Genetics of the peloponnesian populations and the theory of extinction of the medieval peloponnesian Greeks

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Peloponnes has been one of the cradles of the Classical European civilization and an important contributor to the ancient European history. It has also been the subject of a controversy about the ancestry of its population. In a theory hotly debated by scholars for over 170 years, the German historian Jacob Philipp Fallmerayer proposed that the medieval Peloponnesians were totally extinguished by Slavic and Avar invaders and replaced by Slavic settlers during the 6th century CE. Here we use 2.5 million single-nucleotide polymorphisms to investigate the genetic structure of Peloponnesian populations in a sample of 241 individuals originating from all districts of the peninsula and to examine predictions of the theory of replacement of the medieval Peloponnesians by Slavs. We find considerable heterogeneity of Peloponnesian populations exemplified by genetically distinct subpopulations and by gene flow gradients within Peloponnes. By principal component analysis (PCA) and ADMIXTURE analysis the Peloponnesians are clearly distinguishable from the populations of the Slavic homeland and are very similar to Sicilians and Italians. Using a novel method of quantitative analysis of ADMIXTURE output we find that the Slavic ancestry of Peloponnesian subpopulations ranges from 0.2 to 14.4%. Subpopulations considered by Fallmerayer to be Slavic tribes or to have Near Eastern origin, have no significant ancestry of either. This study rejects the theory of extinction of medieval Peloponnesians and illustrates how genetics can clarify important aspects of the history of a human population.

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INTRODUCTION

Peloponnes was peopled by a series of migrations that spanned almost nine millennia. Early migrants arrived from Anatolia ca 9000 B.C.E.1,2 and established in the peninsula several Neolithic sites.2 The Mycenaeans3 who developed an advanced Bronze Era civilization, either migrated from the north around 2200 B.C.E.4,5 or were the descendants of the original Neolithic migrants.1 The next migration took place at the beginning of the first millennium B.C.E with the invasion of Peloponnes by the Dorian Greeks, referred in the Greek tradition as the return of the Heraclids.6 The subsequent 14 centuries were marked by quantitative changes of the population due to wars and epidemics but no qualitative effects from migrations of new population groups. Changes in population structure started in the beginning of the medieval period with the migrations of the Slavs to the Balkans.7,8 The effects of these migrations have dominated the historiography of Peloponnes during the last 170 years.

In 1830 CE, the German historian Jacob Philipp Fallmerayer presented his theory of disappearance of the Greek nation and its substitution by Slavs.9 Fallmerayer proposed that during the 6th century CE, large armies of Avars and Slavs overran the Balkans and eliminated the populations of the Hellas, who up to that period had successfully survived the attacks of barbarians and the religious suppression by the Byzantines. The Peloponnesian Greeks, except for few remnants enclosed in coastal castles, were slaughtered or forced to leave and Peloponnes was inhabited by Slavic tribes. The Slavs kept their identity for few centuries but eventually they were Hellenized under the influence of the Orthodox Church and interactions with Hellenized Asia Minor populations who were settled in Peloponnes by the Byzantines. Since the time Fallmerayer’s theory was published, a debate on the question of the ancestry of Peloponnesians has raged among historians (reviewed in Curta).10 Of note is that in spite of their diametrically different views, all historians have been using the same medieval written sources.

Controversies are rather common in historiography and result to endless debates among scholars. Controversies concerning the ancestry of populations can potentially be resolved by genetic analysis. In this paper, we use genome-wide data to study the genetic structure of the Peloponnesian populations and compare them with other populations of the world. We observe

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Figure 1 Substructure of the Peloponnesian populations. (a) Map of Peloponese showing the populations studied. Each dot corresponds to the origin of a participant. (b) PCA illustrating the separation of Peloponnesians in three groups. On the left is placed the population of Tsakones (north: open circles, south: green dots). On the right are placed the populations of Maniots (Deep Mani, East and West Tayetos.). All the remaining Peloponnesians are clustered in the center. (c) PCA analysis without the Maniot and Tsakones populations showing a partial separation of the population of Laconia. (d) IBD analysis illustrating the separation of Peloponnesians in three subpopulations. The color indicates the \( \log_{10} \) of the average proportion of genome shared IBD between individuals in two populations, where higher values (less negative; red color) indicates a higher pairwise proportion of genome shared IBD, and lower values (more negative; yellow color) indicates a lower proportion of genome shared IBD. (e) ADMIXTURE analysis. Notice the distinct structure of the Maniots and the Tsakones and their clear cut separation from all other Peloponnesians in all values of \( K \).
characteristic patterns of genetic differentiation within Peloponnesian, we examine their possible causes and we focus on the question of the impact of Slavic migrations on the genetic structure of the Peloponnesian populations. Our results reject the theory of extinction of medieval Peloponnesian Greeks and their replacement by Slavic and Asia Minor settlers.

SUBJECTS AND METHODS

Design of the study and populations studied

The study has been reviewed by the Institutional Review Board of the University of Washington and the ethical committees of several provisional hospitals. We focused on the rural population. We analyzed a total of 241 samples genotyped with the Illumina Infinium Omni 2.5–8 arrays. This is a figure illustrating the genetic similarity of Peloponnesians, Sicilians, and Italians.

Figure 2 Genetic similarity of Peloponnesians, Sicilians and Italians. PCA analysis of several European populations. (a) Notice the north to south distribution of the populations and that the Peloponnesians are placed to the far right of the graph and overlap with the Sicilians. (b) PCA analysis of Southern European populations illustrating the close relationship between Peloponnesians Sicilians and Italians (TSI is an Italian population) (c) Network analysis illustrating the high connectivity between the Peloponnesian populations as well as between the Peloponnesians, the Sicilians and the Italians. Notice the distance between Peloponnesians and the Slavic, and Near Eastern populations. Peloponnesians are connected with the Near Eastern populations through Crete and Dodecanese.
novel data set collected under the auspices of our study. Subjects were included in
the study if all four grandparents originated from the same village or from
villages that were <10 kilometers apart. The ages of most participants ranged
between 70 and 90 years (the oldest subject was 107 years old); hence their
grandparents were born between 1860 and 1880. In the 1861 census the
population of Peloponnese was 578,598 individuals. At that time the economy
of Peloponnese was exclusively agricultural and over 85% of the population was
living in small villages and hamlets. We sampled all the districts of Peloponnese
(Figure 1a and Supplementary Table 1) and also focused on two culturally
distinct subpopulations, the Tsacones and the Maniots. To compare the
Peloponnesians with other populations we analyzed samples from published
data sets and data sets generated by our studies (Supplementary Table 2 and
Supplementary Figure 1). Merging genotypes from different sources and quality
control were done as described.11

Identity by descent
The data used for the identity by descent (IBD) analysis were derived from data
sets listed and referenced in Supplementary Table 2. Two individuals share a
genomic segment identical by descent if they have inherited the segment
intact from a common ancestor, without intervening recombination. Allele
strand, reference and alternate alleles were aligned with the 1000 Genomes
European populations (CEU, GBR, TSI, FIN and IBS) using the conform-gt
utility (http://faculty.washington.edu/browning/conform-gt.html) prior to run-
ing Beagle. Marker exclusion criteria were unconfirmed strand orientation,
minor allele frequency <1%, Hardy–Weinberg P-value <10−6, and >2% missing data. After applying these criteria, 560,891 SNPs were used for the IBD
analysis.

We used the Refined IBD algorithm implemented in Beagle 4.110,11 to
phase the data and infer IBD segments. We set niterations = 160 and used
default values for all for all other parameters, including requiring a LOD score
of 3, which means that the probability of the observed genotype data for a pair
of samples in the inferred IBD segment is at least 1000 times greater under an
IBD model than under a non-IBD model. Based on the software recommenda-
tions and marker density, we set ibdtrim = 22. We used the HapMap geneti-
c map to infer genetic length and excluded IBD segments with length <2 cM,
since refined IBD has a low false-positive rate when using this threshold.12

R13 was used to generate plots and summarize IBD distributions. The heat
map summarizes average pairwise IBD between chromosomes from different
individuals in the populations being compared (see footnote 4 in Supplementary Table 3 for details), where higher average pairwise IBD indicates a
closer genetic relationship between populations.

Principal Component Analysis (PCA)
We used Eigenstrat14 as well as our own MatLab implementation of PCA.15,16

Estimating population admixture
We used the ADMIXTURE v1.22 software for all our admixture analyses.17
Cross validation errors of ADMIXTURE analysis ranged between 0.42–0.47
(Figure 1e), 0.55–0.58 (Figure 2c) and 0.56–0.62 (Figure 4d) with smaller error
at K = 2.

Quantitative analysis of ADMIXTURE output
Given a target population X and reference populations Y, Z and so on, we were
interested in quantifying the amount of ancestry of population X that is
captured by populations Y, Z and so on. Towards that end we devised a new
approach to quantitatively analyze the output of ADMIXTURE. Recall that
ADMIXTURE, for a particular value of K, will represent each sample using K
coordinates. Thus, for a particular value of K and for a particular population Y
with n samples, we can represent the output of ADMIXTURE for this
population as an n-by-K table. Then, for each reference population Y, we
summarize this n-by-K matrix using its top right singular vector only; in all our
analyses, the top singular value corresponding to the top right singular vector
captured at least 80% of the reference population variance as represented
by ADMIXTURE. Let vY be the top right singular vector (a K-dimensional
vector) for population Y; similarly, let vZ be the top right singular vector
(a K-dimensional vector) for population Z and so on. Now that we have
represented the ADMIXTURE output for each population as a K-dimensional
signature vector, we can apply standard vector space calculus in order to answer
our original question: how much of the ancestry of population X is captured by
population Y, or population Z and so on. More specifically, in order to
compute the percentage of the ancestry of population X that is captured by
population Y, we compute the percentage of the norm of VX that is captured
(in projection sense) by vY. Formally, we compute

\[ \frac{\|V_X - v_Y v_Y^T V_X\|_F}{\|V_X\|_F} \]

which returns a value between zero and one. In the above, VX denotes the
m-by-K matrix representing the m samples of population X with respect to
the K coordinates returned by ADMIXTURE. The notation v_Y^T indicates
the pseudo inverse of the vector vY, which is equal to the transpose of the vector vY,
suitably normalized. It is also worth noting that the norm used in the above
equation is the standard matrix Frobenius norm. In order to quantify the
amount of ancestry of population X that is captured by both populations Y and
Z, we form the K-by-2 matrix V = [vY vZ] whose columns are the vectors vY and
vZ and we compute

\[ \frac{\|V_X - V V^T V_X\|_F}{\|V_X\|_F} \]

In the above equation, V^T denotes the pseudo inverse of the matrix V; the
matrix V V^T is a projector on the subspace spanned by the column space of V.
Thus, we basically extract from the matrix VX the part of VX that is captured by
the (subspace spanned by the) vectors vY and vZ.

Network analysis
To better visualize and understand the connection between the populations
included in our study, we performed a network analysis on the results of
ADMIXTURE, using a method presented in reference.15

RESULTS
The substructure of the Peloponnesian populations
On PCA analysis the populations are arranged in the form of an
inverted capital letter V (Figure 1b). The left of this formation is
occupied by the population of Tsacones who inhabit the east slopes of
Mount Parnon and the adjacent costal area (Figure 1a). The right of
the formation is occupied by the populations of Maniots who inhabit
the east and west slopes of mount Tayetos and the southern area of
the promontory, the so called Deep Mani (Figure 1a). All other
Peloponnesians cluster in the tip of the letter V (Figures 1b and c).
The ADMIXTURE analysis of Figure 1e shows that the Maniots and
Tsacones are clearly separated from each other and from all other
Peloponnesians populations. Correlations between geographic coordi-
nates and the two principal components (Table 1) are compatible with

Table 1 Correlations between geographic coordinates and principal components

| Populations                      | Latitude correlation | Longitude correlation |
|---------------------------------|----------------------|-----------------------|
| All Peloponese                  | 0.50 (PC1)           | 0.41 (PC2)            |
| Peloponese minus Tsaconia and Mani | 0.49 (PC1)           | 0.09 (PC2)            |
| Arcadia                         | 0.60 (PC1)           | 0.12 (PC2)            |
| Laconia                         | 0.45 (PC1)           | 0.07 (PC2)            |
| Ionian Sea Coast                | 0.31 (PC2)           | 0.06 (PC1)            |
| Elis                             | 0.17 (PC1)           | 0.10 (PC2)            |
| Arcadia and Messenia            | 0.34 (PC2)           | 0.16 (PC1)            |
| Arcadia and Laconia             | 0.36 (PC2)           | 0.20 (PC1)            |
| Deep Mani                       | 0.15 (PC2)           | 0.21 (PC1)            |
| East Tayetos and Deep Mani      | 0.67 (PC1)           | 0.10 (PC2)            |
| West Tayetos and Deep Mani      | 0.73 (PC1)           | 0.42 (PC2)            |

*The respective principal component is indicated in parenthesis.
gradients in gene frequencies from north to south across all Peloponnese, along the Ionian coast, across Arcadia, as well as within Laconia and between the slopes of Tayetos and Deep Mani.

IBD analysis (Figure 1d) confirms the existence of distinct Peloponnesian subpopulations. In the populations of Maniots, individuals share on average 0.25% of their genome (or 35–36 cM) identical by descent, with 95% of pairs of individuals sharing at least one IBD segment (Supplementary Table 3). Similarly, the two Tsakones populations have a very high average pairwise IBD sharing of 0.66% of their genome, or 94 cM; every pair of individuals shares at least one IBD segment. Laconia is fairly closely related to both Deep Mani and to the Tsakones, based on the darker blue for those pairings in the heat map of Figure 1d. The heat map also revealed a larger separation between the southern and northern Peloponnesian populations, with the populations of Arcadia, Messenia, Elis, Achaea, Corinthia and Argolis forming their own subgroup. Corinthia and Argolis are the most strongly related (by IBD) of this subset of populations, sharing nearly 10 cM IBD on average.

Genetic similarity with Sicilians and Italians
As anticipated from the results of previous studies, the Peloponneseans are genetically placed very close to the Sicilians and Italians (Figures 2a and b) but they differ from several other populations we compared them (see Supplementary Figure 2). Network analysis (Figure 2c), highlighted the interconnections of Peloponnesean populations as well as the connections between Peloponneseans, Italians and Sicilians; in this network analysis, Sicilians and Italians serve as a bridge between Peloponneseans and other European populations (Basque, Andalusians and French). Slavic populations are placed far away from the Peloponneseans as are the Near Eastern populations. The latter are connected to the Peloponnesus via the islands of Crete and the Dodecanese.

Testing the theory of extinction of the medieval Peloponnesian Greeks
This theory allows a specific prediction about the genetic ancestry of the Peloponneseans: the great majority, if not all, of Peloponnesian ancestry should be Slavic. We compared, the Peloponneseans (except...
for Maniots and Tsakones) with populations of the Slavic homeland from which the sixth century Slavs should have originated. The exact location of the Slavic homeland is debated, but it is placed north of the Danube, between the Oder and Dnieper rivers and includes areas inhabited by Polish, Ukrainian, Russian and Belarusian populations. PCA analysis showed a clear separation of Peloponneseans from the Slavic populations (Figure 3a). By ADMIXTURE analysis (Figure 3c) the Peloponneseans and the Slavic populations form separate clusters with a small degree of gene flow from the Slavic to the Peloponnesean cluster.

Fallmerayer hypothesized that the Hellenization of the Peloponnesian Slavs was accelerated by the transfer to the Peloponnesse of Hellenized populations from Asia Minor. We tested this hypothesis by comparing the Peloponnesians with three Greek-speaking populations from Asia Minor: a western-coastal population sample extending from the Propontis in the north to Alikarnassos (Bodrum) in the south; a northern population from Pontus, that is, the coast of Black Sea and the Asia Minor interior corresponding to the current northern Turkey; and a central Anatolian population from Cappadocia. All these populations are separated from the Peloponnesians by PCA (Figure 3b). The small degree of overlap between Peloponnesse and the population of the Asia Minor coast (Figure 3b) is expected for neighboring Greek populations. The Byzantines frequently moved Armenians to achieve political objectives and Fallmerayer proposed that Armenians were included in the populations transferred to Peloponnesse by the Byzantines. Peloponnesians differ from the Armenians by PCA and ADMIXTURE analysis (Supplementary Figure 3). Collectively, these results are incompatible with the theory of extinction of the medieval Peloponnesians and their replacement by Slavic and Asia Minor settlers.

The medieval ancestry of the populations of Mani

The Maniots differ from all other Peloponnesians by PCA (Figure 1b) and ADMIXTURE (Figure 1c) analysis. They also differ from mainland, island and Asia Minor Greek populations (data not shown) and from all the other populations of Supplementary Figure 4, which have been compared by PCA analysis, but they partially overlap with the Sicilians and the Italians.
the Mellingi and the Ezernita, were forced by the Byzantines to withdraw to the slopes of Tayetos. The Mellingi continued to be mentioned in the Peloponnesian history until the 15th century. Because of the writings of Porphyrogenitus we sampled separately the populations of the East and the West slopes of the Tayetos and the Deep Mani. By PCA (Figure 4a) and ADMIXTURE (Figure 4d) the populations of Tayetos are distinct from the populations of the Slavic homeland. Fallmayer argued that the inhabitants of Deep Mani are of Slavic origin. PCA (Figure 4b) and ADMIXTURE (Figure 4d) analyses make this hypothesis unlikely.

As an alternative origin of the Maniots, Fallmayer proposed that they are descendants of Mardaites. This medieval warrior tribe used to inhabit the mountainous regions between Asia Minor and Syria but in late seventh century CE was resettled by the Byzantines in Asia Minor and other areas of the Empire. The Mardaites have disappeared from history but oral tradition claims that they are the ancestors of the Maronites of Lebanon, although this claim has been disputed. PCA and ADMIXTURE analyses failed to show any close relationship between Maniots and the Maronites (Supplementary Figure 5).

The question of Slavic ancestry of Tsakones

The Tsakones of the eastern slopes of Mount Parnon differ from all other Peloponnesians (Figures 1b and c) and from all other populations of Supplementary Figure 6 we have compared them. They used to speak a dialect of Doric origin which was not comprehended by the other Peloponnesians. Their name was considered by medieval authors to represent a corruption of the word Lacones (Tsakones=Lacones). Fallmayer argued against a Doric origin of the Tsakones and, instead, proposed that they were the descendants of a Slavic tribe that had migrated to Peloponnes before the flood of the Slavic settlers reached the peninsula. PCA (Figure 4c) and ADMIXTURE (Figure 4d) analyses argue against this hypothesis.

Quantitative assessment of the ancestry of Peloponnesians

To quantify the findings of the ADMIXTURE analyses, we employed a method for the meta-analysis of the ADMIXTURE output that treats the output as a set of vectors in a K-dimensional space (for a particular value of K between four and eight). Each population is then summarized by a single vector (using PCA) and vector space calculus is used in order to identify the percentage of ancestry of a target population that is captured by one or more reference populations. Notice that our choice to summarize each population by a single vector is akin to computing the mean ADMIXTURE output for a particular population. In most cases, ADMIXTURE returns a homogenous structure for a particular population and thus the top principal component is a good summary of the sample vectors returned by ADMIXTURE.

First we focused on the ADMIXTURE analysis of Figure 3c which includes seven Peloponnesian populations (Argolis, Corinthia, Achaia, Elis, Arcadia, Messenia and Laconia), four Slavic populations (Belarusians, Russians, Polish and Ukrainians), three Southern European populations (Italians, Basque and Andalusians) and the French. The results of Table 2 show that there is considerably more shared ancestry between the Peloponnesians and the French, Andalusians and Italians compared to the shared ancestry between the Peloponnesians and the Slavic populations. The average shared ancestry with French ranges from 39 to 42%; with Andalusians from 53 to 62%; and with the Italians from 85 to 96%. In contrast, the average shared ancestry with the Slavic populations is always <15%. Therefore, the Peloponnesians are genetically much more distinct from the Slavic populations and are much more similar to Southern European populations. We also observe that the Basques, (a population that is well-known to be isolated and genetically different from even its neighboring Iberian populations) are very distinct from all populations in our analysis. This is precisely why we included them in these ADMIXTURE meta-analyses: on average Basques share even its neighboring Iberian populations) are very distinct from all populations in our analysis. This is precisely why we included them in these ADMIXTURE meta-analyses: on average Basques share

Table 2 Shared ancestry between Peloponnesian populations and Slavic, Italian and other European populations

|                  | Belorussians | Russians | Polish | Ukrainians | French | Italians | Basque | Andalusians |
|------------------|--------------|----------|--------|-----------|--------|----------|--------|-------------|
| Argolis          | 0.7 (0.1)    | 1.6 (0.7)| 0.9 (0.4)| 1.0 (0.3)| 6.4 (3.5)| 25.3 (21.7)| 0.3 (0.2)| 7.6 (5.1)   |
| Corinthia        | 4.9 (5.1)    | 8.6 (6.9)| 6.8 (5.4)| 6.5 (5.7)| 16.4 (12.7)| 41.5 (32.5)| 0.6 (0.5)| 15.2 (11.1)|
| Achaia           | 5.7 (3.4)    | 10.9 (4.0)| 7.9 (3.7)| 8.0 (3.7)| 27.7 (4.8)| 58.0 (20.7)| 2.0 (1.4)| 27.0 (4.3)  |
| Arcadia          | 3.9 (1.7)    | 8.2 (2.1)| 5.0 (2.2)| 6.0 (2.2)| 26.7 (3.5)| 51.2 (4.6)| 1.5 (1.1)| 26.9 (3.5)  |
| Elis             | 0.2 (0.0)    | 0.9 (0.4)| 0.4 (0.1)| 0.6 (0.2)| 4.1 (2.9)| 14.2 (11.0)| 0.2 (0.1)| 5.3 (3.8)   |

*The first number for each pair of populations indicates the average shared ancestry for values of K between 4 and 8, while the number in parenthesis indicates the s.d.

Table 3 Shared ancestry between the populations of Mani and Tsakonia and Slavic, Italian and other European populations

|                  | Belorussians | Russians | Polish | Ukrainians | French | Italians | Basque | Andalusians |
|------------------|--------------|----------|--------|-----------|--------|----------|--------|-------------|
| Argolis          | 0.2 (0.1)    | 1.2 (1.2)| 5.8 (0.8)| 6.8 (1.1)| 39.1 (19.2)| 94.7 (4.8)| 2.8 (1.4)| 60.5 (5.9)  |
| Corinthia        | 5.9 (1.7)    | 13.0 (1.3)| 6.3 (1)| 7.5 (1.3)| 41.2 (18.5)| 94.9 (4.0)| 3.1 (1.7)| 62.0 (5.9)  |
| Achaia           | 6.5 (1.7)    | 13.8 (1.1)| 7.0 (0.8)| 8.1 (1.1)| 41.4 (18.4)| 94.8 (4.0)| 2.7 (1.4)| 61.3 (5.8)  |
| Arcadia          | 5.3 (1.8)    | 10.9 (2.4)| 5.2 (1.2)| 6.2 (1.5)| 39.1 (18.2)| 85.4 (14.6)| 2.4 (1.4)| 53.8 (9.1)  |
| Elis             | 6.1 (1.3)    | 13.1 (1.2)| 6.5 (0.8)| 7.6 (1.1)| 41.4 (18.3)| 95.0 (3.3)| 3.3 (1.7)| 61.6 (5.6)  |
| Messenia         | 6.7 (1.7)    | 14.4 (1.2)| 7.3 (0.9)| 8.5 (1.2)| 42.6 (18.4)| 95.2 (4.0)| 2.7 (1.3)| 61.8 (5.7)  |
| Laconia          | 4.8 (1.2)    | 11.4 (1.5)| 5.2 (0.9)| 6.4 (1.1)| 41.1 (14.6)| 96.1 (2.3)| 2.3 (1.4)| 59.8 (5.6)  |

*The first number for each pair of populations indicates the average shared ancestry for values of K between 4 and 8, while the number in parenthesis indicates the s.d.*
We next determined the shared ancestry between the five distinct Peloponnesian populations (Deep Mani, West and East Tayetos, North and South Tsakonia), the Slavs, the southern European populations, the French and the Basque. The ADMIXTURE plot of Figure 4d and the data of Table 3 show that the amount of shared ancestry between these five Peloponnesian populations and the Slavic populations is very low. The ancestry Deep Mani shares with Belarusians, Polish and Ukrainians ranges from 0.7 to 1.0%, East and West Tayetos share from 4.9 to 8.6% ancestry with the three Slavic populations which is five to eight times higher than that of Deep Mani but lower to the ancestry the other Peloponnesians share with the Slavs. Slightly lower, compared to the other Peloponnesians, is the ancestry shared between West/East Tayetos and the Russians (8.6–10.9%).

DISCUSSION

Our analysis of the genetic ancestry of the Peloponnesian populations and their relationships with the Slavs and other Europeans settles a historical controversy that has persisted for over 170 years. This controversy illustrates the problems historians face in their efforts to reconstruct population history on the basis of inadequate written sources. Fallmerayer based his theory of extinction of the medieval Peloponnesian Greeks on the writings of few early and two middle-medieval Byzantine authors. The early sources were very short comments in texts of 6th and 7th century historians and ecclesiastic authors. The middle-medieval documents were a letter by an 11th century Patriarch of Constantinople and the writings of tenth century Emperor Constantine Porphyrogenitus. Fallmerayer’s theory created sensation among historians. An early rebuttal was published by the Greek historian Paparrigopoulos who examined the same sources Fallmerayer have used to construct his theory and reached different conclusions, that is, that the Slavs did not reach the Greek proper during the sixth century and, when they arrived, they did not slaughter the local population. The many historians who have contributed to the extensive literature on this topic during the last century (partially due to isolation by distance. Geographic isolation explains the retention of their dialect. In ancient times the land of Tsakones, then called Cynouria, was inhabited by Doric speaking Ionians and because of its geography was most likely isolated from the other Peloponnesians.

Isolation by distance is also the likely explanation of the findings in the populations of Mani. Porphyrogenetus in his writing about the Slavs of Tayetos also asks what happened to the ancient inhabitants of Laconia, the Hellenes, who continued to adhere to the ancient Greek religion. He finds them withdrawn in the inhospitable, agriculturally poor and rocky area of southern Tayetos, the area which we refer to here as the Deep Mani. Ancient DNA studies could perhaps test whether there is any relationship between Maniots and ancient Laconians or between Tsakones and ancient Ionians.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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