Underground isoleucine biosynthesis pathways in *E. coli*

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Abstract

The promiscuous activities of enzymes provide fertile ground for the evolution of new metabolic pathways. Here, we systematically explore the ability of *E. coli* to harness underground metabolism to compensate for the deletion of an essential biosynthetic pathway. By deleting all threonine deaminases, we generated a strain in which isoleucine biosynthesis was interrupted at the level of 2-ketobutyrate. Incubation of this strain under aerobic conditions resulted in the emergence of a novel 2-ketobutyrate biosynthesis pathway based upon the promiscuous cleavage of *O*-succinyl-*L*-homoserine by cystathionine γ-synthase (MetB). Under anaerobic conditions, pyruvate formate-lyase enabled 2-ketobutyrate biosynthesis from propionyl-CoA and formate. Surprisingly, we found this anaerobic route to provide a substantial fraction of isoleucine in a wild-type strain when propionate is available in the medium. This study demonstrates the selective advantage underground metabolism offers, providing metabolic redundancy and flexibility which allow for the best use of environmental carbon sources.

Introduction

The patchwork model suggests that novel metabolic pathways emerge from the promiscuous activities of enzymes participating in diverse metabolic processes (Lazcano and Miller, 1999; Jensen, 1976; Khersonsky and Tawfik, 2010; Noda-Garcia et al., 2018). According to this, underground metabolism – the network of metabolic conversions which are catalyzed as side reactions of enzymes that have evolved to support other activities (D’Ari and Casadesús, 1998) – provides fertile ground for the evolution of new pathways. A computational analysis suggested that about half of all underground reactions generate metabolites that already exist in the endogenous metabolic network, thus enabling the emergence of metabolic bypasses for the production of key cellular building blocks (Notebaart et al., 2014). For example, promiscuous activities of the arginine biosynthesis enzymes enabled proline production in an *E. coli* strain deleted in the canonical proline biosynthesis route (Itikawa et al., 1968); several pathways, each based on the promiscuous activities of different enzymes, have been shown to relieve pyridoxal phosphate auxotrophy in *E. coli* (Kim et al., 2010; Oberhardt et al., 2016); and adaptive evolution of *E. coli* harnessed enzyme promiscuity to enable growth on the non-natural feedstock 1,2-propanediol (Lee and Palsson, 2010).

The biosynthesis of isoleucine provides multiple examples of enzyme promiscuity and structural similarity to other pathways, suggesting underground metabolism as a likely origin (Jensen, 1976). Cellular production of isoleucine and valine are catalyzed by the same enzymes; in valine biosynthesis, pyruvate is self-condensed whereas in isoleucine biosynthesis, pyruvate is condensed with 2-ketobutyrate (2 KB). In most organisms, 2 KB is produced from threonine cleavage. Despite this, different biosynthetic routes, which mirror other metabolic pathways, are known to support the
production of 2 KB in specific lineages (Figure 1). For example, the citramalate route mirrors the TCA cycle, where pyruvate, rather than oxaloacetate, is condensed with acetyl-CoA, and 2 KB, rather than 2-ketoglutarate, is the pathway product (Charon et al., 1974). Similarly, in several anaerobic microorganisms, propionate is converted to 2 KB in a reductive pathway that mirrors acetate conversion to pyruvate via ligation with CoA and ferredoxin-dependent carboxylation (Buchanan, 1969; Monticello et al., 1984; Eikmanns et al., 1983). In plants, methionine γ-lyase, catalyzing a reaction similar to that of cystathionine γ-lyase, is a complementary source of cellular 2 KB (Joshi and Jander, 2009; Joshi et al., 2010).

Here, we tested whether the underground metabolism of E. coli could compensate for the deletion of the canonical 2 KB production route in isoleucine biosynthesis. Under aerobic conditions, we show that a mutation in a cysteine biosynthesis enzyme decreased its activity and lowered the steady-state concentration of this amino acid; this freed an intermediate of methionine biosynthesis, which usually reacts with cysteine, to be converted to 2 KB. Under anaerobic conditions, E. coli was able to use the enzyme pyruvate formate-lyase (PFL) to generate 2 KB from propionyl-CoA and formate. Surprisingly, we found this pathway to provide a substantial fraction of the cellular 2 KB also in a wild-type (WT) strain. This study thus demonstrates the inherent ability of microorganisms to effectively exploit underground metabolism to generate patchwork pathways.

Results

A latent aerobic isoleucine biosynthesis pathway

Biosynthesis of isoleucine in E. coli starts with the deamination of threonine to give 2 KB, which is then condensed with pyruvate to produce 2-aceto-2-hydroxybutanoate. This intermediate is subsequently isomerized, reduced, dehydrated, and aminated to generate isoleucine. To generate an isoleucine auxotrophy, we constructed a strain in which the two genes encoding for threonine deaminase were deleted (ilvA and tdcB). When provided with isoleucine, this strain grew identically to the WT strain (dark red line in Figure 2). Without the addition of isoleucine, we observed no growth in the first 70 hr, but within 70–120 hr, different replicates started growing with a growth rate somewhat lower than that of the WT strain (purple lines in Figure 2).

One explanation of the observed growth without isoleucine is an increased activity of serine deaminases, which are known to accept threonine as a substrate (Cicchillo et al., 2004). We, therefore, deleted all the genes encoding for serine deaminase (sdaA, sdaB, and tdcG). We termed the
resulting strain Δ5 (ΔilvA ΔtdcB ΔsdaA ΔsdaB ΔtdcG). However, strain Δ5 displayed similar behavior to the ΔilvA ΔtdcB strain, that is, while growth without isoleucine was not observed in the first 70 hr, after 70 hr, the replicates started growing (green lines in Figure 2). This strongly suggests the emergence of a latent threonine-independent isoleucine biosynthesis pathway.

To check whether this route still depends on the generation of 2 KB or rather bypasses this metabolic intermediate altogether, we constructed a strain deleted in the gene ilvC. This gene encodes a ketol-acid reductoisomerase that operates downstream of 2 KB in the isoleucine biosynthesis pathway. While cultivation of the ΔilvC strain with valine and isoleucine enabled growth similar to that of the WT strain (pink line in Figure 2; valine is required as IlvC participates also in valine biosynthesis), growth without isoleucine was not observed even after 150 hr (orange line in Figure 2). As a further confirmation, we observed that addition of 2 KB to the cultivation medium rescued the growth of the ΔilvA ΔtdcB strain and the Δ5 strain but not the ΔilvC strain (Figure 2). This confirms that 2 KB still serves as a metabolic intermediate in the underground isoleucine biosynthesis pathway.

In some microorganisms, 2 KB is produced via the citramalate pathway (Xu et al., 2004; Drevland et al., 2007; Hochuli et al., 1999; Risso et al., 2008). This route is a catalytic parallel to the first half of the TCA cycle – from citrate synthase to isocitrate dehydrogenase – where pyruvate replaces oxaloacetate, reacting with acetyl-CoA to give citramalate, which is subsequently metabolized to 2 KB. We wondered whether the enzymes of the TCA cycle, or their isozymes, catalyze the reactions of the citramalate pathway in E. coli. To check if this were the case, we performed a carbon labeling assay, feeding the WT strain, the ΔilvA ΔtdcB strain, and the Δ5 strain with either glucose-1-13C or glucose-3-13C. The expected labeling pattern of isoleucine should completely change depending on the biosynthesis route of 2 KB (Figure 3A). In the WT strain, isoleucine is expected to be roughly half labeled upon feeding with glucose-3-13C and follow a 1:2:1 pattern (unlabeled: once labeled: double labeled) with glucose-1-13C. By contrast, if 2 KB is produced via the citramalate pathway, then isoleucine would be completely unlabeled when feeding with glucose-3-13C and
would roughly follow a 1:2:2:1 pattern (unlabeled: once labeled: double labeled: triple labeled) with glucose-1-\(^{13}\)C. The labeling pattern of isoleucine in the \(\Delta\)ilvA\(\Delta\)tdcB and the \(\Delta\)5 strains is almost identical to that of the WT strain (Figure 3B, the small deviations from the expected labeling can be attributed to the ambient abundance of \(^{13}\)C and the shuffling of labeled carbon by the activity of the pentose phosphate pathway and the TCA cycle). This effectively rules out the activity of the citramalate pathway and indicates that the biosynthesis of 2 KB in both gene deletion strains is closely related to the natural production pathway.

**2-Ketobutyrate biosynthesis from O-succinyl-L-homoserine**

In several organisms, the biosynthesis of isoleucine is at least partially tied to that of the sulfur-containing amino acids cysteine and methionine. For example, in organisms in which cysteine biosynthesis depends on \(\gamma\)-elimination of cystathionine, 2 KB serves as a byproduct (Steegborn et al., 1999). Also, in plants, 2 KB can be produced from \(\gamma\)-elimination of methionine (Joshi and Jander, 2009; Joshi et al., 2010). To test whether the production of 2 KB in the \(\Delta\)5 strain is related to methionine...
biosynthesis, we deleted metA, encoding for homoserine O-succinyltransferase, which catalyzes the first committed step of the methionine biosynthesis pathway. The ΔS ΔmetA strain could grow with the addition of methionine and isoleucine (Figure 4, pink line), while no growth was observed without isoleucine (Figure 4, orange line). This indicates that the production of 2 KB in the ΔS strain is indeed dependent on the methionine biosynthesis route.

At first glance, the biosynthesis of sulfur-containing amino acids in E. coli would not be expected to give rise to 2 KB. Specifically, in this bacterium, cystathionine does not undergo γ-elimination but rather β-elimination which generates homocysteine and pyruvate (Figge, 2006). Also, E. coli lacks a characterized annotated methionine γ-lyase. While an E. coli gene was previously suggested to encode for such an enzyme (Manukhov et al., 2005), the inability of the ΔS ΔmetA strain to grow when supplemented with methionine but not isoleucine confirms that E. coli lacks methionine γ-lyase activity.

We, therefore, searched the literature for reactions that could convert intermediates of the methionine biosynthesis pathway to 2 KB. We found multiple such candidate reactions from a range of organisms (Figure 5). First, rather than catalyzing the condensation of cysteine with O-succinyl-L-homoserine, cystathionine γ-synthase (MetB) can cleave the latter intermediate to succinate and 2 KB (reaction one in Figure 5; \( k_{\text{cat}} = 7.7 \text{ s}^{-1} \) and \( K_M(\text{O-succinyl-L-homoserine})=0.33 \text{ mM} \) for E. coli enzyme [Figge, 2006; Holbrook et al., 1990]). MetB can also condense O-succinyl-L-homoserine and homocysteine to give homolanthionine (reaction two in Figure 5; \( K_M(\text{homocysteine})=0.54 \text{ mM} \) for C. glutamicum enzyme [Krömer et al., 2006]), which can be subsequently cleaved by cystathionine-β-lyase (MetC) to regenerate homocysteine and produce 2 KB (reaction three in Figure 5; e.g. \( k_{\text{cat}} \approx 180 \text{ s}^{-1} \) and \( K_M(\text{homolanthionine})=4.5 \text{ mM} \) for E. coli enzyme [Figge, 2006; Krömer et al., 2006; Alting et al., 1995; Dwivedi et al., 1982]). In addition, MetC from various organisms can catalyze both the β-elimination and the γ-elimination of cystathionine, giving rise to both pyruvate and 2 KB (reaction four in Figure 5; e.g. \( k_{\text{cat}}(\gamma\text{-elimination}) \approx 0.9 \text{ s}^{-1} \) and \( K_M(\text{cystathionine})=0.20 \text{ mM} \) for Streptomyces phaeochromogenes enzyme where β-elimination proceeds at \( \approx 1/7 \) rate of γ-elimination [Alting et al., 1995; Nagasawa et al., 1984]). As MetC can also cleave cysteine to pyruvate...

Figure 4. Methionine biosynthesis is implicated in promiscuous 2-ketobutyrate production. The ΔS strain further deleted in homoserine O-succinyltransferase (ΔS ΔmetA) did not grow when supplemented with 2 mM methionine (orange line) but mutants of the cystathionine β-lyase (ΔS ΔmetC) grew rapidly and consistently on 10 mM glucose alone. A mutation in serine acetyltransferase (ΔS cysE*) resulted in near WT growth on glucose (purple line). Both isoleucine (Ile) and methionine (Met) were added at a concentration of 2 mM where indicated. All experiments were performed in replicates in a 96-well plate reader. Replicates showed an identical growth profile (±5%) and hence are represented by a single curve. Experiments were repeated at least three times for all growth experiments shown.
It is possible that it could also cleave homocysteine to produce 2 KB (reaction five in Figure 5). Finally, some MetC variants can act as methionine γ-lyase, releasing 2 KB by directly cleaving methionine (reaction six in Figure 5; e.g. $k_{cat} \approx 0.01 \text{ s}^{-1}$ for Lactococcus lactis subsp. cremoris B78 enzyme [Alting et al., 1995; Amarita et al., 2004; Irmler et al., 2008]).

To identify which of the possible biosynthesis routes is responsible for the production of 2 KB in the $\Delta$5 strain, we performed several enzymatic experiments (Materials and methods and Figure 5—figure supplement 1). We found that MetC from E. coli did not catalyze the degradation of cystathionine, homocysteine, or methionine to 2 KB (reaction five in Figure 5). Finally, some MetC variants can act as methionine γ-lyase, releasing 2 KB by directly cleaving methionine (reaction six in Figure 5; e.g. $k_{cat} \approx 0.01 \text{ s}^{-1}$ for Lactococcus lactis subsp. cremoris B78 enzyme [Alting et al., 1995; Amarita et al., 2004; Irmler et al., 2008]).

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More importantly, this strain could directly grow without isoleucine and without the need for adaptation (green line in Figure 4).

Supporting previous studies (Figge, 2006; Holbrook et al., 1990), we found MetB to catalyze the cleavage of O-succinyl-L-homoserine to 2 KB (reaction one in Figure 5) with $k_{cat} = 9.3 \pm 0.4 \text{ s}^{-1}$, $K_M$(O-succinyl-L-homoserine)$ = 0.60 \pm 0.08 \text{ mM}$, and thus $k_{cat}/K_M$(O-succinyl-L-homoserine)$ = 16 \pm 2 \text{ mM}^{-1} \text{s}^{-1}$ (Figure 5—figure supplement 2A). Yet, it was previously reported that the formation of 2 KB is suppressed in the presence of cysteine (Holbrook et al., 1990). We, therefore, characterized the cleavage reaction in the presence of either cysteine or homocysteine (as an alternative substrate), at physiological concentrations of $\approx 0.3 \text{ mM}$ (Wheldrake, 1967; Bennett et al., 2009; Guo et al., 2013) and at artificially high concentrations of 3–6 mM. We found that formation of 2 KB was indeed suppressed at physiological concentrations of cysteine, while homocysteine suppressed 2 KB formation only at high concentrations (Figure 5—figure supplement 2B–E). Nonetheless, as the deletion of metB in the $\Delta S$ strain completely abolishes growth without isoleucine (brown line in Figure 4), it seems very likely that MetB is involved in the production of 2 KB.

**Disruption of MetC or a mutation in serine acetyltransferase enable steady 2-ketobutyrate production from O-succinyl-L-homoserine**

Next, we aimed to understand the genetic basis underlying 2 KB biosynthesis in the $\Delta S$ strains. We sequenced the genomes of several strains isolated from the cultures growing without isoleucine. We found that in most of the sequenced strains (11 strains out of the 16 sequenced) metC was either deleted or mutated (Supplementary file 1). This is in line with the findings reported above, that is, $\Delta S$ $\Delta$metC strain is able to directly grow without isoleucine and without lag time (green line in Figure 4). It therefore seems that a lower metabolic flux toward methionine biosynthesis, as expected by the deletion of metC, enhances the side reactivity of the pathway enzymes and results in a higher conversion rate of O-succinyl-L-homoserine to 2 KB. Indeed, we found that, as compared to the WT strain and the $\Delta S$ strain, in the $\Delta S$ $\Delta$metC strain the concentration of methionine was $\approx 3$-fold lower (Figure 6A), while the concentration of O-succinyl-L-homoserine was $\approx 3$-fold higher (Figure 6A). The side reactivity of MetB toward O-succinyl-L-homoserine cleavage therefore seems to be enhanced by the high concentration of this metabolite in the $\Delta S$ $\Delta$metC strain.

Another mutated strain harbored a single mutation in the gene coding for serine acetyltransferase (CysE): Ala33Thr (Supplementary file 1). We used Multiplex Automated Genomic Engineering (MAGE [Wang et al., 2009]) to introduce this mutation into the $\Delta S$ strain. The resulting strain ($\Delta S$ cysE*) grew immediately (with no lag time) on a medium without isoleucine and with a growth rate identical to the WT strain (purple lines in Figure 4; note that the growth of the $\Delta S$ cysE* strain was considerably faster than that of the $\Delta S$ $\Delta$metC strain). Therefore, it seems that the mutation in cysE enabled a steady, efficient production of 2 KB.

CysE catalyzes the first committed step in the biosynthesis of cysteine (Denk and Bock, 1987). To understand the effect of the Ala33Thr mutation on enzyme activity we measured the kinetics of the purified enzyme (Materials and methods and Figure 5—figure supplements 1 and 3). Despite the mutation occurring far from the active site, we found that the apparent $k_{cat}$ of the enzyme decreased 2-fold, from $350 \pm 30 \text{ s}^{-1}$ to $170 \pm 30 \text{ s}^{-1}$, while the $K_M$ for acetyl-CoA increased by more than 8-fold, from $0.6 \pm 0.2 \text{ mM}$ to $5.0 \pm 2.0 \text{ mM}$ (interestingly, $K_M$ for serine changed only slightly from $0.8 \pm 0.3 \text{ mM}$ in the WT to $0.5 \pm 0.2 \text{ mM}$ in the mutant). As the concentration of acetyl-CoA in *E. coli* lies in the range of $0.6$–$0.75 \text{ mM}$ (Bennett et al., 2009), the increase in $K_M$ for acetyl-CoA directly affects the reaction rate. Overall, the Ala33Thr mutation is expected to decrease the rate of the CysE reaction by more than 17-fold under physiological conditions.

The decreased activity of CysE halved the intracellular cysteine concentration relative to that of the WT strain and the $\Delta S$ strain (Figure 7). This deprived MetB of its co-substrate, thus lowering the methionine biosynthesis flux and resulting in more than a 2-fold reduction in methionine concentration (Figure 6A). The lower availability of methionine led to overexpression of genes participating in its biosynthesis, increasing the transcript level of metA and metB more than 3-fold relative to the WT strain and the $\Delta S$ strain (Figure 6B; interestingly, such increase in transcript level was not observed in the $\Delta S$ $\Delta$metC strain). This higher expression, as well as the partial lifting of MetA inhibition by methionine (Born and Blanchard, 1999), led to a dramatic 30- to 60-fold increase in the concentration of O-succinyl-L-homoserine (Figure 6A). Overall, the higher concentrations of both MetB and O-succinyl-L-homoserine, coupled with the lower concentration of cysteine, resulted in the
diversion of more O-succinyl-L-homoserine toward cleavage and 2 KB production. Interpreted so, the mutation in CysE – an enzyme that is not directly involved in 2 KB biosynthesis – enhances a previously negligible underground reaction for 2 KB production, thus awakening a latent isoleucine biosynthesis route (Figure 7).

Anaerobic 2-ketobutyrate biosynthesis from a reversible 2-ketobutyrate formate-lyase activity

Next, we aimed to explore underground isoleucine biosynthesis routes under anaerobic conditions (Figure 8). The evolved ΔilvA ΔdcdB cysE* strain and the constructed Δ5 cysE* strain grew anaerobically without isoleucine (Figure 8—figure supplement 1). However, we did not observe the reemergence of the metC deletion, the cysE* mutation, or any other mutation: the Δ5 strain failed to grow without the addition of isoleucine, even after 120 hr (black lines in Figure 8A). We wondered whether we could enable 2 KB production by the addition of small metabolites that E. coli might encounter in its natural habitat. We focused on propionate, a short-chain fatty acid abundant in the mammalian intestine (McNeil et al., 1978), which is known to be naturally activated to propionyl-CoA (Hesslinger et al., 1998; Liu et al., 2014). As we had previously found that PFL (encoded by pflB) supported the in vivo condensation of acetyl-CoA and formate to produce pyruvate (Zelcbuch et al., 2016), we hypothesized that this enzyme could also catalyze a 2 KB formate-lyase (KBFL) reaction, that is, accepting propionyl-CoA to produce 2 KB. We indeed found that a purified PFL catalyzes the condensation of propionyl-CoA and formate to give 2 KB with a specific activity of 0.6–1.1 μmol/min/mg, $k_{\text{cat}}$ of 0.9–1.7 s$^{-1}$, $K_{\text{M}}$(propionyl-CoA)=0.83 ± 0.6 mM and $K_{\text{M}}$(formate) =69 ± 11 mM (Table 1, Materials and methods, Figure 5—figure supplement 1, and Figure 8—figure supplements 2 and 3).
We found that the Δ5 strain could grow anaerobically when propionate and formate were added to the medium instead of isoleucine (red line in Figure 8A). As formate is naturally produced by the PFL-dependent cleavage of pyruvate, we reasoned that the addition of this compound to the medium might be redundant. Indeed, we found that propionate alone, when added to the medium, supported the growth of the Δ5 strain, albeit with longer lag time and lower growth rate than with the addition of formate (purple line in Figure 8A, all replicates displayed an identical growth phenotype, thus ruling out genetic mutations). To confirm that propionate is assimilated via the activity of PFL, rather than via a reductive carboxylation route (Buchanan, 1969; Monticello et al., 1984; Eikmanns et al., 1983), we deleted pflB. When provided with isoleucine and acetate, the Δ5ΔpflB strain was able to grow (green line in Figure 8B, acetate was added as the deletion of pflB disrupted the endogenous acetyl-CoA biosynthesis route under anaerobic conditions [Hasona et al., 2004]). However, replacement of isoleucine with propionate, with or without further addition of formate, resulted in no growth (red and purple lines in Figure 8B). This confirms that PFL is indeed responsible for 2 KB biosynthesis.

To provide a further validation that 2 KB is produced from propionate, we followed the 13C labeling of isoleucine upon addition of propionate-1-13C to the growth medium under anaerobic conditions. Surprisingly, we found that even within a WT strain, the addition of labeled propionate resulted in 30% labeled isoleucine (Figure 9), suggesting that the KBFL route operates naturally if propionate is present in the medium. When formate was also added to the medium, the fraction of labeled isoleucine increased to almost 60% (Figure 9), making the KBFL-dependent pathway the

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**Figure 7.** A mutation in serine acetyltransferase (cysE*) enables 2-ketobutyrate production by MetB. (A) A schematic representation of endogenous cysteine, methionine, and isoleucine biosynthesis in E. coli. Thickness of arrows indicates flux intensity. (B) Schematic representation of cysteine, methionine, and isoleucine biosynthesis when threonine deaminases are deleted and with the cysE* mutation. Thickness of the arrows indicates higher flux via O-succinyl-L-homoserine cleavage to 2-ketobutyrate. As indicated by the red arrows, the transcript levels of metA and metB as well the concentration of O-succinyl-L-homoserine increase, while the concentrations of cysteine and methionine decrease. Numbers next to the arrows correspond to the factor by which the metabolite concentration or transcript level increased or decreased (as derived from Figure 6). Cysteine concentrations in the Δ5ΔcysE* strain were ≈50% of that in WT and Δ5 strains.

The online version of this article includes the following source data for figure 7:

**Source data 1.** Relative concentrations of cysteine.

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dominant isoleucine biosynthesis route within a WT strain. In the Δ5 strain, in which the canonical isoleucine biosynthesis pathways are deleted, isoleucine was almost 80% labeled upon addition of formate and labeled propionate (Figure 9). Deletion of the metA increased this labeling to 90% (Figure 9), indicating that the MetB-dependent route operates anaerobically although with a low flux. As 10% of the isoleucine was not labeled even in the Δ5ΔmetA strain, it seems that there might be another undiscovered yet marginal route for isoleucine biosynthesis.

We wondered if the KBFL reaction is supported solely by PFL or whether TdcE, which was previously shown to catalyze the 2 KB formate-lyase reaction (Sawers et al., 1998), also contributed to the production of 2 KB. We found that purified TdcE is catalytically superior to PFL, generating 2 KB.

Table 1. Kinetic parameters of PFL and TdcE for catalyzing the condensation of propionyl-CoA with formate to generate 2-ketobutyrate.

| Enzyme     | Propionyl-CoA | Formate |
|------------|---------------|---------|
|            | $V_{\text{max}}$ (U/mg) | $k_{\text{cat}}$ (s$^{-1}$) | $K_{\text{M}}$ (mM) | $K_{i}$ (mM) | $V_{\text{max}}$ (U/mg) | $k_{\text{cat}}$ (s$^{-1}$) | $K_{\text{M}}$ (mM) |
| TdcE       | 2.1 ± 0.1     | 3.2 ± 0.2 | 520 ± 81 | -         | 1.8 ± 0.07 | 2.7 ± 0.1 | 20 ± 3   |
| PFL (PflB) | 1.1 ± 0.6     | 1.7 ± 0.9 | 830 ± 580| 690 ± 500 | 0.59 ± 0.03 | 0.9 ± 0.05 | 69 ± 11  |

See Materials and methods for assay conditions and Figure 8—figure supplements 2 and 3. Note that the $k_{\text{cat}}$ values were calculated assuming that the enzymes are fully activated.
with a specific activity of 1.8–2.1 μmol/min/mg, $k_{\text{cat}}$ of 2.7–3.2 s$^{-1}$, $K_M$(propionyl-CoA)=$0.52 \pm 0.08$ mM, and $K_M$(formate)=$20 \pm 3$ mM (Table 1, Materials and methods, Figure 5—figure supplement 1, and Figure 8—figure supplements 2 and 3). Yet, while the deletion of pflB (in which case acetate was further added to the medium) abolished isoleucine labeling, deletion of tdcE resulted in identical labeling to that of the WT strain (Figure 9). This indicates that PFL is solely responsible for the production of 2 KB from propionate and formate. Still, it could be that TdcE does not contribute to 2 KB biosynthesis only because it is not expressed under the tested conditions. Hence, we explored the consequences of overexpressing tdcE on a plasmid. We found that overexpression of tdcE in a Δ5 ΔpflB strain enabled fast growth with propionate (purple line in Figure 8C), indicating that TdcE can replace PFL both in catalyzing pyruvate cleavage to generate formate (as noted previously [Sawers et al., 1998]) and in catalyzing 2 KB production from propionyl-CoA and formate. Moreover, overexpression of tdcE in a WT strain fed with propionate-1,13C resulted in a high proportion of labeled isoleucine: more than 70% without externally added formate and ~90% with externally added formate, both are considerably higher than without tdcE overexpression (Figure 9). Overexpression of tdcE in the Δ5 ΔmetA strain resulted in 95% labeled isoleucine, the highest labeling we observed (Figure 9). Overall, these results confirm that TdcE can support in vivo 2 KB synthesis from propionyl-CoA in a comparable manner to that of PFL.

Discussion

This study serves to demonstrate the selective advantage underground metabolism may offer in nature, providing metabolic redundancy and flexibility which allows for the best use of
environmental carbon sources (Notebaart et al., 2014). We demonstrated two metabolic pathways for the biosynthesis of isoleucine which are based on the activity of promiscuous enzymes. The first pathway, uncovered under aerobic conditions, is based on rerouting the metabolism of O-succinyl-L-homoserine from methionine biosynthesis toward the production of 2 KB. This represents a rather classic case of underground metabolism: a promiscuous reaction, the activity of which normally has a negligible physiological contribution, is enhanced by the rewiring of cellular metabolism under conditions which couple it to growth. Interestingly, a change in the steady-state concentration of a seemingly unrelated metabolite – cysteine – greatly enhances this promiscuous activity and enables a new biosynthetic route.

Under anaerobic conditions, we uncovered a pathway that relies on the highly expressed PFL (Neidhardt et al., 1996) to condense propionyl-CoA and formate to 2 KB. This route, unlike the aerobic cleavage of O-succinyl-L-homoserine, was highly active also in a WT strain, providing a substantial fraction of cellular isoleucine when propionate was present in the medium. As short-chain fatty acids, such as propionate, are abundant in the mammalian intestine (McNeil et al., 1978), the native habitat of E. coli, the KBFL pathway is likely to play a key role in isoleucine biosynthesis under physiological conditions. Hence, this pathway does not actually represent underground metabolism, but rather should be regarded an auxiliary biosynthesis route based on promiscuous enzyme activities. The use of the KBFL pathway is favorable over the canonical threonine cleavage route as it better uses the carbon sources available in the environment. As PFL supports the KBFL reaction under physiological conditions, we suggest renaming it ‘pyruvate formate-lyase/2-ketobutyrate formate-lyase’, in a similar manner to TdcE, which was named as ‘2-ketobutyrate formate-lyase/pyruvate formate-lyase’.

Several previous studies have suggested that, in various microbial lineages, 2 KB is derived from propionate via the activity of 2 KB:ferredoxin oxidoreductase (Buchanan, 1969; Monticello et al., 1984; Eikmanns et al., 1983; Sauer et al., 1975). The findings presented here indicate that, in at least some of these microorganisms, propionate might be converted to 2 KB via the KBFL route, where formate is produced internally via pyruvate cleavage, or even from the reduction of CO₂ via formate dehydrogenase (Maia et al., 2017). Even the microorganism in which the reductive carboxylation of propionyl-CoA was first demonstrated, that is, Clostridium pasteurianum, harbors a genomic PFL (Kanehisa, 2000) and thus may also assimilate propionate into isoleucine via the KBFL pathway. Moreover, in several microorganisms, propionyl-CoA is produced from succinyl-CoA as part of anaerobic fermentation (Swick and Wood, 1960; Stams et al., 1984; Schink et al., 1987; Tholozan et al., 1994); these organisms might be able to use the KBFL route without the need for an external supply of propionate. The prevalence of both the genes and the substrates involved in the KBFL pathway indicates that this route may be widespread.

Many cellular building blocks have alternative biosynthesis pathways. Nevertheless, isoleucine seems unusually rich in the sheer number of alternative routes for its bioproduction. There appear to be at least nine routes for 2 KB production: threonine cleavage (canonical pathway), the citramalate pathway (Charon et al., 1974), the glutamate mutase pathway (Abramsky et al., 1962), cystathionine cleavage (Steegborn et al., 1999), methionine cleavage (Joshi and Jander, 2009; Joshi et al., 2010), homolanthionine cleavage (Krömer et al., 2006), O-succinyl-L-homoserine cleavage (this study), propionyl-CoA reductive carboxylation (Buchanan, 1969; Monticello et al., 1984; Eikmanns et al., 1983), and propionyl-CoA condensation with formate (this study). The large repertoire of alternative pathways makes isoleucine biosynthesis the epitome of metabolic flexibility.

**Materials and methods**

| Reagent type (species or resource) | Designation | Source or reference | Identifiers | Additional information |
|----------------------------------|-------------|---------------------|-------------|------------------------|
| Strain, strain background (Escherichia coli) | E. coli strains | This study | |

Continued on next page
Table 2. Strains used in this study.

| Strain name | Deletions/Genotype | Description | Source |
|-------------|--------------------|-------------|--------|
| BL21 (DE3) | *E. coli* thiA2 [lon] ompT galA DE3 [dcm] ΔhsdS λ DE3 = λ sβamHio ΔEcoRI-B int: lacI:PlacUV5::T7 gene1 i21 ΔninS | Strain for protein production | Lab collections |
| SU488 | *E. coli* K-12 MG1655Tn7:: pAra-exo-beta-gam; pRha-FLP; xylSpm-lsce | MG1655 derivative with genome integrated recombinase and flippase genes | 57 |
| ΔilvA ΔtdcB | SU488 ΔilvA ΔtdcB::kan | Threonine deaminase deletion strain | This study |
| ΔS | SU488 ΔilvA ΔtdcB ΔsdaA ΔsdaB ΔtdcG::kan | Threonine deaminase and serine deaminase deletion strain | This study |
| ΔS ΔmetA | SU488 ΔilvA ΔtdcB ΔsdaA ΔsdaB ΔtdcG ΔmetA::kan | Threonine deaminase, serine deaminase and homoserine O-succinyltransferase deletion strain | This study |
| ΔS ΔmetC | SU488 ΔilvA ΔtdcB ΔsdaA ΔsdaB ΔtdcG ΔmetC::kan | Threonine deaminase, serine deaminase and cystathionine β-lyase deletion strain | This study |
| ΔS cysE* | SU488 ΔilvA ΔtdcB ΔsdaA ΔsdaB ΔtdcG ΔmetC::kan cysE | Threonine deaminase, serine deaminase and homoserine O-succinyltransferase deletion strain with cysE point mutation G97A | This study |
| ΔilvC | SU488 ΔilvC::cap | Ketol-acid reductoisomerase deletion strain | This study |
| ΔapfB ΔtdcE | ΔilvA ΔtdcB ΔsdaA ΔsdaB ΔtdcG ΔapfB ΔtdcE::cap | Threonine deaminase, serine deaminase, pyruvate formate-lyase and 2-ketobutyrate formate-lyase deletion strain | This study |
| ΔapfB | SU488 ΔapfB::kan | Pyruvate formate-lyase deletion strain | This study |
| ΔtdcE | SU488 ΔtdcE::cap | 2-Ketobutyrate formate-lyase deletion strain | This study |

Continued

| Reagent type (species) or resource | Designation | Source or reference | Identifiers | Additional information |
|-----------------------------------|-------------|--------------------|-------------|-----------------------|
| Sequence-based reagent | pZASS-tdcE (plasmid) | This study | PCR primers | Overexpression vector of TdcE. |
| Recombinant DNA reagent | pCA24N-metB (plasmid) | This study | | Overexpression vector of MetB. |
| Recombinant DNA reagent | pCA24N-metC (plasmid) | This study | | Overexpression vector of MetC. |
| Recombinant DNA reagent | pCA24N-cysE (plasmid) | This study | | Overexpression vector of CysE. |
| Recombinant DNA reagent | pCA24N-cysE_A33T (plasmid) | This study | | Overexpression vector of CysE with a mutation A33T. |
| Recombinant DNA reagent | pORTMAGE | *Nyerges et al., 2016* | Addgene catalog no. 72680 |
| Chemical compound, drug | 13C-1-glucose | Sigma Aldrich |

Continued on next page
Strains and plasmids

All strains used in this study are listed in Table 2. The E. coli SIJ488 strain based upon K-12 MG1655 (Jensen et al., 2016) was used for the generation of deletion strains. SIJ488 is engineered to carry the gene deletion machinery in its genome (inducible recombinase and flippase). All gene deletions were carried out by successive rounds of λ-Red recombineering using kanamycin cassettes (FRT-PGK-gb2-neo-FRT [KAN], Gene Bridges, Heidelberg, Germany) or chloramphenicol cassettes (pKD3 [Datsenko and Wanner, 2000]) as described by Baba et al., 2006. Homologous extensions (50 bp) for the deletion cassettes were generated by PCR using oligonucleotides listed in Supplementary file 2.

For the overexpression of the 2-ketobutyrate lyase (tdcE), endogenous genes were amplified from E. coli genomic DNA using a two-step PCR (to remove cloning system relevant restriction sites [Wenk et al., 2018] – in this case a single site) using in the first round of PCR: primer pairs tdcE_-HIS.A + tdcE_B and tdcE_C + tdcE_CAP.D. The corresponding PCR products were combined in a second PCR using primer pairs tdcE_HIS.A and tdcE_CAP.D. The tdcE gene was subsequently cloned into cloning vector pNivC (Wenk et al., 2018) using restriction enzymes Mph1103I and XhoI, generating pNivC-tdcE. The tdcE gene was subsequently cloned from pNiv-tdcE into expression vector pZ-ASS using enzymes EcoRI and PstI, resulting in pZASS-tdcE (Wenk et al., 2018).

Plasmids for the overexpression of MetB, MetC, and CysE (pCA24N-metB, pCA24N-metC, and pCA24N-cysE) were isolated from the ASKA collection (Kitagawa et al., 2005). The CysE Ala33Thr point mutation was generated by linear amplification of the pCA24N-cysE plasmid by PCR using following mismatch primers: CysE_A33T_fw and CysE_A33T_rv (Supplementary file 2). The template plasmid was digested by DpnI (Thermo Fisher Scientific, Darmstadt, Germany) at 37°C and E. coli DH5α was transformed with the reaction mixture to amplify the pCA24N-cysE_A33T plasmid. The mutation was confirmed by sequencing.

Cultivation conditions

For strain maintenance, generation of deletion strains, and for growth during cloning LB medium (1% NaCl, 1% tryptone, 0.5% yeast extract) was used. Antibiotics for selection were used at the following concentrations: chloramphenicol, 30 μg/ml; kanamycin, 50 μg/ml; ampicillin, 100 μg/ml; streptomycin, 100 μg/ml. M9 minimal media was used for growth experiments (50 mM Na₂HPO₄, 20 mM KH₂PO₄, 1 mM NaCl, 20 mM NH₄Cl, 2 mM MgSO₄, 100 μM CaCl₂, 134 μM EDTA, 13 μM FeCl₃·6H₂O, 6.2 μM ZnCl₂, 0.76 μM CuCl₂·2H₂O, 0.42 μM CoCl₂·2H₂O, 1.62 μM H₃BO₃, 0.081 μM MnCl₂·4H₂O) with carbon sources added to the concentrations specified in the text and figures.

For growth experiments, overnight cultures were incubated in 4 mL LB medium with no antibiotics for deletion strains, with strains having been previously plated on antibiotic media to ensure genotype. In the event a strain carried a plasmid, the relevant antibiotic was also added to the pre-culture medium. Before inoculation of the experiment, cultures were harvested and washed four times in M9 medium without carbon source by centrifugation (6000 rpm, 3 min) to remove residual carbon sources from the cells. Plate reader experiments were inoculated with a starting OD₆₀₀ of...
0.01 using the washed culture. Plate reader experiments were carried out in 96-well microtiter plates (Nunclon Delta Surface, Thermo Fisher Scientific). Each well contained 150 μL of cell culture covered with 50 μL mineral oil (Sigma-Aldrich, Taukirchen, Germany), to avoid evaporation. An Infinite M200 Pro plate reader (Tecan) was used for incubation (37˚C), shaking, and OD₆₀₀ measurements. Three cycles of four shaking phases, each of 1 min were used (1. linear shaking at an amplitude of 3 mm, 2. orbital shaking at an amplitude of 3 mm, 3. linear shaking at an amplitude of 2 mm, and 4. orbital shaking at an amplitude of 2 mm). Optical density (OD 600 nm) was measured after each round of shaking (~12.5 min). Plate reader OD measurements were converted to cuvette values according to the formula ODₐₙ₃₃₅ = OD₆₀₀/0.23. Growth curves were processed in MATLAB and represent averages of (technical) triplicate measurements where the variability between triplicate measurements was less than 5% (unless explicitly stated, as in Figure 2). Within a given plate experiment, technical replicates averaged are replicates of a single genetic strain; biological replicates were generated for the Δ5 ΔmetC strain and Δ5 cysE* strains, these were engineered in parallel and were shown to behave in an identical manner.

Anaerobic growth experiments were performed in an Infinite M200 Pro plate reader (Tecan) inside a vinyl anaerobic chamber (N₂ with 10% CO₂, 2.5% H₂, model B, Coy Laboratory Products, Grass Lake, MI). M9 mediù for growth experiments were placed for at least 24 hr in the anaerobic chamber to allow for the exchange of dissolved oxygen from the media.

Whole-genome sequencing
Genomic DNA was extracted using the NucleoSpin Microbial DNA kit (MACHERY-NAGEL, Düren, Germany) following manufacturer's instructions. Library construction and genome sequencing were performed by Novogene (Cambridge, United Kingdom) using the paired-end Illumina sequencing platform. Analysis of the sequencing data was performed using the bresq pipeline (Deatherage and Barrick, 2014) and the SIJ488 reference genome sequence, which is derived from the MG1655 (NC_000913, GenBank). Supplementary file 1 shows the identified mutations, as compared to the parent strains.

Multiplex automated genome engineering (MAGE)
Introducing genomic point mutations was achieved by using multiplex automated genome engineering (MAGE; Wang et al., 2009). A single colony of desired strain(s) transformed with pORTMAGE (Nyerges et al., 2016; Addgene catalog no. 72680) was incubated in LB medium supplemented with 100 mg L⁻¹ of ampicillin at 30˚C in a shaking incubator. To start the MAGE cycle, overnight cultures were diluted by 100 times in the same medium and cultivated to an optical density of 0.4–0.5 at 600 nm. 1 mL of each culture was transferred to sterile microcentrifuge tubes, and then transferred to 42˚C theromomixer (Thermomixer C, Eppendorf, Hamburg, Germany) to express λ-Red genes by heat shock for 15 min at 1000 rpm. After induction, cells were quickly chilled on ice for at least 15 min, and then made electrocompetent by washing three times with ice-cold ddH₂O. 40 μL of electrocompetent cell was mixed with 2 μL of 50 μM of oligomer stock solution and the final volume of the suspension was adjusted to 50 μL. The oligomer used to introduce the mutation was CysE_A33T_MAGE (Supplementary file 2). Electroporation was done on Gene Pulser XCell (Bio-Rad) set to 1.8 kV, 25 μF capacitance, and 200 Ω resistance for 1 mm gap cuvette. Immediately after electroporation, 1 mL of LB was added to cuvette and the electroporation mixtures in LB were transferred to sterile culture tubes and cultured with shaking at 30˚C, 240 rpm for 1 hr to allow for recovery. After recovery, 2 mL of LB medium supplemented with ampicillin was added and then further incubated in the same condition. When the culture reached an OD₆₀₀ of 0.4–0.5, cells were either subjected to additional MAGE cycles or analyzed for genotype via PCR and sequencing.

Isotopic-labeling of proteinogenic amino acids
Stationary isotope tracing experiments using ¹³C-1-labeled glucose, ¹³C-3-labeled glucose, and labeled sodium propionate (Sigma-Aldrich) were performed in order to understand the metabolic activities responsible for isoleucine biosynthesis. All experiments were performed in duplicate. Strains were grown in M9 with 10 mM labeled glucose as sole carbon source in the case of labeled glucose experiments, and 10 mM unlabeled glucose with 5 mM sodium propionate-1-¹³C in the case of the anaerobic experiments (extra carbon sources added where relevant at concentrations
indicated). Anaerobic experiments were performed also with an anaerobic pre-culture in M9 medium. The equivalent of 1 mL of culture at OD_{600} one was harvested by centrifugation and washed twice in water. Cellular biomass was hydrolyzed by boiling in acid (95°C in 6 M HCl for 24 hr [You et al., 2012]). After drying the samples at 95°C, the samples were resuspended in ultra-pure water and the masses of amino acids analyzed with UPLC–ESI–MS as previously described (Giavalisco et al., 2011). Chromatography was performed with a Waters Acquity UPLC system (Waters), using an HSS T3 C_{18} reversed phase column (100 × 2.1 mm², 1.8 μm; Waters). 0.1% formic acid in H₂O (A) and 0.1% formic acid in acetonitrile (B) were the mobile phases. Flow rate was 0.4 mL/min and the gradient was: 0 to 1 min – 99% A; 1 to 5 min – linear gradient from 99% A to 82%; 5 to 6 min – linear gradient from 82% A to 1% A; 6 to 8 min – kept at 1% A; 8 to 8.5 min – linear gradient to 99% A; 8.5 to 11 min – re-equilibrate. Mass spectra were acquired using an Exact mass spectrometer (Thermo Fisher Scientific) in positive ionization mode, with a scanning range of 50.0 to 300.0 m/z. Spectra were recorded during the first 5 min of the LC gradients. Retention times for amino acids under these conditions were determined by analyzing amino-acid standards (Sigma-Aldrich) under the same conditions. Data analysis was performed using Xcalibur (Thermo Fisher Scientific).

### Enzyme assays for MetB, MetC, and CysE

_E. coli_ BL21 (DE3) was transformed with the overexpression plasmids pCA24N-metB, pCA24N-metC, pCA24N-cysE, or pCA24N-cysE_A33T, and grown on LB agar containing 34 μg/mL chloramphenicol. An expression culture in TB containing 34 μg/mL chloramphenicol was inoculated from the plate and grown at 37°C while shaking at 110 rpm until an OD_{600} of roughly 0.9 was reached. The culture was shaken for another 4 hr at 37°C or overnight at 20°C and 110 rpm for gene expression. The culture was harvested by centrifugation at 4°C and 5000 g. Alternatively, the pellet was stored at –20°C. The pellet was resuspended in Buffer A (50 mM Tris-HCl pH 7.9, 500 mM NaCl) containing 10 μg/mL DNaseI and 5 mM MgCl₂. The cells were lysed by ultrasonication and the lysate was cleared by ultracentrifugation at 50,000 g and 4°C for 45 min followed by filtration through a 0.45 μm syringe filter. The lysate was loaded onto a 1 mL HisTrap FF column (GE Healthcare, Freiburg, Germany) equilibrated in Buffer A. Unspecifically bound protein was washed with Buffer A containing 50 mM imidazole. The protein was eluted with Buffer A containing 500 mM imidazole and subsequently desalted over a HiTrap 5 mL desalting column (GE Healthcare) using 20 mM Tris-HCl pH 7.9, 50 mM NaCl. The protein was concentrated using Amicon Ultra-4 centrifugal filters (Merck Millipore, Darmstadt, Germany). The protein was stored in 50% (v/v) glycerol at –20°C.

The MetB elimination reaction (red arrow, reaction one in Figure 5) was kinetically characterized using a spectrophotometric assay. The assays were performed at 30°C in 10 mm quartz cuvettes (Hellma Analytics, Germany) on a Cary-60 UV/Vis spectrometer (Agilent Technologies Inc, Santa Clara, CA). In the elimination reaction, MetB cleaves O-succinyl-L-homoserine to succinate and 2-ketobutyrate. The latter can be reduced by the coupling enzyme lactate dehydrogenase using NADH as reducing agent. The NADH consumption was followed at 340 nm (ε_{NADH} = 6.22 mM⁻¹ cm⁻¹). The reaction was started by adding O-succinyl-L-homoserine to 22 nM MetB in 200 mM Tris-HCl pH 8.0 containing ~1 U of lactate dehydrogenase (Sigma-Aldrich Chemie GmbH, Germany) and 300 μM NADH. Kinetic parameters were determined from a Michaelis-Menten fit of 18 data points.

In the presence of O-succinyl-L-homoserine and cysteine or homocysteine, MetB can catalyze the replacement reaction yielding succinate and cystathionine or homolanthionine, respectively (MetB black arrow and reaction two in Figure 5). However, the presence of cysteine or homocysteine does not necessarily exclude the elimination reaction activity of the MetB enzyme (reaction one in Figure 5). To determine the reaction route, MetB was incubated with O-succinyl-L-homoserine and cysteine or homocysteine in separate experiments. To investigate the MetB reaction route at physiological cysteine or homocysteine concentrations, we ran an assay with 100 nM of MetB in 200 mM Tris-HCl pH 8.0 and 50 μM pyridoxal-5’-phosphate in the presence of 6 mM O-succinyl-L-homoserine and 300 μM of either cysteine or homocysteine. The assay mixture was incubated at 30°C and samples were quenched with 5% formic acid at specific time points. To investigate the MetB reaction route at high concentrations of cysteine and homocysteine, 90 nM of MetB was assayed in 200 mM Tris-HCl pH 8.0 in the presence of 6 mM O-succinyl-L-homoserine and either 3 mM L-cysteine or 6 mM L-homocysteine. The assay mixture was incubated at 30°C and samples were quenched with 10% formic acid at specific time points. In both experiments, the formation of 2 KB and succinate
was quantitatively determined by HPLC-MS/MS analysis. The levels of succinate shown in Figure 5—
figure supplement 1B–E are the levels of succinate generated from the replacement reaction of
MetB exclusively; the succinate generated by the elimination reaction has already been taken into
account by subtracting the concentration of 2 KB in the sample (elimination reaction produces an
equimolar amount of 2 KB and succinate).

To test whether MetC can produce 2 KB, we incubated 24 μM of MetC and 10 mM of homocyste-
ine or 87 μM MetC and 33 mM of methionine in 50 mM Tris-HCl pH 8.5 at 30°C (reaction 5 or 6 in
Figure 5). No 2 KB generation could be detected when following the NADH consumption of the
coupling enzyme lactate dehydrogenase (~1 U of lactate dehydrogenase and 300 μM NADH). Some
MetC variants from other organisms have also been reported to catalyze both the β-elimination
(canonical E. coli) and the γ-elimination of cystathionine resulting in pyruvate and 2 KB, respectively.
In this case, as both these products are accepted by the coupling enzyme lactate dehydrogenase
(Kim and Whitesides, 1988), the reaction route could not be determined spectrophotometrically.
Instead, 20 nM of MetC was incubated with 3 mM cystathionine in 50 mM Tris-HCl pH 8.5. The assay
mixture was incubated at 30°C and samples were quenched with 10% formic acid at specific time
points. Formation of 2 KB and pyruvate was quantitatively determined using HPLC-MS/MS analysis.
Only pyruvate could be detected in these samples indicating no γ-elimination of cystathionine by the E.
coli MetC enzyme.

The activity of CysE and its variant CysE Ala33Thr were assayed by photospectrometrically follow-
ning the reaction of released CoA with 5,5′-dithiobis-(2-nitrobenzoic acid) (DTNB) at 412 nm using an
extinction coefficient of 14.4 mM⁻¹ cm⁻¹. The assays were performed at 30°C in 10 mm quartz cuv-
ettes (Hellma Analytics) on a Cary-60 UV/Vis spectrometer (Agilent Technologies). The reaction was
started by adding the CysE variant to 0.2 mM DTNB in 100 mM potassium phosphate buffer pH 8.
The dependence of the turnover number on acetyl-CoA concentration (synthesized as previously
described [Peter et al., 2016]) was measured with 0.3 nM CysE or 1.3 nM CysE A33T at a saturating
L-serine concentration of 20 mM. The dependence of the turnover number on L-serine concentra-
tion was measured with 1.0 nM CysE or 2.0 nM CysE A33T at a non-saturating acetyl-CoA concentration
of 0.6 mM.

Succinate, pyruvate, and 2-ketobutyrate in the enzymatic assays were quantitatively determined
using LC-MS/MS based on external calibration curves. The calibration curves were generated with at
least seven concentrations within the linear range of the analyte in the matrix of the assay samples.
The compounds were separated over a Kinetex EVO C18 chromatography column (50 x 2.1 mm²;
3.5 μm, 100 Å, Phenomenex) equipped with a 20 x 2.1 mm² guard column of similar specificity on an
Agilent Infinity II 1290 HPLC system. The column was heated to 55°C and 5 μL of sample or stan-
dard were injected. A constant flow rate of 0.2 mL/min was applied. Mobile phase A (0.1% formic
acid in water [Honeywell, Morristown, New Jersey]) and mobile phase B (0.1% formic acid in acetoni-
trile [Honeywell]) were mixed according to the following mobile phase profile: 0–1.6 min constant at
0% B; 1.6–5 min from 0% to 95% B; 5–6 min constant at 95% B; 6–6.5 min from 95% to 0% B; 6.5–10
min constant at 0% B. An Agilent 6495 ion funnel mass spectrometer equipped with an electrospray
ionization source in negative mode was used for analysis under the following conditions: ESI spray
voltage 1500 V, sheath gas 200˚ C at 12 L/min, nebulizer pressure 30 psig, and drying gas 180˚ C at
11 L/min. Mass transition and retention time of the compounds were compared to standards for
identification. The MassHunter software (Agilent, Santa Clara, CA, USA) was used for chromatogram
integration.

Anaerobic production of 2-ketobutyrate from PFL (i.e. PflB) and TdcE
Propionyl-CoA was synthesized and purified as described previously (Peter et al., 2016). The dried
powder was kept at −20°C and dissolved in 50 mM acetate (pH 4.5) before use. The concentration
was determined by two independent methods. First, a NanoDrop 2000 Spectrophotometer
(Thermo Fisher Scientific) was used with the extinction coefficient of saturated CoA-esters at 260 nm
(ε260nm = 16.4 mM⁻¹ cm⁻¹). Second, the depletion of the compound when incubated with the
enzyme PduP (CoA-acylating propionaldehyde dehydrogenase; Zarzycki et al., 2017), following the
oxidation of NADH at 340 nm (ε340nm = 6.22 mM⁻¹ cm⁻¹). Both methods gave the same results
within a 5% error.

Overexpression plasmids for PFL-AE (PFL-activating enzyme), PFL, and TdcE were obtained from the
ASKA collection. PFL-AE, PFL, and TdcE were produced in E. coli BL21(DE3). 1 L terrific broth
(TB) containing 34 μg/mL chloramphenicol was inoculated with freshly transformed cells and incubated at 37°C. After reaching an OD_{600} of 0.8, the expression was induced by adding IPTG to a final concentration of 0.5 mM and the incubation temperature was lowered to 25°C. Cells were harvested after 16 hr by centrifugation (4500 g, 10 min) and resuspended in buffer A (50 mM HEPES-KOH pH 7.8, 500 mM KCl) containing 10 μg/mL DNaseI and 5 mM MgCl₂. If not used immediately, cell pellets were flash-frozen in liquid nitrogen and stored at −20°C. The cell lysate obtained by sonication was clarified by centrifugation 75,000 g at 4°C for 45 min. The supernatant was filtered through a 0.4 μm syringe tip filter (Sartorius). Ni-affinity purification was performed with an Äkta FPLC system from GE Healthcare. The filtered soluble lysate was loaded onto a 1 ml Ni-Sepharose Fast Flow column (HisTrap FF, GE Healthcare) that had been equilibrated with 10 mM buffer A. After washing with 20 ml 85% buffer A, 15% buffer B (50 mM HEPES-KOH pH 7.8, 500 mM KCl, 500 mM imidazole), the protein was eluted with 100% buffer B. Fractions containing purified protein were pooled and the buffer was exchanged to storage buffer (50 mM HEPES-KOH pH 7.8, 150 mM KCl) using a desalting column (HiTrap, GE Healthcare). Proteins were concentrated by ultrafiltration (Amicon Ultra). Protein concentration was determined on a NanoDrop 2000 Spectrophotometer (Thermo Fisher Scientific) using the extinction coefficient at 280 nm, as calculated by ProtParam (Gasteiger, 2005). Enzyme purity was confirmed using SDS-PAGE. The purified proteins were flash-frozen in liquid nitrogen and stored at −80°C.

PFL-AE was expressed and purified as PFL and TdcE. However, after the HisTrap purification step, the protein was transferred to an anaerobic glovebox. The reconstitution of the iron-sulfur cluster was performed following the protocol of Broderick et al. with a few adaptations (Byer et al., 2018). Using a PD-10 desalting column (GE Healthcare), the buffer was exchanged to an anaerobic reconstitution buffer (50 mM Tris-SO₄ pH 7.5, 150 mM KCl, 5 mM dithiothreitol [DTT]). The DTT was added fresh to the buffer right before use. After buffer exchange the protein was diluted with reconstitution buffer to a concentration of ~3 mg/mL (~100 μM) and transferred to a glass vial with a magnetic stirring bar. The vial was placed on ice on a magnetic stirrer. A 10 mM stock solution of FeCl₃ was added in 10 aliquots over the course of 1 hr to a final concentration of 600 μM. Then, a 50 mM stock of Na₂S·(H₂O)₉ was added in 10 aliquots over the course of 1 hr to a final concentration of 600 μM. This mixture was incubated on ice with stirring for 6 hr. Then the protein was desalted with a PD-10 column (equilibrated with reconstitution buffer), and concentrated on an Amicon to 30 mg/mL. UV-Vis spectra were recorded before and after the reconstitution with a Cary 4000 UV-Vis spectrometer (Agilent Technologies) using a FiberMate2 Fiber Optic Coupler (Harrick Scientific Products) with a path length of 1 cm (Figure 8—figure supplement 2). The reconstituted PFL-AE was transferred in aliquots to rubber stoppered HPLC vials, flash frozen in liquid nitrogen and stored at −80°C.

All anaerobic steps were performed in an anaerobic glovebox (Coy Laboratories) under an N₂ atmosphere containing 3% to 3.5% H₂. Residual O₂ was removed by palladium catalysts. O₂ concentration was monitored and maintained below 5 ppm at all times. All stocks were prepared under anaerobic conditions or prepared outside of the glovebox and equilibrated for at least 3 hr inside the glovebox. Dithionite and DTT stocks were freshly prepared under anaerobic conditions each day.

To activate PFL, 100 mM Tris-HCl pH 7.5, 50 mM KCl, 10 mM DTT, 50 mM sodium oxamate, 125 μM SAM, and 25 μM PFL-AE were incubated for 10 min at 30°C. Then, 25 μM PFL and 62.5 μM sodium dithionite were added. After 30 min at 30°C, the activation mix was placed on ice.

To activate TdcE, 100 mM Tris-HCl pH 7.5, 50 mM KCl, 10 mM DTT, 50 mM sodium oxamate, 50 μM SAM, and 10 μM PFL-AE were incubated for 10 min at 30°C. Then, 10 μM TdcE and 25 μM sodium dithionite were added. After 30 min at 30°C, the activation mix was placed on ice.

Glycyl radical formation was confirmed by exposing the activation mix to oxygen followed by SDS-PAGE analysis. The glycyl radical rapidly reacts with oxygen, cleaving off a ~3.5 kDa fragment from the C-terminus. PFL and TdcE were activated to a similar degree, as demonstrated in Figure 8—figure supplement 2.

Activation and stability of the activation mix was tested by the pyruvate cleavage assay (Figure 8—figure supplement 2). For this, 100 mM Tris-HCl pH 7.5, 50 mM KCl, 10 mM malate, 1 mM NAD, 20 mM pyruvate, 250 μM CoA, 4.5 U/mL citrate synthase from the porcine heart (Sigma-Aldrich), 27.5 U/mL malic dehydrogenase from the porcine heart (Sigma-Aldrich) were mixed and
the reaction initiated by adding the activation mix in 100-fold dilution. The reduction of NAD was followed at 340 nm.

2-Ketobutyrate formation kinetics were measured in a coupled enzyme assay: L-lactate dehydrogenase reduces 2-ketobutyrate to 2-hydroxybutyrate with the consumption of NADH. Absorption at 340 nm was recorded with a Cary 4000 UV-Vis spectrometer using a FiberMate2 Fiber Optic Coupler with a path length of 1 cm. Each substrate concentration was measured in triplicate. One activation mix was used up to 1.5 hr storage on ice, then a fresh activation mix was prepared for kinetic measurements. PFL and TdcE kinetic parameters take the half-of-the-sites activity (Unkrig et al., 1989) into account, that is, the dimer mass was used to calculate the protein concentration. Because the exact active site concentration (i.e. the glycy radical content after activation) was not determined, \( V_{\text{max}} \) is reported as specific activity. For Michaelis-Menten parameter calculation and fitting, GraphPad Prism 8 was used. For PFL with propionyl-CoA as substrate the data was fit to the substrate inhibition equation.

Propionyl-CoA kinetics: 100 mM Tris-HCl pH 7.5, 50 mM KCl, 500 mM sodium formate, 200 µM NADH, 25 U/mL L-lactate dehydrogenase from bovine heart (LDH; Sigma-Aldrich). The reaction was started by adding an activation mix in 100-fold dilution. Thus, the final concentration was 250 nM PFL and 100 nM TdcE, respectively. For PFL, the propionyl-CoA concentrations were 100 µM, 250 µM, 500 µM, 1 mM, 2.5 mM and for TdcE, 150 µM, 350 µM, 550 µM, 800 µM, 1.6 mM, 1.9 mM. In both cases, there was no activity in the absence of propionyl-CoA.

Formate kinetics: 100 mM Tris-HCl pH 7.5, 50 mM KCl, 200 µM NADH, 25 U/mL LDH. Propionyl-CoA was added to a final concentration of 800 µM (PFL) or 1.9 mM (TdcE). The reaction was started by adding an activation mix in 100-fold dilution. Thus, the final concentration was 250 nM PFL and 100 nM TdcE, respectively. For PFL, the formate concentrations were 12 mM, 40 mM, 100 mM, 250 mM, 500 mM, and for TdcE, 5 mM, 12 mM, 40 mM, 100 mM, 500 mM. In both cases, there was no activity in the absence of formate.

### Transcript level analysis by reverse transcriptase quantitative PCR

To determine mRNA levels, total RNA was extracted from three biological replicates from cells in exponential phase growing on M9 minimal medium with 10 mM glucose (supplemented with 2 mM isoleucine in case of the \( \Delta s \) strain). Total RNA was purified using the RNeasy Mini Kit (Qiagen, Hilden, Germany) as instructed by the manufacturer. In brief, \( \sim 2.5 \times 10^8 \) cells (0.5 mL of \( \text{OD}_{600} \) 0.5) were mixed with 2 volumes of RNAprotect Bacteria Reagent (Qiagen) and pelleted, followed by enzymatic lysis, on-column removal of genomic DNA with RNase-free DNase (Qiagen) and spin-column-based purification of RNA. Concentration and integrity of the isolated RNA were determined by NanoDrop and gel electrophoresis. cDNA was synthesized via reverse transcription of 500 ng RNA with the qScript cDNA Synthesis Kit (QuantaBio, Beverly, MA). Quantitative real-time PCR was performed two times in technical triplicates using the Maxima SYBR Green/ROX qPCR Master Mix (Thermo Fisher Scientific). An input corresponding to 25 pg total RNA/cDNA was used per reaction. Non-specific amplification products were excluded by melting curve analysis. The gene encoding 16S rRNA (\( \text{rrsA} \)) was chosen as a well-established reference transcript for expression normalization (Zhou et al., 2011). Two primer pairs for amplification of \( \text{metA} \) and \( \text{metB} \), respectively, were used for qPCR (Supplementary file 2). Equal amplification efficiencies between the primers for the genes of interest and the reference gene were assumed. Differences in expression levels were calculated according to the \( 2^{-\Delta\Delta CT} \) method (Livak and Schmittgen, 2001). Reported data represents the average of \( 2^{\Delta\Delta CT} \) values that were calculated for each sample individually relative to the average of all biological WT replicate \( \Delta CT_{(\text{GOL})-\text{Ctrl} (\text{metA})} \) values. Negative control assays with the direct input of RNA (without previous reverse transcription) confirmed that the observed fold change of \( \text{metA} \) and \( \text{metB} \) in the \( \Delta s \) \( \text{cysE}^+ \) sample was not due to an increase in genomic DNA contamination, that is, it accounted for less than \( \sim 10\% \) of the signal (\( \Delta CT \) between +RT/–RT samples >3).

### Cellular concentrations of methionine and succinyl-homoserine

To assess the relative intracellular concentration changes of succinyl-homoserine and methionine among different strains, we cultivated the strains in four independent replicates of 20 mL media in 100 mL Erlenmeyer flasks (\( n = 4 \)) to mid-exponential phase (\( \text{OD}_{600} \) approx. 0.4–0.8). After measuring \( \text{OD}_{600} \), 12 mL culture were immediately harvested and quenched with 36 mL precooled 60%...
methanol (v/v). Cells were spun down by centrifugation (Beckman Allegra 25R refrigerated centrifuge): 5000 g, −10°C, 10 min, and stored at −80°C before extraction. On ice, the cells were added to precooled extraction fluid (10 mM Tris-HCl, 1 mM EDTA, 50% MeOH (v/v), pH 7.0) and chloroform 0.5 mL of each, briefly vortexed and shaken at 4°C for 2 hr for extraction. After centrifugation (5000 g, −10°C, 10 min), the upper phase of the extracts were filtered (Fisher syringe filter, PTFE, 0.2 μm, 13 mm diameter), and stored at −80°C until further processing.

Quantitative determination was performed using an HRES-LC-MS. The chromatographic separation was performed on a Vanquish HPLC system using a ZicHILIC SeQuant column (150 × 2.1 mm², 3.5 μm particle size, 100 Å pore size) connected to a ZicHILIC guard column (20 × 2.1 mm², 5 μm particle size) (Merck KgAA) with a constant flow rate of 0.3 mL/min with mobile phase A being 0.1% formic acid in 99:1 water:acetonitrile (Honeywell) and phase B being 0.1% formic acid 99:1 water:acetonitrile (Honeywell) at 45°C.

The injection volume was 2 μL. The mobile phase profile consisted of the following steps and linear gradients: 0–1 min constant at 90% B; 1–8 min from 90% to 20% B; 8–9 min constant at 20% B; 9–9.1 min from 20% to 90% B; 9.1–11 min constant at 90% B. An ID-X Orbitrap mass spectrometer equipped with a HESI electrospray ion source (Thermo Fisher Scientific) was used at the following conditions: ESI spray voltage 3500 V, sheath gas at 50 AU, auxiliary gas at 10 AU and sweep gas at 1 AU, with the vaporizor temperature being 350°C and the ion transfer tube temperature being 325°C. Compounds were identified based on their accurate mass and retention time compared to standards. Chromatograms were integrated using Xcalibur software (Thermo Fisher Scientific). Absolute concentrations were calculated based on an external calibration curve.

Relative cellular concentrations of cysteine and homocysteine

The concentration of cysteine could not be determined using the above method due to its instability during the measurement process. Therefore, we used another method to assess the relative intercellular concentration of these amino acids.

To assess the relative intracellular concentration changes of cysteine, homocysteine, and 2 KB among different strains, we cultivated the strains in four independent replicates of 4 mL media in glass test tubes (n = 4) to mid-exponential phase (OD₆₀₀ approx. 0.4–0.8). About 10⁹ cells (equivalent of 1 mL of OD₆₀₀ = 1 culture) were collected on 0.45 μm Durapore membrane filters (Merck Millipore, Ireland). After washing the filters with 1 mL of fresh medium in under 10 s, the filters with cells were placed in 5 mL of −20°C 40:40:20 (v/v/v) acetonitrile/methanol/water overnight. 4 mL of debris-free extract per sample was dried by vacuum centrifugation at 35°C overnight. The dried metabolites were stored at −20°C until further processing.

Metabolites were methoxyaminated and trimethylsilylated before gas chromatography-atmospheric pressure chemical ionization-quadrupole time of flight mass spectrometry (GC/APCI-qTOF-MS) as described earlier (Kopka et al., 2017). GC/APCI-qTOF-MS analyses were performed with an Agilent 7890B gas chromatograph (Agilent Technologies Deutschland GmbH, Germany) hyphenated to a Bruker Impact II mass spectrometer (Bruker Daltonik GmbH, Bremen, Germany). The machine parameters and analysis procedures were exactly as detailed in Kopka et al., 2017. Data mining was performed with Profile Analysis Version 2.2 software (Bruker Daltonik GmbH).

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Charles AR Cotton, Conceptualization, Data curation, Formal analysis, Investigation, Visualization, Methodology, Writing - original draft, Writing - review and editing; Iria Bernhardsgrütter, Formal analysis, Investigation, Visualization, Methodology; Hai He, Nicole Paczia, Alexander Erban, Stepan Toman, Marian Dempfle, Joachim Kopka, Steffen N Lindner, Formal analysis, Investigation, Methodology; Simon Burgener, Luca Schulz, Alberto De Maria, Investigation, Methodology; Beau Dronsella, Formal analysis, Validation, Investigation, Methodology; Tobias J Erb, Supervision; Arren Bar-Even, Conceptualization, Supervision, Funding acquisition, Investigation, Visualization, Writing - original draft, Project administration, Writing - review and editing

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#### Additional files

**Supplementary files**

- **Supplementary file 1.** Summary of identified mutations in 16 strains lacking threonine deaminases (ΔsilVA ΔtdcB) or threonine and serine deaminases (ΔsilVA ΔtdcB ΔsdaA ΔsdaB ΔtdcG) that were evolved to grow on 10 mM glucose after 70 hr without isoleucine or 2 KB.
- **Supplementary file 2.** List of all primers used in this study.
Data availability

All data generated or analysed during this study are included in the manuscript and supporting files. Source data files have been provided for Figures 2 and 7 as well as for the metabolomic analysis.

References

Abramsky T, Rowland LP, Shemin D. 1962. The formation of isoleucine from beta-methylaspartic acid in Escherichia coli W. The Journal of Biological Chemistry 240:PC265–PC266.

Alting AC, Engels W, van Schalkwijk S, Exterkate FA. 1995. Purification and characterization of cystathionine (beta)-lyase from Lactococcus lactis subsp cremoris B78 and its possible role in flavor development in cheese. Applied and Environmental Microbiology 61:4037–4042. DOI: https://doi.org/10.1128/AEM.61.11.4037-4042.1995, PMID: 16535166

Amarita T, Yvon M, Nardi M, Chambellon E, Delettre J, Bonnarme P. 2004. Identification and functional analysis of the gene encoding methionine-gamma-lyase in Brevibacterium linens. Applied and Environmental Microbiology 70:7348–7354. DOI: https://doi.org/10.1128/AEM.70.12.7348-7354.2004, PMID: 15574935

Awano N, Wada M, Kohdoh A, Oikawa T, Takagi H, Nakamori S. 2003. Effect of cysteine desulphhydrase gene disruption on L-cysteine overproduction in Escherichia coli. Applied Microbiology and Biotechnology 62:239–243. DOI: https://doi.org/10.1007/s00253-003-1262-2, PMID: 12883870

Baba T, Ara T, Hasegawa M, Takai Y, Okumura Y, Baba M, Datsenko KA, Tomita M, Wanner BL, Mori H. 2006. Construction of Escherichia coli K-12 in-frame, single-gene knockout mutants: the keio collection. Molecular Systems Biology 2:2006–2008. DOI: https://doi.org/10.1038/msb4100050, PMID: 16738554

Bennett BD, Kimball EH, Gao M, Osterhout R, Van Dien SJ, Rabinowitz JD. 2009. Absolute metabolite concentrations and implied enzyme active site occupancy in Escherichia coli. Nature Chemical Biology 5:593–599. DOI: https://doi.org/10.1038/nchembio.186, PMID: 19561621

Born TL, Blanchard JS. 1999. Enzyme-catalyzed acylation of homoserine: mechanistic characterization of the Escherichia coli metaA-encoded homoserine transsuccinylase. Biochemistry 38:14416–14423. DOI: https://doi.org/10.1021/bi991710o, PMID: 10572016

Buchanan BB. 1969. Role of ferredoxin in the synthesis of alpha-ketoobutyrate from propionyl coenzyme A and carbon dioxide by enzymes from photosynthetic and nonphotosynthetic Bacteria. The Journal of Biological Chemistry 244:4218–4223. PMID: 5800441

Byer AS, McDaniel EC, Impano S, Broderick WE, Broderick JB. 2018. Mechanistic studies of radical SAM enzymes: pyruvate Formate-Lyase activating enzyme and lysine 2,3-Aminomutase case studies. Methods in Enzymology 606:269–318. DOI: https://doi.org/10.1016/bs.mie.2018.04.013, PMID: 30097096

Charon NW, Johnson RC, Peterson D. 1974. Amino acid biosynthesis in the spirochete Leptospira: evidence for a novel pathway of isoleucine biosynthesis. Journal of Bacteriology 117:203–211. DOI: https://doi.org/10.1128/JB.117.1.203-1.1974, PMID: 4808901

Cicchillo RM, Baker MA, Schnitzer EJ, Newman EB, Krebs C, Booker SJ. 2004. Escherichia coli L-serine deaminase requires a [4Fe-4S] cluster in catalysis. Journal of Biological Chemistry 279:32418–32425. DOI: https://doi.org/10.1074/jbc.M404381200, PMID: 15155761

D’Ari R, Casadesús J. 1998. Underground metabolism. BioEssays 20:181–186. DOI: https://doi.org/10.1002/(SICI)1521-1878(199802)20:2<181::AID-BIES10>2.0.CO;2-M, PMID: 9631663

Datsenko KA, Wanner BL. 2000. One-step inactivation of chromosomal genes in Escherichia coli K-12 using PCR products. PNAS 97:6640–6645. DOI: https://doi.org/10.1073/pnas.97163297, PMID: 10829079

Deatherage DE, Barrick JE. 2014. Identification of mutations in laboratory-evolved microbes from next-generation sequencing data using breseq. Methods in Molecular Biology 1151:165–188. DOI: https://doi.org/10.1007/978-1-4939-0554-6_12, PMID: 24838886

Denk D, Bock A. 1987. L-Cysteine biosynthesis in Escherichia coli: nucleotide sequence and expression of the serine acetyltransferase (cysE) Gene from the Wild-type and a Cysteine-excreting mutant. Microbiology 133:515–525. DOI: https://doi.org/10.1099/00221287-133-3-515

Drevland RM, Waheed A, Graham DE. 2007. Enzymology and evolution of the pyruvate pathway to 2-oxobutyrate in Methanocaldococcus jannaschii. Journal of Bacteriology 189:4391–4400. DOI: https://doi.org/10.1128/JB.00116-07, PMID: 17449626

Dwiwidri CM, Ragin RC, Uren JR. 1982. Cloning, purification, and characterization of beta-cystathionase from Escherichia coli. Biochemistry 21:3064–3069. DOI: https://doi.org/10.1021/bi00256a005, PMID: 7049234

Elkmans B, Jaenchen R, Thauer RK. 1983. Propionate assimilation by methanogenic Bacteria. Archives of Microbiology 136:106–110. DOI: https://doi.org/10.1007/BF00404782

Figge RM. 2006. Methionine Biosynthesis in Escherichia coli and Corynebacterium glutamicum. In: Wendisch VF (Ed). Amino Acid Biosynthesis - Pathways, Regulation and Metabolic Engineering. Springer. p. 163–193. DOI: https://doi.org/10.1007/978-3-540-48596-4

Flint DH, Tumlinell JF, Miller TJ. 1996. Studies on the synthesis of the Fe-S cluster of dihydroxy-acid dehydratase in Escherichia coli crude extracts. Isolation of O-acetylserylserine sulphydrylases A and B and beta-cystathionase based on their ability to mobilize sulfur from cysteine and to participate in Fe-S cluster synthesis.
Cotton et al. eLife 2020;9:e54207. DOI: https://doi.org/10.7554/eLife.54207

The Journal of Biological Chemistry 271:16053–16067. DOI: https://doi.org/10.1074/jbc.271.27.16053, PMID: 8663055

Gasteiger E. 2005. The Proteomics Protocols Handbook. In: Walker J. M (Ed). Protein Identification and Analysis Tools on the ExPASy Server. Humana Press. p. 571–607. DOI: https://doi.org/10.1385/1-59259-890-0:571

Giavalisco P, Li Y, Matthes A, Eckhardt A, Hubberten HM, Hesse H, Segu S, Hummel J, Köhl K, Willmitzer L. 2011. Elemental formula annotation of polar and lipophilic metabolites using (13)C, (15)N and (34)S isotope Labelling, in combination with high-resolution mass spectrometry. The Plant Journal 68:364–376. DOI: https://doi.org/10.1111/j.1365-313X.2011.04682.x, PMID: 21699588

Guo AC, Jewison T, Wilson M, Liu Y, Knox C, Djoumbou Y, Lo P, Mandal R, Krishnamurthy R, Wishart DS. 2013. ECMDB: the E. coli Metabolite Database. Nucleic Acids Research D625–D630. DOI: https://doi.org/10.1093/nar/gks992

Hasona A, Kim Y, Healy FG, Ingram LO, Shanmugam KT. 2004. Pyruvate formate lyase and acetate kinase are essential for anaerobic growth of Escherichia coli on xylose. Journal of Bacteriology 186:7593–7600. DOI: https://doi.org/10.1128/JB.186.22.7593-7600.2004, PMID: 15516572

Hesslinger C, Fairhurst SA, Sawers G. 1998. Novel keto acid formate-lyase and propionate kinase enzymes are components of an anaerobic pathway in Escherichia coli that degrades L-threonine to propionate. Molecular Microbiology 27:477–492. DOI: https://doi.org/10.1046/j.1365-2958.1998.00696.x, PMID: 9484901

Hochuli M, Patzelt H, Oesterhelt D, Wüthrich K, Syperski T. 1999. Amino acid biosynthesis in the halophilic archaeon Haloarcula hispanica. Journal of Bacteriology 181:3226–3237. DOI: https://doi.org/10.1128/JB.181.10.3226-3237.1999, PMID: 10322026

Holbrook EL, Greene RC, Krueger JH. 1990. Purification and properties of cystathionine gamma-synthase from overproducing strains of Escherichia coli. Biochemistry 29:435–442. DOI: https://doi.org/10.1021/bi00454a019, PMID: 2405903

Irmler S, Raboud S, Beisert B, Rauhut D, Berthoud H. 2008. Cloning and characterization of two Lactobacillus casei genes encoding a cystathionine lyase. Applied and Environmental Microbiology 74:99–106. DOI: https://doi.org/10.1128/AEM.00745-07, PMID: 17993563

Itikawa H, Baumberg S, Vogel HJ. 1968. Enzymic basis for a genetic suppression: accumulation and decylation of N-acetylglutamic γ-semialdehyde in enterobacterial mutants. Biochimica Et Biophysica Acta (BBA) - Enzymology 159:547–550. DOI: https://doi.org/10.1016/0005-2744(68)90142-3

Jensen RA. 1976. Enzyme recruitment in evolution of new function. Annual Review of Microbiology 30:409–425. DOI: https://doi.org/10.1146/annurev.mi.30.100176.002205, PMID: 791073

Jensen SI, Lennen RM, Herrgård MJ, Nielsen AT. 2016. Seven gene deletions in seven days: fast generation of Escherichia coli strains tolerant to acetate and osmotic stress. Scientific Reports 5:17874. DOI: https://doi.org/10.1038/srep17874

Joshi V, Joung JG, Fei Z, Jander G. 2010. Interdependence of threonine, methionine and isoleucine metabolism in plants: accumulation and transcriptional regulation under abiotic stress. Amino Acids 39:933–947. DOI: https://doi.org/10.1007/s00726-010-0505-7, PMID: 20186554

Joshi V, Jander G. 2009. Arabidopsis methionine gamma-lyase is regulated according to isoleucine biosynthesis needs but plays a subordinate role to threonine deaminase. Plant Physiology 151:367–378. DOI: https://doi.org/10.1104/pp.109.138651, PMID: 19571310

Kaneshia M. 2000. KEGG: kyoto encyclopedia of genes and genomes. Nucleic Acids Research 28:27–30. DOI: https://doi.org/10.1093/nar/28.1.27

Khersonsky O, Tawfiq DS. 2010. Enzyme promiscuity: a mechanistic and evolutionary perspective. Annual Review of Biochemistry 79:471–505. DOI: https://doi.org/10.1146/annurev-biochem-030409-143718, PMID: 20235827

Kim J, Kershner JP, Novikov Y, Shoemaker RK, Copley SD. 2010. Three serendipitous pathways in E. coli sp. PCC 7002. Synechococcus 10:291–299. DOI: https://doi.org/10.1128/JB.181.10.3226-3237.1999, PMID: 10322026

Kim MJ, Whitesides GM. 1988. L-Lactate dehydrogenase: substrate specificity and use as a catalyst in the synthesis of homochiral 2-hydroxy acids. Journal of the American Chemical Society 110:2959–2964. DOI: https://doi.org/10.1021/ja00217a044

Kitagawa M, Ara T, Arifuzzaman M, Ioka-Nakamichi T, Inamoto E, Toyonaga H, Mori H. 2005. Complete set of ORF clones of Escherichia coli ASKA library (a complete set of E. coli K-12 ORF archive): unique resources for biological research. DNA Research 12:291–299. DOI: https://doi.org/10.1093/dnares/dsi012, PMID: 16769691

Kopka J, Schmidt S, Dethloff F, Pade N, Berendt S, Schottkowskii M, Martin N, Dühring U, Kuchmina E, Enke H, Kramer D, Wilde A, Hagemann M, Friedrich A. 2017. Systems analysis of ethanol production in the genetically engineered Cyanobacterium Synechococcus sp. PCC 7002. Biotechnology for Biofuels 10:56. DOI: https://doi.org/10.1186/s13068-017-0741-0, PMID: 28286551

Krömer JO, Heinzel E, Schröder H, Wittmann C. 2006. Accumulation of homolanthionine and activation of a novel pathway for isoleucine biosynthesis in Corynebacterium glutamicum McbR deletion strains. Journal of Bacteriology 188:609–618. DOI: https://doi.org/10.1128/JB.188.2.609-618.2006, PMID: 16385051

Lazcano A, Miller SL. 1999. On the origin of metabolic pathways. Journal of Molecular Evolution 49:424–431. DOI: https://doi.org/10.1007/s007240050538, PMID: 10486000

Lee DH, Palsson BD. 2010. Adaptive evolution of Escherichia coli K-12 MG1655 during growth on a nonnative carbon source, L-1,2-propanediol. Applied and Environmental Microbiology 76:4158–4168. DOI: https://doi.org/10.1128/AEM.00733-10, PMID: 20435762
Cotton et al. eLife 2020;9:e54207. DOI: https://doi.org/10.7554/eLife.54207

Liu F, Gu J, Wang X, Zhang XE, Deng J. 2014. Acsc is essential for propionate utilization in Escherichia coli. Biochemical and Biophysical Research Communications 449:272–277. DOI: https://doi.org/10.1016/j.bbrc.2014.05.015, PMID: 24835953

Livak KJ, Schmittgen TD. 2001. Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta delta C(T)) Method. Methods 25:402–408. DOI: https://doi.org/10.1016/S1046-323X(01)00036-8, PMID: 11846609

Maia LB, Moura I, Moura JGG. 2017. Molybdenum and tungsten-containing formate dehydrogenases: aiming to inspire a catalyst for carbon dioxide utilization. Inorganica Chimica Acta 455:350–363. DOI: https://doi.org/10.1016/j.ica.2016.07.010

Manukhov IV, Mamaeva DV, Rastorguev SM, Faleev NG, Morozova EA, Demidkina TV, Zavilgelsky GB. 2005. A gene encoding L-methionine gamma-lyase is present in Enterobacteriaceae family genomes: identification and characterization of Citrobacter freundii L-methionine gamma-lyase. Journal of Bacteriology 187:3889–3893. DOI: https://doi.org/10.1128/JB.187.3889-3893.2005, PMID: 15901718

McNeil NJ, Cummings JH, James WP. 1978. Short chain fatty acid absorption by the human large intestine. Gut 19:819–822. DOI: https://doi.org/10.1136/gut.19.9.819, PMID: 30683

Monticello DJ, Hadjioetroto RS, Costlow RW. 1984. Isoleucine synthesis by Clostridium sporogenes from propionate or alpha-methylbutyrate. Journal of General Microbiology 130:309–318. DOI: https://doi.org/10.1099/00221287-130-2-309, PMID: 6726176

Nagasawa T, Kanzaki H, Yamada H. 1984. Cystathionine gamma-lyase of Streptomyces phaeochromogenes the occurrence of cystathionine gamma-lyase in filamentous Bacteria and its purification and characterization. The Journal of Biological Chemistry 259:10393–10403. PMID: 6432781

Neidhardt FC, Curtiss IR, Ingraham JL, Low JK, Magasanik B, Riley M, Schaechter M, Umbarger HE. 1996. Escherichia coli and Salmonella. Cellular And Molecular Biology. ASM Press.

Noda-Garcia L, Liebermeister W, Tawfik DS. 2018. Metabolite-Enzyme coevolution: from single enzymes to metabolic pathways and networks. Annual Review of Biochemistry 87:187–216. DOI: https://doi.org/10.1146/annurev-biochem-062917-012023, PMID: 29925259

Notebaart RA, Sappanos B, Kintses B, Pál F, Győrkei Á, Bogos B, Lázár V, Spohn R, Csörgő B, Wagner A, Ruppin E, Pál C, Papp B. 2014. Network-level architecture and the evolutionary potential of underground metabolism. PNAS 111:11762–11767. DOI: https://doi.org/10.1073/pnas.1406102111, PMID: 25071190

Nyerges Á, Csörgő B, Nagy I, Bálint B, Bihari P, Lázár V, Apjok G, Umenhoffer K, Bogos B, Pósfai G, Pál C. 2016. A highly precise and portable genome engineering method allows comparison of mutational effects across bacterial species. PNAS 113:2502–2507. DOI: https://doi.org/10.1073/pnas.1520040113, PMID: 26884157

Oberhardt MA, Barektsi R, Reshef L, Xia F, Duran-Frigola M, Schreiber R, Henry CS, Ben-Tal N, Dwyer DJ, Gopchina U, Ruppin E. 2016. Systems-Wide prediction of enzyme promiscuity reveals a new underground alternative route for pyridoxal 5'-Phosphate production in E. coli. PLOS Computational Biology 12:e1004705. DOI: https://doi.org/10.1371/journal.pcbi.1004705, PMID: 26821166

Patrick WM, Quandt EM, Swartzlander DD, Matsumura I. 2007. Multicopy suppression underpins metabolic evolvability. Molecular and Biology Evolution 24:2716–2722. DOI: https://doi.org/10.1093/molbev/msm204, PMID: 17884825

Peter DM, Vögeli B, Cortina NS, Erb TJ. 2016. A Chemo-Enzymatic road map to the synthesis of CoA esters. Molecules 21:517. DOI: https://doi.org/10.3390/molecules21040517, PMID: 27104508

Risso C, Van Dien SJ, Orloff A, Lovley DR, Coppi MV. 2008. Elucidation of an alternate isoleucine biosynthesis pathway in Geobacter sulfurreducens. Journal of Bacteriology 190:2266–2274. DOI: https://doi.org/10.1128/JB.01841-07, PMID: 18245290

Sauer FD, Erffle JD, Mahadevan S. 1975. Amino acid biosynthesis in mixed rumen cultures. Biochemical Journal 150:357–372. DOI: https://doi.org/10.1042/bj1500357, PMID: 1212197

Sawers G, Hessling C, Muller N, Kaiser M. 1998. The glycol radical enzyme TdcE can replace pyruvate formate-lyase in glucose fermentation. Journal of Bacteriology 180:3509–3516. DOI: https://doi.org/10.1128/JB.180.14.3509-3516.1998, PMID: 9657990

Schink B, Kremer DR, Hansen TA. 1987. Pathway of propionate formation from ethanol in Pelobacter propionicus. Archives of Microbiology 147:321–327. DOI: https://doi.org/10.1007/BF00406127

Stams AJM, Kremer DR, Nicolay K, Weenk GH, Hansen TA. 1984. Pathway of propionate formation in Desulfobulbus propionicus. Archives of Microbiology 139:167–173. DOI: https://doi.org/10.1007/BF00401994

Steegborn C, Clausen T, Sondermann P, Jacob U, Werbs M, Marinovic S, Huber R, Wahl MC. 1999. Kinetics and inhibition of recombinant human cystathionine gamma-lyase toward the rational control of transsulfuration. The Journal of Biological Chemistry 274:12675–12684. DOI: https://doi.org/10.1074/jbc.274.18.12675, PMID: 10212249

Swick RW, Wood HG. 1960. The role of transcarboxylation in propionic acid fermentation. PNAS 46:28–41. DOI: https://doi.org/10.1073/pnas.46.1.28, PMID: 16590594

Tholozan J-L, Grivet J-P, Vallet C. 1994. Metabolic pathway to propionate of Pectinatus frisingensis, a strictly anaerobic beer-spoilage bacterium. Archives of Microbiology 162:401–408. DOI: https://doi.org/10.1007/BF00282104, PMID: 7893481

Unkriag V, Neugebauer FA, Knappe J. 1989. The free radical of pyruvate formate-lyase characterization by EPR spectroscopy and involvement in catalysis as studied with the substrate-analogue hypophosphite. European Journal of Biochemistry 184:723–728. DOI: https://doi.org/10.1111/j.1432-1033.1989.tb15072.x, PMID: 2553398.
Wang HH, Isaacs FJ, Carr PA, Sun ZZ, Xu G, Forest CR, Church GM. 2009. Programming cells by multiplex genome engineering and accelerated evolution. Nature 460:894–898. DOI: https://doi.org/10.1038/nature08187, PMID: 19633652

Wenk S, Yishai O, Lindner SN, Bar-Even A. 2018. An engineering approach for rewiring microbial metabolism. Methods in Enzymology 608:329–367. DOI: https://doi.org/10.1016/bs.mie.2018.04.026, PMID: 30173769

Wheldrake JF. 1967. Intracellular concentration of cysteine in Escherichia coli and its relation to repression of the sulphate-activating enzymes. Biochemical Journal 105:697–699. DOI: https://doi.org/10.1042/bj1050697, PMID: 4868875

Xu H, Zhang Y, Guo X, Ren S, Staempfli AA, Chiao J, Jiang W, Zhao G. 2004. Isoleucine biosynthesis in Leptospira interrogans serotype lai strain 56601 proceeds via a threonine-independent pathway. Journal of Bacteriology 186:5400–5409. DOI: https://doi.org/10.1128/JB.186.16.5400-5409.2004, PMID: 15292141

You L, Page L, Feng X, Berla B, Pakrasi HB, Tang YJ. 2012. Metabolic pathway confirmation and discovery through (13)C-labeling of proteinogenic amino acids. Journal of Visualized Experiments 26:e3583. DOI: https://doi.org/10.3791/3583

Zarzycki J, Sutter M, Cortina NS, Erb TJ, Kerfeld CA. 2017. In vitro characterization and concerted function of three core enzymes of a glycol radical enzyme - Associated bacterial microcompartment. Scientific Reports 7:42757. DOI: https://doi.org/10.1038/srep42757, PMID: 28202954

Zdych E, Peist R, Reidl J, Boos W. 1995. MalY of Escherichia coli is an enzyme with the activity of a beta C-S lyase (cystathionase). Journal of Bacteriology 177:5035–5039. DOI: https://doi.org/10.1128/JB.177.17.5035-5039.1995, PMID: 7665481

Zelcbuch L, Lindner SN, Zegman Y, Vainberg Slutskin I, Antonovsky N, Gleizer S, Milo R, Bar-Even A. 2016. Pyruvate Formate-Lyase enables efficient growth of Escherichia coli on Acetate and Formate. Biochemistry 55:2423–2426. DOI: https://doi.org/10.1021/acs.biochem.6b00184, PMID: 27093333

Zhou K, Zhou L, Lim O’, Zou R, Stephanopoulos G, Too HP. 2011. Novel reference genes for quantifying transcriptional responses of Escherichia coli to protein overexpression by quantitative PCR. BMC Molecular Biology 12:18. DOI: https://doi.org/10.1186/1471-2199-12-18, PMID: 21513543