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Metabolic and proteomic analyses of product selectivity and redox regulation in *Clostridium pasteurianum* grown on glycerol under varied iron availability

Christin Groeger[†], Wei Wang[†], Wael Sabra, Tyll Utesch and An-Ping Zeng^{*}

Abstract

Background: Clostridium pasteurianum as an emerging new microbial cell factory can produce both *n*-butanol (BuOH) and 1,3-propanediol (1,3-PDO), and the pattern of product formation changes significantly with the composition of the culture medium. Among others iron content in the medium was shown to strongly affect the products selectivity. However, the mechanism behind this metabolic regulation is still unclear. For a better understanding of such metabolic regulation and for process optimization, we carried out fermentation experiments under either iron excess or iron limitation conditions, and performed metabolic, stoichiometric and proteomic analyses.

Results: 1,3-PDO is most effectively produced under iron limited condition (Fe—), whereas 1,3-PDO and BuOH were both produced under iron rich condition (Fe+). With increased iron availability the BuOH/1,3-PDO ratio increased significantly from 0.27 mol/mol (at Fe—) to 1.4 mol/mol (at Fe+). Additionally, hydrogen production was enhanced significantly under Fe+ condition. Proteomic analysis revealed differentiated expression of many proteins including several ones of the central carbon metabolic pathway. Among others, pyruvate: ferredoxin oxidoreductase, hydrogenases, and several electron transfer flavoproteins was found to be strongly up-regulated under Fe+ condition, pointing to their strong involvement in the regeneration of the oxidized form of ferredoxin, and consequently their influences on the product selectivity in *C. pasteurianum*. Of particular significance is the finding that H₂ formation in *C. pasteurianum* is coupled to the ferredoxin-dependent butyryl-CoA dehydrogenase catalyzed reaction, which significantly affects the redox balance and thus the product selectivity.

Conclusions: The metabolic, stoichiometric and proteomic results clearly show the key roles of hydrogenases and ferredoxins dependent reactions in determining the internal redox balance and hence product selectivity. Not only the NADH pool but also the regulation of the ferredoxin pool could explain such product variation under different iron conditions.

Keywords: n-Butanol, 1,3-Propanediol, C. pasteurianum, Proteomics, Product selectivity, Metabolic analysis

Background

Clostridium pasteurianum is an emerging and promising microbial cell factory for the production of chemicals and fuels because of some unique features, e.g. utilization

of a wide range of substrates [1–3], production of a wide spectrum of products [4, 5] and robust growth in simple media even under unsterile conditions [6]. Recently, *C. pasteurianum* was shown to accept electrons from the cathode by direct electron transfer [7], which make it an attractive candidate for new bioelectrical systems. Therefore, *C. pasteurianum* has received considerable interests for the production of chemicals and fuels such as 1,3-propanediol (1,3-PDO) and *n*-butanol (BuOH),

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which represents attractive bioprocesses for the use of renewable resources, like biodiesel-derived glycerol or glucose from biomass hydrolysates [8-14]. In such bioprocesses, several other fermentation products like gases (carbon dioxide and hydrogen), ethanol as well as acetic, butyric and lactic acid are produced [2, 4, 15], in addition to 1,3-PDO and BuOH. Even though the formation of organic acids is inevitable for the maintenance of the intracellular redox balance, it represents a loss of carbons at the expense of the target products. Moreover, the resulting product distribution, especially the selectivity of either 1,3-PDO or BuOH, is mainly influenced by the cultivation conditions and/or media supplements. For instances, several studies analyzed the effect of pH [16], inoculum conditions [17], supplementations of yeast extracts and ammonia [18], or acetic and butyric acid [2, 19], or phosphate and iron [18, 20]. Among others, iron seems to have extensive effects, since its absence lead to a strongly reduced BuOH formation [16, 20]. In real fermentation processes, especially under conditions more relevant to industrial applications, with raw substrates and high concentrations of products, the product selectivity and yield often strongly fluctuate and are hardly reproducible. The underlying mechanism(s) of selectivity and regulation of intracellular metabolic pathways are still unclear, even though a combined effect of many iron-related enzymes has been assumed [4]. Indeed, several iron containing enzymes are involved in clostridia metabolism, e.g. nitrogenases, ferredoxin coupled enzymes, and alcohol dehydrogenases. These enzymes play key roles in the maintenance of intracellular redox balance and a limited functionality of them, e.g. due to iron limitation, will be reflected by a metabolic shift and thus change of product selectivity.

In this work, the variations of product selectivity and the underlying mechanisms of pathway regulation in *C. pasteurianum* DSMZ 525 grown on glycerol under varied iron availability are studied with an integrated systems biology approach, particularly with stoichiometric, kinetic and proteomic analyses.

Methods

Microorganism, medium and cultivation

Clostridium pasteurianum DSMZ 525 was routinely maintained as cryoculture at -80 °C in Reinforced Clostridial Medium (RCM, Oxoid Deutschland GmbH) containing 20% (v/v) glycerin. The cryoculture was used for the pre-culture carried out in anaerobic bottles with RCM medium at 35 °C and pH 7 without shaking. The RCM contained 1 mg/L resazurin (7-hydroxy-10-oxidophenoxazin-10-ium-3-one) as a redox indicator for anaerobiosis and 2 g/L CaCO₃ as pH-buffering agent. After 24 h this pre-culture was used as inoculum for

bioreactor fermentation. The bioreactor medium contained the following ingredients in 1 L of distilled water (modified from [4]): glycerol, 80 g; yeast extract, 1 g; K₂HPO₄, 0.5 g; KH₂PO₄, 0.5 g; MgSO₄·7H₂0, 0.2 g; (NH₄)₂SO₄, 5 g; CaCl₂ 2H₂O, 0.02 g; cysteine-HCl, 0.5 g; resarzurin, 0.005 g; trace element solution SL-7 (DSMZ), 2 mL. Iron concentrations were varied in the bioreactor medium as follows: Iron excess (Fe+) condition means the addition of 10 mg/L FeSO₄·7H₂O (2 mg Fe²⁺/L) into the medium and iron limitation (Fe-) condition means no iron addition. Iron originally present in the preculture (0.07 mg Fe²⁺/L) and those present in the yeast extract (up to $0.05 \text{ mg Fe}^{2+}/\text{L}$) were the sole iron sources in the Fe- cultivations. Cultivations were run at 35 °C, pH 6 and 500 rpm agitation in a stirred tank bioreactor (Bioengineering) with a working volume of 1.2 L. During the fermentation pH was maintained at 6 with 5 M KOH. To achieve anaerobic condition prior to the inoculation, the autoclaved medium was sparged with sterile O2-free N₂. The experiments were performed in duplicates. Total volume of the effluent fermentation gas was determined with a Milli-Gascounter (Dr.-Ing. Ritter Apparatebau GmbH & Co. KG), and its composition was measured with the mass spectrometer OmniStar 300 (Balzer Instruments/Pfeiffer Vacuum GmbH). The MS took samples in an interval of 0.5 mL/min for the concentration analysis of H₂, CO₂, O₂, N₂ and Ar.

Analytical methods and calculations

The optical density of cell suspension was measured turbidometrically at 600 nm and correlated with cell dry weight: biomass BM (g/L) = $OD_{600} \times 0.336$. The specific growth rate μ (h⁻¹) was determined from biomass data (smoothed using the software Origin 8.5.1 G SR1, OriginLab Corporation, Northampton, USA) according to Eq. 1, where x_1 and x_2 are the concentrations of biomass (g/L) at the times t_1 and t_2 , respectively. The substrate and product titers in the supernatant were analyzed via HPLC equipped with a refractive index detector and an ultraviolet detector. HPLC was performed on an Aminex HPX-87H column (300 \times 7.8 mm) at 60 °C, with 0.005 M H₂SO₄ as mobile phase at a flow rate of 0.6 mL/min.

For the measurement of 3-HPA, the method described by Oehmke and Zeng [21] was used, in which 3-HPA is converted into acrolein and the concentration of acrolein is determined spectrometrically by external calibration. Briefly, 100 μL of cell free culture supernatant were mixed with 200 μL of HCl (37%) and 50 μL of tryptophan solution in a cooled 96 well plate. The tryptophan solution consisted of 10 mM DL-tryptophan, 0.05 M HCl and 24 mM toluene. After 40 min incubation at 37 °C, the absorbance of the mixtures were determined with a

Multiskan[®] Spectrum plate reader (Thermo Fisher Scientific) at 560 nm.

The yield coefficient (Y) for either product or substrate (i) was calculated according to Eq. 2. Based on the stoichiometric equations for glycerol utilization in *C. pasteurianum* [11], carbon and redox recovery were calculated according to Eqs. 3 and 4, respectively. Here C [—] is the number of carbon atoms in the products and substrate, c is the concentration of products in (mmol/L) and biomass (BM) in (g/L).

$$\mu = \frac{\ln x_2 - \ln x_1}{t_2 - t_1} \tag{1}$$

$$Y_{i/X} = \frac{i_2 - i_1}{x_2 - x_1} \tag{2}$$

$$C_{\text{recovery}} [\%] = \frac{\sum C_{\text{products}}}{\sum C_{\text{substrate}}}$$
 (3)

$$NADH_{recovery} [\%] = \frac{c_{1,3-PDO}}{2c_{acetate} + 2c_{butyrate} + c_{lactate} + 13.2c_{BM}}$$

$$(4)$$

Comparative proteomic analysis

Samples for proteomics were taken in the exponential growth phase and stationary phase during parallel fermentations of *C. pasteurianum* DSMZ 525 under iron excess and iron limited conditions. The detailed methodical procedure for comparative proteomic analysis was previously described by Sabra et al. [11].

Results and discussion

Effects of iron availability on the growth and product formation of *C. pasteurianum*

Different concentrations of iron have been reported for the optimization of butanol or 1,3-PDO formation using C. pasteurianum [2, 18, 20]. Using a fractional factorial experimental design, Moon et al. used 60 mg/L FeSO₄·7H₂O for optimum butanol formation in C. pasteurianum in serum anaerobic bottle experiments, while no iron sulphate was supplemented for a better 1,3-PDO production [18]. In controlled bioreactor, we have found that 10 mg/L FeSO₄·7H₂O is enough to support a similar butanol productivities (0.9 g/L×h) and an almost doubled butanol concentration (21 g/L butanol) by the same strain [2]. Therefore, in the current investigation, 0 and 10 mg/L FeSO₄·7H₂O were chosen, respectively, to describe the growth and product formation under iron limited and iron excess conditions in our glycerol fermentation. The same pre-culture was used to inoculate two bioreactors containing the growth medium supplemented either with or without 10 mg/L FeSO₄·7H₂O (hereinafter termed as iron excess (Fe+) condition or iron limitation (Fe-) condition, respectively). One of the main differences observed was the relatively shorter lag phase under Fecondition, which was accompanied by an early growth cessation (Fig. 1a). With excess iron in the medium, a higher biomass production with a maximum concentration of 5.1 \pm 0.09 g/L and μ_{max} of 0.31 \pm 0.01 h^{-1} were reached, whereas under iron limitation condition only a biomass concentration of 3.2 \pm 0.01 g/L and a μ_{max} of $0.23 \pm 0.01 \; h^{-1}$ could be achieved (Table 1). The cessation of growth under iron limitation was obviously not due to butanol toxicity, as the highest titer of BuOH reached did not exceed 3.7 g/L (Table 1), which was lower than the toxic concentration level of BuOH for C. pasteurianum (>5 g/L [11]). Depletion of the intracellular iron pool and/or the accumulation of 3-hydroxypropionaldehyde (3-HPA), a very toxic intermediate in the formation of 1,3-PDO [22, 23], may cause the relatively earlier growth cessation under Fe- condition. As shown in Fig. 1b, under Fe+ condition the 3-HPA concentration did not exceed 8 mg/L, whereas under Fe- condition up to 30 mg/L 3-HPA were produced. In this time range of relatively high concentrations of 3-HPA a growth cessation was observed. Indeed, it has been reported that the growth of vegetative cells of C. tyrobutyricum was completely inhibited at 38 mg/L externally added 3-HPA [24].

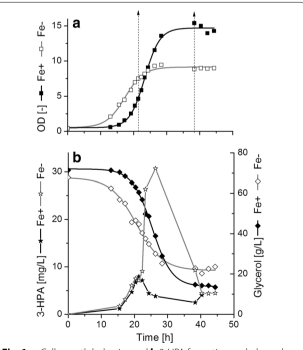


Fig. 1 a Cell growth behavior and **b** 3-HPA formation and glycerol consumption in cultivation of *C. pasteurianum* DSMZ 525 under iron excess (Fe+) and iron limitation (Fe-) conditions. *Arrows* indicate time points of sampling for proteomic analysis

Table 1 Product formation during the growth of C. pasteurianum DSMZ 525 under iron excess (Fe+) and iron limitation (Fe-) conditions. Fermentations were performed in duplicates

| | Biomass | | 1,3-PDO | | Вион | | Етон | | Acetate | | Butyrate | | Lactate | | Formate | |
|---|-------------------------------------|-------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|--------------------------------|-----------------|-----------------|-----------------|---|-----------------------|---|-----------------|
| | Fe+ | Fe- | Fe+ | Fe- | Fe+ | Fe- | Fe+ | Fe- | Fe+ | Fe- | Fe+ | Fe- | Fe+ | Fe- | Fe+ | Fe- |
| Final titer (g/L |) 5.1 ± 0.09 | 3.2 ± 0.01 | 9.4 ± 1.44 | 16.6 ± 2.26 | 12.3 ± 0.06 | 4.4 ± 0.70 | 1.0 ± 0.12 | 0.9 ± 0.60 | 0.8 ± 0.01 | 1.5 ± 0.02 | 1.7 ± 0.45 | 2.8 ± 0.13 | inal titer (g/L) 5.1 ± 0.09 3.2 ± 0.01 9.4 ± 1.44 16.6 ± 2.26 12.3 ± 0.06 4.4 ± 0.70 1.0 ± 0.12 0.9 ± 0.60 0.8 ± 0.01 1.5 ± 0.02 1.7 ± 0.45 2.8 ± 0.13 0.1 ± 0.00 | 7.5 ± 2.58 0.5 ± 0.07 | 0.5 ± 0.07 | 1.1 ± 0.11 |
| Production rate (g/L*h) at exponential growth phase | | 0.39±0.00 0.16±0.03 0.66±0.17 | 0.66 ± 0.17 | 0.74 ± 0.02 | 0.53 ± 0.11 | 0.16 ± 0.04 | 0.06 ± 0.02 | 0.03 ± 0.003 | 0.04±0.004 0.07±0.03 0.14±0.02 | 0.07 ± 0.03 | 0.14 ± 0.02 | 0.13 ± 0.01 | 0.007 ± 0.003 | 0.28 ± 0.02 | 0.03 ± 0.01 | 0.04 ± 0.02 |
| | Y S/X | | Y P/X | | | | | | | | | | | | | |
| Y 1/X (9/9 BM) | Y _{1/X} (g/g BM) 9.6 ± 0.3 | 13.9 土 1.2 | 1.16 ± 0.09 | 4.40 ± 0.52 | 1.88 ± 0.25 | 0.97 ± 0.13 | 0.15 ± 0.06 | 0.17 ± 0.13 | 0.10 ± 0.01 | 0.39 ± 0.09 | 0.39 ± 0.08 | 0.80 ± 0.17 | 139±1.2 1.16±0.09 440±0.52 1.88±0.25 0.97±0.13 0.15±0.06 0.17±0.13 0.10±0.01 0.39±0.09 0.39±0.08 0.80±0.17 0.02±0.00 | 1.64 ± 0.37 | 1.64 ± 0.37 0.08 ± 0.02 0.25 ± 0.05 | 0.25 ± 0.05 |

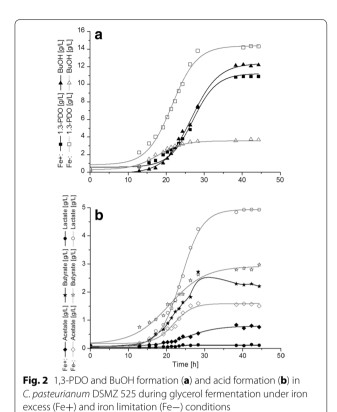


Figure 2 shows the formation of fermentation products in C. pasteurianum DSMZ 525 under iron excess and iron limitation conditions. With higher iron concentration 12 g/L of BuOH and 11 g/L of 1,3-PDO were produced. In comparison, at iron limitation significantly less BuOH, i.e. 3.7 g/L, was formed, accompanied with the formation of 14.5 g/L of 1,3-PDO. In fact, the molar ratio of BuOH to 1,3-PDO decreased from 1.34 mol/mol under Fe+ condition to 0.27 mol/mol under Fe- condition. The yield of 1,3-PDO per biomass increased nearly 4 times from 1.16 \pm 0.09 g/g at Fe+ to 4.4 \pm 0.52 g/g at Fe-, whereas the specific yield of BuOH was halved from 1.88 ± 0.25 g/g at Fe+ to 0.97 ± 0.13 g/g at Fe-. Next to this, the acid formation changed remarkably, especially the lactate production, which was shown to increase significantly in the Fe- culture (Table 1). The specific lactate yield increased significantly from 0.02 \pm 0.0 g/g in the Fe+ culture to 1.6 \pm 0.4 g/g in the Fe- culture. Also acetate and butyrate yield increased under Fe- condition, but to less extent than that of lactate (Table 1). The reason(s) for these dramatic changes of metabolism are not clear yet, but of fundamental importance for the development of C. pasteurianum as an emerging microbial cell factory for the production of chemicals and fuels. Therefore in the following redox regulation and comparative proteomic analysis were performed.

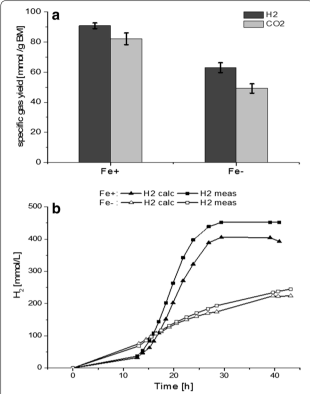


Fig. 3 a Measured cumulative hydrogen and carbon dioxide production and **b** calculated (calc) vs. measured (meas) hydrogen production under iron excess (Fe+) and iron limitation (Fe-) conditions in *C. pasteurianum* DSMZ 525 cultures

Redox regulation and H₂ production in *C. pasteurianum* DSMZ 525

For the growth and metabolism of *C. pasteurianum*, particularly when growing on a more reduced substrate like glycerol, the maintenance of intracellular redox balance is crucial. The shift of metabolism under conditions of iron excess and limitation shown above is postulated to be strongly related to the redox regulation which is addressed below first from a stoichiometric point of view.

To check the stoichiometry and consistency of the fermentation data, fermentation balance analysis was first done. A very good consistency in carbon recovery was observed for the fermentations. The carbon recovered as fermentation products represented approximately 98% of the carbon source consumed. On the other hand, the calculated recovery of the reducing equivalents according to Eq. 4 reached 91% at Fe+ and 94% at Fe-, indicating a lower consistency in reducing equivalent recovery according to the assumed pathways of redox regulation. *C. pasteurianum* contains ferredoxin-dependent hydrogenases, which catalyze the re-oxidation of reduced ferredoxin with the formation of H₂. Reduced ferredoxins are generally formed in the enzymatic step of forming

acetyl-CoA from pyruvate catalyzed by pyruvate: ferredoxin oxidoreductase (PFOR). Hence, under the assumption that the formation of one mole acetyl-CoA from pyruvate is accompanied with the formation of one mole H_2 , the theoretical H_2 production would be calculated according to Eq. 5, where q is the formation rate of each compound (mmol/g×h):

$$q_{H2} = q_{ethanol} + q_{acetate} + 2q_{butvrate} + 2q_{butanol}$$
 (5)

In repeated fermentations to those shown in Fig. 1 under similar conditions we measured the evolution of CO2 and H2 in effluent gas. The results are given in Fig. 3a. Under Fe+ conditions a cumulative amount of 452 mmol/L H₂ and 399 mmol/L CO₂ were produced, compared to 245 mmol/L H₂ and 177 mmol/L CO₂ produced under Fe- condition. Referred to the biomass formed, H2 and CO2 production increased significantly from 63 (± 3.3) and 49 (± 3.1) mmol/g _{biomass} at Fecondition to 91 (±1.9) and 82 (±3.9) mmol/g $_{biomass}$ at Fe+ condition, respectively. Interestingly, the theoretically calculated H₂ production values were lower than the measured ones (Fig. 3b), particularly under Fe+ conditions. Obviously, the re-oxidation of reduced ferredoxin generated in the enzymatic step catalyzed by PFOR was not the only source of hydrogen formation. Similar behavior was also noticed previously in cultures of C. butyricum or Klebsiella pneumoniae [25, 26].

It is known that butyryl-CoA is generally formed from crotonyl-CoA by the NADH dependent trans-2-enoyl-CoA reductase (Ter) enzyme (Eq. 6) [27].

Crotonyl-CoA + NADH
$$\stackrel{\text{Ter}}{\rightarrow}$$
 NAD + Butyryl-CoA (6)

But recently, Buckel and Thauer [28] proposed a new indirect route of $\rm H_2$ formation in *C. pasteurianum* from NADH and ferredoxin in two steps, catalyzed successively by the ferredoxin-dependent butyryl-CoA dehydrogenase/electron transferring flavoprotein complex (BCdH-ETF) (Eq. 7) and a hydrogenase (Eq. 8).

$$Fd_{ox} + 2NADH + Crotonyl-CoA$$

$$\xrightarrow{BCdH-ETF} Fd_{red} + 2NAD + Butyryl-CoA$$
(7)

$$Fd_{red} + 2H^{+} \xrightarrow{H_2 - ase} Fd_{ox} + H_2$$
 (8)

Since the measured $\rm H_2$ production values were significantly higher than the theoretically calculated ones based on Eqs. 7 and 8 (see Fig. 3b), it is reasonable to assume that in *C. pasteurianum* DSMZ 525, BCdH-ETF together with Ter is actively involved in the step of converting crotonyl-CoA to butyryl-CoA, giving rise to an additional source of $\rm H_2$ formation. Thus, with this new suggested

butyryl-CoA formation route one mole NADH $_2$ is additionally required for the formation of one mol butanol or one mol butyrate, accompanied with the formation of one mole more H $_2$, in addition to the H $_2$ formation counted in Eq. 5. Consequently, the calculation of reducing equivalent recovery should be modified as follows (Eq. 9), by also taking into account the difference of calculated and measured H $_2$ values ($c_{\Delta H2}$), representing the additionally consumed NADH $_2$:

$$NADH_{recovery} (\%) = \frac{c_{1,3-PDO} + c_{\Delta H2}}{2c_{acetat} + 2c_{butyrat} + c_{lactate} + 13.2c_{BTM}}$$
(9)

Using Eq. 9, a more satisfying reducing equivalent recovery of 105% under Fe+ condition and 100% under Fe- condition was obtained, giving a strong support for the involvement of the BCdH-ETF complex. Particularly, the results from Fe+ condition are in agreement with the corrected Eq. 9, where more BuOH and hydrogen were produced, and the deviation between the calculated and measured H₂ was higher. However, this is in contrary to what was reported for *C. acetobutylicum*. For a more effective butanol production, a lower hydrogenase activity and hydrogen production was favored in *C. acetobutylicum* [29]. To shed more light on the mechanisms underlying the effect of iron on the regulation of glycerol metabolism in *C. pasteurianum*, comparative proteomic studies were performed as described below.

Comparative proteomic analysis of the iron effect

For proteomic analysis of the effects of iron concentration on the metabolism of *C. pasteurianum*, samples were taken from the two bioreactors in the exponential growth phase (termed as Fe+ early and Fe- early, respectively) and the stationary growth phase (Fe+ late and Fe- late, respectively) (Fig. 1). Each sample was analyzed in triplicates. After 2-DE separation of the intracellular proteins, protein spots showing statistically significant changes between Fe+ early and Fe- early, Fe+ late and Fe- late, Fe+ early and Fe+ late, as well as Fe- early and Fe- late were further identified by LC-MS/MS. Proteins which were identified as single protein present in a spot on the 2-D gels are summarized in Table 2 according to their functional categories and accession numbers, together with the information of their expression changes. The existence of more than one values of fold change for a single protein indicates that this protein appeared as multiple spots on the 2-D gels.

The pyruvate acetyl-CoA node: a focal point in the metabolism of *C. pasteurianum*

The conversion of pyruvate to acetyl-CoA linking glycolysis to TCA cycle is a fundamental metabolic step

Table 2 Proteins showing significant changed expression levels, compared between iron excess (Fe+) and iron limitation (Fe-) conditions, as well as between the exponential growth phase (early) and the stationary phase (late)

| | | | | | | ייסיים | Fold change Iron-related | | Fold Cn | ange grov | Fold change growth-related | þ |
|----------------------------------|--|-----------------------|-------------------|-----|--------------------------|--------|--------------------------|------------|---------------|-----------|----------------------------|---------|
| | | ogous groups (COG) | protein domain | | Early phase higher at | ase | Late phase higher at | iase at | Fe+ higher at | her at | Fe– higher at | yher at |
| | | | | | Fe | Fe+ | Fe- | Fe+ | Early | Late | Early | Late |
| Amino acid tran | Amino acid transport and metabolism | | | | | | | | | | | |
| F502_05097 | Amino peptidase 1 | COG1362 | Lap4 | 298 | 1.9 | | 1.7 | | | | | |
| F502_05412 | Carbamoyl phosphate synthase large subunit | COG0458 | CarB | 68 | | 1.8 | | | 1.9 | | | |
| F502_07028 | Cysteine synthase a | COG0031 | CysK | 449 | 1.8 | | | | | 2.1 | | |
| F502_17572 | Glutamine synthetase type III | COG3968 | GInA3 | 187 | 1.6 | | | | | 1.9 | | |
| | | | | 190 | 1.6 | | | | | 1.6 | | |
| F502_18676 | Threonine synthase | COG0498 | ThrC | 312 | | 1.6 | | | 2.2 | | | |
| Carbohydrate tr | Carbohydrate transport and metabolism | | | | | | | | | | | |
| F502_03412 | Propanediol dehydratase small | COG4910 | PduE | 594 | 1.9 | | 1.9 | | | | | |
| F502_07638 | Subunit flavodoxin | COG0716 | FIdA | 594 | | | | | | | | |
| F502_03417 | Glycerol dehydratase reactivation factor large subunit | No COG | | 220 | | | | | 2.1 | | | |
| F502_03937 | Glycogen synthase | COG0297 | GlgA | 329 | 2.7 | | | | | 2.3 | | |
| F502_06067 | Enolase | COG0148 | Eno | 695 | 4.1 | | 4.0 | | | 2.0 | | 2.0 |
| | | | | 969 | 4.8 | | 1.8 | | | 2.5 | | |
| F502_06077 | Triosephosphate isomerase | COG0149 | TpiA | 491 | | | | 1.5 | | 2.6 | | |
| F502_06087 | Glyceraldehyde 3-phosphate dehydrogenase | COG0057 | GapA | 409 | | 1.6 | | | 2.4 | | | |
| F502_07098 | Glycoside hydrolase | COG1543 | | 277 | 2.0 | | | | | 1.8 | | |
| F502_12758 | Dihydroxyacetone kinase | COG2376 | DAK1 | 158 | | 1.7 | | 2.0 | 1.7 | | 2.1 | |
| | | | | 179 | | 1.7 | | 1.7 | 2.2 | | 2.3 | |
| Cell cycle control/cell division | ol/cell division | | | | | | | | | | | |
| F502_08238 | Cell division protein | COG3599 | DivIVA | 503 | | | | | 2.7 | | 2.0 | |
| Cell wall/memb | Cell wall/membrane/envelope biogenesis | | | | | | | | | | | |
| F502_00655 | Peptidoglycan-binding protein | COG1388 | LysM | 233 | 7.9 | | | | | 6.1 | 1.6 | |
| F502_01965 | Spore coat protein Frelated protein | COG5577 | CotF | 549 | 1.5 | | 2.1 | | | | | 1.6 |
| Coenzyme trans | Coenzyme transport and metabolism | | | | | | | | | | | |
| F502_07578 | Pyridoxal biosynthesis lyase | COG0214 | PdxS | 460 | | 2.5 | | | 2.2 | | | |
| Energy product | Energy production and conversion | | | | | | | | | | | |
| F502_05017 | NifU-related domain containing protein | COG0822 | IscU | 519 | 9.0 | 2.4 | 9.0 | 1.6 | 1.3 | 0.8 | 6.0 | 1.1 |
| | | | | 520 | | 3.7 | | 2.9 | 1.9 | | 1.5 | |
| F507 06787 | Flectron transfer flavonrotein subunit alpha | COG2025 | FixB | 440 | | | | 1.5 | | | | |

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| | Accession no. | Protein name | Cluster of orthol- | Conserved | Spot no. | Fold change iron-related | nge iron- | related | | Fold ch | ange gro | Fold change growth-related | D. |
|---|---------------|--|-----------------------|-------------------|----------|--------------------------|-----------|-------------------|------------|---------|----------|----------------------------|---------|
| Prince transfer flow opportern subunit alpha COCCOOR Fixe Activation COCCOOR Fixe Activation Activation | | | ogous groups (COG) | protein domain | | Early pha higher at | se | Late pl higher | nase at | Fe+ hig | lher at | Fe – hiç | jher at |
| Electron rander flavoprotein subunit alpha / crosobes Fish 487 17 17 17 17 17 17 18 23 33 47 18 13 47 18 17 18 23 | | | | | | | -e+ | Fe- | Fe+ | Early | Late | Early | Late |
| Bifunctional acealdehyde-CoAdischold dehyde-CoAdischold dehyde-CoA | F502_06287 | Electron transfer flavoprotein subunit alpha/ beta-like protein | COG2086 | FixA | 487 | | | | 1.7 | | | | |
| drogenise 715 15 29 53 Nitroreductase homodiment CCGO778 Nirkle 377 24 24 15 47 Pytwater femedoann (flavodaxin) oxidoneductise CCG0674 PonA 70 24 24 15 47 Pytwater femedoann (flavodaxin) oxidoneductise CCG0674 PonA 40 24 24 15 18 Pytwater femedoann oxidoneductise CCG0674 PonA 40 22 23 23 15 15 Pytwater femedoann oxidoneductase CCG1048 PonA 107 24 24 15 16 17 15 16 17 15 16 16 17 17 16 17 16 17 16 17 16 17 17 16 17 | F502_06447 | Bifunctional acetaldehyde-CoA/alcohol dehy- | COG1012 | AdhE | 119 | | | | 2.1 | | 3.7 | | 1.6 |
| Pytuvate ferredoxin flavodoxin) oxidored uccess COCG078 NnhB 537 24 24 47 Pytuvate ferredoxin flavodoxin) oxidored uccess COCG074 PorA 70 24 15 18 Fase, homodiment for the proper trace homodiment for the proper trace homodiment for the proper trace homodiment for the properties or COCG074 PorA 76 24 16 18 Pytuvate ferredoxin foxidoreductase COCG074 PorA 40 22 23 15 15 Butyate first for the properties for the properties oxidoreductase (COCG1048 COCG1048 NorV 131 17 24 21 20 Butyate first first carboxin to be a cocg1048 COCG1048 NorV 156 24 27 17 20 Port ARP synthase subunit be a cocg1048 Arpb 645 21 24 27 17 20 Roll ARP synthase subunit be a cocg1048 Arpb 645 21 24 27 17 17 Roll Agenyate oxidate oxidate by a cocg104 COCG1048 NorD 528 24 24 | | drogenase | | | 715 | 1.5 | | | 2.9 | | 5.3 | | |
| Ningedeenderasee COG0778 Ninfigedeenderasee S37 23 29 1.5 2.4 1.5 2.4 1.5 2.4 1.5 2.4 1.5 2.4 1.5 2.4 1.5 1.5 2.4 1.5 <t< td=""><td></td><td></td><td></td><td></td><td>717</td><td>1.6</td><td></td><td></td><td>3.0</td><td></td><td>4.7</td><td></td><td></td></t<> | | | | | 717 | 1.6 | | | 3.0 | | 4.7 | | |
| Pytruater ferrendoxin (Navodoxint) oxidoreduct COG0674 PorA 70 24 15 15 15 15 19 18 | F502_07493 | Nitroreductase | COG0778 | NfnB | 537 | 2.3 | | 2.9 | | | | | 1.8 |
| tase homodimetric | F502_07643 | Pyruvate: ferredoxin (flavodoxin) oxidoreduc- | COG0674 | PorA | 70 | | 2.4 | | 2.4 | 1.5 | | | |
| Pyruvater ferredoxin oxidoreductase COG0674 PorA 40 27 15< | | tase. homodimeric | | | 75 | | | | 2.4 | 9.0 | 8.1 | | |
| Pyruvate ferredoxin oxidoreductase CGG05/4 PorA PorA PorA PorA PorA PorA PorA PorA | | | | | 9/ | | | | 2.7 | | 1.9 | | |
| Provide Ferredoxin oxidoreductase CXG0674 PorA PorA A | | | | | 77 | | 1.5 | | 2.3 | | | | |
| Pytywate: ferredoxin oxidoreductase COGO674 PorA 40 22 28 1.6 20 Pytywate: ferredoxin oxidoreductase COGO674 PorA 40 23 1.5< | | | | | 87 | | 2.0 | | 2.8 | | | 1.5 | |
| Pynvater ferredoxin oxidoreductase COG0674 PorA 40 2.3 1.6 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.5 1.7 | | | | | 06 | | 2.2 | | 2.8 | 1.6 | | 2.0 | |
| Spanning of the down full and down | F502_07648 | Pyruvate: ferredoxin oxidoreductase | COG0674 | PorA | 40 | | | | 2.3 | | 1.6 | | |
| Rubredoxin/flavodoxin/oxidoreductase COG0426 NovV 131 1.7 2.4 2.4 2.1 2.9 2.1 2.9 2.1 2.9 2.0 | | | | | 57 | | | | 2.3 | | 2.1 | | |
| Rubredoxin/flavodoxin/oxidoreductase COG0426 NorV 131 1.7 2.8 2.0 Hydratase (aconitase A) COG1048 AcnA 156 24 2.5 1.9 2.0 Butyrate kinase COG1038 Butk 383 1.6 2.1 2.5 1.7 2.0 Pyruvate carboxylase COG1038 PycA 7.1 1.5 2.7 1.7 1.7 FOFI ATP synthase subunit beta COG0038 ApD 645 2.1 1.9 1.7 1.7 Havodoxin COG0716 FIdA 575 14.3 8.0 2.8 1.5 1.7 Hell hydrogenase COG0716 FIdA 575 1.43 8.0 2.5 1.6 1.6 Rubrerythrin COG0716 NorD 558 2.1 2.1 2.5 1.8 2.2 Hydrogenase-1 COG1034 NuoG 2.2 2.1 4.5 2.2 2.2 2.1 2.2 2.2 2.2 2.2 <td< td=""><td></td><td></td><td></td><td></td><td>58</td><td></td><td>1.5</td><td></td><td>2.4</td><td></td><td>2.1</td><td></td><td></td></td<> | | | | | 58 | | 1.5 | | 2.4 | | 2.1 | | |
| Rubredoxin/flavodoxin/oxidoreductase COG0426 NorV 131 1.7 2.0 2.0 Hydratase (aconitase A) COG1048 AcnA 156 2,4 2.5 1.9 2.0 Butyrate kinase COG326 Buk 383 1.6 2.7 1.7 1.7 Pyruvate carboxylase COG1038 PycA 71 1.5 2.7 1.7 1.7 FOF1 ATP synthase subunit beta COG1038 AtpD 645 2.1 1.9 2.7 1.7 1.7 FOF1 ATP synthase subunit beta COG2033 SorL 611 2.2 2.4 2.2 2.4 1.7 1.7 Pavodoxin COG0716 FidA 575 14.3 8.0 2.8 1.6 1.7 1.6 1.6 1.6 1.6 | | | | | 59 | | 2.1 | | 2.8 | | | | |
| Hydratase (aconitase A) COG1048 AcnA 156 24 25 1.9 2.0 Butyrate kinase COG3426 Buk 383 1.6 27 1.9 1.7 2.0 Pyruvate carboxylase COG1038 PycA 71 1.5 2.7 1.7 1.7 1.7 FOF1 ATP synthase subunit beta COG0055 AtpD 645 2.1 2.4 2.7 1.7 1.7 1.7 Desulfo ferrodoxin COG2033 SorL 611 2.2 2.4 2.8 1.5 1.7 1.7 Havodoxin Flel hydrogenase COG4624 Narl PurB 303 2.1 2.1 2.8 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.7 1.7 1.6 1.7 1.7 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.7 1.6 1.6 1.6 1.6 | F502_09238 | Rubredoxin/flavodoxin/oxidoreductase | COG0426 | NorV | 131 | 1.7 | | | | | 2.0 | | |
| Hydratase (aconitase A) COG1048 AcnA 156 2.4 2.5 1.9 2.0 2.0 Butyrate kinase COG3256 Buk 383 1.6 2.7 1.7 1.7 Pyruvate carboxylase COG1038 ApD 645 2.1 1.9 1.7 1.7 F0F1 AFP synthase subunit beta COG2033 SorL 611 2.2 2.4 1.7 1.7 Desulfo ferrodoxin COG2033 SorL 611 2.2 2.4 2.8 1.6 1.7 1.7 1.7 Havodoxin COG4624 Narl PurB 373 1.43 2.0 2.3 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.6 1.7 1.6 1.6 1.6 1.6 1.6 1.7 1.6 1.7 1.6 1.6 1.7 1.6 1.6 1.7 1.6 1.6 1.7 1.6 1.6 1.7 1.6 1.6 1.6 | | | | | 730 | | | | | | 2.0 | | 1.6 |
| Butynate kinase COG3426 Buk 383 1.6 1.7 1.8 | F502_09488 | Hydratase (aconitase A) | COG1048 | AcnA | 156 | | 2.4 | | 2.5 | 1.9 | | 2.0 | |
| Pytruvate carboxylase COG1038 PycA 71 1.5 2.7 1.7 F0F1 ATP synthase subunit beta COG2035 AtpD 645 2.1 1.9 7 1.7 Desulfo ferrodoxin COG2033 Sort 611 2.2 2.4 2.8 1.6 Flavodoxin COG4624 Nar1 PurlB 303 2.1 8.0 1.6 1.6 Adenylosuccinate lyase COG015 YotD 558 2.1 4.5 2.5 1.8 2.3 Alydrogenase-1 COG1034 NuoG 223 4.5 1.8 2.3 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 2.3 Abb-dependent glyceralde hyde-3-phosphate COG1013 AdhE 7.6 7.5 1.9 3.6 2.3 Abb-dependent glyceralde hyde-3-phosphate COG1013 7.3 7.5 7.5 | F502_11871 | Butyrate kinase | COG3426 | Buk | 383 | 1.6 | | | | | 1.7 | | |
| FOF1 ATP synthase subunit beta COG0055 AtpD 645 2.1 1.9 Attraction | F502_11976 | Pyruvate carboxylase | COG1038 | PycA | 71 | | 1.5 | | 2.7 | | 1.7 | | |
| Desulfo ferrodoxin COG2033 SorL 611 22 2.4 2.8 1.6 Flavodoxin COG0716 FldA 575 14.3 8.0 1.6 1.6 Fel hydrogenase COG4624 Narl PurB 303 2.1 2.1 2.5 1.6 Adenylosuccinate lyase COG0015 YotD 558 2.2 4.5 2.0 2.0 Rubrerythrin COG1592 YotD 558 2.2 4.5 2.3 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.8 2.3 Adhe-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 4.5 1.6 2.3 Adhe-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 2.3 2.1 | F502_12091 | F0F1 ATP synthase subunit beta | COG0055 | AtpD | 645 | 2.1 | | 1.9 | | | | | |
| Desulfo ferrodoxin COG2033 Sort 611 22 2.8 1.6 Flavodoxin COG4624 Nar1 PurB 303 2.1 7 1.6 Fel hydrogenase COG4624 Nar1 PurB 303 2.1 2.5 1.6 Adenylosuccinate Jyase COG1592 YotD 558 2.2 4.5 2.0 Rubrerythrin COG1034 NuoG 315 1.8 2.3 2.3 Hydrogenase-I COG1034 NuoG 223 4.5 1.6 2.3 NADP-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 3.6 | | | | | 646 | 2.2 | | 2.4 | | | | | |
| Flavodoxin COG0716 FIdA 575 14.3 8.0 1.6 Fel hydrogenase COG4624 Nar1 PurB 303 2.1 2.5 1.6 Adenylosuccinate lyase COG01592 YotD 558 2.2 4.5 2.0 Bubrerythrin COG134 NuoG 315 1.8 2.3 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 Adhe-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 3.6 | F502_12878 | Desulfo ferrodoxin | COG2033 | SorL | 611 | | 2.2 | | 2.8 | | | | |
| [Fe] hydrogenase COG4624 Nar1 PurB 303 2.1 2.5 Adenylosuccinate lyase COG0015 YotD 558 4.5 4.5 2.0 Rubrerythrin COG1592 YotD 558 2.2 4.5 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 Hydrogenase-1 COG1012 AdhE 736 2.7 1.5 3.6 NADP-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 2.1 | F502_13493 | Flavodoxin | COG0716 | FIdA | 575 | 14.3 | | 8.0 | | | 1.6 | | |
| Adenylosuccinate Jyase COG0015 YotD 558 2.2 4.5 2.0 Rubrerythrin COG1592 YotD 315 1.8 2.3 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 Hydrogenase-1 COG1012 AdhE 736 2.7 1.5 3.6 Adhedographate COG1012 AdhE 736 2.7 1.5 2.1 | F502_14390 | [Fe] hydrogenase | COG4624 | Nar1 PurB | 303 | | 2.1 | | | 2.5 | | | |
| Rubrerythrin COG1592 YotD 558 2.2 4.5 2.0 Glycolate oxidase COG1034 GlcD 315 1.8 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 NADP-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 2.1 | F502_04707 | Adenylosuccinate Iyase | COG0015 | | 303 | | | | | | | | |
| Glycolate oxidase COG0277 GlcD 315 1.8 2.3 Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 NADP-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 2.1 | F502_15080 | Rubrerythrin | COG1592 | YotD | 558 | | 2.2 | | 4.5 | | 2.0 | | |
| Hydrogenase-1 COG1034 NuoG 223 4.5 1.6 2.3 NADP-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 | F502_16610 | Glycolate oxidase | COG0277 | GlcD | 315 | | | | 1.8 | | 2.3 | | 1.9 |
| NADP-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 | F502_18287 | Hydrogenase-1 | COG1034 | NuoG | 223 | | 4.5 | | 1.6 | 2.3 | | | |
| NADP-dependent glyceraldehyde-3-phosphate COG1012 AdhE 736 2.7 1.5 | | | | | 224 | -, | 5.1 | | 1.9 | 3.6 | | | |
| | F502_18651 | NADP-dependent glyceraldehyde-3-phosphate | | AdhE | 736 | 2.7 | | 1.5 | | | 2.1 | | |

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| Figure 2015 Figure 3 Figur | Accession no. | Protein name | Cluster of orthol- | Conserved | Spot no. | Fold change iron-related | le iron-r | elated | | Fold change growth-related | ge grow | rth-related | |
|--|------------------|---|-----------------------|-------------------|----------|--------------------------|-----------|---------------------|-----|----------------------------|---------|-------------|-------|
| GOG1882 PRID 174 45 Fe Fe+ Fe+< | | | ogous groups (COG) | protein domain | | Early phase higher at | d) | Late ph higher a | ase | Fe+ highe | ır at | Fe- high | er at |
| COG1882 PRD 174 4.5 5.7 3.4 175 80 6.5 3.7 3.4 181 3.7 1.7 7.4 181 3.7 1.7 7.4 No COG 400 2.1 2.2 3.0 No COG 535 2.0 2.1 2.2 3.0 No COG 537 1.7 3.4 3.8 1.7 2.0 No COG 537 1.7 2.8 1.5 2.0 1.5 2.0 No COG 600 637 2.3 2.0 1.5 2.1 1.9 1.5 2.0 COG1039 You 6.0 2.3 2.1 1.5 2.1 1.9 1.6 2.1 1.9 1.6 2.1 1.9 1.6 2.1 1.9 1.6 2.1 1.6 2.1 1.6 2.1 1.9 1.6 2.1 1.6 2.1 1.6 2.1 1.6 2.1 1.6 | | | | | | | <u>+</u> | Fe- | Fe+ | | ate. | Early | Late |
| 175 180 65 175 181 | F502_19556 | Formate acetyltransferase | COG1882 | PfID | 174 | 4.5 | | 5.7 | | | | | |
| 178 2.5 2.6 2.5 2.6 | | | | | 175 | 8.0 | | 6.5 | | | | | |
| COGI453 32 26 27 27 28 29 20 20 20 20 20 20 20 | | | | | 178 | 2.5 | | | | | | 3.4 | |
| COC61453 362 26 29 30 15 30 15 30 15 30 15 15 30 15 15 30 15 15 15 15 15 20 10 15 20 20 15 15 15 15 20 | | | | | 181 | 3.7 | | | | 1.7 | | 7.4 | |
| COG1453 362 26 29 30 15 < | Function unkno | wn/general function prediction only | | | | | | | | | | | |
| No COG No COG Ano Ano | F502_02435 | Aldo/keto reductase | COG1453 | | 362 | 2.6 | .0 | | | 2.9 | | | |
| No COG A | F502_05012 | Hypothetical protein (GGGtGRT protein) | No COG | | 400 | 7. | _ | | 2.2 | 3.0 | | | |
| COGGO07 318 34 38 1,7 20 No COG 597 1,7 2,8 1,5 2,6 1,5 2,0 1,5 2,0 1,5 2,0 1,5 2,0 1,5 </td <td>F502_05962</td> <td>Hypothetical protein</td> <td>No COG</td> <td></td> <td>635</td> <td>2.0</td> <td></td> <td></td> <td></td> <td>-</td> <td>.5</td> <td></td> <td></td> | F502_05962 | Hypothetical protein | No COG | | 635 | 2.0 | | | | - | .5 | | |
| No COG 197 1,7 28 1,5 </td <td>F502_06682</td> <td>Hypothetical protein</td> <td>COG2607</td> <td></td> <td>318</td> <td>3.4</td> <td>₹+</td> <td></td> <td>3.8</td> <td>1.7</td> <td></td> <td>2.0</td> <td></td> | F502_06682 | Hypothetical protein | COG2607 | | 318 | 3.4 | ₹+ | | 3.8 | 1.7 | | 2.0 | |
| COG1024 CaiD 472 2.1 2.2 2. | F502_15420 | Hypothetical protein | No COG | | 597 | 1.7 | | 2.8 | | 1.5 | | | |
| COG1024 CaiD 472 2.1 1.6 2.2 1.9 COG0439 AccC 302 2.3 1.6 2.1 1.9 GOG043 PurH 281 1.6 1.7 1.6 1.8 1.8 COG043 Dank 217 1.6 2.3 1.6 2.1 1.8 1.8 COG043 Dank 107 3.6 1.6 2.1 1.8 2.1 1.8 1.8 2.1 1.8 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.4 2.1 2.2 2.4 2.1 2.4 2.1 2.2 2.4 2.1 2.2 2.4 2.1 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 2.2 | F502_16320 | Hypothetical protein | COG0393 | YbjQ | 637 | 2.3 | | 2.0 | | | | | |
| COG1024 CaiD 472 2.1 1.6 2.2 2.1 1.9 COG0439 Acc 302 2.5 1.6 2.1 1.9 1.9 GOG138 PurH 281 1.6 1.2 1.2 1.6 1.6 COG043 Dank 217 1.6 2.3 1.6 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.9 1.8 1.9 1.8 1.9 1.9 1.8 1.9 1.9 1.8 1.9 | Lipid transport | and metabolism | | | | | | | | | | | |
| COG0439 AccC 302 2.5 1.6 2.1 1.9 COG0138 PurH 281 1.7 1.6 1.9 COG0543 GrpE 481 1.6 2.3 1.6 1.9 COG0542 CrpC Crp | F502_06297 | 3-Hydroxybutyryl-CoA dehydratase | COG1024 | CaiD | 472 | 2. | _ | | | 2.2 | | | |
| GroE Gribe 481 2.1 2.2 1.8 1.8 1.8 1.8 1.8 1.8 1.8 1.8 2.1 2.2 1.8 1.8 1.8 1.8 2.1 2.1 2.1 1.8 2.1 1.8 2.1 1.8 2.1 2.2 2.1 2.2< | F502_10483 | Biotin carboxylase | COG0439 | AccC | 302 | 2.5 | | | 1.6 | 2 | 1. | 1.9 | |
| COG0138 Purh 281 1.7 1.5 | Nucleotide tran: | sport and metabolism | | | | | | | | | | | |
| GrpE GrpE 481 2.1 2.1 2.2 COG0443 Dank 217 1.6 2.3 1.8 1.8 COG1026 Cym1 106 2.3 1.6 2.1 COG0542 ClpA 165 1.8 2.1 2.4 COG0459 GroEL 258 1.6 1.7 2.0 COG0693 Thi 553 1.8 1.6 2.5 2.6 COG0693 Thi 553 1.8 1.6 2.5 2.4 COG0693 Thi 553 1.8 1.6 2.5 2.4 COG0693 Thi 1.6 2.5 2.5 2.4 COG0693 Thi 5.3 2.4 2.4 COG0693 Thi 2.5 2.4 2.4 COG0693 Thi 2.6 2.4 2.4 COG0693 Thi 2.6 2.4 2.4 COG0694 Thi 2.6 2.4 | F502_17300 | Bifunctional phosphoribosylaminoimida- zolecarboxamide formyltransferase/IMP cyclohydrolase | COG0138 | PurH | 281 | | | | | 1.7 | | 1.6 | |
| Heat shock protein (molecular chaperone GrpE) COG043 GrpE 481 1.6 2.1 2.2 Molecular chaperone DnaK COG1026 Cym1 106 2.3 1.6 1.8 1.8 1.1 1.2 1.8 1.9 1.2 1.2 1.2 1.3 2.1 1.2 | Posttranslationa | Il modification/Protein turnover/Chaperones | | | | | | | | | | | |
| Molecular chaperone DnaK COG0443 Dank 217 1.6 2.3 1.6 2.1 ATP-dependent Clp protease ATP-binding COG1026 ClpA 165 1.8 1.16 2.1 ATP-dependent Clp protease ATP-binding COG0459 CroR 1.6 1.9 1.7 2.4 Chaperonin Cochaperonin COG0234 GroEL 2.8 1.6 1.5 2.0 Thill/Pfol family protein COG0326 Thil 533 1.8 1.5 2.6 Heat shock protein (molecular chaperone lbpA) COG0326 HtyG 106 2.2 2.5 2.5 Heat shock protein 90 COG0326 HtyG 106 2.2 2.5 2.4 Clbb protein COG0326 HtyG 106 2.2 2.5 2.4 Clbb protein COG0326 HtyG 106 2.2 2.5 2.4 Clbb protein COG0326 HtyG 1.6 2.5 2.5 2.4 Clbb protein COG0326 <t< td=""><td>F502_03242</td><td>Heat shock protein (molecular chaperone GrpE)</td><td></td><td>GrpE</td><td>481</td><td></td><td></td><td></td><td>2.1</td><td>2</td><td>2.2</td><td></td><td></td></t<> | F502_03242 | Heat shock protein (molecular chaperone GrpE) | | GrpE | 481 | | | | 2.1 | 2 | 2.2 | | |
| peptidase COG1026 Cym1 106 2.3 1.6 2.1 2.4 ATP-dependent Clp protease ATP-binding COG0542 ClpA 165 1.8 2.1 2.4 ATP-dependent Clp protease ATP-binding COG0459 GroEL 2.8 1.6 1.7 1.7 2.0 Chaperonin CO-chaperonin COG0234 GroES 613 1.8 1.5 2.6 2.6 Thij/Pfpl family protein COG0693 Thij 553 1.8 1.6 2.5 2.6 Heat shock protein (molecular chaperone lbpA) COG071 lbpA 600 2.5 1.6 2.5 2.4 Heat shock protein (molecular chaperone lbpA) COG0326 HtpG 196 2.2 2.5 2.4 2.4 Clpb protein COG0542 ClpA 136 1.6 2.5 2.4 2.5 Clpb protein COG0542 ClpA 136 1.6 2.5 2.5 2.5 Clpb protein COG0542 ClpA 1 | F502_03247 | Molecular chaperone DnaK | COG0443 | Dank | 217 | 1.6 | | | | _ | ∞. | | |
| ATP-dependent Clp protease ATP-binding COG0542 CIPA 165 1.6 2.1 2.4 subunit 168 1.8 2.1 1.9 1.7 2.0 Chaperonin Co-chaperonin COG0234 GroEL 258 1.6 2.0 2.0 Thill/PfpI family protein COG0693 Thil 553 1.8 1.6 2.5 3.0 Heat shock protein (molecular chaperone lbpA) COG0326 HtpG 196 2.2 3.0 2.4 Heat shock protein 90 COG0326 HtpG 196 2.2 3.0 2.4 Clpb protein COG0326 HtpG 196 2.2 2.4 1.5 Clpb protein COG0326 HtpG 18 1.6 2.4 2.4 Clpb protein COG0326 HtpG 1.8 1.5 2.4 2.4 Clpb protein COG0326 HtpG 1.8 1.5 2.4 2.4 | F502_03987 | peptidase | COG1026 | Cym1 | 106 | 2.3 | | | | 2 | 1.1 | | |
| ATP-dependent Clp protease ATP-binding COG0542 CIpA 165 1.8 2.1 1.7 1.7 2.0 Subunit Chaperonin COG0459 GroEL 258 1.6 1.5 2.0 Co-chaperonin COG0693 ThiJ 553 1.8 1.6 2.6 Thij/Pfolf family protein COG06071 IbpA 600 2.5 3.0 Heat shock protein (molecular chaperone lbpA) COG0326 HtpG 196 2.2 3.0 Heat shock protein 90 COG0326 HtpG 196 2.2 2.5 1.5 Clpb protein COG0542 ClpA 18 1.6 2.2 2.4 Thib 206 1.6 2.2 2.4 2.5 2.8 Clpb protein COG0542 ClpA 138 1.8 1.5 2.8 Thib 2.5 2.4 2.4 4.4 4.4 4.4 | | | | | 107 | 3.6 | | 1.6 | | 2 | 4. | | |
| Chaperonin COG0459 GroEL 258 1.6 2.0 Co-chaperonin COG0234 GroES 613 1.6 2.6 Thij/PfpI family protein COG0693 Thi 553 1.8 1.6 2.5 Heat shock protein (molecular chaperone lbpA) COG0376 HtpG 196 2.2 3.0 Heat shock protein 90 COG0326 HtpG 196 2.2 2.4 Clpb protein COG0542 ClpA 138 1.8 1.5 | F502_05557 | ATP-dependent CIp protease ATP-binding subunit | COG0542 | ClpA | 165 | ∞. | | 2.1 | | 17 | | | |
| Co-chaperonin COG0234 GroEs 613 1.6 1.5 2.6 Thij/Pfpl family protein COG0693 Thi 553 1.8 1.6 2.5 2.5 Heat shock protein (molecular chaperone lbpA) COG0326 HtpG 196 2.2 2.2 2.4 Heat shock protein 90 COG0342 ClpA 136 1.6 2.7 1.5 1.5 Clpb protein COG0542 ClpA 138 1.8 1.5 2.8 716 2.5 2.4 4.4 | F502 06242 | Chaperonin | COG0459 | GroEL | 258 | 1.6 | | <u> </u> | | | 5.0 | | |
| Thij/Pfpl family protein COG0693 Thij Thij 553 1.6 1.6 3.0 Heat shock protein (molecular chaperone lbpA) COG0326 HtpG 196 2.2 2.4 2.4 Heat shock protein 90 COG0342 CIpA 1,6 1.6 1.5 1.5 Clpb protein COG0542 CIpA 138 1,8 1.5 2.8 716 2.5 24 4.4 | F502_06247 | Co-chaperonin | COG0234 | GroES | 613 | | | | 1.5 | 2 | 5.6 | | |
| Heat shock protein (molecular chaperone lbpA) COG0326 HtpG 600 2.5 3.0 Heat shock protein 90 COG0326 HtpG 196 2.2 2.4 Clpb protein COG0542 ClpA 138 1.8 1.5 T16 2.5 2.4 4.4 4.4 | F502_07608 | Thij/Pfpl family protein | COG0693 | Thij | 553 | 1.8 | | 1.6 | | | | | |
| Heat shock protein 90 COG0326 HtpG 196 2.2 206 1.6 1.6 1.5 Clpb protein COG0542 ClpA 138 1.8 1.5 716 2.5 2.4 | F502_10228 | Heat shock protein (molecular chaperone lbpA) | | IbpA | 009 | | | 2.5 | | m | 3.0 | | 6.9 |
| 206 1.6 Clpb protein COG0542 ClpA 138 1.8 1.5 716 2.5 2.4 | F502_15425 | Heat shock protein 90 | COG0326 | HtpG | 196 | 2.2 | | | | 2 | 4.2 | | |
| Clpb protein COG0542 ClpA 138 1,8 1.5 716 2.5 2.4 | | | | | 206 | 9.1 | | | | _ | 5 | | |
| 2.5 2.4 | F502_18446 | Clpb protein | COG0542 | ClpA | 138 | 7.8 | | | 1.5 | 2 | 8: | | |
| | | | | | 716 | 2.5 | | | 2.4 | 4 | 4.4 | | |

Table 2 continued

| Accession no. | Protein name | Cluster of orthol- | Conserved | Spot no. | Fold ch | Fold change iron-related | -related | | Fold ch | ange grov | Fold change growth-related | pe |
|-----------------|--|-----------------------|-------------------|----------|--------------------------|--------------------------|-------------------------|-----|---------------|-----------|----------------------------|---------|
| | | ogous groups (COG) | protein domain | | Early phase higher at | hase at | Late phase higher at | at | Fe+ higher at | her at | Fe- higher at | yher at |
| | | | | | Fe- | Fe+ | Fe | Fe+ | Early | Late | Early | Late |
| F502_18743 | ATPase with chaperone activity clpC, two ATP- | COG0542 | ClpA | 147 | 2.1 | | 2.1 | | | 2.0 | | 2.0 |
| | binding domain protein | | ClpA | 149 | 2.5 | | 1.7 | | | 2.4 | | 1.6 |
| Signal transduc | Signal transduction/stress response/defense mechanism | | | | | | | | | | | |
| F502_04082 | GTP-binding protein | COG1217 | ТурА | 47 | | | | | 1.7 | | . 8. | |
| | | | | 148 | | 1.7 | | | 2.4 | | 2.1 | |
| F502_07703 | Chemotaxis histidine kinase. CheA (contains CheW-like adaptor domain) | COG0643 | CheA | 155 | 3.6 | | 2.2 | | | | 1.6 | |
| F502_10768 | Lipid hydroperoxide peroxidase | COG2077 | Трх | 572 | 3.9 | | 2.2 | | | 2.5 | | |
| F502_13258 | CBS domain-containing protein | COG0517 | CBS | 909 | 4.5 | | 4.8 | | | 1.7 | | 1.9 |
| F502_14770 | Serine protein kinase | COG2766 | PrkA | 193 | 4.7 | | 1.5 | | | 3.4 | | |
| | | | | 197 | 3.7 | | 2.6 | | | 2.1 | | |
| | | | | 200 | 6.2 | | 1.9 | | | 3.6 | | |
| | | | | 201 | 3.8 | | | | | 2.6 | | |
| F502_16565 | Nitrogen regulatory protein P-II | COG0347 | GlnK | 638 | 3.1 | | <u>6</u> | | | | | |
| F502_17612 | Alkyl hydroperoxide reductase | COG0450 | AhpC | 267 | 5.3 | | 2.9 | | | 2.4 | | |
| F502_17637 | Spore coat protein | COG3546 | CotlC | 548 | 2.3 | | 3.9 | | | | | 2.1 |
| F502_18092 | Stage V sporulation protein T | COG2002 | AbrB | 544 | 6.1 | | 4.5 | | | 1.7 | | |
| Transcription/D | Transcription/Defense mechanisms | | | | | | | | | | | |
| F502_12326 | Transcription accessory protein | COG2183 | Tex | 160 | | 1.7 | 1.8 | | 4.7 | | 1.5 | |
| Translation/Rib | Translation/Ribosomal structure and biogenesis | | | | | | | | | | | |
| F502_04537 | 30S ribosomal protein S2 | COG0052 | RpsB | 497 | | 2.1 | | 2.1 | 2.7 | | 2.7 | |
| | | | | 500 | | 2.4 | | 1.9 | | | | |
| F502_06817 | Ribosomal 5S rRNA E-loop binding protein Ctc/ L25/TL5 | COG1825 | RplY | 504 | 3.5 | | | | | 3.9 | | |
| F502_12196 | Ribosome-associated protein Y (PSrp-1) | COG1544 | RaiA | 292 | 2.4 | | | | | 2.4 | | |
| F502_18808 | Elongation factor Tu | COG0050 | TufB | 327 | | 1.7 | | 1.5 | 1.5 | | 4. | |
| F502_18833 | 50S ribosomal protein L1 | COG0081 | RpIA | 681 | | 2.4 | | 1.5 | 2.0 | | | |
| F502_18948 | 50S ribosomal protein L5 | COG0094 | RplE | 752 | | 2.1 | | | 8. | | | |
| F502_18963 | 50S ribosomal protein L6 | COG0097 | RpIF | 753 | | 2.4 | | | 2.3 | | | |

*! COG: according to the annotation for C. pasteurianum DSMZ 525 by BioCyc database collection (http://www.biocyc.org/organism-summary?object=CPAS1262499)

^{*2} Conserved Protein Domain Family: according to the definition by NCBI Conserved Domains and Protein Classification (http://www.ncbi.nlm.nih.gov/Structure/cdd/cdd.shtml)

of living organisms in general. In anaerobes, pyruvate can be metabolized through a variety of pathways but it is often oxidized to CO2 and acetyl-CoA with the concomitant reduction of a low-potential redox protein, like ferredoxin or flavodoxin. The enzyme responsible for this oxidative decarboxylation of pyruvate in many anaerobic bacteria and archaea is pyruvate: ferredoxin oxidoreductase (PFOR). PFORs contain thiamin pyrophosphate (TPP) for the cleavage of carbon-carbon bonds next to a carbonyl group, as well as iron-sulfur clusters for electron transfer (see [30, 31] and references therein). For example, PFOR from C. pasteurianum W5 (ATCC 6013) was characterized to be an air-sensitive homodimer with each subunit containing eight iron atoms in two [4Fe-4S] clusters, for which pyruvate is the best substrate found among several α -ketoacids [30]. In the genome of C. pasteurianum DSMZ 525, three homologue enzymes of PFOR are present, namely two pyruvate ferredoxin oxidoreductases (F502_01955 and F502_07648), and one pyruvate:ferredoxin (flavodoxin) oxidoreductase (F502_07643) [32]. In this study, F502_07648 (termed as PFOR1) and F502 07643 (termed as PFOR2) were identified among the proteins showing significant expression changes. On the 2-D gels both PFORs appeared as a chain of spots (Fig. 4). Protein identification showed that the spots 70, 75, 76, 77, 87, 90 are pI isoforms of the homodimeric protein pyruvate: ferredoxin (flavodoxin) oxidoreductase (F502_07643), whereas the spots 40, 57, 58 and 59 are pI isoforms of pyruvate:ferredoxin oxidoreductase (F502_07648). F502_07643 and F502_07648 are homologous proteins with a sequence identity match of 66% and positive match of 81%. They have nearly identical molecular weights but the pI value of F502 07648 is more basic than that of F502_07643, which was also obvious on the 2-D gels. Although previous studies showed that under conditions of iron limitation many anaerobes synthesize flavodoxins as substitution of ferredoxins for many enzymatic reactions [28], all the isoforms of the two PFORs showed, in general, higher expression at iron excess than at iron limitation (Fig. 4). The expression patterns of the isoforms of PFOR1 were similar to each other, with the highest expression at Fe+ late. In contrast, the expression patterns of the isoforms of pyruvate:ferredoxin (flavodoxin) oxidoreductase (PFOR2) were different to each other. While the expression level of the spot 87 in the middle of the spot chain did not change between Fe+ early and Fe+ late, the more basic isoforms (spots 70 and 90) showed higher expression at Fe+ early, and the more acidic isoforms (spots 75, 76 and 77) were up-regulated at Fe+ late which was similar to the expression pattern of PFOR1. Therefore, based on the proteomic results alone, it is not clear whether these two PFORs function in synchronization or are differently regulated in response

to iron availability. Furthermore, whether PFOR2 transfers the electrons generated during the decarboxylation reaction to a ferredoxin or flavodoxin remains elusive.

Beside the ferredoxin (flavodoxin)-dependent PFORs, acetyl-CoA can be synthesized from pyruvate through the pyruvate formate-lyase (PFL) with the formation of formate. There are three genes (F502_19556, F502_15690, F502_15710) in the genome of C. pasteurianum DSMZ 525 being annotated to encoding enzymes functioning as pyruvate formate lyase (PFL). Only one of these PFLs (F502_19556, also named formate acetyltransferase) was unambiguously identified in four protein spots (Fig. 4). Interestingly, the molecular weight of the two acidic isoforms (spots 178 and 181) appeared lower than that of the two basic isoforms (spots 174 and 175). Compared to the iron excess condition, where the expression of PFL was nearly not detectable, all these four PFL isoforms were significantly but differently up-regulated under the iron limitation condition. While the two acidic isoforms showed 2.5 and 3.7 folds increased abundances only in the Fe- early sample, the expression of the two basic isoforms were up to eightfold strongly up-regulated in the Fe- early sample and about sixfold in Fe- late sample. Thus, under iron limitation, it was apparently favorable for C. pasteurianum to use the PFL-catalyzed reaction for the conversion of pyruvate to acetyl-CoA. Correspondingly, under this condition the formate yield was clearly higher than that under the Fe+ condition, namely 0.25 ± 0.05 g/g biomass in contrast to 0.08 ± 0.02 g/g biomass. Nevertheless, the expression levels of the two PFORs, especially the pyruvate:ferredoxin (flavodoxin) oxidoreductase (PFOR2), were visibly higher than that of PFL. Since protein synthesis is an energy-demanding process, cells usually do not produce useless enzymes in noticeable amounts. The presence of the two PFORs under Fe- condition may point to a fact that, in the absence of iron, the two PFORs, especially the pyruvate: ferredoxin (flavodoxin) oxidoreductase (PFOR2), probably use flavodoxin instead of ferredoxin as the electron acceptor. Indeed, the expression of a flavodoxin (F502_13493) was strongly up-regulated under iron limitation for 14.3 folds in the exponential growth phase and remained high even after entering the stationary phase (8.0 fold higher in Fe- late than in Fe+ late). However, whether or not the up-regulated expression of this flavodoxin was coupled to the functionality of the PFORs remains to be verified. In case it is, it did not help much in sustaining the production of butanol under the Fe- condition.

Regulation of the ferredoxin pool

For the proper function of PFORs, ferredoxin_(red), which is generated in the PFOR-catalyzed pyruvate oxidative

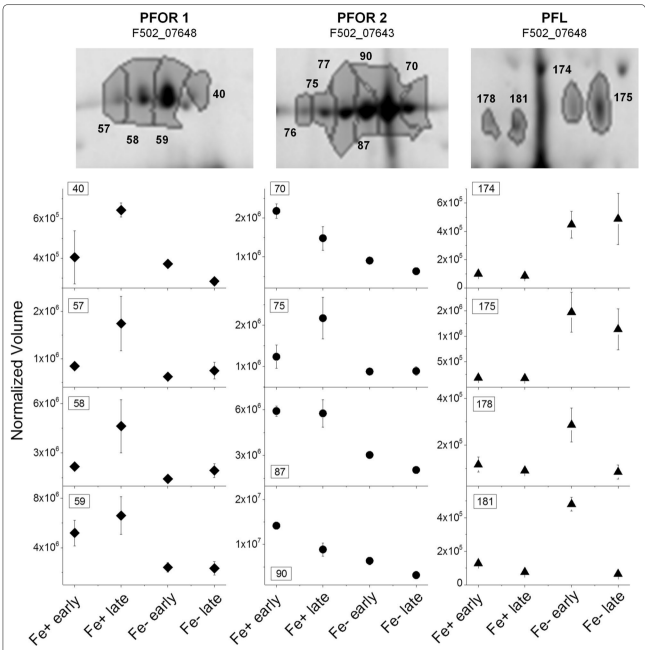


Fig. 4 Expression patterns of enzymes catalyzing the conversion of pyruvate to acetyl-CoA under Fe+ and Fe- conditions at exponential growth phase (early) and stationary phase (late). PFOR 1 (F502_07648, pyruvate: ferredoxin oxidoreductase) and PFOR 2 (F502_07643, pyruvate: ferredoxin (flavodoxin) oxidoreductase] are up-regulated at Fe+; PFL (pyruvate formate lyase) is up-regulated at Fe-

decarboxylation reaction, must be oxidized to regenerate ferredoxin $_{(ox)}$. *C. pasteurianum* DSMZ 525 possesses a big number of ferredoxins and the regeneration of ferredoxin $_{(ox)}$ can be achieved using different electron acceptors. The fact that the redox potentials of ferredoxins ($-400 \, \text{mV}$) are in the range of H_2 electrodes ($-414 \, \text{mV}$, at pH 7) reveals that in most energy metabolisms where ferredoxins are active, H_2 is also involved,

either as substrate or as product. In general, nitrogenases and hydrogenases are the two enzyme classes capable of hydrogen production in Clostridia [33]. But Hallenbeck and Benemann [34] reported that hydrogenases are much more efficient, with more than 1000 times higher turnover than nitrogenases. Hydrogenases are divided into two main groups in *Clostridia* based on their metaollocenter composition, namely [NiFe] and

[FeFe] hydrogenases [33]. In this study, the expression of hydrogenase-1 (F502 18287), which belongs to the [FeFe] group, was highly up-regulated under iron excess, showing up to fivefolds higher expression level in the exponential growth phase under the Fe+ condition compared to the Fe- condition. After entering the stationary phase the expression of hydrogenase-1 (H2-ase) was downregulated for two to threefolds under the Fe+ condition, which could be possibly in response to a depletion of the intracellular iron pool in the Fe+ late sample required for this [FeFe]-hydrogenase. However, it was still nearly twofolds higher than its expression level under the Fe- condition. An additional [FeFe] hydrogenase (F502 14390) was also identified which showed expression regulations similar to that of the hydrogenase-1 (F502 18287). Nevertheless this hydrogenase did not appear as a spot containing only a single protein on the 2-D gels and therefore could not be quantified for comparison. The higher expression of hydrogenase-1 (F502_18287) coincided with the higher H₂ production in the fermentation culture under iron excess and should have significantly contributed to the regeneration of ferredoxin(red) to ferredoxin_(ox).

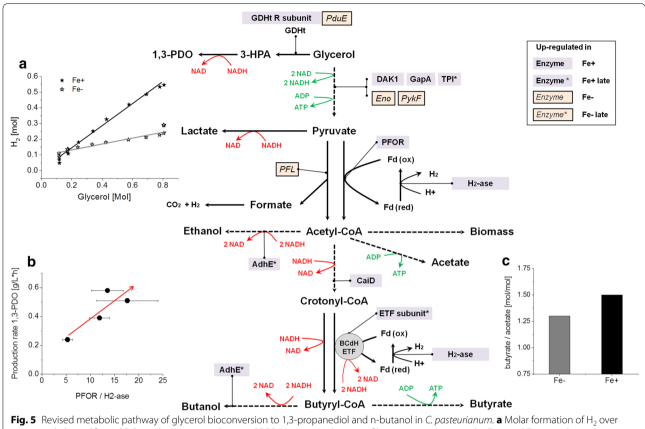
However, it is to notice that the regulation of hydrogenase-1 (F502_18287) expression is rather in agreement with that of the basic iso-forms of the ferredoxin (flavodoxin)-dependent PFOR2 than with the expression patterns of the ferredoxin-dependent PFOR1. Therefore, it is tempting to suggest that under the given experimental conditions the hydrogenase-1 catalyzed reaction should not be the only route of ferredoxin_(ox) regeneration. The PFORs-catalyzed pyruvate oxidation to acetyl-CoA might be coupled with other but yet unknown ferredoxin_(ox) regenerating reaction(s) catalyzed either by other unidentified hydrogenases (at least 5 genes in the genome of C. pasteurianum DSMZ 525 encode hydrogenases) or ferredoxin reductases. In addition, it has also been reported that PFOR can transfer the electrons generated in the decarboxylation reaction directly to protons to generate molecular hydrogen [35].

Within the cells of anaerobes including Clostridia, 90% of ferredoxins were reported to be present in reduced form, allowing them to serve as electron donors in different reactions [28]. In general, this is achieved in *C. pasteurianum* by the following three ferredoxin-dependent redox reactions: the oxidation of pyruvate to acetyl-CoA and $\rm CO_2$ (-500 mV), the oxidation of formate to $\rm CO_2$ (-430 mV, [36]) and the flavoprotein based electron bifurcation involved in the reduction of crotonyl-CoA to butyryl-CoA (Eq. 7). On the other hand, the oxidation of Fd_{red} by NAD is excluded due to the absence of the ferredoxin: NAD oxidoreductase activity [28]. Therefore, hydrogen production via hydrogenase should be

a main route of $\mathrm{Fd}_{\mathrm{ox}}$ regeneration in *C. pasteurianum* DSMZ 525. Based on this assumption, we compared the hydrogen yield from glycerol between the Fe- and the Fe+ conditions. As shown in Fig. 5a, hydrogen yield decreased significantly from 0.75 mol/mol glycerol under Fe+ condition to 0.21 under Fe- condition, which was in agreement with the higher expression of hydrogenase-1 under the Fe+ condition.

In addition, as described in the above redox balance analysis, a possible involvement of a ferredoxin-dependent butyryl-CoA dehydrogenase/electron transferring flavoprotein complex (BCdH-ETF) in H₂ production was proposed. In the BCdH-ETF catalyzed reaction electron transfer flavoproteins (ETFs) are involved in the reduction of crotonyl-CoA to butyryl-CoA, coupled with ferredoxin_(ox) reduction by bifurcating electrons from NADH (Fig. 5) [28, 37]. In this proteomic study, two ETFs, namely ETFs subunit alpha (F502 06282) and subunit alpha/beta-like protein (F502 06287), were identified among the most abundant proteins regardless of the iron availability; however, their abundances were slightly higher (1.5–1.7 folds) in the late phase of the Fe+ culture compared to the Fe- late condition. This might indicate a relative increase in the oxidized ferredoxin pool necessary to carry out the BCdH-ETF reaction and also contributed to the stronger H₂ production in the late fermentation phase under Fe+ condition.

Nevertheless, not only the hydrogenase-1 but also the two PFORs were down-regulated under Fe- condition compared to that under Fe+ condition. Therefore, the relative changes of the expression levels of the two enzymes might be indicative of the overall Fd_{ox} regeneration state (Fig. 5). The expression levels shown as the protein spot intensities of both enzymes under Fe- and Fe+ conditions at the two time points were thus compared. As shown in Fig. 5b, the relative expression of PFOR to hydrogenase-1 (H₂-ase) showed a positive correlation with the 1,3-PDO production rate. The decrease in the Fd_{ox} fraction under Fe – condition due to reduced H₂-ase presence will decrease the Fd_{ox} coupled synthesis of butyryl-CoA catalyzed by the BCdH-ETF complex, a crucial step in butyrate and especially butanol biosynthesis. Moreover, the intermediate acetyl-CoA will be favorably channeled into the Fd_{ox}-independent acetate formation route than the Fd_{ox} -dependent butyrate formation route, as shown by the increase in the butyrate/acetate ratio from 1.3 at Fe- to 1.5 at Fe+ (Fig. 5c). Consequently, it seems that under Fe- condition, Fdox dependent conversion steps are reduced and the resulting free reducing power, usually needed for butanol formation, could be redirected for the sake of redox balance to the production of 1,3-PDO and lactate. Indeed, the overall yield of 1,3-PDO and lactate were much higher under iron limitation than under



consumed glycerol **b** 1,3-PDO productivity over the ratio PFOR/H₂-ase **c** Molar ratio of butyrate to acetate under Fe+ and Fe- conditions

iron excess (Table 1). Nevertheless, lactate dehydrogenase catalyzing the conversion of pyruvate to lactate or 1,3-propanediol dehydrogenase catalyzing the formation of 1,3-propanediol from 3-HPA were not found among the proteins showing significant changes in expression level. Both enzymes are not involved in energy production but constitute the cell's back-up for stabilizing an internal redox balance, and hence their constitutive production may be a mechanism to withstand sudden perturbations in the NADH/NAD ratio. Nevertheless, it should also bear in mind that higher or lower protein level does not always means higher or lower enzyme activity.

Glycerol conversion to 1,3-propanediol

In general the bioconversion of glycerol to 1,3-PDO takes place in two steps, catalyzed successively by glycerol dehydratase (GDHt) and 1,3-propanediol dehydrogenase (PDOR) (Fig. 5). It is known that glycerol dehydratase is the rate-limiting enzyme in this bioconversion. All the three subunit of GDHt encoded by pduC (F502_03402), pduD (F502_03407) and pduE (F502_03412) were identified but unfortunately not as single protein spots and, therefore, could not be quantified. Instead, the large subunit of glycerol dehydratase reactivating factor (GDHt reactivase, GDHtR) was identified in the spot containing this single protein (Fig. 6). GDHtR is a molecular chaperone participating in the reactivation of inactivated GDHt in the presence of ATP and Mg^{2+} [38, 39]. The expression pattern of GDHtR indicates rather a correlation of GDHtR expression to cell growth phase than to iron availability. Among the four samples compared by proteomics, the highest expression level of GDHtR was present in the iron excess culture in the middle exponential growth phase (Fe+ early) showing the highest specific growth rate ($\mu = 0.22$). At this sampling time point, the culture under iron limitation (Fe- early) already entered late exponential growth phase with reduced specific growth rate ($\mu = 0.07$), accompanied with lower GDHtR level. The GDHtR abundance was further reduced to merely detectable levels in the stationary phase (Fe+ late and Fe- late), where the production of 1,3-PDO stagnated. 1,3-propanediol dehydrogenase (PDOR), the responsible enzyme for the conversion of 3-HPA to 1,3-PDO, was one of the highly abundant proteins on the 2-D gels and did not show significant expression changes under the different conditions (data not shown).

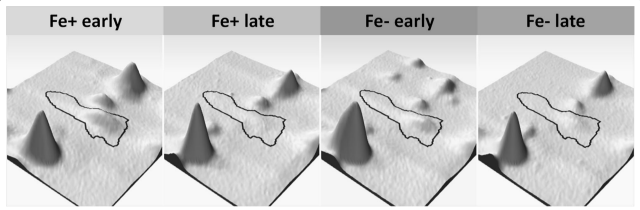


Fig. 6 Expression pattern of the large subunit of glycerol dehydratase reactivating factor (GDHtR) under Fe+ and Fe- conditions as well as early and late sampling point

Conclusion

The iron content in the fermentation medium was shown to influence the product formation, especially the 1,3-propanediol and butanol distribution, in *Clostridium* pasteurianum DSMZ 525 grown on glycerol. Compared to the fermentation under iron limitation, it was shown that the butanol to 1,3-propanediol ratio increased almost fivefold in fermentation under iron excess. To better understand the effect of iron on the regulation of the cell metabolism, physiological and proteomic analyses were performed. Several enzymes like pyruvate: ferredoxin oxidoreductase (PFOR), hydrogenase and bifunctional acetaldehyde-CoA/alcohol dehydrogenase among others were found to be up-regulated under iron excess conditions. The differential expression of PFORs and a pyruvate formate lyase (PFL) in response to iron availability highlighted the impact of iron on the crucial step of the central carbon metabolism in C. pasteurianum. The importance of a hydrogenase in the regeneration of oxidized ferredoxin and therefore the maintaining of the redox balance was confirmed by its strong upregulation under the iron excess condition. Beside the release of molecular hydrogen in the pyruvate to acetyl-CoA step catalyzed by PFORs, stoichiometric analysis showed a possible H₂ production coupled to the reaction catalyzed by the ferredoxin-dependent butyryl-CoA dehydrogenase/electron transfer flavoprotein complex (BCdH-ETF). Indeed, proteomic analysis revealed the up-regulation of two electron transfer flavoproteins which may be involved in this metabolic conversion step. Since both 1,3-propanediol and butanol can be used as sink for NADH, we suggest that the ratio of oxidized ferredoxin to reduced ferredoxin in addition to the NADH availability contributes to the selectivity of the products.

Authors' contributions

CG designed and performed the bioreactor experiments, interpreted the results and wrote the manuscript. WW carried out the proteomic studies, interpreted the results and wrote the manuscript. WS and TU helped in bioreactor experiments and results interpretation. APZ supervised the research. All authors read and approved the final manuscript.

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Competing interests

The authors declare that they have no competing interests.

Availability of data and materials

All data generated or analyzed during this study are included in this published article.

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