

The Etruscans: A Population-Genetic Study

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The origins of the Etruscans, a non-Indo-European population of preclassical Italy, are unclear. There is broad agreement that their culture developed locally, but the Etruscans' evolutionary and migrational relationships are largely unknown. In this study, we determined mitochondrial DNA sequences in multiple clones derived from bone samples of 80 Etruscans who lived between the 7th and the 3rd centuries B.C. In the first phase of the study, we eliminated all specimens for which any of nine tests for validation of ancient DNA data raised the suspicion that either degradation or contamination by modern DNA might have occurred. On the basis of data from the remaining 30 individuals, the Etruscans appeared as genetically variable as modern populations. No significant heterogeneity emerged among archaeological sites or time periods, suggesting that different Etruscan communities shared not only a culture but also a mitochondrial gene pool. Genetic distances and sequence comparisons show closer evolutionary relationships with the eastern Mediterranean shores for the Etruscans than for modern Italian populations. All mitochondrial lineages observed among the Etruscans appear typically European or West Asian, but only a few haplotypes were found to have an exact match in a modern mitochondrial database, raising new questions about the Etruscans' fate after their assimilation into the Roman state.

Introduction

Analysis of genetic data in modern populations has proved to be a powerful tool for reconstructing crucial aspects of human evolutionary history (Cavalli-Sforza et al. 1994; Ingman et al. 2000; Barbujani and Bertorelle 2001; Tishkoff and Verrelli 2003). It is now technically feasible to validate these analyses by directly studying the DNAs of ancient people, but so far only a handful of European sequences have been published (Handt et al. 1994; Di Benedetto et al. 2000). In this study, we present the first extensive genetic data on a European population of the preclassical period, the Etruscans.

The Etruscan culture developed in central Italy (Etruria) in the first millennium B.C. The oldest-known inscriptions in Etruscan, a non-Indo-European language isolate, date back to the end of the 8th century, right after the shift from the rural Villanovian culture, doc-

umented in the same area in the 9th century B.C., to an urban society (Bartoloni 1989). The Etruscan cities were independent states that shared a language and a religion but never formed a political unit. However, between the 7th and the 5th centuries, leagues of Etruscan cities established their political and cultural leadership over an area spanning from the Po Valley to *Magna Graecia* (fig. 1), including, during part of the 6th century, Rome. Military defeats, the Roman expansion, and progressive assimilation caused a decline of the Etruscan cities, which lost their autonomy when Roman citizenship was granted to the Roman allies (90–89 B.C.). Immediately afterward, the language disappeared (Pallottino 1975; Barker and Rasmussen 1998).

Paleoanthropological studies have only proved broad similarities between the Etruscans and their neighbors of the Iron Age (Barker and Rasmussen 1998). Archaeological evidence suggests that the Etruscan culture developed locally, with some features pointing to an Eastern influence (Pallottino 1975; Barker and Rasmussen 1998). However, it is not clear if such influence reflects only trading and cultural exchange or rather some sort of shared biological ancestry. That is a long-lasting controversy. Dionysius of Halicarnassus (1.30.2) favored local development, whereas, according to Herodotus (1.94), the Etruscans were Lydians of Anatolia who were fleeing from famine (Barker and Rasmussen 1998). No modern

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Figure 1 Map of Italy showing the area of Etruscan influence (gray) in the 7th and 6th centuries B.C., from Barker and Rasmussen (1998). A solid line identifies the boundaries of Etruria proper. Solid circles are sampling locations: A, Adria (17 samples, 5 DNA sequences used for statistical analyses); V, Volterra (6, 3); S, Castelfranco di Sotto (2, 1); P, Castelluccio di Pienza (1, 1); M, Magliano and Marsiliana (25, 6); T, Tarquinia (18, 5); C, Capua (8, 6). Additional samples that yielded no amplifiable DNA were from Castelnuovo Berardenga (1, 0) and Pitigliano (2, 0).

archaeologist supports the latter view, but some affinities between the Lydian and the Etruscan languages have been recognized (Beekes 2002), and gene flow from the eastern Mediterranean area is impossible to rule out on archaeological grounds (Tykot 1994). Unfortunately, no original documents are available to help solve this question. Indeed, although the Etruscan alphabet and language are largely understood (Bonfante and Bonfante 2002), the written record is limited to short inscriptions of religious or funerary content.

By themselves, DNA sequences cannot tell us who the Etruscans were and where they came from, but they can provide crucial information on two related questions:

1. Were the Etruscans a single population, or were they simply a set of individuals who shared a language and a culture but not a common ancestry?
2. What are the genetic relationships between the Etruscans and modern populations, and do these relationships suggest any genealogical or migrational links between the Etruscans and other Eurasians?

To address the above questions, we obtained from museums and public collections fragments of 80 well-preserved skeletons from 10 Etruscan necropoleis (fig. 1), covering much of Etruria in terms of both chronology (7th to 2nd centuries B.C.) and geography. All human remains analyzed come from sites where (1) the material culture has been identified as Etruscan by archaeologists, and (2) all inscriptions, if any, are in Etrus-

can. Two cities, Adria and Capua, were at the fringes of Etruscan territory, in the Po River valley and in Campania, respectively. Historical documents and archaeological evidence, such as inscriptions in Etruscan and decorations on the pottery, show that these were indeed Etruscan settlements (Barker and Rasmussen 1998; Bonfante 1999; Haynes 2000), although they were both locations where hybridization may have occurred—with Veneti (in Adria) and with Samnium natives or Greek colonizers (in Capua).

Material and Methods

DNA is generally present only in small amounts in ancient samples, and it is often damaged. Therefore, extreme precautions are needed to minimize the risk of amplifying and typing modern contaminating DNA molecules. This is especially important when dealing with relatively recent human samples, whose genetic material may not be very different from that of the archaeologists and biologists who manipulated it. To maximize the probability of extracting and sequencing authentic DNA from the ancient samples, the strictest available standards (Cooper and Poinar 2000) were followed throughout this study. All samples that did not comply with any of Cooper and Poinar's nine criteria were discarded. In most cases, no information was available on the archaeologists and museum staff who had previous contact with the bone material, but all people who manipulated the specimens for this project had their mtDNA typed and compared with the sequences obtained from the ancient specimens.

Authentication Methods for Ancient DNA

Physically isolated work area.—All work was performed in isolated areas of the Florence and Barcelona laboratories, where no modern DNA has ever been introduced (criterion 1 of Cooper and Poinar [2000]).

Amino acid racemization.—In living individuals, all amino acids are in the L-enantiomeric form, but after death a racemization process begins. Poinar et al. (1996; Poinar and Stankiewicz 1999) described a correlation between the amount of D-forms of three amino acids (Asp, Glu, and Ala) and the presence of amplifiable DNA in the sample. To test for biochemical preservation as indirect evidence for DNA survival, we thus estimated the degree of amino acid racemization in each sample, using ~5 mg of bone and following the procedures described in the work of Vernesi et al. (2001) (criterion 7). In addition, to better estimate the preservation of organic material, 3 mg of bone powder from each individual was also analyzed by a thermogravimetric assay (Peters et al. 2000).

DNA extraction.—After brushing and irradiating each

bone surface (1 hr under ultraviolet light), DNA was extracted from powdered bone by means of a silica-based protocol (modified from Krings et al. 1997; see Di Benedetto et al. 2000). For each individual, we performed two independent extractions from different bones, usually a rib and a fragment of long bone (criterion 2). For each individual, we also attempted to amplify longer mtDNA fragments (360 bp and 700 bp), which have been reported to be unusual in ancient specimens; we sequenced only the samples in which that attempt failed (criterion 3). A negative control was included in each extraction (criterion 2).

Quantitation of ancient DNA.—Sporadic contamination is considered unlikely when the number of molecules that PCR will use as template, or “target DNA,” is >1,000. We estimated the amount of target DNA by competitive PCR, as described in the work of Handt et al. (1996) (criterion 8). A competitor was used containing a 95-bp deletion (nucleotide positions 131–225; positions were numbered according to Anderson et al. [1981], –16,000) in the mitochondrial hypervariable region I (HVR-I) (Caramelli et al. 2003). PCR components were the same as described below for the sequencing of HVR-I, and the primers were the same as those used for the amplification of the second of three HVR-I fragments. A negative control was included in each amplification (criterion 2). Thermal cycler conditions consisted of an initial 10 min incubation at 95°C, followed by 45 cycles of 50 s at 94°C, 50 s at 48°C, and 50 s at 72°C, with a final extension step at 72°C for 5 min.

Amplification.—The following profile was used to amplify 2 μ l of DNA extracted from the bone: 94°C for 10 min (*Taq* polymerase activation), followed by 50 cycles of PCR (denaturation 94°C for 45 s; annealing 53°C for 1 min; and extension 72°C for 1 min) and a final step at 72°C for 10 min. The 50- μ l reaction mix contained 2 U of *AmpliTaq* Gold (Applied Biosystems), 200 μ M of each dNTP, and 1 μ M of each primer. The 360-bp-long HVR-I was subdivided in three overlapping fragments by use of the following primer pairs: L15995/H16132; L16107/H16261; and L16247/H16402 (Caramelli et al. 2003). Each extract was amplified at least twice (criterion 4). Since overlapping primers were used throughout the PCR amplifications, it is highly unlikely that we amplified a nuclear insertion rather than the organellar mtDNA (Handt et al. 1996; Greenwood et al. 1999).

Cloning and sequencing.—PCR products were cloned (criterion 5) using the TOPO TA Cloning kit (Invitrogen), according to the manufacturer’s instructions. Screening of white recombinant colonies was accomplished by PCR. The colonies were transferred into a 30- μ l reaction mix (67 mM Tris HCl [pH 8.8], 2 mM MgCl₂, 1 μ M of each primer, 0.125 mM of each dNTP, and 0.75 U of *Taq* polymerase) containing M13 forward and reverse universal primers. After 5 min at 92°C, 30

cycles of PCR (30 s at 90°C, 1 min at 50°C, 1 min at 72°C) were performed and clones with an insert of the expected size were identified by agarose-gel electrophoresis. After purification of these PCR products with Microcon PCR devices (Amicon), a volume of 1.5 μ l was cycle-sequenced, according to the BigDye Terminator kit (Applied Biosystems) supplier’s instructions. The sequence was determined using an Applied BioSystems 377 DNA sequencer.

Independent replication.—To test for contamination within the laboratory, three bone fragments were subjected to DNA extraction, amplification, cloning, and sequencing in Barcelona (criterion 6). In this lab, the following primer pairs were used: L16055/H16218 and L16209/H16401.

Associated faunal remains.—The presence of human mtDNA sequences in extracts from nonhuman bones proves that those bones have been contaminated by human DNA and, hence, that sequences obtained from human remains in the same burial may also result from contamination. Cattle (*Bos taurus*) remains retrieved in Magliano, in the tomb of the individual with the 5AM sequence, gave us an opportunity to test for this type of contamination. We tried to amplify the DNA extracted from the cattle bones by use of both human-specific and bovine-specific primers. We used primers for a fragment of 152 bp of the *Bos taurus* mtDNA hypervariable region and for a fragment of the human HVR-I (the primers were L16030/H16137 and L16107/H16261, respectively). The PCRs were performed with 2 μ l of DNA, 1 μ M of each primer, 200 μ M of each dNTP, 1 \times reaction buffer (Applied Biosystems), 1.5 mM MgCl₂, and 2 U of *AmpliTaq* Gold (Applied Biosystems) in a total volume of 50 μ l. The cattle hypervariable region was amplified using the following thermal cycle: initial denaturation at 94°C for 10 min, followed by 40 cycles of 94°C for 1 min, 57°C for 1 min, 72°C for 1 min, and final extension at 72°C for 5 min. The conditions described above for human mtDNA amplification were used in the amplification of bovine DNA with human-specific primer. PCR products were obtained using only the primers specific for cattle. After run on a 1.5% agarose gel, bands of the appropriate size were excised from the gel and purified with Ultra Free DNA (Amicon). Cloning and sequencing of the PCR products were performed as described above (criterion 9).

RFLP: typing of the 14766 site.—A short fragment of the coding mitochondrial region around nucleotide position 14766 was amplified in a 50- μ l reaction volume containing 2 μ l of template DNA, 1.5 mM MgCl₂, 200 μ M of each dNTP, 2 U *AmpliTaq* Gold (Applied Biosystems), and 1 μ M of each primer (L14695/H14792), with the following thermal cycles: initial denaturation at 94°C for 10 min, followed by 40 cycles of 94°C for 1 min, 57°C for 1 min, 72°C for 1 min, and final ex-

tension at 72°C for 5 min. Fifteen μ l of PCR product were digested with 10 U of *Mse*I (Celbio Italy), using the recommended buffer with overnight incubation at 37°C. Digestion products were visualized by electrophoresis on 2.5% agarose gels. To confirm the results of the restriction analysis, the nucleotide at position 14766 was also determined in each sample by independent amplification (with the same primer pair and under the same conditions as those described above), cloning, and sequencing.

Database of Modern Mitochondrial Sequences

The data in the database analyzed by Simoni et al. (2000) were updated with the following populations: Haviks of India (Mountain et al. 1995), Egyptians (Krings et al. 1999), Syrians and Greeks (Vernesi et al. 2001), Uighurs, Kazakhs, and Kirghiz of Central Asia (Comas et al. 1998), Armenians and Azerbaijanis (Nasidze and Stoneking 2001), and Ladin speakers of Italy (Vernesi et al. 2002), which replaced the previous sample from the same area. Populations and sample sizes are listed in the legend to figure 4.

Statistical Analysis

Haplotype variation was summarized by a reduced median network (Bandelt et al. 1999). Sites with lower mutation rates were given greater weight. Allele sharing was estimated by counting the occurrences of the haplotypes observed among the Etruscans in the 34 populations of the database.

Pairwise genetic distances (F_{ST}) between populations were estimated by the ARLEQUIN software package (Schneider et al. 2000), considering both haplotype-frequency differences and numbers of substitutions between haplotypes. Kimura's two-parameter method was used (assuming a gamma distribution for rate variation among sites, with $\alpha = 0.26$), which allows for multiple substitutions at a site and for different rates of transitions and transversions. F_{ST} distances calculated in this way are equivalent to $(t_m - t_0)/t_m$, where t_m is the mean coalescence time of two random haplotypes drawn from the two populations, and t_0 is the mean coalescence time of two random haplotypes drawn from the same population (Slatkin 1991). From the matrix of F_{ST} distances, we obtained a two-dimensional representation of the data by multidimensional scaling (MDS) (Kruskal 1964), using the software STATISTICA.

Patterns of genetic variation within the Etruscan and the Italian populations were described by analyses of molecular variance (AMOVA) (Excoffier et al. 1992), as implemented in the ARLEQUIN package. AMOVA estimated indexes of molecular diversity among members of the same populations, among populations, and (in the description of Etruscan diversity across time periods)

among groups thereof, and tested for their statistical significance using a Monte Carlo randomization approach.

The genetic relationships between the Etruscans and modern Italian populations were quantified by estimating admixture coefficients for each of them, representing the relative contribution of potential parental populations. The purpose of this exercise was not to identify the historical admixture process giving rise to the Etruscan gene pool, for which the necessary data will be available only when several ancient populations have been described at the mitochondrial level. Rather, we wanted to obtain figures allowing us to quantitatively compare the composition of modern and ancient gene pools. Thus, admixture coefficients were inferred from differences in haplotype frequencies, considering the Etruscans and the modern Italian populations as hybrids among up to four potential parents (Dupanloup and Bertorelle 2001). The mitochondrial features of the parental populations were approximated assuming that the best available estimates of allele frequencies in past (and unknown) populations is found in their modern counterparts, as is customary in admixture studies (e.g., Chakraborty 1986; Alves-Silva et al. 2000; Chikhi et al. 2002). We chose the Basques as representative of western Europe, the Turks as representative of the eastern Mediterranean region, Karelians and Volga Finns as representative of northeastern Europe, and Egyptians and Algerians as representative of North Africa. Admixture coefficients can also be estimated from the coalescence times between haplotypes. However, mitochondrial coalescence times go back to several tens of thousand years ago in Eurasia (Richards et al. 2000). The specimens in this study are ~2,500 years old, and in this period changes of haplotype frequencies resulting from drift certainly had a greater effect on genetic diversity than the onset of new mutations. Therefore, estimates based only on haplotype frequencies seemed preferable.

Results

The 80 samples in this study underwent nine tests for authentication of ancient DNA, and at each step those that did not comply with the relevant criterion were eliminated. In particular, three samples yielded a D/L ratio for aspartic-acid enantiomers >0.1, which is considered incompatible with retrieval of sufficient endogenous DNA from ancient remains (Poinar et al. 1996) and, hence, were discarded. In the remaining 77 samples, not only the content of racemized amino acids was low but also the relative extent of racemization was Asp>Glu>Ala, both findings generally associated with good preservation of endogenous macromolecules (Poinar et al. 1996). Low temperatures are known to facilitate the preservation of DNA in ancient samples (Kumar et al. 2000; Reed et al. 2003), and, indeed, all samples of this study come from

room burials or caves, where temperatures are low and constant throughout the year.

Competitive PCR showed that 48 of these 77 samples had sufficient amounts (initial number >1,000 copies) of template DNA (Handt et al. 1996); 29 samples were discarded at this stage. Thus, 48 samples were selected for which all tests indicated a good probability to obtain DNA sequences without artifacts. In all of them, the PCR amplification strength appeared inversely related to the size of the product, and PCR products larger than 200 bp were not observed, both findings consistent with the expected behavior of ancient (as opposed to contaminating modern) DNA molecules (Hofreiter et al. 2001). Each extract was subjected to two amplifications of three overlapping fragments, each PCR product was cloned, and multiple clones were sequenced, so that each of the three fragments was sequenced at least eight times (two extracts \times two PCRs \times two clones or more). Eighteen individuals yielded incomplete or multiple sequences and were discarded at this stage. Thirty individuals gave complete PCR products whose sequences were the same across clones, except for sporadic misincorporation of nucleotides in single clones. The overall misincorporation rate was a low 0.27% over the 92,104 nucleotides sequenced (compared, for example, with 0.39% in the “Iceman” specimen of Handt et al. 1994), which confirms that there was a fairly large amount of target DNA in the 30 samples.

The entire procedure was repeated in the Barcelona laboratory, under double-blind conditions, for three bone specimens, and identical sequences were obtained. Finally, the DNA from cattle remains found in the burial site of the individual with the 5AM sequence could be amplified only using bovine-specific (but not human) primers. The sequences, determined from multiple clones, match with those of the *Bos taurus* sequence deposited in GenBank (accession number NC_001567), showing no trace of human sequences, and so once again failing to provide evidence of modern human DNA contamination.

We observed 23 different sequences with 26 variable sites (25 transitions, 1 transversion) (table 1). (Sequences of each clone in this study can be accessed at the authors' Web site.) All identical sequences came from different necropoleis, except for three individuals from the same tomb in Tarquinia. To remove possible effects of consanguinity on the estimated population statistics, we excluded two of them from further analyses. A different problem was represented by sequence 2V, which showed six substitutions—the 069-186-189-223-319-362 motif—that usually occur in distant and mutually exclusive branches of the mtDNA network (Richards et al. 2000). Independent replicates in two laboratories under double-blind conditions confirmed the sequence, showing that the motif does not result from experimental artifacts. At

this stage, we can only speculate that some kind of oxidative damage, possibly resulting from the specific microclimatic condition of the burial site (Lindahl 1993, 1997), has altered the DNA molecules of that specimen. At any rate, for the sake of phylogenetic consistency, it seemed prudent not to consider this sequence in successive statistical analyses.

Exclusion of the 2V sequence brought the final sample size to 27 individuals with 22 different sequences. Only seven such sequences (4V, 5AM, 6AM, 8A, 9A, 16S, and 19M) occur in a database of 34 modern populations comprising 2,481 individuals from western Eurasia and the southern Mediterranean shores.

The Etruscan sequences show substitutions at sites (069, 126, 223, 270, and 356) known to be prone to recurrent mutation or postmortem damage (Gilbert et al. 2003). Substitutions at these mutational hotspots may cause misassignment of sequences to haplogroups (Bandelt et al. 2002). To achieve a better resolution, we then typed the restriction site of the coding region 14766 *MseI*, which is characteristic of the HV superhaplogroup (Richards et al. 2000). Restriction cuts on ancient DNA molecules are not always reproducible (but see Endicott et al. 2003), and so we chose not to use these markers extensively for assigning each haplotype to a haplogroup. However, the 14766 *MseI* restriction cuts were confirmed by sequencing of multiple clones in the region of interest. The results were unambiguous across replicates and make phylogenetic sense (with two exceptions; in these individuals, this position was left undefined) (table 1). Therefore, typing of the 14766 *MseI* site both clarified and further validated the results of HVR-I sequencing.

In the reduced median network based on HVR-I sequences and on the 14766 *MseI* polymorphism (fig. 2), several clusters are evident. Two lineages, characterized by substitutions 193-219 and 356, respectively, have a rather high internal diversity. The former substitutions are documented in Cornwall, England (but in association with 186-260-362), and the latter along the eastern and central Mediterranean shores, including Tuscany, with some derivatives in northern Europe. The only lineage containing a transversion, 095G-189, also occurs in Turkey, while the lineage with the 129 substitution is present in the eastern Mediterranean region and in northern Europe. The lineages with 126 or 126-362 presumably belong to the pre-HV haplogroup and have been observed in southeastern Europe and in the Levant.

Six Etruscan sequences show the 126-193 motif. In modern individuals, these substitutions occur in lineages attributed to the J2, and occasionally T, haplogroups, accompanied by substitutions at 069 or 294, respectively, which were not observed in these six sequences. Therefore, on the basis of the HVR-I motifs, these sequences could belong to either the HV or to the JT

Table 1**Consensus HVR-I Mitochondrial Sequences in 28 Etruscan Individuals**

Site	Century (B.C.)	Haplotype Label	HVR-I Motif (16024–16384)	14766 <i>MseI</i>	N_{SH}
Volterra	6th–5th	1V	193-219	–	0
Volterra ^a	3rd–2nd	2V	069-186-189-223-319-362	–	0
Volterra	2nd–1st	3V	189-274-334-356	–	0
Volterra	6th–5th	4V	261	+	7
Adria	5th–4th	5AM	CRS	–	32
Adria	5th–4th	6AM	126	+	8
Adria	5th–4th	7AC	126-193-278	+	0
Adria	5th–4th	8A	129	–	10
Adria	5th–4th	9A	223	NA	9
Capua ^a	3rd	10C	189-311-356	–	0
Capua	3rd	11C	069-095-223-261	–	0
Capua	3rd	12C	126-274-356	–	0
Capua	3rd	13C	193-219-356	+	0
Capua	3rd	7AC	126-193-278	+	0
Capua	3rd	14CMT	126-193	+	0
Castelluccio di Pienza	?	15P	193-219-256-270-291	–	0
Castelfranco di Sotto	?	16S	189-356	–	4
Magliano/Marsiliana	6th	17M	095G -126-189	–	0
Magliano/Marsiliana	7th	18M	066-126-193-219	–	0
Magliano/Marsiliana	6th	19M	311	–	26
Magliano/Marsiliana	6th	6AM	126	–	0
Magliano/Marsiliana	6th	14CMT	126-193	+	0
Magliano/Marsiliana ^a	7th–6th	5AM	CRS	NA	0
Tarquinia	3rd	20T	126-229-362	+	0
Tarquinia	5th	14CMT	126-193	+	0
Tarquinia	3rd	21T	126-193-228-229-278	+	0
Tarquinia	5th	22T	278-334	+	0
Tarquinia	3rd	23T	098-311-327	+	0

NOTE.—CRS is the Cambridge reference sequence (Anderson et al. 1981). The HVR-I motif is the position (–16,000) where substitutions were observed, with respect to the CRS; the only observed transversion is in boldface italic type. In the haplotype labels, capital letters indicate the site(s) where the haplotype was observed: A, Adria; C, Capua; M, Magliano and Marsiliana; P, Castelluccio di Pienza; S, Castelfranco di Sotto; T, Tarquinia; V, Volterra. The designation “14766 *MseI*” indicates the presence (+) or absence (–) of a diagnostic restriction cut. N_{SH} is the number of modern populations sharing that haplotype, among the 34 in the database. Haplotype 2V was excluded from the statistical analyses. NA = not available.

^a Samples for which DNA was independently reextracted and retyped in Barcelona.

haplogroups. However, in all of them, the *MseI* site was present at 14766, which rules out the hypothesis that they could belong to the HV haplogroup. Considering that postmortem changes lead to false positives (i.e., substitutions observed in the ancient sample that did not exist in the living individual) at 069 or 294 but are not known to generate false negatives there (Gilbert et al. 2003), we see no compelling reason to imagine that these results are due to postmortem damage. In addition, these sequences were determined in multiple clones for each individual, which makes a sequencing error unlikely. Therefore, despite the absence of substitutions at 069 or 294, the sequences with the 126-193 motif appear to belong to the JT haplogroup.

Internal genetic diversity within the Etruscans (gene diversity 0.98 ± 0.01 ; mean number of pairwise sequence differences 3.90 ± 2.02) is close to the average in the database (0.97 and 4.68, respectively). AMOVA

showed no heterogeneity among Etruscan sites (1.11% of the total [NS]), whereas differences are significant among contemporary Italian populations (1.72% of the total, $P < .0001$); in addition, we could not demonstrate any significant differences among time periods (7th–6th centuries vs. 5th–4th vs. 3rd–2nd). None of these tests suggests that the Etruscans are more of a cultural assemblage than a biological population.

Only two Etruscan haplotypes (5AM and 6AM, carried by 13.7% of the individuals) occur in a sample of modern Tuscans who were selected to represent inhabitants of former Etruria (Francalacci et al. 1996). The average value in comparisons of pairs of modern European populations is $27.9\% \pm 12.0\%$, showing that the genetic resemblance between the Etruscans and their modern counterparts is much less than observed between random European populations with no special evolutionary ties. Allele sharing is higher not only with

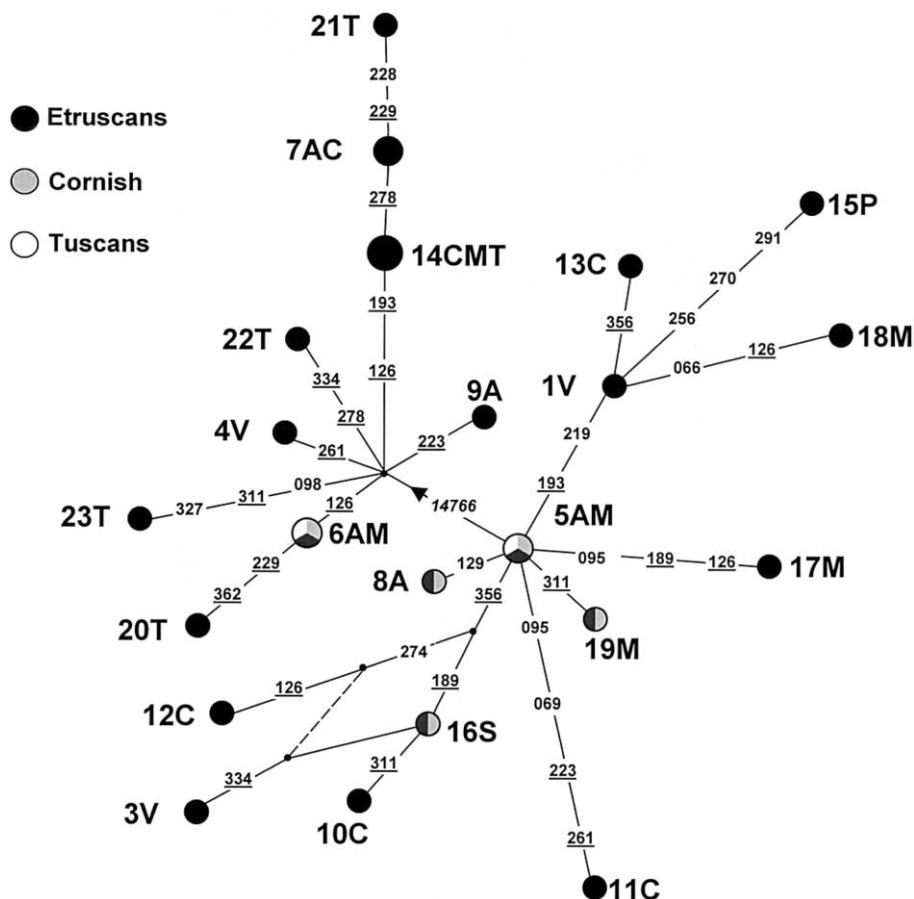


Figure 2 Reduced median network of the haplotypes identified among the Etruscans. Haplotype 5AM is the Cambridge reference sequence (CRS). Circle sizes are proportional to frequencies. Transitions are numbered relative to the CRS (-16,000); 095G is the only transversion. Sites undergoing recurrent mutations are underlined. The alleles shared with modern Tuscans and modern Cornish are highlighted.

the Turks (four haplotypes in common) but also with other, presumably unrelated, populations, such as the Cornish or the Germans (five and seven haplotypes in common, respectively). However, allele sharing may not be the best statistic summarizing the evolutionary relationships between the Etruscans and modern populations. Indeed, many haplotypes that so far have been observed only in the Etruscans differ by just one substitution from haplotypes that are present, or even common, among modern Europeans. Therefore, pairwise sequence differences, as well as genetic distances between populations, are more informative. The shortest genetic distances between the Etruscan and modern populations are with Tuscans ($F_{ST} = 0.036$; $P = .0017$) and Turks ($F_{ST} = 0.037$; $P = .0001$); values of $F_{ST} < 0.050$ were also observed for other populations of the Mediterranean shores and for the Cornish (fig. 3).

The mean pairwise sequence differences between the Etruscans and other European populations range from 3.79 to 6.26 substitutions, with a global average of

4.64. The Tuscans differ from the Etruscans by 4.65 substitutions—that is, almost exactly the value that would be expected with a random population from the database. However, these values are also affected by the internal diversity of both samples being compared. If we subtract the internal diversity, thus obtaining Nei's (1987) d_{AB} distance, the genetic distance between Etruscans and Tuscans becomes the shortest, with other southern and eastern Mediterranean populations also showing relatively close genetic relationships (fig. 3). In the MDS plot (fig. 4), the Etruscans fall out of an unstructured cluster comprising most European and Caucasus populations, including the Turks.

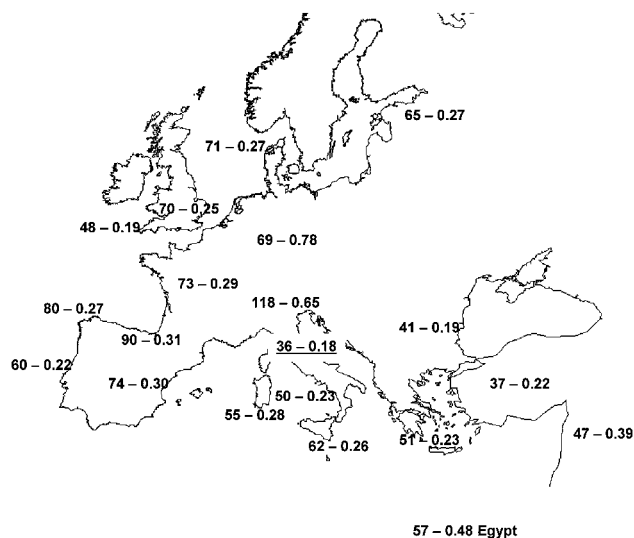
To better compare the Etruscan gene pool with those of contemporary Italy, we treated these populations as hybrids among four potential parental populations, from the four corners of the area considered in this study (table 2). The likely contributions of each parental population, or admixture coefficients, are similar for the three modern Italian populations, but Etruscans differ

in two aspects: they show closer relationships both to North Africans and to Turks than any contemporary population. In particular, the Turkish component in their gene pool appears three times as large as in the other populations. These admixture estimates are not to be taken at their face value, for numerous assumptions underlie their estimation. Here they only serve to show that, with respect to modern Italian gene pools, the Etruscan one contains an excess of haplotypes suggesting evolutionary ties with the populations of the southern and eastern Mediterranean shores.

Discussion

The first question we wanted to address is whether there is any evidence that the Etruscans were not a biological population, but rather an assemblage of biologically heterogeneous people who shared many cultural traits. Neither their internal genetic diversity, which is less than in comparable modern populations, nor the insignificant heterogeneity among Etruscan sites or time periods supports that view. In agreement with dental evidence (Moggi-Cecchi et al. 1997), the genetic data of this study suggest that either the people whom we call “Etruscans” shared a set of ancestors (and therefore can be considered a biological population as much as can current European populations) or they were mixed with people whose mitochondrial features did not differ from theirs.

As for the second question, which concerned the genetic relationships between the Etruscans and modern populations, various tests show that the Tuscans are the Etruscans’ closest neighbors in terms of genetic distances. Despite that broad similarity, however, Etruscans and Tuscans share only two haplotypes. This finding is difficult to interpret in the absence of data on any other European population of the preclassical period. One possible interpretation is that *all or most* European populations of that time period were as different from their modern counterparts as the Etruscans appear to be. This would imply either extensive gene flow or a high rate of extinction of mitochondrial haplotypes,



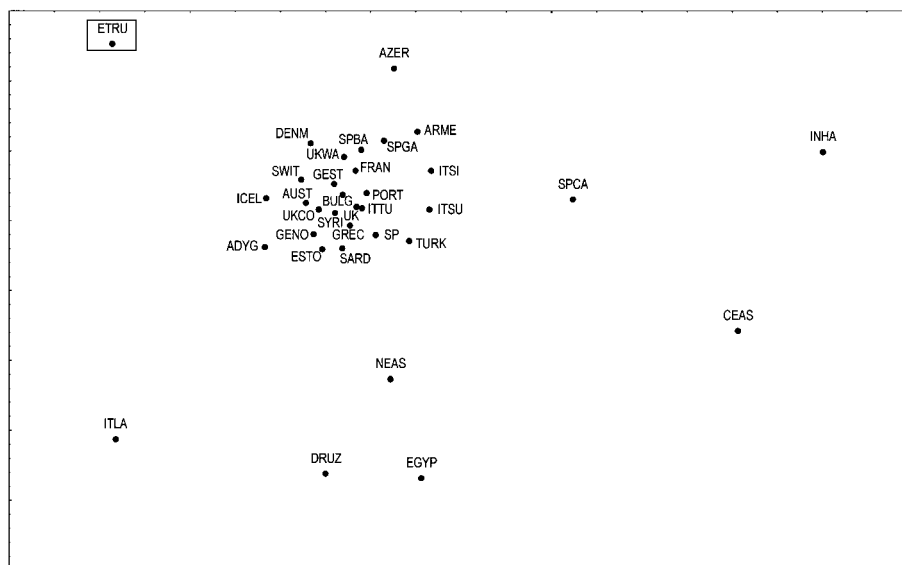


Figure 4 Two-dimensional representation of the relationships between populations (MDS plot) based on F_{ST} distances. Population labels and sample sizes: ETRU: Etruscans, 27; AUST: Austrians, 117; BULG: Bulgarians, 30; DENM: Danes, 32; ESTO: Estonians, 28; FRAN: French, 111; GENO, 108: Northern Germans; GEST: Southern Germans 249; GRC: Greeks, 48; ICEL: Icelanders, 53; ITLA: Ladins of Italy, 20; ITSI: Sicilians, 63; ITSU: Southern Italians, 37; ITTU: Tuscans, 49; PORT: Portuguese, 54; SARD: Sardinians, 72; SP: Spaniards, 74; SPBA: Basques, 106; SPCA: Catalans, 15; SPGA: Gallegos, 92; SWIT: Swiss 72; UK: English, 100; UKCO: Cornish, 69; UKWA: Welsh, 92; ADYG: Adigheians, 50; ARME: Armenians, 42; AZER: Azerbaijanis, 40; CEAS: Central Asians, 100; DRUZ: Druzes, 45; EGYPT: Egyptians, 91; INHA: Haviks of India, 48; NEAS: Near Easterners, 42; SYRI: Syrians, 49; and TURK: Turks, 96.

tifacts that could be attributed with confidence to the Etruscan culture. Those tombs typically belong to the social elites (Barker and Rasmussen 1998), and so the individuals we studied may represent a specific social group, the upper classes. We do not know whether that group differed genetically from the rest of the population, which might be the case when a foreign elite imposes its rule, and often its language, over a region (Renfrew 1989). If the upper class had indeed somewhat distinct DNAs, our results could mean that this elite class became largely extinct, while the rest of the population, whose DNA we do not know, may well have contributed to the modern gene pool of Tuscany. This would be the likely effect of a process of assimilation, from which the social elites were excluded, more or less deliberately.

To summarize, only a few Etruscan sequences find an exact match in the modern database, but all belong to lineages that are still present in Europe. Some genetic affinities with modern people from western Europe reflect the sharing of lineages (5AM, 6AM, and 19M) that are widespread over the whole continent, and that therefore do not seem to point to any migrational contact but rather to a common origin of various European gene pools. On the contrary, the similarity between the Etruscan and Turkish gene pools may indeed reflect some degree of gene flow. Commercial exchanges are docu-

mented between the Etruscan harbours and Asia Minor (Tykot 1994) and trading is often accompanied by interbreeding, ultimately leading to detectable levels of genetic affinity (see Relethford and Crawford 1995). Thus, the present study suggests that gene flow from the eastern (and possibly southern) Mediterranean shores, not necessarily from Lydia as proposed by Herodotus, left a mark in the Etruscan gene pool, above and beyond what is observed in contemporary Italy.

The limited genealogical continuity between the Etruscans (be they representative of the upper class or of the entire population) and their modern counterparts of Tuscany calls for an explanation. An approximate estimate of the Etruscan population size in the 6th century B.C. is probably somewhere between 150,000 and 250,000 women (Rasmussen 2004). In the future, we plan to quantify the probability of extinction of all mitochondrial haplotypes but two in a subdivided population of that size, by simulating both neutral evolution and a disadvantage representing ethnic discrimination after the Roman assimilation. We shall also ask whether reasonable rates of gene flow for 2,500 years can cause a dramatic replacement of mitochondrial lineages in a population of that size. But a clearer picture is likely to emerge only when genetic data on other European populations of the same age become available for comparison.

For the time being, however, this is the first large-

scale study of a pre-Roman European population in which all the strictest criteria for the validation of ancient DNA sequences have been followed. Within the limits imposed by the sample size (although our sample is large for ancient DNA studies, compared with 12 samples in Endicott et al. 2003, 19 samples in Lalueza-Fox et al. 2001, and 17 samples in Lalueza-Fox et al. 2003), the Etruscan sites appear to have rather homogeneous genetic characteristics. Their mitochondrial haplotypes are very similar, but rarely identical, to those commonly observed in contemporary Italy and suggest that the links between the Etruscans and eastern Mediterranean region were in part associated with genetic, and not only cultural, exchanges.

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Electronic-Database Information

Accession numbers and URLs for data presented herein are as follows:

Authors' Web Site, <http://web.unife.it/progetti/genetica/pdata/Etruscan.txt> (for sequences of each clone in this study)
GenBank, <http://www.ncbi.nlm.nih.gov/Genbank/> (*Homo sapiens* sequences [accession numbers AY530759–AY530781] and *Bos taurus* sequence [accession number NC_001567])

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