# Journal of Bacteriology

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M D Hulanicka, S G Hallquist, N M Kredich and T Mojica-A *J. Bacteriol.* 1979, 140(1):141.

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# Regulation of O-Acetylserine Sulfhydrylase B by L-Cysteine in Salmonella typhimurium

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Received for publication 6 July 1979

A technique based on resistance to azaserine was used to isolate mutants lacking O-acetylserine sulfhydrylase B, one of two enzymes in Salmonella typhimurium capable of synthesizing L-cysteine from O-acetyl-L-serine and sulfide. The mutant locus responsible for this defect has been designated cysM, and genetic mapping suggests that cysM is very close to and perhaps contiguous with cysA. Strains lacking either O-acetylserine sulfhydrylase B or the second sulfhydrylase, O-acetylserine sulfhydrylase A (coded for by cysK), are cysteine prototrophs, but cysK cysM double mutants were found to require cysteine for growth. O-Acetylserine sulfhydrylase B was derepressed by growth on a poor sulfur source, and derepression was dependent upon both a functional cysB regulatory gene product and the internal inducer of the cysteine biosynthetic pathway, Oacetyl-L-serine. Furthermore, a cysB° strain, in which other cysteine biosynthetic enzymes cannot be fully repressed by growth on L-cystine, was found to be constitutive for O-acetylserine sulfhydrylase B as well. Thus O-acetylserine sulfhydrylase B is regulated by the same factors that control the expression of Oacetylserine sulfhydrylase A and other activities of the cysteine regulon. It is not clear why S. typhimurium has two enzymes whose physiological function appears to be to catalyze the same step of L-cysteine biosynthesis.

The final step of L-cysteine biosynthesis in Salmonella typhimurium and Escherichia coli consists of the sulfhydration of O-acetyl-L-serine by either free sulfide (3, 12) or by protein-bound hydrodisulfide (17). In S. typhimurium two enzymes have been described which are capable of carrying out this reaction using free sulfide as a substrate. O-Acetylserine sulfhydrylase A is the better characterized of the two and is coded for by the cysK gene (8). Several lines of evidence indicate that O-acetylserine sulfhydrylase A participates in cysteine biosynthesis in vivo. In addition to its high in vitro enzyme activity, this protein readily binds to serine acetyltransferase (EC 2.3.1.30) (which is coded for by cysE and synthesizes O-acetyl-L-serine [12]) to form a complex designated cysteine synthetase (11). The importance of serine acetyltransferase itself in in vivo cysteine biosynthesis has been clearly established (10, 12), and its physical association with O-acetylserine sulfhydrylase A strongly implies a similar physiological role for the latter. Furthermore, O-acetylserine sulfhydrylase A is a part of the cysteine regulon, since its full expression requires the same factors that are necessary for derepression of other activities of the biosynthetic pathway, i.e., growth on a poor sulfur source such as L-djenkolate or glutathione, the presence of the internal "inducer" Oacetyl-L-serine, and a functional cysB gene (10). The cysB gene (5) codes for a regulatory protein (18) of positive control (15) which is necessary for the derepression of sulfate permease, ATP sulfurylase (EC 2.7.7.4), adenylylsulfate kinase (EC 2.7.1.25), 3'-phosphoadenosine-5'-phosphosulfate reductase, and sulfite reductase (EC 1.8.1.2), in addition to O-acetylserine sulfhydrylase A (9, 10).

Although a large number of cysK mutants have been isolated and characterized in our laboratories, none has shown a nutritional requirement for cysteine, even though many totally lack O-acetylserine sulfhydrylase A activity. However, all such strains, as well as wild type, have small amounts of an enzyme activity which by chromatographic behavior and lack of reactivity with antibody specific for O-acetylserine sulfhy-

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drylase A has been identified as O-acetylserine sulfhydrylase B (4). It is the presence of this enzyme that is thought to be responsible for the Cys<sup>+</sup> phenotype of all cysK mutants. In this communication we describe the isolation and characterization of mutant strains lacking Oacetylserine sulfhydrylase B activity, and we present evidence indicating that the normal physiological function of this enzyme too is one of cysteine biosynthesis.

### MATERIALS AND METHODS

Bacterial strains and genetic methods. The strains of S. typhimurium LT2 used in these studies are listed in Table 1. The cycB1352 allele causes highlevel derepression of all the enzymes of the cysteine regulon regardless of the sulfur source used for growth (10). Therefore, it is designated a  $cysB^{c}$  (constitutive) allele and gives a Cys<sup>+</sup> phenotype. cysB484 does not allow expression of the enzymes of the sulfate reduction pathway and gives a Cys phenotype, but this allele is cysB° for O-acetylserine sulfhydrylase A, causing constitutive expression of this enzyme (10). The cysA locus is required for sulfate permease activity (6), and cysK is the structural gene for O-acetylserine sulfhydrylase A (8). Mutations in cysA and cysK are selected for by resistance to chromate (14) and 1,2,4triazole (7, 8), respectively.

Transduction was performed as described (7) using

Table 1. Bacterial strains

Strain	Genotype	Source		
cysA3	cysA3	Demerec collection"		
cysA20	cysA20	Demerec collection"		
cvsA197	cysA197	Demerec collection"		
cysA272	cysA272	Demerec collection"		
cysB403	cysB403	Demerec collection"		
DW130	cysK1751	(8)		
DW18	cysE2	(10)		
DW42	trpA160 cysB15	(10)		
DW48	trpA160 cysB1352	(10)		
SB3751	trpB223 Δ(cysK- ptsHI182)	C. Cordaro		
TK181	trpC109 cysK1772	Triazole-resistant (7, 8) derivative of <i>trpC109</i>		
DW378	trpC109 cysK1772 cysM1770	Cys azaserine-resistant derivative of TK181		
DW379	trpC109 cysK1772	Cys azaserine-resistant		
	cys <b>M</b> 1771	derivative of TK181		
DW384	trpC109 cysM1770	Cys <sup>+</sup> revertant of DW378		
DW385	trpC109 cysM1771	Cys <sup>+</sup> revertant of DW379		
DW393	trpC109 cysK1772 cvsB1773	Nitrosoguanidine-induced Cvs <sup>-</sup> from TK181		
DW164	pyrE125 cysA	Chromate resistant (14) Cys <sup>-</sup> from pyrE125		
DW382	pvrE125 cysK1751	$DW130 \times DW164^{h}$		
DW383	cvsE2 cvsK1751	$DW18 \times DW382^{b}$		
TK1115	pyrF cysA	Chromate-resistant (14) Cys <sup>-</sup> from pyrF		
TK1129	pyrF cysK1751	$DW130 \times TK1115^b$		
TK1190	cysB1352 cysK1751	$DW48 \times TK1129^b$		
TK1191	cysB484 cysK1751	$cysB484 \times TK1129^b$		

<sup>&</sup>quot;Obtained from either P. Hartman or K. E. Sanderson.

the IA strain of phage P22 (16). Growth on mannitol was used to select for Pts+.

Media and culture conditions. Bacteria were grown at 37°C with rotary shaking in the minimal salts medium C of Vogel and Bonner (19) in which the sulfur source was either 0.5 mM L-cystine, 0.5 mM Ldjenkolate, 0.5 mM reduced glutathione, or 0.8 mM sulfate (10). Glucose (5 g/liter) was added as the usual carbon source; mannitol (2 g/liter) was substituted for selection of Pts+ recombinants. L-Tryptophan (40 mg/ liter) and uracil (10 mg/liter) were included when required for the growth of auxotrophs, and agar (15 g/ liter) was added for solid media. Cells grown for enzyme assays were harvested by centrifugation during exponential growth, washed once with 0.1 M Trishydrochloride (pH 7.6), and usually frozen overnight as a pellet before processing.

Azaserine selection for cysteine auxotrophs. Approximately 108 cells are spread on a minimal agar plate containing 40 mg of L-tryptophan per liter and 0.5 mM glutathione. A 1-cm filter-paper disk containing 1 µmol of freshly prepared azaserine is placed in the center of the plate, and after 48 h resistant colonies appearing in the zone of inhibition are picked and cloned several times on plates lacking azaserine. L-Tryptophan itself gives considerable resistance to azaserine, presumably owing to competition with the aroP permease system (1, 2). For this reason we routinely use L-tryptophan even with Trp+ strains, since in its absence a large percentage of resistant colonies appear to be aroP mutants rather than cysteine auxotrophs. With this technique, the majority of azaserine-resistant isolates are cysteine auxotrophs of various different genotypes. Nutritional and genetic tests have shown that these carry mutations either in the regulatory gene cysB or in any of the structural genes involved in the pathway of sulfate reduction. The rationale for the use of azaserine in selecting for cysteine auxotrophs is discussed in Results.

Assay for O-acetylserine sulfhydrylase. O-Acetylserine sulfhydrylase activity was measured by a previously described method (3) which was adapted for use in an autoanalyzer. The enzyme reaction mixture was formed by mixing 0.3 M Tris-hydrochloride (pH 7.2) containing 1 mM Na<sub>2</sub>EDTA (0.32 ml/min) with air bubbles (0.23 ml/min), 10 mM  $Na_2S$  (0.10 ml/  $\,$ min), 0.1 M O-acetyl-L-serine (0.10 ml/min), and enzyme (0.05 ml/min) in a 2.4-mm-ID glass coil at 23°C. After a transit time of 5 min, the reaction was terminated by the addition of 10 mM NaNO2 in 2 N H2SO4 at 0.16 ml/min. After retention in a mixing coil for 4 min, 0.17 M ammonium sulfamate was added at 0.23 ml/min, and 3.4 min later a solution composed of 4 parts of 0.4 M sulfanilamide in 0.4 N HCl, 2 parts of 7.7 mM N-naphthylethylenediamine dihydrochloride in 0.4 N HCl, and 1 part of 74 mM HgCl<sub>2</sub> in 0.4 N HCl was added at 1.00 ml/min. Four minutes later the stream was debubbled and directed through a 1-cmlight path flow cell in which absorbance at 540 nm was measured in a Gilford model 2000 recording spectrophotometer. A 1-U/ml solution of O-acetylserine sulfhydrylase, as determined by our manual assay in which a unit catalyzes the formation of 1 µmol of Lcysteine per min, gives an absorbancy at 540 nm of 2.2 in the autoanalyzer assay.

<sup>&</sup>lt;sup>b</sup> Constructed by transduction (donor × recipient).

The two O-acetylserine sulfhydrylases were resolved by a modification of the method of Becker and Tomkins (4). Cells (1 to 2 g) were suspended in 3 volumes of 0.1 M Tris-hydrochloride (pH 7.6) and disrupted by sonic oscillation. After centrifugation at  $27,000 \times g$  for 30 min, the supernatant was mixed with 0.5 volume of neutralized 10% streptomycin sulfate, and the resulting precipitate was removed by centrifugation and discarded. The supernatant was then treated with solid ammonium sulfate to obtain the protein fraction precipitating between 0.40 and 0.70 saturation. This material, containing virtually all the O-acetylserine sulfhydrylase activity, was dissolved in a small volume of 0.1 M Tris-hydrochloride (pH 7.6) containing 1 mM 2-mercaptoethanol (buffer A) and desalted by gel filtration on Sephadex G-25 equilibrated in the same buffer. From 10 to 20 mg of protein was then applied to a column (8 cm by 1.8 cm<sup>2</sup>) of DE23 anion-exchange cellulose equilibrated in buffer A, and protein was eluted at 23°C with a 250-ml gradient of 0 to 0.3 M NaCl in buffer A at a flow rate of 0.15 ml/min. Fractions of 3 ml were collected and assayed with the autoanalyzer for O-acetylserine sulfhydrylase both before and after treatment (8) with a 5- to 20-fold excess of rabbit antibody, which inactivates O-acetylserine sulfhydrylase A but not O-acetylserine sulfhydrylase B (4).

With this technique, cystine-grown wild type gives a profile of enzyme activity with three distinct but incompletely resolved peaks (Fig. 1). The first peak is free O-acetylserine sulfhydrylase A; the second is cysteine synthetase, a complex of serine acetyltransferase and O-acetylserine sulfhydrylase A; and the third is O-acetylserine sulfhydrylase B. After treatment with antibody to O-acetylserine sulfhydrylase A, only the O-acetylserine sulfhydrylase B peak is noted. O-Acetylserine sulfhydrylase levels quantified in this manner are expressed in terms of units per milligram of protein

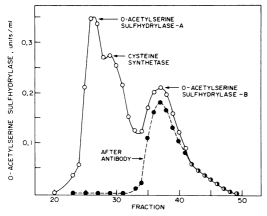


Fig. 1. Elution of O-acetylserine sulfhydrylase activities from a DE23 anion-exchange column. Enzyme was assayed before and after preincubation with a 5-to 20-fold excess of antibody, which specifically inactivates O-acetylserine sulfhydrylase A but not O-acetylserine sulfhydrylase B. Cysteine synthetase is a complex of O-acetylserine sulfhydrylase A and serine acetyltransferase.

applied to the DE23 column, giving specific activities that are about twice that expected in unfractionated crude extracts.

Although this method for assaying O-acetylserine sulfhydrylase B is useful for repressed wild-type cells, it is unsatisfactory for derepressed cells, where an almost 100-fold increase in O-acetylserine sulfhydrylase B peak. Antibody is of no help in this regard because it is capable of inactivating no more than 95% of O-acetylserine sulfhydrylase A. Therefore, to examine the regulation of O-acetylserine sulfhydrylase B, many of our experiments were performed in strains carrying either the cysK1751 or cysK1772 alleles, which code for normally regulated O-acetylserine sulfhydrylase A proteins having less than 2% of wild-type specific activity (8).

**Materials.** Azaserine was purchased from Calbiochem, and autoanalyzer glassware and tubing were obtained from Evergreen Scientific. Rabbit antibody to purified *O*-acetylserine sulfhydrylase A was prepared as described (11). The sources of other chemicals have been cited (8).

### RESULTS

Isolation of mutant strains lacking Oacetylserine sulfhydrylase B. Mutant strains lacking O-acetylserine sulfhydrylase B activity were first isolated by mutagenizing a Cys<sup>+</sup> cysK strain lacking detectable O-acetylserine sulfhydrylase A activity and then scoring for Cys colonies after penicillin selection. Several of these mutants grew on L-cystine but would not grow on sulfate, sulfite, sulfide, O-acetyl-L-serine, or combinations of the latter compounds. Enzyme assays confirmed that these isolates contained no O-acetylserine sulfhydrylase activity. A more convenient method based on azaserine resistance was used for the isolation of the strains described here. Azaserine (O-diazoacetyl-L-serine) is an analog of O-acetyl-L-serine, and we find that it is an excellent substrate for both O-acetylserine sulfhydrylase A and O-acetylserine sulfhydrylase B, giving L-cysteine and diazoacetic acid as products of its reaction with sulfide (data not shown). Mutant strains unable to reduce sulfate to sulfide are resistant to azaserine when grown on minimal agar containing sulfate and a poor sulfur source such as L-djenkolate or glutathione. Presumably, azaserine toxicity in S. typhimurium results in large part from the generation of diazoacetic acid, which in wild type is derived from the reaction of azaserine with sulfide. cysK mutants are also relatively resistant to azaserine, and we assume that this is because they lack O-acetylserine sulfhydrylase A, one of the two enzymes which catalyze the reaction between azaserine and sulfide. By selecting for a higher level of azaserine resistance in a cysK strain on plates containing L-djenkolate (to permit growth of Cys mutants) it is

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relatively easy to obtain mutants which lack both O-acetylserine sulfhydrylase A and O-acetylserine sulfhydrylase B activities.

Strains DW378 and DW379 were derived from the cysK strain TK181 by this technique, and they were found by both nutritional phenotype and enzyme assay to lack any O-acetylserine sulfhydrylase activity. Their Cys' revertants DW384 and DW385 were selected for further study because they were found to contain normal levels of O-acetylserine sulfhydrylase A and no detectable O-acetylserine sulfhydrylase B. We have given the designation cysM to the mutant locus responsible for the absence of Oacetylserine sulfhydrylase B activity in these strains. The trpC cysK<sup>+</sup> cysM strains DW384 and DW385 have no nutritional requirements other than tryptophan and grow as well on sulfate as sole sulfur source as do wild type or cvsK  $cvsM^+$  strains. Identical results have been found in  $trpC^+$  cysM derivatives, which are  $Trp^+$ . It appears, therefore, that either of the two Oacetylserine sulfhydrylase enzymes is sufficient for cysteine prototrophy under our usual conditions of aerobic growth and that neither enzyme is required for some other essential metabolic function.

Genetic mapping of cysM. Preliminary experiments indicated that cysK and cysM are cotransducible by P22. Since cysK is known to be very close to the pts locus we measured the linkage between cysM and pts. Using the Cys<sup>+</sup>  $\Delta(cysK-ptsHI)$  strain SB3751 as a recipient and two different Cys<sup>-</sup> cysK cysM donors, we selected for Pts<sup>+</sup> and scored for Cys. A cysK<sup>+</sup> recombinant is expected very rarely or not at all from such a cross, depending on whether the recipient's cysK deletion includes the site in which the donor cysK locus is mutated. Therefore all or virtually all Cys' recombinants should be  $cysM^+$  cysK, and all Cys recombinants of course are cysK cysM. In the experiments summarized in Table 2, cysM was cotransduced with pts at a frequency of 52 to 55%.

The linkage between pts and cysM is very close to that known for pts and cysA (8). Three-point tests could not be performed to determine the relative order of these three loci owing to the lack of a reliable scoring method for cysM in Cys' strains. However, direct assay of a strain carrying cysA20, a deletion which covers all tested cysA point mutations (13), showed complete absence of O-acetylserine sulfhydrylase B activity. Three cysA point mutants, cysA3, cysA197, and cysA272, had normal levels of O-acetylserine sulfhydrylase B, suggesting that the loss of this enzyme activity in cysA20 is due to a deletion extending through cysA into cysM. O-

Table 2. Genetic mapping of cysM<sup>o</sup>

	Recombinants			
$\mathbf{Donor}^h$	Total no. of Pts <sup>+</sup>	No. of Cys Pts*	Percent co- transduc- tion	
DW378 (cysM1170 cysK1772)	94	49	52	
DW379 (cysM1171 cysK1772)	97	53	55	

<sup>&</sup>quot;Transduction was performed with the L4 strain of P22. Recipient was SB3751 [Δ(cysK-ptsHI)]. Selection was for Pts<sup>+</sup> on plates containing mannitol as a carbon source and L-cystine.

Acetylserine sulfhydrylase A was present in all four *cysA* strains.

Regulation of O-acetylserine sulfhydrylase B activity. In cystine-grown, repressed wild type, O-acetylserine sulfhydrylase B comprises about 30% of the total O-acetylserine sulfhydrylase activity; the remainder is O-acetylserine sulfhydrylase A, approximately 25% of which is found complexed to serine acetyltransferase as cysteine synthetase. Levels of free Oacetylserine sulfhydrylase A are increased 50- to 100-fold during derepression by growth on a limiting sulfur source such as L-djenkolate or glutathione, making it impossible to estimate accurately the O-acetylserine sulfhydrylase B peak from a DE23 column even with the aid of antibody to O-acetylserine sulfhydrylase A. Therefore, to study the regulation of the latter enzyme it was often necessary to use strains with mutant cysK alleles which code for O-acetylserine sulfhydrylase A proteins having little or no activity. In the cysK strains DW130, TK181, and TK1129, growth on a poor sulfur source gives Oacetylserine sulfhydrylase B levels 4- to 14-fold higher than those found in L-cystine-grown cells (Table 3). This derepression of enzyme activity also requires a functional cysB gene product, as deduced from the fact that sulfur-deprived cultures of cysB403, DW42, and DW393 (carrying three separate cysB alleles) all show little or no increase in O-acetylserine sulfhydrylase B activity over the low levels found in L-cystine-grown cultures.

In contrast, the  $cvsB^c$  allele cvsB1352, which is present in TK1190, causes high-level expression of O-acetylserine sulfhydrylase B in both L-cystine-grown and sulfur-deprived cells. This effect is similar to that observed for other enzymes of the cysteine regulon (10). cysB484, which is cysB (negative) for the enzymes of sulfate reduction but  $cysB^c$  for O-acetylserine sulfhydrylase A, was found to be cysB (negative)

<sup>&</sup>lt;sup>b</sup> Only the relevant genotypes are given.

Table 3. O-Acetylserine sulfhydrylase B expression in various cysteine mutants

	Relevant geno- type	O-Acetylserine sulfhydrylase B (units/mg of protein)		
Strain		Grown on L-cystine	Grown on L- djenkolate	
Wild type		0.199	a	
cysA20	$\Delta cysA$	< 0.02	_	
DW384	cys <b>M</b>	< 0.02	_	
DW385	cys <b>M</b>	< 0.02	_	
DW130	cysK	0.158	0.580	
TK181	cysK	0.045	0.713	
TK1129	cysK	0.104	0.512	
cysB403	cysB	0.072	0.079	
DW42	cysB	0.062	0.122	
DW393	cysB cysK	0.115	0.121	
TK1191	cysB484 cysK	0.097	0.092	
TK1190	$cysB^c$ $cysK$	0.542	0.492	
DW18	cysE	0.148	0.076	
DW383	cysE cysK	0.046	$0.188^b (0.414)^c$	

<sup>&</sup>lt;sup>a</sup> -, Not measured owing to large excess of O-acetylserine sulfhydrylase A.

with respect to O-acetylserine sulfhydrylase B expression.

O-Acetyl-L-serine is also required for derepression of the cysteine regulon in cysB<sup>+</sup> strains. Therefore cysE strains are not derepressed for these enzymes owing to the lack of serine acetyltransferase, the enzyme required for O-acetyl-L-serine synthesis. DW18, carrying cysE2, could not be derepressed for O-acetylserine sulfhydrylase B by growth on glutathione, and DW383, also carrying cysE2, was only partially derepressed for this enzyme. The addition of 2 mM O-acetyl-L-serine to a glutathione culture of the latter strain, however, gave a greater than twofold rise in O-acetylserine sulfhydrylase B activity.

### DISCUSSION

The lack of measurable O-acetylserine sulfhydrylase B activity in cysM mutants and the Cys<sup>-</sup> phenotype of cysK cysM strains suggests that cysM is the structural gene for this enzyme. A more definite designation, however, must await the demonstration of an altered enzyme protein in a cysM strain. Although our genetic mapping data are insufficient to establish the precise chromosomal location of cysM, its linkage to pts and the absence of O-acetylserine sulfhydrylase B in the deletion strain cysA20 suggest close linkage and perhaps even contiguity between cysM and cysA.

Since cysM mutants display no nutritional requirements other than the Cys phenotype of cysK cysM double mutants, it seems unlikely

that O-acetylserine sulfhydrylase B is simply a spurious activity of an enzyme with another physiological function. Of course the same argument might be made against the role of this enzyme in L-cysteine biosynthesis because  $cysK^+$  cysM strains grow as wild type on sulfate. However, the apparent close linkage of cysM with cysA, and our finding that O-acetylserine sulfhydrylase B is regulated by the same nutritional and genetic factors that control expression of other cysteine biosynthetic activities, indicate that this enzyme is a part of the cysteine regulon and plays a role in cysteine biosynthesis other than that imposed upon it in a cysK strain.

Having used similar reasoning in concluding that the physiological function of O-acetylserine sulfhydrylase A is to synthesize L-cysteine, we are now confronted with the perplexing question of why there should exist two sulfhydrylase enzymes, either one of which is sufficient for normal growth on sulfate. In general it is very unusual for an organism such as S. typhimurium to carry and express genes for two different enzymes which carry out the same biochemical reaction. One obvious explanation for this situation is that they may not catalyze exactly the same reaction in vivo. For example, even though both enzymes use free sulfide as a reactant with O-acetyl-L-serine in our in vitro assay, the actual sulfur donor in vivo may be different for each enzyme. Perhaps one enzyme prefers free sulfide and the other is required for the direct utilization of carrier-bound sulfide, such as the thioredoxin hydrodisulfide intermediate proposed by Tsang and Schiff in Escherichia coli (17).

An interesting observation (our unpublished data) which may bear on this problem is that although cysM strains grow well aerobically on sulfate, they are cysteine bradytrophs under anaerobic conditions. In contrast, cysK strains, which lack the major portion of total O-acetylserine sulfhydrylase activity as measured by our in vivo assay, grow on sulfate as well as wild type both aerobically and anaerobically. Perhaps a careful examination of the pathway of L-cysteine biosynthesis and its regulation during anaerobiosis will eventually lead to an explanation for the existence of two different O-acetylserine sulfhydrylases in S. typhimurium.

# **ACKNOWLEDGMENTS**

This work was supported by Public Health Service grant AM-12828 from the Institute of Arthritis, Metabolism and Digestive Diseases and by the Polish Academy of Sciences within the Project: 09.7. N.M.K. is an Investigator of the Howard Hughes Medical Institute.

We thank Giovanna Ames, who informed us of a possible relationship between azaserine toxicity and cysteine biosynthesis.

<sup>&</sup>lt;sup>b</sup> Grown on 0.5 mM glutathione because of very poor growth on L-djenkolate.

Grown on 0.5 mM glutathione plus 2 mM O-acetyl-L-

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