloride, polyacrylates) (Celińska and Grajek, 2009; Ji et al., 2011). BDO can also be used as an antifreezing agent and octane booster for petrol as is or can be converted to useful derivatives such as 1,3butadiene, and diacetyl and methyl ethyl ketone.

Unlike 3-HP and PDO, BDO is a product of oxidative metabolism. Of the three 693 694 stereoisomers of BDO, K. pneumoniae mainly produces the meso form (Ji et al., 2011). BDO 695 is synthesized from pyruvate via α -acetolactate and acetoin. The C5 intermediate, α acetolactate, is formed from two molecules of pyruvate by self-condensation catalyzed by α -696 acetolactate synthase (ALS; *budB*). α -Acetolactate is decarboxylated to acetoin by α -697 698 acetolactate decarboxylase (budA), and acetoin is reduced to BDO by 2,3-butanediol dehydrogenase/acetoin reductase (budC) using NADH as a reductant. In the presence of 699 700 oxygen, α -acetolactate is spontaneously decarboxylated to diacetyl, further reduced to acetoin

701 by action of diacetyl reductase, and then reduced to BDO. α -Acetolactate is also the precursor 702 of branched-chain amino acids (valine, leucine, and isoleucine); thus, elimination of the bud operon can arrest K. pneumoniae growth when cultured on glycerol minimal medium (Kumar 703 704 et al., 2016). In the anaerobic process, one NADH molecule is required for the conversion of two pyruvates to one BDO, while the aerobic process requires two NADH (Fig. 5). The 705 706 enzyme DhaD may also contribute to BDO formation (see Section 2.2.1). In the BDO pathway itself, the conversion of two pyruvates to BDO regulates or is regulated by the 707 intracellular NADH/NAD⁺ ratio, similar to other fermentative pathways. Another important 708 709 role of the BDO pathway is to reduce intracellular acidification by converting acids to the neutral metabolite, BDO (Celińska and Grajek, 2009; Ji et al., 2011). 710 711 Although BDO is one of the main byproducts of glycerol fermentation by *K*. 712 pneumoniae (Petrov and Petrova, 2009; Durgapal et al. 2014), its production from glycerol 713 has not been studied extensively. In few studies using glycerol as the carbon source, Petrov and Petrova (2009) focused on the effects of pH and aeration with their K. pneumoniae 714 715 isolate G31. BDO production was greatest at alkaline initial pH when tested at a pH range of 5–8. Without pH control a sharp pH drop occurred initially due to the production of organic 716 717 acids, and this pH drop triggered the synthesis of BDO, which is considered the only noninhibitory compound produced by the oxidative metabolism of glycerol. Once neutral pH is 718 719 restored with the rising concentration of BDO, catabolism shifted toward the production of 720 ATP-generating acetic acid. The amount of BDO generated in each cycle (pH drop and neutralization) was dependent on the extent of the pH drop. In one study, a BDO 721 concentration of 49.8 g/L was reached when the initial pH was set at 8.0 and no pH control 722 was given during the rest of fermentation. PDO was the major byproduct (11.6 g/L) (Table 4) 723 and its production decreased by exclusion of Co^{2+} , an essential component of coenzyme B_{12} . 724 They also found that intensive aeration led to a significant increase in BDO production; an 725

726	increase in aeration rate from 1.1 to 2.2 vvm enhanced the yield from 0.38 to 0.46 mol/mol
727	and productivity from 0.24 to 0.36 g/L h. BDO production was also increased by forced pH
728	fluctuations of the culture medium with discrete ΔpH values (1.0, 2.0, and 3.0) at
729	predetermined time intervals (6, 12, and 24 h). The highest BDO concentration (70.0 g/L)
730	was produced by increasing the pH by one unit ($\Delta pH = 1.0$) every 12 h (Petrov and Petrova,
731	2010). The mechanism behind the effect of this peculiar pH change on BDO production has
732	not been elucidated. Although BDO is readily produced from glycerol by K. pneumoniae, the
733	volumetric productivity is low due to the long fermentation time (generally >150 h). Huang et
734	al. (2013b) isolated two Klebsiella strains that produce BDO plus acetoin with a total yield of
735	0.44–0.45 (g/g). One isolate utilized 64.5 g/L glycerol in 78 h and produced 24.9 g/L BDO,
736	3.9 g/L acetoin, and 3.2 g/L ethanol. Again, the volumetric productivity was not high.
737	Metabolic engineering efforts with K. pneumoniae to improve BDO production from
738	glycerol are limited. Using glucose as carbon source, Kim et al. (2014b) reported that
739	overexpression of the BDO pathway (budA, budB) in the ldhA deficient K. pneumoniae strain
740	KCTC2242 improved BDO production and resulted in 90 g/L BDO, with a productivity of
741	2.75 g/L. h. Because both glucose and glycerol are converted through the common
742	intermediate pyruvate, BDO production from glycerol is also expected to increase by
743	amplification of the bud operon. Further work with glycerol as carbon source is required to
744	prove this effect. Recently, co-production of BDO and PDO from glycerol has also been
745	attempted (Park et al., 2017). In glycerol fermentation, both alcohols always appear
746	simultaneously and co-production seems to be beneficial for cells to maintain redox balance
747	during anaerobic growth. Park et al. (2017) found that by using the K. pneumoniae mutant
748	deficient of <i>ldhA</i> and <i>mdh</i> , 50.1 g/L BDO was produced along with 63.8 g/L PDO in 75 h.
749	The ratio of the two alcohols varied according to aeration; with increasing aeration, the ratio
750	of [BDO]/[PDO] increased as observed by Petrov and Petrova (2010). In the 5,000 L pilot-

scale fed-batch fermentation using crude glycerol, they could produce 114 g/L diols (70 g/L

PDO and 44 g/L BDO) with the yield of 0.60 diols/glycerol (g/g) and a productivity of 2.2

g/L. h of diols. If separation is not an issue, co-production of the two diols can be a goodoption.

755 **3.4 Lactic acid**

Lactic acid has applications in the food, chemical, cosmetic, and pharmaceutical industries. Lactic acid is a monomer that can be polymerized to yield the biodegradable plastics polylactic acid (PLA) and poly(3-hydroxybutyrate-*co*-lactate). Two optical isomers of lactic acid, D- and L-lactic acid, are produced in racemic mixtures depending on the chemical route used. Microbial fermentation can yield optically pure isomers. Currently, nearly all commercial lactic acid comes from microbial fermentation (Abdel-Rahman et al., 2013; Abdel-Rahman and Sonomoto, 2016).

763 Because of the availability of highly efficient lactic-acid-producing microbes derived from E. coli, lactic acid bacteria, and even acid-tolerant yeasts (Abdel-Rahman et al., 2013), 764 765 K pneumoniae has received little attention as a lactic acid producer. However, as previously described, lactic acid is a major byproduct of glycerol fermentation by K. pneumoniae, and its 766 potential as a producer is high. This bacterium contains metabolic pathways for the 767 production of both L- and D-isomers (Fig. 7). They are synthesized through the pyruvate 768 769 and/or methylglyoxal pathways in K. pneumoniae (Ashok et al., 2011). When produced from 770 pyruvate, lactate dehydrogenase (*ldhA*) reduces pyruvate using NADH as reductant. In the methylglyoxal route, DHAP is converted to methylglyoxal, which can be transformed to both 771 D- and L-lactic acid. For the production of L-lactic acid, methylglyoxal is reduced to L-772 773 lactaldehyde, which is subsequently oxidized to L-lactic acid. Using glutathione, D-lactic acid is obtained through simultaneous aldehyde group oxidation and keto group reduction of 774 775 methylglyoxal (Booth et al., 2003; Booth, 2005). The methylglyoxal route has several

776 disadvantages. Methylglyoxal is highly toxic, and its accumulation can severely impair other 777 metabolisms and even lead to cell death. Second, the route is energetically inefficient, with an ATP yield of -1. The L- or D-lactic acid product can be converted back into pyruvate by 778 779 respiratory lactate dehydrogenase (Mazumdar et al., 2010, 2013). Several studies showing the potential of K. pneumoniae as a lactic acid producer are 780 available (Table 4). For example, K. pneumoniae DSMZ is reported to accumulate 22.7 g/L 781 lactic acid in 24 h during co-production of 3-HP and PDO from glycerol. The titer of lactic 782 acid was more than that of any of the targeted products, and the yield of lactic acid on 783 784 glycerol alone was 0.31 (Ashok et al., 2011). Similarly, fed-batch cultivation of K. 785 pneumoniae BLh-1 under oxygen-limited conditions gave lactic acid as the main product, 786 outcompeting PDO; 59 g/L lactic acid was produced in 40 h when crude glycerol from 787 biodiesel synthesis without any purification was used (Rossi et al., 2013). Durgapal et al. 788 (2014) also found that K. pneumoniae J2B and DSMZ strains produced lactic acid at 37.0 and 47.8 g/L, respectively (Table 4); in both strains, lactic acid levels surpassed those of the target 789 790 product PDO. Recently, Feng et al. (2014) engineered K. pneumoniae ATCC25955 by overexpressing *ldhA* and deleting *dhaT* and *yqhD* (recall that these two genes are mainly 791 792 responsible for PDO production). They found that the recombinant produced 142.1 g/L of optically pure D-lactic acid from glycerol in fed-batch cultivation under microaerobic 793 794 conditions (2.5 vvm and 400 rpm). This result is the highest lactic acid concentration 795 produced from glycerol reported to date. E. coli has also been engineered for the production of D- and L-lactate from glycerol (Mazumdar et al., 2010, 2013). For diverting glycerol flux 796 toward L-lactic acid in E. coli, the chromosomal copy of D-lactate dehydrogenase was 797 798 replaced with Streptococcus bovis L-lactate dehydrogenase (Mazumdar et al., 2013). Moreover, gene encoding *lldD* (respiratory L-lactate dehydrogenase) was inactivated to 799 800 prevent the consumption of L-lactic acid, and the methylglyoxal route ($\Delta mgsA$) was blocked

801 to prevent production of toxic methylglyoxal and formation of racemic mixtures of D- and Llactic acid (Fig. 7). Furthermore, the respiratory route for glycerol assimilation was 802 overexpressed (glpK and glpD) to improve the ATP yield of the metabolic pathway through 803 804 transfer of electrons from glycerol-3-phosphate to oxygen (i.e., oxidative phosphorylation). 805 The engineered E. coli strain ($\Delta pflB \Delta pta \Delta adhE \Delta frdA \Delta mgsA \Delta lldD \Delta ldhA:: ldhA_glpK-$ _glpD) produced 50 g/L of L-lactate from 56 g/L of crude glycerol, at a yield 93% of the 806 theoretical maximum and with high optical (99.9%) and chemical (97%) purity (Mazumdar et 807 al., 2013). The approaches proven successful in *E. coli* should be applicable to *K*. 808 809 pneumoniae. These studies, along with the results by Feng et al. (2014), suggest that K. pneumoniae is a good host for the production of D- and L-isomers of lactic acid, with high 810 optical purity, from glycerol. 811

812 **3.5 Ethanol and other metabolites**

K. pneumoniae has also been studied for the production of ethanol, succinic acid, 2-813 ketogluconic acid, catechol, *cis, cis*-muconic acid and 2-butanol although its potential for such 814 production has not been fully explored (Cheng et al., 2013; Jung et al., 2015; Oh et al., 2014; 815 Wang et al. 2015; Wei et al., 2013). Of these chemicals, ethanol production has been 816 relatively well studied. Ethanol is a renewable energy source and is widely used as a fuel 817 additive for partial gasoline replacement. Currently, commercial ethanol is produced from the 818 819 fermentation of sugar and starch. However, glycerol is one of the cheapest substrates for 820 ethanol production, estimated as nearly 40% cheaper than sugar-based production (Yazdani 821 and Gonzalez, 2007). This factor has encouraged the development of microbial strains for fermentative conversion of glycerol to ethanol. The metabolic route by which ethanol is 822 823 synthesized from glycerol is shown in Fig. 6. Oh et al. (2011, 2012b) constructed a mutant 824 strain of K. pneumoniae (termed GEM167) through γ -irradiation. PDO synthesis in this mutant strain was lower than that of wild type (0.2 vs. 7.93 g/L, respectively), while ethanol 825

826 accumulation was higher (8.6 vs. 1.1 g/L, respectively). The mechanism underlying these changes remains unclear. Introduction of the pyruvate decarboxylase (pdc) and aldehyde 827 dehydrogenase (*adhII*) genes into mutant strains lacking lactate dehydrogenase (*ldhA*) 828 829 resulted in the production of 31.0 g/L (673.9 mM) ethanol, with a yield of 0.89 (mol/mol) and productivity of 1.2 g/L h (Oh et al., 2012b), the highest level of ethanol production from 830 glycerol reported to date. More extensive studies including estimation of the maximum 831 832 achievable titer and microbial tolerance to high-concentration ethanol, both of which are key factors for commercialization, are required to assess the potential to produce ethanol from 833 834 glycerol by the use of K. pneumoniae.

4. Challenges to bioconversion of glycerol using *K. pneumoniae* as a biocatalyst

The design of microbial cell factories is gaining unprecedented momentum as 836 837 metabolic engineering is progressively aided by advances in synthetic biology and multi-838 omics analyses. Several bio-based chemicals have already been launched successfully in the marketplace, and others are in the pipeline. For a bulk chemical to be considered for 839 commercial production, the production process should achieve a product concentration of 100 840 g/L, carbon yield of 0.50 (g/g), and volumetric productivity of 2.0 g/L/h. The aforementioned 841 products produced by K. pneumoniae from glycerol are no exception. Challenges to the use 842 of K. pneumoniae as a microbial cell factory using glycerol as a carbon source are described. 843

844 **4.1 Pathogenicity**

The pathogenicity of *K. pneumoniae* is one of the major obstacles to its commercial application. *K. pneumoniae* is a Bio-safety Level 2 microorganism and causes nosocomial and urinary tract infections. Moreover, capsulated *K. pneumoniae* cells are difficult to separate from culture broth, complicating the downstream processing. The biotechnological application of this bacterium will require attenuation of its pathogenicity and reduction of biosafety concerns. *K. pneumoniae* produces a number of virulence factors that contribute to

its pathogenicity, including lipopolysaccharide (LPS), capsular antigens, fimbrial adhesins,
siderophores, and O antigens (Albertí et al., 1993; Fang et al., 2004; Podschun and Ullmann
1998). The mechanism by which this bacterium causes disease is still obscure because most
studies have been conducted on a limited number of virulence factors (El Fertas-Aissani et al.,
2013).

In the last decade, efforts have been made to overcome the pathogenicity of K. 856 857 pneumoniae (Shrivastav et al., 2013), suggesting hope for the generation of nonpathogenic K. pneumoniae. The two major pathogenic determinants in K. pneumoniae are the capsular 858 859 polysaccharides (CPS), which cover the entire bacterial surface, and LPS, expressed on the outer bacterial membrane. LPS consists of lipid A, a core oligosaccharide, and an O-antigenic 860 polysaccharide. In a highly virulent strain of K. pneumoniae, mutations in genes involved in 861 862 LPS biosynthesis (*waaC*, *waaF*, and *wabG*) resulted in a decrease in bacterial colonization and virulence (Izquierdo et al., 2003; Jung et al., 2013). Mutant strains devoid of the outer 863 core LPS were also found to be deficient in the "cell-attached" capsular polysaccharides 864 covering the bacterial surface. These mutations abolished the highly virulent characteristics 865 of pathogenic K. pneumoniae when tested in different animal models. Furthermore, these 866 mutants were more sensitive to several hydrophobic compounds than were wild-type strains. 867 Reintroduction of the waaC, waaF, and wabG genes into K. pneumoniae rescued the 868 869 pathogenic properties, confirming the role of these genes in pathogenicity.

Another important study was conducted by Lin et al. (2012), in which eight highly conserved residues of the MagA protein (G308, G310, G334, G337, R290, P305, H323, and N324) encoded by the mucoviscosity-associated gene A (*magA*) of *K. pneumoniae* were subjected to site-directed mutagenesis. The *magA* gene contributes to the biosynthesis of K1 capsular polysaccharide (CPS), which correlates with pathogenic phenotypes including mucoviscosity, serum and phagocytosis resistance, and virulence. Alanine substitutions at
R290A or H323A abolished MagA function, with annihilation of CPS production, serum
resistance, and virulence. Some *K. pneumoniae* strains showing less pathogenicity were also
isolated. For example, *K. pneumoniae* strain J2B produced less LPS and demonstrated a high
sensitivity to many antibiotics and lower pathogenicity. Moreover, its biomass was readily
separated from fermentation broth by centrifugation (Arasu et al., 2011). Future studies
should include further engineering of these less pathogenic strains to make them completely
nonpathogenic, followed by careful evaluation of their toxicity.

883 **4.2 Complex glycerol metabolism**

884 Although glycerol metabolism has been extensively studied, the mechanism and regulation of its gene expression in K. pneumoniae have not been fully elucidated. The 885 diversity of glycerol metabolism should be a great advantage for K. pneumoniae survival in 886 887 different environments and for its biotechnological utilization as well. However, a high level of complexity poses difficulty in the engineering of a strain to suit our purposes. Metabolic 888 complexity is likely responsible in part for the fact that, despite serious attempts, the molar 889 890 yield of many important products such as PDO and 3-HP is still far below the theoretical maximum. Efforts based on our current knowledge, such as the overexpression of DhaB, 891 892 DhaT, YqhD and AldH and inactivation/deletion of GlpK, DhaD, DhaK and DhaKLM, were not satisfactory (Maervoet et al. 2011; Kumar et al. 2013a; Kumar et al., 2016). Another 893 894 challenge is to understand how the ratio of glycerol distribution between respiratory and 895 fermentative routes is controlled. Because the $K_{\rm M}$ of glycerol kinase toward glycerol is low, it has been speculated that a major fraction of glycerol flows through the respiratory route when 896 oxygen is present. However, some experimental results do not support this hypothesis. For 897 898 example, deletion of *glpK* does not decrease the glycerol flow through the oxidative pathway in the presence of oxygen (Ashok et al. 2013a). In fact, wild-type and mutant ($\Delta g l p K$) strains 899 900 of K. pneumoniae demonstrate similar cell growth, glycerol consumption, and PDO

901 production profiles under (micro)aerobic conditions. These observations suggest the presence902 and/or involvement of other kinases.

Another challenging issue for the use of K. pneumoniae is its inefficient TCA cycle 903 904 (Cabelli, 1955). Several genes, including isocitrate dehydrognease (*icd*), fumarase (*fumA*), and malate dehydrogenase (mdh), in K. pneumoniae are significantly less transcribed than 905 those in E. coli (unpublished data). Furthermore, the activity of isocitrate dehydrogenase was 906 907 more than sevenfold lower in K. pneumoniae than in E. coli. If the TCA cycle is inefficient, achieving a high cell density, which is essential for improving the productivity of target 908 909 metabolites, is difficult. Moreover, achieving high-yield generation of NADH with little consumption of glycerol carbon is difficult. Furthermore, the overflow metabolism that 910 911 produces highly toxic acetate becomes more serious. These problems have been well 912 documented in PDO production. In a recent study, Kumar et al. (2016) attempted to increase 913 PDO production yield by blocking the lactate and BDO pathways, while diverting carbon flux through the TCA cycle for efficient regeneration of NADH. However, this effort resulted 914 915 in the detrimental accumulation of pyruvate, acetate, and ethanol, causing low NADH regeneration and low PDO production. Neither increased aeration nor the addition of good 916 nitrogen sources (complex nitrogen sources) alleviated the accumulation of these 917 intermediates or activated the TCA cycle. Furthermore, the accumulation of pyruvate quickly 918 919 terminated glycerol assimilation. K. pneumoniae has a well-established BDO production 920 pathway that likely compensates for its inefficient TCA cycle, avoiding the overflow metabolism and carbon traffic at the pyruvate node. If this is the case, co-production of BDO 921 along with PDO or 3-HP, rather than single production of PDO or 3-HP, should be more 922 923 appropriate, which suits the nature of K. pneumoniae. In fact, when BDO and PDO were coproduced, more PDO was generated than during PDO production alone. More studies are 924 925 required to gain a better understanding of glycerol metabolism, the slow operation of the

TCA cycle in *K. pneumoniae*, and its effect on the selection of target products and/or
development of suitable strains and processes. Otherwise, it will be difficult to exploit this
potential biocatalyst for commercial manufacturing of valuable chemicals.

929 **4.3** Toxicity of intermediate metabolites and end-products

A general but major challenge to the use of microbial cell factories for producing bulk 930 chemicals is the toxic effect of target products and metabolic intermediates at high 931 932 concentrations. In particular, during the production of PDO or 3-HP from glycerol, the highly toxic intermediate 3-HPA is inevitably generated. Aldehydes generally damage DNA, 933 934 inactivate enzymes, and inhibit DNA synthesis. 3-HPA causes irreversible cessation of metabolic activity and cellular growth, even at very low concentrations of 15-30 mM (Hao et 935 al., 2008; Zheng et al., 2008). Native K. pneumoniae does not accumulate 3-HPA at toxic 936 937 levels during PDO production, but in engineered strains that overexpress DhaB, high 3-HPA 938 accumulation is often experienced. To avoid 3-HPA accumulation, enzymes that consume 3-HPA, ALDH (for 3-HP production), and PDOR (for PDO production) are expressed at 939 940 sufficiently (sometimes excessively) high levels compared with that of DhaB (Lim et al., 2016). Intermediates such as methylglyoxal, DHA, DHAP, and glyceraldehyde-3-phosphate 941 are also known to be toxic. 942

Organic acids and alcohols are toxic at high concentrations. Generally, organic acids 943 944 are more toxic than their corresponding alcohols because they disturb the intracellular pH and 945 have anion-specific effects on metabolism (Chun et al., 2014; Warnecke and Gill, 2005). When accumulated in cell membranes, organic compounds damage membrane integrity and 946 inhibit the activity of membrane-bound enzymes. In K. pneumoniae, cell growth and glycerol 947 948 assimilation are severely inhibited by high concentrations of PDO or 3-HP. Furthermore, enzymes that play major roles in 3-HP and PDO production, such as DhaB, DhaT, and AldH, 949 950 are inhibited by 3-HP and PDO (Kumar et al. 2013b). Furthermore, K. pneumoniae is

sensitive to 24 g/L acetic acid, 26 g/L lactic acid, and 17 g/L ethanol under aerobic conditions
and 15, 19, and 26 g/L, respectively, under anaerobic condition (Celińska, 2012; Cheng et al.,
2005). The inhibition of pathways by target product(s) and/or unwanted byproducts often
places a limit on the maximum attainable titer of bulk chemicals.

Several studies have reported engineered improvements in microbial tolerance against
biochemicals and biofuels, especially in *E. coli*. Modifications of exporter proteins, heat
shock proteins, membrane composition, and stress responses have been attempted. Moreover, *in situ* recovery of target products has been attempted for hydrophobic targets. Some tolerant
strains have been developed, but unfortunately, without appreciable increases in productivity
(Dunlop, 2011; Dunlop et al., 2011; Jarboe et al., 2011). Such efforts to improve tolerance to
chemicals have not been reported for *K. pneumoniae*.

962 **4.4 Availability of glycerol**

The global market for glycerol is known to be unpredictable and complex. Glycerol in 963 the current market mainly is from the biodiesel industry, and its supply is directly affected by 964 965 the use of biodiesel. In contrast, glycerol demand depends on many industries that use it as feedstock. Between 2001 and 2011, the price varied significantly, from \$0.70/kg to \$1.70/kg 966 for pure glycerol and from \$0.04/kg to \$0.33/kg for crude glycerol (Ayoub and Abdullah, 967 2012; Ciriminna et al., 2014; Kong et al., 2016; Quispe et al., 2013). Nevertheless, glycerol 968 969 prices, for pure as well as crude, have come down significantly since 2004, when many 970 biodiesel production plants initiated operation. The countries producing biodiesel in large quantities (in million liters) include Germany (2900), Brazil (2300), Argentina (2100), France 971 (2000), USA (1200), Spain (1100), Italy (800), Indonesia (700), and Thailand (600) (Kong et 972 973 al., 2016). The Asia-Pacific region overtook Europe as the largest glycerol market in the world in 2009 and has remained so ever since. Market growth is driven by increases in the 974

975 uses for glycerol in such sectors as pharmaceuticals, personal hygiene, and food and beverage
976 production (Quispe et al., 2013).

The production of crude glycerol is expected to reach 6 million tons by 2025 977 978 (Ciriminna et al., 2014). At present, the surplus of glycerol from biodiesel and oleochemicals 979 is assumed to be sufficient to meet these new demands. However, the scenario may change in the future with rapid growth of glycerol-based industries and a limited supply of crude 980 981 glycerol. For example, if 3-HP produced from glycerol were used as a substitute for the chemical acrylic acid (global production, 5.85 million tons as of 2014), approximately 7 982 983 million tons of glycerol/year would be needed. If we include other platform chemicals such as PDO, BDO, and ethanol, the demand will be even higher, much more than the expected 984 supply. The growth of glycerol-based biotechnology will eventually be constrained by the 985 986 supply and price of crude glycerol as a carbon source.

987 5. Concluding remarks

Glycerol is an excellent substrate for the production of biochemicals and biofuels. 988 989 Although the biodiesel industry has grown rather slowly in recent years, crude glycerol is still cheap and generated in large quantities. K. pneumoniae has already been successfully 990 991 employed for the production of PDO, 3-HP, BDO, lactic acid, and ethanol from glycerol at industrially attractive levels. Other metabolites such as acetic acid, pyruvic acid, malic acid, 992 993 fumaric acid, alanine, n-butanol, 1,2-propanediol, propionic acid, formic acid, and hydrogen 994 can also be produced from glycerol by K. pneumoniae, although these have not yet been 995 extensively explored. At present, sugar-based bioprocesses are prevalent, and no bioprocesses 996 employing K. pneumoniae and glycerol are commercially available. The pathogenicity of K. 997 pneumoniae and other technical issues are existing challenges to its use. However, advancements in the fields of metabolic engineering, synthetic biology, systems biology, and 998 999 evolutionary engineering will enable us to further exploit the advantageous nature of K.

- 1000 *pneumoniae* as a biocatalyst and glycerol as a carbon source. A better understanding of
- 1001 glycerol metabolism in *K. pneumoniae* and improvements in its performance as a microbial
- 1002 cell factory should make this organism an attractive alternative to current sugar-based
- 1003 methods for the production of biochemicals and biofuels.

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1462 **Figure captions**

Figure 1: Transformation of glycerol into a number of valuable chemicals. The solid linerepresent single step while broken one indicate multiple steps.

1465 **Figure 2:** Oxidative metabolism of glycerol in *K. pneumoniae* (Ashok et al., 2011; Kumar et

al., 2013a). Gene names are shown in italics on arrows. Solid lines indicate single steps;

1467 broken lines indicate multiple steps.

Figure 3: Metabolic pathways for the production of PDO and 3-HP from glycerol (Ashok et al., 2011; Kumar et al., 2013a). Gene names are shown in italics on arrows.

1470 Figure 4 (A): Arrangement of genes in the *dha* regulon of *K. pneumoniae*. 1, *dhaK* (ATP-

1471 dependent dihydroxyacetone kinase); 2, *dhaK3/dhaM* (large subunit of ATP-dependent

1472 dihydroxyacetone kinase); 3, *dhaK2/dhaL* (small subunit of ATP-dependent

1473 dihydroxyacetone kinase); 4, *dhaK1/dhaK* (medium subunits of ATP-dependent

1474 dihydroxyacetone kinase); 5, *orfW* (hypothetical protein); 6, *dhaD* (glycerol dehydrogenase);

1475 7, *dhaR* (transcription regulator); 8, *orfW*/CdAT [cob(I)yrinic acid a,c-diamide

adenosyltransferase]; 9, *orfX/dhaG* (small subunit of reactivating factor of glycerol

1477 dehydratase); 10, *dhaT* (1,3-propanediol oxidoreductase); 11, *orfY* (hypothetical protein); 12,

1478 *dhaB/B1* (large subunits of glycerol dehydratase); 13, *dhaC/B2* (medium subunit of glycerol

1479 dehydratase); 14, *dhaE/B3* (small subunit of glycerol dehydratase); 15, *orfZ/dhaF* (large

subunit of reactivating factor of glycerol dehydratase); 16, *glpF* (glycerol uptake facilitator);

1481 17, 22- *hdeB*; 18, *hdeD* (*hdeBD*-genes related to global regulation); 19, *yfdX*; 20, *kvgS*; and

1482 21, *kvgA* (*kvgS* and *kvgA* encode for a two-component signal transduction system) (Celińska,

1483 2012; Sun et al., 2003; Wei et al., 2014). (B): Glycerol dehydratase gene arrangements and

1484 reactivation factors in different organisms (Martins-Pinheiro et al., 2016).

Figure 5: Metabolic pathway for the synthesis of BDO from glycerol (Ashok et al., 2011;

1486 Celińska and Grajek, 2009; Ji et al., 2011). Gene names are shown in italics on arrows. Solid

1487 lines indicate single steps; broken lines indicate multiple steps.

- 1488 Figure 6: Biochemical routes for ethanol and acetic acid production from glycerol (Ashok et
- 1489 al., 2011; Kumar et al., 2013a). Gene names are shown in italics on arrow. Solid lines
- 1490 indicate single steps; broken lines indicate multiple steps.
- 1491 Figure 7: Lactic acid production from glycerol via methylglyoxal and pyruvate routes
- 1492 (Ashok et al., 2011; Mazumdar et al., 2010; 2013). Gene names are shown in italics on arrow.
- 1493 Solid lines indicate single steps; broken lines indicate multiple steps.
- 1494 **Fig. S1** Effect of deletion of glycerol transporters, *glpF1* and *glpF2*, on glycerol consumption
- and PDO production in *K. pneumoniae* J2B3.

- 1 Potential and Limitations of *Klebsiella pneumoniae* as a Microbial Cell Factory Utilizing
- 2 Glycerol as the Carbon Source
- 3
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- 5
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18 Abstract

Klebsiella pneumoniae is a Gram-negative facultative anaerobe that metabolizes 19 glycerol efficiently under both aerobic and anaerobic conditions. This microbe is considered 20 21 an outstanding biocatalyst for transforming glycerol into a variety of value-added products. Crude glycerol is a cheap carbon source and can be converted by K. pneumoniae into useful 22 compounds such as lactic acid, 3-hydroxypropionic acid, ethanol, 1,3-propanediol, 2,3-23 butanediol, and succinic acid. This review summarizes glycerol metabolism in K. 24 pneumoniae and its potential as a microbial cell factory for the production of commercially 25 26 important acids and alcohols. Although many challenges remain, K. pneumoniae is a promising workhorse when glycerol is used as the carbon source. 27 Key words: Klebsiella pneumoniae; Glycerol; 3-Hydroxypropionic acid; 1,3-Propanediol; 28

29 2,3-Butanediol; Lactic acid; Ethanol

30 **1. Introduction**

Crude glycerol is produced in large quantities as a byproduct of the biodiesel industry. 31 Global biodiesel production was 22.7 million tons in 2012 and is forecasted to reach 36.9 32 33 million tons by 2020. As a result, worldwide production of glycerol climbed from 1 million 34 tons in 2000 to 3 million tons in 2011 and is expected to reach 6 million tons by 2025 (Ciriminna et al., 2014; OECD, 2011; Katryniok et al., 2013; Sun et al. 2017). A significant 35 36 amount of glycerol is also generated during bioethanol fermentation by yeast. The liquid fraction of the stillage obtained from ethanol separation contains up to 2% glycerol. Another 37 38 source of glycerol is industrial waste generated from vegetable oils and animal fats. For example, the concentration of glycerol in waste streams from the oleochemical industry is 39 55%-90% (da Silva et al., 2009; Yazdani and Gonzalez, 2007). The increased availability of 40 41 glycerol has caused a substantial reduction in its cost. According to a recent report, the price 42 of crude glycerol is \$0.24/kg and that of pure USP grade glycerol is \$0.9/kg (Kong et al., 2016). Extensive research has been conducted to investigate ways to utilize this surplus crude 43 44 glycerol. The annual number of research articles addressing the use of glycerol has increased to >7000, doubling in number from the year 2000 to 2007. Numerous chemical conversions 45 to valuable products have been reported (Ciriminna et al., 2014). 46 Glycerol can be used for the microbial production of 1,3-propanediol (PDO), ethanol, 47 48 2,3-butanediol (BDO), butanol, lactic acid, 3-hydroxypropionic acid (3-HP), and succinic

49 acid (Fig. 1) (Clomberg and Gonzalez, 2013; Mattam et al., 2013; Yazdani and Gonzalez,

50 2007). Because of its reduced nature, glycerol generates twice the number of reducing

51 equivalents per carbon than do traditional carbohydrates (glucose, xylose, and sucrose) when

- 52 converted into the glycolytic intermediates phosphoenolpyruvate (PEP) or pyruvate
- 53 (Dharmadi et al., 2006). As a consequence, glycerol gives higher yields of reduced
- 54 metabolites (e.g., succinate, ethanol, PDO, and butanol) than does glucose (Yazdani and

Gonzalez, 2007). Table 1 compares the maximum theoretical yield of various metabolites,
ATP, and redox balances between glycerol and glucose.

Despite its great potential as a carbon source, glycerol is not efficiently metabolized 57 by many microorganisms under anaerobic or oxygen-limited conditions. The ability of K. 58 pneumoniae to assimilate glycerol under oxygen-limited conditions is outstanding and has 59 been studied extensively. Several review articles addressing this issue have also been 60 61 published (Clomberg and Gonzalez, 2013; Mattam et al., 2013; Yazdani and Gonzalez, 2007), but the biotechnological potential and physiological aspects of this strain have not been fully 62 63 elucidated. This review explores and discusses glycerol metabolism in K. pneumoniae and the production of value-added chemicals from glycerol. The potential and challenges of using K. 64 pneumoniae as a microbial cell factory are also discussed. 65

66 **2. G**

2. Glycerol metabolism in K. pneumoniae

Many microorganisms can metabolize glycerol in the presence of external electron 67 acceptors (respiratory metabolism), but relatively few under non-respiratory conditions 68 69 (without exogenous electron acceptors). Among the latter, genera including *Citrobacter*, Enterobacter, Bacillus, Propionibacterium, Anaerobiospirillum, Klebsiella, Clostridium, and 70 71 Lactobacillus are well documented (da Silva et al., 2009; Yazdani and Gonzalez, 2007). Some Lactobacillus species including L. reuteri, L. brevis, and L. bucheneri convert glycerol 72 73 to PDO but only in the presence of a major carbon source for growth. Glycerol does not 74 support cell growth, limiting its biotechnological applications. In contrast, *Clostridium sp.* grow well on glycerol as the sole carbon source. However, the use of *Clostridium* is limited 75 by its slow growth and requirement of strict anaerobic conditions. The lack of a convenient 76 77 genetic tool box for large-scale metabolic engineering is another serious drawback of Clostridium (Branduardi et al., 2014; Lütke-Eversloh and Bahl, 2011). 78

From a biotechnological standpoint, Klebsiella sp., especially K. pneumoniae, have 79 many advantages over the aforementioned microbes. They grow on glycerol as the sole 80 carbon source under both aerobic and anaerobic conditions (Arasu et al., 2011; Kumar et al., 81 82 2012). Their growth is fast, typically growing at a rate >0.9/h, even on glycerol minimal medium (Arasu et al., 2011). They naturally produce coenzyme B₁₂, an essential cofactor for 83 the synthesis of the commercially important chemicals PDO and 3-HP (Huang et al., 2013a; 84 85 Oh et al., 2012a). Furthermore, their evolutionary and biochemical proximity to E. coli allows for the application of most of gene manipulation methods developed for E. coli without much 86 87 modification (Celińska, 2012; Kumar et al., 2013a).

88

2.1 Overview of glycerol metabolism in K. pneumoniae

In *K. pneumoniae*, glycerol is metabolized oxidatively and/or reductively (Ashok et
al., 2011, Kumar et al., 2013a). The oxidative pathway provides energy and carbon
constituents for the synthesis of cell biomass, while the reductive pathway reduces glycerol
and regenerates NAD⁺, enabling the oxidative assimilation of glycerol under anaerobic
conditions.

Glycerol oxidation proceeds in either a respiratory or a fermentative manner. 94 Respiration requires an exogenous electron acceptor such as oxygen, nitrate, or fumarate 95 (Ashok et al., 2013a,b) and is carried out by enzymes encoded by the glp regulon. During 96 97 respiration, glycerol is first phosphorylated by glycerol kinase (glpK) to yield sn-glycerol-3-98 phosphate using ATP or phosphoenolpyruvate (PEP) as a phosphate donor. Next, snglycerol-3-phosphate is oxidized to dihydroxyacetone phosphate (DHAP) with electron 99 100 transfer to a quinone molecule, which is linked to the electron transport chain (ETC). This 101 transformation is catalyzed by glycerol-3-phosphate dehydrogenase, which is expressed under both aerobic (glpD) and anaerobic (glpABC) conditions. In contrast, fermentation of 102 103 glycerol is conducted without a supply of external electron acceptors, and the reactions are

104 catalyzed by enzymes encoded by the *dha* regulon. Here, glycerol is oxidized to

105 dihydroxyacetone (DHA) by glycerol dehydrogenase (mainly encoded by *dhaD*) using NAD⁺

106 as an electron acceptor. DHA then is phosphorylated by dihydroxyacetone kinase, an enzyme

107 encoded by the ATP-dependent *dhaK1/dhaK* and/or PEP-dependent

108 *dhaK2/dhaKLM/dhaK123*. DHAP obtained from respiration and/or fermentation is channeled

109 into the glycolytic pathway (Fig. 2). A variety of organic acids and alcohols, including acetic

110 acid, lactic acid, succinic acid, ethanol, BDO, and formic acid, are produced by the

downstream glycolytic pathway (Ashok et al., 2011; Kumar et al., 2013a).

112 The anaerobic growth capability of *K. pneumoniae* using glycerol as a carbon source

is attributed to its well-established reductive metabolism. In the reductive pathway, glycerol

114 first undergoes a difficult rearrangement reaction catalyzed by glycerol dehydratase (GDHt)

to yield 3-hydroxypropioaldehyde (3-HPA). Catalysis by GDHt requires coenzyme B_{12} , the

116 *de novo* synthesis of which is limited to only few genera of microorganisms. The 3-HPA is

subsequently reduced to PDO by at least two 1,3-propanediol oxidoreductases (1,3-PDORs),

118 including NADH-dependent DhaT (PDOR) and NADPH-dependent hypothetical

119 oxidoreductase (HOR), along with the regeneration of $NAD(P)^+$ (Fig. 3). *E. coli* and other

120 *enterobacter sp.* have the oxidative (respiratory) pathway of glycerol assimilation. However,

121 these organisms lack the reductive pathway and cannot grow on glycerol under anaerobic

122 conditions (Dharmadi et al., 2006; Kumar et al., 2013a).

123 2.2 Genes and enzymes involved in fermentative metabolism of glycerol

124 The *dha* regulon, induced by the DHA molecule and expressed in the absence of

125 exogenous electron acceptors, encodes numerous genes needed for both the oxidative

126 (fermentative) and reductive pathways (Celińska, 2012; Forage and Foster, 1982) (Fig. 4A).

127 The *dha* regulon of *K*. *pneumoniae* contains the following genes arranged in the order as

128 appears: *Kpk_0615/dhaK*, *dhaM*, *dhaL*, *dhaK*, *dhaD*, *dhaR*, *orfW*, *CdAT*, *orfX* (*dhaG*), *dhaT*,

orfY, dhaB, dhaC, dhaE, orfZ (dhaF), and glpF. Of these, dhaB, dhaC, dhaE, orfX, orfZ, and 129 *dhaT* are relatively well documented for their functions in the reductive metabolism of 130 glycerol. The *orfW* and *orfY* genes, although commonly found in the *dha* regulons of many 131 organisms, are not as well described. The proteins encoded by the *dha* regulon of *K*. 132 pneumoniae have high similarity (80%–95%) with those of Citrobacter species but not with 133 those of *Clostridium* species (30%–80%). Surprisingly, the sequence similarity of *dha* 134 135 proteins among *Clostridium* species is very low in some cases, even lower than their similarity to the corresponding proteins in K. pneumoniae or C. freundii (Celińska, 2012; Sun 136 137 et al., 2003; Wei et al., 2014).

138 The complete *dha* regulon is found in only a few dozen species belonging to five

139 different taxonomic groups, including four bacterial species [Actinobacteria, Firmicutes,

140 Fusobacteria, and Proteobacteria (Gammaproteobacteria and Deltaproteobacteria)] and one

141 archaea species, *Halalkalicoccus jeotgali* (Martins-Pinheiro et al., 2016). Interestingly, an

142 incomplete *dha* regulon is present in more than 100 prokaryotes, suggesting that the enzymes

143 of the regulon have other functions not restricted to the anaerobic growth on glycerol. A well-

144 documented example is the one found in *Lactobacillus* species. *Lactobacillus* cannot grow on

145 glycerol as the sole carbon source because it lacks the enzymes for oxidative assimilation of

146 glycerol. However, the enzymes encoded in the incomplete *dha* regulon enable the

147 production of PDO from glycerol, which accelerates cell growth in *Lactobacillus* grown on

148 glucose (Årsköld et al., 2008; Martins-Pinheiro et al., 2016).

In the following sections, the major enzymes involved in fermentative glycerolmetabolism and their genes are described in detail.

151 2.2.1 Glycerol dehydrogenase

152 Encoded by the *dhaD* and/or *gldA* gene, glycerol dehydrogenase is an oxidoreductase
153 that converts glycerol to DHA. The enzyme extracts electrons from glycerol and transfers

them to NAD⁺. DhaD plays a major role during the anaerobic growth of *K. pneumoniae* on
glycerol. Quantification of mRNA expression indicates that *dhaD* is highly induced when
glycerol is used as carbon source. GldA expression is lower than that of DhaD and is thought
to serve as a back up to DhaD during anaerobic growth on glycerol. However, DhaD and
GldA exhibit a high level of similarity in amino acid sequence and have the same substratebinding site (Wang et al., 2014).

160 Recently, DhaD was reported to catalyze the interconversion of acetoin and BDO for NADH disposal and recovery (Wang et al., 2014). The promiscuity of DhaD is responsible 161 162 for BDO production in the $\Delta budC$ mutant of K. pneumoniae. The contribution by DhaD to or its participation in BDO production is also supported by the observed increase in BDO 163 synthesis and increased concentration of NADH in the presence of the *dhaD* gene. NADH is 164 165 generated by the oxidation of glycerol and consumed by the production of BDO from acetoin. 166 The dual physiological functions of DhaD allow K. pneumoniae to control the intracellular redox level. In response to the intracellular level of NADH, DhaD is able to switch its role 167 from NADH production (glycerol oxidation) to NADH consumption (BDO production) 168 (Wang et al., 2014). The diversion of glycerol flux from organic acids toward alcohols such 169 170 as BDO also helps the cells to counteract intracellular acidification (Petrov and Petrova, 2009). Thus, this promiscuity confers different physiological roles to the enzyme, which 171 include regulation of the intracellular NADH/NAD⁺ ratio, prevention of acidification, and 172 173 storage of carbon and energy (Wang et al., 2014).

174 **2.2.2 Dihydroxyacetone kinases**

Dihydroxyacetone kinase (DHAK) phosphorylates DHA to DHAP, an intermediate of
the glycolytic pathway. Sun et al. (2003) analyzed the genome of *K. pneumoniae* MGH
78578 and found, in addition to ATP-dependent DhaK I, a PEP-dependent DhaK II. DhaK I
is encoded by the *dhaK* (*Kpk* 0615) gene, while DhaK II is encoded by three genes, *dhaK*,

179 *dhaL* and *dhaM*, which correspond to the medium, small, and large subunits of the enzyme, respectively (Celińska, 2012; Sun et al., 2003; Wei et al., 2014). The names dhaKLM and 180 181 *dhaK123* are synonyms used without distinction (Martins-Pinheiro et al., 2016; Sun et al., 182 2003; Wei et al., 2014). DhaK enzymes are found in many microorganisms. PEP-dependent DhaK II is present in E. coli, L. lactis, and C. butyricum, whereas ATP-dependent DhaK I is 183 present in Citrobacter freundii (Garcia-Alles et al., 2004; Raynaud et al., 2011). The PEP-184 dependent DhaK II of E. coli, encoded by dhaKLM, is homologous to dhaK123 (dhaKLM) of 185 K. pneumoniae (Gutknecht et al., 2001). In a recent study, Wei et al. (2014) further elucidated 186 187 the physiological roles of DhaK I and DhaK II of the *dha* regulon of *K. pneumoniae*. They found that disruption of ATP-dependent DhaK I had no significant effect on glycerol uptake, 188 indicating that PEP-dependent DhaK II is the major contributor to the conversion of DHA to 189 190 DHAP. As in E. coli, the subunits of DhaK II regulate expression of the dha regulon; dha 191 regulon expression was suppressed by the disruption of *dhaK1* (*dhaK*) and *dhaK2* (*dhaL*). However, in contrast to E. coli, mutation of dhaK3 (dhaM) in K. pneumonia did not 192

upregulate the *dha* regulon but rather decreased cell growth and glycerol uptake.

194 2.2.3 Glycerol dehydratase and reactivating factor

Glycerol dehydratase (GDHt) is the first enzyme in the reductive pathway and 195 196 catalyzes the dehydration of glycerol to 3-HPA. The same reaction can be carried out by diol dehydratase (DDHt), although the latter has a higher activity on 1,2-propanediol than on 197 glycerol. GDHt requires coenzyme B₁₂ or S-adenosyl methionine (SAM) as a cofactor. K. 198 pneumoniae has B₁₂-dependent GDHt only, while strict anaerobes such as clostridia sp. have 199 200 SAM-dependent GDHt. The three subunits of GDHt are encoded by the genes dhaB1 (~1670 bp), dhaB2 (~590 bp) and dhaB3 (~430 bp), for which the gene products are the α (large, 201 60.7 kDa), β (medium, 21.3 kDa), and γ (small, 16.1 kDa) subunits, respectively (Xu et al., 202 2009a). GDHt of K. pneumoniae is known to be present as a dimer of heterotrimers, $(\alpha\beta\gamma)_2$. 203

The subunits of GDHt have high homology among *K. pneumoniae* strains; for example, *K. pneumoniae* XJPD-Li and U30909 strains showed >99% identity.

Coenzyme B_{12} is present in the reaction center and plays an essential role in the 206 207 catalysis of GDHt. The catalytic cycle often results in the mechanism-based inactivation of coenzyme B₁₂ due to the irreversible breakage of the chemical bond between cobalt (Co) and 208 a carbon of the adenosyl moiety. Oxygen accelerates breakage of the Co-C bond. Oxygen is 209 210 also known to inactivate GDHt, even in the absence of the substrate glycerol. The gene products of orfZ/dhaG (small subunit) and orfX/dhaF (large subunit) act as reactivating 211 212 factors of GDHt (GdrAB) (Celińska, 2012; Sun et al., 2003). They reactivate GDHt by catalyzing the exchange of damaged for intact coenzyme B_{12} in the presence of ATP and 213 Mg²⁺/Mn²⁺ (Celińska, 2012; Kumar et al., 2013a; Maervoet et al., 2011; Sun et al., 2003; Wei 214 215 et al., 2014). The reactivase is a molecular chaperone that functions as a heterotetramer 216 containing two elongated α subunits (63 kDa) and two globular β subunits (14 kDa). Structurally, the α subunit resembles both GroEL and Hsp70 chaperones, while the β subunit 217 resembles that of the β subunit of glycerol dehydratase, except that it lacks some of the amino 218 acids responsible for coenzyme B_{12} binding (Liao et al., 2003). 219 In some organisms, the GDHt enzyme varies in terms of the number of subunits and 220 their copies (Fig. 4B) (Liu et al., 2010). For example, in Mesorhizobium loti and 221 222 Mesorhizobium opportunistum, the large and medium subunits (dhaB1 and dhaB2) are fused together and encoded by a single gene. The gene fusion may arise by frameshift mutations. 223 224 The active site of the fused enzyme differs slightly from that of unfused enzyme. Moreover, these microorganisms lack the genes encoding the reactivation factor, indicating that the 225 226 reactivation process may not exist or is carried out by a different route. Mycobacterium smegmatis has two subunits (fused dhaB1 + dhaB2 and dhaB3) that are similar to those of M. 227 *loti* and *M. opportunistum*; however, the reactivating factor differs in that it has one (large) 228

229 subunit. No small subunit has been identified in *M. smegmatis*. Fusobacterium *Ilyobacter polytropus* is unique in that it has genes coding for both B₁₂-independent (SAM-dependent) 230 and B₁₂-dependent GDHt. The genes for B₁₂-independent GDHt are present in the genomic 231 232 DNA, while those encoding B₁₂-dependent GDHt along with its reactivation factor are present in two copies, one in the chromosome and the other in a plasmid. Surprisingly, even 233 with this gene redundancy, no significant production of 3-HPA and/or PDO has been reported 234 235 in *I. polytropus* (Stieb and Schink, 1984). The B₁₂-independent proteins showed no homology with B₁₂-dependent GDHt but significant similarity with the pyruvate formate lyases (PFL) 236

and PFL-activating enzymes and their homologues (Raynaud et al., 2003).

238 2.2.4 1,3-Propanediol oxidoreductase and hypothetical oxidoreductase

The 1,3-propanediol oxidoreductase (PDOR), responsible for the second step of the 239 240 reductive pathway of glycerol, catalyzes the conversion of 3-HPA to PDO with the oxidation 241 of NAD(P)H. 3-HPA is a toxic intermediate, and PDOR protects the cell from deleterious damage by reducing 3-HPA to PDO (Celińska, 2012). In K. pneumoniae, PDOR is encoded 242 by the *dhaT* gene and is strictly NADH dependent. Active PDOR is a homo-octamer with a 243 monomeric molecular mass of 43 kDa. Despite its importance, the kinetic properties of DhaT 244 were elucidated only recently (Lama et al., 2015). DhaT has high specificity for 3-HPA 245 among various aldehydes. Under physiological conditions at pH 7.0 and 37°C, the catalytic 246 247 efficiency (k_{cat}/K_m) of the forward reaction (3-HPA to PDO) was found to be 77.2/s. mM, 248 which is 57-fold higher than that of the reverse reaction (1.3/s. mM; PDO to 3-HPA). The $K_{\rm m}$ (mM) values for 3-HPA and NADH (forward reaction) are 0.77 and 0.03, respectively, while 249 those for PDO and NAD⁺ (backward reaction) are 7.4 and 0.23, respectively. Furthermore, 250 251 the maximum activity of DhaT for the reverse reaction was obtained at pH 9.0 and 55°C. These results strongly suggest that, under physiological conditions, DhaT catalyzes the 252 reduction of 3-HPA to PDO rather than the oxidation of PDO to 3-HPA. Nevertheless, the 253

reverse reaction (PDO→3-HPA) becomes significant as the PDO concentration increases.
This factor is considered a serious drawback of using DhaT for PDO production from
glycerol.

K. pneumoniae has several oxidoreductases in addition to PDOR that can reduce 3-257 HPA to PDO. One such enzyme, designated as hypothetical oxidoreductase (HOR), has been 258 studied extensively and shares 89% identity with YqhD of E. coli (Zhu et al., 2009; Zhuge et 259 al., 2010). Expression of yqhD in wild-type K. pneumoniae is not high, with an mRNA level 260 ~20-fold lower than that of *dhaT*. However, deletion of *dhaT* increases the transcription of 261 262 *yqhD* by more than 10-fold (Ko et al., 2015). This observation suggests that in addition to its wide-ranging activity on short-chain aldehydes, YqhD is a detoxification enzyme for many 263 aldehydes (Jarboe, 2011). YqhD exhibits negligible activity on PDO (backward reaction), 264 265 which is an important advantage over DhaT for use in the production of PDO from glycerol 266 (Celińska, 2012; Chen et al., 2011; Zheng et al., 2006). However, YqhD is kinetically inferior to DhaT because of its low catalytic activity (k_{cat}/K_m) on 3-HPA and the requirement of 267 NADPH as a cofactor. The k_{cat}/K_m value of YqhD on 3-HPA (pH 7.0 and 37°C) is estimated 268 to be 2.1/s. mM, which is 36-fold lower than that of DhaT (unpublished data). The use of 269 270 NADPH as a cofactor prohibits this enzyme from being directly coupled to the reactions catalyzed by DhaD and other enzymes in glycolysis and the TCA cycle, by which NADH is 271 mainly produced. Furthermore, excessive use of NADPH for PDO production can 272 273 significantly reduce the cellular NADPH level and this can disturb many anabolic reactions requiring NADPH (Wang et al., 2003; Jarboe, 2011). NADH and NADPH differ in a 274 phosphate group only, and it has been suggested that electrostatic interaction of this 275 276 phosphate is the major factor distinguishing NADH from NADPH for enzymes using these cofactors. Comparison of the amino acid sequences of DhaT and HOR from K. pneumoniae 277 278 and YqhD from E. coli has revealed that DhaT contains aspartic acid (Asp) at position 41,

while HOR and YqhD have valine (Val) at this position. The steric hindrance and
electrostatic repulsion between Asp in the active site and the phosphate group of NADPH
impede their binding. In contrast, the hydrophobic amino acid Val poses no such obstruction;
thus, YqhD uses NADPH as a cofactor (Ma et al., 2010).

283 Despite their difference in cofactor preference, DhaT and YqhD are similar (44%) in 284 amino acid sequences. *K. pneumoniae* has five enzymes that have at least 40% similarity in 285 amino acid sequence with DhaT and YqhD: putative alcohol dehydrogenase (PaldH), L-1,2-286 propanediol oxidoreductase (FucO), ethanolamine-utilization enzyme (EutG), propanediol-287 utilization propanol dehydrogenase (PduQ), and bifunctional alcohol dehydrogenase (AdhE) 288 (Ko et al., 2015). The roles of these enzymes in PDO production have been investigated.

289 2.2.5 Regulatory protein

290 The *dhaR* gene product is an important transcription factor that is responsible for 291 expression of the *dhaT* and *dhaB* genes (Sun et al., 2003). This gene is found in K. pneumoniae, C. freundii, and Clostridium botulinum but not in every organism containing a 292 293 complete *dha* regulon (Martins-Pinheiro et al., 2016). The presence of DhaR is thought to allow the *dha* operon to function as a separate regulatory system, independent of fumarate 294 295 nitrate reduction (FNR). Under anaerobic conditions, FNR acquires DNA-binding properties and induces the expression of many anaerobiosis-related genes. DhaR contains domains for 296 GAF (52-199), PAS (203-267), σ^{54} factor interaction, and histidine HTH-8. The PAS senses 297 internal levels of energy charge, light, oxygen, and redox potential. The HTH-8 domain has a 298 helix-turn-helix conformation and acts as a DNA binding structure. The σ^{54} factor interaction 299 domain interacts with the σ^{54} factor of RNA polymerase and activates RNA transcription 300 from the σ^{54} promoters (Buck et al., 2000; Taylor and Zhulin, 1999). Thus, the DhaR protein 301 triggers the transcription of *dha* genes in response to intracellular levels of several important 302 physiological parameters (Celińska, 2012; Sun et al., 2003). Zheng et al. (2006) investigated 303
the effect of overexpression of the putative regulatory gene dhaR in the dha regulon on 304 glycerol metabolism in K. pneumoniae. dhaR overexpression increased PDOR activity up to 305 6.7-fold over that of the control wild-type strain, confirming the role of DhaR as a positive 306 regulator of the *dhaT* gene. However, *dhaR* overexpression did not increase the activity of 307 glycerol dehydratase (DhaB). Furthermore, despite the significant increase in PDOR activity, 308 PDO production was unexpectedly low compared with that of the wild type (440 vs. 656.23 309 310 mM, respectively). More studies are required to determine the precise role of DhaR and its regulatory function in the *dha* regulon. 311

312 DhaR in *E. coli* (70% identity to that of *K. pneumoniae*) has also been studied. This

313 protein stimulated transcription of the *dhaKLM* operon from a σ^{70} promoter. Interestingly, in

E. coli, DhaL was a positive regulator of the *dhaKLM* operon while phosphorylated DhaM

and DhaK were negative regulators of the same operon. DhaK contains the DHA-binding site.

316 DhaL uses ADP as a cofactor for the double displacement of phosphate from DhaM to DHA.

317 DHA is the oxidation product of glycerol metabolism; binding of DHA to DhaK reduces the

affinity of DhaK for DhaR. In the presence of DHA, DhaL::ATP is dephosphorylated,

leading to the displacement of DhaK by DhaL::ADP, which stimulates DhaR activity. In the

320 absence of DHA, DhaL::ADP is phosphorylated back to DhaL::ATP by the phototransferase

321 system (PTS), thereby inhibiting the binding of the phosphorylated complex (DhaL::ATP) to

322 DhaR. This phosphorylation is mediated by DhaM, providing a phospho-histidine relay

between the PTS and DhaL::ADP. This double-check mechanism of binding and turnover

324 increases the selectivity such that the binding of nonphosphorylated compounds does not

induce the *dha* operon (Bächler et al., 2005).

326 **2.3 Genes and enzymes involved in respiratory assimilation of glycerol**

The *glp* regulon comprises the genes encoding proteins required for the conversion of glycerol to DHAP via sn-glycerol-3-phosphate. The five operons that constitute the *glp*

regulon are *glpFK*, *glpD* (anaerobic glycerol-3-phosphate dehydrogenase), *glp ACB* (aerobic 329 glycerol-3-phosphate dehydrogenase), *glpTQ* (glycerol-phosphate 330 permease/glycerophosphodiesterase), and *glpEGR*. The *glpF* gene encodes a cytoplasmic 331 332 membrane protein that facilitates the diffusion of glycerol into the cell. The *glpE* gene encodes an acidic, cytoplasmic protein of 108 amino acids with a molecular weight of 333 12,082. The glpG gene encodes a basic, cytoplasmic membrane-associated protein of 276 334 335 amino acids with a molecular weight of 31,278 (Lin, 1976; Yang and Larson, 1998; Zeng et al., 1996). The biochemical functions of GlpE and GlpG are unknown. The *glpR* gene 336 337 encodes the GlpR repressor protein. These five operons are located at three different positions on the chromosome. Transcription of these operons is subject to multiple controls, including 338 catabolite repression mediated by cAMP-CRP and respiratory control mediated by the FNR 339 340 and ArcA/ArcB systems. Moreover, each of the operons is negatively controlled by a 341 repressor specific for the regulon (Lin, 1987; Zeng et al., 1996).

342 2.3.1 GlpR and other regulatory proteins

The GlpR repressor protein, predicted to contain 252 amino acids with a molecular 343 weight of 28,048, belongs to the DeoR bacterial regulatory factor super family. Although not 344 studied extensively, GlpR of K. pneumonia is expected to have the same biochemical 345 properties and regulatory functions as that of E. coli. GlpR negatively controls all the glp 346 347 operons by binding to operators that overlap or are close to the *glp* promoters. In case of E. 348 *coli*, glycerol-3-phosphate acts as an inducer and has a high affinity for GlpR ($K_d = 31 \mu M$) (Lin, 1987; Weissenborn et al., 1992; Zeng et al., 1996). When complexed with glycerol-3-349 350 phosphate, GlpR cannot bind to the operator sites, and repression is relieved. Glycerol-3-351 phosphate is a product of GlpK and a substrate of GlpD/GlpABC; thus, the absence of GlpD causes a substantial increase in the level of glycerol-3-phosphate, resulting in induction of the 352 353 glp regulon (Fig. 2). The GlpR protein exhibits differential binding affinity for the operators

354 of corresponding operons, as follows: glpFK > glpD > glpTQ > glpACB. The glpFK promoter appears to be the strongest among the *glp* promoters. Therefore, the maximum strength of the 355 *glpFK* promoter combined with the high affinity of GlpK for glycerol ($K_m = 10 \mu M$) should 356 357 result in sufficient accumulation of the inducer, glycerol-3-phosphate, in the presence of glycerol to relieve repression by GlpR. The relatively lower strength of the *glpD* promoter 358 along with the low affinity of the GlpD enzyme for glycerol-3-phosphate ($K_m = 1 \text{ mM}$) 359 prevents excessive degradation of glycerol-3-phosphate that is needed for phospholipid 360 synthesis when the exogenous source of glycerol-3-phosphate becomes limiting, until after 361 362 accumulation of the inducer (Lin, 1987; Weissenborn et al., 1992). Recently, Jung et al. (2014) investigated the role of GlpR in the aerobic production of 3-HP from glycerol in E. 363 coli. They found that elimination of this regulatory factor caused increased assimilation of 364 365 glycerol and higher production of 3-HP and suggested that the deletion of glpR led to the 366 upregulation of genes involved in glycerol transport and assimilation. K. pneumoniae has several other regulatory genes in addition to *glpR*: the global regulatory genes *hdeB*, *hdeD*, 367 and *yfdX*, with unknown functions and *kvgS* and *kvgA*, encoding two proteins comprising a 368 two-component signal transduction system (Sun et al., 2003). 369

370 2.3.2 Glycerol uptake/transport facilitator and other components

The glycerol facilitator GlpF is an integral membrane protein that forms aqueous 371 372 pores. The GlpF pores selectively allow passive transport of glycerol and other molecules 373 such as DHA and urea across the cytoplasmic membrane, but not water or ions (Heller et al., 1980; Reizer et al., 1993). The GlpF protein contains six membrane-spanning helices that are 374 unique among prokaryotic transport proteins. These transmembrane helices are arranged in 375 376 two bundles. The GlpF protein may function as a homodimer with the two six-member domains arranged in the membrane to form a channel (Weissenborn et al., 1992). K. 377 pneumoniae has two glycerol transporter genes, one next to orfZ/dhaF in the fermentative 378

379 *dha* regulon (see Fig. 4A) and the other in the respiratory *glpFK* operon. Both *glpF* genes have a high similarity (85 %) to each other. When one but not both of the glpF genes is 380 disrupted, the glpF deletion mutants of K. pneumoniae can consume glycerol and convert it to 381 382 PDO at a high rate, similar to the wild type (unpublished data; Supplementary Fig. S1). This observation indicates that both *glpF* genes encode active glycerol transporter proteins. 383 However, their presence is not essential for glycerol utilization in K. pneumoniae because the 384 385 double-deletion mutant for both facilitators can still uptake glycerol and convert it to PDO. In *E. coli*, only one *glpF* gene is present in the *glpFK* operon; the GlpF protein shows a high 386 387 similarity (80.9%) to that (encoded from *glpFK*) of *K. pneumoniae* (Sun et al., 2003). The importance of GlpF in *E. coli* also seems to be limited only at low levels of glycerol (<8 388 mM), as glycerol itself may pass through the membrane by diffusion at high concentrations 389 390 (Richey and Lin, 1972).

2.3.3 Simultaneous operation of fermentative and respiratory pathways

The presence of the *dha* and *glp* regulons affords *K*. *pneumoniae* metabolic flexibility. 392 393 The *dha* regulon enables fermentative utilization of glycerol, while the *glp* regulon enables respiratory metabolism of glycerol. During fermentative metabolism of glycerol, redox 394 395 constraints force the production of PDO from glycerol, with a low ratio of carbon conversion yield to cell biomass. Moreover, ATP is formed through substrate-level phosphorylation, and 396 397 NADH oxidation occurs through the formation of reduced metabolites. In the presence of 398 oxygen, the redox constraints forcing the production of PDO are relieved; therefore, the carbon loss required for regeneration of NAD⁺ is greatly reduced. As a consequence, glycerol 399 utilization, biomass synthesis, and even the formation of many metabolites increase (Durnin 400 401 et al., 2008). Interestingly, the fermentative route (dha regulon) is in operation under a wide range of oxygen levels, from microaerobic to highly aerobic conditions, as evidenced by 402 PDO formation. For example, a high titer PDO (102.7 g/L) was produced under the condition 403

of 2.0 vvm aeration and even at 3.5 vvm, although the production was a little lowered in the
latter case (Oh et al., 2012a). These results suggest that the fermentative and respiratory
pathways might have evolved to complement each other for optimal control of cell growth
under a wide range of aeration conditions except the strictly anaerobic one.

408

2.4 Carbon catabolite repression

Carbon catabolite repression (CCR) refers to the suppression of the use of less-409 preferred carbon sources, when a more preferred carbon, most prominently glucose, is 410 present. Cyclic AMP, cAMP receptor protein (CRP), and the enzyme EIIA^{Glc}, an 411 412 intermediate in the phosphorylation cascade of the PTS, are key players in the CCR of enterobacteria. The EIIA^{Glc}, a cytosolic protein, exists in phosphorylated and 413 414 unphosphorylated forms. The phosphorylated form stimulates membrane-bound adenylate 415 cyclase, which catalyzes the formation of cAMP; the cAMP-CRP complex then induces 416 expression of the catabolic genes of less-preferred carbon sources. The unphosphorylated form of EIIA^{Glc} is also responsible for inhibiting the transport of less-preferred carbon 417 418 sources by inhibitory binding to permeases, causing so-called inducer exclusion (Eppler and Boos, 1999; Eppler et al., 2002). Glycerol assimilation in K. pneumoniae is greatly inhibited 419 by CCR in the presence of glucose. This repression is related to not only low cAMP and 420 cAMP-CRP but also inhibition of GlpK (which functions mainly under aerobic conditions) 421 by fructose 1,6-bisphosphate, a metabolite produced from glucose. The unphosphorylated 422 EIIA^{Glc} is also known to allosterically inhibit GlpK (Holtman et al., 2001). Anaerobic 423 glycerol metabolism mediated by the *dha* operons is also suppressed by the presence of 424 glucose. Enzyme-level inhibition similar to that of the aerobic GlpK has not been reported, 425 426 but we have observed that, in K. pneumoniae, transcription of the dha operons is greatly reduced in the presence of glucose (Suman et al., 2017). 427

Interestingly, glycerol exerts catabolite repression on the assimilation of maltose in E. 428 *coli*. To exert catabolite repression, glycerol must be phosphorylated to glycerol-3-phosphate 429 (Hogema et al., 1998; Eppler et al. 2002). However, unlike other carbon sources, further 430 431 metabolism of glycerol is not required. According to Eppler and Boos (1999), glycerol-3phosphate reduces expression of MalT, a positive activator of all mal genes. The key players 432 in this repression are adenylate cyclase, EIIA^{Glc}, and CRP. The growth on maltose of the 433 mutants lacking EIIA^{Glc} or containing truncated adenylate cyclase was no longer repressed by 434 glycerol, and CRP-independent transcription of malT was also not influenced by glycerol 435 either. In addition, stimulation of adenylate cyclase by phosphorylated EIIA^{Glc} was controlled 436 by glycerol-3-phosphate. The mutation in adenylate cyclase relieved the repression caused by 437 glycerol-3-phosphate (Eppler et al., 2002). Although not investigated, we assume that the 438 439 glycerol repression on maltose catabolism also hold true in K. pneumoniae. 440 Glycerol also exerts catabolite repression on the assimilation of citrate in K. pneumoniae. Citrate can be used as the sole carbon and energy source during anaerobic 441 growth of K. pneumoniae, but the presence of glycerol suppresses the expression of the 442

citrate fermentation genes. A reduced concentration of the cAMP–CRP complex has been
postulated to be the cause of this repression (Meyer et al., 2001). Further research is needed
to test the hypothesis.

3. *K. pneumoniae* as a microbial cell factory for the production of commodity chemicals During glycerol fermentation, *K. pneumoniae* produces many metabolites, including PDO, BDO, and lactic acid. This section describes the biotechnological production of bulk chemicals from glycerol in *K. pneumoniae*. 3-HP is not a natural product during the growth of *K. pneumoniae* on glycerol. However, its production is possible using a minor modification of the PDO synthetic pathway and is included here. The biochemical pathways

used for the synthesis of these chemicals are explained, and the metabolic engineering used toimprove their production is discussed.

454 **3.1 1,3-Propanediol**

PDO, also known as trimethylene glycol, is a promising platform chemical that has
two hydroxyl groups. With terephthalic acid, it can be co-polymerized to the novel copolymer polytrimethylene terephthalate (PTT). PDO also has applications in the food and
cosmetic industries. Currently, commercial PDO is mainly produced by recombinant *E. coli*(Maervoet et al., 2011; Saxena et al., 2009). The global demand for PDO was 60.2 kilotons in
2012 and is predicted to reach approximately 150 kilotons by 2019. The market for PDO is
growing rapidly, from \$157 million in 2012 with projections up to \$560 million in 2019

462 (MarketsANDMarkets, 2012; Lee et al., 2015).

463 K. pneumoniae is one of the best native producers of PDO from glycerol. GDHt and PDOR are two important enzymes for the conversion of glycerol to PDO (Fig. 3). 464 Physiologically, the synthesis of PDO from glycerol is essential for the anaerobic growth of 465 K. pneumoniae, as described previously. The production of PDO requires two cofactors, 466 coenzyme B₁₂ and NAD(P)H. K. pneumoniae has a de novo pathway for the biosynthesis of 467 coenzyme B₁₂ that comprises more than 20 genes. NAD(P)H is generated through the 468 oxidative metabolism of glycerol, during the production of cell biomass and/or oxidized 469 470 (pyruvate and acetate) or partially reduced (BDO, lactic acid) metabolites. The production 471 yield of PDO from glycerol varies depending on the cost of carbon for NAD(P)H production; the lower the cost, the higher the yield of PDO from glycerol. NADPH can also be the 472 electron donor for the conversion of 3-HPA to PDO when YqhD is present. However, the 473 474 contribution of NADPH to PDO production in native K. pneumoniae seems to be limited because the PP pathway, which is the main source of NADPH, is not active when glycerol is 475

476	the sole carbon source. Furthermore, K. pneumoniae has very low transhydrogenase activity
477	for the conversion of NADH to NADPH (Zhang and Xiu, 2009; Chen et al., 2011).
478	Several metabolic engineering strategies have been employed to improve PDO
479	production by K. pneumonia (Table 2). Amplification of the reductive pathway by
480	overexpressing DhaB, PDOR, or both has been attempted by many research groups. As
481	indicated previously, 3-HPA is highly toxic, and its accumulation is detrimental to cell
482	viability and toxic to PDO production. To reduce 3-HPA accumulation, overexpression of
483	dhaT has been attempted. In one study, 3-HPA accumulation was reduced by dhaT
484	overexpression, but no improvement in PDO production resulted (Hao et al., 2008).
485	Overexpression of both DhaT and DhaD increased PDO production (56.3%) in batch culture
486	but no increase in fed-batch culture (Chen et al., 2009). With another K. pneumoniae strain,
487	DSM 2026, the same experiments were repeated but also with no increase in PDO production
488	(Zheng et al., 2006). These results suggest that the reductive pathway in K. pneumoniae is
489	evolutionarily well established (perhaps in concert with the oxidative pathway) and that PDO
490	production is not limited by expression of the <i>dhaB</i> and <i>dhaT</i> genes of the reductive pathway.
491	However, in one study where the <i>E</i> . <i>coli</i> yqhD gene was highly overexpressed to give $\sim 10^{-10}$
492	fold higher PDOR activity (110 IU/mg total protein), a significant increase in PDO titer (25%;
493	67.6 g/L) and yield (from 0.53 to 0.62 mol/mol), along with less 3-HPA accumulation, was
494	achieved (Zhu et al., 2009). This observation suggests that in the recombinant strain, NADPH
495	can play an important role as a cofactor and that PDO production can be increased by
496	modifying the reductive pathway. It is also probable that the effect of DhaB and PDOR on
497	PDO production is highly dependent on the strain and/or culture conditions adopted.
498	Lactic acid, BDO, ethanol, and acetic acid are the major byproducts of PDO synthesis
499	in K. pneumoniae. Inactivation of the metabolic pathways that give rise to these products has
500	resulted in increased PDO synthesis. In particular, deletion of the <i>ldhA</i> gene encoding lactate

501 dehydrogenase has been highly beneficial. For example, by disrupting the *ldhA* gene, Xu et al. (2009b) significantly increased PDO production without lactic acid production. They 502 observed increases in the PDO titer (95.4–102.1 g/L), conversion yield (0.48–0.52 mol/mol), 503 504 and productivity (1.98–2.13 g/L h; Table 2). Similarly, Oh et al. (2012a) achieved the high PDO titer of 102.7 g/L by deleting ldhA in their own K. pneumoniae isolate. When lactic acid 505 production was eliminated, more ethanol and succinic acid were generated from the pyruvate 506 507 node. Reduction of these compounds has also been attempted by deleting *adhE* (encoding alcohol dehydrogenase) and *frdA* (encoding fumarate reductase). Although the production of 508 509 ethanol and succinate was substantially reduced in the triple mutant ($\Delta ldhA\Delta adhE\Delta frdA$), PDO production was only marginally increased compared with that of the single $\Delta ldhA$ 510 mutant. The failure of the triple mutant ($\Delta ldhA\Delta adhE\Delta frdA$) to increase PDO production was 511 512 mainly due to an increase in BDO production. Disruption of the BDO pathway encoded by the *bud* operon (*budA*, *budB*, and *budC*) has also been studied (Fig. 5). Although partially 513 effective, individual inactivation of each of the three genes did not successfully decrease 514 BDO production (Oh et al., 2012a; Wu et al., 2013). Deletion of the entire bud operon 515 completely eliminated BDO synthesis (Kumar et al., 2016), but seriously hampered cell 516 growth and glycerol consumption. Consequently, no increase in PDO production was resulted. 517 Wu et al. (2013) attempted to decrease BDO production while increasing PDO production by 518 519 inserting the *fdh* gene from *Candida boidinii* (NADH-forming formate dehydrogenase) into 520 the budC locus. This protocol increased the PDO titer (62.3-72.2 g/L) and yield (0.47-0.57 mol/mol) and decreased the production of BDO by 52.2% and formic acid by 73.4%. 521 Acetic acid is one of the most toxic metabolites, accumulating in large quantities 522 523 during glycerol fermentation by K. pneumonaie (Celińska, 2012). The toxic effects of acetate is realized even at the low concentration of 20 mM. Physiologically, acetate is a preferred 524 metabolite for many Enterobacter sp. because its production is accompanied by ATP 525

generation. In a typical bioreactor run for PDO production by K. pneumoniae, the 526 accumulation of acetic acid to >300 mM by the end is not uncommon, which often 527 completely stops glycerol fermentation. Acetate is produced by Pta-Ack and PoxB when 528 529 pyruvate formation is faster than its consumption, which is so-called 'overflow metabolism' (Fig. 6) (De Mey et al., 2007). In K. pneumoniae, the contribution of the poxB gene is not 530 significant and its deletion does not much affect acetate production. In comparison, deletion 531 532 of the *pta-ack* genes greatly reduces cell growth and results in high accumulation of pyruvate and pyruvate-based metabolites (Ko et al., 2017). Extensive studies to reduce or eliminate 533 534 acetate production have been conducted in E. coli. Compared to poxB, ackA and/or pta were much more significant in reducing acetate production. However, deletion of the latter genes 535 was not beneficial because the glycolytic flux and cell yield were seriously reduced and the 536 537 accumulation of other byproducts such as pyruvate, lactate, and formate was greatly increased (Causey et al., 2004; De Mey et al., 2007). Several indirect approaches to 538 decreasing pyruvate/acetate accumulation were investigated: stimulation of the anaplerotic 539 pathway by overexpression of pyruvate carboxylase, PEP carboxylase, or PEP carboxykinase 540 (Abdel-Hamid et al., 2001; Causey et al., 2004; Chao and Liao, 1993; Sanchez et al., 2005); 541 overexpression of acetyl-CoA synthetase (ACS); increase of the TCA cycle throughput by 542 deleting the transcriptional repressor arcA gene; and stimulation of the glyoxylate shunt by 543 544 deleting the transcriptional repressor *iclR* gene (De Mey et al., 2007; Jeong et al., 2004; Lin et 545 al., 2006). Moreover, the use of a mutant pyruvate dehydrogenase which is less sensitive to inhibition by NADH has also been examined (Kim et al., 2008). In E. coli, these approaches 546 proved to be effective to varying extents under properly selected culture conditions. However, 547 548 they have not yet been fully investigated in K. pneumoniae.

To eliminate byproduct formation, reduction of glycerol flux through the oxidativepathway at the glycerol node has also been attempted. The deletion of glycerol

551 dehydrogenase (dhaD) and/or dihydroxyacetone kinase (dhaK) resulted in decrease of the oxidative flux and increase of the conversion yield of glycerol to PDO. In addition, 552 production of the byproducts lactate, ethanol, and succinate but not acetate decreased 553 significantly, even without blocking the pathways leading to these byproducts (Seo et al. 554 2009; Horng et al. 2010). However, cell growth and glycerol consumption rate were also 555 reduced. Interestingly, in these mutant strains, deletion of *dhaT* has almost no effect on PDO 556 production. In the absence of *dhaT*, expression of an HOR, highly homologous to YqhD, was 557 upregulated (see section 2.2.4). 558

559 The theoretical maximum yields of PDO on glycerol are 0.875 and 0.844 mol/mol under aerobic and anaerobic conditions, respectively (Celińska, 2012). However, 560 experimentally, the yields are reported between 0.35 and 0.65 mol/mol (Saxena et al., 2009; 561 562 Sun et al., 2008). The low yield is mainly attributed to the accumulation of unnecessary byproducts and inefficient metabolism for generating reducing equivalents (NADH) and ATP. 563 If acetyl-CoA is fully oxidized through the TCA cycle (without byproducts formation) and 564 the activity of the electron transport chain (ETC) oxidizing NADH is properly controlled, a 565 significant increase in PDO yield is expected. Attaining this goal requires extensive pathway 566 engineering of cellular metabolism. In bioreactor operation, the aeration rate should be 567 optimized. Oxygen regulates gene expression and the enzymatic reactions of glycerol 568 metabolism and determines the diversion of DHA to the PP pathway. Regeneration of NADH 569 570 by the TCA cycle and its oxidation by ETC are also controlled by oxygen levels.

571 **3.2 3-Hydroxypropionic acid**

As with PDO, 3-HP is an important platform chemical and has been selected by the US Department of Energy (DOE) as one of the top 12 chemicals obtainable from biomass. 3-HP is a C3 bifunctional molecule and a structural isomer of lactic acid (Werpy and Petersen, 2004). 3-HP can be converted into a variety of useful chemicals such as acrylic acid and acryl

576	amide (Pina et al., 2011; Kumar et al., 2013a). 3-HP can be synthesized by chemical routes
577	from acrylic acid, 3-propiolactone, 3-hydroxypropionitrile, allyl alcohol, vinyl acetate, and
578	PDO. However, none of the chemical processes is commercially feasible at present due to the
579	high cost of the starting materials, toxicity of intermediates, and/or the environmental
580	incompatibility of the processes (Jiang et al., 2009; Pina et al., 2011; Kumar et al., 2013a).
581	Many microorganisms, including both prokaryotes and eukaryotes, can naturally synthesize
582	3-HP as either an intermediate or end product through a range of metabolic pathways.
583	However, the production of 3-HP by these native microorganisms is too low to be
584	commercially meaningful (Jiang et al., 2009; Kumar et al., 2013a).
585	In studies pursuing commercial production of 3-HP, glycerol is first converted to 3-
586	HPA by GDHt. 3-HPA then is converted to 3-HP via the coenzyme A (CoA)-dependent or
587	CoA-independent pathway (Fig. 3). In the CoA-independent pathway, 3-HPA is oxidized to
588	3-HP by aldehyde dehydrogenase (ALDH); in the CoA-dependent pathway, 3-HPA is
589	converted to 3-HP via 3-HP-CoA and 3-HP-phosphate. In K. pneumoniae, both the CoA-
590	dependent and CoA-independent pathways are present. However, the titer of 3-HP produced
591	by the CoA-dependent pathway is low as <5.0 g/L (Luo et al., 2011 and 2012). High
592	production of 3-HP requires the disruption of oxidoreductases (e.g., DhaT, YqhD) and the
593	overexpression of a proper aldehyde dehydrogenase (ALDH). Ashok et al. (2013a) developed
594	a recombinant <i>K. pneumoniae</i> in which <i>dhaT</i> and <i>yqhD</i> were deleted, and the homologous
595	<i>puuC</i> gene encoding the NAD ⁺ -dependent γ -glutamyl- γ -aminobutyraldehyde dehydrogenase
596	was overexpressed. The recombinant <i>K. pneumoniae</i> produced 3.8 g/L 3-HP in 12 h of flask
597	culture, but only under appropriate aeration (i.e., microaerobic) conditions. Under anaerobic
598	conditions, PDO (instead of 3-HP) was obtained as the main product, even though two
599	oxidoreductases were disrupted and $puuC$ was highly overexpressed. In contrast, highly
600	aerobic conditions produced a high cell mass without much accumulation of either 3-HP or

601	1,3-PDO. In a glycerol-fed batch bioreactor experiment under a constant dissolved oxygen
602	(DO) concentration of 5% (considered "proper" microaerobic condition), the recombinant K .
603	<i>pneumoniae</i> $\Delta dhaT \Delta yqhD$ overexpressing both PuuC and DhaB produced >28 g/L 3-HP in
604	48 h, with a yield of >40%. In contrast, the same strain produced only 10.5 g/L 3-HP when
605	cultivated under the higher 10% DO conditions. Several important conclusions regarding 3-
606	HP production by K. pneumoniae were made, as follows. First, in addition to DhaT and
607	YqhD, K. pneumoniae has more unidentified oxidoreductases that can produce PDO from 3-
608	HPA. Second, to produce 3-HP rather than PDO, proper aeration to regenerate NAD^+ is
609	essential. Third, excessively high aeration decreases the expression of the Dha regulon and
610	synthesis of coenzyme B ₁₂ , an essential cofactor for GDHt catalysis.
611	To eliminate PDO production even under limited aeration conditions, Ko et al.
612	attempted to identify and disrupt other potential PDORs from K. pneumoniae (Ko et al.,
613	2015). A mutant strain devoid of <i>dhaT</i> , <i>yqhD</i> , <i>ahpF</i> , and <i>adhE</i> genes was developed;
614	however, the mutant neither eliminated 1,3-PDO production nor increased 3-HP production.
615	Again, this result suggests that K. pneumoniae has more unidentified oxidoreductases, and
616	thus, the complete elimination of 1,3-PDO production during 3-HP production is highly
617	challenging. To address the important problem, of NAD^+ regeneration and coenzyme B_{12}
618	production, Ashok et al. (2013b) studied anaerobic respiration using nitrate as an external
619	electron acceptor. They attempted to regenerate NAD ⁺ from NADH by nitrate reduction
620	while maintain anaerobic conditions to synthesize and stabilize the oxygen-sensitive
621	coenzyme B_{12} . Disruption of the <i>glpK</i> gene (encoding for glycerol kinase) was also necessary
622	because with $glpK$ intact, the rate of anaerobic respiration was too fast, and most of the
623	glycerol was metabolized through GlpK to produce biomass, as in the case of highly aerobic
624	cultivation. The anaerobic glycerol fermentation in the presence of nitrate was successful:
625	1.74 g/L 3-HP was obtained from a 12-h flask culture and 22 g/L 3-HP was produced in a 48-

626 h fed-batch bioreactor culture (Ashok et al., 2013b) (Table 3). However, disadvantages were noted, including the requirement for large amounts of nitrate and the toxic effect of nitrite 627 generated as a reduced intermediate of nitrate. In a recent study, Li et al. (2016) optimized a 628 629 promoter for ALDH expression, culture conditions, and metabolic flux to achieve high-level production of 3-HP in their K. pneumoniae isolate. One recombinant strain, K. pneumoniae 630 (pTAC-puuC) expressing puuC under the IPTG-inducible tac promoter, produced 73.4 g/L 3-631 632 HP in a bioreactor with a molar yield of glycerol of 0.52 and productivity of 1.53 g/L h. Further, elimination of the *ldh1*, *ldh2*, and *pta* genes elevated the titer and molar yield to 83.8 633 634 g/L 3-HP and 0.54 mol/mol, respectively, with decreased cell growth and productivity. This titer of 3-HP is the highest produced by K. pneumoniae to date. 635

As another approach to addressing the problem associated with NAD⁺ regeneration 636 637 and coenzyme B₁₂ synthesis in 3-HP production, co-production of 3-HP and PDO has been investigated. The simultaneous production of 3-HP and PDO can eliminate the dependency 638 on oxidative metabolism of glycerol (and electron transport chain), as the cofactor required 639 for 3-HP production is regenerated by PDO production or vice versa. Furthermore, the 640 problems associated with the expression of genes for the assimilation of vitamin B_{12} and 641 glycerol can be alleviated substantially because co-production can be performed under 642 anaerobic or microaerobic conditions (Kumar et al., 2012, 2013b). Ashok et al. (2011) 643 developed a recombinant strain of K. pneumoniae DSMZ by overexpressing ALDH and 644 645 deleting *dhaT*. With *dhaT* intact, PDOR activity was too high compared with that of ALDH, even with overexpression of the latter enzyme from a multicopy plasmid. The recombinant K. 646 pneumoniae DSMZ accumulated 16.0 g/L 3-HP and 16.8 g/L PDO in 24 h, and the 647 648 cumulative yield of these two metabolites on glycerol was 51% (Table 3). Huang et al. (2012) studied the same co-production using their K. pneumoniae isolate overexpressing ALDH and 649 obtained 24.4 3-HP and 49.9 g/L PDO, respectively, after 24 h under anaerobic conditions. 650

651 The cumulative molar yield of the two metabolites reached 0.61 (0.18 for 3-HP and 0.43 for PDO). In a following study, they showed that the maximum titer of 48.9 g/L 3-HP (along 652 with 25.3 g/L PDO) was achieved at 1.5 vvm, while the highest concentration of 38.4 g/L 653 654 PDO (along with 16.6 g/L 3-HP) was obtained at 0.1 vvm (Huang et al., 2013a). During 3-HP production from glycerol by K. pneumoniae, the accumulation of such 655 byproducts as lactic acid, ethanol, acetic acid, formic acid, and 2,3-BDO is a serious 656 657 consideration, as in the case of 1,3-PDO production (Ashok et al., 2013a,b; Li et al., 2016). Deletion of *ldhA* reduces lactic acid formation and increases the 3-HP yield. However, 658 659 deletion of other genes such as *adhE*, *pfl*, *budBAC*, and *pta-ack* had little desired effect and seriously hampered cell growth and glycerol assimilation (unpublished data). Even with the 660 co-production of 3-HP and 1,3-PDO, where oxidative glycerol assimilation was greatly 661 662 reduced, accumulation of these byproducts was a serious drawback (Ko et al., 2017). To prevent the formation of lactic acid along with other byproducts, Kumar et al. (2013b) 663 employed resting cells of recombinant K. pneumoniae J2B overexpressing ALDH and devoid 664 of lactate dehydrogenase (*ldhA*). For this strain, final titers of 3-HP and PDO of 22.7 and 23.5 665 g/L, respectively, were obtained without lactic acid accumulation. The cumulative product 666 yield increased to 0.77. 667 In addition to K. pneumoniae, E. coli has been extensively studied for 3-HP 668

669 production from glycerol. Several successful results have been reported by Samsung Ltd. Co.

670 Korea. One study (Chu et al., 2015) used an engineered strain (*E. coli* W3110 ΔackA-pta

671 *ΔyqhD dhaB mutant gabD4*) harboring an active ALDH mutant (designated as GabD4) from

672 *Cupriavidus necator*, producing 71.9 g/L 3-HP with a productivity of 1.8 g/L h. However,

673 unlike *K. pneumoniae*, *E. coli* does not synthesize vitamin B₁₂ naturally, and it was necessary

to add the coenzyme externally. Furthermore, a significant amount of glucose for cell growth

675 was required because for *E. coli*, glycerol is a less-preferred carbon source than glucose.

676 Consequently, the researchers at Samsung initially grew cells to a high density (40-300 OD_{600}) on glucose and then produced 3-HP by feeding glycerol (Chu et al., 2015; Kim et al., 677 2014a). In comparison, K. pneumoniae can produce a high concentration of 3-HP growing 678 purely on glycerol at a much lower cell concentration (OD₆₀₀, <20) (Huang et al., 2013a,b; Li 679 et al., 2016). Together, the use of glucose for cell proliferation, exogenous addition of 680 expensive cofactor vitamin B_{12} , and low cumulative yield of 3-HP (on glucose plus glycerol) 681 682 increase the production cost. Thus, K. pneumoniae can be considered a better biocatalyst than E. coli. However, for the commercial production of 3-HP, many other factors are important, 683 684 including biosafety, process stability, and downstream processing. Further studies to assess the potential of these strains as hosts for 3-HP production are needed. 685

686 **3.3 2,3-Butanediol**

BDO has many applications in the pharmaceutical, biomedical, and other chemical industries for the production of printing inks, perfumes, fumigants, spandex, moistening and softening agents, and plasticizers (e.g., cellulose nitrate, polyvinyl chloride, polyacrylates) (Celińska and Grajek, 2009; Ji et al., 2011). BDO can also be used as an antifreezing agent and octane booster for petrol as is or can be converted to useful derivatives such as 1,3butadiene, and diacetyl and methyl ethyl ketone.

Unlike 3-HP and PDO, BDO is a product of oxidative metabolism. Of the three 693 694 stereoisomers of BDO, K. pneumoniae mainly produces the meso form (Ji et al., 2011). BDO 695 is synthesized from pyruvate via α -acetolactate and acetoin. The C5 intermediate, α acetolactate, is formed from two molecules of pyruvate by self-condensation catalyzed by α -696 697 acetolactate synthase (ALS; *budB*). α -Acetolactate is decarboxylated to acetoin by α -698 acetolactate decarboxylase (*budA*), and acetoin is reduced to BDO by 2,3-butanediol dehydrogenase/acetoin reductase (budC) using NADH as a reductant. In the presence of 699 700 oxygen, α -acetolactate is spontaneously decarboxylated to diacetyl, further reduced to acetoin

701 by action of diacetyl reductase, and then reduced to BDO. α -Acetolactate is also the precursor 702 of branched-chain amino acids (valine, leucine, and isoleucine); thus, elimination of the bud operon can arrest K. pneumoniae growth when cultured on glycerol minimal medium (Kumar 703 704 et al., 2016). In the anaerobic process, one NADH molecule is required for the conversion of two pyruvates to one BDO, while the aerobic process requires two NADH (Fig. 5). The 705 706 enzyme DhaD may also contribute to BDO formation (see Section 2.2.1). In the BDO pathway itself, the conversion of two pyruvates to BDO regulates or is regulated by the 707 intracellular NADH/NAD⁺ ratio, similar to other fermentative pathways. Another important 708 709 role of the BDO pathway is to reduce intracellular acidification by converting acids to the neutral metabolite, BDO (Celińska and Grajek, 2009; Ji et al., 2011). 710 711 Although BDO is one of the main byproducts of glycerol fermentation by *K*. 712 pneumoniae (Petrov and Petrova, 2009; Durgapal et al. 2014), its production from glycerol 713 has not been studied extensively. In few studies using glycerol as the carbon source, Petrov and Petrova (2009) focused on the effects of pH and aeration with their K. pneumoniae 714 715 isolate G31. BDO production was greatest at alkaline initial pH when tested at a pH range of 5–8. Without pH control a sharp pH drop occurred initially due to the production of organic 716 717 acids, and this pH drop triggered the synthesis of BDO, which is considered the only noninhibitory compound produced by the oxidative metabolism of glycerol. Once neutral pH is 718 719 restored with the rising concentration of BDO, catabolism shifted toward the production of 720 ATP-generating acetic acid. The amount of BDO generated in each cycle (pH drop and neutralization) was dependent on the extent of the pH drop. In one study, a BDO 721 concentration of 49.8 g/L was reached when the initial pH was set at 8.0 and no pH control 722 was given during the rest of fermentation. PDO was the major byproduct (11.6 g/L) (Table 4) 723 and its production decreased by exclusion of Co^{2+} , an essential component of coenzyme B_{12} . 724 They also found that intensive aeration led to a significant increase in BDO production; an 725

726	increase in aeration rate from 1.1 to 2.2 vvm enhanced the yield from 0.38 to 0.46 mol/mol
727	and productivity from 0.24 to 0.36 g/L h. BDO production was also increased by forced pH
728	fluctuations of the culture medium with discrete ΔpH values (1.0, 2.0, and 3.0) at
729	predetermined time intervals (6, 12, and 24 h). The highest BDO concentration (70.0 g/L)
730	was produced by increasing the pH by one unit ($\Delta pH = 1.0$) every 12 h (Petrov and Petrova,
731	2010). The mechanism behind the effect of this peculiar pH change on BDO production has
732	not been elucidated. Although BDO is readily produced from glycerol by K. pneumoniae, the
733	volumetric productivity is low due to the long fermentation time (generally >150 h). Huang et
734	al. (2013b) isolated two Klebsiella strains that produce BDO plus acetoin with a total yield of
735	0.44–0.45 (g/g). One isolate utilized 64.5 g/L glycerol in 78 h and produced 24.9 g/L BDO,
736	3.9 g/L acetoin, and 3.2 g/L ethanol. Again, the volumetric productivity was not high.
737	Metabolic engineering efforts with K. pneumoniae to improve BDO production from
738	glycerol are limited. Using glucose as carbon source, Kim et al. (2014b) reported that
739	overexpression of the BDO pathway (budA, budB) in the ldhA deficient K. pneumoniae strain
740	KCTC2242 improved BDO production and resulted in 90 g/L BDO, with a productivity of
741	2.75 g/L. h. Because both glucose and glycerol are converted through the common
742	intermediate pyruvate, BDO production from glycerol is also expected to increase by
743	amplification of the bud operon. Further work with glycerol as carbon source is required to
744	prove this effect. Recently, co-production of BDO and PDO from glycerol has also been
745	attempted (Park et al., 2017). In glycerol fermentation, both alcohols always appear
746	simultaneously and co-production seems to be beneficial for cells to maintain redox balance
747	during anaerobic growth. Park et al. (2017) found that by using the K. pneumoniae mutant
748	deficient of <i>ldhA</i> and <i>mdh</i> , 50.1 g/L BDO was produced along with 63.8 g/L PDO in 75 h.
749	The ratio of the two alcohols varied according to aeration; with increasing aeration, the ratio
750	of [BDO]/[PDO] increased as observed by Petrov and Petrova (2010). In the 5,000 L pilot-

scale fed-batch fermentation using crude glycerol, they could produce 114 g/L diols (70 g/L

PDO and 44 g/L BDO) with the yield of 0.60 diols/glycerol (g/g) and a productivity of 2.2

753 g/L. h of diols. If separation is not an issue, co-production of the two diols can be a good

754 option.

755 **3.4 Lactic acid**

Lactic acid has applications in the food, chemical, cosmetic, and pharmaceutical industries. Lactic acid is a monomer that can be polymerized to yield the biodegradable plastics polylactic acid (PLA) and poly(3-hydroxybutyrate-*co*-lactate). Two optical isomers of lactic acid, D- and L-lactic acid, are produced in racemic mixtures depending on the chemical route used. Microbial fermentation can yield optically pure isomers. Currently, nearly all commercial lactic acid comes from microbial fermentation (Abdel-Rahman et al.,

762 2013; Abdel-Rahman and Sonomoto, 2016).

763 Because of the availability of highly efficient lactic-acid-producing microbes derived from E. coli, lactic acid bacteria, and even acid-tolerant yeasts (Abdel-Rahman et al., 2013), 764 765 K pneumoniae has received little attention as a lactic acid producer. However, as previously described, lactic acid is a major byproduct of glycerol fermentation by K. pneumoniae, and its 766 potential as a producer is high. This bacterium contains metabolic pathways for the 767 production of both L- and D-isomers (Fig. 7). They are synthesized through the pyruvate 768 769 and/or methylglyoxal pathways in K. pneumoniae (Ashok et al., 2011). When produced from 770 pyruvate, lactate dehydrogenase (*ldhA*) reduces pyruvate using NADH as reductant. In the methylglyoxal route, DHAP is converted to methylglyoxal, which can be transformed to both 771 D- and L-lactic acid. For the production of L-lactic acid, methylglyoxal is reduced to L-772 773 lactaldehyde, which is subsequently oxidized to L-lactic acid. Using glutathione, D-lactic acid is obtained through simultaneous aldehyde group oxidation and keto group reduction of 774 775 methylglyoxal (Booth et al., 2003; Booth, 2005). The methylglyoxal route has several

776 disadvantages. Methylglyoxal is highly toxic, and its accumulation can severely impair other 777 metabolisms and even lead to cell death. Second, the route is energetically inefficient, with an ATP yield of -1. The L- or D-lactic acid product can be converted back into pyruvate by 778 779 respiratory lactate dehydrogenase (Mazumdar et al., 2010, 2013). Several studies showing the potential of K. pneumoniae as a lactic acid producer are 780 available (Table 4). For example, K. pneumoniae DSMZ is reported to accumulate 22.7 g/L 781 lactic acid in 24 h during co-production of 3-HP and PDO from glycerol. The titer of lactic 782 acid was more than that of any of the targeted products, and the yield of lactic acid on 783 784 glycerol alone was 0.31 (Ashok et al., 2011). Similarly, fed-batch cultivation of K. 785 pneumoniae BLh-1 under oxygen-limited conditions gave lactic acid as the main product, 786 outcompeting PDO; 59 g/L lactic acid was produced in 40 h when crude glycerol from 787 biodiesel synthesis without any purification was used (Rossi et al., 2013). Durgapal et al. 788 (2014) also found that K. pneumoniae J2B and DSMZ strains produced lactic acid at 37.0 and 47.8 g/L, respectively (Table 4); in both strains, lactic acid levels surpassed those of the target 789 790 product PDO. Recently, Feng et al. (2014) engineered K. pneumoniae ATCC25955 by overexpressing *ldhA* and deleting *dhaT* and *yqhD* (recall that these two genes are mainly 791 792 responsible for PDO production). They found that the recombinant produced 142.1 g/L of optically pure D-lactic acid from glycerol in fed-batch cultivation under microaerobic 793 794 conditions (2.5 vvm and 400 rpm). This result is the highest lactic acid concentration 795 produced from glycerol reported to date. E. coli has also been engineered for the production of D- and L-lactate from glycerol (Mazumdar et al., 2010, 2013). For diverting glycerol flux 796 toward L-lactic acid in E. coli, the chromosomal copy of D-lactate dehydrogenase was 797 798 replaced with Streptococcus bovis L-lactate dehydrogenase (Mazumdar et al., 2013). Moreover, gene encoding *lldD* (respiratory L-lactate dehydrogenase) was inactivated to 799 800 prevent the consumption of L-lactic acid, and the methylglyoxal route ($\Delta mgsA$) was blocked

801 to prevent production of toxic methylglyoxal and formation of racemic mixtures of D- and Llactic acid (Fig. 7). Furthermore, the respiratory route for glycerol assimilation was 802 overexpressed (glpK and glpD) to improve the ATP yield of the metabolic pathway through 803 804 transfer of electrons from glycerol-3-phosphate to oxygen (i.e., oxidative phosphorylation). 805 The engineered E. coli strain ($\Delta pflB \Delta pta \Delta adhE \Delta frdA \Delta mgsA \Delta lldD \Delta ldhA:: ldhA_glpK-$ _glpD) produced 50 g/L of L-lactate from 56 g/L of crude glycerol, at a yield 93% of the 806 theoretical maximum and with high optical (99.9%) and chemical (97%) purity (Mazumdar et 807 al., 2013). The approaches proven successful in *E. coli* should be applicable to *K*. 808 809 pneumoniae. These studies, along with the results by Feng et al. (2014), suggest that K. pneumoniae is a good host for the production of D- and L-isomers of lactic acid, with high 810 optical purity, from glycerol. 811

812 **3.5 Ethanol and other metabolites**

K. pneumoniae has also been studied for the production of ethanol, succinic acid, 2-813 ketogluconic acid, catechol, *cis, cis*-muconic acid and 2-butanol although its potential for such 814 production has not been fully explored (Cheng et al., 2013; Jung et al., 2015; Oh et al., 2014; 815 Wang et al. 2015; Wei et al., 2013). Of these chemicals, ethanol production has been 816 relatively well studied. Ethanol is a renewable energy source and is widely used as a fuel 817 additive for partial gasoline replacement. Currently, commercial ethanol is produced from the 818 819 fermentation of sugar and starch. However, glycerol is one of the cheapest substrates for 820 ethanol production, estimated as nearly 40% cheaper than sugar-based production (Yazdani 821 and Gonzalez, 2007). This factor has encouraged the development of microbial strains for fermentative conversion of glycerol to ethanol. The metabolic route by which ethanol is 822 823 synthesized from glycerol is shown in Fig. 6. Oh et al. (2011, 2012b) constructed a mutant 824 strain of K. pneumoniae (termed GEM167) through γ -irradiation. PDO synthesis in this mutant strain was lower than that of wild type (0.2 vs. 7.93 g/L, respectively), while ethanol 825

826 accumulation was higher (8.6 vs. 1.1 g/L, respectively). The mechanism underlying these changes remains unclear. Introduction of the pyruvate decarboxylase (pdc) and aldehyde 827 dehydrogenase (*adhII*) genes into mutant strains lacking lactate dehydrogenase (*ldhA*) 828 829 resulted in the production of 31.0 g/L (673.9 mM) ethanol, with a yield of 0.89 (mol/mol) and productivity of 1.2 g/L h (Oh et al., 2012b), the highest level of ethanol production from 830 glycerol reported to date. More extensive studies including estimation of the maximum 831 832 achievable titer and microbial tolerance to high-concentration ethanol, both of which are key factors for commercialization, are required to assess the potential to produce ethanol from 833 834 glycerol by the use of K. pneumoniae.

4. Challenges to bioconversion of glycerol using *K. pneumoniae* as a biocatalyst

The design of microbial cell factories is gaining unprecedented momentum as 836 837 metabolic engineering is progressively aided by advances in synthetic biology and multi-838 omics analyses. Several bio-based chemicals have already been launched successfully in the marketplace, and others are in the pipeline. For a bulk chemical to be considered for 839 commercial production, the production process should achieve a product concentration of 100 840 g/L, carbon yield of 0.50 (g/g), and volumetric productivity of 2.0 g/L/h. The aforementioned 841 products produced by K. pneumoniae from glycerol are no exception. Challenges to the use 842 of K. pneumoniae as a microbial cell factory using glycerol as a carbon source are described. 843

844 **4.1 Pathogenicity**

The pathogenicity of *K. pneumoniae* is one of the major obstacles to its commercial application. *K. pneumoniae* is a Bio-safety Level 2 microorganism and causes nosocomial and urinary tract infections. Moreover, capsulated *K. pneumoniae* cells are difficult to separate from culture broth, complicating the downstream processing. The biotechnological application of this bacterium will require attenuation of its pathogenicity and reduction of biosafety concerns. *K. pneumoniae* produces a number of virulence factors that contribute to

its pathogenicity, including lipopolysaccharide (LPS), capsular antigens, fimbrial adhesins,
siderophores, and O antigens (Albertí et al., 1993; Fang et al., 2004; Podschun and Ullmann
1998). The mechanism by which this bacterium causes disease is still obscure because most
studies have been conducted on a limited number of virulence factors (El Fertas-Aissani et al.,
2013).

In the last decade, efforts have been made to overcome the pathogenicity of K. 856 857 pneumoniae (Shrivastav et al., 2013), suggesting hope for the generation of nonpathogenic K. pneumoniae. The two major pathogenic determinants in K. pneumoniae are the capsular 858 859 polysaccharides (CPS), which cover the entire bacterial surface, and LPS, expressed on the outer bacterial membrane. LPS consists of lipid A, a core oligosaccharide, and an O-antigenic 860 polysaccharide. In a highly virulent strain of K. pneumoniae, mutations in genes involved in 861 862 LPS biosynthesis (*waaC*, *waaF*, and *wabG*) resulted in a decrease in bacterial colonization and virulence (Izquierdo et al., 2003; Jung et al., 2013). Mutant strains devoid of the outer 863 core LPS were also found to be deficient in the "cell-attached" capsular polysaccharides 864 covering the bacterial surface. These mutations abolished the highly virulent characteristics 865 of pathogenic K. pneumoniae when tested in different animal models. Furthermore, these 866 mutants were more sensitive to several hydrophobic compounds than were wild-type strains. 867 Reintroduction of the waaC, waaF, and wabG genes into K. pneumoniae rescued the 868 869 pathogenic properties, confirming the role of these genes in pathogenicity.

Another important study was conducted by Lin et al. (2012), in which eight highly conserved residues of the MagA protein (G308, G310, G334, G337, R290, P305, H323, and N324) encoded by the mucoviscosity-associated gene A (*magA*) of *K. pneumoniae* were subjected to site-directed mutagenesis. The *magA* gene contributes to the biosynthesis of K1 capsular polysaccharide (CPS), which correlates with pathogenic phenotypes including mucoviscosity, serum and phagocytosis resistance, and virulence. Alanine substitutions at

R290A or H323A abolished MagA function, with annihilation of CPS production, serum
resistance, and virulence. Some *K. pneumoniae* strains showing less pathogenicity were also
isolated. For example, *K. pneumoniae* strain J2B produced less LPS and demonstrated a high
sensitivity to many antibiotics and lower pathogenicity. Moreover, its biomass was readily
separated from fermentation broth by centrifugation (Arasu et al., 2011). Future studies
should include further engineering of these less pathogenic strains to make them completely
nonpathogenic, followed by careful evaluation of their toxicity.

883 **4.2 Complex glycerol metabolism**

884 Although glycerol metabolism has been extensively studied, the mechanism and regulation of its gene expression in K. pneumoniae have not been fully elucidated. The 885 diversity of glycerol metabolism should be a great advantage for K. pneumoniae survival in 886 887 different environments and for its biotechnological utilization as well. However, a high level of complexity poses difficulty in the engineering of a strain to suit our purposes. Metabolic 888 complexity is likely responsible in part for the fact that, despite serious attempts, the molar 889 890 yield of many important products such as PDO and 3-HP is still far below the theoretical maximum. Efforts based on our current knowledge, such as the overexpression of DhaB, 891 892 DhaT, YqhD and AldH and inactivation/deletion of GlpK, DhaD, DhaK and DhaKLM, were not satisfactory (Maervoet et al. 2011; Kumar et al. 2013a; Kumar et al., 2016). Another 893 894 challenge is to understand how the ratio of glycerol distribution between respiratory and 895 fermentative routes is controlled. Because the $K_{\rm M}$ of glycerol kinase toward glycerol is low, it has been speculated that a major fraction of glycerol flows through the respiratory route when 896 oxygen is present. However, some experimental results do not support this hypothesis. For 897 898 example, deletion of *glpK* does not decrease the glycerol flow through the oxidative pathway in the presence of oxygen (Ashok et al. 2013a). In fact, wild-type and mutant ($\Delta g l p K$) strains 899 900 of K. pneumoniae demonstrate similar cell growth, glycerol consumption, and PDO

901 production profiles under (micro)aerobic conditions. These observations suggest the presence902 and/or involvement of other kinases.

Another challenging issue for the use of K. pneumoniae is its inefficient TCA cycle 903 904 (Cabelli, 1955). Several genes, including isocitrate dehydrognease (*icd*), fumarase (*fumA*), and malate dehydrogenase (mdh), in K. pneumoniae are significantly less transcribed than 905 those in E. coli (unpublished data). Furthermore, the activity of isocitrate dehydrogenase was 906 907 more than sevenfold lower in K. pneumoniae than in E. coli. If the TCA cycle is inefficient, achieving a high cell density, which is essential for improving the productivity of target 908 909 metabolites, is difficult. Moreover, achieving high-yield generation of NADH with little consumption of glycerol carbon is difficult. Furthermore, the overflow metabolism that 910 911 produces highly toxic acetate becomes more serious. These problems have been well 912 documented in PDO production. In a recent study, Kumar et al. (2016) attempted to increase 913 PDO production yield by blocking the lactate and BDO pathways, while diverting carbon flux through the TCA cycle for efficient regeneration of NADH. However, this effort resulted 914 915 in the detrimental accumulation of pyruvate, acetate, and ethanol, causing low NADH regeneration and low PDO production. Neither increased aeration nor the addition of good 916 nitrogen sources (complex nitrogen sources) alleviated the accumulation of these 917 intermediates or activated the TCA cycle. Furthermore, the accumulation of pyruvate quickly 918 919 terminated glycerol assimilation. K. pneumoniae has a well-established BDO production 920 pathway that likely compensates for its inefficient TCA cycle, avoiding the overflow metabolism and carbon traffic at the pyruvate node. If this is the case, co-production of BDO 921 along with PDO or 3-HP, rather than single production of PDO or 3-HP, should be more 922 923 appropriate, which suits the nature of K. pneumoniae. In fact, when BDO and PDO were coproduced, more PDO was generated than during PDO production alone. More studies are 924 925 required to gain a better understanding of glycerol metabolism, the slow operation of the

TCA cycle in *K. pneumoniae*, and its effect on the selection of target products and/or
development of suitable strains and processes. Otherwise, it will be difficult to exploit this
potential biocatalyst for commercial manufacturing of valuable chemicals.

929 **4.3** Toxicity of intermediate metabolites and end-products

A general but major challenge to the use of microbial cell factories for producing bulk 930 chemicals is the toxic effect of target products and metabolic intermediates at high 931 932 concentrations. In particular, during the production of PDO or 3-HP from glycerol, the highly toxic intermediate 3-HPA is inevitably generated. Aldehydes generally damage DNA, 933 934 inactivate enzymes, and inhibit DNA synthesis. 3-HPA causes irreversible cessation of metabolic activity and cellular growth, even at very low concentrations of 15-30 mM (Hao et 935 al., 2008; Zheng et al., 2008). Native K. pneumoniae does not accumulate 3-HPA at toxic 936 937 levels during PDO production, but in engineered strains that overexpress DhaB, high 3-HPA 938 accumulation is often experienced. To avoid 3-HPA accumulation, enzymes that consume 3-HPA, ALDH (for 3-HP production), and PDOR (for PDO production) are expressed at 939 940 sufficiently (sometimes excessively) high levels compared with that of DhaB (Lim et al., 2016). Intermediates such as methylglyoxal, DHA, DHAP, and glyceraldehyde-3-phosphate 941 are also known to be toxic. 942

Organic acids and alcohols are toxic at high concentrations. Generally, organic acids 943 944 are more toxic than their corresponding alcohols because they disturb the intracellular pH and 945 have anion-specific effects on metabolism (Chun et al., 2014; Warnecke and Gill, 2005). When accumulated in cell membranes, organic compounds damage membrane integrity and 946 inhibit the activity of membrane-bound enzymes. In K. pneumoniae, cell growth and glycerol 947 948 assimilation are severely inhibited by high concentrations of PDO or 3-HP. Furthermore, enzymes that play major roles in 3-HP and PDO production, such as DhaB, DhaT, and AldH, 949 950 are inhibited by 3-HP and PDO (Kumar et al. 2013b). Furthermore, K. pneumoniae is

sensitive to 24 g/L acetic acid, 26 g/L lactic acid, and 17 g/L ethanol under aerobic conditions
and 15, 19, and 26 g/L, respectively, under anaerobic condition (Celińska, 2012; Cheng et al.,
2005). The inhibition of pathways by target product(s) and/or unwanted byproducts often
places a limit on the maximum attainable titer of bulk chemicals.

Several studies have reported engineered improvements in microbial tolerance against
biochemicals and biofuels, especially in *E. coli*. Modifications of exporter proteins, heat
shock proteins, membrane composition, and stress responses have been attempted. Moreover, *in situ* recovery of target products has been attempted for hydrophobic targets. Some tolerant
strains have been developed, but unfortunately, without appreciable increases in productivity
(Dunlop, 2011; Dunlop et al., 2011; Jarboe et al., 2011). Such efforts to improve tolerance to
chemicals have not been reported for *K. pneumoniae*.

962 **4.4 Availability of glycerol**

The global market for glycerol is known to be unpredictable and complex. Glycerol in 963 the current market mainly is from the biodiesel industry, and its supply is directly affected by 964 965 the use of biodiesel. In contrast, glycerol demand depends on many industries that use it as feedstock. Between 2001 and 2011, the price varied significantly, from \$0.70/kg to \$1.70/kg 966 for pure glycerol and from \$0.04/kg to \$0.33/kg for crude glycerol (Ayoub and Abdullah, 967 2012; Ciriminna et al., 2014; Kong et al., 2016; Quispe et al., 2013). Nevertheless, glycerol 968 969 prices, for pure as well as crude, have come down significantly since 2004, when many 970 biodiesel production plants initiated operation. The countries producing biodiesel in large quantities (in million liters) include Germany (2900), Brazil (2300), Argentina (2100), France 971 (2000), USA (1200), Spain (1100), Italy (800), Indonesia (700), and Thailand (600) (Kong et 972 973 al., 2016). The Asia-Pacific region overtook Europe as the largest glycerol market in the world in 2009 and has remained so ever since. Market growth is driven by increases in the 974

975 uses for glycerol in such sectors as pharmaceuticals, personal hygiene, and food and beverage976 production (Quispe et al., 2013).

The production of crude glycerol is expected to reach 6 million tons by 2025 977 978 (Ciriminna et al., 2014). At present, the surplus of glycerol from biodiesel and oleochemicals 979 is assumed to be sufficient to meet these new demands. However, the scenario may change in the future with rapid growth of glycerol-based industries and a limited supply of crude 980 981 glycerol. For example, if 3-HP produced from glycerol were used as a substitute for the chemical acrylic acid (global production, 5.85 million tons as of 2014), approximately 7 982 983 million tons of glycerol/year would be needed. If we include other platform chemicals such as PDO, BDO, and ethanol, the demand will be even higher, much more than the expected 984 supply. The growth of glycerol-based biotechnology will eventually be constrained by the 985 986 supply and price of crude glycerol as a carbon source.

987 5. Concluding remarks

Glycerol is an excellent substrate for the production of biochemicals and biofuels. 988 989 Although the biodiesel industry has grown rather slowly in recent years, crude glycerol is still cheap and generated in large quantities. K. pneumoniae has already been successfully 990 991 employed for the production of PDO, 3-HP, BDO, lactic acid, and ethanol from glycerol at industrially attractive levels. Other metabolites such as acetic acid, pyruvic acid, malic acid, 992 993 fumaric acid, alanine, n-butanol, 1,2-propanediol, propionic acid, formic acid, and hydrogen 994 can also be produced from glycerol by K. pneumoniae, although these have not yet been 995 extensively explored. At present, sugar-based bioprocesses are prevalent, and no bioprocesses 996 employing K. pneumoniae and glycerol are commercially available. The pathogenicity of K. 997 pneumoniae and other technical issues are existing challenges to its use. However, advancements in the fields of metabolic engineering, synthetic biology, systems biology, and 998 999 evolutionary engineering will enable us to further exploit the advantageous nature of K.

- 1000 *pneumoniae* as a biocatalyst and glycerol as a carbon source. A better understanding of
- 1001 glycerol metabolism in *K. pneumoniae* and improvements in its performance as a microbial
- 1002 cell factory should make this organism an attractive alternative to current sugar-based
- 1003 methods for the production of biochemicals and biofuels.

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1462 **Figure captions**

- 1463 Figure 1: Transformation of glycerol into a number of valuable chemicals. The solid line
- 1464 represent single step while broken one indicate multiple steps.
- 1465 **Figure 2:** Oxidative metabolism of glycerol in *K. pneumoniae* (Ashok et al., 2011; Kumar et
- al., 2013a). Gene names are shown in italics on arrows. Solid lines indicate single steps;
- 1467 broken lines indicate multiple steps.
- **Figure 3:** Metabolic pathways for the production of PDO and 3-HP from glycerol (Ashok et al., 2011; Kumar et al., 2013a). Gene names are shown in italics on arrows.
- 1470 Figure 4 (A): Arrangement of genes in the *dha* regulon of *K. pneumoniae*. 1, *dhaK* (ATP-
- 1471 dependent dihydroxyacetone kinase); 2, *dhaK3/dhaM* (large subunit of ATP-dependent
- 1472 dihydroxyacetone kinase); 3, *dhaK2/dhaL* (small subunit of ATP-dependent
- 1473 dihydroxyacetone kinase); 4, *dhaK1/dhaK* (medium subunits of ATP-dependent
- 1474 dihydroxyacetone kinase); 5, *orfW* (hypothetical protein); 6, *dhaD* (glycerol dehydrogenase);
- 1475 7, *dhaR* (transcription regulator); 8, *orfW*/CdAT [cob(I)yrinic acid a,c-diamide
- adenosyltransferase]; 9, *orfX/dhaG* (small subunit of reactivating factor of glycerol
- 1477 dehydratase); 10, *dhaT* (1,3-propanediol oxidoreductase); 11, *orfY* (hypothetical protein); 12,
- 1478 *dhaB/B1* (large subunits of glycerol dehydratase); 13, *dhaC/B2* (medium subunit of glycerol
- 1479 dehydratase); 14, *dhaE/B3* (small subunit of glycerol dehydratase); 15, *orfZ/dhaF* (large
- subunit of reactivating factor of glycerol dehydratase); 16, *glpF* (glycerol uptake facilitator);
- 1481 17, 22- *hdeB*; 18, *hdeD* (*hdeBD*-genes related to global regulation); 19, *yfdX*; 20, *kvgS*; and
- 1482 21, *kvgA* (*kvgS* and *kvgA* encode for a two-component signal transduction system) (Celińska,
- 1483 2012; Sun et al., 2003; Wei et al., 2014). (B): Glycerol dehydratase gene arrangements and
- 1484 reactivation factors in different organisms (Martins-Pinheiro et al., 2016).
- **Figure 5:** Metabolic pathway for the synthesis of BDO from glycerol (Ashok et al., 2011;
- 1486 Celińska and Grajek, 2009; Ji et al., 2011). Gene names are shown in italics on arrows. Solid
- 1487 lines indicate single steps; broken lines indicate multiple steps.

- 1488 Figure 6: Biochemical routes for ethanol and acetic acid production from glycerol (Ashok et
- 1489 al., 2011; Kumar et al., 2013a). Gene names are shown in italics on arrow. Solid lines
- 1490 indicate single steps; broken lines indicate multiple steps.
- 1491 Figure 7: Lactic acid production from glycerol via methylglyoxal and pyruvate routes
- 1492 (Ashok et al., 2011; Mazumdar et al., 2010; 2013). Gene names are shown in italics on arrow.
- 1493 Solid lines indicate single steps; broken lines indicate multiple steps.
- **Fig. S1** Effect of deletion of glycerol transporters, *glpF1* and *glpF2*, on glycerol consumption
- and PDO production in *K. pneumoniae* J2B3.

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Key enzymes: 1- Glycerol kinase; 2- Glycerol-3-phosphate dehydrogenase (aerobic); 3- Glycerol-3-phosphate dehydrogenase (anaerobic); 4- Glycerol dehydrogenase; 5- Putative glycerol dehydrogenase; 6- Dihydroxyacetone kinase I; 7- Dihydroxyacetone kinase II; 8- Pyruvate kinase

Fermentative route: C ₃ H ₈ O ₃ + 2NAD ⁺ + ADP + Pi	$\blacktriangleright C_3H_4O_3 + 2NADH + 2H^+ + ATP$
Respiratory route: C ₃ H ₈ O ₃ + NAD ⁺ + Q + ADP + Pi	$\longrightarrow C_3H_4O_3 + NADH + H^+ + QH_2 + ATP$





Key enzymes: 1- Glycerol dehydratase; 2- NADH-dependent PDOR/Propanol dehydrogenase; 3- Hypothetical oxidoreductase; 4- Aldehyde dehydrogenase; 5- Propionaldehyde dehydrogenase; 6- Phosphotransacylase; 7- Propionate kinase

CoA-independent route:
$$C_3H_8O_3 + NAD^+ \longrightarrow C_3H_6O_3 (3-HP) + NADH + H^+$$

CoA-dependent route: $C_3H_8O_3 + NAD^+ + ADP + Pi \longrightarrow C_3H_6O_3 (3-HP) + NADH + H^+ + ATP$
 $C_3H_8O_3 + NAD(P)H + H^+ \longrightarrow C_3H_8O_2 (PDO) + NAD(P)^+ + H_2O$

A)



B)







Key enzymes: 1- Pyruvate oxidase; 2-Pyruvate-formate lyase, putatative formate acetyltransferanse 3- Pyruvate dehydrogenase complex; 4- Phosphoacetyl transferase 5- Acetate kinase; 6 and 7- Bifunctional (acetaldehyde/alcohol) dehydrogenase/putative alcohol dehydrogenase; 8-Formate dehydrogenase, part of the formate hydrogenlyase complex

> PFL route: $C_3H_8O_3 + ADP + Pi$ PDHC route: $C_3H_8O_3 + NAD^+ + ADP + Pi$ C₂H₆O + HCOOH + ATP C₂H₆O + NADH + H⁺ + CO₂ + ATP





Key enzymes :1-D-Lactate dehydrogenase (fermentative); 2-D-Lactate dehydrogenase (respiratory); 3-Methylglyoxal synthase; 4- Glyoxalase I and II; 5-Glycerol dehydrogenase; 6-Aldehyde dehydrogenase isozymes; 7-L-Lactate dehydrogenase (respiratory).

> Pyruvate route: $C_3H_8O_3 + NAD^+ + ADP + Pi$ Methylglyoxal route: $C_3H_8O_3 + NAD^+ + ATP$ $C_3H_6O_3 + NADH + H^+ + ATP$

Supplementary Figure S1



**glpF1* designates the gene located at the respiratory *glpFK* operon, while *glpF2* indicates a separate operon in the fermentative *dha* regulon located between *orfZ/ dhaF* operon and *hdeB* operon (see Fig. 4A). For the growth of *K. pneumoniae* J2B3 which is deleted for *glpK*, *glpD*, *glpA* and *dhaD* (see Fig. 2 for their roles), gluco se was added at 100 mM because the strain cannot grow on glycerol as the sole carbon source (Lama et al., 2017)

Metabolite	Maximum theor	Maximum theoretical yield**		ted/Metabolite	Redox constrained			
			(mo	l/mol)	[NAD(P)H generated &	consumed/ Metabolite]		
	Glucose	Glycerol	Glucose	Glycerol	Glucose	Glycerol		
	(mol/0.5 mol)	(mol/mol)						
1,3-propanediol	0.75	0.88	-1	0	Yes (0 & 2)	Yes (0 & 1)		
$C_3H_8O_2$								
3-hydroxypropionic acid	1.0	1.17	1/0/-1/-0.33	0/1	No (1 & 1;2 & 2)	Yes (1 & 0)		
$C_3H_6O_3$								
2,3-butanediol	0.55	0.64	2	2	Yes/No (2 & 1/2)	Yes (4 & 1/2)		
$C_4H_{10}O_2$								
Ethanol	1.0	1.17	1	1	Yes/No (1/2 & 2)	Yes/No (2/3 & 2)		
C_2H_6O								
Succinic acid	0.86	1.0	0/1	0/1	Yes (1 & 2)	No (2 & 2)		
$C_4H_6O_4$								
Lactic acid	1.0	1.17	1/-1	1/-1	No (1 & 1; 0 & 0)	Yes (2 & 0; 1 & 0)		
$C_3H_6O_3$								
Pyruvic acid	1.2	1.4	1	1	Yes (1/2 & 0)	Yes (2/3 & 0)		
$C_3H_4O_3$								
Acetic acid	1.5	1.75	1/2	1/2	Yes (1/2 & 0)	Yes (2/3 & 0)		
$C_2H_4O_2$								
1-Butanol	0.5	0.58	2	2	No (4 & 4)	Yes (6 & 4)		
$C_4H_{10}O$								

Table 1: Comparison of maximum theoretical yield, ATP yield and redox balance during microbiological production of different metabolites from glycerol and glucose*

*The variation in ATP yield, NADH generated and consumed is due to formation of metabolite through different routes.

**The theoretical yields of metabolites were calculated on the basis of degree of reduction (Dugar and Stephanopoulos, 2011).

Strain	Mode of	Carbon Aeration		Titer	Yield	Productivity	Reference
	operation	source(s)	condition	(g/L)	(mol/mol)	(g/L. h)	
			Wild type strain	ns			
K. pneumoniae M5al	Batch	Glycerol	Anaerobic	58.8	0.53	0.92	Cheng et al. 2007
K. pneumoniae XJPD-Li	Fed-batch	Glycerol	Aerobic	65.3	0.56	3.16	Ma et al. 2009
K. pneumoniae AC 15	Fed-batch	Glycerol	Aerobic	74.1	0.62	3.08	Zheng et al. 2008
K. pneumoniae DSM 2026	Fed-batch	Glycerol	Anaerobic	81.0	0.64	3.00	Liu et al. 2007
K. pneumoniae DSM 2026	Fed-batch	Glycerol	Microaerobic	77.0	0.62	2.70	Liu et al. 2007
K. pneumoniae HR526	Fed-batch	Glycerol	Aerobic	95.4	0.48	1.98	Xu et al. 2009b
K. pneumoniae KG1	Fed-batch	Glycerol	Aerobic	98.8	0.51	3.29	Zhao et al. 2009
			Mutant strain	s			
K. pneumoniae HR526	Fed-batch	Glycerol	Aerobic	102.1	0.52	2.13	Xu et al. 2009b
$\Delta ldhA$							
K. pneumoniae Cu ΔldhA	Fed-batch	Glycerol	Aerobic	102.7	0.50	1.53	Oh et al. 2012a
K. pneumoniae YMU2	Fed-batch	Glycerol	Anaerobic	70.6	0.70	1.1	Zhang et al. 2006
$\Delta aldA$							
K. pneumoniae YMU2	Fed-batch	Glycerol	Anaerobic	72.2	0.57	2.41	Wu et al. 2013

Table 2: Summary of PDO production from glycerol using different K. pneumoniae strains

$\Delta budC_fdh$

Recombinant strains (Overexpression of Genes from the Reductive Branch)							
K. pneumoniae TUAC01_	Batch	Glycerol	Aerobic	15.0	0.61	1.36	Hao et al. 2008
dhaT							
K. pneumoniae KG1_ dhaT	Fed-batch	Glycerol	Aerobic	90.9	0.64	2.16	Zhao et al. 2009
K. pneumoniae ME-	Fed-batch	Glycerol	Microaerobic	67.6	0.62	1.69	Zhu et al. 2009
308_yqhD							
K. pneumoniae $\Delta dhaD$	Batch	Glycerol	Aerobic	7.7	0.53	0.26	Seo et al. 2010
$\Delta dhaK \Delta dhaT_yqhD$							

Strain	Carbon source(s)	Aeration condition	Titer* (g/L)	3-HP yield on glycerol (mol/mol)	Productivity (g/L. h)	Reference
K. pneumoniae DSM 2026 $\Delta dhaT_puuC$	Glycerol	Microerobic	16.0 (16.8)	0.23	0.67	Ashok et al. 2011
K. pneumoniae WM3 pUC18kan_aldHec	Glycerol	Anaerobic	24.4 (49.3)	0.18	1.02	Huang et al. 2012
K. pneumoniae J2B _ KGSADH	Glycerol	Anaerobic	11.3 (15.9)	0.27	0.94	Kumar et al. 2012
K. pneumoniae ∆glpK∆dhaT_puuC	Glycerol	Anaerobic	22.0 (5.9)	0.30	0.46	Ashok et al. 2013a
K. pneumoniae ∆dhaT∆yqhD_dhaB_puuC	Glycerol	Aerobic	28.1 (3.3)	0.40	0.58	Ashok et al. 2013b
K. pneumoniae WM3 pUC18kan_aldHec	Glycerol	Microaerobic	48.9 (25.3)	0.41	1.75	Huang et al. 2013a
K. pneumoniae J2B Δ ldhA _ KGSADH	Glycerol	Microaerobic	22.7 (23.4)	0.35	0.38	Kumar et al. 2013b
E. coli W3110 ∆ackA-pta ∆yqhD_dhaB_mutant gabD4	Glucose & glycerol	Aerobic	71.9	-	1.8	Chu et al. 2015
K. pneumoniae DSM 2026 $\Delta ldh1\Delta ldh2\Delta pta_puuC$ (pTAC)	Glycerol	Microaerobic	83.8 (22.1)	0.54	1.16	Li et al. 2016

Table 3: Different strategies carried out for 3-HP production through fed-batch mode

*The values shown in the bracket are the PDO concentration obtained along with 3-HP.

Strain	Mode of	Carbon	Aeration	Titer	Yield (mol/mol)	Productivity	Reference
	operation	500100(5)	BDO	(g/12)		(g, L, II)	
K. pneumoniae G31	Fed-batch	Glycerol	Microaerobic	49.2	0.37	0.17	Petrov and Petrova, 2009
K. pneumoniae G31	Fed-batch	Glycerol	Aerobic	70.0	0.40	0.47	Petrov and Petrova, 2010
Klebsiella sp. 2	Fed-batch	Glycerol	Aerobic	24.9	0.40	0.34	Huang et al. 2013b
K. pneumoniae SDM	Fed-batch	Glucose	Aerobic	150.0	0.84	3.93	Ma et al. 2009
K. pneumoniae KCTC2242	Fed-batch	Glucose	Aerobic	90.0	0.76	2.75	Kim et al. 2014b
$\Delta ldhA_budA_budB$							
			Ethanol				
K. pneumoniae GEM167 mutant	Fed-batch	Glycerol	Microaerobic	21.5	0.62	0.93	Oh et al. 2011
K. pneumoniae GEM167 mutant	Fed-batch	Glycerol	Microaerobic	31.0	0.89	1.20	Oh et al. 2012b
$\Delta ldhA_pdc_adhII$							
E. coli (nine gene knockout strain)	Batch	Glycerol	Microaerobic	18.0	0.90	0.38	Trinh and Srienc,
							2009
			Lactic acid				
K. pneumoniae DSM 2026	Fed-batch	Glycerol	Microaerobic	22.7	0.32	0.95	Ashok et al. 2011
$\Delta dhaT_puuC$							
K. pneumoniae J2B	Fed-batch	Glycerol	Microaerobic	28.2	0.28	0.39	Kumar et al. 2013b

Table 4: Comparison of BDO, ethanol and lactic acid production by different strains of *K. pneumoniae* and other organisms

Δ ldhA _ KGSADH							
K. pneumoniae BLh-1	Fed-batch	Glycerol	Microaerobic	59.0	0.48	1.48	Rossi et al. 2013
K. pneumoniae J2B	Fed-batch	Glycerol	Microaerobic	37.0	0.32	0.77	Durgapal et al. 2014
K. pneumoniae DSMZ	Fed-batch	Glycerol	Microaerobic	47.8	0.39	1.00	Durgapal et al. 2014
K. pneumoniae ATCC25955	Fed-batch	Glycerol	Microaerobic	142.1	0.84	2.96	Feng et al. 2014
$\Delta dhaT_\Delta yqhD_ldhA$				(D-isomer)			
E. coli MG1655	Fed-batch	Glycerol	Microaerobic	32.0	0.82	0.54	Mazumdar et al. 2010
$\Delta pta \Delta adh E \Delta frdA$ -				(D-isomer)			
Δdld_glpK_glpD							
E. coli MG1655	Fed-batch	Glycerol	Microaerobic	50.0	0.92	0.60	Mazumdar et al. 2013
$\Delta pflB_\Delta pta_\Delta adhE_\Delta frdA$				(L-isomer)			
$\Delta mgsA \Delta lldD \Delta ldhA:: ldhA glpK-$							
_glpD							

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Potential and limitations of Klebsiella pneumoniae as a microbial cell factory utilizing glycerol as the carbon source

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