Inactivation of *E. coli* pyruvate formate-lyase: Role of AdhE and small molecules

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Abstract

*Escherichia coli* AdhE has been reported to harbor three distinct enzymatic activities: alcohol dehydrogenase, acetaldehyde-CoA dehydrogenase, and pyruvate formate-lyase (PFL) deactivase. Herein we report on the cloning, expression, and purification of *E. coli* AdhE, and the re-investigation of its purported enzymatic activities. While both the alcohol dehydrogenase and acetaldehyde-CoA dehydrogenase activities were readily detectable, we were unable to obtain any evidence for catalytic deactivation of PFL by AdhE, regardless of whether the reported cofactors for deactivation (Fe(II), NAD, and CoA) were present. Our results demonstrate that AdhE is not a PFL deactivating enzyme. We have also examined the potential for deactivation of active PFL by small-molecule thiols. Both β-mercaptoethanol and dithiothreitol deactivate PFL efficiently, with the former providing quite rapid deactivation. PFL deactivated by these thiols can be reactivated, suggesting that this deactivation is non-destructive transfer of an H atom equivalent to quench the glycyl radical.

Keywords: Pyruvate formate-lyase; PFL; AdhE; Glycyl radical quenching; Deactivation of glycyl radical enzyme

Pyruvate formate-lyase (PFL) is one of the central enzymes in anaerobic metabolism in *Escherichia coli* (*E. coli*), and is responsible for catalyzing the conversion of pyruvate and CoA to formate and acetyl-CoA by a non-oxidative route (Scheme 1). Consistent with its central role in anaerobic metabolism, *pfl* transcription and PFL biosynthesis increase 10- to 12-fold when *E. coli* are grown under anaerobic conditions [1,2]. Induction of *pfl* gene expression under anaerobic conditions has been shown to involve both the FNR and the ArcA/ArcB regulatory proteins, which regulate gene expression in response to O₂ and various metabolites, respectively [3,4]. In addition to this transcriptional regulation, PFL requires post-translational activation by the pyruvate formate-lyase activating enzyme (PFL-AE) [5]. PFL-AE is constitutively expressed under both aerobic and anaerobic conditions [6], however *in vitro* studies have demonstrated that it is catalytically active only under anaerobic reducing conditions [7,8]. There are, therefore, at least two levels of control for regulating PFL activity: transcriptional up-regulation and post-translational activation. A potential third level of control involving PFL deactivation is addressed herein.

PFL-AE activates PFL by generation of a stable and catalytically essential radical at PFL G734 (Scheme 2) [9]. The activation reaction requires a cluster bound to PFL-AE [10], as well as S-adenosylmethionine (SAM or AdoMet) and a reductant, which *in vivo* is reduced flavodoxin [8]. These requirements place PFL-AE in the radical AdoMet superfamily [11]. The radical AdoMet enzymes use a reduced [4Fe-4S]⁺ cluster to reductively cleave AdoMet, producing methionine and a putative adenosyl radical intermediate; the adenosyl radical intermediate is believed to be responsible for H atom abstraction from substrate in all characterized members of the superfamily [12–15].
radical AdoMet superfamily has hundreds of putative members spanning the phylogenetic kingdom; functions include cofactor biosynthesis, rearrangement reactions, DNA repair, and enzyme activation. In addition to PFL-AE, *E. coli* harbors another radical AdoMet enzyme that is central to its anaerobic metabolism, the anaerobic ribonucleotide reductase (ARR) activating enzyme (AE). ARR plays a central role in nucleotide metabolism in anaerobic bacteria; like PFL, it requires post-translational activation to generate its catalytically essential glycyl radical [16, 17].

*Escherichia coli* thus utilizes post-translational glycyl radical generation as part of its regulatory mechanism for two of its most central metabolic functions: glucose and nucleotide metabolism. Once generated, the glycyl radicals in PFL [18] and ARR [19] are stable under anaerobic conditions; further, the radical is not consumed during catalysis but rather is a true cofactor. Thus once the respective activating enzymes activate PFL and ARR, the glycyl radicals are present and the enzymes are active indefinitely. The glycyl radicals in both PFL and ARR are, however, extremely sensitive to oxygen; oxygen reacts rapidly with the radical *in vitro*, resulting in protein cleavage at the site of the radical [9, 20]. This oxygen sensitivity raises the question of the fate of activated PFL and ARR when anaerobic *E. coli* are subjected to aerobic conditions, a situation that can occur routinely in facultative anaerobes such as *E. coli*.

The report of PFL deactivation catalyzed by AdhE seemed to resolve the puzzle of the fate of this catalytically essential radical: rather than undergoing oxygenolytic cleavage, as occurs *in vitro*, the glycyl radical was reported to be catalytically deactivated, or quenched, by AdhE [21, 22]. This intriguing deactivation reaction resulted in the conversion of active PFL back to its activatable form, thereby circumventing the irreversible protein cleavage observed *in vitro*, and presumably allowing the protein to be re-activated upon return to anaerobic conditions. The deactivation of PFL by AdhE was reported to be dependent on Fe(II), NAD, and CoA, with both pyruvate and NADH acting as competitive inhibitors of deactivation (Scheme 3). The effects of NAD and NADH led the authors to propose that the intracellular NAD/NADH ratio controls the state of PFL *in vivo*, resulting in activation under anaerobic conditions and deactivation as conditions become more aerobic [22]. Their proposal was further supported by *in vivo* studies showing that PFL is deactivated by AdhE under microaerophilic conditions [22].

AdhE, like PFL, is expressed under anaerobic conditions, as it plays an essential role in alcoholic fermentation. That AdhE should also function to deactivate PFL was a very intriguing proposal as it suggests that AdhE somehow acts as both a microaerophilic sensor and a deactivator of PFL, thus protecting it from oxygenolytic degradation. The proposed deactivation reaction, which involves the addition of a hydrogen atom equivalent to the glycyl radical of active PFL using the cofactors Fe(II), NAD, and CoA, is intriguing and unprecedented in biology. We therefore set out to investigate the PFL deactivative activity of AdhE in more detail, in order to gain some understanding of the catalytic mechanism. Described herein are our results showing that AdhE does not, in fact, catalyze the deactivation of PFL. We also provide data to show that some small-molecule thiols are quite efficient deactivators of PFL activity.

### Experimental methods

#### Materials

Unless otherwise stated, all chemicals were obtained from commercial sources and used without further purification. Deazariboflavin was synthesized according to literature procedures [23–25]. Cloning vectors (PET-21a+ and PET-Blue1) were obtained from Novagen. *E. coli* strains BL21(DE3)pLysS (Stratagen), Epicurian XL1 Blue (Novagen), NovaBlue (Novagen), and Tuner (DE3)pLacI (Novagen) were purchased from the indicated companies. Genomic *E. coli* (BL21(DE3)) DNA was purified using the Wizard genomic DNA extraction kit from Promega. Plasmid purification was performed using a purification kit from Stratagen. PCR primers were purchased from Integrated DNA Technologies, and PCRs were done using Pfu Turbo DNA polymerase (Stratagene). Immobilized metal ion affinity resin was purchased from Qiagen.

#### Cloning of His-tagged and wild type AdhE

For cloning of AdhE with a His6 affinity tag, PCR was performed using genomic *E. coli* BL21(DE3) DNA as the template with two primers (sense, 5’-CGCCGCGCATCCAGCTTGTTATCTATAGTC-3’ and antisense, 5’-CGCCGCGCATCCAGCTTGTTATCTATAGTC-3’) that incorporated appropriate restriction sites (*Bam*HI and *Xho*, respectively) for cloning into pET-21a+. The PCR product and vector were digested with these enzymes and then ligated to produce pMN1. Restriction digests verified the presence of a correctly sized insert, and the resulting vector was subjected to DNA sequencing (MSU Biotechnology Facility) to verify the presence of *adhE*. This plasmid is hereafter referred to as pET-21a+/*adhE*. For cloning of wild type *adhE*, PCR was performed using genomic *E. coli* DNA as the template with two primers (sense, 5’-GGCGCGCATCCAGCTTGTTATCTATAGTC-3’ and antisense, 5’-GGCGCGCATCCAGCTTGTTATCTATAGTC-3’).
TTTTTTGCTTGGGTTTTCTCAGCTTT-3’). The resulting PCR product was subjected to an end- conversion reaction prior to ligation into pET-Blue-1 blunt vector. Isolation of plasmid with the correct insert was confirmed by restriction digest analysis and DNA sequencing; this plasmid is hereafter referred to as pET-Blue-1/AdhE.

**Purification of AdhE-His6**

The plasmid PET-21a(+)/AdhE was transformed into BL21(DE3)pLysS host strain. Cultures were grown in LB/Amp media to an OD600 of 0.5–0.7, and then induced by addition of isopropyl β-D-thiogalactopyranoside (IPTG) to a final concentration of 1 mM. After an additional 2 h growth, the cells were harvested by centrifugation. BL21(DE3)pLysS/PET-21a(+)/AdhE cells were lysed in buffer (3 mM/L wet weight) containing 50 mM NaH2PO4, 10 mM imidazole, and 300 mM NaCl, pH 8, to which was added lysozyme (0.5 mg/mL), PMSF (1 mM), and trace quantities of DNAse and RNase. After two hours at 4 °C with gentle agitation, the suspension was centrifuged at 9000 g for 30 min. The supernatant (7 mL) was loaded onto an immobilized metal affinity chromatography (IMAC) column (30 mL, Qiagen) in 50 mM NaH2PO4, 10 mM imidazole, and 300 mM NaCl, pH 8. The column was washed with 50 mM NaH2PO4, 20 mM imidazole, and 300 mM NaCl, pH 8, and then the AdhE-His6 was eluted with 50 mM NaH2PO4, 250 mM imidazole, and 300 mM NaCl, pH 8. The purified protein was dialyzed into 20 mM Hepes, pH 7.2, and stored at −80 °C.

**Purification of wild-type AdhE**

The plasmid PET-Blue-1/AdhE was transformed into Tuner (DE3)pLacI cells. Using this vector/host combination, AdhE was found to be overexpressed constitutively, so growth and overexpression was carried out simply by growing a culture to its maximum optical density (OD600 ~ 2.5) prior to harvesting by centrifugation. BL21(DE3)pLysS/PET-Blue-1/AdhE cells were lysed at 4 °C for 2–3 h in lysis buffer (1.5 mL per gram of cells) containing 50 mM Tris-sulfate, pH 7.5, 200 mM NaCl, 1% Triton X-100, 5% glycerol, 10 mM MgCl2, 1 mM DTT, 1 mM PMSF, 0.25 mM lysozyme, and RNase and DNAse (~0.1 mg each per 10 mL), prior to centrifugation to remove cell debris. The crude extract (30–40 mL) was loaded onto a Superdex 75 gel filtration column (5 × 60 cm) equilibrated with 50 mM Tris, pH 7.5, and eluted with the same buffer. Fractions were analyzed using SDS–PAGE, and those containing >80% total protein as AdhE were pooled, concentrated to 30–40 mL, and dialyzed against 20 mM Hepes, pH 7.2, and stored at −80 °C.

**Activation of PFL**

PFL activation reactions were performed in an anaerobic chamber (Mbraun) maintained at <2 ppm O2. All reagents were thoroughly degassed prior to beginning the reaction. A typical PFL activation mix contained 150 mM Tris–Cl, pH 7.6, 0.1 M KCl, 10 mM oxamate, 8 mM DTT, 0.03–15 μM PFL-AE, 2–150 μM PFL, 0.2 mM AdoMet, and 50 μM 5-deazaribovlin, added in the order given. After mixing, activation was initiated by illuminating the sample with a 300 W halogen lamp. The activation sample was kept at approximately 5 cm from the lamp in a water bath to which ice was added periodically to keep the temperature at approximately 20–23 °C. Illumination was typically carried out for 30–45 min, until maximal PFL activity was obtained.

**PFL activity assay**

The PFL activity assay was carried out in an anaerobic chamber (Mbraun) maintained at <2 ppm O2 and in sealed quartz cuvettes. PFL activity was measured by using a modification of the coupled enzymatic assay previously described [7]. The coupling assay mix contained 150 mM Tris–Cl, pH 8.5, 3 mM NAD+, 55 μM CoA, 0.1 mg BSA, 10 mM pyruvate, 10 mM malate, 6 U citrate synthase, and 14 U malic dehydrogenase in a total volume of 700 μL. PFL activity was assayed by adding a small volume of the activated PFL (typically 1–5 μL) to the coupling assay mix, and monitoring the rate of production of NADH at 340 nm.

**AdhE deactivation activity assays**

The ability of AdhE to deactivate active PFL was investigated using a modification of previously reported assays; these assays were carried out in an anaerobic chamber and sealed cuvettes. To activated PFL (typically approximately 200 μM final concentration) was added AdhE (generally to 10–15 μM final concentration) NAD+ (100–200 μM final), CoA (50–100 μM final), and Fe(II) (0.5–1 mM final). Some assays also included 5 mM MgSO4 and 2 mM NAD. The mixture was incubated under anaerobic conditions (in the dark unless stated otherwise, in order to prevent competing re-activation by PFL-AE during deactivation experiments), and aliquots were removed periodically to assay for residual PFL activity. For some assays, one or more of the cofactors or AdhE was left out of the mixture, in order to determine the effect of each component individually.

**Other AdhE activity assays**

In order to assay alcohol dehydrogenase activity, an aliquot of AdhE was added to a solution containing 300 mM K2CO3 (pH 10), 0.25 mM NAD+, and 170 mM ethanol. The production of NADH was monitored at 340 nm. To assay acetaldehyde reductase activity, an aliquot of AdhE (generally 2–4 μM final concentration) was added to a solution containing 50 mM Tris–Cl, pH 7.6, 2.5 μM NADH, and 0.2–0.7 M acetaldehyde, and
NADH oxidation was monitored at 340 nm. To assay acetyl-CoA reductase activity, an aliquot of AdhE (1–10 μM) was added to a solution containing 50 mM CHES, pH 9.5, 0.2 mM dithiothreitol, 0.075 mM NAD, 0.1 mM CoA, and 10 mM acetaldehyde, and the absorbance at 340 nm was monitored. For each of the activities of AdhE, the specific activity is defined as 1 U = 1 μmol NADH/min.

Other PFL deactivation assays

The ability of a series of small molecules to inactivate active PFL was examined by adding the small molecules at a range of concentrations to active PFL in Tris–Cl buffer pH 8.5. The residual PFL activity was measured using the coupling assay as described previously.

Electron microscopy

Electron microscopy was performed using a Zeiss 100 CA microscope and a LEO 912 AB Zeiss microscope. AdhE (50 μg/mL) was in either 50 mM anaerobic MOPS/KOH, pH 7.5, or in the same buffer containing 3 mM DTT, 3 mM MgSO₄, and 0.3 mM Fe(NH₄)₂(SO₄)₂, or in the same buffer containing 3 mM DTT, 3 mM MgSO₄, and 0.3 mM Fe(NH₄)₂(SO₄)₂, 5 mM NAD, and 5 mM CoA. Samples were adsorbed on carbon film supported by a 200 mesh copper grid, and then stained by 1% uranyl acetate, pH 4.5.

PFL cleavage as a function of redox potential

BL21(DE3)pLys cells containing the PFL or AdhE expression vectors were grown in 50 mL of Luria Broth media at 37 °C overnight. A 5 mL aliquot of each of the overnight cultures was transferred into 850 mL of LB media or a de

Overexpression and purification

Both AdhE and AdhE-His₆ were overexpressed efficiently using the constructs and host cells described in Experimental methods. The AdhE-His₆ overexpressed at a higher level and was easier to purify due to the presence of the His-tag. Typical yields were 25 mg purified AdhE-His₆ per liter of culture. The untagged AdhE was partially purified using gel filtration and ion exchange chromatography and was used primarily to confirm results obtained with the AdhE-His₆. Yields of purified wt-AdhE were typically on the order of 9 mg per liter of culture. PFL overexpressed efficiently and could be purified in reasonably large quantities (generally 120–160 mg purified PFL per L of culture).

Alcohol dehydrogenase activities

AdhE catalyzes the generation of ethanol from acetyl-CoA by two sequential NADH-dependent reactions, as shown in Scheme 4. Each of the reactions shown in Scheme 4, the acetyl-CoA reductase and the acetaldehyde reductase reactions, can be assayed independently in either the forward or reverse direction by applying the appropriate substrates. Both activities have been identified in our purified AdhE, both the wild type and the hexahistidine-tagged versions, with specific activities similar to those reported in the literature. For example, we have found the specific activity of AdhE for the ethanol dehydrogenase reaction to be 9.5 U/mg (for both wild type and histidine-tagged enzyme), as compared to values of 10 U/mg reported previously for wild-type AdhE [21]. All other specific activities were also

![Scheme 4](image-url)
similar to those reported in the literature, thereby verifying that we have isolated AdhE in its fully functional form.

**Deactivation of PFL by AdhE**

AdhE has been reported to harbor PFL deactivation as a third enzymatic activity, with a requirement for NAD, Fe\(^{2+}\), and CoA for full PFL deactivation activity. We have re-investigated this report through a series of detailed experiments as outlined below. In general, our approach was to fully activate PFL using PFL activating enzyme and the procedures described in the Experimental methods. Then, under strictly anaerobic conditions (due to the extreme air sensitivity of the glycyl radical present in active PFL), we incubated PFL alone or with AdhE and/or a range of possible cofactors. Residual PFL activity was monitored as a function of time in order to probe the rate of deactivation, if any, resulting from the added components.

Initial experiments involved incubating PFL with AdhE alone, with the putative cofactors NAD, Fe\(^{2+}\), and CoA, or with both AdhE and the putative cofactors. The results of representative experiments are shown in Fig. 1. Over the time course of the experiment, the activity of the PFL incubated with AdhE alone increased slightly and then remained essentially constant, suggesting that AdhE in the absence of any added cofactors is unable to deactivate PFL. PFL incubated alone under identical conditions exhibited identical behavior (data not shown). It is noteworthy that the PFL activity is stable over 8 h in this experiment, as previous reports estimated AdhE in the absence of any added cofactors is unable to deactivate PFL. PFL incubated alone under identical conditions exhibited identical behavior (data not shown).

Fig. 1. Lack of deactivation of PFL by AdhE. Activated PFL (193 \(\mu\)M) was incubated with AdhE (3 \(\mu\)M), with AdhE and its reported “PFL deactivase” cofactors NAD\(^+\) (0.2 mM), CoA (0.1 mM), and Fe(II) (1 mM), with the cofactors alone, or with no additives; experiments were performed in the dark. The results clearly demonstrate that AdhE has no effect on the deactivation of PFL, although some of the cofactors appear to display a modest effect. (*) PFL + AdhE; (■) PFL + AdhE, NAD, CoA, and Fe(II); (▲) PFL + NAD, CoA, and Fe(II).

Kessler and Knappe previously showed that in *E. coli* K12 cells, protection of PFL against oxygenolytic cleavage was dependent on both the presence of AdhE and the redox potential of the culture, such that deactivation was turned on only under microaerophilic conditions (+100 mV). Therefore we examined the PFL deactivation activity of our overexpressed AdhE as a function of culture and lysate redox potential. Briefly, anaerobic cultures expressing AdhE and PFL separately were adjusted to a range of redox potentials and then lysed together prior to exposure to oxygen to cleave any remaining active PFL. Alternatively, anaerobic cultures expressing AdhE and PFL separately were lysed alone or in combination prior to adjusting the redox potential of the lysate; the lysate was then exposed to oxygen to cleave any remaining active PFL. In all experiments, complete cleavage of PFL is observed, as evidenced by the equal intensity bands corresponding to the uncleaved PFL polypeptide (85 kDa) and the large fragment of the oxygenolytically cleaved polypeptide (82 kDa), as seen in Fig. 2. The cleavage of one-half of all PFL polypeptides is considered complete PFL cleavage due to the half-of-sites reactivity of PFL. It should be noted that even if our cloned AdhE was somehow defective in PFL deactivation, the wt AdhE of the host cells would be induced to fairly high levels (3 \(\times\) 10^6 copies/cell) [28] under the anaerobic growth conditions used in these experiments. Thus the redox shifting should have resulted in PFL deactivation (even in the absence of our overexpressed AdhE). No evidence for inactivation of PFL by wt AdhE is observed, however, providing further support for our conclusion that AdhE is not a PFL deactivating enzyme.

**Oligomerization properties of overexpressed AdhE**

Kessler and Knappe were the first to observe the unusual oligomerization properties of AdhE, which forms rod-like
oligomeric particles that change conformation, as observed by electron microscopy, upon addition of Fe$^{2+}$, CoA, and NAD. In order to further address the similarities between our overexpressed AdhE and the wt protein characterized by Kessler and Knappe, we performed electron microscopy on our purified AdhE in the absence and presence of cofactors (data not shown). Similar to the observations of Kessler and Knappe, we see rod-like particles (average length 56 nm and average width of 13 nm) for the protein in MOPS buffer with or without Fe$^{2+}$. In the presence of NAD and CoA, further aggregation occurs, and the rods change shape (average length 100 nm and average width 13 nm).

Inactivation of PFL by small molecule reductants

An observation made consistently in these deactivation experiments was that reactions that included either Fe$^{2+}$ or CoA exhibited faster rates of loss of PFL activity than the control reactions, regardless of other components present in the mix (data not shown). These observations suggested that reducing agents such as thiols or reduced metals might be able to slowly quench the glycyl radical of PFL. To investigate this further, we have examined the inactivation of active PFL by small-molecule thiols and other reducing agents. Both β-mercaptoethanol (β-ME) and dithiothreitol (DTT) were observed to inactivate PFL, with the former doing so at a much faster rate (Fig. 3). β-ME inactivation is quite rapid, with complete loss of activity observed in just over one hour under the conditions of this experiment. DTT inactivation is almost imperceptible over the same time period, but is complete within 7 h under these conditions. Loss of PFL activity is also accompanied by a loss of the EPR signal due to the PFL glycyl radical (data not shown), demonstrating that these thiols inactivate PFL by destroying the glycyl radical found in active PFL. That these two thiols, commonly used in biological buffers, should inactivate PFL is curious, since both thiols have been used in the past in various stages of PFL purifications.
and activations reported in the literature. In fact, it has been observed in our laboratory that PFL must be treated with DTT prior to activation in order to achieve maximal activity (unpublished observation). However since DTT inactivation is so slow, it presumably would not have interfered with previous experiments in which PFL activation was carried out for one hour or less.

Other thiols, including ethanethiol and cysteine, were also observed to inactivate PFL (Fig. 4). Ethanethiol, like βME, inactivates PFL rapidly, while cysteine caused much slower inactivation. In contrast, larger thiols (such as homocysteine and glutathione) or non-thiolic compounds (such as methionine) showed no detectable inactivation. The results suggest that thiols must access the active site glycyl radical directly in order to efficiently promote inactivation, since the smallest thiols provide the most efficient inactivation. Non-thiolic reductants, such as dithionite and ascorbic acid, also inactivate PFL, with dithionite showing a rate similar to that of DTT and ascorbate being significantly slower (data not shown).

Reversibility of inactivation

Observation of a loss of PFL activity over time could be due to deactivation, i.e., conversion of the glycyl radical to a non-radical glycyl residue without damaging the protein, or to inactivation, in which the glycyl radical is quenched in a destructive manner. The former process is reversible and would allow re-generation of the glycyl radical, while the latter process involves irreversible destruction of the protein. The latter process, inactivation, is commonly observed when working with active PFL. For example, exposure of active PFL to even trace quantities of oxygen in vitro results in oxygenolytic cleavage at the site of the glycyl radical. The former process, deactivation, was previously ascribed only to AdhE, and was proposed to be the means by which PFL is conserved in vitro when E. coli cycle between anaerobic and aerobic conditions. However in our hands AdhE does not bring about a loss of PFL activity. Since small molecule thiols and other reductants do bring about a loss of PFL activity, we proceeded to investigate whether the loss of activity mediated by thiols was reversible, i.e., whether these thiols were mediating deactivation.

Active PFL was incubated with β-ME as previously described, and essentially complete loss of activity was observed, while essentially no loss of activity was observed for the control reaction containing no β-ME (Fig. 5). Additional PFL activating enzyme, 5-deazariboflavin, and AdoMet were added to both reactions at this point, and both reactions were exposed to a 300 W halogen lamp for 30 min. As can be seen in Fig. 5, this treatment had essentially no effect on the control (−βME) reaction, other than a slight (~5–10%) increase in PFL activity; this is consistent with the fact that the PFL used in the experiment had been essentially completely activated prior to beginning the experiment. The addition of PFL-AE, 5-deazariboflavin, and AdoMet and exposure to light did, however, have a dramatic effect on the +βME reaction, in which the PFL activity returned to that of the control. Removal of the light source again leads to loss of PFL activity in the +βME reaction, while the activity of the control again remains constant. That full PFL activity was restored to the β-ME-treated PFL clearly demonstrates that the β-ME-mediated deactivation is a reversible process that does not destroy G734 or the PFL protein structure. That is, β-ME inactivates PFL by the simple addition of a hydrogen atom equivalent to the glycyl radical. Similar experiments performed with DTT, ethanethiol, and other thiols showed similar results, in which PFL inactivation by these thiols was completely reversible (data not shown).

Fig. 4. Inactivation of active PFL by thiols. Activated PFL (370 μM) was incubated with the indicated reagents (4 mM) under anaerobic conditions in the dark, and residual PFL activity was monitored after 4 h.
The nature of PFL inactivation by βME was further investigated by examining the effect of dioxygen on βME-inactivated PFL. Oxygenolytic cleavage of activated PFL can be readily observed by SDS-PAGE (Fig. 6). Unactivated PFL migrates as an 85 kDa band on SDS-PAGE. If activated PFL is fully inactivated by exposure to oxygen, the resulting PFL migrates as two observed bands of equal intensity, one at 85 kDa and one at 82 kDa (Fig. 6, lane 1). The latter band is the larger fragment resulting from oxygenolytic cleavage at G734; the smaller 3 kDa fragment is not observed. Observation of both 85 and 82 kDa fragments at equal intensities is consistent with the previously reported half-of-sites reactivity for the PFL dimer; one glycyl radical per dimer results in oxygenolytic cleavage of exactly half of the PFL molecules. When activated PFL is deactivated with βME and then exposed to oxygen, no oxygenolytic cleavage is observed (Fig. 6, lane 2), consistent with the quenching of the glycyl radical by βME.

**PFL activation using ambient light**

We observed during these experiments that the rate of inactivation could be perceptibly slowed if the reactions were not protected from ambient light; in fact, if PFL had not been fully activated prior to deactivation experiments, we could observe activation of PFL, particularly certain controls, during the deactivation experiments (Fig. 7). We surmised that this slowing of deactivation, or further activation, of PFL was due to a competition between deactivation and re-activation (by the trace PFL-AE, SAM, and the photoreductant 5-deazariboflavin, which was not removed from PFL prior to these inactivation experiments). In order for this to be true, the relatively dim ambient lighting in the anaerobic chamber would have to be sufficient to promote the 5-deazariboflavin-mediated photoreduction of PFL-AE. This possibility was independently tested by incubating inactive PFL with PFL-AE, AdoMet, and 5-deazariboflavin in the ambient lighting in the anaerobic chamber. Although activation was slower than that observed using the intense 300 W halogen lamp as described in Experimental methods, full activation could be achieved with ambient lighting (data not shown). Activation using ambient lighting was found to be preferable in many instances for these experiments, as it avoided the temperature-control problems associated with intense illumination. In addition, the PFL activity appeared to be more reproducible and more stable when activation was done with ambient lighting.

**Discussion**

Our laboratory has a long-standing interest in the mechanism of radical generation by which PFL is activated,
therefore we were naturally intrigued by the report of an enzyme that catalytically deactivates PFL. Although several other glycyl radical enzymes and their corresponding activating enzymes are known, no potential deactivases (other than AdhE) have been identified. Despite this, the possibility of an enzymatic means by which to deactivate PFL (or other anaerobic glycyl radical enzymes such as the anaerobic ribonucleotide reductase) during cycling between anaerobic and aerobic conditions is compelling, as oxygenolytic cleavage upon transition to aerobic conditions followed by de novo protein synthesis and activation would be energetically costly and inefficient.

The results presented herein, however, demonstrate that AdhE and AdhE-His\textsubscript{6} purified from overexpressing cultures do not catalyze the deactivation of active PFL, despite previous reports to the contrary for the wt enzyme [21,22]. Repeated deactivation experiments in the presence of the purported cofactors Fe(II), CoA, and NAD\textsuperscript{+} show that AdhE has no effect on the rate of deactivation of PFL, although both Fe(II) and CoA appear to increase the rate of deactivation. Small-molecule thiols were found to have an even more dramatic effect on the rate of loss of PFL activity, with βME causing essentially complete inactivation within one hour. Thiol-dependent inactivation of PFL was shown to be a non-destructive process that allowed regeneration of the active, glycyl radical-containing PFL by incubation with the activation components.

In contrast to the results of Kessler and Knapp, redox shifting of overexpressing cultures or lysates did not have an effect on the amount of activated PFL, as evidenced by SDS-PAGE gels showing fully cleaved PFL in all cases, regardless of the presence of overexpressed AdhE or the potential of the culture or lysate. It should be noted that in the cultures overexpressing PFL, anaerobic incubation of the culture allows the wt PFL-AE present in the host cells to fully activate the overexpressed PFL, such that subsequent aerobic lysis of those cells results in full PFL cleavage. Likewise, it is expected that the wt AdhE present in the host cells would be able to deactivate some of the overexpressed PFL during redox-potential shifting of the PFL cultures, as previously reported by Knappe and Kessler; however, no evidence for deactivation is observed by SDS-PAGE (Fig. 2). The overexpressed AdhE, when combined with the PFL at different potentials, also provides no hint of PFL deactivation.

We are unable at this time to resolve the discrepancies between our results and the AdhE-catalyzed deactivation reported by Kessler and Knappe [21,22]. Our data suggests that we have isolated fully functional AdhE; this is evidenced by the presence of the other two AdhE activities (alcohol dehydrogenase and acetaldehyde-CoA dehydrogenase) and by our ability to reproduce the cofactor-mediated change in oligomeric state of AdhE. Yet in hundreds of experiments in our laboratory, no evidence for AdhE-catalyzed PFL deactivation can be found. Our results are, however, consistent with a recent report on YfID, a PFL homolog in E. coli that can be activated to generate a glycyl radical by PFL-AE, but cannot be deactivated by AdhE [29].

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