



# Domestication and Genetic Diversity Analysis of Several Sheep Breeds in China

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## ABSTRACT

Sheep is the main livestock species in northern China. Under diverse climatic conditions, a gene pool of local sheep resources is now forming. There are many sheep breeds in northern China, including local breeds and hybrid breeds. However, little is known about the genetic background and genetic variation of local sheep populations in northern China. Therefore, our aim is to provide a characterization of the population structure and genetic relationship within sheep breeds in northern China and to study its genetic relationship with other sheep breeds. We collected blood samples from 390 sheep for genetic diversity analysis using reduced-representation genome sequencing (RRGS). This study used bioinformatics methods to analyze linkage disequilibrium, population structure, evolutionary tree, selective sweep *etc.* The mixed model revealed in this study may be formed in the process of sheep breeding, partially overlapping with historical and archaeological findings. New information on the domestication and genetic variety of sheep is provided by this study, which will help sheep in the future be improved through genomics.

**Key words:** Domestication, Genetic diversity, Sheep, Whole-genome resequencing.

This study uses whole genome sequencing (WGS) to re-sequence different individuals or tissues of sheep with known genomic information to discover differences between individuals. Through this method, genome-wide SNP and InDel molecular markers were developed, and SNP/InDel markers were used for population genetics research.

## Animals and genome sequencing

Two sheep populations-NKY and THT - were analyzed in this study. A total of 5 mL jugular vein blood was collected from 390 individuals (185 NKY and 205 THT population) in 2022 at Bayannaoer Academy of Agriculture and Animal Husbandry Sciences, Inner Mongolia, China. After the sample's genomic DNA was qualified, the DNA sequence was fragmented by ultrasound to form a random fragment.

## Bioinformatic analysis workflow

After the Illumina NovaSeq<sup>TM</sup> sequencing data (Raw Data) is offline, the quality control of the offline data is performed, and the low-quality data is filtered to obtain high-quality data (Clean Data). Clean Data was aligned to the reference genome sequence using BWA software (Li and Durbin 2009) to obtain the sequence location attribution (*i.e.*, BAM file). BAM files were corrected using GATK's Best Practices process (McKenna *et al.* 2010) and SNP and Small InDel markers were detected. The functional annotation information of SNP and InDel was obtained by using SNPEff software (Cingolani *et al.*, 2012) and gene prediction information of reference genome. Based on the obtained SNP and InDel molecular markers, genetic diversity, population structure, linkage disequilibrium and selective sweep were further studied (Table 1 for

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bioinformatics analysis tools).

## Linkage disequilibrium

In a population, the frequency of simultaneous inheritance of two genes at different loci is significantly higher than the expected random frequency, which is called linkage disequilibrium. The minimum genetic unit of species can be obtained by linkage disequilibrium analysis.

The linkage of SNP in all samples was analyzed by SNP combination on the same chromosome. Natural populations are represented by a linkage disequilibrium coefficient ( $r^2$ ). The closer  $r^2$  is to 1, the stronger the linkage. The distance between SNPs in the genome was fitted to  $r^2$ . Generally, the closer the distance between SNPs, the larger

the  $r^2$ , and the farther the distance between SNPs, the smaller the  $r^2$ . Generally, the distance between the corresponding sites when  $r^2$  decays to half is used as the value of LD-decay. The longer the LD-decay, the slower the LD decay, the greater the probability of linkage between the SNP of the representative species. The shorter the LD-decay, the faster the LD decay, the smaller the probability of linkage between the SNP of the representative species. It is generally believed that the species with fast LD decay are relatively primitive. The LD-decay of each subpopulation is shown in Fig 1.

Genome-wide linkage disequilibrium (LD) research showed that the NKY and THT populations had nearly the

same level of LD and decay, with a lower  $R^2$  correlation coefficient becoming stable at a distance of about 100 kb.

### Population structure

Principal component analysis (PCA) is a pure mathematical operation method, which can select a small number of important variables through linear transformation of multiple related variables. Based on SNP, principal component analysis (PCA) was performed by GCTA software to obtain the principal component clustering of 36 samples. Through PCA analysis, we can know which samples are relatively close and which samples are relatively distant, which can assist evolutionary analysis.

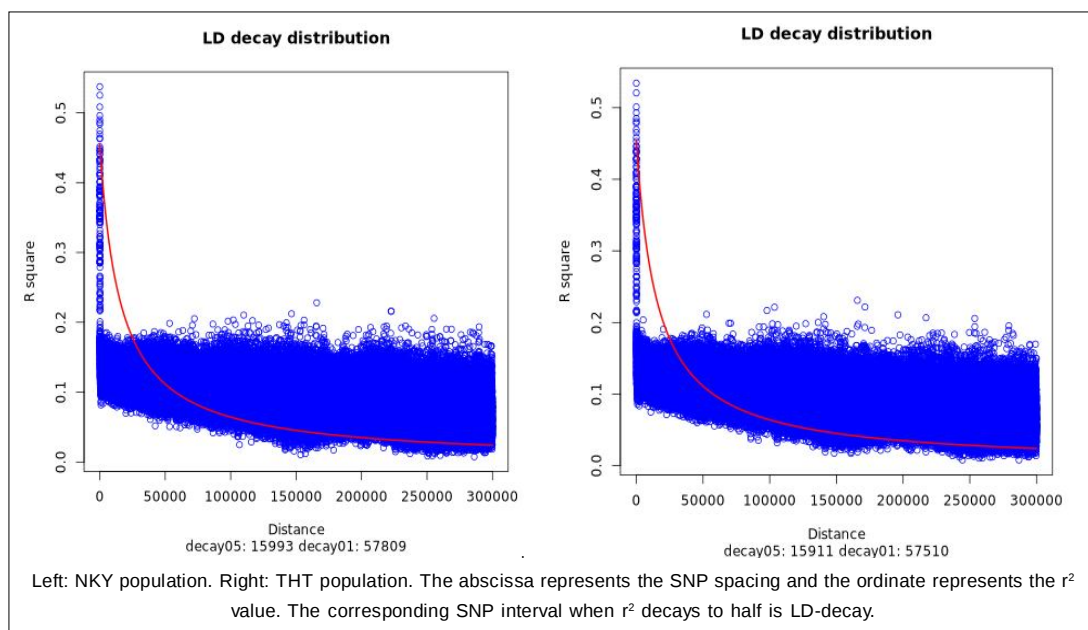
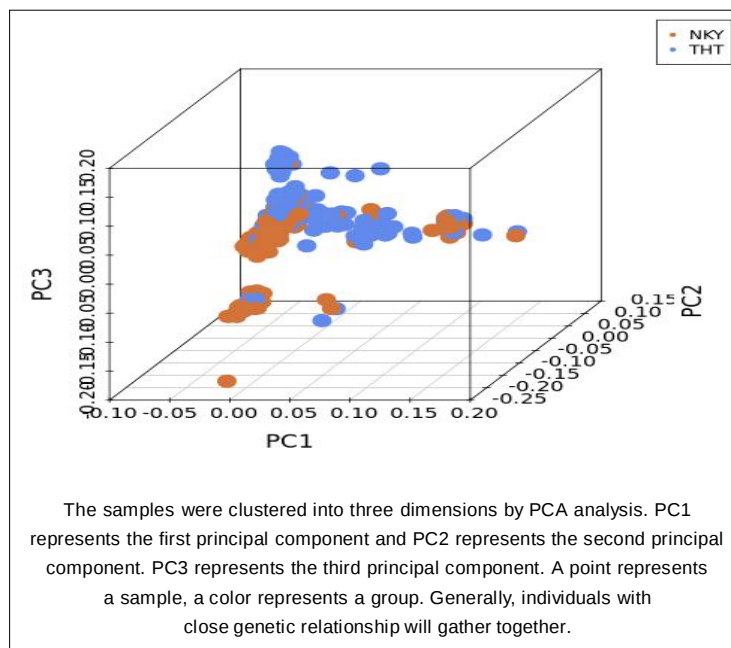


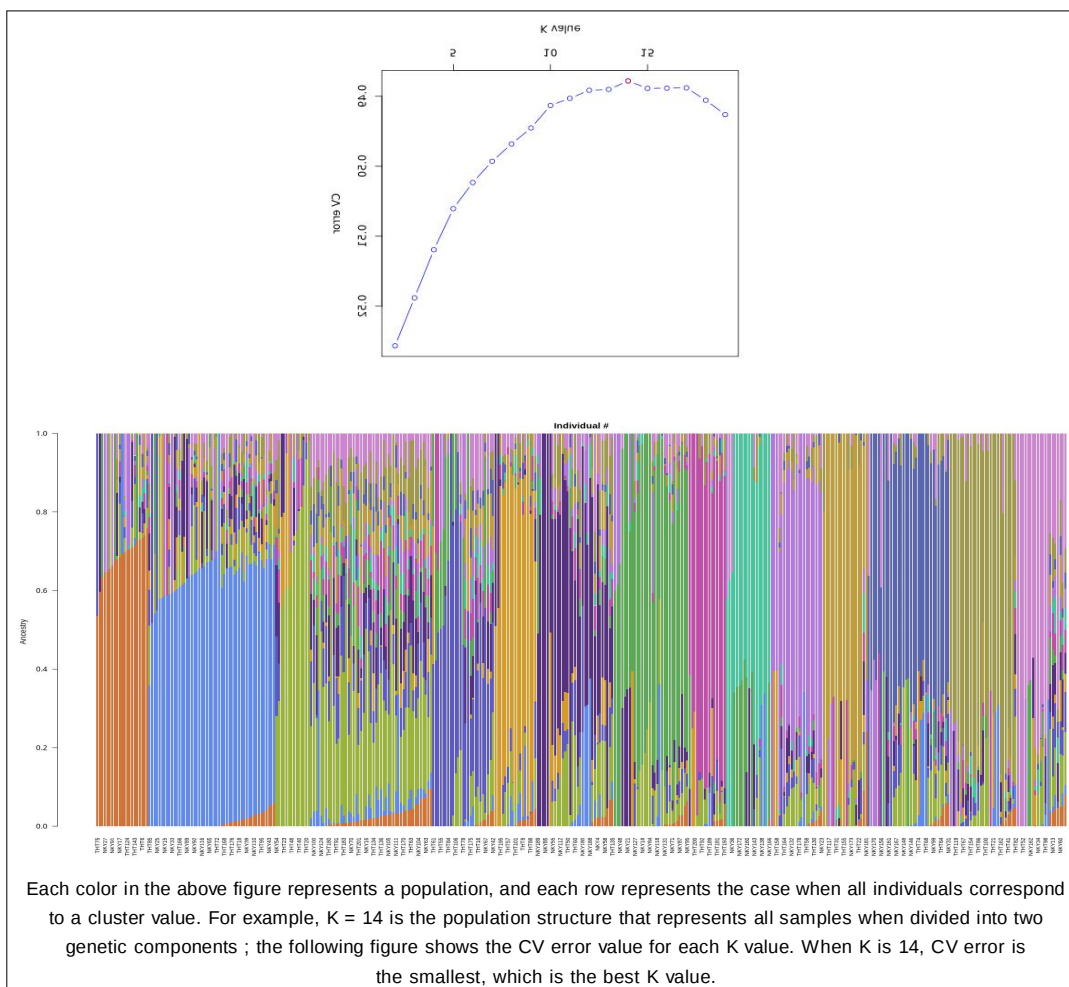
Fig 1: LD of the two populations.

Table 1: Bioinformatics analysis tools.

| Tool                   | Version              | Purpose                         | Availability                                                                                                                                                                                                                          |
|------------------------|----------------------|---------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Fastp                  | 0.23.0               | Raw data quality control        | <a href="https://github.com/OpenGene/fastp">https://github.com/OpenGene/fastp</a>                                                                                                                                                     |
| BWA                    | 0.7.17-r1198-dirty   | Mapping to reference genome     | <a href="http://bio-bwa.sourceforge.net/">http://bio-bwa.sourceforge.net/</a>                                                                                                                                                         |
| GATK/SAMtools/sentieon | 4.2.4.1/1.14/2021.12 | SNP, InDel detection            | <a href="https://software.broadinstitute.org/gatk/http://www.htslib.org/">https://software.broadinstitute.org/gatk/http://www.htslib.org/</a> <a href="https://www.sentieon.com/products/">https://www.sentieon.com/products/</a>     |
| SnpEff                 | 5.0e                 | Variation functional annotation | <a href="http://snpeff.sourceforge.net/">http://snpeff.sourceforge.net/</a>                                                                                                                                                           |
| Blast                  | 2.12.0+              | Gene functions annotation       | <a href="https://blast.ncbi.nlm.nih.gov/Blast.cgi">https://blast.ncbi.nlm.nih.gov/Blast.cgi</a>                                                                                                                                       |
| RAXML/Fast tree        | v8.2/2.1.11          | Evolutionary tree analysis      | <a href="https://cme.h-its.org/exelixis/web/software/raxml/index.html">https://cme.h-its.org/exelixis/web/software/raxml/index.html</a> <a href="http://www.microbesonline.org/fasttree/">http://www.microbesonline.org/fasttree/</a> |
| Admixture              | 1.3.0                | Population structure analysis   | <a href="http://software.genetics.ucla.edu/admixture/">http://software.genetics.ucla.edu/admixture/</a>                                                                                                                               |
| GCTA                   | 1.93.2               | PCA                             | <a href="http://cnsgenomics.com/software/gcta/#Overview">http://cnsgenomics.com/software/gcta/#Overview</a>                                                                                                                           |
| Vcftools               | v0.1.16              | selective sweep                 | <a href="https://vcftools.github.io/">https://vcftools.github.io/</a>                                                                                                                                                                 |
| PopLDdecay             | v3.41                | LD-decay                        | <a href="https://github.com/BGI-shenzhen/PopLDdecay">https://github.com/BGI-shenzhen/PopLDdecay</a>                                                                                                                                   |



**Fig 2:** PCA clustering diagram of samples.



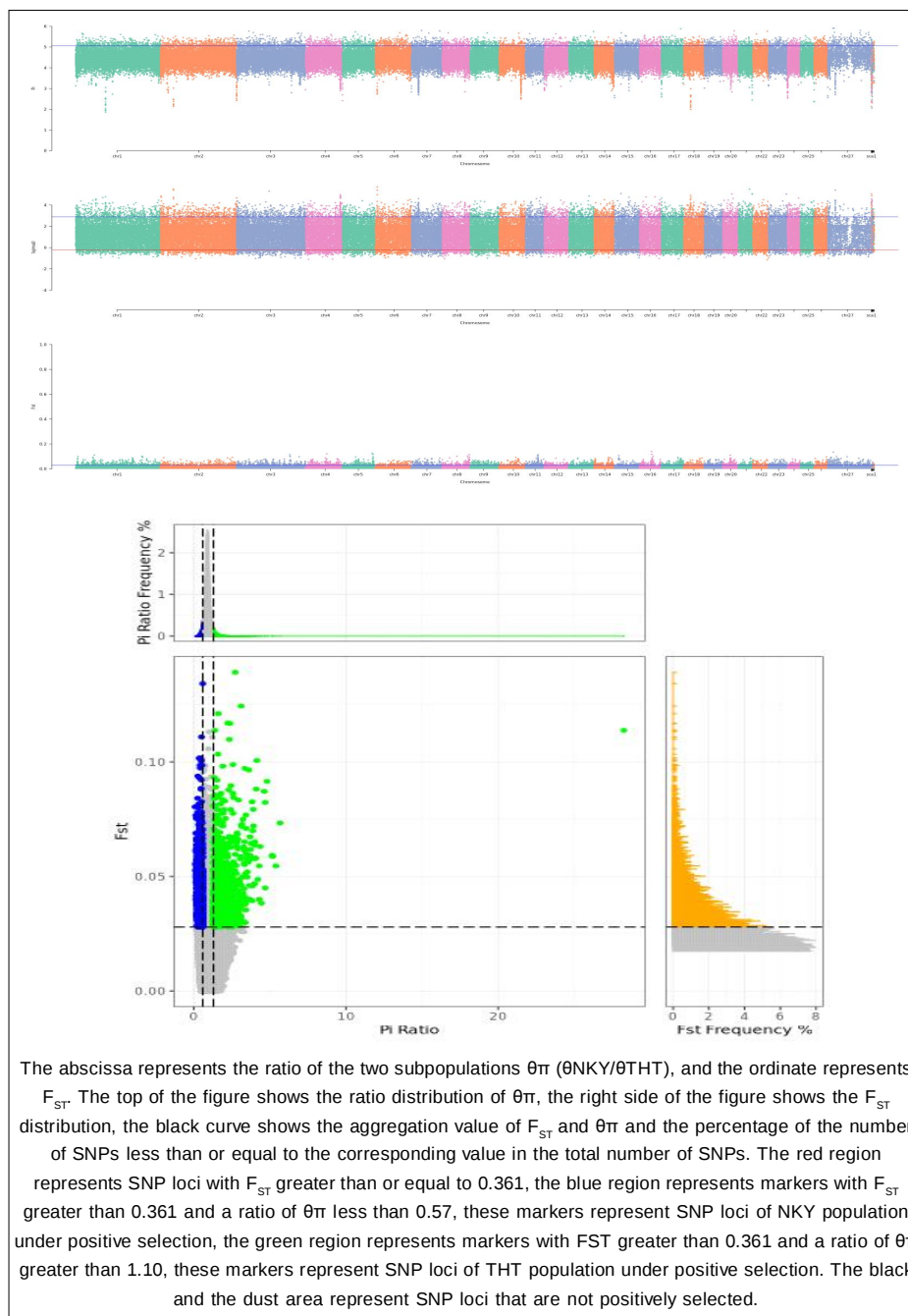
**Fig 3:** Population genetic structure based on whole genomes.

The results of the principal component analysis that was carried out on the genomic relatedness matrix between individuals are illustrated in Fig 2.

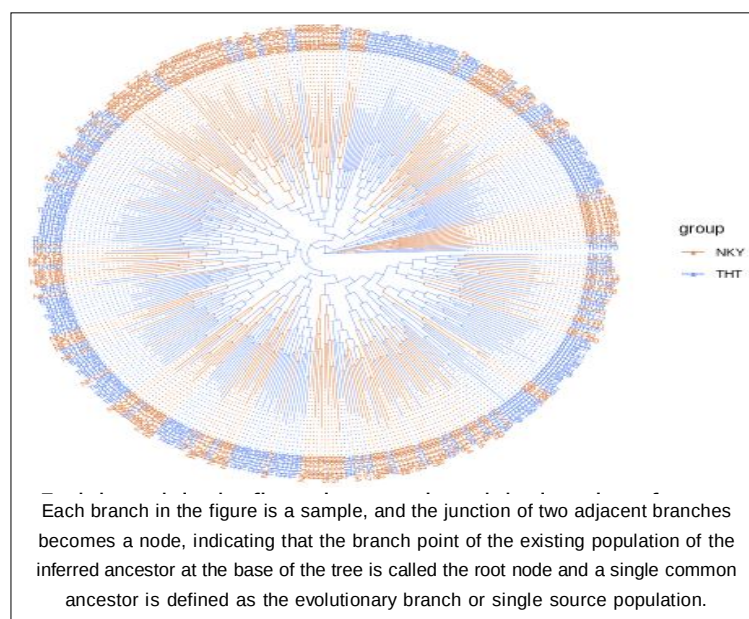
The population structure of the samples was analyzed. The number of clusters (K value) was assumed to be 1-20. The optimal number of clusters was determined to be 14 according to the K value corresponding to the lowest point of CV error (Cross validation error). It reflects that all our samples may come from 14 original ancestors. The population structure with K value of 14 is shown in Fig 3.

Through whole genome resequencing, it can be determined that there are regions with higher  $F_{ST}$  values and lower  $\pi/\theta$  values on the genome. The q chromosome results are shown in Fig 4.

$F_{ST}$  is an important index to measure the degree of genetic differentiation between populations. If an allele in a population experiences adaptive selection due to its high fitness to a specific environment, the increase of its frequency will increase the level of population differentiation, which is reflected in the higher  $F_{ST}$  value in F statistics.



**Fig 4:** Selective sweep analysis (Manhattan plot).



**Fig 5:** Phylogenetic tree.

The  $\pi/\theta$  represents the nucleotide polymorphism within the population, and the selective sweep will cause the selected genomic region to show that the polymorphism within the population is significantly lower than the average level of the genome, which corresponds to a lower  $\pi/\theta$  value.

The evolutionary tree is used to represent the evolutionary relationship between species. According to the genetic relationship between various organisms, various organisms are placed on a branched tree-like figure to concisely represent the evolutionary history and genetic relationship of organisms. The phylogenetic tree is shown in Fig 5.

Before human-mediated specialization for wool and milk began some 4000-5000 years ago, sheep were initially raised for access to meat. One of the earliest morphological modifications that likely occurred alongside domestication and is now a trait shared by many modern breeds is the removal of horns, which has recently been demonstrated to be one of those regions of the genome that contain strong evidence for accelerated change in response to artificial selection (Mohamadipoor *et al.*, 2021). Additionally, sheep's other genomic areas that are being selected for comprise genes that regulate body size, reproduction and color. The separation of animals into breeds, followed by the discovery of better rams and their disproportionate genetic contribution through artificial insemination, has increased the rate of genetic gain for productive attributes during the previous few hundred years. Numerous genetic exchanges have taken place during the evolution of contemporary breeds, as evidenced by the high haplotype sharing and relatively short divergence times across breeds (Missohou *et al.* 2022). A population size of more than 300 has been maintained by around 75% of contemporary sheep breeds. There are between 850 and

1409 different breeds. But several sheep breeds, such as the Shetland, Soay and Herdwick breeds, that were first chosen for their superior performance in a particular, sometimes remote, geographic location are now regarded as uncommon. It has become difficult to maintain genetic variety by preserving these historic breeds as generalist-type sheep have mostly taken over intensive sheep farming (Alberto *et al.*, 2018).

## CONCLUSION

Domestication is complex and entails multiple and parallel events and population admixtures. There are many breeds of sheep in China that have complex genetic exchanges. Native sheep breeds are abundant in China and some of them have unique characteristics including procreation, good lamb pelts, numerous horns, fat tails and great stress tolerance. The focus of the research has been on all of these. However, there is still debate regarding the origins of sheep breeds and there have been few thorough investigations on the genetic diversity of local sheep in China. Their population dynamics and genetic variety were subsequently discovered, which was beneficial for our comprehension and preservation of these genetic resources. The current research, which for the first time examined the genome-wide diversity and population structure of sheep breeds with a Northern Chinese origin, advanced knowledge of the intricate genetic relationships and anticipated shared ancestry of the many sheep breeds in China.

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### Author contribution

Bujun Mei designed the study and analyzed the RNA-seq data and drafted the manuscript. ZH, YP and Rong Liu joined the research and improved the manuscript.

### Competing interest

The authors declare that there are no competing interests associated with the manuscript.

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