



Resilience to Historical Human Manipulations in the Genomic Variation of Italian Wild Boar Populations

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Human activities can globally modify natural ecosystems determining ecological, demographic and range perturbations for several animal species. These changes can jeopardize native gene pools in different ways, leading either to genetic homogenization, or conversely, to the split into genetically divergent demes. In the past decades, most European wild boar (*Sus scrofa*) populations were heavily managed by humans. Anthropogenic manipulations have strongly affected also Italian populations through heavy hunting, translocations and reintroductions that might have deeply modified their original gene pools. In this study, exploiting the availability of the well-mapped porcine genome, we applied genomic tools to explore genome-wide variability in Italian wild boar populations, investigate their genetic structure and detect signatures of possible introgression from domestic pigs and non-native wild boar. Genomic data from 134 wild boar sampled in six areas of peninsular Italy and in Sardinia were gathered using the Illumina Porcine SNP60 BeadChip (60k Single Nucleotide Polymorphisms – SNPs) and compared with reference genotypes from European specimens and from domestic pigs (both commercial and Italian local breeds), using multivariate and maximum-likelihood approaches. Pairwise F_{ST} values, multivariate analysis and assignment procedures indicated that Italian populations were highly differentiated from all the other analyzed European wild boar populations. Overall, a lower heterozygosity was found in the Italian population than in the other European regions. The most diverging populations in Castelporziano Presidential Estate and Maremma Regional Park can be the result of long-lasting isolation, reduced population size and genetic drift. Conversely, an unexpected similarity was found among Apennine populations, even at high distances. Signatures of introgression from both non-Italian wild boar and domestic breeds were very limited. To summarize, we successfully applied genome-wide procedures to explore, for the first time, the genomic diversity of Italian wild boar,

demonstrating that they represent a strongly heterogeneous assemblage of demes with different demographic and manipulation histories. Nonetheless, our results suggest that a native component of genomic variation is predominant over exogenous ones in most populations.

Keywords: *Sus scrofa*, SNPs, endemic diversity, genetic structure, admixture, genomic introgression, translocation, wild pigs

INTRODUCTION

Animal species can be strongly managed by humans across their natural ranges, being harvested, translocated or sometimes admixed with alien or domesticated forms. All these direct actions can amplify the effects of other ecological perturbations, such as habitat or landscape modifications, contributing to jeopardizing the gene pool of wild populations and their adaptive capacities (Allendorf et al., 2008; Laikre et al., 2010; Boyce et al., 2011; Apollonio et al., 2014).

In Europe, large mammals, that have survived as metapopulations after experiencing demographic and range fluctuations, mainly due to human impacts, are recently spreading to form larger continuous populations (Apollonio et al., 2010a; Chapron et al., 2014; Martin et al., 2020). This apparent new range continuity may underlie genetic gaps which are the legacy of a past fragmentation and that are tricky to be detected. On the opposite, admixture between species, subspecies or populations can take place at suture zones or at release sites, leading to a local increase of genetic variation and to a potential loss of local adaptations.

In the past decades, most genetic investigations to identify cryptic population structure or local admixture in different species were conducted relying on limited sets of autosomal microsatellites (Frantz et al., 2006; Dellicour et al., 2011; Hindrikson et al., 2013; Olano-Marin et al., 2014; Bioss et al., 2015; Silva et al., 2018). However, the genomic era has provided new tools and a higher diagnostic power to detect gene pool differences and admixture signals, creating the conditions to disentangle complicated patterns due to high levels of human disturbance.

The European wild boar (*Sus scrofa*) population is a good example of the aforementioned changes: after a strong decline resulting in a fragmented and scattered distribution, it is now covering almost continuously all the continent up to the Baltic countries (Apollonio et al., 2010a) with a steadily increasing trend (Massei et al., 2015). In Italy, the species went through demographic declines and range contractions until the first half of the XX century (Ghigi, 1911, 1950), followed by a continuous re-expansion (Apollonio et al., 1988) that was sustained by socio-economic changes, leading to habitat recovery, and by the release of captive-reared or imported individuals (Apollonio et al., 2010b). After a progressive increase, the wild boar today represents one of the most invasive and impacting animals in the Country, likely due also to its genetic make-up which might have played a crucial role in its adaptation to a variety of different environmental and ecological contexts.

During the major range contraction, Italian wild boar survived only in the coastal Tyrrhenian region of Central Italy (named Maremma), in a few central-southern areas of the peninsula, where a relict isolated population persisted within a formerly royal hunting estate that became presidential estate since 1948 (Castelporziano), and in the island of Sardinia (Ghigi, 1911).

Morphometric analyses induced De Beaux and Festa (1927) to propose the Maremma and Sardinian populations as belonging to two different endemic subspecies, respectively, *Sus scrofa majori* and *Sus scrofa meridionalis*. However, the population recovery, which mostly followed the World War II, involved both the natural spread from existing populations (like the recolonization of the Western Alps from France) and artificial processes such as reintroductions and population restocking (Monaco et al., 2007; Apollonio et al., 2010b). These latter operations implied a massive release of boars from different sources, that possibly admixed with local wild stocks. In the assessment of the status of Eurasian *Sus scrofa* populations dated 1993, the IUCN Specialist Group reported the Castelporziano population as likely representing the “only surviving pure-bred native stock” in Italy (Oliver et al., 1993). The first genetic studies on Italian wild boar, based on mitochondrial DNA (mtDNA) sequencing, revealed the existence of an endemic Italian mtDNA lineage, occurring both in historical and modern samples (Randi, 1995; Larson et al., 2005; Scandura et al., 2008, 2009), that ended up to be isolated in Italy during glacial periods (Vilaça et al., 2014; Maselli et al., 2016; Khederzadeh et al., 2019). Further studies confirmed the occurrence of native genetic variation also in autosomal regions, although detecting signatures of local admixture with non-native stocks (Vernesi et al., 2003; Scandura et al., 2008; Veličković et al., 2016). Nonetheless, this information was only partial and not conclusive, since these studies were based on a limited number of molecular markers and the number of sampled populations and individuals analyzed were poorly representative of the variety of situations that are likely to occur in the Italian peninsula. More refined details were obtained on the Sardinian populations, which, despite the intricate pattern of genetic structure and local introgression from non-native wild boar and domestic pigs, still retain a meaningful proportion of endemic diversity, well-distinguishing them from the related peninsular conspecifics (Scandura et al., 2008, 2009, 2011; Iacolina et al., 2016).

However, all studies so far conducted on this species in Italy have experienced the same difficulties to untangle local effects of human manipulation, including the admixture among individuals from multiple wild and domestic sources (Vernesi et al., 2003; Scandura et al., 2011; Veličković et al., 2016; Canu et al., 2018).

The advent of genome-wide single nucleotide polymorphisms (SNPs) can offer practical solutions. Such markers have, indeed, the advantage to be the most widespread source of standing variation at both neutral and non-neutral genomic regions, and being bi-allelic, they permit to easily gather fully comparable data, which can be also merged with already published and publicly available datasets.

Additionally, the development of genome-wide genotyping assays in the domestic pig (Porcine SNP60 BeadChip; Ramos et al., 2009) and the availability of comprehensive genome data for this model species provide a powerful tool to explore genomic variation in wild boar populations and to investigate population history, as well as patterns of differentiation and introgression (Goedbloed et al., 2013a,b; Manunza et al., 2013; Iacolina et al., 2016; Alexandri et al., 2017; Smyser et al., 2020).

Therefore, in this study we genotyped a set of Italian wild boar collected from different peninsular and insular areas using the Porcine SNP60 BeadChip, which were then combined and analyzed together with publicly available genotypes of domestic pigs and wild boar from several European populations. We built an exhaustive genome-wide dataset including a good representation of the overall genomic variation so far available for the Italian population, which was analyzed by multivariate and maximum-likelihood approaches to specifically: (i) assess levels of genomic variation in the Italian wild boar populations; (ii) detect local population structure and quantify the extent of genomic differentiation; (iii) investigate the genomic make-up of the most differentiated local populations, distinguishing between multiple sources of standing variation (native vs. non-native wild boar, and commercial vs. local domestic pig breeds).

MATERIALS AND METHODS

Sampling, Genotyping and Dataset Building

We analyzed 98 wild boar (WB) individuals collected in six administrative regions spanning the Italian peninsula (Aosta Valley, Liguria, Tuscany, Latium and Calabria, **Figure 1**). Sampling areas differ in latitude, orography, habitat, human density, and WB presence. Moreover, for comparative purposes we included in the analyses 7 rural domestic pigs (DP) from the same area in Calabria where WB were sampled.

DNA was extracted from muscular tissue samples using the Qiagen DNAeasy Blood and Tissue kit (Qiagen Inc., Hilden, Germany) according to the manufacturer's instructions, quantified using the QubitTM Fluorometer (Invitrogen Thermo Fisher Scientific, Waltham, MA, United States) and normalized to a final concentration of 25–50 ng/μl.

DNA samples were genotyped using the Illumina Porcine SNP60 BeadChip v2 (Illumina Inc., San Diego, CA, United States; Ramos et al., 2009) at AIA¹ and the results were combined with 164 additional homologous WB genotypes, including 36 samples from the Italian populations of Sardinia and Tuscany and 128 spanning most of the European distribution range. All

these additional individuals were previously analyzed in a study focused on the Sardinian WB (Iacolina et al., 2016). A total of 103 porcine genotypes (56 from Italian local breeds – LocDP – and 47 from commercial breeds – ComDP) previously analyzed from Iacolina et al. (2016) and AIA were added to verify possible genomic introgression into WB populations (**Table 1**).

An overall dataset of 372 individuals (262 WB and 110 DP, **Table 1**) genotyped at 48,296 autosomal SNPs was thus obtained.

The number of available genotypes to be included in the dataset was selected so as to limit bias in the analyses introduced by unequal sample sizes. Based on previous studies (Goedbloed et al., 2013b; Alexandri et al., 2017; Iacolina et al., 2018), we grouped European (non-Italian) samples into four macro-regions: Iberia (WIbe), Central-Western Europe (WCW), Central-Eastern Europe (WCE), South-Eastern Europe (WSE, **Table 1**).

Dataset Filtering

The initial dataset was processed for quality control and filtering in the SNP & Variation Suite 8.0.1 (SVS, Golden Helix Inc., Bozeman, MT, United States) keeping only samples and loci with high genotyping rates (>0.95), and checking for related individuals through Identity By Descent (IBD) pairwise tests to remove one individual per pair where IBD > 0.5. After data filtering for quality and IBD, our dataset consisted of 258 WB, 134 from Italy and 124 from other European countries, and 109 DP from commercial and local breeds, giving a total of 367 individuals genotyped at 47,809 autosomal SNPs (**Table 1**).

This dataset (hereafter called 48K) was then pruned for linkage disequilibrium (LD, $r^2 > 0.5$ calculated along sliding windows of 50 SNPs) to delve into overall population structure, leaving 30,218 SNPs (30K).

We further explored WB population structure and composition. In PLINK 1.9 (Chang et al., 2015) we created a dataset including all WB ($n = 249$) and another dataset including only the Italian WB ($n = 125$) but excluding Aosta Valley because of its genetic composition highlighted by the first analyses (see section “Results”). Both datasets contained SNPs with a minimum minor allele frequency (MAF) of 0.05, in order to exclude rare alleles which can bias structure analysis, in linkage (LE, $r^2 < 0.3$, windows of 50 SNPs) and Hardy-Weinberg equilibrium (HWE, p -value > 0.001), in order to safely make inferences on neutral loci (Jungerius et al., 2005; Purcell et al., 2007; Benestan et al., 2016), thus resulting in 13,218 SNPs (13K) and 14,155 SNPs (14K), respectively.

Finally, specifically for migration analysis in TREEMIX (see below), WB data were merged with 11 publicly available *Sus barbatus* (SB), used as outgroup, genotyped with the same SNP chip (Yang et al., 2017). We kept only SNPs with no missing values that were pruned for high LD ($r^2 > 0.5$) in PLINK (see **Supplementary Material**).

Genetic Variation and Differentiation

Genetic variability in our data (48K) was investigated by computing with PLINK observed (H_o) and expected (H_e) heterozygosity, minor allele frequency (MAF), and number of polymorphic loci for each geographic region and separately

¹ www.aia.it

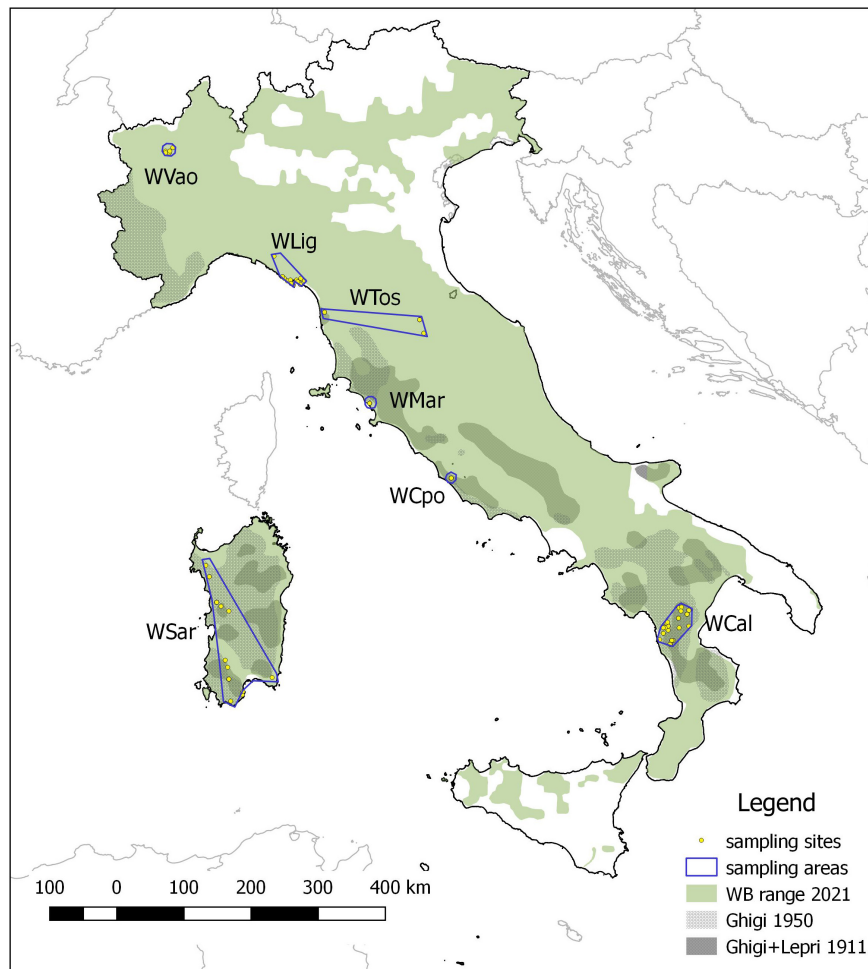


FIGURE 1 | Wild boar historical and current distribution in Italy and sampling areas. The historical range refers to the minimum extension reached at the beginning (Ghigi, 1911; Lepri, 1911) and at the middle of the XX century (Ghigi, 1950), which was followed by a progressive expansion leading to the current distribution.

for each Italian WB population. All available genotypes were used, aiming at verifying the real genomic variability within the sampled populations, regardless of the cause of their similarities/dissimilarities (i.e., including possible hybrids).

Subsequently, we assessed levels of genetic differentiation within the Italian populations, and between Italian samples and reference WB and DP populations using the 30K dataset filtered for high LD. Among the different parameters that can be estimated for this purpose, we selected two indexes, a fixation index (pairwise F_{ST}) and a measure of allelic differentiation (Jost's D , Jost, 2008), as they can be considered complementary in comparing allele distributions between populations (see Jost et al., 2018). We used ARLEQUIN v3.5 (Excoffier and Lischer, 2010) for pairwise F_{ST} calculations and the R package *mmom* (Winter, 2012) for Jost's D . With the same dataset, we also explored the overall genetic structure in our samples by a Principal Component Analysis (PCA) performed in SVS, using all WB and DP together and only WB genotypes.

Then, we focused on the Italian WB, using the 14K dataset and running a Discriminant Analysis of Principal Components

(DAPC) in the R package *ade4* v2.1.1 (Jombart, 2008; Jombart and Ahmed, 2011); the function `find.clusters` was used to identify the most likely grouping and a cross-validation approach allowed to determine the best number of Principal Components (PCs) to retain. Furthermore, we evaluated the contribution of different populations and groupings to the overall genomic differentiation among Italian populations performing an Analysis of Molecular Variance (AMOVA; Excoffier et al., 1992) using the R package *poppr* v2.8.5 (Kamvar et al., 2014, 2015) and the method implemented in R package *ade4* (Dray and Dufour, 2007).

Admixture Analyses

Once obtained an overview of the overall genetic variation and local differentiation among Italian samples, we investigated the possible impact of anthropogenic drivers on the observed genetic make-up. Indeed, hybridization with DPs or artificial gene flow with distant populations (i.e., translocations) can generate admixed WB populations, either geographically localized or more widespread in case of high levels of human-mediated gene flow;

TABLE 1 | List of Italian and European wild boar (WB) and domestic pig (DP) samples used in the study.

Population	Code	Macro-region	Wild/ Domestic	N	Source
Castelporziano	WCpo	WIta	Wild	19	This study
Liguria	WLig	WIta	Wild	20	This study
Pollino N.P.	WCAl	WIta	Wild	36	This study
Aosta Valley	WVao	WIta	Wild	9	This study
Maremma	WMar	WIta	Wild	14	This study
North Tuscany	WTos	WIta	Wild	13	Iacolina et al., 2016
Sardinia	WSar	–	Wild	23	Iacolina et al., 2016
Total Italian WB				134	
Spain	WSpa	Wlbe	Wild	7	Iacolina et al., 2016
Portugal	Wpor	Wlbe	Wild	10	Iacolina et al., 2016
France	Wfra	WCW	Wild	15	Iacolina et al., 2016
Luxembourg	Wlux	WCW	Wild	4	Iacolina et al., 2016
Germany	WGer	WCW	Wild	5	Iacolina et al., 2016
Netherlands	Wned	WCW	Wild	15	Iacolina et al., 2016
Greece	Wgre	WSE	Wild	9	Iacolina et al., 2016
Bulgaria	WBul	WSE	Wild	5	Iacolina et al., 2016
Croatia	Wcro	WCE	Wild	16	Iacolina et al., 2016
Serbia	Wser	WCE	Wild	4	Iacolina et al., 2016
Slovenia	WSln	WCE	Wild	13	Iacolina et al., 2016
Slovakia	WSlk	WCE	Wild	5	Iacolina et al., 2016
Poland	Wpol	WCE	Wild	9	Iacolina et al., 2016
Hungary	WHun	WCE	Wild	11	Iacolina et al., 2016
Greece	Wgre	WSE	Wild	9	Iacolina et al., 2016
Bulgaria	WBul	WSE	Wild	5	Iacolina et al., 2016
Total European WB				128	
Casertana	DCas	LocDP	Domestic	10	Iacolina et al., 2016
Nera Siciliana	DNsi	LocDP	Domestic	10	Iacolina et al., 2016
Mora Romagnola	DMro	LocDP	Domestic	8	Iacolina et al., 2016
Calabrese	DCal	LocDP	Domestic	10	Iacolina et al., 2016
Cinta Senese	DCse	LocDP	Domestic	10	Iacolina et al., 2016
Sardinia rural	DSar	LocDP	Domestic	8	Iacolina et al., 2016
Pollino rural	DPol	LocDP	Domestic	7	This study
Berkshire	DBer	ComDP	Domestic	10	Iacolina et al., 2016
Duroc	DDur	ComDP	Domestic	10	Iacolina et al., 2016
Large White	DLwh	ComDP	Domestic	8	Iacolina et al., 2016
Yorkshire	DYor	ComDP	Domestic	9	Iacolina et al., 2016
Pietrain	DPie	ComDP	Domestic	10	Iacolina et al., 2016
Total DP				110	

For each population the adopted code, geographic macro-region (WIta, Italian peninsula; Wlbe, Iberian peninsula; WCW, Central-Western Europe; WCE, Central Eastern Europe; WSE, South Eastern Europe) or pig group (LocDP, local breed; ComDP, commercial breed), sample size (N) and source are reported.

this latter case can lead to “genetic homogenization” (*sensu* Olden et al., 2004).

A specific analysis to ascertain possible differences in the local levels of admixture was performed using the software ADMIXTURE v1.2.3 (Alexander et al., 2009). We ran the program with the WB + DP dataset (30K) for *K* ranging from 1 to 30 and determined the best *K* according to the lowest cross-validation error (Alexander et al., 2009). At the best *K* we checked for domestic and

extra-Italian genetic components in the Italian populations. As a signature of genomic admixture might appear also in the reference DP breeds, we run the cluster analysis twice, with and without DP, to ascertain whether possible past hybridization events were masked by the variegated nature of some DPs.

Since the interpretation of ADMIXTURE results can be cumbersome (Lawson et al., 2018), on account of the available historical information, we considered the possibility that past introductions of European WB (from multiple sources) to the Italian peninsula had produced a composite genomic make-up, not associable to any of the reference populations. Therefore, we further verified the origin and composition of the Italian populations putatively carrying signs of past admixtures with European WB (see section “Admixture Analyses” in “Results”) using two different approaches. Specifically, we performed a PCA with *ade4* on the WB 13K dataset, leaving out the populations that were suspected to be admixed; we subsequently projected their coordinates on the first two axis (i.e., PC1 and PC2) with the function *suprow* in *ade4*. This approach eliminates the possible bias in PCA calculation introduced by admixed populations and allows to evaluate their actual origin according to their projection on the PCs estimated from the reference populations (McVean, 2009; see also Smyser et al., 2020).

We also tested for past gene flow from other European populations into one or more Italian WB populations with the program TREEMIX (Pickrell and Pritchard, 2012). The approach used in TREEMIX consists first in drawing a maximum likelihood (ML) tree, and then to add migration edges to verify if the data are better explained by a network-like model. As the program corrects for LD by dividing the whole dataset into blocks of *k* SNPs (where *k* is specified by the user), using PLINK we first computed the LD decay as the distance at which $r^2 < 0.2$ (option *-r2*), and then we obtained the mean number of SNPs in our dataset within that distance with a custom R script. We rooted the ML tree with no migration events on WSE (see **Supplementary Material** for details) and built the ML tree sampling blocks of 200 SNPs (option *-k*) and activating the *-global* option for a global rearrangement after adding all the populations to the tree. We then started to add migration events from 1 to 21 (option *-m*), supplying at each run the tree with *n*-1 migration edges (option *-g*) to save time. We calculated the variance explained by each model thus obtained with the function *get_f* that comes with the program in an R script; after identifying in *m* = 5 the knee point (i.e., where the increment of the variance explained started to be less pronounced) we ran 200 bootstrap replicates (option *-bootstrap*), and then we used the functions provided by Zecca et al. (2020) to visualize the best ML tree and to calculate the migration support (MS) and the extended migration support (MSE) across our replicates for the migration events involving the Italian populations (see Zecca et al., 2020 for the description of the indexes). We finally calculated the *f4* statistic with the program *fourpop*, included in TREEMIX, to formally test for admixture. It requires four populations (A, B, C, and D) and tests if the tree topology (A, B; C, D) is correct; if the Z score is significantly different from zero it means that the history of

the four populations is not strictly tree-like (Reich et al., 2009; Pickrell and Pritchard, 2012).

RESULTS

Genomic Diversity

Minor allele frequencies were comparable among the Italian populations (ranging from 0.147 in WCpo to 0.193 in WVao) and appeared slightly lower in Italy than in other European geographic regions (0.169 vs. 0.186–0.194). The number of polymorphic loci varied from the lowest value observed in WCpo ($n = 18,544$) to the highest in WCal ($n = 28,401$) for Italy, whereas in the rest of Europe it ranged between $n = 26,944$ in Wlbe and $n = 28,598$ in WCE (Table 2). Overall, heterozygosity levels were generally higher in DP than in WB. In particular, observed and expected heterozygosity differed significantly, especially in WB, with all Italian populations showing a remarkable deficit of heterozygotes (Wilcoxon test p -value < 0.001), which was less pronounced in the other WB populations. The lowest values of H_o were observed in WLig and WCpo (0.135 and 0.139, respectively), whilst WSar showed the highest value (0.215, see Table 2).

Genomic Differentiation

The exploratory PCA clearly separated WB from DP along PC1, and the Italian WB from the European populations along PC2 (Figure 2A). Individuals from the Aosta Valley population (WVao, North Italy) clearly clustered with the European

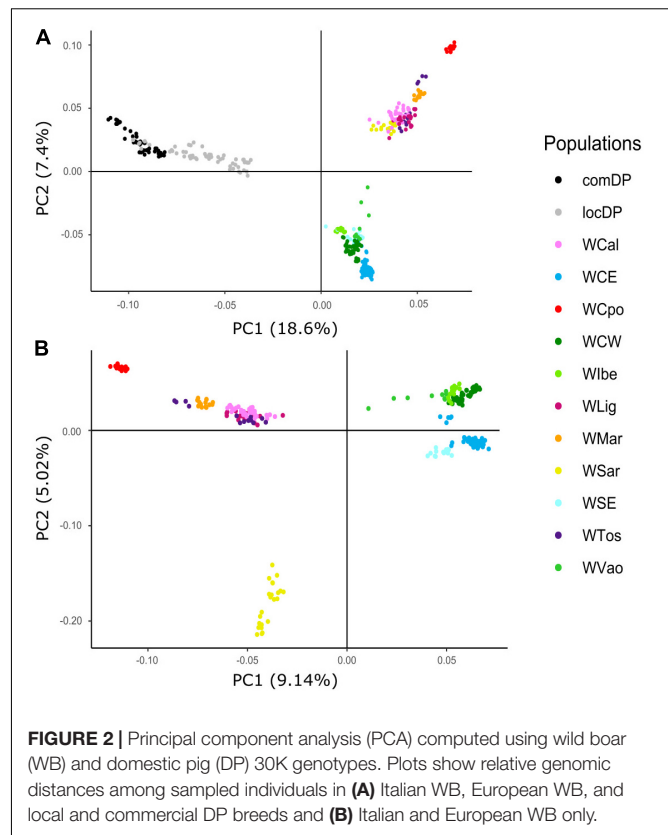


TABLE 2 | Genetic diversity in Italian wild boar populations compared to the reference European macro-regions and domestic pig groups (local and commercial breeds).

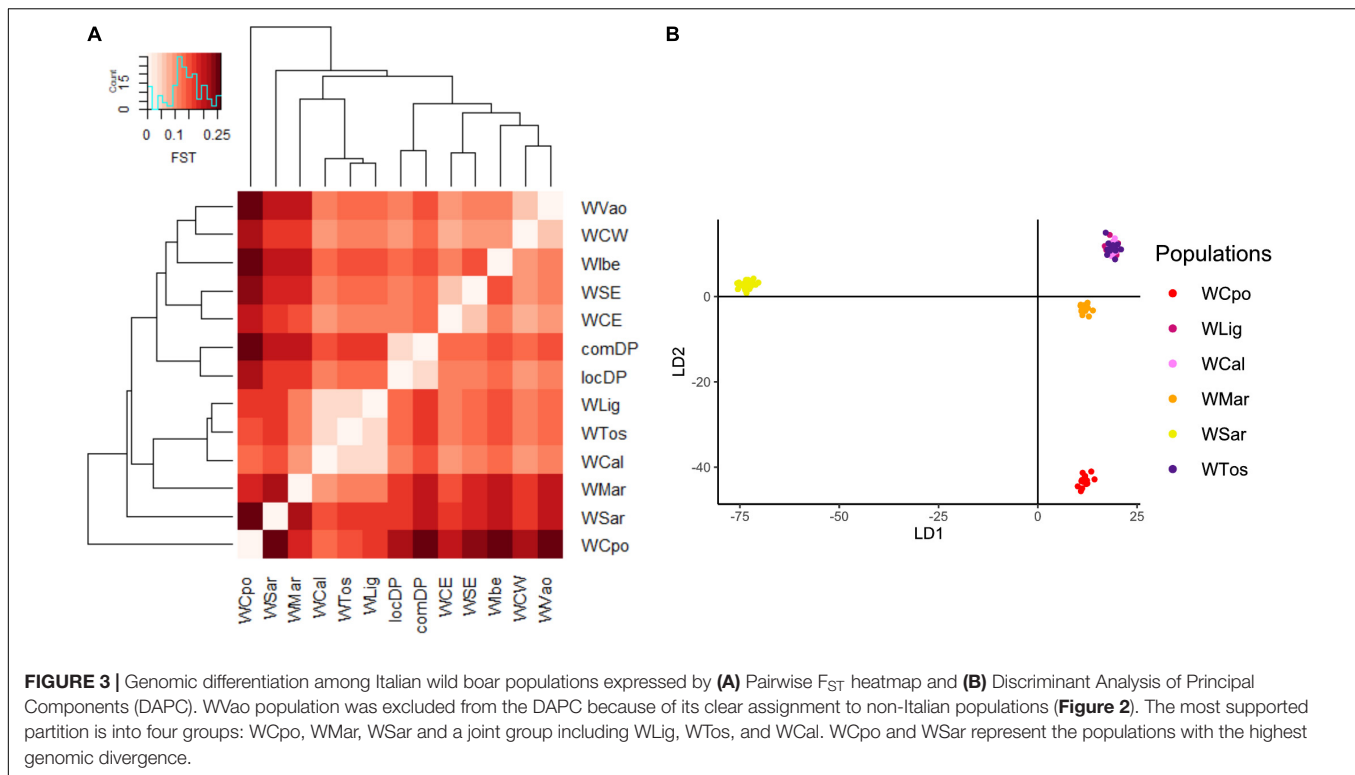
	Population	N	MAF (SD)	NP	Ho	He
Wild Boar (WB)	WCpo	19	0.147 (0.241)	18,544	0.139	0.312
	WLig	20	0.169 (0.204)	27,079	0.135	0.314
	WCal	36	0.173 (0.189)	28,401	0.190	0.282
	WMar	14	0.161 (0.229)	21,776	0.198	0.276
	WTos	13	0.165 (0.206)	24,440	0.173	0.224
	WVao	9	0.193 (0.238)	27,852	0.210	0.258
	WSar	23	0.173 (0.236)	22,973	0.215	0.263
	Wlta	111	0.169 (0.185)	27,697	0.186	0.212
	Wlbe	16	0.193 (0.228)	26,944	0.189	0.207
	WCW	36	0.186 (0.203)	28,056	0.194	0.221
	WCE	58	0.186 (0.200)	28,598	0.206	0.223
	WSE	14	0.194 (0.216)	27,895	0.201	0.219
	LocDP	62	0.264 (0.193)	40,791	0.250	0.314
Domestic Pig (DP)	ComDP	47	0.306 (0.203)	42,868	0.267	0.343

For each population, sample size (N), minor allele frequency (MAF) and corresponding standard deviation (SD), number of polymorphic sites (NP), mean observed heterozygosity (H_o) and mean expected heterozygosity (H_e) are reported. Areas/breeds corresponding to the population codes are reported in Table 1. Values referring to European macro-regions and domestic pig groups are in bold.

populations (see also ADMIXTURE results). When DP were removed from the analyses, WB population structure became clearer (Figure 2B): the most diverging Italian populations were WCpo and WSar. WMar, WTos, WLig, and WCal clustered in close proximity, but WTos was split into two separate groups, corresponding, respectively, to the western (coastal) and eastern (Apennine) areas in Figure 1. These results are also mirrored by pairwise F_{ST} (Figure 3A) and Jost's D (Supplementary Table 1) values, as well as by the DAPC restricted to the Italian populations (WVao excluded, Figure 3B). The most supported partition of the Italian populations on the basis of BIC values corresponded to four discrete groups (Supplementary Figures 1A,B), specifically: WSar (group 1), WMar (group 2), WCpo (group 3), and WLig-WTos-WCal (group 4). Coherently, the AMOVA showed that this partition was associated with the highest percentage of variance between groups explaining 15.26% of the total variance in the Italian sample ($p < 0.001$), more than double compared to the percentage of variance between populations within groups (6.88%, Supplementary Table 2).

Admixture Analyses

A contribution to clarify the nature of the identified groupings arose from the admixture analyses. ADMIXTURE results depicted a variegated picture of the Italian populations. The lowest cross validation error was reached at $K = 19$ but local minima were observed at $K = 12$ and $K = 16$ (Figure 4B). At these K values, WVao clearly clustered with the western European populations, consistently with its geographical proximity. In



agreement with PCA and F_{ST} results, starting from $K = 12$, the other Italian populations were roughly assigned each to their own clusters, except the group WLig-WCal-WTos (**Figure 4A**). This group formed a single cluster at $K = 12$ and was split into two clusters at higher K values. Striking signals of introgression from domestic pigs or from other European populations did not emerged in the Italian samples, with a few exceptions only in WCal and some individuals from WLig and WMar.

Based on the results above and on historical information, we hypothesized that WLig, WTos, and WCal appeared to be similar while diverging from other Italian populations, because they were carrying a unique genomic assemblage influenced by a past introgression from non-Italian gene pools (see section “Discussion”). The projected PCA confirmed, indeed, a much higher proximity of the three tested Italian populations to the European clusters (with respect to **Figure 2**) and a central position in the plots, providing an indirect support to the hypothesis of past admixture events (**Figure 5**). Like in the previous PCA and DAPC, most of the genotypes from WLig, WTos, and WCal overlapped, confirming their genomic homogeneity despite their geographical location.

The ML tree obtained in TREEMIX (**Figure 6** and see also **Supplementary Figure 2**) confirms the overall WB structure. With no migration edges, the ML tree explained 97.7% of the variance. The long branches of WCpo, WMar, and WSar confirmed their high genomic differentiation from the other Italian populations, under the effect of drift. WVao fell in a separate branch and appeared as a sister group to WCW. With five migration edges the percentage of variance explained by the model increased to 99.7%. According to the MS and MSE

(**Supplementary Tables 3, 4**), both WSar and WVao received a migration from populations of central Italy, whereas WCal showed a signature of immigration from the WMar-WCpo lineage. The f_4 statistic confirmed that WVao is admixed with the Italian populations ($Z > 2$, **Supplementary Table 5**).

DISCUSSION

In the present study, we explored, for the first time, the genome-wide diversity of the Italian WB, analyzing a set of populations sampled from several locations in the Italian peninsula and Sardinia. We expected that the present overall levels of diversity and the genomic make-up of the Italian populations were largely influenced by their demographic histories and human manipulations.

The progressive decline and range contraction of the species in Italy that occurred between the end of the XIX and the beginning of the XX century caused its fragmentation into small isolated demes (Ghigi, 1917, 1950). These nuclei have lasted for decades until the more recent re-expansion, that started after World War II, and have likely suffered bottlenecks accompanied by a loss of genetic variation and changes in allele frequencies. This historical process might explain why the levels of genomic diversity are lower in Italy than in most European populations (**Table 2**), particularly those recorded in areas that suffered prolonged and strong isolation (e.g., Presidential Estate of Castelporziano).

However, the translocations of wild animals and the release of captive specimens that occurred in the last 60 years, have taken

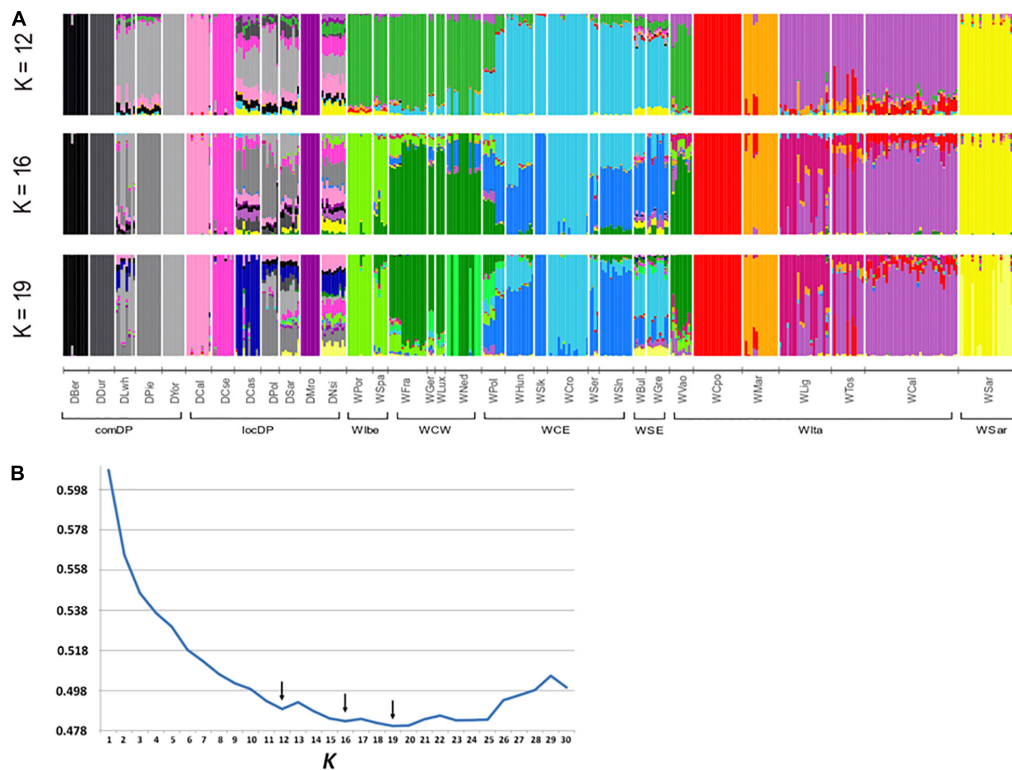


FIGURE 4 | Admixture analyses in wild boar (WB) populations conducted using the software ADMIXTURE. **(A)** Membership proportions of every genotyped individual to each of the K inferred clusters are shown as different colors. Reference European WB and domestic pig (DP) genotypes are included to test for genomic introgression in Italian WB. Histograms are reported for the best K values indicated by the cross-validation error **(B)**. Arrows in **(B)** indicate local minima. WB and DP population codes are shown in **Table 1**.

place with no homogeneity throughout the country and have determined a human-mediated gene flow that can be responsible for a fairly higher variation in some zones, where individuals of different origins may have mixed up.

Indeed, for a long time the Italian legislation allowed the release of wild boar for restocking or reintroduction. Yet, since the end of 1980s a ban was introduced, first in single regions and then, only in 2015, at national level.

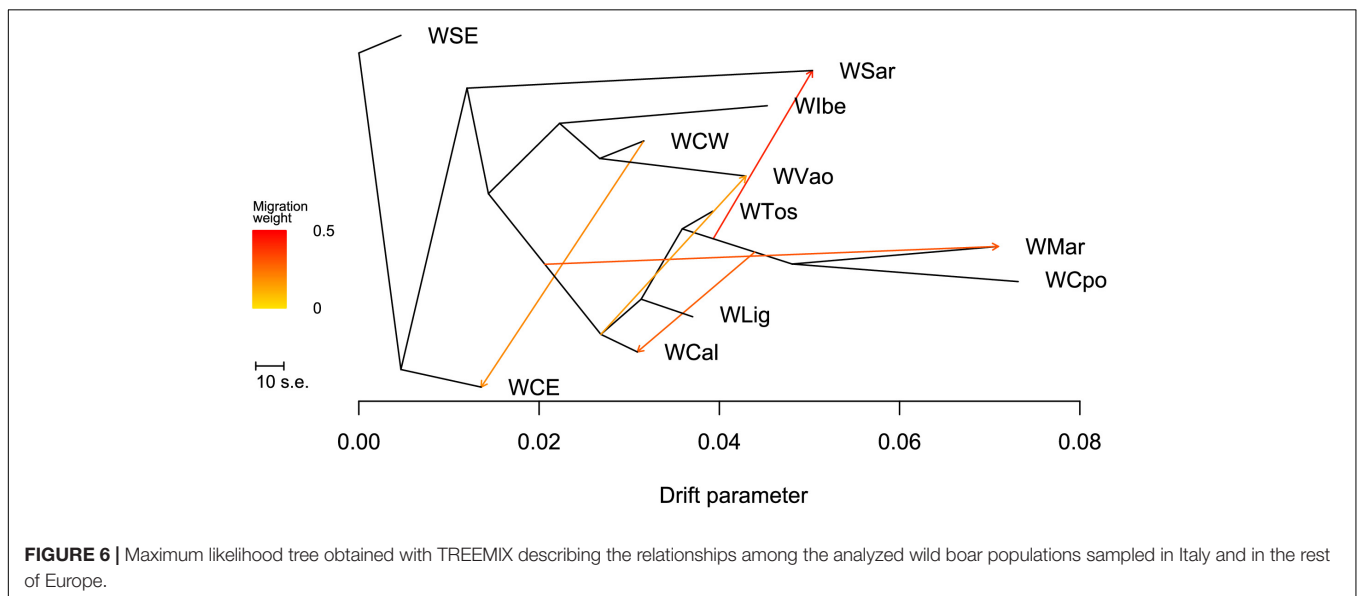
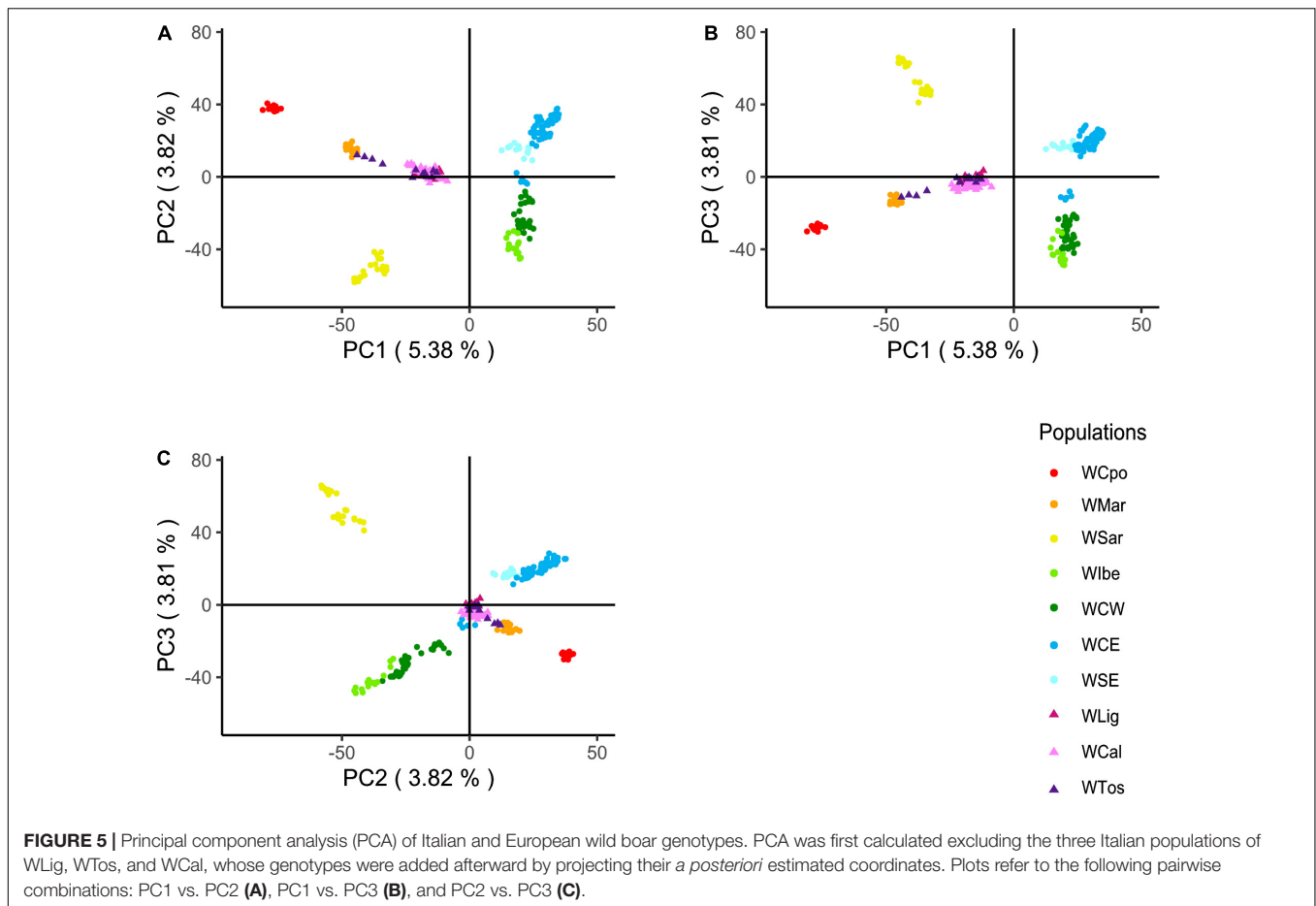
Genomic Diversity and Differentiation of Italian Wild Boar Populations

Our results on genome-wide heterozygosity levels in WB and DP were consistent with findings from other studies, based on the analyses of similar types and number of genomic markers (Goedbloed et al., 2013b; Alexandri et al., 2017; Iacolina et al., 2018), showing a considerable heterogeneity within and among groups.

However, the low levels of polymorphism we observed in Italian WB, even though comparable with genome-wide values from other European WB populations, should be treated with caution due to the possible ascertainment bias from the applied SNP array. The majority of the SNPs were indeed discovered in commercial pig breeds, and only a pooled set of 31 Central European WB was included in the discovery

panel (Ramos et al., 2009). Thus, polymorphisms of the Italian population are likely to be under-represented in the SNPs chip.

The most diverging population turned out to be the one living in the Castelporziano Presidential Estate (WCpo), which was deemed in the past the only pure indigenous nucleus of Italian WB remaining in the peninsula (Oliver et al., 1993). Such a pronounced divergence of this population can be the result of long-lasting isolation and limited effective size, which have likely shaped its allele frequencies by genetic drift. A similar interpretation can be given for WB living in the Maremma Regional Park (WMar), which also showed a certain level of divergence from the other Italian populations, yet lower than WCpo. Similar results had been already found by Vernesi et al. (2003), who included these two populations in their analyses based on microsatellite markers, detecting a certain level of differentiation that was interpreted as a possible expression of two native lineages. These populations were also included in previous phylogeographic analyses (Scandura et al., 2008; Vilaça et al., 2014), which detected a high proportion of endemic diversity both at autosomal microsatellite and mitochondrial D-loop regions. Specifically, Castelporziano samples showed exclusively mtDNA haplotypes belonging to the Italian lineage E2, which were also detected in museum specimens of Maremma WB (Larson et al., 2005). The observed divergence of WCpo and WMar might thus,



in principle, be determined also by a different level of introgression with respect to other (Apennine) populations. The artificial selection for “native morphotypes” that was carried out in the Maremma Natural Park in the past

might have contributed to the lower introgression of the individuals sampled in this area, while the isolation of WCpo should have prevented any genetic contamination (see **Supplementary Material**).

All analyses confirmed the high genomic differentiation of Sardinian WB from all mainland populations. This situation had been already described by Iacolina et al. (2016), who were also able to detect a major proximity with the mainland Italy populations by analyzing a larger sample of Sardinian specimens. This was attributed to a likely shared ancestry, but also at least in part to a recent genetic introgression from mainland populations, due to translocations for hunting purposes. Our results are also coherent with recent findings described by Petrelli et al. (2022) who investigated population genomic and selection patterns in WB living on the island.

The uniqueness of Castelporziano and Sardinian WB had been detected also in a study using microsatellite loci to assess the differentiation among European WB populations (Veličković et al., 2016). Both populations showed significant evidences of past bottlenecks, thus calling into question the main role of genetic drift in determining their differentiation.

The Aosta Valley population clearly showed a different make-up from the other Italian samples. It clustered with Central-Western European populations (WCW), which is consistent with an origin by range expansion from neighboring areas, as known for the Piedmont population (De Beaux and Festa, 1927; Monaco et al., 2007) and suggested by spatial information on the recolonization process in the region (Peracino and Bassano, 1995). This result, however, contrasts with the presumed release of captive hybrid stocks in the region (Apollonio et al., 2010b). Our genomic data thus suggest that natural processes (i.e., expansion from surrounding areas) have prevailed over human-mediated causes (i.e., releases).

An outstanding unexpected outcome of our study was represented by the mismatch between genetic and spatial distance we found. Wild boar from WTos, WLig, and WCal populations showed indeed a high similarity in all the analyses conducted, despite their geographical position. These populations are likely to represent the most manipulated WB populations in our sample. Their genomic composition could be affected by a number of human interventions (reintroductions, translocations) that have mixed up and spread an admixed gene pool, appearing as a separate cluster in ADMIXTURE analyses. This interpretation is confirmed by the central position of these three WB populations in the projected PCA (Figure 5). Within the WTos population, we found a different genetic composition between localities: a group of individuals sampled in the San Rossore estate, on the Tyrrhenian coast, slightly deviated from the other WTos individuals from eastern Tuscany and also from the nearby WLig population, showing similarities with individuals from WMar (Figures 2, 5). According to its history, this coastal nucleus has probably a higher proportion of native variation than the other individuals sampled in northern Tuscany (see Supplementary Material).

Genomic Introgression in Italian Wild Boar Populations

Despite human intervention has to be invoked to interpret the observed patterns of genomic variation in Italy, the detected

signals of admixture with DP and non-Italian WB in the sampled populations were lower than expected based on prior knowledge (Oliver et al., 1993; Apollonio et al., 2010b). Both ADMIXTURE and f_3 analyses did not show relevant signals of introgression from external sources into the Italian populations, contrasting with patterns emerging from other studies based on microsatellite and MC1R analyses (Vernesi et al., 2003; Scandura et al., 2008, 2011; Canu et al., 2016).

Despite close proximity of WTos, WLig, and WCal to European populations in the projected PCA, only weak signatures of introgression from other European macro-areas were detected by ADMIXTURE and f_3 analyses. Two possible explanations could be proposed for our findings: (1) introgression from multiple European WB and DP populations was relevant but it occurred many generations ago and was followed by a local divergence of the admixed gene pool from all parental populations; (2) releases of WB of foreign origin or WB $\times$ DP hybrids took place locally but involved a limited number of individuals, which were outnumbered by translocated individuals of Italian origin and/or by naturally expanding native populations (so non-native alleles were diluted by the native ones). In the former situation, the novel gene pool would appear as a different exclusive cluster in the ADMIXTURE analyses (see Figure 4A, $K = 12$), as it would not be possible to track back the contribution of the parental populations. In the second case, the prevailing native origin of the individuals would explain the observed lack of introgression; however, the origin by translocation from multiple areas in Italy would still generate an admixed and unique gene pool.

Historical information available for our sampling areas (see Supplementary Material) and results of f_3 analyses support the second line of interpretation, considering the expansion of remnant native populations sustained by human restocking with native boars, as main drivers of WB recovery in the peninsula. Both past and more recent translocations in fact, although partly undocumented, were carried out using free-ranging or captive animals from central Italy, that derived in large part from the Tyrrhenian populations (like Castelporziano estate or Maremma).

The low rate of introgression from DP detected in this study is consistent with previous findings. Using the same set of markers, no hybrid was indeed detected by Iacolina et al. (2018) in a subsample of 19 Italian WB (from Tuscany) among those considered in the present study, whereas 3 hybrids out of 25 individuals (12%) were detected in a random sample of Sardinian WB. Conversely, using a multiple-marker approach (coat color, MC1R, SNPs), (Petrelli et al., 2022) found a complete overlap among the genomes of Southern Italian known hybrids with those of the sympatric WB, likely as a result of deep introgression into the latter.

Interestingly, though in the present work signals of human-caused introgression were not expected for WSar, as the used Sardinian genotypes had been selected among the less introgressed individuals detected in a previous study focused on the insular population (Iacolina et al., 2016), f_3 analyses detected a migration from mainland

WB to Sardinia (**Figure 6**), thus supporting a recent gene flow.

Evidence of a recent introgression from mainland Italy, as well as from DPs, had been also detected by microsatellite analyses (Scandura et al., 2011), while reciprocal introgression with Sardinian DPs was proven by the analysis of allelic variation at nuclear genes (MC1R and NR6A1, Canu et al., 2016).

CONCLUSION

Despite the prevalent opinion among hunters and managers depicted the native Italian WB as almost genetically extinct, the present study reveals that Italian populations may still maintain a high proportion of native genomic diversity. The strong divergence that Italian populations showed from all other European WB and from DPs, as well as the occurrence of an exclusive mtDNA lineage, are clear indicators of a retained endemic variation.

Our results suggest that the impact of long-term natural processes (such as Quaternary climatic fluctuations, range expansions, natural selection), as well as the prolonged geographic isolation of Italy determined by the Alps and the Mediterranean Sea, might still affect the phylogeographic structure of a number of species, prevailing over the effects of human manipulations in shaping WB genetic diversity patterns. Similar results were recently found in a genomic study on managed populations of red-legged partridges (*Alectoris rufa*, Forcina et al., 2021).

The most preserved Italian WB populations appear to occur along the central Tyrrhenian coast of the peninsula (WCpo and WMar). The high genomic differentiation of these WB populations, together with their morphometric uniqueness (Apollonio et al., 1988; Tinelli et al., 1999), would call for a re-evaluation of the subspecies *Sus scrofa majori* (De Beaux and Festa, 1927), as recently already proposed for other highly differentiated Italian mammal populations, such as the Mesola red deer (*Cervus elaphus italicus*; Zachos et al., 2014), the Italian roe deer (*Capreolus capreolus italicus*; Randi et al., 2004), and the Apennine wolf (*Canis lupus italicus*; Montana et al., 2017).

However, the Italian WB populations we investigated, yet contributing to the overall genetic variation of the Italian WB, represent only part of it. Populations of north-eastern Italy, not included in the present study, showed a different composition from those living in central Italy, revealing higher similarities with eastern European populations (Scandura et al., 2008). Similarly, in southern Italy, a highly detectable rate of introgression from DPs was found in WB living north to WCal population (Cilento area), where a high morphological variation was also described (Fulgione et al., 2016; Petrelli et al., 2022). Unfortunately, no genetic nor phenotypical information is instead available for reintroduced WB that are spreading across Sicily, as well as for other areas of more recent recolonization.

It is therefore evident that the Italian WB population represents a strongly heterogeneous assemblage of demes with different histories and genomic make-up, determined by a combination of human-mediated factors such as reintroductions, translocations, restocking activities with farmed (sometimes hybrid) individuals, range fragmentation and demographic events, resulting in a genomic heterogeneity much higher than in any other European region.

Evidence from the present work clearly acknowledges the endemic value of Italian WB native genomic diversity. Thus our findings strongly recommend its conservation (a) by preventing further admixture among WB populations, strictly regulating WB farming and permanently banning wild boar import (as currently regulated due to sanitary reasons, i.e., African Swine Fever prevention), as well as (b) by minimizing domestic introgression by carefully preventing the interaction with pig breeds, especially when raised in a semi-natural state, and with released captive wild boar stocks, possibly deliberately introgressed by domestic swine.

Future studies based on whole genome sequencing could contribute to definitively clarifying the evolutionary histories and divergence timing of Italian WB populations through the analyses of demographic trajectories and past gene flow patterns with other European wild populations and domestic breeds. Paleogenomic and ancient DNA approaches could additionally help assess how much of the current diversity in WB local stocks can be ascribed to a native gene pool comparing modern and historical samples (i.e., museum specimens or stuffed trophies) dated before 1950, thus prior to most human-mediated gene flow.

Moreover whole-genome and transcriptome analyses could help identify further evidences of genetic differentiation and adaptive variation in Italian WB, which could not be detected by the Porcine SNP60 chip due to limitations in its development (not including Italian animals). Such additional genomic regions could allow us to investigate possible selection or adaptive patterns in Italian WB populations, potentially hosting genes responsible for their local adaptation to the Mediterranean environment, and, theoretically, for their high phenotypic divergence. Finally, the availability of entire genomes could also shed lights on the long-term evolutionary potential of Italian WB populations, their ability to actively face the ongoing climate changes and their future ecological role in the Anthropocene.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available. This data can be found here: doi: 10.5061/dryad.hmqnq9jm.

ETHICS STATEMENT

Ethical review and approval was not required for this study in accordance with the local legislation and institutional requirements.

AUTHOR CONTRIBUTIONS

MS, NM, and MA conceived the study. NM, CM, and LI contributed to sampling and lab analyses. GF, RC, FM, GP, and MS performed the data analyses. MS, GF, RC, FM, and NM wrote the manuscript. All authors contributed to the discussion of results, manuscript revision, read, and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fevo.2022.833081/full#supplementary-material>

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