Virulence of *Streptococcus mutans*: An intrafamilial cohort study on transmission of genotypes

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Received: 01/07/2019
Accepted: 04/11/2019

Abstract
Background: The main aims of this cohort study were to measure the intrafamilial risk of transmission, sharing and stability of the most virulent *S. mutans* genotypes.

Material and Methods: A total of 392 clinical isolates of *S. mutans* obtained from caries-active adults and genotyped to evaluate their transmissibility over time. After extraction of the chromosomal DNA, PCR were performed to detect the genes involved in the production of GbpA (gbpA) and mutacin types I, II, III and IV (mutAI, mutAII, mutAIII and mutAIV).

Results: The gbpA, mutAI, mutAII, mutAIII and mutAIV genes were detected in 77.3, 12.5, 51, 16.6 and 89.8% of *S. mutans* isolates, respectively. The virulence of *S. mutans* was associated with its transmission (P < 0.01) and stability (P = 0.01), with the most virulent genotypes having higher transmissibility (RR = 1.83, 95% CI 1.44 to 2.32) and higher stability in the oral cavity (RR = 1.52, 95% CI 1.06 to 2.19).

Conclusions: Genotypes with the genetic information to synthesize GbpA and mutacins present an important ecological advantage in the process of colonization by *S. mutans*; they remain stable among the oral microbiota of the host and favor intrafamilial transmission.

Key words: Streptococcus mutans, virulence factors, transmission, dental caries.

Introduction
Due to its virulence factors, *Streptococcus mutans (S. mutans)* is considered the most cariogenic microorganism to colonize the human oral cavity (1). Consequently, for decades, the scientific community has been dedicated to investigating *S. mutans* using various biochemical, serological and genetic techniques (2). The virulence of *S. mutans* is related, among other mechanisms, to its ability to synthesize glucan-binding proteins (GBPs), a heterogeneous group of proteins that...
promote cell adhesion to tooth surfaces (3). GBPs are known to influence the maintenance of the dental biofilm architecture by joining bacteria to extracellular glucan molecules (4), which contribute to plaque formation and to the subsequent development of dental caries (5). GbpA, a GBP that depends on the quantity of environmentally available glucan (6), contributes to S. mutans cariogenicity through its role in bacterial adhesion and cohesion during dental biofilm formation (7).

Mutacins also represent an important virulence factor associated with the risk of dental caries. The adsorption capacity of active molecules of mutacins on the surface of sensitive bacteria through specific or not specific receptors can increase their antimicrobial efficiency (8). Furthermore, when a child is exposed to infection by an S. mutans strain exhibiting an increased level of mutacin production, it can be presumed that under favorable circumstances the strain will colonize, especially if the flora has not yet reached stability (9).

Previous research (thesis available at: http://dx.doi.org/10.11606/T.25.2007.tde-09112007-094447) revealed that not all S. mutans genotypes detected in parents remained stable over time and colonized the oral cavity of their children. In addition, a systematic review (10) showed that caries risk assessment in children generally remained stable over time and colonized the oral cavity of their children. In addition, a systematic review (10) showed that caries risk assessment in children generally is based in case-control or cross-sectional and, cohort studies with adequate follow-up are needed. The aims of this cohort study were 1) to determine the frequencies of genes involved in the production of GbpA (gbpA) and mutacin types I, II, III and IV (mutAI, mutAII, mutAIII and mutAIV) in S. mutans and 2) to measure the intrafamilial risk of transmission, sharing and stability of genotypes of virulent S. mutans genotypes.

Material and Methods
- Study Population and Bacterial Isolates
This cohort study used 392 samples of S. mutans isolated from the saliva of 20 caries-active adults (with one or more cavitated lesions) and stored at -86°C in brain heart infusion (BHI) broth containing 20% glycerol. The participants were members of eight Brazilian families with low socioeconomic status living in areas with sub-optimal concentrations of fluoride (0.60 to 0.79 mg F/L). All mothers were primiparous and had salivary levels of streptococci from the mutans group ≥ 106 CFU/mL. The isolates had already been identified as S. mutans by checkerboard DNA-DNA hybridization and genotyped by arbitrarily primed PCR (AP-PCR) using the arbitrary primer OPA-02, in a previous research (thesis available at: http://dx.doi.org/10.11606/T.25.2007.tde-09112007-094447). Microbiological examinations were performed on participants at baseline (T1) when the eldest son was 7-8 months old and after 22 months (T2) of follow-up. In the present work were used the adults S. mutans isolated (mothers, fathers and grandparents) since the genotypes transmitted from parents to children were identical. A pilot study was conducted with a randomized subsample (n = 20) from computer-generated random sets. This research was approved by the Research Ethics Committee of the Bauru School of Dentistry at the University of São Paulo (FOB/USP; Protocol 073/2011).

- Cultivation and Transfer of S. mutans
In a laminar flow hood, 10 μL of each thawed and homogenized sample was streaked in Petri dishes (90×15 mm) containing blood agar medium (BioCen do Brasil, Campinas, São Paulo State (SP), Brazil) and on plates (Interlab Distribuidora de Produtos Científicos SA, São Paulo, SP, Brazil) containing mitis salivarius bacitracin sucrose selective medium (11) (Difco Laboratories, Detroit, Michigan (MI), USA). The plates were stored in anaerobic jars (Difco Laboratories, Detroit, MI, USA) and incubated under microaerophilic conditions (candle flame method) at 37°C for 72 hours. After this period, a putative identification of S. mutans species was conducted based on the morphology and purity of the microbial culture (11) with a stereo microscope (Olympus, Model SZ 40, Shinjuku-ku, Tokyo, Japan).

A representative colony of S. mutans from each mitis salivarius bacitracin sucrose agar plate was selected and aseptically transferred to a culture tube containing 5 mL of BHI broth (Difco Laboratories, Detroit, MI, USA). The tubes were incubated at 37°C in anaerobic jars for 48 hours under microaerophilic conditions. Once bacterial growth was observed, 500 μL was suspended in 20% glycerol and frozen at -80°C. Likewise, aliquots of 1,000 μL were distributed in 2 mL Eppendorf tubes and stored at room temperature for a maximum of 24 hours until the next step was performed.

- Bacterial DNA Extraction
The QIAamp DNA mini kit (Qiagen, Valencia, California (CA), USA) was used for the extraction of bacterial chromosomal DNA following the protocol for manipulating gram-positive bacteria. Sequentially, the purity and concentration of the extracted genetic material were determined by spectrophotometry (Nanodrop spectrophotometer ND-1000, Thermo Fisher Scientific, Wilmington, Delaware, USA). The extract purity was considered adequate when the DNA samples showed an A260 nm/A280 nm ratio between 1.8 and 2.0. Through DNA dilution in ultrapure water, the samples were standardized to a final concentration of 15 ng/μL and stored at -20°C.

- Genetic-Molecular Analysis: GbpA
Bacterial chromosomal DNA samples were amplified by PCR using specific primers flanking the gbpA gene that encodes the GbpA protein (5’-TAGATATCCGACACATTGCAAGTAAATAGAGT-3’ and 5’-TAGATATCCGTTATCATACGACACATACAAA-3’ (6). Each PCR mixture, in a reaction volume of 25 μL, consisted of 2.5 μL of 10X buffer (10 mM Tris-HCl, 50 mM KCl, pH 8.0), 0.75 μL of MgCl₂ (1.5 mM), 0.5 μL of dN-
The specific primers PCR amplified at least one and at maximum four virulence genes from all of S. mutans isolates, corresponding to most and least virulent genotypes isolated. The gbpA, mutAI, mutAII, mutAIII and mutAIV genes were detected in 303/392 (77.3%, 95% CI 72.9 to 81.2), 49/392 (12.5%, 95% CI 9.6 to 16.1), 200/392 (51.0%, 95% CI 46.1 to 55.9), 65/392 (16.6%, 95% CI 13.2 to 20.6) and 352/392 (89.8%, 95% CI 86.4 to 92.4) of isolates, respectively.

The set of clinical isolates used (N = 392) included all 24 different S. mutans genotypes previously genotyped and represented by different letters; 11/24 (45.8%) genotypes were identified as the most virulent (Table 2). Table 2 also shows the qualitative virulence ratings for all the identified S. mutans genotypes and also identifies those transmitted from adults to children in each family, those shared by at least two subjects in a family and those that remained stable in the oral cavity of at least one family member over 22 months of follow-up. High virulence was observed in 6/11 transmitted genotypes, 10/23 shared genotypes and 8/17 stable genotypes. The virulence of S. mutans was associated with its tran-
mission (P < 0.01) and stability (P = 0.96). The most virulent genotypes having higher transmissibility (RR = 1.83, 95% CI 1.44 to 2.32) and higher stability in the oral cavity (RR = 1.52, 95% CI 1.06 to 2.19). With a 95% confidence interval, between 14.7% to 32.2% of transmission of most virulent S. mutans genotypes would be prevented if the risk factors were eliminated (Table 3).

Discussion
The present work provides insight into the mechanism by which bacterial virulence contributes to the coloni-

Table 1: Distribution of number of S. mutans isolates from each family member at baseline (T1) and after 22 months (T2) of follow-up and general characteristics.

<table>
<thead>
<tr>
<th>Family</th>
<th>Mother (n = 8)</th>
<th>Father (n = 8)</th>
<th>Grandparent (n = 4)</th>
<th>Total (n = 20)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>20 (10/10)</td>
<td>20 (10/10)</td>
<td>19 (9/10)</td>
<td>59 (29/30)</td>
</tr>
<tr>
<td>2</td>
<td>20 (10/10)</td>
<td>20 (10/10)</td>
<td>20 (10/10)</td>
<td>60 (30/30)</td>
</tr>
<tr>
<td>3</td>
<td>19 (9/10)</td>
<td>19 (9/10)</td>
<td>20 (10/10)</td>
<td>58 (28/30)</td>
</tr>
<tr>
<td>4</td>
<td>19 (9/10)</td>
<td>20 (10/10)</td>
<td>NA</td>
<td>39 (20/19)</td>
</tr>
<tr>
<td>5</td>
<td>20 (10/10)</td>
<td>20 (10/10)</td>
<td>NA</td>
<td>40 (20/20)</td>
</tr>
<tr>
<td>6</td>
<td>20 (10/10)</td>
<td>20 (10/10)</td>
<td>NA</td>
<td>40 (20/20)</td>
</tr>
<tr>
<td>7</td>
<td>20 (10/10)</td>
<td>18 (8/10)</td>
<td>18 (8/10)</td>
<td>56 (26/30)</td>
</tr>
<tr>
<td>8</td>
<td>20 (10/10)</td>
<td>20 (10/10)</td>
<td>NA</td>
<td>40 (20/20)</td>
</tr>
</tbody>
</table>

Total | 158 (79/79)    | 157 (77/80)   | 77 (37/40)          | 392 (193/199) |

Age (± SD)*
19.37 ± 2.66  23.25 ± 4.06  44.75 ± 6.75  32.9 ± 6.9

DMFS (± SD)*
20.25 ± 10.85  18.5 ± 9.05  6.75 ± 16.80  43.7 ± 12.8

NA—Not available; n—Subject numbers; X̄—Mean; SD—Standard deviation; DMFS—Number of decayed, missing, and filled tooth surfaces; * At baseline.

Table 2: Characterization of S. mutans genotypes detected.

<table>
<thead>
<tr>
<th>Family</th>
<th>Genotypes (Number of isolates)</th>
<th>Transmitted genotypes</th>
<th>Shared genotypes</th>
<th>Stable genotypes§</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>A' (40), B (19)</td>
<td>A'</td>
<td>A', B</td>
<td>A', B</td>
</tr>
<tr>
<td>2</td>
<td>C' (27), D' (16), E' (17)</td>
<td>C', D', E'</td>
<td>C', D', E'</td>
<td>C', D', E'</td>
</tr>
<tr>
<td>3</td>
<td>F' (20), G' (19), H' (19)</td>
<td>F', G', H'</td>
<td>F', G', H'</td>
<td>F', G', H'</td>
</tr>
<tr>
<td>4</td>
<td>I' (23), J (16)</td>
<td>I'</td>
<td>I', J</td>
<td>I', J</td>
</tr>
<tr>
<td>5</td>
<td>K' (17), L' (6), M' (12), N' (5)</td>
<td>K', L', M', N'</td>
<td>K', L', M', N'</td>
<td>K', L', M', N'</td>
</tr>
<tr>
<td>6</td>
<td>O' (23), P (17)</td>
<td>O'</td>
<td>O', P</td>
<td>O'</td>
</tr>
<tr>
<td>7</td>
<td>Q' (8), R (25), S (14), T' (8), U' (1)</td>
<td>Q', R, S, T', U'</td>
<td>Q', R, S, T', U'</td>
<td>Q', R, S, T', U'</td>
</tr>
<tr>
<td>8</td>
<td>V' (17), X' (14), Y' (9)</td>
<td>V', X', Y'</td>
<td>V', X', Y'</td>
<td>V', X', Y'</td>
</tr>
</tbody>
</table>

Total | 24 (392) | 54.5% (11/24) | 95.8% (23/24) | 70.8% (17/24) |

* Most virulent S. mutans genotypes isolated; † Least virulent S. mutans genotypes isolated; § Genotypes detected at baseline (T1) and after 22 months (T2) of follow-up; Genotypes A and Q: gbpA, mutAI, mutAIII, mutAIV genes; Genotype D: gbpA, mutAIII, mutAIV genes; Genotypes E, F, I, M, O, T and X: gbpA, mutAI, mutAIV genes; Genotype U: mutAI, mutAII, mutAIV genes; Genotypes B, C, H, S and Y: gbpA, mutAII, mutAIII, mutAIV genes; Genotypes G, J and L: mutAI, mutAIII, mutAIV genes; Genotypes K and P: gbpA, mutAII genes; Genotypes R and V: mutAIV gene; Genotype N: mutAII gene.
However, the mutAI gene was found to be the most prevalent in previous research (17). The low detection frequencies of some of these structural genes reveal the existence of a wide genetic diversity in the mutA gene locus or even its absence in the samples tested (18). Due to the broad detection spectrum shown by the analyzed genes, we chose to group the S. mutans genotypes into different virulence classes. This approach made it possible to identify those genotypes with the genetic potential to function as highly virulent colonizers.

The detection of these genotypes in 48% of the analyzed S. mutans isolates could indicate an important biological role of their encoded proteins in the formation of dental biofilm. S. mutans genotypes that produce a broad spectrum of mutacins tend to become predominant over time in most of the oral cavity sites (19).

There is evidence of vertical transmission of S. mutans, i.e., from mother to child. A systematic review and meta-analysis demonstrated an association between S. mutans in mothers and their respective children (20). From this study, the possible intrafamilial transmission between the mother-child pairs was also evidenced among the children and their respective parents and grandparents. This intense sharing of genotypes within each family made it difficult to accurately identify the transmission routes and indicates the need for a reassessment of antimicrobial preventive models focused only on the maternal role (e.g., if the S. mutans genotype was shared by all family members, the child may have acquired the genotype from the mother, the father or even the grandmother). Our results for etiologic fraction in population; P-value—Probability value from the chi-square or Mid-P exact test; ¥ Percentage and absolute frequency (n/N) for the number of S. mutans isolates; * Statistically significant association (P ≤ 0.01); # Added 0.5 to each cell for calculation; § At least one expected value was < 5 and Mid-P exact test was used.

In contrast with the present study, other researchers (18) found that not all the isolates transmitted from mothers to their respective children carry the mutAI gene. It is necessary to consider that the transmission of S. mutans is a process influenced not only by microbial factors but also by host and environmental factors, which modulate host immune defenses and bacterial competitiveness (21). In addition to the salivary level of S. mutans (22), transmission fidelity has been associated with the type of delivery (18), the duration and intensity of breastfeeding (23) and the race of the host (24).

The oral cavity is an open growth system. This means that nutrients and microorganisms are repeatedly introduced and removed, and only those that are able to adhere to a surface or otherwise find refuge in the grooves, cracks or interproximal spaces can overcome the removal forces imposed by salivary flow (25). S. mutans may derive benefit from the availability of specific adhesion mechanisms, particularly those mediated by GbpA. Since the inactivation of the gbpA gene tends to reduce sucrose-dependent adhesion in vitro (26) and in vivo (7), its detection in the analyzed bacterial isolates could explain the persistence of the genotypes in the oral cavity of the family members. There was wide variability in the genetic determinants associated with the detection of S. mutans virulence factors. This variability was demonstrated by the broad spectrum of gbpA, mutAI, mutAII, mutAIII and mutAIV genes detected among the samples studied. These genotypes were more likely to be virulent, with increased transmission from adults to children and persistence in the oral cavity. Thus, the present study suggested that S. mutans genotypes with the genetic information to synthesize GbpA and mutacins show an important ecological advantage during the colonization process, remaining stable in the oral microbiota of the host and favoring bacterial transmission among individuals.

**Table 3: Virulence of S. mutans isolates and the transmission, sharing and stability of genotypes.**

<table>
<thead>
<tr>
<th>S. mutans genotypes isolated [% (n/N)]</th>
<th>Point Estimates (95% CI)</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most virulent</td>
<td>Least virulent</td>
<td>RR</td>
</tr>
<tr>
<td>Risk for transmission</td>
<td>69.1 (130/188)</td>
<td>42.2 (86/204)</td>
</tr>
<tr>
<td>Risk for sharing</td>
<td>99.5 (187/188)</td>
<td>100 (204/204)</td>
</tr>
<tr>
<td>Risk for stability</td>
<td>88.8 (167/188)</td>
<td>79.4 (162/204)</td>
</tr>
</tbody>
</table>

95% CI—95% confidence interval; RR—Relative risk; EFp—Etiologic fraction in population; P-value—Probability value from the chi-square or Mid-P exact test; ¥ Percentage and absolute frequency (n/N) for the number of S. mutans isolates; * Statistically significant association (P ≤ 0.01); # Added 0.5 to each cell for calculation; § At least one expected value was < 5 and Mid-P exact test was used.
References

Acknowledgments

We thanks the Laboratory of Microbiology of Bauru Dental School from Universidade de São Paulo, which provided optimized PCRs with a standard strain of S. mutans, UA159 (ATCC 700610).

Grant support

Grant awarded by Programa de Apoio à Pós-Graduação da Coordenação de Aperfeiçoamento de Nível Pessoal (PROAP/CAPES).

Conflicts of interest

None.

Virulence factors of S. mutans