Passive Transfer of Immunoglobulin Y Antibody to *Streptococcus mutans* Glucan Binding Protein B Can Confer Protection against Experimental Dental Caries

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Active immunization with *Streptococcus mutans* glucan binding protein B (GBP-B) has been shown to induce protection against experimental dental caries. This protection presumably results from continuous secretion of salivary antibody to GBP-B, which inhibits accumulation of *S. mutans* within the oral biofilm. The purpose of this study was to explore the influence of short-term (9- or 24-day) passive oral administration of antibody to *S. mutans* GBP-B on the longer-term accumulation and cariogenicity of *S. mutans* in a rat model of dental caries. Preimmune chicken egg yolk immunoglobulin Y (IgY) or IgY antibody to *S. mutans* GBP-B was supplied in lower (experiment 1) and higher (experiment 2) concentrations in the diet and drinking water of rats for 9 (experiment 1) or 24 (experiment 2) days. During the first 3 days of IgY feeding, all animals were challenged with 5 × 10⁶ streptomycin-resistant *S. mutans* strain SJ-r organisms. Rats remained infected with *S. mutans* for 78 days, during which rat molars were sampled for the accumulation of *S. mutans* SJ-r bacteria and total streptococci. Geometric mean levels of *S. mutans* SJ-r accumulation on molar surfaces were significantly lower in antibody-treated rats on days 16 and 78 of experiment 2 and were lower on all but the initial (day 5) swabbing occasions in both experiments. Relative to controls, the extent of molar dental caries measured on day 78 was also significantly decreased. The decrease in molar caries correlated with the amount and duration of antibody administration. This is the first demonstration that passive antibody to *S. mutans* GBP-B can have a protective effect against cariogenic *S. mutans* infection and disease. Furthermore, this decrease in infection and disease did not require continuous antibody administration for the duration of the infection period. This study also indicates that antibody to components putatively involved only in cellular aggregation can have a significant effect on the incorporation of mutans streptococci in dental biofilm.

The molecular pathogenesis of mutans streptococcus-associated dental caries involves a series of binding events that eventually lead to the accumulation of sufficient numbers of these cariogenic bacteria to cause disease (6). The initial binding event appears to involve the interaction of bacterial cell surface adhesins (antigen I/II) with receptors in the salivary pellicle. These cariogenic streptococci then accumulate in the dental biofilm following enzymatic synthesis of extracellular glucans which provide multiple binding sites for glucan binding proteins (GBPs) associated with the bacterial cell. At least six glucans with glucan binding properties (GBPs) have been identified in proteins with glucan binding properties (i.e., the ability to bind glucans which provide multiple binding sites for glucan binding proteins in the accumulation phase of cariogenic microbiota in dental biofilms. At present, *S. mutans* GBP-B is of the greatest immunological interest in this regard. This protein bears little antigenic similarity to other *S. mutans* or *S. sobrinus* GBPs. Saliva of young children often contains immunoglobulin A (IgA) antibody to *S. mutans* GBP-B (25), indicating that initial infection with *S. mutans* can lead to induction of immune responses to this protein in humans. GBP-B appears to be more immunogenic than GBP-A in rodents (24). Furthermore, active immunization with *S. mutans* GBP-B induces immune responses that result in lower levels of *S. mutans* colonization and in reduced dental caries caused by experimental infection with *S. mutans* (27). Similar active immunization with GBP-A did not demonstrate these protective effects (24). Thus, antibody to *S. mutans* GBP-B appears to have the potential to modulate infection and disease caused by *S. mutans*.

Passive immunization approaches have been used with success to interfere with mutans streptococcal infection and resulting dental caries. Experiments using intravenous, suckling (17; reviewed in reference 28), dietary (18), or mouthwash (3, 7) transfer of antibody raised to intact mutans streptococcal
cells have shown each of these methods to result in protection. Passive administration of chicken egg yolk IgY antibody to
mutans streptococcal GTF (5) has protected rats from experi-
mental dental caries. Topical administration of mouse (11, 13, 15) or transgenic (14) monoclonal antibody to antigen I/II
has also been shown to modify mutans streptococcal infection
of humans or nonhuman primates. Since active immunization
with S. mutans GBP-B has been shown to induce a protective
response in rats, and since this component is theoretically
implicated only in the accumulation phase of mutans strepto-
coccal colonization, it was of interest to explore the effect
of passive administration of antibody to GBP-B in this model. In
the present set of experiments, we show that short-term dietary
administration of IgY antibody to S. mutans GBP-B diminishes
the accumulation of cariogenic mutans streptococci and the
resulting dental disease on molar surfaces of rats.

MATERIALS AND METHODS

GBP. GBP-B was purified from culture supernatant of S. mutans strain SJ by
anion-exchange chromatography. Bacteria were cultivated in sucrose-free chem-
ically defined medium (S. Socransky, C. Smith, and A. Manguellelo, J. Bacteriol.
52:88, 1973) and subsequently removed by centrifugation. The supernatant
was clarified by 0.45-μm-pore-size filtration as previously described (23, 27). The
filtrate was brought to pH 6.5 with sodium hydroxide, and 0.02% sodium azide
was added. This filtrate was concentrated by tangential flow ultrafiltration using
a Pellicon cassette (Millipore Corp., Bedford, Mass.) and further concentrated
on a 30-kDa-cutoff Minipart tangential flow ultrafilter (Millipore Corp.). The
low-molecular-weight components were removed by dialysis on the Minitan
with 0.02 M sodium phosphate (pH 6.5), 0.02% sodium azide, and 3.3 mM
sodium carbonate, (pH 9) to 10 mM, and concentrated to 4.6 liters by
nium sulfate precipitation (29). Briefly, approximately 4.8 liters of yolk was
nlysed with 6 volumes of water at 4°C. Precipitated material was allowed to
done after exposure to the preimmune IgY. Observed after exposure to the preimmune
mixture of components was eluted with 3 M guanidine HCl. The latter preparation contained all com-
ponents with specificity for α-1,6-linked glucan, including GBP-A, GBP-B,
FT-C, and GTF-D. Proteins were first separated by SDS-PAGE, which was performed for 1 h at 17 mA/gel on 7% polyacrylamide gels containing 0.01%
SDS with a 4% stacking gel in an air-cooled slab-gel apparatus (Mighty Small,
 Hoefer Scientific Instruments, San Francisco, Calif.) as previously described (23).
For Western blot analysis, SDS-PAGE-separated proteins were electro-
phoretically transferred to nitrocellulose for 1 h at 200 mA. After blocking, the
blotted proteins were incubated for 3 h with preimmune and postimmune
chicken IgY reagents (30 μg/ml in 1× PBS/1% Tween 20 and 0.1% gelatin). Reactive bands were revealed with biotin-
rabbit anti-chicken IgG (Zymed), followed by streptavidin-horseradish
zymoxygenase (Zymed), followed by the addition of 0.05% 4-chloro-1-naphthol,
16.7% methanol, and 0.015% hydrogen peroxide. Only one band of reactivity was
observed when the purified GBP-B or the guanidine-eluted mixture of compo-
ents was exposed to IgY antibody. No reaction with either preparation was
observed after exposure to the preimmune IgY.

ELISA. Antibody titers were measured by sandwich ELISA. Purified GBP-B
(100 ng in 0.1 ml) was used to coat Maxisorp microtiter plates (Nalge Nunc,
Kingston, N.Y.) for 1 h at 4°C. Plates subsequently were blocked with 1% gelatin
type B; Sigma) in PBS for 1 h at room temperature and washed with 0.1% Tween 20 in PBS. Anti GBP-B antibody was added at 200, 100, 50, and 25
ng/ml in PBS containing 0.1% Tween 20 and 0.1% gelatin. After washing, the
plates were incubated for 1 h at room temperature with alkaline phosphatase-
labeled goat anti-chicken IgY at 0.5 μg/ml in PBS/0.1% Tween 20-0.1% gelatin
(Kirkegaard and Perry Laboratories, Inc., Gaithersburg, Md.). The plates were
washed again and incubated with Blue Phos phosphatase substrate (Kirkegaard
and Perry Laboratories). The anti GBP signal was twofold above background
at 100 ng/ml, whereas preimmune IgY is typically unreactive up to 25 μg/ml.

Experimental protocol. Two experiments were performed to test the effects on
S. mutans colonization, accumulation, and resulting dental caries of short-term
passive oral administration of IgY antibody to S. mutans GBP-B (Fig. 2). The two
experiments differed in the amount of IgY in the diet and in the duration of IgY
supplementation. Both experiments used male Sprague-Dawley rats from
Charles River Laboratories (Kingston, N.Y.) that were 28 days old at the start of
the experiments. In preliminary experiments, no indigenous mutans streptococci

FIG. 1. Western blot analysis of preimmune IgY (lanes 1 and 5; 30 μg/ml) or GBP-B IgY (lanes 2 and 4; 30 μg/ml) preparations with purified GBP-B in lanes 4 and 5 (0.04 μg/ml) and Sephadex-purified
components from S. mutans SJ culture supernatant in lanes 1 and 2 (10 μl of the concentrated eluate per lane from 500 μl of culture). This eluate includes GBP-A, GBP-B, GTF-C, and GTF-D. Lane 3 contains
molecular mass standards, whose sizes (in kilodaltons) are indicated at left.
could be cultivated from rats obtained from this facility and maintained on high-sucrose Diet 2000 (9).

Both experiments used for infection an *S. mutans* strain (SJ-r) that was resistant to streptomycin sulfate at a concentration of 0.2 mg/ml. This strain was obtained by plating 0.12 ml of a 10-fold-concentrated anaerobic Todd-Hewitt broth culture of strain SJ (wild type) onto mitis salivarius (MS) agar (Becton Dickinson) containing 0.2 mg of streptomycin sulfate (Sigma Chemical Co., St. Louis, Mo.) per ml. After 2 days of anaerobic incubation, resistant colonies were recultured in broth and purified by streaking onto tryptic soy agar containing 5% sheep blood (Becton Dickinson). Single colonies were then recultured in broth and stored in 50% glycerol at −20°C. Immediately prior to the experiment, broth cultures of *S. mutans* SJ-r were streaked onto MS agar plates with or without streptomycin and cultured anaerobically for 2 days at 37°C to confirm streptomycin resistance. Preliminary experiments indicated that the acquired antibiotic resistance did not diminish the cariogenicity of this bacterial strain in the Sprague-Dawley rat.

Rats were housed individually in suspended cages. Additional preliminary experiments using differing doses of *S. mutans* SJ-r to infect rats of this age revealed that a dose of $5 \times 10^6$ organisms was required to establish an infection in all animals. This level of infection was thus selected to examine the effect of dietary antibody to GBP-B on *S. mutans* accumulation.

Two groups of 10 rats each were used in the first experiment. Upon arrival, the 28-day-old rats were placed on Diet 2000-IgY. The control group (cIgY) received Diet 2000 supplemented with 0.17% preimmune IgY. The anti-GBP-B group (GBP-IgY) received Diet-2000 supplemented with 0.17% IgY containing antibody to *GBP-B*. Rats were fed ad libitum for 9 days, consuming approximately 10 to 15 g of diet each day. On day 10, the supplemented diet was removed and rats in both groups were placed on Diet 2000 for the remaining 69 days of the experiment. On the first 3 days of the experiment, the cIgY and GBP-IgY rat groups received distilled drinking water that contained 75 μg of control or immune IgY/ml, respectively, together with 0.1% sodium benzoate. On day 5, the supplemented drinking water was removed and rats in both groups were placed on distilled water alone. All rats in the first experiment were orally infected with $5 \times 10^6$ *S. mutans* SJ-r bacteria on days 0, 1, and 2. The subsequent accumulations of the infecting strain, as well as total streptococcal microbiota, were sampled on days 5, 9, and 16. All animals were sacrificed on day 78 to measure the extent of dental caries.

The second experiment included two groups of 13 rats each and followed a format similar to that of experiment 1 (Fig. 2) except for the amount and duration of IgY feeding. Rats were placed on Diet 2000-IgY 1 day after arrival. For the first 9 days of experiment 2, the control group (cIgY) received Diet 2000 supplemented with 0.44% preimmune IgY. During this 9-day period, the anti-GBP-B group (GBP-IgY) received Diet 2000 supplemented with 0.44% IgY containing antibody to *S. mutans* GBP-B. On day 10, the amounts of IgY in the control and immune diets were decreased to 0.30% preimmune IgY or anti-GBP-B IgY, respectively. The animals were maintained on this diet until day 24, at which time the supplemented diet was removed and replaced by Diet 2000 for the remaining 54 days of the experiment. On the first 5 days of the experiment, the cIgY and GBP-IgY rat groups received distilled drinking water that contained 75 μg of control or immune IgY/ml, respectively, together with 0.1% sodium benzoate. On day 5, the supplemented drinking water was removed and rats in both groups were placed on distilled water alone. As in the first experiment, all rats in the second experiment were orally infected with $5 \times 10^6$ *S. mutans* SJ-r organisms on days 0, 1, and 2. The subsequent accumulations of the infecting strain, as well as total streptococcal microbiota, were sampled on days 5, 13, 20, and 78. All animals were sacrificed on day 78 to measure the extent of dental caries.

**Bacterial recoveries.** Systematic swabbing of teeth, sonication, and plating to determine bacterial cell counts were all performed as previously described (27). Appropriate dilutions were placed on MS agar to count total streptococci and on MS agar with 0.2 mg of streptomycin sulfate (MS)/ml to count the experimentally inoculated *S. mutans* SJ-r organisms. The numbers of *mutans* streptococci on these plates, as identified by colony morphology, are presented as percentages of the total number of streptococci.

**Caries assessment.** The extent and depth of carious lesions in all rat molar teeth (caries scores) were microscopically evaluated by a modified Keyes’ method as previously described (27). These combined caries scores were determined separately on buccal, lingual, and occlusal dental surfaces.

**Statistical analysis.** The differences in the median values among the treatment groups were analyzed by one-way analysis of variance (ANOVA), followed by the
RESULTS

Two experiments using different dietary concentrations of antibody were performed to test the hypothesis that short-term exposure to antibody to \textit{S. mutans} GBP-B can influence the accumulation of \textit{S. mutans} in the rodent oral flora. In the first passive immunization experiment (experiment 1), rats were fed a dietary concentration of 1.7% preimmune control IgY (cIgY) or IgY antibody to \textit{S. mutans} GBP-B (GBP-IgY) for 9 days. In addition, distilled drinking water was supplemented with cIgY or GBP-IgY in control or antibody-fed rats, respectively (Fig. 2). Rats were infected with $5 \times 10^6$ \textit{S. mutans} SJ-r bacteria on days 0, 1, and 2 of the first experiment. The molars of all rats were systematically sampled for streptomycin-resistant \textit{S. mutans} and for total cultivable streptococci twice during the 9-day dietary administration of antibody (days 5 and 9) and once soon after IgY was removed from the diet (day 16). The percentages of \textit{S. mutans} (CFU on MSS agar) relative to total streptococcal flora (CFU on MS agar) are shown in Fig. 3 for both groups on all sampling occasions.

Five days after the start of infection, \textit{S. mutans} comprised less than 1% of the total streptococcal flora in both cIgY-fed (geometric mean = 0.07%) and GBP-IgY-fed (geometric mean = 0.17%) groups. Streptomycin-resistant \textit{S. mutans} could be detected in all but three rats at this time. Differences between groups were not statistically significant.

Four days later (experimental day 9), \textit{S. mutans} SJ-r showed appreciable accumulation in all cIgY- and GBP-IgY-fed rats (Fig. 3). The geometric mean percentage \textit{S. mutans} relative to total streptococcal flora was 16.1% in the cIgY-fed rats versus 5.5 percent in the GBP-IgY-fed rats. Accumulation of \textit{S. mutans} continued in both groups such that on experimental day 16, \textit{S. mutans} SJ-r represented a geometric mean of 33.8% of the total streptococcal flora in the cIgY group, in contrast to a geometric mean of 24.2% in the GBP-IgY group. Thus, relative to the total streptococcal flora, \textit{S. mutans} appeared to accumulate to lower levels in the GBP-IgY-fed rats than in the cIgY-fed rats on days 9 and 16, although these differences did not achieve statistical significance.

Figure 4 illustrates the extent of dental caries on buccal, lingual, and occlusal surfaces after 78 days of infection of cIgY-fed and GBP-IgY-fed rats in experiment 1. Mean caries scores on each of these surfaces of GBP-IgY-fed rats were approximately half of those observed among controls. These differences were highly significant ($P < 0.005$) on lingual surfaces and were nearly significant ($P = 0.061$) on buccal surfaces. Taken together, these data suggest that early exposure to dietary IgY antibody against \textit{S. mutans} GBP-B may diminish the extent of eventual dental disease and possibly of bacterial accumulation prior to observable disease.

To test whether increased exposure to GBP-IgY can further inhibit \textit{S. mutans} accumulation or disease progression, a second passive immunization experiment was performed (experiment 2). In this experiment, the duration of exposure to cIgY or GBP-IgY was increased to 24 days, and the concentration of dietary IgY was increased to 4.4% during the initial 9 days of feeding and to 3.0% during the next 15 days of feeding. As in the first experiment, 28-day-old rats were infected with $5 \times 10^6$
S. mutans SJ-r organisms on days 0, 1, and 2. The molars of all rats were systematically sampled for streptomycin-resistant S. mutans and for total cultivable streptococci three times during the dietary administration of antibody (days 5, 13, and 20) and at the end of the experiment (day 78). The geometric mean percentage of S. mutans (CFU on MSS agar) relative to total streptococci (CFU on MS agar) are shown in Fig. 3 for both groups on all sampling occasions.

Five days after the start of infection, the geometric mean levels of S. mutans infection within the two groups were similar (cIgY = 0.43%; GBP-IgY = 0.27%), although in two of the cIgY rats, S. mutans comprised more than 10% of the total streptococcal flora. Streptomycin-resistant S. mutans could not be detected in one antibody-fed rat at this time.

Eight days later (day 13), S. mutans SJ-r showed appreciable accumulation in all cIgY- and GBP-IgY-fed rats. The mean percentage of S. mutans relative to total streptococcal flora was 24.0% in the cIgY-fed rats and 6.6% in the GBP-IgY-fed rats. These differences were significant at a P value of 0.05. S. mutans accumulation continued in both groups during dietary administration of cIgY or GBP-IgY. On day 20, S. mutans SJ-r represented a mean of 48.9% in the cIgY group in contrast to a mean of 20.4% in the GBP-IgY group. Although differences on day 20 did not achieve statistical significance, the presence of dietary IgY antibody to S. mutans GBP-B appeared to inhibit the accumulation of S. mutans SJ-r in the dental biofilm.

Most rats were heavily colonized with S. mutans at the end of the experiment (day 78). However, the percentage of S. mutans relative to total streptococcal flora was still significantly lower (P < 0.04) in the group that had been initially fed GBP-IgY (geometric mean = 63.4%) than in the cIgY-fed group (mean = 80.1%), despite the fact that no dietary antibody supplements were given during the last 54 days of the experiment.

Figure 4 illustrates the extent of dental caries on buccal, lingual, and occlusal surfaces after 78 days of infection in cIgY- and GBP-IgY-fed rats. Mean caries scores on each of these surfaces of GBP-IgY-fed rats were approximately half of those observed on all comparable surfaces of rats that were fed IgY. These differences were highly significant (P < 0.003) on occlusal surfaces. Differences on smooth surfaces did not reach significance at a P value of 0.05 (buccal, P = 0.051; lingual, P = 0.086).

Rats in both experiments were infected 29 days after birth. The third molars of the rat erupt into the oral cavity on day 35 (22). Thus, the colonization potential of these nascent surfaces by S. mutans SJ-r may be different from that of the earlier-erupting first and second molars, which had been initially colonized in the absence of this strain. To examine whether molar eruption in the presence or absence of S. mutans infection and/or antibody ultimately influenced the extent of disease, the caries scores of all molar teeth of the rats in experiments 1 and 2 were calculated with respect to individual teeth. The mean caries scores for individual teeth in both experiments were lower in the group given dietary GBP-IgY, supporting the modulating effect of antibody on disease seen when data were calculated with respect to surface (Fig. 4). Differences in caries scores were significant in 2 of 12 teeth in experiment 1 and 8 of 12 teeth in experiment 2 (Fig. 5), indicating that increasing the amount and duration of dietary antibody application increased the protective effect. Significant effects were most often observed in mandibular molars in both experiments. The earlier-erupting first (day 19) and second (day 22) molars were more...
significantly protected by dietary antibody than were the third molars, which erupted during infection.

**DISCUSSION**

The results of this study indicate that passive administration of antibody to *S. mutans* GBP-B can inhibit the accumulation of *S. mutans* in the dental biofilm and can provide a level of protection from dental caries caused by these organisms. This finding parallels the effectiveness of active immunization with GBP-B, via systemic (27) or mucosal (24) routes, in modulating *mutans streptococcal* accumulation and disease. The present study, however, is the first to demonstrate that antibody to GBP-B can be protective via passive administration. Furthermore, the protective effect was achieved with the administration of antibody for less than one-third of the infection period, in contrast to other rat model studies in which antibody to intact cells (18) or to GTF (5) was continuously administered throughout the infection period. The protective effect of GBP-B antibody was improved by increasing the concentration and duration of dietary exposure to antibody. Hamada and coworkers (5) found that the protective effects of IgY antibody to cell-associated GTF occurred at or above IgY dietary concentrations of 0.1%, consistent with our observations that protection could be observed with 0.17 or 0.44% IgY antibody in the diet.

Previous studies by Ma and coworkers (13, 14) also demonstrated a prolonged protective effect from infection and disease well after exposure to antibody had ceased. In those studies, the teeth were made aseptic using topical antibacterial mouthwash. The surfaces were then discontinuously exposed for 3 weeks to high concentrations of mouse (13) or plant (14) antibody to *S. mutans* antigen I/II adhesin during *mutans streptococcal* recolonization. Detectable recolonization with indigenous *mutans streptococci* was prevented for up to 4 months after the topical antibody treatment. Although the mechanism for the long-term effects of the 3-week topical monoclonal antibody exposure is unclear, the inhibition of recolonization in the presence of monoclonal antibody can be assumed to result from blocking of *mutans streptococcal* adhesin, thus preventing the initial colonization phase in which antigen I/II participates. Accessory immune effects of monoclonal antibody treatment were unlikely, since F(ab')2 fragments of the mouse IgG monoclonal antibody were also effective, as was the transgenic secretory IgA monoclonal antibody, which is not expected to fix complement (14).

The present study also demonstrated a long-term protective effect after limited (9- or 24-day) exposure to IgY antibody against *S. mutans* GBP-B, another component putatively involved in the molecular pathogenesis of dental caries. This study lends further support to the concept that short-term passive treatment with antibody of appropriate specificity may have immunotherapeutic efficacy for dental caries. Others have suggested that passive immunization can increase active immune responses to infective antigens. Ramisse and coworkers (19) reported that passive intranasal administration of polyvalent human immunoglobulin to mice, given during a *Streptococcus pneumoniae* infection, resulted in a convalescent immune response to *S. pneumoniae* antigens that was higher than that in infected-only controls.

The present IgY anti-GBP-B study differs from the monoclonal antiadhesin studies of Ma and coworkers (13–15) on two main points. Firstly, antibody interactions with newly colonizing *mutans streptococci* were far more favorable in the mono-
clonal study. The total bacterial load, including the indigenous mutans streptococci, had been drastically lowered by chlorhexidine treatment prior to antibody exposure. Subsequent bacterial challenge either by external exposure or by surviving indigenous mutans streptococci would be expected to be quite modest, thus favoring the inhibiting effect of monoclonal antibody that periodically bathed the chemically treated teeth during the 3-week period. In contrast, in the present experiments dietary IgY antibody to GBP-B functioned in an untreated oral environment and in the presence of a fairly vigorous (three doses of 5 × 10^6 bacteria) mutans streptococcal challenge.

In the present experiments, passive antibody was present during the eruption of the rats’ third molars, which were exposed both to the indigenous microbiota and to the exogenous S. mutans SJ-r infecting strain. This condition may be expected to favor the inhibitory effects of dietary IgY antibody if uncolonized surfaces are more readily protected than those surfaces on which a dental biofilm is already fully developed. However, analysis of the effect of passive antibody treatment with respect to dental caries on individual teeth did not support this notion (Fig. 5). Rather, these results suggested a greater influence on the first molars, which had erupted earliest and were thus exposed to the indigenous microbiota prior to treatment with IgY. This observation would then suggest that interfering with the colonization or accumulation of superinfecting mutans streptococcal challenge may be easier than interfering with the initial challenge, at least in this model system.

Secondly, the mechanism(s) of protection with antibody to GBP-B is likely to be different from that with antibody to antigen I/II (13–15) or other components of S. mutans. The antiadhesin specificity of the monoclonal antibodies presumably blocks initial colonization events. Other studies, using passive immunization with antibody to GTf, have shown a protective effect on bacterial accumulation and subsequent dental caries in the rat model (5). Since GTf has both catalytic and glucan binding domains (1, 12, 20, 30), antibody to this component could be expected to block the synthetic ability as well as other aspects of protein-glucan interactions. The glucan binding domains of GTf may function primarily in glucan branching and chain lengthening, rather than in attachment of mutans streptococci to the glucan matrix. Antibody to S. mutans GBP-B, used in the present study, has specificity for a glucan binding protein whose primary sequence (8a) is quite different from those found in the putative glucan binding domains of GTf or of the constitutively secreted S. mutans GBP-A, which has sequence similarity with the putative glucan binding domains of GTf (2). Thus, the inhibitory activity of antibody to GBP-B on mutans streptococcal accumulation would be expected to be independent of that due to antibody to antigen I/II or GTf. It cannot be excluded, however, that antibody to GTf, GBP-A, GBP-B, or antigen I/II functions in whole or in part by promoting immune aggregation of planktonic mutans streptococci, thus decreasing or eliminating their ability to incorporate into the dental biofilm. Support for this concept derives from the observation that S. mutans grown in the presence of monoclonal antibody forms long chains and is associated with clumping of cells (15). Similarly, Hamada and coworkers (5) found that IgY antibody to cell-associated GTf also aggregates S. mutans in vitro.

These studies support the use of oral passive immunization to interfere with S. mutans accumulation and subsequent development of dental caries. Furthermore, S. mutans GBP-B is identified as a new target, thus increasing the number of pathways by which passive immunization can achieve a protective effect.

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