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Comparative analysis of bacterial profiles in unstimulated and stimulated saliva samples

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Background and objective: The microbial profiles of stimulated saliva samples have been shown to differentiate between patients with periodontitis, patients with dental caries, and orally healthy individuals. Saliva was stimulated to allow for easy and rapid collection; however, microbial composition may not reflect the more natural, unstimulated state. The purpose of this study was to validate whether stimulated saliva is an adequate surrogate for unstimulated saliva in determining salivary microbiomes.

Design: Unstimulated (n = 20) and stimulated (n = 20) saliva samples were collected from 20 orally and systemically healthy, non-smoking participants. Salivary bacterial profiles were analyzed by means of the Human Oral Microbe Identification using Next Generation Sequencing (HOMING NGS), and statistical analysis was performed using Mann–Whitney test with Benjamini–Hochberg’s correction for multiple comparison, cluster analysis, principal component analysis, and correspondence analysis.

Results: From a total of 40 saliva samples, 496 probe targets were identified with a mean number of targets per sample of 203 (range: 146–303), and a mean number of probe targets of 206 and 200 in unstimulated and stimulated saliva samples, respectively (p = 0.62). Based on all statistical methods used for this study, the microbial profiles of unstimulated and stimulated saliva samples collected from the same person were not statistically significantly different.

Conclusions: Analysis of bacterial salivary profiles in unstimulated and stimulated saliva samples collected from the same individual showed comparable results. Thus, the results verify that stimulated saliva is an adequate surrogate of unstimulated saliva for microbiome-related studies.

Keywords: saliva; HOMING NGS; bacteria

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The composition of oral microbiota is highly complex, diverse, and site specific (1, 2), and interactions between the resident microbiota and the host is believed to be critically involved in maintenance of oral health. In addition, local alterations of the oral microbiota in relation to ecological perturbations have been considered a prerequisite for the development of periodontitis and dental caries (3–5).

The salivary microbiota is composed of bacteria shed from different oral surfaces being in direct contact with saliva and contains more than 100 million bacteria/mL of saliva (6), which accounts for approximately 1% of the total amount of DNA present in saliva (7). Using molecular methods, the salivary microbiota in oral health has been characterized (8, 9) and has been shown to be mostly comparable with the microbiota of the throat, the tonsils, and the dorsum of the tongue (8).

Saliva is an attractive medium for studies of biomarkers associated with oral health and disease for several reasons including that it is non-invasive, fast, safe to
Material and methods

Study population
In April 2015, 20 adults who fulfilled the inclusion criteria were enrolled in this study. Inclusion criteria were individuals aged \( \geq 18 \) years seeking dental treatment at the Department of Odontology, University of Copenhagen, Copenhagen, Denmark. Exclusion criteria were as follows: presence of periodontitis or dental caries according to pre-defined criteria, current daily smokers, presence of any systemic disease, use of any kind of medication, and treatment with local or systemic antibiotics within the past 3 months prior to participation.

Smoking habit, systemic health, use of medication, and previous antibiotic therapy were addressed by questionnaire. Clinical screening of periodontitis and dental caries was performed by the same examiner (DB) in combination with the analysis of bite-wing X-rays. Based on our previous studies, periodontitis was defined as bleeding on probing \( \geq 25 \% \) of total sites + minimum two teeth with clinical attachment level \( \geq 4 \) mm + minimum two teeth with probing depth \( \geq 6 \) mm. Pronounced dental caries was defined as untreated caries \( \geq 3 \) surfaces (9, 12–14).

Thus, the study population comprised 20 adults, mean age 29 (range: 20–53 years), 10 male and 10 female Caucasians, all seeking dental care at the Department of Odontology, University of Copenhagen. All participants signed an informed consent prior to participation, and the study was approved by the regional ethical committee (H-15000856) and reported to the Danish Data Authorization (2015-54-0970).

Collection of saliva samples
All saliva samples were collected between 8 AM and 11 AM, and to avoid bleeding or any other contamination of saliva, samples were collected before dental treatment. Participants were instructed not to brush the morning before sampling. Initially, an unstimulated saliva sample was collected, followed immediately by a stimulated saliva sample from the same individual.

Unstimulated saliva samples were collected according to a protocol, modified from a previous study (20). Participants were initially asked to rinse their mouth with tap water prior to sampling, followed by collection of at least 5 ml unstimulated saliva in a plastic cup. Finally, the sample was transferred to a plastic tube and stored at \(-80^\circ C\). Collection of stimulated saliva samples was performed according to a previously described protocol (21). In brief, participants started by flushing thoroughly with tap water followed by chewing of paraffin gum for 1 min. Participants expectorated for 1 min and the saliva was discarded. Participants then continued to expectorate for an additional 3 min in a plastic cup. Finally, the stimulated saliva sample was transferred to a plastic tube and stored at \(-80^\circ C\) as above.

Human Oral Microbe Identification using Next Generation Sequencing
DNA isolation was performed as previously described (9, 12–14), and HOMINGs, the successor to the Human Oral Microbe Identification Microarray (HOMIM), was used for microbial analysis (22–24). In brief, HOMINGs is an Illumina-based next-generation sequencing technique, which in combination with a customized BLAST program (called ProbeSeq for HOMINGs) developed at the Forsyth Institute, enables simultaneous identification of approximately 600 oral taxa at the species level.

The laboratory procedures of HOMINGs were performed using a modified protocol previously described (25). Initially, 10–50 ng of sample DNA were PCR-amplified using V3-V4 Forward (341F) AATGATACGGCGAC CACCGAGATCTACACTATGGTAATTGTCCTACGG and stored at \(-80^\circ C\) as above.

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GAGGCAGCAG and reverse (806R) CAAGCAGAAG
ACGGCATACGAGATNNNNNNNNNNAGTCA
GTCAGCCGGACTACHVGGGTWTCTAAT primers,
followed by purification (AMPure beads), gel-purification,
and quantification (qPCR). Finally, 12 pM of the library
mixture, spiked with 20% PhiX (Illumina, San Diego, CA),
was sequenced by use of MiSeq (Illumina). In this study, an
average of 34,091 sequences (441 bp long) was obtained
from each sample. Bad sequences were determined as
based on Q scores of less than 25 and were not considered
for stitching of forward and reverse reads. For this study,
chimeric sequences after stitching were not removed for
analyses. However, in a separate analysis, chimeric se-
quences ranged from 10 to 15% of the total reads. Relative
proportions of detected taxa did not vary significantly
(data not shown).

Bacterial identification using ProbeSeq was performed
as all generated sequences were BLASTed against 598
species-specific and 94 genus-specific (sequences that
recognizes two or more closely related taxa within
the same bacterial genus) 16S rDNA-based reference
sequences (17 to 40 bases long), many of which were
originally designed for HOMIM. A complete list of probes
present in ProbeSeq is presented in Supplementary file 1).

Fig. 1. Relative abundance of predominant species-level and genus-level probe targets. (a) Relative abundance of the 20 most
predominant species-level probe targets in each group. (b) Relative abundance of the 10 most predominant genus-level probe
targets in each group.
Statistical analysis

Comparisons of unstimulated saliva samples (n = 20) and stimulated saliva samples (n = 20) at probe level were analyzed using the Mann–Whitney test with Benjamini–Hochberg correction for multiple comparisons. Benjamini–Hochberg correction was used for control of false-positive discoveries in positive-dependent assumptions (26). For this analysis, an adjusted p-value of <0.01 was considered statistically significant. Analysis of comparability of microbial community profiles in unstimulated and stimulated saliva samples collected from the same individual was performed using cluster analysis, principal component analysis, and correspondence analysis. For hierarchal clustering, Spearman Rank correlations with average linkage were used to construct dendrograms. The alpha diversity for microbial profiles was compared by Shannon index calculations using the following formula. $H' = - \sum p_i \ln(p_i)$, where $p_i$ is the relative abundance. Student t-tests, paired and unpaired, were used to determine statistical differences of alpha diversity. Graphpad prism 5 (San Diego, CA) and MeV 4_8_1 (27) were used as statistical software.

Results

General findings

Of a total of 40 saliva samples (20 unstimulated and 20 stimulated), positive identification for targets of 496 probe sequences were observed (421 identifying a bacterial taxon and 75 identifying a bacterial genus, for example, two or more members of the same genus) corresponding to a coverage of 65% of the 768 probe sequences present in the ProbeSeq database (a complete list of bacterial probes detected are presented in Supplementary file 2). The mean number of probe targets identified per sample was 203 (range: 146–303), with a mean number of probe targets of 206 and 200 in unstimulated and stimulated saliva samples, respectively ($p = 0.62$). No difference between distributions of predominant genera was observed between unstimulated and stimulated saliva samples (adjusted $p$-value = 1). An average of 34,091 sequences (range: 10,237–65,108) was
generated out of which 55.3% (range: 46.2–65.6%) and 17.9% (range: 8.2–30.5%) were identified at species-level and genus-level, respectively. In addition, an average of 26.8% (range: 14.8–39.3%) of the sequences generated could not be assigned to either a species-specific or a genus-specific probe sequence based on blast against the ProbeSeq database. No differences between number of sequences (p = 0.51), proportions of sequences identified at species (p = 0.74) and genus levels (p = 0.39), as well as proportions of unmatched sequences (p = 0.60) were observed between unstimulated and stimulated saliva samples.

No difference between unstimulated and stimulated saliva samples at probe level

Based on comparison at probe level, six out of 496 probe targets were present with a significantly different frequency and a different mean proportion, respectively, in unstimulated (n = 20) or stimulated saliva samples (n = 20) (p < 0.05). However, when adjusting for multiple assumptions no probe targets were present with a different frequency or with different mean proportions in unstimulated or stimulated saliva samples (adjusted p-value = 1, Supplementary file 2).

Strong concordance of bacterial salivary profiles in unstimulated and stimulated saliva samples collected from the same individual

Comparison of bacterial community profiles in unstimulated and stimulated saliva samples using cluster analysis demonstrated tight clustering of sample pairs (unstimulated vs. stimulated saliva sample) collected from the same individual (Fig. 2). In addition, principal component analysis and correspondence analysis illustrated clustering between the majority of sample pairs (Figs. 3 and 4). Alpha diversity as calculated using the Shannon index demonstrated no statistically significant differences between microbial profiles of unstimulated versus stimulated saliva samples collectively among subjects (unpaired t-test, p = 0.74, Fig. 5a) or within the same subject (paired t-test, p = 0.60, Fig. 5b).

Discussion

The purpose of this study was to determine whether stimulated saliva was a valid surrogate for the more natural, unstimulated saliva, as we hypothesized that the bacterial component of stimulated saliva might be diluted as compared with unstimulated saliva samples. The main finding was that there was essentially no difference in stimulated versus unstimulated saliva in the same individual with respect to microbiome analysis. There was likely a dilution effect, but this could not be determined in this study. Nevertheless, the overall composition and proportions remained constant, which is why based on data from this study, stimulated saliva samples might be employed as an adequate surrogate of unstimulated saliva used for oral microbiome studies.

Saliva-based analysis has been proposed as an approach for population-based screening of oral health and disease,
and salivary concentrations of several proteins (28, 29), as well as alterations of salivary bacterial profiles (12, 13) have been suggested as candidate biomarkers associated with oral health and disease. The main advantage of using saliva for biomarker analysis is that saliva can be easily and non-invasively collected routinely in the dental office (10, 11, 30). For practical reasons, however, stimulated saliva samples may be preferred over unstimulated saliva samples, as these can be collected in higher volumes and considerably faster than unstimulated saliva samples. Not surprisingly, the proteomic profile of stimulated saliva samples has been reported to be diluted when compared with unstimulated saliva samples, which is why unstimulated saliva samples are preferred for proteomic analysis of saliva (15, 16). A few studies have previously compared the ability to detect specific oral bacteria, in unstimulated versus stimulated saliva samples by means of different culture-dependent microbial techniques (18, 19). Interestingly, both studies reported stimulated saliva samples as superior to unstimulated saliva samples for bacterial identification (18, 19). On the contrary, a recently published report based on pyrosequencing of microbial samples from two healthy individuals suggested that the microbiota of stimulated saliva samples might be diluted compared with unstimulated saliva samples (17). Thus, to the best of our knowledge, a systematic comparative analysis between saliva microbiomes in unstimulated and stimulated saliva samples, using contemporary molecular methods, remains to be performed. Essentially, from a practical point of view, this is crucial when evaluating the feasibility of using salivary bacterial profiles as a potential biomarker for screening of oral health and disease status.

We demonstrated similarities between the microbiomes of stimulated versus unstimulated saliva by several means. For example, essentially no differences in the proportional presence of the most predominant species-level and genus-level probe targets were observed (Fig. 1a and b), and comparative analysis was very similar based on cluster analysis (Fig. 2), principal component analysis (Fig. 3), and correspondence analysis (Fig. 4). In addition, alpha diversity, as measured using the Shannon index, demonstrated further that there were no statistically significant differences in the microbial profiles of stimulated and unstimulated saliva (Fig. 5a and b). Furthermore, no differences at probe level based on proportional presence between unstimulated saliva samples and stimulated saliva samples were evident (adjusted p-value = 1, Supplementary file 2). Notably, these findings suggest that even though the bacterial component of stimulated saliva samples may be diluted when compared with unstimulated saliva samples, comparable information on the proportional presence of the salivary microbiota could be obtained from both stimulated and unstimulated saliva samples by means of HOMING analysis. This indicates that it is feasible to use salivary bacterial profiles as biomarkers of oral health and disease. Consequently, data from this study highlight that salivary microbiomes can be accurately characterized by analysis of stimulated saliva samples. Thus, based on these findings, we propose that longitudinal screening of the salivary microbiota may be performed using stimulated saliva samples, and speculate that, at the individual level, alterations of the salivary microbiota might hold the potential to identify local bacterial alterations associated with oral diseases at preclinical stages.

In summary, the data suggest that in microbiome-based studies, stimulated saliva samples can serve as an adequate surrogate for unstimulated saliva. Use of stimulated saliva in such studies will significantly reduce time spent, and hence cost, on sample collection. Furthermore, the shorter duration of collection time is easier and more comfortable for patients.

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