Excretion of Enterochelin by exbA and exbB Mutants of Escherichia coli

S. K. GUTERMAN, 1 AND L. DANN²

Biology Department, Brandeis University, Waltham, Massachusetts 02154

Received for publication 18 January 1973

Escherichia coli mutants that are insensitive to colicins B and I hyperproduce and excrete the iron chelator enterochelin, which is an inhibitor of these colicins. These mutants are classified as exbA and exbB. The exbA mutants are chromium sensitive and require iron for growth, and the mutations are located in the tonB region at min 25 of the *E. coli* chromosome. tonB mutants in which the genome of phage lambda is inserted into the bacterial chromosome within the tonB gene also exhibit enterochelin excretion. The exbB mutants require methionine and probably result from deletions which are located between min 56 and 58. Colicin insensitivity, enterochelin excretion and methionine auxotrophy are recessive in exbB merodiploids. The methionine requirement of exbB strains is satisfied by cystathionine or homocysteine, and exbB mutants are sensitive to ethionine.

Colicin B- and I-insensitive mutants were described by Gratia (6). Some of the mutations conferred resistance to phage T1, were mapped in the tonB gene near the tryptophan operon, and were frequently deletions (7). The tonBgene has recently been involved in iron transport: tonB mutants are chromium sensitive (16) and show decreased binding constants for iron in kinetic studies of iron transport (17). Gratia (6) described another class of mutants insensitive to colicins B and I which required methionine. The corresponding mutation was located in the section of the *E. coli* chromosome between *his* at min 39 and *str* at min 65.

We present evidence concerning the excretion of enterochelin by a series of newly isolated mutants insensitive to colicins B and I, as well as the physiology of these mutants and the location of the corresponding genes.

MATERIALS AND METHODS

Bacteria and phage. Bacterial strains are listed in Table 1 or have been described previously (8). Phage T5 and T6 were obtained from S. E. Luria; $\phi 80vir$, from E. R. Signer; $\lambda c/857$ and $\lambda c/60$, from R. Sussman; P1 and R17, from R. Schleif. Isolation of *exb* mutants from colicin-sensitive strains has been described (8).

Media. OM minimal medium contains per liter of

¹Present address: Department of Molecular Biology and Microbiology, Tufts University School of Medicine, 136 Harrison Avenue, Boston, Mass. 02111.

² Present address: Mount Sinai School of Medicine, 1200 Fifth Avenue, New York, N.Y. 10029. distilled water: 10.5 g of K_2 HPO₄; 4.5 g of KH_2 PO₄; 0.05 g of MgSO₄; 1.0 g of (NH₄)₂SO₄, and 1.6 mM sodium citrate. Other media have been described (8). Minimal media were supplemented with 0.2% glucose and 1 μ g of thiamine per ml.

Phage P1 was titered on LB agar with 2.5×10^{-4} M CaCl₂ and 0.1% glucose. Chromium sensitivity was determined on LB agar supplemented with 4×10^{-4} M CrCl₃.

Colicin techniques. Preparation and titration of colicin B and colicin inhibitor have been described (8).

Bacterial conjugation. Matings were performed according to Nagel de Zwaig and Luria (12). A recA derivative of strain LD60 exbB metL serA tsx was constructed by conjugation with Hfr 111 recA nalB and selection for recombinants resistant to T6 and to nalidixic acid and the recA phenotype (sensitivity to ultraviolet light or methylmethane sulfonate). Into one such recombinant, LD63 recA nalB exbB serA metL tsx, the episome F' KLF16 containing genes from metC to fuc was introduced by mating with strain KL110/KLF16, and merodiploids were selected by plating on minimal agar with methionine. The exconjugants that grew were the merodiploids LD63/ KLF16.

Transductions. Recipient cells for P1 transductions were grown in LB broth to log phase, and CaCl₂ was added to 5×10^{-3} M. Phage P1 grown on the appropriate donor was added at a multiplicity of 0.1 to 0.2 phage per cell. After 15 min at 37 C, samples of the mixture or controls were plated on selective media.

Selection of λ -induced tonB mutants. Log-phase cells of strain B(583) $\Delta 24$ (gal bio λatt) were infected with $\lambda cI857$ (temperature inducible) and grown 4 h at

Strain	Relevant characteristics	Source
GUC6	exbA ^b from C600	this paper
GUC12	exbA from C600	this paper
GUC41	exbB metL from C600	this paper
GUC49	colicin B insensitive, from C600	this paper
C600	thr leu tonA	L104
W4032	pro met tsx Hfr Cav	LA235
GUW5	exbA from W4032	this paper
PA309	thr leu trp his arg thi str F⁻	LA435
WA5028a	<i>trpA</i> double-point mu- tant	E. Signer
X5050	<i>lac-pro</i> deletion (<i>φ</i> 80d <i>lac</i>)	E. Signer
VXII	deletion from <i>trp</i> to <i>lac</i> in prophage, from X5050	E. Signer
M107	sul	E. Signer
B(583) Δ24	gal λ att bio deletion	D. Freifelder
3LH9	tonB (λcI857) from B(583) Δ24	this paper
P10	thr leu malB lac str ^R Hfr	C. Kennedy
LD28	exbB metL from P10	this paper
A2325	argE his thi pro trp mtl gal tsx F ⁻	D. Boyd
LD54	exbB metL from A2325	this paper
KL16	Hfr	LA761
AB856	serA6	B. Bachmann
LD60	serA exbB metL tsx from AB856	this paper
LD63	serA exbB metL recA nalB tsx from LD60 and Hfr 111	this paper
Hfr 111	recA nalB	R. Schleif
KL110/KLF16	chromosome: argG6 metB1 his-1 thy-23 leu-6 recA1; F': from AB312, metC ⁺ to fuc ⁺	B. Bachmann

 TABLE 1. Bacterial strains, E. coli K-12

^a Symbols refer to Luria stock collection.

^bGenetic abbreviations are described (15) or defined in this paper.

30 C. Samples were cospread with sufficient colicin B, $\phi 80\nu ir$ and $\lambda cI60$ so that, on control plates with any one of these three agents, most of the cells were killed. Surviving colonies were picked to duplicate grids on LB plates which were incubated at 30 or 42 C. Twelve ts isolates were obtained, three of which produced phage lambda and were considered to be tonB ($\lambda cI857$).

High-titer λ for transduction was prepared from one such strain 3LH9 and tested for ability to transduce Trp⁺ to strain WA5028a *trpA*. Transductants appeared at a frequency of 3×10^{-6} .

RESULTS

Isolation and phenotype of exb mutants. Independent spontaneous colicin-insensitive mutants of *E. coli* K-12 strain C600 were obtained (8) by selection with colicin produced by strains C1 139 (Col B, M), CA53 (Col Ia), C600 (Col Ib), or K94 (Col V). Mutants fell into various classes: insensitive to colicin B, I, or V; or to B and I, I and V, and B and V; or to all three colicins B, I, and V. Response of mutants to colicins Ia and Ib was identical. Some of these excrete large quantities of enterochelin, an inhibitor of colicins B and I, and have been named *exb* mutants (8). Some of these mutants have a requirement for methionine.

Enterochelin production in wild-type strains is repressed by exogenous iron (3,18). In *exb* strains, enterochelin is produced in high quantities in media containing up to 2 mM FeCl₃. Of the two classes of *exb* mutants distinguished genetically below, the *exbA* type produces a larger quantity of colicin inhibitor than the *exbB* (Fig. 1). For example, GUC6 *exbA* and GUC41 *exbB* produced 1,000- and 250-fold more enterochelin, respectively, than strain C600, as judged by titration of culture supernatant fluids against colicin B (8).

Map location of the exbA mutation. Strain GUW5exb was isolated from strain W4032 Hfr Cav pro met and crossed with the multiauxotrophic recipient PA309 (Table 2). The results, although not fully consistent, suggested that the exb gene might be most close to the trp locus. This in turn suggested that some exb mutants might map in the tonB gene, adjacent to the trp operon (15).

The tonB phenotype, resistance to phages T1 and ϕ 80, cannot be observed in mutants of C600, a tonA strain resistant to T1, T5, and ϕ 80.

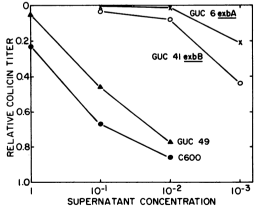


FIG. 1. Production of colicin inhibitor by insensitive mutants. Supernatant fluids of overnight cultures of the indicated strains grown in M9 medium with amino acids plus 200 μ M FeCl, were diluted in the same medium and assayed for colicin inhibitory content, plotted as relative colicin titer. Low colicin titers indicate high inhibitory power.

Selected No.	No.	Unselected markers (%)							
markers	tested	Thr ⁺ Leu ⁺	Arg ⁺	His ⁺	Trp+	Met-	Pro-	Exb	
Thr ⁺ Leu ⁺	100		14	0	0	32	72	15	
Arg ⁺	100	45		0	0	88	43	30	
His ⁺	9 8	15	19		14	17	22	56	
Trp+	98	14	8	16		5	16	88	

TABLE 2. Linkage of exbA and trp^a

^a Strain GUW5 Hfr Cav met pro exb-5 str^a was crossed with PA309 F⁻ thr leu arg his trp str^R. All selective media contained 50 μ g of streptomycin per ml.

To determine if exb mutants from C600 were tonB we used transduction of the trpA gene adjacent to tonB. Strain WA5028a, a doublepoint mutant in trpA, was infected with P1 grown on GUC12 exbA, GUC41exbB, or C600, and trp^+ transductants were selected and tested for phage and colicin sensitivity (Table 3). Eighty-four percent of trp^+ transductants from P1-GUC12 were tonB and exb. All transductants from C600 and GUC41 were wild type. Thus strain GUC12 exb is tonA tonB.

tonB mutants require supplementary iron for growth in minimal medium and are inhibited by chromium (16, 17). Growth of strain GUC12exbA was totally inhibited by CrCl₃ (Fig. 2) as were the $trpA^+$ tonB transductants from this strain. Strain GUC12 grew poorly in minimal medium, but grew at wild-type rates if supplemented with 100 μ M FeCl₃ (Fig. 3).

tonB mutants are exb. The above data indicate that certain mutants which we identify as exbA are tonB. To determine if this relationship is reciprocal. tonB mutants from other laboratories were tested. Strain VXII (provided by E. R. Signer) was prepared from X5050 by selection with colicin V and phage $\phi 80vir$ and contains a deletion extending from trp to lac in prophage $\phi 80 dlac$. Strain VXII was found to be colicin B insensitive and exb. Mu-induced tonB mutants (provided by M. Howe) were obtained from cells of strain M107 infected with phage mu by selection with colicin B and $\phi 80vir$. Phage mu was inserted in the tonB gene since $trpA^+$ tonB derivatives by P1 transduction were lysogenic for mu. Mu-induced tonB as well as spontaneous tonB mutants from infected cells produced high levels of colicin inhibitor (Table 4).

tonB phenotype can be produced by insertion of lambda. Since tonB mutants frequently carry deletions, the various effects of the mutation might reflect a loss of different genes rather than pleiotropic effects of a lesion in a single gene. One test for pleiotropic effects was to isolate chromium-resistant revertants from an exbA strain and test for the other properties.

TABLE 3. Co-transduction of exbA and trpA

Donor	Trp ⁺ transduc-	Se	tonB		
Donor		Col B	ø80vir	T5	(%)
C600	35	+	+	+	0
GUC12 exbA	27ª	-	_	+	84
	5°	+	+	+	0
GUC41 exbB	50	+	+	+	0

^a Six strains were tested and found to excrete inhibitor comparable to parent GUC12 exbA.

^o Two strains were tested and found to excrete inhibitor comparable to recipient parent WA5028a.

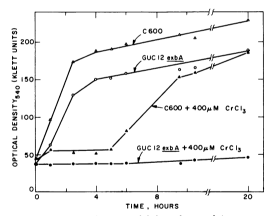


FIG. 2. Chromium sensitivity of an exbA mutant. Chromium chloride was added to nutrient broth cultures in early log phase.

Twelve such revertants from GUC12 exbAproved to be colicin insensitive, iron requiring, and exb, and yielded chromium-sensitive tonBstrains by transduction of $trpA^+$ to WA5028a. In these revertants, as possibly also in other tonBrevertants selected for chromium resistance, the resistance may have arisen by extragenic suppression.

To test in another way if all the tonB effects can result from disfunction within a single cistron, lambda-induced tonB mutants were obtained in a strain of *E. coli* K-12 with a

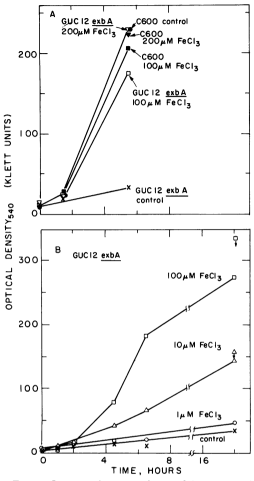


FIG. 3. Iron requirement of an exbA mutant. A, Strains were grown in M9 medium supplemented with FeCl_s as indicated. B, Strain GUC12, an exbA mutant from C600, was grown in M9 medium with 0, 1, 10, or 100 μ M FeCl_s added. Arrows indicate values corrected for pink color of the culture fluid.

deletion of the normal attachment site. Shimada et al. (13) demonstrated that infection of a strain lacking the normal lambda attachment site yielded some lysogens in which the prophage had become inserted inside bacterial genes inactivating them. Reversion to wild type results in simultaneous loss of the prophage, indicating that insertion and excision of lambda occur with exact precision. Lambda-induced tonB mutants were obtained in strain B(583) $\Delta 24(gal \ bio \ \lambda att)$ by infection with $\lambda cI857$ (temperature inducible) and selection with colicin B, $\phi 80vir$ and $\lambda cI60$. Three temperature-sensitive phage-producing strains were obtained. High-titer lambda, prepared from strain 3LH9, mediated specialized trp transduction, indicating that the prophage was inserted next to the trpA gene. These lambda-induced tonB mutants were insensitive to colicins B, Ia, and Ib, required iron for growth in minimal medium, and were chromium sensitive and exb (Table 4). These phenotypes therefore can result from a mutation in a single cistron, assuming that the insertion of lambda does not result in polarity effects.

Genetic analysis of exbB mutants. Exb Met⁻ strains are T1 sensitive, chromium resistant, and grow on minimal medium without supplementary iron. The ExbB phenotype is probably due to a deletion since no Met⁺ revertant colonies could be obtained by plating exbB cells on medium lacking methionine.

To locate the exbB gene on the E. coli chromosome, strain LD28 (an exbB mutant of Hfr strain P10 which inserts from min 79 in the order arg, mtl, his) was crossed with A2325 F⁻ argE his mtl; Arg⁺ and Mtl⁺ recombinants were selected at specific time intervals. The colicininsensitive phenotype appeared between 30 and 40 min after mixing the strains, and was more closely linked to *mtl* than to *arg*. Correction for a lag in entry of markers located the exbB gene between 49 and 59 min. Conjugation of strain LD54 F⁻ his exbB met with Hfr KL16, which inserts from min 55 with his^+ , an early marker, yielded his^+ recombinants that were exbB like the F^- parent. These data suggest that exbB is located between min 55 and 59.

Transduction of the exbB gene by phage P1 was performed to determine linkage to serA at min 57. Nine percent of $serA^+$ transductants (6 out of 66) with P1 grown on an exbB donor acquired colicin insensitivity and methionine

TABLE 4. Production of colicin inhibitor by mu-induced or lambda-induced tonB mutants

Supernatant fluid ^e	Inhibition of colicin (%)	
<i>tonB</i> (mu)	96	
ara-20 (mu)		
tonB spontaneous		
M107, parent		
3LH4 tonB (lambda)		
3LH5 tonB (lambda)	99	
3LH9 tonB (lambda)		
B(583) \triangle 24, parent		

^a Mu lysogens and controls were grown at 32 C in nutrient broth, and lambda lysogens and controls were grown at 30 C in M9 medium with 200 μ M FeCl_a and 1 μ g of biotin per ml. Supernatant fluids were diluted 1:10 and titered for ability to inhibit colicin B. auxotrophy. These transductants hyperexcreted colicin inhibitor in amounts comparable to the donor strain (Table 5). Thus, the exbB gene is located within about 1 min of *serA*.

The exbB phenotype is recessive. Strain LD63 recA tsx exbB met serA was mated with KL110/KLF16, which contains the F' episome F116 including genes from metC at min 59 to fuc at min 54. Presumed merodiploid recombinants were selected on minimal medium supplemented only with methionine. These recombinants were sensitive to colicin B, produced 100 times less colicin inhibitor than the exbB parent, and approximately 3 times more than the wild type as shown in Table 6, and did not require methionine.

Characteristics of the methionine auxotrophy of exbB mutants. exbB strain LD28 required at least 10 μ g of methionine per ml of minimal medium for optimal growth. The methionine requirement of strain GUC41 was compared to those of a *metA* and a *metE* mutant. GUC41 grew well with either homocysteine or cystathionine, suggesting that the auxotrophy is probably not due to a defect of conversion of cystathionine to methionine.

TABLE 5. Co-transduction of exbB and serA^a

Supernatant fluid	Phenotype	Inhibition of colicin (%)
Transductant-1	Col ¹ Met ⁻	96
Transductant-2	Col ¹ Met ⁻	97
Transductant-3	Col ⁹ Met ⁺	7
AB856 serA (recipient)	Col ⁹ Met ⁺	0.5
LD28 exbB (donor)	Col ¹ Met ⁻	99

^a Phage P1 grown on strain LD28 exbB was used to transduce SerA⁺ to strain AB856 serA. Transductants were tested for colicin sensitivity (Col¹ or Col⁸), for methionine requirement and for ability to excrete colicin inhibitor.

TABLE 6. Production of colicin inhibitor by exbB/exbB⁺ merodiploids^a

	Inhibition of colicin (%)				
Supernatant fluid	Undiluted	1:10	1:100		
Merodiploid-1	85	30			
Merodiploid-2	83	21			
Merodiploid-4	89	28			
LD63 exbB parent		97	84		
KL110/KLF16 parent	62	5			

^a Merodiploids were isolated from a mating of LD63 serA exbB metL recA nalB tsx with KL110/KLF16 F' episome from min 54 to 59. Merodiploids were tested for production of colicin inhibitor in M9 medium with 200 μ M FeCl₃ and 20 μ g of methionine per ml. Strains of *E. coli* and *S. typhimurium* known as *metK* mutants are characterized by loss of sensitivity to methionine analogs including ethionine and norleucine (9). The *exbB* mutant LD28 and its parent P10 were tested in minimal medium containing 5 mg of ethionine and 1 μ g of methionine per ml, and were fully sensitive to ethionine (Fig. 4). Analogous experiments with norleucine were inconclusive since the sensitive control strain grew at all concentrations in the presence of methionine.

DISCUSSION

In this paper we have presented evidence concerning two types of mutations which have been mapped in different regions of the *E. coli* chromosome and cause excretion of large quantities of enterochelin in the presence of sufficient iron to repress synthesis of this compound in wild-type cells. *exbA* mutants excrete 1,000 times more enterochelin than wild-type strains, and the corresponding mutations are located in

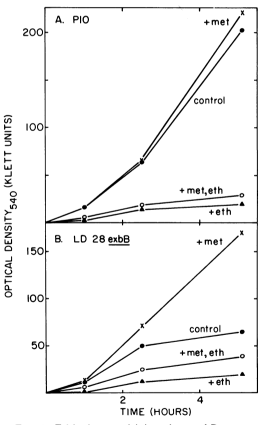


FIG. 4. Ethionine sensitivity of an exbB mutant. Cultures were diluted into M9 medium with or without methionine (1 μ g per ml) and ethionine (5 mg per ml). A, Parent strain P10. B, Strain LD28exbB.

the tonB gene, and tonB mutants isolated independently are exb. The exbB mutations that cause a 250-fold increase in enterochelin production are located between min 56 and 58, may be mainly deletions, and are recessive with respect both to colicin sensitivity and enterochelin excretion. Neither exb trait could result from an operator constitutive mutation since these mutations are not linked to the ent genes for enterochelin synthesis, which map at min 14 (5, 10).

The tonB gene is involved in iron uptake, hence enterochelin excretion by exbA mutants may be related to the internal iron pools of these cells. Bryce and Brot (3) have calculated that synthesis of 2,3-dihydroxybenzolyserine occurs in cells when the intracellular iron level falls below 22×10^{-19} mol per cell. Assuming an *E*. coli cellular volume of 1 μ m³, the internal iron level necessary to repress synthesis is 2.2 mM. It is possible that exbA strains cannot accumulate enough iron to repress enterochelin biosynthesis and are therefore acting as constitutive. Excretion by exbB mutants, which are not defective in iron transport, may be due to failure to synthesize an iron-sensitive aporepressor of the enterochelin operon.

The methionine lesion of exbB mutants is satisfied by homocysteine and cystathionine. This observation and the map position rule out a mutation in known methionine biosynthetic enzymes or a defect in metG which maps near the histidine operon in Salmonella typhimurium (14) and probably at a comparable position in E. coli (1). Mutants insensitive to the methionine analog ethionine, known as metK, map at min 57 (11). The sensitivity of exbB strains to ethionine rules out the identity of exbB with metK. One possible basis of the exbB methionine auxotrophy is that these mutants lack a positive controlling substance for transcription or translation of one or more of the genes involved in methionine biosynthesis. The methionine auxotrophy associated with the mutation may be called *metL*.

The tonB region is one of the classical deletion systems in E. coli (4), and the exact relationship of fine structure to function has remained obscure. Our data on the lambda-containing tonB strains indicate that a defect in a single cistron can cause loss of all the properties attributed to the tonB gene, assuming no polarity effects from the inserted prophage. Gratia (6) found that different combinations of T1 and the colicin sensitivity mapped in this region and that these phenotypes are recessive. The recessive character indicates that an operator defect cannot be invoked to explain the pleiotropic effects of tonB. Further work is needed to clarify the relationship of the *tonB* gene product to the mutant phenotypes.

ACKNOWLEDGMENTS

We thank D. Boyd and R. Schleif for suggestions, S. E. Luria for discussions and criticism of the manuscript, and C. L. Howitt for expert technical assistance. This investigation was initiated while one of us (S. K. G.) held NIH predoctoral fellowship 5-F1-GM-34, 383, and was further supported by NIH Biomedical Sciences Support Grant from Brandeis University (RR07044).

LITERATURE CITED

- Blumenthal, T. 1972. P1 transduction: formation of heterogenotes upon cotransduction of bacterial genes with a P2 prophage. Virology 47:76-93.
- Brot, N., J. Goodwin, and H. Fales. 1966. In vivo and in vitro formation of 2, 3-dihydroxybenzoylserine by E. coli K12. Biochem. Biophys. Res. Commun. 25:454-461.
- Bryce, G. F., and N. Brot. 1971. Iron transport in *E. coli* and its relation to the repression of 2,3-dihydroxy-*N*benzoyl-L-serine synthetase. Arch. Biochem. Biophys. 142:399-406.
- Coukell, M. B., and C. Yanofsky. 1971. Influence of chromosome structure on the frequency of tonB trp deletions in Escherichia coli. J. Bacteriol. 105:864-872.
- Cox, G. B., F. Gibson, R. K. J. Luke, N. A. Newton, I. G. O'Brien, and H. Rosenberg. 1970. Mutations affecting iron transport in *Escherichia coli*. J. Bacteriol. 104:219-226.
- Gratia, J. P. 1964. Résistance a la colicine B chez E. coli. Annu. Inst. Pasteur 107(Suppl. 5):132-151.
- Gratia, J. P. 1966. Studies on defective lysogeny due to chromosomal deletion in *Escherichia coli*. I. Single lysogens. Biken J. 9:77-87.
- Guterman, S. K. 1973. Colicin B: mode of action and inhibition by enterochelin. J. Bacteriol. 114:1217-1224.
- Lawrence, D. A., D. A. Smith, and R. J. Rowbury. 1968. Regulation of methionine synthesis in Salmonella typhimurium: mutants resistant to inhibition by analogues of methionine. Genetics 58:473-492.
- Luke, R. K. J., and F. Gibson. 1971. Location of three genes concerned with the conversion of 2,3-dihydroxybenzoate into enterochelin in *Escherichia coli* K-12. J. Bacteriol. 107:557-562.
- Maas, W. K. 1972. Mapping of genes involved in synthesis of spermidine in *Escherichia coli*. Mol. Gen. Genet. 119:1-9.
- Nagel de Zwaig, R., and S. E. Luria. 1967. Genetics and physiology of colicin-tolerant mutants of *Escherichia* coli. J. Bacteriol. 94:1112-1123.
- Shimada, K., R. A. Weisberg, and M. E. Gottesman. 1972. Prophage lambda at unusual locations. I. Location of the secondary attachment sites and the properties of the lysogens. J. Mol. Biol. 63:483-503.
- Smith, D. A., and J. D. Childs. 1966. Methionine genes and enzymes of Salmonella typhimurium. Heredity 21: 265-286.
- Taylor, A. L. 1970. Current linkage map of *Escherichia* coli. Bacteriol. Rev. 34:155-175.
- Wang, C. C., and A. Newton. 1969. Iron transport in Escherichia coli: relationship between chromium sensitivity and high iron requirement of mutants of Escherichia coli. J. Bacteriol. 98:1135-1141.
- Wang, C. C., and A. Newton. 1971. An additional step in the transport of iron defined by the tonB locus of Escherichia coli. J. Biol. Chem. 246:2147-2151.
- Young, I. G., and F. Gibson. 1969. Regulation of the enzymes involved in the biosynthesis of 2, 3-dihydroxybenzoic acid in Aerobacter aerogenes and Escherichia coli. Biochem. Biophys. Acta 177:401-411.