Understanding the microbial biogeography of ancient human dentitions to guide study design and interpretation

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Abstract

The oral cavity is a heterogeneous environment, varying in factors such as pH, oxygen levels, and salivary flow. These factors affect the microbial community composition and distribution of species in dental plaque, but it is not known how well these patterns are reflected in archaeological dental calculus. In most archaeological studies, a single sample of dental calculus is studied per individual and is assumed to represent the entire oral cavity. However, it is not known if this sampling strategy introduces biases into studies of the ancient oral microbiome. Here, we present the results of a shotgun metagenomic study of a dense sampling of dental calculus from four Chalcolithic individuals from the southeast Iberian peninsula (ca. 4500–5000 BP). Interindividual differences in microbial composition are found to be much larger than intraindividual differences, indicating that a single sample can indeed represent an individual in most cases. However, there are minor spatial patterns in species distribution within the oral cavity that should be taken into account when designing a study or interpreting results. Finally, we show that plant DNA identified in the samples is likely of postmortem origin, demonstrating the importance of including environmental controls or additional lines of biomolecular evidence in dietary interpretations.

Keywords: ancient DNA, microbiome, dental calculus, metagenomics, biogeography, archaeology

Introduction

Dental calculus forms when the dental plaque biofilm mineralizes during life (Jin and Yip 2002), a periodic occurrence that encapsulates microbes, host biomolecules, food residues, and particles from the environment (Velsko and Warinner 2017). After the death of an individual, biomolecules within dental calculus can be preserved for tens of thousands of years (Fellows Yates et al. 2021), largely protected from environmental processes within the mineral matrix. Studies of archaeological dental calculus have rapidly increased in number in recent years, in part due to an elevated interest in the evolution of the oral microbiome and a growing understanding of the plethora of ancient biomolecules and information that can be recovered from this semi-fossilized microbial biofilm. However, there are still many unknown factors regarding the formation and preservation of archaeological dental calculus, and further method development is, therefore, necessary. In carrying out comparative studies of ancient dental calculus, researchers aim to set up a sampling strategy that mitigates biases caused by intraindividual variability of the studied individuals. However, as archaeological dental calculus is often found in small quantities, especially in individuals dating far back in time, and is not always present on the same teeth across individuals, it may not always be possible to adhere to such a sampling scheme. Pre- and post-mortem tooth loss can further complicate sampling designs, as does working with calculus samples that were dislodged from the teeth during handling or storage, such that the precise tooth of origin is unknown. Due to such sampling constraints, some studies have pooled and homogenized calculus from several teeth for analysis (Warinner et al. 2014), which may partly mitigate spatial biases, but this approach requires the presence and collection of larger amounts of calculus, which is a finite archaeological substrate. In light of these challenges, most ancient oral...
microbiome studies implicitly assume that a single sample can be representative of the entire dentition, regardless of the tooth niche from which the calculus sample is obtained, and analyze only a single dental calculus deposit per individual. The oral cavity, however, is not a uniform environment, and thus microbial communities may vary across the dentition, potentially leading to bias when comparing across individuals from whom different teeth were sampled.

Differences in the microbial composition of different oral tissues, such as buccal mucosa, keratinized gingiva, saliva, and teeth, have been reported in present-day humans (Aas et al. 2005, Ding and Schloss 2014, Eren et al. 2014, Mark Welch et al. 2016, Proctor et al. 2018, Utter et al. 2020). Further, differences in dental plaque microbial communities have been previously reported between mandibular and maxillary teeth (Haffajee et al. 2009, Simon-Soro and Tomás 2013), between tooth position (e.g. anterior vs. posterior teeth; Haffajee et al. 2009, Proctor et al. 2018), between tooth surfaces (e.g. buccal vs. lingual; Simon-Soro and Tomás 2013, Proctor et al. 2018), and between supragingival and subgingival plaque (Simon-Soro and Tomás 2013, Eren et al. 2014). Local variations in oral physiological conditions, such as salivary flow rate, salivary composition, oxygen availability, and mechanical abrasion during mastication, may contribute to these subtle spatial microbial differences in dental plaque. However, while such spatial differences have been detected in the microbial composition of dental plaque, it is not known whether these patterns are also reflected in dental calculus. Dental calculus represents a fully matured stage of oral biofilm development that is often disrupted in living individuals practicing oral hygiene, leading to a distinct microbial profile between dental plaque and dental calculus (Velsko et al. 2019, Kazarina et al. 2021). Overall, dental calculus typically contains higher proportions of late colonizer taxa that thrive in the anaerobic environment created as the biofilm matures, and thus its composition may be less spatially variable than developing plaque biofilms, which are more dynamic and subject to periodically disruptive forces such as toothbrushing (Velsko et al. 2019).

However, evaluating intraindividual microbial variation in dental calculus across the dental arcade, and thus determining the degree to which a single sample can represent an individual, is challenging. Dense sampling of calculus is often hindered by missing teeth or a lack of calculus deposits distributed across the entire dental arcade. Consequently, previous studies have attempted to identify microbial spatial patterns across the dentition by instead sampling diverse individual teeth from a large number of individuals (Farrer et al. 2018), but this introduces a number of uncontrolled variables, such as individual differences, different biological and absolute ages of samples, different postmortem conditions, and differing degrees of preservation and degradation, which may introduce biases or otherwise alter the observable spatial patterns. Further, this approach does not allow for comparisons of how much of the variation in the dental calculus microbiome stems from intra- vs. inter-individual differences.

To determine the degree to which tooth selection matters in dental calculus sampling for comparative ancient microbiome studies, we conducted a systematic analysis of microbial spatial variation in four nearly complete human dentitions with low to heavy dental calculus deposits from the Iberian Chalcolithic site of Camino del Molino (ca. 4500–5000 BP). With dense sampling across tooth types (incisor, canine, premolar, and molar) and tooth surfaces (buccal, labial, interproximal, and occlusal), we performed shotgun metagenomic analysis of 87 dental calculus samples. We find that the main source of variation in the oral microbiome is the sampled individual, and therefore, one randomly selected sample can, for most purposes, be used to represent an individual in population-level comparative studies. However, minor intraindividual patterns in community composition, functional potential, and species abundances are detectable with respect to tooth position (anterior vs. posterior), dental calculus deposit size, and tooth surface, although with low effect sizes. Only occlusal calculus, which is uncommon and may indicate injury or physiological dysfunction, considerably differed in composition.

We found that ancient human DNA is randomly distributed across the dentition, and no spatial patterns were observed with respect to postmortem environmental contamination. Finally, we found that ancient grapevine (Vitis vinifera) DNA was present in the dental calculus we analyzed, however, it was also present in mandibular bone, suggesting a postmortem origin.

### Materials and methods

#### Samples

Dental calculus was collected from four Chalcolithic (4500–5000 BP) individuals from the southeastern Iberian archaeological site of Camino del Molino near the city of Caravaca de la Cruz in Murcia, Spain, excavated during a salvage excavation in 2008 (Lomba Maurandi et al. 2009a,b, Haber-Uriarte et al. 2011, Diaz Navarro et al. 2019). The Camino del Molino communal burial is a natural pit with a 7 m diameter circular base and a depth of 4 m (of which only the lower 2 m were used for burial), which was likely covered and sealed by a perishable structure (Lomba Maurandi et al. 2009a). The upper layers of the site were destroyed in the early 20th century as a result of agricultural terracing, but the damage did not extend to the burial deposits. Approximately 1300 human individuals representing a broad demographic profile were buried at the site (Haber-Uriarte et al. 2011). The site was chosen for this study because prior dental calculus research at the site had shown excellent oral microbiome preservation (Ziesemer et al. 2015, Mann et al. 2018), and microfossil studies of the dental calculus had been conducted (Power et al. 2014), and because the large number of individuals excavated from the site made it possible to select suitable individuals with nearly complete dentitions and sufficient dental calculus for this study. The four selected individuals were adults and had dental calculus present on most teeth (Fig. 1; Figure S1, Supporting Information), allowing near comprehensive sampling. Dental notation below follows the FDI World Dental Federation standard (Peck and Peck 1993); molar enamel wear is reported as a Brothwell score from 1 (none) to 7 (obliteration of crown and wear of roots; Brothwell 1972), and dental calculus deposits are graded from 1 (slight) to 4 (gross) according to Dobney and Brothwell (1987).

#### Individual CM55

Individual CM55 (35–39-year-old female) had a complete mandible and a partial, fragmented maxilla, with a total of 22 teeth (Fig. 1). Alveolar bone loss and reactive bone formation was observed throughout the mandibular periodontium, suggesting generalized periodontitis. Gross carious lesions were present in teeth 17, 35, 37, 45, and 47. Molar enamel wear was low (Brothwell stage 2). Dental calculus deposits were grades 1–2 in size, except on left premolars and molars, where they reached grade 4. The excessive calculus accumulation on the left posterior teeth, including on the occlusal surfaces, suggests that this individual had experienced pain on the left side of the mouth and avoided mastication on this side. Although no skeletal trauma...
were analyzed: teeth 16, 18, 45, and 46. Gross carious lesions were present in molars, and healing was incomplete for four molars that had been lost antemortem. Alveolar recession was pronounced around the healed alveolar bone where teeth 37 and 38 had been lost. Alveolar recession was not pronounced. Molar enamel wear was minimal (Brothwell stages 1 and 2), and no gross carious lesions were observed. Dental calculus deposits were grades 1–2 in size. Alveolar recession was slight across the periodontium, and in general the individual exhibited good dental health.

**Individual CM59**

Individual CM59 (25–35-year-old male) had an intact mandible and a partial, fragmented maxilla, with a total of 25 teeth (Fig. 1). Molar enamel wear was minimal (Brothwell stages 1 and 2), and no gross carious lesions were observed. Dental calculus deposits were grades 1–2 in size. A large bone abscess was present adjacent to the healed alveolar bone where teeth 37 and 38 had been lost antemortem. Alveolar recession was pronounced around the molars, and healing was incomplete for four molars that had been lost antemortem. Gross carious lesions were present in teeth 16, 18, 45, and 46.

**Individual CM82**

Individual CM82 (35–45-year-old female) had a complete mandible and a partial, fragmented maxilla, with a total of 23 teeth (Fig. 1). Heavy enamel wear (Brothwell stage 4) was observed on the molar teeth. Dental calculus deposits were grades 1–2 in size. A large bone abscess was present adjacent to the healed alveolar bone where teeth 37 and 38 had been lost antemortem. Alveolar recession was pronounced around the molars, and healing was incomplete for four molars that had been lost antemortem. Gross carious lesions were present in teeth 16, 18, 45, and 46.
eight interproximal sites in CM59 and CM165 yielded sufficient calculus for analysis and were also sampled. In total, 87 calculus samples were selected from the four individuals for metagenomic analysis (Dataset S1A).

**Laboratory methods**

Surface contamination was reduced by UV irradiation (30 s on both sides), followed by a washing step in 1 ml of 0.5 M EDTA (without incubation). DNA was extracted from the calculus and bone samples using a modified version of (Dabney et al. 2013a) adapted for dental calculus (Mann et al. 2018, Aron et al. 2020a) and allowing for potential future protein extraction from the same samples (Fagernäs et al. 2020). Briefly, the samples were decalcified in 1 ml 0.5 M EDTA for 3 days, after which the cell debris pellet and 100 μl of the supernatant was frozen at −20°C and set aside for future analyses (Fagernäs et al. 2020). To the remaining 900 μl supernatant, proteinase K (Sigma-Aldrich) was added, and the samples were incubated at room temperature overnight. The supernatant was then mixed with binding buffer (5 M guanidine hydrochloride, 0.12 M sodium acetate, and 40% isopropanol) and DNA was purified using a High Pure Viral Nucleic Acid kit (Roche Life Science) according to the manufacturer’s instructions. DNA was eluted in Qiagen EB buffer, to which Tween 20 had been added to a final concentration 0.05%. DNA was quantified using a Qubit HS assay (Thermo Fisher Scientific). Extraction blanks (one per batch) were processed alongside the samples. The full extraction protocol is available at (Arton et al. 2020a).

Extracted DNA was processed with a partial uracil–DNA–glycosylase treatment (Rohland et al. 2015, Aron et al. 2020b) and was prepared into double-stranded libraries with dual indexing (Meyer and Kircher 2010, Kircher et al. 2012, Stahl et al. 2019). Library blanks were processed alongside the samples, one per batch. The DNA libraries were shotgun sequenced on an Illumina MiSeq (Thermo Fisher Scientific). Exclusion blanks were processed alongside the samples. The full extraction protocol is available at (Aron et al. 2020a).

The EAGER v.1.92.56 (Peltzer et al. 2016) pipeline was used for preprocessing of the raw data. Adapter removal and merging of reads were performed using AdapterRemoval v.2.3.1 (Schubert et al. 2016). The reads were mapped to the human reference genome HG19 using BWA aln v. 0.7.12 (Li and Durbin 2009) with default settings (-t 32, -m 0.01), and unmapped reads were extracted with SAMtools v.1.3 (Li et al. 2009) for downstream microbiome analyses. The unmapped reads were aligned to a custom RefSeq database (Fellows Yates et al. 2021) using MALT v. 0.4.0 (Herbig et al. 2016) (settings -id 85.0 -top 1 -supp 0.01). This database contains all bacterial and archaeal assemblies at scaffold/chromosome/complete levels (as of November 2018), with max 10 randomly selected genomes per species (prioritizing more complete genomes), as well as the human HG19 reference genome. A preliminary screening for eukaryotic DNA was performed as described above, using the NCBI full nt database (as of October 2017), but the custom RefSeq database was chosen for further analyses, as it has been shown to yield a higher percentage aligned sequences for dental calculus (Fellows Yates et al. 2021). OTU tables with summarized read counts at species and genus level were exported through MEGAN v. 6.17.0 (Huson et al. 2016, Dataset S1B and S1C). The R-package decontam v. 1.6.0 (Davis et al. 2018) was used to identify putative laboratory and environmental contaminants from OTU tables, using the prevalence method with two sets of controls (cutoff 0.8 for each): mandibular bone from the sampled individuals in this study and previously published bone samples from Bronze Age Mongolia (Jeong et al. 2018, Fellows Yates et al. 2021), and laboratory extraction and library preparation blanks.

**Preservation assessment**

A genus-level OTU table was used as input for SourceTracker v. 0.1.1 (Knights et al. 2011). Included were also comparative samples from published shotgun microbiome studies, including 10 nonindustrialized gut samples (Obregon-Tito et al. 2015, Rampelli et al. 2015), 11 industrialized gut samples (Gevaerts et al. 2012, Sankaranarayanan et al. 2015), 10 skin samples (Oh et al. 2016), 11 subgingival and 10 supragingival plaque samples (Gevaerts et al. 2012), 10 archaeological bone samples (Fellows Yates et al. 2021), 10 modern dental calculus samples (Fellows Yates et al. 2021), and 10 archaeological sediment samples (Slon et al. 2017). During the SourceTracker analysis, the samples were rarefied to 10 000 reads, with a training data rarefaction of 5000. A principal component analysis was conducted on summarized genus level read counts of all samples, blanks, and sources (including an additional nine modern dental calculus samples). Multiplicative zero replacement was conducted using the R-package zCompositions v. 1.3.4 (Palarea-Albaladejo and Martín-Fernández 2015) and the data was CLR-transformed (Gloor et al. 2017). The nonhuman DNA sequences were also mapped to the Tannerella forsythia representative genome (strain 9212) using EAGER v 1.92.38 as described above. The output from DamageProfiler v. 0.3.10 (Neukamm et al. 2021) was used to visualize damage curves for the samples, and fragment length was extracted from the output table from EAGER.

**Community composition**

Analyses of community composition were conducted on the MALT taxon tables, where putative contaminants had been removed, following recommendations for compositional data (Gloor et al. 2017). Significant differences in community composition of samples in selected metadata groups were tested using a PERMANOVA with the R-package vegan v. 2.5.6 (Oksanen et al. 2019), using euclidean distance and 9999 permutations, and individuals as strata when needed. A PCA was conducted as described above. Alpha diversity was analyzed using a species-level OTU table, and Shannon Index and Inverse Simpson Index were computed using the R-package microbiome v. 1.8.0 (Lahti and Shetty 2012).

**Differential abundance**

Differential abundance of species was calculated using Songbird v1.0.1 (Morton et al. 2019) (--formula ‘Jawbone+ToothSurface+ToothPosition+DepositMass_scaled+Individual’, –epochs 10 000 and –differential-prior 0.5). Tensorboard v. 1.14.0 was used for model checking. Input was a species-level OTU table, where putative contaminants were removed. Further, taxa present in fewer than three samples per individual were removed, and thereafter taxa absent in one or more of the individuals. This stringent filtering was applied in order to avoid any potential remaining contaminants or mismapping to influence the results. A total of two separate analyses were conducted, one without occlusal samples and one including occlusal samples.
**Functional analysis**

The functional profiles of the microbial communities were extracted from the nonhuman DNA sequences using HUMAnN v. 2.8.0 (Franzosa et al. 2018), using the CocoPhlAn nucleotide database and the UniRef90 protein database. The output was normalized to copies per million, and translated into KEGG orthologies. Gene families were analyzed, without taking into account species assignments, and putative contaminants were removed from the dataset using decontam as described above (threshold 0.5 for both blanks and bones). A PCA was conducted, and drivers of variation identified using PERMANOVA, all as described above for community composition.

**Human reads**

In order to investigate the amount of host human DNA in the samples, while controlling for contaminating human DNA, the raw reads were mapped to the human HG19 genome as described above, with the exception of filtering for mapping quality (-q 37). Duplicates were removed using DeDup v. 0.12.2 (Peltzer et al. 2016), and the reads were filtered for a PMD (postmortem damage) score of 3 using PMDTOols v.0.6 (Skoglund et al. 2014), thereby only retaining damaged ancient reads. This is likely an underrepresentation of the number of ancient reads, as not all DNA fragments will have damage. However, assuming a consistent rate of postmortem damage accumulation over the dental arcade, the bias will be even across all sampling sites, and the patterns of damaged reads can be assumed to also represent patterns of total endogenous human reads. It was noted that occlusal samples generally have a higher percentage damage than other samples, and were therefore, excluded from this analysis, as they break the assumption of equal damage. Deposit mass was accounted for in the analysis, as a positive correlation was found between deposit mass and DNA damage.

**Plant DNA**

During preliminary eukaryotic screening of the dental calculus samples, it was observed that the samples contain a considerable amount of DNA mapping to grapevine (*V. vinifera*). To further explore this pattern, the complete experimental dataset of dental calculus, mandibular bone controls, and blanks were mapped to the grapevine representative genome (GCA_000003745.212X) using EAGER as described above, with mapping quality set to 37. Damage profiles, specifically cytosine to thymine (C–T) transitions typical for ancient DNA, were created using DamageProfiler v.0.3.10 (Neukamm et al. 2021).

**General statistics**

Unless otherwise stated, data was processed in R v. 3.6.1 (R Core Team 2019), using packages tidyverse v.1.3.0 (Wickham et al. 2019), ggpubr v.0.3.0 (Kassambara 2018), readxl v.1.3.1 (Wickham and Bryan 2019), janitor v.2.0.1 (Firke 2018), and ggeffects v.0.14.3 (Lüdecke 2018). In order to investigate patterns across the dentition, linear mixed-effects models (LME) were fitted to the variables in question using lmerTest v.3.1.2 (Kuznetsova et al. 2017) and Box–Cox transformations identified using MASS v.7.3.51.4 (Venables and Ripley 2002). Explanatory variables in all tests are: jawbone (mandible/maxilla), tooth surface (lingual/buccal/interproximal/occlusal), tooth position (anterior/posterior), and mass of the original calculus deposit (scaled and centered continuous variable). Incisors and canines were treated as anterior teeth; premolars and molars were treated as posterior teeth. Unless otherwise noted, occlusal calculus, which was only obtained from a single individual, was excluded from most analyses because these samples were found to break the assumption of homogeneous distribution of variance (euclidean distances, ANOVA, \( P = .001 \)).

**Results**

**Preservation and authentication**

Total DNA yield from a sample, normalized by the mass of the dental calculus sample used for DNA extraction, may vary depending on preservation and organic matter content of the sample, and may bias downstream taxonomic profiles (Fagernäs et al. 2020). Occlusal samples were excluded from this analysis, as it was noted during sampling that their consistency was different from all other samples. Using linear mixed effects modelling, we tested whether tooth surface, tooth position, jawbone, or deposit mass influenced the mass-normalized DNA yield from a sample. We found that none of these factors outperformed the null model (LME, individual as random effect), and therefore, normalized DNA yield cannot be predicted by these variables (Fig. 2A).

Prior to oral microbiome analysis, the archaeological dental calculus in this study was evaluated for preservation and authenticity of the ancient oral microbiome. This is important because poor dental calculus preservation and contamination with environmental microbes can bias or interfere with downstream analyses. A PCA on genus level read counts shows that all the archaeological dental calculus samples cluster together with modern dental plaque samples, and are clearly separated from archaeological bone, gut, and sediment samples (Fig. 2D). To further assess preservation of the dental calculus samples, the contribution of different source environments to the composition of the samples was estimated using SourceTracker (Knights et al. 2011). All samples were estimated to have a majority contribution from oral microbiome sources, indicating good preservation of the oral microbiome (Figure S2, Supporting Information). Some samples were estimated to have a minor contribution from the skin microbiome. Minor estimated contributions from the gut microbiome and sediment are also present, but are expected because gut and oral taxa are similar and can be difficult to distinguish using short read data, and because archaeological samples typically contain some soil contamination, even after washing. After taking these factors into consideration, all dental calculus samples were determined to be sufficiently well-preserved for inclusion in downstream analyses.

We next assessed DNA damage patterns in the dental calculus as an indicator of authenticity. DNA from archaeological samples accumulates specific forms of damage over time, which can be seen as C–T transitions at the ends of DNA fragments and a high degree of DNA fragmentation (Dabney et al. 2013b). We generated a damage plot for fragments mapping to the prevalent oral bacterium *T. forsythia* (Fig. 2B), and all four individuals exhibit damage patterns typical for ancient DNA that has undergone partial UDG-treatment (Röhlund et al. 2015). The fragment length distributions of reads mapping to *T. forsythia* show that most samples have a median length < 50 bp, as is expected for ancient sam-
Figure 2. Preservation assessment of dental calculus samples. (A) normalized DNA yield (in ng DNA per mg calculus) across the dental arcade averaged across individuals. (B) C-T transitions at the 5′ end of DNA fragments aligning to T. forsythia, consistent with ancient DNA. Note that the sharp drop from the first to the second base is due to treatment with uracil–DNA–glycosylase. (C) DNA aligning to T. forsythia has short median fragment lengths, consistent with ancient DNA. (D) PCA on genus level read counts of samples from this study, before removing putative contaminants, as well as published metagenomic datasets; dental calculus from this study forms a cluster overlapping with modern plaque and calculus, indicating good oral microbiome preservation.

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account evenness, and is less influenced by rare species than the Shannon index, indicating that the generally large number of rare species in archaeological dental calculus may erode any spatial patterns in alpha diversity.

**Differential taxonomic abundance**

Due to different local environmental conditions in different areas of the oral cavity, small differences in microbial composition have been reported across the dentition in present-day dental plaque (Haffajee et al. 2009, Simon-Soro and Tomás 2013, Proctor et al. 2018). It is, however, not known if such patterns can be detected in archaeological samples, after both biofilm maturation during life and postmortem degradation over time. Here, we find that there are slight taxonomic differences with respect to tooth surface and initial deposit mass. First, we observe differences in taxa between anterior (incisors and canines) and posterior (premolars and molars) teeth, where the taxa that are more abundant in the anterior teeth are more often aerobic or facultatively anaerobic, while the taxa that are most associated with posterior teeth are anaerobic (Fig. 4A). Second, interproximal spaces seem to be enriched in species belonging to the genera *Methanobrevibacter* and *Olsenella* (Fig. 4B), which are both acid tolerant anaerobes. Finally, the species *Actinobaculum* sp. oral taxon 183 and *Fusobacterium* sp. oral taxon 203 are found at a higher abundance in low mass dental calculus deposits, as compared to high mass deposits (Fig. 4C). However, little is known about their physiology or the role they play in the dental plaque biofilm. Fusobacteria are generally secondary colonizers in the dental plaque biofilm, and bind to several other bacterial taxa (Kolenbrander 1988). A 2D model showing the spatial distributions of the taxa in Fig. 4A–C across the dentition can be found at [https://tinyurl.com/eyjcs674](https://tinyurl.com/eyjcs674).

**Functional profile**

In addition to their taxonomic composition, microbial communities may also differ in their gene content, and therefore functional potential, and this can have important implications for understanding oral microbiome evolution (Fellows Yates et al. 2021). To evaluate whether there are potential functional differences across the dental arcade, we analyzed the genes present in the dental calculus metagenomes. In total, 2791 gene families were identified in the dataset, after removing putative contaminants that were identified from blanks and bone samples. The individual was found to be the strongest driver of variation (PERMANOVA, $P = .001$ and $R^2 = 0.12$; Fig. 5A), and after accounting for this, tooth surface ($P = .020$, $R^2 = 0.049$), tooth position ($P = .039$, $R^2 = 0.029$), and deposit mass ($P = .013$, $R^2 = 0.036$) were found to significantly drive functional variation (PERMANOVA, individual as strata, Fig. 5B). However, these factors explain only a very minor part of the variation, as can be seen by the low $R^2$ values. It should also be noted that the tooth surface variable breaks the assumption of homogeneity of variance for this analysis, which may affect the results of the PERMANOVA.

**Human genetic content**

Human DNA from dental calculus is mainly derived from a single source—the host (Ozga et al. 2016). Although human dental calculus generally contains a very low proportion of human DNA (Mann et al. 2018), different enrichment approaches have been used to increase the human DNA fraction enough to study the human genome (Ozga et al. 2016, Ziesemer et al. 2019). Human DNA may in theory be differentially incorporated into dental calculus across the dental arcade, depending on salivary flow, inflammation, or disease, among other factors. We investigated the presence and relative abundance of ancient human DNA in our samples to assess potential spatial patterning of human host DNA in calculus. To focus our analysis on host ancient DNA, we restricted our analysis to only DNA fragments with C–T DNA damage. As a slight positive correlation was found between deposit mass and damage (Figure S4, Supporting Information), deposit mass was accounted for in this analysis. We found that the best fitting model for predicting the proportion of human reads in the dental calculus samples is a null model, indicating that the distribution of human DNA in dental calculus does not significantly vary according to tooth surface, tooth position, or jawbone (LME, deposit mass as random effect; Fig. 6A).

**Postmortem environmental colonization**

Whether contamination by infiltration of environmental microbes from the burial context is introduced in a nonrandom way across the oral cavity is not known. Because different properties of cal-
Figure 4. Differential abundance of species across the dental arcade. (A) Species associated with posterior (premolars and molars) vs. anterior (incisors and canines) teeth. (B) Species associated with interproximal spaces vs. all other tooth sites. (C) Species associated with high vs. low initial deposit mass. Only the top 10 taxa most associated with each factor are shown.

Figure 5. Functional profile of dental calculus samples. (A) PCA of gene families, normalized to copies per million, with colour indicating individual. (B) Same data as (A), but with shape fill indicating tooth position, shape representing tooth surface, and samples coloured by initial deposit mass (scaled variable).

Occlusal calculus

The occlusal dental calculus analyzed in this study differed from the calculus from other tooth surfaces in several ways, and was therefore, excluded from most analyses. During sampling, occlusal calculus was found to have a different consistency from the other calculus, being less dense and having less structural integrity. Occlusal calculus was found to have a higher amount of DNA damage than other calculus. For reads mapping to T. forsythia, a model including tooth surface and deposit mass best predicted damage at the first base at the 5' end of the fragment (LME, individual as random effect, $P = .018$), with occlusal samples having higher levels of damage than other samples (Figure S4, Supporting Information). Further, occlusal calculus samples broke the assumption of homogeneity of dispersion for the community composition, which may be due to the fact...
that they were only collected from a single individual, and from only posterior teeth on the same side of the mouth. Overall, we found that despite forming on posterior teeth, occlusal calculus samples are somewhat enriched in aerotolerant species, possibly due to their more exposed location on the tooth, compared to the lingual and buccal surfaces of the posterior teeth that directly abut the tongue and buccal mucosa, respectively (Fig. 7).

Plant DNA

Ancient dental calculus is a potentially valuable source of information about ancient diets, as it is possible to directly study diet-related biomolecules and microfossils incorporated in the calculus during an individual’s lifetime. Researchers have previously attempted to identify dietary sources using DNA from dental calculus (Warinner et al. 2014, Weyrich et al. 2017), an approach that also has many difficulties due to the exceptionally low number of dietary DNA sequences typically found in dental calculus (Mann et al. 2020). The dental calculus samples in this study contained trace amounts of plant DNA (between 170 and 1578 reads per individual, or 0.002%–0.011% of total reads) that map to the genome of grapevine (V. vinifera), a currently and historically widely cultivated plant in the region. From the present data, however, it is not possible to determine if this DNA originated from domesticated V. vinifera or one of its less studied wild relatives. However, it was noticed that similar numbers of grapevine reads (205–2119 reads, or 0.003%–0.034%) were also recovered in the mandibular bone control samples (Fig. 8A). Both sets of reads were found to have C–T damage typical of ancient DNA (7%–9% for bones and 3%–14% for calculus; Fig. 8B), but at lower levels than observed for the oral bacterium T. forsythia (11%–21%; Fig. 2B). The presence of grapevine reads in both dental calculus and bone, together with the lower amount of damage, suggests a likely postmortem origin of the grapevine DNA. However, a dietary origin of the grapevine DNA cannot completely be excluded, as a wild variety has been documented in the region since the Palaeolithic (Aura et al. 2005, Iriarte-Chiapusso et al. 2017).

Discussion

A potentially uneven distribution of microbes in microbiomes can cause biases in downstream analyses if spatial variation is not taken into account during sampling design and data interpretation. Archaeological dental calculus provides a valuable window into the evolution of the oral microbiome, but to date it has not been clear to what degree microbial taxa are spatially patterned across the dentition, and thus, to what degree sampling strategy might impact comparative studies of dental calculus microbial communities. The results of present-day dental plaque studies cannot be directly applied to dental calculus because the two substrates reflect different levels of biofilm maturity and have slightly different composition (Velsko et al. 2019), and in previous studies of spatial variation in archaeological dental calculus, which sampled diverse individual teeth from a large number of individuals (Farrer et al. 2018), potentially confounding factors such as individual, temporal, environmental, and taphonomic differences were not controlled for. Here, we have presented a systematic study of intra-individual variation in archaeological dental calculus by focusing on intensive, comprehensive sampling of the dentitions of four contemporaneous individuals from the same burial context.

Overall, we find that although there are small differences in the spatial distribution of anaerobic and aerotolerant taxa, as well as minor associations between taxonomic composition and initial calculus deposit size, these factors account for very little of the overall microbial and functional variation within dental calculus. Spatial patterns in the oral microbiome that have been identified in studies of modern dental plaque (Haffajee et al. 2009, Simon-Soro and Tomás 2013, Proctor et al. 2018) are not obvious in this study. Such patterns may have been present during life but were subsequently lost over time due to taphonomy, or these patterns may simply not be present in calculus. Although taphonomic processes, such as C–T damage accumulation and DNA fragmentation, as well as postmortem colonization of the body by environmental taxa, may obscure oral microbiome spatial patterns, we did not find these factors to correlate with the microbial patterns we observed. A study of modern dental calculus that investigates species spatial patterning will be needed to determine if the patterns observed in dental plaque are maintained as the biofilm matures and calcifies into dental calculus.

Although this study investigated a small number of individuals from a single archaeological site, the purpose of this study design was to limit the number of potentially confounding factors, such as different sample ages, different burial conditions, and different storage and handling practices after excavation. Microbial spatial patterning may differ in other populations or at other archaeological sites, and this warrants further investigation. However, as the species profiles of human dental calculus appear to be more consistent across time, space, and health status than dental plaque (Velsko et al. 2019, Fellows Yates et al. 2021), it is possible that any variation will be very minor.

Although we observed few spatial patterns in archaeological dental calculus, we find that occlusal calculus may represent a special exception. Dental calculus rarely accumulates on the occlusal surfaces of teeth, in part due to the abrasive forces of mastication, and large deposits of occlusal calculus are generally indicative of physiological injury or dysfunction. Here, only one individual had occlusal calculus, but this calculus had a distinct tex-
Differential abundance of taxa in occlusal samples compared to other tooth surfaces. Only the top 10 taxa are shown, and the bars are coloured by aerotolerance of the taxa.

Presence of grapevine DNA in bones and dental calculus samples. (A) The percentage of reads that aligned to the grapevine genome per sample. (B) C–T miscoding lesions at the 5′ end of the read, for each sample with > 500 reads aligning to grapevine.

Figure 7

Figure 8

Contrary, higher DNA damage, and different levels of taxonomic dispersion compared to other dental calculus in the study, even from the same individual. Although further research on a larger number of individuals is necessary, occlusal calculus is likely not representative of oral microbiome communities, and therefore, should be avoided in comparative studies of microbial variation across individuals. In addition to occlusal samples having an increased amount of DNA damage, an association was also found between deposit mass and DNA damage for other sampling sites. This pattern may be related to a greater immune response at sites with large dental calculus deposits, or possibly due to the lower density that is common for large deposits, but more research is necessary in order to understand this association.

Beyond microbes, dental calculus is also valuable because it entraps dietary and other environmental debris during life, and thus can provide clues about the foods and activities of past societies (Hardy et al. 2009, Leonard et al. 2015, Power et al. 2015, Radini et al. 2017). Although dietary proteins have been shown to preserve within dental calculus (Hendy et al. 2018, Wilkin et al. 2020, Scott et al. 2021), the metagenomic recovery of dietary DNA from calculus has yielded more equivocal results (Mann et al. 2020). The recovery and authentication of eukaryotic DNA in metagenomic datasets is not trivial due to complicating factors such as the very low number of nonhost eukaryotic DNA fragments typically found in dental calculus and the problem of microbial contamination in eukaryotic reference genomes, which can lead to false positives (Mann et al. 2020). Here, we show that other unknown postmortem processes may also introduce eukaryotic DNA, for example through mortuary practices or by contamination from the local environment, either recently or in the distant past. It is, therefore, advisable to include environmental controls, such as bone or sediment samples, in metagenomic studies of diet. Another authentication aid may be the use of complementary dietary identification methods, such as microfossil analysis or palaeoproteomics. Through proteomic analyses, for example, it is possible to deduce the part of an organism from which the biomolecules originate, such as seed proteins from plant seeds, or milk proteins from dairy products. Combining methods may thus aid researchers in establishing the plausibility of a given organism being incorporated into dental calculus as a food as opposed to environmental contamination.

To conclude, we find that in most applications a single sample of archaeological dental calculus can be used to represent an individual in comparative studies of the ancient oral microbiome, as the main source of variation generally is the sampled individual. The use of a single sample instead of multiple samples, either pooled or studied separately, reduces the destructive demands on this finite archaeological material. However, as there are minor spatial patterns present, care should be taken to record the sampling location within the oral cavity for each dental calculus sample, whenever possible. This makes it possible to later reevaluate findings if systematic biases are suspected. Occlusal calculus is found to differ from other sampling locations and should, thereby, not directly be compared to other sampling sites. Further, includ-
ing environmental controls or multiple lines of evidence is highly recommendable in studies of diet from archaeological dental calculus. This study contributes to an awareness of spatial variation in dental calculus microbial community composition that aims to aid researchers in developing robust study designs and valid interpretations for ancient oral microbiome studies.

**Supplementary data**

Supplementary data are available at FEMSMC online.

**Accession numbers**

Genetic data have been deposited in the ENA under the accession PRJE46022. R Markdown files and illustration files have been archived with Zenodo (DOI: 10.5281/zenodo.6034699).

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