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Isolation and Partial Characterization of an argR Mutant of Salmonella typhimurium

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An arginine regulatory mutant (i.e., mutated in the argR gene) has been isolated from a strain of $Salmonella\ typhimurium\ LT2$. The argR mutant was found to excrete arginine into the growth medium with glycerol but not glucose as carbon source. Constitutive synthesis of arginine biosynthetic enzymes was observed. Whereas previous results (A. T. Abd-El-Al and J. L. Ingraham, Abstr. Annu. Meet. Am. Soc. Microbiol. 1975, K169, p. 175) have shown constitutive synthesis of carbamyl phosphate synthetase in the argR mutant, the regulation of the synthesis of the last five enzymes of the pyrimidine pathway was unaffected. However, in pyrH mutants, known to exhibit derepressed synthesis of the pyrimidine enzymes, a 10-fold derepression of ornithine transcarbamylase was observed.

The arginine biosynthetic system of *Escherichia coli* has received extensive study for 20 years (for a recent review, see reference 22). However, arginine metabolism in the closely related bacterium *Salmonella typhimurium* has not been as intensively investigated. In both of these genera, a single carbamyl phosphate synthetase (CPSase) (encoded by the gene *pyrA*) catalyzes the synthesis of carbamyl phosphate (CP) (18, 23). CP is considered the first intermediate of pyrimidine biosynthesis and is also required at the sixth step of arginine biosynthesis (Fig. 1).

To date, no true regulatory mutants in which the pyrimidine pathway is affected have been isolated, but mutants are available that mimic constitutive synthesis of all six pyrimidine enzymes. These mutants do not have a regulatory dysfunction per se, but rather contain a partial impairment of the enzyme uridine monophosphate kinase, encoded by the gene *pyrH* (9).

Regulatory mutants (mutated in the *argR* gene) altered in control of the arginine pathway of *E. coli* were isolated in 1961 by Maas (15), and a brief report of a method for the isolation of arginine regulatory mutants of *S. typhimurium* has been published (10). A variety of methods have been established for the isolation of arginine regulatory mutants of *E. coli*, such as selection for canavanine resistance (11, 15), *N*-acetyl amino acid utilization in the presence of arginine by various amino acid auxotrophs (3), and phenotypic suppression of proline auxotrophy in the presence of arginine (10).

Auxotrophs for certain amino acids are capa-

ble of utilizing the *N*-acetyl derivative of the required amino acid; such utilization is precluded when arginine is added to the medium because the added arginine represses the synthesis of the enzyme that deacylates the *N*-acetyl amino acid, namely *N*-acetylornithinase (NAOase). Thus, mutants selected for the ability to use the *N*-acetyl amino acid in the presence of arginine are, generally, *argR* mutants.

Phenotypic revertants of proA, proB, or proAB deletion mutants can be obtained (4, 10, 14); however, such proline-independent strains do not result from true back-mutation, but rather occur from a mutation in the arginine pathway in the argD gene (encoding the enzyme N-acetylornithine- δ -transaminase [AMTase]) of E. coli (21); the homologous locus in S. typhimurium is argG (19). The biochemical basis for the phenotypic suppression of proline auxotrophy, termed indirect suppression (7, 14), is outlined in Fig. 2. These double mutants are able to grow in the absence of arginine because a nonspecific transaminase partially substitutes for the missing enzyme (10). When arginine is added, arginine biosynthesis is repressed (22) and the indirectly suppressed mutants become proline requiring. Arginine regulatory mutants can be derived from indirectly suppressed proline auxotrophs by selecting for strains capable of growth in the presence of arginine without added proline (10).

We undertook our studies on arginine metabolism with the primary aim of exploring the interrelationships of the arginine and pyrimidine pathways of *S. typhimurium*. For such a

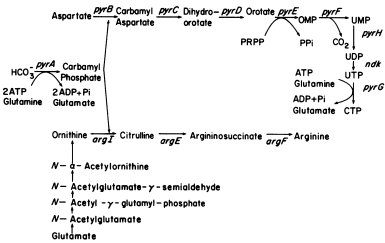


Fig. 1. Pyrimidine and arginine biosynthetic pathways of Salmonella typhimurium. Genetic symbols for the enzymes are shown in italics. The gene designations are as follows: pyrA, carbamyl phosphate synthetase (EC 2.7.2.5); pyrB, aspartate transcarbamylase (EC 2.1.3.2); pyrC, dihydroorotase (EC 3.5.2.3); pyrD, dihydroorotate dehydrogenase (EC 1.3.3.1); pyrE, orotidine monophosphate (OMP) pyrophosphorylase (EC 2.4.2.10); pyrF, OMP decarboxylase (EC 4.1.1.23); pyrH, uridine monophosphate (UMP) kinase (EC 2.7.4.4); ndk, nucleoside-diphosphokinase (EC 2.7.4.6); pyrG, cytidine triphosphate (CTP) synthetase (EC 6.3.4.2); argI, ornithine transcarbamylase (EC 2.1.3.3); argE, argininosuccinate synthetase (EC 6.3.4.5); and argF, argininosuccinase (EC 4.3.2.1).

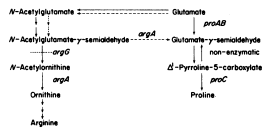


Fig. 2. Biosynthesis of arginine and proline from glutamate in E. coli and S. typhimurium. In argG mutants of S. typhimurium (equivalent to argD mutants of E. coli), the argA enzyme (argE in E. coli) mediates the deacylation of N-acetylglutamate- γ -semialdehyde to glutamate- γ -semialdehyde. Symbols: \rightarrow , Normal pathways; $- \rightarrow$, proline-suppressed pathway in proAB argG double mutants. The genetic designations represent the following enzymes: argG, N-acetylornithine- δ -transaminase (EC 2.6.1.11); argA, N-acetylornithinase (EC 3.5.1.16); proAB, glutamate- γ -semialdehyde dehydrogenase; proC, pyrroline carboxylate reductase (EC 1.5.1.2).

project, it was important to isolate arginine regulatory mutants. The studies described herein report the isolation and partial characterization of an argR mutant and the effect of a pyrH mutation on the arginine pathway. The argR gene has been located at 106 min on the S. typhimurium chromosome (12), which is analogous to the position on the E. coli K-12 chromosome (11).

MATERIALS AND METHODS

Organisms. Most of the bacterial strains used in this study were derivatives of S. typhimurium strain LT2. Strain P110 is derived from S. typhimurium strain LT7. Table 1 lists the strains used, their genotypes, and the source. Bacteriophage P22-L4 was used for transductions and was received from K. E. Sanderson.

Media and growth conditions. The composition of the defined medium, AG medium, has been reported (12). Nutritional supplements were added at $100~\mu g/$ ml unless stated otherwise. The complex medium was nutrient broth (Difco). Solid media were prepared by the addition of agar to 1.5%. Growth procedures were as previously described (12).

Preparation of cell extracts and enzyme assays. The method of preparing cell extracts and determining protein concentration has been reported (13). Assays for the last five enzymes of the pyrimidine pathway and for ornithine transcarbamylase (OT-Case) were conducted according to previously published procedures (13). The assays of NAOase and AMTase were carried out as previously described (5). Specific activities are defined as nanomoles of product formed (or substrate utilized) per minute per milligram of protein.

N-acetylhistidine utilization. The procedure described by Baumberg (3) was used as a possible method of selecting arginine regulatory mutants.

Selection of indirectly suppressed mutants. The basic procedure for the isolation of indirectly suppressed mutants of proline auxotrophs and the subsequent selection of arginine regulatory mutants from indirectly suppressed strains has been described (10). The following is a detailed presentation

Table 1. Bacterial strains

Strain	Genotype	Source
LT2	Wild type	P. E. Hartman
JL84	HfrA, metB406	J. L. Ingraham
P21	hisD23 gal-50 proAB21	K. E. Sanderson
B75	argB75	K. E. Sanderson
F87	argF87	K. E. Sanderson
SU453	his-1009 trpE2 metA22 xyl-1	K. E. Sanderson
D	strA201	W D 0 1
P110	proC110 (LT7)	K. E. Sanderson
TR192	ser-791 his-644	J. Roth
JL1269	pyrH1609 cdd-7	J. L. Ingraham
KD1272	pyrH1609 cdd-7 pro-5	This study
KD1275	cdd-7	This study

of the protocol used in this study. Strain P21 was mutagenized with N-methyl-N'-nitro-N-nitrosoguanidine (2), and samples of the mutagen-treated culture were spread on AG plates. After 72 h of incubation at 37°C, some of the developing colonies were picked and purified on AG medium. These colonies were then tested for the phenotype of being proline requiring in the presence of arginine.

Selection of arginine regulatory mutants. Those indirectly suppressed isolates showing arginine sensitivity were grown to stationary phase in nutrient broth medium. Samples were spread on AG plates containing arginine and incubated. Resulting colonies were picked and purified on AG medium supplemented with arginine and then tested for the ability to grow on unsupplemented AG medium.

Genetic techniques. (i) Conjugation. The method for introducing the *pyrH*⁺ allele into a mutant *pyrH* strain has been described (13).

(ii) Transduction. The methods used with P22-L4 were as previously published (6). For cotransfer of $argG^+$ and strA, we used strA as the selective marker and screened for the cotransfer of $argG^+$ to indirectly suppressed strains as described (14). Selection for transfer of $argG^+$ was also performed. Kuo and Stocker (14) have reported that $argG^+$ transductants of indirectly suppressed strains are easily recognizable as fast-growing colonies on a background of slow-growing recipient cells when plated on defined medium supplemented with proline. Their basic procedure was followed except that the plates were incubated at 20°C rather than 36°C. The $argG^+$ (or strA $argG^+$) transductants were then tested for arginine excretion on indicator plates (12) and assayed for AMTase activity.

Syntrophic studies. The testing of various strains for the excretion of arginine or proline was carried out as described by Berg and Rossi (4) with slight modification. The test and indicator strains were grown in nutrient broth medium, and testing was carried out on a medium consisting of the basal salts of AG medium and 0.4% glycerol as carbon source.

Quantitation by bioassay of arginine excreted. A standard curve relating the amount of growth of strain F87 to various limiting concentrations of argi-

nine was prepared. The argR mutant test strain was grown in various media to a cell density of 6 imes 108/ ml. The cells were pelleted by centrifugation, and the supernatants were collected and filter sterilized. A 5-ml volume of a sterile culture supernatant was added to a flask containing 5 ml of AG medium. The flask was inoculated with 0.05 ml of a strain F87 culture (grown in arginine limiting medium). The flasks were incubated and growth was monitored. In the flasks in which no growth occurred within 12 h, a known, limiting concentration of arginine was added. The resulting growth was followed, and the amount of growth was checked against the standard curve to insure that it was consistent with that expected for the particular arginine concentration. This procedure was necessary in order to eliminate the possibility that a lack of growth was due to the accumulation of a toxic product in the culture fluid rather than to an actual deficiency in arginine ex-

Chemicals. Substrates and reagents for the various enzymes were purchased from Sigma Chemical Co. (St. Louis, Mo.). All other chemicals were reagent grade.

RESULTS

N-acetylhistidine utilization and arginine sensitivity. A histidine auxotroph of S. typhimurium (strain TR192) was tested for the ability to use *N*-acetylhistidine as histidine source. The strain grew with *N*-acetylhistidine as histidine source, but growth was not inhibited by arginine as occurs with E. coli (3). Measurement of the level of NAOase in cells grown in the presence and absence of arginine explained the lack of arginine sensitivity. In the presence of arginine, NAOase was repressed from a specific activity of 18.4 to 8.3, and thus an adequate level of NAOase was still present in the cells to carry out the deacylation of *N*-acetylhistidine. Various intermediates of the arginine pathway were then tested for their repressive effect on the synthesis of NAOase, but none was as effective as arginine (R. A. Kelln, unpublished observations).

Indirectly suppressed proline auxotrophs and arginine-resistant derivatives. After nitrosoguanidine mutagenesis, derivatives of strain P21 were isolated that grew in the absence of proline, although at a slow rate. Eleven selected isolates were then tested for their growth responses on various media. All of the isolates showed excellent growth on medium supplemented with proline and arginine. Nine of the eleven isolates failed to grow with arginine as the sole supplement. These nine isolates were assumed to be argG mutants (Fig. 2). A number of the indirectly suppressed isolates showing arginine sensitivity were chosen, and arginine-resistant derivatives of these isolates were selected spontaneously.

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Enzymes activities of various mutants. The activities of two enzymes of the arginine pathway (encoded by unlinked genes) were assayed from isolates grown in the presence of both proline and arginine (Table 2). The most significant finding was the high levels of OTCase in extracts of arginine-resistant derivatives of isolate 11.

Growth rates in various media. The growth rates in defined media of the parent strain (P21), an indirectly suppressed isolate, and of an arginine-resistant strain derived from it are given in Table 3. Growth of strain P21 occurred only in the presence of proline. Isolate 11 grew slowly in AG medium or in AG medium plus proline; it did not grow at all when arginine was the sole supplement. In the presence of both amino acids, the growth rate of isolate 11 was comparable with that of strain P21. The arginine-resistant derivative of isolate 11

Table 2. NAOase and OTCase activities in various isolates ^a

Isolate ^b	Sp act ^c		
isolate	NAOase	OTCase	
4	42.8	240	
8	29.1	56.4	
11	40.3	16.2	
4-7	94.8	242	
8-4	58.3	268	
11-5	175	1138	
11-6	138	992	
11-9	124	1214	
11-10	125	1142	
11-11	111	1102	
11-12	108	1110	

^a Cells were grown in AG medium supplemented with proline and arginine.

^c As defined in the text.

Table 3. Growth rates of various strains in different media

Strain	Relevant geno- type	Generation time (min) in various media ^a				
		No addi- tions	Pro- line	Argi- nine	Pro- line + argi- nine	
P21	proAB	_ b	55	_	52	
11	proAB argG	185	175	_	52	
11-5	proAB argG argR	150	150	144	54	

^a The strains were grown in AG medium with supplements as indicated.

showed a similar growth pattern to isolate 11 except that it grew in the presence of arginine, but at a slow rate. This result indicated that the supply of glutamate- γ -semialdehyde for the cross-feeding of the proline pathway was limiting.

Transductions. Kuo and Stocker (14) have demonstrated the cotransduction of argG and strA in S. typhimurium LT7. A lysate of P22-L4 was prepared on an strA $argG^+$ donor strain (strain SU453) and used to infect the recipient strain, isolate 11-5. A second lysate of phage P22-L4 was prepared on wild-type LT2 and used to infect the recipient strain. The $argG^+$ strAtransductants were assayed for AMTase activity. These transductants showed high levels of AMTase, whereas the recipient strain was devoid of measurable activity (Table 4). The argG+ transductants were also assayed for AM-Tase activity, and similar observations were made. The strA $argG^+$ and $argG^+$ transductants were tested for arginine excretion on suitable indicator plates; both were found to excrete arginine.

Syntrophic studies. Berg and Rossi (4) have reported that proAB argG double mutants of S. typhimurium LT7 excrete proline. Table 5 presents our results of syntrophic studies with LT2 strains. The wild-type LT2 did not excrete proline or arginine. The argG proAB mutant strain excreted proline, and this excretion was independent of the argR mutation. Arginine was excreted only by the argR mutant strain with the wild-type $argG^+$ allele. This finding suggests that in an argR mutant the other transaminase(s) synthesizes insufficient N-acetylornithine for adequate arginine biosynthesis.

Pyrimidine and arginine enzyme levels in the argR mutant. The last five enzymes of the pyrimidine pathway (along with three enzymes of the arginine pathway) were assayed to determine whether any of these pyrimidine biosynthetic enzymes were derepressed in an argR mutant (Table 6). With strain P21, repression of the three arginine enzymes occurred when the strain was grown in the presence of arginine. In the argR derivative, very high levels of the same three arginine enzymes were observed. Moreover, the levels were unaffected by growth in the presence of arginine. Indeed, a 300-fold derepression of OTCase was found in the argR mutant in comparison to the repressed level of strain P21. The other two arginine enzymes assayed in the argR mutant were not as markedly derepressed.

Even with this high level of OTCase, the pyrimidine enzyme levels were unaffected. It was anticipated that the marked increase in

^b The first number denotes the original indirectly suppressed proline auxotroph derived from strain P21. The number after the hyphen designates the arginine-resistant subisolate.

 $^{^{}b}$ -, No growth.

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OTCase would result in excessive utilization of affected the enzyme levels of the arginine path-

CP, causing a limitation of CP for pyrimidine biosynthesis. However, NAOase was derepressed to a far lesser degree than OTCase, and the possibility that the synthesis of ornithine may be a limiting factor was considered. The basic experiment was repeated with ornithine added to the medium. The results were unchanged; no effect on the pyrimidine enzyme levels was observed (R. A. Kelln, unpublished observations).

Enzyme levels in pyrH and pyrH⁺ strains. In the argR mutant, the loss of the arginine regulatory gene product affected the synthesis of only one pyrimidine enzyme, namely CPSase. As mentioned earlier, pyrH mutants show derepressed pyrimidine enzyme levels (9, 13, 20), and it was in our interest to observe whether derepression of pyrimidine enzymes

Table 4. Activity of AMTase of various strains^a

Strain b	Relevant genotype	Sp act		
11-5	proAB argG argR	0		
11-5 G ₁		93		
11-5 G ₂	proAB argR strA	111		
11-5 G ₃		105		
11-5 G ₄		107		

^a Cells were grown in AG medium plus proline.

Table 5. Syntrophic properties of different strains

Relevant genotype of test	Growth response of indicator strain				
strain -	proAB ^b	proC	argF		
Wild type (LT2)	_	_	_		
proAB $argG$	+	_	-		
proAB argG argR	+	+	_		
proAB argR	ND	ND	+		

^a Symbols: -, No growth; +, growth; ND, not done

affected the enzyme levels of the arginine pathway (Table 7). The pyrH mutant had a 10-fold higher OTCase level than the $pyrH^+$ strain.

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Quantitation of excreted arginine. The argR mutant strain was grown under four different conditions to the same cell density. The amount of arginine present in the culture fluid was determined by bioassay (Table 8). The parent strain excreted a small amount of arginine only with glycerol as carbon source and ornithine added to the medium. The argR mutant likewise excreted an appreciable amount of arginine (7.5 μ g/ml) with glycerol as carbon source, and ornithine supplementation increased the amount of arginine excreted. With glucose as the carbon source, the argR mutant failed to excrete arginine even when ornithine was added to the medium.

DISCUSSION

Our initial attempt to isolate an arginine regulatory mutant of S. typhimurium LT2 by the conventional canavanine resistance method (15) was unsuccessful. S. typhimurium was found to be naturally resistant to canavanine (12) even when AF medium (16) was used. The next approach was based on the selection of mutants from histidine-requiring strains that are capable of utilizing N-acetylhistidine as histidine source in the presence of arginine (3). This approach also proved inappropriate, because arginine supplementation resulted in only a twofold repression of the deacylating enzyme, NAOase. Various attempts to increase the repression of this enzyme were unsuccessful.

Consequently, a third method was used, an approach that had apparently already been successful with $S.\ typhimurium\ LT2\ (10).$ As our starting strain, we chose a multisite deletion mutant in the proAB region to eliminate any possibility of true back-mutation when selecting for proline-independent derivatives. A number of proline-independent strains were isolated. Those isolates showing arginine sensi-

Table 6. Comparison of enzyme levels in an argR+ strain and an argR mutant

Strain	Sp act of enzymes of the arginine and pyrimidine pathwa						pathways	.ys ^c		
	Relevant genotype	Additions to medium ^b	ATCase	DHOase	DHO- dehase	OMP- ppase	OMP- decase	AMTase	NAOase	OTCase
P21	proAB	None Arginine	14.7 11.6	130 129	13.5 11.8	46 45	14.6 14.6	17.4 7.5	18.7 8.2	63.6 4.8
KD2111	proAB argR	None Arginine	11.8 13.7	148 132	13.0 13.1	46 46	12.1 11.0	95.8 110	191 216	1393 1423

^a Strain KD2111 is an arg⁺ transductant of isolate 11-5 (proAB21 argG11 argR5).

^b Strains 11-5 G_1 through 11-5 G_4 are strA $argG^+$ transductants of strain 11-5.

 $^{^{}b}$ Denotes the relevant genotype of the indicator strain.

^b The medium was AG medium with added proline.

Enzyme abbreviations: ATCase, aspartate transcarbamylase; DHOase, dihydroorotase; DHOdehase, dihydroorotate dehydrogenase; OMPppase, orotidine monophosphate pyrophosphorylase; OMPdecase, orotidine monophosphate decarboxylase; AMTase, N-acetylornithine-8-transaminase; NAOase, N-acetylornithinase; OTCase, ornithine transcarbamylase.

Table 7. Comparison of enzyme levels in a pyrH⁺ strain and a pyrH mutant^a

Relevant gen-	Sp act				
otype	ATCase	DHOdehase	OTCase		
pyrH cdd	606	34.5	476 50.4		
	otype	otype ATCase pyrH cdd 606	otype ATCase DHOdehase pyrH cdd 606 34.5		

^a The strains were grown in AG medium.

Table 8. Quantitation of arginine overproduction: amount of arginine excreted by an argR⁺ strain and an argR mutant under various growth conditions^a

Strain		Amt of arginine excreted					
	Relevant gen- otype	AG	AG + orni- thine	A- glyc- erol	A- glyc- erol + orni- thine		
P21 KD2111	proAB proAB argR	<1 ^b <1	<1 <1	<1 7.5	2 11		

^a The cells were grown in AG medium or in modified AG medium where glycerol was the carbon source instead of glucose. The quantitation of arginine in the culture supernatants was carried out as described in Materials and Methods.

tivity were assumed to be argG mutants and were used for the selection of arginine regulatory mutants.

Though all of the arginine enzymes were not assayed, those enzymes chosen for study were encoded by unlinked genes, thus effectively eliminating operator mutations. Only one isolate, isolate 11, yielded arginine-resistant mutants of the desired phenotype. These mutants had highly elevated enzyme levels when grown in the presence of arginine. Kadner and Maas (11) have reported that selection of argR mutants of E. coli using canavanine resistance gives rise to at least three classes of argR mutants. Class I mutants show fully derepressed enzyme synthesis under all growth conditions. Class II mutants are fully derepressed in the absence of arginine, but when arginine is present they are partially or fully repressed. Class III mutants show partial derepression that remains constant under all growth conditions. Comparisons of our mutants to these various classes were not carried out, but from later experiments it would seem reasonable that the arginine regulatory mutant of isolate 11 is representative of class I.

Indirectly suppressed proline auxotrophs have been described as having slow growth rates on minimal medium or on proline-supplemented medium, as well as being sensitive to arginine (10, 14). With isolate 11, selection for

the argR mutation completely overcame the sensitivity to arginine, but the growth rate in the presence of arginine alone was still very slow (Table 3). Thus, even in the argR mutant, when grown in the presence of arginine alone, N-acetylglutamate- γ -semialdehyde synthesis appeared impaired and consequently partially prevented the cross-feeding of the proline pathway. With the constitutive synthesis of the arginine enzymes, it would appear that the reason for impaired N-acetylglutamate-y-semialdehyde synthesis was a consequence of feedback inhibition of the first enzyme of the arginine pathway. It should be possible, therefore, to isolate mutants that are insensitive to feedback inhibition by selecting derivatives that grow rapidly in the presence of arginine.

The regulation of proline biosynthesis is a subject of controversy at present (4). Evidence has been provided for regulation at both the first and last steps and, in addition, a pleiotropic effect on proline regulation resulting from the introduction of an argD mutation in $E.\ coli$ has been demonstrated. Our results lend further support to the previous observation (4) that indirectly suppressed proline auxotrophs of $S.\ typhimurium$ excrete proline. Thus, the accumulated evidence has shown that a bypass of a regulatory point in proline biosynthesis occurs in indirectly suppressed proline auxotrophs.

CPSase has been designated as the first enzyme of pyrimidine biosynthesis in *E. coli* (21) and S. typhimurium (19). However, when an argR mutation was introduced into S. typhimurium, this enzyme was found to be constitutively synthesized (A.T. Abd-El-Al and J.L. Ingraham, Abstr. Annu. Meet. Am. Soc. Microbiol. 1975, K169, p. 175). In an E. coli K-12 strain harboring an argR mutation, the repression of CPSase by arginine and/or uracil was markedly reduced, but some repression did occur (17). Pierard et al. (17) made the suggestion (based on the decreased repressibility of CPSase) that repression of the enzyme may be exerted by a complex repressor molecule, composed of the argR gene product and another gene product related to the pyrimidine pathway. In the S. typhimurium mutant, evidence points to an involvement of only the argR gene product, which presumably forms a complex with arginine, and some pyrimidine compound that alone results in repression of CPSase. An interaction of the arginine repressor and a pyrimidine compound is supported by the finding that OTCase was repressed to a greater extent by the addition of both pyrimidines and arginine than by the addition of arginine alone (13). The differences observed in CPSase regulation

b Values are expressed as micrograms of arginine per milliliter of culture filtrate.

in *E. coli* K-12 and *S. typhimurium* LT2 may be a function of the nature of the *argR* mutation (either class I, II, or III). However, inherent differences in CPSase regulation in these two genera of bacteria may also account for the change.

The common feature of the two transcarbamylases ATCase and OTCase is the requirement of CP as a substrate. When OTCase was constitutively synthesized in the argR mutant, the ATCase level was unaffected; when ATCase was derepressed in the pyrH mutant, OTCase was derepressed as well. ATCase may preferentially use CP and may create a partial arginine starvation that should result in derepression of OTCase and the other enzymes of the arginine pathway. In support of this statement is the isolation of a pyrA mutant (1) having an arginine requirement but not a pyrimidine requirement. It seems pertinent to point out that the genes encoding OTCase and ATCase are located adjacent to each other on the $E.\ coli\ (21)$ and S. typhimurium (19) chromosomes and that these two enzymes exhibit the greatest degree of derepression of the enzymes of the two pathways.

Determination of the amount of arginine excreted produced two interesting results. First, glucose-grown cells did not excrete arginine even when supplied with ornithine. Gorini and Gundersen (8) have reported a form of catabolite repression for glucose-grown cells of $E.\ coli$ B; the arginine enzyme levels are much lower in glucose-grown cells than in glycerol-grown cells. Perhaps even in the absence of the argRproduct, the level of derepression in glucosegrown cells of S. typhimurium is markedly lower than that in glycerol-grown cells. In view of such a glucose effect, a study on enzyme levels in glycerol-grown cells appears warranted. Second, the amount of arginine excreted with glycerol-grown cells could be increased when ornithine was added to the medium, showing that in the argR mutant ornithine biosynthesis was a limiting factor in arginine overproduction. Even in the parent strain, arginine was excreted when ornithine was added. Thus, in a strain with the wild-type argR allele, it appears that the rate of ornithine synthesis plays an important role in the maintenance of balanced arginine synthesis.

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