Formate hydrogen lyase mediates stationary-phase deacidification and increases survival during sugar fermentation in acetoin-producing enterobacteria

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Two fermentation types exist in the Enterobacteriaceae family. Mixed-acid fermenters produce substantial amounts of lactate, formate, acetate, and succinate, resulting in lethal medium acidification. On the other hand, 2,3-butanediol fermenters switch to the production of the neutral compounds acetoin and 2,3-butanediol and even deacidify the environment after an initial acidification phase, thereby avoiding cell death. We equipped three mixed-acid fermenters (Salmonella Typhimurium, S. Enteritidis and Shigella flexneri) with the acetoin pathway from Serratia plymuthica to investigate the mechanisms of deacidification. Acetoin production caused attenuated acidification during exponential growth in all three bacteria, but stationary-phase deacidification was only observed in Escherichia coli and Salmonella, suggesting that it was not due to the consumption of protons accompanying acetoin production. To identify the mechanism, 34 transposon mutants of acetoin-producing E. coli that no longer deacidified the culture medium were isolated. The mutations mapped to 16 genes, all involved in formate metabolism. Formate is an end product of mixed-acid fermentation that can be converted to H2 and CO2 by the formate hydrogen lyase (FHL) complex, a reaction that consumes protons and thus can explain medium deacidification. When hycE, encoding the large subunit of hydrogenase 3 that is part of the FHL complex, was deleted in acetoin-producing E. coli, deacidification capacity was lost. Metabolite analysis in E. coli showed that introduction of the acetoin pathway reduced lactate and acetate production, but increased glucose consumption and formate and ethanol production. Analysis of a hycE mutant in S. plymuthica confirmed that medium deacidification in this organism is also mediated by FHL. These findings improve our understanding of the physiology and function of fermentation pathways in Enterobacteriaceae.

Keywords: mixed-acid fermentation, 2,3-butanediol fermentation, acid stress, hydrogenase 3, formate hydrogen lyase

INTRODUCTION
Within the Enterobacteriaceae family, a distinction is made between mixed-acid (e.g., Escherichia, Salmonella, and Shigella) and 2,3-butanediol fermenters (e.g., Klebsiella, Serratia, and Enterobacter) based on their fermentation end products produced during sugar fermentation. Mixed-acid fermenters ferment sugars to ethanol and a range of organic acids, including lactate, succinate, acetate, and formate. Formate can be further converted to H2 and CO2 by the formate hydrogen lyase (FHL) complex (White, 2000). Mixed-acid fermentation generally leads to rapid and strong medium acidification and even cell death. On the other hand, 2,3-butanediol fermenters use the mixed-acids pathway only during the early growth phase, and switch in the late exponential phase to a different fermentation pathway, in which pyruvate is converted to the neutral end products acetoin or 2,3-butanediol, thereby preventing excessive acidification (Van Houdt et al., 2006; Xiao and Xu, 2007). Moreover, after the initial decline of medium pH, 2,3-butanediol fermenters typically deacidify the medium toward more neutral values during stationary phase (Johansen et al., 1975; Yoon and Mekalanos, 2006; Van Houdt et al., 2007; Moons et al., 2011). This is in contrast to mixed-acid fermenters or 2,3-butanediol fermenters with an inactivated 2,3-butanediol pathway, where a sustained pH decrease is usually observed during sugar fermentation (Yoon and Mekalanos, 2006; Moons et al., 2011). Thus, 2,3-butanediol fermentation is apparently associated with stationary-phase deacidification. Synthesis of 2,3-butanediol from pyruvate requires three steps. First, the conversion of two molecules of pyruvate to α-acetolactate is catalyzed by the α-acetolactate synthase (α-ALS). Next, α-acetolactate is decarboxylated to acetoacetanilide by the α-acetolactate deacylase
(α-ALD). In a last step, acetoin is reduced to 2,3-butanediol by the 2,3-butanediol dehydrogenase (BDH), which can also catalyze the reversed reaction. Each of these three reactions consumes an intracellular proton, and this potentially explains the observed stationary-phase deacidification. In *Serratia plymuthica* RVH1, a strain previously isolated from a food processing environment (Van Houdt et al., 2005), α-ALS and α-ALD are encoded by the *budB* and *budA* genes, respectively, which are located on the *budAB* operon (Moons et al., 2011). We previously showed that transfer of the *S. plymuthica* RVH1 *budAB* operon conveys the capacity to produce acetoin, to prevent lethal medium acidification and to reverse acidification (Vivijs et al., 2014a). In the present study, we transferred the *budAB* operon to some additional mixed-acid fermenting enterobacteria, *Salmonella* Typhimurium, *Salmonella* Enteritidis, and *Shigella flexneri*, and show that these also acquire the capacity to produce acetoin. However, acetoin production was not associated with stationary-phase deacidification in *S. flexneri*. This observation is remarkable since *Shigella* and *E. coli* are considered as a single species based on DNA homology (Fukushima et al., 2002). Thus, our results suggested the involvement of a deacidification mechanism different from proton consumption during acetoin production. To identify this mechanism, we performed random transposon mutagenesis in *budAB*-containing *E. coli* searching for mutants that lost their stationary-phase deacidification capacity but still produced acetoin. This led us to identify the FHL complex as the primary deacidification mechanism in 2,3-butanediol-fermenting *Enterobacteriaceae*.

### MATERIALS AND METHODS

#### BACTERIAL STRAINS, PLASMIDS, OLIGONUCLEOTIDES, AND GROWTH CONDITIONS

The bacterial strains and plasmids used in this study are listed in **Table 1**. All bacteria were cultured in lysogeny broth (LB; 10 g/l tryptone, 5 g/l yeast extract, 5 g/l NaCl) or on LB agar (15 g/l agar) at 37°C except *Serratia plymuthica*, which was grown at 30°C. Media were supplemented with the following chemicals (Applichem, Darmstadt, Germany) when appropriate: 5 g/l

### Table 1 | Strains and plasmids used in this study.

| Strain or plasmid | Relevant features | Reference |
|-------------------|------------------|-----------|
| **Strains**       |                  |           |
| *Escherichia coli*|                  |           |
| S17-1 λpir        | *pro thi recA hsdR− hsdM+ RP4: 2-Tc:Mu: Km Tn7 λpir* | Simon et al. (1983) |
| DH5∞              | F− *endA1 hsdR17 (r−, m−) supE44 thi−1 h− recA1 gyrA96 relA1 deoR ΔlacZYA-argF(U169 Pho80 lacZΔM15* | Grant et al. (1990) |
| MG1655            | F− λ− rph-1 Δ*hycE* | Guyer et al. (1981) |
| MG1655 hycE       | Δ*hycE*          |           |
| Typhimurium LT2   | Wild-type        | McClelland et al. (2001) |
| Enteritidis ATCC 13076 | Wild-type | Tindall et al. (2005) |
| *Shigella flexneri*|                  |           |
| ATCC 12022        | Wild-type; serotype 2b | Daligault et al. (2014) |
| *Serratia plymuthica*|                |           |
| RVH1              | Wild-type; biofilm isolate from food processing plant | Van Houdt et al. (2005) |
| RVH1 budAB        | Δ*budAB:cat, CmR* | Vivijs et al. (2014b) |
| RVH1 hycE         | Δ*hycE*          |           |
| **Plasmids**      |                  |           |
| pTrc99A           | Cloning vector carrying IPTG-inducible trc promoter (P*trc*); Ap*R* | Amann et al. (1988) |
| pTrc99A-P*trc-budAB| pTrc99A carrying the *S. plymuthica* RVH1 budAB operon downstream of P*trc*; Ap*R* | Moons et al. (2011) |
| pKD3              | Template plasmid containing *cat* gene flanked by FRT sites; Cm*R* Ap*R* | Datsenko and Wanner (2000) |
| pKD46             | Plasmid expressing γ, β, and exo recombination genes of phage λ under control of P*BAD*; temperature-sensitive replicon; Ap*R* | Datsenko and Wanner (2000) |
| pCP20             | Plasmid expressing the FLP (flippase) gene, directing recombination of FRT sites; Ap*R* Cm*R* | Datsenko and Wanner (2000) |
| pUC18             | Cloning vector; Ap*R* | Laboratory collection |
| pUCGmlox          | pUC18-based vector containing thelox-flanked aacC7 gene; Ap*R* Gm*R* | Quénéé et al. (2005) |
| pSF100            | pGP704 suicide plasmid; *pir* dependent; Ap*R* Km*R* | Rubirés et al. (1997) |
| pCM157            | cre expression vector; Tc*R* | Marx and Lidstrom (2002) |
glucose; 100 µg/ml ampicillin (Ap); 200 µg/ml carbenicillin (Cb); 30 µg/ml chloramphenicol (Cm); 5 µg/ml gentamicin (Gm); 10 µg/ml tetracycline (Tc); 50 µg/ml kanamycin (Km); and 1 mM isoprpyl-β-D thiogalactopyranoside (IPTG). Plasmids pTrc99A and pTrc-99A-Ptrc-budAB were introduced into the mixed-acid fermenters by electroporation. All oligonucleotides used in this work are listed in Table 2, and were purchased from IDT (Haasrode, Belgium).

SCREENING FOR MUTANTS THAT HAVE LOST STATIONARY-PHASE DEACIDIFICATION CAPACITY

A random knockout library of E. coli MG1655 containing pTrc99A-P-rc-budAB was constructed using λNK1324, which carries a mini-Tn10 transposon with a Cm resistance gene, according to the protocol described by Kleckner et al. (1991). The mutants were subsequently grown in 300 µl LB medium with glucose, IPTG, Ap, and Cm in a 96-well plate. The plates were sealed with an oxygen impermeable cover foil and incubated without shaking at 37°C. After 24 h, medium acidification was analyzed by adding 5 µl of a 0.06% w/v methyl red solution in 60% v/v ethanol to 200 µl culture (MR test). For mutants that no longer acidified the medium, the remaining 100 µl culture was subjected to the Voges–Proskauer (VP) test by adding 30 µl of 5% w/v α-naphthol and 10 µl of 40% w/v KOH to 100 µl of culture. To quantify acetoin production, the mixture was stirred vigorously after 1 h and the optical density at 550 nm (OD550) was measured. Acetoin concentrations were determined using a standard curve relating the OD550 with the acetoin concentration in LB medium. From mutants that no longer acidified the medium, the remaining 100 µl culture was subjected to the lambda red recombinase system described by Datsenko and Wanner (2000), followed by removal of the introduced antibiotic resistance cassette using the FRT/FLP recombination system. Briefly, 70-bp PCR primers were designed comprising a 50-bp part complementary to the region down- or upstream of hycE and a 20-bp 3′ part allowing amplification of the FRT-flanked Cm resistance cassette present in the plasmid pKD3. The purified PCR product was electrotransformed into E. coli MG1655 containing the pKD46 plasmid providing the lambda red recombinase. The resistance cassette was subsequently removed by expression of the flippase recombination enzyme (FLP) of the FRT/FLP recombination system on the temperature-sensitive pCP20 plasmid.

To delete the hycE gene in S. plymuthica RVH1, a fragment encompassing 643 bp upstream and 559 bp downstream of the gene was PCR-amplified using primers SP_HycE_1(XbaI) and SP_HycE_2(XbaI), cut with XbaI, ligated into a XbaI-digested pUC18 vector and transformed into E. coli DH5α. The resulting plasmid pUC18-hycE was used as a template for PCR using the outward-oriented primers SP_HycE_3(XhoI) and SP_HycE_4(XhoI). In a separate reaction, the loxP flanked Gm resistance cassette from plasmid pUCGmlox was amplified using primers LoxP_Gm_1(XhoI) and LoxP_Gm_2(XhoI). Both PCR products were then cleaved with XhoI and ligated together, generating pUC18-hycE::aacC1, which was transformed in E. coli DH5α. The hycE::aacC1 insert from this plasmid was then amplified using primers SP_HycE_1(XbaI) and SP_HycE_2(XbaI), cut with XbaI, ligated into a XbaI-digested pSF100 vector and transformed into E. coli S17-1 λpir. After conjugation of the resulting plasmid pSF100-hycE::aacC1 into S. plymuthica RVH1 (which does not support replication of this suicide plasmid), transconjugants were selected on LB agar with Km at 15°C. This temperature allows good growth of S. plymuthica but prevents growth of E. coli S17-1 λpir. Loss of Km resistance (pSF100 marker) was assessed by replica plating on LB agar with Km. The Gm resistance cassette was then spliced out using the cre recombinase on plasmid pCM157, which catalyzes site specific recombination between loxP sites. Restriction endonucleases and T4 DNA ligase were purchased from Thermo Scientific (St. Leon Rot, Germany) and used according to the supplier’s instructions.

CHARACTERIZATION OF FERMENTATIVE GROWTH AND FERMENTATION END PRODUCTS

Strains were first grown overnight at the appropriate incubation temperature in 4 ml LB. For strains containing pTrc99A or pTrc99A-P-rc-budAB, Ap was added to ensure plasmid

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Table 2 | Oligonucleotides used in this study.

| Primer          | Sequence (5′-3′)                                      |
|-----------------|------------------------------------------------------|
| Linker 1        | TTTCTGTCGGAATTCAGCTTCTAACGATGTAC                    |
| Linker 2        | TGTCCCCGTACATCGTTAGAACTACTCTGTACC                   |
| Y linker primer | ATCCACAT                                            |
| NK_Cm_DWN      | CCTCCCAAGACGCTGATA                                  |
| EC_HyeE_pKD3_1  | GCGTGGCGGTGGTTAGAAGCTGACTTTTTTTGAAAGGTT             |
| EC_HyeE_pKD3_2  | TTTTATTGGCCGTGCTGTCCTCTGGGGCGGGTGATTA               |
| SP_HyeE_1(XbaI) | GCAATCTGAAATACGCTGTTGGTTCTGGAT                    |
| SP_HyeE_2(XbaI) | AACTCTGTAGTTACTGCTGCGGTTGAC                       |
| SP_HyeE_3(XhoI) | GGGACCGAGCATGTGTTACCTACTGTGAAAGATG                 |
| SP_HyeE_4(XhoI) | GACACTGAGCGGAAAAACGCACCATGCC                      |
| LoxP_Gm_1(XhoI) | AACCTGAGCGTCAAGCTAGTTAAC                           |
| LoxP_Gm_2(XhoI) | AACCTGAGCGGAAAAACGCACCATGCC                      |
maintenance. Since *S. plymuthica* RVH1 is somewhat Ap resistant, Cb was used instead of Ap. Next, the cultures were diluted 1:1000 in tubes containing 30 ml LB with glucose and, when appropriate, IPTG and Ap or Cb. Five ml of paraffin oil was layered on top of the cultures to create anaerobic conditions and the tubes were incubated at the appropriate incubation temperature for 48 h. The cultures were sampled at regular time points to determine cell concentrations, medium pH and acetoin concentration, and for analysis of fermentation end products. Plate counts were determined by spot-plating (5 μl) a decimal dilution series in potassium phosphate buffer (10 mM; pH 7.00) on LB agar. Gas production was evaluated qualitatively using Durham tubes. Fermentation end products were analyzed in 600 μl culture supernatants stored at –20°C. Succinic, lactic, formic, and acetic acid, ethanol, and glucose were determined via high-performance liquid chromatography (HPLC; Agilent 1200 series) using an ion exclusion column (Aminex® HPX-87H) maintained at 55°C, and with 5 mM H2SO4 as the mobile phase (0.6 ml/min). The system was equipped with a refractive index detector operating at 35°C and a diode array detector set at 210 nm.

**STATISTICAL ANALYSIS**

All experiments were carried out in triplicate using independent cultures, and results are presented as the mean values ± SD. Statistical significance between mean values were determined by Student’s *t*-test analysis using the Microsoft Excel statistical package. Results were reported as significant when a *p*-value of <0.05 was obtained, based on a two-sided *t*-test with unequal variance.

**RESULTS AND DISCUSSION**

**INTRODUCTION OF ACETOIN SYNTHESIS PATHWAY IN MIXED-ACID FERMENTERS**

Previously, we introduced the *budAB* operon from *S. plymuthica* RVH1, encoding the α-ALS and α-ALD of the acetoin synthesis pathway, in *E. coli* MG1655 and observed that this attenuated lethal medium acidification during fermentative growth on glucose (Vivijs et al., 2014a). Here, we extended this experiment to *S. Typhimurium*, *S. Enteritidis*, and *S. flexneri* by introducing the pTrc99A-Ptrc-*budAB* plasmid into these organisms to see whether other mixed-acid fermenters would show a similar behavior. Figure 1 shows the growth curves and medium pH during fermentative growth in glucose-containing LB medium of these (squares) and medium pH (triangles) of strains harboring pTrc99A (gray) or pTrc99A-Ptrc-*budAB* (black) are shown. Pictures below the figures show the results of the VP test, with a red color indicating the presence of acetoin. Error bars represent SD.
bacteria with and without the budAB genes. As expected, _E. coli_, both _Salmonella_ strains and _S. flexneri_ without budAB strongly acidified the medium (to pH 4.50–4.70 after 48 h) and this resulted in cell death during the stationary phase. Introduction of the budAB genes did not change growth of the bacteria until stationary phase was reached, but it changed the pH profile of the _E. coli_ and _Salmonella_ cultures in two aspects. Firstly, the acidification during the growth phase was less strong, reaching a minimum pH of about 5.60. Secondly, the pH increased again during stationary phase, up to 6.60–7.00 after 48 h. As a result, plate counts remained almost constant once they had reached their maximal stationary phase level (10–48 h).

Surprisingly, a different pattern was observed in _S. flexneri_. Introduction of the budAB genes also attenuated medium acidification during the growth phase (pH 5.60 after 10 h), but no deacidification occurred during stationary phase. As a result, this culture reached a final pH of 4.80 after 48 h and the plate counts decreased to a similar extent as those of the strain without budAB genes. The strain with the budAB genes produced acetoin in similar amounts as the _E. coli_ and _Salmonella_ strains carrying these genes, so that poor expression of the acetoin pathway could be ruled out to explain the different behavior of _S. flexneri_. Therefore, proton consumption in the acetoin production pathway cannot fully explain the deacidification during stationary phase in _E. coli_ and _Salmonella_, and it can be concluded that other deacidification mechanisms must be involved.

**SCREENING FOR LOSS OF DEACIDIFICATION CAPACITY IN _E. coli_ CONTAINING A FUNCTIONAL ACETOIN PATHWAY**

In order to identify additional mechanisms involved in stationary-phase deacidification, we performed random transposon mutagenesis in _E. coli_ MG1655 containing the pTrc99A-P<sub>trc</sub>-budAB plasmid and searched for mutants that were unaffected in acetoin production (VP test), yet were no longer able to increase the pH of glucose-containing LB medium at 37°C after 24 h (MR test), thus having a MR+/VP+ phenotype. Although in most _Enterobacteriaceae_ a positive VP test is usually associated with a negative MR test (MR-/VP+, e.g., _Enterobacter aerogenes_), and vice versa (MR+/VP−, e.g., _E. coli_), there are also some species in this family (e.g., _Enterobacter intermedius_, _Klebsiella planticola_, or _Serratia liquifaciens_) reported to be positive for both tests (MR+/VP+; Holt et al., 1994).

Out of 6,048 mutants screened, 34 MR+/VP+ mutants were identified and their phenotype was confirmed after transferring the mutation to a native MG1655 strain by P1-transduction, followed by transformation of pTrc99A-P<sub>trc</sub>-budAB. Identification of the transposon insertion sites of these 34 mutants led to 16 different genes (Table 3). Interestingly, all genes were related to the metabolism of formate, one of the acids formed by mixed-acid fermentation. Formate is produced by the pyruvate formate lyase (PFL) enzyme, which catalyzes the CoA-dependent cleavage of pyruvate to formate and acetyl-CoA (Sawers and Böck, 1988). An overview of the fermentation routes present in _E. coli_ containing pTrc99A-P<sub>trc</sub>-budAB is shown in Figure 2. The formate that is produced and secreted can also be reimported in the cell through the FocA channel and become disproportionated to CO<sub>2</sub> and H<sub>2</sub> by the membrane-associated FHL complex (Sawers, 2005; Lü et al., 2012; Beyer et al., 2013). This complex consists of the formate dehydrogenase H (FDH-H), a selenoprotein carrying a molybdenum cofactor, and hydrogenase 3, a nickel-containing protein complex (Bagramyan and Trchounian, 2003). FDH-H catalyzes the oxidation of formate (HCOO<sup>−</sup>), generating CO<sub>2</sub> and H<sup>+</sup>. The electrons from this reaction are transferred via several subunits of the FHL complex to hydrogenase 3, where they combine with two cytoplasmic protons to form dihydrogen. This pathway is thus a net consumer of protons and is used by _E. coli_ to counteract acidification (Leonhartsberger et al., 2002). All gene products found in our screening could be linked to this particular pathway: FdhF (FDH-H), HycB, HycD, and HycE are part of the FHL complex (Bagramyan and Trchounian, 2003); HycI and HycE are both involved in maturation of the large subunit of hydrogenase 3 (Forzi and Sawers, 2003); ModC is the ATP binding subunit of the molybdate ABC transporter (Leonhartsberger et al., 1988; Driscoll and Copeland, 2003); MoeA and MoeB are involved in the biosynthesis of selenocysteine, and mutants lacking these gene products fail to synthetize FDH-H (Leinfelder et al., 2001; Nichols and Rajagopalan, 2002); FdhD is an accessory protein functioning as a sulfurtransferase between IscS and FdhF and is required for FDH activity (Thomé et al., 2012); FocA and PflB are coexpressed from a single operon and form a bidirectional formate channel and the PFL enzyme, respectively.

**Table 3** List of genes knocked out in transposon insertion mutants of _E. coli_ MG1655 containing pTrc99A-P<sub>trc</sub>-budAB that had lost the stationary-phase deacidification capacity but still produced acetoin (MR+/VP+).

| Gene      | Description                                                                 |
|-----------|-----------------------------------------------------------------------------|
| fdhD      | Redox enzyme maturation protein (REMP) for FdnG/FdoG; required as a sulfurtransferase for FDH activity |
| fdhF      | FDH-H                                                                       |
| hflA      | FHL system activator                                                        |
| focA      | Formate channel                                                             |
| hycB      | FHL complex iron−sulfur protein                                             |
| hycD      | FHL complex inner membrane protein                                          |
| hycE      | Hydrogenase 3 large subunit                                                 |
| hycI      | Maturation endoprotease for hydrogenase 3 large subunit HycE                |
| hypE      | Maturation protein required for the assembly of the CN ligand of the NiFe metal center of hydrogenase 1, 2, and 3. |
| modC      | ATP binding subunit of the molybdate ABC transporter                        |
| moeA      | Molybdopterin molybdenumtransferase                                          |
| moeB      | Molybdopterin-synthase adenylytransferase                                   |
| mog       | Molybdochelatase incorporating molybdenum into molybdopterin               |
| pfkB      | Pyruvate formate lyase                                                       |
| selA      | Selenocysteine synthase                                                      |
| selD      | Selenophosphate synthase                                                     |
Vivijs et al. Deacidification by acetoin-producing enterobacteria

hydrogenases catalyzing the reversible reaction $2H_2 \leftrightarrow H_2O$ when compared to the wild-type strain (Noguchi et al., 2010). showed a 20-fold loss in survival of an extreme acid stress (2 h at pH 6.0) when compared to the wild-type strain. This finding suggested that the FHL complex supports survival of extreme acid challenge by counteracting intracellular acidification. Our results now show that the complex can also assist in the reaction carried out by the FHL complex since deletion of hycE resulted in loss of acidification. This explanation is also supported by the observed gas production. Since CO2 is very soluble in water, gas accumulation in a Durham tube can be mainly ascribed to H2 production, and is thus indicative of the action of the FHL complex (White, 2000). Both strains with an intact FHL complex produced more or less the same amount of gas at 12 h, filling approximately half of the Durham tube with gas (Figure 3). However, no additional gas production was seen in case of wild-type *E. coli* after 12 h, while the Durham
FIGURE 3 | Cell numbers (squares) and medium pH (triangles) during fermentative growth of *E. coli* MG1655 wild-type (solid lines) or ΔhycE (dashed lines) containing pTrc99A (gray) or pTrc99A-P_trc-budAB (black) in LB medium with 5 g/l glucose, 1 mM IPTG and 100 μg/ml Ap at 37°C for 48 h. Error bars represent SD. Gas production (expressed as % of the volume of the Durham tube) is shown at the bottom.

Table 4 | Acetoin production (in mM) by *E. coli* MG1655 containing pTrc99A-P_trc-budAB or *E. coli* MG1655 ΔhycE containing pTrc99A-P_trc-budAB during fermentative growth in LB with 5 g/l glucose, 1 mM IPTG, and 100 μg/ml ampicillin (Ap) for 48 h.

| Time (h) | *E. coli* MG1655 | *E. coli* MG1655 ΔhycE |
|---------|-----------------|----------------------|
|         | pTrc99A-P_trc-budAB | pTrc99A-P_trc-budAB |
| 4       | 0.7 ± 0.2        | 0.5 ± 0.2            |
| 6       | 5.3 ± 0.7        | 5.0 ± 0.4            |
| 8       | 9.8 ± 0.9        | 7.1 ± 1.3            |
| 10      | 21.8 ± 0.8       | 18.4 ± 0.5           |
| 12      | 19.4 ± 1.2       | 12.5 ± 1.1           |
| 24      | 18.7 ± 0.7       | 12.3 ± 0.5           |
| 48      | 20.5 ± 2.3       | 11.1 ± 2.3           |

The evolution of plate counts during stationary phase in this experiment was generally in line with the observed pH changes, with cell death taking place in the strongly acidified cultures. In particular, lethal acidification could not be prevented by acetoin fermentation in a *budAB*-containing ΔhycE mutant since plate counts of this strain significantly decreased after the stationary phase, as was also the case for the two *budAB*-less strains performing a mixed-acid fermentation. Cell death can be explained by the combination of the low pH environment and the toxic accumulation of organic acids.

**ANALYSIS OF METABOLITES PRODUCED DURING FERMENTATIVE GROWTH OF *E. coli***

To provide more direct evidence for the involvement of formate disproportionation in the deacidification capacity of *budAB*-containing *E. coli*, glucose consumption and the production of metabolites were determined by HPLC during fermentative growth in LB with glucose (*Figure 4*). Succinate concentrations (*Figure 4E*) remained low for all strains during the course of the experiment. On the other hand, the *budAB* genes caused a marked shift in the production of two of the major acids of the mixed-acid fermentation pathway, especially in the late exponential and stationary growth phase, with no more acetate and much less lactate being produced (*Figures 4C,D*, respectively). With
FIGURE 4 | Time profiles of glucose consumption (A) and production of the metabolites ethanol (B), acetate (C), lactate (D), succinate (E), and formate (F) during fermentative growth of E. coli MG1655 wild-type (solid lines) or ΔhycE (dashed lines) containing pTrc99A (gray) or pTrc99A-Ptrc-budAB (black) in LB medium with 5 g/l glucose, 1 mM IPTG and 100 μg/ml Ap at 37°C for 48 h. Error bars represent SD.

regard to formate (Figure 4F), the highest formate accumulation was seen in the ΔhycE mutants, probably because these have lost their major route to convert formate to H₂ and CO₂. During the stationary growth phase (up to 48 h), the formate concentrations remained almost constant in the hyc⁻ strains, but strongly decreased in the hyc⁺ strains, indicating the reuptake and conversion of formate to CO₂ and H₂. Interestingly, a close look at the formate accumulation curves of the ΔhycE mutants reveals a transient decline in the late exponential growth phase (onset at 4 h of growth). Also in the hyc⁺ background a decline (budAB-less strain) or a diminished accumulation (budABContaining strain) of formate was observed in this phase. A possible explanation for this is the activity of the FDH-N, which also catalyzes the oxidation of formate to CO₂ (Sawers, 1994). However, FDH-N transfers the electrons to nitrate (via a nitrate reductase) instead of protons and has a much higher affinity for formate than the FDH-H (Leonhartsberger et al., 2002), which could explain why it is active in an earlier growth stage. The activity of FDH-N is limited, however, because LB medium contains only a small amount of nitrate. The disproportionation of formate (Figure 4F) by the hyc⁺ strains lasted longer when the budAB genes were present (48 h) than when they were absent (24 h), probably because a higher amount of formate was produced. This was also reflected by an increased gas production in the presence of the budAB genes during this phase, as reported above (Figure 3). Finally, ethanol was produced in higher quantities by the budAB-containing strains (Figure 4B).

As a final experiment to demonstrate that formate conversion causes medium deacidification during stationary phase, 5 or 10 mM formate from a 1 M solution (pH 5.50) was added to the
medium after 10 h of fermentative growth of budAB-containing E. coli MG1655 and the pH was subsequently measured after 10, 24, and 48 h. As expected, the addition of formate in the medium resulted in a significantly stronger pH increase during stationary phase (Table 5).

| Concentration of formate added at 10 h | 10 h | 24 h | 48 h |
|---------------------------------------|------|------|------|
| 0 mM                                  | 5.64 ± 0.03a | 5.81 ± 0.04a | 6.14 ± 0.02a |
| 5 mM                                  | 5.66 ± 0.05a | 5.96 ± 0.04b | 6.35 ± 0.04bc |
| 10 mM                                 | 5.63 ± 0.08a | 6.05 ± 0.04c | 6.48 ± 0.05c |

E. coli MG1655 harboring pTrc99A-Ptrc-budAB was grown in LB medium with 5 g/l glucose, 1 mM IPTG, and 100 μg/ml Ap. Formate was added at 10 h and pH was measured at 10, 24, and 48 h. a-c pH values with different superscripts in the same column are significantly different (p < 0.05).

**FIGURE 5** | Cell numbers (squares) and medium pH (triangles) during fermentative growth of S. plymuthica RVH1 wild-type (black solid lines), budAB::cat (gray solid lines), budAB::cat containing pTrc99A-Ptrc-budAB (dashed line), or ΔhycE (dotted line) in LB medium with 5 g/l glucose at 30°C for 48 h. For the strain containing pTrc99A-Ptrc-budAB, 1 mM IPTG and 200 μg/ml carbenicillin (Cb) were added to the medium. Error bars represent SD.

**ROLE OF HYDROGENASE 3 IN FERMENTATIVE GROWTH OF S. plymuthica RVH1**

Finally, we investigated whether the FHL complex also attenuates acid formation and drives deacidification during fermentative growth of a natural 2,3-butanediol fermenter, using S. plymuthica RVH1 as a model. To this end, we constructed a ΔhycE mutant in this strain. The evolution of medium pH for S. plymuthica RVH1 wild-type shows three phases (Figure 5). There was a decrease during the first 8 h, followed by a rapid increase between 8 and 10 h, and then a slower increase until 48 h. The initial pH increase is probably due to the switch to 2,3-butanediol production in the late exponential phase since it was lost upon knockout of the 2,3-butanediol pathway (budAB::cat) but not by knockout of hydrogenase 3 (ΔhycE). In contrast, the deacidification during stationary phase required both an active 2,3-butanediol pathway and an active hydrogenase 3. Genetic complementation of the budAB mutant restored its pH profile to that of the wild-type strain. However, since this complemented strain produces acetoin under the control of the plasmid Pinc promoter right from the start of the experiment, its deacidification is more attenuated than in the wild-type strain. Cell numbers declined after 48 h in the mutant strains that had lost or reduced deacidification capacity and, as a result, there was a clear correlation between cell numbers and medium pH at the end of the experiment.

Previously, the pH profile during glucose fermentation in the 2,3-butanediol fermenter Enterobacter aerogenes was divided into three phases (Johansen et al., 1975). The first phase was characterized by a rapid drop to about pH 5.8, in the second phase the pH remained almost constant at pH 5.6 and in the third phase the pH increased again to about 6.5. However, during the last phase, the total amount of acetoin and 2,3-butanediol remained constant and 2,3-butanediol was reoxidized to acetoin, indicating that the 2,3-butanediol pathway is not involved in this deacidification (Johansen et al., 1975). Our results demonstrate that—at least in S. plymuthica—the FHL complex is responsible for stationary-phase deacidification since the final pH was about 1.3 pH units lower in a S. plymuthica RVH1 ΔhycE mutant compared to the wild-type.

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