Analysis of an Agmatine Deiminase Gene Cluster in 
Streptococcus mutans UA159

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Received 19 September 2003/Accepted 3 December 2003

An operon encoding enzymes of the agmatine deiminase system (AgDS) has been identified in the cariogenic bacterium Streptococcus mutans UA159. The AgDS is regulated by agmatine induction and carbohydrate catabolite repression. Ammonia is produced from agmatine at low pH, suggesting that the AgDS could augment acid tolerance.

Streptococcus mutans is the etiological agent of dental caries (12). A key virulence attribute of this organism is acid tolerance, since catabolism of dietary carbohydrates lowers the pH of dental plaque to values below 4.0 (5). A primary determinant of acid tolerance is the membrane-bound F1F0-ATPase, although reduction in the proton permeability of the cell membrane and induction of DNA repair pathways and stress proteins also contribute to the acid tolerance of S. mutans (17).

Another widely used mechanism for acid resistance in less acid-tolerant oral streptococci is the generation of ammonia by the arginine deiminase system (ADS) (6). S. mutans metabolizes a wider variety of carbohydrates than many gram-positive bacteria, but the ADS is not present in this organism (1, 15). Consequently, when a cluster of genes with similarity to ADS genes (1) was identified in the S. mutans UA159 genome, it was proposed that the genes encoded enzymes of the agmatine deiminase system (AgDS) (6). The AgDS is analogous to the ADS but is involved in catabolism of agmatine (Fig. 1) (1), a decarboxylated derivative of arginine (19).

In lactic acid bacteria and Enterococcus faecalis, the AgDS resembles the ADS. However, in Pseudomonas aeruginosa and other organisms, agmatine is produced from arginine via an intracellular arginine decarboxylase, and putrescine is converted to spermidine or broken down to succinate. The identification of an ADS-like gene cluster in S. mutans is significant because agmatine catabolism may contribute to acid tolerance. Production of alkali from agmatine may have critical implications for the pathogenicity of S. mutans and may profoundly affect the ecology of oral biofilms. Thus, we have initiated an analysis of the AgDS gene cluster.

Analysis of the sequence of the AgD gene cluster. The first gene in the operon, otcA, encodes a putative putrescine carbamoyltransferase (PTC) and has been redesignated aguB. AguB is 80% identical to ArgF-2, annotated as an ornithine carbamoyltransferase (OTC) in E. faecalis V583 (Fig. 2) (14). ArgF-2 is one of two putative OTCases encoded by the E. faecalis genome (14), although argF-2 is located in an operon encoding an apparent AgD, carbamate kinase, and antipporter.

The other OTC gene is in the ADS operon (4), so ArgF-2 may be a PTC (18). The PTC amino acid sequence is not available, but the enzyme is well characterized and distinct from OTC (21).

According to Prosite (http://www.expasy.org/prosite), the carbamoyltransferase consensus is F-X-(E/K)-X-S-(G/T)-R-T, with the third residue differentiating aspartate (E) and ornithine (K) carbamoyltransferases. This consensus is present in the amino acid sequence of the S. mutans aguB product, as well as E. faecalis ArgF-2, Lactobacillus lactis subsp. lactis OtcA, Listeria monocytogenes LMO0036, and Lactobacillus sakei ArgF, with a highly conserved Q in the third position, perhaps reflecting a preference for putrescine. The HPTQ residues at positions 143 to 146, which are involved in carbamoylphosphate binding, are present in AguB, as are the HCLP residues at positions 281 to 284, which facilitate ornithine binding (11). Since the structures of ornithine and putrescine are identical at the position of cleavage, this conservation is not surprising. The second gene in the operon, SMU.263, encodes a putative amino acid antipporter and has been redesignated aguD. Similar to the case of other amino acid antipporters, 11 transmembrane helices were predicted for the amino acid sequence of AguD by using the “DAS” - Transmembrane Prediction server at http:

![FIG. 1. Comparison of the ADS and AgDS pathways. Agmatine can be produced from arginine by arginine decarboxylase. AD, arginine deiminase; CK, carbamate kinase.](http://jb.asm.org/)
The third gene, SMU.264, encodes AgD and has been redesignated aguA. The GGGNI- HCITQQ sequence (13) was identified at the C terminus of AguA. The final gene, arcC, encodes a carbamate kinase and has been redesignated aguC to reflect its association with the AgDS.

In addition to aguBDAC, a transcriptional regulator of the LuxR family is located 239 bp upstream of aguB and transcribed in the opposite direction. LuxR-type proteins belong to the FixJ-NarL superfamily, mainly composed of two-component response regulators (10). Homologs of this regulatory protein were identified 399 and 234 bp upstream of the AgDS gene clusters in E. faecalis and L. lactis subsp. lactis, respectively. Involvement of this protein in AgDS regulation is under investigation.

Cotranscription of aguBDAC. No terminator-like sequences were identified in the agu intergenic regions. To demonstrate cotranscription of aguBDAC, a polar mutation in aguB was constructed by insertion of an Owon cassette harboring strong transcription/translation termination signals (16). Northern blotting was performed (3) with 10 μg of total RNA extracted (8) from wild-type and aguB mutant cells grown in low-carbo-
AgDS expression in *S. mutans*. To quantify mRNA under different growth conditions, slot blot analysis was used. Total RNA extracted from cells grown in TV broth plus 0.5% glucose or 0.5% galactose, with or without 10 mM agmatine, was transferred to a 0.45-μm nitrocellulose membrane. The RNA was hybridized to an *aguB* probe. AgDS-specific mRNAs were detected under all conditions, although expression was severalfold higher when cells were grown in medium containing agmatine (Fig. 4).

AgD activity in permeabilized cells grown under different growth conditions was measured by colorimetric determination of N-carbamoylputrescine production from agmatine (Fig. 5A) (2). In wild-type *S. mutans*, peak AgD activity was observed in cells grown in TV containing the non-catabolite-repressing sugar galactose and agmatine, whereas activity was 65% lower in cells grown in the repressing sugar glucose and agmatine. Thus, the operon is inducible by agmatine and under the control of carbohydrate catabolite repression. No AgD activity was observed only in cells grown with agmatine (Fig. 5B).

Ammonia production was measured after cells were incubated with buffer and 10 mM agmatine, using an ammonia detection kit (Diagnostic Chemicals Limited, Charlottetown, Canada). Consistent with AgD activity, ammonia production was observed only in cells grown with agmatine (Fig. 5B). Ammonia production increased two-thirds in cells grown in galactose and agmatine, compared to glucose and agmatine. The optimum pH for ammonia production from agmatine was 4 (Fig. 6).

Summary. AgDS of *S. mutans* is functional and under tight genetic control. The system is expressed at a low level relative to urease and arginine deiminase of other oral streptococci, and it is unlikely that agmatine catabolism would cause significant environmental alkalization. However, ammonia production by the AgDS at low pH would increase ΔpH and provide ATP, which could be used for growth or to extrude protons, and the system is capable of functioning at pH values that are considered extreme in dental plaque (20). Therefore, the AgDS may represent a significant contributor to acid tolerance and thus to the virulence of *S. mutans* in vivo.

Nucleotide sequence accession number. The sequences of these genes have been deposited with GenBank and bear accession number BK004003.

This research was supported by Public Health Service grant DE10362 from the National Institute of Dental and Craniofacial Research.

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