Manipulation of the Anoxic Metabolism in Escherichia coli by ArcB Deletion Variants in the ArcBA Two-Component System Gonzalo N. Bidart, a† Jimena A. Ruiz, b,c† Alejandra de Almeida, b Beatriz S. Méndez, b and Pablo I. Nikela,b# Instituto de Investigaciones Biotecnológicas "Dr. Rodolfo A. Ugalde", Universidad Nacional de San Martín,ª Departamento de Química Biológica, Facultad de Ciencias Exactas y Naturales, Universidad de Buenos Aires, INQUIBICEN-CONICET, and Instituto de Biociencias Agrícolas y Ambientales, CONICET, c Buenos Aires, Argentina Running headline: ArcB deletions and anoxic fluxome in E. coli **Section:** Biotechnology Keywords: Escherichia coli, ArcB, anoxic fluxome, redox balance, metabolic flux analysis, reduced biochemicals # Address correspondence to Pablo I. Nikel, Instituto de Investigaciones Biotecnológicas "Dr. Rodolfo A. Ugalde", Campus Miguelete UNSAM, Av. 25 de Mayo & Francia, 1650 Buenos Aires, Argentina. Phone: +54 11 4006 1500. Fax: +54 11 4724 1500. E-mail: pnikel@iib.unsam.edu.ar † Ex aeguo contribution

Bioprocesses conducted under conditions with restricted oxygen supply are increasingly exploited for the synthesis of reduced biochemicals using different biocatalysts. The model facultative anaerobe Escherichia coli has elaborate sensing and signal transduction mechanisms for redox control in response to the availability of O2 and other electron acceptors. The ArcBA twocomponent system consists of ArcB, a membrane-associated sensor kinase, and ArcA, the cognate response regulator. The tripartite hybrid kinase ArcB possesses a transmembrane, a PAS, a primary transmitter (H1), a receiver (D1), and a phosphotransfer (H2) domain. Metabolic fluxes were compared under anoxic conditions in a wild-type E. coli strain, its $\triangle arcB$ derivative, and two partial arcB deletion mutants in which ArcB lacks either the H1 domain or the PAS-H1-D1 domains. These analysis revealed that elimination of different segments in ArcB determines a distinctive distribution of D-glucose catabolic fluxes, different from that observed in the $\triangle arcB$ background. Metabolite profiles, enzyme activity levels, and gene expression patterns were also investigated in these strains. Relevant alterations were observed at the P-enol-pyruvate/pyruvate and acetylcoenzyme A metabolic nodes, and the formation of reduced fermentation metabolites such as succinate, D-lactate, and ethanol was favored in the mutant strains to different extents as compared to the wild-type strain. These phenotypic traits were associated to altered levels of the enzymatic activities operating at these nodes, as well as to elevated NADH/NAD+ ratios. Thus, targeted modification of global regulators to obtain different metabolic flux distributions under anoxic conditions emerges as an attractive tool for metabolic engineering purposes.

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INTRODUCTION

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Anoxic fermentation of different carbon sources by *Escherichia coli* is increasingly gaining momentum in biotechnological setups aimed at obtaining reduced biochemicals. Relevant examples in this sense include (but are not limited to) the production of ethanol (47, 57), succinate (52), D-lactate (35), and polyhydroxyalkanoates (30); often using redox and/or regulatory *E. coli* mutants as the biocatalyst. These metabolic engineering approaches clearly underscore the need of a complete understanding of cell

physiology and metabolic network operativity under anoxic growth conditions. Metabolic fluxes through the central carbon pathways constitute the backbone of cell metabolism and represent the *in vivo* reaction rates of cognate enzymatic steps (51). The observed fluxome is the phenotypic consequence of both gene transcription and translation, as well as enzymatic activity and the regulation exerted at the metabolite level (38). As such, fluxome analysis is a useful approach to undertake the description of phenotypes associated with mutations in global regulators and constitutes a helpful strategy to harness the full biotechnological potential of redox mutants.

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In E. coli, the metabolic regulation in response to changes in O₂ availability is mainly orchestrated by the ArcBA (anoxic redox control) two-component system (1, 7, 26, 48), which is composed of ArcB, the tripartite membrane-associated sensor kinase, and ArcA, the cognate response regulator. The ArcB sensor has a small transmembrane domain comprising 16 amino-acid residues followed by a leucine zipper and a PAS domain, that connects the transmembrane domain to the catalytic domains. ArcB also possesses three catalytic domains (Fig. 1A): a primary transmitter domain (H1), containing a conserved His²⁹²; a receiver domain (D1), containing a conserved Asp⁵⁷⁶; and a phosphotransfer (secondary transmitter) domain (H2), containing a conserved His⁷¹⁷ (18, 20, 21). ArcA phosphorylation takes place through a phosphorelay process comprising the activity of all three domains (24). Under micro-oxic conditions, ArcB undergoes auto-phosphorylation (39) followed by intramolecular phosphate transfer and transphosphorylation of ArcA (17, 27). ArcA~P modulates the expression of ca. 135 genes (43), mainly acting as a negative transcriptional regulator of genes encoding enzymes involved in oxic pathways, such as the major dehydrogenase enzymes of the tricarboxylic acid (TCA) cycle and the glyoxylate shunt (49). At the same time, genes encoding enzymes belonging to fermentation pathways become activated by ArcA~P under micro-oxic or anoxic conditions (26, 34, 40). Respiratory pathways are also affected by the ArcBA system, since expression of cyoABCDE and cydAB is repressed and activated by ArcA~P, respectively (26, 49).

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Genome-wide patterns of gene expression proved helpful to understand the complex ArcBA-dependent transcriptional regulation (41, 43). Unfortunately, and due to the diverse regulatory mechanisms that operate both at the post-translational and enzymatic activity levels, it is sometimes difficult to directly infer

phenotypic traits in mutant strains from these analysis. In particular, the biochemical mechanism of this regulatory system has been deciphered *in vitro* (2, 18), yet little is known about the *in vivo* effects of different *arcB* mutations on the central metabolic pathways of *E. coli* under anoxic growth conditions. In this study, the phenotypic and metabolic effects of targeted *arcB* deletions on the central carbon metabolism of *E. coli* was systematically evaluated under anoxic growth conditions through the analysis of growth parameters and the pattern of fermentation metabolites. The information was integrated in an *in silico* stoichiometric model of the central catabolic pathways, and was further substantiated by studying the transcription pattern of selected genes as well as *in vitro* measurements of relevant enzymatic activities. Taken together, the results show an incremental impact of partial deletions in ArcB on the distribution of metabolic fluxes under anoxic growth conditions that can be traced to the redox state. Furthermore, the incremental differences observed in the redox homeostasis among the mutant strains appear attractive for biotechnological purposes.

MATERIALS AND METHODS

Bacterial strains, oligonucleotides, and plasmids. All *Escherichia coli* strains are listed in Table 1 along with oligonucleotides and plasmids used in this work.

DNA manipulations and mutant construction. Standard DNA procedures followed well established protocols (44) and specific recommendations from manufacturers. Different protein domains in ArcB were eliminated by using the λ -Red recombination technology (12) in wild-type *E. coli* K1060. *E. coli* GNB1061, GNB1062, and GNB1063 were generated using amplification products (*i.e.*, *FRT-aphA-FRT*) obtained by PCR with oligonucleotides Δ H1-F and Δ H1-R, Δ PAS-F and Δ D1-R, and Δ arcB-F and Δ arcB-R, respectively, and plasmid pKD4 as the template. A DNA fragment encompassing nucleotides 802 to 1560 of *arcB* was deleted in *E. coli* GNB1061, resulting in an ArcB derivative in which the entire H1 domain was removed (*i.e.*, ArcB²⁶⁸⁻⁵²⁰). In *E. coli* GNB1062, the PAS, H1, and D1 domains (*i.e.*, ArcB¹⁷⁷⁻⁶⁴⁰), were eliminated by deletion of the DNA segment comprised between nucleotides 529 and 1920 of *arcB* (Fig. 1A). An *arcB* deletion mutant, termed *E. coli* GNB1063, was also constructed using the same methodology. Antibiotic resistant determinants were eliminated by FLP-mediated recombination using plasmid pCP20

(10). A suitable pairwise combination of the oligonucleotides *arcB1*-C-F, *arcB1*-C-R, *arcB2*-C-F, *arcB2*-C-R, PDH-C-F, PAS-C-R, H1-C-R, and D1-C-R (Table 1), followed by DNA sequencing of the corresponding amplicons, were used to confirm that the correct deletions were introduced into the *arcB* locus.

Growth media and culture conditions. All cultures were incubated at 37°C at the indicated rotary agitation. During mutant construction and inocula preparation, cells were grown in LB medium (44) under oxic conditions. Pre-cultures and working cultures were carried out under anoxic conditions in 100-ml bottles containing 90 ml of M9 minimal medium containing 3% (wt/vol) D-glucose as the sole carbon source and supplemented with 1 g \cdot l-¹ Na₂S as a reducing agent. Bottles were incubated with rotary shaking at 125 rpm to avoid cell sedimentation, and anoxic conditions were qualitatively checked by adding 50 μ g \cdot l-¹ resazurin to the culture medium. Whenever needed, antibiotics were used at the following concentrations: ampicillin, 100 μ g \cdot ml-¹; kanamycin, 50 μ g \cdot ml-¹; and chloramphenicol, 30 μ g \cdot ml-¹. The Dye phenotype of the strains, growing under oxic conditions on toluidine blue O agar medium, was evaluated as previously described (9, 42).

Analytical procedures. Biomass concentration was determined as the cell dry weight (CDW) fraction of washed pellets from valorated broth aliquots dried at 65°C to constant weight. Dried samples were allowed to cool and held *in vacuo* until weighed.

Extracellular metabolic products and residual D-glucose in culture supernatants were determined by HPLC (using a HPX-87H column; BioRad Labs., Hercules, CA) as previously described (13). In some experiments, H₂ evolution was measured by head-space gas chromatography using a column packed with a divinylbenzene porous polymer (HayeSep DB; HayeSeparations Inc., Bandera, TX) and a thermal conductivity detector in a Varian3000 gas chromatograph and a 4000MS ion trap mass spectrometer (Bruker Daltonik GmbH, Bremen, Germany).

The intracellular content of NADH, NADPH, NAD+, and NADP+ was estimated by using *in vitro* procedures based on rapid inactivation of the metabolism of growing cells followed by acid or alkaline extraction of

nucleotides. Nucleotide content was determined by means of spectrophotometric cycling assays with 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide as the final electron acceptor (5, 31).

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Metabolic flux analysis based on metabolic pathway balances. A metabolic network designed for anaerobic E. coli cells was drafted by including the biochemical reactions corresponding to the Embden-Meyerhof-Parnas pathway, biomass generation from D-glucose-6-P, and fermentation pathways from Penol-pyruvate, pyruvate, and acetyl-CoA (Fig. 2). Fermentation stoichiometry was derived from metabolic pathway balances. Cell composition was assumed to be the same in all experimental strains, and it was derived from Neidhardt et al. (28). Time-averaged concentrations in batch cultures, in which the specific rates of synthesis vary between two successive sampling periods (t and $t + \Delta t$), were estimated based on the average cell density according to the procedure described by Aristidou et al. (4). This methodology provides a good estimate when the specific rate of synthesis and the specific growth rate are more or less constant during consecutive sampling points, conditions which are assumed to be met under balanced growth (55). Metabolic fluxes (in mmol · g-1 · h-1) were calculated from time-averaged concentrations of secretion metabolites and carbon source, and represent the average of 4-5 independent sampling points about the mid-exponential growth phase (i.e., within balanced growth). Calculations that span the entire fermentation period indicated that indeed the time dependency of the fluxes was smallest during this interval (data not shown). The metabolic matrix was constructed based on the law of mass conservation and on the pseudo-steady state hypothesis on the intracellular intermediate metabolites (32, 45, 53). This formulation resulted in a set of linear equations that can be expressed as a stoichiometric matrix A of dimension $m \times n$, which was in turn related to vectors for net accumulation, $\mathbf{r}(m \times 1)$, and for metabolic fluxes, $\mathbf{v}(n \times 1)$. Considering the 13 reactions of the metabolic network (Fig. 2), and based solely on the measured extracellular metabolites and pseudo-steady state hypothesis balances of the intracellular intermediate metabolites, A becomes a square matrix (m = v = 13). The number of degrees of freedom equals the number of variables that were actually measured to describe the system, or, in some cases, were derived from experimental measurements. Standard mathematical methods were applied for the resolution of A·v = r = 0, and all fluxes within the network were derived from these stoichiometric constraints. Carbon balances were calculated from all the experimental and in-silico calculated fluxes, as well as the specific rate of CO₂ evolution, and biomass synthesis.

Quantitative reverse transcription (RT)-PCR. Genes studied in this work and the oligonucleotides used to analyze their expression are listed in Table 1. Culture samples were centrifuged at $10,000 \times g$ for 5 min at 4°C, washed once in ice-cold 50 mM Tris HCl buffer (pH = 7.5), and then pelleted by centrifugation under the same conditions before total RNA was isolated using a commercially available kit (TRIzol Max™ Bacterial RNA Isolation Kit; Life Technologies Corp., Grand Island, NY). gRT-PCR reactions were carried out in an Applied Biosystems 7900HT Fast Real-Time PCR System™ (Life Technologies Corp.) using the Promega Reverse Transcription System™ (Promega Corp., Madison, WI). In RT reactions, the reaction mixture (containing 1 µg of RNA in 60 µl reaction mixtures), was incubated for 10 min at room temperature, followed by 30 min at 50°C for reverse transcription, 5 min at 94°C, and 10 min at 65°C for reverse transcriptase inactivation. Amplification controls (containing RNA but not reverse transcriptase) were included in each run. The first strand cDNA was diluted to 250 µl with nuclease-free H₂O and further diluted 10-fold as needed. Quantitative PCR reactions were carried out in 96-well plates; each well contained 20 µl of a reaction mixture consisting of 2 μl diluted cDNA, 2 μl pre-mixed primers (1.25 pmol · μl-1), 10 μl SYBR Green™ PCR Master mix, and 6 µl nuclease-free H₂O. Reaction mixtures were incubated 2 min at 50°C, 10 min at 94°C (Taq DNA polymerase activation), followed by 40 cycles of 15 s at 94°C (denaturation), and 1 min at 60°C (annealing and extension). Each plate included amplification controls as well as samples without template. The comparative C_T (threshold cycle) method ($\Delta\Delta C_T$) was applied for relative quantification of gene expression, and rrsA, encoding the 16S subunit of ribosomal RNA, was used as the control gene.

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Enzyme activity measurements. Cells were harvested at $OD_{600} = 0.5$ -0.7 by centrifugation at $10,000 \times g$ for 15 min at 4°C, washed twice with ice-cold 50 mM Tris·HCl buffer (pH = 7.5) containing 20 mM KCl, 5 mM MgSO₄, 2 mM 1,4-dithio-D-threitol, and 0.1 mM EDTA; and finally resuspended in the same buffer (*ca*. 0.5 g wet cells in 1 ml of buffer solution). The resulting cell suspension was stored at -20° C for 3 h, thawed in an ice bath, and disrupted by sonication. Cell-free extracts were obtained by centrifugation at $15,000 \times g$ for 30 min and 4°C. Crude enzyme extracts were immediately used for determinations as well as to measure protein concentration by means of the Bradford method (6). Enzyme activities were assayed spectrophotometrically in a thermostated recording Beckman DU 650 spectrophotometer (Beckman Coulter Inc., Fullerton, CA). The components of the reaction mixture were pipetted into a cuvette of 1-cm light path,

and the reaction was initiated by adding the cell-free extract or the appropriate substrate to give a final volume of 1 ml. Standard *in vitro* protocols for pyruvate kinase (50), D-glucose-6-*P* 1-dehydrogenase (16), pyruvate-formate lyase (23), D-lactate dehydrogenase (8), succinate dehydrogenase (56), citrate synthase (14), alcohol dehydrogenase (25), and acetate kinase (15) were used to estimate the enzymatic activities at stake. The extinction coefficient at 340 nm used for NADH and NADPH was 6.22 mM⁻¹ cm⁻¹, and 1 unit of the specific enzyme activity was defined as the amount of enzyme required to convert 1 μmol of the substrate into the specific product per minute per milligram of protein.

Statistical analysis. All reported experiments were independently repeated at least twice, and the mean values of the corresponding parameter \pm standard deviation (and, in the case of metabolic flux analysis, 90% confidence intervals) are presented. Determination of statistical significance between multiple comparisons was assessed using analysis of variance (ANOVA, with α = 0.01 and 0.05) followed by a Bonferroni post-test using transformed data whenever necessary.

212 RESULTS

Evaluation of the Dye phenotype in *E. coli* strains expressing different ArcB variants. The Dye phenotype, *i.e.*, sensitivity to the redox dye toluidine blue O (3, 42), was analyzed as a coarse estimation of the phenotypic effects caused by the deletions implemented in arcB. To this end, cultures of *E. coli* K1060 (wild-type strain), GNB1061 (ArcB²⁶⁸⁻⁵²⁰), GNB1062 (ArcB¹⁷⁷⁻⁶⁴⁰), and GNB1063 ($\Delta arcB$) were spotted onto plates containing this dye and their growth was scored after overnight incubation (Fig. 1B). *E. coli* K1060 grew well on toluidine blue O plates, whereas the $\Delta arcB$ strain formed much smaller colonies. However, *E. coli* GNB1061 and GNB1062 differed noticeably in their ability to grow on this medium. While *E. coli* GNB1061 formed colonies somewhat bigger than those of the $\Delta arcB$ strain (yet smaller than those of the wild-type strain), *E. coli* GNB1062 exhibited a phenotype similar to that observed for the $\Delta arcB$ strain. These traits prompted us to further evaluate the metabolic properties in the mutants that caused the alterations in colony size.

E. coli strains expressing different ArcB variants exhibit incremental phenotypic features. Growth kinetic parameters of the strains under study were next analyzed in anoxic batch cultures. All experiments were conducted in M9 minimal medium supplemented with 3% (wt/vol) D-glucose in the absence of O_2 as the ArcBA two-component system is known to be active under these culture conditions (7). Interestingly, biomass concentration obtained after 24 h of cultivation was lower for mutants expressing the ArcB variants in comparison with the wild-type strain (*i.e.*, 36% and 60% lower for *E. coli* GNB1061 and GNB1062, respectively; Table 2). We also observed a reduction in the specific growth rate (μ_{max}) of the mutant strains (*i.e.*, 21% and 53% for GNB1061 and GNB1062, respectively; P < 0.05). As expected, *E. coli* GNB1063 showed the highest reduction in both final biomass concentration and μ_{max} (*i.e.*, 72% and 68% lower, respectively, when compared to the wild-type strain; P < 0.05), which is much lower than that observed in the strains bearing ArcB variants.

Specific rates of D-glucose consumption and yields of biomass on D-glucose ($Y_{X/S}$) were also determined in these cultures. All strains carrying ArcB variants showed a reduction in both parameters that qualitatively correlates with the results obtained for the specific growth rate and final biomass concentration (Table 2). The more marked differences among the partial mutants were observed in *E. coli* GNB1062, which showed a 27% and 62% reduction in both the specific rate of D-glucose consumption and $Y_{X/S}$, respectively, when compared to *E. coli* K1060 (P < 0.05). On the other hand, the $\Delta arcB$ strain exhibited a 40% and 73% reduction in the specific rate of D-glucose consumption and $Y_{X/S}$, respectively, when compared to the wild-type strain (P < 0.01).

In order to evaluate the metabolic state of the cells, the concentration of key fermentation metabolites was analyzed in culture supernatants of *E. coli* K1060 and the *arcB* mutant strains growing during 24 h under anoxic conditions and the corresponding yields on the carbon substrate were calculated. Metabolites detected in all cultures were formate, D-lactate, succinate, ethanol, and acetate. Higher concentrations of metabolic products in which carbon atoms are more reduced than in D-glucose were observed in cultures of the *arcB* strains as compared to those of *E. coli* K1060. The most significant differences in terms of yields on D-glucose were observed in acetate and ethanol. The acetate yield on D-glucose in the wild-type strain reached 0.89 ± 0.07 mol·mol-1. While *E. coli* GNB1061 had a similar acetate yield $(0.91 \pm 0.07 \text{ mol} \cdot$

mol⁻¹), *E. coli* GNB1062 and GNB1063 have significantly lower values for this parameter than the wild-type strain (0.68 \pm 0.05 and 0.59 \pm 0.04 mol \cdot mol⁻¹, respectively; *P* < 0.01). On the other hand, the yield of ethanol (the most reduced metabolite) on D-glucose followed the opposite trend. While in *E. coli* K1060 it reached 0.52 \pm 0.02 mol \cdot mol⁻¹, the values for *E. coli* GNB1061, GNB1062, and GNB1063 were, respectively, 0.59 \pm 0.01, 0.81 \pm 0.04, and 0.89 \pm 0.03 mol \cdot mol⁻¹.

The pattern of secreted fermentation metabolites correlates with differences in the redox state in the arcB mutants. As the fate of carbon atoms at the acetyl-CoA node seemed to be markedly different in the strains analyzed, we next studied the ethanol/acetate molar ratios under anoxic growth conditions. Given the difference in the oxidation state of carbon atoms in these fermentation products, their molar ratio provides a good estimation of the redox state of the cell (32, 48). All arcB mutant strains showed significantly higher ethanol/acetate ratios than that obtained for E. coli K1060 (Table 3), which offers a strong indication of an altered redox metabolism in the mutants. Among the partial deletion strains studied, E. coli GNB1062 had the highest ethanol/acetate ratio (i.e., 40% higher than that of the wild-type strain). Again, the $\Delta arcB$ strain was the most severely affected in terms of redox balance, as the observed ethanol/acetate ratio was 53% higher than that of the wild-type strain. In order to further correlate the synthesis of oxidized and reduced fermentation metabolites with the redox state, we measured the actual cellular content of nicotinamide dinucleotides (Table 3). In full accordance with the ethanol/acetate ratios, a significant bias towards higher NADH/NAD+ ratios was observed in the arcB mutants. In particular, E. coli GNB1062 and GNB1063 had similar NADH/NAD+ ratios, which were 44% and 69% higher than that of the wild-type strain, respectively. Importantly, the NAD+ + NADH content of the cells remained almost constant (ca. 6 μ mol · g⁻¹) among the strains under study.

Metabolic flux distribution in central catabolic pathways is differentially affected by the redox state in the *arcB* mutants. Metabolic flux analysis based on stoichiometric constraints constitutes a straightforward way to visualize the operativity of the entire metabolic network in anaerobic *E. coli* cells (4, 45, 55). Under these growth conditions, the carbon source is mostly converted into fermentation products and, to a lesser extent, into biomass (11, 28). We applied this methodology to study the distribution of metabolic fluxes in the *arcB* deletion mutants derived from wild-type K1060, using extracellular fluxes and

specific rates of D-glucose consumption as input for a simplified stoichiometric model of central carbon catabolism (Fig. 2). The actual flux values were normalized to the specific rate of D-glucose consumption of each strain, allowing us to establish comparisons between different strains (Table 4). The values of the carbon balance for each set of fluxes was close to the unit, suggesting a balanced closure between the carbon source consumed by the cells and the formation of biomass, end fermentation products, and CO₂.

In agreement with the experimental $Y_{X/S}$ values (Table 2), the flux towards biomass (from D-glucose-6-P) was consistently lower in all mutant strains when compared to that of $E.\ coli\ K1060\ (P<0.05)$. Fluxes through the Embden-Meyerhof-Parnas pathway up to $P-\ enol$ -pyruvate (v_2 and v_3) were higher in the arcB mutants (in particular for $E.\ coli\ GNB1062$ and GNB1063) than in $E.\ coli\ K1060$. Significant differences were also observed in the fluxes at the $P-\ enol$ -pyruvate/pyruvate and acetyl-CoA metabolic nodes (see below). The NADH-dependent conversion of $P-\ enol$ -pyruvate into succinate through the reductive branch of the TCA cycle (lumped in a single flux in the stoichiometric model, v_5) had a significantly higher value in the arcB mutants than in the wild-type strain. The increment in this flux in the mutants with respect to the wild-type strain ranged between 1.5-fold in $E.\ coli\ GNB1061$ to 2.8-fold in $E.\ coli\ GNB1063$ (P<0.01). Considering that several genes of the TCA cycle are regulated by the ArcBA system (37, 49), and as citrate is the first intermediate of this metabolic pathway, we also quantified its concentration as a coarse estimation of the TCA cycle operativity towards the oxidative branch. As expected for anoxic cultures, the oxidative activity of the TCA cycle was almost null. Citrate concentrations were 0.18 ± 0.01 and 0.37 ± 0.02 mM for $E.\ coli\ GNB1061$ and $E.\ coli\ GNB1062$, respectively, compared to <0.05 mM for the parental strain. In contrast, the citrate concentration in culture supernatants of the $E.\ coli\ GNB1061$ at $E.\ coli\ GNB1062$ at $E.\ coli\ GNB1063$ at $E.\ coli\ GNB1064$ and $E.\ coli\ GNB1065$ and $E.\ coli\ GNB1066$ are specified by the $E.\ coli\ GNB1066$ and $E.\ coli\ GNB1066$ and $E.\ coli\ GNB1066$ and $E.\ coli\ GNB1066$ and $E.\ coli\ GNB1066$ are specified by the $E.\ coli\$

Pyruvate can be either converted into acetyl-CoA and formate through the activity of pyruvate-formate lyase (PfIB), or transformed into D-lactate by the fermentative, NADH-dependent D-lactate dehydrogenase (LdhA) (11). Surprisingly, we detected a small but clearly discernible rate of pyruvate secretion (v_6) in the wild-type strain and E. coli GNB1061, which in both cases accounted for <2% of the specific rate of D-glucose consumption, probably implying a limited processing of this metabolic intermediate via PfIB/LdhA under anoxic conditions. D-lactate synthesis was higher in all arcB strains, which is in good agreement with their unregulated redox state that favors NADH oxidation. In particular, the flux through LdhA (v_7) was the

highest for *E. coli* GNB1063, the $\Delta arcB$ strain; closely followed by that of *E. coli* GNB1062 and representing a 2.9- and 2.1-fold increase, respectively, to the same flux in the wild-type strain (P < 0.01). The rate of pyruvate conversion into acetyl-CoA was similar in all strains, though the corresponding PflB flux (v_8) was slightly higher in the arcB mutants when compared to that of *E. coli* K1060 (P = 0.085). Subsequent processing steps of formate from this reaction were also markedly different among the experimental strains. The evolution of H₂ in these cultures was measured as a direct estimation of the formate-hydrogen lyase activity, which converts formate in H₂ and CO₂ (11). Judging by the values of the v_{10} flux derived from these measurements, the formate-hydrogen lyase activity was the highest in the wild-type strain, and decreased significantly in all the mutant strains. The lowest v_{10} value was observed in *E. coli* GNB1062, and was 1.2-fold lower than the flux in the wild-type strain (P < 0.05).

As explained in the preceding sections, acetyl-CoA can be either converted into acetate or ethanol under anoxic conditions. In *E. coli* K1060, the rate of ethanol synthesis (v_{11}) represents a 65.9% of that for D-glucose uptake, whereas acetate formation (v_{12}) accounts for 82.1%. This distribution between acetate and ethanol reflects the need to reoxidize reducing equivalents formed through the Embden-Meyerhof-Parnas pathway in order to achieve both redox and carbon balances. In line with this rationale, ethanol synthesis was favored over acetate accumulation in the mutants (P < 0.05), which also corresponded qualitatively to the high ethanol/acetate and NADH/NAD+ ratios measured in these strains (Table 3). Again, the more evident differences were observed in *E. coli* GNB1062 and GNB1063. As a quantitative estimation of the molar fraction of carbon diverted from acetyl-CoA towards acetate and ethanol (f), the corresponding split ratios were calculated as $f_{\text{ethanol}} = v_{11}/(v_{11} + v_{12})$ and $f_{\text{acetate}} = v_{12}/(v_{11} + v_{12})$. In *E. coli* K1060, the values for f_{ethanol} and f_{acetate} were 44.5 and 55.5; whereas in *E. coli* GNB1062 and GNB1063 the split ratios were 50.9 and 49.1, and 54.2 and 45.8, respectively. The acetyl-CoA split ratio for ethanol in *E. coli* GNB1061 was slightly higher than that obtained for the wild-type strain, but the difference was barely significant (P = 0.072).

Finally, the availability of reducing power was also deduced at the fluxome level by considering the overall molar NADH availability per equivalent of consumed D-glucose, *i.e.*, $f_{NADH/G} = (2v_5 + v_7 + 2v_{11})/v_0$. The experimental value obtained for this parameter in the wild-type strain was 1.45 ± 0.08 , and the increment in

 $f_{\text{NADH/G}}$ in the mutants ranged from 1.1-fold for *E. coli* GNB1061 to 1.4-fold for *E. coli* GNB1063, thus confirming the *in vitro* measurements of nucleotide content.

Transcriptional analysis and measurement of key enzymatic activities support the observed differences at the fluxome level. In order to study the incremental effects of these deletions on the cell physiology at different regulatory levels, we next determined the transcriptional activity of relevant genes by quantitative RT-PCR as well as the activities of selected enzymes which showed significant differences using the metabolic flux analysis approach (Fig. 3) under the same growth conditions used for those experiments. These two levels of regulation were investigated at key points in different metabolic blocks within the proposed anoxic biochemical network (*i.e.*, Embden-Meyerhof-Parnas pathway, pentose phosphate pathway, TCA cycle, and fermentation pathways) to provide a complete snapshot of the metabolic landscape in each strain.

D-Glucose is split into glycolysis and pentose phosphate pathway at the D-glucose-6-P branching point (Fig. 2). A slight but consistent increase in the expression of pfkA (encoding the glycolytic enzyme 6-phosphofructokinase I) was observed for E. coli GNB1062 and GNB1063 (P < 0.05, Fig. 3A), thus suggesting a higher activity through the initial steps in D-glucose catabolism in the mutant strains when compared to E. coli K1060 and in good agreement with the experimental values of v_3 . The activity of pyruvate kinase (PykAF, that converts P-enol-p-pyruvate into pyruvate) was analyzed to estimate carbon routing through the lower Embden-Meyerhof-Parnas pathway (Fig. 3B). No significant differences were observed in the specific PykAF activity among the strains, which correlates well with the flux values through v_4 (Table 4). In stark contrast, the activity of D-glucose-6-P 1-dehydrogenase (encoded by zwf, and the key enzyme of the oxidative pentose phosphate pathway) was >74% lower in all the mutant strains when compared to E. coli K1060 (Fig. 3B), demonstrating that the flux towards pentoses formation was very low in these strains. Moreover, the enzymatic activity observed in the wild-type strain was <50% than that observed in oxic cultures of E. coli (data not shown).

The next metabolic node studied was the pyruvate branching point. In accordance with the expected transcriptional activation of *focA-pflB* by the ArcBA system during the oxic/micro-oxic transition (46), the

level of pflB mRNA in $E.\ coli$ GNB1063 was 26% lower than that of $E.\ coli$ K1060 (P < 0.05, Fig. 3A). The difference in this parameter for the other mutants and the wild-type strain resulted not significant. In vitro enzymatic activity of PflB demonstrated that this activity is indeed affected by the different arcB mutations tested (Fig. 3B). In particular, the PflB activity was 36% and 27% lower in $E.\ coli$ GNB1062 and GNB1063, respectively, than that measured in $E.\ coli$ K1060 (P < 0.05). The level of activity detected in $E.\ coli$ GNB1061 and $E.\ coli$ K1060 was very similar. To also evaluate the contribution of the pyruvate dehydrogenase complex in the flux conducive to acetyl-CoA formation, we measured the transcriptional activity of aceE, which encodes the E1 component of this enzymatic complex (Fig. 3A). While no differences were observed for $E.\ coli$ GNB1061 as compared to the wild-type strain, the transcriptional level of aceE was incremented 2.4- and 3.6-fold in $E.\ coli$ GNB1062 and GNB1063, respectively (P < 0.01). Even when it is known that pyruvate dehydrogenase could contribute to acetyl-CoA formation under micro-oxic conditions (22), no significant enzymatic activity was detected in cell-free extracts of any of the strains under study (data not shown).

Fermentation pathways that operate at the P-enol-pyruvate/pyruvate metabolic node also showed significant differences among the mutants. Both IdhA (encoding D-lactate dehydrogenase) and frdA (encoding one of the fumarate reductase subunits, the enzymatic complex that converts fumarate into succinate) were strongly upregulated in all arcB mutants (Fig. 3A), attaining mRNA levels >2-fold higher than those observed in E. coli K1060 (P < 0.01). In the $\Delta arcB$ mutant, IdhA expression peaked at ca. 6-fold higher than in the wild-type strain. In contrast, frdA expression was the highest in E. coli GNB1062 (4.5-fold increment, P < 0.01). LdhA activity followed the same trend as that observed at the transcriptional level, being higher in all arcB mutants when compared to E. coli K1060 (P < 0.05, Fig. 3B). However, at least at this level of regulation, no significant differences were observed among mutant strains. Total succinate dehydrogenase (Sdh) activity was also higher in the mutants as compared to E. coli K1060, and the highest activity was observed in E. coli GNB1062, in accordance with the transcriptional regulation results obtained by means of quantitative RT-PCR. The behavior in the strain expressing ArcB¹⁷⁷⁻⁶⁴⁰, quantitatively different from that observed in that context.

The *in vitro* activity of citrate synthase was also evaluated under anoxic conditions to substantiate the results of citrate synthesis discussed above. GltA activity was higher in *E. coli* GNB1062 and GNB1063 than in the wild-type strain and in *E. coli* GNB1061, qualitatively reproducing the results obtained by measuring the extracellular citrate concentration. Although some residual activity of the oxidative branch of the TCA cycle is present in the *arcB* backgrounds analyzed in this study under anoxic conditions, our results clearly show a much higher activity towards succinate formation (*i.e.*, reducing branch).

The regulation at the acetyl-CoA branching point was then assessed by measuring the expression of *adhE* and *ackA*, as well as the corresponding alcohol dehydrogenase and acetate kinase activities. A good correlation between transcriptional regulation and enzymatic activity was observed for *adhE*. Indeed, a significant increment in both parameters was observed in the mutant strains in comparison with wild-type K1060 (*P* < 0.05), which was even more evident for *E. coli* GNB1062 and GNB1063 as compared to *E. coli* GNB1061. In particular, *adhE* transcription levels attained the maximum fold change among all the genes evaluated, being *ca*. 8-fold higher in *E. coli* GNB1063 than in *E. coli* K1060. While in *E. coli* GNB1062 and GNB1063 the levels of *ackA* did not differ significantly from those in wild-type K1060, *ackA* expression had a *ca*. 2.9-fold increment in *E. coli* GNB1061 (Fig. 3A). However, these sharp differences at the transcriptional level did not translate into a similar pattern at the enzymatic activity level. Moreover, AckA activity was lower in all the *arcB* mutants than in *E. coli* K1060, which fits well with the observed fluxes through this pathway (Fig. 3B). Both AdhE and AckA showed a regulatory pattern that qualitatively correlated with the changes observed at the flux level, thus reflecting the crucial competence between the corresponding pathways at the acetyl-CoA branching point in terms of both precursors and reducing power availability.

Finally, the transcriptional level of *arcA* was also evaluated, and no significant differences were observed in its transcription level among the experimental strains. Although the mechanism whereby the ArcB variants determine phenotypic differences in the mutants cannot be directly concluded from the present results and remains to be exposed, the transcriptional analysis suggests that the pattern of regulation of *arcA* is not significantly affected by the deletions introduced in *arcB*.

429 DISCUSSION

Metabolic manipulations to enhance the synthesis of metabolic products include several approaches to increase the availability of substrates needed for its formation or to eliminate competing pathways, which sometimes leads to undesired phenotypes. An alternative strategy that has been scarcely exploited is the network-wide manipulation of metabolic fluxes by means of mutations in global regulators. In this sense, the modularity of the ArcB sensor of the ArcBA two-component system (19), a prototypal member of the bacterial global regulatory network in the model facultative anaerobe *E. coli*, represents an ideal model to explore this approach. The present study analyzed some of the complex genotype-phenotype relationships in mutants of this regulatory system under anoxic growth conditions, which also permits to foresee their potential applicability for the synthesis of reduced biochemicals.

The relative lack of knowledge on the cellular wiring of these regulatory networks under conditions relevant to both laboratory and industrial applications represents a significant hurdle that has to be overcome for the efficient design of industrial processes. A complete analysis of the cellular physiology in different *arcB* mutants is needed in order to understand (and take advantage of) the metabolic effects of *arcB* mutations *in vivo*. Metabolic and physiological variations resulting from different deletions in the global regulator ArcB were systematically analyzed by combined physiological, transcriptomic, and fluxome analysis. This combined approach is relevant since deletion of global regulatory genes were observed to affect the entire cellular and metabolic landscape in a rather difficult-to-predict fashion.

The Dye phenotype served as an *a priori* indication of differences at the metabolic level among the mutants. Indeed, the overall colony morphology under these growth conditions was in good agreement with the altered metabolic patterns observed in each strain; while $E.\ coli$ GNB1062 showed a Dye phenotype compatible with that of the $\Delta arcB$ strain, $E.\ coli$ GNB1061 presented an intermediate phenotype, somewhat closer to that of the wild-type strain. These morphological alterations are likely to arise from differences in the redox homeostasis, as previously suggested (3, 42). In line with this hypothesis, all the mutants had a high NADH/NAD+ ratio and the most significant differences were observed in the NADH content, as the NAD+ content did not show changes in the different strains. In close connection with this trait, the split of

the acetyl-CoA between ethanol and acetate was predictably affected by the redox state measured in each strain (see below).

The regulatory pattern of some fluxes (also observed at both the transcriptional and enzymatic activity levels) significantly departs from that reported for arcA mutants under conditions with restricted O₂ supply (32, 59), thus evidencing that elimination of the entire arcB coding sequence (or sequences encoding different ArcB domains) has a different effect on the cell physiology as compared to the absence of the cognate response regulator. For instance, the high glycolytic fluxes observed in the mutants (especially in E. coli GNB1062 and GNB1063) result in an elevated NADH generation rate that cells need to recycle to efficiently continue D-glucose catabolism; therefore, anoxic fermentation pathways have to fulfill the requirement for NADH regeneration under these culture conditions (11). Although the increase in NADH in the mutants can be mostly attributed to the activity of the Embden-Meyerhof-Parnas pathway, it is worth considering that it can also arise from either a high activity of the TCA cycle enzymes and/or a low activity of the electron transfer chain. Our results from transcriptomic analysis as well as enzymatic activity measurements support some residual contribution of the TCA cycle to the redox balance, and the repression of cyoABCDE (that encodes cytochrome o) by the ArcBA system is well known (26, 54). However, the activity of the oxidative branch of the TCA cycle under the experimental conditions explored here is expected to be low, considering the inhibition exerted by the high NADH/NAD+ ratios themselves on both GltA and 2-ketoglutarate dehydrogenase activities (37, 58).

The pyruvate/acetyl-CoA branching points showed the most striking alterations among the mutant strains. PflB is known to convert pyruvate into acetyl-CoA under conditions with restricted O₂ supply (1), but recent studies suggest that some activity of the pyruvate dehydrogenase complex also contributes to anoxic acetyl-CoA formation (22). Under the working conditions tested in this study, pyruvate produced during glycolysis seems to be mainly processed by pyruvate-formate lyase to generate acetyl-CoA. Since the activity of the pyruvate dehydrogenase complex (the other source of acetyl-CoA from pyruvate besides PflB) is inhibited by NADH (22), and higher ethanol/acetate and NADH/NAD+ redox ratios were detected in all *arcB* strains compared to the wild-type strain, the flux through pyruvate dehydrogenase would contribute to acetyl-CoA only marginally. In turn, the acetyl-CoA metabolic node, at which carbon atoms from D-

glucose catabolism can be either converted into an oxidized or a reduced fermentation metabolite (*i.e.*, acetate or ethanol, respectively), is of paramount interest for the synthesis of various heterologous metabolites of industrial interest. By evaluating the flux values conducing to ethanol and acetate, as well as the flux split ratios for these two metabolites, it can be seen that the synthesis of reduced biochemicals is favored in all the mutants.

At this point, it is relevant to consider that the actual phenotype of each strain (*i.e.*, the macroscopic distribution of metabolic fluxes and other phenotypic traits) is the final consequence of multiple (and very complex) regulatory processes that act hierarchically at different levels (38, 40). Several fine-tuning mechanisms for metabolic modulation, such as allosteric regulation of enzyme activity, might well operate differently in the strains analyzed. However, one can safely assume that the gross regulatory mechanisms, other than that exerted by the ArcBA system itself, are similar in the parental strain and its *arcB* derivatives. The mechanism underlying the phenotypic differences observed could lie beyond the known phosphorelay process described *in vitro* for the cellular signaling mediated by the ArcBA system (24, 27). The possibility that similar (or radically different) effects on central metabolism could be observed in partial-deletion mutants of genes encoding components of other signal transduction systems in *E. coli* is an exciting scenario that remains to be explored.

Different types of mutations in components of global regulatory systems, such as *arcA* and *creB* (29-31, 33), have been shown to influence both carbon and redox balances in *E. coli*, mainly under micro-oxic growth conditions. The results of the current study focus on the manipulation of global regulators as a relevant tool to modulate central metabolic fluxes under anoxic conditions and to harness reducing power availability for biotechnological purposes. The unregulated redox state of the mutant strains provides diverse metabolic backgrounds for the synthesis of reduced biochemicals both native to *E. coli*, such as ethanol, D-lactate, and succinate; and those resulting from heterologous pathways, such as poly(3-hydroxybutyrate). Moreover, the use of targeted deletions in the ArcB tripartite sensor protein enabled us to obtain increasing phenotypic effects that could be further exploited for the synthesis of reduced biochemicals, such as those mentioned above.

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FIGURE LEGENDS

FIG. 1 (A) Schematic representation of the ArcB protein. The different modules in the sensor protein are indicated with boxes, along with the amino acid coordinates they span. Individual amino acids relevant in the intramolecular phosphorelay that passes the phosphate residue among the different ArcB domains (which ultimately leads to phosphorylation of ArcA) are shown below the corresponding modules in the protein. Note that the elements in this outline are not drawn to scale. TM, transmembrane domain; LZ, leucine zipper. (B) Dye phenotype of the strains under study on toluidine blue O medium. Strains were firstly grown in LB medium under oxic conditions and adequate dilutions in cold saline were spotted onto the plates as previously described (9, 42). Plates were photographed after overnight incubation at 37°C, and representative colonies resulting from the 10-6 dilution are shown at a 13.4× magnification.

FIG. 2 Metabolic network used to study central catabolism of D-glucose in different E. $coli\ arcB$ mutants under anoxic conditions. Fluxes within the network are codified as v_0 (which corresponds to the specific rate of D-glucose consumption) to v_{12} . Note that in this representation some biochemical reactions (e.g., those generating biomass precursors from D-glucose-6-P) are lumped into a single flux in order to simplify the representation of the biochemical network. Enzymes that were assayed *in vitro* are highlighted in gray in the corresponding flux at which they are involved, and genes quantified at the transcriptional level are shown in a gray box beside their associated flux. The subscript res indicates accumulation of either residual pyruvate or formate in the culture medium. CoA, coenzyme A.

FIG. 3 Transcriptional analysis of relevant genes and *in vitro* enzymatic activity measurements during exponential growth of E. coli K1060 (wild-type strain) and its arcB mutant derivatives [GNB1061 (ArcB²⁶⁸⁻⁵²⁰), GNB1062 (ArcB¹⁷⁷⁻⁶⁴⁰), and GNB1063 ($\Delta arcB$)] in anoxic batch cultures developed in M9 minimal medium with 3% (wt/vol) D-glucose as the carbon source. (A) Results from quantitative RT-PCR analysis in the mutant strains are normalized to the corresponding transcript levels in E. coli K1060, and represent the mean value \pm standard deviation of sextuplicate measurements from at least three independent cultures. (B) Results from determinations of specific enzymatic activities represent the mean value \pm standard deviation of triplicate measurements from at least two independent cultures. Note that the activity of

pyruvate kinase (PykAF) is separately represented as the corresponding values were much higher than those obtained for the rest of the enzymes assayed. Sp, specific.

TABLE 1 *E. coli* strains, oligonucleotides, and plasmids used in this study.

_	\sim	\sim
•	11	11

Strain,	Relevant characteristics or sequence (5'→3')	Source or reference				
oligonucleotide,						
or plasmid						
E. coli						
K1060ª	Considered wild-type in this study; F- fadE62 lacI60 tyrT58(AS)	(36)				
	fabB5 mel-1					
GNB1061	Same as K1060, but ArcB ²⁶⁸⁻⁵²⁰ ; confined deletion between 802-	This study				
	1560 nt of arcB					
GNB1062	Same as K1060, but ArcB ¹⁷⁷⁻⁶⁴⁰ ; confined deletion between 529-	This study				
	1920 nt of arcB					
GNB1063	Same as K1060, but ∆ <i>arcB</i>	This study				
Oligonucleotide ^t						
$\Delta PAS ext{-}F$	TCC TTG ATG CTT CAC CCG ACC TGG TTT TTT ATC GTA	This study				
	ACG AAG ATA AAG AG G TGT AGG CTG GAG CTG CTT C					
ΔD1-R	TCT TCT GTC GTC ACC GTA CTC TCC TCA TCA TCC TGG	This study				
	GTA TCC CAG AAT TT C ATA TGA ATA TCC TCC TTA G					
∆H1-F	ACC GCG TGG GTA AAC GTC ACG GTT TGA TGG GCT TTG	This study				
	GTC GCG ACA TTA CC G TGT AGG CTG GAG CTG CTT C					
∆H1-R	TTC AGT TCA ATG TCT TCC ACC AGC AGC ACA TTC AGC	This study				
	GCC GGT AAA GGC AT C ATA TGA ATA TCC TCC TTA G					
∆arcB-F	AAC GTA ACT GTC AGA ATT GGG TAT TAT TGG GGC AGG	This study				
	TTG TCG TGA AGG AA G TGT AGG CGT GAG CTG CTT C					
<i>∆arcB</i> -R	ATA ATA ATT TAC GGC CGA GCC AAG ATT TCC CTG GTG	This study				
	TTG GCG CAG TAT TC C ATA TGA ATA TCC TCC TTA G					
arcB1-C-F	GGG TAT TAT TGG GGC AGG TT	This study				
arcB1-C-R	GTC TAG CCG GGG TCA TTT TT	This study				
arcB2-C-F	AAT GAT TCG CCA TAC GCC AC	This study				

arcB2-C-R	GTG CTG TGC CCT TGT AAC TC	This study
PDH-C-F	GAG CAA CTG GAG GAG TCA CG	This study
PAS-C-R	GCA AGG AAG CTG GTG AAA TC	This study
H1-C-R	GGT GAT CAG CTT CGG TCC TA	This study
D1-C-R	CGG AAG GTC AGG AGA CTG AA	This study
rrsA-RT-F	AGG CCT TCG GGT TGT AAA GT	This study
rrsA-RT-R	ATT CCG ATT AAC GCT TGC AC	This study
pfkA-RT-F	GGT GCC TTA CGA CCG TAT TC	This study
pfkA-RT-R	GGA CGC TTC ATG TTT TCG AT	This study
IdhA-RT-F	AGT CCG TGT TCC AGC CTA TG	This study
IdhA-RT-R	CGG TCA GAC CTT CCA GAG AG	This study
pflB-RT-F	GCG AAA TAC GGC TAC GAC AT	This study
pflB-RT-R	CAT CCA GGA AGG TGG AGG TA	This study
ackA-RT-F	CGT TGA CGC AAT CAA CAA AC	This study
ackA-RT-R	GGT GGC AGT AAA CG TCC ATT	This study
adhE-RT-F	CTG GCA GGC TTC TCT GTA CC	This study
adhE-RT-R	TAC CGC GTC TTC GAA ATC TT	This study
frdA-RT-F	CGA TAA GAC CGG CTT CCA TA	This study
frdA-RT-R	CCT TCC ATC ATG TTC ATT GCT	This study
fdhF-RT-F	AAA AAG TCG TCA CGG TTT GC	This study
fdhF-RT-R	TTC GCG CAG GCG AGT TTT	This study
arcA-RT-F	TGT TTT CGA AGC GAC AGA TG	This study
arcA-RT-R	GAA CAT CAA CGC AAC ATT CG	This study
Plasmid		
pKD46	Vector containing the λ -Red (γ , β , and exo) recombination	(12)
	functions under control of the P _{araB} promoter; oriR101,	
	repA101(Ts), Apr	
pKD4	Vector used as template for amplification of FRT-aphA-FRT;	(12)
	oriR6Kγ, Ap ^r Km ^r	

pCP20	Vector expressing the FLP recombinase from Saccharomyces	(10)
	cerevisiae; λ cl857 λ P _R FLP repA(Ts), Apr Cmr	

- ^a Strain obtained through the E. coli Genetic Stock Center, University of Yale, CT.
- b Sequences with homology to FRT-aphA-FRT in the template plasmid pKD4 are shown in boldface.
 Oligonucleotides used for deletions are preceded by the symbol Δ, and those used to check deletions are
 followed by the letter C. Oligonucleotides used for quantitative RT-PCR experiments are codified according
 to target gene-RT-F/R.

TABLE 2 Fermentation and growth parameters^a for 24-h anoxic batch cultures in M9 minimal medium with 3% (wt/vol) D-glucose as the carbon source.

E. coli strain	Relevant	Biomass	$\mu_{max}{}^b$	Sp rate of D-glucose	$Y_{X/S}$
	characteristics	(g · liter-1)	(h ⁻¹)	consumption ^b	(mg · g-1)
				$(mmol \cdot g^{-1} \cdot h^{-1})$	
K1060	Wild-type strain	0.65 ± 0.08	0.34 ± 0.02	7.22 ± 0.05	79 ± 2
GNB1061	ArcB ²⁶⁸⁻⁵²⁰	0.43 ± 0.04	0.27 ± 0.03	6.07 ± 0.04	54 ± 2
GNB1062	ArcB ¹⁷⁷⁻⁶⁴⁰	0.26 ± 0.03	0.16 ± 0.01	5.31 ± 0.08	27 ± 1
GNB1063	∆arcB	0.18 ± 0.02	0.11 ± 0.02	4.85 ± 0.06	21 ± 3

 $[^]a$ Reported results represent the mean value \pm standard deviation of triplicate measurements from at least two independent cultures.

^b The specific growth rate and the specific rate of D-glucose consumption were determined during balanced growth.

TABLE 3 Redox parameters^a for 24-h anoxic batch cultures in M9 minimal medium with 3% (wt/vol) D-glucose as the carbon source.

E. coli strain	Relevant	Intracellular content		NADH/NAD+ ratio	Ethanol/acetate
	characteristics	μ mol · g ⁻¹) of:		(mol·mol-1)	ratio (mol·mol-1)
		NADH	NAD+		
K1060	Wild-type strain	2.03 ± 0.09	3.96 ± 0.07	0.51 ± 0.05	0.53 ± 0.05
GNB1061	ArcB ²⁶⁸⁻⁵²⁰	2.14 ± 0.08	3.45 ± 0.21	0.62 ± 0.04	0.63 ± 0.03
GNB1062	ArcB ¹⁷⁷⁻⁶⁴⁰	2.91 ± 0.12	3.93 ± 0.15	0.74 ± 0.08	$\textbf{0.74} \pm \textbf{0.03}$
GNB1063	Δ arcB	2.61 ± 0.05	3.02 ± 0.09	0.86 ± 0.06	0.81 ± 0.02

 $[^]a$ Reported results represent the mean value \pm standard deviation of duplicate measurements from at least two independent cultures.

Metabolic flux distribution in anoxic batch cultures of E. coli K1060 and its arcB mutant TABLE 4 derivatives during balanced growth in M9 minimal medium with 3% (wt/vol) D-glucose as the carbon source.

Flux	Flux to:	Relative flux ^a for <i>E. coli</i> :			
		K1060	GNB1061	GNB1062	GNB1063
		(wild-type strain)	(ArcB ²⁶⁸⁻⁵²⁰)	(ArcB ¹⁷⁷⁻⁶⁴⁰)	(∆arcB)
V 0 ^b	D-Glucose-6-P	100	100	100	100
V 1	Biomass	21.1 ± 0.9	19.9 ± 0.2	16.7 ± 0.4	13.8 ± 0.6
/ 2	D-Glyceraldehyde-3-P	78.9 ± 1.3	80.1 ± 1.4	83.3 ± 1.5	86.2 ± 1.9
′ 3	P-enol-pyruvate	157.8 ± 2.5	160.3 ± 2.9	166.7 ± 1.8	172.4 ± 3.6
′ 4	Pyruvate	152.9 ± 0.8	153.1 ± 0.9	156.5 ± 0.9	159.2 ± 0.4
/ 5	Succinate	4.8 ± 0.5	7.2 ± 0.6	10.2 ± 0.7	13.2 ± 0.3
′ 6	Residual pyruvate	1.7 ± 0.3	1.8 ± 0.1	0.0 ± 0.1	0.0 ± 0.2
/ 7	D-Lactate	3.2 ± 0.4	4.6 ± 0.5	6.6 ± 0.4	8.5 ± 0.1
′ 8	Acetyl-CoA	148.1 ± 0.7	148.4 ± 0.3	149.9 ± 0.9	150.7 ± 0.9
′ 9	Residual formate	79.5 ± 0.9	86.7 ± 1.1	91.3 ± 0.7	91.5 ± 1.2
′ 10	H ₂	68.6 ± 1.2	61.8 ± 0.9	58.6 ± 1.4	59.2 ± 0.6
/11	Ethanol	65.9 ± 0.8	68.7 ± 0.4	76.3 ± 0.5	81.6 ± 0.3
/12	Acetate	82.1 ± 0.7	79.7 ± 0.7	73.6 ± 0.2	69.1 ± 0.2
	Carbon balance	0.93 ± 0.04	0.96 ± 0.02	0.91 ± 0.12	0.92 ± 0.09

⁷²⁹ 730

725

726

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731

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^a The relative flux values were normalized to the specific rate of D-glucose consumption in each strain (v_0), and the reported results represent the mean value \pm 90% confidence intervals calculated using triplicate measurements of extracellular fluxes in at least two independent cultures. Fluxes were codified according to the biochemical reactions as shown in Fig. 2.

 $[^]b$ Absolute flux values for v_0 are as shown in Table 2.

 c Carbon balances were calculated from experimental fluxes to extracellular metabolites, inferred rate of CO₂ evolution, and biomass generation. The reported results represent the mean value \pm standard deviation calculated from the flux values within the metabolic network.





