The ancestral origin of the critically endangered Quadricorna sheep as revealed by genome-wide analysis

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Abstract

Livestock European diffusion followed different human migration waves from the Fertile Crescent. In sheep, at least two diffusion waves have shaped the current breeds’ biodiversity generating a complex genetic pattern composed by either primitive or fine-wool selected breeds. Among primitive breeds, aside from sharing common ancestral genomic components, they also show several traits such as the policeraty, large horns in the ram, short tail, and a molting fleece, considered as ancestral. Although most of the primitive breeds characterized by these traits are confined on the very edge of Northern Europe, several residual populations are also scattered in the Mediterranean region. In fact, although in Italy a large number of local breeds are already extinct, others are listed as critically endangered, and among these there is the Quadricorna breed which is a four-horned sheep characterized by several ancestral traits. In this context we genotyped 47 individuals belonging to the Quadricorna sheep breed, a relict and endangered breed, from Central and Southern Italy. In doing so we used the Illumina OvineSNP50K array in order to explore its genetic diversity and to compare it with other 33 primitive traits-related, Mediterranean and Middle-East breeds, with the specific aim to reconstruct its origin. After retaining 35,680 SNPs following data filtering, the overall genomic architecture has been explored by using genetic diversity indices, Principal Component Analysis (PCA) and admixture analysis, while the genetic relationships and migration events have been inferred using a neighbor-joining tree based on Reynolds’ distances and by the maximum likelihood tree as implemented in treemix. Multiple convergent evidence from all our population genetics analyses, indicated that the two Quadricorna populations differ from all the other Italian breeds, while they resulted to be very close to the Middle Eastern and primitive European breeds. In addition, the genetic diversity indices highlighted values comparable with those of most of the other analyzed breeds, despite the two populations exhibit slightly different genetic indices suggesting different levels of genomic inbreeding and drift (FIS and FROH). The admixture analysis does not suggest any signal of recent gene exchange with other Italian local breeds, highlighting a rather ancestral
purity of the two populations, while on the other hand the treemix analysis seems to suggest an ancient admixture with other primitive European breeds. Finally, all these evidences seem to trace back the residual Quadricorna sheep to an early Neolithic spread, probably following a Mediterranean route and that urgent conservation actions are needed in order to keep the breed and all related cultural products alive.

**Introduction**

Since Neolithic revolution, domestic animals followed human migration routes and spread throughout the world. For many livestock species, this processes have been shown to have occurred in distinct historical periods by following different colonization routes leading to a vast array of breed diversity well adapted to specific local environments [1–3], that is to say with a high "capacity for constructivism", understood as the tendency of organisms to actively participate in the construction of a specific "bioterritory" rather than simply "adapting", establishing a vital relationship with it to achieve maximum fitness [4–7].

Compared to other domestic species such as cattle, goat and dogs, sheep (Ovis aries) have been shown to have retained much higher genetic diversity conform with wider heterogeneous pre-domestication populations, combined with less severe post-domestication bottlenecks [8, 9]. Moreover, after sheep domestication, at least two principal waves from the Fertile Crescent prompted the diffusion of an early hair-type (~10,500 YBP) followed by a later wool-type diffusion into Europe [10, 11]. An additional migration wave involving fat-tailed and fat-rumped sheep probably occurred about 3,500 YBP determining their expansion across North Africa, Middle-east and Asia [12–14].

In Europe, most of the genome of the first migration wave (hair type sheep), has been later replaced, generating a well-defined geographic cline from Middle-East to Balkan and continental Europe [15]. However, several relict primitive populations are still reared in scattered areas of Northern Europe [10] while different insular mouflon populations have also retained ancestral genetic diversity. These peripheral breeds apart from sharing a similar genotype and retrotype, are also recognized as primitive by possessing several morphological traits considered ancestral such as large horns in the ram, a short tail, and a moulting fleece [16]. Moreover, the common presence of four-horned phenotypes has also been argued to be associated with primitive sheep breeds being present in short-tailed prehistoric sheep in Northwest-Europe (Iceland and British Isles) and in primitive breeds of Southern Europe (Cyprus and Spain), while the four-horned Jacob sheep is believed to be possibly linked to Middle-East [10, 17–19].

Concerning the Italian peninsula, a high level of genetic diversity with a strong phylogeographic structuring has been observed [20]. All genetic studies to date, identified a complex background compatible with a late migration triggered by wool production [21, 22]. For example, the presence of fine wool sheep was already documented in the Greek and Roman times and it is believed that starting from this period, they spread throughout Europe and crossed with other native breeds giving rise to the Spanish Merino breeds [12, 20]. Nowadays, Italy is home to more than 60 breeds, many of them reduced to small local populations and listed as critically endangered [23]. Instead, autochthonous breeds and their relative products, represent an important resource not only for their cultural and economical heritage but also as a source of genetic diversity important to face the new millennium challenges of livestock. So far, local breeds are experiencing an intense crisis due to their replacement with exotic and
more productive breeds and as a consequence, many native livestock populations have already become extinct [24]. In particular, it is estimated that as regards the Italian sheep populations, as many as 48% fall into the damaged risk category and 30% into the critical one [25]. In such a context, there is a growing urgency to perform genomic assessments of local breeds in order to explore their level of genetic diversity and eventually to improve adequate conservation and breeding strategies. Moreover, local breeds being well adapted to marginal areas characterized by poor quality feed and low input conditions, they potentially function as a reservoir of genes able to confer high "capacity of constructivism", important to counteract all related deleterious effects caused by climate and environmental changes. Indeed, many recent studies focusing on local breeds have highlighted a number of potential genes involved in disease resistance, tolerance to harsh conditions and adaptation to low input and sustainable systems [26].

Among the most critically endangered Italian local breed is the Quadricorna sheep which is constituted by a very residual population reared in Central-Southern Italy, counting in 2022 only 2 farms, one located in Colliano (SA, Campania) and the other in Monte San Giovanni Campano (FR, Lazio) with 33 (20 females, 13 males) and 23 (21 females, 2 males) heads respectively. The sheep population called "Pecora Quadricorna" has been registered since 2006 in the Regional Voluntary Register of Lazio established and from 2018 to the National Registry, but currently, the Herd Book is still lacking. The most characteristic trait of the breed is the polyceraty, hence the name Quadricorna which means four-horned. Worldwide, there are numerous sheep populations showing polycerism [27] being present in both Northern European populations (sheep of Ireland, Felsen and Soa, Faröe, Shetland, Orkney and Man islands) and in populations of Tartaria, Nepal, Himalaya, Caucasus, Tunisia, Algeria, and in the Americas (Navajo-Churro). In Italy, four-horned sheep was mainly common among the Pagliarola breed in Abruzzo and in those of the Emilian Apennines [27]. Although still debated, several hypotheses have been argued to explain the origin of four-horned sheep. According to Marchi [27], the presence of multiple horns was common in all sheep derived from the paleo-Egyptian sheep and referable to Ovis longipes. On the other hand, other authors, primarily Sanson [28], argued that the origin of polycerate sheep can be traced back to the breed of Syria (Ovis aries asiatica), as described by Lemoigne [29].

To disentangle such a complexity and to shed light on the origin of the critically endangered Quadricorna sheep, we genotyped two populations from Central and Southern Italy using the Illumina OvineSNP50K BeadChip. The genomic architecture was therefore compared with other primitive and Mediterranean breeds, in order to gather insights into a more complete post-domestication evolutionary picture.

**Materials and methods**

**Ethics statement**

Blood samples were collected from sheep during normal health prophylaxis procedures by the Istituto Zooprofilattico Sperimentale del Lazio e della Toscana, in agreement with the recommendations of European Union Directive 2010/63/EU, to ensure appropriate animal care.

**Sampling and genetic diversity indices**

A total of 47 blood samples belonging to the Quadricorna sheep breed (31 females, 12 males and 4 individuals with no sex information) were stored and selected for the genotyping. These samples included 22 individuals belonging to Colliano (QUAD_SA) and 25 to Monte San Giovanni Campano (QUAD_FR). The two examined farms pursue similar breeding conditions in a semi-wild state with small additions of flaked feed when fresh forage is scarce.
The Illumina OvineSNP50K BeadChip was used to genotype the Quadricorna sheep population while the raw genotyped data for 33 breeds (944 individuals) were retrieved from previous works [8, 11, 20, 30–33] (S1 Table).

After merging all the genotyped samples, 53,854 SNPs were obtained. Using the software plink v1.9 [34], the genotypes were further filtered for missing call rate (--geno 0.1, removing 18,155 SNPs) and minor allele frequency (--maf 0.05, removing 19 SNPs), obtaining a final dataset consisting of 35,680 SNPs.

For each breed expected (He) and observed (Ho) heterozygosity and the inbreeding coefficient (FIS) was calculated using the R package dartR [35]. In addition, inbreeding coefficient based on runs of homozygosity (FROH) was calculated for each breed using the following parameters: the minimum length of the ROH was set to 4,000 (--homozyg-kb), the SNPs density was set to 100 (--homozyg-density) with a maximum gap between consecutive homozygous SNPs of 1,000 (--homozyg-gap), one missing SNP (--homozyg-window-missing), up to one possible heterozygous genotype was allowed in the ROH (--homozyg-window-het) and the minimum number of SNPs within a ROH was set to 50 (--homozyg-window-snp). Thus, the final inbreeding coefficients (FROH), have been calculated using the autosomal length as implemented in the genome-wide function of the R package detectRUNS [36].

Finally, the contemporary effective population size (Ne) for each breed was calculated using the software GONe [37] which uses the linkage disequilibrium (LD) between markers to infer very recent demographic changes. All parameters except the recombination rate (hc = 0.01) were set by default.

**Genetic structure and breed relationships**

To visualize the genetic differentiation among breeds, a principal component analysis (PCA) was performed using plink. The PCA was performed through a supervised approach using the pca-cluster flag in plink v1.9, in order to account for breed heterogeneity and outliers. Results of the centroids were then plotted using the R package ggplot2 [38].

The maximum clustering likelihood approach as implemented in the software admixture [39], was used to estimate the genetic structure, testing K values ranging from 1 to 50. For each K, we run three independent iterations, setting the cross-validation error procedure to 10-fold in order to choose the most informative number of K repartitions. The number of K before the separation of the two Quadricorna populations, was also used to plot the mean of the ancestral components in a pie fashion into a geographic map.

The phylogenetic relationships among breeds were inferred using an in-house R script which calculates the genetic Reynolds’ distance matrix and uses a neighbor-joining approach to compute the tree.

To assess the genetic drift and migration events among breeds, the maximum likelihood dendrogram was inferred using the software treemix [40]. The analysis was performed taking into account LD over blocks of 500 contiguous SNPs, using the Soay breed as outgroup, while allowing 10 migration events over 5 iterations. To evaluate the best number of migration edges, the R package OptM [41] was used to fit the parametric models to the likelihood values across runs.

**Results**

All genetic diversity indices are reported in Table 1 and Fig 1. The minimum value of He and Ho was observed in the Soay breed, while the highest values were observed in Bagnolese and Comisana breeds respectively. Concerning the FIS, the highest value was observed in the Lecce breed while the most negative value was for Sakiz. In general, we observed a higher
amount of negative FIS values in ether breeds considered primitive (SOAY, AFSH, SAKZ and SRMF) or with known recent history of admixture (BASC, SARB, SARW and NACH). Interestingly both Quadricorna populations showed negative Fis values (Fig 1B). The inbreeding estimated using runs of homozygosity (FROH) showed the highest values in Jacob and Soay, while the minimum value was observed in the Iranian LORB breed. The two Quadricorna populations showed different values of inbreeding, lower in the QUAD_FR than in the QUAD_SA (Fig 1A).

The contemporary effective population size (Ne) is reported in Table 1. The minimum value was observed in BASC while the maximum value in BIELL. The two Quadricorna

Table 1. Genetic diversity indices for all the analyzed breeds.

| Breed code | Location | N   | Ho    | SD   | He    | SD   | Fis   | Froh  | Ne |
|------------|----------|-----|-------|------|-------|------|-------|-------|----|
| 1          | SOAY     | Scozia | 110  | 0.269 | 0.189 | 0.266 | 0.184 | -0.008 | 0.127 | 40 |
| 2          | JACO     | USA   | 124  | 0.298 | 0.176 | 0.298 | 0.172 | 0.001  | 0.165 | 32 |
| 3          | NACH     | USA   | 36   | 0.370 | 0.166 | 0.354 | 0.145 | -0.032 | 0.053 | 26 |
| 4          | SRMF     | Sardinia | 24  | 0.330 | 0.188 | 0.314 | 0.165 | -0.030 | 0.084 | 25 |
| 5          | SARB     | Sardinia | 20  | 0.358 | 0.187 | 0.333 | 0.155 | -0.048 | 0.057 | 26 |
| 6          | SARW     | Sardinia | 24  | 0.361 | 0.168 | 0.352 | 0.146 | -0.003 | 0.036 | 33 |
| 7          | PINZ     | Sicily | 24   | 0.362 | 0.155 | 0.370 | 0.132 | 0.041  | 0.055 | 40 |
| 8          | VALB     | Sicily | 24   | 0.341 | 0.165 | 0.347 | 0.149 | 0.038  | 0.094 | 32 |
| 9          | BASC     | Sicily | 24   | 0.350 | 0.191 | 0.327 | 0.163 | -0.047 | 0.077 | 24 |
| 10         | COMI     | Sicily | 24   | 0.375 | 0.162 | 0.364 | 0.137 | -0.009 | 0.012 | 42 |
| 11         | SAMB     | Piemonte | 24  | 0.363 | 0.165 | 0.362 | 0.139 | 0.020  | 0.038 | 35 |
| 12         | DELL     | Piemonte | 24  | 0.345 | 0.174 | 0.343 | 0.152 | 0.015  | 0.077 | 30 |
| 13         | BIEL     | Piemonte | 23  | 0.361 | 0.162 | 0.362 | 0.138 | 0.025  | 0.030 | 71 |
| 14         | BERG     | Lombardy | 24  | 0.364 | 0.162 | 0.363 | 0.138 | 0.017  | 0.021 | 39 |
| 15         | ALPG     | Veneto | 24   | 0.353 | 0.172 | 0.352 | 0.147 | 0.017  | 0.048 | 33 |
| 16         | MASS     | Toscana | 24   | 0.357 | 0.170 | 0.351 | 0.147 | 0.002  | 0.048 | 34 |
| 17         | FABR     | Marche | 23   | 0.370 | 0.173 | 0.353 | 0.145 | -0.025 | 0.040 | 29 |
| 18         | APEN     | Umbria | 24   | 0.371 | 0.165 | 0.360 | 0.140 | -0.009 | 0.035 | 32 |
| 19         | SOPR     | Marche | 24   | 0.368 | 0.153 | 0.372 | 0.131 | 0.029  | 0.057 | 38 |
| 20         | MERI     | Abruzzo | 20  | 0.371 | 0.163 | 0.366 | 0.135 | 0.013  | 0.036 | 35 |
| 21         | GEPR     | Puglia | 24   | 0.365 | 0.170 | 0.352 | 0.146 | -0.015 | 0.057 | 28 |
| 22         | ALFG     | Puglia | 24   | 0.345 | 0.187 | 0.334 | 0.158 | -0.010 | 0.093 | 25 |
| 23         | ALTA     | Puglia | 22   | 0.365 | 0.165 | 0.360 | 0.142 | 0.009  | 0.061 | 32 |
| 24         | LECE     | Puglia | 23   | 0.352 | 0.154 | 0.367 | 0.134 | 0.062  | 0.077 | 39 |
| 25         | BAGN     | Campania | 23  | 0.368 | 0.154 | 0.373 | 0.130 | 0.033  | 0.047 | 40 |
| 26         | LATI     | Campania | 24  | 0.366 | 0.158 | 0.366 | 0.135 | 0.023  | 0.047 | 36 |
| 27         | QUAD_SA  | Campania | 22  | 0.345 | 0.200 | 0.318 | 0.169 | -0.060 | 0.088 | 34 |
| 28         | QUAD_FR  | Lazio | 25   | 0.372 | 0.176 | 0.349 | 0.148 | -0.045 | 0.053 | 26 |
| 29         | LESV     | Grecia  | 6    | 0.370 | 0.225 | 0.346 | 0.153 | 0.021  | 0.027 | 38 |
| 30         | CHIO     | Grecia  | 23   | 0.327 | 0.177 | 0.326 | 0.159 | 0.018  | 0.079 | 49 |
| 31         | SAKZ     | Turchia | 22  | 0.337 | 0.204 | 0.309 | 0.174 | -0.064 | 0.077 | 45 |
| 32         | KYMI     | Grecia  | 6    | 0.363 | 0.227 | 0.341 | 0.157 | 0.024  | 0.050 | 33 |
| 33         | LOAW     | Israele | 24   | 0.374 | 0.154 | 0.371 | 0.129 | 0.013  | 0.012 | 41 |
| 34         | AFSH     | Iran   | 25   | 0.369 | 0.165 | 0.355 | 0.142 | -0.018 | 0.008 | 32 |
| 35         | LORB     | Iran   | 30   | 0.371 | 0.155 | 0.364 | 0.135 | -0.001 | 0.005 | 38 |

Observed (Ho) and expected heterozygosity and their relative standard deviation (SD), inbreeding coefficients (Fis and Froh) and effective population size (Ne).

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populations showed different Ne estimates with the QUAD_FR displaying a very low value (26), while the QUAD_SA showed an intermediate value (34).

The supervised PCA is reported in Fig 2. The first principal component which accounted for 42.2% of the total variance, clearly distinguishes the two Iranian (AFSH and LORB), Navajo-Churro, Soay, Jacob and the two Quadricorna populations, from all the other breeds. On the other hand, the second principal component which accounted for 6.8% of the total variance, highlights a genetic gradient in accordance with the geography from Western Mediterranean areas (including Turkey and Greece) to the Italian Peninsula.

Fig 1. Inbreeding coefficients based on Froh (a) and Fis (b). Colours of the breeds are depicted according to the main genomic repartition at K = 8.

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The genetic repartition as highlighted by the admixture analysis showed a best K value at 40 (S1 Fig) where almost all breeds showed their own genetic pattern while for other breeds a marked sub-structuring is observable (Fig 3A). The admixture model, assuming two ancestral
populations (K = 2), first separated all the primitive breeds (JACO, SOAY) including Iranian (AFSH and LORB), Navajo-Churro and the two Quadricorna populations from all the other

**Fig 3.** Results of the admixture analysis for K values ranging from 2 to 8, 35 and 40 (a). For K = 8, the genomic components are shown on a geographic map (b). To simplify map representation, the Navajo-Churro breed has been placed in the ancestral origin area (Spain), rather than in the current reared region (Americas). See Table 1 for a full definition of the breeds.

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breeds. At this K value, a slight admixture is observable for the Iranian, Navajo-Churro and Quadricorna breeds which instead assume their own component at K = 3. The two Quadricorna populations separated with all the other breeds at K = 9, however, at higher K value, they display a different genetic admixture pattern (Fig 3B). Indeed, while the QUAD_SA population showed a rather homogeneous pattern, the QUAD_FR showed more mixed components up to K = 40.

The neighbor-joining tree based on Reynolds’ distances, basically confirms all the aforementioned results, showing a basal group which includes the Quadricorna breed nested within the Middle-East and primitive European breeds (AFSH, LORB, NACH, SOAY and JACO). A second group includes all the South-West Mediterranean breeds which do not form a monophyletic. In this analysis the Local Awassi clusters in a basal position of this group instead of grouping with the South-West Mediterranean breeds as highlighted in both the PCA and the admixture analyses (Figs 2 and 3). Finally, all the Italian breeds form a third monophyletic group in a derived position (Fig 4A).

These main topological clusters are also supported by the treemix analysis which showed the presence of these three main genetic groups according to the geography (Italian, South-West Mediterranean and Middle-East and primitive European breeds). Also, in this analysis the two Quadricorna populations fall close to the Middle-East breeds (Fig 4B). According to the likelihood fitted using the optM function, the most supported number of migration edges was between 4 and 6, thus, 5 migration events were considered and plotted (Fig 4B). Interestingly, the two strongest signals were inferred from NACH to the two Quadricorna populations and from BASC to the basal node of the South-West Mediterranean breeds. Other migration events were assigned from LATI to LOAW, from AFSH to LOAW and from ALFG to DELL.

**Discussion**

The genetic indices for the two Quadricorna populations in terms of expected and observed heterozygosity resulted in line with values of other local breeds in the Italian Peninsula (Table 1).

Also, the effective population size and inbreeding coefficients are comparable with those observed in most of the analyzed breeds. However, the two Quadricorna populations displayed slightly different genetic indices with the QUAD_SA showing higher diversity indices (Ho and He), effective population size (Ne) and lower inbreeding coefficients (FIS and FROH). Different patterns of genetic diversity indices between populations of the same breed are not surprising and have been commonly reported in many livestock species including local [26] and cosmopolitan breeds [42, 43]. Especially when considering residual populations at extinction risk and composed by few herds, the genetic drift can quickly act by reducing allele frequencies and increasing homozygosity. In this context, the genetic assessment is of primary importance in order to establish conservation and recovery plans that take into account the maintenance of adequate genetic diversity.

In all the population genetics analyses and phylogeographic reconstructions, the two Quadricorna populations from Central and Southern Italy cluster close to those breeds currently diffused near the sheep domestication centre (Fig 3B). Moreover, this genetic relationship is even more evident than those observed between the primitive breeds from Northern Europe (SOAY and JACO) and Middle-East breeds (AFSH and LORB). These results are particularly in line with several authors [28, 29], which argued that the origin of polycerate sheep can be traced back to the breed of Syria (Ovis aries asiatica). It is also interesting to note that our analysis does not seem to suggest any signal of recent admixture of the Quadricorna breed with other Italian breeds reared in neighbouring areas, pointing out a rather ancestral purity of the
Fig 4. Genetic relationships among breeds as inferred by the neighbor-joining tree based on Reynold’s distances (a) and by the maximum likelihood treemix analysis (b). The best number of migration events is shown with relative arrows. Colours of the breeds are depicted according to the main genomic repartition at K = 8. For a full definition of the breeds see Table 1.

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two populations. The genetic isolation of these residual Quadricorna populations might have been reinforced by the traditional vertical transhumance, that in turn, is based on the proximity between winter and summer pastures. Such a marginal pastoral system has been shown to have a Neolithic origin [44] and might have prevented large genetic exchanges as occurred for Merino-derived breeds in which the horizontal transhumance involving movements covering long distances, have favoured strong genetic connections among herds in the so called “Merinization” processes [45].

Although our analysis seems to trace back those residual populations to an early Neolithic diffusion probably following a Mediterranean route, several alternative hypotheses cannot be denied. First of all, a more recent diffusion from the Middle-East probably as a consequence of the increased marine trade from the Hellenistic to the Roman periods cannot be completely excluded. For example, written sources about livestock trade are reported in the De re Rustica of Lucius Junius Moderatus Columella (AD 4 – c.70), documenting the presence in the Italian Peninsula of livestock varieties from the Middle-East [46 and references therein]. Additional evidence coming from zooarchaeological and ancient DNA, showed similar outcomes, not only concerning the introduction of fine-fleeced sheep [47, 48]. Secondly, although we do not find evidence of recent introgression with other local Italian breeds, the inferred gene flow analysis suggested a strong migration edge from the Navajo-Churro to the Quadricorna populations (Fig 4B). This result is not straightforward since the southern Italian Peninsula has had a centuries-old Spanish domination. Therefore, it is not surprising that in parallel with the introduction of the Merino breeds in Italy, other not fine wool breeds, such as the ancestor of the current Churra, might have admixed with some local flocks giving rise to the current Quadricorna breed. In this regard it should be noted that the ancestors of the Navajo-Churro reached the Southern United States in parallel with the first Spanish settlers which brought with them small Iberian nuclei to supply for meat and dairy. During the eighteen century, some flocks were then selected by native Americans for textile purposes giving rise to the current Navajo-Churro. The close genomic relationship between the Navajo-Churro and the current Spanish Churra has been reported in several studies [49–51] and also assessed by us (data not shown).

From a morphological point of view, it is interesting to note that both Quadricorna and primitive traits-related breeds, aside from sharing a common genomic ancestral background, are also characterized by the frequent presence of four-horned individuals. Four-horned sheep were historically widely distributed in Europe, due to advantages in the size and arrangement of the horns in anti-predatory strategies [27]. However, this trait has subsequently become absent in ewes and rare in rams due to the long-term artificial selection [18, 19]. The common presence in the Quadricorna breed, of four-horned individuals in both sexes seems to support an early introduction likely occurred alongside the first wave of post-Neolithic human diffusion. Finally, another common trait between the Quadricorna breed and other primitive traits-related breeds with high frequency of four-horned individuals (Jacob and Navajo-Churro), is the presence of widespread eyelid abnormality with variable severity. However, a recent study seems to deny a pleiotropic effect of the genes underlying these two morphological traits [49].

Future studies would be desirable in order to shed light into the mode and the time of diffusion of this ancestral breed, moreover we believe that our results should be considered in light to plan adequate conservation measures in order to recover the population and to valorise their related cultural and commercial products.

**Conclusion**

Sheep are among the first animals to be domesticated in the Fertile Crescent and as initially reared mainly for meat production, they reached Europe and Mediterranean areas around
8,000 YBP. However, subsequent migration waves mainly addressed at satisfying the wool production, have probably replaced the ancestral genetic signature. In spite of this, several scattered primitive populations characterized by both genetic and morphological ancestral traits are still scattered on the very edge of North Europe. Our analysis showed that a comparable ancestral genetic background is still present also in the Quadricorna sheep breed, indicating that the genetic diversity in terms of ancestral components of Southern Europe might be underestimated. This outcome is of considerable interest when considering the potential of genes that confer high “capacity for constructivism” to specific environments and to preserve and valorise local products alongside with their long-standing history in the Italian Peninsula.

Supporting information
S1 Fig. Cross-validation error for the admixture analysis for K values ranging from 2 to 50. (PDF)
S1 Table. Names, codes, geographic origin and all references of the analyzed breeds. (DOCX)

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