The Effect of Xylitol on the Composition of the Oral Flora: A Pilot Study

Eva Söderlinga
Aino Hirvonenb
Sara Karjalainenb
Margherita Fontanac
Diana Cattc
Liisa Seppäa

ABSTRACT

Objectives: Our aim was to investigate the effect of short-term xylitol consumption on the microbial composition of plaque and saliva.

Methods: Twelve volunteers [22–38 yrs] harboring mutans streptococci (MS) participated in the randomized, double-blind, cross-over study. The experimental chewing gum contained 65% xylitol while the control gum contained 63% sorbitol and 2% maltitol w/w. The polyol dose was approximately 6 g/day. Stimulated saliva and plaque samples were collected before and after the two four-week test periods. The samples were cultured for MS, total streptococci, lactobacilli, and total facultatives. A part of the samples were subjected to DNA-DNA hybridizations of 14 microbial plaque species: Actinomyces naeslundii, A. viscosus, Fusobacterium nucleatum, Lactobacillus acidophilus, L. fermentum, L. paracasei, L. rhamnose, L. plantarum, Streptococcus gordonii, S. oralis, S. parasanguis, S. salivarius, S. sanguinis, Veillonella parvula.

Results: The MS counts of the plaque samples collected from “caries-prone” tooth sites decreased significantly (P<.01) in the xylitol gum group but not in the sorbitol gum group. Also the plaque MS percentage decreased significantly in the xylitol gum group (P<.01). The salivary MS counts did not decrease either in the xylitol or in the sorbitol gum groups. Nor were changes detected in the salivary levels of total streptococci or lactobacilli. The DNA-DNA hybridization assay revealed no study-induced changes in the microbial composition of the dental plaque.

Conclusions: Within the limitations of this pilot study, xylitol consumption reduced MS counts in plaque but appeared not to affect the microbial composition of plaque or saliva in general. [Eur J Dent 2011;5:24-31]

Key words: Xylitol; Mutans streptococci; Plaque; Oral flora; Chewing gum.
**INTRODUCTION**

Xylitol is a five-carbon polyol sweetener with specific, beneficial effects on oral health. Xylitol promotes mineralization by increasing the flow of saliva, which is a common effect of all polyol sweeteners. What is unique in xylitol is that it is practically nonfermentable by oral bacteria which counteracts low pH-values in the oral cavity. The caries-preventive effect of xylitol in clinical studies could not be explained by the exclusion of fermentable sugars from the diet. In controlled clinical trials xylitol has performed better than sorbitol. Recently, it was shown that xylitol syrup administration to children prevented early childhood caries, while no such effect was detected in the control group receiving high-sorbitol syrup. This study also demonstrated that not only chewing gum but also other xylitol vehicles can be effective in delivering xylitol.

The mechanism of action of xylitol on mutans streptococci (MS) is not fully known but habitual xylitol consumption, at high enough doses reduces counts of MS, apparently making plaque and mutans streptococci less adhesive to teeth. This appears to be reflected in a significant reduction in mother-child transmission of MS; in three of four published mother-child studies, habitual xylitol consumption by mothers resulted in reduced MS colonization in their children. According to in vitro studies MS can be regarded as target organisms of xylitol. Loesche et al showed that consumption of 5–7 g of xylitol in chewing gum reduced MS in both plaque and saliva but not counts of S. sanguis. Very little is actually known about the effects of xylitol on the oral flora apart from MS.

The hypothesis of the study was that in subjects showing xylitol-induced decreases in the counts of mutans streptococci no effects on the oral flora in general would be observed.

**MATERIALS AND METHODS**

**Subjects**

The clinical study was carried out in Oulu which is in Ostrobothnia, Finland. Twelve healthy dental and medical students, six males and six females with an age range of 22–38 years were recruited for the present study. Based on previous xylitol studies habitual xylitol consumption with daily xylitol doses >5 g should reduce counts of mutans streptococci in all subjects. Thus twelve subjects were estimated to give enough power for statistical analyses. All subjects showed normal salivary flow rates of paraffin-stimulated saliva (>1 ml/min) in all examinations. The subjects were screened for salivary MS (Dentocult SM Strip mutans test, Orion Diagnostica, Espoo, Finland), and all but three harbored high MS counts. All subjects had used xylitol chewing gum before the study started. Two of the subjects used xylitol gum approximately twice a week, the others on daily basis. In Finland, about half of the young adults have used xylitol chewing gums during most of their adolescence on a daily basis. Among university students this figure is even higher. The majority of the commercially available xylitol gums are pellet gums.

The Ethics Committee of the Northern Ostrobothnia Hospital District approved the study. Written consent was obtained from all subjects.

**Experimental and control chewing gums**

The experimental chewing gum (1.5 g/pellet) contained 65% xylitol w/w. The control gum was identical in pellet size but the main sweetener was sorbitol, with a concentration of 63%. Maltitol (2%) was used in the control gum to make the texture of the gums similar. Both gums contained ascorbate which was used to compensate for the differences in sweetness. The two gums had the same mild flavor and color. They were packed in identical plastic containers which were color-coded, blue and red. The codes of the test gums were kept by the gum manufacturer until the study was completed. The gum used during the washout periods was identical with the control gum but was packed in containers coded in green. The gums were manufactured and donated to the study by Karl Fazer Ab, Vantaa, Finland.

**Study design**

The study lasted 18 weeks altogether. During this period the subjects were instructed not to use antimicrobial medications, mouthwashes, or xylitol products, but to consume their normal, habitual diet, and continue their usual tooth brushing.

The study had a double-blind, crossover design as shown in Figure 1. The subjects were randomly allocated into two groups: half of the subjects used blue-coded gum first, and then the red-coded gum for four weeks, while the other half used the gums in reverse order. On the first appoint-
ment the subjects were given instructions concerning the study. They were also given ordinary fluoridated toothpaste to be used during the study (free-of-charge) as well as the green-coded wash-out gum to be used during the wash-out periods. Since all subjects were gum users to start with it was considered that gum use also during the wash-out periods would ensure compliance. The subjects were told that all gums were sugar-free gum with benefits for dental health. Professional tooth-cleaning and dental examination were carried out for all participants (AH).

Before the test periods the subjects received the test chewing gums. During the test periods, the subjects were instructed to use two chewing gum pieces at a time, three times a day, preferably after meals. The recommended chewing time was 5 min. The gum consumption resulted in a daily polyol dose of approximately 6 g. The recommendation for gum usage during the wash-out periods was less strict than for the test periods, i.e. 2–3 times a day. The subjects were instructed to refrain from tooth brushing and other oral hygiene procedures for 24 hours before sample collection and not to eat or drink for 1 hour before the sampling appointment. Compliance with the instructions provided was verified at the beginning of each appointment. At the appointments the subjects were also interviewed about amount and frequency of chewing gum use, and confounding factors like acute infectious diseases and use of antibiotics.

The subjects and researchers in Oulu were blinded throughout the study. Also the microbiological analyses in Turku were carried out blinded. The test code was revealed after the results had been fed into SPSS files. Thereafter one of the authors (ES) randomly chose the samples of five subjects from those subjects who harbored MS throughout the study to be sent to the US for DNA-DNA hybridization. The microbial analyses in the US were performed in a blinded fashion and the results were analyzed in Turku.

Saliva/plaque sampling

Paraffin-stimulated saliva was collected for 5 minutes. 100 μl of the saliva was transferred to a transport tube containing 900 μl Tryptic Soy Broth (Difco, MI, USA) with 10% glycerol v/v. The tube was stored at -70˚C before analyses. After the saliva collection dental floss attached to a floss-holder was used to collect plaque from “caries-prone” tooth sites, the mesial surfaces of two first molars (D16, D36). The mid-parts of the flosses with the plaque samples were cut into a transport tube containing 1 ml of transport medium. All available plaque was then collected with curettes, first from the left side of the mouth, and the plaque was suspended in EDTA buffer (0.25 M NaOH, 5 mM Tris, 0.5 mM EDTA), stored at -70˚C, and mailed for microbiological analysis to Indiana University, Indianapolis, USA on dry ice. The plaque from the right side of the mouth was suspended in 1 ml saline, and dispersed by pumping the suspension back-
and-forth with a disposable pipette. Then, 500 μl of the suspension was pipetted into a transport tube containing 2 ml of transport medium and stored frozen at -70°C until analysed. Except for the analyses performed in the US, all microbiological analyses were performed at the University of Turku, Turku, Finland, 620 km south of Oulu. The samples were mailed from Oulu to Turku frozen, on dry ice. The transport medium and the transport conditions have been tested earlier and they have shown excellent recoveries of streptococci and total facultatives.8

Microbiological analyses

For salivary analyses the transport tubes were thawed and vortexed for 1 minute. The plaque samples were further dispersed with a mild 10-second sonication at 4°C. After 10-fold serial dilutions the samples were plated on Mitis salivarius agars containing bacitracin [MSB; Difco, Detroit, MI, USA14], TYCSB agars [Difco15], Mitis salivarius agars [Difco], Rogosa agars [Difco] and blood agars (Orion Diagnostica). The MS grown on MSB agar were incubated for 2–3 days in a 7% CO₂ atmosphere, and the TYCSB plates anaerobically for 3 days at 37°C. The identification of MS has been described in detail earlier.8 The numbers of MS were identified on the basis of colony morphology and counted by means of a stereomicroscope. The identification of S. mutans was based on consistent findings of "rough" colony morphology on the MSB plate, positive fermentation with sorbitol, mannitol, raffinose and melibiose, and negative dextran agglutination. Identification of S. sobrinus was based on "smooth" colonies on the MSB plate, positive fermentation with mannitol but negative with raffinose and melibiose, and positive dextran agglutination. The detection limit of the MS assay for saliva was 200 CFU/ml and for plaque 200 CFU/sample. Total streptococci were grown for 2 days on Mitis salivarius agars at 37°C. "Lactobacilli-like" colonies were counted on the Rogosa agars and total facultatives on blood agars grown anaerobically for 2 days at 37°C.

From those subjects who harbored MS throughout the study, five were randomly chosen for the DNA-DNA hybridizations16 performed at the Indiana University. Since the analyses are laborious and expensive to perform, it was decided that five samples would be analyzed first and then, depending on the results, the rest of the samples. Briefly, plaque DNA samples [Master Pure Gram Positive DNA Purification Kit, Epicentre Biotechnologies, Madison, WI, USA] were applied to a membrane using a vacuum slot blot, then UV crosslinked and hybridized to DIG-labelled species-specific probes using standard techniques (Roche Applied Sciences, Indianapolis, IN, USA) for hybridization and chemiluminescent detection. Specific probe sequences, Tm [°C] values and ref.

Table 1. Species - specific probes used for DNA hybridization.

<table>
<thead>
<tr>
<th>Microorganism</th>
<th>ATCC</th>
<th>Specific sequence</th>
<th>Tm [°C]</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>S. gordonii</td>
<td>35105</td>
<td>5’-AGTTCAAACGATCTCACGATTG-3’</td>
<td>56,3</td>
<td>17</td>
</tr>
<tr>
<td>S. sanguinis</td>
<td>10556</td>
<td>5’-GACACAGGAGTTGCATTG-3’</td>
<td>60,2</td>
<td>18</td>
</tr>
<tr>
<td>S. mitis/oralis</td>
<td>9811</td>
<td>5’-GTGCACACGAGTTGCATTG-3’</td>
<td>66,2</td>
<td>19</td>
</tr>
<tr>
<td>S. paraestruitis</td>
<td>15912</td>
<td>5’-ACACATGATACACGATTG-3’</td>
<td>54,2</td>
<td>18</td>
</tr>
<tr>
<td>S. salivarius</td>
<td>13419</td>
<td>5’-CATGATACATCATACTATAGATG-3’</td>
<td>56,4</td>
<td>18</td>
</tr>
<tr>
<td>L. fermentum</td>
<td>14932</td>
<td>5’-ATCAAGTTGACGCAACAAC-3’</td>
<td>55,8</td>
<td>20</td>
</tr>
<tr>
<td>L. rhamnose</td>
<td>7469</td>
<td>5’-CTGCGACACGAGTTGCATTG-3’</td>
<td>53,4</td>
<td>21</td>
</tr>
<tr>
<td>L. paracasei</td>
<td>11582</td>
<td>5’-TGGCGGCACCAG-3’</td>
<td>61,1</td>
<td>21</td>
</tr>
<tr>
<td>L. plantarum</td>
<td>8014</td>
<td>5’-ACATCTCTGGAAACATTTG-3’</td>
<td>51,5</td>
<td>21</td>
</tr>
<tr>
<td>L. acidophilus</td>
<td>43164</td>
<td>5’-TACACATCTTGAGCTCTACA-3’</td>
<td>58,4</td>
<td>21</td>
</tr>
<tr>
<td>A. viscosus</td>
<td>43146</td>
<td>5’-ATCCAAGTTGACGCAACAAC-3’</td>
<td>69,9</td>
<td>19</td>
</tr>
<tr>
<td>A. naeslundii</td>
<td>19039</td>
<td>5’-CATCAGCATACTACAGAGGCG-3’</td>
<td>57,2</td>
<td>20</td>
</tr>
<tr>
<td>F. nucleatum</td>
<td>25586</td>
<td>5’-CAACGCTATAGGGAAGGGGACG-3’</td>
<td>67,4</td>
<td>19</td>
</tr>
<tr>
<td>V. parvula</td>
<td>10790</td>
<td>5’-ATTCCCTCCTCAGTA-3’</td>
<td>57,6</td>
<td>20</td>
</tr>
</tbody>
</table>
ferences are listed in Table 1. The following microorganisms were assessed: Actinomyces naeslundii, A. viscosus, Fusobacterium nucleatum, Lactobacillus acidophilus, L. fermentum, L. paracasei, L. rhamnose, L. plantarum, Streptococcus gordonii, S. oralis, S. parasanguis, S. salivarius, S. sanguinis, Veillonella parvula. The results were analyzed using Un-Scan-It software (Silk Scientific, Inc, Orem, UT, USA) and were expressed as pixels reflecting the number of microorganisms per sample.

Statistical analysis

The analysis of variance of repeated measures was used for longitudinal comparisons and for pair-wise comparisons the paired-samples t-test. Means were compared using independent samples t-test. The statistical package used was SPSS 14.0 for Windows. The level of statistical significance was set at P<.05.

RESULTS

In the plaque samples collected from the “caries-prone” tooth surfaces a decrease in the MS counts was seen for all ten subjects in the xylitol gum group (P<.01; Figure 2). No change was seen in the MS counts of the control/sorbitol gum group (Figure 2). Also the MS percentage of plaque decreased significantly in the xylitol gum group (P<.01), while no change took place in the sorbitol gum group (Figure 2). Concerning salivary MS counts, no “xylitol effects” were detected.

The subjects were invited to participate in the study on the basis of a prescreening for the presence of MS. Two of the subjects showed rather low levels of MS at the prescreening and in the study proper they did not harbor detectable counts of MS in either plaque or stimulated saliva. Only one of the subjects harbored both S. mutans and S. sobrinus, with S. mutans being the dominant MS. MS culturing on MSB and TYCSB gave similar results: for some subjects the MS counts were higher on TYCSB, but in other respects the results were similar. In the counts of salivary total streptococci and lactobacilli no changes were seen during the study (Figure 3). The cross-over design did not affect the results. The two subgroups did not differ from each other when baseline counts of MS were compared before the two test periods. Also within the same subgroup the baseline values before the test periods did not differ.

All available plaque collected from the right side of the mouth was cultured for total facultatives at the University of Turku. There were no differences in the counts of total facultatives in plaque indicating that sampling was consistent and no changes in the amount of plaque took place during the study. The plaque from the left side of the mouth of five subjects was subjected to DNA-DNA hybridization at Indiana University. A. naeslundii, L. acidophilus, L. fermentum and S. parasanguis were detected in so few samples that they could not be included in the statistical analysis. L. rhamnose and L. plantarum were not detected in any of the samples. Only A. viscosus, S. gordonii, S. oralis and S. sanguinis were detected in the samples of all five subjects. The mean counts of the microor-
organisms detectable in the samples of at least four subjects are shown in Table 2. No study-induced effects were detected in microbial counts of *A. viscosus*, *F. nucleatum*, *L. paracasei*, *S. gordonii*, *S. oralis*, *S. salivarius*, *S. sanguinis* or *V. parvula*. Since not even trends for study-induced effects on the flora could be observed in these analyses, the rest of the samples were omitted from the analyses.

**DISCUSSION**

The present short-term pilot study demonstrated that xylitol consumption did not affect the composition of oral flora except for a decrease in plaque MS. MS form only a small percentage of oral flora, thus it is surprising that so few studies have addressed this topic. Xylitol inhibits the growth of MS via the inducible fructose transport system.22 The degree of xylitol inhibition varies among MS strains.23-25 In addition to MS, *S. gordonii* possesses an inducible fructose pathway26 and could thus be inhibited by xylitol. Inhibition with xylitol is often addressed as the main mechanism by which xylitol consumption may reduce MS counts.7 Moreover, xylitol-induced changes in the virulence of MS have been proposed to contribute to this phenomenon.7 Recently, however, it was demonstrated in vitro that xylitol could decrease polysaccharide-mediated adhesion of MS contributing to plaque accumulation with a mechanism not dependent on growth inhibition.25

**In vitro** studies have suggested that MS are target organisms of xylitol.12,23 Loesche et al13 demonstrated a xylitol-associated decrease in MS counts, but no effects of lactobacilli. A recent clinical trial showed a xylitol-associated decrease in salivary lactobacilli. This result may, however, reflect a xylitol-induced elevation of the oral pH and thus an indirect effect on lactobacilli.32 Our results support the idea that xylitol consumption does not reduce counts of lactobacilli.

*A. viscosus*, *F. nucleatum* and *V. parvula* apparently do not possess the inducible fructose pathway but could, in theory, be affected for example by the changes xylitol consumption may have on oral pH. All subjects harbored these organisms; but no trends to any study-induced changes could be detected in the counts of the microorganisms. Thus, xylitol appeared not to affect the levels of these microorganisms.

### Table 2. The effect of chewing xylitol vs. sorbitol gum on the composition of plaque flora. The counts of each microorganism per plaque sample are expressed as pixels (mean±SD of logarithmic values). The pixels can be compared intraindividually within one subject but not interindividually. No statistically significant differences were found.

<table>
<thead>
<tr>
<th>Microorganism</th>
<th>Xylitol chewing gum</th>
<th>Sorbitol chewing gum</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>After gum use</td>
</tr>
<tr>
<td><em>A. viscosus</em></td>
<td>4.2±0.7</td>
<td>4.4±0.4</td>
</tr>
<tr>
<td><em>F. nucleatum</em></td>
<td>1.4±3.1</td>
<td>2.4±2.5</td>
</tr>
<tr>
<td><em>L. paracasei</em></td>
<td>3.1±2.9</td>
<td>3.1±2.8</td>
</tr>
<tr>
<td><em>S. gordonii</em></td>
<td>4.1±0.5</td>
<td>4.4±0.5</td>
</tr>
<tr>
<td><em>S. oralis</em></td>
<td>3.7±0.4</td>
<td>3.8±0.5</td>
</tr>
<tr>
<td><em>S. salivarius</em></td>
<td>0.4±3.3</td>
<td>1.5±3.2</td>
</tr>
<tr>
<td><em>S. sanguinis</em></td>
<td>4.2±0.4</td>
<td>4.3±0.4</td>
</tr>
<tr>
<td><em>V. parvula</em></td>
<td>3.6±0.6</td>
<td>4.0±0.4</td>
</tr>
</tbody>
</table>
Rather poor "xylitol effects" on MS were demonstrated in the present study compared to earlier studies employing similar xylitol doses and consumption frequencies. The study subjects were highly motivated dental/medical students. The fact that significant decreases were seen in the MS of the plaque samples from the caries-prone areas and plaque MS percentages indicates good compliance. Ten subjects were daily xylitol consumers and two used xylitol approximately twice a week before the study started. In spite of the long washout period before the study and the professional tooth cleaning, habitual xylitol consumption may cause long-term changes in the MS counts as discussed recently. In this study, the cross-over design was not the reason for the poor "xylitol effects"; the confounding factor was most probably the fact that the subjects were xylitol consumers to start with. Long-term consumption of xylitol has been connected with the selection of naturally occurring, so-called "xylitol-resistant" MS, not inhibited by xylitol. This phenomenon may at least partly explain why the "xylitol effects" were relatively poor in this study. The "resistancy" phenomenon has, however, never been demonstrated for any other oral microorganisms than MS. Thus, within the limitations caused by the small number of subjects, our results concerning the composition of the oral flora in general should be valid.

CONCLUSIONS
Within the limitations of this pilot study, xylitol consumption reduced MS counts but appeared not to affect the microbial composition of plaque or saliva in general.

ACKNOWLEDGEMENTS
The excellent technical assistance provided by biomedical research technician Oona Hällfors is gratefully acknowledged.

REFERENCES


