Genomic Characterization of the Three Balkan Livestock Guardian Dogs

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Abstract: Balkan Livestock Guardian Dogs (LGD) were bred to help protect sheep flocks in sparsely populated, remote mountainous areas in the Balkans. The aim of this study was genomic characterization (107,403 autosomal SNPs) of the three LGD breeds from the Balkans (Karst Shepherd, Sharplanina Dog, and Tornjak). Our analyses were performed on 44 dogs representing three Balkan LGD breeds, as well as on 79 publicly available genotypes representing eight other LGD breeds, 70 individuals representing seven popular breeds, and 18 gray wolves. The results of multivariate, phylogenetic, clustering (STRUCTURE), and FST differentiation analyses showed that the three Balkan LGD breeds are genetically distinct populations. While the Sharplanina Dog and Tornjak are closely related to other LGD breeds, the Karst Shepherd is a slightly genetically distinct population with estimated influence from German Shepard (TreeMix analysis). Estimated genomic diversity was high with low inbreeding in Sharplanina Dog (Ho = 0.315, He = 0.315, and FROH>2Mb = 0.020) and Tornjak (Ho = 0.301, He = 0.301, and FROH>2Mb = 0.033) breeds. Low diversity and high inbreeding were estimated in Karst Shepherds (Ho = 0.241, He = 0.222, and FROH>2Mb = 0.087), indicating the need for proper diversity management. The obtained results will help in the conservation management of Balkan LGD dogs as an essential part of the specific grazing biocultural system and its sustainable maintenance.

Keywords: canine; livestock guardian dogs; conservation management; genomic diversity and characterization

1. Introduction

Livestock Guardian Dogs (LGDs, Canis familiaris) are a specialized type of dog breeds kept with grazing livestock as they are able to protect them from large predators, including gray wolves, brown bears, foxes, lynxes, and others. Together with the livestock, mostly sheep flocks, LGDs typically live under harsh mountain environments in the open and free-range grazing production systems. They are large-bodied dogs that are locally adapted and selected to demonstrate intelligence, trustworthiness, attentiveness, and protectiveness [1]. While most LGD breeds are common in Europe and Asia, they can be found all over the world. Today, LGDs are a very important component in the sustainable maintenance of the specific grazing biocultural systems. Despite several hypotheses that can be traced from the Tibetan Mastiff to the Molossers given to Alexander the Great by an Indian king, little is known about their precise origin. The oldest link between dogs and sheep can only be traced back to 5600 BP [2]. However, it would not be surprising if their functional role and interaction with humans date back to the beginnings of the Neolithization process.
Karst Shepherd (KRA), Sharplanina Dog (SAR), and Tornjak (TOR) are breeds recognized by the Fédération Cynologique Internationale (FCI), belonging to the LGD subgroup, geographically linked to the Balkans and considered molosser-like mountain dogs. Originally the Karst Sheperd, formerly also called the Illyrian Shepherd, was native to the Slovenian region of the Karst, and the Sharplanina Dog comes from Sharplanina Massif, while the name Tornjak comes from the Croatian word “tor”, which means a fenced area for keeping livestock in the mountainous part of Croatia and Bosnia and Herzegovina. The ancestors of KRA and SAR were listed as subtypes of the same breed, but in 1968 they were internationally recognized as separate breeds. By then the number of KRAs had dropped to the critical point, whereupon two Newfoundland males and some German Shepherds were added to the breed. SAR became a rare breed after World War II, probably mixed with several breeds, including Caucasian Shepherd dogs, but still representing a native population of herding dogs. In 2007, Tornjak, the Bosnian and Herzegovinian-Croatian Shepherd dog was provisionally recognized by the FCI. During the recognition process, the breed was listed in two subtypes (Bosnian-Herzegovinian subtype and Croatian subtype), a practice that was later abandoned. These breeds are large-framed, often around 70 cm at the withers and more than 45 kg body weight. They were formed to protect livestock from wild predators in the absence of a human shepherd, as shown in Figure 1.

The first comprehensive genetic variability was studied by Irion et al. [6], who used 100 autosomal microsatellite markers in the study of genotypic variation within and between globally representative samples of 28 breeds. As the authors noted, “While the set

Figure 1. Tornjak dogs guarding sheep on a mountain pasture. The photo was provided by Ljuban Gudelj and Ilija Smiljanić (Tomislavgrad, Bosnia & Herzegovina).
of autosomal microsatellites was useful in describing genetic variation within breeds, establishing the genetic relatedness between breeds was less conclusive. In addition, the use of microsatellites was limited because it was difficult to link genotypes obtained in different laboratories/machines. For this response, results from a number of studies focusing on small native breeds were underutilized. The genetic diversity of LGD by microsatellites has been analyzed in several studies. For example, genetic diversity and relatedness of Balkan LGD breeds were analyzed by Ceh and Dovc (2014) [9], Dimitrijević et al. (2020) [10] analyzed genetic diversity and population structure within SAR sampled in Serbia, while the genetic diversity and relatedness of Italian LGD breeds have been analyzed in several studies [11,12]. Recent advances in next-generation sequencing provide affordable access to high-throughput genotypic information and enable more comprehensive estimates of genetic diversity, population structure, genetic admixture, inbreeding levels, and effective population size at global scales [13,14]. At the same time, developments in the isolation and analysis of ancient DNA have led to a better understanding of the origin and domestication process [15] and the interrelationship between dogs and wolves [16]. In general, the genetic diversity of popular dog breeds (PDB) has been frequently analyzed [17–19], although there are several studies that have investigated the genomic diversity of local native breeds [20]. At the same time, the number of studies analyzing the genetic variability of LGDs is far from its matching their importance [9,11], especially those using high-throughput molecular information [21].

The aim of this study was to provide information relevant to the genomic characterization and conservation of the three Balkan LGD breeds (KRA, SAR, and TOR) by hypothesizing the following: (i) serious depletion of genetic diversity and high inbreeding is present in KRA, SAR and TOR; (ii) using genomic arrays, we can classify KRA, SAR, and TOR with other LGD breeds; (iii) KRA, SAR, and TOR are genetically pure breeds with little admixture with other LGD breeds; and (iv) we can detect genomic introgression of gray wolves within KRA, SAR, and TOR. To answer these hypotheses, we performed several high-resolution population genomic analyses, such as principal component discriminant analyses, phylogenetic network graph reconstruction, unsupervised clustering, estimation of migrations, and estimation of conservation genetics parameters (diversity and genomic inbreeding). In our analyses, we made the comparisons of Balkan LGDs with other eight LGD breeds, seven popular dog breeds, and a gray wolf as a wild reference population.

2. Materials and Methods

2.1. Sampling, DNA Extraction, Breed Origin, and Genomic Information

2.1.1. Sampling, DNA Extraction, and Genotyping

Overall, 52 dogs registered to the three Balkan LGD breeds (KRA, 14; SAR, 14; and TOR, 24) were sampled as buccal swabs or hair follicles at dog shows or were obtained from breeders and owners. Geographical origin and distribution, together with illustrated physical appearance (photo), of the sampled dogs, is shown in Figure 2. An additional four gray wolves representing wild reference populations from Croatia were provided (tissue) by Ana Galov and Haidi Arbanasić (the University of Zagreb, Faculty of Science). DNA from buccal swabs or hair follicles was extracted using the standard protocol of the DNeasy Blood & Tissue kit reagents (Qiagen, Germany). The quantity of DNA was evaluated using NanoVue (GE Healthcare Life Sciences, USA). All 56 DNA samples were genotyped using Illumina CanineHD BeadChip (Illumina Inc., San Diego, CA, USA) containing 172,115 SNPs. We performed quality control using SNP & Variation Suite v8.7.0 (Golden Helix, Inc., Bozeman, MT, www.goldenhelix.com), and after filtering (SNP call rate <95%) and excluding nonautosomal and unassigned markers, a total of 137,725 markers remained. We also excluded four KRA and four TOR with an individual call rate < 90%.
2.1.2. Publicly Available Genomic Information

For this study, we obtained publicly available SNP data [8,16], from which we selected a representative sample of LGD, PDB, and gray wolves (WOL). Publicly available SNP data were merged with the Balkan LGD dataset described above. The final dataset contained 107,403 autosomal SNPs that were common to both previously described datasets and 211 dogs representing 18 breeds with a population of 18 gray wolves (Table 1).

Table 1. Genomic diversity and inbreeding in the dog breeds and gray wolves (reference wild population).

| Breed (code)                     | Animals | Ho ± SE     | He ± SE     | Fis ± SE    | FROH±2Mb ± SE |
|---------------------------------|---------|-------------|-------------|-------------|---------------|
| Livestock guarding dogs (LGD)   | 123     | 0.290 ± 0.003 | 0.296 ± 0.003 | 0.015 ± 0.010 | 0.035 ± 0.003 |
| Anatolian Shepherd (ANA)        | 6       | 0.294 ± 0.018 | 0.329 ± 0.000 | 0.105 ± 0.056 | 0.026 ± 0.013 |
| Caucasian Shepherd (CAU)        | 5       | 0.278 ± 0.009 | 0.251 ± 0.000 | -0.107 ± 0.034 | 0.038 ± 0.004 |
| Fonni’s Dog (FON)               | 6       | 0.300 ± 0.018 | 0.319 ± 0.000 | 0.073 ± 0.056 | 0.025 ± 0.010 |
| Great Pyrenees (GPY)            | 13      | 0.277 ± 0.011 | 0.328 ± 0.000 | 0.157 ± 0.032 | 0.043 ± 0.007 |
| Karst Shepherd (KRA)            | 10      | 0.241 ± 0.003 | 0.222 ± 0.000 | -0.081 ± 0.014 | 0.087 ± 0.012 |
| Kuvazs (KUV)                    | 10      | 0.279 ± 0.007 | 0.288 ± 0.000 | 0.033 ± 0.024 | 0.045 ± 0.006 |
| Maremmano-Abruzzese Sheepdog (MAR) | 14  | 0.291 ± 0.006 | 0.296 ± 0.000 | 0.016 ± 0.022 | 0.036 ± 0.005 |
| Sharpalina Dog (SAR)            | 14      | 0.315 ± 0.005 | 0.315 ± 0.000 | 0.000 ± 0.017 | 0.020 ± 0.007 |
| Pastore della Sila (SIL)        | 14      | 0.302 ± 0.007 | 0.284 ± 0.000 | -0.064 ± 0.025 | 0.022 ± 0.004 |
| Tibetan Mastiff (TIB)           | 11      | 0.289 ± 0.005 | 0.300 ± 0.000 | 0.036 ± 0.017 | 0.018 ± 0.005 |
| Tornjak (TOR)                   | 20      | 0.301 ± 0.003 | 0.301 ± 0.000 | 0.000 ± 0.010 | 0.033 ± 0.005 |
| Popular dog breeds (PDB)        | 70      | 0.208 ± 0.004 | 0.220 ± 0.004 | 0.052 ± 0.010 | 0.105 ± 0.005 |
| Cavalier King Charles Spaniel (CKC) | 10   | 0.192 ± 0.004 | 0.199 ± 0.000 | 0.034 ± 0.019 | 0.119 ± 0.005 |
| Rough Collie (COL)              | 10      | 0.173 ± 0.005 | 0.181 ± 0.000 | 0.044 ± 0.028 | 0.140 ± 0.012 |
| Doberman Pinscher (DOB)         | 10      | 0.187 ± 0.004 | 0.193 ± 0.000 | 0.032 ± 0.023 | 0.120 ± 0.016 |
| German Shepherd Dog (GSD)       | 10      | 0.208 ± 0.005 | 0.221 ± 0.000 | 0.058 ± 0.023 | 0.118 ± 0.007 |
| Husky (HUS)                     | 10      | 0.023 ± 0.007 | 0.237 ± 0.001 | 0.061 ± 0.030 | 0.077 ± 0.009 |
| Labrador Retriever (LAB)        | 10      | 0.256 ± 0.006 | 0.277 ± 0.000 | 0.074 ± 0.020 | 0.065 ± 0.005 |
| Pekingese (PEK)                 | 10      | 0.215 ± 0.009 | 0.229 ± 0.000 | 0.061 ± 0.038 | 0.097 ± 0.008 |
2.2. Genomic Diversity, Inbreeding, and Molecular Coancestry

2.2.1. Genomic Diversity and Inbreeding

Heterozygosity (observed and expected) and genomic individual inbreeding coefficient \( (F_{IS}) \), were calculated in SNP & Variation Suite v8.7.0 (Golden Helix, Inc., Bozeman, MT, www.goldenhelix.com) using the “inbreeding coefficient” option on autosomal markers of the respective population. The \( F_{IS} \) indicates the deviation from Hardy-Weinberg heterozygosity equilibrium and can range from -1 to +1, indicating mating randomness. While negative values usually indicate outbreeding, positive values result from inbreeding [22].

2.2.2. Runs of Homozygosity Inbreeding

ROH segments were detected using SNP & Variation Suite v8.7.0 (Golden Helix, Inc., Bozeman, MT, www.goldenhelix.com (accessed on 19 February 2021)). According to the recommendations in Ferenčaković et al. (2013) [23], ROH segments with different length categories (from 2 to 5 Mb, from 5 to 10 Mb, and longer than 10 Mb) were identified based on different criteria. Thus, ROH segments were named in the category from 2 to 5 Mb if there were at least 15 consecutive homozygous SNPs covering more than 2 Mb, with one missing SNP allowed. In the 5 to 10 Mb category, two missing SNPs and one heterozygous SNP were allowed. For segments greater than 10 Mb, four missing and two heterozygous SNPs were allowed. Where more than one missing or heterozygous call was allowed, they were not allowed to be consecutive. The maximum gap between the two SNPs was set at 1 Mb, and the minimum required density was 1 SNP per 75 kb. The number of missing and heterozygous calls allowed was set following Ferenčaković et al. (2013) [23]. Next, the genomic inbreeding coefficient \( F_{ROH} \) was calculated following McQuillan et al. (2008) [24] and Curik et al. (2014) [25], where \( F_{ROH} \) is defined as genome length in ROH/autosomal genome length covered by the SNP chip (here 2,200 Mb distributed over 38 chromosomes). Inbreeding coefficients with different ROH lengths (\( F_{ROH>2Mb} \), \( F_{ROH2-5Mb} \), \( F_{ROH5-10Mb} \), and \( F_{ROH>10Mb} \)) were calculated as they correspond to a different base population and allow decomposition of the \( F_{ROH>2Mb} \) into its remote \( (F_{ROH2-5Mb}) \), intermediate \( (F_{ROH5-10Mb}) \), or recent \( (F_{ROH<10Mb}) \) origin. Boxplots and stacked-bar-plots were created using the ggplot2 package for R software [26].

2.2.3. Molecular Coancestry

Molecular (marker-by-marker) coancestry coefficients were estimated for each population using the “Relationship Matrix” option in JMP Genomics and IBS statement (JMP® Genomics, Version 9.1. SAS Institute Inc., Cary, NC, 1989-2019). Pairwise molecular coancestry coefficients, all unique (lower) off-diagonal elements from the coancestry matrix, were further analyzed within each breed/population, with mean values presenting the probability that two randomly sampled alleles, each from one individual, are the same copy of an allele. Note that molecular coancestry refers to alleles that are identical-by-state, in contrast to genealogical coancestry coefficients which refer to identical-by-descent alleles inherited from the same ancestor [27].

2.3. Population Structure and Gene Flow

2.3.1. Discriminant Analysis of Principal Components

We performed a discriminant analysis of principal components (DAPC) to identify and visualize clusters of genetically related individuals [28]. All calculations and visualizations required for DAPC were performed in the R package adegenet [29].
2.3.2. Phylogenetic Analyses and Gene Flow

Inference of the phylogenetic relationship between 18 dog breeds and WOL was represented as the NeighborNet network inferred from pairwise Nei’s genetic distances [30]. Nei’s distances were calculated using the R package StAMPP R [31]. A NeighborNet network was created and represented using SplitsTree5 software [32].

In addition, patterns of splits and mixtures of a subset of 18 dog breeds, and the gray wolf population as a rooted outgroup, were detected by the maximum-likelihood algorithm implemented in the TreeMix program [33]. First, we constructed a maximum-likelihood tree of the populations with no migration events considered. We then constructed a phylogenetic network for all selected populations, sequentially increasing migration events up to 12 migrations. Residuals from fitting the model to the data were visualized using the R script implemented in TreeMix.

2.3.3. Unsupervised Clustering with STRUCTURE Analyses

Population genetic structure was inferred by the algorithm implemented in the STRUCTURE program [34,35]. We had to reduce the number of SNP genotypes in the dataset to 22,220 to enable the complex calculations that would not otherwise have been possible with our computational resources. The reduction of the dataset (pruning) was based on the threshold of linkage disequilibrium between SNPs, here defined as the pairwise $r^2$, which was equal to 0.1, with the window size equal to 100 SNPs and a step size equal to 20. A model assuming admixture and correlated allele frequencies was used for all STRUCTURE runs, with a burn-in of $10^5$ iterations followed by $10^4$ MCMC iterations. Runs were repeated ten times for each assumed K, starting with $K = 1$ to $K = 24$. The resulting individual genotype membership coefficients were displayed using DISTRUCT [36].

To gain additional insight into population structure and quantify genetic differentiation among the analyzed populations, we also estimated pairwise Wright’s $F_{ST}$ fixation indices [37] using the R package StAMPP [31].

3. Results

3.1. Genetic Diversity, Inbreeding, and Molecular Coancestry

3.1.1. Genomic Diversity and Inbreeding

The estimated observed (Ho) and expected (He) heterozygosity, $F_{IS}$, and $F_{ROH>2Mb}$ means in 18 dog breeds and gray wolves are shown in Table 1. Compared to PDB and gray wolves, significantly higher heterozygosity (Ho and He) and significantly lower $F_{ROH>2Mb}$ were observed in LGD breeds. For example, the estimated 95% confidence intervals for Ho ranged from 0.284 to 0.296 for LGD, from 0.200 to 0.216 for PDB, and from 0.155 to 0.196 for WOL. Because the confidence intervals did not overlap, the estimated Ho were significantly higher in LGD than in PDB and/or WOL. Considerably lower $F_{IS}$, although the difference was not significant, was observed in LGD than in PDB. The values observed in KRA, with exception of $F_{IS}$, were within the PDB estimates (see Table 1). Thus, in LGD breeds with exception of KRA, estimates of observed (expected) heterozygosity ranged from 0.277 in Great Pyrenees (GPY) to 0.315 in SAR (from 0.251 in Caucasian Shepherd (CAU) to 0.329 in Anatolian Shepherd (ANA)) while $F_{ROH>2Mb}$ ranged from 0.018 in Tibetan Mastiff (TIB) to 0.045 in Kuvaszs (KUV). Interestingly, almost the lowest negative $F_{IS}$ value ($-0.081$) was observed in KRA, indicating strong avoidance of inbreeding among recent genealogical relatives. The contrast between high $F_{ROH>2Mb}$ and low $F_{IS}$ suggests that low diversity and relatively high ROH inbreeding in KRA resulted from small population size and genetic drift. Very high heterozygosity and low ROH inbreeding, even among LGD, were observed in SAR and TOR, indicating importance for genetic diversity. Zero $F_{IS}$ values observed in these two breeds show complete randomness in mating which could be improved by pedigree management and mating avoidance of genealogical relatives. At the same time, the lowest heterozygosity and the highest inbreeding values ($F_{IS}$) were observed in WOL.
Ho is observed heterozygosity, He is expected heterozygosity (gene diversity), F<sub>s</sub> is Wright’s inbreeding coefficient, and F<sub>RHOH>2Mb</sub> is ROH inbreeding coefficient with the assumption that 2Mb long ROHs are autozygous, SE is the standard error.

3.1.2. Runs of Homozygosity, Inbreeding, and Molecular Coancestry

The distribution of F<sub>RHOH>2Mb</sub> and its partitioning with respect to the origin of autozygosity (remote, intermediate, and recent) is shown in Figure 3. While LGD breeds have a much lower inbreeding in several breeds (ANA, KUV, Maremmano-Abruzzese Sheepdog (MAR), and Pastore della Sila (SIL)), outlining dogs with much higher inbreeding has been observed, a similar pattern was also observed in PGD breeds (Husky (HUS) and Pekingese (PEK)), see Figure 3a. This points to the lack of pedigree inbreeding management or intentional mating with respect to the desired ancestor.

Assuming that the expected length of an IBD haplotype segregating in a population follows an exponential distribution, we can infer the number of generations from the inbred individual to the common ancestor. It is generally assumed that if 1 cM is approximated by the length of 1 Mb, we would expect ROH that are 16.6, 10.0, or 5.0 Mb in length to be derived from a common ancestor 3 generations back (6 meioses), 5 generations back (10 meioses), or 10 generations back (20 meioses), respectively. Further details can be found in Curik et al. (2014) [25]. Thus, with the exception of KRA, which exhibits the PDB pattern, the differences in the degree of inbreeding between LGD and PDB are mainly the result of distant and intermediate inbreeding, originating from 25 to 10 (F<sub>RHOH>2Mb</sub>) and from 10 to 5 (F<sub>RHOH>10Mb</sub>) generations distant ancestors, respectively (Figure 3b).

Mean molecular coancestry in a breed is functionally related to expected heterozygosity, which is also defined as the probability that two randomly sampled alleles are not identical-by-state. It is evident that much higher molecular coancestry was observed in
PDB and WOL compared to LGD (Table 2). However, we also analyzed the range of all pairwise molecular coancestry coefficients and found that the mean range (the difference between the minimum and maximum molecular coancestry averaged over LGD or PDB) in LGD was 0.088 compared to 0.043 in PDB. By comparing the mean range of molecular coancestry, we were able to analyze the genetic heterogeneity (or compactness) of each breed. WOL had a fairly large molecular coancestry range (0.172), but this was a sampling effect as gray wolves were sampled from several different populations.

3.2. Population Structure and Gene Flow
Discriminant Analysis of Principal Components and Phylogenetic Analyses with Gene Flow

Presentation of the DAPC analysis, variation in five principal components of the DAPC method performed on 14 dog breeds and a gray wolf population, is shown in Figure 4. We have omitted Cavalier King Charles Spaniel (CKC), Rough Collie (COL), Doberman Pinscher (DOP), and PEK from this illustration (Figure 4) as these four breeds are distant from the LGD breeds and did not help to clarify their genetic relationship. From the first two principal components of DAPC (Figure 4a), it was clear that LGDs form a distinct group of dogs, although a slight separation was observed for KRA, while KUV was quite distinct. The first two principal components of DAPC explained 40.1% of the variation, while most of the variation remained in the third, fourth, fifth, and further principal components of DAPC (Figure 4b,c,d,e).

However, the NeighborNet network inferred from pairwise Nei’s genetic distances was able to complement the DAPC approach in presenting genetic relationships between the analyzed breeds and gray wolves. According to the NeighborNet network (Figure 5) CAU, SAR, TOR, and KUV form a closely related cluster within the main reticulation. TIB slightly separated towards WOL. MAR and SIL are clustered on a separate reticulation, in a similar way as GPY and Fonni’s Dog (FON). This clustering pattern can be explained to some extent by the geographical origin of LGD. KUV, SAR, and TOR are geographically very close to each other, while CAU is further away but in the same direction on the continent. However, the genetic relatedness of KUV, SAR, and TOR with CAU could be explained by human migrations, but this aspect is not considered in our analysis. The clustering of MAR and SIL is logical since both are Italian LGD breeds. Since there is a historical link between Sardinia and Spain, we were not surprised that FON and GPY share the same cluster. In contrast, ANA was separated from other LGD breeds while KRA was clustered with German Shepherd Dog (GSD) indicating the documented influence of GSD on the breeding history of KRA.
Figure 4. Graphical illustrations of the population genetic structure of 14 dog breeds and gray wolves, generated by the “supervised” DAPC algorithm: (a) scatter-plot presenting a variation of the first two principal components of the DAPC; (b) distribution of the third principal components of the DAPC; (c) distribution of the fourth principal components of the DAPC; (d) distribution of the fifth principal components of the DAPC and (e) Barr plot visualization of the contribution of discriminant functions explaining sample variation. The breeds shown are Anatolian Shepherd (ANA), Caucasian Shepherd (CAU), Fonni’s Dog (FON), Great Pyrenees (GPY), Karst Shepherd (KRA), Kuvazs (KUV), Maremmano-Abruzzese Sheepdog (MAR), Sharplanina Dog (SAR), Pastore della Sila (SIL), Tibetan Mastiff (TIB), and Tornjak (TOR) as livestock guarding dogs, German Shepherd Dog (GSD), Husky (HUS), Labrador Retriever (LAB) as popular dog breeds, and gray wolves (WOL) as a reference wild population.
Figure 5. NeighborNet network inferred from pairwise Nei’s genetic distances, estimated from the CanineHD BeadChip information, among in 10 livestock guarding dog breeds, 8 popular dog breeds, and gray wolves (wild reference population). Reticulations on the graph indicate past hybridization events among populations. The breeds shown are Anatolian Shepherd (ANA), Caucasian Shepherd (CAU), Fonni’s Dog (FON), Great Pyrenees (GPY), Karst Shepherd (KRA), Kuvazs (KUV), Maremmano-Abruzzese Sheepdog (MAR), Sharplanina Dog (SAR), Pastore della Sila (SIL), Tibetan Mastiff (TIB), and Tornjak (TOR) as livestock guarding dogs, Cavalier King Charles Spaniel (CKC), Rough Collie (COL), Doberman Pinscher (DOB), German Shepherd Dog (GSD), Husky (HUS), Labrador Retriever (LAB), and Pekingese (PEK) as popular dog breeds, and gray wolf (WOL) as a reference wild population.

Genetic relatedness and migration patterns, allowing 12 migration events (that best fit the model), were observed using TreeMix analysis (Figure 6). While some differences from the NeighborNet network analysis were observed, much of the relatedness was logical and in agreement with other analyses. For example, KRA was clustered slightly differently, but the migration influence of GSD and the root of LGD breeds was clearly identified (Figure 5). Interestingly, the migration pattern of TOR and KUV root towards CAU may explain why CAU was clustered with KUV, SAR, and TOR in NeighborNet network analysis (Figure 5). Analyses implemented in the TreeMix algorithm revealed no migration flow between dog breeds and gray wolves.
Figure 6. TreeMix graph visualizing the phylogenetic relationship among 18 dog breeds and gray wolf as a root, in a model allowing 12 migration events. Migration arrows are colored according to their weights, while branch lengths are proportional to the estimated genetic drift. The breeds shown are Anatolian Shepherd (ANA), Caucasian Shepherd (CAU), Fonni’s Dog (FON), Great Pyrenees (GPY), Karst Shepherd (KRA), Kuvazs (KUV), Maremmano-Abruzzese Sheepdog (MAR), Sharplanina Dog (SAR), Pastore della Sila (SIL), Tibetan Mastiff (TIB), and Tornjak (TOR) as livestock guarding dogs, Cavalier King Charles Spaniel (CKC), Rough Collie (COL), Doberman Pinscher (DOB), German Shepherd Dog (GSD), Husky (HUS), Labrador Retriever (LAB), and Pekingese (PEK) as popular dog breeds and gray wolves (WOL) as a reference wild population.

3.3. Population Structure and Admixture

We estimated population structure and admixture by the algorithm implemented in the STRUCTURE software. The STRUCTURE can reveal “hidden structure” without a priori assigning individual breed or population membership. According to recommendations [34,35], the most likely K is the one where Ln Pr(G|K) is maximized or where the smallest value of K captures the main structure in the data. In our analyses, it was challenging to decide which was the optimal number of clusters in the dataset (the most likely parameter K), as the Ln Pr(G|K) values increased continuously with the number of clusters (Ks) used in the model (Figure S1A in the Supplement). We further analyzed the rate of change in the Ln Pr(G|K) between successive K values as suggested by Evanno et al. (2005) [38]. In the presence of a hierarchical island population structure, this approach could reveal a more subtle structure within populations. In our dataset, see Figure S1B in the Supplement, several peaks were observed at K = 4, K = 11, K = 16, and K = 19, indicating the presence of a complex hierarchical structure. Therefore, we decided to present from the STRUCTURE analyses (Figure 7), only the results that are in concordance with the observed divergent change in \( \Delta K \) [38].
Figure 7. Graphical presentation of the Bayesian unsupervised clustering of individual 193 dogs and 18 gray wolves performed with the STRUCTURE algorithm. The selection of the meaningful and relevant clusters (K = 19, K = 16, K = 11, and K = 4) is concordant with the procedure suggested by Evanno et al. (2005) [38]. The breeds shown are Anatolian Shepherd (ANA), Caucasian Shepherd (CAU), Fornis’s Dog (FON), Great Pyrenees (GPY), Karst Shepherd (KRA), Kuvasz (KUV), Maremmano-Abruzzese Sheepdog (MAR), Sharplanina Dog (SAR), Pastore della Sila (SIL), Tibetan Mastiff (TIB), and Tornjak (TOR) as livestock guarding dogs (*), Cavalier King Charles Spaniel (CKC), Rough Collie (COL), Doberman Pinscher (DOB), German Shepherd Dog (GSD), Husky (HUS), Labrador Retriever (LAB), and Pekingese (PEK) as popular dog breeds, and gray wolves (WOL) as a reference wild population.

At K = 19, the STRUCTURE algorithm distinguished three Balkan LGD breeds, although considerable admixture was observed within SAR and TOR. Notable admixture patterns were observed in a majority of LGD breeds (GPY, FON, KUV, MAR, SAR, SIL, and TOR). The presence of admixture patterns in LGD breeds is undoubtedly consistent with the explanations for the estimated higher heterozygosity and lower ROH inbreeding values, especially in those of intermediate and remote ancestry. Gene exchange, in both directions, was observed in all STRUCTURE results presented but was most pronounced in K = 4. Very strong introgression of dog genes was observed in some gray wolves, while the migration signal was not specific to a particular dog breed (Figure 7). We also recorded the introgression of gray wolf genes into some dog breeds. The strongest WOL introgression was observed in HUS and TIB, weak in SAR and ANA, and barely visible in GPY, PEK, and TOR.

Using pairwise Wright’s coefficients, we also estimated population differentiation between the analyzed breeds and the WOL, as well as average genetic relatedness to other dog breeds in the dataset (Table 2). According to the mean $F_{ST}$ values across all other dog breeds, SAR ($MF_{ST} = 0.13$) and TOR ($MF_{ST} = 0.15$) were in the group of LGD breeds (GPY, $MF_{ST} = 0.11$; ANA, $MF_{ST} = 0.12$, and FON, $MF_{ST} = 0.14$) with the lowest mean $F_{ST}$ values, occupying a central position among the analyzed dogs. In contrast, KRA ($MF_{ST} = 0.27$) was the most pronounced among the other LGD breeds with $MF_{ST}$ values ranging from 0.11 to 0.23, which could be the consequence of the strong influence of GSD ($MF_{ST} = 0.27$) migrations.
Table 2. Population differentiation among 18 dog breeds and gray wolves (wild reference population) based on genome-wide $F_{ST}$ estimates.

| Breed     | KRA* | SAR* | TOR* | ANA* | CAU* | COL | FON* | GPY* | GSD | KUV* | MAR | SIL* | TIB* | CKC | DOB | HUS | LAB | PEK | MFst |
|-----------|------|------|------|------|------|-----|------|------|-----|------|-----|------|------|-----|-----|-----|-----|-----|------|
| KRA*      | 0.27 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.27 |
| SAR*      | 0.18 | 0.07 |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.13 |
| TOR*      | 0.20 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.15 |
| ANA*      | 0.19 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.12 |
| CAU*      | 0.30 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.23 |
| COL       | 0.40 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.33 |
| FON*      | 0.20 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.33 |
| GPY       | 0.17 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.14 |
| GSD       | 0.32 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.14 |
| KUV*      | 0.23 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.17 |
| MAR*      | 0.21 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.17 |
| SIL*      | 0.23 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.18 |
| TIB*      | 0.23 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.17 |
| CKC       | 0.37 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.30 |
| DOB       | 0.38 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.31 |
| HUS       | 0.33 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.26 |
| LAB       | 0.24 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.19 |
| PEK       | 0.33 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.26 |
| WOL       | 0.35 |      |      |      |      |     |      |      |     |      |     |      |      |     |     |     |     |     | 0.28 |

Genomic $F_{ST}$ estimates [37]. MFst is mean overall pairwise comparisons with all other dog breeds. The breeds shown are: Anatolian Shepherd (ANA), Caucasian Shepherd (CAU), Fonni’s Dog (FON), Great Pyrenees (GPY), Karst Shepherd (KRA), Kuvasz (KUV), Maremmano-Abruzzese Sheepdog (MAR), Sharplanina Dog (SAR), Pastore della Sila (SIL), Tibetan Mastiff (TIB), and Tornjak (TOR) as livestock guarding dogs (*), Cavalier King Charles Spaniel (CKC), Rough Collie (COL), Doberman Pinscher (DOB), German Shepherd Dog (GSD), Husky (HUS), Labrador Retriever (LAB), and Pekingese (PEK) as popular dog breeds, and gray wolves (WOL) as a reference wild population.
4. Discussion

Although they do not achieve the popularity of some other dog breeds, LGD breeds have specific functional roles and behavioral abilities that were historically selected by humans during the domestication process. Today, their ability to protect sheep from large wild predators, usually wolves, ensures the livelihood of people in remote rural areas with harsh environmental conditions. Thus, LGD breeds, together with sheep, are important participants in the sustainable maintenance of specific biocultural grazing systems [39].

Three well-known Balkan LGD breeds that are used in mountainous areas of Bosnia and Herzegovina (TOR), Croatia (TOR), North Macedonia (SAR), Serbia (SAR), and Slovenia (KRA) were analyzed for the first time using high-throughput genomic information. The availability and use of high-density genome-wide arrays and the development of some population genomic methods, e.g., estimation of ROH-based inbreeding, enabled a more precise estimation of genetic conservation parameters compared to microsatellite-based studies [9,10]. More so, the open data sharing policy (samples were taken from Parker et al. (2017) [8] and Talenti et al. (2018) [21]) allowed comparison, with the same methodology used, of the targeted breeds (KRA, SAR, and TOR) with representative samples of LGDs and other popular dog breeds, as well as with gray wolves.

Sample sizes (from 5 to 20 in TOR) were very small for some breeds (ANA, CAU, and FON), but still comparable to other similar analyses [8,13,21]. However, we stand on the conclusions made because in many methods used (DAPC, NeighborNet, TreeMix analyses, STRUCTURE) the high number of informative SNPs can compensate for the small sample size since the mosaicism of the segregating segments is high. For example, looking back 5 or 10 generations, each individual inherits haplotypes from 32 or 1024 parents, respectively. When we have presented estimates related to genetic diversity and the degree of inbreeding, we have provided standard errors to allow the calculation of confidence intervals. For example, the same order of genetic diversity, SAR (highest), TOR (second highest), and KAR (lowest), was estimated in Ceh and Dovc (2014) [9] in 471 dogs (KRA = 326, SAR = 109, and TOR = 36) using 18 microsatellite loci.

On the other hand, we were not confident in estimating effective population size or selection signatures because these estimates are very sensitive to sample size [40]. A small sample size may be the reason for not detecting a subtle subdivision of the population. Dimitrijević et al. (2020) [10] detected the presence of two clusters when analyzing 94 SAR dogs based on 10 nuclear microsatellites. Therefore, it could be possible that we only sampled individuals from a single SAR cluster.

In our analysis, SAR dogs, like the majority of LGD breeds, showed some degree of admixture that was not detected by Dimitrijević et al. (2020) [10] using microsatellites. A large proportion of SAR dogs was assigned to the CAU breed in Ceh and Dovc (2014) [9]. The same authors (Ceh and Dovc, 2014 [9]) were able to show the presence of two subpopulations within the breed TOR when they performed a separate STRUCTURE analysis only for TOR individuals. We could not detect this subtle substructure since we did not perform a separate STRUCTURE analysis. TOR dogs are bred in two countries separated by the border (non-EU member Bosnia and Herzegovina versus EU member Croatia), which is a strong factor for modern breed differentiation (a good example in Druml et al. 2007 [41]). Therefore, we would not be surprised if the small level of subtle population structure is present in TOR. However, we do not believe that supporting breeding to separate populations is the sensible option, as these are dogs kept under strong natural selection pressure, and reduced genetic diversity would certainly reduce their ability to respond to natural selection. Identification of adaptive genes in Balkan LGDs is desirable but requires a large sample size.

Dreger et al. (2016) [42] found that the FON is present in the genetic profiles of other Mediterranean breeds and that the breed originated from the interbreeding of sighthound and mastiff breeds. It was further speculated that the relationship of FON with breeds
from the eastern (Komondor) and southern (Saluki) Mediterranean parallels human migration to Sardinia [14,21,43]. We observed that the predominant cluster present in GPY and FON is also present in varying degrees of admixture in KUV, MAR, SAR, SIL, and TOR. It would be interesting to investigate whether the common origin present within LGD breeds is related to migrations in the Neolithization process. For example, the existence of hunting dogs similar to FON dates back to the Bronze Age [43]. Although this is speculation, it may be worthwhile to conduct further research on a large dataset, supplemented by additional archaeological evidence and evidence observed in analyses of ancient human and sheep DNA.

5. Conclusions

Using high-throughput genomic information and a series of analyses, we were able to answer all four hypotheses posed in this study and reached the following conclusions: (i) higher genetic diversity and low inbreeding, following the pattern observed in other LGD breeds, was observed in SAR (95% confidence interval; for Ho from 0.305 to 0.325, for He from 0.313 to 0.317, for Fs from −0.033 to 0.033, and for FROH>2Mb from 0.006 to 0.034) and TOR (95% confidence interval; for Ho from 0.295 to 0.307, for He from 0.299 to 0.302, for Fs from −0.020 to 0.020, and for FROH>2Mb from 0.023 to 0.043). In contrast, much lower genetic diversity and higher inbreeding, following the pattern observed in PDB, was observed in KRA (95% confidence interval; for Ho from 0.235 to 0.247, for He from 0.220 to 0.224, for Fs from −0.108 to −0.054, and for FROH>2Mb from 0.063 to 0.111). (ii) Population genetic structure analyses (multivariate, phylogenetic, and unsupervised clustering) showed that SAR and TOR are closely related to other LGD breeds, whereas KRA is a slightly genetically divergent population with estimated influence from GSD. (iii) Analysis of STRUCTURE revealed the presence of admixture in SAR and TOR, which seems to be characteristic of most LGD breeds, while genetic purity was observed in KRA. (iv) Traces of admixture from dogs to gray wolves and from gray wolves to dogs were most pronounced at K = 4 in the STRUCTURE analysis. The strongest introgression of gray wolf genes was observed in HUS and TIM, while signals were much lower in SAR and ANA and negligible in GPY, PEK, and TOR.

Supplementary Materials: The following are available online at www.mdpi.com/2071-1050/13/4/2289/s1, Figure S1. The choice of the representative number of clusters (K): (A) Plot of Ln Pr(G|K) values presented as a function of the number of clusters [34,35], 10 runs presented. (B) Plot of ΔKs presented as a function of the number of clusters [38].

Author Contributions: V.C.-C., I.C., and P.D. conceived and designed the study; M.J., V.C.-C., and P.D. organized sample collection, extraction, and genotyping; M.J., M.Z., and M.F. performed the analyses with the supervision of I.C.; M.J., I.C., and V.C.-C. wrote the manuscript; V.C.-C. and P.D. funding acquisition. All authors have read and agreed to the published version of the manuscript.

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