Do dental nonmetric traits actually work as proxies for neutral genomic data? Some answers from continental- and global-level analyses

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Abstract

Objectives: Crown and root traits, like those in the Arizona State University Dental Anthropology System (ASUDAS), are seemingly useful as genetic proxies. However, recent studies report mixed results concerning their heritability, and ability to assess variation to the level of genomic data. The aim is to test further if such traits can approximate genetic relatedness, among continental and global samples.

Materials and Methods: First, for 12 African populations, Mantel correlations were calculated between mean measure of divergence (MMD) distances from up to 36 ASUDAS traits, and $F_{ST}$ distances from >350,000 single nucleotide polymorphisms (SNPs) among matched dental and genetic samples. Second, among 32 global samples, MMD and $F_{ST}$ distances were again compared. Correlations were also calculated between them and inter-sample geographic distances to further evaluate correspondence.

Results: A close ASUDAS/SNP association, based on MMD and $F_{ST}$ correlations, is evident, with $r_m$ values between .72 globally and .84 in Africa. The same is true concerning their association with geographic distances, from .68 for a 36-trait African MMD to .77 for $F_{ST}$ globally; one exception is $F_{ST}$ and African geographic distances, $r_m$ = 0.49. Partial MMD/$F_{ST}$ correlations controlling for geographic distances are strong for Africa (.78) and moderate globally (.4).

Discussion: Relative to prior studies, MMD/$F_{ST}$ correlations imply greater dental and genetic correspondence; for studies allowing direct comparison, the present correlations are markedly stronger. The implication is that ASUDAS traits are reliable proxies for genetic data—a positive conclusion, meaning they can be used with or instead of genomic markers when the latter are unavailable.

KEYWORDS

Africa, Arizona State University Dental Anthropology System, geographic distance, population affinities, single nucleotide polymorphisms
Nonmetric traits of the human permanent dentition, as in those from the Arizona State University Dental Anthropology System (ASUDAS), have a significant genetic component in expression. At least this was suggested in earlier research (e.g., Sofaer, Niswander, MacLean, & Workman, 1972; Scott, 1973; Brewer-Carias, Le Blanc, & Neel, 1976; Scott, Yap Potter, Noss, Dahlberg, & Dahlberg, 1983; Turner II, 1985a; Sofaer, Smith, & Kaye, 1986; Scott & Turner II, 1988, 1997; Turner II, Nichol, & Scott, 1991; Pinkerton, Townsend, Richards, Schwerdt, & Dempsey, 1999; Rightmire, 1999; although see Harris, 1977). Therefore, phenetic affinities based on these traits approximate genetic relatedness, at levels of comparison ranging from local to global. At least this was assumed in previous work (Irish, 1997; Scott et al., 1983; Scott & Turner II, 1988, 1997; Turner II, 1985a; Turner II et al., 1991). These and other attributes, such as their ease of recording, observer replicability—especially if dichotomized (below), evolutionarily conservative nature, lack of sexual dimorphism, and a low likelihood of selection in expression (Scott & Irish, 2017; Scott & Turner II, 1997; Scott, Turner II, Townsend, & Martinón-Torres, 2018; Turner II et al., 1991), are part of a common refrain in ASUDAS studies, including most cited here. Recently, however, the degree of genetic contribution and concordance of dental and genetic data, in particular, have come under renewed scrutiny—several examples of which are summarized. Before proceeding, it of course goes without saying that non-ASUDAS traits are also used in dental research (e.g., Bailey & Hublin, 2013; Martinón-Torres et al., 2007); however, because few have been universally accepted or formally tested (below), such traits are not considered further.

Concerning genetic input, narrow-sense heritability ($h^2$) estimates in samples of Australian twins for several ASUDAS molar traits surpass 0.60, with UM1 Carabelli’s reaching 0.80 and the UM1 and UM2 hypocone 0.87 and 0.93, respectively (Higgins, Hughes, James, & Townsend, 2009; Hughes & Townsend, 2011, 2013; Hughes, Townsend, & Bockmann, 2016). Yet, analyses of an African American Gullah sample (Stojanowski, Paul, Seidel, Duncan, & Guatelli-Steinberg, 2018, 2019) yielded some estimates that are more modest. For eight distinct ASUDAS crown traits recorded across incisor, canine, and premolar fields and between antimeres and isomerines, the average $h^2$ for traits with significant $p$-values is $\geq 0.33$. The range is $0.00–0.82$, depending on whether the rank-scale traits were treated as continuous data or dichotomized using alternate breakpoints (defined in Turner II et al., 1991; Scott & Turner II, 1997; Scott & Irish, 2017; Scott, Turner II, et al., 2018; below). In the same Gullah sample, Stojanowski et al. (2019) then looked at 14 distinct ASUDAS premolar and molar traits in maxillary and mandibular dental fields. Heritabilities are nearer the Australian findings, as UM1 Carabelli’s reached 0.85 and, on average, all statistically significant trait estimates are higher than for Gullah anterior teeth. Still, several premolar and molar values are unexpectedly low. The practice of pooling sexes, in this sample at any rate, was also questioned contra one of the abovementioned ASUDAS attributes. This $h^2$ range is $0.00–1.00$, depending on if the traits were treated as continuous or dichotomized; the latter provided much higher estimates. Socioeconomic stress and reproductive isolation in the Gullah, along with small samples, are acknowledged that may account for the low to moderate $h^2$ estimates (Stojanowski et al., 2018, 2019). In support, prior research also suggests stress affects expression of certain crown traits (Riga, Belcastro, & Moggi-Cecchi, 2014). Lastly, several of the above authors contributed to another study analyzing Australian twins (Paul, Stojanowski, Hughes, Brook, & Townsend, 2020). Results for permanent teeth reflect those of the Gullah studies, but with higher $h^2$ estimates (mostly 0.4–0.8), greater trait heritability when appropriately dichotomized, and less concern for sexual dimorphism in expression. Nevertheless, the above results offer mixed signals about genetic input to expression across a range of ASUDAS traits.

With regard to correspondence of dental and neutral genetic data in appraising relatedness, four recent studies are referenced. First, in a model-free analysis of four samples of living Kenyans ($n = 295$ individuals), correlations were calculated between pseudo-Mahalanobis D$^2$ distances based on nine ASUDAS crown traits, and delta-mu squared distances from 42 short tandem repeats (STRs) (Hubbard, 2012; Hubbard, Guatelli-Steinberg, & Irish, 2015). To date, this is the only research to compare phenotypic and genomic data in the same individuals. A moderate to strong (per Cohen, 1988) positive, though nonsignificant correlation, .50, resulted from Mantel ($p = 0.21$) and bivariate Pearson tests ($p = 0.31$) (Hubbard, Guatelli-Steinberg, & Irish, 2015). Second, 19 dental and up to 19 genetic global samples were matched for comparison by population or similar provenience and ethno-linguistic affinity (Rathmann et al., 2017). Most are recent—although the dates are not listed, two samples are from medieval to Victorian times, and one includes prehistoric material (>2000 BP). The model-bound R-matrix method, initially derived to compare allele frequencies, was used to produce pairwise population kinships from both dental (methods in Relethford, 1991; Konigsberg, 2006) and genomic data, all of which had been published previously: 12 ASUDAS crown traits (19 samples, 1872 individuals), 28 crown measurements (19 samples, 1,016 inds), 645 STR loci (13 samples, 265 inds), and 1,778 SNPs, that is, single-nucleotide polymorphisms (19 samples, 1,652 inds). Focusing just on ASUDAS results, Mantel tests yielded higher correlations than Hubbard et al. (2015), with an $r_{max}$-value for STRs in 13 samples of .55 (10,000 random permutations, $p < .001$) and for SNPs in 19 samples, .64 ($p < .001$). The dental-STR correlation is strong (per Cohen, 1988), prompting Rathmann et al. (2017:3) to suggest dental data may be used as genetic proxies, although they "reason that a substantial portion of the variation can be explained by natural selection on dental morphology." Indeed, the presence and expression of several traits have been linked with selection (Bryk et al., 2008; Hlusko et al., 2018; Kimura et al., 2009; Park et al., 2012); this idea challenges yet another perceived attribute of the ASUDAS (see Scott & Irish, 2017; Turner II et al., 1991). Third, somewhat tangentially, Delgado et al. (2019) calculated phenetic distances from 16 ASUDAS traits in 477 living Colombians to those in dental samples of Europeans, Native Americans, and Africans (Irish, 1993, 1997; Scott & Turner II, 1997); these three are said to represent principal ancestors of admixed Latin Americans. The same Colombians had also been
genotyped, and 93,328 SNPs were compared with samples of the same populations to quantify admixture proportions. Dental-based affinities revealed the closest link with Europeans, and average ancestry estimates based on genetic and dental data generally concur. However, dental traits were not useful in assessing individual ancestries (Delgado et al., 2019). Lastly, to again explore whether dental and genetic data return similar information on admixture, Gross and Edgar (2019) employed Fisher Information in samples from West Africa, Europe, and North America. A model-bound clustering method, for multi-locus genotype data in the program STRUCTURE, was then applied to consider correspondence in estimating ancestry of individuals, that is, African, European/European American, and African American, with 53 unspecified crown traits (797 inds), up to 992,601 SNPs (271 inds), and 645 STRs (177 inds). Like most all dental studies many trait data are missing, which was suggested to affect the performance. Still, SNPs, followed by STRs delivered superior results in “detecting differences in admixture proportions between individuals within admixed populations;” dental data were, however, deemed to be useful to investigate population-level variation (Gross & Edgar, 2019:528). As above, outcomes of these four studies offer mixed support for ASUDAS traits, in this case pertaining to use as genetic proxies for analyses of populations and, in the latter two cases, individuals.

Today, it is patent that neutral genomic markers are the definitive choice in population (and individual) studies, and the standard to which all phenotypic data are and should be compared (see Rathmann et al., 2017). That said, on the above bases expression of the latter is minimally heritable for some, while analyses based on 9, 12, 16 and 53 ASUDAS traits failed to account fully for variation among population samples and/or individuals. Two other long held attributes—lack of sexual dimorphism and minimal selection—were also called into question. Nonetheless, given its long successful run, the aim here is to give the ASUDAS another chance to demonstrate its once-possited potential, through enhanced comparative analyses at both continental and global levels; heritability is not considered directly, but rather the capacity for these traits to approximate genetic relatedness.

All of these recent studies provide a foundation on which to build. The global approach of Rathmann et al. (2017), specifically, is used as a baseline. However, the first author’s (JDI) African and other recent ASUDAS data (>100 samples, >6,000 individuals), and vast CG Turner II database (>300 samples, >23,000 inds; parts of which are presented in Scott & Irish, 2017) accessed here offer additional comparative choices. As such, many more dental samples could be matched with their genetic counterparts by population at a level of concordance not possible before (below). So, in 12 populations across the continent of Africa, correlations were calculated between the matrices of phenetic distances based on 36 and 25 ASUDAS traits, respectively, and genetic distances from >350,000 SNPs. The latter were selected over STRs because (a) published SNP data are available for more global populations, (b) they afford superior differentiation among such populations, and (c) these markers seemingly correspond more closely with dental nonmetric data (Gross & Edgar, 2019; Rathmann et al., 2017). Matrices calculated from the same dental traits and SNPs were then tested for correlation in an expanded analysis of 32 total global populations. Finally, it is assumed genetic and, by extension (as above), phenetic distances among populations increase exponentially as geographic distances increase (Wright, 1943; Relethford, 2004); thus, correlations between the latter distances among samples and those from dental and genetic data were calculated to explore the influence of geographic structure (e.g., extreme isolation) on the two datasets.

2 | MATERIALS AND METHODS

Given the large dataset and familiarity of JDI with the post-Pleistocene peopling of Africa, the latter was the clear choice for continental-level analyses; dental and genetic samples of three North and nine sub-Saharan African populations were compared. Other than one dental sample (Riet River San, Table 1) with a few earlier historic specimens, all data were recorded in recent, 19-20th century crania and hardstone casts to match close as possible the existing genomic data from living individuals (Table 2). Next, combined with Africans for global analyses were 20 dental samples from Europe, Asia, Australia, Melanesia, and the Americas. With two earlier historic exceptions (Table 1), the latter comprise similarly recent specimens to compare with 20 matched genetic samples (Table 2; Figure 1). In all following tables and figures, samples are abbreviated with a prefix of “D” (dental) or “G” (genetic), followed by sample number (1–32), and three letters for the name. For instance, the Bedouin dental sample D1_BED (Table 1) corresponds with genetic sample G1_MOR from Morocco, and so forth (Table 2). An extensive anthropological literature review facilitated sample matching based on (a) shared language and ethnic groups (e.g., Turner II, 1985a; Scott & Turner II, 1997; Irish, 1993, 1997, 2000; Irish et al., 2014; Irish, 2016; see below), and (b) similar geographic locations. The average distance in km between the matched African dental and genetic samples, as determined from latitude and longitude coordinates (Tables 1 and 2), is 286.6 with a low of 64.2 and high of 505.1. The mean for all 32 samples is 347.9 km with a range, excluding the African low, of 89.6 to 1,361.3 km. The latter is the distance between the dental and genetic Aleut samples—one from the American and the other from the Russian side of the island chain. These and the two Pima samples separated by 581.9 km were included to provide some level of New World coverage. That is, recent Native North American dental data are ample in Turner’s database, but matching genetic data are not (Reich et al., 2012; Skoglund et al., 2015), with the reverse true for recent dental and genetic data in Meso- and South Americans.

Thirty-six crown, root, and intra-oral osseous nonmetric traits (refer to list in Table 3) used in previous affinity studies (Irish, 1993, 1997, 1998a, 1998b, 2000, 2005, 2006, 2010, 2016; Irish et al., 2014; Irish et al., 2017) were initially compared with the SNP data for the African analysis. Beyond the abovementioned ASUDAS attributes, the rationale for choosing, and the standard approach in recording these specific traits are detailed in the preceding references and elsewhere (Scott & Irish, 2017; Scott & Turner II, 1997; Scott,
| Dental sample Abbreviation | Region | Country/Area | Data source | n | Lat  | Lon  |
|---------------------------|--------|--------------|-------------|---|------|------|
| Africa                    |        |              |             |   |      |      |
| Bedouin (Arab)            | D1_BED | North Africa | Morocco and Algeria | Irish, 1993, 1998a | 49 | 34.8 | −5.2 |
| Kabyle (Berber)           | D2_KAB | North Africa | Algeria      | Irish, 1993, 1998a | 32 | 36.6 | 3.7  |
| Kikuyu                    | D3_KKU | East Africa  | Kenya        | Irish unpublished data | 60 | −0.3 | 36.1 |
| Riet River (San; >12-19th Cent) | D4_RRI | South Africa | South Africa | Irish, Black, Sealy, & Ackermann, 2014 | 66 | −29.3 | 24.8 |
| San                       | D5_SAN | South Africa | Botswana, South Africa | Irish, 1993, 1997 | 99 | −22.4 | 24.6 |
| Senegambia (Wolof)        | D6_SEN | West Africa  | Senegambia   | Irish, 1993, 1997 | 42 | 15.2 | −16.7 |
| Shawia (Berber)           | D7_SHA | North Africa | Algeria      | Irish, 1993, 1998a | 26 | 35.4 | 6.7  |
| Somalia                   | D8_SOM | East Africa  | Somalia      | Irish, 2010        | 77 | 9.0  | 46.4 |
| Sotho                     | D9_SOT | South Africa | South Africa | Irish, 2016        | 66 | −29.4 | 28.3 |
| Tswana                    | D10_TSW| South Africa | South Africa | Irish, 2016        | 63 | −25.8 | 23.0 |
| Yoruba                    | D11_YOR| West Africa  | Benin (Dahomey) | Irish unpublished data | 28 | 6.6  | 2.6  |
| Zulu                      | D12_ZUL| South Africa | South Africa | Irish, 2016        | 67 | −28.0 | 32.4 |
| Total                     |        |              |              | 675 |      |      |
| America, Asia, Australia, Melanesia, Europe |        |              |              |     |      |      |
| Pima 94                   | D13_PIM| North America | Salt River–Maricopa, Arizona | Turner unpublished data | 165 | 33.3 | −111.5 |
| Aleut (Western US)        | D14_ALE| North America | Attu, Atka plus Western Aleut Historic | Turner unpublished data; Scott & Irish, 2017 | 95 | 52.0 | −174.0 |
| Kazak 94 (17-19th Cent)² | D15_KAZ| Central Asia  | East Kazakhstan | Turner unpublished data | 204 | 47.0 | 76.0 |
| Mongol 2 and 3 Pooled     | D16_MON| Central Asia  | Northeast Mongolia | Turner unpublished data; Turner 1990 | 82 | 48.0 | 110.0 |
| Lower Ob Khanty           | D17_LOK| Central Asia  | Khant-Mansi (Ugrian), Central Russia | Turner unpublished data | 49 | 63.0 | 70.0 |
| Chukchi plus Eastern Siberia | D18_CHU| Central Asia  | Northeast Russia | Turner unpublished data | 126 | 67.5 | 170.0 |
| Recent Thailand           | D19_THA| East Asia      | Central Thailand | Turner unpublished data; Turner 1990 | 189 | 13.0 | 101.0 |
| Recent Tonkin, Historic Annam | D20_VIE| East Asia      | North Vietnam | Turner unpublished data | 76 | 20.0 | 107.0 |
| Recent Japanese           | D21_JAP| East Asia      | Central Japan | Turner unpublished data; Scott & Irish, 2017 | 131 | 36.0 | 138.0 |
| Malay Composite           | D22MAL | Southeast Asia | Central Malaysia | Turner unpublished data; Scott & Irish, 2017 | 58 | 1.0  | 102.5 |
| Philippines no 2 Calatagan BP | D23_PHI| Southeast Asia | Central Philippines | Turner unpublished data; Scott & Irish, 2017 | 58 | 12.3 | 122.0 |
| Borneo 94                 | D24_BOR| Southeast Asia | Central Borneo | Turner unpublished data; Scott & Irish, 2017 | 144 | 1.5  | 114.5 |
| Australia–North BP        | D25_AUN| Australia      | Northeast Australia | Turner unpublished data; Scott & Irish, 2017 | 57 | −20.8 | 139.5 |
| New Britain 1_4 738 BP, no 3 | D26_NBR| Melanesia      | New Britain | Turner unpublished data; Scott & Irish, 2017 | 238 | −6.0 | 150.0 |
| Nepal 94 BP               | D27_NEP| South Asia     | Central Nepal | Turner unpublished data | 97 | 28.0 | 84.0 |
| Greek Recent              | D28_GRK| South Europe   | South Greece  | Irish, Lillios, Waterman, & Silva, 2017 | 70 | 37.5 | 22.3 |
| Italy Modern              | D29_ITY| South Europe   | Central Italy | Irish et al., 2017 | 55 | 42.0 | 14.0 |
Turner II, et al., 2018; Turner II et al., 1991). As well, these traits were found to be largely independent of one another (Nichol, 1990), a key factor in avoiding data redundancy that adds little additional discriminatory value in biodistance analyses (see below); independence of these 36 traits was subsequently supported with a range of pairwise Kendall's tau-b correlations among the rank-scale data in 1625 Africans of just |0.00–0.33| (Irish, 1993). A succession of later studies did identify some strong correlations, $\tau_b \geq 0.5$, but the lack of patterning suggests the affected trait pairs vary across populations (Irish, 2005, 2006, 2010, 2016; Irish et al., 2014). Next, as mentioned, the 12 dental and 12 genetic samples were compared again after dropping 11 traits: UI1 labial curvature, palatine torus, UC distal accessory ridge, UI2 peg-reduced, UI1 midline diastema, LM1 anterior fovea, mandibular torus, rocker jaw, LM1 deflecting wrinkle, LM1 C1-C2 crest, and LM2 torsomolar angle. This reduction allowed direct comparison of the Africans with 20 additional global samples (Tables 1 and 2), 18 of which are in Turner’s dental database; he regularly recorded just 25 traits (e.g., Turner II, 1985a; and below). Any dissimilarities in the African results relating to different dental trait numbers were then quantified.

All SNP data, except whole genome sequences of the West African Wolof sample (Table 2), were genotype with the Affymetrix Human Origins Array (AHOA) (in Patterson et al., 2012; Pickrell & Pritchard, 2012; Lazaridis et al., 2014; Pickrell et al., 2014; Skoglund et al., 2016; or by request from these authors via signed letters). These data are high density, that is, 593,124 SNPs, and ascertained for all modern populations. Specifically, because the AHOA was built from 13 different global samples, ascertainment bias (i.e., systematic distortion of true allele frequencies) is limited, to reliably represent demographic history. The five low coverage Wolof sequences (9X) were produced by the Gambian Genome Variation Project. To merge them with the AHOA dataset, all reads were mapped against the reference genome (Hg19/GRC37, 1,000 Genome release) by the second author (AM), using the Burrow-Wheeler Aligner-MEM with $-M$ option (BWA-MEM) (Li & Durbin, 2009). Any duplicates reads were removed with markdup, in Samtools Release 1.9 (http://www.htslib.org/) (Li et al., 2009). The genotypes were called only on SNPs in the AHOA. Samtools 1.9 (Li et al., 2009) mpileup was used to generate genotype likelihoods for each SNP, and post-genotype filtering was employed to remove bases with a phred score (base quality) of <30, reads with a mapping quality of <30 (probability of correctly mapping a read >.999), and for sequences with mismatches of >50% (when this percentage of the read’s bases differ from the reference). Then, genotypes were called using bcftools default method (–m option) (Danecek, Schaffels, & Durbin, 2014), omitting insertions-deletions (indels). Filtered out with the bcftools filter were all variants having a calling quality of <20% and depth of $>18$—where a general rule for maximum coverage depth is to filter position depth as $DP > 2*DP$ (for the Wolof genomes $DP = 59$). SNPs (n = 144) that did not match Human Origins allele codes (i.e., a “new” allele was discerned in a sequence) or had minor allele frequencies (<.05) were removed. This progression yielded a final total of 353,091 SNPs for the African and global comparative analyses.

At both geographic levels a model-free approach (Hubbard et al., 2015), standard in most phenotype affinity studies, was conducted (though see below). For example the R-matrix method, to estimate between-population kinship coefficients (Rathmann et al., 2017), may not be the best suited for SNP data—particularly the large numbers, and correlation of results (below) based on dental and genomic data is unlikely to be dependent on the distance measures (Relethford, personal communication, 2019). An advantage of the R-matrix method is that it can correct for genetic drift among samples, but this weighting procedure is only possible when effective population sizes are known (Leigh, Relethford, Park, & Konigsberg, 2003; Relethford & Crawford, 1995), a difficult proposition with premodern peoples (Irish, 2016). So, to evaluate correspondence, common field-specific measures of divergence based on ASUDAS and SNP data among respective dental and genetic samples were obtained here, where low distance values indicate similitude and vice versa between samples.

For dental data, the mean measure of divergence (MMD) was chosen relative to others, for example, pseudo-Mahalanobis $D^2$ (Konigsberg, 1990). It is a robust statistic that yields reliable results even with problematic traits, such as those, that are highly intercorrelated or invariant across samples; it is also less affected by missing data that characterize most dental studies and, while not necessary for the present comparative analyses, has a significance test (Irish, 2010; Nikita, 2015; Sjøvold, 1973, 1977). Finally, it was

### TABLE 1 (Continued)

| Dental sample                  | Abbreviation | Region          | Country/Area       | Data source                  | n   | Lat  | Lon  |
|--------------------------------|--------------|-----------------|--------------------|------------------------------|-----|------|------|
| Kamberla 1.2.3 (13th–17th Cent) | D30_KBR      | North Europe    | North Estonia      | Turner unpublished data      | 160 | 59.5 | 25.3 |
| Ladoga Finns                   | D31_FIN      | North Europe    | Finland/Western Russia | Turner unpublished data      | 51  | 61.0 | 30.0 |
| Lapps (Kola Peninsula)         | D32_LAP      | North Europe    | Lapland/Northwest Russia | Turner unpublished data; Scott & Irish, 2017 | 64  | 67.0 | 40.0 |

aSamples contain some pre-19th century specimens.

bSample specimens are all pre-19th century.
| Sample name               | Abbreviation | Region         | Country/area          | Data source                     | n  | Lat  | Lon  |
|---------------------------|--------------|----------------|-----------------------|---------------------------------|----|------|------|
| **Africa**                |              |                |                       |                                 |    |      |      |
| Moroccan                  | G1_MOR       | North Africa   | Morocco, Casablanca   | Lazaridis et al., 2014          | 10 | 33.5 | −7.6 |
| Algerian                  | G2_ALG       | North Africa   | Algeria               | Lazaridis et al., 2014          |  7 | 36.8 |  3.0 |
| Kikuyu                    | G3_KKU       | East Africa    | Kenya                 | Lazaridis et al., 2014          |  4 | −0.4 | 36.9 |
| Khomani (San)             | G4_KHO       | South Africa   | South Africa          | Lazaridis et al., 2014          | 11 | −27.8| 21.1 |
| Ju_hoan_North (San)       | G5_JUH       | South Africa   | Namibia               | Patterson et al., 2012; Pickrell & Pritchard, 2012 | 21 | −18.9| 21.5 |
| Wolof                     | G6_WOL       | West Africa    | Gambia                | Gambian Genome Variation Project |  5 | 13.4 | −16.7|
| Mozabite                  | G7_MOZ       | North Africa   | Algeria               | Patterson et al., 2012          | 21 | 32.0 |  3.0 |
| Somalia                   | G8_SOM       | East Africa    | Somalia               | Lazaridis et al., 2014          | 13 |  5.6 | 48.3 |
| Sotho                     | G9_SOT       | South Africa   | South Africa          | Patterson et al., 2012          |  1 | −29.0| 29.0 |
| Tswana                    | G10_TSW      | South Africa   | South Africa/ Botswana/ Namibia | Patterson et al., 2012; Pickrell & Pritchard, 2012 |  7 | −28.0| 24.0 |
| Yoruba                    | G11_YOR      | West Africa    | Nigeria               | Lazaridis et al., 2014          | 70 |  7.4 |  3.9 |
| Zulu                      | G12_ZUL      | South Africa   | South Africa          | Patterson et al., 2012          |  1 | −28.0| 31.0 |
| **Total**                 |              |                |                       |                                 | 171|      |      |
| America, Asia, Australia, Melanesia, Europe | | | | | |
| Pima                      | G13_PIM      | Mesoamerica    | Chihuahua, Mexico     | Patterson et al., 2012          | 14 | 29.0 |−108.0|
| Aleut (Nikolskoye)        | G14_ALE      | East Russia    | Bering Island, Russia | Lazaridis et al., 2014          |  2 | 55.2 | 166.0|
| Kyrgyz                    | G15_KRG      | Central Asia   | North Kyrgyzstan      | Lazaridis et al., 2014          |  9 | 42.9 |  74.6|
| Mongola                   | G16_MON      | Central Asia   | East Mongolia         | Patterson et al., 2012          |  6 | 45.0 | 111.0|
| Mansi                     | G17_MAN      | Central Asia   | Central Russia (Konda River) | Lazaridis et al., 2014          |  3 | 62.5 |  63.3|
| Chukchi                   | G18_CHU      | Central Asia   | Northeast Russia      | Lazaridis et al., 2014          | 10 | 69.5 | 168.8|
| Thai                      | G19_THA      | East Asia      | Central Thailand      | Lazaridis et al., 2014          | 10 | 13.8 | 100.5|
| Kinh_Vietnam_KHV          | G20_KIN      | East Asia      | North Vietnam         | Lazaridis et al., 2014          |  8 | 21.0 | 105.9|
| Japanese                  | G21_JAP      | East Asia      | Central Japan         | Patterson et al., 2012          | 29 | 38.0 | 138.0|
| Malays                    | G22_MAL      | Southeast Asia | Central Malaysia      | Skoglund et al., 2016           |  9 |  4.2 |102.0 |
| Visayan, Kankanaey, Ilocano, Tagalog | G23_PHI | Southeast Asia | Central Philippines | Skoglund et al., 2016 | 21 | 9.8  | 125.5|
| Lebbo                     | G24_LEB      | Southeast Asia | Central Borneo        | Skoglund et al., 2016 (signed letter) |  8 |  0.0 | 115.0|
| CAI - North Australia/ Queensland, WPA - North Australia/Queensland, Australian_ECCAC | G25_AUN | Australia | Northeast Australia | Lazaridis et al., 2014 | 3  | −16.9| 145.0|
| All HO New Britain from Skoglund 2016 | G26_NBR | Melanesia | New Britain | Skoglund et al., 2016 (signed letter) | 156 | −5.8 | 150.8|
| Kusunda                   | G27_KUS      | South Asia     | Central Nepal         | Lazaridis et al., 2014          | 10 | 28.1 | 82.5 |
| Greek_Coriell             | G28_GRK      | South Europe   | East Greece           | Lazaridis et al., 2014          | 20 | 38.0 |  23.7|
| Italian_Tuscan            | G29_ITY      | South Europe   | Central Italy         | Patterson et al., 2012          | 20 | 43.0 |  11.0|
| Estonian                  | G30_EST      | North Europe   | West Estonia          | Lazaridis et al., 2014          | 10 | 58.5 |  24.9|
| Finnish_FIN               | G31_FIN      | North Europe   | South Finland         | Lazaridis et al., 2014          |  8 | 60.2 |  24.9|
found that MMD values are more highly correlated with geographic distances (Irish, 2010, 2016; Schillaci, Irish, & Wood, 2009). The formula used here has a bias correction, the Freeman and Tukey angular transformation, to correct for very low or high trait frequencies and small sample sizes (Green & Suchey, 1976; Sjøvold, 1973, 1977). As required by the MMD and to simplify presentation of dental trait frequencies, rank-scale ASUDAS data were dichotomized into categories of present and absent using standard breakpoints (refer to Table 3) (Irish, 1993, 1997, 2005, 2006; Scott & Irish, 2017; Scott & Turner II, 1997; Scott, Turner II, et al., 2018). A few workers suggest rank-scale data would give better results (Gross & Edgar, 2019; Nikita, 2015; Rathmann et al., 2017)—a concept that is not novel (see Sjøvold, 1977; Turner II, 1985b). However, suitably dichotomized trait data, perhaps surprisingly, hold several advantages

(a) importantly, $h^2$ estimates were demonstrated to be higher (see Stojanowski et al., 2019 for details), (b) weighting bias from different grade numbers across ASUDAS traits is avoided, (c) proven distance statistics like the MMD (and $D^2$) can be applied, and (d) residual intra- and inter-observer error is reduced further. That said, the latter should be negligible, at least relative to the above studies, because data were recorded by Turner—the ASUDAS designer, and DJI, who was directly instructed by and calibrated with him (Haeussler, Turner II, & Irish, 1988; Irish & Turner II, 1990).

Many measures of divergence are available for genomic data. However, only $f_{st}$, outgroup-$f_{st}$, and $F_{st}$ were considered due to their ubiquitous application, the availability of online programs and, importantly, the capability of these programs to process large numbers of markers (Holsinger & Weir, 2009; Patterson et al., 2012; Peter, 2016;

| Sample name | Abbreviation | Region | Country/area | Data source | n  | Lat | Lon |
|-------------|--------------|--------|--------------|-------------|----|-----|-----|
| Saami_WGA   | G32_SAM      | North Europe | North Finland | Lazaridis et al., 2014; Mallick et al., 2016 | 3  | 68.4 | 23.6 |
| Total       |              |         |              |             | 359|     |     |
| Grand Total |              |         |              |             | 530|     |     |

FIGURE 1 Origin locations of the 32 dental and matched 32 genetic samples. Modified version originally created using Google My Maps (https://www.google.com)
| Trait/grades present            | D1_BED | D2_KAB | D3_KKU | D4_RRI | D5_SAN | D6_SEN | D7_SHA | D8_SOM | D9_SOT | D10_TSW | D11_YOR | D12_ZUL |
|--------------------------------|--------|--------|--------|--------|--------|--------|--------|--------|--------|---------|---------|---------|
| Winging UI1 (+ = ASU 1)        | 5.4    | 0.0    | 0.0    | 8.1    | 10.5   | 1.9    | 6.0    | 4.0    | 6.0    | 4.0     | 0.0     | 0.0     |
| Labial curvature UI1 (+ = ASU 2) | 37.1  | 50.0   | 50.0   | 65.1   | 50.0   | 28.6   | 27.8   | 60.6   | 68.6   | 42.9    | 27.8    | 77.8    |
| Palatine torus (+ = ASU 2)     | 2.5    | 3.5    | 0.0    | 8.1    | 10.5   | 0.0    | 0.0    | 0.0    | 0.0    | 0.0     | 0.0     | 0.0     |
| Shoveling UI1 (+ = ASU 3)      | 12.5   | 12.5   | 0.0    | 8.3    | 25.0   | 5.9    | 2.7    | 0.0    | 0.0    | 0.0     | 0.0     | 0.0     |
| Interruption groove UI2 (+ = ASU 3) | 37.5  | 21.4   | 14.3   | 3.7    | 14.4   | 21.4   | 46.2   | 0.0    | 4.7    | 0.0     | 10.0    | 15.8    |
| Tuberculum dentale UI2 (+ = ASU 3) | 43.5  | 50.0   | 33.3   | 42.3   | 44.2   | 58.3   | 25.0   | 39.1   | 36.6   | 37.5    | 50.0    | 29.7    |
| Bushman canine UC (+ = ASU 1)  | 0.0    | 0.0    | 11.5   | 31.0   | 38.8   | 10.5   | 0.0    | 3.1    | 24.4   | 37.8    | 22.2    | 23.4    |
| Distal accessory ridge UC (+ = ASU 3) | 12.0  | 27.3   | 29.6   | 16.7   | 21.8   | 41.7   | 22.2   | 34.5   | 39.0   | 60.6    | 14.3    | 26.3    |
| Hypocone UM2 (+ = ASU 3)       | 58.8   | 63.6   | 79.0   | 96.0   | 86.4   | 64.5   | 68.4   | 71.0   | 93.7   | 77.6    | 73.9    | 91.1    |
| Parastyle UM3 (+ = ASU 3)      | 0.0    | 0.0    | 4.4    | 2.7    | 0.0    | 0.0    | 7.7    | 2.1    | 0.0    | 2.3     | 4.0     | 0.0     |
| Enamel extension UM1 (+ = ASU 3) | 5.6    | 0.0    | 0.0    | 2.3    | 5.9    | 4.8    | 4.4    | 4.8    | 3.9    | 10.7    | 1.7     | 0.0     |
| Root number UP1 (+ = ASU 3)    | 50.0   | 52.2   | 69.2   | 44.7   | 39.5   | 69.0   | 52.2   | 75.0   | 74.4   | 57.1    | 53.9    | 69.0    |
| Root number UM2 (+ = ASU 3)    | 69.0   | 68.4   | 87.5   | 80.8   | 84.9   | 78.3   | 72.2   | 82.9   | 83.3   | 65.0    | 100.0   | 84.6    |
| Peg-reduced UI2 (+ = ASU 3)    | 0.0    | 6.3    | 0.0    | 1.6    | 7.0    | 0.0    | 0.0    | 0.0    | 3.3    | 1.9     | 4.2     | 3.2     |
| Trait/grades present | D1_BED* | D2_KAB | D3_KKU | D4_RRI | D5_SAN | D6_SEN | D7_SHA | D8_SOM | D9_SOT | D10_TSW | D11_YOR | D12_ZUL |
|----------------------|---------|--------|--------|--------|--------|--------|--------|--------|--------|---------|---------|---------|
| (+ = ASU P or R) n  | 27      | 16     | 55     | 64     | 115    | 20     | 13     | 73     | 61     | 54      | 24      | 63      |
| Odontome P1-P2 %    | 0.0     | 0.0    | 4.1    | 0.0    | 0.0    | 0.0    | 0.0    | 0.0    | 1.6    | 1.6     | 4.2     | 0.0     |
| (+ = ASU +) n       | 40      | 22     | 49     | 39     | 105    | 32     | 23     | 68     | 64     | 61      | 24      | 63      |
| Congenital absence UM3 % | 21.1   | 3.5    | 5.1    | 7.3    | 2.0    | 13.5   | 23.1   | 4.1    | 4.6    | 7.1     | 3.7     | 6.8     |
| (+ = ASU -) n       | 38      | 29     | 59     | 69     | 98     | 37     | 26     | 74     | 66     | 56      | 27      | 59      |
| Midline diastema UI1 % | 8.8    | 12.0   | 7.6    | 5.1    | 9.7    | 7.7    | 0.0    | 3.0    | 8.5    | 2.1     | 15.8    | 10.2    |
| (+ ≥ 0.5 mm) n      | 34      | 25     | 53     | 59     | 114    | 26     | 23     | 67     | 59     | 47      | 19      | 59      |
| Lingual cusp LP2 %  | 64.3    | 69.2   | 80.8   | 74.2   | 64.6   | 58.3   | 92.3   | 63.2   | 66.7   | 68.6    | 50.0    | 54.7    |
| (+ = ASU 2–9) n     | 28      | 13     | 26     | 31     | 96     | 12     | 13     | 38     | 57     | 51      | 8       | 53      |
| Anterior fovea LM1 % | 37.5    | 60.0   | 84.8   | 70.0   | 70.6   | 50.0   | 29.4   | 50.0   | 68.8   | 74.4    | 70.0    | 54.0    |
| (+ = ASU 2–4) n     | 24      | 10     | 46     | 20     | 68     | 12     | 17     | 52     | 48     | 43      | 10      | 50      |
| Mandibular torus %  | 2.9     | 0.0    | 0.0    | 18.0   | 0.0    | 0.0    | 4.2    | 0.0    | 1.5    | 0.0     | 0.0     | 0.0     |
| (+ = ASU 2–3) n     | 35      | 19     | 55     | 55     | 116    | 15     | 24     | 64     | 65     | 61      | 14      | 63      |
| Groove pattern LM2 % | 46.9    | 27.8   | 75.0   | 67.4   | 74.1   | 58.3   | 36.8   | 58.3   | 64.8   | 75.6    | 42.9    | 72.0    |
| (+ = ASU Y) n       | 32      | 18     | 48     | 46     | 112    | 12     | 19     | 60     | 54     | 45      | 14      | 50      |
| Rocker jaw %        | 9.4     | 10.5   | 11.1   | 5.7    | 8.9    | 6.7    | 8.3    | 9.4    | 1.5    | 5.2     | 14.3    | 3.2     |
| (+ = ASU 1–2) n     | 32      | 19     | 54     | 53     | 45     | 15     | 24     | 64     | 66     | 58      | 14      | 62      |
| Cusp number LM1 %   | 12.5    | 31.3   | 13.0   | 8.0    | 4.7    | 7.1    | 9.5    | 14.0   | 13.8   | 6.3     | 0.0     | 1.9     |
| (+ = ASU 6+) n      | 32      | 16     | 46     | 25     | 107    | 14     | 21     | 57     | 58     | 48      | 11      | 54      |
| Cusp number LM2 %   | 42.9    | 33.3   | 82.0   | 96.7   | 92.6   | 54.6   | 31.6   | 57.9   | 83.6   | 76.6    | 83.3    | 90.0    |
| (+ = ASU 5+) n      | 28      | 18     | 50     | 30     | 108    | 11     | 19     | 57     | 55     | 47      | 12      | 50      |
| Deflecting wrinkle LM1 % | 15.6  | 6.7   | 34.0   | 34.6   | 25.6   | 27.3   | 5.0    | 27.1   | 30.0   | 26.7    | 20.0    | 31.4    |
| (+ = ASU 2–3) n     | 32      | 15     | 47     | 26     | 78     | 11     | 20     | 59     | 50     | 45      | 10      | 51      |
| C1-C2 crest LM1 %   | 3.0     | 0.0    | 0.0    | 0.0    | 3.1    | 8.3    | 0.0    | 0.0    | 1.9    | 2.2     | 0.0     | 2.0     |
| (+ = ASU +) n       | 33      | 14     | 47     | 25     | 65     | 12     | 20     | 60     | 53     | 45      | 11      | 51      |
| Protostylid LM1 %   | 0.0     | 0.0    | 0.0    | 5.6    | 7.8    | 0.0    | 0.0    | 0.0    | 0.0    | 0.0     | 0.0     | 0.0     |
| (+ = ASU 3–6) n     | 33      | 16     | 46     | 18     | 103    | 13     | 21     | 58     | 59     | 49      | 13      | 56      |
| Cusp 7 LM1 %        | 5.9     | 5.9    | 31.9   | 22.0   | 27.9   | 28.6   | 4.8    | 28.8   | 37.9   | 37.5    | 18.2    | 49.1    |
| (+ = ASU 2–4) n     | 34      | 17     | 47     | 41     | 111    | 14     | 21     | 59     | 58     | 48      | 11      | 55      |
| Tome's root LP1 %   | 6.3     | 5.3    | 14.9   | 5.3    | 2.6    | 14.3   | 10.5   | 18.4   | 10.5   | 5.7     | 33.3    | 7.4     |
| (+ = ASU 3–5) n     | 32      | 19     | 47     | 38     | 39     | 14     | 19     | 49     | 38     | 35      | 12      | 27      |
| Root number LC %    | 0.0     | 20.0   | 0.0    | 0.0    | 0.0    | 0.0    | 0.0    | 1.9    | 0.0    | 0.0     | 0.0     | 0.0     |
| (+ = ASU 2+) n      | 26      | 10     | 50     | 37     | 42     | 12     | 16     | 53     | 32     | 29      | 13      | 33      |
Reich, Thangaraj, Patterson, Price, & Singh, 2009; Skoglund et al., 2015). Of these, $F_{ST}$ was selected as most appropriate for the SNP data, samples, and overall approach (Supporting information Text S1.1). Calculated under the Hardy–Weinberg model, it is theoretically not model-free; however, it is used in that capacity here, like the MMD, to "describe overall patterns of variation that can be interpreted in light of population history and structure" (Relethford & Harpending, 1994:251). The motivation is that, beyond its descriptive attributes and use for identifying genomic regions under selection, $F_{ST}$ yields a more generalized estimate of genetic differentiation among population pairs (but see Séré, Thévenon, Belem, & De Meeûs, 2017). Further, it (a) works with small samples if, as here, many loci are included, (b) does well on a broad geographic scale, (c) is not associated with population divergence time, (d) is reliable regardless of the population structure model, and (e) like the MMD, maintains constant pairwise values if any samples are added (Diniz-Filho et al., 2013; Holsinger & Weir, 2009; Nelis et al., 2009; Ortega-Del Vecchyo & Slatkin, 2019; Peter, 2016; Tian et al., 2009; Weir, Cardon, Anderson, Nielsen, & Hill, 2005; Willing, Dreyer, & van Oosterhout, 2012). Both the Weir and Cockerham (1984) and Hudson $F_{ST}$ estimators (Hudson, Slatkin, & Maddison, 1992) were used, with the results of the latter detailed below. The Hudson estimator is stated to return more accurate distances with genomic data, and is less affected by very small sizes, that is, $n < 4$, that affect several genetic samples (see Table 2) (Bhatia, Patterson, Sankararaman, & Price, 2013; Ortega-Del Vecchyo & Slatkin, 2019).

Geographic distances among the 32 dental and 32 genetic samples were determined based on associated latitudes and longitudes in decimal degrees submitted to the Similarity and Distances Indices module in PAST 3.23 (http://folk.uio.no/ohammer/past; Hammer, Harper, & Ryan, 2001; Hammer, 2019). The default PAST output is meters, which were used for all quantitative analyses; however, for convention they were converted to km in the corresponding tables and figures (below and Supporting information). Without reference to hypothesized migration routes they measure, simply, straight-line distances along a great circle over the surface of the earth between coordinates of sample pairs in the WGS84 datum. This method also accounts for variance between longitudes at high and low latitudes (Hammer, 2019, personal communication, 2019). Equivalent results were returned with the Geographic Distance Matrix Generator (version 1.2.3) (Ersts, 2014).

Finally, as mentioned, correlations between dental, genetic, and geographic distances were calculated. Like prior studies (Hubbard et al., 2015; Rathmann et al., 2017) Mantel tests were used, with a null hypothesis of no association between matrices (Mantel, 1967; Smouse & Long, 1992; Smouse, Long, & Sokal, 1986; Sokal & Rohlf, 1995). To explore dental vs. genetic correspondence irrespective of geographic separation, partial Mantel tests were also conducted; for these, the third variable consists of "midpoint" distances calculated from mean latitude and longitude coordinates between the matching pairs of dental and genetic samples. Of course, Mantel tests are not without criticism. It was said that $r_m$ values "obtained by permutations do not display enough variability,"
TABLE 4  MMD distance matrix (bottom diagonal) for the three North and nine sub-Saharan African dental samples based on 36 ASUDAS traits, and FST distance matrix (top diagonal) for the three North and nine sub-Saharan African genetic samples based on 353,091 SNPs

|          | G1_MOR | G2_ALG | G3_KKU | G4_KHO | G5_JUH | G6_WOL | G7 MOZ | G8_SOM | G9_SOT | G10_TSW | G11_YOR | G12_ZUL |
|----------|--------|--------|--------|--------|--------|--------|--------|--------|--------|---------|---------|---------|
| D1_BED  | 0.016  | 0.053  | 0.122  | 0.171  | 0.082  | 0.009  | 0.031  | 0.091  | 0.094  | 0.086   | 0.095   | G1_MOR  |
| D2_KAB  | 0.000  | 0.070  | 0.138  | 0.188  | 0.099  | 0.024  | 0.046  | 0.110  | 0.112  | 0.103   | 0.112   | G2_ALG  |
| D3_KKU  | 0.095  | 0.078  | 0.063  | 0.101  | 0.026  | 0.066  | 0.020  | 0.016  | 0.020  | 0.015   | 0.022   | G3_KKU  |
| D4_RRI  | 0.136  | 0.145  | 0.025  | 0.078  | 0.136  | 0.087  | 0.039  | 0.038  | 0.070  | 0.055   |         | G4_KHO  |
| D5_SAN  | 0.149  | 0.150  | 0.050  | 0.000  | 0.111  | 0.186  | 0.131  | 0.072  | 0.068  | 0.105   | 0.089   | G5_JUH  |
| D6_SEN  | 0.000  | 0.000  | 0.015  | 0.050  | 0.046  | 0.096  | 0.053  | 0.028  | 0.032  | 0.021   | 0.031   | G6_WOL  |
| D7_SHA  | 0.000  | 0.000  | 0.113  | 0.177  | 0.205  | 0.006  | 0.043  | 0.106  | 0.110  | 0.100   | 0.110   | G7_MOZ  |
| D8_SOM  | 0.048  | 0.031  | 0.018  | 0.087  | 0.115  | 0.000  | 0.067  | 0.051  | 0.057  | 0.049   | 0.057   | G8_SOM  |
| D9_SOT  | 0.117  | 0.096  | 0.000  | 0.009  | 0.032  | 0.008  | 0.146  | 0.025  | 0.002  | 0.011   | 0.004   | G9_SOT  |
| D10_TSW | 0.150  | 0.127  | 0.017  | 0.026  | 0.038  | 0.023  | 0.178  | 0.048  | 0.000  | 0.016   | 0.008   | G10_TSW |
| D11_YOR | 0.049  | 0.045  | 0.000  | 0.032  | 0.041  | 0.000  | 0.085  | 0.007  | 0.001  | 0.043   | 0.014   | G11_YOR |
| D12_ZUL | 0.133  | 0.139  | 0.020  | 0.029  | 0.033  | 0.033  | 0.187  | 0.075  | 0.000  | 0.024   | 0.024   | G12_ZUL |

aG1_MOR = Moroccan, G2_ALG = Algerian, G3_KKU = Kikuyu, G4_KHO = Khomani San, G5_JUH = Ju-hoan North San, G6_WOL = Wolof, G7 MOZ = Mozabite, G8_SOM = Somalia, G9_SOT = Sotho, G10_TSW = Tswana, G11_YOR = Yoruba, G12_ZUL = Zulu. Details in Table 2.
bD1_BED = Bedouin, D2_KAB = Kabyle, D3_KKU = Kikuyu, D4_RRI = Riet River San, D5_SAN = San, D6_SEN = Senegambia, D7_SHA = Shawia, D8_SOM = Somalia, D9_SOT = Sotho, D10_TSW = Tswana, D11_YOR = Yoruba, D12_ZUL = Zulu. Details in Table 1.
and spatial autocorrelation, which is often inherent with biological data, can lead to inaccurately low \( p \)-values (Legendre & Fortin, 2010; Guillot & Rousset, 2013:341). Nevertheless, Mantel tests have been shown to be robust (Séré et al., 2017), are easy to interpret, remain widely used (above) to facilitate between-study comparisons, and various alternative methods are neither unreservedly accepted nor, in this case, applicable, for example, comparing directly the non-dichotomized ASUDAS and SNP data. In any event, results based on applications of Mantel and alternative methods were demonstrated to largely converge (Diniz-Filho et al., 2013). Therefore, distance matrices were submitted to the Mantel Test module of PAST 3.23. The resulting \( r_m \)-values are Pearson’s correlation coefficients, with one-tailed \( p \)-values from 10,000 random permutations (Hammer, 2019). A \( p \)-value adjustment such as the Bonferroni procedure is often used to address Type I error from multiple testing (see Rathmann et al., 2017) and/or spatial autocorrelation. Obviously, a lower alpha level can increase “a Type II error [some say] to unacceptable levels,” and because of the magnitude of the resulting \( r_m \)-values in the present study (see below) all significant \( p \)-values are exceptionally low, that is, \( p = 0.001 \) to 9.999E-05. Thus, correction was not to be deemed necessary (Nakagawa, 2004:1045; Gelman, Hill, & Yajima, 2012). As above, identical results were achieved with an alternate program, in this case Function Mantel in Vegan R-package 2.5–6 (https://cran.r-project.org; https://github.com/vegandevs/vegan; Oksanen et al., 2013).

3 | RESULTS

3.1 | African analyses

Table 3 lists percentages of expression for the 36 traits in the North and sub-Saharan African dental samples, and the total number of individuals recorded. All 36 ASUDAS breakpoints are provided as well. As mentioned, SNP data in the corresponding African genetic samples are available from the literature (Lazaridis et al., 2014; Patterson et al., 2012; Pickrell & Pritchard, 2012) and via the Gambian Genome Variation Project.

To maximize the comparison, all 36 dental traits were submitted first to the MMD. The resulting inter-sample distances are listed in the bottom diagonal of Table 4. Those from Hudson \( F_{ST} \) for the genetic samples are in the top diagonal. To visualize the inter-sample variation, multi-dimensional scaling (MDS) in SPSS 25.0 Procedure Alscal was used to generate three-dimensional spatial representations
of each matrix (Figures 2 and 3). Interval-level MDS solutions provide good representations of the MMD, with a Kruskal’s stress formula 1 value of .099 and $r^2$ of .943 and, particularly, $F_{ST}$ distances with a stress value of .044 and $r^2$ of .992. It is apparent that the dental sample locations correspond roughly with geographic origins, where north-to-south is along the X- and west-to-east on the Y-axis (Figure 2). A similar, yet less marked geographic distribution can be seen among the genetic samples (Figure 3).

These qualitative observations are sustained by Mantel test results. The correlation between 36-trait MMD and geographic distances (Table 5) for the dental samples is strongly positive, $r_m = 0.682$ ($p = 0.000$). Between $F_{ST}$ and geographic matrices for the 12 matching genetic samples (Table 5) the $r_m$-value is moderately positive, .486 ($p = 0.001$). Yet, the $r_m$-value between the 36-trait MMD and $F_{ST}$ matrices indicates a very strong correlation, .786 ($p = 0.000$). This correspondence is supported further in controlling for the geographic midpoint distances (Supporting information Table S1); that is, between MMD and $F_{ST}$ residuals the partial Mantel correlation remains strong, $r_m = 0.699$ ($p = 0.000$). Scatterplots depicting these correlations are presented in Figure 4.

Next, the 36 ASUDAS traits were reduced to 25, matching those in Turner’s database (refer to list in Table 7, below). However, before that, the 12 dental and 12 genetic samples were again compared to quantify any effect that reduced trait number has on the results. So as before, this trait set was submitted to the MMD. The new values are listed in Table 6 and depicted via MDS in Figure 5; for this solution the stress increased slightly (.107), and $r^2$ decreased (.939). Minor differences are evident between the 36- and 25-trait matrices (compare Tables 5 and 6) and in relative sample locations (Figures 2 and 5). Yet, not unexpectedly, the correlation between the two MMD matrices is almost perfect, $r_m = 0.977$ ($p = 0.000$), as seen in Figure 6a. The remaining correlations based on 25 traits all increased: (a) $r_m = 0.696$ ($p = 0.000$) for the MMD and geographic distances among dental samples (Figure 6b), (b) $r_m = 0.838$ ($p = 0.000$) for MMD and $F_{ST}$ (Figure 6c), and (c) $r_m = 0.782$ ($p = 0.000$) for the partial correlation between MMD and $F_{ST}$ residuals, relative to the geographic midpoint distances (Figure 6d).

### 3.2 | Global analyses

The 25 ASUDAS percentages for the 20 non-African dental samples using the same breakpoints as before are listed in Table 7. To include the maximum number of traits, plus dental samples—to match with...
### Table 5
Geographic distance matrix in km for the three North and nine sub-Saharan African dental samples (bottom diagonal), and for the three North and nine sub-Saharan African genetic samples (top diagonal)

|        | G1_MOR  | G2_ALG  | G3_KKU  | G4_KHO  | G5_JUH  | G6_WOL  | G7_MOZ  | G8_SOM  | G9_SOT  | G10_TSW | G11_YOR | G12_ZUL |
|--------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| D1_BED | 1.02917 | 5.97447 | 7.43269 | 6.57297 | 2.41347 | 1.00529 | 6.53046 | 7.92245 | 7.58208 | 3.12561 | 7.93443 | G1_MOR  |
| D2_KAB | 825.32  | 5.40140 | 7.39740 | 6.46435 | 3.25116 | 5.3246  | 5.75243 | 7.77422 | 7.50196 | 3.25730 | 7.74819 | G2_ALG  |
| D3_KKU | 5.79278 | 5.28023 | 3.64945 | 2.64897 | 6.10437 | 5.05771 | 1.43034 | 3.27461 | 3.34988 | 3.76384 | 3.11840 | G3_KKU  |
| D4_RRI | 7.76995 | 7.62170 | 3.42322 | 986.52  | 6.11556 | 6.89049 | 4.71168 | 785.35  | 286.36  | 4.31225 | 974.67  | G4_KHO  |
| D5_SAN | 7.07309 | 6.88677 | 2.73892 | 768.10  | 5.50625 | 5.96831 | 3.99792 | 1.35341 | 1.03961 | 3.49184 | 1.39806 | G5_JUH  |
| D6_SEN | 2.46048 | 3.11113 | 6.05185 | 6.64564 | 6.12867 | 2.87733 | 7.17123 | 6.79278 | 6.34260 | 2.34989 | 6.88524 | G6_WOL  |
| D7_SHA | 1.08238 | 301.77  | 4.99302 | 7.40745 | 6.66488 | 3.22573 | 5.53732 | 7.28991 | 7.00308 | 2.72516 | 7.72764 | G7_MOZ  |
| D8_SOM | 5.97089 | 5.28283 | 1.54328 | 4.83098 | 4.20969 | 6.88666 | 4.98115 | 4.35088 | 4.54235 | 4.91317 | 4.15631 | G8_SOM  |
| D9_SOT | 7.92751 | 7.73839 | 3.31939 | 332.20  | 855.29  | 6.90096 | 7.51318 | 4.67107 | 501.86  | 4.84440 | 224.99  | G9_SOT  |
| D10_TSW | 7.34369 | 7.19910 | 3.15283 | 426.27  | 415.69  | 6.25234 | 6.98720 | 4.61404 | 650.00  | 4.47394 | 688.44  | G10_TSW |
| D11_YOR | 3.22950 | 3.32414 | 3.80064 | 4.63190 | 4.00404 | 2.30922 | 3.21419 | 4.84475 | 4.83491 | 4.21120 | 4.88228 | G11_YOR |
| D12_ZUL | 8.00521 | 7.75948 | 3.09251 | 7.5207  | 1.00421 | 7.13266 | 7.52031 | 4.36811 | 430.16  | 964.81  | 4.99472 | G12_ZUL |

aG1_MOR = Moroccan, G2_ALG = Algerian, G3_KKU = Kikuyu, G4_KHO = Khomani San, G5_JUH = Ju-hoan North San, G6_WOL = Wolof, G7_MOZ = Mozabite, G8_SOM = Somalia, G9_SOT = Sotho, G10_TSW = Tswana, G11_YOR = Yoruba, G12_ZUL = Zulu. Details in Table 2.

bD1_BED = Bedouin, D2_KAB = Kabyle, D3_KKU = Kikuyu, D4_RRI = Riet River San, D5_SAN = San, D6_SEN = Senegambia, D7_SHA = Shawia, D8_SOM = Somalia, D9_SOT = Sotho, D10_TSW = Tswana, D11_YOR = Yoruba, D12_ZUL = Zulu. Details in Table 1.
the most available genetic sample counterparts, a few individual counts (n ≤ 6), and/or samples (D17_LOK, D18_CHU, D25_AUN) are unavoidably small. In these cases, results should be interpreted with caution. SNP data for the 20 genetic samples are again from published sources (Lazaridis et al., 2014; Mallick et al., 2016; Patterson et al., 2012; Pickrell & Pritchard, 2012; Skoglund et al., 2016) or were obtained by request.

The 32 x 32 25-trait MMD and Hudson $F_{ST}$ matrices are too large to fit with the main text, so are included in Supplemental Information Tables S2-S3. However, the resulting MDS plots are presented in Figures 7 and 8, respectively. The three-dimensional solutions yielded good resolution for the MMD (stress = 0.088; $r^2 = 0.960$) and $F_{ST}$ distances (stress = 0.071; $r^2 = 0.978$). The dental and genetic samples cluster relative to their geographic origins.

The Mantel correlation between MMD and geographic distances for the 32 dental samples (Supporting information Table S4) is strongly positive, $r_m = 0.710$ ($p = 0.000$), as it is now also for $F_{ST}$ and genetic sample geographic distances, $r_m = 0.768$ ($p = 0.000$) (Supporting information Table S5). The $r_m$-value between the MMD and $F_{ST}$ matrices again indicates a very strong correlation, .720 ($p = 0.000$). Unlike the African findings, when controlling for the global geographic midpoint distances (Supporting information Table S6), the partial Mantel correlation between MMD and $F_{ST}$ residuals is only moderately positive, $r_m = 0.400$ ($p = 0.000$). Scatterplots depicting these various correlations are provided in Figure 9. Lastly, because the abovementioned dental samples affected most by small numbers are from Turner’s database, it was decided to rerun these same Mantel tests with only his 18 samples to explore whether correlations alter substantially. Doing so also serves to quantify if the relatively large number of African samples artificially inflated the $r_m$-value, while identifying indirectly major inter-observer error with JDI. Although slightly lower, as would be expected because of fewer more geographically limited samples, the Mantel correlations
remain similar in magnitude to those of the 32-sample analyses (Supporting information Text S1.2).

4 | DISCUSSION

4.1 | Interpretations

Some unpublished data were included (D3_KKU and D11_YOR), but the 36-trait MMD distances (Table 4) and MDS plot of 12 African dental samples (Figure 2) parallel prior results (Irish, 1993, 1997, 1998a, 2010, 2016; Irish et al., 2014). That is, based on documented population history these ASUDAS traits provided reliable phenetic affinities among samples, and patterning indicative of geographic provenience. Reliability is why the ASUDAS has had such a long run in population studies worldwide, as evidenced by hundreds of publications (https://scholar.google.com/) and as summarized in several compendia (Scott & Irish, 2013, 2017; Scott & Turner II, 1997; Scott, Turner II, et al., 2018). Now, however, what had only been assumed from earlier research about the genetic component of trait expression (Scott et al., 1983; Sofaer et al., 1972; Turner II, 1985a), but recently queried (Hughes et al., 2016; Stojanowski et al., 2019; etc.)—that dental affinities approximate genetic relatedness, is readily testable empirically with genomic data (Delgado et al., 2019; Gross & Edgar, 2019; Hubbard et al., 2015; Rathmann et al., 2017). In the present analysis, the MDS plot of $F_{ST}$ distances (Figure 3) is somewhat akin to that from dental data, but the correlation between 36-trait MMD and $F_{ST}$ matrices (Table 4) is most telling (Figure 4; Table 8). Stronger than $r_{M}$-values in Hubbard et al. (2015) and Rathmann et al. (2017), it affords additional support for using ASUDAS traits as genetic proxies. So too does the relation between MMD and dental geographic distances (Table 5), and the partial MMD-$F_{ST}$ correlation controlling for geographic midpoint distances (Supplemental Information Table S1). The $r_{M}$-value of the latter infers that geographic separation is not an overriding factor in the African dental-genetic correspondence. The lower, yet still moderately positive correlation between $F_{ST}$ and geographic distances (Table 5) may, on this continental level, indicate that $F_{ST}$ does not specifically detect increased gene flow between geographically remote populations—particularly since the 19th–20th century dates of the present dental samples, along with extreme reproductive isolation, as in those populations who may be geographically proximate but genetically divergent (Jay, Sjödin, Jakobsson, & Blum, 2012; Ramachandran et al., 2005).

Seemingly contrary to purpose, reducing the number of dental traits from 36 to 25 (Table 6) increased the $r_{M}$-values between MMD and geographic distances, and MMD and $F_{ST}$ distances among the African samples. The partial MMD-$F_{ST}$ correlation controlling for geographic midpoint distances increased most (Figure 6; Table 8). This is all despite the minimal change in 36- and 25-trait distances, as indicated by a correlation near 1.0 and similitude in MDS configurations.
Deleting the 11 traits essentially functioned to emulate the editing process typically used prior to submitting data to the MMD and other similar statistics (Irish, 2010). That is, reliable results are attainable with problematic traits as noted, but it is prudent to delete the same. If this study focused only on ASUDAS-based affinities (Irish, 2005, 2006, 2016; etc.), then nine of these traits would have been deleted in any event as standard practice, for being: 1) mostly invariant (palatine torus, UI2 peg-reduced, mandibular torus), 2) otherwise unimportant for driving inter-sample variation based on low loadings (<.5) in principal component analysis (UC distal accessory ridge, rocker jaw), and 3) highly inter-correlated ($\tau_b \geq |0.5|$) with other traits (UI1 labial curvature; LM1 anterior fovea, LM1 deflecting wrinkle, LM1 C1-C2 crest). Thus, while one idea was to include more traits than similar studies to maximize comparative analyses, it is apparent here that “more” is not always “better.”

Finally, for the global-level analyses, the MDS plot of 32 dental samples (Figure 7) based on 25-trait MMD distances (Supporting information Table S2), and the matching plot (Figure 8) of the $F_{ST}$ matrix (Supporting information Table S3) are comparable. Explicitly, from sub-Saharan Africa to the Americas and Melanesia, the samples cluster by regional origin and evidence overall geographic patterning. The latter is quantified by correlations >.7 (Table 8; Figure 9a,b) for both MMD and $F_{ST}$ with dental and genetic sample geographic distances (Supporting information Table S4-S5). As first suggested >20 years ago (Irish, 1997:463), the MMD variation depicted in Figure 7 may be detecting the vestiges of “an expansive dental morphological cline.” Recent sub-Saharan Africans were revealed to possess high frequencies of ancestral dental nonmetric traits, while other world populations transition toward higher frequencies of derived traits with increasing geographic separation (Irish, 1998b; Irish & Guatelli-Steinberg, 2003). The idea was later revisited using other dental data (Hanihara, 2013; Reyes-Centeno, Rathmann, Hanihara, & Harvati, 2017). Therefore, if beginning with the two samples of southern African San (D4_RRI, D5_SAN) in the plot, this presumedcline can be seen to extend from sub-Saharan to North Africa and into the Mediterranean region and farther north in Europe. Toward what would be east on the right side of the X-axis, are Northeast and Southeast Asians (also see Jay et al., 2012). The New World samples (D13_PIM, D14_ALU) are then farther to the north and east, and Australians (D25_AUN) and Melanesians (D26_NBR) south, near the bottom of the Y-axis. Naturally, neutral genomic data are increasingly being used to investigate ancient migrations, including routes out of Africa (e.g., Jay et al., 2012; Kantz, Guillot, Antoniazza, Neuenschwander, & Goudet, 2018;
Ramachandran et al., 2005). And here, though a more linear trajectory is illustrated from $F_{ST}$ distances (Figure 8), signs of such a cline are sustained—again starting with the southern African San samples (G4_KHO, G5_JUH) and ending with Australians (G25_AUN) and Melanesians (G26_NBR).

Considerably expanded from continental to global in scale with an additional 20 samples, the $r_m$-value between 25-trait MMD and $F_{ST}$ matrices is still $>0.70$ (Table 8; Figure 9c)—despite the caution that results may be influenced by low counts or small samples. Thus, for these 32 dental samples and MMD distances from ASUDAS data, and these 32 genetic samples and $F_{ST}$ distances from SNP data, the correlation further supports use of dental traits for population affinity research, with or instead of neutral genomic data if the latter are unavailable. The higher correlation between $F_{ST}$ and geographic distances relative to the African results is likely linked to the expanded scale. Irrespective of whether $F_{ST}$ is or is not unequivocally suited to detect isolation by distance (above), it is perhaps picking up on ancient among-region affinities rather than that of recent within-region (or -continent) population movements and genetic exchange. This possibility is also likely why the global partial MMD-$F_{ST}$ correlation controlling for geographic midpoint distances (Supplemental Information Table S6; Figure 9d), though positive (Table 8), is much lower than the African $r_m$-value; geographic separation does appear to be a contributing factor to the overall dental-genetic association in this broad-scale example.

4.2 | Implications

The results cannot be equated directly with ancestry estimates in individuals (Delgado et al., 2019; Gross & Edgar, 2019) but anecdotally, and at least relative to some population-level analyses also presented in these two articles, the present MMD-$F_{ST}$ correlations suggest similar if not greater correspondence of dental and genomic data. More direct comparisons are possible, as above, with articles describing

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**FIGURE 6** Scatterplots of Mantel correlations between matrices from the second African analysis for (a) 25- and 36-trait MMD distances, (b) 25-trait MMD and dental sample geographic distances, (c) 25-trait MMD and Hudson $F_{ST}$ distances, and (d) 25-trait MMD and $F_{ST}$ residuals, controlling for geographic midpoint distances.
outcomes on regional and global levels (Hubbard et al., 2015; Rathmann et al., 2017); the current Mantel correlations are decidedly stronger. Why? And what is the significance?

Why? In answer, it is stressed that the intent is to explore potential reasons for the enhanced dental-genetic correspondence, not to critique prior studies (Delgado et al., 2019; Gross & Edgar, 2019; Hubbard et al., 2015; Rathmann et al., 2017) that provide the foundation for this research. Multiple explanations are possible vis-à-vis differences in data, samples, and methods. First, the numbers of traits, 36 and 25, are larger than in three of the four studies, and unlike all four include root and root-related traits that are highly diagnostic in characterizing populations (Scott & Irish, 2017; Scott & Turner II, 1997; Scott, Turner II, et al., 2018; Turner II et al., 1991). Moreover, when the 11 traits were dropped from analyses, testing revealed that nine are problematic for the specific 32 samples under study, including four that are highly inter-correlated; this "editing" further enhanced the dental data for comparative purposes. And, dental data were recorded by Turner and JDI. Though standardized, the ASUDAS is not intuitive to the point where trait recording can be undertaken without requisite training, including quantification of inter-observer error (Scott & Irish, 2017); examples of suspect affinities and misidentified traits illustrate this potential issue (e.g., Irish & Morris, 1996). With regard to the >350,000 SNPs, among other attributes these genomic data have been shown to better differentiate among populations, and they appear to correspond more closely with dental traits than, for example, STRs (Gross & Edgar, 2019; Rathmann et al., 2017). These markers are also substantially greater in number than in three of the recent studies, including 1,718 SNPs in Rathmann et al. (2017).

Second, 32 dental samples consisting of 2,844 individuals were compared with 32 genetic samples comprised of 530 individuals—all considerably more than the previous studies. But most importantly, though not the same individuals (Hubbard et al., 2015), creation of the dental samples and matching them with their genetic counterparts were less subjective. To illustrate, given the focus of one admixture study, recent African Americans, the authors were obliged to use casts of individuals whose "race was assigned by the orthodontists who assembled the collections" (Gross & Edgar, 2019:522). For global population comparisons, Rathmann et al. (2017) did not have the present luxury of choice among dental samples, so several of their matched pairs are characterized by ethnic, as well as significant linguistic differences, including (a) dental sample Kikuyu of Niger-Congo Language Superfamily versus a genetic sample with some Masai and Luo of Nilo-Saharan Superfamily, (b) dental Zulu of Nguni Branch vs. genetic
Pedi, Sotho, and Tswana in the Sotho Branch of the Bantu Language Family, and (c) dental Haya of Niger-Congo Superfamily vs. genetic sample with Hadza who, if not a language isolate belong to the Khoisan Superfamily (Greenberg, 1963). Further, the overall lack of matches for the Americas necessitated their pairing of a recent dental sample of varied Mexican ethnicities with a genetic sample of archeological specimens, including 2,500 year-old Zapotec (Rathmann et al., 2017).

Third, methodological differences are likely a key factor in correspondence of dental and genomic data, especially their performance in estimating individual ancestry (Delgado et al., 2019; Gross & Edgar, 2019) relative to population affinities (Hubbard et al., 2015; Rathmann et al., 2017; this study). Again, comparing results between these two types of study is impractical, but another matter is the nature of ASUDAS data. While progress has been made using dental nonmetric traits to estimate individual ancestry and affinities (Edgar, 2013, 2015; Irish, 2015; Scott, Pilloud, et al., 2018; Stojanowski & Paul, 2015; Stojanowski & Schillaci, 2006), the ASUDAS was designed to analyze samples and variation therein (Scott & Turner II, 1997; Turner II et al., 1991); thus, the “low predictive power for genetic ancestry of individuals” compared with SNPs is not surprising (Delgado et al., 2019:439). In the Gross and Edgar (2019) study, as recognized, a contributing factor is the genetic program used to estimate ancestry, which is affected by missing data; thus, the poorer performance of dental traits is also related to this issue relative to more complete genomic datasets. Concerning population studies (Hubbard et al., 2015; Rathmann et al., 2017), the justification for instead using MMD and $F_{ST}$ distances in a model-free approach, of course with much larger datasets, has already been discussed. This too may have played a role in stronger Mantel correlations calculated here.

So, what are the implications of this study? It did not address directly questions concerning such perceived attributes of the ASUDAS as low sexual dimorphism, minimal selection, and high heritability in trait expression. These matters have been dealt with elsewhere. Sexual dimorphism could relate simply to tooth size where, for example, larger crowns in males may promote the formation of additional, later-forming cusps, all other developmental factors being equal (Jernvall & Jung, 2000). That said, prior dental nonmetric studies found few or no statistically significant differences for cusp number or other trait by sex (Bermudez de Castro, 1989; Hanihara, 1992; Irish, 1993; Smith & Shegev, 1988). Significant differences that may occur appear random, in that different traits are affected among studies depending on the population (Irish, 2016); for example, it was a factor in the Gullah

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**FIGURE 8** Three-dimensional MDS plot of the Hudson $F_{ST}$ distances among the 32 global genetic samples. The sample abbreviations are defined in Tables 2, 4-5.
heritability paper (Stojanowski et al., 2019), though not so much in the most recent Australian study (Paul et al., 2020). Concerning selection, it was established to impact some traits (Bryk et al., 2008; Kimura et al., 2009; Park et al., 2012; Hlusko et al., 2018), but indirectly as a consequence of pleiotropy. Recent trait variation, at any rate, is seemingly more “a product of random processes (i.e., genetic drift and founder effect) rather than genetic adaptation” (Scott, Turner II, et al., 2018: 223). Lastly, trait heritability was shown to vary across studies and dental fields. The effects of various stressors and other issues on the populations under study may play a role, along with methodological factors like appropriate dichotomization of traits (Higgins et al., 2009; Hughes et al., 2016; Hughes & Townsend, 2011, 2013; Paul et al., 2020; Riga et al., 2014; Stojanowski et al., 2018, 2019). However, the exact answers will require additional research, which is beyond the present scope of study.

Of course, dental and genetic data correspondence was addressed, and they do correspond to a much greater degree than before, based on comparing distances calculated from them. The $r_m$ values, however, do not approach 1.0 (Table 8). Consequently, under the assumption that neutral genomic markers are indeed the definitive choice in affinity study, dental nonmetric traits are not about to supplant them. That said, these correlations may be considered minimum values (also see Rathmann et al., 2017). The data were not collected from the same individuals, and though able to match at a high level of concordance, the paired dental and genetic samples are of different ages and several come from similar, not identical, populations. Some sample size issues, inter-observer error in the ASUDAS data, and other stochastic and nonstochastic factors (Rathmann et al., 2017), also cannot be ruled out totally. In any event, the results speak for themselves. If sufficient attention is paid to the data, samples, and methods, it does appear ASUDAS traits can serve as highly reliable proxies for neutral genomic markers—regardless of potential issues mentioned, including sexual dimorphism, selection, and heritability. If the latter could be identified on individual trait and/or sample bases, which was not the case here, data correspondence may conceivably be even higher.
TABLE 7  Percentages (%) of 25 dental traits considered present and number of individuals scored (n) in the 20 additional (non-African) global samples

| Trait/grades present | D13 _PIM a | D14 _ALE | D15 _KAZ | D16 _MON | D17 _LOK | D18 _THA | D19 _CHU | D20 _VIE | D21 _JAP | D22 _MAL | D23 _PHI | D24 _BOR | D25 _AUN | D26 _NBR | D27 _NEP | D28 _GRK | D29 _ITY | D30 _KBR | D31 _FIN | D32 _LAP |
|----------------------|------------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|----------|
| Winging UI1 %        | 25.5       | 21.8     | 14.6     | 25.0     | 28.6     | 37.0     | 27.6     | 36.4     | 23.3     | 4.2      | 18.2     | 28.6     | 28.6     | 23.2     | 33.3     | 1.5      | 2.1      | 4.3      | 4.3      | 54.5     |
| (+ = ASU 1) n        | 161        | 78       | 48       | 28       | 7        | 27       | 76       | 11       | 86       | 24       | 11       | 14       | 7        | 155      | 12       | 68       | 48       | 70       | 23       | 11       |
| Shoveling UI1 %      | 84.2       | 53.3     | 27.9     | 71.5     | 40.0     | 61.5     | 32.5     | 55.6     | 71.9     | 28.6     | 47.4     | 29.4     | 12.5     | 8.9      | 18.8     | 0.0      | 5.9      | 8.7      | 9.5      | 16.7     |
| (+ = ASU 3–6) n      | 159        | 75       | 43       | 21       | 5        | 13       | 80       | 9        | 89       | 14       | 19       | 17       | 8        | 124      | 16       | 5        | 17       | 69       | 21       | 6        |
| Double shoveling UI1 | 38.5       | 14.5     | 9.4      | 7.7      | 33.3     | 25.0     | 7.7      | 22.2     | 13.2     | 8.3      | 14.3     | 11.8     | 0.0      | 3.3      | 5.9      | 0.0      | 0.0      | 19.4     | 23.8     | 16.7     |
| (+ = ASU 2–6) n      | 156        | 76       | 43       | 13       | 3        | 8        | 65       | 9        | 83       | 12       | 7        | 17       | 5        | 123      | 17       | 5        | 17       | 67       | 21       | 6        |
| Interruption groove UI2 % | 54.7       | 46.5     | 32.8     | 36.4     | 33.3     | 46.2     | 33.7     | 25.0     | 44.6     | 35.3     | 27.8     | 30.0     | 0.0      | 19.2     | 25.0     | 35.0     | 17.7     | 31.0     | 45.8     | 45.0     |
| (+ = ASU 2–6) n      | 117        | 71       | 67       | 22       | 6        | 26       | 86       | 8        | 83       | 17       | 18       | 20       | 5        | 146      | 12       | 20       | 17       | 71       | 24       | 20       |
| Tuberculum dentale UI2 % | 38.2       | 31.5     | 22.1     | 24.3     | 33.3     | 33.3     | 21.2     | 11.1     | 9.6      | 33.4     | 38.9     | 22.6     | 20.0     | 36.8     | 16.7     | 5.3      | 50.0     | 17.6     | 125      | 33.5     |
| (+ = ASU 2–6) n      | 141        | 73       | 68       | 33       | 6        | 21       | 85       | 9        | 83       | 18       | 18       | 22       | 5        | 144      | 12       | 19       | 18       | 74       | 24       | 21       |
| Bushman canine UC %  | 0.0        | 0.0      | 2.4      | 3.7      | 0.0      | 0.0      | 9.3      | 0.0      | 4.7      | 4.3      | 0.0      | 111      | 0.0      | 19       | 0.0      | 8.7      | 0.0      | 2.7      | 0.0      | 4.2      |
| (+ = ASU 1–3) n      | 96         | 69       | 85       | 27       | 9        | 24       | 97       | 19       | 86       | 23       | 24       | 27       | 7        | 154      | 24       | 23       | 23       | 73       | 29       | 24       |
| Hypocone UM2 %       | 82.1       | 38.4     | 73.4     | 68.5     | 73.0     | 63.6     | 87.3     | 80.8     | 72.7     | 80.9     | 84.4     | 89.8     | 93.1     | 87.4     | 75.4     | 50.0     | 59.6     | 66.0     | 65.7     | 54.0     |
| (+ = ASU 3–5) n      | 112        | 73       | 124      | 54       | 26       | 55       | 157      | 52       | 99       | 42       | 45       | 79       | 29       | 214      | 73       | 54       | 47       | 94       | 35       | 37       |
| Cusp 5 UM1 %         | 5.4        | 1.2      | 3.0      | 15.0     | 12.9     | 0.0      | 23.2     | 37.5     | 12.2     | 35.0     | 23.9     | 6.6      | 77.0     | 6.3      | 5.7      | 23.3     | 3.2      | 0.0      | 12.9     |
| (+ = ASU 2–5) n      | 94         | 82       | 101      | 40       | 31       | 21       | 112      | 40       | 90       | 40       | 46       | 60       | 13       | 183      | 64       | 53       | 43       | 94       | 22       | 31       |
| Carabelli’s trait UM1 % | 47.8       | 11.5     | 31.4     | 24.0     | 23.1     | 10.4     | 39.9     | 30.6     | 27.0     | 41.8     | 40.4     | 44.2     | 33.4     | 37.8     | 27.4     | 58.3     | 512      | 40.2     | 40.9     | 14.7     |
| (+ = ASU 3–7) n      | 157        | 78       | 118      | 54       | 39       | 48       | 143      | 49       | 100      | 43       | 47       | 70       | 15       | 207      | 66       | 48       | 43       | 87       | 22       | 34       |
| Parastyle UM3 %      | 0.0        | 0.0      | 0.0      | 0.0      | 0.0      | 0.0      | 1.7      | 2.7      | 2.1      | 2.9      | 0.0      | 5.2      | 3.3      | 20.0     | 0.0      | 0.0      | 0.0      | 4.5      | 3.7      |
| Trait/grades present | D13  | D14 | D15  | D16   | D17   | D18  | D19 | D20  | D21  | D22  | D23  | D24  | D25  | D26 | D27 | D28  | D29  | D30 | D31 | D32 |
|---------------------|------|-----|------|-------|-------|------|-----|------|------|------|------|------|------|-----|-----|------|------|-----|-----|-----|
|                     | _PIM | _ALE| _KAZ | _MON  | _LOK  | _THA | _VIE| _JAP | _MAL | _PHI | _BOR | _AUN | _NBR | _NEP | _GRK| _ITY | _KBR | _FIN | _LAP|
| (+ = ASU 3–5)       | n    | 26  | 35   | 79    | 38    | 8    | 27  | 116  | 37   | 48   | 35   | 31   | 39   | 30  | 151 | 33   | 33  | 28  | 66  | 22  | 27  |
|                     | %    | 59.2| 55.1 | 40.6  | 83.3  | 69.1 | 78.1| 57.8 | 66.1 | 71.3 | 52.1 | 74.5 | 63.6 | 62.1 | 21.5| 62.7 | 16.7 | 9.3 | 24.1| 16.7| 47.3 |
| Enamel extension UM1|      |     |      |       |       |      |     |      |      |      |      |      |      |      |     |     |      |      |     |     |     |
| (+ = ASU 1–3)       | n    | 27  | 49   | 165   | 66    | 42   | 96  | 166  | 62   | 108  | 48   | 51   | 85   | 37  | 195 | 91   | 54  | 43  | 116 | 30  | 55  |
| Root number UP1     |      |     |      |       |       |      |     |      |      |      |      |      |      |      |     |     |      |      |     |     |     |
| (+ = ASU 2 +)       | n    | 28  | 47   | 170   | 58    | 41   | 115 | 168  | 62   | 93   | 50   | 49   | 133  | 38  | 175 | 94   | 63  | 37  | 100 | 35  | 61  |
| Root number UM2     |      |     |      |       |       |      |     |      |      |      |      |      |      |      |     |     |      |      |     |     |     |
| (+ = ASU 3 +)       | n    | 22  | 30   | 141   | 64    | 34   | 58  | 144  | 58   | 94   | 44   | 50   | 105  | 34  | 184 | 86   | 36  | 22  | 85  | 25  | 49  |
| Odontome P1–P2      | %    | 8.6 | 3.2  | 1.9   | 2.5   | 11.1 | 0.0 | 13   | 2.9  | 7.8  | 0.0  | 5.3  | 2.7  | 0.0 | 18  | 1.9  | 0.0 | 2.2 | 3.3 | 0.0 | 0.0 |
| (+ = ASU +)         | n    | 128 | 94   | 107   | 41    | 9    | 21  | 152  | 34   | 103  | 45   | 38   | 37   | 13  | 168 | 54   | 44  | 46  | 61  | 36  | 27  |
| Congenital absence UM3| %    | 10.3| 30.6 | 29.5  | 33.4  | 44.2 | 18.3| 15.9 | 13.6 | 39.1 | 18.4 | 8.5  | 25.0 | 8.1 | 7.7 | 30.0 | 17.7| 20.0| 19.8| 11.4| 13.0|
| (+ = ASU -)         | n    | 29  | 62   | 156   | 72    | 43   | 104 | 182  | 66   | 110  | 49   | 47   | 104  | 37  | 207 | 90   | 68  | 50  | 111 | 44  | 54  |
| Lingual cusp LP2    | %    | 34.2| 40.3 | 71.4  | 40.0  | 50.0 | 25.0| 69.7 | 54.5 | 63.3 | 81.8 | 71.4 | 83.8 | 100.0| 90.7| 58.3| 60.0| 27.6| 65.1| 48.5| 66.6|
| (+ = ASU 2–9)       | n    | 120 | 72   | 105   | 15    | 2    | 8   | 119  | 22   | 75   | 33   | 28   | 37   | 11  | 172 | 24   | 10  | 29  | 63  | 33  | 45  |
| Groove pattern LM2  | %    | 27.2| 24.1 | 20.7  | 16.0  | 19.2 | 15.4| 19.6 | 26.1 | 38.7 | 38.5 | 31.8 | 30.4 | 43.1 | 15.4| 43.5| 27.9 | 28.4| 13.8| 22.2|
| (+ = ASU Y)         | n    | 82  | 54   | 140   | 25    | 26   | 13  | 148  | 46   | 75   | 39   | 44   | 55   | 23  | 211 | 52   | 23  | 43  | 88  | 29  | 45  |
| Cusp number LM1     | %    | 46.0| 26.2 | 12.9  | 29.4  | 3.2  | 40.0| 36.5 | 29.4 | 34.8 | 34.8 | 35.1 | 34.9 | 77.8| 52.9| 33.3 | 0.0 | 2.6 | 5.9 | 5.6 | 20.0|
| (+ = ASU 6 +)       | n    | 150 | 65   | 116   | 17    | 31   | 5   | 96   | 34   | 69   | 23   | 37   | 43   | 9  | 170 | 39   | 19  | 38  | 85  | 18  | 30  |
| Trait/grades present | D13_PIM | D14_ALE | D15_KAZ | D16_MON | D17_LOK | D18_CHU | D19_THA | D20_VIE | D21_JAP | D22_MAL | D23_PHI | D24_BOR | D25_AUN | D26_NBR | D27_NEP | D28_GRK | D29_ITY | D30_KBR | D31_FIN | D32_LAP |
|----------------------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|---------|
| Cusp number LM2      | %       |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |         |
| (+ = ASU 5+)         | n       | 107     | 68      | 121     | 21      | 23      | 8       | 134     | 45      | 72      | 39      | 42      | 52      | 16      | 193     | 48      | 21      | 35      | 87      | 28      | 39      |
| Protostylid LM1      | %       | 24.4    | 7.6     | 1.6     | 19.1    | 0.0     | 0.0     | 0.7     | 0.0     | 13.0    | 0.0     | 2.4     | 8.1     | 0.0     | 0.0     | 1.9     | 0.0     | 0.0     | 0.0     | 0.0     | 2.4     |
| (+ = ASU 3–6)        | n       | 148     | 66      | 125     | 21      | 38      | 10      | 135     | 39      | 77      | 31      | 42      | 49      | 18      | 45      | 52      | 19      | 38      | 68      | 21      | 38      |
| Cusp 7 LM1           | %       | 6.8     | 3.8     | 2.0     | 0.0     | 10.5    | 0.0     | 2.0     | 4.5     | 1.2     | 3.1     | 7.0     | 11.6    | 0.0     | 9.1     | 3.6     | 5.6     | 2.6     | 2.0     | 0.0     | 0.0     |
| (+ = ASU 2–4)        | n       | 162     | 78      | 149     | 21      | 38      | 12      | 147     | 44      | 82      | 32      | 43      | 60      | 12      | 221     | 56      | 18      | 38      | 99      | 22      | 44      |
| Tome’s root LP1      | %       | 20.0    | 0.0     | 14.1    | 14.3    | 33.3    | 0.0     | 9.9     | 37.5    | 34.9    | 25.8    | 42.9    | 45.1    | 26.1    | 143     | 36.9    | 7.1     | 132     | 13.9    | 100     | 8.8     |
| (+ = ASU 3–5)        | n       | 30      | 27      | 163     | 7       | 6       | 6       | 91      | 16      | 43      | 31      | 21      | 31      | 23      | 126     | 19      | 28      | 38      | 101     | 40      | 45      |
| Root number LC       | %       | 0.0     | 0.0     | 3.4     | 0.0     | 0.0     | 0.0     | 0.6     | 0.0     | 1.7     | 0.0     | 2.3     | 1.7     | 0.0     | 0.0     | 3.4     | 3.5     | 5.1     | 10.6    | 2.3     | 3.2     |
| (+ = ASU 2+)         | n       | 38      | 20      | 175     | 14      | 41      | 17      | 157     | 49      | 58      | 30      | 44      | 58      | 29      | 130     | 59      | 29      | 39      | 104     | 44      | 62      |
| Root number LM1      | %       | 19.2    | 53.1    | 13.9    | 25.0    | 2.4     | 50.0    | 5.9     | 15.4    | 24.2    | 10.0    | 21.6    | 14.1    | 9.1     | 36      | 27.9    | 0.0     | 0.0     | 0.9     | 0.0     | 0.0     |
| (+ = ASU 3+)         | n       | 26      | 49      | 180     | 28      | 42      | 20      | 186     | 52      | 95      | 40      | 51      | 85      | 33      | 193     | 61      | 22      | 25      | 116     | 30      | 60      |
| Root number LM2      | %       | 89.5    | 72.5    | 57.8    | 52.0    | 65.9    | 55.6    | 18.9    | 67.9    | 69.4    | 73.2    | 68.0    | 79.5    | 90.6    | 95.6    | 72.9    | 91.3    | 100.0   | 79.4    | 84.0    | 63.3    |
| (+ = ASU 2+)         | n       | 19      | 40      | 159     | 25      | 41      | 18      | 180     | 53      | 85      | 41      | 50      | 73      | 32      | 184     | 59      | 23      | 38      | 92      | 25      | 60      |

D13_PIM = Pima, D14_ALE = Aleut, D15_KAZ = Kazak, D16_MON = Mongol, D17_LOK = Lower Ob Khanty, D18_CHU = Chukchi, D19_THA = Thailand, D20_VIE = Tonkin-Annam Viet Nam, D21_JAP = Japanese, D22_MAL = Malay, D23_PHI = Philippines, D24_BOR = Borneo, D25_AUN = Australia North, D26_NBR = New Britain, D27_NEP = Nepal, D28_GRK = Greek, D29_ITY = Italy, D30_KBR = Kaberla, D31_FIN = Ladoga Finns, D32_LAP = Lapps. Details in Table 1.

ASUDAS rank-scale trait breakpoint information in Irish (1993, 1997, 2005, 2006), Scott & Turner II (1997), Scott & Irish (2017) and Scott, Turner II, et al., 2018.
The capability of using dental traits as proxies is not insignificant given the cost, destructive sampling, and processing time of genetic analyses. More critically, the traits can be substituted if DNA and, more likely, ancient DNA is not recoverable. Specifically, degradation is of particular concern in tropical and sub-tropical environments, like Africa, Southeast Asia, and other equatorial regions; in more temperate climates time is also a factor, though to a lesser degree, for example, negatively affecting recovery in specimens of Middle Pleistocene age and older (Kistler, Ware, Smith, Collins, & Allaby, 2017; Pinhasi et al., 2015; Smith, Chamberlain, Riley, Stringer, & Collins, 2003). In the latter instance, "paleoanthropologists [currently do] consider teeth the "safe box" of the genetic code" (Martínón-Torres et al., 2007:7), as evidenced by many studies using dental nonmetric traits (Bailey & Hublin, 2013; Bailey, Weaver, & Hublin, 2017; Irish & Guatelli-Steinberg, 2003; Irish, Guatelli-Steinberg, Legge, de Ruiter, & Berger, 2013; Martínón-Torres et al., 2007, 2008, 2013; among others). So, while there is no assurance heritability estimates in recent humans apply to our Plio-Pleistocene ancestors, these dental traits are likely as close as possible to genomic data for determining hominin origins and affinities (also see Irish, Bailey, Guatelli-Steinberg, Delezene, & Berger, 2018).

Lastly, while genomic markers are largely seen as the definitive data for population studies some caution, like that with the dental traits, should be exercised in choice and interpretation. For example, one limitation with the present SNP data is that each locus can only be represented by two alleles, as genetic software programs (see above) generally do not permit input of multi-allelic states. The result is a decrease in among-population variability, especially on a broad global basis. Another potential limitation is bias introduced when using coding positions potentially influenced by selection, that is, not all markers are neutral. Such is the case for rare variant positions like those, that code for diseases. In the present study, such bias is limited because of the large SNP number. Specifically, drift, unlike selection, influences the whole genome so selection effects are relatively few (Kimura, 1968). Moreover, the Human Origins ascertainment used in this study, unlike some other arrays, is not biased by the inclusion of SNPs with medical/clinical interest.

### Analysis

| Matrices compared: | Analysis: | Africa 1 36 traits | Africa 2 25 traits | Global 25 traits |
|--------------------|-----------|-------------------|-------------------|-----------------|
| MMD + DGGeo        | $r_{MMD}$ | .682              | .696              | .710            |
|                    | $p_{MMD}$ | .000              | .000              | .000            |
| $F_{ST} + GGeo$    | $r_{F_{ST}}$ | .486              | .486              | .768            |
|                    | $p_{F_{ST}}$ | .001              | .001              | .000            |
| MMD + $F_{ST}$     | $r_{MMD}$ | .786              | .838              | .720            |
|                    | $p_{MMD}$ | .000              | .000              | .000            |
| MMD + $F_{ST}$ - DGGeo | $r_{MMD}$ | .700              | .782              | .400            |
|                    | $p_{MMD}$ | .000              | .000              | .000            |

Abbreviations: DGGeo, matrix of geographic distances among dental samples; DGGeo, matrix of geographic midpoint distances for matching dental and genetic samples; $F_{ST}$, Hudson $F_{ST}$ distance matrix; GGeo, matrix of geographic distances among genetic samples; MMD, mean measure of divergence matrix.

### 5 CONCLUSION

In sum, the present study is the most comprehensive to date comparing dental nonmetric traits and neutral genomic markers, in terms of the amount of data and number of samples at continental and global levels. The correspondence of these datasets based on comparison of MMD and $F_{ST}$ distance matrices is greater in than any prior studies, likely because of the data and samples, as well as the methods used. Mantel correlations between these distances are all strongly positive, ranging from .72 globally to .84 within Africa. These and inter-sample geographic distance matrices are also strongly correlated, ranging from .68 for the 36-trait MMD in Africa to .77 for $F_{ST}$ globally; the only exception is between $F_{ST}$ and African geographic distances, which though moderate, remains positive and significant, $r_{MMD} = 0.49$. This, and partial correlations between MMD and $F_{ST}$ controlling for geographic distances, namely, high in Africa (.78) and moderate globally (.4), suggest that the genetic distance measure is better able to pick up on recent within-continent variation, while recognizing ancient relationships on a global level.

That said, as mentioned, all correlations may be seen as minimum values in light of several recognized limitations, notably, ASUDAS and SNP data are not from the same individuals across samples. Though prohibitive concerning cost and time, future researchers could follow and expand upon the approach of Hubbard et al. (2015); ideally, both sets of data would be collected from the same skeletal remains in local- through global-level samples. Among other potential advantages over living individuals, actual teeth instead of casts would promote more detailed trait recording and include key root data. The bottom-line then, in conjunction with the recent heritability studies and population analyses, and despite potential concerns (sex dimorphism, selection, etc.), is that dental nonmetric traits actually can and do work well as proxies for neutral genomic data.

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DATA AVAILABILITY STATEMENT
The ASUDAS frequency data used in the analyses are available in Tables 3 and 7 of this article, as well as the relevant references cited in the main text. Many ASUDAS nondichotomized data are available in Scott & Irish (2017). The remaining ASUDAS data from the study are in preparation for publication, and/or are available upon reasonable request from the first or fifth authors. As noted, all SNP data, except the whole genome sequences of the West African Wolof sample, were genotyped with the Affymetrix Human Origins Array (AHOA), available in the relevant references cited in the text.

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REFERENCES
Bailey, S. E., & Hublin, J. J. (2013). What does it mean to be dentally “modern”? In G. R. Scott & J. D. Irish (Eds.), Anthropological Perspectives on Tooth Morphology: Genetics, Evolution, Variation (pp. 222–249). Cambridge: Cambridge University Press.

Bailey, S. E., Weaver, T. D., & Hublin, J. J. (2017). The dentition of the earliest modern humans: How ‘modern’ are they? In A. Marom & E. Hovers (Eds.), Human Paleontology and Prehistory (pp. 215–232). Cham: Springer.

Bermudez de Castro, J. M. (1989). The Carabelli trait in human prehistoric populations of the Canary Islands. Human Biology, 61, 117–131.

Bhatia, G., Patterson, N., Sankararaman, S., & Price, A. L. (2013). Estimating and interpreting FST: the impact of rare variants. Genome Research, 23, 1514–1521.

Brewer-Carias, C. A., Le Blanc, S., & Neel, J. V. (1976). Genetic structure of a tribal population, the Yanomama Indians. XIII. Dental microdifferentiation. American Journal of Physical Anthropology, 44, 5–14.

Bryk, J., Hardouin, E., Pugach, I., Hughes, D., Strotmann, R., Stoneking, M., & Myles, S. (2008). Positive selection in East Asians for an EDAR allele that enhances NF-kB activation. PLoS One, 3, e2209.

Cohen, J. (1988). Set correlation and contingency tables. Applied Psychological Measurement, 4, 425–434.

Danecek, P., Schifflers, S., & Durbin, R. (2014). Multiallelic calling model in bcftools (–m).

Delgado, M., Ramirez, L. M., Adhikari, K., Fuentes-Guajardo, M., Zanoli, C., Gonzalez-José, R., … Rothhammer, F. (2019). Variation in dental morphology and inference of continental ancestry in admixed Latin Americans. American Journal of Physical Anthropology, 168, 438–447.

Diniz-Filho, J. A. F., Soares, T. N., Lima, J. S., Dobrovoljski, R., Landeiro, V. L., Telles, M. P. D. C., … Bini, L. M. (2013). Mantel test in population genetics. Genetics and Molecular Biology, 36, 475–485.

Edgar, H. J. H. (2013). Estimation of ancestry using dental morphological characteristics. Journal of Forensic Sciences, 58, S3–S8.

Edgar, H. J. H. (2015). Dental morphological estimation of ancestry in forensic contexts. In G. E. Berg & S. C. Ta’ala (Eds.), Biological Affinity in Forensic Identification of Human Skeletal Remains (pp. 191–208). Boca Raton, FL: CRC Press.

Ersts, P. J. (2014). Geographic Distance Matrix Generator (version 1.2.3). New York: American Museum of Natural History, Center for Biodiversity and Conservation. Available at: http://biodiversityinformatics.amnh.org/open_source/gdmg, accessed on September 7, 2019.

Gelman, A., Hill, J., & Yajima, M. (2012). Why we (usually) don’t have to worry about multiple comparisons. Journal of Research on Educational Effectiveness, 5, 189–211.

Green, R. F., & Suchey, J. M. (1976). The use of inverse sine transformations in the analysis of non-metric cranial data. American Journal of Physical Anthropology, 45, 61–68.

Greenberg, J. H. (1963). The languages of Africa. The Hague: Mouton & Co.

Gross, J. M., & Edgar, H. J. (2019). Informativeness of dental morphology in ancestry estimation in African Americans. American Journal of Physical Anthropology, 168, 521–529.

Guillot, G., & Rousset, F. (2013). Dismantling the Mantel tests. Methods in Ecology and Evolution, 4, 336–344.

Haeussler, A. M., Turner, C. G., & Irish, J. D. (1988). Concordance of American and Soviet methods in dental anthropology. American Journal of Physical Anthropology, 75, 219.

Hammer, Ø. (2019). PAST: Paleontological Statistics, Version 3.23 Reference Manual. Oslo: Natural History Museum, University of Oslo.

Hammer, Ø., Harper, D. A. T., & Ryan, P. D. (2001). PAST: Paleontological Statistics Software Package for Education and Data Analysis. Palaeontologia Electronica, 4, 1–9.

Hanifa, T. (1992). Dental and cranial affinities among populations of East Asia and the Pacific: the basic populations in East Asia, IV. American Journal of Physical Anthropology, 88, 163–182.

Hanifa, T. (2013). Geographical structure of dental variation in the major human populations of the world. In G. R. Scott & J. D. Irish (Eds.), Anthropological perspectives on tooth morphology: genetics, evolution, variation (pp. 479–509). Cambridge: Cambridge University Press.

Harris, E. F. (1977). Anthropologic and genetic aspects of the dental morphology of Solomon Islanders, Melanesia. PhD dissertation, Arizona State University, Tempe.

Higgins, D., Hughes, T. E., James, H., & Townsend, G. C. (2009). Strong genetic influence on hypocone expression of permanent maxillary molars in South Australian twins. American Journal of Physical Anthropology, 22, 1–7.

Hlusko, L. J., Carlson, J., Chaplin, G., Elias, S. A., Hoffecker, J. F., Huffman, M., … Scott, G. R. (2018). Evidence of environmental selection on the mother-to-infant transmission of vitamin D and fatty acids during the last ice age in Beringia. Proceedings of the National Academy of Sciences, 115, e4426–e4432.
Holsinger, K. E., & Weir, B. S. (2009). Genetics in geographically structured populations: defining, estimating and interpreting Fst. Nature Reviews Genetics, 10, 639.

Hubbard, A.R. (2012). An examination of population history, population structure, and biological distance among regional populations of the Kenyan coast using genetic and dental data. Ph.D. Dissertation. Ohio State University, Columbus.

Hubbard, A. R., Guatelli-Steinberg, D., & Irish, J. D. (2015). Do nuclear DNA and dental nometric data produce similar reconstructions of regional population history? An example from modern coastal Kenya. American Journal of Physical Anthropology, 157, 295–304.

Hudson, R. R., Slatkin, M., & Maddison, W. P. (1992). Estimation of levels of gene flow from DNA sequence data. Genetics, 132, 583–589.

Hughes, T.E., & Townsend, G.C. (2011). Twin studies of dental crown morphology: genetic and environmental determinants of the Cusp of Carabelli. In: Program 15th International Symposium on Dental Morphology, 37, Newcastle, UK, 25 August.

Hughes, T. E., & Townsend, G. C. (2013). Twin and family studies of human dental crown morphology: genetic, epigenetic, and environmental determinants of the modern human dentition. In G. R. Scott & J. D. Irish (Eds.), Anthropological perspectives on tooth morphology: genetics, evolution, variation (pp. 31–68). Cambridge: Cambridge University Press.

Hughes, T., Townsend, G., & Bockmann, M. (2016). An overview of dental genetics. In J. D. Irish & G. R. Scott (Eds.), A companion to dental anthropology (pp. 123–141). New York: Wiley-Blackwell.

Irish, J.D. (1993). Biological affinities of late Pleistocene through modern African aboriginal populations: the dental evidence. Ph.D. Dissertation. Arizona State University, Tempe.

Irish, J. D. (1997). Characteristic high- and low-frequency dental traits in sub-Saharan African populations. American Journal of Physical Anthropology, 102, 455–467.

Irish, J. D. (1998a). Diachronic and synchronic dental trait affinities of Late and post-Pleistocene peoples from North Africa. Homo, 49, 138–155.

Irish, J. D. (1998b). Ancestral dental traits in recent sub-Saharan Africans and the origins of modern humans. Journal of Human Evolution, 34, 81–98.

Irish, J. D. (2000). The Iberomaurusian enigma: North African progenitor or dead end? Journal of Human Evolution, 39, 392–410.

Irish, J. D. (2005). Population continuity versus discontinuity revisited: dental affinities among Late Paleolithic through Christian era Nubians. American Journal of Physical Anthropology, 128, 520–535.

Irish, J. D. (2006). Who were the ancient Egyptians? Dental affinities among Neolithic through post-dynastic peoples. American Journal of Physical Anthropology, 129, 529–543.

Irish, J. D. (2010). The mean measure of divergence (MMD): its utility in model-free and model-bound analyses relative to the Mahalanobis D2 distance for nonmetric traits. American Journal of Human Biology, 22, 378–395.

Irish, J. D. (2015). Dental nonmetric variation around the world: Using key traits in populations to estimate ancestry in individuals. In G. E. Berg & S. C. Ta’ala (Eds.), Biological Affinity in Forensic Identification of Human Skeletal Remains (pp. 165–190). Boca Raton, FL: CRC Press.

Irish, J. D. (2016). Who were they really? Model-free and model-bound dental nonmetric analyses to affirm documented population affiliations of seven South African “Bantu” samples. American Journal of Physical Anthropology, 159, 655–670.

Irish, J. D., Bailey, S. E., Guatelli-Steinberg, D., Delezene, L. K., & Berger, L. R. (2018). Ancient teeth, phonetic affinities, and African hominins: Another look at where Homo naledi fits in. Journal of Human Evolution, 122, 108–123.

Irish, J. D., Black, W., Sealy, J., & Ackermann, R. R. (2014). Questions of Khoesan continuity: dental affinities among the indigenous Holocene peoples of South Africa. American Journal of Physical Anthropology, 155, 33–44.

Irish, J. D., & Guatelli-Steinberg, D. (2003). Ancient teeth and modern human origins: an expanded comparison of African Plio-Pleistocene and recent world dental samples. Journal of Human Evolution, 45, 113–144.

Irish, J. D., Guatelli-Steinberg, D., Legge, S. S., de Ruiter, D. J., & Berger, L. R. (2013). Dental morphology and the phylogenetic “place” of Australopithecus sediba. Science, 340, 1233062.

Irish, J. D., Lillios, K. T., Waterman, A. J., & Silva, A. M. (2017). “Other” possibilities? Assessing regional and extra-regional dental affinities of populations in the Portuguese Estremadura to explore the roots of Iberia’s Late Neolithic-Copper Age. Journal of Archaeological Science: Reports, 11, 224–236.

Irish, J. D., & Morris, D. H. (1996). A supplemental description of the Bushman mandibular canine polymorphism. Journal of Human Evolution, 31, 351–353.

Irish, J. D., & Turner, C. G., II. (1990). West African dental affinity of late Pleistocene Nubians: peopling of the Eurafrican-South Asian triangle II. Homo, 41, 42–53.

Jay, F., Sjödin, P., Jakobsson, M., & Blum, M. G. (2012). Anisotropic isolation by distance: the main orientations of human genetic differentiation. Molecular Biology and Evolution, 30, 513–525.

Jernvall, J., & Jung, J.-S. (2000). Genotype, phenotype, and developmental biology of molar tooth characters. Yearbook of Physical Anthropology, 48, 171–190.

Kanitz, R., Guillot, E. G., Antoniazza, S., NeuenSchwander, S., & Goudet, J. (2018). Complex genetic patterns in human arise from a simple range-expansion model over continental landmasses. PLoS One, 13, e0192460.

Kimura, M. (1968). Evolutionary rate at the molecular level. Nature, 217, 624–626.

Kimura, R., Yamaguchi, T., Takeda, M., Kondo, O., Toma, T., Haneji, K., ... Osawa, M. (2009). A common variation in EDAR is a genetic determinant of shovel-shaped incisors. The American Journal of Human Genetics, 85, 528–535.

Küster, L., Ware, R., Smith, O., Collins, M., & Allaby, R. G. (2017). A new model for ancient DNA decay based on paleogenomic meta-analysis. Nucleic Acids Research, 45, 6310–6320.

Konigsberg, L.W. (2006). A post-Neumann history of biological and genetic distance studies in bioarchaeology. In J.E. Buikstra & L.A. Beck LA (Eds.), Bioarchaeology: the contextual analysis of human remains (pp. 262–279). New York: Academic Press.

Konigsberg, L. W. (1990). Analysis of prehistoric biological variation under a model of isolation by geographic and temporal distance. Human Biology, 62, 49–70.

Lazaridis, I., Patterson, N., Mittnik, A., Renaud, G., Mallick, S., Li, H., Lipson, M., Mathieson, I., Gymrek, M., Racimo, F., ... Skoglund, P. (2016). The Simons genome diversity project: 300 genomes from 142 diverse populations. Nature, 538, 201.
Smith, C. I., Chamberlain, A. T., Riley, M. S., Stringer, C., & Collins, M. J. (2003). The thermal history of human fossils and the likelihood of successful DNA amplification. *Journal of Human Evolution, 45*, 203–217.

Smith, P., & Shegev, M. (1988). The dentition of Nubians from Wadi Halfa, Sudan: an evolutionary perspective. *Journal of the Dental Association of South Africa, 43*, 539–541.

Smouse, P. E., & Long, J. C. (1992). Matrix correlation analysis in anthropology and genetics. *American Journal of Physical Anthropology, 35* (S15), 187–213.

Smouse, P. E., Long, J. C., & Sokal, R. R. (1986). Multiple regression and correlation extensions of the Mantel test of matrix correspondence. *Systematic Zoology, 35*, 627–632.

Sofaer, J. A., Niswander, J. D., MacLean, C. J., & Workman, P. L. (1972). Population studies on Southwestern Indian tribes V. Tooth morphology as an indicator of biological distance. *American Journal of Physical Anthropology, 37*, 357–366.

Sofaer, J. A., Smith, P., & Kaye, E. (1986). Affinities between contemporary and skeletal Jewish and non-Jewish groups based on tooth morphology. *American Journal of Physical Anthropology, 70*, 265–275.

Sokal, R. R., & Rohlf, F. J. (1995). Biometry. New York: Freeman and Company.

Stojanowski, C. M., & Paul, K. S. (2015). Performance analysis of deciduous morphology for detecting biological siblings. *American Journal of Physical Anthropology, 157*, 615–629.

Stojanowski, C. M., Paul, K. S., Seidel, A. C., Duncan, W. N., & Guatelli-Steinberg, D. (2018). Heritability and genetic integration of anterior tooth crown variants in the South Carolina Gullah. *American Journal of Physical Anthropology, 167*, 124–143.

Stojanowski, C. M., Paul, K. S., Seidel, A. C., Duncan, W. N., & Guatelli-Steinberg, D. (2019). Quantitative genetic analyses of postcanine morphological crown variation. *American Journal of Physical Anthropology, 168*, 606–631.

Stojanowski, C. M., & Schillaci, M. A. (2006). Phenotypic approaches for understanding patterns of intracemetery biological variation. *Yearbook of Physical Anthropology, 13*(suppl. 43), 49–88.

Tian, C., Kosoy, R., Nasir, R., Lee, A., Villoslada, P., Klareskog, L., ... Gregersen, P. K. (2009). European population genetic substructure: further definition of ancestry informative markers for distinguishing among diverse European ethnic groups. *Molecular Medicine, 15*, 371–383.

Turner, C. G., II (1985a). The dental search for Native American origins. In R. Kirk & E. Szathmary (Eds.), *Out of Asia* (pp. 31–78). Canberra: The Journal of Pacific History.

Turner, C. G., II. (1985b). Expression count: a method for calculating morphological dental trait frequencies by using adjustable weighting coefficients. *American Journal of Physical Anthropology, 68*, 263–268.

Turner, C. G. (1990). Major features of Sundadonty and Sinodonty, including suggestions about East Asian microevolution, population history, and late Pleistocene relationships with Australian aboriginals. *American Journal of Physical Anthropology, 82*, 295–317.

Turner, C. G., II, Nichol, C. R., & Scott, G. R. (1991). Scoring procedures for key morphological traits of the permanent dentition: The Arizona State University Dental Anthropology System. In M. A. Kelley & C. S. Larsen (Eds.), *Advances in dental anthropology* (pp. 13–31). New York: Wiley-Liss.

Weir, B. S., & Cockerham, C. C. (1984). Estimating F-statistics for the analysis of population structure. *Evolution, 38*, 1358–1370.

Weir, B. S., Cardon, L. R., Anderson, A. D., Nielsen, D. M., & Hill, W. G. (2005). Measures of human population structure show heterogeneity among genomic regions. *Genome Research, 15*, 1468–1476.

Willing, E.-M., Dreyer, C., & van Oosterhout, C. (2012). Estimates of genetic differentiation measured by FST do not necessarily require large sample sizes when using many SNP markers. *PLoS One, 7*, e42649.

Wright, S. (1943). Isolation by distance. *Genetics, 28*, 114.

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of this article.

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