Insights of the dental calculi microbiome of pre-Columbian inhabitants from Puerto Rico

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Background. The study of ancient microorganisms in mineralized dental plaque or calculi is providing insights of microbial evolution, as well as lifestyles and disease states of extinct cultures; yet, little is still known about the oral microbial community structure and function of pre-Columbian Caribbean cultures. In the present study, we investigated the dental calculi microbiome and predicted function of one of these cultures, known as the Saladoid. The Saladoids were horticulturalists that emphasized root-crop production. Fruits, as well as small marine and terrestrial animals were also part of the Saladoid diet.

Methods. Dental calculi samples were recovered from the archaeological site of Sorcé, in the municipal island of Vieques, Puerto Rico, characterized using 16S rRNA gene high-throughput sequencing, and compared to the microbiome of previously characterized coprolites of the same culture, as well modern plaque, saliva and stool microbiomes available from the Human Microbiome Project.

Results. Actinobacteria, Proteobacteria and Firmicutes comprised the majority of the Saladoid dental calculi microbiome. The Saladoid dental calculi microbiome was distinct when compared to those of modern saliva and dental plaque, but showed the presence of common inhabitants of modern oral cavities including Streptococcus sp., Veillonella dispar and Rothia mucilaginosa. Cell motility, signal transduction and biosynthesis of other secondary metabolites may be unique features of the Saladoid microbiome.

Discussion. Results suggest that the Saladoid dental calculi microbiome structure and function may possibly reflect a horticulturalist lifestyle and distinct dietary habits. Results also open the opportunity to further elucidate oral disease states in extinct Caribbean cultures and extinct indigenous cultures with similar lifestyles.
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Short title: Oral microbiome of Saladoids

ABSTRACT

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INTRODUCTION

Paleomicrobiology or the study of ancient microorganisms is providing insights into microbial evolution, the lifestyles of extinct cultures, as well as ancient diseases. Mummified gut remains (Ubaldi, Luciani et al. 1998, Cano, Tiefenbrunner et al. 2000, Santiago-Rodriguez, Fornaciari et al. 2015), coprolites (Tito, Knights et al. 2012, Santiago-Rodriguez, Narganes-Storde et al. 2013, Cano, Rivera-Perez et al. 2014), bones (Yang, Eng et al. 1998) and dental calculi (Adler, Dobney et al. 2013, Warinner, Rodrigues et al. 2014, Warinner, Speller et al. 2015, Ziesemer, Mann et al. 2015) are known to harbor ancient microbial DNA. Dental calculi, particularly, is known to be commonly recovered from archaeological excavations across diverse geographical regions, and studies have shown that it is an important bacterial reservoir (Vandermeersch, Arensburg et al. 1994, Arensburg 1996, Preus, Marvik et al. 2011). Previous paleomicrobiology studies relied heavily on techniques such as microscopy and amplification of genes from particular bacterial species using the Polymerase Chain Reaction (PCR) (Pääbo, Higuchi et al. 1989, Cano and Borucki 1995, Pap, Tillier et al. 1995, Willerslev and Cooper 2005). Recent advances in high-throughput sequencing and bioinfomatic tools have enabled the study of bacterial communities by simultaneously amplifying and sequencing the 16S ribosomal RNA (16S rRNA) gene (Caporaso, Kuczynski et al. 2010).

The modern human oral microbiome includes diverse communities composed of >1,000 bacterial species belonging to the Actinobacteria, Bacteroidetes, Firmicutes and Proteobacteria (Dewhirst, Chen et al. 2010). Unlike the human gut microbiome, little is still known about the potential effect that lifestyles, dietary habits and oral hygiene may exert on the oral microbiome (Adler, Dobney et al. 2013, Clemente, Pehrsson et al. 2015). A previous study characterizing the
oral microbiome of individuals from three distinct geographical regions found profound bacterial
taxonomic differences that may be attributed to culture (Li, Quinque et al. 2014). Another study
found that the oral microbiome of uncontacted Amerindians is distinct and more diverse
compared to subjects with westernized lifestyles (Clemente, Pehrsson et al. 2015). Similarly, a
study characterizing the gut microbiome of hunter-gatherers from South America showed that it
is different and more diverse compared to that of westernized cultures (Obregon-Tito, Tito et al.
2015). These studies indicate that modern, isolated societies have distinct microbiomes
compared to modern westernized societies, and that these may better resemble the ancestral or
original state of the human microbiome.

Microbial characterization of ancient human societies may potentially provide direct
insights into the ancestral state of the oral microbiome (Warinner, Speller et al. 2015). Ancient
human oral microbiomes have been studied through dental calculi, which have been found to be
dominated by Actinobacteria, Firmicutes, Proteobacteria and Bacteroidetes, and also harbor oral
pathogens (Warinner, Rodrigues et al. 2014). Previous studies have characterized the oral
microbiome of ancient European cultures (Adler, Dobney et al. 2013, Warinner, Rodrigues et al.
2014). Results have shown that changes in dietary habits associated with specific time periods
affect the structure and function of the human oral microbiome (Adler, Dobney et al. 2013). Pre-
Columbian cultures also had a significant impact on modern western societies, and
characterization of their oral microbiome may also provide insights of the ancestral state of the
human oral microbiome throughout part of human history (Ziesemer, Mann et al. 2015). It is
feasible to hypothesize that the recovered dental calculi may resemble taxonomic and functional
profiles of modern dental plaque, and also possibly reflect pre-Columbian lifestyles.
Archaeological evidence suggests that pre-Columbian cultures came in close contact with each other, suggesting long-distance migrations possibly due to knowledge in seafaring (Curet 2005). The arrangement of the Caribbean archipelago may have facilitated the migration of pre-Columbian cultures from diverse regions across the Americas (Fig 1A). Migration throughout the Caribbean resulted in the settlement and development of a diverse mosaic of cultures. Puerto Rico, particularly, was an important site of ancient human migrations in the Caribbean, representing the longest continuous span of occupation in the Antilles (Curet 2005).

Archaeological evidence in Puerto Rico suggest two major pre-Columbian migrations, that included the Archaic people and the Saladoids (Siegel 2005, Siegel 2010). The Saladoids migrated from South America (lower Orinoco River) and established at the site of Sorcé in the municipal island of Vieques, Puerto Rico around 180 B.C., and the main island of Puerto Rico by 430 B.C. While a major hypothesis suggests that the Saladoid culture was divided in two subseries, the Huecan and the Cedrosan Saladoid, an alternate hypothesis suggests that each of the subseries were unique and distinct cultures (Siegel 2005, Ramos 2010). The alternate hypothesis is mainly supported by differences in semi-precious stone ornaments, and ceramic paintings and decorations, the latter which still remain the primary analytical units of pre-Columbian Caribbean cultures. Based on the alternative hypothesis, the Saladoids co-inhabited with the Huecoids in Sorcé for over 1,000 years (5 A.D. to 1170 A.D.) (Siegel 2010).

Archaeological and isotope evidence have suggested that the Saladoids relied on terrestrial resources and brought to Puerto Rico a developed horticultural economy, which may have been modified with interactions with Archaic groups (Siegel 2005). The Saladoid diet included manioc or yuca, sweet potatoes, yautia, squashes, maize, nuts, as well as fruits such as soursop,
papaya, guava and passion fruit (Siegel 2005). Proteins were mainly acquired from the ingestion
of fish, crustaceans, bivalves, sea turtles, birds and small terrestrial mammals (Storde 1982, Baik
2013). While food preparation methods remain largely unknown (Siegel 2005), thoroughly
boiling of food items, such as tubers, prior to consumption would have been a feasible practice.
Food boiling may be consistent with the absence of nucleic acids from food items, such as
manioc, in ancient human samples.

Recently, microbial community analyses of coprolites found in Saladoid deposits from
Sorcé have shown marked differences when compared to coprolites found in Huecoid deposits
from the same site, as well as modern Amazonian cultures, suggesting that the gut microbiome
may reflect culture-specific dietary habits (Santiago-Rodriguez, Narganes-Storde et al. 2013,
Cano, Rivera-Perez et al. 2014). Teeth with dental calculi from the Saladoid culture at Sorcé
have also been recovered, representing an opportunity to provide insights into their oral
microbiome. Unfortunately, Huecoid human remains, including teeth, have not been recovered
from the archaeological site of Sorcé and were not included in the present study; yet, the
previously characterized coprolite microbiomes of the Saladoid culture, as well as the
microbiomes of supragingival and subgivival plaque, saliva, stool and soil from the
archaeological site of Sorcé have been included in the present study. It is hypothesized that the
oral microbiome of the Saladoids may be distinct from modern individuals, possibly due to
differences in culture, lifestyles, dietary habits, and lack of oral hygiene; therefore, the aims of
the present study were to characterize the taxonomic and predicted functional categories of
Saladoid dental calculi using 16S rRNA gene high-throughput sequencing, and compare them to
the microbiomes of modern oral, gut and soil from the archaeological site of Sorcé.
MATERIALS AND METHODS

Archaeological site and sample description

The archaeological excavation site was located at the Sorcé Estate in La Hueca, Vieques, Puerto Rico (Fig 1B). The excavation was directed by archeologists Luis Chanlatte-Baik and Yvonne Narganes-Storde from the Center for Archaeological Investigations at the University of Puerto Rico. A total of 12 loose dental samples were included in the present study and were excavated from the Saladoid deposit YTA-2 at different depths. Three dental calculi samples were recovered from teeth attached to bone fragments that enabled gender or age determination by forensic archeologist Dr. Edwin Crespo from the University of Puerto Rico (Table 1). Age-at-death and sex were determined by the application of morphological and quantitative methods as described in (Bass 1971, Krogman and Işcan 1986, Scheuer and Black 2000). Teeth for this study were acquired with all necessary field permits complying with all relevant regulations for the collection of skeletal remains from archeological sites. Repository information, including the nomenclature for precise identification containing geographical location, excavation site and archaeological depth are described in Table 1. ^14C dating was conducted by Teledyne Isotopes (Westwood, NJ) and BETA Analytic, Inc. (Miami, FL, USA) using standard methods. Most recovered teeth samples correspond to adult subjects, with the exception of sample H6, which was identified as an infant, showing moderate to severe wear and carious lesions. Fig 1C shows examples of the teeth included in the present study.

Dental calculi recovery and DNA extraction
Dental calculi recovery was performed in a class II hood at the Center for Applications in Biotechnology at the California Polytechnic State University, San Luis Obispo, dedicated for ancient DNA analyses. Hoods were sterilized with UV-radiation at least 15 min before and after every use. Dental calculi were separated from teeth using a dental scalar kindly provided by Dr. Rodney Hiltbrandt, D.D.S. that was sterilized before and after every use with UV-radiation, and bleach between samples. Protective clothing and gloves were worn at all times, and were also changed between samples. The recovered dental calculi were then placed in sterilized tubes and pulverized using custom-made micropestles. Micropestles were sterilized using UV-radiation, and bleach between samples. A mixture of supragingival and subgingival calculi was obtained from each tooth as it was not possible to make a clear distinction. In order to reduce environmental contamination, DNA extraction was performed in dedicated class II hood, where the calculi was also recovered, using the PowerSoil DNA Isolation Kit following manufacturer’s instructions with the following modifications: (i) the PowerBead tube was placed in a Qbiogene Fast Prep Instrument (Carlsbad, CA) at 4.5 m/s for 30 s and then centrifuged at 10,000×g for 30 s; and (ii) incubation periods at -20 °C for 15 min (Mo Bio Laboratories, Carlsbad, CA, USA). The PowerSoil DNA Isolation Kit can recover DNA fragments as short as 60 bp (Qiagen technical support personal communication). All necessary equipment for DNA extraction, including micropipets, was sterilized before and after every use with a 10 % bleach solution and 70 % ethanol. Soil from the archaeological site of Sorcé surrounding the maxillary fragment of sample M8 (Table 1) was recovered as a control for further analyses. A blank control was also included throughout the DNA extraction, PCR amplification and sequencing as described below.

16S rRNA gene amplification and sequencing
DNA amplification of the 16S rRNA gene was performed at Molecular Research Laboratory (MRDNA) (www.mrdnalab.com; Shallowater, TX, USA). All DNA samples were handled in exclusive areas for PCR amplification, which are sterilized before and after every use using DNAaway and UV-radiation to eliminate cross-contamination with modern samples. Template manipulations are handled in separate hoods that are sterilized before and after every manipulation using DNAaway and UV-radiation. Negative PCR controls were included in all amplification reactions, showing no amplification. The 16S rRNA gene V4 variable region was amplified using the universal PCR primers 515f (GTGCCAGCMGCCGCGGTAA)/806r (GGACTACHVGGGTWTCTAAT), as described previously (Caporaso, Lauber et al. 2012). PCR amplifications were conducted using a single step 30 cycle PCR using the HotStarTaq Plus Master Mix Kit (Qiagen, USA) under the following conditions: 94 °C for 3 min, followed by 28 cycles of 94 °C for 30 seconds, 53 °C for 40 seconds and 72 °C for 1 min, after which a final elongation step at 72 °C for 5 min was performed. Two PCR amplifications following the conditions described above were performed due to the limiting starting material and to obtain higher yields to continue with library preparation. After amplification, PCR products were checked in 2 % agarose gel to determine the success of amplification and the relative intensity of the bands. All amplicon products from each sample were mixed or pooled after tagmentation in equal concentrations and purified using Agencourt AMPure beads (Agencourt Bioscience Corporation, MA, USA). The pooled and purified PCR products were used to prepare the DNA library following Illumina MiSeq DNA library preparation protocol using the MiSeq reagent kit v2 (2X150 bp) on a MiSeq following the manufacturer’s guidelines. Preprocessing of reads including removal of chimeras and singletons was performed using proprietary tools from
MRDNA. Phred scores were checked using the FastQC tool

http://www.bioinformatics.babraham.ac.uk/projects/fastqc/

16S rRNA gene analyses

Due to data availability, modern oral and gut samples used for comparison were obtained from the Human Microbiome Project (HMP) (http://hmpdacc.org/). Data correspond to overall healthy individuals, age 18-40, from the USA. Further information about these subjects is available at http://www.ncbi.nlm.nih.gov/projects/gap/cgi-bin/study.cgi?study_id=phs000228.v3.p1. Stool samples corresponded to an Amazonian culture. These samples were chosen over USA individuals because that may better resemble that of the Saladoid culture due to similarities in lifestyles. All 16S rRNA gene sequence analyses were performed using the Quantitative Insights Into Microbial Ecology (Qiime) (Caporaso, Kuczynski et al. 2010). Reads were assigned to samples based on their corresponding barcode using split_libraries.py with default filtering parameters. Operational taxonomic units (OTUs) were selected using the pick_closed_reference_otus.py workflow in Qiime because different regions of the 16S rRNA gene were being characterized. 16S rRNA taxonomy was defined by 97 % similarity to reference sequences. OTU biom files were created for the modern and ancient oral and gut microbiome sequences, and checked using biom summarize-table. OTU biom files were also created separately for the blank control and soil from the archeological site of Sorcé, and checked using biom summarize-table. OTUs present in the blank control (Data Set S1) and soil from the archaeological site of Sorcé (Data Set S2) were removed from all the samples using the workflow described in http://qiime.org/tutorials/filtering_contamination_otus.html to eliminate
the potential contamination effect of laboratory reagents and post-mortem environment to the results. All subsequent analyses were also performed with unfiltered soil and blank control OTUs.

Bayesian Source Tracker or SourceTracker analyses were then performed to identify possible sources of contamination and/or microbiomes sharing some resemblance with the dental calculi microbiomes (Knights, Kuczynski et al. 2011). Microbiomes included in the Bayesian Source Tracker analyses were supragingival plaque (n=5), subgingival plaque (n=5), saliva (n=5), stool (n=5), coprolite (n=5) (described below), soil from the archaeological site of Sorcé (n=1), and a blank control (n=1). Coprolite sequences were included, since certain microorganisms may be part of both oral and gut microbiomes. The mentioned ancient and modern oral, as well as gut microbiomes were also considered in further analyses (Table S1).

The lowest sequence count was 245 after removing from the dental calculi samples OTUs corresponding to the blank control and soil from the archaeological site of Sorcé; thus, data were rarefied to 245 sequences to minimize the effect of disparate sequence number to the results. Rarefaction allows the calculation of species richness through the construction of rarefaction curves. Taxonomy, as well as alpha- and beta-diversity plots were determined using the core_diversity_analyses.py workflow. Alpha diversities, including chao1 and observed OTUs were computed from the average of ten iterations using the collate_alpha.py workflow (Chao 1984, Gotelli and Colwell 2001). Statistical analyses were performed using the alpha_compare.py workflow with default parameters. Since subsampled reads for each comparison were used, a Bonferroni correction was applied. 2D plots were constructed as
described http://qiime.org/scripts/make_2d_plots.html. Group significance analyses were performed using the group_significance.py script in Qiime with default parameters, including Kruskal-Wallis as the statistical test and 1,000 permutations. The test compares OTU frequencies between sample groups to see if these are differentially represented. The category considered in this analysis was sample type including dental calculi, dental calculi from teeth attached to bone, supragingival plaque, subgingival plaque, saliva, stool, coprolite and soil from the archaeological site of Sorcé. The core microbiome between the dental calculi and modern oral microbiomes (i.e. supragingival and subgingival plaque, as well as saliva) was computed using the compute_core_microbiome.py after filtering soil and blank OTUs in order to identify shared OTUs between ancient and modern oral samples. The script identifies the core OTUs in a biom table. Minimum fraction for core was set to 0.25 or 25%. Analyses stopped at 0.40 or 40%, indicating that no OTUs were shared in over 40% of the samples.

Prediction of functional categories based on 16S rRNA gene data was performed using the package Phylogenetic Investigation of Communities by Reconstruction of Unobserved States (PICRUSt) available online at https://huttenhower.sph.harvard.edu/galaxy/ (Langille, Zaneveld et al. 2013). Functional profiles predicted at level 2 were visualized using a heatmap that was constructed using the function heatmap.2 available in the R package gplots. The relative abundances of the functional profiles were also visualized using Linear Discriminatory Analysis Effect size (LEfSe) plots. LEfSe determines the features most likely to explain differences between categories (e.g. sample type). Analyses were also performed with unfiltered soil and blank control OTUs.
RESULTS

16S rRNA gene diversity of dental calculi

Fifteen dental calculi samples from the Saladoid culture were recovered from the archaeological site of Sorcé, Puerto Rico, and the DNA characterized using 16S rRNA gene high-throughput sequencing (Table 1). An average of 692±429 dental calculi sequences with an average size of 272 bp (after quality filtering including chimeras and singletons removal), were analyzed. Average Phred score was 40. An average of 5,097±1,007 (supragingival plaque); 4,094±372 (subgingival plaque); 5,686±1369 (saliva); 2,640,130±548,433 (stool); 48,100±32,461 (coprolites); and 22,945 (soil from the archaeological site of Sorcé) sequences were also analyzed. OTUs corresponding to a blank control and soil from the archaeological site of Sorcé are included in Data Sets S1 and S2, respectively, and were filtered from the dental calculi OTUs, as described above. Prior any further analyses, we performed SourceTracker analyses to identify possible sources of contamination, and modern microbiomes that could potentially share resemblance with the dental calculi microbiomes. Results showed that > 99.0 % of the dental calculi sequences did not resemble any of the modern or ancient human microbiomes (shown as unknown in grey) (Fig S1 Panels A-O). SourceTracker analyses were also performed prior filtering soil and blank control OTUs. Results showed a percentage of OTUs in the dental calculi samples that resemble soil and blank controls prior filtering (Fig S1 Panels A-O). The dental calculi microbiomes were then compared to those of coprolites from the same culture, and modern dental plaque, saliva, stool and soil from the archaeological site of Sorcé. Data were rarefied to 245 sequences (given that this was the lowest sequence count after filtering OTUs from soil and blank control) in order to minimize the effect of disparate sequence number on the results (Table 1 and Table S1). Samples reached or were close to reach a plateau
when data were rarefied, as demonstrated with the chao1 (Fig S2A) and observed OTUs (Fig S2B) rarefaction plots. Rarefaction curves of chao1 (Fig S2C) and observed OTUs (Fig S2D) indices prior filtering OTUs corresponding to soil and blank control from the dental calculi samples are also shown. Two different alpha diversity indices including chao1 (Fig 2A) and observed OTUs (Fig 2B) were plotted after filtering OTUs corresponding to soil and blank control. Chao1 (Fig S3A) and observed OTUs (Fig S3B) indices prior filtering soil and blank control OTUs are also shown. chao1 (Table S2) and observed OTUs (Table S3) values exhibiting statistically significant differences are highlighted in bold. Values shown in Tables S2 and S3 correspond to samples after filtering soil and blank control OTUs. Analyses were also performed with unfiltered soil and blank control OTUs, showing differing p-values in both chao1 (Table S4) and observed OTUs (Table S5) when compared to filtered soil and blank control OTUs.

Principal Coordinate Analyses (PCoA) plots were constructed to visualize beta-diversity indices of the modern and ancient oral and gut microbiomes, as well as that of soil from the archaeological site of Sorcé (Fig 3). Results show clustering based on sample type. Analyses were also performed with unfiltered soil and blank control OTUs, showing a separation based on sample type. Interestingly, the soil sample clustered closely to stool samples from Amazonians (Fig S4). On average, Actinobacteria (14.7 %), Proteobacteria (28.4 %) and Firmicutes (26.5 %) comprised the majority of the Saladoid dental calculi microbiome at the phylum level. Notably, the taxonomic composition of the dental calculi recovered from bone that enabled gender or age determination exhibited variations in these relative abundances. For instance, sample H6, identified as an infant, had 55.7 % of the OTUs classified as Bacteroidetes, from which 43.2 %
where \textit{Bacteroides} sp. Actinobacteria comprised an average of 29.4 \%, 12.7 \%, 2.6 \%, 2.3 \% and 63.0 \% of the bacterial communities in modern supragingival and subgingival plaque, saliva and stool, and coprolites, respectively (\textbf{Fig 4}). The majority of the modern supragingival plaque microbiome was comprised by Firmicutes (27.3 \%), Bacteroidetes (17.6 \%), Fusobacteria (13.5 \%) and Proteobacteria (10.3 \%). The majority of the modern subgingival plaque microbiome was also composed of the same phyla as supragingival plaque, but in differing proportions, depending on the subject (\textbf{Fig 4}). Notably, the microbiome of soil from the archaeological site of Sorcé was mostly comprised by Firmicutes (57.3 \%) and Actinobacteria (34.8 \%) (\textbf{Fig 4}). Taxa identified in the blank control are shown in \textbf{Data Set S3}. Taxonomy analyses at the phylum level with unfiltered soil and blank control OTUs showed differing proportions of Proteobacteria and Firmicutes (\textbf{Fig S5}). Taxonomic analyses at the genus level after filtering soil and blank control OTUs enabled the identification of 869 taxonomic groups, where 345 could not be classified at the genus level (\textbf{Data Set S4}). Genera including \textit{Corynebacterium} sp., \textit{Lactobacillus} sp., \textit{Rothia} sp., \textit{Staphylococcus} sp., \textit{Streptococcus} sp., and \textit{Treponema} sp. were identified in both ancient and modern oral microbiomes. Taxonomic classification at the genus level prior filtering soil and blank OTUs showed 840 taxonomic groups, where 325 could not be classified at the genus level (\textbf{Data Set S5}). A number of unclassified taxonomic groups that were present in soil were also identified in both ancient and modern microbiomes.

\begin{enumerate}
\item Group significance analyses after filtering soil and blank control OTUs showed a number of unique OTUs in the dental calculi microbiomes that were not identified in the modern and ancient gut or oral microbiomes (\textbf{Data Set S6}). There were also a number of shared OTUs between modern oral and dental calculi microbiomes including, but not limited to \textit{Streptococcus}
(OTU 1082294), *Veillonella dispar* (OTU 962249), *Rothia mucilaginosa* (OTU 1017181) and *Porphyromonas* sp. (OTU 4301737) (*Table 2* and *Data Set S6*). Group significance analyses prior filtering soil and blank OTUs showed OTUs shared between soil and the dental calculi samples (*Data Set S7*). Interestingly, OTUs 172063 (*Bacillus* sp.), 961009 (*Acinetobacter johnsonii*), 1088265 (*Propionibacterium acnes*) and 3453734 (*Acinetobacter* sp.) were identified in the soil, dental calculi and Amazonian stool samples. Core microbiome analyses of the dental calculi samples after filtering soil and blank OTUs, and supragingival and subgingival plaque, as well as saliva showed *Rothia mucilaginosa* in both ancient (*Data Set S8*) and modern oral (*Data Set S9*) microbiomes.

**Predicted functional categories**

Functional categories in ancient and modern oral and gut microbiomes were predicted using PICRUSt. In principle, PICRUSt does not predict functional categories from 16S rRNA gene data, since the originating organisms are not being detected, but rather the “bleed over” of putative pathways from the underlying catalog due to cross-annotation of gene families, computationally detected sequence homology, or very rarely from sequence misclassification. Most often this is due to conserved genes being annotated to those pathways in KEGG and other databases, although the actual underlying gene family is something more basic (e.g. cell cycle or structural) (Curtis Huttenhower personal communication). The relative abundances of the functional categories at level 2 were visualized using heatmap plots (*Fig 5*). Overall, the modern gut and oral microbiomes clustered separately from the coprolites, dental calculi and soil from the archaeological site of Sorcé (*Fig 5*). In the cluster including the modern microbiomes, there was a separation based on biogeographical site. In the cluster including the ancient microbiomes,
there was, in turn, a separation between the coprolites and dental calculi microbiomes (Fig 5).

Functional categories were also predicted prior filtering soil and blank OTUs. Results also showed a separation of the dental calculi samples from the modern microbiomes, but expectedly, functional profiles exhibited differing proportions (Fig S6).

Differences in the functional categories predicted using PICRUSt were visualized using LEfSe plots. Dental calculi microbiomes were characterized by functional categories associated with cell motility, signal transduction and biosynthesis of other secondary metabolites. Modern oral microbiomes were characterized by functional categories associated with replication and repair; translation; nucleotide metabolism; metabolism of cofactors and vitamins; glycan biosynthesis and metabolism; folding, sorting and degradation; cell growth and death; as well as signaling molecules and interaction. The microbiome of soil from the archaeological site of Sorcé was characterized by metabolism of other amino acids (Fig 6). Unique functional categories of the coprolites and modern gut microbiomes are also shown in Fig 6. LEfSe analyses of unfiltered soil and blank control OTUs showed that cell motility and signal transduction were still unique features of the dental calculi samples. Categories unique to each sample type were similar prior and after filtering soil and blank control OTUs (Fig S7).

**DISCUSSION**

Contamination of human samples by the *post-mortem* environment is an issue when working with ancient oral samples that have been in direct contact with soil as microbes may penetrate the dentition. This issue has previously been raised for dental calculi found in archaeological sites in the Virgin Islands (Ziesemer, Mann et al. 2015). One possible approach to
authenticate ancient microbiomes is utilizing Bayesian Source Tracker analyses (Tito, Knights et al. 2012, Warinner, Rodrigues et al. 2014, Santiago-Rodriguez, Fornaciari et al. 2015). In the present study, Saladoid’s dental calculi microbiomes did not resemble the modern or ancient microbiomes included in the analyses. SourceTracker analyses showed a number of OTUs in the dental calculi samples resembling soil from the archaeological site of Sorcé and a blank control; thus, these OTUs were filtered from the dental calculi sequences prior main analyses. Filtering OTUs corresponding to soil from the archaeological site of Sorcé and a blank control possibly increased the potential of analyzing sequences associated exclusively with dental calculi. A previous study showed that even after decontamination of the exterior of dental calculi, a number of soil sequences may still be detected; therefore, filtering soil OTUs may give a better resolution of the authentic dental calculi microbiome (Ziesemer, Mann et al. 2015). The present, as well as previous studies have found bacterial DNA in commercially available extraction kits (Glassing, Dowd et al. 2016); therefore, it is of importance to include a blank control in sequencing reactions that could be used to filter potential contaminating OTUs prior further analyses.

While a previous study characterizing the dental calculi microbiomes of ancient European cultures showed that these resembled dental plaque microbiomes of people of European ancestry (Warinner, Rodrigues et al. 2014), our data showed that the dental calculi microbiome of the Saladoid culture is taxonomically different at the phylum level and generally more diverse compared to modern dental plaque. It has been argued that this is due to contamination with soil microbes (Warinner, Rodrigues et al. 2014). While this is a possibility that is often visited by other researchers in the field of ancient DNA, it should also be considered that differences in dietary habits and hygiene practices may have an impact on the taxonomic
structure of the human oral microbiome. Studies characterizing the dental plaque microbiome of modern isolated and industrialized cultures with differences in ancestry, dietary habits and oral hygiene are needed to provide better insights of the evolution of the human oral microbiome throughout part of human history. For instance, a recent study characterizing the saliva microbiome of the Tsimane, indigenous people from the Amazon, showed that the Firmicutes, Bacteroidetes, Actinobacteria and Proteobacteria comprised the majority of the bacterial taxa in adults. The salivary microbiome of Tsimane children was characterized by not resembling that of their mothers and having a higher relative proportion of Firmicutes (Han, Martin et al. 2016). Although researchers sampled saliva, results resemble some our findings. This resemblance may be due to the Tsimane diets consisting primarily of cultivated carbohydrates and fish, similar to the diet of the Saladoid culture from Puerto Rico. We also found evidence of the presence of bacteria known to inhabit the oral cavity of modern subjects in the Saladoid’s dental calculi, including *Streptococcus* sp., *Porphyromonas* sp., and *Gemellaceae* sp., supporting previous findings that bacterial members of the human oral microbiome are ubiquitous across diverse geographical regions and time periods (Adler, Dobney et al. 2013, Warinner, Rodrigues et al. 2014, Clemente, Pehrsson et al. 2015).

Even though the DNA extraction and sequencing methods differed, our data are consistent with another study characterizing the dental calculi microbiome of a pre-Columbian Caribbean culture inhabiting the Virgin Islands, where Actinobacteria comprised a large proportion of the bacterial taxa (Ziesemer, Mann et al. 2015). This same study showed that tropical climates may not have the same degree of ancient bacterial DNA preservation as colder climates. Given that these samples have been in contact with soil for extended periods of time, it
is expected to find 16S rRNA gene sequences matching soil microbiomes, even after external
decontamination of the dental calculi; yet, one advantage of the bioinformatic tools developed
for 16S rRNA gene analyses is that reads corresponding to soil contaminants can be filtered.
Bioinformatic tools available to discriminate potential sources of contamination, such as
SourceTracker, may not conclusively indicate that results in the present study were influenced by
soil contamination. While bioinformatic analyses do not truly support that the presence of
Actinobacteria is due to contamination with the post-mortem environment, it is feasible to
hypothesize that the OTUs present in the dental calculi samples reflect, to some extent, dietary
habits of the Saladoid culture. It is hypothesized that the Saladoid diet was mainly composed of
root crops and fruits, whose residues are known to get entrapped in dental calculi (Mickleburgh
and Pagán-Jiménez 2012). It is also possible that intake of root crops and fruits contaminated
with soil may have had some impact in the bacterial taxa being detected in the dental calculi
samples. Differences in lifestyles and dietary habits may explain some of the lack of similarity
between the Saladoid and ancient European oral microbiomes characterized previously, even
when both of these ancient cultures lacked oral hygienic conditions. In addition, some members
of the Actinobacteria, including Actinomyces have been associated with dental caries in modern
individuals. Detection of Actinomyces DNA in the Saladoid is intriguing as it supports
observations that pre-Columbian cultures from the Caribbean also suffered from oral diseases
(Chalmers, Oh et al. 2015, Ziesemer, Mann et al. 2015). Our data may also be consistent with a
previous study suggesting that dental caries in ancient human populations may have resulted
from the intake of fermentable carbohydrates (Adler, Dobney et al. 2013), as there is
archaeological evidence suggesting that the Saladoid ingested fermentable carbohydrates
(Widstrom, Carr et al. 1987, Humphrey, De Groote et al. 2014) that are part of the root crops.
Results may have also been influenced by the amplified 16S rRNA gene variable region. Recent evidence suggests that amplification of different 16S rRNA gene variable regions may also have profound effects on ancient microbiome studies due to PCR amplification biases (Ziesemer, Mann et al. 2015). This study also suggested that amplification of the V4 region may be impractical for ancient DNA studies; however, <300 bp is still within what would be considered the norm in the field of ancient DNA (Drancourt, Aboudharam et al. 1998, von Wurmb-Schwark, Schwark et al. 2004, Roberts and Ingham 2008, Paéibo 2012). In addition, the V4 region has been better characterized and has shown to have a better phylogenetic resolution (Yang, Wang et al. 2016) than other 16S rRNA gene variable regions.

Due to the small number of recovered sequences after filtering soil from the archeological site of Sorcé and blank control OTUs, special consideration should be taken when interpreting results from the prediction of functional categories in the dental calculi samples. In addition, while PICRUSt was utilized to predict functional profiles based on 16S rRNA gene data, which has shown to work more efficiently with stool microbiomes, results may still add valuable information regarding the potential effect of culture, lifestyle, dietary habits, oral hygiene and disease states to the ancestral state of human oral microbiomes. Ancient and modern oral microbiomes showed shared functional profiles, suggesting that certain predicted functional pathways may be ubiquitous to the human oral microbiome. While taphonomic conditions may have influenced the predicted functional profiles of the dental calculi included in the present study, it is also feasible to hypothesize that some of the predicted functional profiles were altered due to differences in culture, lifestyle, dietary habits, oral hygiene and disease
states, which are known to affect the community structure and function of modern human oral microbiomes (Zarco, Vess et al. 2012).

CONCLUSIONS

Skepticism is met when studying the oral microbiomes of ancient Caribbean cultures as there is always the risk that soil bacteria may have penetrated the dentine; thus, the reliability of the microbiome structure and function of dental calculi excavated from tropical regions may be challenged. Tools such as Bayesian SourceTracker analyses may be used to authenticate ancient microbiomes by providing insights into the contribution of possible contaminating sources, including soil from the archaeological site, as well as reagents. Our data may provide insights of the oral microbiome of the Saladoid culture from Sorcé, Puerto Rico, although a deeper sequencing depth is needed. Comparative studies would ideally include plaque and saliva samples from modern indigenous individuals, but access to this type of samples is challenging. Given that most human microbiomes studies have been performed with subjects from the US limits our ability to interpret some of the results. Bacteria present in the oral cavities of modern subjects were also present in the dental calculi microbiome of the Saladoid culture. Advantages and disadvantages exist when using 16S rRNA gene high-throughput sequencing to characterize ancient human oral microbiomes.

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Figure legends

Fig 1. Location and example of the samples obtained. Panel A shows the organization of the Caribbean as a site for ancient human migrations. Vieques Island, which is part of the main island of Puerto Rico, is highlighted by the red circle. Panel B shows the location of the
archaeological site of Sorcé on Vieques. Panel C shows examples of the teeth recovered from the
archaeological site of Sorcé.

**Fig 2.** Bar plots of alpha diversity indices. Bar plots representing the chao 1 (Panel A) and
observed OTUs (Panel B) indices for the bacterial taxonomy based on 16S rRNA gene of the
dental calculi, modern supragingival and subgingival plaque, saliva, coprolites, stool and soil
from the archaeological site of Sorcé microbiomes. Alpha diversity indices were computed from
the average of ten iterations using the collate_alpha.py workflow. Soil from the archeological
site of Sorcé and blank control OTUs were filtered from the dental calculi prior analyses.

**Fig 3.** Principal Coordinates Analysis (PCoA) 2D plots of ancient and modern oral and gut
microbiomes, as well as that of soil from the archaeological site of Sorcé. Dental calculi, dental
calculi of teeth attached to bones that enabled the identification of gender or age (Dental calculi
(Bone)), coprolites, stool, soil from the archaeological site of Sorcé, supragingival (grey) and
subgingival plaque (black) and saliva (white). Soil from the archeological site of Sorcé and blank
control OTUs were filtered from the dental calculi prior analyses.

**Fig 4.** Barplots representing the bacterial taxonomy based on 16S rRNA gene. Data are shown at
the phylum level for dental calculi, supragingival plaque, subgingival plaque, saliva, coprolites,
stool and soil from the archaeological site of Sorcé. Soil from the archaeological site of Sorcé and
blank control OTUs were filtered from the dental calculi prior analyses.

**Fig 5.** Heatmap of the normalized relative abundances of the predicted functional categories
(level 2) of the microbiomes of dental calculi, coprolites, supragingival and subgingival plaque,
saliva, stool and soil from the archeological site of Sorcé. Functional categories were predicted
using PICRUST. Soil from the archeological site of Sorcé and blank control OTUs were filtered
prior analyses.
Fig 6. Linear discriminatory analyses Effect size (LEfSe) plots of predicted functional categories (level 2). Functional categories of dental calculi (ancient oral microbiomes), supragingival plaque, subgingival plaque and saliva (modern oral microbiomes), coprolites (ancient gut microbiomes), stool (modern gut microbiomes) and soil from the archeological site of Sorcé (tropical soil) were predicted using PICRUSt. Soil from the archeological site of Sorcé and blank control OTUs were filtered prior analyses.

Supplementary Figure legends

Fig S1. SourceTracker analyses of the dental calculi samples including D21 (Panel A), E19 (Panel B), E26 (Panel C), F20 (Panel D), G18 (Panel E), G21 (Panel F), G22 (Panel G), I19 (Panel H), I23A (Panel I), I23B (Panel J), I24A (Panel K), I24B (Panel L), F24 (Panel M), H6 (Panel N) and M8 (Panel O). Dental calculi microbiomes were compared to stool, coprolite, saliva, soil from the archaeological site of Sorcé, subgingival plaque, and supragingival plaque microbiomes. Dental calculi sequences not matching any of the microbiomes included were classified as unknown. Figure shows results prior and after filtering soil and blank control OTUs from the dental calculi.

Fig S2. Alphararefaction curves of chao1 (Panel A), and observed OTUs values (Panel B) after filtering soil and blank control OTUs from the dental calculi, and chao1 (Panel C), and observed OTUs (Panel D) values prior filtering soil and blank control OTUs. Samples included dental calculi from loose teeth samples (yellow), dental calculi from teeth attached to bone (red), supragingival plaque (pink), subgingival plaque (green), saliva (orange), coprolites (light blue), stool (blue), soil from the archaeological site of Sorcé (brown), and blank control (purple).

Fig S3. Bar plots of alpha diversity indices. Bar plots representing the chao 1 (Panel A) and observed OTUs (Panel B) indices for the bacterial taxonomy based on 16S rRNA gene of the
dental calculi, modern supragingival and subgingival plaque, saliva, coprolites, stool and soil from the archaeological site of Sorcé microbiomes. Alpha diversity indices were computed from the average of ten iterations using the collate_alpha.py workflow. Soil and blank control OTUs were not filtered from the dental calculi prior analyses.

**Fig S4.** Principal Coordinates Analysis (PCoA) 2D plots of ancient and modern oral and gut microbiomes, as well as that of soil from the archaeological site of Sorcé. Dental calculi, dental calculi of teeth attached to bones that enabled the identification of gender or age (Dental calculi (Bone)), coprolites, stool, soil from the archaeological site of Sorcé, supragingival (grey) and subgingival plaque (black), and saliva (white). Soil and blank control OTUs were not filtered from the dental calculi prior analyses.

**Fig S5.** Barplots representing the bacterial taxonomy based on 16S rRNA gene. Data are shown at the phylum level for dental calculi, supragingival plaque, subgingival plaque, saliva, coprolites, stool, soil from the archaeological site of Sorcé, and a blank control. Soil and blank control OTUs were not filtered from the dental calculi prior analyses.

**Fig S6.** Heatmap of the relative abundances of the predicted functional categories (level 2) of microbiomes of dental calculi, coprolites, supragingival and subgingival plaque, saliva, stool and soil from the archaeological site of Sorcé. Functional categories were predicted using PICRUSSt. Soil and blank control OTUs were not filtered from the dental calculi prior analyses.

**Fig S7.** Linear discriminatory analyses Effect size (LEfSe) plots of predicted functional categories (level 2). Functional categories of dental calculi (ancient oral microbiomes), supragingival plaque, subgingival plaque and saliva (modern oral microbiomes), coprolites (ancient gut microbiomes), stool (modern gut microbiomes) and soil from the archeological site
of Sorcé (tropical soil) were predicted using PICRUSt. Soil and blank control OTUs were not filtered from the dental calculi prior analyses.

**Supplementary Table legends**

**Table S1.** Modern oral or stool microbiomes subjects’ information. Sequences were downloaded from the Human Microbiome Project (HMP).

**Table S2.** Chao1 statistical analyses. Statistically significant results are shown in bold. Soil and blank control OTUs were filtered from the dental calculi prior analyses.

**Table S3.** Observed OTUs statistical analyses. Statistically significant results are shown in bold. Soil and blank control OTUs were filtered from the dental calculi prior analyses.

**Table S4.** Chao1 statistical analyses. Statistically significant results are shown in bold. Soil and blank control OTUs were not filtered from the dental calculi prior analyses.

**Table S5.** Observed OTUs statistical analyses. Statistically significant results are shown in bold. Soil and blank control OTUs were not filtered from the dental calculi prior analyses.

**Supplementary Data Sets legends**

**Data Set S1.** OTU_table_Blank_Control.txt

**Data Set S2.** OTU_table_Soil.txt

**Data Set S3.** Taxa_found_in_Blank_Control.xls

**Data Set S4.** Dental_Calculi_Genus_Level_Filtered.xls

**Data Set S5.** Dental_Calculi_Genus_Level_Unfiltered.xls

**Data Set S6.** Group_Significance_Dental_Calculi_Filtered.xls

**Data Set S7.** Group_Significance_Dental_Calculi_Unfiltered.xls

**Data Set S8.** Core_OTUs_25_Dental_Calculi.txt

**Data Set S9.** Core_OTUs_25_Modern_Oral_Microbiome.txt
Figure 1

Map Samples

Location and example of the samples obtained. Panel A shows the organization of the Caribbean as a site for ancient human migrations. Vieques Island, which is part of the main island of Puerto Rico, is highlighted by the red circle. Panel B shows the location of the archaeological site of Sorcé on Vieques. Panel C shows examples of the teeth recovered from the archeological site of Sorcé.
Figure 2

Alpha Diversity Filtered

Bar plots of alpha diversity indices. Bar plots representing the chao 1 (Panel A) and observed OTUs (Panel B) indices for the bacterial taxonomy based on 16S rRNA gene of the dental calculi, modern supragingival and subgingival plaque, saliva, coprolites, stool and soil from the archaeological site of Sorcé microbiomes. Alpha diversity indices were computed from the average of ten iterations using the collate_alpha.py workflow. Soil from the archeological site of Sorcé and blank control OTUs were filtered from the dental calculi prior analyses.
Figure 3

PCoA Filtered

Principal Coordinates Analysis (PCoA) 2D plots of ancient and modern oral and gut microbiomes, as well as that of soil from the archaeological site of Sorcé. Dental calculi, dental calculi of teeth attached to bones that enabled the identification of gender or age (Dental calculi (Bone)), coprolites, stool, soil from the archaeological site of Sorcé, supragingival (grey) and subgingival plaque (black) and saliva (white). Soil from the archeological site of Sorcé and blank control OTUs were filtered from the dental calculi prior analyses.
Manuscript to be reviewed
Figure 4

Taxa Plots Filtered

Barplots representing the bacterial taxonomy based on 16S rRNA gene. Data are shown at the phylum level for dental calculi, supragingival plaque, subgingival plaque, saliva, coprolites, stool and soil from the archeological site of Sorcé. Soil from the archeological site of Sorcé and blank control OTUs were filtered from the dental calculi prior analyses.
Figure 5

PICRUST

Heatmap of the normalized relative abundances of the predicted functional categories (level 2) of the microbiomes of dental calculi, coprolites, supragingival and subgingival plaque, saliva, stool and soil from the archeological site of Sorcé. Functional categories were predicted using PICRUST. Soil from the archeological site of Sorcé and blank control OTUs were filtered prior analyses.
Figure 6

LEfSe Filtered

Linear discriminatory analyses Effect size (LEfSe) plots of predicted functional categories (level 2). Functional categories of dental calculi (ancient oral microbiomes), supragingival plaque, subgingival plaque and saliva (modern oral microbiomes), coprolites (ancient gut microbiomes), stool (modern gut microbiomes) and soil from the archeological site of Sorcé (tropical soil) were predicted using PICRUSt. Soil from the archeological site of Sorcé and blank control OTUs were filtered prior analyses.
Sample description.

All samples were collected from the archeological site YTA-2 at the Sorcé Estate in La Hueca, Vieques, Puerto Rico. Samples date from 10 A.D. to 385 A.D.
Table 1. Sample description. All samples were collected from the archeological site YTA-2 at the Sorcé Estate in La Hueca, Vieques, Puerto Rico. Samples date from 10 A.D. to 385 A.D.

| Sample | Depth (m) | Description | Weight (mg) | Sequences prior filtering | OTUs prior filtering | Sequences after filtering | OTUs after filtering |
|--------|-----------|-------------|-------------|---------------------------|---------------------|--------------------------|----------------------|
| D21    | 0.60-0.80 | Maxillary alveolar bone fragment with (6) teeth. Moderate wear. | 2.50 | 2,677 | 497 | 799 | 304 |
| E19    | 0.20-0.40 | Loose tooth with severe wear. | 2.78 | 4,309 | 571 | 1,297 | 371 |
| E26    | 0.60      | Loose molar tooth with severe wear. | 3.02 | 5,933 | 490 | 604 | 288 |
| F20    | 0.80      | Loose tooth with moderate wear. | 4.10 | 3,345 | 334 | 337 | 178 |
| G18    | 0.40-0.60 | Loose teeth (2) with wear | 3.89 | 14,303 | 363 | 374 | 196 |
| G21    | 0.20-0.40 | Loose molar tooth with abscess, and teeth (3). Moderate wear. | 2.92 | 24,371 | 480 | 879 | 300 |
| G22    | 0.20-0.40 | Loose tooth with enamel detachment. Moderate wear. | 4.35 | 5,227 | 707 | 1,863 | 505 |
| I19    | 0.00-0.20 | Loose teeth (2). Moderate wear. | 2.37 | 3,523 | 499 | 877 | 339 |
| I23A   | 0.20-0.40 | Molar tooth and tooth. Moderate wear. | 4.28 | 2,424 | 495 | 673 | 308 |
| I23B   | 0.40-0.60 | Maxillary alveolar bone fragment with (2) teeth and loose tooth. Moderate wear. | 3.17 | 2,031 | 439 | 602 | 278 |
| I24A   | 0.20-0.40 | Loose tooth. Moderate wear. | 1.95 | 2,125 | 418 | 625 | 265 |
| I24B   | 0.60-0.80 | Maxillary alveolar bone fragment with (3) teeth and (2) loose teeth. Moderate wear. | 2.02 | 2,321 | 376 | 602 | 228 |
| F24    | 0.60      | Mandibular alveolar bone fragment with attached teeth (11). Male. Moderate wear. | 5.25 | 3,005 | 110 | 245 | 33 |
| H6     | 0.70      | Infant maxillary alveolar bone fragment with fully erupted tooth and deciduous teeth (3) | 5.14 | 56,503 | 74 | 334 | 31 |
| M8     | 0.00-0.20 | Mandibular alveolar bone fragment with (4) teeth. Male. Moderate wear | 4.98 | 2,214 | 46 | 267 | 23 |
| Soil*  | -         | Archaeological site of Sorcé | - | 22,945 | 336 | - | - |
| Blank* | -         | - | - | 34,392 | 956 | - | - |

3  Dental calculi samples highlighted in bold were attached to bone fragments that enabled the gender or approximate age determination.

4  *Soil and blank sequences were filtered from calculi samples for main analyses.
Table 2 (on next page)

Group Significance

Group significance analyses of dental calculi, dental calculi (Bone), coprolites, supragingival and subgingival plaque, saliva, stool and soil from the archaeological site of Sorcé. Selected OTUs present in both dental calculus and modern oral samples are shown. Group significance analyses were performed after filtering soil and blank OTUs.
Table 2. Group significance analyses of dental calculi, dental calculi (Bone), coprolites, supragingival and subgingival plaque, saliva, stool and soil from the archaeological site of Sorcé. Selected OTUs present in both dental calculus and modern oral samples are shown. Group significance analyses were performed after filtering soil and blank OTUs.

| OTU      | Test Statistic P | FDR P | Bonferroni P | Soil* | Saliva* | Coprolite* | Calculi* | Subgingival plaque* | Calci (Bone)* | Supragingival plaque* | Stool* | Taxonomy                      |
|----------|------------------|-------|---------------|-------|---------|------------|-----------|---------------------|---------------|------------------------|--------|------------------------------|
| 349024  | 29.790           | 0.000 | 0.000         | 1.000 | 0.000   | 0.833      | 0.000     | 0.000               | 0.250         | 32427.600               |        | *Streptococcus sp.*          |
| 1090059 | 26.416           | 0.000 | 0.002         | 1.000 | 3.500   | 0.250      | 1.000     | 0.000               | 2.750         | 174.400                 |        | *Granulicatella sp.*         |
| 139056  | 5.396            | 0.612 | 0.735         | 1.000 | 0.250   | 0.250      | 0.250     | 0.000               | 0.000         | 0.000                  |        | *Aggregatibacter sp.*       |
| 966331  | 5.396            | 0.612 | 0.735         | 1.000 | 0.000   | 0.250      | 0.250     | 0.000               | 0.250         | 0.000                  |        | *Dermacoccus sp.*           |
| 1082294 | 36.401           | 0.000 | 0.000         | 0.087 | 72.750  | 0.167      | 18.250    | 0.000               | 207.000       | 0.000                  |        | *Streptococcus sp.*         |
| 1065974 | 22.882           | 0.002 | 0.005         | 1.000 | 5.250   | 0.167      | 1.500     | 0.000               | 4.500         | 46.800                 |        | *Vagococcus sp.*            |
| 131775  | 5.646            | 0.603 | 0.725         | 1.000 | 0.000   | 0.167      | 0.000     | 0.000               | 1.250         | 0.000                  |        | *Micrococcaceae sp.*        |
| 535196  | 5.123            | 0.645 | 0.759         | 1.000 | 0.000   | 0.167      | 0.250     | 0.000               | 0.000         | 0.000                  |        | *Rhodobacteraceae sp.*      |
| 962249  | 34.888           | 0.000 | 0.000         | 0.167 | 18.500  | 0.083      | 12.250    | 0.000               | 12.250        | 0.000                  |        | *Veillonella dispar*         |
| 1017181 | 29.704           | 0.000 | 0.000         | 1.000 | 8.750   | 0.083      | 4.500     | 0.000               | 4.000         | 20.400                 |        | *Rothia mucilaginosa*       |
| 1059729 | 28.524           | 0.000 | 0.000         | 1.000 | 6.500   | 0.083      | 1.000     | 0.000               | 5.000         | 4.400                  |        | *Granulicatella sp.*        |
| 73471   | 22.635           | 0.002 | 0.006         | 1.000 | 0.000   | 0.083      | 7.750     | 0.000               | 16.250        | 1.200                  |        | *Actinomyces sp.*           |
| 1084417 | 19.585           | 0.007 | 0.018         | 1.000 | 5.000   | 0.083      | 12.000    | 0.000               | 9.250         | 2.800                  |        | *Lautropia sp.*             |
| 925707  | 15.381           | 0.031 | 0.065         | 1.000 | 1.000   | 0.083      | 12.750    | 0.000               | 0.750         | 2.000                  |        | *Gemellaceae sp.*           |
| 499378  | 14.442           | 0.044 | 0.090         | 1.000 | 0.000   | 0.083      | 0.500     | 0.000               | 0.750         | 0.000                  |        | *Actinomyces sp.*           |
| 509773  | 10.181           | 0.179 | 0.334         | 1.000 | 18.500  | 0.083      | 2.000     | 0.000               | 0.250         | 0.000                  |        | *Streptococcus sp.*         |
| 577170  | 9.319            | 0.231 | 0.398         | 1.000 | 0.000   | 0.083      | 0.250     | 0.000               | 0.000         | 4.400                  |        | *Bacteroides sp.*           |
| 542066  | 5.629            | 0.584 | 0.702         | 1.000 | 0.250   | 0.083      | 0.000     | 0.000               | 0.250         | 0.000                  |        | *Actinomycetales sp.*       |
| 432284  | 5.464            | 0.603 | 0.725         | 1.000 | 0.500   | 0.083      | 0.000     | 0.000               | 0.000         | 0.000                  |        | *Streptophyta sp.*          |
| 1005406 | 5.293            | 0.624 | 0.736         | 1.000 | 0.250   | 0.083      | 0.000     | 0.000               | 0.000         | 0.000                  |        | *Staphylococcus aurues*     |
| 898207  | 5.293            | 0.624 | 0.736         | 1.000 | 0.250   | 0.083      | 0.000     | 0.000               | 0.000         | 0.000                  |        | *Granulicatella sp.*        |
| 336228  | 5.293            | 0.624 | 0.736         | 1.000 | 0.250   | 0.083      | 0.000     | 0.000               | 0.000         | 0.000                  |        | *Dialister sp.*             |
| 4301737 | 19.335           | 0.007 | 0.020         | 1.000 | 0.500   | 0.000      | 0.000     | 13.000              | 0.250         | 0.000                  |        | *Porphyromonas sp.*         |
| 341460  | 35.743           | 0.000 | 0.000         | 0.115 | 180.750 | 0.000      | 29.500    | 0.667               | 40.500        | 0.000                  |        | *Haemophilus sp.*           |
| 530164  | 20.981           | 0.004 | 0.011         | 1.000 | 102.750 | 0.000      | 41.000    | 0.667               | 1.000         | 0.000                  |        | *Porphyromonas sp.*         |

*Sample means are shown.