

# The *kdp* System of *Clostridium acetobutylicum*: Cloning, Sequencing, and Transcriptional Regulation in Response to Potassium Concentration

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The complete sequence of the *kdp* gene region of *Clostridium acetobutylicum* has been determined. This part of the chromosome comprises two small open reading frames (*orfZ* and *orfY*), putatively encoding hydrophobic peptides, and the genes *kdpA*, *kdpB*, *kdpC*, and *kdpX*, followed by an operon encoding a pair of sensor-effector regulatory proteins (KdpD and KdpE). Except for *orfZ*, *orfY*, and *kdpX*, all genes showed significant homology to the *kdp* genes of *Escherichia coli*, encoding a high-affinity potassium transport ATPase and its regulators. The complete genome sequence of *Synechocystis* sp. strain PCC 6803 and a recently published part of the *Mycobacterium tuberculosis* genome indicate the existence of a *kdp* system in these organisms as well, but all three systems comprise neither a second *orf* upstream of *kdpA* nor an additional *kdpX* gene. Expression of the clostridial *kdp* genes, including the unique *kdpX* gene, was found to be inducible by low potassium concentrations. A transcription start point could be mapped upstream of *orfZ*. A promoter upstream of *kdpD* was active only under noninducing conditions. Lowering the potassium content of the medium led to formation of a common transcript (*orfZYkdpABCXDE*), with a putative internal RNase E recognition site, which could be responsible for the instability of the common transcript. Except for the two small peptides, all gene products could be detected in *in vitro* transcription-translation experiments.

Regulation of the potassium content of the cytoplasm is the primary response of bacterial cells to osmotic stress (11, 23). A number of different K<sup>+</sup> transporters are known; these include uptake systems such as TrkG, TrkH, Kup, and Kdp (in *Escherichia coli*) as well as efflux systems such as the enterobacterial KefB, KefC, antiporters, colicin, and stretch-activated channels (for reviews, see references 7, 8, and 49).

Of these, Kdp of *E. coli* is the best-studied bacterial K<sup>+</sup> uptake system (for reviews, see references 2 and 46). It is an atypical P-type ATPase, consisting of three cytoplasmic membrane proteins, KdpA, KdpB, and KdpC, and probably a small hydrophobic peptide (KdpF). The transporter is induced by potassium limitation and changes in medium osmolarity. It is a high-affinity system that uses ATP as the driving force for K<sup>+</sup> uptake. The genes encoding the four different subunits are organized in a common operon, whose expression is controlled by a sensor kinase and response regulator (KdpD and KdpE). The respective genes, *kdpD* and *kdpE*, are organized in an operon directly downstream of *kdpFABC* (40, 57). However, little is known about *kdp*-analogous systems from other bacteria. DNA-DNA hybridization experiments indicated its presence in other enterobacteria, cyanobacteria, *Pseudomonas aeruginosa*, and *Spiroplasma citri* (47, 56). The solubilization of Kdp-ATPase complexes is described for *E. coli* (45), for *Alicyclobacillus acidocaldarius* (formerly *Bacillus acidocaldarius* [29]), and *Rhodobacter sphaeroides* (1), both including immunological cross-reactivity of KdpB with antibodies against the enterobacterial ATPase, and there is no evidence for a fourth subunit within the complexes.

Recently, genes of a two-component regulator system from

the gram-positive, obligately anaerobic spore former *Clostridium acetobutylicum* have been cloned and sequenced. They showed significant homology to *kdpD* and *kdpE* of *E. coli* (52). This report describes sequencing and analysis of the adjacent DNA region, which was found to comprise a *kdp*-homologous system with, however, several significant differences from the enterobacterial complex and from the sequences of *Mycobacterium tuberculosis* (*Mycobacterium tuberculosis* sequencing project [<http://www.sanger.ac.uk/pathogens/>]) and *Synechocystis* sp. strain PCC 6803 (30).

## MATERIALS AND METHODS

**Bacterial strains, plasmids, and growth conditions.** *C. acetobutylicum* DSM 792 was used as a source of genomic DNA and total RNA. The organism was grown under strictly anaerobic conditions at 37°C in clostridial basal medium (38) or in a minimal medium which contained, per liter, the following: glucose, 40 g; 2-[*N*-morpholino]ethanesulfonic acid (MES), 21.3 g; NaH<sub>2</sub>PO<sub>4</sub>, 1.4 g; (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 2 g; MgSO<sub>4</sub>, 0.1 g; NaCl, 10 mg; Na<sub>2</sub>MoO<sub>4</sub> · 2H<sub>2</sub>O, 10 mg; CaCl<sub>2</sub> · 2H<sub>2</sub>O, 10 mg; MnSO<sub>4</sub> · H<sub>2</sub>O, 15 mg; FeSO<sub>4</sub> · 7H<sub>2</sub>O, 15 mg; *p*-aminobenzoic acid, 2 mg; thiamine hydrochloride, 2 mg; biotin, 0.1 mg; and Na<sub>2</sub>S<sub>2</sub>O<sub>4</sub>, 35 mg. The fermentation medium for the potassium-limited chemostat experiments was the same as above except that no MES was added and a pH of 2 was adjusted with H<sub>2</sub>SO<sub>4</sub>. The chemostat experiments were performed in a fermentor with 1,000-ml culture volume, the temperature was held at 37°C by an external water bath, the pH was measured with a glass electrode (Dr. W. Ingold KG, Frankfurt, Germany), maintained at the desired value (6.0) by automatic addition of 2 N NH<sub>3</sub>, and the culture volume was maintained constant by providing an overflow pipe for gases and cell suspension. In the first limitation experiment, the starting potassium concentration was approximately 6 mM, using KCl as the potassium source. By addition of minimal medium without potassium at a dilution rate of 50 ml/h, the potassium concentration was slowly decreased to 50 μM within 100 h. The second limitation experiment started at a concentration of 1.5 mM KCl, which was decreased to 0.13 mM potassium at a dilution rate of 150 ml/h within 16 h.

*E. coli* JM109 (58), *E. coli* JM83 (58), and *E. coli* SURE (Stratagene GmbH, Heidelberg, Germany) were used as hosts, and pUC9 (53) and pEcoR252 (kindly supplied by D. R. Woods, University of Cape Town, Cape Town, South Africa) were used as vectors for cloning experiments and genomic library constructions. *E. coli* was grown aerobically in Luria broth or on Luria agar (41), supplemented with ampicillin (100 μg/ml) when required for plasmid selection and maintenance.

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**Measurement of potassium concentration.** Cells from 1 ml of culture were separated by centrifugation, and the potassium concentration of the supernatant was determined by flame photometry.

**Measurement of optical density.** The optical density at 600 nm ( $OD_{600}$ ) was measured in a spectrophotometer (Spectronic 601; Milton Roy, Ostende, Belgium). The light path was 1 cm.

**Nucleic acid isolation and manipulation.** Chromosomal DNA of *C. acetobutylicum* was isolated by the method of Marmor (35), with the modifications described by Bertram and Dürre (10). For isolation of plasmids from *E. coli*, a Qiagen Midi kit (Qiagen GmbH, Hilden, Germany) was used. Total RNA of *C. acetobutylicum* was isolated with the hot phenol-chloroform procedure of Oel-müller et al. (39) as modified by Gerischer and Dürre (28). DNA and RNA were manipulated by standard methods (41); restriction endonucleases and other modifying enzymes used were obtained from a variety of sources and used according to the manufacturers' instructions.

**PCR.** PCR amplifications were performed in 100- $\mu$ l volumes containing primers (0.5  $\mu$ M each) and chromosomal DNA (50 to 500 ng). When *Taq* polymerase (AGS GmbH, Heidelberg, Germany) was used, reactions were performed with 2 U of polymerase in 20 mM Tris hydrochloride (pH 8.55)–16 mM  $(NH_4)_2SO_4$ –1.5 mM  $MgCl_2$ –deoxynucleoside triphosphates (dNTPs; 50  $\mu$ M each) covered with paraffin oil. For amplifications of fragments that were to be used for further cloning procedures, 2 to 4 U of Vent polymerase was used in 10 mM KCl–10 mM  $(NH_4)_2SO_4$ –20 mM Tris hydrochloride (pH 8.8)–2 to 6 mM  $MgSO_4$ –0.1% Triton X-100 covered with paraffin oil. Temperature cycling was performed by a programmable thermocycler (Trio-Thermoblock; Biometra biomedizinische Analytik GmbH, Göttingen, Germany) with the following conditions: 95°C for 5 min (without enzyme), cooling to 80°C, and addition of enzyme, 95°C for 30 s,  $T_d$  (primer 1 or 2) – 5°C (50) for 30 s, and 72°C for 1 min/1,000 bp for 35 cycles.

**Primers.** Synthetic oligodeoxynucleotides as primers for sequencing and PCR were prepared with a Gene Assembler Plus (Pharmacia Biotech Europe GmbH, Freiburg, Germany) as instructed by the manufacturer. The following primers (in 5'-to-3' direction; numbers indicate the position of each primer within the *kdp* region) for PCR amplification, dot blot hybridization, and reverse transcriptase PCR (RT-PCR) were used: for RT-PCR, CTTTAAATGTGTAATGC (*kdpA*<sub>9</sub>, 1106 to 1122), CTATTGCTAATTTTGC (*kdpA*<sub>20</sub>, 2834 to 2850), TAAGTGC ATCTGCTTGA (*kdpA*<sub>21</sub>, 2897 to 2881), TAGCAATACCAAAAATG (*kdp9.1*, 4386 to 4402), CCCACAAAACCAATAC (*kdpB*<sub>2</sub>, 4579 to 4562), GAAAA TTCACITTTGTA (*kdp9.9*, 5299 to 5315), GGATTGATAATAATAT (*kdp9.10*, 5539 to 5555), TAGATAAGGAAGAAAA (*kdp9.11*, 5755 to 5771), GATGTGTACACCATT (*kdp9.13*, 6126 to 6142), GTCTGTAAATCTCT TTA (*kdpD*<sub>6</sub>, 6296 to 6280), ATTATTCTTGCCGAAGA (*kdpD*<sub>7</sub>, 6517 to 6501), ATGGATGATATTATCCC (*kdp9.22*, 7081 to 7065), TACATTCCTATAAA AAT (*kdpD*<sub>5</sub>, 7545 to 7562), CATCTATAAGTCTCCA (*kdp9.15*, 7836 to 7820), TCGCTATATTATAGCC (*kdp9.5*, 8519 to 8503), and TTTTACTC TATTGGC (*kdp9.2*, 8914 to 8898); for the *kdpCXD* probe, CTAGCTCGGGC TCTGGA (*kdp9.8*, 5066 to 5082) and *kdpD*<sub>6</sub>; for the *kdpA* probe, *kdpA*<sub>9</sub> and TTTATTCTTTGCCITC (*kdpA*<sub>22</sub>, 2145 to 2129); for the *kdpX* probe, *kdp9.9* and CCCTTGAAAAATGCCTT (*kdpD*<sub>2</sub>, 5597 to 5581); for the *kdpDE* probe, GGGATCCGAACATAAAATCAGTT (*kdpE*<sub>6</sub>, 9161 to 9145) and CTTTGTG ATTAATGCTC (*kdpE*<sub>4</sub>, 8103 to 8119); for cloning of pTC, CGGGATCCCGT TAATTTAGGATAGA (*kdpC*<sub>BamHI linker</sub>, 4644 to 4658) and CCGTCTCGAGC TGAATTTCTCATTA (*kdpC*<sub>XhoI linker</sub>, 5307 to 5293); and for quantitative dot blot hybridizations, ACACCCGCTGCATAGCC (*kdpD*<sub>3</sub>, 5821 to 5805), *kdpD*<sub>2</sub>, and TGCCCTACTGCCGTAT (*kdpC*<sub>12</sub>, 4769 to 4753).

**Genomic libraries of *C. acetobutylicum*.** Chromosomal DNA of strain DSM 792 was partially digested with *Sau3AI* and ligated in *Bam*HI-digested pUC9 (27) or in *Bgl*III-digested pEcoR252 (25). A third library used resulted from digestion of chromosomal DNA of strain DSM 792 with *Hind*III, fractionation by sucrose density centrifugation, and ligation of the fraction containing fragments of 0.8 to 3.5 kbp into *Hind*III-digested pUC9 (42). Colony hybridization using Hybond N nylon filters (Amersham Buchler GmbH, Braunschweig, Germany) was used for screening the genomic libraries. The colony filters were prepared as described by Buluwela et al. (13).

**Nucleic acid hybridizations.** For dot blots, 10  $\mu$ g of RNA was incubated at 65°C for 5 min in 3 volumes of a solution consisting of 500  $\mu$ l of formamide, 162  $\mu$ l of formaldehyde (37%, vol/vol), and 100  $\mu$ l of 10 $\times$  MOPS buffer (0.2 M 3-[*N*-morpholino]propanesulfonic acid, 0.5 M sodium acetate [pH 7.0], 0.01 M  $Na_2EDTA$ ). After addition of 1 volume of cold 20 $\times$  SSC (3.0 M NaCl, 0.3 M sodium citrate; 1 $\times$  SSC is 0.15 M NaCl plus 0.015 M sodium citrate) the RNA samples were spotted onto a Hybond N nylon membrane (Amersham Buchler), pretreated in 10 $\times$  SSC. RNA was fixed onto the membranes by UV cross-linking (direct contact of the RNA-containing side for 2 min at 254 nm on a Cromax 41 UV transilluminator [Vetter, Germany] or for 5 min at 312 nm on a dual light transilluminator [MWG-Biotech GmbH, Ebersberg, Germany]). For Northern blots, 10  $\mu$ g of RNA was separated in 0.8% (wt/vol) denaturing formaldehyde agarose gels and transferred to Hybond N nylon membranes (Amersham Buchler).

For hybridization with DNA fragments of >100 bp, membranes were prehybridized in 0.2% (wt/vol) polyvinylpyrrolidone–0.2% (wt/vol) Ficoll 400–0.2% (wt/vol) bovine serum albumin–50 mM Tris hydrochloride (pH 7.5)–0.1% (wt/vol) sodium pyrophosphate–10% (wt/vol) dextran sulfate–1% (wt/vol) sodium dodecyl sulfate (SDS)–1 M NaCl for 0.5 to 2 h at 55°C. The DNA fragments were

radiolabeled with [ $\alpha$ -<sup>32</sup>P]dATP (Hartmann Analytic GmbH, Braunschweig, Germany) by using a random primers labeling kit (GIBCO/BRL GmbH, Eggenstein, Germany). The labeled probes were purified by using Sephadex G-25 columns (NAP-5; Pharmacia Biotech Europe). For hybridization with oligonucleotides, membranes were prehybridized in 0.15% (wt/vol) polyvinylpyrrolidone–0.15% (wt/vol) Ficoll 400–0.15% (wt/vol) bovine serum albumin–90 mM Tris hydrochloride (pH 7.5)–6 mM  $Na_2EDTA$ –1% (wt/vol) SDS–salmon sperm DNA (1 mg/ml)–1 M NaCl for 0.5 to 2 h at  $T_d$  – 5°C (50). The oligonucleotides (17 pmol, 17-mer) were radiolabeled with 50  $\mu$ Ci of [ $\gamma$ -<sup>32</sup>P]dATP (Amersham Buchler) and purified with a QIAquick nucleotide removal kit (Qiagen). After addition of radiolabeled probes to the prehybridization solution, hybridization was continued for 10 to 12 h. Membranes were then washed in 2 $\times$  SSC (DNA probes of >100 bp, 45°C) or 6 $\times$  SSC (oligonucleotides,  $T_d$  – 10°C) and subjected to autoradiography.

**Quantification of hybridization results.** For quantification of hybridization signals, the membranes were subjected to autoradiography using Fujifilm BAS Phosphor Images plates (Fuji Photo Film Europe GmbH, Düsseldorf, Germany). The image plates were scanned with a BAS 1000 Image Reader (Fuji Photo Film Europe), and the scanned images were analyzed in the quantifications mode of the Mac Bas version 2.3 software (Fuji Photo Film Europe). Another method used was autoradiography with conventional X-ray films, scanning the films with a Umax Power Look scanner using the corresponding plug-in image-scanning MagicScan program that runs under Photoshop on a Macintosh computer (UMAX Data Systems Inc., Taipei, Taiwan). The TIFF-formatted images were analyzed with the Intelligent Quantifier software (Bio Image U.K., Cheshire, England). Pict-formatted images were analyzed with the Mac Bas Version 2.3 program (Fuji Photo Film Europe).

**DNA sequencing and sequence analysis.** The nucleotide sequence of the double-stranded plasmid DNA was determined by primer walking according to the dideoxy-chain termination method, using a T7 sequencing kit (Pharmacia Biotech Europe). All nucleotide data presented were confirmed by sequencing both strands of the inserts. The dideoxy-terminated fragments were separated on 55-cm wedge-shaped thickness gradient gels (0.2 to 0.4 mm, 6% [wt/vol] polyacrylamide) with a MacroPhor sequencing unit (Pharmacia Biotech Europe) as recommended by the manufacturer. The obtained sequence data were analyzed by using the DNA Strider (34) and GeneWorks (IntelliGenetics, Inc., Mountain View, Calif.) programs. Sequence comparison results and Codon Preference results were obtained with the Bestfit program of the Genetics Computer Group sequence analysis software package (Wisconsin Package version 9.0; Genetics Computer Group, Madison, Wis.) (20).

**Nonradioactive determination of the transcription start point.** The IRD41-labeled oligonucleotides *kdpA*<sub>9</sub> (GTACGAACAATAAGAATA [950 to 931]) and *orfZ*<sub>9</sub> (AAAAGCCTAGAACTATTAAT [738 to 719]) were obtained from MWG-Biotech and were used to determine a transcription start point upstream of *kdpA*. The annealing of 0.2 pmol of each primer was performed in a total volume of 10  $\mu$ l containing 10  $\mu$ g of total RNA, 10 mM Tris hydrochloride (pH 7.9), 0.5 M KCl, and 12.5 U of human placenta RNase inhibitor (GIBCO/BRL). After heating for 5 min at 80°C, the annealing reactions were incubated for 3 h at 30°C. For a 50- $\mu$ l primer extension reaction, 10  $\mu$ l of 5 $\times$  reverse transcription buffer (250 mM Tris hydrochloride [pH 8.3], 125 mM KCl, 50 mM dithiothreitol, 15 mM  $MgCl_2$ ), 500  $\mu$ M dNTPs, 2.5  $\mu$ g of actinomycin D, 200 U of SuperScript II (GIBCO/BRL), and  $H_2O$  were added to the annealing reaction mixtures and incubated for 1 h at 37°C. The primer extension products were phenol-chloroform extracted, ethanol precipitated, and analyzed on a 6% polyacrylamide sequencing gel running on a LI-COR model 4000L DNA sequencer (MWG-Biotech) as instructed by the manufacturer. The length of each of the primer extension reaction products was determined by running sequence reactions with the same primers on the same gel. The sequencing reactions were performed with the SequiTherm cycle sequencing kit for LI-COR (Biozym Diagnostik GmbH, Hessisch Oldendorf, Germany), using a MultiCycler PTC 200 (Biozym Diagnostik).

**RT-PCR.** cDNA synthesis was performed by reverse transcription of RNA primed with different sequence-specific oligonucleotides. The reverse transcriptase used was SuperScript II, obtained from GIBCO/BRL. After priming of the oligonucleotide (1 to 20 pmol) in 0.5 M KCl–10 mM Tris hydrochloride (pH 7.9), in the presence of 12.5 U of human placenta RNase inhibitor and 5 to 10  $\mu$ g of RNA, for 3 h at 30°C, the reaction mixture was divided into two parts. The first aliquot was brought to a final concentration of 50 mM Tris hydrochloride (pH 8.3)–25 mM KCl–3 mM  $MgCl_2$ –10 mM dithioerythritol–0.5 mM dNTPs–0.5 mg of actinomycin D. The second aliquot was treated the same way but received additionally 200 U of reverse transcriptase. Both mixtures were incubated for 1 h at 37°C and were used directly as templates in the following PCRs. As a further control, additional reactions were performed with plasmid or chromosomal DNA as a template, using the same master mix containing all necessary components for all assays.

**In vitro transcriptions and translations.** <sup>35</sup>S-labeled proteins were synthesized by using an *E. coli* S30 extract system for circular DNA (Promega Corp., Madison, Wis.) with <sup>35</sup>S-labeled methionine (Hartmann Analytic). After an acetone precipitation, the labeled translation products were analyzed on SDS–4 to 20% and 12 to 20% (wt/vol) polyacrylamide gradient gels and visualized by autoradiography. Protein size was determined by using <sup>14</sup>C-methylated protein molecular weight markers in a range of 14,300 to 200,000 (Amersham Buchler) and the

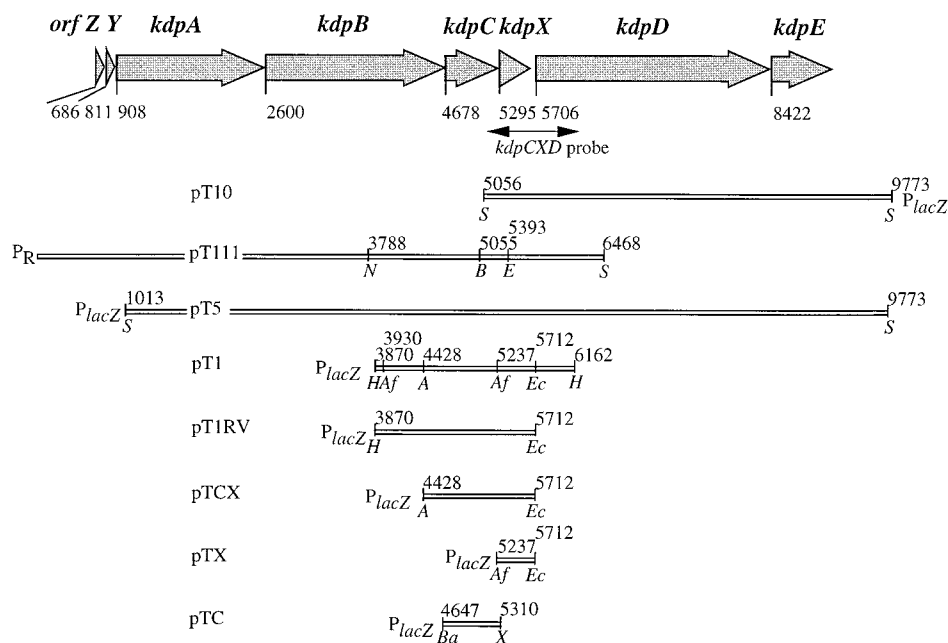


FIG. 1. Schematic representation of the *kdp* genes from *C. acetobutylicum*. The positions of the ATG start codons of the *kdp* genes (grey arrows) are indicated by numbers. The line with arrowheads at both ends indicates the *kdpCXD* probe used for screening of the different gene libraries of *C. acetobutylicum*. The 4,718-bp insert of pT10 (52) is shown, as well as the insert positions of the new positive clones pT111, pT5, and pT1. Clones pT1RV, pTCX, and pTX were constructed by different deletions from pT1. pTC is the product of cloning a PCR fragment with a *Bam*HI and an *Xho*I linker. Positions of the promoters of pUC9 ( $P_{lacZ}$ ) and of pEcoR252 ( $P_R$ ) are indicated. Numbers and letters represent restriction sites and their positions within the *kdp* gene region. Abbreviations: A, *Acc*I; Af, *Afl*II; Ba, *Bam*HI; B, *Bgl*II; E, *Eco*NI; Ec, *Eco*RV; H, *Hind*III; N, *Nde*I; S, *Sau*3AI; X, *Xho*I.

positive control reaction of the kit, showing the 60.7-kDa luciferase, a 48-kDa band due to an internal start site in the luciferase sequence, and the 31.5-kDa  $\beta$ -lactamase.

**Nucleotide sequence accession number.** The sequence data shown in Fig. 2 were submitted to the EMBL database and assigned accession no. U44892.

## RESULTS

**Cloning and sequencing of the complete *kdp* DNA.** For cloning of the upstream region of *kdpDE* (52), a DNA probe was generated by PCR using a primer pair (*kdp*<sub>9.8</sub>-*kdp*<sub>6</sub>) from the 5' end of the available *kdpDE* inserts. This 1,214-bp probe was used for screening of three different genomic libraries of *C. acetobutylicum*. Hybridization experiments allowed us to identify three new clones (pT1, pT5, and pT11) in addition to the already sequenced clone pT10 (52). pT1 contained a 2,298-bp *Hind*III fragment in the vector pUC9 (Fig. 1). The 8,368-bp pT5 insert consisted of a partially *Sau*3AI-digested fragment in the vector pUC9. Sequence analysis of the 5' and 3' ends of pT5 showed that the pT5 sequence contained two open reading frames (ORFs) in addition to the genes *kdpC*, *kdpX*, *kdpD*, and *kdpE* (52). The first ORF was truncated; the deduced amino acid sequence showed significant homology to the KdpA protein of *E. coli* and was named accordingly. The second ORF was identified as the *kdpB*-homologous gene of *C. acetobutylicum* and was thus designated *kdpB*. To determine the missing part of *kdpA* and the promoter region, the pT11 clone was sequenced by primer walking. The 7,940-bp pT11 insert represented a *Sau*3AI fragment ligated in the vector pEcoR252. The 3' end of the insert was located within the *kdpD* gene and contained approximately 2,400 additional bp upstream of the *kdpA* ATG start codon. For sequence analysis, a subclone (pT111) was constructed by deletion of a 1,562-bp *Sty*I fragment, creating blunt ends, and ligation into pEcoR252. The resulting plasmid, pT111, contained a 6,474-bp insert (Fig.

1). The nucleotide sequence of the *kdp* region was determined, and the heretofore unpublished sequence (starting with the *Sty*I restriction site of pT111 and including the complete *kdpC* gene) is presented in Fig. 2. The *kdp* region of *C. acetobutylicum* consists of eight ORFs (Fig. 1), all of which start with ATG start codons that are preceded by putative ribosome-binding sites (Fig. 2) (52). The designations of the ORFs and their positions within the *kdp* region, lengths, and G+C contents are summarized in Table 1. The DNA sequences of *orfZ*, *orfY*, and *kdpX* had a significantly lower G+C content than the genes *kdpA*, *kdpB*, *kdpC*, *kdpD*, and *kdpE*.

**Amino acid similarities and features of the deduced Kdp proteins from *C. acetobutylicum*.** The deduced amino acid sequences encoded by the *C. acetobutylicum* *kdpA*, *kdpB*, *kdpC*, and *kdpX* genes were used to search the different nucleic acid and protein databases with the BLAST computer program (3). As expected, there was considerable sequence similarity between the different KdpA proteins (Table 2) and the clostridial KdpA, consisting of 556 amino acids with a deduced  $M_r$  of 60,077. The KdpA protein of *E. coli* is predicted to span the membrane with 12  $\alpha$  helices based on the hydrophobicity of the protein (22) or with at least 10  $\alpha$  helices (14). Hydrophobicity plots (31) of the clostridial KdpA and the enterobacterial KdpA showed almost identical patterns, including hydrophobic regions long enough to be membrane-spanning segments (data not shown). Nine amino acid residues, suggested to form a periplasmic  $K^+$ -binding site in the *E. coli* KdpA (14), are conserved in the clostridial KdpA. However, of the four residues assumed to be part of the cytoplasmic  $K^+$ -binding site in *E. coli* (14), only two are conserved in the protein of *C. acetobutylicum*.

The 2,055-bp *kdpB* gene encodes a protein with 685 amino acids yielding a calculated  $M_r$  of 73,751. KdpB of *C. acetobutylicum* showed significant homology to a number of other

CCTTGGAACTCTTCTTATAG GAGCTTGGACTCTGGCAT AACAGAGATTTTGGTTAATT CGATACCATCGTAAATATTC TCGTATCTTGATACAAAAATC 100  
 TTTTGGAGAAATAGATTTTT TACTATCTGTACTCAAGATA CTATACATGGTTTTATAGTC ATTTTTATTCCAAGCATTTA TGTATTTTTCAAAAGAGGCT 200  
 TTTGGAGTGTGAGAAGAACT GCATCCAAATGTAATGAAA TTGTTAAAAGAAGGTTAAA AGCAACAAGTATTTCTTTT CATAATTTATCTCCTTGATA 300  
 ATTAATATTTTTTATATTTG ATAAAAAGCTTTTCATAGGT ATACGGTACTTAATATAAAT GTATAGTTTATATACAATAA TTATAGTATTATTTTTAAA 400  
 ATTGGGAATAAAAACCTGTG TTGTTATTTAATAAACCTGA AAGATATATGTAAGAGCTTC TTAATTTGCACAGAGATAAT ATATTTCTGGGAAACAAGA 500  
 ATAGTTTTTAATAATTAAT GAAATATCAAGGGAAGATAA AGTGAATTTTATCTCTCT TTTTTGCGTAAAGCTTGAA AACATTATCATTGAAATCT 600  
 TTATACTTTCTTAACACCTG CTCTTAAAATCTTAACTCAA ATTTTATATATATGGTGTA AGATGAAGGCATAAGCGATA AQAGGAGGGATAGATATGCT 700  
 TGATGTTATTTTCTAGTAT TAATAGTCTTAGGCTTTTA TTTTAAAGTATTTTATAAA CTGGTGTGAAGGAACCATAA ATAAGAAGTAGTTTTTAGG 800  
 D V I F L V L I V L G F L F L R Y F I N W C E G T I N K K  
GGGATTCAATATGACTTT TAGCTATTATAATCATTTTT CTATTTATATATTTGTGCTA TGCATTATTTAATCCTGAAA AATTTTAGGTAACAGGAGG 900  
 M I L L A I I I I F L F I Y L C Y A L F N P E K F  
*orfY*  
 TTAAGATATGGAAATATAC AAATAGCAATTAFTCTTATT GTGTTCTGACTTCTTTGTAT ACCTATAGGAAGATATATGT ACAAGTTTCAGAGCACAAA 1000  
 M E I L Q I A I I L I V F V L L C I P I G R Y M Y K V S E H K  
*kdpA*  
 AAAACTTTATAGTCCAGT ATTAGATAAGATTGATGGCT TTATATATAGCTTTCAGGT ATACAAAAAGAAGAGGAAAT GAACTGGAAGCAATATATTT 1100  
 K T L L D P V L D K I D G F I Y K L S G I Q K E E E M N W K Q Y I F  
 TTGCACCTTTAATGTGAAT GCAGTCCAGCAATTATAGG GTATATAATTTAAGAATTC AAGCAGTAGGTATTTTTAAT CCTAATCATGTAAGGGAAT 1200  
 A L L M C N A V P A I I G Y I I L R I Q A V G I F N P N H V K G M  
 GGAACAAGGACTCACCTTTA ACACAATAAAGCTTTTTA ACTAATACAAATCTGCAAGA TTATGCAGGAGAACTGGAG CCTCTTATTTATCGCAAATG 1300  
 E Q G L T F N T I I S F L T N T N L Q D Y A G E T G A S Y L S Q M  
 ATAGTAATTACATTTTTTAT GTTCTTTGCTGCTGCAACAC CAATAGCAGTTGCATTAGCA TTTATAAGAGCACTTTCAGG CAAGAAGAAATTAGGAACT 1400  
 I V I T F F M F F A A A T P I A V A L A F I R A L S G K K K L G N F  
 TCTATGTTGATCTTGTGAAGA ATTACAACAAGAATTAFTACT TCCTCTATCTATAATTTGTCG CTATATTTTATATTTGGACAG GGAGTACCACAAACACTTTTC 1500  
 Y V D L V R I T T R I L L P L S I I V A I F Y I G Q G V P Q T L S  
 GGCAAAATAGACAGTTACGA CAATAGAAGGAAAGCTTCAA AATATTCCTACTTGGTCCAGT TGCAAGCCTTGAAGCAATAA AGCTTATTGGAACAAATGGG 1600  
 A N K T V T T I E G K L Q N I P L G P V A S L E A I K L I G T N G  
 GGAGGCTTTTTTAGTGCTAA TTCATCTCATCTTTTGAAA ATCCAACACCGCTTACCAAT TCAGTGCAGATAATAACCCCT GCTTTTACTAGCAGGATCAA 1700  
 G G F F S A N S S H P F E N P T P L T N S V Q I I T L L L L A G S M  
 TGGTAGTATGTTTTGGACAC ATGATAAAAAAGAAAAACA GGCAGTAGCATATTTGAG CTATGATGTTACTACTTTTA GCAGGAGCAGTATATGTTT 1800  
 V V C F G H M I K K K K Q A V A I F A A M M V L L L A G A A I C F  
 TTCGCTGAGAAAGCAGGAAA TCAGCACTCTCAGCTATAGG CTTAAGTCAGAGCATGGGAA ACTGGAGGGGAAAGAAGAGA GGTTTGGAATAGCAGGGTCT 1900  
 S L R K Q E I S T L T Y R L K S E H G K L E G K E E R F G I A G S  
 AGTTTATTTACCACAGTAC AACGGATACCTCCTGTGGAG CGGTTAATAATATGACAGAT TCGCTAACACCAATTTGGAGG AGCTGTACCTCTTATAAATA 2000  
 S L F T T V T T D T S C G A V N N M H D S L T P I G G A V P L I N M  
 TGATGCTAAATGTAATTTTT GGAGGCGTTGGAGTTGGCTT TATGAACATGATAATGTACG CCATTTTAAACAGTTTTTCCTC TGCGGACTTATGTTAGGAAG 2100  
 M L N V I F G G V G V G F M N M I M Y A I L T V F L C G L M V G R  
 AACTCCAGAGTTTTTGAATA AGAAAATTGAAGCAAAGAA ATAAAGCTAGTAGCTTTTGC TATAATTTGTCATCCATTTT TAATATTAATGTCTTCAGCT 2200  
 T P E F L N K K I E G K E I K L V A F A I I V H P F L I L M S S A  
 TTGGCGCTTACAACAAAACA GGGACTAGCAGGAATATCGA ATCCAGGTTTTTACAGCACTT ACACAAGTTTTTATATCAATT TACCAGTTACAGCAGTAATA 2300  
 L A L T T K Q G L A G I S N P G F H G L T Q V L Y Q F T S S A A N N  
 ATGGTTCTGGATTTGAAGGG CTTATAGATAACACGATGTT TTGGAATGCTCAGCAGGTG TGGTTATGTTCTTAGGAAGA TATTTATCTATAATAACT 2400  
 G S G F E G L I D N T M F W N V S A G V V M F L G R Y L S I I I L  
 TTTAGCAGTAGCAAGTTCTT TTGGCGCTAAAAGAGCAGTA CCGGCAACGCAAGGAACCTT TAAAACGACAACACTATTT TACTGTAACGTTAATAGTT 2500  
 L A V A S S F A A K R A V P A T Q G T F K T D N T I F T V T L I V  
 ATTATAGTTATAATTTGGAGC ACTTACATTTCTTCCAGCAG TTGCACTTGGACCTATTTCA GAGTATCTAACCGTATAAAG CATAGGGGTGAGGTCATTA 2600  
 I I V I I G A L T F L P A V A L G P I S E Y L T L \*  
*kdpB*  
 TGAAGCAAAAAGTCAAAA TTTATTACAAAGGATATATT AAAGGAAGCCATAATTGAGT CTTTTAAAAAATTAACCTT AAATATATGATGAAAAATCC 2700  
 K S K K S K F I T K D I L K E A I I E S F K K I K P K Y M M K N P  
 GGTATGTTTGGTTGAGG TTGGATTCTTCGTTACAATT TTATTAACCATTTTCCAAG TATATTTGGAGATAAGGGAC ACAATTTAAGAGTATATAAC 2800  
 V M F V V E V G F F V T I L L T I F P S I F G D K G H N L R V Y N  
 TTAATGTAAACATCATTTT ATTTATAACGGTCTATTG CTAATTTTGCAGAGTCTGTA GCTGAAGGACGCGGAAAAGC TCAAGCAGATGCACTTAAAA 2900  
 L I V T I I L F I T V L F A N F A E S V A E G R G K A Q A D A L K K

FIG. 2. Nucleotide sequence of *orfZ*, *orfY*, *kdpA*, *kdpB*, and *kdpC* of *C. acetobutylicum*. The deduced amino acid sequences are provided in single-letter code below the DNA sequence. Putative ribosome-binding sites are boxed. The putative promoter region is marked by thick solid bars above the -10 and -35 regions joined by a line. The determined mRNA start point is marked by an arrow. Translation stop signals are marked by asterisks below the codon.

AGACCCGTAAGATACAATA GCAAAGCTCATAGGAAAAGA TGGTAGTATAAAAACATATA ATGCAAATGAGCTTAAAAAG GGTGATGATGTTCTTGATAGA 3000  
 T R K D T I A K L I G K D G S I K T I N A N E L K K G D V V L V E

AAATGGAGATGTAATACCAA ACGACGGTGAAGTGGTTGAC GGAGTTGCATCTGTAGATGA ATCAGCAATAACAGGAGAAT CAGCACCTGTTATGAAGGAG 3100  
 N G D V I P N D G E V V D G V A S V D E S A I T G E S A P V M K E

CCAGGAGGAGATTTGTCATC AGTTACAGGAGGAACAAAGG TTGTAAGCGATTGGTTAAAG GTTGAATAACAGCAACACC AGGAGAATCCTTCCTTGATA 3200  
 P G G D F A S V T G G T K V V S D W L K V E I T A T P G E S F L D K

AAATGATTAATCTTGATAGAA GGTGCTTCAAGGCAAAAAAC TCCTAATGAAATGCACTTA ATACAATACTTGTAGTCTT ACTTTGATATTTTTAATTGT 3300  
 M I N L V E G A S R Q K T P N E I A L N T I L V S L T L I F L I V

CTTGGTTGCACTTTACCCTA TGGCAACATACACAGGTGTA AAGATTCTATGTCAACCTT GATACGACTTTTAGTTTGTG TTATTCCAACAACCATAGGA 3400  
 L V A L Y P M A T Y T G V K I P M S T L I R L L V C L I P T T I G

GCACCTTTATCAGCAATAGG TATAGCAGGAATGGATAGAG TTACAAGATTTAATGTAATA GCAATGTCAGGAAAAGCAGT AGAGGCTTGTGGTGTGTTG 3500  
 A L L S A I G I A G M D R V T R F N V I A M S G K A V E A C G D V D

ATACAATGATCTTGATAAA ACAGGAACTATAACCTATGG AAATAGACTAGCAGCTGATT TTATAACGGTTGGAGGTGCA GATAAACAAAAATTAATAGA 3600  
 T M I L D K T G T I T Y G N R L A A D F I T V G G A D K Q K L I D

TTACTCCGTTATGTGTTCTT TAAAAGATGATACCCCTGAG GGTAAAGTCAATAGTTGAAC TGGAAAACAGTTAGGTATAA CAATAGATACTAAAAAATAT 3700  
 Y S V M C S L K D D T P E G K S I V E L G K Q L G I T I D T K K Y

GAGAGTATAGAATTTGAAGA GTTACAGCTCAAAACAAGA TGAGCGAAATAAGCTAGAA AATGGAATGCAAGTTAAAAA AGGAGCATATGATGCCATAA 3800  
 E S I E F E E F T A Q T R M S G I K L E N G T A V K K G A Y D A I K

AGAAAAGAGTACAGGAGTTA AAAGGAGTTATCCTAAAGA TTTAGATGAAGCTGTAAACA AGGTAGCAAAGCTTGGAGGA ACGCCACTTGTAGTATGTGT 3900  
 K R V Q E L K G V I P K D L D E A V N K V A K L G G T P L V V C V

TGATAATAAAATTTATGGAG TTATATATCTTAAGGATACA GTAAAGCCAGGCTTAGTTGA GAGATTTGAAAGGCTTAGGG AAATAGGTATAAAGACAATA 4000  
 D N K I Y G V I Y L K D T V K P G L V E R F E R L R E I G I K T I

ATGTGTACAGGGGATAATCC TTTAACAGCCGCAACTATAG CAAAGGAAGCTGGTGTGGAT GGATTTATAGCTGAGTGAA ACCTGAAGATAAGATAGAAG 4100  
 M C T G D N P L T A A T I A K E A G V D G F I A E C K P E D K I E A

CTATAAAAAGGAACAGGAC GAAGGAAAACCTTGTGCAAT GACAGGTGATGGAAC TAACG ATGCACCAGCACTTGCTCAG GCAGATGTTGGTCTTGCAAT 4200  
 I K K E Q D E G K L V A M T G D G T N D A P A L A Q A D V G L A M

GAATAGTGAACAACCCGAC CTAAGAGGCTGCTAACATG GTAGATTTGGATTCCGATCC TACAAAAGTCTTGAGGTTG TAGAAATCGGAAAGCAACTT 4300  
 N S G T T A A K E A A N M V D L D S D P T K V L E V V E I G K Q L

TTAATAACAAGAGGGCGCT TACTACCTTTAGTATAGCAA ATGATGTTGCTAAATATTTT GCTATAATACCAGCTATATT TACAATAGCAATACCAAAAA 4400  
 L I T R G A L T T F S I A N D V A K Y F A I I P A I F T I A I P K M

TGCAGCTAATGAATATAATG CACCTGTCTACTCCTTATAG TGCAATACTATCGGCACCTA TATTTAATGCGATAATAA CCGGCATTAATACCTATTGC 4500  
 Q L M N I M H L S T P Y S A I L S A L I F N A I I I P A L I P I A

AATGAAGGGTAAAGTACA GACCTATGAAATCAGAAGCT CTCTTTTAAAGAAATATGAT TGTATTTGGTTTTGGTGGAA TTATAGTCCGTTTGGTTGGA 4600  
 M K G V K Y R P M K S E A L L L R N M I V F G F G G I I V P F V G

ATTAAGATAATGATATGAT AATAACCCCAATGGTTAGAA TCCTTAATTTAGGATAGAG GAGGCATGGTTAATTTGATG AAATATTTTAAAGTGTCTCT 4700  
 I K I I D M I I T P M V R I L N L G \* M K Y F K S A L  
*kdpC*

TAGATTAGGTATTGTTTTAA TAATAATATGTGGACTTATA TATCCACTTTTTATAACGGC AGTAGGGCAGACAGTTTTTC ATAATAAGCAAAATGGAAGC 4800  
 R L G I V L I I I C G L I Y P L F I T A V G Q T V F H N K A N G S

ATAGTTACCTTTAAGGGTAA GGAGGTTGGCTCTGCTCTTT TAGGACAAAACCTTACGGAT AAAAGATTTTTTAGAGGAAG AGTTTCTTCTGTAAATTATA 4900  
 I V T F K G K E V G S A L L G Q N F T D K R F F R G R V S S V N Y N

ATACCTACACTAAAATGAC TCAATAAGGATGAAGTGGC CTCTGGTTCACAGAACCTAG CTCCATCCAATAAGGATTTA AAAAATAGGGTTAAAAGGA 5000  
 T Y T K N D S N K D E V A S G S Q N L A P S N K D L K N R V K K D

TATAGATGATTTCTTAAAAA CTCATCCAGGAGTGAAGAAG GATGAGATACTACAGATCT TTTAACTAGCTCGGGCTCTG GATTAGATCCAGATATAAGC 5100  
 I D D F L K T H P G V K K D E I P T D L L T S S G S G L D P D I S

CCTAAAGCAGTGAAATTTCA AGTGCCTTCTGTATCAAAAG CAACAGGCATAAGCCAAAGT AAACCTAAACAATAATAAA AAAATGTACAGAAGGTAGGA 5200  
 P K A A E I Q V P S V S K A T G I S Q S K L K Q I I K K C T E G R T

CTTGGGAGTACTTGGAGAG GAAAGAGTAAATGTTCTTAA GGTTAATCTTGAGGTAGCTT CAATGCTAAAGAATAGTAAA ATAGGTGAGTAATAATGAGA 5300  
 L G V L G E E R V N V L K V N L E V A S M L K N S K I G E \* \*

FIG. 2—Continued.

TABLE 1. Features of the different ORFs of the *kdp* system of *C. acetobutylicum*

ORF	Position (nucleotides)	Length (bp)	G+C content (mol%)
<i>orfZ</i>	696–789	93	26
<i>orfY</i>	811–886	75	19
<i>kdpA</i>	908–2576	1,668	35
<i>kdpB</i>	2600–4655	2,055	35
<i>kdpC</i>	4678–5290	612	33
<i>kdpX</i>	5295–5637	342	24
<i>kdpD</i>	5706–8406	2,700	33
<i>kdpE</i>	8422–9118	696	35

proteins. The highest value could be determined for the KdpB proteins of *Synechocystis* sp. strain PCC 6803, *M. tuberculosis*, and *E. coli* (Table 2). The next matches with decreasing homology are all characterized as P-type ATPases or E<sub>1</sub>-E<sub>2</sub> ATPases (Table 2). The highest similarities within the different P-type ATPases were found within cluster 4 described by Fagan and Saier (24), including all bacterial P-type ATPases except a Mg<sup>2+</sup>-ATPase of *S. typhimurium* (48). The three most conserved regions of P-type ATPases (24) could be identified in the clostridial KdpB. The motif (DNS)(QENR)(SA)(LIVAN)(LIV)(TSN)GE(SN) (24) is conserved in the clostridial KdpB with the amino acid sequences DESAITGES. Like KdpB of *E. coli* (32), the clostridial KdpB has a short C-terminal region and only one pair of possible transmembrane segments before the TGES sequence. The motif (LIV)(CAML)(STFL)DKTGT(LI)T, which is the essential phosphorylation site, was represented by the clostridial sequence MILDKTGTIT. The last very well conserved motif, (TIV)GDGXND(ASG)P(ASV)L, belongs to the ATP-binding site (24), and in *C. acetobutylicum*, the corresponding sequence TGDGTNDAPAL could be found.

The *kdpC* gene was found to consist of 612 bp and encodes a protein of 204 amino acids with a calculated *M<sub>r</sub>* of 22,187. KdpC of *C. acetobutylicum* showed similarities with the other KdpC proteins as expected (Table 2). Hydrophathy analysis of the KdpC proteins led to predicting strongly hydrophobic N-terminal transmembrane segments (data not shown).

The *kdpX* gene would encode a protein of 114 amino acids with a calculated *M<sub>r</sub>* of 13,465. KdpX did not show significant homology to other known protein sequences, but it resembles some prokaryotic and eukaryotic proteins shown in Table 2, including KdpC of *E. coli* in an N-terminal hydrophobic region. Nothing is known about the function of the predicted *orf27* product from *Helicobacter pylori*. In a distance of ca. 4,000 bp, another ORF (*orf32*), encoding a protein which resembles the potassium efflux system protein KefC of *E. coli*, has been detected. However, any possible functional relationship would be purely speculative without further experimental evidence. The three ATPase synthase proteins 8 of the different *Drosophila* spp. (Table 2) are all described as a membrane-bound nonenzymatic component of the mitochondrial ATPase complex. Within the first N-terminal amino acids, there are several conserved residues in KdpC of *E. coli* and in KdpC of *C. acetobutylicum* as well as in the clostridial KdpX (Fig. 3). KdpC in the Kdp-ATPase of *E. coli* seems to play a role in assembly of the complex or the subunits KdpA and KdpB, because KdpA/B aggregates are not formed without KdpC (49).

The two ORFs upstream of *kdpA* seem to have peptide-coding function because of the translational signals (Fig. 2). The first ORF, as designated *orfZ*, consists of 93 bp and would encode a small, rather hydrophobic peptide with 31 amino acids. The second ORF was designated *orfY*. The 75-bp *orfY*

would also encode a small, rather hydrophobic peptide. The KdpF peptide of *E. coli* is likewise encoded by an ORF of 29 codons just upstream of *kdpA* (46). Upstream of *kdpA* of *M. tuberculosis*, we found another small ORF, encoding likewise a small hydrophobic peptide with 32 amino acids. In Table 2, this ORF is designated KdpF, in analogy to the Kdp system of *E. coli*. In the genome sequence of *Synechocystis* sp. strain PCC 6803, we could find no similarly small ORF.

**In vitro transcription-translation experiments.** Proteins of the sizes expected for KdpB, KdpD, and KdpE were produced in a coupled in vitro transcription-translation assay of clones pT11 and pT5 (Fig. 4). In case of the 60,077-Da clostridial KdpA, a protein band with an apparent *M<sub>r</sub>* of approximately 50,000 was detectable, resembling the situation in *E. coli*, where the 59-kDa KdpA runs as a diffuse band with an apparent *M<sub>r</sub>* of 49,000 in an SDS-gel (45). No distinct protein that might represent the proteins KdpC and the putative KdpX could be detected. In a second in vitro transcription-translation experiment, the constructed subclones pTC, pTCX, and pTX

TABLE 2. Amino acid similarity of the Kdp peptides and proteins from *C. acetobutylicum* to corresponding proteins of other organisms

Organism	Protein (aa)	% Identity	% Similarity	Accession no.
OrfZ (31 aa)				
<i>E. coli</i>	KdpF (29)	45.5 <sup>a</sup>	72.3 <sup>a</sup>	P36937
<i>M. tuberculosis</i>	OrfY (32)	45.0 <sup>b</sup>	60.0 <sup>b</sup>	Z92539
OrfY (25 aa)				
<i>M. tuberculosis</i>	KdpF <sup>c</sup> (32)	45.0	60.0	Z92359
<i>E. coli</i>	KdpF (29)	41.7	70.8	P36937
KdpA (555 aa)				
<i>Synechocystis</i> sp.	KdpA (559)	55.1	45.3	D90910
<i>E. coli</i>	KdpA (557)	43.8	56.2	P03959
<i>M. tuberculosis</i>	KdpA (571)	41.8	51.8	Z92539
KdpB (685 aa)				
<i>Synechocystis</i> sp.	KdpB (691)	61.9	71.7	D90910
<i>M. tuberculosis</i>	KdpB (708)	55.9	68.8	Z92539
<i>E. coli</i>	KdpB (682)	55.4	67.5	P03960
<i>M. tuberculosis</i>	P-type ATPase (770)	32.0	41.6	1524193
<i>Proteus mirabilis</i>	P-type ATPase (829)	31.1	40.7	1353678
<i>Synechococcus</i> sp.	P-type ATPase (747)	29.5	41.8	584792
KdpC (204 aa)				
<i>Synechocystis</i> sp.	KdpC (191)	44.8	53.6	D90910
<i>M. tuberculosis</i>	KdpC (189)	40.1	50.3	Z92539
<i>E. coli</i>	KdpC (190)	37.7	48.6	P03961
KdpX (114 aa)				
<i>Helicobacter pylori</i>	Orf 27 (255)	31.0 <sup>d</sup>	55.2 <sup>d</sup>	1800178
<i>Drosophila yakuba</i>	ATP synthase protein 8 (53)	50.0 <sup>e</sup>	60.7 <sup>e</sup>	114481
<i>D. melanogaster</i>	ATP synthase protein 8 (53)	46.4 <sup>e</sup>	60.7 <sup>e</sup>	114480
<i>D. simulans</i>	ATP synthase protein 8 (53)	46.4 <sup>e</sup>	60.7 <sup>e</sup>	1703632
<i>E. coli</i>	KdpC (191)	23.7 <sup>f</sup>	39.5 <sup>f</sup>	P03961

<sup>a</sup> Result of an alignment over a length of only 11 amino acids (aa).

<sup>b</sup> Result of an alignment over a length of only 12 amino acids.

<sup>c</sup> This name was chosen in analogy to the Kdp system of *E. coli*, missing also the second Orf of the Kdp system of *C. acetobutylicum*.

<sup>d</sup> Result of an alignment over a length of only 29 amino acids.

<sup>e</sup> Result of an alignment over a length of only 28 amino acids.

<sup>f</sup> Result of an alignment over a length of only 38 amino acids.



FIG. 3. Amino acid alignment of the proteins KdpC of *E. coli* and *C. acetobutylicum* and the clostridial KdpX. Amino acids identical among the sequences are boxed.

as well as pT1 (Fig. 1) and the vector control were used. Because of the small expected  $M_r$  of 22,187 (KdpC) and 13,465 (KdpX), we used an SDS-12 to 20% gradient gel for the separation (Fig. 5). We detected two bands which were not produced in the vector control. The putative KdpC showed some deviation from the expected mobility; the protein band corresponded to a molecular mass of ca. 27 kDa (Fig. 5).

**K<sup>+</sup> limitation experiments and mRNA analysis.** With total RNA from *C. acetobutylicum* cells growing in clostridial basal medium with ca. 38 mM potassium and harvested in the exponential acidogenic growth phase and the stationary solventogenic growth phase, it was not possible to determine a 5' end of an mRNA upstream of *orfZ*, *orfY*, or *kdpA*, and there were no detectable Northern signals by hybridization experiments with different *kdp* probes. Since the genes *kdpA*, *kdpB*, and *kdpC* of *E. coli* encode an inducible high-affinity K<sup>+</sup>-ATPase, which is induced only under K<sup>+</sup>-limiting conditions, it seemed necessary to find conditions in which the clostridial *kdp* genes were induced. In two different chemostat experiments, the K<sup>+</sup> concentration was continuously decreased and RNA samples were taken at different stages of the limitation. K<sup>+</sup> concentration was determined in the cell-free supernatant of all samples. In the first limitation experiment, the starting potassium concentration was 6.04 mM K<sup>+</sup> (Fig. 6). Down to a potassium concentration of 600  $\mu$ M, the optical density and thereby the population in the chemostat was not influenced by the limitation and consistently yielded values of 2.3. However, continued decreasing of the potassium concentration led to reduced optical density, and the growth of the population could no longer balance the dilution. A subsequent increase of K<sup>+</sup> to 5.24 mM restored the original OD<sub>600</sub> of 2.3. Dot blot hybridization with a *kdpA* probe revealed that induction of the clostridial *kdpA* gene occurred when the potassium concentration was decreased. Highest induction occurred at potassium concentrations below 200  $\mu$ M. Increasing the amount of K<sup>+</sup> again led to

disappearance of the hybridization signals (Fig. 6). Growth impairment by the limitation paralleled the appearance of *kdpA* mRNA transcripts. Hybridization with *kdpB* and *kdpC* probes showed an identical pattern (data not shown). In a second limitation experiment, the potassium concentration was decreased from 1.5 mM to 130  $\mu$ M (Fig. 7). Despite some fluctuations of the potassium concentration in this chemostat, it is evident that the optical density of the culture started to decrease at 0.49 mM K<sup>+</sup>. *kdpA* and *kdpX* were clearly induced, but the hybridization signals with the *kdpX* probe yielded weaker signals although the same amount of RNA was used.

Using RNA samples 7 (first limitation; Fig. 6) and 4 (second limitation; Fig. 7), we determined the site of transcription initiation upstream of *kdpA* with two different primers. Primer extension reactions revealed in all cases a 5' end of the mRNA 236 bp upstream of the *kdpA* start codon. The primer extension signals revealed with the *orfZ*<sub>p</sub> primer are shown in Fig. 8. This mRNA start point was preceded by a possible promoter sequence. The putative promoter (5'-TCAAT-16 bp-TAA GAT-3' [Fig. 2]) showed only little homology to the consensus promoter of clostridia (5'-TTGACA-17 bp-TATAAT-3' [59]). The determined transcription start point of the *kdp* genes would allow transcription of the two small ORFs upstream of *kdpA*. The same RNA samples were used to confirm the mRNA start point upstream of *kdpD* (52) by primer extension (*kdpD*<sub>2</sub> served as a primer for the primer extension and sequencing reaction). No distinct signal of the expected size of 128 bp could be detected with these RNA samples, but there were a number of signals with larger and smaller sizes in the gel (data not shown). The same result was obtained when the adjacent oligonucleotide *kdpD*<sub>2</sub> was used.

RNA samples of the first limitation experiment were used for two identically prepared Northern blots. The first blot was hybridized with a *kdpA* probe (1,039 bp); the second was hybridized with a *kdpDE* probe (1,058 bp, last 300 bp of the *kdpD*

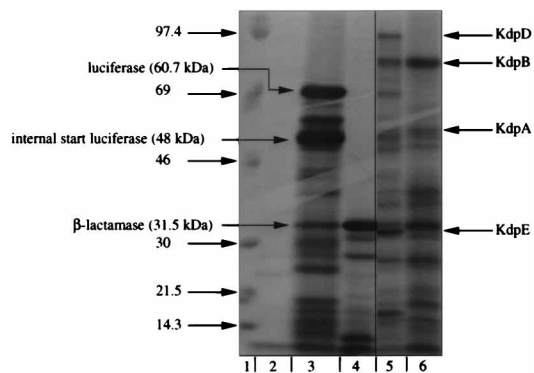


FIG. 4. Protein analysis of in vitro transcription-translation reactions by SDS-4 to 20% gradient gel electrophoresis and subsequent autoradiography. Lane 1, molecular weight markers; lane 2, negative control (no DNA added to the otherwise complete reaction mixture); lane 3, proteins synthesized from positive control plasmid pBestluc; lane 4, proteins synthesized from pUC9; lane 5, proteins synthesized from pT5; lane 6, proteins synthesized from pT111.

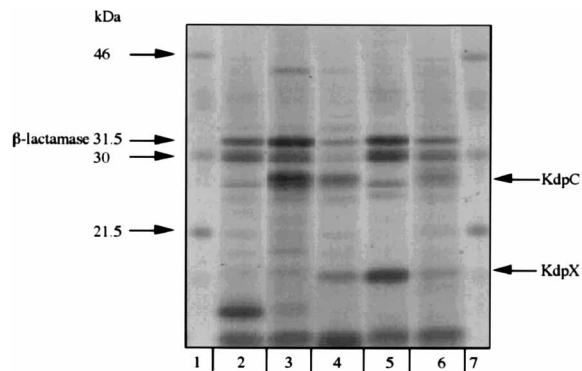


FIG. 5. Protein analysis of in vitro transcription-translation reactions in an SDS-12 to 20% gradient gel, with subsequent autoradiography. Lanes 1 and 7, molecular weight markers; lane 2, proteins synthesized from pUC9; lane 3, proteins synthesized from pTC; lane 4, proteins synthesized from pTCX; lane 5, proteins synthesized from pTX; lane 6, proteins synthesized from pT1.

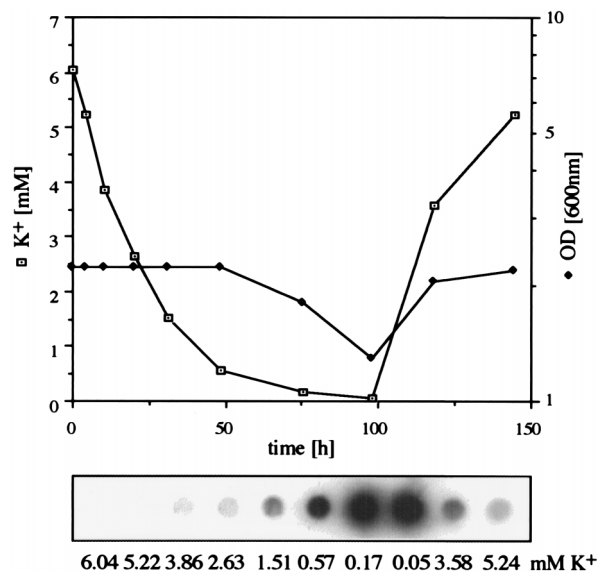


FIG. 6. Influence of decreasing the potassium concentration on growth in a continuous culture of *C. acetobutylicum*. Below, dot blot hybridization signals of RNA samples with a *kdpA* probe are presented. Potassium concentrations of the supernatants of the respective samples are indicated below the autoradiography signals.

sequence). In both cases, the same dependence of hybridization signals on the potassium concentration could be observed as in the dot blots, but it was not possible to determine the length of the mRNA transcript (data not shown). There were no distinct bands on the blots, but there was a large smear, especially at K<sup>+</sup> concentrations of 570 and 50  $\mu$ M. This was not due to the quality of the RNA preparation since the blots

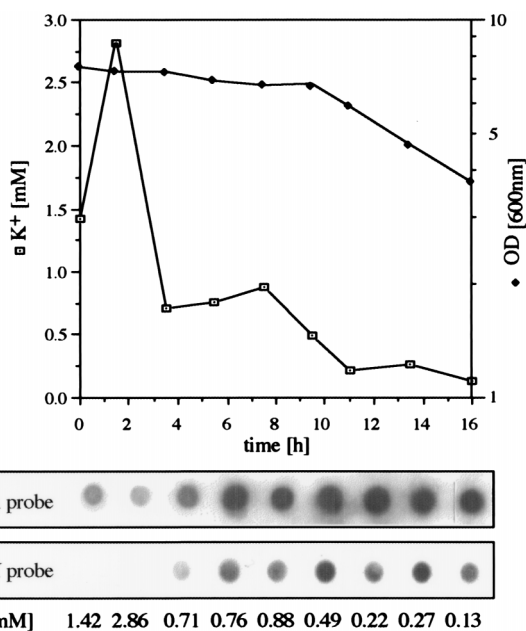


FIG. 7. Influence of decreasing the potassium concentration on growth in a continuous culture of *C. acetobutylicum*. Below, dot blot hybridization signals of RNA samples with *kdpA* and *kdpX* probes are presented. Potassium concentrations of the supernatants of the respective samples are indicated below the autoradiography signals.

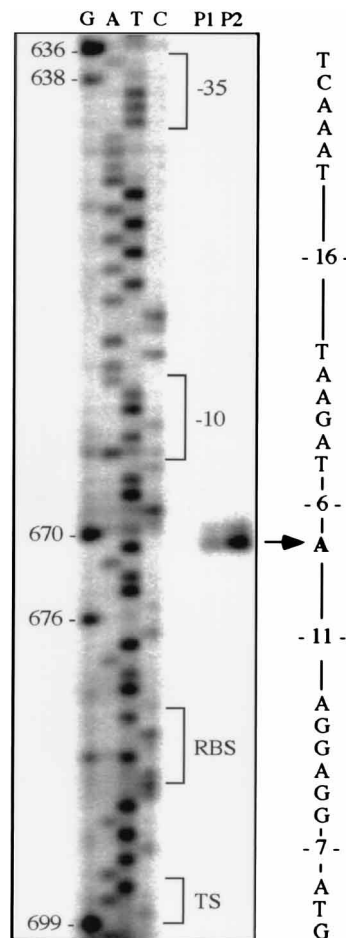
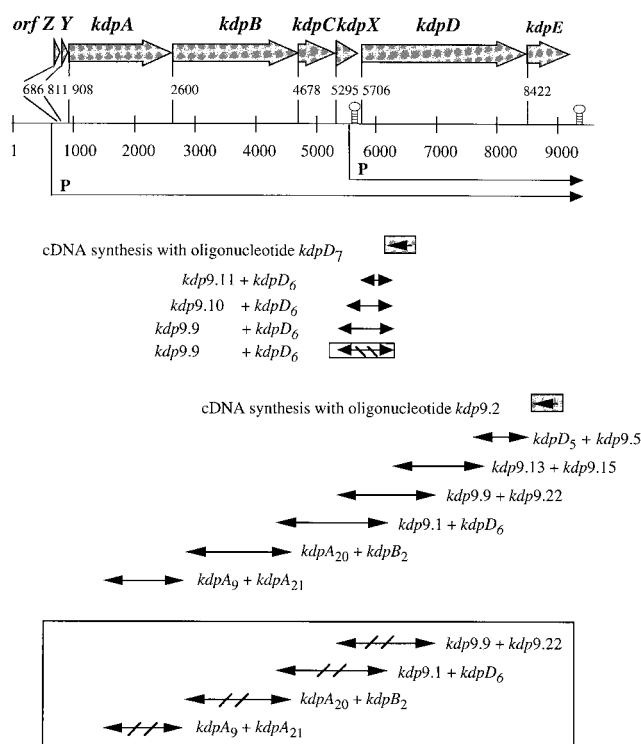


FIG. 8. Identification of the 5' end of the *C. acetobutylicum kdpA* transcript by primer extension. The primer extension products (P1 and P2) were run on a 6% polyacrylamide gel together with a sequencing reaction with the same primer (*orfZ<sub>p</sub>*). The sequence depicted at the right side indicates the proposed -10 and -35 promoter region, the ribosome-binding site, and the translation start; the arrow marks the transcription start point. Numbers on the left indicate the positions of the corresponding G nucleotide in the *kdp* sequence.

were washed and hybridized again with a gene probe of a constitutive gene (*ptb*, encoding phosphotransbutyrylase) of *C. acetobutylicum*. A distinct band of the expected size became visible in each lane of the blots (data not shown). The failure to determine the length of the *kdp* mRNAs by Northern blot hybridizations led to the usage of the more sensitive RT-PCR method.

Oligonucleotide *kdpD<sub>7</sub>* was hybridized to RNA of *C. acetobutylicum* cells grown in CBM medium (no detectable *kdpA* mRNA transcripts, thus designated uninduced RNA) and with RNA of the first limitation experiment (sample 7, detectable *kdpA* transcripts in cells growing at 170  $\mu$ M K<sup>+</sup>, thus designated induced RNA). The cDNAs produced (designated accordingly as uninduced and induced cDNA) were used as templates in three different PCR assays (Fig. 9). The oligonucleotides pairs *kdpD<sub>6</sub>-kdp9.11* and *kdpD<sub>6</sub>-kdp9.10* revealed the expected fragments of 541 and 757 bp with both cDNAs. On the other hand, using the oligonucleotide pair *kdpD<sub>6</sub>-kdp9.9*, we could amplify only the expected 997-bp fragment with the induced cDNA as a template (Fig. 9). PCR with the uninduced cDNA revealed no visible fragment. Oligonucleotide *kdp9.9* hybridized to a region upstream of the *kdpD*





promoter (52), indicating that the uninduced RNA contained only mRNA transcripts starting at the *kdpD* promoter. With both RNA samples, two other cDNAs were produced with oligonucleotide *kdp9.2*, which hybridized to the 3' end of the *kdpE* gene. Six different PCRs were performed with the produced cDNAs. PCR with the oligonucleotide pairs *kdpD*<sub>5</sub>-*kdp9.5* and *kdp9.13*-*kdp9.15* revealed the expected fragments of 974 and 1,710 bp. DNA amplifications with the oligonucleotide pairs *kdp9.9*-*kdp9.22*, *kdpD*<sub>6</sub>-*kdp9.1*, *kdpA*<sub>20</sub>-*kdpB*<sub>2</sub>, and *kdpA*<sub>9</sub>-*kdpA*<sub>21</sub> proved to be possible only with the induced cDNA, indicating that under induced conditions, transcription of the clostridial *kdp* genes did not stop at the putative terminator structure downstream of *kdpX* but continued with the transcription of the genes *kdpD* and *kdpE*. The uninduced RNA contained only mRNA transcripts of the *kdpD* and *kdpE* genes (initiated at the promoter upstream of *kdpD*), since all attempts to amplify DNA encoding the upstream *kdp* genes failed (Fig. 9). Although the results of the RT-PCR experiments showed clearly that a common *kdpZYABCXDE* transcript was synthesized under inducing conditions, it could not be excluded that some transcripts starting upstream of *kdpA* were terminated downstream of *kdpX*. To quantify the amounts of *kdpDE* transcripts and of *kdpZYABCXDE* or *kdpZYABCX* transcripts, we analyzed the above-described Northern blots, hybridized with a *kdpA* probe and a *kdpDE* probe. The two probes were almost equal in length and were used in the same concentration on the two identically prepared Northern blots. In both cases, the strongest hybridization signals were found with RNA sample 8 of the first limitation experi-

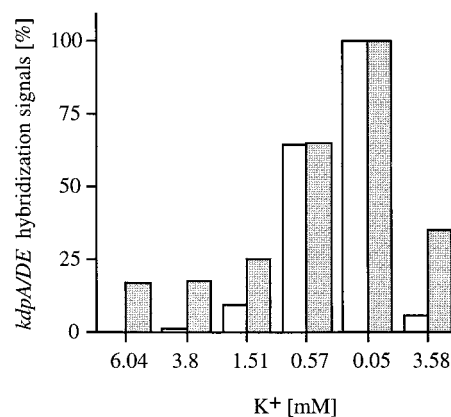


FIG. 10. Quantification of Northern blot hybridizations performed with a *kdpA* probe (white bars) and a *kdpDE* probe (grey bars) and different RNA samples from the first limitation experiment. The potassium concentrations of the supernatants of the respective samples are indicated below the bars.

ment (Fig. 6). With the thus obtained density values of the different lanes, a background subtraction including unspecific rRNA background was done. The new values ( $D$  [density] -  $B$  [background]) were finally corrected by division with the scanned area value ( $D - B/\text{pixel}^2$ ). Since the value of the strongest *kdpDE* hybridization signal with sample 8 was still higher than that of the *kdpA* hybridization signal with sample 8 ( $D - B/\text{pixel}^2 = 119.10$  with the *kdpDE* probe and 88.50 with the *kdpA* probe), both values were set as 100%, and the corresponding values of the other samples were accordingly calculated. The data are shown in Fig. 10.

In a second quantification experiment, three different oligonucleotides were used for dot blot hybridizations with three identically prepared membrane strips with 10  $\mu\text{g}$  of plasmid DNA (pT5 [Fig. 1]) as a hybridization control and two different RNA samples (first limitation, sample 7) and an RNA sample, prepared from a batch culture of *C. acetobutylicum* inoculated five times in K0 medium. The oligonucleotides were chosen to quantify the region upstream (primer *kdpD*<sub>2</sub>) and downstream (primer *kdpD*<sub>3</sub>) of the putative terminator in comparison with a region in *kdpC* (primer *kdpC*<sub>12</sub>). Data were obtained in terms of phosphostimulated luminescence and corrected as described above. With both RNA samples, we found weaker hybridization signals in the region upstream of the stem-loop structure than in the downstream region (Fig. 11).

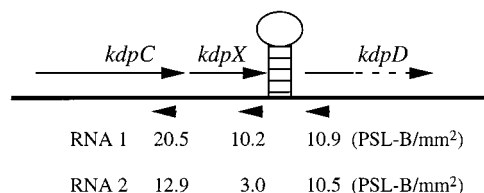


FIG. 11. Schematic representation of the *kdpCXD* region (arrows indicate the genes) with the stem-loop between *kdpX* and *kdpD*. The small leftward-directed arrows represent the oligonucleotides *kdpC*<sub>12</sub>, *kdpD*<sub>2</sub>, and *kdpD*<sub>3</sub>. The numbers represent the quantification results revealed by hybridization of these oligonucleotides with RNA 1 (first limitation, sample 7) and RNA 2 (K<sup>+</sup>-limited batch culture). PSL, phosphostimulated luminescence.

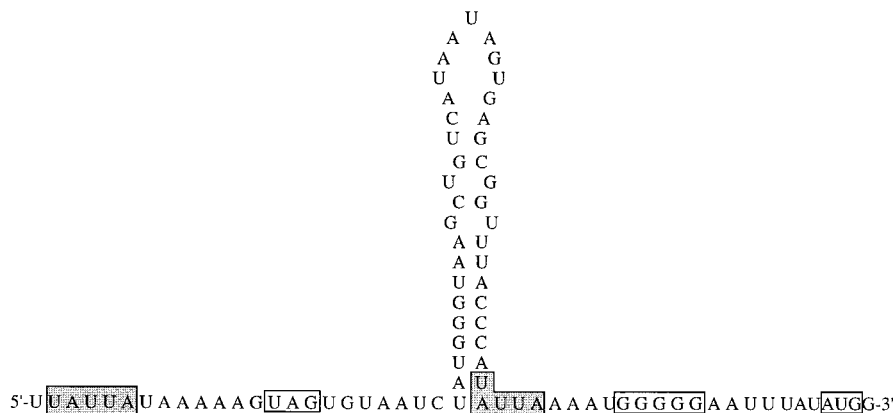


FIG. 12. Nucleotide sequence between the end of *kdpX* and the beginning of *kdpD*. The grey boxes indicate the putative recognition sites of RNase E. The white boxes represent the UAG stop codon of *kdpX*, the ribosome-binding site upstream of *kdpD*, and the AUG start codon of *kdpD*. For the stem-loop, a free energy of  $-18$  kJ/mol (26) was calculated.

## DISCUSSION

Recently, two regulatory genes of *C. acetobutylicum* have been cloned and sequenced, representing the first report on a sensor histidine kinase/response regulator system in this organism (52). Significant homology to the respective *E. coli* genes led to their identification as *kdpD* and *kdpE*. In this study, the remainder of the *kdp* system of *C. acetobutylicum* has been cloned and sequenced. Upstream of *kdpX*, a gene that has no counterpart in *E. coli*, *M. tuberculosis*, and *Synechocystis* sp. strain PCC 6803 and precedes *kdpDE*, homologous genes to *kdpA*, *kdpB*, and *kdpC* could be found. Upstream of *kdpA* are two small ORFs which might be part of the *kdp* system of *C. acetobutylicum* in that their deduced products resemble the KdpF peptide of *E. coli* (2) in size and hydrophathy. Thus, *C. acetobutylicum* possesses a *kdp* DNA region very similar to that of *E. coli*. With the exception of the two small ORFs (*orfZ* and *orfY*) instead of *kdpF* and an additional ORF (*kdpX*) between *kdpC* and *kdpD*, all *kdp* genes are conserved and organized accordingly. The clostridial KdpA comprises very hydrophobic regions and a conserved region which in *E. coli* is described as a putative periplasmic  $K^+$ -binding site, suggesting a role as a integral membrane  $K^+$ -transporting protein.

KdpB of *C. acetobutylicum* showed the typical features of P-type ATPases; the most conserved motifs of this family could be found in the clostridial amino acid sequence. The description for  $P_3$  ATPases (32) fits the clostridial KdpB in that KdpB probably is part of a protein complex consisting of KdpA, KdpB, KdpC, and possibly KdpX and does not function as a single unit like most other P-type ATPases (43). The clostridial KdpB thus represents only the second member of the  $P_3$  group, the enterobacterial protein being so far the only known representative. Kdp from *M. tuberculosis* and *Synechocystis* sp. strain PCC 6803, identified only by sequence similarity from genome sequencing projects, might be further candidates. On one hand, KdpB resembles the non-heavy-metal-transporting  $P_2$  ATPases because of a single pair of transmembrane fragments before the well conserved TGES sequence (24, 43, 44), which is thought to participate in vanadate and  $P_i$  binding (44) and is proposed to be responsible for phosphatase activity (44). On the other hand, the KdpB of *E. coli* is similar to the heavy-metal-transporting  $P_1$  ATPases because of the shorter C-terminal region (32). All these features are conserved in the clostridial KdpB.

The clostridial KdpC showed no high similarity to other

proteins except for KdpC of *E. coli*, *M. tuberculosis*, and *Synechocystis* sp. strain PCC 6803, and so its physiological function remains unclear. The putative membrane-spanning segment of the KdpC proteins near the N terminus is similar to the structure of the glycosylated  $\beta$  subunits of the  $\alpha/\beta$ -heterodimeric, cation-exchange ATPase subfamily (16) which includes the ubiquitous  $Na^+/K^+$ -ATPases and the gastric  $H^+/K^+$ -ATPases. For the  $\beta$  subunits, an involvement in  $K^+$ -dependent reactions of the enzymes such as  $K^+$  occlusion is suggested, and the extracellular domain might play an important role in determining the kinetics of  $K^+$  interaction (16). The deglycosylated core proteins of the glycosylated  $\beta$  subunits usually have molecular masses of ca. 34 kDa, whereas the  $M_r$  of the four KdpC proteins is in the range of 22,000.

The results of the different hybridization experiments together with the results of the in vitro transcription-translation experiments revealed that *kdpX* is indeed transcribed and translated. The similarity of the hydrophobicity plots of KdpC and KdpX of *C. acetobutylicum* and KdpC of *E. coli* make it tempting to speculate that in *C. acetobutylicum*, a heterodimer of KdpC and KdpX is formed in the active complex, whereas in *E. coli*, a homodimer of KdpC serves this function. This speculation might be supported by a possible function of the small hydrophobic peptides as a "glue," of which two (*OrfZ* and *OrfY*) have been found in *C. acetobutylicum*, whereas only one (KdpF) is present in *E. coli*.

Induction of the genes is clearly potassium dependent. Transcription occurred from two different promoters. In media of high potassium concentration, only the genes *kdpD* and *kdpE* are transcribed from the *kdpD* promoter (52). This transcription ensures a constitutive but very low expression of the *kdpDE* operon (52). As soon as the potassium concentration decreased and became a growth-limiting factor, transcription of the *kdp* genes started upstream of *orfZ*, and all *kdp* genes, including *kdpD* and *kdpE*, which encode for the *kdp* sensor histidine kinase/response regulator system, were transcribed together, as indicated by the results of the RT-PCR. Induction by lowering the potassium concentration obviously prevented transcription from the *kdpD* promoter, since under these conditions determination of the respective mRNA start point failed. This also explains why Northern hybridization with a *kdpDE* probe revealed the same dependence on the potassium concentration as experiments using a *kdpA* probe. In *E. coli*, similar experiments have not been performed. A promoter has

been identified upstream of *kdpD*, in the *kdpC* gene (40). Transcriptional fusions of *lacZ* to *kdp* genes indicated increased expression, attributed to read-through from transcription of *kdpFABC* (2). However, the authors assumed that about 90% of all transcripts, regardless of where they begin, terminate early in *kdpD* (40). In contrast, experiments aiming at quantitation of KdpD by antibodies clearly showed that the amount of KdpD was increased under potassium-limiting conditions (54). Thus, the situation in *E. coli* might be the same as in *C. acetobutylicum*, where the *kdp* genes are transcribed together with *kdpD* and *kdpE*. At first glance, it does not make sense that transcription of the regulator genes, whose products control expression of the Kdp system, is also increased by low potassium concentrations. However, it could be that KdpD and KdpE not only activate transcription of the *kdp* system but together serve as a master unit playing a pivotal role in general osmoregulation. Future experiments will try to solve this question.

Northern hybridizations revealed that the *kdp* mRNAs are very unstable, whereas other mRNA transcripts were identified on the same blots without problems. The fast degradation of *kdp* mRNA seems to be part of the complex regulation of this system. The common *kdpZYABCXDE* transcript is detectable only with RT-PCR; on a Northern blot, many degradation products in a range of 8,000 to 1,000 ribonucleotides become visible. The results of the different hybridization experiments indicate that the region upstream of the stem-loop structure between *kdpX* and *kdpD* is much more unstable than the regions within the *kdpA*, *kdpC*, and *kdpD* genes. In recent years, much progress has been made in identification of enzymes which are involved in mRNA degradation, and it seems generally accepted that mRNA turnover plays a central role in determining levels of protein synthesis (4, 9, 21). Many reports describe specific sequences, and structural features of an mRNA probably determine its stability (for examples, see references 5, 12, 15, 17, 37, 55, and 60). A recent review (18) summarizes features of RNase E, which is known to play a general role in mRNA decay. Naureckiene and Uhlin (37) describe RNase E as a single-strand-specific endoribonuclease and a high A/U content as a common feature of the cleaved sequences. A pentanucleotide consensus sequence (G/A)'A'UU(A/U) followed by a stable secondary structure has been proposed as the cleavage site of RNase E (21). Recent data have shown that RNase E can cleave oligoribonucleotides lacking a stem-loop structure (36), but it also has been suggested that stem-loops in the vicinity of the RNase E cleavage site could stabilize mRNA structure and maintain the single-strandedness of the cleavage site (19, 33). It has also been shown that destabilization of such a stem-loop structure reveals probably cryptic RNase E cleavage sites, suggesting a shielding role of these structures in RNase E recognition (37). Taking into account all of these recent findings, it is possible to propose an RNase E cleavage site in the clostridial nucleotide sequence between the end of *kdpX* and the beginning of *kdpD* (Fig. 12). In accordance with the consensus sequence prepared by Ehretsmann et al. (21), we found two possible cleavage sites, one single stranded and the other partially folded in the stem-loop. A cleavage of the mRNA transcript at the single-stranded site would explain the weaker hybridization signals of the *kdpX* probes in comparison to those of *kdpA* and *kdpD*. The downstream cleavage product would be more stable than the upstream mRNA because of the stem-loop structure, whereas the free 3' end of the *kdpZYABCXDE* transcript could be degraded by the action of 3'-to-5' exonucleases. A cleavage of the mRNA at this position could also be possible in the *kdpDE* transcript, which is formed under uninducing conditions, since

none of the translation signals would be deleted by this cleavage (Fig. 12). Nevertheless, this RNase E cleavage model is only a speculation since we know nothing about RNA-degrading enzymes in *C. acetobutylicum*.

For the DNA region upstream of *kdpA* in *E. coli*, a KdpE-binding sequence, a 21-bp target site of the *trans*-acting activator protein KdpE, has been described as an essential sequence for activation of the *kdpA* promoter. Five stretches of T's occur periodically around this bending center (51). The distance between the -35 region of the *kdpA* promoter and the KdpE-binding sequence is 9 bp. The KdpE-binding sequence was shown to be 5'-TTTATACTTTTTTACACCC-3'. Upstream of the clostridial *orfZY/kdpA* promoter, a similar region with the sequence 5'-TTTATACTTTCTTAACACCTG-3' and a distance of 16 bp to the -35 box was identified. However, stretches of T's are common in DNA of *C. acetobutylicum*, since this organism has a G+C content of 28 to 29 mol% (6). Thus, it remains to be determined whether this region represents a KdpE-binding site.

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