# 1 The genetic origin of Daunians and the Pan-Mediterranean 

## southern Italian Iron Age context

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## Summary

The geographical location and shape of Apulia, a narrow land stretching out in the sea at the South of Italy, made this region a Mediterranean crossroads connecting Western Europe and the Balkans. Such movements culminated at the beginning of the Iron Age with the Iapygian civilization which consisted of three cultures: Peucetians, Messapians and Daunians. Among them, the Daunians left a peculiar cultural heritage, with one-of-a-kind stelae and pottery, but, despite the extensive archaeological literature, their origin has been lost to time. In order to shed light on this and to provide a genetic picture of Iron Age Southern Italy, we collected and sequenced human remains from three archaeological sites geographically located in Northern Apulia (the area

[^0]historically inhabited by Daunians) and radiocarbon dated between 1157 and 275 calBCE. We find that Iron Age Apulian samples are still distant from the genetic variability of modern-day Apulians, they show a remarkable genetic heterogeneity, even though a few kilometers and centuries separate them, and they are well inserted into the Iron Age Pan-Mediterranean genetic landscape. Our study provides for the first time a window on the genetic make-up of pre-imperial Southern Italy, whose increasing connectivity within the Mediterranean landscape, would have contributed to laying the foundation for modern genetic variability. In this light, the genetic profile of Daunians may be compatible with an autochthonous origin, with plausible contributions from the Balkan peninsula.

## Introduction

The Mediterranean's Iron Age populations, between 1100 and 600 BCE , lived in a time of previously unprecedented connectivity ${ }^{1}$. Although the technological advances in seafaring had allowed great opportunities for long-distance mobility just a few millennia earlier, it was during the Iron Age that the cosmopolitan role of Mare Nostrum arose, by promoting the spreading of cultures, goods, languages, technical advances as well as heterogeneous ancestral genetic components coming from far and wide ${ }^{1,2}$. Notable examples of such distant connections are the Greek and Phoenician settlements across the Central and Western Mediterranean shores beginning from the ninth and eighth centuries ${ }^{3,4}$.

The Italian peninsula and its Tyrrenian islands, where the Iron Age is conventionally marked as beginning $950 \mathrm{BCE}^{1,5}$, joined as well the cosmopolitan wave sweeping across Southern Europe, by hosting numerous trading posts along its shores. A patchwork of communities appeared in this period within the Italian borders, each characterized by unique and well-defined cultures and identities, which were later encapsulated and blurred by the Roman colonization. The shift
from Republican to Imperial Rome, with the consequent inclusion of non-Roman civilizations from and beyond the Italian peninsula, has already been shown to be connected with a major eastward genetic shift in Central Italian samples ${ }^{6}$. Genetic material from Imperial Romans from Central Italy is indeed the first to co-localize with contemporary Italians in a Principal Component space, pointing to this time period ( $\sim 200 \mathrm{BCE})$ as a crucial one in shaping contemporary Italian genetic makeup. Whether such a shift can also be observed in the rest of the Italian peninsula and how the Republic to Imperial transition may have impacted on the genetic landscape of local, preRoman populations remains an open question.

Despite the numerous written records and archaeological findings, questions about Iron Age populations, their origins and mutual relationships remain. Among the many groups occupying Italy in the Iron Age, the Daunians, a Iapygian population from northern Apulia, were first mentioned in the 7th-6th century $\mathrm{BCE}^{7,8}$. Similarly to their neighbouring populations, Peucetians and Messapians (living in central and southern Apulia, respectively), the name of the Daunians comes from ancient Greek documents and, given the absence of written Daunian records, the scant information we have on their social, political and religious life are wholly reliant upon the material record, such as their one-of-a-kind stelae ${ }^{8}$. For instance, we know that they were mainly farmers, animal breeders, horsemen and maritime traders with an established trade network extending across the sea with Illyrian tribes ${ }^{8-10}$. A fascinating aspect of this population, as opposed to their neighbours in Apulia, was their tenacious resistance to external influences. For instance, they did not acquire either social or cultural Hellenic elements and no Greek alphabet inscriptions have been found in their settlements. Indeed, they retained a strong cultural identity and political autonomy until the Roman arrival in the late 4th - early 3 rd century $\mathrm{BCE}^{8}$.

Despite the extensive archaeological literature, their origin has been lost to time and, as early as in the Hellenistic period, various legends already existed connecting them to either Illyria (an ancient region broadly identifiable with the Balkan peninsula), Arkadia (present-day Peloponnesus) or Crete ${ }^{7,8}$.

To shed light on this population and provide a glance of the Iron Age genetic landscape of the Southern Italian peninsula, we collected human remains from three Iron Age necropoleis geographically located within the area historically inhabited by Daunians: Ordona (ancient Herdonia), one of the largest settlements estimated at 600 hectares $^{8}$, Salapia, and San Giovanni Rotondo (Figure 1A). Here, we provide for the first time a genetic investigation of Iron Age Apulian associated with the Daunian culture, offering insight into the genetic landscape of preImperial Southern Italy.

## Results

We extracted DNA from 34 human skeletal remains (petrous bone $=23$ and teeth $=11$ ) from three necropoleis $($ Ordona $=19 ;$ Salapia $=12 ;$ San Giovanni Rotondo $=3$; hereafter ORD, SAL and SGR, respectively) at the Ancient DNA Laboratory of the Institute of Genomics, University of Tartu in Estonia (Data S1A). All three necropoleis are geographically located less than 50 km from each other in modern Apulia, Southeastern Italy. Ordona and Salapia are archaeologically dated to the Daunian period (6th-3rd century BCE) except one individual (ORD010) archaeologically dated to the Medieval time period (which in Europe dates approximately from the 5 th to the late 15 th century CE) (Figure 1B). Based on the museum record, the samples from the San Giovanni Rotondo necropole have been archaeological inferred to be from the Iron Age period. After screening 21 libraries at a low depth ( $\pm 20 \mathrm{M}$ reads per library), we
additionally sequenced to higher depth 16 libraries with endogenous DNA between $1.81-38.82 \%$ and mitochondrial DNA-based (mtDNA) contamination estimated at less than 2.89\% (Data S1A).

The sequencing runs were merged resulting in 16 individuals for genome-wide analysis: eight from Ordona, five from Salapia, and three from San Giovanni Rotondo. The final dataset includes individuals with an average genomic coverage between $0.031-0.995 \times$ and a number of single nucleotide polymorphisms (SNPs) overlapping with Human Origins 1240k between $\sim 40,000$ and $\sim 810,000$ (Data S1A).

Out of those 16 individuals, we selected 10 individuals based on their proximity within the principal component analysis (PCA) space (Figure 1C) for radiocarbon dating and estimated their age between 1157 and 275 calBCE with a median date of 521 calBCE (Data S1B). The radiocarbon dates confirm the archaeological dates of individuals from Salapia and Ordona as well as the Iron Age affiliation of the two San Giovanni Rotondo samples (SGR002 and SGR003). Two additional samples (ORD010 and SGR001) with a shift towards the Near East in the PCA (Figure 1C) were radiocarbon dated to $1078-1156$ calCE ( $95.4 \%$ ) and 670-774 calCE (95.4\%) respectively (Data S1B). Based on their dates, the samples were used as an external control for further analyses focusing on Iron Age Apulia (IAA).

We determined the mitochondrial DNA (mtDNA) haplotype of each individual and the Y chromosome (Ychr) haplogroup for 9 male individuals (See STAR methods and Data S1A and S1C-D). We found that the mtDNA haplotypes mostly belong to the mtDNA lineages H1, H5, K1, and U5, haplotypes found in previous studies of individuals from this time period in the Italian Peninsula ${ }^{6}$. Besides the Ychr haplogroup R1b, which is the most frequent haplogroup during the Bronze Age in the Italian Peninsula and on the islands Sardinia and Sicily ${ }^{4,6,11-13}$, we found the Ychr lineages I1-M253, I2d-M223, and J2b-M241. The haplogroup I2d-M223 was one of the main

Y chromosome lineages in Western Europe until the Late Neolithic whereas J2b-M241 first appears in the Bronze Age ${ }^{3,4,12,14,15}$. We found one Early Medieval individual (SGR001 (670-774 calCE ( $95.4 \%$ )) belonging to haplogroup I1-M253, which is common in Northern Europe and previously also detected in a 6th Century Langobard burial from North Italy ${ }^{16}$.

We used READ ${ }^{17}$ to identify pairs with first or second degree of genetic relatedness from autosomal data (See STAR methods and Data S1E-H for details, Figure S11). We found one firstdegree relationship between the two female individuals ORD001 (age: 9-11 years, 651-544 calBCE (95.4\%)) and ORD009 (age: 40-45 years, 570-406 calBCE (95.4\%)) from Ordona sharing the identical mtDNA haplogroup H5c and indicating a mother-daughter relationship (Data S1, Figure S1A-B).

## The making of modern Italians

To explore the genetic make-up of the IAA population, we performed a PCA projecting the ancient individuals onto the genetic variation of modern Eurasian samples (Figure 1C, Data S2 and Data S3). Our samples are largely scattered between modern peninsular Italians and Sardinians, and, in contrast to what was generally described ${ }^{18,19}$ for other European Iron Age populations (e.g., Northern_Europe_IA, Western_Europe_IA and Levant_IA in Figure 1C), they are still clearly distant from the genetic variability of modern-day inhabitants of Apulia. The downward shift of Iron Age Apulians from the present-day ones is further confirmed by the significantly negative $f 4$ (Modern Apulians, IAA; $X$, Mbuti), where $X$ is a Neolithic/Chalcolithic/Copper Age population (Figure 2A, Data S4). Within the remarkable heterogeneity reported by the PCA, which does not mirror the archaeological sites, the two medieval individuals are shifted towards modern Middle Eastern and Caucasus populations (ORD010 and SGR001), while the others are stretched along the PC2. This pattern partially
mirrors the chronological date with the most recent being more similar to present-day Southern Europeans, and is further strengthened when considering the PC3 distribution (Figure S2). Three samples located at the bottom of the PCA (ORD004, ORD019, SAL007) and one (SAL010) falling in the middle did not include modern Apulians among the top 25 results of an $f 3$ outgroup analysis (Figure S3). All of them showed an affinity to Copper and Bronze Age Italians ${ }^{11}$ as well as the Aegean and the Mediterranean worlds (including Minoans, Greece, Croatians, and Gibraltar). A similar distribution is mirrored in the Multi-Dimensional scaling (MDS) built from the $f 3$ outgroup measures, where the oldest IAA individual (SAL001; 1235-1048 calBCE (95.4\%)) lies farthest from the modern samples, while the medieval ones (ORD010: 1078-1156 cal CE (95.4\%) and SGR001: 670-774 cal CE (95.4\%)) are the closest (Figure S4).

The peculiar positioning of the IAA individuals casts doubt on when the major population shift resulting in modern Italian genetic composition took place. The shift towards the modern Italian genetic variability can be seen with the Republican-Imperial Roman samples ${ }^{6}$, the latter being more "similar" to modern Italians (Figure 1C). Whether Apulian individuals dating back to the Imperial phase would also show a repositioning towards modern genetic variability remains an open question, although the later, medieval samples of this study point in that direction.

## The pan-Mediterranean genetic landscape of Iron Age Apulia

The geographic location of Apulia, a narrow peninsula stretching out in the sea at the South of Italy, has made this region an important Mediterranean crossroads connecting Western Europe, the Balkans, the Aegean, and Levant worlds. This is reflected in the PCA where IAA individuals are closely related to other Iron Age populations from the Mediterranean and surrounding areas (e.g., Montenegro, Bulgaria and Sardinia) (Figure 1C and Figure S4). Nomadic or cosmopolitan
groups scatter like IAA: three Punic individuals from Sardinia (Italy_Sardinia_IA_Punic ${ }^{3}$ ), three Moldova Scythians already reported to be genetically similar to Southern Europeans ${ }^{20}$, Spanish individuals from the Hellenistic and the Romans periods ${ }^{21}$ and an individual from the $12^{\text {th }}$ century Iron Age Ashkelon ${ }^{22}$ which clusters with ORD001.

In order to shed light onto the genetic composition of the IAA individuals, we modelled them as a combination of the main ancestries documented across Western Europe at that time: Western Hunter-Gatherers (WHG), Anatolian Neolithic (AN), Steppe-related and, interchangeably, Caucasus Hunter-Gatherers (CHG) or Iranian Neolithic (IN) using the qpWave/qpAdm framework (Figure 2B, Figure S5A, STAR Methods). Broadly, the contributions of such ancestries to the genetic variability of ancient European populations vary according to their geographical positions: in particular, northernmost locations received higher proportions of WHG, Steppe-related ancestry and, consequently, CHG ancestries, while Southern European groups carried variable Iranian Neolithic or CHG traces ${ }^{23}$. In view of this, we observed that while the IAA individuals could generally be modelled as a two-way admixture between AN and Steppe ( $0.63 \pm 0.08$ and $0.37 \pm 0.08$, respectively), the alternative model $\mathrm{AN}+\mathrm{CHG} / \mathrm{IN}$ could also fit for a subset of them, particularly in case of the samples ORD004, ORD010 and SAL010 with higher or comparable p-values (Figure S5A, first row with two sources). When three or four sources were tested, the presence of WHG ancestry in the majority of our individuals emerges, which, together with AN, Steppe and CHG/IN, forms a supported model for IIA samples (Figure 2B, Figure S5A and Data S5). Notably, for the individuals stretching downwards in the PCA (ORD004, ORD019, SGR002 and the Medieval ORD010) a three-way admixture involving AN, Steppe and CHG/IN is generally preferable. To better understand the putative contribution of more recent populations, we modelled our samples with base sources (WHG, AN, Steppe-related and CHG/IN) and,
alternatively, Minoans (Odigitria and Lasithi), Amhara_NAF and Roman Republicans (Figure 2C, Figure S5B-D, STAR Methods). Amhara_NAF can be used as a proxy for the Non-African component in modern Ethiopian individuals that was tentatively linked to the Sea People, a Bronze Age nomadic seafaring population ${ }^{22,24}$. Together with Minoans and Roman Republicans, this component can be broadly modelled as a Pan-Mediterranean population (constituted by AN and IN/CHG components) with the addition of WHG and Steppe-related ancestry in Roman Republicans. When modelled also with Minoans and Amhara_NAF, which roughly proxies the same ancestral signature, the majority of the samples required an additional $\mathrm{CHG} / \mathrm{IN}$ contribution (two-way admixtures in Figure S5B,C) as well as Steppe-related and WHG. We further observed that, as previously seen, the WHG contribution is less clear in those samples stretching downwards in the PCA. While the CHG/IN additional contribution may simply proxy the presence of Stepperelated ancestry in IAA, the absence of which in Minoans has already been reported ${ }^{25}$, the same can not be said about Roman Republicans (two-way admixtures in Figure S5D), which harboured a considerable amount of Steppe component ${ }^{6}$. However, this signature is not confirmed with $f 4$ analyses (Figure S6 B,C), where just Mycenaean groups report less CHG ancestry than our samples.

The broader picture emerging from qpAdm analyses is the pervasive presence of coeval Italian ancestries (Rome Republicans), as well as previous Bronze Age sources (Minoans and Sea People) spread far and wide across the Mediterranean Sea. In such a melting pot scenario, the peculiar genetic heterogeneity of IAA individuals, who lived in close temporal and geographical proximity, stand out (Figure 1C, Figure S4, Figure S7). An example of the cosmopolitan nature of Iron Age Mediterranean is the first-degree relationship between ORD009 (mother) and ORD001 (daughter), whose positions in the PCA strikingly differ with the individual ORD001 being
stretched towards Middle Eastern and Caucasus modern populations, as a consequence of the foreign origin of the father (Figure 1C and Figure S4). F4 analyses in the form $f 4($ ORD009, ORD001; X, Mbuti) report a slightly significant (Z-score between -2 and -3) excess of Greece_N, Portugal_LN_C, Lebanon_Roman and Italy_Sicily_EBA in ORD001, which may explain its eastward shift (Figure S8A). Moreover, ORD001, together with SGR002, but not ORD009 harboured more CHG when compared to Lebanon_Hellenistic and Lebanon_IA3 samples, respectively (Figure S8C), as well as an increase in Lebanon_Hellenistic traces when compared with modern Apulians (data not shown).

We also investigated whether the PCA scattering was due to varying African or Levantine contributions with $f 4$ (Rome Republican, IAA, Levant_N/YRI, Mbuti) and tried the same on Medieval ancient Apulians (ORD010 and SGR001). However, none of the tested ancient Apulians shows a significant excess of YRI ancestry when compared to the contemporary Roman Republicans, even though ORD014, SAL007 and SAL011 show negative $f 4$ values with a Z-score between 2 and 3 (Figure S8B).

## The origin of Daunians

The genetic heterogeneity of IAA individuals connected to the Daunian population and the cosmopolitan landscape of Iron Age Mediterranean populations hinder a full reconstruction of the demographic processes leading to the Daunians. Nevertheless, a few milestones can be spotted.

When we performed $f 3$ analyses to investigate the nearest possible source for each IAA individual using Minoans, Iron Age Croatians and the local Roman Republicans (Figure 3A), we found that none of the IAA individuals shows higher affinities with Minoans. Three of them, clustered close to modern Italians in the PCA (ORD001, ORD014 and SGR003, Figure 1C), show
higher affinity with the Iron Age Croatian sample (ORD004 followed this pattern too, but with lower $f 3$ values). However, the remaining majority are closest to the Roman Republicans, which can be interpreted as representative of local Iron Age peninsular Italy ancestry.

Moreover, the WHG contribution, which was a necessary component to explain IAA and Roman Republicans according to qpAdm output, is absent from Minoans and Iron Age Croatians, thus making it a putative signature of an at least partial autochthonous origin (Figure 2B,C and Figure S5). These results are confirmed for Minoans, but not for Croatia_EIA, by the $f 4$ analyses (Figure 3B, Figure S6A and Data S6). Indeed, the significantly negative $f 4$ (Minoans, IAA; WHG, Mbuti) (Figure 3B and Figure S6A), as well as the f4(Greece_Minoan_Lassithi/Greece_BA_Mycenaean/Greece_BA_Mycenaean_Pylos, IAA; X, Mbuti), reported a significant excess of WHG ancestry ( $X$ being Tagliente2, Loschbour, Italy_Central_M, Switzerland_Bichon and so on) in all IAA individuals, with the exception of ORD004, SAL001 and SGR003 (also the medieval samples ORD010 and SGR001 deviate from this pattern, Data S6). An excess of Bronze Age Steppe-related component, which at that time was already present along the Italian peninsula ${ }^{11}$, was clearly present in SGR002, ORD006 and ORD009 by the $f 4$ (IAA, Greece_Minoan_Lassithi; X, Mbuti) (Data S6).

The excess of WHG ancestry, tentatively suggesting a local origin, is somewhat blurred by the genetic similarity of the two most probable sources - Illyrians (Croatia EIA) and an autochthonous one (Roman Republicans), which together make part of the same Mediterranean continuum. Indeed, while no significantly negative f4(Croatia_EIA, IAA; X, Mbuti) values have been found (with the exception of $X$ being Italy_C_BA, Italy_Iceman_CA, Portugal_MBA and Anatolia_MLBA in SAL007, which retains the highest proportion of AN in qpAdm (Figure 2B and Figure S5), the same pattern is obtained when IAA individuals are compared with Roman

Republicans (f4(Roman Republicans, IAA; X, Mbuti), (Data S6)). Imperial individuals from a few centuries later (f4(Rome_Imperial, IAA; X, Mbuti)) show quite the opposite pattern: no positive results have been produced with the exception of ORD004 (X=Lebanon_Roman), SAL001 (X=Italy_Sardinia_Roman_o), SAL010 (X=Russia_LateMaikop) and ORD010. Conversely, the negative $f 4$ values point toward WHG, Neolithic and Bronze Age Steppe-related ancestries (Data S6).

Another signal coming from qpAdm analyses is the apparent excess of CHG ancestry in IAA; however, the predominant contribution of CHG to the Steppe-related ancestry that, by the Iron Age, had already spread to the Mediterranean area makes it hard to properly detect a CHG signature independent from the Steppe wave, possibly brought by pan-Mediterranean influxes. When directly investigated with an $f 4$ framework, IAA shows generally more CHG than Mycenaean, less CHG than contemporary Croatian_EIA and, in some cases (ORD019, SGR002 and the Medieval SGR001 with Z-scores higher than 2) more CHG than older Croatian samples (_N and _MN) (Figure S6).

## Discussion

The new genomic sequences Daunian samples reveal that Iron Age (pre-Imperial) Southern Italy (Apulia) can be placed within a Pan-Mediterranean genetic continuum that stretches from Crete (Minoans ${ }^{25}$ ) and the Levant (Sea People ${ }^{22,24}$ ) to the Republican Rome and the Iberian Peninsula ${ }^{6}$, mainly composed by AN and IN/CHG genetic features with the addition of WHG and Steppe-related influences in Continental Italy. Pre-Imperial Italian populations, being part of this broader landscape, are not directly superimposable with contemporary Italians which instead seem to be influenced by the homogenizing effect of Imperial Roman and late Antiquity events.

Within the described Pan Mediterranean landscape, the IAA/Daunians show a compelling heterogeneity, and the highest genetic affinity to Republican Romans and Iron Age Croatians, while Minoans and other Iron Age Greek samples show absent or reduced WHG contribution when compared to IAA. This makes a Cretan or Arkadian origin less likely, even though some tales have connected them with the Greek hero Diomedes and many ancient historians have claimed such origins for their neighbour Messapians and Peucetians.

The Daunians maintained strong commercial and political relations with the Illyrian people, controlling together the area spanning from the Dalmatia to the Gargano peninsula ${ }^{10}$ and had many cultural affinities with them ${ }^{26}$. The material culture, involving peculiar anthropomorphic statue stelae, has provided some information on Daunian culture and may also help in unravelling their mysterious origin. In particular, the forearm decorations on a female stela have been interpreted as tattoos and, while tattooing practices were considered barbarian among the Greeks ${ }^{27}$, they were customary in populations from Tracia and Illyria and, more generally, among the women of status from the Balkans ${ }^{8,28}$.

It is not clear whether these connections indicate a movement of people or a sharing of cultural ideas and a conclusive answer to the origin of the Daunians remains elusive. From a parsimony perspective, the genetic results point to an autochthonous origin, although we cannot exclude additional influences from Croatia (ancient Illyria), as described by available historical sources and by the material remains ${ }^{8,29}$.

## Table

Table 1. Archaeological information, dating, genome coverage, genetic sex, mtDNA, and Y chromosome haplogroups of the individuals of this study selected for genome-wide analysis. Dates in BP are raw radiocarbon dates, calibrated dates are the $95.4 \%$ probability and were calibrated using IntCal2030. Gen.genetic; HG-haplogroup. See also Data S1.

|  |  |  | Genome | Gen. | mtDNA | $\mathbf{Y}$ <br> chromosome | no. SNPs |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Individual | Site | Date | Coverag <br> e | Sex | HG | HG |  |  |
| ORD001 | Ordona | $2516 \pm 25 \mathrm{BP} ; 651-$ <br> 544 calBCE | 0.0396 | XX | H5c | - | 53,277 |  |
| ORD004 | Ordona | $\begin{aligned} & 2374 \pm 21 \mathrm{BP} ; 490- \\ & 394 \text { calBCE } \end{aligned}$ | 0.0429 | XX | U8b1b1 | R1b-M269 | 56,070 |  |
| ORD006 | Ordona | $\begin{aligned} & 2476 \pm 29 \mathrm{BP} ; 770- \\ & 476 \mathrm{calBCE} \end{aligned}$ | 0.12 | XY | H+16291T | - | 158,562 |  |
| ORD009 | Ordona | $\begin{aligned} & 2433 \pm 27 \mathrm{BP} ; 570- \\ & 406 \mathrm{calBCE} \end{aligned}$ | 0.995 | XX | H5c | - | 805,966 |  |
| ORD010 | Ordona | $\begin{aligned} & 984 \pm 25 \mathrm{BP} ; 1078- \\ & 1156 \text { calCE } \end{aligned}$ | 0.91 | XX | X2i | - | 769,336 |  |
| ORD011 | Ordona | not dated | 0.0889 | XY | H1e | R1b-P312 | 116,800 |  |
| ORD014 | Ordona | $\begin{aligned} & 2438 \pm 25 \text { BP; } 570- \\ & 408 \text { calBCE } \end{aligned}$ | 0.108 | XY | I5a2+16086C | J2b2-L283 | 141,736 |  |
| ORD019 | Ordona | not dated | 0.0484 | XY | T2e | I2d- <br> Z2093/Y3670 | 61,292 |  |
| SAL001 | Salapia | $\begin{aligned} & 2946 \pm 30 \mathrm{BP} ; 1235- \\ & 1048 \text { calBCE } \end{aligned}$ | 0.049 | XY | H1+16189! | J2b-M241 | 64,574 |  |
| SAL003 | Salapia | $\begin{aligned} & 2287 \pm 24 \mathrm{BP} ; 401- \\ & 354 \text { calBCE } \end{aligned}$ | 0.113 | XX | K1a2a | - | 144,490 |  |


| SAL007 | Salapia | not dated | 0.0529 | XX | K1a+195T! | - | 64,034 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| SAL010 | Salapia | not dated | 0.031 | XY | U5a1 | J2b-M241 | 39,174 |
| SAL011 | Salapia | $\begin{aligned} & 2241 \pm 23 \mathrm{BP} ; 313- \\ & 206 \text { calBCE } \end{aligned}$ | 0.0496 | XY | U5b1 | I2d-M223 | 62,155 |
| SGR001 | San <br> Giovanni <br> Rotondo | $\begin{aligned} & 1285 \pm 23 \mathrm{BP} ; 670- \\ & 774 \mathrm{calCE} \end{aligned}$ | 0.044 | XY | U3a | I1-M253 | 59,213 |
| SGR002 | San <br> Giovanni <br> Rotondo | $\begin{aligned} & 2451 \pm 22 \mathrm{BP} ; 591- \\ & 415 \mathrm{calBCE} \end{aligned}$ | 0.102 | XY | U5b1d1 | R1b-M269 | 131,437 |
| SGR003 | San <br> Giovanni <br> Rotondo | $\begin{aligned} & 2205 \pm 26 \mathrm{BP} ; 368- \\ & 195 \text { calBCE } \end{aligned}$ | 0.0572 | XX | H1+16311T! | - | 76,687 |

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## STAR Methods

## Resource availability

## Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact: Serena Aneli (serena.aneli@unipd.it)

## Materials Availability

This study did not generate new unique reagents.

## Data and Code Availability

The DNA sequences generated during this study are available at the European Nucleotide Archive (ENA; [link]) at the accession number [accession number]. The data are also available in PLINK format through the data depository of the EBC (http://evolbio.ut.ee).

## Experimental model and subject details

The ancient human remains analysed in this work belong to the collections of the Museum of Anthropology, University of Padua and have been found and catalogued by different archaeological campaigns carried out during the twentieth century.

## Ordona

The necropolis of Herdonia (today's Ordona, within the Apulian Foggia province in Italy) was studied by different archaeological campaigns interested in the Daunians, Roman as well as the medieval settlements in 1978 and $1981^{31}$. The human remains collected on such occasions were later extensively catalogued and studied and new paleopathological evidence was also brought back to light ${ }^{32}$. Inhabited starting from the Neolithic period, Herdonia became an important Daunian center from the 6th century BCE.

For this study, in collaboration with the University of Padova and Museum of Anthropology of Padova, samples of human remains (petrous bone=19) were taken. The samples ORD001, ORD004, ORD006, ORD009, ORD010, and ORD014 were dated at ${ }^{14} \mathrm{CHRONO}$ Centre for Climate, the Environment, and Chronology in Belfast, UK (Data S1B).

## Salapia

The necropolis of Salapia, an ancient town located 10 km from contemporary Cerignola, within the Apulian Foggia province in Italy. The osteological samples were brought by Prof. Santo Tiné to the Museum of Anthropology of the University of Padova with no further information on the archaeological context, and osteological studies were carried out by Cleto Corrain and colleagues in $1971^{33}$. From this site, samples of 12 human remains (petrous bone $=5$, teeth=7) were taken. The samples SAL001, SAL003, and SAL011 were dated at ${ }^{14} \mathrm{CHRONO}$ Centre for Climate, the Environment, and Chronology in Belfast, UK (Data S1B).

## San Giovanni Rotondo

The San Giovanni Rotondo samples come from the osteo-archaeological collection of the Museum of Anthropology of the University of Padova and are not associated to any further record with the exception of a broad "Iron Age" archaeological label and may be part of the samples brought to the Museum by Prof. Santo Tiné in the 1960s. From this site, samples of human remains (petrous bone=3) were taken. The samples SGR001, SGR002, and SGR003 were dated at ${ }^{14} \mathrm{CHRONO}$ Centre for Climate, the Environment, and Chronology in Belfast, UK (Data S1B).

## Method Details

All of the laboratory work was performed in dedicated ancient DNA laboratories at the Estonian Biocentre, Institute of Genomics, University of Tartu, Tartu, Estonia. The library quantification
and sequencing were performed at the Core Facility of the Institute of Genomics, Tartu, Estonia. The main steps of the laboratory work are detailed below.

## DNA extraction

In total 34 samples from human remains were extracted for DNA analysis (Data S1).

The first layer of pars petrous was removed with a sterilised drill bite to avoid exogenous contamination. A 10 mm core of the inner ear was sampled from the pars petrous. The drill bits and core drill were sterilised in between samples with $6 \%(\mathrm{w} / \mathrm{v})$ bleach followed by distilled water and then ethanol rinse. Root portions of teeth were removed with a sterile drill wheel.

The root and the petrous portions were soaked in $6 \%(\mathrm{w} / \mathrm{v})$ bleach for 5 minutes. Samples were rinsed three times with $18.2 \mathrm{M} \Omega \mathrm{cm} \mathrm{H}_{2} \mathrm{O}$ and soaked in $70 \%$ (v/v) Ethanol for 2 minutes. The tubes were shaken during the procedure to dislodge particles. The samples were transferred to a clean paper towel on a rack inside a class IIB hood with the UV light on and allowed to dry for two to three hours.

Afterwards, the samples were weighed to calculate the accurate volume of EDTA (20x EDTA [ $\mu \mathrm{l}$ ] of sample mass [mg]) and Proteinase K ( 0.5 x Proteinase $\mathrm{K}[\mu \mathrm{l}$ ] of sample mass [mg]). EDTA and Proteinase K were added into PCR-clean 5 ml or 15 ml conical tubes (Eppendorf) along with the samples inside the IIB hood and the tubes were incubated 72 h on a slow shaker at room temperature.

The DNA extracts (of root portions and pars petrous portions) were concentrated to $250 \mu \mathrm{l}$ using the Vivaspin® Turbo 15 (Sartorius) and purified in large volume columns (High Pure Viral Nucleic Acid Large Volume Kit, Roche) using 2.5 ml of PB buffer, 1 ml of PE buffer and $100 \mu \mathrm{l}$
of EB buffer (MinElute PCR Purification Kit, QIAGEN). For the elution of the endogenous DNA, the silica columns were transferred to a collection tube to dry and followed in 1.5 ml DNA lo-bind tubes (Eppendorf) to elute. The samples were incubated with $100 \mu \mathrm{~EB}$ buffer at 37 C for 10 minutes and centrifuged at $13,000 \mathrm{rpm}$ for two minutes. After centrifugation, the silica columns were removed and the samples were stored at -20 C . Only one extraction was performed per extraction for screening and $30 \mu \mathrm{l}$ used for libraries.

## Library preparation

Sequencing libraries were built using NEBNext ${ }^{\circledR}$ DNA Library Prep Master Mix Set for $454^{\mathrm{Tm}}$ (E6070, New England Biolabs) and Illumina-specific adaptors ${ }^{34}$ following established protocols ${ }^{34-}$ ${ }^{36}$. The end repair module was implemented using $18.75 \mu \mathrm{l}$ of water, $7.5 \mu \mathrm{l}$ of buffer and $3.75 \mu \mathrm{l}$ of enzyme mix, incubating at $20^{\circ} \mathrm{C}$ for 30 minutes. The samples were purified using $500 \mu \mathrm{~PB}$ and $650 \mu l$ of PE buffer and eluted in $30 \mu \mathrm{~EB}$ buffer (MinElute PCR Purification Kit, QIAGEN). The adaptor ligation module was implemented using $10 \mu \mathrm{l}$ of buffer, $5 \mu \mathrm{l}$ of T4 ligase and $5 \mu \mathrm{l}$ of adaptor mix ${ }^{34}$, incubating at $20^{\circ} \mathrm{C}$ for 15 minutes. The samples were purified as in the previous step and eluted in $30 \mu$ l of EB buffer (MinElute PCR Purification Kit, QIAGEN). The adaptor fillin module was implemented using $13 \mu \mathrm{l}$ of water, $5 \mu \mathrm{l}$ of buffer and $2 \mu \mathrm{l}$ of Bst DNA polymerase, incubating at $37{ }^{\circ} \mathrm{C}$ for 30 and at $80^{\circ} \mathrm{C}$ for 20 minutes. Libraries were amplified using the following PCR set up: $50 \mu \mathrm{l}$ DNA library, 1 X PCR buffer, $2.5 \mathrm{mM} \mathrm{MgCl} 2,1 \mathrm{mg} / \mathrm{ml} \mathrm{BSA}, 0.2 \mu \mathrm{M}$ inPE1.0, 0.2 mM dNTP each, $0.1 \mathrm{U} / \mu \mathrm{l}$ HGS Taq Diamond and $0.2 \mu \mathrm{M}$ indexing primer. Cycling conditions were: $5^{\prime}$ at 94 C , followed by 18 cycles of 30 seconds each at $94 \mathrm{C}, 60 \mathrm{C}$, and 68 C , with a final extension of 7 minutes at 72C. The samples were purified and eluted in $35 \mu l$ of EB buffer (MinElute® PCR Purification Kit, QIAGEN). Three verification steps were implemented to make sure library preparation was successful and to measure the concentration of dsDNA/sequencing
libraries - fluorometric quantitation (Qubit, Thermo Fisher Scientific), parallel capillary electrophoresis (Fragment Analyser, Agilent Technologies) and qPCR.

## DNA sequencing

DNA was sequenced using the Illumina NextSeq500/550 High-Output single-end 75 cycle kit. As a norm, 15 samples were sequenced together on one flow cell; additional data was generated for 16 samples to increase coverage (Data S1).

## Quantification and statistical analysis

## Mapping

Before mapping, the sequences of the adapters, indexes, and poly-G tails occuring due to the specifics of the NextSeq 500 technology were cut from the ends of DNA sequences using cutadapt$1.11^{37}$. Sequences shorter than 30 bp were also removed with the same program to avoid random mapping of sequences from other species. The sequences were aligned to the reference sequence GRCh37 (hs37d5) using Burrows- Wheeler Aligner (BWA 0.7.12) ${ }^{38}$ and the command mem with re-seeding disabled.

After alignment, the sequences were converted to BAM format and only sequences that mapped to the human genome were kept with samtools $1.3^{39}$. Afterwards, the data from different flow cell lanes were merged and duplicates were removed using picard 2.12 (http://broadinstitute.github.io/picard/index.html). Indels were realigned using GATK $3.5^{40}$ and reads with a mapping quality less than 10 were filtered out using samtools $1.3^{39}$.

## aDNA authentication

As a result of degradation over time, aDNA can be distinguished from modern DNA by certain characteristics: short fragments and a high frequency of $\mathrm{C}=>\mathrm{T}$ substitutions at the $5^{\prime}$ ends of sequences due to cytosine deamination. The program mapDamage $2.0^{41}$ was used to estimate the frequency of $5^{\prime} \mathrm{C}=>\mathrm{T}$ transitions. Rates of contamination were estimated on mitochondrial DNA by calculating the percentage of non-consensus bases at haplogroup-defining positions as detailed in ${ }^{42}$. Each sample was mapped against the RSRS downloaded from phylotree.org and checked against haplogroup-defining sites for the sample-specific haplogroup (Data S1A).

Samtools $1.9^{39}$ option stats was used to determine the number of final reads, average read length, average coverage etc. The final average endogenous DNA content of all individuals (proportion of reads mapping to the human genome) was $10.39 \%(0.09-38.82 \%)$ (Data S1A).

## Calculating genetic sex estimation

Genetic sex was calculated using the methods described $\mathrm{in}^{43}$, estimating the fraction of reads mapping to Y chromosome out of all reads mapping to either X or Y chromosome. Additionally, sex was determined using a method described in ${ }^{44}$, calculating the X and Y ratio by the division of the coverage by the autosomal coverage. Here, the sex was calculated for samples with a coverage $>0.01 \times$ and only reads with a mapping quality $>10$ were counted for the autosomal, X, and Y chromosome (Data S1A).

## Determining mtDNA haplogroups

Mitochondrial DNA haplogroups were determined using Haplogrep2 on the command line. For the determination, the reads were re-aligned to the reference sequence RSRS and the parameter -
rsrs were given to estimate the haplogroups using Haplogrep2 ${ }^{45,46}$ (Data S1A). Subsequently, the identical results between the individuals were checked visually by aligning mapped reads to the reference sequence using samtools $0.1 .19^{39}$ command tview and confirming the haplogroup assignment in PhyloTree. Additionally, private mutations were noted for further kinship analysis. The polymorphisms were estimated using the online platform of haplogrep2. Here, the variant calling files (vcf) were uploaded to the online platform and the known polymorphism in the RSRS were converted to rCRS (Data S1D).

## $Y$ chromosome variant calling and haplotyping

A total of 161,140 binary Y chromosome SNPs that have been detected as polymorphic in previous high coverage whole $Y$ chromosome sequencing studies ${ }^{47-49}$ were called in 9 male individuals with more than $0.01 \times$ autosomal coverage using ANGSD-0.916 ${ }^{47-49}$ '-doHaploCall’ option. A subset of 826 sites had at least one of the 9 individuals with a derived allele (DATA C- Table Y). Basal haplogroup affiliations of each sample were determined by assessing the proportion of derived allele calls in a set of primary haplogroup defining internal branches, as defined in ${ }^{47}$, using 1677 informative sites.

## Kinship analysis

All newly generated individuals with an average human coverage more than $0.03 \times$ were selected to assess kinship relationships up to the 3rd degree. We divided the individuals in different groups regarding their geographically area and resulting relationships. First, we were called all individuals using ANGSD- $0.916^{50}$ command --doHaploCall to sample a random base for the positions that are present at MAF>0.1 in the 1000 Genomes GBR population ${ }^{51}$ giving a total of $4,045,514$ SNPs for autosomal kinship analysis. The ANGSD output files were converted to .tped format and used as
an input for kinship analyses with READ ${ }^{17}$. First, we tested all individuals together and found one 1st degree relationship between ORD001 and ORD009 and two 2nd degree relationships between ORD004, SAL007, and SAL011. We tested those relationships using all individuals from Ordona including SAL007 and SAL011 and all individuals from Salapia including ORD004. We compared the estimated mtDNA haplogroups, radiocarbon dates, genetic sexes, average human coverages, and the age of death from the individuals to confirm or dismiss relationships. (Data S1, Figure S1). Based on the outputs, we dismissed the possible 2nd degree relationships.

## Dataset preparation and pre-processing for autosomal analysis

We assembled a genome-wide dataset of ancient and modern samples by converting the following datasets, where needed, in PLINK format using convertf from the EIGENSOFT 8.0.0 ${ }^{52}$ and merging them together with PLINK $1.9^{53}$ : i) the ancient individuals and the Mbuti Congolese individuals $\left(\mathrm{HGDP}^{54}\right)$ from the " 1240 K " dataset from Dr David Reich laboratory (https://reich.hms.harvard.edu/allen-ancient-dna-resource-aadr-downloadable-genotypes-present-day-and-ancient-dna-data, version 44.3 release), ii) the modern samples from " $1240 \mathrm{~K}+\mathrm{HO}$ " of the same release, iii) the Italian Chalcolithic and Bronze Age samples produced by a recent study ${ }^{11}$, iv) the genotypes of an Italian hunter-gatherer from the Late Epigravettian site of Riparo Tagliente dated around $17 \mathrm{kya}^{55}, \mathrm{v}$ ) additional genome-wide data of 129 modern-day Italian samples covering the entire country ${ }^{56}$, vi) genome-wide data of 217 present-day Apulian individuals (available at this link https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE44974) and 48 haploid genomes representing the Eurasian component of modern Ethiopians (here called Amhara_NAF for "Non African Fraction" ${ }^{24}$ ). We excluded the ancient samples showing issues in their group assignment (i.e., "Ignore" in the " 1240 K " dataset) or in their DNA quality (e.g, those not containing the string "PASS", with the exception of the Iceman individual, or those with less than
$5,000 \mathrm{SNPs}$ ) and one of a pair of related samples. We further refined the list of ancient samples by retaining only those coming from geographical regions relevant for this study (Data S2) 3,4,6,12,14-16,18-23,25,42,44,57-102. The same geographical filter was applied on the modern samples from " $1240 \mathrm{~K}+\mathrm{HO}$ " and only those flagged as "PASS (genotyping)" have been selected in order to minimize variation due to different genotyping techniques (Data S3) ${ }^{23,54,72,103,104}$. In case of duplicated samples, we selected the item showing the highest number of SNPs among the 1240K SNP set. Finally we added the newly generated ancient Apulian samples to the merge.

We used two different SNPs set for the following analyses. The autosomal "1240K" SNP set ( $1,150,639$ SNPs) was used for the analyses involving just ancient samples, such as the $f$-statistics and qpAdm analyses (see below). Conversely, the autosomal "HO" SNP set was used for those analyses relying also on modern populations (e.g., PCA and ADMIXTURE). In the latter case, we further refine the "HO" SNP set by removing monomorphic SNPs and those with more than $5 \%$ missing count in modern populations with PLINK, thus obtaining 503,062 SNPs.

## Principal component analysis

We performed principal component analysis using the program smartpca implemented in EIGENSOFT software 8.0.0 $0^{52}$ with the parameters "lsqproject" and "shrinkmode". In particular, we projected the ancient samples' genetic variation onto the principal components inferred from the modern samples of the " $1240 \mathrm{~K}+\mathrm{HO}$ " dataset. Modern Italian samples coming from ${ }^{56}$, the newly genotyped ones from modern Apulia and the Amhara_NAF were also projected.

## Admixture

We used the same dataset of modern and ancient individuals presented in the PCA to explore the genetic components of each individual. For the analysis, we exploited the model-based algorithm implemented in ADMIXTURE ${ }^{105}$ projecting ancient individuals (-P flag) into the genetic structure
calculated on the modern dataset, due to missing data in the ancient individuals. We performed unsupervised Admixture for $\mathrm{K} \in\{2 . .10\}$ for modern individuals and used the "per-cluster" inferred allele frequencies to project the ancient individuals. We visualised the Q output using R $3.6^{106}$.

## f4 statistics

We performed a series of $f 4$ statistics using the program qpDstat (option f4mode) implemented in the software ADMIXTOOLS 7.0.2 ${ }^{54}$ in different forms. In particular, we grouped together ancient published samples from the same cultural/archeological assemblages (labels used for f 3 and f 4 analyses can be found in the column "Analysis.Label" of Data S2). Conversely, ancient Apulian samples were both analysed as individual samples and grouped together according to their archeological age (Iron Age samples were grouped together under the acronym "IAA", Iron Age Apulians). 10 individuals from the Mbuti population from Congo coming from the " 1240 K " dataset were used as an outgroup.

## Outgroup f3 statistics

To investigate genetic affinities between ancient Apulian samples with other ancient and modern human populations, we performed a series of Outgroup $f 3$ analyses in the form $f 3(\mathrm{X}, \mathrm{Y}$; outgroup=Mbuti) using the qp3Pop function implemented in ADMIXTOOLS 7.0.2 ${ }^{54}$. As abovedescribed we used the "Analysis.Label" groups for the ancient samples while we used the "Group.Label" already present in the " $1240 \mathrm{~K}+\mathrm{HO}$ " annotation file for modern samples.

We also computed the distance matrices from the Outgroup $f 3$ results comparing ancient Apulian samples (both Iron and Middle Age samples) with other published ancient samples by subtracting the f 3 values from 1 and we performed a classical (metric) Multi-Dimensional Scaling (MDS) with default options using the function $c m d s c a l e$ in the stats package of R (version 4.0.4 ${ }^{107}$ ).

## qpAdm

We exploited the $q p W a v e / q p A d m$ framework within the software ADMIXTOOLS 7.0 .2 to investigate the genetic composition of ancient Apulian samples and other ancient samples of interest with respect to human ancestries that contributed to coeval European population in the Mediterranean area. In particular, we tested a number of left sources from 2 to 5 in combination with a list of reference sources (right). We used the options 'allsnps=YES'" and "inbreed=NO" (the last option was used because some populations were composed of just one haploid individual). We tested four different sets of left sources. The "base" set included Western hunter-gatherers (Loschbour_published.DG), Anatolian Neolithic (26 samples flagged as "Anatolia_N" in the Analysis.Label column of Data S2), Iranian Neolithic (samples Iran_GanjDareh_N), Caucasus hunter-gatherers (KK1.SG and SATP.SG) and Yamnaya (10 Russia_Samara_EBA_Yamnaya samples). To this set of base sources we separately added as left sources: Minoans (both Greece_Minoan_Lassithi and Greece_Minoan_Odigitria, but taken individually), Amhara_NAF and Roman Republican samples (Italy_Central_Republic_IA). In each run, we consistently used the same set of right sources (if present in the left set, they were removed from the right): Russia_Ust_Ishim_HG_published.DG (which was always kept as the first of the list), Russia_Kostenki14, Russia_AfontovaGora3, Eastern hunter-gatherers (I0061=Russia_HG_Karelia, I0124=Russia_HG_Samara, I0211=Russia_HG_Karelia and UzOO77=Russia_EHG), Spain_ElMiron, Belgium_UP_GoyetQ116_1_published_all, Levant_N (Israel_PPNB), Russia_MA1_HG.SG, Israel_Natufian_published, Czech_Vestonice16 and Caucasus hunter-gatherers.

## Additional resources

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## Author contributions

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Investigation $=$ T.S. $($ aDNA data generation $)$, A.S. (quality control)
Project administration $=$ C.L.S.
Resources = C.S., N.C., L.M.

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Visualization $=$ S.A., T.S.

Writing - original draft = S.A., T.S., L.P.
Writing - review \& editing = All authors

## Declaration of interests

The authors declare no competing interests.

## Supplementary Data

Data S1. Sample information (Related to Table 1). (A) Summary archaeological, mapping, uniparental marker, genetic sex estimation, and damage statistics for samples sequenced in this study. (B) Results of radiocarbon dating of newly generated samples in this study. (C) Binary $Y$ chromosome haplogroup-informative variants found in derived state in at least one of the Iron Age Italian genomes. (D) Mitochondrial variants present in the newly generated sample in this study. Polymorphisms are estimated with haplogrep2. Here, the variant calling file (vcf) is uploaded on the online platform and the input file with the known polymorphism in chrRSRS is converted to $r C R S$. (E) Comparison of the mean PO values when running READ on different groupings of all individuals. (F) READ results for individuals when running all individuals with coverage over $0.03 x$ together. (G) READ results for individuals when running all Ordona individuals including SAL007 and SAL011 with coverage over $0.03 X$ together. (H) READ results for individuals when running all Salapia individuals including ORD004 with coverage over 0.03 X together.

Data S2. List and available information of published ancient samples used in this work.
Data S3. List and available information of published modern samples used in this work.
Data S4. $f 4$ investigating the genetic relationships among Iron Age Apulians (IAA) and modern Apulians. 14 Apulian samples from Raveane et al. 2019 (Modern_Apulia_GP) were used as Modern Apulians (Data S3).

Data S5. qpAdm results with 2, 3 and 4 possible sources for each test model (see STAR methods).
Data S6. $f 4$ investigating the genetic relationships among Iron Age Apulians (IAA) and the two Middle Age Apulian samples, the putative Daunians'population of origin (ancient populations coming from Crete, Peloponnese, Croatia and Iron Age Italians) and other ancient human groups. $f 4$ analyses were computed in the form f4(IAA, X; Y, Mbuti) where $X$ is a putative Daunians source and $Y$ is another ancient population.

## References

1. Hodos, T. (2020). The Archaeology of the Mediterranean Iron Age: A Globalising World c.1100-600 BCE (Cambridge University Press).
2. Abulafia, D. (2011). The Great Sea: A Human History of the Mediterranean (Oxford University Press).
3. Marcus, J.H., Posth, C., Ringbauer, H., Lai, L., Skeates, R., Sidore, C., Beckett, J., Furtwängler, A., Olivieri, A., Chiang, C.W.K., et al. (2020). Genetic history from the Middle Neolithic to present on the Mediterranean island of Sardinia. Nat. Commun. 11, 939.
4. Fernandes, D.M., Mittnik, A., Olalde, I., Lazaridis, I., Cheronet, O., Rohland, N., Mallick, S., Bernardos, R., Broomandkhoshbacht, N., Carlsson, J., et al. (2020). The spread of steppe and Iranian-related ancestry in the islands of the western Mediterranean. Nat Ecol Evol.
5. Nijboer, A.J. (2006). La cronologia assoluta dell'età del Ferro nel Mediterraneo, dibattito sui metodi e sui risultati. Oriente e Occidente: Metodi e Discipline a Confronto.
6. Antonio, M.L., Gao, Z., Moots, H.M., Lucci, M., Candilio, F., Sawyer, S., Oberreiter, V., Calderon, D., Devitofranceschi, K., Aikens, R.C., et al. (2019). Ancient Rome: A genetic crossroads of Europe and the Mediterranean. Science 366, 708-714.
7. Lombardo, M. (2014). Iapygians: The Indigenous Populations of Ancient Apulia in the Fifth and Fourth Centuries B.C.E. In The Italic People of Ancient Apulia: New Evidence from Pottery for Workshops, Markets, and Customs, T. Carpenter, K. Lynch, \& E. Robinson, ed. (Cambridge: Cambridge University Press), pp. 3668.
8. Norman, C. (2016). Daunian women: costume and actions commemorated in stone. In Women in antiquity. Real women across the Ancient World, Lynn Budin, Stephanie and Macintosh Turfa, Jean, ed., pp. 901-912.
9. Tagliente, M. (1986). I signori dei cavalli nella Daunia di età arcaica. Annali della Facoltà di Lettere e Filosofia dell'Università degli Studi di Perugia 23, 1985-1986.
10. Kirigin, B., Miše, M., and Barbarić, V. (2010). Palagruža-the Island of Diomedes: summary excavation report 2002-2008. Palagruža-the Island of Diomedes: summary excavation report 2002-2008, 65-91.
11. Saupe, T., Montinaro, F., Scaggion, C., Carrara, N., Kivisild, T., D’Atanasio, E., Hui, R., Solnik, A., Lebrasseur, O., Larson, G., et al. (2021). Ancient genomes reveal structural shifts after the arrival of Stepperelated ancestry in the Italian Peninsula. Curr. Biol. 31, 2576-2591.e12.
12. Allentoft, M.E., Sikora, M., Sjögren, K.-G., Rasmussen, S., Rasmussen, M., Stenderup, J., Damgaard, P.B., Schroeder, H., Ahlström, T., Vinner, L., et al. (2015). Population genomics of Bronze Age Eurasia. Nature 522, 167-172.
13. Haak, W., Lazaridis, I., Patterson, N., Rohland, N., Mallick, S., Llamas, B., Brandt, G., Nordenfelt, S., Harney, E., Stewardson, K., et al. (2015). Massive migration from the steppe was a source for Indo-European languages in Europe. Nature 522, 207-211.
14. Mathieson, I., Lazaridis, I., Rohland, N., Mallick, S., Patterson, N., Roodenberg, S.A., Harney, E., Stewardson, K., Fernandes, D., Novak, M., et al. (2015). Genome-wide patterns of selection in 230 ancient Eurasians. Nature 528, 499-503.
15. Schuenemann, V.J., Peltzer, A., Welte, B., van Pelt, W.P., Molak, M., Wang, C.-C., Furtwängler, A., Urban, C., Reiter, E., Nieselt, K., et al. (2017). Ancient Egyptian mummy genomes suggest an increase of SubSaharan African ancestry in post-Roman periods. Nat. Commun. 8, 15694.
16. Amorim, C.E.G., Vai, S., Posth, C., Modi, A., Koncz, I., Hakenbeck, S., La Rocca, M.C., Mende, B., Bobo, D., Pohl, W., et al. (2018). Understanding 6th-century barbarian social organization and migration through paleogenomics. Nat. Commun. 9, 3547.
17. Monroy Kuhn, J.M., Jakobsson, M., and Günther, T. (2018). Estimating genetic kin relationships in prehistoric populations. PLoS One 13, e0195491.
18. Schiffels, S., Haak, W., Paajanen, P., Llamas, B., Popescu, E., Loe, L., Clarke, R., Lyons, A., Mortimer, R., Sayer, D., et al. (2016). Iron Age and Anglo-Saxon genomes from East England reveal British migration history. Nat. Commun. 7, 10408.
19. Saag, L., Laneman, M., Varul, L., Malve, M., Valk, H., Razzak, M.A., Shirobokov, I.G., Khartanovich, V.I., Mikhaylova, E.R., Kushniarevich, A., et al. (2019). The Arrival of Siberian Ancestry Connecting the Eastern Baltic to Uralic Speakers further East. Curr. Biol. 29, 1701-1711.e16.
20. Krzewińska, M., Kılınç, G.M., Juras, A., Koptekin, D., Chyleński, M., Nikitin, A.G., Shcherbakov, N., Shuteleva, I., Leonova, T., Kraeva, L., et al. (2018). Ancient genomes suggest the eastern Pontic-Caspian steppe as the source of western Iron Age nomads. Sci Adv 4, eaat4457.
21. Olalde, I., Mallick, S., Patterson, N., Rohland, N., Villalba-Mouco, V., Silva, M., Dulias, K., Edwards, C.J., Gandini, F., Pala, M., et al. (2019). The genomic history of the Iberian Peninsula over the past 8000 years.

Science 363, 1230-1234.
22. Feldman, M., Master, D.M., Bianco, R.A., Burri, M., Stockhammer, P.W., Mittnik, A., Aja, A.J., Jeong, C., and Krause, J. (2019). Ancient DNA sheds light on the genetic origins of early Iron Age Philistines. Sci Adv 5, eaax0061.
23. Lazaridis, I., Patterson, N., Mittnik, A., Renaud, G., Mallick, S., Kirsanow, K., Sudmant, P.H., Schraiber, J.G., Castellano, S., Lipson, M., et al. (2014). Ancient human genomes suggest three ancestral populations for present-day Europeans. Nature 513, 409-413.
24. Molinaro, L., Montinaro, F., Yelmen, B., Marnetto, D., Behar, D.M., Kivisild, T., and Pagani, L. (2019). West Asian sources of the Eurasian component in Ethiopians: a reassessment. Sci. Rep. 9, 18811.
25. Lazaridis, I., Mittnik, A., Patterson, N., Mallick, S., Rohland, N., Pfrengle, S., Furtwängler, A., Peltzer, A., Posth, C., Vasilakis, A., et al. (2017). Genetic origins of the Minoans and Mycenaeans. Nature 548, 214-218.
26. Nava, M.L., and Descoeudres, J.P. (1990). Greek and Adriatic influences in Daunia in the Early Iron Age. Greek Colonists and Native Populations, 560-578.
27. Jones, C.P. (1987). Stigma: Tattooing and Branding in Graeco-Roman Antiquity. Journal of Roman Studies 77, 139-155.
28. Norman, C. (2011). The Tribal Tattooing of Daunian Women. European Journal of Archaeology 14, 133-157.
29. De Juliis, E.M. (1988). Gli Iapigi: storia e civiltà della Puglia preromana.
30. Reimer, P.J., Austin, W.E.N., Bard, E., Bayliss, A., Blackwell, P.G., Ramsey, C.B., Butzin, M., Cheng, H., Lawrence Edwards, R., Friedrich, M., et al. (2020). The IntCal20 Northern Hemisphere Radiocarbon Age Calibration Curve ( $0-55$ cal kBP). Radiocarbon 62, 725-757.
31. Corrain, C. (1986). Resti scheletrici umani da Ordona (Foggia), Secc. VII--IV aC In Iker R (ed) Ordona VII/2. Les Tombes Dauniennes. Etudes de Philologie, d'Archéologie et d'Histoire Ancienne Publiées par l'Institut Historique Belge de Rome 24, 787-807.
32. Scaggion, C., and Carrara, N. (2016). New studies on human skeletal remains from the ancient Herdonia (southeast Italy). Evidences of tuberculosis and brucellosis: two diseases connected with farm animals. Antrocom: Online Journal of Anthropology 12.
33. Corrain, C., Capitanio, M., and Erspamer, G. (1972). I resti scheletrici della necropoli di Salapia (Cerignola), secoli IX-III aC (Società Cooperativa Tipografica).
34. Meyer, M., and Kircher, M. (2010). Illumina Sequencing Library Preparation for Highly Multiplexed Target Capture and Sequencing. Cold Spring Harbor Protocols 2010, db.prot5448-pdb.prot5448.
35. Orlando, L., Ginolhac, A., Zhang, G., Froese, D., Albrechtsen, A., Stiller, M., Schubert, M., Cappellini, E., Petersen, B., Moltke, I., et al. (2013). Recalibrating Equus evolution using the genome sequence of an early Middle Pleistocene horse. Nature 499, 74-78.
36. Malaspinas, A.-S., Lao, O., Schroeder, H., Rasmussen, M., Raghavan, M., Moltke, I., Campos, P.F., Sagredo, F.S., Rasmussen, S., Gonçalves, V.F., et al. (2014). Two ancient human genomes reveal Polynesian ancestry among the indigenous Botocudos of Brazil. Curr. Biol. 24, R1035-7.
37. Martin, M. (2011). Cutadapt removes adapter sequences from high-throughput sequencing reads. EMBnet.journal 17, 10-12.
38. Li, H., and Durbin, R. (2009). Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 25, 1754-1760.
39. Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., Marth, G., Abecasis, G., and Durbin, R. (2009). 1000 Genome Project Data Processing Subgroup. 2009. The sequence alignment/map format and samtools. Bioinformatics 25, 2078-2079.
40. McKenna, A., Hanna, M., Banks, E., Sivachenko, A., Cibulskis, K., Kernytsky, A., Garimella, K., Altshuler, D., Gabriel, S., Daly, M., et al. (2010). The Genome Analysis Toolkit: a MapReduce framework for analyzing next-generation DNA sequencing data. Genome Res. 20, 1297-1303.
41. Jónsson, H., Ginolhac, A., Schubert, M., Johnson, P.L.F., and Orlando, L. (2013). mapDamage2.0: fast approximate Bayesian estimates of ancient DNA damage parameters. Bioinformatics 29, 1682-1684.
42. Jones, E.R., Zarina, G., Moiseyev, V., Lightfoot, E., Nigst, P.R., Manica, A., Pinhasi, R., and Bradley, D.G. (2017). The Neolithic Transition in the Baltic Was Not Driven by Admixture with Early European Farmers. Curr. Biol. 27, 576-582.
43. Skoglund, P., Storå, J., Götherström, A., and Jakobsson, M. (2013). Accurate sex identification of ancient human remains using DNA shotgun sequencing. J. Archaeol. Sci. 40, 4477-4482.
44. Fu, Q., Posth, C., Hajdinjak, M., Petr, M., Mallick, S., Fernandes, D., Furtwängler, A., Haak, W., Meyer, M., Mittnik, A., et al. (2016). The genetic history of Ice Age Europe. Nature 534, 200-205.
45. Weissensteiner, H., Pacher, D., Kloss-Brandstätter, A., Forer, L., Specht, G., Bandelt, H.-J., Kronenberg, F.,

708

Salas, A., and Schönherr, S. (2016). HaploGrep 2: mitochondrial haplogroup classification in the era of highthroughput sequencing. Nucleic Acids Res. 44, W58-63.
46. van Oven, M., and Kayser, M. (2009). Updated comprehensive phylogenetic tree of global human mitochondrial DNA variation. Hum. Mutat. 30, E386-94.
47. Karmin, M., Saag, L., Vicente, M., Wilson Sayres, M.A., Järve, M., Talas, U.G., Rootsi, S., Ilumäe, A.-M., Mägi, R., Mitt, M., et al. (2015). A recent bottleneck of Y chromosome diversity coincides with a global change in culture. Genome Res. 25, 459-466.
48. Poznik, G.D., Xue, Y., Mendez, F.L., Willems, T.F., Massaia, A., Wilson Sayres, M.A., Ayub, Q., McCarthy, S.A., Narechania, A., Kashin, S., et al. (2016). Punctuated bursts in human male demography inferred from 1,244 worldwide Y-chromosome sequences. Nat. Genet. 48, 593-599.
49. Hallast, P., Batini, C., Zadik, D., Maisano Delser, P., Wetton, J.H., Arroyo-Pardo, E., Cavalleri, G.L., de Knijff, P., Destro Bisol, G., Dupuy, B.M., et al. (2015). The Y-chromosome tree bursts into leaf: 13,000 highconfidence SNPs covering the majority of known clades. Mol. Biol. Evol. 32, 661-673.
50. Korneliussen, T.S., Albrechtsen, A., and Nielsen, R. (2014). ANGSD: Analysis of Next Generation Sequencing Data. BMC Bioinformatics 15, 356 .
51. 1000 Genomes Project Consortium, Auton, A., Brooks, L.D., Durbin, R.M., Garrison, E.P., Kang, H.M., Korbel, J.O., Marchini, J.L., McCarthy, S., McVean, G.A., et al. (2015). A global reference for human genetic variation. Nature 526, 68-74.
52. Patterson, N., Price, A.L., and Reich, D. (2006). Population structure and eigenanalysis. PLoS Genet. 2, e190.
53. Chang, C.C., Chow, C.C., Tellier, L.C., Vattikuti, S., Purcell, S.M., and Lee, J.J. (2015). Second-generation PLINK: rising to the challenge of larger and richer datasets. GigaScience 4.
54. Patterson, N., Moorjani, P., Luo, Y., Mallick, S., Rohland, N., Zhan, Y., Genschoreck, T., Webster, T., and Reich, D. (2012). Ancient admixture in human history. Genetics 192, 1065-1093.
55. Bortolini, E., Pagani, L., Oxilia, G., Posth, C., Fontana, F., Badino, F., Saupe, T., Montinaro, F., Margaritora, D., Romandini, M., et al. (2021). Early Alpine occupation backdates westward human migration in Late Glacial Europe. Curr. Biol.
56. Raveane, A., Aneli, S., Montinaro, F., Athanasiadis, G., Barlera, S., Birolo, G., Boncoraglio, G., Di Blasio, A.M., Di Gaetano, C., Pagani, L., et al. (2019). Population structure of modern-day Italians reveals patterns of ancient and archaic ancestries in Southern Europe. Science Advances 5, eaaw3492.
57. Brace, S., Diekmann, Y., Booth, T.J., van Dorp, L., Faltyskova, Z., Rohland, N., Mallick, S., Olalde, I., Ferry, M., Michel, M., et al. (2019). Author Correction: Ancient genomes indicate population replacement in Early Neolithic Britain. Nat Ecol Evol 3, 986-987.
58. Broushaki, F., Thomas, M.G., Link, V., López, S., van Dorp, L., Kirsanow, K., Hofmanová, Z., Diekmann, Y., Cassidy, L.M., Díez-Del-Molino, D., et al. (2016). Early Neolithic genomes from the eastern Fertile Crescent. Science 353, 499-503.
59. Cassidy, L.M., Martiniano, R., Murphy, E.M., Teasdale, M.D., Mallory, J., Hartwell, B., and Bradley, D.G. (2016). Neolithic and Bronze Age migration to Ireland and establishment of the insular Atlantic genome. Proc. Natl. Acad. Sci. U. S. A. 113, 368-373.
60. Damgaard, P. de B., Marchi, N., Rasmussen, S., Peyrot, M., Renaud, G., Korneliussen, T., Moreno-Mayar, J.V., Pedersen, M.W., Goldberg, A., Usmanova, E., et al. (2018). 137 ancient human genomes from across the Eurasian steppes. Nature 557, 369-374.
61. Damgaard, P. de B., de Barros Damgaard, P., Martiniano, R., Kamm, J., Víctor Moreno-Mayar, J., Kroonen, G., Peyrot, M., Barjamovic, G., Rasmussen, S., Zacho, C., et al. (2018). The first horse herders and the impact of early Bronze Age steppe expansions into Asia. Science 360, eaar7711.
62. Gamba, C., Jones, E.R., Teasdale, M.D., McLaughlin, R.L., Gonzalez-Fortes, G., Mattiangeli, V., Domboróczki, L., Kővári, I., Pap, I., Anders, A., et al. (2014). Genome flux and stasis in a five millennium transect of European prehistory. Nature Communications 5.
63. González-Fortes, G., Jones, E.R., Lightfoot, E., Bonsall, C., Lazar, C., Grandal-d'Anglade, A., Garralda, M.D., Drak, L., Siska, V., Simalcsik, A., et al. (2017). Paleogenomic Evidence for Multi-generational Mixing between Neolithic Farmers and Mesolithic Hunter-Gatherers in the Lower Danube Basin. Current Biology 27, 1801-1810.e10.
64. González-Fortes, G., Tassi, F., Trucchi, E., Henneberger, K., Paijmans, J.L.A., Díez-del-Molino, D., Schroeder, H., Susca, R.R., Barroso-Ruíz, C., Bermudez, F.J., et al. (2019). A western route of prehistoric human migration from Africa into the Iberian Peninsula. Proceedings of the Royal Society B: Biological Sciences 286, 20182288.
65. Günther, T., Valdiosera, C., Malmström, H., Ureña, I., Rodriguez-Varela, R., Sverrisdóttir, Ó.O., Daskalaki,
E.A., Skoglund, P., Naidoo, T., Svensson, E.M., et al. (2015). Ancient genomes link early farmers from Atapuerca in Spain to modern-day Basques. Proc. Natl. Acad. Sci. U. S. A. 112, 11917-11922.
66. Haber, M., Doumet-Serhal, C., Scheib, C., Xue, Y., Danecek, P., Mezzavilla, M., Youhanna, S., Martiniano, R., Prado-Martinez, J., Szpak, M., et al. (2017). Continuity and Admixture in the Last Five Millennia of Levantine History from Ancient Canaanite and Present-Day Lebanese Genome Sequences. Am. J. Hum. Genet. 101, 274-282.
67. Harney, É., May, H., Shalem, D., Rohland, N., Mallick, S., Lazaridis, I., Sarig, R., Stewardson, K., Nordenfelt, S., Patterson, N., et al. (2018). Publisher Correction: Ancient DNA from Chalcolithic Israel reveals the role of population mixture in cultural transformation. Nat. Commun. 9, 3913.
68. Hofmanová, Z., Kreutzer, S., Hellenthal, G., Sell, C., Diekmann, Y., Díez-Del-Molino, D., van Dorp, L., López, S., Kousathanas, A., Link, V., et al. (2016). Early farmers from across Europe directly descended from Neolithic Aegeans. Proc. Natl. Acad. Sci. U. S. A. 113, 6886-6891.
69. Järve, M., Saag, L., Scheib, C.L., Pathak, A.K., Montinaro, F., Pagani, L., Flores, R., Guellil, M., Saag, L., Tambets, K., et al. (2019). Shifts in the Genetic Landscape of the Western Eurasian Steppe Associated with the Beginning and End of the Scythian Dominance. Current Biology 29, 2430-2441.e10.
70. Jones, E.R., Gonzalez-Fortes, G., Connell, S., Siska, V., Eriksson, A., Martiniano, R., McLaughlin, R.L., Gallego Llorente, M., Cassidy, L.M., Gamba, C., et al. (2015). Upper Palaeolithic genomes reveal deep roots of modern Eurasians. Nat. Commun. 6, 8912.
71. Kılınç, G.M., Omrak, A., Özer, F., Günther, T., Büyükkarakaya, A.M., Bıçakçı, E., Baird, D., Dönertaş, H.M., Ghalichi, A., Yaka, R., et al. (2016). The Demographic Development of the First Farmers in Anatolia. Curr. Biol. 26, 2659-2666.
72. Lazaridis, I., Nadel, D., Rollefson, G., Merrett, D.C., Rohland, N., Mallick, S., Fernandes, D., Novak, M., Gamarra, B., Sirak, K., et al. (2016). Genomic insights into the origin of farming in the ancient Near East. Nature 536, 419-424.
73. Lipson, M., Szécsényi-Nagy, A., Mallick, S., Pósa, A., Stégmár, B., Keerl, V., Rohland, N., Stewardson, K., Ferry, M., Michel, M., et al. (2017). Parallel palaeogenomic transects reveal complex genetic history of early European farmers. Nature 551, 368-372.
74. Martiniano, R., Caffell, A., Holst, M., Hunter-Mann, K., Montgomery, J., Müldner, G., McLaughlin, R.L., Teasdale, M.D., van Rheenen, W., Veldink, J.H., et al. (2016). Genomic signals of migration and continuity in Britain before the Anglo-Saxons. Nat. Commun. 7, 10326.
75. Martiniano, R., Cassidy, L.M., Ó’Maoldúin, R., McLaughlin, R., Silva, N.M., Manco, L., Fidalgo, D., Pereira, T., Coelho, M.J., Serra, M., et al. (2017). The population genomics of archaeological transition in west Iberia: Investigation of ancient substructure using imputation and haplotype-based methods. PLoS Genet. 13, e1006852.
76. Mathieson, I., Alpaslan-Roodenberg, S., Posth, C., Szécsényi-Nagy, A., Rohland, N., Mallick, S., Olalde, I., Broomandkhoshbacht, N., Candilio, F., Cheronet, O., et al. (2018). The genomic history of southeastern Europe. Nature 555, 197-203.
77. Mittnik, A., Wang, C.-C., Pfrengle, S., Daubaras, M., Zariņa, G., Hallgren, F., Allmäe, R., Khartanovich, V., Moiseyev, V., Tõrv, M., et al. (2018). The genetic prehistory of the Baltic Sea region. Nat. Commun. 9, 442.
78. Narasimhan, V.M., Patterson, N., Moorjani, P., Rohland, N., Bernardos, R., Mallick, S., Lazaridis, I., Nakatsuka, N., Olalde, I., Lipson, M., et al. (2019). The formation of human populations in South and Central Asia. Science 365.
79. Olalde, I., Schroeder, H., Sandoval-Velasco, M., Vinner, L., Lobón, I., Ramirez, O., Civit, S., García Borja, P., Salazar-García, D.C., Talamo, S., et al. (2015). A Common Genetic Origin for Early Farmers from Mediterranean Cardial and Central European LBK Cultures. Mol. Biol. Evol. 32, 3132-3142.
80. Olalde, I., Brace, S., Allentoft, M.E., Armit, I., Kristiansen, K., Booth, T., Rohland, N., Mallick, S., SzécsényiNagy, A., Mittnik, A., et al. (2018). The Beaker phenomenon and the genomic transformation of northwest Europe. Nature 555, 190-196.
81. Omrak, A., Günther, T., Valdiosera, C., Svensson, E.M., Malmström, H., Kiesewetter, H., Aylward, W., Storå, J., Jakobsson, M., and Götherström, A. (2016). Genomic Evidence Establishes Anatolia as the Source of the European Neolithic Gene Pool. Curr. Biol. 26, 270-275.
82. Saag, L., Varul, L., Scheib, C.L., Stenderup, J., Allentoft, M.E., Saag, L., Pagani, L., Reidla, M., Tambets, K., Metspalu, E., et al. (2017). Extensive Farming in Estonia Started through a Sex-Biased Migration from the Steppe. Curr. Biol. 27, 2185-2193.e6.
83. Sikora, M., Pitulko, V.V., Sousa, V.C., Allentoft, M.E., Vinner, L., Rasmussen, S., Margaryan, A., de Barros Damgaard, P., de la Fuente, C., Renaud, G., et al. (2019). The population history of northeastern Siberia since
the Pleistocene. Nature 570, 182-188.
84. Unterländer, M., Palstra, F., Lazaridis, I., Pilipenko, A., Hofmanová, Z., Groß, M., Sell, C., Blöcher, J., Kirsanow, K., Rohland, N., et al. (2017). Ancestry and demography and descendants of Iron Age nomads of the Eurasian Steppe. Nat. Commun. 8, 14615.
85. Villalba-Mouco, V., van de Loosdrecht, M.S., Posth, C., Mora, R., Martínez-Moreno, J., Rojo-Guerra, M., Salazar-García, D.C., Royo-Guillén, J.I., Kunst, M., Rougier, H., et al. (2019). Survival of Late Pleistocene Hunter-Gatherer Ancestry in the Iberian Peninsula. Curr. Biol. 29, 1169-1177.e7.
86. Wang, C.-C., Reinhold, S., Kalmykov, A., Wissgott, A., Brandt, G., Jeong, C., Cheronet, O., Ferry, M., Harney, E., Keating, D., et al. (2019). Ancient human genome-wide data from a 3000 -year interval in the Caucasus corresponds with eco-geographic regions. Nat. Commun. 10, 590.
87. Agranat-Tamir, L., Waldman, S., Martin, M.A.S., Gokhman, D., Mishol, N., Eshel, T., Cheronet, O., Rohland, N., Mallick, S., Adamski, N., et al. (2020). The Genomic History of the Bronze Age Southern Levant. Cell 181, 1146-1157.e11.
88. Brunel, S., Bennett, E.A., Cardin, L., Garraud, D., Barrand Emam, H., Beylier, A., Boulestin, B., Chenal, F., Ciesielski, E., Convertini, F., et al. (2020). Ancient genomes from present-day France unveil 7,000 years of its demographic history. Proc. Natl. Acad. Sci. U. S. A. 117, 12791-12798.
89. Fregel, R., Méndez, F.L., Bokbot, Y., Martín-Socas, D., Camalich-Massieu, M.D., Santana, J., Morales, J., Ávila-Arcos, M.C., Underhill, P.A., Shapiro, B., et al. (2018). Ancient genomes from North Africa evidence prehistoric migrations to the Maghreb from both the Levant and Europe. Proc. Natl. Acad. Sci. U. S. A. 115, 6774-6779.
90. Furtwängler, A., Rohrlach, A.B., Lamnidis, T.C., Papac, L., Neumann, G.U., Siebke, I., Reiter, E., Steuri, N., Hald, J., Denaire, A., et al. (2020). Ancient genomes reveal social and genetic structure of Late Neolithic Switzerland. Nat. Commun. 11, 1915.
91. Gokhman, D., Nissim-Rafinia, M., Agranat-Tamir, L., Housman, G., García-Pérez, R., Lizano, E., Cheronet, O., Mallick, S., Nieves-Colón, M.A., Li, H., et al. (2020). Differential DNA methylation of vocal and facial anatomy genes in modern humans. Nat. Commun. 11, 1189.
92. Haber, M., Doumet-Serhal, C., Scheib, C.L., Xue, Y., Mikulski, R., Martiniano, R., Fischer-Genz, B., Schutkowski, H., Kivisild, T., and Tyler-Smith, C. (2019). A Transient Pulse of Genetic Admixture from the Crusaders in the Near East Identified from Ancient Genome Sequences. The American Journal of Human Genetics 104, 977-984.
93. Haber, M., Nassar, J., Almarri, M.A., Saupe, T., Saag, L., Griffith, S.J., Doumet-Serhal, C., Chanteau, J., Saghieh-Beydoun, M., Xue, Y., et al. (2020). A Genetic History of the Near East from an aDNA Time Course Sampling Eight Points in the Past 4,000 Years. Am. J. Hum. Genet. 107, 149-157.
94. Keller, A., Graefen, A., Ball, M., Matzas, M., Boisguerin, V., Maixner, F., Leidinger, P., Backes, C., Khairat, R., Forster, M., et al. (2012). New insights into the Tyrolean Iceman's origin and phenotype as inferred by whole-genome sequencing. Nat. Commun. 3, 698.
95. Margaryan, A., Lawson, D.J., Sikora, M., Racimo, F., Rasmussen, S., Moltke, I., Cassidy, L.M., Jørsboe, E., Ingason, A., Pedersen, M.W., et al. (2020). Population genomics of the Viking world. Nature 585, 390-396.
96. Rivollat, M., Jeong, C., Schiffels, S., Küçükkalıpçı, İ., Pemonge, M.-H., Rohrlach, A.B., Alt, K.W., Binder, D., Friederich, S., Ghesquière, E., et al. (2020). Ancient genome-wide DNA from France highlights the complexity of interactions between Mesolithic hunter-gatherers and Neolithic farmers. Sci Adv 6, eaaz5344.
97. Skourtanioti, E., Erdal, Y.S., Frangipane, M., Balossi Restelli, F., Yener, K.A., Pinnock, F., Matthiae, P., Özbal, R., Schoop, U.-D., Guliyev, F., et al. (2020). Genomic History of Neolithic to Bronze Age Anatolia, Northern Levant, and Southern Caucasus. Cell 181, 1158-1175.e28.
98. Valdiosera, C., Günther, T., Vera-Rodríguez, J.C., Ureña, I., Iriarte, E., Rodríguez-Varela, R., Simões, L.G., Martínez-Sánchez, R.M., Svensson, E.M., Malmström, H., et al. (2018). Four millennia of Iberian biomolecular prehistory illustrate the impact of prehistoric migrations at the far end of Eurasia. Proceedings of the National Academy of Sciences 115, 3428-3433.
99. van den Brink, E.C.M., Beeri, R., Kirzner, D., Bron, E., Cohen-Weinberger, A., Kamaisky, E., Gonen, T., Gershuny, L., Nagar, Y., Ben-Tor, D., et al. (2017). A Late Bronze Age II clay coffin from Tel Shaddud in the Central Jezreel Valley, Israel: context and historical implications. Levant 49, 105-135.
100. Veeramah, K.R., Rott, A., Groß, M., van Dorp, L., López, S., Kirsanow, K., Sell, C., Blöcher, J., Wegmann, D., Link, V., et al. (2018). Population genomic analysis of elongated skulls reveals extensive female-biased immigration in Early Medieval Bavaria. Proc. Natl. Acad. Sci. U. S. A. 115, 3494-3499.
101. Zalloua, P., Collins, C.J., Gosling, A., Biagini, S.A., Costa, B., Kardailsky, O., Nigro, L., Khalil, W., Calafell, F., and Matisoo-Smith, E. (2018). Ancient DNA of Phoenician remains indicates discontinuity in the
settlement history of Ibiza. Sci. Rep. 8, 17567.
102. Feldman, M., Fernández-Domínguez, E., Reynolds, L., Baird, D., Pearson, J., Hershkovitz, I., May, H., Goring-Morris, N., Benz, M., Gresky, J., et al. (2019). Late Pleistocene human genome suggests a local origin for the first farmers of central Anatolia.
103. Jeong, C., Balanovsky, O., Lukianova, E., Kahbatkyzy, N., Flegontov, P., Zaporozhchenko, V., Immel, A., Wang, C.-C., Ixan, O., Khussainova, E., et al. (2019). The genetic history of admixture across inner Eurasia. Nat Ecol Evol 3, 966-976.
104. Biagini, S.A., Solé-Morata, N., Matisoo-Smith, E., Zalloua, P., Comas, D., and Calafell, F. (2019). People from Ibiza: an unexpected isolate in the Western Mediterranean. Eur. J. Hum. Genet. 27, 941-951.
105. Alexander, D.H., Novembre, J., and Lange, K. (2009). Fast model-based estimation of ancestry in unrelated individuals. Genome Res. 19, 1655-1664.
106. Core Team, R., and Others (2013). R: a language and environment for statistical computing. R Foundation for statistical computing, Vienna.
107. R Core Team (2021). R: A language and environment for statistical computing.
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Figure 1: Geographical location, dating and genetic structure. (A) Geographical location of the newly generated Apulian ancient individuals (green, red and blue diamonds and triangles) and other published Iron Age samples from Italy and surrounding areas (color scale mirroring their "years before present" dates). See also Table 1, Data S1, and Data S2. (B) Dating and number of SNPs from 1240K set covered. Samples with thick edges have been radiocarbon dated (mean presented), while for the others we represented their archaeological dating (Table 1, Data S1B). (C) Principal component analysis (PCA) of newly generated individuals with reference ancient individuals projected onto the genetic variation inferred from modern-day Western Eurasian populations (Data S2 and Data S3). For the ancient reference populations which have not been attributed to the Iron Age, we plotted their centroids.


Figure 2: Genetic relationship of Iron Age and Middle Age Apulians and their ancestral composition. (A) Heatmap representing the Z -score values of f4(Modern Apulia, IAA; X , Mbuti) where X is an ancient Italian population. We also added the two Middle Age samples for comparison. Tests with Z-scores between -3 and 3 or with less than 5,000 SNPs were not included (P: Palaeolithic, M: Mesolithic, N: Neolithic, C: Chalcolithic, CA: Copper Age, BA: Bronze Age, IA: Iron Age, A: Antiquity). (B) qpAdm proportions of ancient Apulian samples and other reference populations (Modern Apulians, Minoans, Croatia_EIA and the Roman individuals from the Republican and the Imperial period) using "base" sources: Caucasus and Western hunter-gatherer component (HG), Anatolia Neolithic (Anatolia_N), Iranian Neolithic (Iran_N), Stepperelated ancestry. (C) qpAdm proportions using also the Minoans as source (STAR methods). In B and C, we plotted the model with the highest number of sources and p-value (complete results may be seen in Figure S5).


Figure 3: Genetic affinities of Iron Age Apulian samples with the putative populations of origin: Minoans (Minoan Lassithi), Illyrians (here proxied by the Croatia_EIA individual) and the Roman Republicans (here proxying the autochthonous Iron Age Italian ancestry). (A) Outgroup f3 statistics of IAA samples compared with the putative Daunian sources. Samples have been sorted and grouped according to the source whose f 3 values were higher. (B) Heatmap showing the Z -score values of f4(putative sources, IAA_P; X, Mbuti) where X is an ancient Italian population and IAA_P is the entire set of Iron Age Apulian samples taken together. Tests with Z-scores between -3 and 3 or with less than 5,000 SNPs were not included (P: Palaeolithic, M: Mesolithic, N: Neolithic, C: Chalcolithic, CA: Copper Age, BA: Bronze Age, IA: Iron Age, A: Antiquity).
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## Supplementary Materials

## This PDF file includes:

- Supplementary Figures 1 to 7 .


Figure S1. Graphical output of READ. A) All newly generated individuals run together regardless of geography or chronology. B) All newly generated individuals from the archaeological site Ordona plus SAL007 and SAL011. C) All newly generated individuals from the archaeological site Salapia plus ORD004.


Figure S2. Correlation of radiocarbon dating and principal components for the ancient Apulian samples. A) Pearson correlation of dates with the first three principal components for the radiocarbon dated ancient Apulian samples. B) Same as A, after removing the Middle Age samples ORD010 and SGR001. C), D) and E) show three principal components analyses based on PC1 vs PC2, PC2 vs PC3 and PC1 vs PC3, respectively.


Figure S3. Outgroup $f 3$ analyses in the form $f 3$ (Ancient Apulian samples, X ; Mbuti). A) Outgroup $f 3$ analyses comparing the newly generated ancient Apulian samples with other ancient human groups (see STAR Methods and Data S1). B) Outgroup $f 3$ analyses comparing the newly generated ancient Apulian samples with present-day populations (see STAR Methods and Data S2). The number of the SNPs used in each run is reported near each $f 3$ value. The ancient Apulian samples are plotted according to their age (SAL001 is the oldest) and Middle Age samples (ORD010 and SGR001) are reported in blue.


Figure S4. outgroup-f3 multidimensional scaling (MDS) of ancient Apulian samples and published ancient and modern reference populations. A pairwise matrix of outgroup $f 3$ (Ancient Apulian samples, X; Mbuti) distance was used to compute the MDS (STAR Methods) after removing samples belonging to Palaelolithic and Mesolithic periods. Ancient and modern groups were aggregated according to their published label (Data S1, Data S2 and STAR Methods).


Figure S5. Ancestral composition of ancient Apulian samples inferred by qpAdm. A) qpAdm proportions of ancient Apulian samples and other reference populations (Modern Apulians, Minoans, Croatia_EIA and the Roman individuals from the Republican and the Imperial period) using base sources: Caucasus and Western hunter-gatherer component (HG), Anatolia Neolithic (Anatolia_N), Iranian Neolithic (Iran_N), Stepperelated ancestry. B), C) and D) Same as A, using also the Minoans, Amhara_NAF and Rome Republicans as source (STAR methods). Only the models with p-values higher than 0.05 are shown.

A $\quad f 4(\mathrm{X}, \mathrm{IAA} ;$ WHG, Mbuti)















C


Figure S6. Genetic relationships among Iroan Age Apulians (IAA), their putative sources (ancient populations coming from Crete, Peloponnese, Croatia and Iron Age Italians) and Caucasus, Western hunter-gatherer (CHG and WHG, respectively) and Iranian Neolithic (IN) components. A) $f 4$ (X, IAA; WHG, Mbuti) values, where X is a putative source of the Daunian population. B) and C) Same as A, using CHG and IN, respectively. Purple points represent $f 4$ values with significant Z-scores ( $|Z| \geq 3$ ), pink points represent almost significant $f 4$ values $(2 \leq|Z|<3$ ), while dashed lines indicate $f 4$ runs involving less than 5,000 SNPs. Also the Middle Age Apulian samples (ORD010 and SGR001) were added for comparison.


Figure S7. Model-based clustering on modern and ancient individuals as inferred by ADMIXTURE. A) The ancestral components were inferred in a set of 90 modern Eurasian populations. B) Using the allele frequency for ancestral components inferred in A. we projected a subset of ancient Europasian individuals.


Figure S8. $f 4$ analyses investigating the genetic heterogeneity of IAA samples. A) Almost significant $f 4$ values involving a putative first-degree relationship between ORD009 and ORD001 (possibly, mother and daughter). B) $f 4$ analyses involving Rome Republicans, IAA and Levant_N/YRI. C) and D) show the $f 4$ analyses involving IAA, Levant_N and, alternatively, CHG and IN. In all plots, purple points represent $f 4$ values with significant Z -scores $(|Z| \geq 3$ ), pink points represent almost significant $f 4$ values $(2 \leq|Z|<3)$, while dashed lines indicate $f 4$ runs involving less than 5,000 SNPs. Also the Middle Age Apulian samples (ORD010 and SGR001) were added for comparison.


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