University of Nebraska - Lincoln DigitalCommons@University of Nebraska - Lincoln

Papers in Veterinary and Biomedical Science

Veterinary and Biomedical Sciences, Department of

March 1989

Impairment of Melibiose Utilization in *Streptococcus mutans* Serotype c *gtfA* Mutants

Raul G. Barletta University of Nebraska - Lincoln, rbarletta1@unl.edu

Roy Curtiss III Washington University, St. Louis, Missouri

Follow this and additional works at: https://digitalcommons.unl.edu/vetscipapers

Part of the Veterinary Medicine Commons

Barletta, Raul G. and Curtiss, Roy III, "Impairment of Melibiose Utilization in *Streptococcus mutans* Serotype c *gtfA* Mutants" (1989). *Papers in Veterinary and Biomedical Science*. 13. https://digitalcommons.unl.edu/vetscipapers/13

This Article is brought to you for free and open access by the Veterinary and Biomedical Sciences, Department of at DigitalCommons@University of Nebraska - Lincoln. It has been accepted for inclusion in Papers in Veterinary and Biomedical Science by an authorized administrator of DigitalCommons@University of Nebraska - Lincoln.

Impairment of Melibiose Utilization in Streptococcus mutans Serotype c gtfA Mutants

RAUL G. BARLETTA^{1,2}[†] and ROY CURTISS III^{2*}

Department of Microbiology, University of Alabama at Birmingham, Birmingham, Alabama 35294,¹ and Department of Biology, Washington University, St. Louis, Missouri 63130²

Received 16 September 1988/Accepted 21 November 1988

The Streptococcus mutans serotype c gtfA gene encodes a 55-kilodalton sucrose-hydrolyzing enzyme. Analysis of S. mutans gtfA mutants revealed that the mutant strains were specifically impaired in the ability to use melibiose as a sole carbon source. S. mutans gtfA mutant strains synthesized less α -galactosidase activity inducible by raffinose than wild-type strains. Melibiose (an inducer in wild-type strains) failed to induce significant levels of α -galactosidase in the mutant strains. We hypothesize that melibiose use by S. mutans requires the interaction of the GtfA enzyme, or another gene product under the control of the gtfA promoter, with other gene product(s) involved in melibiose transport or hydrolysis.

Sucrose metabolism plays a critical role in caries formation by *Streptococcus mutans* (15). Recently, the genes for several of these enzymes have been cloned and thoroughly analyzed (1, 10, 14, 21), yet there is no full description of the importance, function, and interactions of these enzymes (5).

We have previously reported the cloning, characterization, and properties of a unique S. mutans sucrose-hydrolyzing activity, the GtfA enzyme (21). It was hypothesized that the GtfA enzyme synthesizes a glucan primer for the synthesis of larger glucan products, perhaps for the synthesis of water-insoluble glucans (19-21). Further analysis of the product formed from sucrose by purified enzyme encoded by the ggg gene indicates that the GtfA enzyme is most likely a sucrose phosphorylase, which catalyzes the transfer of a glucosyl moiety from sucrose to P_i to generate α -D-glucose-1-phosphate and free fructose (22). Since both glucan and glucose-1-phosphate have similar mobilities in paper and thin-layer chromatography (21, 22), and they are also methanol insoluble, this new finding (22) is consistent with previous reports (19-21). However, the identity of the reaction products of the GtfA enzyme with sucrose as the substrate has not been fully solved. We have previously reported that the product made from sucrose had an apparent molecular mass of 1,500 kilodaltons (kDa) by gel filtration (21). Russell et al. (22) suggested that charge interaction between glucose-1-phosphate and the column matrix led us to an erroneous estimate of size. However, we used charged standards to calibrate the column, and most importantly, we analyzed the product synthesized early (30 min) in the reaction kinetics (21). Furthermore, the product synthesized late (18 h) in the reaction kinetics (Russell et al. analyzed the product of an overnight incubation) had a lower apparent molecular mass (less than 0.75 kDa; unpublished observations). Therefore, it is still possible for the GtfA enzyme to synthesize a phosphorylated glucan product with P_i as the initial acceptor. This phosphorylated glucan could then serve as the primer for the synthesis of other extracellular (19-21) or storage (22) glucan products or be degraded to glucose-1-phosphate (22).

On the other hand, the final elucidation of the function of

۱

the GtfA enzyme requires the biochemical analysis of S. *mutans* mutants with specific lesions in the gtfA gene. Such mutants have been constructed and partially analyzed (2, 19, 20). In this regard, we have shown that these mutants were fully virulent in the rat model system (2). This result suggests that either the GtfA enzyme is not a major virulence determinant or its function is or can be supplied by another gene. In this manuscript, we describe and analyze the unexpected inability of S. *mutans* gtfA mutants to use melibiose as a carbon source. We hypothesize that the GtfA enzyme, or a gene product regulated by the gtfA promoter, interacts with other proteins involved in melibiose transport or hydrolysis.

Bacterial strains are listed in Table 1. Growth of *Escherichia coli* and streptococci, complex media and their components (Difco Laboratories, Detroit, Mich.), and nutritional supplements (Sigma Chemical Co., St. Louis, Mo.) have been described before (21, 24).

Wild-type serotype c S. mutans strains are able to ferment and grow on various sugars as sole carbon sources (8). Therefore, we tested the ability of S. mutans gtfA mutants to utilize different carbohydrates as carbon sources. Carbohydrate utilization by S. mutans was evaluated by fermentation tests in thioglycolate medium plus purple agar base (16), supplemented with 1% fructose, 1% galactose, 1% lactose (4-O- β -D-galactopyranosyl-D-glucoside), 1% maltose (4-O- α -D-glucopyranosyl-D-glucoside), 1% melibiose (6-O- α -D-galactopyranosyl-D-glucoside), 1% raffinose (6-O- α -D-galactopyranosyl-D-glucoside), 1% raffinose (5- α -D-galactopyranosyl-D-glucoside),

Pfanstiehl Laboratories, Inc., Waukegan, Ill.), or 1% sucrose (β -D-fructofuranosyl- α -D-glucopyranoside; ultrapure; Schwarz/Mann, Cambridge, Mass., or Sigma). Plates were incubated anaerobically for 36 to 48 h at 37°C. Fermentation of the sugar was evidenced by a color change from purple to yellow. We also tested the ability of S. mutans strains to grow on defined FMC medium (24) with the same carbohydrates as sole carbon sources. Wild-type strains MT8148, UA101, UA130, and V403 fermented and grew on 1.0% (wt/vol) fructose, galactose, lactose, maltose, mannitol, melibiose, raffinose, sorbitol, and sucrose. On the other hand, S. mutans gtfA insertion-duplication mutant strain UAB751 and deletion mutant strain V1362 fermented and grew on all carbon sources except melibiose. Five other independent insertion-duplication mutant strains gave the same results (data not shown). The mutant strains were

^{*} Corresponding author.

[†] Present address: Department of Microbiology and Immunology, Albert Einstein College of Medicine, Bronx, NY 10461.

Strain	Phenotype or genotype	Source, reference, or derivation	
E. coli K-12			
M1900	F ⁻ lac Y96 glnV44? rpsL45 or rpsL110 metB1 melB4	R. Schmitt ^{a} (23)	
M2701	F ⁻ galK rpsL metBl melA7	R. Schmitt (23)	
Streptococci ^b			
MT8148	S. mutans serotype c, rough, Mel ⁺	18	
UA101	S. mutans serotype c, rough, Mel ⁺	P. Caufield ^c (17)	
UA130	S. mutans serotype c, smooth, Mel ⁺	P. Caufield (17)	
UAB751	S. mutans serotype c, smooth, gtfA Mel ⁻	Derived from UA130 (2)	
V403	S. mutans serotype c, smooth, GtfA ⁺	F. L. Macrina ^d (12)	
V1362	S. mutans serotype c, smooth, $\Delta gtfA$	Derived from V403 (20)	

TABLE 1. Bacterial strains

^a Institut für Biochemie Genetik und Mikrobiologie, Universität Regensburg, Regensburg, Federal Republic of Germany.

^b Rough or smooth phenotype refers to the colony morphology on brain-heart agar plates after 48 h of incubation at 37°C under anaerobic conditions. It is indicated for identification purposes and does not correlate with other traits described.

^c School of Dentistry, University of Alabama at Birmingham.

^d Department of Microbiology and Immunology, Virginia Commonwealth University, Richmond, Va.

incapable of fermenting melibiose when the sugar was added at higher concentrations (2 to 8% [wt/vol]). The inability of mutant strains to utilize melibiose suggested that the S. mutans gtfA gene, or a gene under the control of the gtfA promoter, was involved in melibiose metabolism. S. mutans gtfA mutants also displayed a reduced ability to grow on FMC-raffinose agar, suggesting a common pathway for melibiose and raffinose catabolism in S. mutans.

We performed reversion tests to verify the tight linkage between the presence of the GtfA enzyme and melibiose use by S. mutans. Insertion-duplication mutant strains were stable in the absence of antibiotic selective pressure (2). However, melibiose-utilizing (Mel⁺) true revertants were obtained at frequencies between 1.1×10^{-6} and 3.0×10^{-5} by plating S. mutans gtfA mutant cells onto FMC agar supplemented with 1% melibiose. Revertants were also obtained by repeated passages on thioglycolate medium plus purple broth (or agar) base with 1% melibiose. All revertants displayed the same patterns of carbohydrate utilization as wild-type strains and gave the wild-type colony size on FMC-raffinose agar. Furthermore, analysis of one revertant by Southern blotting analysis showed the loss of the DNA sequence inserted to inactivate the gtfA gene (data not shown). Six independent revertants analyzed by Western blot had a wild-type GtfA protein (data not shown).

Since S. mutans gtfA mutants were unable to utilize melibiose, we investigated whether the GtfA enzyme was able to transport and hydrolyze melibiose. We tested these properties of the GtfA enzyme in vivo by the use of appropriate Escherichia coli gtfA clones. Fermentation of E. coli Mel⁻ strains M1900 and M2700 (Table 1) and their transformants with pYA601 (GtfA⁺ [21]) was tested at 25 and 37°C in MacConkey base agar supplemented with various carbohydrates: 1% (wt/vol) galactose, 1% glucose, 1% lactose, 1% melibiose, 1% raffinose, or 1% sucrose. The wild-type gtfA gene failed to complement either a mutation in the α -galactosidase gene (melA) or a mutation in the thiomethylgalactoside permease II gene (melB). Since the E. coli melA mutant has all the machinery necessary to transport melibiose inside the cells, we concluded that the GtfA enzyme cannot hydrolyze melibiose. Furthermore, the GtfA enzyme is partly accessible to the periplasmic space (21) (see below), where melibiose can diffuse freely and react directly with the enzyme. The lack of complementation of the melB mutation shows that the GtfA enzyme, by itself, cannot function as a melibiose permease, at least not in E. coli. This result should be viewed with caution, since the S. mutans gene product might not behave in *E. coli* exactly as it would in *S. mutans*.

E. coli melA GtfA⁺ clones were also unable to ferment raffinose. This result indicated that fermentation of raffinose by E. coli GtfA⁺ clones is dependent on the hydrolysis of raffinose (into sucrose and galactose) by α -galactosidase, as we have suggested before (21). Furthermore, strains M1900(pYA601) and M2701(pYA601) grew on sucrose as the sole carbon source. Colonies on sucrose minimal agar were readily visible after 2 to 3 days of incubation at 37°C. No growth was observed when the host strains were incubated for 7 days in the same conditions. This result is consistent with the localization of the GtfA enzyme on the external side of the E. coli cytoplasmic membrane in a functional way so as to utilize sucrose as a substrate (21).

We investigated whether melibiose had any effect on sucrose hydrolysis by the GtfA enzyme. If the GtfA enzyme binds melibiose and sucrose through the same site, melibiose should competitively inhibit product (glucose-1-phosphate or a phosphorylated glucan) formation from sucrose. If it binds elsewhere, it should inhibit product formation noncompetitively. For the inhibition experiment, sucrose was used at 1.0 mM, a concentration slightly below the K_m of 1.25 mM (21). Melibiose was added to reaction mixtures at concentrations ranging from 12.5 to 100 mM. The unrelated sugar lactose was also tested at 100 mM. Quantitation of product synthesis for the various reaction mixtures (data not shown) showed that neither melibiose nor lactose had any inhibitory effect on product synthesis by the GtfA enzyme. This result suggests that the GtfA enzyme does not bind melibiose, at least not at a site that would impair sucrose hydrolysis.

Use of melibiose and raffinose by *S. mutans* requires their uptake and subsequent hydrolysis of the product of the transport reaction to monosaccharides. Alternatively, melibiose could be hydrolyzed by an extracellular α -galactosidase into galactose and glucose and metabolized via the homolactic fermentation pathway, which is prevalent in streptococci (6). The initial step in raffinose catabolism may involve various alternative routes: uptake of raffinose followed by intracellular hydrolysis, hydrolysis by an extracellular α -galactosidase into sucrose and galactose, or hydrolysis by extracellular invertase (4) or fructosyltransferase (11) into melibiose and fructose. The latter route may explain the reduced ability of *S. mutans* Mel⁻ mutants to grow on raffinose, as we have observed. So far, it remains unknown whether the transport of raffinose and melibiose also in-

Strain	Dhanatura	Sugar(s) added and concn (wt/vol)	Mean α -galactosidase activity (nmol of <i>o</i> -nitrophenol/ml per min) \pm SD	
	глепотуре		Intact cells	Permeabilized cells
UA130	GtfA ⁺ Mel ⁺	Galactose (1%)	1.9 ± 0.2	1.0 ± 0.2
		Galactose (1%), melibiose (0.1%)	4.0 ± 0.2	2.2 ± 0.2
		Galactose (1%), raffinose (0.1%)	6.2 ± 0.2	5.0 ± 0.2
		Melibiose (1%)	7.0 ± 0.2	5.8 ± 0.2
		Raffinose (1%)	11.5 ± 0.2	8.9 ± 0.2
UAB751	GtfA ⁻ Mel ⁻ (insertion-duplication mutant	Galactose (1%)	1.8 ± 0.2	0.6 ± 0.2
	derived from UA130)	Galactose (1%), melibiose (0.1%)	2.1 ± 0.2	0.7 ± 0.2
		Galactose (1%), raffinose (0.1%)	3.3 ± 0.21	1.6 ± 0.2
		Raffinose (1%)	6.0 ± 0.2	2.6 ± 0.2
V403	GtfA ⁺ Mel ⁺ (wild type)	Galactose (1%)	0.8 ± 0.2	<0.2
		Galactose (1%), melibiose (0.1%)	1.7 ± 0.2	0.2 ± 0.2
		Galactose (1%), raffinose (0.1%)	3.1 ± 0.2	0.6 ± 0.2
		Raffinose (1%)	4.9 ± 0.2	1.2 ± 0.2
V1362	GtfA ⁻ Mel ⁻ (deletion mutant derived	Galactose (1%)	0.4 ± 0.2	0.2 ± 0.2
	from V403)	Galactose (1%), melibiose (0.1%)	0.6 ± 0.2	< 0.2
		Galactose (1%), raffinose (0.1%)	1.5 ± 0.2	0.3 ± 0.2
		Raffinose (1%)	3.0 ± 0.2	0.7 ± 0.2

TABLE 2. Measurement of α -galactosidase activity in S. mutans wild-type and gtfA mutant strains^a

^a S. mutans cells were grown at 37°C to the stationary phase as standing cultures in modified FMC medium (containing 0.5% casein hydrolysate instead of the standard amino acid mixture) supplemented with various sugars as indicated. Erythromycin (25 μ g/ml) was added for growth of strain UAB751 (2). Cells (from 50-ml cultures) were harvested at 4°C and washed once with 5 ml and resuspended in 2 ml of buffered saline with 0.01% gelatin. Cells were permeabilized as described by LeBlanc et al. (13). All samples were kept at 2°C pending assay. α -Galactosidase activity was measured by the hydrolysis of σ -nitrophenyl- α -D-galactoside (α -ONPG; Sigma) to the chromogenic product σ -nitrophenol. All assays were run in duplicate, and the final data were calculated from averages for at least two time points when the hydrolysis of α -ONPG was linear with time and cell concentration. The assay mixture contained 50 mM Tris hydrochloride (pH 7.5), 5.0 mM MnCl₂, 5.0 mM dithiothreitol, 5.0 mM α -ONPG, 15 µg of chloramphenicol per ml (to prevent α -galactosidase to permeabilized cells at an optical density of approximately 1.0 at 600 nm. Reactions were terminated by adding stop reagent (166 mM sodium carbonate and 8.3 mM EDTA at final concentrations). The amount of σ -nitrophenol was determined by measuring the A_{420} .

volves the phosphoenolpyruvate:sugar phosphotransferase system, responsible for the translocation of all major sugars catabolized by *S. mutans* (9).

Since α -galactosidase levels might be an important parameter in melibiose and raffinose use by S. mutans, we measured α -galactosidase activity in S. mutans wild type, gtfA insertion-duplication mutants (2), and gtfA deletion mutants (20). The results (Table 2) showed that (i) wild-type and mutant strains displayed approximately the same level of basal activity in FMC medium with 1% galactose; (ii) wildtype strains displayed a moderate but significant increase (about four to nine times the basal levels) of α -galactosidase activity when grown in the presence of 1% melibiose or 1% raffinose (raffinose was a slightly better inducer); (iii) addition of 0.1% melibiose or 0.1% raffinose to FMC medium with 1% galactose also raised the α -galactosidase levels in wild-type strains, but only to two to five times the basal levels; (iv) melibiose failed to induce α -galactosidase activity in S. mutans gtfA mutants, and raffinose induced only between 30 and 60% of the enzyme levels of their wild-type parents grown under the same conditions; and (v) intact cells displayed higher (up to three times) enzyme activities than permeabilized cells, suggesting that S. mutans α -galactosidase activity is (at least in part) accessible from the cell surface.

In summary, the effect of a mutation in the gtfA gene seemed to be quite specific on melibiose utilization, since S. *mutans* gtfA mutants were able to grow on various other sugars as sole carbon sources. The tight linkage between the gtfA gene and melibiose use was further verified by the isolation of S. *mutans* Mel⁺ revertants. All revertants had a wild-type GtfA enzyme. The impairment of melibiose use in

S. mutans gtfA mutants could be a direct consequence of inactivation of the gtfA gene or of a polar effect on the expression of another gene. In this regard, Burne et al. identified a gene coding for a 38-kDa protein following the gtfA gene (3). DNA sequence analysis of this region seemed to indicate that the 38-kDa protein gene is under transcriptional control by the gtfA gene (7). However, analysis of S. mutans $gtfA^+/gtfA$ merodiploid strains (unpublished observations), constructed from insertion-duplication mutants, suggests a direct involvement of the gtfA gene. The studies with E. coli Mel⁻ mutants showed that melibiose is not hydrolyzed or transported by the GtfA enzyme. Furthermore, melibiose had no effect on sucrose hydrolysis by the GtfA enzyme. The analysis of α -galactosidase activity in S. mutans gtfA mutant and parent strains showed that S. mutans may have two α -galactosidase activities, one inducible by melibiose and raffinose and another inducible by raffinose only. Induction of the former α -galactosidase activity by melibiose or raffinose might require the cooperation of the GtfA enzyme with the melibiose transport or hydrolysis machinery. In this regard, the finding of Russell et al. (22) that the GtfA enzyme is a sucrose phosphorylase led us to speculate that the GtfA enzyme might phosphorylate melibiose (without hydrolysis of the glycosidic bond) at the reducing carbon of the glucose moiety.

We thank Bruce Chassy, Raul Goldschmidt, Madelon Halula, Francis Macrina, Michael Pucci, and Jonathan Reizer for useful discussions and suggestions. We also thank Michael Pucci and Francis Macrina for generously providing streptococcal strains. We thank Jorge Galan, Paul Gulig, Michael Hudson, and Hank Lockman for critical review of the manuscript. We thank Jolaine Lauridsen and Betty Donovan at the Albert Einstein College of Medicine for expert secretarial assistance in preparing the manuscript.

Research was supported by Public Health Service grant DE06801 from the National Institute of Dental Research.

LITERATURE CITED

- 1. Aoki, H., T. Shiroza, M. Hayakawa, S. Sato, and H. K. Kuramitsu. 1986. Cloning of a *Streptococcus mutans* glucosyltransferase gene coding for insoluble glucan synthesis. Infect. Immun. 53:587-594.
- Barletta, R. G., S. M. Michalek, and R. Curtiss III. 1988. Analysis of the virulence of *Streptococcus mutans* serotype c gtfA mutants in the rat model system. Infect. Immun. 56: 322-330.
- Burne, R. A., B. Rubinfeld, W. H. Bowen, and R. E. Yasbin. 1986. Tight genetic linkage of a glucosyltransferase and dextranase of *Streptococcus mutans* GS-5. J. Dent. Res. 65:1392–1401.
- Chassy, B. M., R. M. Bielawski, J. R. Beall, E. V. Porter, M. I. Krichevsky, and J. A. Donkersloot. 1974. Extracellular invertase in *Streptococcus mutans*. Life Sci. 15:1173–1180.
- 5. Curtiss, R., III. 1985. Genetic analysis of *Streptococcus mutans* virulence. Curr. Top. Microbiol. Immunol. 118:253–277.
- Drucker, D. B., and T. H. Melville. 1968. Fermentation endproducts of cariogenic and non-cariogenic streptococci. Arch. Oral Biol. 13:565-570.
- Ferretti, J. J., T.-T. Huang, and R. R. B. Russell. 1988. Sequence analysis of the glucosyltransferase A gene (gtfA) from Streptococcus mutans Ingbritt. Infect. Immun. 56:1585–1588.
- Hamada, S., and H. D. Slade. 1980. Biology, immunology, and cariogenicity of *Streptococcus mutans*. Microbiol. Rev. 44: 331-384.
- 9. Hamilton, I. R., and G. C. Y. Lo. 1978. Coinduction of β galactosidase and the lactose-P-enolpyruvate phosphotransferase system in *Streptococcus salivarius* and *Streptococcus mutans.* J. Bacteriol. 136:900-908.
- Hayakawa, M., H. Aoki, and H. K. Kuramitsu. 1986. Isolation and characterization of the sucrose 6-phosphate hydrolase gene from *Streptococcus mutans*. Infect. Immun. 53:582-586.
- 11. Hojo, S., M. Mitsutomi, and T. Yamada. 1987. Metabolism of glycosylsucrose by oral microorganisms and its hydrolysis by *Streptococcus salivarius* fructosyltransferase. Infect. Immun. 55:698-703.

- 12. Kral, T. A., and L. Daneo-Moore. 1980. Glycerol incorporation in certain oral streptococci. Infect. Immun. 30:759-765.
- LeBlanc, D. J., V. L. Crow, L. N. Lee, and C. F. Garon. 1979. Influence of the lactose plasmid on the metabolism of galactose by *Streptococcus lactis*. J. Bacteriol. 137:878–884.
- 14. Lunsford, R. D., and F. L. Macrina. 1986. Molecular cloning and characterization of *scrB*, the structural gene for the *Streptococcus mutans* phosphoenolpyruvate-dependent sucrose phosphotransferase system sucrose-6-phosphate hydrolase. J. Bacteriol. 166:427-434.
- 15. Michalek, S. M., J. R. McGhee, T. Shiota, and D. Devenyns. 1977. Low sucrose levels promote extensive *Streptococcus mutans*-induced dental caries. Infect. Immun. 16:712-714.
- Murchison, H., S. Larrimore, and R. Curtiss III. 1981. Isolation and characterization of *Streptococcus mutans* mutants defective in adherence and aggregation. Infect. Immun. 34:1044– 1055.
- Murchison, H. H., J. F. Barrett, G. A. Cardineau, and R. Curtiss III. 1986. Transformation of *Streptococcus mutans* with chromosomal and shuttle plasmid (pYA629) DNAs. Infect. Immun. 54:273-282.
- Okahashi, N., H. Asakawa, T. Koga, N. Masuda, and S. Hamada. 1984. Clinical isolates of *Streptococcus mutans* serotype c with altered colony morphology due to fructan synthesis. Infect. Immun. 44:617-623.
- 19. Pucci, M. J., and F. L. Macrina. 1985. Cloned gtfA gene of Streptococcus mutans LM7 alters glucan synthesis in Streptococcus sanguis. Infect. Immun. 48:704-712.
- Pucci, M. J., and F. L. Macrina. 1986. Molecular organization and expression of the gtfA gene of Streptococcus mutans LM7. Infect. Immun. 54:77-84.
- Robeson, J. P., R. G. Barletta, and R. Curtiss III. 1983. Expression of a Streptococcus mutans glucosyltransferase gene in Escherichia coli. J. Bacteriol. 153:211-221.
- Russell, R. R. B., H. Mukasa, A. Shimamura, and J. J. Ferretti. 1988. Streptococcus mutans gtfA gene specifies sucrose phosphorylase. Infect. Immun. 56:2763-2765.
- Schmitt, R. 1968. Analysis of melibiose mutants deficient in α-galactosidase and thiomethylgalactoside permease II in *Escherichia coli*. J. Bacteriol. 96:462-471.
- Terleckyj, B., N. P. Willet, and G. D. Shockman. 1975. Growth of several cariogenic strains of oral streptococci in a chemically defined medium. Infect. Immun. 11:649–655.