Effects of mutation of 2,3-butanediol formation pathway on glycerol metabolism and 1,3-propanediol production by Klebsiella pneumoniae J2B

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Abstract

The current study investigates the impact of mutation of 2,3-butanediol (BDO) formation pathway on glycerol metabolism and 1,3-propanediol (PDO) production by lactate dehydrogenase deficient mutant of Klebsiella pneumoniae J2B. To this end, BDO pathway genes, budA, budB, budC and budO (whole-bud operon), were deleted from K. pneumoniae J2B ΔldhA and the mutants were studied for glycerol metabolism and alcohols (PDO, BDO) production. ΔbudO-mutant-only could completely abolish BDO production, but with reductions in cell growth and PDO production. By modifying the culture medium, the ΔbudO mutant could recover its performance on the flask scale. However, in bioreactor experiments, the ΔbudO mutant accumulated a significant amount of pyruvate (>73 mM) in the late phase and PDO production stopped concomitantly. Glycolytic intermediates of glycerol, especially glyceraldehyde-3-phosphate (G3P) was highly inhibitory to glycerol dehydratase (GDHt); its accumulation, followed by pyruvate accumulation, was assumed to be responsible for the ΔbudO mutant’s low PDO production.

Key words: Klebsiella pneumoniae J2B; Glycerol; 1,3-Propanediol; 2,3-Butanediol; ΔbudO mutant
1. Introduction

1,3-Propanediol (PDO) is an important platform chemical having a wide range of applications in the production of polymers, cosmetics, and lubricants, among others. The most important industrial use of PDO is as a monomer for synthesis of the new polyester polytrimethylene terephthalate (PTT) (Celińska, 2010; Maervoet et al., 2011; Saxena et al., 2009). Many organisms in the Enterobacteriaceae family, such as Klebsiella pneumoniae, Klebsiella oxytoca, Citrobacter freundii, Enterobacter aerogenes and Enterobacter agglomerans, naturally produce PDO from glycerol, among which K. pneumoniae is the most extensively studied (Celińska, 2010; 2012). K. pneumoniae dissimilates glycerol by two parallel “oxidative” and “reductive” pathways (Supplementary Fig. 1). In the oxidative pathway, glycerol is converted to dihydroxyacetone phosphate (DHAP) by a respiratory (aerobic or anaerobic, according to the electron acceptor type) and/or fermentative route, DHAP then being funneled into the glycolytic pathway. In the reductive pathway, glycerol is converted to PDO by a two-step process: first, it is dehydrated to 3-hydroxypropionaldehyde (3-HPA) by the coenzyme B$_{12}$-dependent glycerol dehydratase (GDHt), which is encoded by dhaB; then, 3-HPA, at the expense of NAD(P)H, is reduced to PDO by 1,3-propanediol oxidoreductases (PDORs) such as DhaT (NADH-PDOR) and/or NADPH-dependent oxidoreductase (Celińska, 2012; Kumar et al., 2012; Saxena et al., 2009). The reductive pathway regenerates NAD$^+$, which enables K. pneumoniae to grow on glycerol under limited-O$_2$ conditions.

During oxidative metabolism of glycerol, K. pneumoniae generates a number of metabolites including organic acids (lactic acid, succinic acid, acetic acid, formic acid) and alcohols [2,3-butanediol (BDO), ethanol] (Ashok et al., 2011; Kumar et al., 2013a). These
glycerol-metabolic byproducts subdue the glycerol flux to PDO, thereby significantly reducing PDO production yields; additionally, at high concentrations, they are toxic to cell growth and PDO production. Lactic acid, particularly, is a major byproduct (Ashok et al., 2011; Durgapal et al., 2014; Huang et al., 2012), and much research efforts and resources have been devoted to the elimination of its formation (Kumar et al., 2013b; Oh et al., 2012; Xu et al., 2009). Another major byproduct, especially appearing when lactic acid production has been eliminated, is BDO. BDO production reduces PDO yield more significantly than does lactic acid, because synthesis of one molecule of BDO requires two molecules of glycerol. Moreover, due to their similar boiling points, the presence of BDO in the culture broth complicates PDO purification in downstream processing (Anand et al., 2011; Kaur et al., 2012; Zeng and Biebl, 2002). BDO synthesis begins with self-condensation of two molecules of pyruvate to one molecule of C5 intermediate α-acetolactate (Supplementary Fig. 1), which is catalyzed by α-acetolactate synthase (ALS, budB). In the next step, α-acetolactate is decarboxylated to acetoin (catalyzed by α-acetolactate decarboxylase (ALDC, budA)) and then reduced to BDO by 2,3-butanediol dehydrogenase/acetoin reductase (AR, budC) using NADH as the reductant. In the presence of oxygen, α-acetolactate is spontaneously decarboxylated to diacetyl, which is then reduced to acetoin by the action of diacetyl reductase, and acetoin in turn is reduced to BDO. When diacetyl is converted to acetoin or acetoin is converted to BDO, one NADH, at each step, is required (Celińska and Grazek, 2009; Ji et al., 2011). The genes coding for the three enzymes in the BDO-producing branch are located in one operon (budO) in the order budA (ALDC), budB (ALS) and budC (AR).

The current study investigated the role of the three enzymes of the BDO synthetic pathway in cell growth, glycerol metabolism and BDO production under different aeration
conditions. Also, the possibility of producing PDO at an economically feasible level without BDO production was explored. The newly isolated *K. pneumoniae* J2B (KCCM11213P) strain was used, as it does not produce pathogenic and sticky lipopolysaccharides (Arasu et al., 2011), and also because it has better sedimentation properties and a higher sensitivity to antibiotics than other well-studied *K. pneumoniae* such as the DSMZ2026 strain. Using *K. pneumoniae* J2B $\Delta ldhA$ as the base strain, four mutant strains ($\Delta budA$, $\Delta budB$, $\Delta budC$ and $\Delta budO$) were developed and their performances studied by shake-flask and bioreactor experiments.

2. Materials and methods

2.1 Materials

A genomic DNA isolation kit and pGEM-T vector was purchased from Promega (Madison, WI, USA). High-fidelity *pfx* polymerase was acquired from Invitrogen (Seoul, Korea). Restriction DNA-modifying enzymes were obtained from New England Bio-Labs (Beverly, MA, USA). The miniprep and DNA gel extraction kits were purchased from Qiagen (Mannheim, Germany). The primers were synthesized by Cosmotech Co. Ltd. (Seoul Korea). Yeast extract (Cat. 212750) was obtained from Difco (Becton Dickinson; Franklin Lakes, NJ, USA). Glycerol and all other chemicals and enzymes (unless indicated otherwise) were purchased from Sigma-Aldrich (St. Louis, MO, USA).

2.2 Construction of plasmids and mutant strains

All of the strains, plasmids and primers are listed in Table 1. For the design of the four mutant strains of *K. pneumoniae* J2B ($\Delta budA$, $\Delta budB$, $\Delta budC$ and $\Delta budO$), an in-frame tagged deletion approach was employed according to a slightly modified version of the method
introduced by Link et al. (1997). Briefly, PCR amplification of ~500 bp of the upstream (fragment A) and ~500 bp of the downstream (fragment B) regions of the above-noted genes was performed using the primers listed in Table 1. In the next step, the two amplified fragments A and B were ligated using overlapping PCR methods to synthesize the engineered fragment AB, which subsequently was cloned into the pGEMT vector. After confirming this sequence, the engineered fragment AB was sub-cloned into the pKOV vector between the restriction sites (shown in Table 1). These plasmids were used to knock out, by homologous recombination, the target genes from the chromosomal DNA of *K. pneumoniae* J2B ΔldhA. The mutant strains were screened using PCR and confirmed by sequencing. The developed mutant strains of *K. pneumoniae* J2B ΔldhA, ΔldhAΔbudA, ΔldhAΔbudB, ΔldhAΔbudC and ΔldhAΔbudO, were designated KpΔbudA, KpΔbudB, KpΔbudC and KpΔbudO, respectively. *K. pneumoniae* J2B ΔldhA was designated as the *K. pneumoniae* control (Kp control).

### 2.3 Shake-flask cultivation

Shake-flask experiments were carried out at 37°C with an initial pH of 7. Different strains used in gene-deletion studies were cultured in Luria-Bertani medium (LB medium) containing yeast extract (5 g/L), NaCl (10 g/L), and tryptone (10 g/L). For shake flask production experiments, a primary inoculum was prepared by culturing cells in LB medium from agar plates for 8 h. From the primary inoculum, the cells were transferred to fresh LB medium for 2 h in order to harvest active cells at the mid-log phase. The primary and the secondary inoculum cultures were grown under the same conditions (working volume: 50 mL in 250 mL Erlenmeyer flask; agitation speed: 200 rpm; initial pH: 7.0, temperature: 37°C). The cell OD₆₀₀ at the end of pre-cultures I and II were in the 5.0 - 6.0 and 1.5 - 2.0 ranges, respectively. The
starting OD$_{600}$ in all of the experiments was 0.1 - 0.2, and so the inoculum volume was dependent on the final OD$_{600}$ achieved at the end of each stage. The main cultivation of the designed mutant strains of \textit{K. pneumoniae} J2B (Kp control, Kp$\Delta$bud$A$, Kp$\Delta$bud$B$, Kp$\Delta$bud$C$ and Kp$\Delta$bud$O$) was conducted in 250 mL flasks containing a medium of the following composition: glycerol-20 g/L (220 mM), yeast extract-1.0 g/L, NH$_2$SO$_4$-2.0 g/L, MgSO$_4$·7H$_2$O-0.2 g/L, CaCl$_2$·2H$_2$O-0.02 g/L, K$_2$HPO$_4$-3.4 g/L, KH$_2$PO$_4$-1.3 g/L; Fe solution-1 mL/L, and trace-element solution-1 mL/L (Oh et al., 2012).

Shake-flask cultivation of the different mutant strains was performed under various aeration conditions: aerobic, microaerobic, and anaerobic. In the aerobic and microaerobic cultures, the flasks were plugged with an oxygen-permeable cotton stopper; in the anaerobic cultures, the flasks were closed with a screw cap, and the flask head space was replaced with argon gas prior to cultivation. The working volumes under aerobic, microaerobic, and anaerobic conditions were 25, 100 and 50 mL, respectively. The agitation speed under the aerobic condition was 250 rpm, and under both the microaerobic and anaerobic conditions, 100 rpm. The culture medium was supplemented with different components in order to investigate their effects on cell metabolism and PDO production. The Kp$\Delta$bud$A$ and Kp$\Delta$bud$O$ strains were supplemented with two different concentrations of branched-chain amino acids (leucine, isoleucine, and valine) (1.0 and 2.0 mM) as well as complex nitrogen sources (yeast extract-1.0 g/L, peptone-2.5 g/L, beef extract-2.5 g/L, yeast extract-2.5g/L, peptone-5.0 g/L, beef extract-5.0 g/L). The shake-flask cultivation of Kp$\Delta$bud$O$ was additionally supplemented with exogenous additions of BDO (25 mM) and NaHCO$_3$ (50 mM).
2.4 Bioreactor cultivation

The bioreactor experiments were carried out in a 1.5 L jar bioreactor (KO Biotech, Korea) containing a 1.0 L medium of the same composition as in the shake-flask cultivation (see above). The inoculum culture was prepared in the same way as described for the flask experiment. The pH, temperature, stirrer speed and aeration rate in the bioreactor experiments were maintained at 6.8, 37°C, 250 rpm and 0.5 vvm, respectively, unless specified otherwise. The pH was maintained by addition 5 N NaOH. Fed-batch experiments were performed using Kp\textit{budO} with a feeding solution of glycerol (1,000 g/L; 10.9 M) and yeast extract (25 g/L). In one run, carbon dioxide (CO$_2$) was continuously flushed at an aeration rate of 0.05 vvm, and in another run, NaHCO$_3$ (100 mM) was added to the culture medium.

2.5 Glycerol dehydratase (DhaB) activity and inhibition by glycerol metabolites

\textit{K. pneumoniae} cells grown in LB medium were induced and harvested 3 h from the time of induction. The harvested cells were lysed anaerobically using a bead beater and subjected to a DhaB-activity assay inside an anaerobic chamber. An appropriate amount of cell lysate was pre-incubated with 10 mM of potassium phosphate buffer (pH 7), purified KGSADH and 40 mM 1,2-PDO at 37°C for 3 min. The reaction was initiated by adding 15 \textmu M coenzyme-B$_{12}$, 1.5 mM ATP, 3 mM MgCl$_2$ and 2 mM NAD$^+$ to the reaction mixture for a total volume of 2 mL. In this mixture, 1,2-PDO acts as a substrate for DhaB protein and is converted to propionaldehyde, which, in sequence, is converted to propionic acid by the action of coupling-enzyme KGSADH. The activity of KGSADH was evaluated by measuring the reduction of NAD$^+$ to NADH at 340 nm. The amount of NADH formed was determined using an extinction coefficient of 6.22 mM$^{-1}$ cm$^{-1}$. One unit of enzymatic activity was defined as the amount of
enzyme required to form 1 µmol of propionaldehyde per minute. The metabolites pyruvate, DHAP, MG, G3P and DHA were added to the enzymatic reaction mixture in different concentrations (0.5, 1, 2, 5 and 10 mM, respectively) in order to study their individual inhibitory effects.

2.6 Analytical methods

Cell density was measured in a 10-mm-path-length cuvette using a double-beam spectrophotometer (Lambda 20, PerkinElmer; Norwalk, CT, USA). One unit of absorbance at 600 nm (OD$_{600}$) corresponded to 0.31 g dried cell mass per liter. The concentrations of glycerol and of the other metabolites were determined by HPLC using a slightly modified version developed by Raj et al. (2008). Briefly, the supernatants, obtained by centrifuging the culture samples at 10,000 ×g for 10 min, were filtered through a Tuffryn membrane (Acrodisc, Pall Life Sciences, USA) and eluted through a 300 ×7.8 mm Aminex HPX-87H (Bio-Rad, USA) column at 65°C. The mobile phase and flow rate were 2.5 mM H$_2$SO$_4$ and 0.5 mL/min, respectively. All of the experiments were carried out at least three times to ensure reproducibility. The data reported are the averaged values for three independent measurements; the standard deviation was less than 10% unless otherwise indicated.

3. Results and discussion

3.1 Metabolic changes in Kp control, Kp$\Delta$budA, Kp$\Delta$budB, Kp$\Delta$budC and Kp$\Delta$budO under different aeration conditions during flask cultivation

Five mutant strains, one base strain (Kp control) and four BDO-pathway mutant strains (Kp$\Delta$budA, Kp$\Delta$budB, Kp$\Delta$budC and Kp$\Delta$budO) were developed and cultured under three...
different aeration conditions (aerobic, microaerobic and anaerobic) (Table 2). Cell growth, glycerol consumption and PDO production significantly varied depending on the genes disrupted and aeration conditions adopted for cultivation. Under aerobic condition, the growths of Kp\textit{budB} and Kp\textit{budC} (OD\textsubscript{600}, 7-8) were similar to that of the Kp control (OD\textsubscript{600}, 7.6) whereas those of Kp\textit{budA} and Kp\textit{budO} (OD\textsubscript{600}, 4-5) were lower. Similarly, glycerol consumption was high in Kp\textit{budB} and Kp\textit{budC} (~200 mM) but low in Kp\textit{budA} and Kp\textit{budO} (<70 mM). PDO production was generally low under the aerobic conditions, due mainly to insufficient production of coenzyme B\textsubscript{12}. Among the mutant strains, Kp\textit{budB} and Kp\textit{budC} (38-41 mM) produced much higher PDO than did Kp\textit{budA} or Kp\textit{budO} (2-6 mM); but none of the mutants showed higher PDO production than that of the Kp control (44.7 mM) under the aerobic conditions. BDO production decreased when the enzyme(s) of the BDO pathway were disrupted, but varyingly; Kp\textit{budB} and Kp\textit{budC} showed a small decrease, while Kp\textit{budA} manifested a significant drop. It was noted that \textit{budO}-only completely abolished BDO production. The low impacts of \textit{budB} and \textit{budC} were attributed to the presence (and active expression) of the isozymes of ALS (encoded by \textit{budB}) and AR (encoded by \textit{budC}). Recently it was reported that AR can convert 3-hydroxypropionaldehyde to 1,3-PDO, similar to 3-hydroxypropionaldehyde-specific oxidoreductase, DhaT (Wang et al., 2014). Wu et al. (2013), moreover, earlier reported the presence of several genes having similar sequences to that of \textit{budC}. The low cell growth and glycerol consumption in Kp\textit{budA} indicates that \textit{budA} does not have isozyme(s), and further, that accumulation of the substrate of \textit{budA}, \textit{\alpha}-acetolactate, is highly toxic to cells. The detour route for the conversion of \textit{\alpha}-acetolactate to acetoin via diacetyl can compensate for the lack of \textit{budA} isozymes, but its efficiency seems to be low. The similar behaviors of Kp\textit{budA} and Kp\textit{budO} with respect to cell growth, glycerol consumption and PDO production, among others,
strongly suggest that, under aerobic conditions, $\Delta$bud$A$ is the most influential of all of the
mutations in the BDO operon.

Under the anaerobic condition, cell growth decreased while PDO production increased.
Kp$\Delta$bud$B$ and Kp$\Delta$bud$C$ (final OD$_{600}$, 3-4) showed a higher cell density than Kp$\Delta$bud$A$ or
Kp$\Delta$bud$O$ (final OD$_{600}$, ~1.9). Surprisingly, the Kp control, similarly to the latter two mutants,
showed low cell growth (final OD$_{600}$, ~1.9). Glycerol consumption and PDO production were
high in Kp$\Delta$bud$B$ and Kp$\Delta$bud$C$, and low in Kp$\Delta$bud$A$ and Kp$\Delta$bud$O$, in comparison to Kp
control. As was the case under the aerobic conditions, Kp$\Delta$bud$A$ and Kp$\Delta$bud$O$ showed similar
behaviors in terms of cell growth, glycerol consumption, and production of PDO and other
metabolites; this suggests, again, that bud$A$ deletion has a higher impact on the BDO pathway
than either bud$B$ or bud$C$ deletion. It should be noted, too, that whereas Kp$\Delta$bud$B$ and Kp$\Delta$bud$C$
showed similar cell growth and PDO production, the metabolite production profiles were
significantly different: Kp$\Delta$bud$B$ produced much acetate and little ethanol, while Kp$\Delta$bud$C$
produced much ethanol and little acetate. Also, whereas Kp$\Delta$bud$B$ did not produce BDO,
Kp$\Delta$bud$C$ produced a significant amount of it, even more than the Kp control. It is not clear why
Kp$\Delta$bud$B$ and Kp$\Delta$bud$C$ show such different behaviors, though we speculate that it must be
closely related to NADH balance. As indicated in supplementary Fig. 1, BDO production
generates, from the two moles of pyruvate with consumption of one mole of NADH and
unutilized NADH molecules may be consumed in the production of ethanol (i.e., in Kp$\Delta$bud$C$).
If, as in the case of Kp$\Delta$bud$B$, BDO is not produced, no excessive NADH are generated, and
thus, acetate production is the right venue. In addition, acetate production yields ATP, which
seems always to be beneficial to cells. This NADH effect is observed also under aerobic
conditions, though not as significantly. We assume that, under aerobic conditions, maintenance of
the redox balance can be managed rather easily by active operation of the electron transport
chain. And in fact, under microaerobic conditions, cell growth, glycerol consumption and PDO
production were quite similar to those under anaerobic conditions, as were the metabolite
production profiles. Once again, these results indicate that, among the three genes, budA deletion
has the greatest impact on glycerol metabolism.

BDO production, meanwhile, was suppressed by mutations in the BDO pathway, though
its complete elimination was observed only in Kp∆budO. In most of the strains tested, acetate
and ethanol were produced as major byproducts, towards which, additional carbon flux was
diverted upon suppression of the BDO pathway. In some cases, especially with Kp∆budB and
Kp∆budO, substantial pyruvate excretion was observed; indeed, with Kp∆budO, a significant pH
drop (below 5.0) also was noticed. Such pyruvate excretion and pH loss in Kp∆budO suggests
that the complete elimination of the BDO pathway in the ldhA− background causes heavy
metabolic traffic at the pyruvate node.

3.2 Flask culture of Kp∆budO with supplementation of amino acids, NaHCO₃ and/or BDO

Among the four mutant strains, only Kp∆budO could produce PDO without BDO
accumulation. However, after elimination of the whole-bud operon, glycerol consumption and
PDO production were sharply diminished. In order to explore the reasons for this as well as the
possibility of PDO production without BDO, optimization of Kp∆budO culture conditions was
performed. All of the experiments were carried out under the microaerobic condition, as these,
among the three aeration conditions, had proved the most conducive to cell growth and PDO
production. First, the effect of complex nitrogen was studied. Branched-chain amino acids
(leucine, isoleucine and valine) or larger amounts of complex nitrogen sources (yeast extract,
beef extract, and peptone) were added to the culture medium. It was hypothesized that elimination of the *bud* operon might curtail the availability of the three essential branched-chain amino acids, due to the lack of their precursor, α-acetolactate, the first intermediate of the BDO pathway (see supplementary Fig. 1). The branched-chain amino acids were added to the culture medium at two different concentrations (1 and 2 mM each), as were the mixtures of complex nitrogen sources (1 g/L yeast extract, 2.5 g/L peptone and 2.5 g/L beef extract; 2.5 g/L yeast extract, 5 g/L peptone and 5 g/L beef extract) (data not shown). However, no improvement in glycerol consumption or PDO production was noted for any of the cultures.

Next, the effects of higher phosphate concentration (100 mM) and the addition of BDO and/or NaHCO₃ were studied (Fig. 1). The testing of the high phosphate concentration was carried out owing to the precipitous pH drop (below 5.0) that had been observed in the experiments where 29 mM of potassium phosphate was used (see Table 1 for data). As for BDO supplementation, we sought to determine if BDO itself, the final product of the BDO pathway, has a physiological role (its lack had hampered glycerol consumption and PDO production). In the same context, we hypothesized that the lack of CO₂, which is generated at 2 moles per mole of BDO (see supplementary Fig. 1), can hamper the performance of KpΔ*budO*. In the preliminary results, the increase in the buffering capacity of the culture medium, from 29 to 100 mM, greatly improved glycerol consumption and PDO production (Figs. 1C and 1D): 91.1 mM PDO in a significantly enhanced yield (0.67 mol PDO/mol glycerol) was produced from 135.8 mM glycerol in 12 h. However, the pH nonetheless decreased below 5.3 after 9 h, and a substantial amount of glycerol was left unused. Significantly, the addition of BDO (25 mM) to the high-buffer culture medium barely affected the performance of the strain (Figs. 1E and 1F). The addition of NaHCO₃
(50 mM), on the other hand, further improved glycerol consumption and PDO production (Figs. 1G and 1H): more than 90% of glycerol was consumed in 9 h, and cell growth (3.0 at OD$_{600}$) along with the PDO titer (112 mM) were enhanced by 45.4 and 22.5%, respectively, relative to the case without NaHCO$_3$. Significantly too, when BDO was additionally supplemented to the NaHCO$_3$-containing medium (Figs. 1I and 1J), no further improvement in PDO production was observed. Sodium bicarbonate (NaHCO$_3$) is a source of CO$_2$, but it can also counteract pH drop caused by generation of various acids. When NaHCO$_3$ was added, the final pH increased to ~5.8 (Figs. 1G and 1I) from ~5.3 (Figs. 1C and 1E). The production of metabolites, meanwhile, varied significantly with the culture conditions (Figs. 1B, 1D, 1F, 1H and 1J). Acetate, pyruvate and formate were the major byproducts, with ethanol, lactate and succinate as the minor ones (under 10 mM). When NaHCO$_3$ was added, production of acetate (~40 mM) and formate (~27 mM) was greatly increased, whereas pyruvate excretion was greatly reduced. This suggests that the addition of NaHCO$_3$ and/or the increase of culture pH stimulates pyruvate-formate-lyase (PFL), which converts pyruvate to acetyl-CoA and formate, thereby reducing pyruvate accumulation. As for succinate, its production was slightly increased, suggesting that carboxylation of pyruvate to oxaloacetate also was stimulated. Overall, these results indicate that, in the Kp$\Delta$budO mutant where both the lactic acid and BDO production pathways were eliminated, excessive production of acids (especially the accumulation of pyruvate) and/or subsequent pH drop could be the principal cause of the reduction of PDO production and cell growth, and that their negative impact can be greatly alleviated by increasing medium buffering capacity and/or the addition of NaHCO$_3$ to the culture medium. It should also be noted that Kp$\Delta$budO cultured under microaerobic and modified conditions exhibited a similar performance (in terms of cell growth,
glycerol consumption, PDO production, final pH, production of acetate and ethanol, etc.) to that of KpΔbudB cultured under anaerobic and non-modified conditions.

3.3 Batch bioreactor experiments on KpΔbudO with NaHCO₃ or continuous CO₂ supply

The effects of CO₂ and NaHCO₃ on KpΔbudO performance was further studied using pH-controlled bioreactors in a batch mode (Fig. 2). Carbon dioxide was added in the form of NaHCO₃ (100 mM) or by continuous sparging of CO₂ gas at 0.05 vvm. The pH was maintained at 7.0 and the initial glycerol concentration was set to 600 mM. Despite the pH maintenance, KpΔbudO exhibited a performance much inferior to the Kp control. When NaHCO₃ was added, some improvement in cell growth and PDO production was noticed, but not to the level by the Kp control. Specifically, whereas the Kp control produced ~305 mM PDO, KpΔbudO with NaHCO₃ produced only 265 mM. In the case of the continuous sparging of CO₂ (Fig. 2E), PDO production and cell growth were even lower than in the case without CO₂ (Fig. 2C), suggesting that excessive CO₂ is inhibitory. Cell growth and glycerol consumption also were lower in KpΔbudO. Overall, these results indicate that the negative impact of the deletion of the bud operon, cannot be removed by simple optimization of culture conditions at flask level. These bioreactor results also suggest that the improvement of the performance of KpΔbudO (shown in Fig. 1) was mostly owed to the increase of the culture pH to a more neutral level.

The major end products of glycerol metabolism in K. pneumonia under limited-aeration conditions are lactic acid and BDO (Durgapal et al., 2014). When lactic acid production is blocked, more BDO is produced, and vice versa. If both BDO and lactic acid production are blocked at the same time, other metabolites such as ethanol, acetate, formate, hydrogen, succinate, and others will be produced at higher concentrations to keep the carbon flow at the pyruvate node (see supplementary Fig. 1). Otherwise, due to metabolic traffic at the pyruvate
node, glycerol metabolism through the oxidative pathway is terminated, which seemingly was the case with the current KpΔbudO. There are three strategies for dealing with the carbon traffic problem at the pyruvate node of K. pneumonia that lacks both ldhA and budO. First, the consumption of pyruvate through the TCA cycle can be accelerated. Among the advantages of this strategy is improved cell growth and viability, and enhanced the NADH supply for PDO production. On the other hand, the glycerol-to-PDO yield can be reduced if carbon is mainly used for cell growth. The second strategy for dealing with the carbon traffic problem at the pyruvate node is suppression of pyruvate generation by reduction of glycerol consumption through the oxidative pathways at the glycerol node (by perturbing the expressions of glpK, gldA and/or dhaD; see supplementary Fig. 1). The advantage and disadvantage of the second strategy are exactly the opposite of those indicated for the first strategy. Furthermore, a lowered NADH production rate can lead to a greatly reduced PDO production rate. The third and most desirable strategy is to accelerate the PDO production (reductive) pathway relative to the oxidative pathway at the glycerol node. Zheng et al. (2006) attempted overexpression of dhaB to enhance the reductive pathway in a wild-type strain of K. pneumonia which resulted in accumulation of toxic 3-HPA; it did not improve PDO production (though it should be noted that minimization of BDO production was not their goal). No report on dhaB and dhaT expression enhancement in the background of ΔldhA and ΔbudO has yet been published.

One additional interesting feature of the present bioreactor experiments with KpΔbudO was the re-appearance of lactic acid, which had not been observed in the flask experiments. We speculated that activation of the pathway via methylglyoxal was responsible for this (see supplementary Fig. 1). To prove that hypothesis, the mgsA deletion mutant K. pneumonia J2B
ΔldhAΔbudOΔmgsA was developed and tested (data not shown). The results showed that even though no detectable lactic acid was produced, PDO production was not improved.

3.4 Fed-batch bioreactor experiments on KpΔbudO

Among the above-suggested strategies for alleviation of the carbon traffic at the pyruvate node, the first, which is to say, improvement of TCA-cycle throughput was briefly examined. This goal can be accomplished by increasing aeration and/or supplying additional nitrogen. First, increased yeast-extract concentration (initial concentration: 5 g/L) was explored (Fig. 3). The air-flow rate was set at 0.5 vvm. For comparison, the same experiments but with 1 g/L yeast extract were also performed for both the Kp control and KpΔbudO. The PDO production of the Kp control with 1 g/L yeast extract was 752.9 mM at 36 h; under the same conditions, KpΔbudO produced only ~410 mM PDO. For both strains, the initial (to 6 h) cell growth and PDO production, ~ 5 OD and 130 ~ 150 mM, respectively, were similar; thereafter however, KpΔbudO showed a performance significantly inferior to that of the Kp control. Moreover, from 6 h, KpΔbudO started to accumulate pyruvate and lactate, as in the batch bioreactor experiments. When the initial yeast-extract concentration was elevated to 5 g/L (Fig. 3E and 3F), the cell growth rate and glycerol consumption were greatly accelerated, and the maximum cell density (OD₆₀₀-12) reached a significantly higher level at ~15 h. The final PDO titer also was considerably improved, to 512 mM, at 36 h. However, the level did not match that of the Kp control, as the PDO production rate had declined significantly after 12 h. Interestingly, the two strains’ molar PDO yields with glycerol were comparable (Kp control, 0.43; KpΔbudO, 0.46).

The metabolite profiles of the two strains are compared in Table 3. The cultivation phase was divided into two phases according to the cell-growth pattern and metabolite production.
The carbon recoveries in each phase were higher than 95% when the yeast extract was 1 g/L, but were reduced to ~85% when the yeast extract was elevated to 5 g/L. The reason for this is not clear. With KpΔbudO, glycerol assimilation in the second phase was greatly reduced relative to that in the first phase. With the Kp control, a large amount of BDO, ~274 mM, was produced in 36 h. In KpΔbudO, the glycerol carbon that had been used for BDO formation in the Kp control seemed to be diverted to a variety of metabolites, especially formate, acetate, pyruvate, lactate and ethanol. However, the total amount diverted to these metabolites was far less than that of BDO in the Kp control, indicating that glycerol assimilation and its metabolism was significantly reduced in KpΔbudO. It was also noted that, as in the batch experiments, KpΔbudO produced pyruvate and lactate at high concentrations. The high accumulation of pyruvate, even with a high concentration of yeast extract, indicated that the strategy to alleviate the carbon traffic at the pyruvate node by supplementing additional, readily-usable nitrogen and stimulating the TCA cycle was not successful, especially in the later phase of the bioreactor experiment.

In order to stimulate TCA throughput, another set of bioreactor experiments with an increased aeration rate (to 1 vvm) were performed (data not shown). Cell growth and glycerol consumption were enhanced in all three cases (Kp control with 1 g/L yeast extract; KpΔbudO with 1 g/L yeast extract; KpΔbudO 5 g/L yeast extract), but PDO production was reduced by ~15%. Also, PDO production began at ~6 h, about 3 h later than in the former, 0.5 vvm-aeration-rate case, probably due to lowered or delayed synthesis of coenzyme B_{12}. With KpΔbudO, metabolic traffic at the pyruvate node was not alleviated, >50 mM pyruvate having been accumulated. Intermittent feedings of yeast extract in the later phase did not improve glycerol consumption and/or PDO production. Again, these results show that carbon traffic at
the pyruvate node could not be lessened by increasing aeration and/or addition of a rich nitrogen source.

It has been reported that *K. pneumoniae* has an inefficient or incomplete TCA cycle, unlike other, similar enterobacter sp. such as *E. coli* (Cabelli, 1955). In that study, the growth with fumarate or succinate as the sole carbon source required a much longer lag period than that with TCA intermediates including malate, oxaloacetate, citrate and others, and it was suggested that fumarate does not go to malate. According to our preliminary studies, transcription of some enzymes (e.g., *fumA* and *sdhA*) was low, as were their enzymatic activities, especially under the anaerobic condition. If the TCA cycle does not operate properly, pyruvate utilization cannot be efficient and/or stimulation of TCA-cycle throughput becomes highly challenging. This suggests that in the context of the production of PDO without BDO, *K. pneumoniae* has a serious limitation as a microbial cell factory. Additional studies on the TCA cycle of *K. pneumoniae* are in progress.

### 3.5 Effects of glycerol carbon metabolites on glycerol dehydratase (DhaB) activity

The results obtained thus far indicate that the metabolic traffic at the pyruvate node is critical to glycerol metabolism in KpΔ*budO*. Accumulation of pyruvate causes that of other glycolytic intermediates such as dihydroxyacetone (DHA), dihydroxyacetone phosphate (DHAP), methylglyoxyl (MG) and glyceraldehyde-3-phosphate (G3P), and threatens to terminate glycerol assimilation through the oxidative pathway. Further, if any of these metabolites specifically inhibit the activity of glycerol dehydratase (DhaB) (see supplementary data Fig. 1), the conversion of glycerol to PDO also is greatly reduced. Therefore, the effect of glycolytic intermediates on DhaB activity was studied using crude cell extract expressing DhaB
at a high level (Fig. 4). Among the metabolites tested, G3P was the most highly inhibitory: at 1 mM, DhaB activity was completely eliminated. Other metabolites such as pyruvate, MG and DHAP also showed some inhibition, but much less significantly than G3P. DHA was not inhibitory up to 10 mM. These results suggest that G3P followed by pyruvate accumulation greatly decreases reductive flux in addition to oxidative flux, which eventually halts glycerol consumption.

4. Conclusion

The current study investigated the effects of the deletion of the BDO pathway on glycerol metabolism and PDO production by *K. pneumoniae* J2B ΔldhA. The deletion of *budO* (whole-*bud* operon) could completely eliminate BDO production, but this triggered heavy carbon-metabolic traffic at the pyruvate node. Glycerol dehydratase, which converts glycerol to 3-HPA, was specifically inhibited by G3P which should accumulate along with pyruvate. It is suggested that pyruvate accumulation should be strictly prevented during PDO production by carefully decreasing oxidative flux and/or increasing reductive flux.

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References


Figure legends

Fig. 1 Flask culture of K. pneumoniae J2B ΔldhAΔbudO with 29 mM (low) or 100 mM (high) phosphate (Pi) in presence or absence of BDO (25 mM) and/or NaHCO₃ (100 mM). A and B 29 mM Pi; C and D, 100 mM Pi; E and F, 100 mM Pi with BDO; G and H, 100 mM Pi with NaHCO₃; I and J, 100 mM Pi with BDO and NaHCO₃.

Symbols: In A, C, E, G and I, filled triangle down (residual glycerol), half-filled circle (cell OD), empty circle (PDO), and cross (pH); in B, D, F, H and J, empty squares (ethanol), empty diamond (acetate), filled circle (pyruvate), filled triangle up (formate), empty circle (BDO), filled star (succinate), and empty triangle up (lactate).

Fig. 2 Bioreactor culture of K. pneumoniae J2B ΔldhA and ΔldhAΔbudO with or without continuous purging of CO₂ (0.05 vvm) or addition of NaHCO₃ (100 mM). A and B, ΔldhA; C and D, ΔldhAΔbudO; E and F, ΔldhAΔbudO with CO₂ purging; G and H, ΔldhAΔbudO with NaHCO₃.

Symbols: In A, C, E, and G, filled triangle down (glycerol), half-filled circle (cell OD), and empty circle (PDO); in B, D, F, and H, empty squares (ethanol), empty diamond (acetate), filled circle (pyruvate), filled triangle up (formate), empty circle (BDO), filled star (succinate), and empty triangle up (lactate).

Fig. 3 Fed-batch bioreactor culture of K. pneumoniae J2B ΔldhA and ΔldhAΔbudO. A and B, ΔldhA with 1 g/L yeast extract; C and D, ΔldhAΔbudO with 1 g/L yeast extract; E and F, ΔldhAΔbudO with 5 g/L yeast extract.

Symbols: In A, C, and E, filled triangle down (glycerol), half-filled circle (cell OD), and empty circle (PDO); in B, D, and F, empty squares (ethanol), empty diamond (acetate), filled circle (pyruvate), filled triangle up (formate), empty circle (BDO), filled star (succinate), and empty triangle up (lactate).

Fig. 4 Glycerol dehydratase (DhaB) enzyme activity in presence of dihydroxyacetone (DHA), dihydroxyacetone phosphate (DHAP), glyceraldehyde 3-phosphate (G3P), methylglyoxyl (MG) and pyruvate.
Supplementary Fig. 1 Glycerol metabolism in *K. pneumoniae*. The genes encoding the relevant enzymes are shown in italics on the arrows. The encircled genes had been deleted in the current study. The solid lines represent single steps in the metabolic pathway, while broken lines indicate multiple steps (Ashok et al. 2011; Mazumdar et al. 2013).
**Fig. 1**

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![Graphs showing various data points and trends over time.](image-url)
Fig. 3