Bacterial Composition and Metabolomics of Dental Plaque From Adolescents

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Supragingival dental plaque samples were collected from 40 Swedish adolescents, including 20 with caries lesions (CAR) and 20 caries-free (CF). Fresh plaque samples were subjected to an ex vivo acid tolerance (AT) test where the proportion of bacteria resistant to an acid shock was evaluated through confocal microscopy and live/dead staining, and the metabolites produced were quantiﬁed by 1H Nuclear Magnetic Resonance (1H NMR). In addition, DNA was extracted and the 16S rRNA gene was sequenced by Illumina sequencing, in order to characterize bacterial composition in the same samples. There were no signiﬁcant differences in AT scores between CAR and CF individuals. However, 7 out of the 10 individuals with highest AT scores belonged to the CAR group. Regarding bacterial composition, Abiotrophia, Prevotella and Veillonella were found at signiﬁcantly higher levels in CAR individuals (p=0.0085, 0.026 and 0.04 respectively) and Rothia and Corynebacterium at signiﬁcantly higher levels in CF individuals (p=0.026 and 0.003). The caries pathogen Streptococcus mutans was found at low frequencies and was absent in 60% of CAR individuals. Random-forest predictive models indicate that at least 4 bacterial species or 9 genera are needed to distinguish CAR from CF adolescents. The metabolomic proﬁle obtained by NMR showed a signiﬁcant clustering of organic acids with speciﬁc bacteria in CAR and/or high AT individuals, being Scardovia wiggsiae the species with strongest associations. A signiﬁcant clustering of ethanol and isopropanol with health-associated bacteria such as Rothia or Corynebacterium was also found. Accordingly, several relationships involving these compounds like the Ethanol : Lactate or Succinate : Lactate ratios were signiﬁcantly associated to acid tolerance and could be of predictive value for caries risk. We therefore propose that future caries risk studies would beneﬁt from considering not only the use of multiple organisms as potential microbial biomarkers, but also their functional adaptation and metabolic output.

Keywords: Streptococcus mutans (S. mutans), NMR, metabolomics (OMICS), Rothia, acid tolerance, caries risk (assessment), dental caries, microbiome
INTRODUCTION

Microorganisms in the oral biofilm can metabolize dietary carbohydrates to produce organic acids, which will decrease pH and initiate demineralization of dental hard tissues (Marsh, 1994). *Streptococcus mutans* has for decades been considered the key pathogen for dental caries (Loesche et al., 1975), due to its well-known acidogenic/aciduric properties as well as its ability to attach to enamel. However, during the last decades there has been a switch from explanations focused on specific bacteria to an ecology-centred hypothesis of the disease (Marsh, 1994). In addition, other bacteria, ranging from *Bifidobacterium* to *Scardovia wiggsiae* (Kressirer et al., 2017) have also been found to be involved in tooth decay, and high-throughput sequencing data underline that caries is a polymicrobial disease (Simón-Soro and Mira, 2015). This hampers the strategy of using individual species as predictors of caries risk (Mira, 2018), which has been the classical approach with cultures of mutans streptococci or lactobacilli, and a global analysis of salivary microbiota has been proposed as an alternative (Teng et al., 2015). Other authors have advised that microbial analysis of oral samples should assess not only the taxonomic composition but also their functional features (Takahashi, 2015; Nascimento et al., 2017).

To identify key biomarkers for assessing bacterial composition and metabolic functions related to caries development, metatranscriptomic, proteomic, or metabolomic approaches can be used (Nyvad et al., 2013). For instance, metabolomic analysis of plaque samples after a sugar rinse has been performed by mass spectrometry, identifying key metabolites produced as a consequence of sugar fermentation (Takahashi et al., 2010). Recently, 1H Nuclear Magnetic Resonance spectroscopy (NMR) has been applied to identify a wide range of metabolites in oral samples, especially saliva (Gardner et al., 2018; Pereira et al., 2019). NMR is inherently quantitative and allows the detection and quantification of organic acids, alcohols and other metabolites in a small sample volume (Bertram et al., 2009). However, analyses of saliva samples will show metabolites produced by all oral communities including the tongue, palate, or epithelial mucosa among others, and metabolomic analysis of dental plaque samples would therefore be more informative in relation to caries development. Thus, the recent development of standardized ex vivo tests of acidogenicity and acid tolerance of dental plaque (Senneby et al., 2017) could provide a unique opportunity to study the metabolomic profile of caries-associated microbial communities. Adolescents may have a higher sensitivity for caries compared to other age-groups, as this period in life involves physical and psychological maturation together with a number of new teeth (Silk and Kwok, 2017). To our knowledge, research comparing bacterial composition and metabolomic data in adolescents is lacking, compared to the larger availability of data in adults and in childhood caries.

Thus, the aim of the present study is to compare bacterial composition, acid tolerance and metabolic output of oral bacteria in dental plaque samples, collected from adolescents with and without caries experience. To address this mixed taxonomic and functional analysis, we used a combination of NMR, high-throughput DNA Illumina sequencing and fluorescence microscopy on the same plaque samples, with the aim of understanding the interactions between bacteria and their metabolic products, as well as relating their composition, acidogenicity and acid tolerance to the risk of developing dental caries.

MATERIALS AND METHODS

Study Design and Patient Selection

Forty subjects, aged 14-18 years, were selected (50% males). The participants received routine treatment at two Public Dental Services clinics, which were selected in areas with highest caries prevalence in adolescents, according to the statistics from Public Dental Service, Diver, in the county of Jönköping, Sweden. Ethical approval was obtained from the Regional Ethics Committee for Human Research at Linköping University, Sweden 2017/599-31 and from Swedish Ethical Review Authority, with reference 2019-05656. An intraoral examination included radiographic examination, registration of caries lesions and dental plaque index. All participants had permanent molars in occlusions and exclusion criteria included antibiotic treatment within three months, routine use of oral antiseptics, daily smoking, or regular use of snuff, and systemic or autoimmune diseases.

Study participants included a group of 20 individuals with caries lesions (CAR) and 20 who were caries-free (CF). Inclusion criteria in the caries-free group were no manifest or initial caries lesions and no past history of caries (DMFT index=0). Participants in the caries group had ≥3 surfaces with initial or manifest proximal and/or buccal caries lesions and a past history of caries (fillings). All participants were examined by the same specialist in pediatric dentistry. Initial and manifest caries were registered (clinical and radiographic) according to the criteria by Koch, 1967 and Alm et al., 2007. For intra-individual calibration, all bitewings were reviewed after an interval of three months. The weighted kappa value was 0.61 (CI 0.54-0.68). The percentage agreement was 89.7%.

Sample Collection

Sampling was performed between October 2018 to February 2019. All participants were instructed to avoid tooth brushing from the night before sampling. Plaque was obtained by using a sterile Quickstick (Dab Dental AB) at interproximal sites in molar and premolar areas (3 sites per quadrant, 12 sites per individual). Sampling was avoided if any tooth was missing or if it didn’t exist any contact point between neighboring teeth. Interproximal plaque samples were pooled to give one sample (final). Sampling was repeated if any plaque couldn’t be obtained. Plaque was obtained at each site before using the plaque remover. The material obtained was put into a sterile Quickstick (Dab Dental AB) at interproximal sites in molar and premolar areas (3 sites per quadrant, 12 sites per individual). Sampling was avoided if any tooth was missing or if it didn’t exist any contact point between neighboring teeth. Interproximal plaque samples were pooled to give one sample (final). Sampling was repeated if any plaque couldn’t be obtained. Plaque was obtained at each site before using the plaque remover. The material obtained was put into a sterile Quickstick (Dab Dental AB) at interproximal sites in molar and premolar areas (3 sites per quadrant, 12 sites per individual). Sampling was avoided if any tooth was missing or if it didn’t exist any contact point between neighboring teeth. Interproximal plaque samples were pooled to give one sample (final).
plaque sample was mixed with 75 μl TYE medium (1.7% tryptone, 0.3% yeast) containing 20 mM glucose and 40 mM phosphate/citrate buffer adjusted to pH 3.5 and incubated aerobically at 37°C for 2 hours. Following incubation, the cells were stained with LIVE/DEAT® BacLight™ Fluorescent Stain (Molecular Probes) and transferred into an Ibidi mini flow cells (Ibidi GmbH). Flow-cells were then viewed with confocal laser scanning microscopy (CLSM) using a Nikon Eclipse TE2000 microscope (Nikon Corp.) with an Ar laser (488 nm laser excitation). Images were acquired with a Photometrics Prime 95B camera using Nikon NIS-Elements software. Ten randomly selected images from each sample were saved for further analysis. All confocal images were examined by an experienced oral microbiologist and given a score 1-5 as described by Senneby et al. (2017). This method has been shown to have high intra-rater agreement and the scoring corresponding well to the percentage obtained by manually counting the cells (Senneby et al., 2017).

**Generation of Metabolic End Products**

Two hundred μl of the suspended plaque sample was diluted in 37°C 1,2% NaCl solution to a final concentration of 0.9% NaCl. One molar sterile glucose solution was added giving a final concentration of 100 mM glucose. This was followed by incubation at 37°C for 60 min. After incubation, the sample was cooled on ice for minimum 5 min, then centrifuged at 14000 rpm for 10 min. Supernatants were stored at -80°C until further analysis. Metabolomic analysis of plaque supernatants were obtained by NMR.

**Metabolomic Analysis of Dental Plaque: NMR Data Acquisition and Processing**

Sample supernatants were thawed at room temperature for 20 min and spun down at 2000xg, 4°C for 5 min in a S-4-72 swing-out rotor of a 5804R centrifuge (Eppendorf). 180 µl supernatant was added to a 2 ml cryo vials (Sarstedt) containing 20 µl buffer (1.5M potassium phosphate, pH 6.95, in 100% D2O, 0.5% w/v sodium azide and 0.1% w/v 3-(trimethylsilyl)propionic-2,2,3,3-d4 acid (TSP-d4)). Similarly, one blank sample (0.1 M glucose, 0.9% w/v sodium chloride) was made in the same way and also a buffer-only sample, exchanging sample volume with LC-MS-grade water. The cryovials were put into a rack and shaken at 800 rpm for 2 min at 12°C in a Thermomixer Comfort (Eppendorf). 180 µl of each sample was transferred to a 3 mm SampleJet NMR tube rack using a Bruker SamplePro Tube L liquid handling robot, keeping samples and NMR tube rack at 2°C throughout the preparation. The SampleJet rack was put in a cooled SampleJet sample changer on an 800 MHz Oxford magnet equipped with an Avance III HD console and a 3 mm TCI cryoprobe (Bruker BioSpin). 1D 1H NMR data was acquired using the ‘zgespe’ pulse sequence, encompassing water suppression through excitation sculpting with a perfect echo element. 128 scans were acquired into 65536 data points with a spectral width of 20 ppm, acquisition time of 2.04 s, inter-scan relaxation delay of 3 s and a fixed receiver gain of 18. The low receiver gain setting was necessary to avoid analog/digital converter overflow stemming from the intense glucose signals. TopSpin3.5pl7 (Bruker BioSpin) was used for acquisition and processing of data. Data was Fourier-transformed, including zero-filling twice, 0.3 Hz line-broadening, phasing, baseline correction and referencing to TSP-d4. Processed data was imported into MatLab 2017b (MathWorks Inc.), aligned with icoshift (Savorani et al., 2010) and integrated manually peak-by-peak down to a linear baseline using an in-house developed routine. Peak annotation was accomplished with Chenomx 8.4 (Chenomx Inc.) and the Human Metabolome Database (HMDB) (Wishart et al., 2018). All NMR data on an individual basis are provided as a Supplementary Dataset.

**NMR Data Correlation Analyses**

In order to compare the abundance of detected metabolites, data (arbitrary units, au) were normalized by ng of plaque’s DNA. We used Sparse Partial Least Squares (sPLS) (canonical mode) as a multivariate methodology to perform simultaneous variable selection in the two datasets (the species abundance and the NMR output) based on the importance of the association computed (Lé Cao et al., 2009). These associations were plotted in a heatmap and a network using the mixOmics R package (Rohart et al., 2017).

**Sequence of the 16S rRNA Gene**

DNA from plaque samples was extracted using the MagNa Pure LC DNA Isolation kit II in a MagNa Pure Robot (Roche), following the protocol recommended by the manufacturer with some modifications (Dzidic et al., 2018). DNA concentration was estimated with the Quant-it™ PicoGreen® dsDNA Assay Kit and a Qubit™ 3 Fluorometer (ThermoScientific). The V3-V4 hypervariable region of the 16S rRNA gene was amplified using the universal primers V3-V4 Forward (CCTACGGGNGGCWGCAG) and V3-V4 Reverse (GACTACHVGGGTATCTATCACT) optimized for Illumina sequencing. Library was constructed using the Metagenomic Sequencing Library Preparation Illumina protocol (Part #15044223 Rev. A) and sequenced at the FISABIO Institute (Valencia, Spain) using 2x300 bp paired-end sequencing with an Illumina MiSeq instrument. Sequencing data have been publicly deposited in the SRA database (Bioproject PRJNA681486, Accession number SRR13194555- SRR1319594).

**Bioinformatic Analysis of Sequencing Data**

The software Dada2 v1.16 was used to filter, end-trim, denoise and merge paired reads (Callahan et al., 2016) using default parameters. Adapters and primers were first filtered out from the sequence reads and then end-trimmed in 10 bp windows with quality values <35 and absence of Ns. Singleton reads were removed except for calculating richness and diversity indexes. The remaining reads were merged, clustered and cleaned for host and chimeric reads and finally assigned a taxon at the genus and species level (with Amplified Sequence Variants, or ASVs) using the SILVA non-redundant database v138.1 (Quast et al., 2013).

We used overall R programming language for statistical computing (R Development Core Team, 2016) to perform downstream analyses. Genera with an abundance of <0.01% were removed from all samples. For multivariate analysis, an Adonis test (Permutational Multivariate Analysis of Variance Using Distance Matrices), provided by the Vegan library of R (Oksanen et al., 2015), was used to compare groups. Rarefaction curves, richness and diversity indexes were performed with 20,000 sequences per sample. To visualize groups and their
differences in a two-dimensional map, we computed constrained correspondence analysis (CCA) with the Vegan library (Oksanen et al., 2015). For univariate analyses, non-parametric Wilcoxon tests (wilcox.test function of stats library of R) (R Development Core Team, 2016) were performed to test the differences in genera and species-level taxa. Adjusted p-values, obtained by the FDR method, were used.

We followed Random Forest modelling (Liaw and Wiener, 2002) to address complex interactions between bacterial biomarkers, in order to determine the number of bacteria necessary to efficiently discriminate between CAR and CF groups. We used the bootstrapping algorithm implemented in Boruta R library (Kursa and Rudnicki, 2010) to select biomarkers. The model accuracy was evaluated by the area under curve (AUC) values, when using 30 or less bacteria with the top biomarker scores, ending by the diagnostic value with the top 3 biomarkers (a random forest model needs at least 3 variables). This analysis was performed for biomarkers at the genus and species levels.

RESULTS

The mean age of participants was 16.3 and 15.6 years in CAR and CF groups. Three different analyses were performed for each dental plaque sample: 16S rRNA gene sequencing, acid tolerance test and metabolomics. All participants in the caries group exhibited proximal initial caries lesions (range 1-16 surfaces) and ten participants had proximal manifest cavities (0-12 surfaces). Mean and median carious tooth surfaces including initial and manifest caries lesions and dental restorations (D1+mF) were 17.25 and 15.00 (see Supplementary Table 1 for details).

Oral Microbiota Associated to Dental Plaque

After trimming, quality filtering and annotation of the reads the mean number of sequences per sample was 70,266 ± 6,539. According to the rarefaction curves, all samples showed saturation of diversity after 10,000 reads, suggesting that the sequencing effort was sufficient to cover diversity at species level (Figure 1). Bacterial composition was similar to what had previously been described in other populations (Supplementary Figure 1). We detected higher but not significant mean values for richness (as quantified by the Chao1 index) and statistically significant higher diversity values (Shannon index, p=0.05) in the CAR group (Figure 2A). At genus level, there was a significantly higher proportion of Rothia and Corynebacterium and lower abundances of Abiotrophia, Cryptobacterium, Shuttleworthia, Peptostreptococcus, Prevotella, Veillonella and Ruminococcaceae UCG-014 in CF individuals (Figure 3). At species level, Rothia dentocariosa, Corynebacterium matruchoti,
Corynebacterium durum and Gemella sanguinis were significantly more abundant in CF individuals (Figure 2B). Meanwhile, 14 species were associated to CAR individuals, including a not assigned Prevotella, Leptotrichia buccalis, Abiotrophia defectiva, unassigned Saccharimonadales or Prevotella denticola. In addition, we used a random forest analysis to build a model that predicts how many features (genera or species) are needed to differentiate a CAR from a CF individual (Figures 2C, D). As a result, a minimum of 4 species and 9 genera are necessary to differentiate these two groups with an accuracy of 0.87 or higher. Therefore, not a single species could be detected as a robust biomarker discriminating the caries status of participants, due to overlap between the two groups, even for Abiotrophia which presents the most significant difference. The best studied caries-associated bacterium to date, Streptococcus mutans, was found at extremely low prevalence (8 out of 20 individuals in the CAR group and 2 out of 20 individuals in the CF group). Its mean frequency in the CAR group was 0.16% (and 0.003% in the CF group), showing that although it could be a potential biomarker of disease, it is still absent in more than 50% of the studied CAR adolescents.

**Acid Tolerance Test**

As explained before (Senneye et al., 2017), an Acid Tolerance (AT) test allows us to separate individuals into 5 groups depending on their Acid Tolerance level (1 to 5 AT value) being those with a small proportion of acid tolerant bacteria and 5 those with a large proportion. Although 7 out of 10 individuals with high AT scores were CAR, no significant differences in AT were detected between CAR and CF individuals (Figure 4, left panel). However, we did detect differences in the proportions of some specific bacteria. To clarify the differences, groups with the lowest (AT values 1 and 2) and those with the highest tolerance (AT values 4 and 5) were pooled. When these two clusters were compared, several species showed differences in abundance (Supplementary Figure 2). Fusobacterium periodonticum, Prevotella melaninogenica, Campylobacter concisus, and unassigned species of Veillonella and Alloprevotella were significantly more abundant in the individuals with higher acid tolerance. Streptococcus salivarius, Fusobacterium nucleatum and Campylobacter gracilis were more prevalent in the plaque samples that had lower AT. Interestingly, different Fusobacterium and Campylobacter species were associated with different clusters, underlining the importance of species-level resolution analysis.

**Metabolomic Profile**

A total of 67 peaks or peak clusters were aligned and integrated out of which 39 could be unequivocally annotated. The rest were putatively annotated or not assigned and numbered in ascending order. Using normalized data, the most abundant metabolites detected (besides glucose) in the samples were acetate (14.2 ± 17 Kau/ng), metabolite 16 -putatively assigned to lactate-proline mix- (6.9 ± 3.8 Kau/ng), lactate (2.6 ± 2.2 Kau/ng), metabolite 44 (1.8 ± 2.3 Kau/ng) and ethanol (1.8 ± 1.4 Kau/ng). When the concentration of the metabolites was compared between CAR and CF groups, 3-hydroxyisovalerate appeared to be over-represented in the CF samples, but this difference disappeared when using corrected p-values (Table 1). In addition, metabolites 28 (putatively assigned to glutamate-proline) and 25 showed a statistical trend (0.05<p<0.1) but again only when using uncorrected p-values. However, when the metabolites were compared between Low and High AT clusters the number of metabolites differentially represented increased up to 10 and an additional 4 showed a statistical trend. Interestingly, all these 14 metabolites were more abundant in the Low AT group.

Taking into consideration the metabolites that were significantly associated to Low AT group, we calculated several ratios that could help us to separate individuals with different AT capacity (Figure 4, right panel). Lactate/succinate, ethanol/lactate, ethanol-acetate/lactate and ethanol+succinate/lactate significantly separated High AT from Low AT samples. Moreover, lactate/succinate also had a significantly higher value in High AT than Mid AT.

**Microbiota-Metabolites Correlations**

To integrate available data, we studied the putative correlations between species abundances and metabolites concentrations for the different groups studied (CAR, CF, Low AT, Mid AT, and High AT), clustering those bacteria with a similar metabolomic profile in heatmap plots. Significant correlations for CAR and CF groups are shown in Figure 3. Clear clusters were formed by species that correlate significantly with a given set of metabolites (positively and negatively). For example, when correlations for CAR individuals are shown, several organic acids (acetate, succinate, and lactate) form a cluster with high positive correlations with Streptococcus constellatus, Scardovia wiggiae, Capnocytophaga ochracea, Veillonella tolbetsuensis, Atopobium parvulum, or Actinomyces sp. Interestingly, these organic acids do not seem to correlate significantly with any species in the CF group. However, in caries-free individuals, a cluster was formed with some oral microbes typically associated with oral health such as S. oralis, S. parasanguinis, Corynebacterium durum, Rothia dentocariosa or R. mucilaginosa, which correlate positively with isopropanol, 3-hydroxyisovalerate and succinate (among other unknown metabolites) (Figure 5).

In addition, when a network using metabolites-bacterial correlations was built, several species highly correlated with organic acids among other molecules, and Scardovia wiggiae appears to be central in this network (Figure 6). In CF individuals a cluster formed by Rothia species showed a correlation with isopropanol and a metabolite putatively assigned to butyrate.

Regarding the correlation of bacterial species with metabolites in the AT groups, several clusters were also apparent (Supplementary Figure 3). In the Low AT group, three clusters were formed. In one of them, several health-associated, nitrate-reducing organisms are observed such as Kingella denitrificans, Neisseria lactamica or Rothia aeria which all correlate with acetone and some unknown metabolites. A second cluster correlated with succinate, ethanol, ethanol-isopropanol, dimethylamine, isobutyrate and isovalerate. In the microbiota associated with high AT, lactate, acetate, pyruvate, xanthine, formate, and dimethylamine formed a cluster with positive correlations with some putative oral pathogens such as G. morbillorum, Parvimonas micra, Leptotrichia shahii, Porphyromonas catonae, Selenomonas noxia or Treponema socranskii.
FIGURE 2 | Microbiology of dental plaque samples from caries-experienced (CAR) and caries-free (CF) adolescents, as obtained by sequencing of the 16S rRNA gene. (A) Boxplots show diversity (Shannon) and richness (Chao1) indexes in CAR and CF individuals. The p-values (Wilcoxon test) are indicated. (B) Mean abundance of bacterial species found at significantly different proportions in either CF or CAR groups (p-values <0.05, Wilcox test). Bacteria over-represented in CF individuals are highlighted by a green dot. Lower plots represent the accuracy (Area under Curve) of random forest models to discriminate between CAR and CF individuals according to the number of bacterial species (C) and genera (D).
DISCUSSION

The results obtained in the current manuscript regarding bacterial composition are consistent with the ecological hypothesis of dental caries (Marsh, 1994) and with a polymicrobial etiology of the disease (Simón-Soro and Mira, 2015). The prevalence of S. mutans, which has long been considered the main caries causing agent, was only 40% in our CAR samples, where it accounted for 0.16% of the total. Thus, although their mean proportion was higher than that observed in the CF group, it is not a universal marker for the disease (Mira, 2018). This is consistent with the work by Johansson and collaborators, who concluded that the role of mutans streptococci as primary caries pathogens is less pronounced in populations with prevention programs (Johansson et al., 2016), such as the one studied in our work. However, our data identify other bacteria with higher association to caries experience, including Scardovia wiggsiae, Prevotella denticola or Abiotrophia defectiva, all of which have previously been found at higher levels in caries-active children and to have strong acidogenicity (ElSalhy et al., 2016; Kressirer et al., 2017; Zhang et al., 2020). The higher levels of Veillonella in the caries group is probably related to its dependency on organic acids as carbon source, suggesting that this organism could be a good marker for high lactate levels (Belda-Ferre et al., 2012). Health-associated bacteria included Corynebacterium, which is now identified as a key species in biofilm architecture (Mark Welch et al., 2016), and Rothia. The latter are consistently associated with good oral health in multiple studies, and recent work has demonstrated its capacity to buffer extracellular pH by lactate use and ammonia production from dietary nitrate (Rosier et al., 2020).

Our metabolomic data show that S. wiggsiae has a central role in organic acid production and identify this species as a fundamental player in caries risk in adolescents, consistent with its already-established participation in childhood caries (Kressirer et al., 2017). However, it is not universally distributed in all caries-experienced individuals, again suggesting that caries etiology is not only...
polymicrobial but also highly variable among individuals (Mira, 2018). Previous studies targeting specific bacteria have demonstrated that models based on two organisms, such as S. mutans and Lactobacillus (Featherstone, 2000) or S. mutans and Prevotella pallens (Zhang et al., 2020) improve caries assessment compared to the use of a single species. Our open-ended modelling data, based on random forest classification, indicate that optimal models require at least four oral species or nine genera, and even with this high number of organisms, classification accuracy is below 90% (Figure 1). This suggests that apart from taxonomic classification, functional assessment of bacterial communities could be instrumental to predict caries risk.

A limitation of the current study is a modest sample size (n=20 individuals per group) and the use of incomplete 16S DNA sequences (450 bp in length) derived from Illumina reads. The use of short length reads of the 16S rRNA gene has been shown to limit accurate taxonomic assignment at the species level (Claesson et al., 2010). Thus, although we have used an advanced bioinformatic method for assignment Amplicon Sequence Variants to bacterial species, some of these assignments may contain mistakes or just be impossible to be performed, especially in bacterial genera which present similar sequences among their corresponding species such as streptococci, where S. mitis, S. infantis or S. dentisani contain identical sequences in the amplified segment of the 16S rRNA gene analyzed by standard Illumina library protocols (Dzidic et al., 2018). The taxonomic data assigned at the genus level in the current manuscript are therefore more robust, but species-
level analyses are also included given their congruence with the results from genus-level data and the high-resolution of the 16S rRNA gene sequence for many oral taxa. Future work should also consider performing whole-DNA metagenomic sequencing in order to provide a functional assessment of the gene content of caries and caries-free bacterial communities (Belda-Ferre et al., 2012) and studying the potential role of other members of the microbiome, such as fungi, which were not considered in the current manuscript. Another limitation is given by the high variability in caries experience among CAR individuals, and the absence of past caries experience in some participants from this group, which could make them overlap in physiological or microbiological features with caries-free participants. However, our work aimed at identifying biomarkers of caries in a common population, not only in those individuals at the extreme of the caries landscape, even if this implies that the microbial signals of tooth decay may be weak in those individuals with low caries burden.

Central to caries development is the capacity of supragingival biofilm bacteria to metabolize dietary carbohydrates to organic acids that when accumulated and disassociated disturb the delicate balance between enamel de- and remineralization. In the current study, plaque samples were exposed to an excess of glucose to simulate metabolism taking place at carbohydrate intake. NMR was found to be a powerful methodology for metabolite profiling, with over 60 spectral features from 39 unique metabolites assigned in our samples, despite the difficulty imposed by the high glucose content. A lower glucose concentration would not only enable increase of the receiver gain, allowing detection of less abundant metabolites, but also reduce the amount of overlap from the glucose signals, which cover the 3.3 to 3.9 ppm spectral region. Given that some health-associated peaks in the NMR profile could not be unequivocally identified (e.g. compounds 14, 15 or 17), future work should be aimed at identifying their nature and potential role in caries prevention. When considering caries-experiencing individuals, several species display highly significant correlations with organic acid production, including not only lactate but also propionate, acetate, or pyruvate (Figure 5), and a similar pattern was found in high-AT individuals (Supplementary Figure 3). The combined effect of some of those acids could be instrumental for caries development, as it has been demonstrated that acetic and lactic acid have additive demineralizing properties (Featherstone and Rodgers, 1981) and similar synergistic processes could be taking place for other combinations of organic acids. Interestingly, in both CF and low-AT groups, there are no significant correlations of any bacterial species with lactate. On the contrary, in these health-associated communities we identify significant associations with succinate, ethanol, isopropanol or acetone. Some of these associations are strong in nitrate-reducing species like Neisseria, Rothia or Kingella, supporting an important role of nitrate in oral health homeostasis (Rosier et al., 2018) but significant correlations are also found in other bacteria like...
Streptococcus oralis, Capnocytophaga spp. or Corynebacterium durum. The opposite trends shown for lactate and succinate, as well as for lactate and ethanol-isopropanol, are consistent with alternative fermentation pathways that could drive the selection of acid-tolerant over competing bacterial strains in caries and high-AT individuals (Marsh, 1994). A shift from lactate to ethanol production has also been observed in several oral bacteria under low glucose or low growth rate conditions (Mikx and van der Hoeven, 1975; Yamada and Carlsson, 1975). In agreement with these two metabolic paths driving high and low acidogenicity, lactate:succinate and lactate:ethanol ratios are significantly different between AT levels (Figure 4), in line with the altered metabolism towards lactate production by acid tolerant bacteria even when the environmental pH is lowered (Matsui and Cvitkovitch, 2010). Future studies should evaluate if these ratios could have a diagnostic and predictive value for caries disease.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: https://www.ncbi.nlm.nih.gov/genbank/, SRR13194555- SRR1319594.
ETHICS STATEMENT

The studies involving human participants were reviewed and approved by Regional Ethics Committee for Human Research at Linköping University, Sweden 2017/599-31 and from Swedish Ethical Review Authority, with reference 2019-05656. Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

AUTHOR CONTRIBUTIONS

AM: Performed sequencing work, contributed to conception, design, data acquisition and interpretation, and drafted and critically revised the manuscript. MS, HJ, GS: Contributed to conception, design, data acquisition and interpretation, and drafted and critically revised the manuscript. KH Contributed to conception and design, collected samples, performed dental examinations, and drafted and critically revised the manuscript. JN performed acid tolerance experiments, contributed to data analysis, and critically revised the manuscript. AP performed NMR experimental work, contributed to data analysis, and critically revised the manuscript. MC-D contributed to data analysis and designed all figures, performed statistical analysis, and critically revised the manuscript. All authors contributed to the article and approved the submitted version.

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