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Genetic origins of the Minoans and Mycenaeans

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Data availability

The aligned sequences are available through the European Nucleotide Archive under accession number PRJEB20914. Genotype datasets used in analysis are at <https://reich.hms.harvard.edu/datasets>.

Author Contributions

G.S. conceived the study. D.R. and J.K. co-supervised the ancient DNA work, sequencing, and data analysis. I.L. performed population genetic analysis and wrote the manuscript with input from other authors. P.J.P.McG, E.K, G.K., H.M., M.M, M.Ö, N.Ö, A.Pa., M.R., S.A.R., Y.T, A.V., R.A. P.S., R.P., J.K., G.S. assembled, studied, or described archaeological and osteological material. J.R.H, P.A.N., G.S., D.R. assembled samples for genotyping. A.M., S.P., N.R., A.F., C.P., D.F., J.R.H, D.M.L, Y.M, J.A.S, K.St., R.P., G.S., D.R., J.K. performed wet lab work. A.M., N.P., S.M., A.Pe., performed bioinformatic analyses.

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Abstract

The origins of the Bronze Age Minoan and Mycenaean cultures have puzzled archaeologists for more than a century. We assembled genome-wide data from nineteen ancient individuals, including Minoans from Crete, Mycenaeans from mainland Greece, and their eastern neighbours from southwestern Anatolia. We show that Minoans and Mycenaeans were genetically similar, having at least three quarters of their ancestry from the first Neolithic farmers of western Anatolia and the Aegean^{1,2}, and most of the remainder from ancient populations like those of the Caucasus³ and Iran^{4,5}. However, the Mycenaeans differed from Minoans in deriving additional ancestry from an ultimate source related to the hunter-gatherers of eastern Europe and Siberia^{6–8}, introduced via a proximal source related to either the inhabitants of either the Eurasian steppe^{1,6,9} or Armenia^{4,9}. Modern Greeks resemble the Mycenaeans, but with some additional dilution of the early Neolithic ancestry. Our results support the idea of continuity but not isolation in the history of populations of the Aegean, before and after the time of its earliest civilizations.

Ancient DNA research has traced the principal ancestors of early European farmers to highly similar Neolithic populations of Greece and western Anatolia, beginning in the 7th millennium BCE^{1,2}, but the later history of these regions down to the Bronze Age, a transformational period in the history of Eurasia^{4,6,9}, is less clear. There is limited genetic evidence suggesting migrations from both the east (the area of Iran and the Caucasus), reaching Anatolia by at least ~3,800 BCE⁴, and the north (eastern Europe and Siberia) contributing ‘Ancient North Eurasian’ ancestry^{6,10} to all modern Europeans. The timing and impact of these migrations in the Aegean is, however, unknown.

During the Bronze Age, two prominent archaeological cultures emerged in the Aegean. The culture of the island of Crete, labelled ‘Minoan’ by Arthur Evans¹¹, was Europe’s first literate civilization, and has been described as ‘Europe’s first major experience of civilization’¹². However, the Linear A syllabic ideographic and Cretan hieroglyphic scripts used by this culture remain undeciphered, obscuring its origins. Equally important was the civilization of the ‘Mycenaean’ culture of mainland Greece, whose language, written in the Linear B script, was an early form of Greek¹³. Cretan influence in mainland Greece and the

later Mycenaean occupation of Crete link these two cultures, but the degree of genetic affinity between mainland and Cretan populations is unknown. Greek is related to other Indo-European languages, leading to diverse theories tracing its earliest speakers from the 7th millennium down to ~1,600 BCE, and proposing varying degrees of population change (Supplementary Information, section 1).

Genome-wide ancient DNA data provides a new source of information about the people of the Bronze Age, who were first known through the ancient poetic and historical traditions commencing with Homer and Herodotus, later through the disciplines of archaeology and linguistics, and, more recently, by the limited information from ancient mitochondrial DNA^{14,15}. Here we answer several questions. First, do the labels ‘Minoan’ and ‘Mycenaean’ correspond to genetically coherent populations or do they obscure a more complex structure of the peoples who inhabited Crete and mainland Greece at this time? Second, how were the two groups related to each other, to their neighbours across the Aegean in Anatolia, and to other ancient populations from Europe^{1,2,6,8–10} and the Near East^{2–5,9,16,17}? Third, can inferences about their ancestral origins inform debates about the origins of their cultures? Fourth, how are the Minoans and Mycenaeans related to Modern Greeks, who inhabit the same area today?

We generated genome-wide data from 19 ancient individuals (Fig. 1a; Extended Data Table 1; Supplementary Information, section 1). This includes 10 Minoans from Crete, (~2,900–1,700 BCE; labelled Minoan_Odigitria: from Moni Odigitria near the southern coast of central Crete and Minoan_Lasithi: from the cave of Hagios Charalambos in the highland plain of Lasithi in east Crete). From mainland Greece, 4 Mycenaeans were included (~1,700–1,200 BCE; from the western coast of the Peloponnese, from Argolis, and the island of Salamis). An additional individual from Armenoi in western Crete (~1,370–1,340 BCE; labelled Crete_Armenoi) postdates the appearance of Mycenaean culture on the island of Crete. Our dataset also includes a Neolithic sample from Alepotrypa Cave at Diros bay in the southern Peloponnese (~5,400 BCE) adding to previously published samples from northern Greece² (collectively labelled Greece_N). Finally, it includes 3 Bronze Age individuals (~2,800–1,800 BCE; labelled Anatolia_BA) from Harmanören Göndürle in southwestern Anatolia (Turkey), adding knowledge about genetic variation in Anatolia after the Neolithic/Chalcolithic periods^{1,2,4,17} (Supplementary Information, section 1). We processed the ancient remains, extracted DNA, and prepared Illumina libraries in dedicated clean rooms (Supplementary Data Table 1; Methods), and, after initial screening for mitochondrial DNA, used in-solution hybridization¹⁸ to capture ~1.2 million single nucleotide polymorphisms^{6,19} on the ancient samples. We assessed contamination by examining the rate at which they matched the mitochondrial consensus sequence (Supplementary Data Table 2) and by the rate at which male samples were heterozygous on the X-chromosome (Methods). We combined the dataset of the 19 ancient individuals with 332 other ancient individuals from the literature, 2,614 present-day humans genotyped on the Human Origins array, and 2 present-day Cretans (Methods).

We carried out principal components analysis²⁰ (Methods), projecting ancient samples onto the first two principal components inferred from present-day West Eurasian populations¹⁰ that form two south-north parallel clines in Europe and the Near East along PC2. Minoans

and Mycenaean populations are centrally positioned in the PCA (Fig. 1b), framed to the left by ancient populations from mainland Europe and the Eurasian steppe, to the right by ancient populations from the Caucasus and Western Asia, and to the bottom by Early/Middle Neolithic farmers from Europe and Anatolia. The Neolithic samples from Greece cluster with these farmers and are distinct from the Minoans and Mycenaean populations. The Bronze Age individuals from southwestern Anatolia are also distinct, intermediate between Anatolian and Levantine populations towards the bottom, and populations from Armenia, Iran, and the Caucasus towards the top. ADMIXTURE analysis (Extended Data Fig. 1) shows that both Minoans and Mycenaean populations possess a ‘pink’ genetic component (K=8 and greater) as do Bronze Age southwestern Anatolians, Neolithic Central Anatolians from Tepecik-Çiftlik¹⁷, a Chalcolithic northwestern Anatolian¹, and western Anatolians from Kumtepe¹⁶. This component is maximized in the Mesolithic/Neolithic samples from Iran^{4,5} and hunter-gatherers from the Caucasus³ (Extended Data Fig. 1). It is not found in the Neolithic of northwestern Anatolia, Greece, or the Early/Middle Neolithic populations of the rest of Europe, only appearing in the populations of the Late Neolithic/Bronze Age in mainland Europe⁶, introduced there by migration from the Eurasian steppe^{1,6}.

Beyond the visual impressions of PCA and ADMIXTURE, we formally tested the relationships between populations from our study and the literature, using f_4 -statistics of the form $f_4(X, Y; Test, Chimp)$ that evaluate whether *Test* shares more alleles with *X* or *Y*. We find that *Test* populations from Iran, the Caucasus, and eastern Europe share more alleles with Minoans and Mycenaean populations than with the Neolithic population of Greece (Extended Data Fig. 2a,b). The Minoans from the Lasithi plateau in the highlands of eastern Crete and from the coast of southern Crete (Extended Data Fig. 2c) are consistent with being a homogeneous population. Mycenaean populations differ from these Minoans in sharing significantly fewer alleles with Neolithic people from the Levant, Anatolia, Greece, and mainland Europe (Extended Data Fig. 2d). In comparison, the Bronze Age Anatolians share fewer alleles with ancient Europeans and more with ancient populations of Iran and the Levant (Extended Data Fig. 3). We used f_3 -statistics of the form $f_3(Ref_1, Ref_2; Test)$ which, if negative, show that *Test* is admixed from sources related to the *Ref_1*, *Ref_2* source populations. We do not find significantly negative (*Ref_1*, *Ref_2*) pairs for Minoans or Bronze Age Anatolians ($Z > -2.5$), but do for Mycenaean populations ($-4.9 < Z < -3.0$; Extended Data Fig. 4), involving early farmers from the Levant, Anatolia, Greece, and the rest of Europe as one source, and Iran or the Eurasian steppe or steppe-influenced Europeans as the other.

We modelled Bronze Age populations using *qpAdm*/*qpWave*⁶ framework (Methods; Supplementary Information, section 2) which relates a set of ‘left’ populations (admixed population and ancestral source populations) with a set of ‘right’ populations (diverse outgroups) and allows one to test for the number of streams of ancestry from ‘right’ to ‘left’ and to estimate admixture proportions. This analysis shows that all Bronze Age populations from the Aegean and Anatolia derived most (~62–86%) of their ancestry from an Anatolian Neolithic-related population (Table 1). However, they also had a component (~9–32%) of ‘eastern’ (Caucasus/Iran-related) ancestry. It was previously shown that this type of ancestry was introduced into mainland Europe via Bronze Age pastoralists from the Eurasian steppe who were a mix of both eastern European hunter-gatherers and populations from the Caucasus and Iran^{4,6}; our results show that it also arrived on its own, at least in the Minoans, without

eastern European hunter-gatherer ancestry. This ancestry need not have arrived from regions east of Anatolia, as it was already present during the Neolithic in central Anatolia at Tepecik-Çiftlik¹⁷ (Supplementary Information, section 2). The eastern influence in the Bronze Age populations from Greece and southwestern Anatolia is also supported by an analysis of their Y-chromosomes. Four out of five males belonging to Minoans, Mycenaeans, and southwestern Anatolians (Supplementary Information, section 3) belonged to haplogroup J which was rare or non-existent in earlier populations from Greece and western Anatolia which were dominated by Y-chromosome haplogroup G2^{1,2,17}. Haplogroup J was present in Caucasus hunter-gatherers³ and a Mesolithic individual from Iran⁴ and its spread westward may have accompanied the ‘eastern’ genome-wide influence.

The Minoans could be modelled as a mixture of the Anatolia Neolithic-related substratum with additional ‘eastern’ ancestry, but the other two groups had additional ancestry: the Mycenaeans had ~4–16% ancestry from a ‘northern’ ultimate source related to the hunter-gatherers of eastern Europe and Siberia (Table 1), while the Bronze Age southwestern Anatolians may have had ~6% ancestry related to Neolithic Levantine populations. The elite Mycenaean individual from the ‘royal’ tomb at Peristeria in the western Peloponnese did not differ genetically from the other three Mycenaean individuals buried in common graves. To identify more proximate sources of the distinctive eastern European/north Eurasian-related ancestry in Mycenaeans, we included later populations as candidate sources (Supplementary Information, section 2), and could model Mycenaeans as a mixture of the Anatolian Neolithic and Chalcolithic-to-Bronze Age populations from Armenia (Table 1). Populations from Armenia possessed some ancestry related to eastern European hunter-gatherers⁴, so they, or similar unsampled populations of western Asia, could have contributed it to populations of the Aegean. This model makes geographical sense, since a population movement from the vicinity of Armenia could have admixed with Anatolian Neolithic-related farmers on either side of the Aegean. However, Mycenaeans can also be modelled as a mixture of Minoans and Bronze Age steppe populations (Table 1; Supplementary Information, section 2), suggesting that, alternatively, ‘eastern’ ancestry arrived in both Crete and mainland Greece, followed by ~13–18% admixture with a ‘northern’ steppe population in mainland Greece only. Such a scenario is also plausible: first, it provides a genetic correlate for the distribution of shared toponyms in Crete, mainland Greece, and Anatolia discovered by Kretschmer²¹; second, it postulates a single migration from the east; third, it proposes some gene flow from geographically contiguous areas to the north where steppe ancestry was present since at least the mid-3rd millennium BCE^{6,9}. We validated inferences from *qpAdm* by treating source populations as ‘ghosts’ and re-estimating mixture proportions⁴, by examining the correspondence between *qpAdm* estimates and PCA⁴ (Extended Data Fig. 5), and by comparing simulated individuals of known ancestry against the Mycenaeans (Extended Data Fig. 6).

Geographical structure may have prevented the spread of the ‘northern’ ancestry from the mainland to Crete, contributing to genetic differentiation. Such structure may, in principle, be long-standing, even prior to the advent of the Neolithic in the 7th millennium BCE. Alternatively, both ‘northern’ and ‘eastern’ ancestry may have arrived in the Aegean at any time between the Early Neolithic and the Late Bronze Age. Wider geographical and temporal sampling of pre-Bronze Age populations of the Aegean may better trace the advent

of ‘northern’ and ‘eastern’ ancestry in the region. However, sampled Neolithic samples from Greece, down to the Final Neolithic ~4,100 BCE², do not possess either type of ancestry, suggesting that the admixture we detect probably occurred during the 4th–2nd millennium BCE time window. Other proposed migrations, such as settlement by Egyptian or Phoenician colonists²² are not discernible in our data, as there is no measurable Levantine or African influence in the Minoans and Mycenaeans, thus rejecting the hypothesis that the cultures of the Aegean were seeded by migrants from the old civilizations of these regions. On the other hand, migrants from areas east or north of the Aegean, while numerically less influential than the locals, may have contributed to the emergence of the 3rd–2nd millennium BCE Bronze Age cultures as ‘creative disruptors’ of local traditions, bearers of innovations, or through cultural interaction with the locals, coinciding with the genetic process of admixture.²³ Relative ancestral contributions do not determine the relative roles in the rise of civilization of the different ancestral populations, but, nonetheless, the strong persistence of the Neolithic substratum does suggest a key role for the locals in this process.

Phenotype prediction from genetic data has enabled the reconstruction of the appearance of ancient Europeans^{1,24} who left no visual record of their pigmentation. By contrast, the appearance of the Bronze Age people of the Aegean has been preserved in colourful frescos and pottery, depicting people with mostly dark hair and eyes²⁵. We used the HIrisPlex²⁶ tool (Supplementary Information, section 4) to infer that the appearance of our ancient samples matched the visual representations (Extended Data Table 2), suggesting that art of this period reproduced phenotypes naturalistically.

We estimated F_{ST} of Bronze Age populations with present-day West Eurasians, finding that Mycenaeans are least differentiated from populations from Greece, Cyprus, Albania, and Italy (Fig. 2), part of a general pattern in which Bronze Age populations broadly resemble present-day inhabitants from the same region (Extended Data Fig. 7). Modern Greeks occupy the intermediate space of the PCA along PC1 (Fig. 1b) between ancient European and Near Eastern populations, like the ones of the Bronze Age. They are not, however, identical to Bronze Age populations, as they are above them along PC2 (Fig. 1b). This is due to the fact that Neolithic farmers share fewer alleles with Modern Greeks than with Mycenaeans (Extended Data Fig. 8), consistent with additional later admixture^{27,28}.

The Minoans and Mycenaeans, sampled from different sites in Crete and mainland Greece, were homogeneous, supporting the genetic coherency of these two groups. Differences between them are only relative, viewed against their broad overall similarity to each other and to the southwestern Anatolians, sharing in both the ‘local’ Anatolian Neolithic-like farmer ancestry and the ‘eastern’ Caucasus-related admixture. Two key questions remain to be addressed by future studies. First, when did the common ‘eastern’ ancestry of both Minoans and Mycenaeans arrive in the Aegean? Second, is the ‘northern’ ancestry in Mycenaeans due to sporadic infiltration of Greece, or the result of a rapid migration as in Central Europe⁶? Such a migration would support the idea that Proto-Greek speakers²⁹ formed the southern wing of a steppe intrusion of Indo-European speakers. Yet, the absence of ‘northern’ ancestry in the Bronze Age samples from Pisidia, where Indo-European languages were attested in antiquity, casts doubt on this genetic-linguistic association, with further sampling of ancient Anatolian speakers needed. Whatever the answer to these

questions, the discovery of at least two migration events into the Aegean in addition to the first farming dispersal and before the Bronze Age, and of additional population change since that time, supports the view that the Greeks did not emerge fully-formed from the depths of prehistory, but were, indeed, a people ‘ever in the process of becoming.’³⁰

Methods

No statistical methods were used to predetermine sample size. The experiments were not randomized and the investigators were not blinded to allocation during experiments and outcome assessment.

Ancient DNA

An overview of which steps in processing the ancient samples were undertaken in which lab is provided in Supplementary Data Table 1.

Dublin—The inner ear area of each petrous bone was identified, isolated, and then ground to a fine powder. Cleaning and isolation of the cochlea was performed using aluminum oxide powder in a sandblasting chamber. Once isolated, it was decontaminated by UV irradiation for 7.5 minutes on each side, ground on a mixer mill to a weight of about 50mg, and finally transferred to a sterile Eppendorf tube. All procedures were conducted in clean and dedicated ancient DNA facilities.

Seattle—Teeth processed in this laboratory were decontaminated and pulverized to powder in clean and dedicated ancient DNA facilities following previously published methods¹¹.

Leipzig—As previously described,³² sampling, extraction and preparation of double-indexed, double-stranded libraries took place in the clean room facilities of the Max Planck Institute for Evolutionary Anthropology, Leipzig, Germany (MPI-EVA), followed by enrichment of human mtDNA³³. Enriched libraries were sequenced on an Illumina GAIIx platform for 2×76+7 cycles and resulting data was mapped to the rCRS using the EAGER pipeline to evaluate DNA preservation (Supplementary Table 2). These libraries were then shipped to Boston, where nuclear target enrichment was performed (see below).

Tübingen—Pre-PCR steps took place in the clean room facilities of the Institute for Archaeological Sciences at the University of Tübingen, Germany. After surface irradiation with UV-light, the tooth was sawed apart transversally at the border of crown and root, and dentine powder from the inside the crown was sampled using a sterile dentistry drill. Extraction, library preparation and enrichment of human mtDNA used the same protocols as described for MPI-EVA, with addition of a updated extraction protocol³⁴. Sequencing of shotgun and mtDNA-enriched libraries took place at the facilities of the Frauenklinik of the University of Tübingen, on an Illumina MiSeq for 2×150+8 cycles or on an Illumina HiSeq2500 for 2×101+8 cycles (Supplementary Table 2).

Additional libraries were produced including full or partial³⁵ repair with UDG and endonuclease VIII to remove deaminated bases. In-solution enrichment was performed using previously reported protocols^{6,18}. Two SNP sets of 394,577 SNPs (390k capture⁶) or

1,237,207 SNPs (1240k capture¹) were targeted. Sequencing took place in the facilities of the Frauenklinik, University of Tübingen, on an Illumina HiSeq2500 for 2×101+8 cycles and at the facilities of the University of Kiel on a HiSeq4000 for 2×150+8 cycles. One UDG-treated library (I0071) was sent to Boston for nuclear target enrichment, see below.

Boston—The bone powders, prepared from petrous bones in Dublin, were sent to Boston, where DNA extractions and barcoded library preparations without Uracil-removal was performed in the HMS cleanroom following previously described protocols^{34–36}. At the screening stage, libraries were (a) shotgun sequenced, and (b) sequenced after enriching for the human mitochondrial DNA³⁷ together with some nuclear loci in order to approximate the nuclear coverage and mitochondrial contamination. All four libraries (barcoded) prepared in Boston, three libraries (indexed) prepared in Leipzig and one library (indexed) prepared in Tuebingen, were used to perform 390k⁶ and 840k¹⁹ or 1240k (= 390k+840k) targeted capture of a total of 1,233,013 SNPs, following the in-solution target enrichment protocol in Fu et al.¹⁸ and sequenced on either an Illumina HiSeq2500 or Illumina NextSeq500 (see Supplementary Data Table 1 for details).

For each sample, each SNP position is represented by a randomly chosen sequence, restricting to those with a minimum mapping quality (MAPQ≥10), sites with a minimum sequencing quality (≥20), and removing two bases at the ends of reads⁴.

Testing for contamination

Modern human contamination of the mitochondrial DNA was assessed using the software *schmutz*³⁸ which takes into account that the consensus sequence should be reconstructed from reads showing characteristics of ancient DNA and originating from a single individual (Supplementary Data Table 2). We assessed contamination by examining heterozygosity on the X-chromosome in five males (which possess only one copy of the X chromosome) using ANGSD³⁹ (Supplementary Information, section 3); this was in the range of 0.3–4%. Indirect evidence that the females in our dataset (for which X-chromosome based contamination estimation is impossible) are authentic is furnished by their clustering with male samples and distinctiveness from present-day Greek or central European populations that may have possibly contaminated them (Fig. 1b). We also computed f_4 -statistics of the form $f_4(\text{Males, Females; Test, Chimp})$ for populations that had both male and female individuals for all ancient or present-day *Test* populations in our dataset. If female samples were substantially contaminated from a source related to *Test* these statistics would be significantly negative; however we find that the Z-score of these statistics is $-1.6 < Z < 2.5$. We thus included both male and female samples in our analysis to maximize sample size instead of restricting to damaged molecules for females⁸.

Modern human data

We used a dataset of 2,614 individuals genotyped on the Affymetrix Human Origins array^{4,5,10,31}, including 28 Modern Greek (from Greece and Cyprus) samples previously described¹⁰. We also included data from 2 Modern Greeks from Crete whose whole genome sequences were published as part of the Simons Genome Diversity Project⁴⁰. We also

analyzed Modern Greek data from Thessaly and Central Greece⁴¹ and diverse regions^{27,42} genotyped on Illumina arrays.

Datasets

We analyzed two datasets, *HO* which includes the Affymetrix Human Origins genotyping data together with 351 ancient humans (including samples from the literature^{1–5,7–10,16,17,43–51} and the newly reported data) on 591,642 autosomal SNPs and the *HOIII* dataset which does not include the Human Origins data, but has a larger number of 1,054,671 autosomal SNPs⁴. We did not use previously performed genotype calls of literature data, but re-processed them in-house, beginning with the original data release format (FASTQ or BAM). The main analysis dataset was *HOIII* except for analyses that include modern populations in which case the *HO* dataset was analyzed. For the analysis of Illumina genotype data of Modern Greeks (Extended Data Fig. 6) a total of 489,148 autosomal SNPs were analyzed.

Abbreviations used

For brevity, we used the following abbreviations in population names, following the convention of ref.4: CHG: Caucasus hunter-gatherers, EHG: Eastern European hunter-gatherers, WHG: Western European hunter-gatherers, SHG: Scandinavian hunter-gatherers, N: Neolithic, EN: Early Neolithic, MN: Middle Neolithic, ChL: Chalcolithic, LNBA: Late Neolithic/Bronze Age, BA: Bronze Age, EBA: Early Bronze Age, EMBA: Early/Middle Bronze Age, MLBA: Middle/Late Bronze Age, IA: Iron Age.

Principal components analysis

Principal components analysis was performed in the *smartpca* program of EIGENSOFT²⁰, using default parameters and the *lsqproject*: YES¹⁰ and *numoutlieriter*: 0 options. PCA was performed on 1,029 present-day West Eurasians and 334 ancient samples were projected (Fig. 1b); Upper Paleolithic individuals prior to the appearance of the Villabruna cluster⁸ plot in the middle of present-day West Eurasian variation and are not shown.

ADMIXTURE analysis

ADMIXTURE analysis⁵² of the *HO* dataset was performed after pruning for linkage disequilibrium in PLINK^{53,54} with parameters *indep-pairwise* 200 25 0.4, after which 299,971 SNPs were retained. Twenty replicates of the analysis were performed with different random seeds, and the highest likelihood replicate for each value of *K* was retained. We show the *K*=2 to *K*=17 results for the 351 ancient and 30 Modern Greek samples in Extended Data Fig. 1.

f-statistics

f_3 and f_4 -statistics were computed in ADMIXTOOLS³¹ using programs *qp3Pop*, *qpF4ratio* with default parameters, and *qpDstat* with *f4mode*: YES. Standard errors were computed with a block jack-knife⁵⁵. When an ancient population was the target for f_3 -statistics we set *inbred*: YES parameter, as our data are represented by pseudo-haploid genotypes which introduce artificial genetic drift that masks the negative signal of admixture³¹.

Testing for the number of streams of ancestry and estimating mixture proportions

We used the *qpWave*^{6,56,57}/*qpAdm*⁶ framework which relates a set of ‘left’ populations (the population of interest and candidate ancestral sources) to a set of ‘right’ populations (diverse outgroups), testing for the number of streams of ancestry from ‘right’ to ‘left’ and estimating mixture proportions.

Simulations of admixed individuals

We simulated admixed individuals (Supplementary Information, section 2) given a set of sources and mixture proportions by first sampling (at each SNP) one of the sources (according to the mixture proportions), and then one of the individuals from that population (with equal probability). Due to missingness, the data-generating mixture proportions do not correspond precisely to the actual ancestry of simulated individuals (Supplementary Information, section 2). We note the maximum absolute value of the Z-score of the statistic $f_4(\text{Mycenaean}, \text{Simulated}; A, B)$, where A, B are two outgroup populations to test whether for a particular choice of ancestry of *Simulated* it forms a clade with the sampled Mycenaeans.

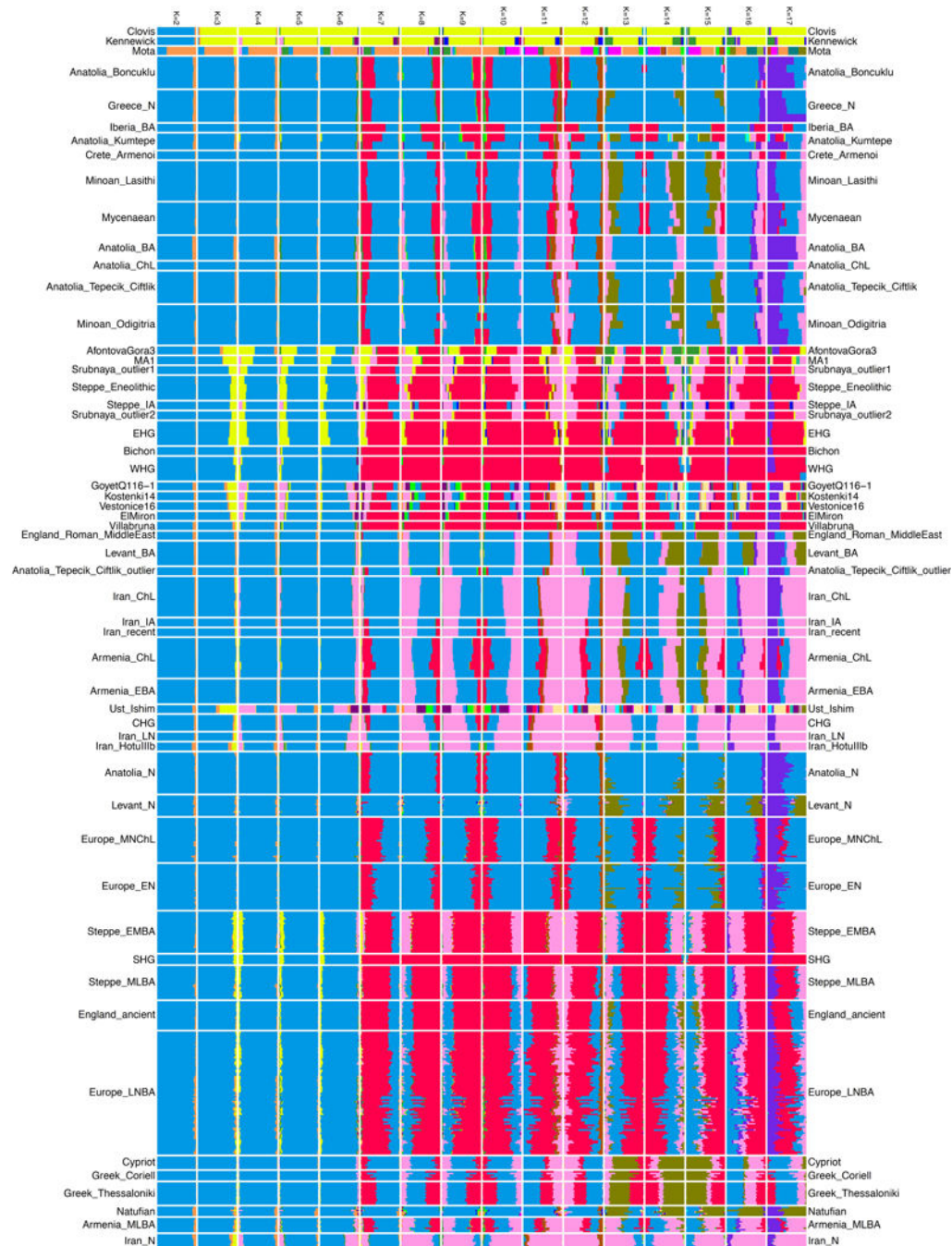
Estimation of F_{ST} coefficients

We estimated F_{ST} in *smartpca*²⁰ with the default parameters, inbreed: YES⁵⁷, and fstonly: YES.

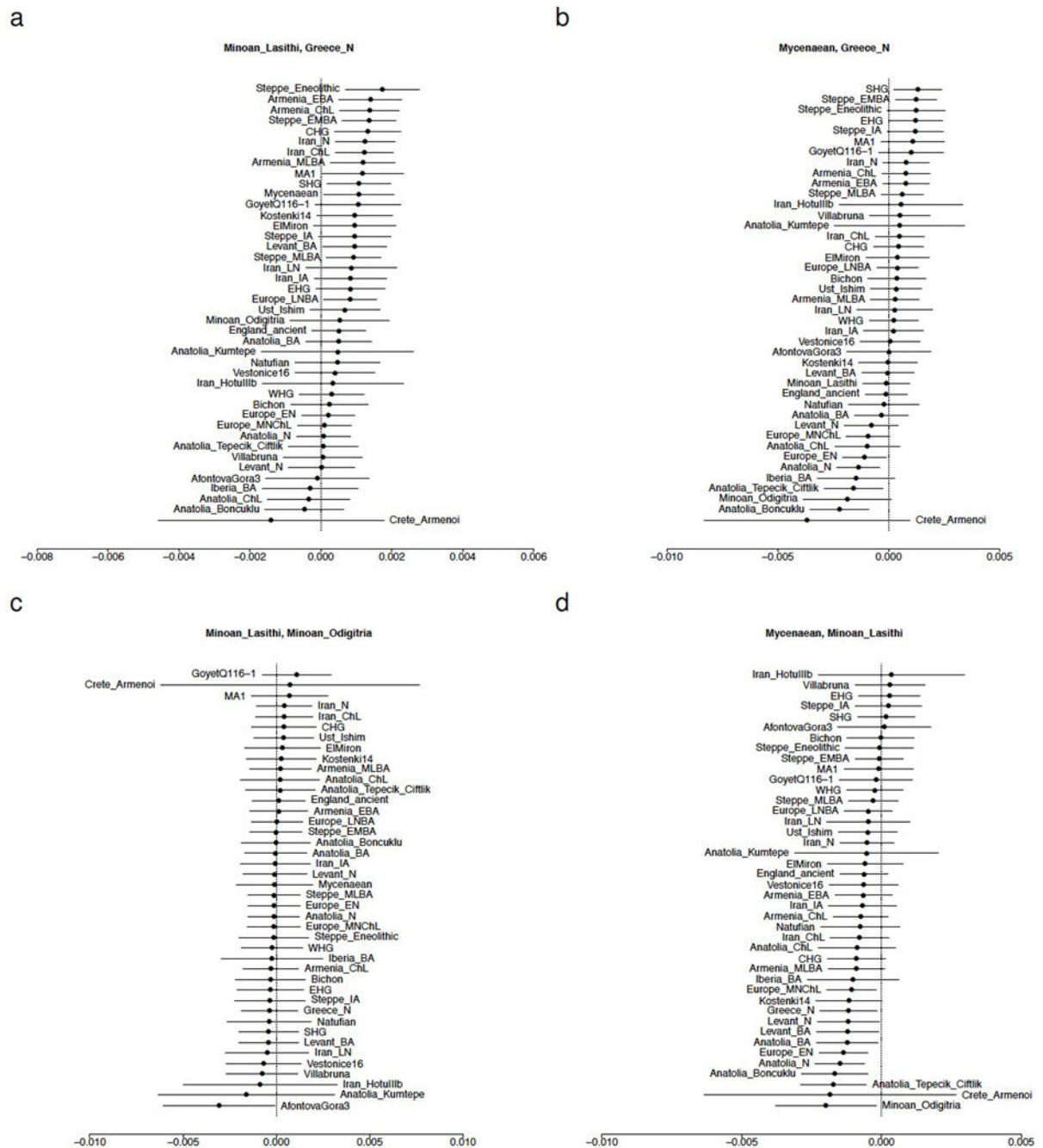
Phenotypic inference

The ancient samples have low coverage (median 0.87×) and thus diploid genotypes cannot be reliably assessed for them. However, we can use the low coverage data to compute allele frequencies in all individuals and the Bronze Age Aegean using likelihood approach¹. We then sample from the posterior distribution of the genotypes g given the read counts r of the reference allele and t of the total reads covering a site. We took 100 random genotype samples per individuals and submitted them to HIRISplex²⁶, obtaining an estimate of the uncertainty of phenotype inference (Supplementary Information, section 4; Extended Data Table 4).

Extended Data

**Extended Data Figure 1. ADMIXTURE analysis**

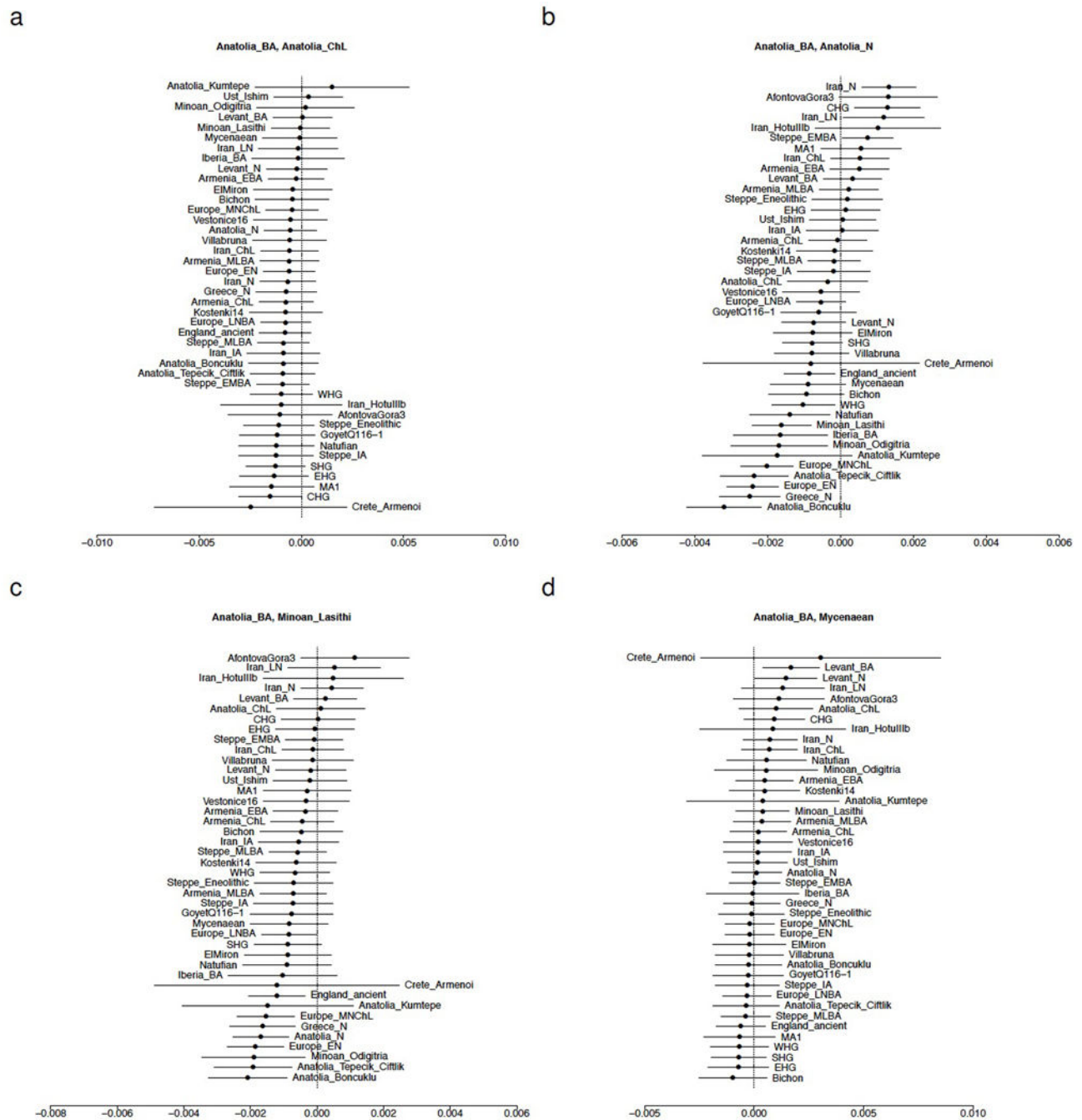
ADMIXTURE analysis with K=2 to K=17 is shown. 351 ancient and 2,616 present-day individuals were used in this analysis; ancient samples and present-day Greeks are displayed. To avoid visual clutter of labels, individuals in populations with sample size ≤ 5 are shown with thicker lines.



Extended Data Figure 2. Symmetry testing of Aegean Bronze Age populations

The statistic $f_4(X, Y; \text{Test}, \text{Chimp})$ is shown with ± 3 standard errors. Each panel is titled with the pair X, Y . Populations are ordered according to the value of the statistic. Positive values indicate that *Test* shares more alleles with X than Y and negative values that it shares more with Y than X . (a) ‘northern’ and ‘eastern’ populations share more alleles with Minoans than with Neolithic Greece. (b) ‘northern’ and ‘eastern’ populations share more alleles with Mycenaeans than with Neolithic Greece. (c) Minoans from Lasithi and Minoan_Odigithria are symmetrically related to diverse populations. (d) Neolithic populations from

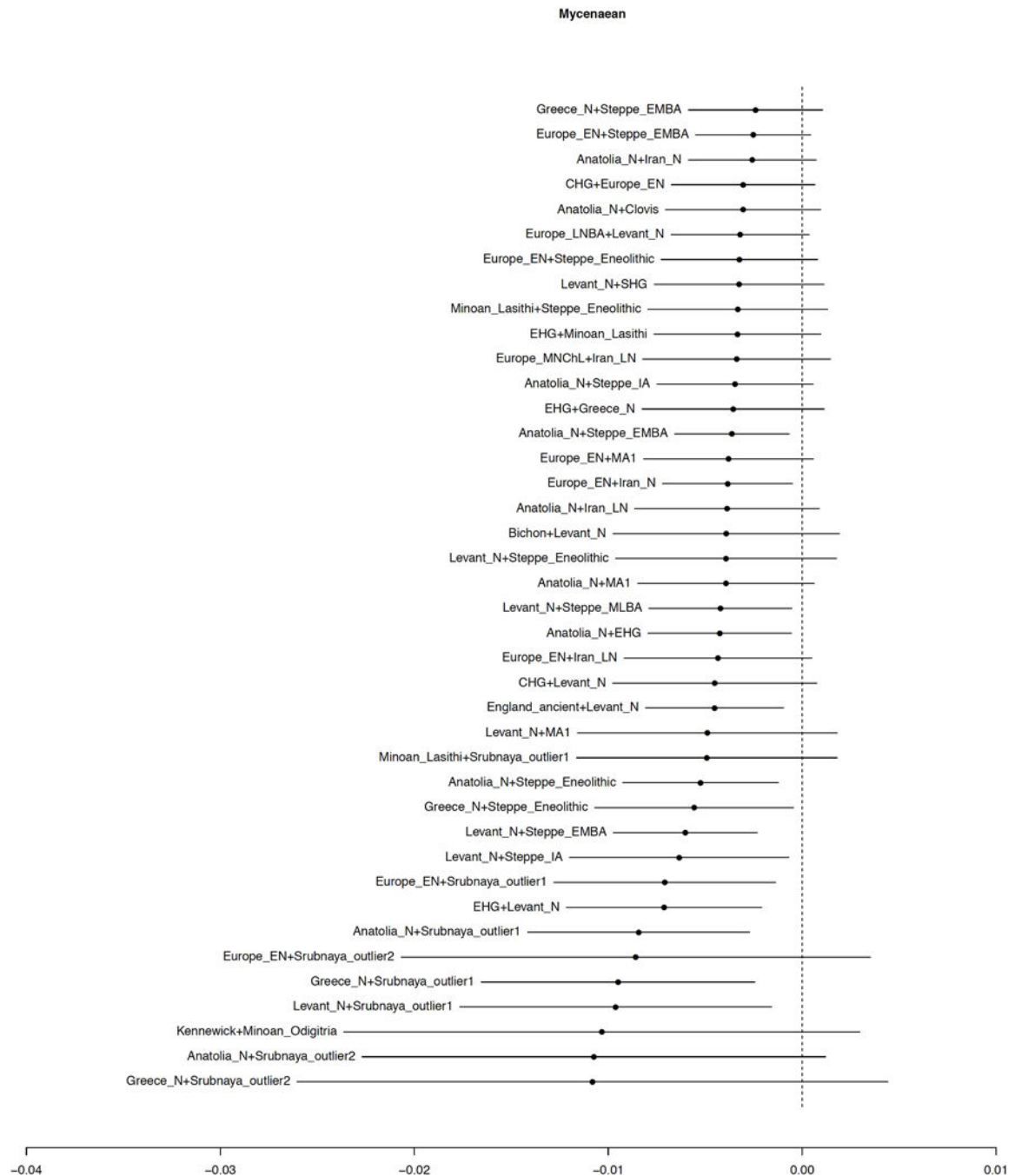
Anatolia, Europe, Greece, and the Levant share fewer alleles with Mycenaeans than with Minoans.



Extended Data Figure 3. Symmetry testing of Anatolian Bronze Age populations

The statistic $f_4(X, Y; \text{Test}, \text{Chimp})$ is shown with ± 3 standard errors. Each panel is titled with the pair X, Y. Populations are ordered according to the value of the statistic. Positive values indicate that *Test* shares more alleles with X than Y and negative values that it shares more with Y than X. (a) European, Siberian, and Caucasus hunter-gatherers share fewer

alleles with Bronze Age Anatolians from Harmanören Gündürle than with a Chalcolithic Anatolian from Barcın. (b) Bronze Age Anatolians differ from Neolithic ones in sharing more alleles with populations of Iran, the Caucasus, and the Steppe than with those of Europe. (c) Bronze Age Anatolians differ from Minoans in sharing more alleles with populations from Neolithic Iran than Neolithic Anatolia and Europe. (d) Bronze Age Anatolians differ from Mycenaeans in sharing more alleles with Neolithic and Bronze Age populations of the Levant.



Extended Data Figure 4. f_3 -statistics of Mycenaean as a target with different pairs of reference populations

We show the value of the statistic $f_3(Ref_1, Ref_2; \text{Mycenaean})$ and ± 3 standard errors; only the population pairs (Ref_1, Ref_2) for which the Z-score of the statistic is < -2 are shown. Negative values indicate that the Mycenaean population is admixed from sources related to the two reference populations.



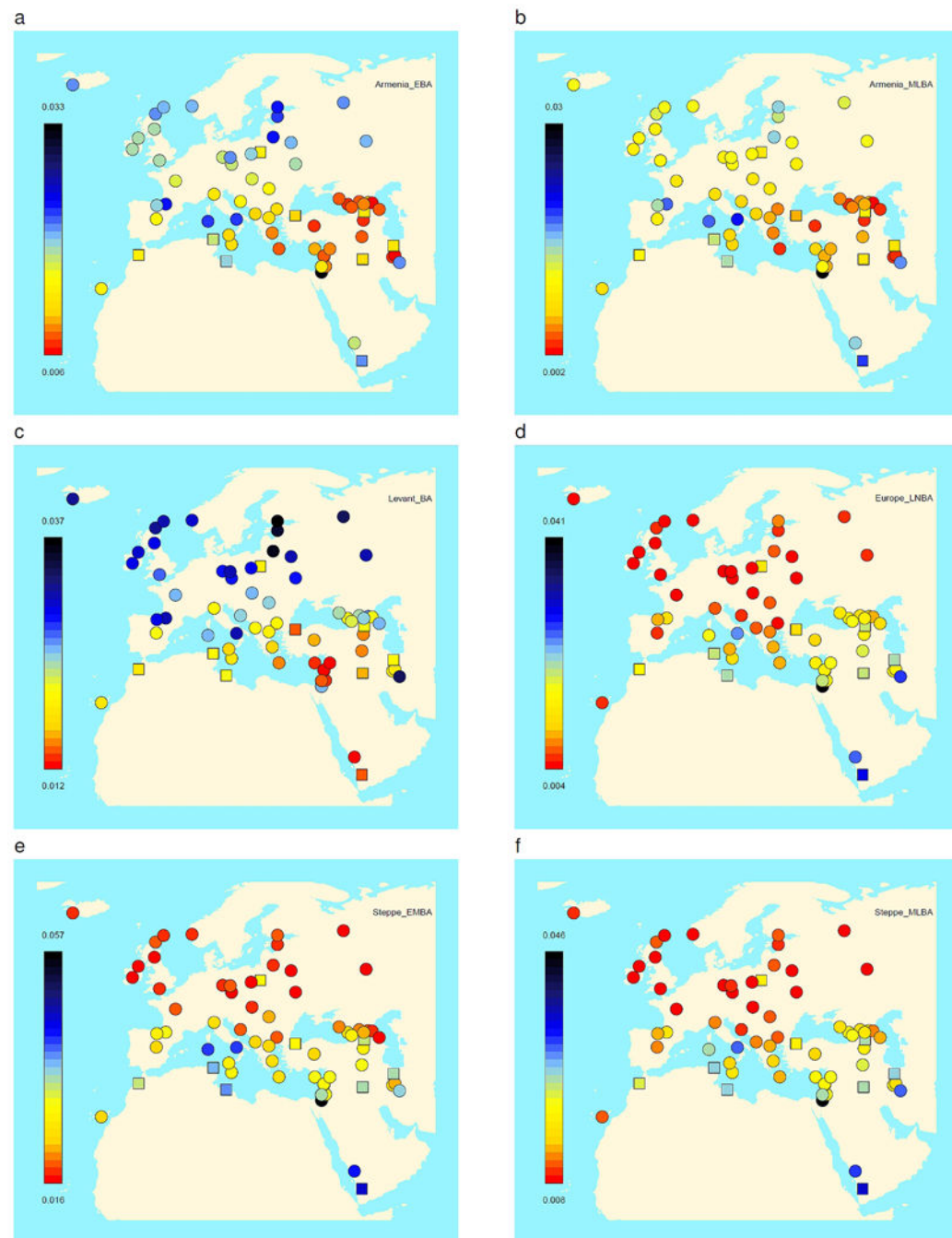
Extended Data Figure 5. Correspondence of $qpAdm$ estimates with PCA

As a way to validate *qpAdm* models of admixture for Mycenaean from three ancestral populations (Anatolia_N or Minoan_Lasithi), (Armenia_ChL or Armenia_MLBA), (Steppe_EMBA, Steppe_MLBA, Europe_LNBA), representing substratum, 'eastern', and 'northern' ancestry respectively (Supplementary Information, section 2), we plot the *qpAdm*-predicted position in the PCA space of Fig. 1 vs. the actual position of the Mycenaean population.



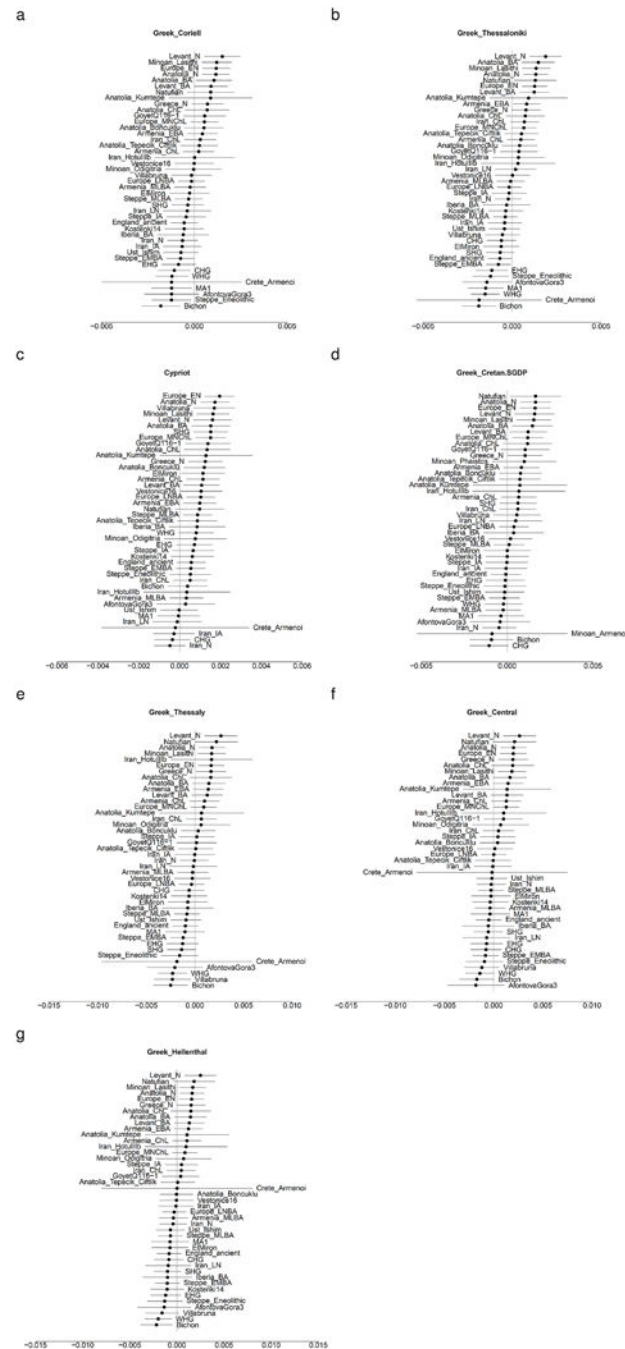
Extended Data Figure 6. Comparison of Mycenaean and simulated admixed populations

We simulate admixed individuals with known ancestry from three ancestral populations (Anatolia_N or Minoan_Lasithi), (Armenia_ChL or Armenia_MLBA), (Steppe_EMBA, Steppe_MLBA, Europe_LNBA), representing substratum, 'eastern', and 'northern' ancestry respectively (Methods; Supplementary Information, section 2). The maximum $|Z|$ -score of statistics $f_4(\text{Mycenaean}, \text{Simulated}, \text{Outgroup}_1, \text{Outgroup}_2)$ is plotted with circles of varying size (proportional to $\log|Z|$) for each assignment of ancestry proportions. The best estimate (red) corresponds to the proportions that minimize $|Z|$, and they are compared against the *qpAdm* estimate for the same ancestral sources (blue).



Extended Data Figure 7. F_{ST} between Bronze Age and present-day West Eurasian populations
 (a) The population of Early Bronze Age Armenia⁴ shows an affinity to present-day populations from Armenia, Anatolia, the Caucasus, and Iran, as does (b) Middle/Late Bronze Age Armenia^{4,9}. (c) The Bronze Age Levant⁴ has an affinity to Levantine and Arabian populations. (d) Late Neolithic/Bronze Age Europeans^{1,6,9,43} most resemble present-day northern/central Europeans, as do (e) Early/Middle Bronze Age steppe populations^{1,6,9}, who also resemble populations of the northeast Caucasus, while (f) Middle/Late Bronze Age steppe populations resemble central/northern Europeans^{1,9}. Jewish

populations are plotted with a square to distinguish them from non-Jewish populations from the same geographical area. The plots for the newly reported populations of Mycenaeans, Minoans, and Bronze Age Anatolians are shown in Fig. 2.



Extended Data Figure 8. Symmetry testing of Mycenaeans with Modern Greek populations
The statistic $f_4(\text{Mycenaean}, \text{Modern Greek}; \text{Test}, \text{Chimp})$ is shown with ± 3 standard errors. Modern Greeks share fewer alleles with Levantine/Anatolian/European Neolithic populations and with Minoans than Mycenaeans do, suggesting a dilution of early Neolithic

ancestry since the Bronze Age. Human Origins genotype data: (a) Greeks from the Coriell repository¹⁰, (b) Greeks from Thessaloniki¹⁰, (c) Cypriots¹⁰. Whole genome data: (d) Cretans⁴⁰. Illumina genotype data: (e) Greeks from Thessaly⁴¹, (f) Greeks from Central Greece⁴¹, (g) Greeks from the study by Hellenthal et al.²⁷

Extended Data Table 1
Information on ancient samples reported in this study

Dates marked simply as BCE are based on the associated archaeology of the samples. Dates marked as calBCE are based on radiocarbon dating of the samples (Supplementary Information, section 1).

Individual_ID	Genotype_ID	Other_ID	Source	Date	Population_Label	Location	Country	Latitude	Longitude	Sex	Coverage	Autosomal_SNPs	mtDNA	Y-chromosome
I2937	I2937	A2197	1240K	5419±41 cal BC	Greece_N	Dion, Alepotrypa Cave	Greece	36.64	22.38	F	0.870	481848	K1a2b	
I0071	I0071	Lasithi4	1240K	2000-1700 BCE	Minoan_Lasithi	Hagios Charalambos Cave, Lasithi, Crete	Greece	35.08	25.83	F	7.312	953157	U5a1	
I0070	I0070	Lasithi2	1240K	2000-1700 BCE	Minoan_Lasithi	Hagios Charalambos Cave, Lasithi, Crete	Greece	35.08	25.83	M	1.267	619767	H13a1	J2a1d
I0073	I0073	Lasithi7	1240K	2000-1700 BCE	Minoan_Lasithi	Hagios Charalambos Cave, Lasithi, Crete	Greece	35.08	25.83	M	1.481	643360	H	J2a1
I0074	I0074	Lasithi9	1240K	2000-1700 BCE	Minoan_Lasithi	Hagios Charalambos Cave, Lasithi, Crete	Greece	35.08	25.83	F	0.874	506434	H5	
I9005	I9005	Lasithi17	1240K	2000-1700 BCE	Minoan_Lasithi	Hagios Charalambos Cave, Lasithi, Crete	Greece	35.08	25.83	F	1.351	388859	H	
I9006	I9006	Salamis31	1240K	1411-1262 cal BCE (3067 ± 25 BP, DEM-2905)	Mycenaean	Agia Kyriaki, Salamis	Greece	37.97	23.50	F	1.387	361193	X2d	
I9123	I9123	S-EVA 1263 Ammenoi 503	1240K	1370-1340 BCE	Crete_Armenoi	Armenoi, Crete	Greece	35.45	24.17	F	0.041	45158	U5a1	
I9127	I9127	12V i2	1240K	2900-1900 BCE	Minoan_Odigirra	Moni Odigirra, Heraklion, Crete	Greece	35.05	24.81	F	0.035	36475	J2b1a1	
I9128	I9128	13V i2	1240K	2900-1900 BCE	Minoan_Odigirra	Moni Odigirra, Heraklion, Crete	Greece	35.05	24.81	F	0.016	17081	I5	
I9129	I9129	14V i2	1240K	2900-1900 BCE	Minoan_Odigirra	Moni Odigirra, Heraklion, Crete	Greece	35.05	24.81	F	0.063	63986	H+163	
I9130	I9130	16V Tholos	1240K	2900-1900 BCE	Minoan_Odigirra	Moni Odigirra, Heraklion, Crete	Greece	35.05	24.81	M	0.086	92186	U3b3	G2a2b2
I9131	I9131	19V i2	1240K	2900-1900 BCE	Minoan_Odigirra	Moni Odigirra, Heraklion, Crete	Greece	35.05	24.81	F	0.095	96946	K1a2	
I9010	I9010	Galatas19	1240K	1700-1200 BCE	Mycenaean	Galatas Apatheia, Peloponnese	Greece	37.50	23.45	F	0.379	242265	X2	
I9033	I9033	Peristeria4	1240K	1416-1280 cal BCE (3064 ± 24 BP, DEM-2903)	Mycenaean	Peristeria Tryfilla, Peloponnese	Greece	36.92	21.70	F	0.439	248912	H	
I9041	I9041	Galatas4	1240K	1700-1200 BCE	Mycenaean	Galatas Apatheia, Peloponnese	Greece	37.50	23.45	M	1.558	417898	X2	J2a1
I2495	I2495	A4-1	1240K	2558-2295 calBCE (3923±35 BP, Poz-811111)	Anatolia_BA	Harmandören-Göndürlü Höyük, Isparta	Turkey	37.92	30.71	M	1.981	637146	H	J1a
I2499	I2499	UC1	1240K	2836-2472 calBCE (4040±35 BP, Poz-82213)	Anatolia_BA	Harmandören-Göndürlü Höyük, Isparta	Turkey	37.92	30.71	F	0.285	243348	K1a2	
I2683	I2683	G3-95	1240K	2500-1800 BCE	Anatolia_BA	Harmandören-Göndürlü Höyük, Isparta	Turkey	37.92	30.71	F	3.695	749308	T2b	

Extended Data Table 2
Phenotypic inference of ancient individuals

We list the probability assignments for different phenotypes by HirisPlex²⁶ and an assessment of the phenotype. We generate 100 random replicates of the genotypes of each individual, listing the standard deviation in parentheses (Supplementary Information, section 4).

ID	Population	PBlueEye	PIntermediateEye	PBrownEye	PBlondHair	PBrownHair	PRedHair	PBlackHair	PLightHair	PDarkHair	Hair Color	Eye Color
I2495	Anatolia_BA	1.6 (4.4)	3.6 (3.9)	94.9 (8.3)	10.7 (6.1)	51.6 (6.4)	0.1 (0.1)	37.6 (9.3)	18.0 (11.7)	82.0 (11.7)	Brown	Brown
I2499	Anatolia_BA	16.6 (28.3)	7.4 (2.2)	76.0 (28.7)	2.2 (2.2)	64.7 (11.8)	2.0 (5.3)	31.1 (13.8)	12.9 (20.1)	87.1 (20.1)	Brown	Blue or Brown
I2683	Anatolia_BA	0.3 (0.9)	1.3 (1.7)	98.4 (2.6)	3.3 (2.5)	33.0 (4.6)	0.0 (0.0)	63.7 (7.0)	4.9 (4.5)	95.1 (4.5)	Black	Brown
I2937	Greece_N	0.3 (1.3)	2.2 (1.9)	97.5 (3.2)	3.6 (1.9)	33.9 (6.2)	0.1 (0.0)	62.4 (7.4)	6.7 (4.3)	93.3 (4.3)	Black	Brown
I0070	Minoan_Lasithi	0.4 (1.8)	2.2 (1.9)	97.4 (3.7)	30.4 (5.1)	66.4 (5.9)	3.2 (0.9)	0.0 (0.0)	100.0 (0.0)	0.0 (0.0)	Brown	Brown
I0071	Minoan_Lasithi	0.0 (0.0)	0.2 (0.0)	99.8 (0.0)	0.4 (0.0)	20.3 (0.0)	0.0 (0.0)	79.3 (0.0)	0.5 (0.0)	99.5 (0.0)	Black	Brown
I0073	Minoan_Lasithi	0.1 (0.7)	1.7 (1.4)	98.2 (2.2)	12.5 (3.4)	61.1 (1.2)	0.2 (0.1)	26.2 (2.7)	32.4 (8.8)	67.6 (8.8)	Brown	Brown
I0074	Minoan_Lasithi	0.0 (0.0)	1.3 (0.3)	98.7 (0.4)	9.3 (3.2)	54.8 (8.5)	0.1 (0.1)	35.8 (10.5)	18.8 (10.3)	81.2 (10.3)	Brown	Brown
I9005	Minoan_Lasithi	5.2 (0.0)	11.6 (0.0)	83.2 (0.0)	49.6 (1.4)	38.8 (1.2)	4.2 (0.5)	7.4 (0.7)	85.6 (1.7)	14.4 (1.7)	Blond or Brown	Brown
I9006	Mycenaean	0.0 (0.0)	1.1 (0.4)	98.9 (0.4)	8.7 (4.9)	59.9 (6.4)	1.8 (2.9)	29.6 (11.8)	25.7 (16.5)	74.3 (16.5)	Brown	Brown
I9033	Mycenaean	0.4 (1.0)	1.6 (1.9)	98.0 (3.0)	4.6 (3.9)	51.0 (6.3)	0.1 (0.5)	44.2 (9.8)	10.5 (13.2)	89.5 (13.2)	Brown	Brown
I9041	Mycenaean	1.4 (0.5)	5.3 (1.0)	93.3 (1.4)	7.8 (0.7)	63.2 (2.0)	0.2 (0.4)	28.7 (2.3)	21.2 (2.5)	78.8 (2.5)	Brown	Brown

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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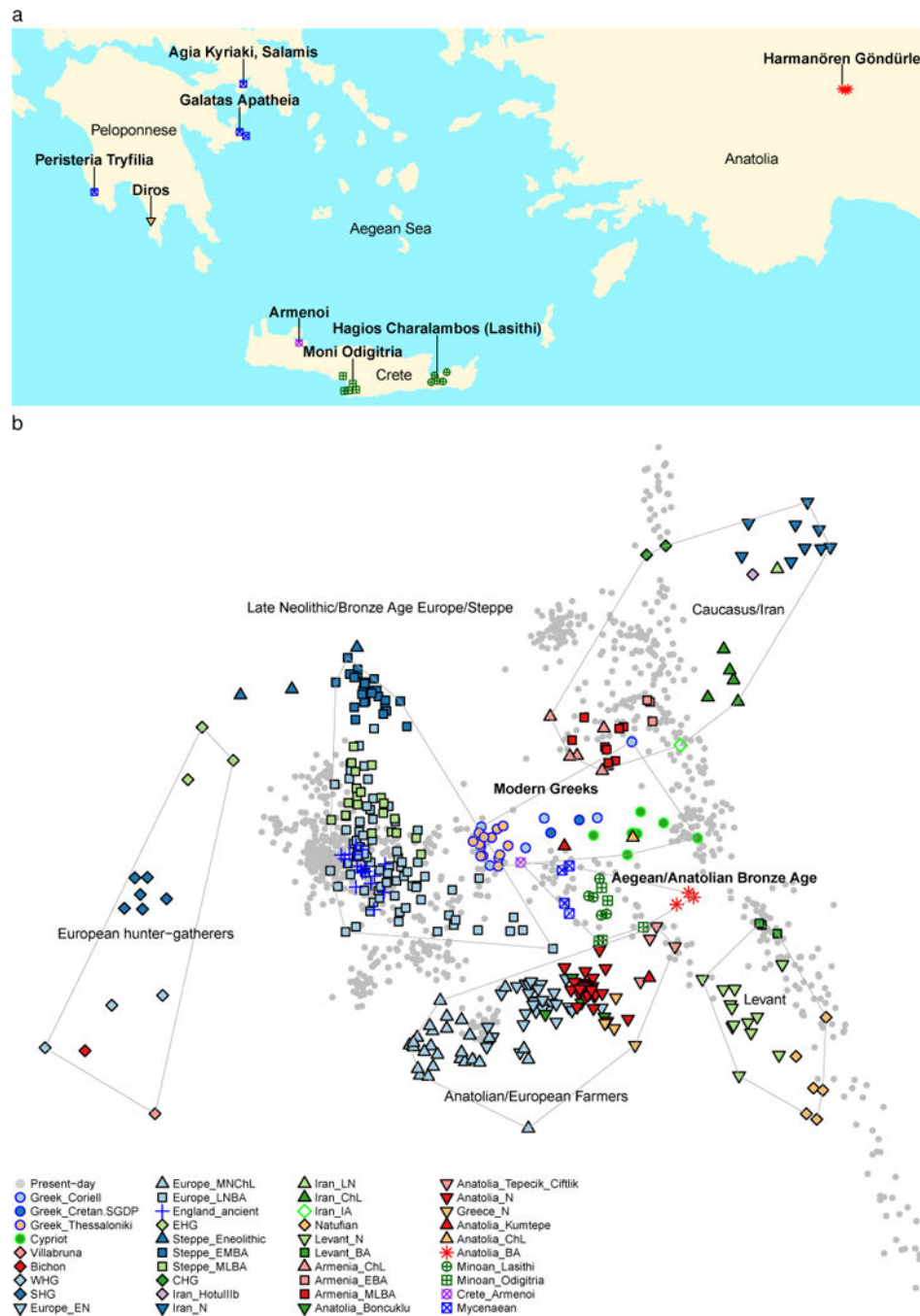


Figure 1. Samples and principal components analysis

(a) Geographical locations of newly reported ancient data. Lines point to sampling locations; jitter is added to show the number of sampled individual per location. (b) 334 ancient individuals projected onto the first two principal components computed on a sample of 1,029 present-day West Eurasians^{4,5,10,31}, including 30 Modern Greek samples from Greece and Cyprus.

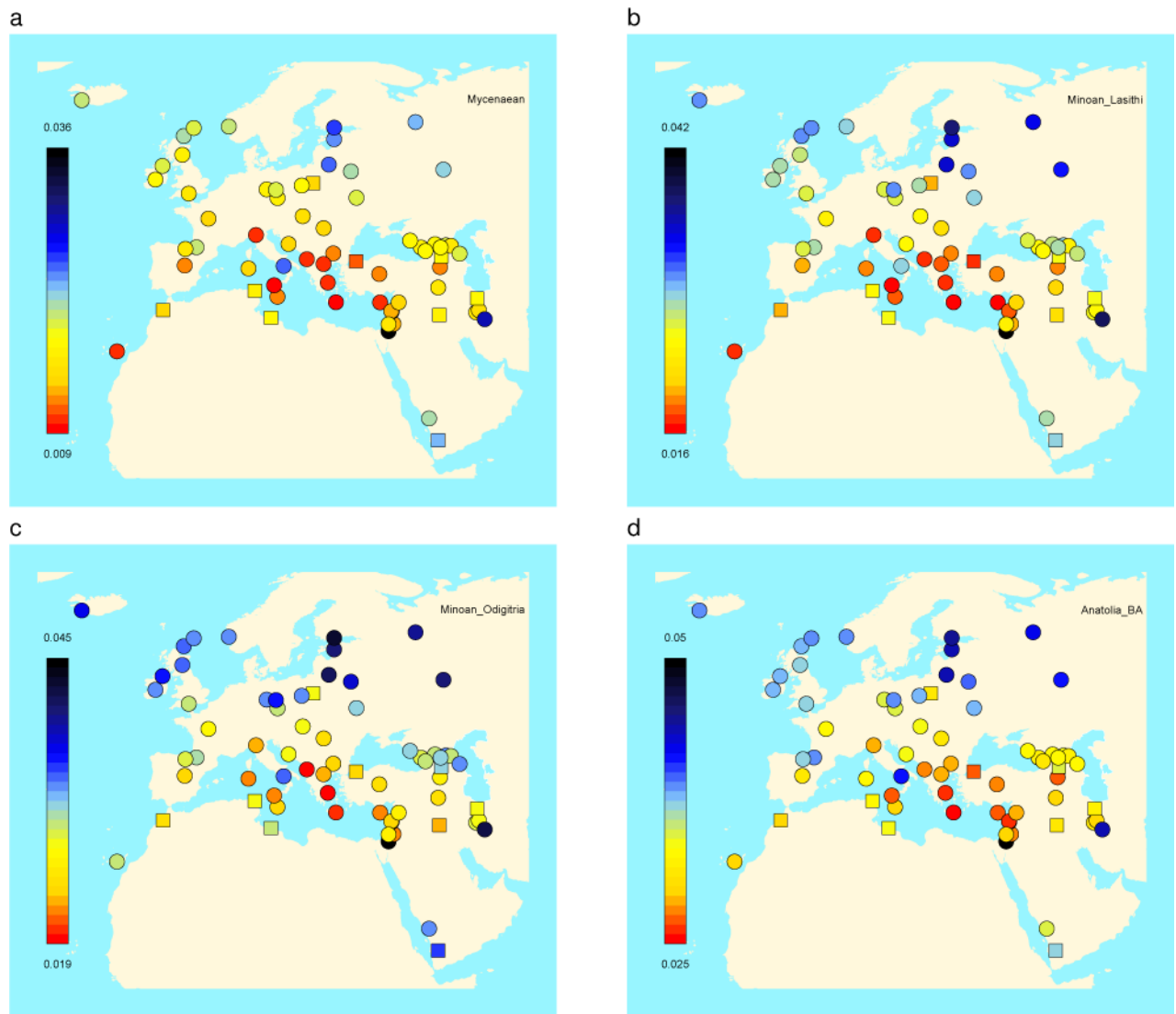


Figure 2. Genetic differentiation of Bronze Age populations to present-day populations

We plot the F_{ST} inbreeding coefficient (Methods) between newly reported populations and present-day West Eurasian populations which shows a pattern of genetic affinity between Bronze Age and present-day populations from the corresponding broad geographical regions. (a) Mycenaean, (b) Minoans from Hagios Charalambos (Lasithi regional unit), (c) Minoans from Moni Odigitria (Heraklion regional unit), (d) southwestern Bronze Age Anatolians. The same pattern also applies to Bronze Age populations from other regions of West Eurasia (Extended Data Fig. 5).

Table 1

Admixture modelling of Bronze Age populations

For each test population, mixture proportions from four source populations with their standard errors are given. Ancestry is inferred from both ‘ultimate’ sources representing the earliest populations, and ‘proximate’ sources representing populations down to the Bronze Age (Supplementary Information, section 2). Column A lists ‘northern’ sources from eastern Europe and Siberia, including the Eurasian steppe; column B lists ‘eastern’ sources from Iran, the Caucasus, and Anatolia (after the Early Neolithic); column C lists ‘local’ sources from Anatolia and the Aegean; column D lists sources from the Levant. For abbreviations of population names see Methods.

Test	Ancestral Sources				Mixture Proportions				Standard Errors			
	A	B	C	D	A	B	C	D	A	B	C	D
Anatolia_BA		CHG	Anatolia_N	Levant_N		0.319	0.618	0.063		0.029	0.078	0.063
Minoan_Odigitria		CHG	Anatolia_N			0.144	0.856			0.031	0.031	
Minoan_Odigitria		Iran_N	Anatolia_N			0.137	0.863			0.032	0.032	
Minoan_Lasithi	MA1	CHG	Anatolia_N		0.001	0.152	0.847		0.015	0.021	0.020	
Minoan_Lasithi	Moia	CHG	Anatolia_N		0.004	0.154	0.842		0.024	0.026	0.020	
Mycenaean	AfontovaGora3	CHG	Anatolia_N		0.133	0.126	0.741		0.027	0.026	0.024	
Mycenaean	AfontovaGora3	Iran_N	Anatolia_N		0.161	0.086	0.754		0.026	0.025	0.024	
Mycenaean	EHG	Iran_N	Anatolia_N		0.065	0.136	0.799		0.016	0.022	0.024	
Mycenaean	EHG	CHG	Anatolia_N		0.044	0.176	0.780		0.016	0.023	0.024	
Mycenaean	MA1	CHG	Anatolia_N		0.052	0.159	0.789		0.019	0.026	0.024	
Anatolia_BA			Anatolia_ChL	Natufian			0.908	0.092		0.039	0.039	
Anatolia_BA			Anatolia_ChL	Levant_BA			0.892	0.108		0.114	0.114	
Anatolia_BA			Anatolia_ChL	Levant_N			0.951	0.049		0.051	0.051	
Anatolia_BA			Anatolia_ChL	Anatolia_N			0.935	0.065		0.062	0.062	
Mycenaean		Armenia_MLBA	Anatolia_N			0.367	0.633			0.020	0.020	
Mycenaean		Armenia_ChL	Anatolia_N			0.441	0.559			0.025	0.025	
Anatolia_BA		Anatolia_ChL	Minoan_Lasithi			0.970	0.030			0.108	0.108	
Mycenaean	Steppe_MLBA		Minoan_Lasithi		0.175		0.825		0.017		0.017	

Test	Ancestral Sources				Mixture Proportions				Standard Errors			
	A	B	C	D	A	B	C	D	A	B	C	D
Mycenaean	Europe_LNBA		Minoan_Lasithi		0.198		0.802		0.019		0.019	
Mycenaean	Steppe_EMBA		Minoan_Lasithi		0.132		0.868		0.014		0.014	