The glyceraldehyde-3-phosphate dehydrogenase of Clostridium acetobutylicum: isolation and purification of the enzyme, and sequencing and localization of the gap gene within a cluster of other glycolytic genes

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Glyceraldehyde-3-phosphate dehydrogenase was purified from Clostridium acetobutylicum by sequential ammonium sulfate precipitation, gel filtration and anion-exchange chromatography (to a specific activity of 27 U mg⁻¹). The enzyme had a molecular mass of 40 kDa as determined by SDS-PAGE and a native molecular mass of 160 kDa as determined by nondenaturing PAGE, indicating that it has a homotetrameric composition. Its pH optimum was between 8.5 and 9.3. The corresponding gene (gap) was cloned and sequenced from C. acetobutylicum DSM 792 and found to cluster with other genes of enzymes from the glycolytic pathway (pgk, phosphoglycerate kinase; tpi, triosephosphate isomerase; pgm(i), 2,3-bisphosphoglycerate-independent phosphoglycerate mutase). No sequences resembling rho-independent transcription terminators were found in the intergenic regions. A plasmid carrying the clostridial gap gene complemented an Escherichia coli gap mutant.

Keywords: GAPDH, glycolytic gene cluster, Clostridium acetobutylicum

INTRODUCTION

Clostridium acetobutylicum is an obligately anaerobic, Gram-positive, spore-forming bacterium which is characterized by a biphasic fermentation pattern. A typical butyric acid fermentation is carried out during the exponential-growth phase with acetate and butyrate as main fermentation products from starch or glucose. At the end of exponential growth, a metabolic transition to the production of organic solvents (acetone, butanol) takes place (Jones & Woods, 1986; Dürre, 1998). During the solventogenic growth phase, glycolytic reactions are most important for energy conservation in this organism, leading to a net production of two moles of ATP per mole of hexose consumed. In contrast to genes of enzymes involved in solvento- and acidogenesis, only limited information is available about the structure and organization of genes encoding glycolytic enzymes in C. acetobutylicum. So far, only the pfk (phosphofructokinase) gene has been cloned and sequenced. Adjacent to pfk, the pyruvate kinase gene (pyk) has been identified and it was shown that the two genes are organized within a single operon (Belouski et al., 1998). One key enzyme controlling glycolytic activity is the NAD⁺-dependent glyceraldehyde-3-phosphate dehydrogenase (GAPDH, EC 1.2.1.12) catalysing the reversible phosphorylation of glyceraldehyde 3-phosphate to 1,3-bisphosphoglycerate. GAPDH is inhibited by a high intracellular NAD⁺/NAD⁺ ratio (Girbal & Soucaille, 1994), which would otherwise favour the biotechnologically desired butanol production.

In this study, the purification of the clostridial NAD⁺-dependent GAPDH is presented. Cloning of the respective gene revealed a clustering with further genes (tpi, pgk, pgm(i)) encoding enzymes catalysing sequential reactions in the Embden–Meyerhof–Parnas pathway of glycolysis. Detailed knowledge of GAPDH and its

Abbreviations: GAPDH, glyceraldehyde-3-phosphate dehydrogenase; PGK, phosphoglycerate kinase; PGM, phosphoglycerate mutase; PGM(i), 2,3-bisphosphoglycerate-independent phosphoglycerate mutase; TPI, triosephosphate isomerase.

The GenBank accession number for the sequence reported in this paper is AF043386.
corresponding gene will offer the opportunity of constructing tailor-made strains that allow high glycolytic flux at high NADH concentrations and thus enhanced butanol production.

**METHODS**

**Bacterial strains, plasmids and growth conditions.** *C. acetobutylicum* DSM 792 (Deutsche Sammlung von Mikroorganismen und Zellkulturen, Braunschweig, Germany) was grown under strictly anaerobic conditions at 37 °C in clostridial basal medium (O’Brien & Morris, 1971). Continuous culture experiments were performed as described by Bahl *et al.* (1982) and were carried out in a Biostat B fermenter (Braun) with a working volume of 8 l. *Clostridium pasteurianum* DSM 525 was grown under anaerobic conditions in yeast/glucose medium (101 mM glucose, 1% yeast extract, 200 mM CaCO₃). *Escherichia coli* was cultivated aerobically at 37 °C in LB medium (Sambrook, 1989) supplemented with ampicillin (100 µg ml⁻¹) when required.

Functional complementation experiments were performed with the *E. coli* gap mutant W3CG (Ganter & Plickthun, 1990), in which this gene was inactivated by transposon Tn10 insertion. The mutant was cultured at 37 °C in M63 medium (Silhavy *et al.*, 1984) supplemented with malate or succinate (0.4%, w/v), glycerol (0.1%, v/v) and tetracycline (20 µg ml⁻¹). In screening experiments for functional complementation, bacteria were grown in M63 medium plus glucose or LB medium supplemented with tetracycline and ampicillin.

**Preparation of cell extracts.** Anaerobic conditions were maintained throughout the whole procedure. Culture aliquots (6 l) were removed from the continuous culture of *C. acetobutylicum* under sterile and strictly anaerobic conditions, employing tightly scaling centrifuge bottles and an anaerobic cabinet (Mecaplex). After harvesting, the cells were washed once in 1/10 0.1 M potassium phosphate buffer with 5 M CaCl₂ (pH 7.4). The cell pellet was suspended in 1/100 buffer, supplemented with DNase I (0.1 mg ml⁻¹) and incubated for 30--60 min at 37 °C. Lysis was achieved by 3--5 passages through a French pressure cell at 170 MPa. Cell debris was removed by centrifugation at 40 000 g for 1 h at 4 °C. The clear supernatant (crude extract) was used immediately or stored at −70 °C.

**Enzyme assays.** The activity of GAPDH was determined spectrophotometrically at 30 °C by monitoring the generation of NADH (physiological direction, oxidation of glyceraldehyde-3-phosphate in the presence of sodium ascorbate according to Ganter & Plickthun (1990) at 365 nm. A coupled assay in which aldolase cleaves fructose 1,6-bisphosphate to dihydroxyacetone phosphate and glyceraldehyde-3-phosphate and the latter product serves as actual substrate, was usually used during enzyme purification (Lovitt *et al.*, 1988). One unit of enzyme activity was defined as the amount of enzyme required to form 1 µmol NADH min⁻¹.

**Purification of GAPDH.** If not otherwise noted, all steps were performed at 4--8 °C. As a first purification step, a 50--90% (NH₄)₂SO₄ fractionation was carried out. The precipitated material was dissolved in 20 mM BTP (Bistrispropane) buffer (pH 6.5) containing 1 mM EDTA, 2 mM DTT and 5% glycerol and applied onto a gel filtration column (Superdex 200 preparatory grade; Amersham Pharmacia Biotech) equilibrated with the same buffer. The active fractions were pooled and concentrated if necessary using Diallo chambers and YM or FM membranes (Amicon). The protein solution was then applied to a Q-Sepharose HP column (C26/60; Amersham Pharmacia Biotech) equilibrated with BTP buffer. After the column had been washed with the same buffer, GAPDH was eluted with 5 column volumes of a linearly increasing NaCl gradient from 0 to 0.25 M NaCl in BTP buffer. For further experiments, the protein solution was desalted using PD10 columns according to the manufacturer’s instructions (Amersham Pharmacia Biotech).

Protein concentrations were determined by the dye-binding method of Bradford (1976) using bovine serum albumin as a standard.

**Gel electrophoresis.** PAGE was performed using the Laemmli buffer system (Laemmli, 1970). The purity and the molecular mass of the subunits of GAPDH were examined under denaturing conditions with SDS-polyacrylamide gels. For calibration, standard proteins in the range 14--98 kDa (Silver Stain SDS-PAGE Standards, Low Range; Bio-Rad) were used. Non-denaturing linear gradient gels (4--27.5%) were used to estimate the molecular mass of the native enzyme. Molecular mass standards in the range 67--660 kDa (HMW Calibration kit; Amersham Pharmacia Biotech) were used. Proteins in gels were silver-stained by the method of Blum *et al.* (1987).

**Nucleic acid isolation and manipulation.** Isolation of *C. acetobutylicum* and *C. pasteurianum* chromosomal DNA was performed according to Bertram & Durre (1989). Plasmid isolation from *E. coli* was done with the Qiagen Midi kit (Qiagen). DNA was manipulated by standard procedures (Sambrook *et al.*, 1989). Enzymes were purchased from commercial suppliers and reactions were carried out under the recommended conditions.

**PCR.** PCR amplifications were performed as described earlier (Treuner-Lange *et al.*, 1997). Synthetic oligonucleotides for PCR and sequencing reactions were prepared by a Gene Assembler Plus (Amersham Pharmacia Biotech) or purchased from commercial suppliers. The following primers were used for PCR amplification of a gap gene fragment of *C. pasteurianum*: 5'-GCTATATAATGTGGTTTTGGAAAT-3' (sense primer) and 5'-TGTTGACCATCACAATCTAAT-3' (antisense primer). These oligonucleotides correspond to the DNA sequence (GenBank accession number X72219) at positions 163--185 and 1061--1039.

**Genomic libraries of *C. acetobutylicum*.** A HindIII genomic library of strain DSM 792 (Sauer *et al.*, 1994) and another genomic library which resulted from partial digestion with Sau3AI and ligation into pEcoR252 (Fischer *et al.*, 1993) were used. Colony hybridization using Hybond-N nylon filters (Amersham) was performed for screening the genomic libraries. The filters were prepared according to Buluwela *et al.* (1989) for the hybridization procedure.

**Hybridization.** Chromosomal DNA of *C. acetobutylicum* was digested to completion with the appropriate restriction enzymes, separated on 0.8% agarose gels and transferred to Hybond-N+ nylon membranes (Amersham) by capillary transfer in 20 × SSC (1 × SSC is 0.15 NaCl, 15 mM sodium citrate). Prehybridization, probe labelling, hybridization, signal generation and detection were performed according to the instructions of the ECL direct nucleic acid labelling and detection system (Amersham).

Colony hybridization was performed with radiolabelled (α-³²P)dATP (Hartmann Analytic) DNA probes. The labelling reaction was performed using the random primer labelling kit (Gibco-BRL Life Technologies), followed by purification of the probe on Sephadex G-25 columns (NAP; Amersham).
Pharmacia Biotech). The filters were prehybridized in 0.2% (w/v) polyvinylpyrrolidone, 0.2% (w/v) Ficoll 400, 0.2% (w/v) BSA, 0.1% (w/v) sodium pyrophosphate, 10% (w/v) dextran sulfate, 1 M NaCl, 1% (w/v) SDS and 50 mM Tris/HCl (pH 7.5) for at least 0.5 h at 55 °C. After addition of the denaturated, radio labelled probe, hybridization was continued for 8–12 h. Filters were then washed twice in 2× SSC and subjected to autoradiography.

DNA sequencing and sequence analysis. A SequiTherm cycle sequencing kit for LI-COR 4000L (Biozym Diagnostik) and IRD41-labelled primers (MWG Biotech) were used for non-radioactive sequencing. The dideoxy-terminated fragments were separated on 41 cm long, 6% polyacrylamide gels or 66 cm long, 4.3% polyacrylamide gels (thickness 0.25 mm) with a LI-COR model 4000L sequencer (MWG Biotech) as instructed by the manufacturer. Data collection and analysis were performed by the programs BASE IMAGIR Data Collection (version 2.31) and BASE IMAGIR Image Analysis (version 2.3).

Nucleotide and deduced amino acid sequences were analysed by using the GeneWorks (IntelliGenetics) program on a Power Macintosh 8500/120 computer (Apple). Additional analysis was carried out on a UNIX computer, using the Genetics Computer Group sequence analysis software package (version 9.0).

RESULTS

Enzyme purification and characterization

The NAD⁺-dependent GAPDH was purified about 56-fold from acid-producing C. acetobutylicum cells using a procedure involving a two-step ammonium sulfate precipitation, gel filtration and anion-exchange chromatography. Results of the purification procedure are summarized in Table 1. The specific activity of the enzyme after anion-exchange chromatography was 27 U mg⁻¹. Dye-ligand chromatography on Reactive Blue 72 as last polishing step resulted in a loss of about 80% of the enzyme activity. Characterization was therefore carried out with the desalted anion chromatography pool.

SDS-PAGE experiments revealed one type of subunit with a relative molecular mass of 40 kDa (Fig. 1). The Mₐ of native GAPDH was determined by non-denaturing gradient PAGE. Analysis of three separate runs gave an average Mₐ of 160000 (±10000; data not shown). In agreement with the structure of many other known NAD⁺-dependent GAPDHs, the enzyme of C. acetobutylicum thus shows a homotetrameric composition. A pH activity profile was determined for the physiological direction with glyceraldehyde 3-phosphate (1 mM) as substrate and overlapping buffer systems in the range 5.8–11.6. The polynomial curve showed a maximum at pH 8.5, whereas highest activity was observed with sodium carbonate buffer at pH 9.3. Above pH 10, stability decreased rapidly. The temperature optimum of the enzyme was 55 °C; a further increase resulted in rapid inactivation. Above 65 °C no activity could be detected. According to the Arrhenius plot, an activation energy of 29 kJ mol⁻¹ was calculated in the range 10–50 °C. GAPDHs of various sources are able to use various aliphatic aldehydes as substrates. Since butyraldehyde is an intermediate in the butanol production of C. acetobutylicum, we were interested to determine if the GAPDH is able to use butyraldehyde as substrate. However, in contrast to other GAPDHs (e.g. Fife & Rikihisa, 1970), the clostridial enzyme was not able to

<table>
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<th>Purification step</th>
<th>Total protein (mg)</th>
<th>Total activity (U)</th>
<th>Specific activity (U mg⁻¹)</th>
<th>Recovery (%)</th>
<th>Purification factor</th>
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<tr>
<td>Ammonium sulfate (50–90%)</td>
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<td>518</td>
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<td>72</td>
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<tr>
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<td>173</td>
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<td>Reactive Blue 72</td>
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<td>35</td>
<td>14</td>
<td>5</td>
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![Fig. 1. SDS-PAGE of samples from the purification steps of GAPDH from C. acetobutylicum DSM 792. The proteins were separated on a 12% acrylamide gel. Lanes: 1, standard proteins (see Methods) with molecular masses on the left; 2, crude extract; 3, ammonium sulfate precipitate; 4, Superdex 200 pool; 5, Q-Sepharose pool; 6, Reactive Blue 72 pool.](image)
oxidize butyraldehyde at a significant rate (less than 1% activity compared to glyceraldehyde 3-phosphate as substrate).

To investigate possible regulation of the enzyme, specific activities were determined in the acidogenic, and early and late solventogenic stages of a continuous culture of *C. acetobutylicum*. The respective values were 0.42, 0.7 and 0.43 U·mg⁻¹ protein, indicating a low level of regulation at the most.

**Cloning and sequencing of the gap gene region from *C. acetobutylicum***

Based on the close relationship of *C. acetobutylicum* to *C. pasteurianum*, chromosomal DNA of *C. pasteurianum* was used to generate a 0.9 kbp probe via PCR encoding the main part of the gap gene. Southern hybridization with this heterologous probe showed positive signals with differently digested chromosomal DNA of *C. acetobutylicum* (data not shown). Two genomic libraries were therefore screened and several positive clones were detected. Two clones were sequenced, one (pWH19) obtained from the HindIII genomic library and the other (pWS10) from the partially digested Sau3A1 genomic library. Fig. 2 shows a schematic map of the gap gene cluster. The gap gene region of *C. acetobutylicum* includes three complete and two incomplete ORFs, all being transcribed in the same direction. The G+C content of the genes varies between 34 and 35.5 mol%.

From the first ORF (orf1') 831 bp were sequenced. Computer alignments (Table 2) of the deduced amino acid sequence revealed significant homology to the C-terminal part of hypothetical proteins of *Bacillus megaterium* (Ur1), *Bacillus subtilis* (YvbQ) and *Lactobacillus delbrueckii* (YgaP), which are also located upstream of their gap genes. The second ORF, encoding a protein of 334 amino acids and a deduced molecular mass of 35.8 kDa, was located 145 bp downstream of orf1'. Because of the high level of similarity of the deduced protein to NAD⁺-dependent GAPDHs of a variety of organisms and especially to the protein of *C. pasteurianum* (95% identity; Table 2), the respective gene was designated gap. A putative Shine–Dalgarno sequence (5'-TGGAGG-3') was located 8 bp upstream of the methionine start codon. Separated by 136 bp, a third ORF of 1194 bp was identified downstream of the gap gene. The deduced protein comprised 397 amino acids (42.3 kDa). A computer search revealed high level similarity between this gene product and those of various phosphoglycerate kinases (PGKs) (Table 2). The gene, designated pgk, is preceded by a putative Shine–Dalgarno sequence (5'-ATGAGG-3') 8 bp upstream of the start codon. Another ORF was located 198 bp downstream of pgk. Its predicted amino acid sequence, encoding a protein of 248 amino acids and a molecular mass of 26.5 kDa, showed high similarity to another glycolytic enzyme, triosephosphate isomerase (TPI) (Table 2). The respective gene was therefore designated tpi. Eight base pairs upstream of tpi, a putative Shine–Dalgarno sequence (5'-AGGAGG-3') was located. Further sequencing revealed a fifth ORF, located 188 bp downstream of tpi. Amino acid comparison showed high similarities of the respective gene product with the N-terminal region of members of the subclass of phosphoglycerate mutase (PGM) 2,3-bisphosphoglycerate-independent phosphoglycerate mutase [PGM(i)] (Table 2). A putative Shine–Dalgarno sequence (5'-TGGAGG-3') of this gene, pgm(i), was identified 8 bp upstream of the methionine start codon.

In the intergenic regions no typical rho-independent terminator structures could be identified, suggesting a common transcriptional unit comprising either at least these five genes or rho-dependent termination mechanisms (Platt, 1986). Nucleotide analysis with the programs Stemloop and FoldRNA (Zuker, 1989) revealed only secondary structures with comparatively
low energy values. The hairpin with the highest energy value (approx. $-20.6$ kJ mol$^{-1}$) was located downstream of the gap gene.

**Conserved functional and structural domains of the glycolytic genes of C. acetobutylicum**

The results of alignments of the deduced amino acid sequences of gap, pgk, tpi and pgm(i) of C. acetobutylicum with the respective gene products of other organisms (Gram-positive and Gram-negative prokaryotes and eukaryotes) are summarized in Table 2. As expected, a comparison of GAPDH of C. acetobutylicum revealed high similarity values (95%) to the closely related C. pasteurianum. Significant homology was also observed with GAPDHs of streptococcal origin (Lottenberg et al., 1992). Even enzymes from eukaryotic organisms showed relatively high homology values. Several regions with important catalytic or structural functions are conserved in all sequences, i.e. residues which are involved in forming and stabilizing the nucleotide-binding site (residues 3–18, especially Asn-7, Arg-11, Ile-12, Asn-32, Asp-47, Glu-76, Glu-94, Thr-96, His-108 and Glu-315 in the C. acetobutylicum sequence), in forming and function of the binding site for inorganic phosphate (Ser-151, Cys-152 and Thr-153), or in other catalytically important residues such as His-180 (Biesecker et al., 1977; Duee et al., 1996; Moras et al., 1975; Skarzynski et al., 1987; Soukri et al., 1989). A less conserved region is the so-called S-loop, which is...
responsible for building the tetrameric structure of GAPDH (residues 182–206).

The mostly monomeric enzyme PGK is also highly conserved among pro- and eukaryotic organisms. The degree of identity of the clostridial enzyme to other PGKs varies between 46% (E. coli) and 62% for the PGK part of the PGK–TPI fusion protein of the hyperthermophilic *Thermotoga maritima* (Schurig et al., 1995). In addition, eukaryotic enzymes such as PGKY from *Nicotiana tabacum* show a comparatively high identity of 59%. Taking into account the results of crystal structure and mutant analysis of pro- and eukaryotic organisms, several conserved sequence regions and certain amino acids which are important for the structure and the catalytic mechanism of PGK (Banks et al., 1979; Davies et al., 1993; Fairbrother et al., 1989; Watson et al., 1982) were identified. The PGK of *Clostridium acetobutylicum* also shows the highly conserved residues 198 to 220, 220 to 224, 297 to 298, 319 to 327 and 352 to 355, especially Lys-205, Leu-297, Glu-327 and Asp-355. His-61, Arg-154 and, in the C-terminal region, amino acids 373 to 378, which are involved in substrate binding. Arg-64, which is conserved in most sequences, is replaced in the clostridial enzyme by another basic amino acid, lysine. ADP-binding regions could be identified in the *Clostridium acetobutylicum* gene product (residues 198 to 202, 220 to 224, 297 to 298, 319 to 327 and 352 to 355, especially Lys-205, Leu-297, Glu-327 and Asp-355). His-61, Arg-154 and His-156 are the counterpart residues in *Clostridium acetobutylicum* which are essential for conformational change of PGK during catalysis (Fairbrother et al., 1989).

TPI is also well characterized by various crystal structure and mutant analyses. The degree of identity of the clostridial TPI and other TPIs varies from 60% (*B. subtilis*) to 42% for the eukaryotic yeast enzyme (Table 2). The catalytically and structurally important residues Lys-11, His-94, Glu-164-Gly 174 (especially Glu-166), Ile-207, residues 209 to 212 and Leu-231-Leu-237 are also found in the *Clostridium acetobutylicum* sequence (Banner et al., 1975; Lolis et al., 1990; Wierenga et al., 1992).

Based on sequence comparison the clostridial PGM was identified as PGM(i). The first PGM(i) was isolated from maize only in 1992 (Graha et al., 1992), explaining the lack of structural information about this enzyme type. A monomeric composition is characteristic for this class of PGMs (Fothergill-Gillmore & Watson, 1989).

**Functional complementation of an E. coli gap mutant**

To determine if the predicted clostridial gap gene really encodes the purified clostridial NAD$^+$-dependent GAPDH, the expression of this gene in the *E. coli* gap mutant (W3CG) was attempted. This mutant carries a transposon insertion within the gap gene so that no enzyme activity can be measured nor a complete protein be produced. It is therefore only able to grow with succinate or malate as carbon source. Mutant cells harbouring the recombinant plasmid pWH19 with the whole clostridial gap gene and parts of orf1 and pgk were able to grow on glucose minimal or LB medium.

Enzyme activities were determined from cell extracts of the mutants and the complemented mutants grown on M63 medium with succinate as carbon source. Activity in *Clostridium acetobutylicum* (0.5 U (mg protein)$^{-1}$) was determined from glucose-grown cultures. Whereas the extracts of W3CG and W3CG harbouring pUC9 showed a specific activity of less than 1 mU (mg protein)$^{-1}$, the complemented mutant had a 100-fold higher activity (110 mU (mg protein)$^{-1}$). The data confirmed that pWH19 really contained the clostridial gap gene and that this gene is expressed in the *E. coli* mutant W3CG.

**DISCUSSION**

The data reported describe the purification and enzymic characterization of the NAD$^+$-dependent GAPDH of *Clostridium acetobutylicum* DSM 792, and cloning and sequencing of the respective gene. Complementation of an *E. coli* gap mutant, and the similarity of the gene product to homologous enzymes, clearly revealed the identity of the gene.

Interestingly, the clostridial GAPDH seems to be a moderately thermotolerant enzyme with a temperature optimum of 50°C. This could be explained by a slightly higher portion of structure-stabilizing amino acids such as alanine, arginine, leucine and proline (Rentier-Delrue et al., 1993; Schläpfer & Zuber, 1992). An increased thermostability is not characteristic for enzymes of the mesophilic *Clostridium acetobutylicum*. GAPDH does not appear to be significantly regulated during the transition from the acidogenic to the solventogenic stage of *Clostridium acetobutylicum*. The less than twofold increase in specific activity corresponds nicely to data obtained from two-dimensional gel electrophoresis analysis, which also show an about twofold higher abundance of the protein in the early solventogenic stage. However, no increase in synthesis rate could be observed (Schaffer & Dürre, unpublished results), indicating that no regulation at the transcriptional level occurred, but rather a decrease in protein turnover or denaturation. This again is in accordance with the transient increase in chaperone synthesis at the onset of solventogenesis in *Clostridium acetobutylicum* (Pich et al., 1990), which also explains the later decrease in the enzyme specific activity to the original level.

The high degree of conservation of the clostridial GAPDH also extends to residues important for NAD$^+$/NADH binding. The invariant Asn-313 of *E. coli* is found in *Clostridium acetobutylicum* at position 314. A change of this amino acid in *E. coli* to threonine resulted in a drastic decrease in NAD$^+$ affinity (Dueé et al., 1996). Another possible way to influence the cofactor effect on the enzyme could be to induce a mutation at Gly-188 Pro-189 (conserved in *E. coli* at positions 187 and 188), which determine the specificity of NAD$^+$ versus NADP$^+$. A change of the proline to a serine in *E. coli* indeed allowed the enzyme to use NADP$^+$ as a coenzyme (Dueé et al., 1996).

The GAPDH of *Clostridium acetobutylicum* shows high homology (72–78% with *S. pyogenes*, Table 2) to members of
this enzyme class that are characterized by additional physiological functions, i.e. adhesion to fibronectin, myosin and actin or a function as plasmin receptor on the cell surface or a role in ADP ribosylation (Lottenberg et al., 1992; Pancholi & Fischetti, 1993). Eukaryotic GAPDHs are also involved in several processes such as endocytosis, DNA repair, tRNA export, programmed death of neurons, etc. (Sirover, 1997). This indicates that GAPDH might possess other functions in addition to its role in energy metabolism.

Nothing is known about the function of the ORF preceding the glycolytic genes in several bacteria (Table 2). It was first detected in B. megaterium (Schläpfer & Zuber, 1992), but there is no unequivocal evidence for its expression. For the corresponding gene in L. delbrueckii it was suggested that it is part of an operon localized directly upstream of the gap–pgk–tpi operon (Brann et al., 1998). Codon usage analysis of Orfl' shows a much weaker bias compared to the following glycolytic genes, indicating a weaker expression of the putative protein in C. acetobutylicum.

Sequence analysis of the incomplete ORF located at the 3' terminus of the insert revealed a high homology to PGM(i)s. Whilst most PGMs show a clear cofactor dependence on bisphosphoglycerate and a monomeric, dimeric or trimeric structure with subunit molecular masses of 23–30 kDa, this newly identified subclass of PGM is characterized by a monomeric structure with a molecular mass of approximately 60 kDa and these enzymes do not show a dependence on bisphosphoglycerate (Fothergill-Cillmore et al., 1998). Interestingly, the well-characterized PGM(i) of B. subtilis has an absolute requirement for Mn2+, which is responsible for the extreme sensitivity of the enzyme to pH variations: lowering of pH values results in the inactivation of PGM (Kuhn et al., 1993). This feature of PGM enables the forespore to accumulate 3-phosphoglycerate as an energy storage compound since sporulation is accompanied by a slow decrease in pH. As realkalinization is one of the first reactions during germination, PGM is partly reactivated and 3-phosphoglycerate can be reutilized within the very first minutes of germination (Kuhn et al., 1993). Clostridia also accumulate 3-phosphoglycerate in late phases of sporulation (Loshon & Setlow, 1994) and it is likely that the PGM(i) of C. acetobutylicum is also Mn2+-dependent and plays an important role in accumulating energy resources for germination. It remains to be shown if a decrease in PGM activity in C. acetobutylicum takes place upon induction of sporulation or solvent formation, which are accompanied by a decrease in pH.

Organization of glycolytic pathway genes in bacteria is characterized by a high degree of variability. Whereas in some bacteria, for example L. lactis (Cancilla et al., 1995a,b), genes like gap or tpi are not clustered with those of other enzymes of the glycolytic pathway, others show one or more glycolytic genes in the adjacent regions. In E. coli, gapA and tpi are not clustered with other genes of the pathway (Branlant & Branlant, 1985; Berlyn, 1998), but pgk, fda (fructose 1,6-bisphosphate aldolase gene) and gapB (erythrose-4-phosphate dehydrogenase) are (Alefounder & Perham, 1989). A bicistronic gap–pgk operon exists in Zymomonas mobilis (Eddy et al., 1989), whereas in Corynebacterium glutamicum, Borrelia burgdorferi and Lactobacillus delbrueckii a cluster of gap–pgk–tpi was detected (Eikmanns, 1992; Gebbia et al., 1997; Brann et al., 1998). Exactly the same arrangement of glycolytic genes as in C. acetobutylicum, including orfl', was found in B. subtilis and B. megaterium (Kunst et al., 1997; Schläpfer & Zuber, 1992). The glycolytic gene cluster of B. subtilis contains additionally the gene for the enolase (eno) downstream of pgm(i). The possibility that the clostralid gene cluster also contains an additional glycolytic gene downstream of pgm(i) cannot be ruled out. Computer analysis revealed no rho-independent terminator structures in this region. Future experiments will aim at a detailed transcript analysis of this region in C. acetobutylicum.

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