

Identification of Essential Glutamates in the Acetate Kinase from *Methanosarcina thermophila*

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Acetate kinase catalyzes the reversible phosphorylation of acetate ($\text{CH}_3\text{COO}^- + \text{ATP} \rightleftharpoons \text{CH}_3\text{CO}_2\text{PO}_3^{2-} + \text{ADP}$). A mechanism which involves a covalent phosphoryl-enzyme intermediate has been proposed, and chemical modification studies of the enzyme from *Escherichia coli* indicate an unspecified glutamate residue is phosphorylated (J. A. Todhunter and D. L. Purich, *Biochem. Biophys. Res. Commun.* 60:273–280, 1974). Alignment of the amino acid sequences for the acetate kinases from *E. coli* (Bacteria domain), *Methanosarcina thermophila* (Archaea domain), and four other phylogenetically divergent microbes revealed high identity which included five glutamates. These glutamates were replaced in the *M. thermophila* enzyme to determine if any are essential for catalysis. The histidine-tagged altered enzymes were produced in *E. coli* and purified to electrophoretic homogeneity by metal affinity chromatography. Replacements of E384 resulted in either undetectable or extremely low kinase activity, suggesting E384 is essential for catalysis which supports the proposed mechanism. Replacement of E385 influenced the K_m values for acetate and ATP with only moderate decreases in k_{cat} , which suggests that this residue is involved in substrate binding but not catalysis. The unaltered acetate kinase was not inactivated by *N*-ethylmaleimide; however, replacement of E385 with cysteine conferred sensitivity to *N*-ethylmaleimide which was prevented by preincubation with acetate, acetyl phosphate, ATP, or ADP, suggesting that E385 is located near the active site. Replacement of E97 decreased the K_m value for acetate but not ATP, suggesting this residue is involved in binding acetate. Replacement of either E32 or E334 had no significant effects on the kinetic constants, which indicates that neither residue is essential for catalysis or significantly influences the binding of acetate or ATP.

Acetate is an end product of most fermentative microbes and is the major growth substrate for the methanoarchaea (22); thus, carbon flow through acetate is of primary importance in anaerobic microbial consortia and the global carbon cycle. Although the metabolisms of fermentatives and acetotrophic methanoarchaea represent the extremes of biochemical divergence in energy-yielding pathways, these microbes have in common the enzymes acetate kinase (reaction 1) and phosphotransacetylase (reaction 2).



These two enzymes produce acetate from acetyl coenzyme A (acetyl-CoA) in the fermentatives, where a major portion of the energy requirements is obtained through substrate-level phosphorylation catalyzed by acetate kinase. The methanosarcinas utilize acetate kinase and phosphotransacetylase to activate acetate to acetyl-CoA in the first step of the pathway for the fermentation of acetate to methane (5). The acetyl-CoA is cleaved by the CO dehydrogenase–acetyl-CoA synthase enzyme complex, yielding methyl and carbonyl groups (5). The methyl group is reduced to CH_4 with electrons derived from oxidation of the carbonyl group to CO_2 . Transport of electrons through a membrane-bound transport chain generates an elec-

trochemical ion gradient driving ATP synthesis. Thus, acetate kinase and phosphotransacetylase are at the interface of energy-yielding metabolism between fermentatives and the acetotrophic methanoarchaea, which are the principal metabolic groups in anaerobic consortia degrading complex organic matter to methane. In addition to a key intermediate in energy metabolism, acetylphosphate acts as a phosphoryl donor to enzyme I of the phosphoenolpyruvate:glucose phosphotransferase system in *Escherichia coli* and *Salmonella typhimurium* via a phosphoenzyme intermediate of acetate kinase (7). Acetylphosphate is also a phosphoryl donor to periplasmic binding proteins (8) as well as to many response regulator proteins of two-component systems (13). It has been proposed that acetylphosphate functions as a global regulatory signal in *E. coli* (13, 20). Thus, acetate kinase influences the physiology of diverse microbes in a variety of ways.

Comparison of deduced amino acid sequences reveals high identity among acetate kinases (Fig. 1) from the Archaea and Bacteria, which suggests that this enzyme was either highly evolved prior to divergence of the domains or that horizontal gene transfer occurred between domains. The high identity suggests similar mechanisms for these acetate kinases. The acetate kinase from *E. coli* is phosphorylated with acetylphosphate or ATP (3, 6, 19), and it has been proposed that a phosphoryl-enzyme intermediate is involved in the catalytic mechanism. Acid hydrolysis of the phosphorylated enzyme reduced with [^3H]borohydride yields [^3H] α -amino- δ -hydroxyvaleric acid, suggesting that an unspecified γ -phosphorylated glutamyl residue is involved in catalysis (19). This hypothesis is novel, as there are no reports of a phosphorylated glutamate in

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M. t.	---MKVLVINAGSSSLKYQLIDMTNESALAVGLCERIGIDNSIITQKKFDGKKLEKLTDLPTHKDALEEVVKALTDDEFGVIKDMGE	84
E. c.	MSSKLVVLNCGSSSLKFAIDAVNGEYLSGLAECFHLPEARIKWKMDGNKQEAALGAGAAHSEALNFIVNTILAQ---KPELSAQ	84
C. a.	---MKNLVINCSSSIKYQFIDMKDETVLAKGLVERIGIKGSVITHKVN-GEKYVTETPMEDHKRAIKLVLDALLNDEYGVIKNIDE	83
B. s.	--MSKIIAINAGSSSLKQFLEMPSETVLTKGLVERIGIADSVFTISVN-GEKNTFVTDIPDHAVAVKMLLNKLT--EFGI IKDLNE	82
H. i.	-MSKLVILNCGSSSLKFAILD PATGEEKLSGLAEAFFLPEARIKWKLNAGEKGNADLGAGAAHTEALNF IASNILN-----DELKNS	81
M. g.	MQSHKILVVNAGSSSIKQFLEFNDK-KQVLAKGLCERIFIDG-FFKLEFN-QKKIEEKVQFNDHNLA VKHFLNALKKNK--IITELSE	82
M. t.	INAVGHRVVGGEKFTTSALYDEGVEKAIKDCFELAPLHNPPNMGISACAEIMPOTP--MVIVFDTAFHQTMPYAYMYALPYDLY	169
E. c.	LTAIGHRIVHGGEKYTSSVVIDESVIQGIKDAASFAPLHNPAHLIGIEEALKSFPQLKDKNVAVFDTAFHQTMPEESYLALPYNLY	171
C. a.	ISAVGHRIVHGGEKYANSVLIDEDVMKSIEDCVSLAPLHNPPHIGINACKELMPNVF--MVAVFDTAFHQTI PDYAYMYAIPYEYY	168
B. s.	IDGIGHRVVHGGEKFSDSVLLTDETIKEIEDISELAPLHN PANIVG IKA FKEVLPNVF--AVAVFDTAFHQTMPEQSYLYSLPYEYY	167
H. i.	IAAIGHRIVHGGEKYTQSVIVTDEVVKGIEDAAQFAPLHNPAHLIGIREAFKAPHLKDKNVVFDTAFHQTMPEEAFLYALPYSLY	168
M. g.	IGLIGHRVVQGANFYTDAVLVDTHSLAKIKEFIKLAPLHNKPEADVIEIFLKEIKTAK--NVAVFDTTFTTIPRENYLYAVPENWE	167
M. t.	KEHGVRKYGFHGTSHKYVAERAALMLGKPAEETKIITCHLGNSSITAVEGGKSVETSMGFTTLEGLAMGTRCGSIDPAIVPFLMEK	256
E. c.	KEHGIRRYGAHGTSHFYVTQEAAKMLNKPVEELNIITCHLGNSSVSAIRNGKCVDTSMGLTPLEGLVMGTRSGDIDPAIIFHLHDT	258
C. a.	DKYKIRKYGFHGTSHKYVSR TAAEFIGKKVEDLKMVVCHMNGASITAVEGGKSVDTSMGFTT PPGGLAMGARS GMDMPAVVTF LMDK	255
B. s.	EKFGIRKYGFHGTSHKYVTERAAELLGRPLKDLRLISCHLGNASIAAVEGGKSIDTSMGFTTLAGVAMGTRSGNIDPALIPYIMEK	254
H. i.	KEHGVRRYGAHGTSHYFVSREVAKYVGPADQVNAIICHLGNSSVSVVRNGQCIDTSMGLTPLEGLVMGTRCGDIDPAIVFYLYKT	255
M. g.	KNNLVRRYGFHGTSYKYINEFLEKKNK--KPLNLIVCHLGNASVCAIKQKSLNTSMGFTTLEGLIMGTRSGDIDPAIVSYIAEQ	252
M. t.	EGLTTREIDTLMNKKSGVLGVSGLSNDFRDLDEAASKG-NRKAELALEIFAYKVKKFIGEYSAVL--NGADAVVFTAGIGENSASIR	340
E. c.	LGMSVDAINKLLTKESGLLGLTEVTSDCRYVEDNY--ATKEDAKRAMDVYCHRLAKYIGAYTALM-DGRLDAVVFTGGIGENAMVR	342
C. a.	LNINASEVNNLLNKKSGIEGLSGISSDMRDIKKNYVDKDPKAMLAYSVFTYKIKQFIGSYTAVM--NGLDCLVFTGGIGENSFENR	340
B. s.	TGQTAEVNLNTLNKKSGLLGISGFSSDLRDI VEATKEG-NERAETALEVFASRIHKYIGSYAARM--SGVD AII FTAGIGENSVEVR	338
H. i.	LGMSMDQIEETLVKKSGLLGLTEVTSDCRYAEDNYDDESKPETRRALNVYSYRLAKYIGAYMAVLGDDHLDAIAFTGGIGENSAHVR	342
M. g.	QKLSNDVNVNELNKKSGMFAITG-SSDMRDI FDKP-----EINDIAIKMYVNRVADYIAKYLNQLS-GEIDSLVFTGGVGENASYCV	332
M. t.	KRILTGLDGIGIKIDDEKN--KIRGQEIDISTPDAKVRVFIPTNEELAIARETKEIVETEVLKRSSIPV	408
E. c.	ELSLGKLVGLGFEVDHERNLAARFGKSGFINKEG-TRPAVVIPITNEELVIAQDASRLTA-----	400
C. a.	REICKNMDYLGKIKIDDKND-ETMGIPMDISAEKSVRVLVFIPTNEELMIARDTKDIVGKLK-----	401
B. s.	ERVLRGLEFMGVYWDPALN--NVRGEEAFISYPHSPVKVMIIPITDEEVM IARDVVR LAK-----	395
H. i.	ELALNHLKLFGIKIDNERNLATRFKGDGVITDSDAFKAI VLPITNEELVIAQDTAKLCF-----	401
M. g.	QLIIEKVASLGFKTNSNLFG--NYQDSSLISTNESKYQIFRVRTNEELMIVEDALRVSTNIKK-----	393

FIG. 1. Alignment of the deduced amino acid sequences of acetate kinases. Abbreviations and GenBank accession numbers: M.t., *M. thermophila* (L23147); E.c., *E. coli* (M22956); C.a., *Clostridium acetobutylicum* (U38234); B.s., *Bacillus subtilis* (L17320); H.i., *Haemophilus influenzae* (L45839); M.g., *Mycoplasma genitalium* (L43967). Amino acids are in the single-letter code. Dashes represent gaps introduced for alignment. Closed circles mark the glutamate residues targeted for replacement. Identical residues are shown in bold.

any enzyme reaction mechanism. The proposed mechanism was challenged when it was shown that phosphorylated acetate kinase from *E. coli* is a phosphoryl donor to enzyme I of the bacterial phosphotransferase system (7), suggesting the possibility of an alternate function for phosphorylated acetate kinase. Furthermore, no evidence has been reported for a glu-

tamate which is essential for kinase activity. Thus, the involvement of glutamate in catalysis of kinase activity by acetate kinases is a matter of controversy.

Despite the broad importance of acetate kinase in microbial physiology, the use of biochemical genetics to probe the catalytic site has not been reported. Here we identify an essential

TABLE 1. Plasmids used in this study

Plasmid	Origin	Genotype	Reference or source
pUC19/ <i>ack</i>	pUC19	<i>ack</i>	12
pTZ18U			Bio-Rad
pTZ <i>ack</i>	pTZ18U	<i>ack</i>	This study
pET15b			Novagen
pET <i>ack</i>	pET15b	<i>ack</i>	This study
pET <i>ack</i> E32A	pET15b	<i>ack</i> _{E32A}	This study
pET <i>ack</i> E97A	pET15b	<i>ack</i> _{E97A}	This study
pET <i>ack</i> E97D	pET15b	<i>ack</i> _{E97D}	This study
pET <i>ack</i> E97Q	pET15b	<i>ack</i> _{E97Q}	This study
pET <i>ack</i> E334A	pET15b	<i>ack</i> _{E334A}	This study
pET <i>ack</i> E384A	pET15b	<i>ack</i> _{E384A}	This study
pET <i>ack</i> E384D	pET15b	<i>ack</i> _{E384D}	This study
pET <i>ack</i> E384Q	pET15b	<i>ack</i> _{E384Q}	This study
pET <i>ack</i> E385A	pET15b	<i>ack</i> _{E385A}	This study
pET <i>ack</i> E385C	pET15b	<i>ack</i> _{E385C}	This study
pET <i>ack</i> E385D	pET15b	<i>ack</i> _{E385D}	This study
pET <i>ack</i> E385Q	pET15b	<i>ack</i> _{E385Q}	This study

glutamate in the *Methanosarcina thermophila* enzyme which provides support for the previously proposed catalytic mechanism. Two other glutamates are implicated in substrate binding.

MATERIALS AND METHODS

Protein sequence analyses. Databases were searched at the National Center for Biotechnology Information by using the BLAST network server (2). CLUSTAL W (18) was used for multiple protein sequence alignment at the Human Genome Center of the Baylor School of Medicine.

Site-directed mutagenesis. Mutagenesis was performed according to the manufacturer's instructions with the Muta-Gene phagemid mutagenesis kit (Bio-Rad), which employs the oligonucleotide-directed in vitro mutagenesis method (11). The plasmids used are shown in Table 1. The *M. thermophila* acetate kinase gene was excised from pUC19/*ack* (12) by *Kpn*I and *Bam*HI digestion and ligated into pTZ18U to generate pTZ*ack*. The mutations were verified by double-stranded sequence analysis by using the dideoxy chain termination method (15) and Sequenase version 2.0 (United States Biochemicals).

Heterologous production and purification of acetate kinase. The unaltered and altered acetate kinase genes were subcloned into the T7-based expression vector pET15b (Novagen) to generate the plasmids listed in Table 1. In these plasmids, a 60-nucleotide leader sequence with six tandem histidine codons was fused in frame to the 5' end of the unaltered and altered *ack* genes. *E. coli* BL21(DE3) was transformed with the expression vectors, inoculated into 50 ml of Luria-Bertani medium containing 100 µg of ampicillin per ml, and grown at 37°C to an A_{600} of 0.6 to 0.9, at which time IPTG (isopropyl-β-D-thiogalactopyranoside) was added to a final concentration of 1 mM. After 1.5 to 2.0 h of induction, the cells were harvested and stored at -70°C. The unaltered and altered acetate kinases were purified using a Ni-nitrilotriacetic acid silica spin kit (Qiagen) according to the manufacturer's instructions. The enzymes were eluted in buffer (pH 7.0) containing 50 mM NaH₂PO₄, 300 mM NaCl, and 250 mM imidazole. Protein concentrations were determined by the Bradford method (4), using protein dye reagent (Bio-Rad) and bovine serum albumin as the standard.

Enzyme activity assays. Acetate kinase activity was determined by the previously described (1) standard hydroxamate assay, which detects the formation of acetyl phosphate from acetate (200 mM) and ATP (10 mM). The acetate concentrations were 0.7 and 1.5 M for determination of the kinetic constants for ATP for the E385A and E385D enzymes.

The ATP-ADP exchange assays were performed as follows. The reaction mixture (100 µl) contained the following: triethanolamine-HCl, 50 mM, pH 7.0; ATP, 1 mM; ADP, 1 mM; MgCl₂, 2 mM; acetate kinase, 10 µg; [α-³²P]ATP (3,000 Ci/mmol), 5 µCi. The reactions were initiated by the addition of enzyme, and mixtures were incubated at 20°C. At various times, 5 µl was withdrawn, and the reaction was stopped by the addition of an equal volume of 1 N HCl. A volume of 5 µl was applied to polyethyleneimine cellulose thin-layer chromatography plates (Bakerflex), which were developed ascendingly in 0.52 M potassium phosphate (pH 3.5). The plates were air dried and autoradiographed to visualize the ATP and ADP spots. The ADP spots were excised and counted in 5 ml of Scintiverse (Fisher Scientific) scintillation cocktail. Initial velocities were calculated as described previously (16).

Inhibition by NEM. *N*-Ethylmaleimide (NEM) (final concentration, 10 µM) was added to the unaltered acetate kinase (5 µg/ml) and to the E385C enzyme (40 µg/ml) in a final volume of 100 µl at 37°C. Aliquots (10 µl) were removed at

the indicated times and assayed for kinase activity. Substrate protection experiments were performed by preincubating the enzyme for 5 min at 37°C with (final concentrations) ADP (10 mM), ATP (10 mM), acetyl phosphate (10 mM), or potassium acetate (200 mM) prior to addition of NEM.

Circular dichroism spectroscopy. Spectra were acquired at 37°C with an Aviv circular dichroism (CD) spectrometer, model 62DS. Samples (1 to 10 µM) of acetate kinase in 20 mM sodium phosphate (pH 7.5) containing 0.1 M NaCl were placed in a cuvette with a 1-mm path length and data points obtained from 205 to 320 nm in 1.0-nm increments. Five spectra were taken for each sample and averaged. The resulting spectra were normalized for direct comparison.

RESULTS AND DISCUSSION

The sequences of acetate kinases deduced from the genes of widely divergent microbes from the *Archaea* and *Bacteria* domains are highly identical, suggesting a similar catalytic mechanism (Fig. 1); for example, the enzymes from *E. coli* (*Bacteria*) and *M. thermophila* (*Archaea*) have 44% identity. It has been proposed that a phosphorylated glutamate functions in a covalent catalytic mechanism for the *E. coli* enzyme (19). Five glutamates in the *M. thermophila* acetate kinase were selected for replacement to determine if any are essential for catalysis. The glutamates selected (Fig. 1) are either highly conserved and within a region of high conservation (E97) or 100% conserved (E32, E334, E384, and E385) with the deduced sequences for five other acetate kinases from extremely diverse microbes. The histidine-tagged altered enzymes were heterologously produced in *E. coli* and purified by one-step metal affinity chromatography. Each was judged to be homogeneous by sodium dodecyl sulfate-polyacrylamide gel electrophoresis, and the subunit molecular masses were indistinguishable from that of the unaltered histidine-tagged acetate kinase (not shown). Native gel filtration chromatography (not shown) indicated that the altered enzymes were dimeric, in accord with the authentic acetate kinase purified from *M. thermophila* (1). The unaltered histidine-tagged acetate kinase displayed a k_{cat} (Table 2) slightly greater than the value previously determined for the authentic enzyme (1) and was most likely a result of the one-step purification procedure yielding more active enzyme. There were no significant differences between the unaltered (Table 2) and authentic acetate kinases with respect to the K_m values for acetate or ATP.

Replacement of E384 and E385. Among the five glutamates targeted (Fig. 1), all except the enzymes altered at E384 had substantial activity (Table 2). Activity in the standard assay was undetectable (less than 1.0 µmol/min/mg of protein) for the E384D and E384Q enzymes. The E384A enzyme had detectable but extremely low specific activity (4.0 ± 2.0 µmol/min/mg of protein) in the standard assay, which was only 0.5% of the specific activity for the unaltered acetate kinase assayed under the same conditions. The low activity precluded reliable determinations of kinetic constants. These results suggest that E384 is the only essential glutamate. This glutamate is 100% conserved with all other deduced sequences for acetate kinases from widely divergent organisms (Fig. 1), which is consistent with an essential function. The CD spectra of the E384Q enzyme (Fig. 2) and the E384D enzyme (not shown) were similar to that of the unaltered acetate kinase. The ellipticity values at 210 nm for the E384Q and E384D enzymes were 3.6 and 1.2% less negative than that for the unaltered acetate kinase. This result suggests no global conformational change on substitution of E384 with Q or D, which is consistent with the high recovery on purification of the enzymes altered at E384 and suggests that they were dimeric in accord with the authentic enzyme purified from *M. thermophila*. These results, combined with the loss of kinase activity for the enzymes altered at E384, suggest that E384 is essential for catalysis, which is consistent with the previously proposed covalent mechanism in which a

TABLE 2. Kinetic constants of unaltered and altered acetate kinases^a from *M. thermophila*

Enzyme	Acetate			ATP		
	K_m (mM)	k_{cat} (s ⁻¹)	k_{cat}/K_m (mM ⁻¹ s ⁻¹)	K_m (mM)	k_{cat} (s ⁻¹)	k_{cat}/K_m (mM ⁻¹ s ⁻¹)
Authentic ^b	22	1,050	47	2.8	1,221	436
Unaltered	19 ± 3	1,596 ± 140	84	1.0 ± 0.3	1,669 ± 288	1,669
E32A	26 ± 2	1,029 ± 97	40	1.1 ± 0.2	1,012 ± 262	920
E97A	2.3 ± 0.5	165 ± 3	72	1.4 ± 0.1	302 ± 49	216
E97D	7.2 ± 3.5	430 ± 60	60	1.4 ± 0.6	567 ± 175	405
E97Q	4.0 ± 0.4	273 ± 56	68	1.4 ± 0.7	276 ± 59	197
E334A	15 ± 6	931 ± 35	62	0.7 ± 0.2	1,198 ± 62	1,711
E385A	86 ± 30 ^c	636 ± 158 ^c	7	6.9 ± 0.5	1,308 ± 112	190
E385C	17 ± 4	637 ± 139	37	1.6 ± 0.5	813 ± 214	508
E385D	337 ± 81 ^c	456 ± 132 ^c	1	4.5 ± 0.5	671 ± 34	149
E385Q	44 ± 19	1,869 ± 479	42	2.0 ± 0.6	1,711 ± 231	856

^a Unaltered and altered acetate kinases were produced in *E. coli*.

^b Published values for the acetate kinase purified from *M. thermophila* (1) are given.

^c Apparent value.

glutamate residue is γ -phosphorylated; however, further experimentation is necessary to prove that E384 is phosphorylated and that the phosphorylated enzyme is kinetically competent to phosphorylate acetate. The complete loss of activity for the E384D enzyme indicates that displacement of the active-site nucleophile by a single methylene carbon is sufficient to completely disrupt catalysis.

Although the features of the CD spectra were identical, the difference at 210 nm between the unaltered acetate kinase and the E384A enzyme was 14% less negative (not shown), suggesting the possibility that this replacement may have produced secondary structural changes. The thermostability of the E384A kinase was greater than that of the unaltered acetate kinase when the enzymes were preincubated either with or without ATP (Fig. 3A), a result which is consistent with changes in secondary structure. The temperature stability profile for the unaltered acetate kinase is shifted approximately 27°C higher in the presence of ATP (Fig. 3A). A similar shift was observed on preincubation of the E384A enzyme with ATP (Fig. 3B), suggesting that ATP interacts with the altered enzyme in the same way as it does with the unaltered acetate kinase. This result is unlikely if substitution of E384 with alanine produced conformational changes in the active site which abolished activity; however, this possibility cannot be ruled out. Nonetheless, these results do not detract from the results obtained for the E384Q and E384D enzymes which indicate that E384 is essential for catalysis. The small amount of activity

recorded for the E384A enzyme is unexplained; however, it is theoretically possible that replacement with alanine may have increased the accessibility of substrate to E385, which functioned as an active-site nucleophile in place of E384.

Mechanistically, an exchange of phosphate between ATP and ADP predicts a phosphoryl-enzyme intermediate; thus, linking exchange activity with kinase activity would provide

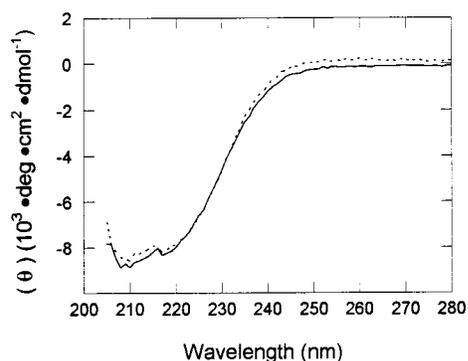


FIG. 2. CD spectra of unaltered (dotted line) and E384Q (solid line) acetate kinase from *M. thermophila*.

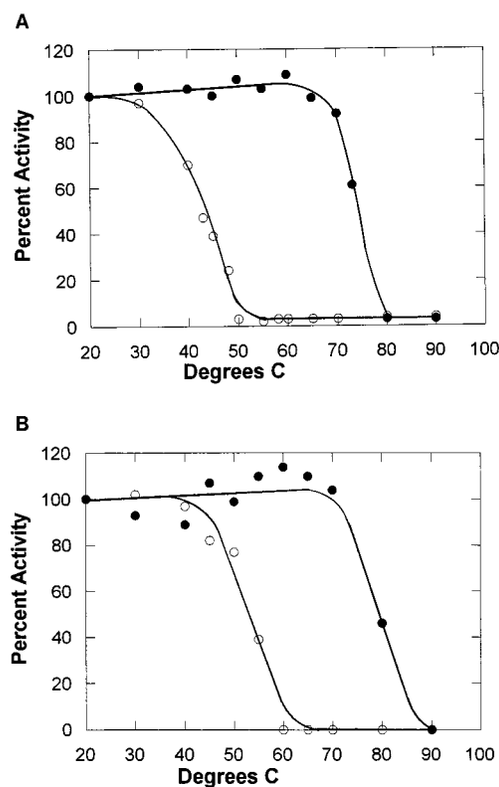


FIG. 3. Thermostability profiles of unaltered and E384A acetate kinase from *M. thermophila*. The enzyme preparations were preincubated without (○) and with (●) 10 mM ATP (final concentration) after which samples were incubated for 15 min at the indicated temperatures, cooled to 4°C, and immediately assayed for kinase activity. (A) Unaltered acetate kinase. One hundred percent activity was 470 μ mol/min/mg of protein. (B) E384A acetate kinase. One hundred percent activity was 4.0 μ mol/min/mg of protein.

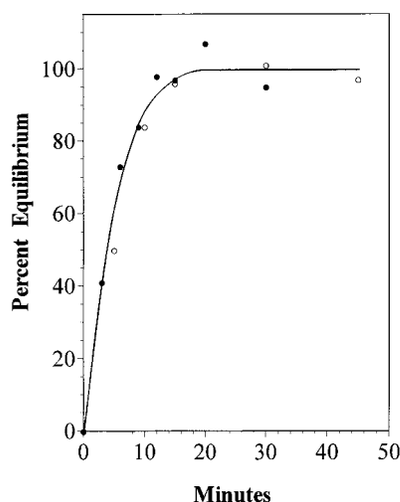


FIG. 4. ATP-ADP exchange activity of acetate kinase from *M. thermophila*. Activity was determined for purified histidine-tagged (○) and untagged (●) *M. thermophila* acetate kinase produced in *E. coli*. The ADP fractions contained 35,700 (○) and 92,000 (●) cpm at equilibrium.

evidence that both activities involve the same intermediate. The authentic and heterologously produced acetate kinase from *M. thermophila* catalyzed an exchange of phosphate between ATP and ADP (Fig. 4), as reported for other acetate kinases (10, 17). Although the initial rate of exchange (250 ± 40 nmol/min/mg of protein) was nearly 3,000-fold lower than kinase activity (705 ± 70 μ mol/min/mg of protein) assayed under the same conditions, the result does not rule out the possibility that a phosphoryl-enzyme intermediate is required for kinase activity. Unknown determinants such as substrate synergism may contribute to relatively low activity of the partial reaction. The previously reported ATP-ADP exchange activity of the *E. coli* enzyme is consistent with a phosphoryl-enzyme intermediate; however, it was not possible to exclude the possibility that kinase activity occurred by a different mechanism (14). Likewise, it cannot be concluded from the exchange activity alone that kinase activity for the enzyme from *M. thermophila* involves a phosphoryl-enzyme intermediate; however, the loss of kinase activity on replacement of E384 was paralleled by the inability to detect significant (limit of detection, 10 nmol/min/mg of protein) ATP-ADP exchange activity for any of the enzymes altered at E384 (data not shown), which is consistent with a linkage between the two activities centering on E384.

Further support for involvement of E384 in the active site of the acetate kinase from *M. thermophila* was obtained by replacing the neighboring E385 residue. E385 is 100% conserved with the acetate kinase sequences from diverse microbes (Fig. 1); however, other residues with different functionality replaced E385 (Table 2) without the reduction in activity observed for the enzymes altered at E384, suggesting that E385 is not essential for catalysis. Nonetheless, replacement of E385 with alanine or aspartate significantly increased the K_m values for acetate and ATP, suggesting that this residue is important for binding these substrates. However, the kinetic constants for the E385Q enzyme were not significantly different from those for the unaltered acetate kinase, suggesting that the negative charge of E385 is inconsequential for substrate binding. The unaltered acetate kinase from *M. thermophila* was not inactivated by NEM (Fig. 5); however, the E385C enzyme was in-

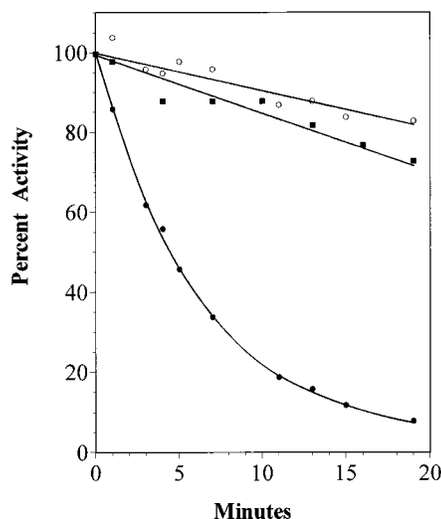


FIG. 5. Time course for NEM inactivation of unaltered and E385C acetate kinase from *M. thermophila*. Kinase activity was determined at the indicated times after addition of NEM to unaltered (■) or E385C (●) acetate kinase and compared to a control where no inhibitor was added to the E385C enzyme (○).

activated in a time-dependent manner, and preincubation with ATP, ADP, acetate, or acetyl phosphate protected against the inactivation (Fig. 6), which suggests that E385 is near the active site.

The acetate kinase isolated from *E. coli* is inactivated by NEM, and the inactivation is protected by preincubation with either ATP or ADP, suggesting that a cysteine residue is located near the active site (21). The inhibition is incomplete, and, therefore, it was concluded that the cysteine is not involved in catalysis. Only one cysteine residue is conserved between the *M. thermophila* and *E. coli* acetate kinases (Fig. 1); thus, either the *M. thermophila* enzyme has no cysteine residue corresponding in location to the modified residue in the *E. coli* enzyme or the only conserved cysteine residue in the *M. ther-*

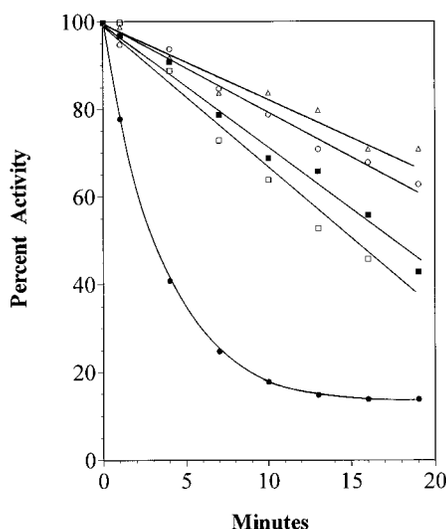


FIG. 6. Substrate protection from NEM inactivation of the E385C-altered acetate kinase from *M. thermophila*. Kinase activity was determined at the indicated times after addition of NEM to the E385C acetate kinase preincubated with ADP (□), ATP (△), acetate (○), acetyl phosphate (■), or no substrate (●).

mophila enzyme (C207) is shielded from attack by NEM. Preincubation with acetate or acetyl phosphate does not protect the *E. coli* acetate kinase from NEM inactivation; however, the pattern by which these substrates prevent protection by AMP, ADP, or ATP suggests that the binding sites for acetate and acetyl phosphate are in close proximity to the phosphoryl binding regions of ADP and ATP (21). The ability of all four substrates to prevent NEM inactivation of the *M. thermophila* E385C acetate kinase (Fig. 6) is consistent with the results obtained for the *E. coli* enzyme.

Replacement of E97. Compared with values obtained with the unaltered acetate kinase, the K_m values for acetate of all the E97-altered enzymes decreased with corresponding decreases in k_{cat} , which resulted in relatively smaller differences in the catalytic efficiencies (k_{cat}/K_m) (Table 2). These results suggest that E97 is not essential for catalysis but influences the binding of acetate. Replacement of E97 with aspartate was the least effective in lowering the K_m for acetate, suggesting that the negative charge of E97 influences binding of the acetate anion by charge repulsion. The results also suggest that displacement of the negative charge by only one methylene carbon significantly influences the binding of acetate. None of the E97 replacements influenced the K_m values for ATP, suggesting no involvement of E97 at the ATP binding site.

The K_m for acetate of the unaltered acetate kinase is relatively high, which appears to place *M. thermophila* at a disadvantage when competing for substrate; thus, the ability to decrease the K_m nearly 10-fold by simple replacement of a single residue suggests that the enzyme evolved to prefer a high K_m , which may be physiologically significant. Indeed, *M. thermophila* and other *Methanosarcina* species predominate in environments where the acetate concentrations are in the millimolar range compared with species from the acetotrophic genus *Methanotrix*, which thrive in habitats with acetate concentrations well below 1 mM (22). The acetotrophic *Methanosarcina* species outcompete *Methanotrix* species with higher growth rates and greater conservation of energy. The acetate-activating enzyme acetate thiokinase from *Methanotrix soehngenii* has a K_m for acetate below 1 mM, which allows growth of this species in habitats with low concentrations of acetate where methanosarcinas are unable to compete (9). Apparently, the *M. thermophila* acetate kinase evolved to favor a high turnover at the expense of a low K_m for acetate which supports a faster growth rate in environments with high concentrations of acetate.

Replacement of E32 and E334. The kinetic constants determined for the E32A and E334A enzymes (Table 2) indicated that neither residue is essential for catalysis or significantly influences the binding of acetate or ATP.

Conclusions. The first site-specific replacement of amino acids for any acetate kinase has identified a glutamate residue (E384) in the enzyme from *M. thermophila* which is essential for catalysis and is consistent with the previously proposed covalent mechanism for catalysis of kinase activity. Replacement of two other glutamates (E385 and E97) suggests that they are involved in substrate binding.

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AUTHOR'S CORRECTION

Identification of Essential Glutamates in the Acetate Kinase from *Methanosarcina thermophila*

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