Role of HtrA in Surface Protein Expression and Biofilm Formation by Streptococcus mutans

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The HtrA surface protease in gram-positive bacteria is involved in the processing and maturation of extracellular proteins and degradation of abnormal or misfolded proteins. Inactivation of htrA has been shown to affect the tolerance to thermal and environmental stress and to reduce virulence. We found that inactivation of Streptococcus mutans htrA by gene-replacement also resulted in a reduced ability to withstand exposure to low and high temperatures, low pH, and oxidative and DNA damaging agents. The htrA mutation affected surface expression of several extracellular proteins including glucan-binding protein B (GbpB), glucosyltransferases, and fructosyltransferase. In addition, htrA mutation also altered the surface expression of enolase and glyceraldehyde-3-phosphate dehydrogenase, two glycolytic enzymes that are known to be present on the streptococcal cell surface. As expected, microscopic analysis of in vitro grown biofilm structure revealed that the htrA-deficient biofilms adopted a much more granular patchy appearance, rather than the relatively smooth confluent layer normally seen in the wild type. These results suggest that HtrA plays an important role in the biogenesis of extracellular proteins including surface associated glycolytic enzymes and in biofilm formation of S. mutans.

Streptococcus mutans has been strongly implicated as the principal etiological agent in human dental caries (44). In addition to dental caries, S. mutans is also an important agent of infective endocarditis (4, 79). More than 20% of cases of viridans streptococcus-induced endocarditis are caused by S. mutans (29, 44). S. mutans expresses plethora surface proteins, and many of them are virulence factors. These include adhesins, specialized transport systems for fermentable sugars, synthesis of soluble and insoluble dextrans and fructans, and factors required for acidogenesis and acidouricity (38). Various cell-surface substances, including serotype-specific polysaccharide antigens, lipoteichoic acid, glucosyltransferases (GTFs), fructosyltransferase (FTF), dextranase, glucan-binding proteins, a 29-kDa protein WapA, and a 190-kDa SpaP (also known as protein P1, PAc, antigen B, and antigen I/II) are thought to play important roles in interaction between the organism and its host (35, 38), and have been considered for vaccine candidates for dental caries (for review, see reference 34).

Biofilms consist of complex mixture of microorganisms that adhere to each other and in most cases to a surface. One of the important virulence properties of mutants streptococci is their ability to form biofilms along with other bacteria (38, 83). This biofilm, known as dental plaque, is one of the best-studied biofilms (20, 36). Biofilm formation is considered to be a two-step sequential process requiring early attachment of the bacterial cells to a surface, followed by growth dependent multilayer accumulation of bacteria involving intra- and intergenic cell-to-cell interactions and adhesions (20). There have been considerable efforts to identify factors of S. mutans and other oral streptococci that are involved in biofilm initiation and development. Surface-associated proteins, such as SpaP and Fap1, function as high-affinity adhesins and are required in the initiation of biofilm (12, 23, 53). Extracellular glucans, synthesized from sucrose by GTFs, are key players in adhesion interactions and accumulation of mutants streptococci on smooth surfaces (38). Glucan-binding proteins (Gbp) such as GbpC are involved in rapid, dextran-dependent aggregation during biofilm formation (70). Interestingly, immunization with GbpB, another glucan-binding protein, induces an immune response in rats that interferes with the accumulation of S. mutans and reduces the level of dental caries (73). There are numerous cell-surface and extracellular proteins that work in concert to successfully establish S. mutans in tooth biofilms (38).

In bacteria, proteolysis plays important roles in many biological processes such as post-translational regulation of gene expression. This includes processing and maturation of various surface associated proteins in the case of gram-positive bacteria (26, 40). In S. mutans, cell surface protein SpaP is found free in culture supernatant, along with a number of lower-molecular-mass forms (65). Another surface associated protein WapA, is also released by proteolytic cleavage (22) and two cell-wall-associated enzymes, dextranase and fructanase, are found predominantly in culture supernatants (14, 31, 32). In addition to the proteolytic events that release these cell-surface-associated proteins and enzymes, extracellular GTFs and FTF are also broken down to lower-molecular-mass-forms (1, 68) and function as glucan-binding proteins (67). Thus, expression of various surface proteins is dependent on proteolysis, which can strongly influence both the level of activity and the cellular localization of these proteins.

There are several different proteases present in various...
streptococci (for reviews, see references 19 and 62). Among them, the trypsin like serine protease HtrA (also known as DegP) (for reviews, see references 18 and 54) is of scientific interest for several reasons including its role in cellular physiology, in particular in helping bacteria to survive environmental stresses such as elevated temperature, oxidative and osmotic stresses. HtrA homologs have been identified in phylogenetically distinct bacteria including many gram-positive pathogens suggesting a fundamental role of HtrA in response to environmental stress and possibly in pathogenesis. The proteins characteristically possess an amino-terminal hydrophobic region, a trypsin-like catalytic domain with conserved His, Ser and Asp residues and a PDZ domain thought to be involved in protein-protein interactions (69). HtrA proteins have been shown to have a housekeeping function, acting as chaperones and degrading misfolded proteins (74). Interestingly, in Lactococcus lactis, a closely related bacterium to S. mutans, HtrA was shown to be involved in the processing of propeptides and in the maturation of a natural extracellular protein (59). HtrA was thus proposed to be involved not only in degradation of misfolded proteins, but also in the processing/maturation of normal proteins. The Streptococcus pyogenes HtrA was also shown to be involved in production of several virulence factors whose biogenesis requires extensive processing. This includes a cysteine protease SpeB and a hemolysin SLS (46). Recently, the Staphylococcus aureus HtrA was also shown to control expression of several secreted virulence factors, including hemolysins (63). Thus, it appears that HtrA is involved in bacterial pathogenesis or by modulating virulence factor expression.

The regulation and organization of the htrA locus are very different in various gram-positive bacteria. For example, Bacillus subtilis and S. aureus both have more than one functional htrA gene and they are located far from one another in the genome (51, 52, 63). In streptococci, a single copy of htrA is present and it is located near the chromosomal replication origin. The immediate downstream genes are highly conserved and involved in cell division (24), but the organization of genes upstream of the htrA locus is not well conserved (Fig. 1).

Because htrA is located near the replication origin, inactivation by insertional mutagenesis may alter the efficiency of cell division by interfering with the downstream dnaA gene and can cause growth defects as proposed for S. pyogenes (46).

Based on the studies in other bacteria, one would expect S. mutans HtrA to be involved in the processing of extracellular (exported/cell-wall-associated) proteins including adhesins that are involved in biofilm formation and pathogenesis. How-
ever, a recent study with a single-crossover htrA mutant of S. mutans showed no significant effect on expression or processing of several extracellular proteins (21). The aim of this present study was to further investigate the role of HtrA on surface protein expression in S. mutans. In particular, the contribution of HtrA to the development of biofilm formation by this organism was examined. As previously observed, a mutant of HtrA showed several stress sensitive phenotypes (21). However, in contrast to the previous study, we found that mutants of HtrA displayed altered expression of several surface proteins including various surface associated glycolytic enzymes as well as formation of a biofilm with altered architecture.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** E. coli strain DH5α was grown in Luria-Bertani medium, and, when necessary, ampicillin (100 μg ml⁻¹), kanamycin (100 μg ml⁻¹), and/or spectinomycin (100 μg ml⁻¹) were included. S. mutans strains were routinely grown in Todd-Hewitt medium (BBL; Becton Dickinson), supplemented with 0.2% yeast extract (THY). When necessary, kanamycin (300 μg ml⁻¹) and/or spectinomycin (300 μg ml⁻¹) was included. For growth experiments involving stress tolerance, THY agar plates (1.5%) containing NaCl (0.5 to 1.5 M), puromycin (0.5 to 2.0 μg ml⁻¹; Sigma-Aldrich), H₂O₂ (1.0 to 2.5 mM; Sigma-Aldrich), mitomycin C (7.5 to 15 ng ml⁻¹; Sigma-Aldrich) were poured (we observed that the activity of puromycin and H₂O₂ varied between batches). Freshly grown overnight cultures were spun down and washed with sterile 0.9% saline. Cultures were resuspended in either the same or 1/10 of the original volume in 0.9% saline and 10-fold serial dilutions were made. A 10-μl aliquot of each dilution was spotted on the THY agar plates containing different stressors and plates were incubated at 37°C aerobically.

For growth experiments involving pH, initial pHs of THY agar were adjusted to pHs 5.5, 6.0, and 7.0 with HCl before sterilization. 50 mM citrate-phosphate buffer of desired pH was added to media after sterilization. Different dilutions of bacterial cultures were spotted as described above and plates were incubated at 37°C anaerobically. Hydrophobicity assay was done as described previously (71).

For some growth experiments and biofilm assay, semidefined biofilm medium (BM) (45) was also used supplemented with 1% glucose or 1% sucrose as the carbon source. Preparation of competent cells and transformation of S. mutans were done as described previously (15, 58).

**Construction of strains.** The htrA gene was insertional inactivated in NG-8 strains by gene-replacement. Based on the UA159 genome sequence information (GenBank accession no. AE015037), a 1.9-kb DNA fragment containing the htrA gene was PCR amplified from NG-8 genomic DNA using primers HtrA-For (5′-GACTACGATTATTTTGGAATTTTCTCATG-3′) and HtrA-Rev (5′-GTCACCGATTGTGCTCATATCTCACCCTC-3′). The DNA fragment was cloned into plasmid pUC4-Km upon digestion by SmaI was then cloned into the pGEMT-Easy TA cloning vector (Promega), and the resulting construct was confirmed by restriction analysis. A kanamycin-resistant cassette (1,000 bp) (57) isolated from plasmid pUC4-Km upon digestion by Smal was then ligated into a unique Pael site (restricted and blunted by T4 polymerase) within the htrA coding sequence (at amino acid position 243), and the resulting construct was named pBlb101. Plasmid pBlb101 was linearized by NotI and then used to transform S. mutans NG-8 strains according to a previously published protocol (15). Kanamycin-resistant transformants were selected on THY agar containing appropriate antibiotic, and PCR analysis was done to confirm that htrA inactivation had occurred by double-crossover recombination.

To express the htrA gene in trans, the full-length htrA (with promoter) was amplified by PCR using primers HtrA-For and HtrA-Rev and cloned into pTA28 (78) that had been restricted by Smal. The resulting plasmid contained the intact htrA gene under its own promoter.

**Preparation of whole-cell extract, cell wall, and supernatant proteins.** Overnight cultures were grown in THY, collected by centrifugation, and washed twice in phosphate-buffered saline (PBS). Cell density was adjusted to 5.0 cell units ml⁻¹. The cells were lysed by adding mutanolysin and lysozyme as described previously (10). For culture supernatant proteins, cells were removed from fresh overnight culture by centrifugation followed by filtration through a 0.2-μm pore size filter. Proteins were obtained by precipitation with trichloroacetic acid (TCA, 20% [wt/vol] final concentration) on ice for 2 to 4 h, washed with acetone, and resuspended in PBS or 1 × gel loading buffer (NEB) to 1/40 volume. Equal amounts of proteins (as measured by Bradford assay) were loaded in each lane for Western analysis.

**Western blot analysis.** Proteins were separated by sodium dodecyl sulfate–4 to 20% polyacrylamide gel electrophoresis (SDS-PAGE) and electroblotted onto a nitrocellulose membrane and blocked with 3% bovine serum albumin (BSA)–0.1% Tween 20 in Tris-buffered saline for 1 h at room temperature. Polyclonal antibodies to S. mutans glyceraldehyde-3-phosphate dehydrogenase (GAPDH) or enolase (kindly provided by Vijay Panholi) and FTF (kindly provided by Gilad Bachrach) were added (1:1,000 dilutions) and incubated for 1 h at room temperature. Membranes were washed thoroughly with 0.1% Tween 20 in Tris-buffered saline, probed with anti-rabbit immunoglobulin G-alkaline phosphatase-conjugated secondary antibodies for 1 h, and washed again. Reactivity was detected using nitroblue tetrAZolium and 5-bromo-4-chloro-3-indolyl phosphate (BCIP) as substrates.

**Detection of proteins on immobilized S. mutans cells.** S. mutans strains grown overnight were collected by centrifugation and washed twice with PBS. Cell density was adjusted to 2.0 cell units ml⁻¹ and serial dilutions were made in PBS. Aliquots of 5 μl were spotted onto nitrocellulose membranes and air dried, and nonspecific sites were blocked with 3% BSA-0.1% Tween 20 in PBS (PBST) at room temperature for 1 h. Binding to digoxygenin (DIG)-labeled proteins was carried out as described previously (10). Binding to anti-GbpA antibody (kindly provided by Daniel Smith) was carried out as described above for Western blot analysis except that anti-rat secondary antibody was used. Binding to antigen P1 was done using a monoclonal antibody to P1 (kindly provided by L. Jeannine Brady). Binding to biotin-dextran was done as described previously (71) with the following modifications. Membranes were blocked in PBS with 3% BSA for one hour, washed three times with PBS plus 0.1% Tween 20 (PBST). Biotin-dextran (200 μg ml⁻¹; Sigma) in PBS with 0.2% BSA were then added and incubated for one hour. Membranes were washed with PBST and then anti-biotin monoclonal antibody (Sigma) was used to detect dextran binding. All the secondary antibodies used were alkaline phosphatase conjugated. Binding of ligand to the cell wall was detected using nitroblue tetrAZolium and BCIP as substrates.

**Biofilm formation assay.** Biofilms of S. mutans were grown in two different media. NG-8 and its derivatives were grown overnight in THY medium at 37°C anaerobically. The culture was diluted 1:10 into fresh THY medium and incubated further for 6 h. The culture was then diluted 1:1,000 into either THY or BM (45) media containing 1% sucrose, 0.4 or 0.8 ml of this cell suspension was added to each well of an eight-well or four-well, respectively, glass chamber slide (Lab-Tek; Nalge Nunc International) for biofilm formation on glass. For biofilm formation on polystyrene surface, flat-bottomed 96-well microtiter plates (Corning Inc.) were used. The slides or the microtiter plates were incubated at 37°C for 20 to 24 h as a static culture to allow biofilm formation.

**Confocal and scanning election microscopy.** Biofilm formation on glass slides was first examined by confocal laser scanning microscopy (CLSM). Culture media was removed from the slide chamber and the chamber was washed with sterile distilled water. Biofilm cells were stained with labeling solution containing 0.2 μM of Light Green (Invitrogen) in PBS, and then through a series of ethanol washes, slides were air dried, sputter coated with gold and analyzed by SEM (ISI-60A; International Scientific Instruments) at several magnifications at the University of South Dakota core facility.

**PM tests.** Phenotype microarray (PM) tests were performed at Biolog, Inc., essentially as described elsewhere. Both the wild type (NG-8) and the htrA mutant (IBS101) were subjected to full 20-panel PM analysis. PM arrays contain different carbon sources, nitrogen sources, phosphorus sources, sulfur sources as well as various inhibitory compounds for stress responses as described previously (11, 84). Readings were recorded for 24 h, and kinetic data were analyzed with OmniLog-PM software. This software generates time course curves for respiration and calculates differences in the areas for mutant and wild-type cells. The differences are averages of values reported for two separate experiments. Detailed information of PM experiments is available at http://www.biolog.com.

**RESULTS**

**Characterization of S. mutans htrA mutants.** The htrA gene of S. mutans strain NG-8 was disrupted by the introduction of a kanamycin-resistant cassette into the coding region of the
gene. The inactivation was confirmed by primers flanking htrA to amplify a diagnostic PCR fragment of 4.0 kb from the kanamycin-resistant transformants and a 1.9-kb fragment of the wild-type copy of htrA (Fig. 1).

We first verified the subcellular localization of HtrA protein in S. mutans strains. Different subcellular fractions from NG-8 (wild type) were prepared as described in the Materials and Methods, fractionated by SDS-PAGE, and probed with anti-HtrA antibody generated against the B. subtilis HtrA (3, kindly provided by Dr. Devine). This antibody reacted with a diagnostic band of ~43-kDa length from whole-cell lysate, cell wall, and culture supernatant fractions from NG-8 strain. This immunoreactive band was absent from any of the fractions from IBS101 (htrA mutant) (data not shown). This suggests that, like B. subtilis HtrA, S. mutans HtrA is also present on the cell surface (extracellular).

Compared to the wild-type strain, the htrA mutant strain (IBS101) was found to grow similarly at 37°C in both rich (THY) and chemically defined (BM) media (data not shown). No differences in colony morphology were noted between the htrA and wild-type strain. However, one striking feature of the htrA mutant was its capability to form longer chains compared to wild-type NG-8 (two to three times longer) (Fig. 2A). Increased chaining was primarily associated with late-exponential- to stationary-phase cultures. In this aspect, the htrA mutant behaved like the biofilm deficient brpA mutant strain described previously (80).

As expected, the S. mutans htrA mutant also showed a reduced temperature range for growth. As shown in Fig. 2B, htrA failed to grow on mitis salivarius agar plate at both 30°C and 42°C, whereas the wild-type grew satisfactorily (Fig. 2). To confirm that the lack of HtrA was responsible for the observed phenotype, the intact htrA gene was provided in trans on plasmid pIB108. Introduction of pIB108 in IBS101 restored the growth capacity both at 30°C and 42°C (Fig. 2B).

In S. pneumoniae, disruption of htrA causes a drastic reduction in transformation efficiency (30). However, when we compared the transformation efficiency for replicated plasmid, integrated plasmid and linear chromosomal DNA, we found no differences between wild-type and htrA strains of S. mutans (data not shown).

Stress response phenotypes of htrA mutant. The HtrA protease in other bacteria has been found to be associated with the ability to survive under different stress conditions. Therefore first we tested the htrA mutant for its ability to withstand treatment with H2O2 for oxidative stresses and treatment with puromycin (which causes premature chain termination during protein synthesis) to mimic thermal stress. In both cases, the htrA mutant was more sensitive to stress conditions compared to the wild-type strain (Fig. 3). Interestingly, there were no significant differences when cells were subjected to superoxide stress generated by paraquat or tert-butyl hydroperoxide (data not shown).

S. mutans is known to adapt to a low pH environment and mount an acid tolerance response (7, 27, 61). Acid tolerance induces several stress-responsive proteins. In order to deter-
mine whether htrA is involved in acid tolerance, we measured growth of the htrA mutant under acidic conditions. As shown in Fig. 3B, the growth of the htrA mutant strain was significantly reduced at pH 5.5 as compared to the wild type. However, the htrA mutant grew as the wild type at pH 6.0.

Since many of the proteins involved in acid tolerance are also involved in DNA damage repair (such as uvrA), we tested htrA mutant for its ability to withstand the DNA damaging agent mitomycin C. Mitomycin C alkylates double stranded DNA and thereby blocks DNA replication. As shown in Fig. 3C, survival of S. mutans was greatly reduced when htrA was absent. This suggests that htrA plays an important role in the DNA damage repair pathway. RecA is one of the key proteins involved in DNA repair, however it is also induced under acid stress (42). We therefore tested the strains for UV sensitivity since RecA is involved in UV-irradiated DNA damage repair. We found no differences between wild-type and htrA mutant strains (data not shown) indicating protein other than RecA is directly or indirectly involved in this phenomenon.

In Listeria monocytogenes, htrA was recently shown to be involved in salt tolerance (82). Therefore, we tested whether S. mutans htrA is also involved in salt tolerance. We found that like L. monocytogenes, the S. mutans htrA mutant was sensitive to salt stress. All the stress sensitive phenotypes could be complemented by introducing plasmid pIB108 into the IBS101 strain. Taken together our results show that S. mutans htrA is indeed involved in various stress response pathways including DNA damage repair.

**PM analysis of the htrA mutant.** PM assay is a relatively new tool which allows testing of mutants for a large number of phenotypes simultaneously (11, 37, 60, 84). The growth of bacteria in different media is measured with tetrazolium redox dye. Reduction of this dye due to respiration during bacterial growth results in purple color formation and accumulates in the well over the incubation period. Total loss of function will result in no growth and no color formation. Thus, colorimetric detection due to respiration can provide a reporter system for phenomic testing. PM assays were performed on the htrA mutant (IBS101), compared with the wild-type strain (NG-8), in a set of 20 96-well microplates containing various nutrients or inhibitors. This allowed testing of nearly 1,900 cellular phenotypes in a sensitive, highly controlled, and reproducible format. As shown in Fig. 4, the metabolic signal was generally low for nitrogen metabolism (PM3, -6, -7, -8), phosphate and sulfate metabolism (PM4), nutrient stimulation (PM5) with the exception of carbon sources (PM1 and -2). However, the result indicated that the mutant (IBS101) showed slower growth using carbon sources such as sucrose (PM1, D11) and lactulose (PM1, D10) than its parental strain (NG-8), while there was no significant change in utilization of other carbon sources (Fig. 4). There was systematically lower signal for the mutant strain (IBS101) in PM9 through -20, suggesting lower levels of growth and/or metabolism relative to the wild type (NG-8) in these panels. Specifically, there were negative differences in salt and osmotic tolerance for 1% sodium chloride (PM9, A1), 3 to 4% sodium sulfate (PM9, D6 and -7), and phosphate toxicity of 100 mM sodium phosphate (PM9, G3). This observation is consistent with our result with osmotic stress (Fig. 3). Also, there were significant differences for other chemical sensitive panels such as hexamine cobalt chloride (PM19, H1 and -2), polymyxin B (PM19, H11 and -12), and amitriptyline (PM20, A1).

**Altered surface protein expression in htrA mutant.** Based on the previous studies in other gram-positive bacteria, it appears that HtrA is involved in the processing and maturation of extracellular proteins. Recently in S. pyogenes it was shown that SpeB protein is processed by HtrA to active form fromzymogen (46). In addition, biogenesis of streptolysin S (SLS) activity is altered by htrA mutation in this bacterium (46). We hypothesized that in S. mutans, HtrA is also involved in processing and maturation of various extracellular proteins. If HtrA affects the processing of pre-proteins or cell-associated proteins, one would expect to find a different protein profile in the culture supernatant of htrA mutant, compared to wild-type strain. When we tested for hydrophobicity, it appeared that the htrA mutant showed a drastic reduction, calculated from the proportion of cells associated with the hexadecane phase (29%
for wild-type NG-8 compare to 10% for the mutant IBS101). This indicates that surface protein expression is probably altered in the mutant. Therefore, we tested expression of several extracellular proteins in the htrA mutant. A simple test for expression of surface proteins is by spotting whole cells onto nitrocellulose membranes and developing the membranes with various DIG-labeled ligands or antibodies. Since glucans play an important role in S. mutans adhesion and accumulation on tooth surfaces, we assayed for expression of glucan-binding proteins (GBPs) first. At least three GBPs are present in S. mutans and these proteins are very important for bacterial adhesion (5). Most of the GBPs are found both on the cell surface and in the intracellular compartment of the bacterium. GBPs can bind to dextran as ligand. As shown in Fig. 5, binding to biotin-dextran was drastically increased (approximately two- to fourfold) in the htrA mutant IBS101. Since, GbpB was shown to bind glucan (72), we tested for surface expression of GbpB. As expected, GbpB expression was increased (more than eightfold) in IBS101 compared to NG-8. Both of these phenotypes can be complemented with pIB108, a plasmid containing full-length htrA. We also tested surface expression of antigen P1 (data not shown), fibrinogen (data not shown) binding protein, and fibronectin-binding protein (Fig. 5) but did not find any differences.

Expression of several exported virulence factors was recently shown to be affected by htrA mutation in S. aureus (62). Therefore, involvement of HtrA in the expression of exported proteins in S. mutans was examined. Protein extracts were prepared from cells and culture supernatants from stationary phase cultures of the wild-type (NG-8) and htrA mutant (IBS101) cells and analyzed by SDS-PAGE. As shown in Fig. 6A, the expression profile of exoproteins was markedly different. In the IBS101 extracts, we consistently observed increased proteins compared to NG-8. Protein profiles of mid-exponen-
tial-phase growing cultures analyzed by SDS-PAGE were essentially the same for wild type and the mutant (data not shown). This was also the case for total cellular fraction profiles from overnight cultures (data not shown), although the resolution in the SDS-PAGE may be likely to mask any differences. Mass spectrometric analysis was performed on several secreted proteins that were present in higher amount in the mutant strain compared to wild-type strain. At least five secreted proteins appear to be accumulated more in the stationary-phase supernatant of IBS101 relative to wild-type strain. Mass spectrometric identification suggests that GtfB, FTF, GbpB, enolase, and GAPDH (Fig. 6A and data not shown) are accumulated more in the IBS101 strain compared to NG-8.

In streptococci, several enzymes involved in glycolysis were known to be present as both extracellular and intracellular forms (9, 55, 56). It was surprising to observe that these enzymes are negatively affected by HtrA. In order confirm our mass-spectrometric identification, Western blot analyses were carried out. Stationary-phase culture supernatants from the wild-type (NG-8), htrA mutant (IBS101), and complemented (IBS101/pIB108) strains were separated by SDS-PAGE and probed with anti-enolase or anti-GAPDH antibodies. As shown in Fig. 6B and C, mutation in htrA caused increased production (approximately two- to threefold) of both enzymes. This increased production is specific to htrA mutation since it can be complemented (Fig. 6B and C, lanes 3). In both cases antibody reacted with two major bands. In both cases the smaller bands probably correspond to the proteolytic products. Taken together, the results described above show that the htrA mutation affects normal exoprotein expression during stationary phase growth.

**Effect of HtrA on FTF expression.** *S. mutans* produces a FTF which synthesizes fructan polymers from sucrose. Enzymatically active FTF can be found in either cell-associated or extracellular form (8). FTF is one of the most abundant proteins in the culture supernatant of NG-8. Since we identified FTF by mass spectrometric analysis as one of the proteins that was accumulated in the culture supernatant of IBS101, we verified our result by Western blot analysis (Fig. 7). Stationary phase culture supernatants from the wild-type (NG-8), htrA mutant (IBS101), and complemented (IBS101/pIB108) strains were separated by SDS-PAGE and probed with anti-FTF antibody. As expected, IBS101 showed increased presence of FTF in the supernatant compared to NG-8. However, we observed only partial complementation when htrA was overexpressed from plasmid pIB108.

**Altered biofilm formation in htrA mutant.** Biofilm formation by *S. mutans* involves two processes: sucrose independent initial attachment mediated by surface proteins (such as SpaP, GbpC), and sucrose-dependent attachment mediated by glu-
cans synthesized by glucosyltransferases from sucrose. For biofilm formation on in vitro surfaces such as glass and polystyrene, sucrose plays an important role. Since surface protein expression was altered in the htrA mutant strain, we wanted to know whether htrA mutation affected biofilm formation. Biofilms were grown in THY or BM media supplemented with sucrose on glass or polystyrene surfaces. Biofilms formed equally well in both media by the wild-type NG-8 after 20 h of growth. After overnight growth in the chamber slides or the microtiter plates, there was a noticeable difference in biofilm structure formed by IBS101 compared to the parental strain (Fig. 8A). Wild-type biofilm had a very confluent appearance while the htrA mutant showed a patchy appearance. This defect in biofilm formation by htrA mutant was complemented when htrA was provided in trans by plasmid pIB108 (Fig. 8A). The architecture of biofilms formed by the htrA mutant appeared to be altered as compared to the wild type (Fig. 8B). The htrA mutant formed large aggregates heterogeneously distributed throughout the biofilm matrix. There is a noticeable large gap in the biofilm matrix and the microcolonies were large dome-like structures. Wild-type biofilms are very uniform with thick layers of cells completely covering the attached surfaces. Thus, HtrA deficiency causes alter biofilm formation in S. mutans.

DISCUSSION

Streptococcal genomes encode only one HtrA protease, unlike other low G+C gram-positive bacteria such as B. subtilis and S. aureus that encodes multiple HtrA paralogues. However, the genomic organization of the htrA locus in streptococci is quite unique (Fig. 1A). In most streptococci, htrA is located near the chromosomal replication origin. While the organization of the downstream region of htrA appears to be highly conserved, the upstream region is not well conserved among streptococci. For example, in S. pneumoniae the com operon is present near the htrA locus (76) whereas in S. mutans the com operon is present elsewhere in the genome (Fig. 1A). Immediately downstream of htrA is a gene with similarity to spo0J. This gene is associated with chromosome partitioning in B. subtilis (33). spo0J is followed by the oriC region and dnaA, whose product is required for initiation of chromosomal replication. In streptococci, the htrA locus also contains several binding sites for DnaA protein (24) and, as suggested by Lyon and Caparon (46), may be involved in chromosome replication and cell division. In other gram-positive bacteria including the closely related L. lactis and Enterococcus faecalis, the htrA locus is not associated with the chromosomal replication origin and therefore may not be involved in cell division.

Like in other bacteria, we have observed stress sensitive phenotypes due to the htrA mutation. S. mutans can grow in temperatures ranging from 28°C to 47°C with optimum growth around 37°C (47). The htrA mutant of S. mutans showed a restricted temperature range and failed to grow at both 30°C
and 42°C. In *S. pyogenes*, a single-crossover insertional mutant showed a temperature sensitive phenotype while a deletion mutant did not (46). Because a large insertion had occurred into the *htrA* locus during single crossover integration, it was suggested that such a large insertion may have interfered with the normal chromosomal replication. However, our inactivated *htrA* mutant was made by gene-replacement and we were able to complement the thermo-sensitive phenotype by providing the *htrA* gene in trans from the pBl108 plasmid. This complementation rules out the possibility of polar effect of mutation on downstream genes. Interestingly, our mutant also failed to grow at low temperature (30°C). Most evidence suggests that *htrA* acts as a housekeeping protease to degrade unfolded proteins during heat shock (54). While this function is necessary for growth at high temperature, *htrA* also acts as chaperon at low temperature (74). Possibly due to loss of this latter function, the *htrA* mutant failed to grow at 30°C.

The *S. mutans* *htrA* mutant also showed sensitivity towards osmotic stress generated by NaCl, acid stress, and oxidative stress. These observations are consistent with HtrA being involved in stress response. Although the role of HtrA in osmotic stress was not characterized previously, in *L. monocytogenes* HtrA was shown to be involved in osmotic stress (82). We also found that in *S. mutans* HtrA plays a role in osmotic stress. We have observed acid sensitivity of our *htrA* mutant which is consistent with the earlier observation (21). During acid stress, several proteins including many heat shock proteins and chaperons are overexpressed in *S. mutans* (42, 81). This general stress induction in *S. mutans* confers cross-protection to other stresses such as thermal and oxidative stress (75). It is possible that *htrA* expression is also induced during acid stress and plays an important role in acid-tolerance. To understand how HtrA participates in osmotic and acid stress responses requires further investigation.

Interestingly, the *htrA* mutant showed sensitivity towards the DNA damaging agent mitomycin C. Mitomycin C induces mainly interstrand DNA cross-links (77) and results in induction of SOS response. RecA protein plays a central role in repair of mitomycin C damaged DNA and HtrA could modulate RecA function. However, we tested *htrA* mutant for UV sensitivity and found no difference compared to the wild type (data not shown). Therefore, a protein other than RecA is probably involved in mitomycin C sensitivity in the *htrA* mutant. In this context, it is important to point out that several DNA repair proteins are also induced during acid stress (42) and thus there may be a direct connection between DNA repair and acid-tolerance mediated by HtrA.

During stationary phase, the expression of surface associated proteins was altered by *htrA* mutation in *S. mutans*. Notably, we found that expression of GbpB was greatly enhanced in the *htrA* mutant. GbpB is a glucan-binding protein which shares extensive homology with peptidoglycan hydrolases from other gram-positive bacteria (50). Interestingly, GbpB is up-regulated during high osmolarity, high temperature, and low pH, and therefore GbpB may act as a general stress protein (17). GbpB is also essential for cell wall integrity and maintenance of cell shape (17). Attempts to knock out the *gbpB* gene have also supported the notion that expression of GbpB is essential for cell survival (49). In the *htrA* mutant, we have found increased expression of GbpB not only in the cell surface fraction but also in the culture supernatant and whole-cell lysate fractions (data not shown). Therefore it appears that overall expression of GbpB is increased in the *htrA* mutant. It is possible that under normal conditions the protease activity of HtrA regulates GbpB expression either directly by degrading the protein or indirectly by degrading a transcriptional regulator necessary for gbpB expression. How HtrA regulates GbpB expression and what is the relationship between GbpB and stress response remains to be explored.

We have tested expression of various other cell surface proteins in the *htrA* mutant. We found that *htrA* mutation had no effect on cell surface expression of antigen P1 (data not shown) or binding to fibronectin and fibrinogen (data not shown) consistent with previous results (6, 21). However, in contrast with the previous report, we found that *htrA* mutation affects GtfB and FTF expression (Fig. 6A and 7). Both GtfB (39) and FTF (64) bind to glucan and play an important role in *S. mutans* adhesion to the tooth surface. Both proteins are found in the extracellular fraction in addition to cell-associated forms (5, 38). The predicted molecular mass of *S. mutans* FTF, after removal of the signal peptide, is ~83 kDa. However, *S. mutans* was found to secrete FTFs with various molecular masses ranging from 59 to 106 kDa. Processing of a single *ftf* gene product at the post-translational level accounts for this diversity (64, 66). The increased production of extracellular FTF in the *htrA* mutant suggests that HtrA plays a role in FTF secretion. However, unlike GbpB, we did not find increased FTF expression on the cell surface (data not shown), suggesting that association of FTF to the cell surface rather than total production is altered by *htrA* mutation. Since HtrA is also cell wall associated (3), it is possible that HtrA facilitates proper folding of FTF needed for surface anchoring during or after translocation from cytoplasm to the cell wall.

Previously, Diaz-Torres and Russell characterized an *htrA* mutant from *S. mutans* (21). They investigated the effect of *htrA* mutation on seven extracellular or wall associated proteins including GTF, FTF and WapA, and found no differences between wild-type and *htrA* mutant strains. However, their *htrA* mutant was generated by single-crossover insertion using an internal fragment of *htrA* gene. Single cross over mutant may generate a partially functional truncated HtrA. Indeed, their *htrA* mutant contained a truncated copy encoding a 392-amino-acid polypeptide, lacking only C-terminal 10 amino acids and therefore might be fully functional. In contrast, our *htrA* mutant was generated by double crossover by disrupting HtrA at amino acid position 243, thereby deleted the conserved catalytic domain. Moreover, recently in *S. pyogenes* it was shown that a single-crossover insertional inactivation of *htrA* resulted in different phenotypes when compared with a clean non-polar deletion mutant (46). Although unlikely, strain differences (LT11 [21] versus NG-8, our strain) might have played a role in observed phenotypic differences. Nevertheless, the single-crossover insertion mutant of *S. mutans* *htrA* formed cohesive clumps in liquid media and showed different colony morphology indicating altered surface properties.

Glycolytic enzymes including GAPDH and enolase were also enriched in the stationary phase culture supernatant of the *htrA* mutant. GAPDH and enolase have been identified on the cell surfaces and culture supernatants of pathogenic streptococci, which raises questions as to their function at these loca-
tions (16). None of these proteins contain signal peptide, LPXTG anchoring motif or other known cell-wall-anchoring motifs and thus how these proteins are secreted or anchored to the cell surface remains to be uncovered. Pancholi and Fischetti (56) found that the surface GAPDH of S. pyogenes binds to fibronectin, lysozyme and to the cytoskeletal proteins and contributes to pathogenesis in this organism. In S. oralis, GAPDH binds to Porphyromonas gingivalis fimbriae and may contribute to its colonization in the periodontal pockets (48). In other streptococci, GAPDH also acts as an adhesin and contributes to virulence (13, 43). Surface enolase is also important for pathogenesis. For example, S. pyogenes enolase binds to plasminogen and plays a role in bacterial invasion of the human host and autoimmune sequelae (55). None of these proteins contain signal peptide, 6932 BISWAS AND BISWAS INFECT. IMMUN.

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HtrA in surface protein expression of S. mutans


