The aim of this study was to locate SSTR5 polymorphisms and evaluate their association with growth traits in Hulun Buir sheep. The study followed up 884 Hulun Buir sheep from birth to 16 months of age, which were born in the same pasture and the same year, and a consistent grazing management strategy was maintained. The birth weight (BRW) was recorded at birth, and body weight (BW), body height (BH), body length (BL), chest circumference (ChC), chest depth (ChD), chest width (ChW), hip width (HW), and cannon circumference (CaC) were measured at 4 and 9 months of age. BW, BH, BL, ChD, HW, and CaC were also recorded at 16 months of age. Based on the growth traits, 233 sheep were selected as experimental animals. Sanger sequencing was performed, and seven single-nucleotide polymorphisms (SNPs) were identified. Association analyses of the SNPs and the growth traits were then conducted. Seven SNPs of the SSTR5 exhibited moderate polymorphism (0.25 < PIC < 0.5) and were consistent with the Hardy–Weinberg equilibrium. SNP7 (T989C, rs601836309) caused a change in amino acid sequences, while others did not cause any change. The genotypes of SNP1 (C186T, rs400914340) were significantly associated with BW, ChW, and ChC at 4 months of age and with HW at 9 months of age (p < 0.05). These genotypes also showed extremely significant association with CaC at 4 months of age (p < 0.01). The genotypes of SNP7 exhibited a significant association with ChW and CaC at 4 and 9 months of age, respectively. Moreover, the genotypes of SNP3 (T384C, rs601836309) and SNP4 (T537C, rs605867745) were significantly associated with CaC at 9 months of age (p < 0.05). Linkage disequilibrium was observed among the seven SNPs with five haplotypes. However, these haplotypes were not associated with growth traits at different ages. In conclusion, SNP1, SNP3, SNP4, and SNP7 may serve as molecular markers for the growth traits of Hulun Buir sheep.

Keywords: SSTR5, association, growth traits, Hulun Buir sheep, haplotypes
1 INTRODUCTION

Hulun Buir sheep is an esteemed local mutton breed in Hulun Buir, Inner Mongolia, China. This breed exhibits outstanding stress resistance, strong adaptability, stable heredity and provides high-quality, low-fat meat with a variety of amino acids. As a traditional mutton sheep breed, Hulun Buir sheep are not selected via advanced breeding methods; therefore, the breed exhibits low productivity, a slow growth rate, and a low slaughter rate. Many candidate genes have been reported to regulate metabolism and control the growth rate of domestic animals (Al-Mamun et al., 2015; Wang et al., 2015; La et al., 2019). Genetic variations in the candidate genes have been widely used as molecular markers, accelerating the breeding process and improving productivity. For example, new breeds of beef cattle with a myostatin mutation receptor subtype 5 (SSTR5) was identified as a differentially expressed gene. In the present study, we explored whether this gene would affect the growth traits of Hulun Buir sheep.

As a somatostatin (SST) receptor, SSTR5 has functions inseparable from those of SST. SST, also known as growth hormone (GH)-inhibiting hormone or somatotropin release-inhibiting factor, is considered as a hypothalamic factor that inhibits the secretion of GH (Brazeau et al., 1973). In mammals, there are five somatostatin receptor subtypes (SSTR5). SST and SSTR are widely distributed in the central nervous system, pancreas, intestines, stomach, kidney, liver, pancreas, lungs, and placenta and has a variety of biological functions (Finley et al., 1981; Maecke and Reubi, 2011; Quan et al., 2020). SST employs diverse mechanisms to regulate growth, but its activity depends on the binding of G-protein-coupled somatostatin receptors (Anzola et al., 2019). In addition to exerting an inhibitory effect on GH release (Lulle et al., 2006), SSTR also represses the secretion of prolactin, thyroid-stimulating hormone (TSH) (Cattaneo et al., 1996; Melmed, 2003). SST and SSTR5 are expressed in the islets of Langerhans and has a variety of biological functions (Strowski et al., 2000).

The nucleotide sequence of SSTR5 is highly conserved among species; ovine SSTR5 shares 85% sequence homology with humans and rats and 87% homology with mice. The ovine SSTR5 transcript has four exons and three introns (ENSOARG00000014478.1) located on chromosome 24 (GenBank, Gene ID: 443,210), encoding 1,044-bp base (rs 812,728–813866) and 347-amino acid residues. As a receptor for somatostatin, SSTR5 plays an important role in many physiological processes, such as GH release, cell anti-proliferation, and regulating a variety of signal transduction pathways (Cattaneo et al., 1996; Melmed, 2003). SSTR5 is also one of the major SSTRs expressed in the islets of Langerhans and plays an essential role in mediating the inhibitory effect of SST on insulin expression, secretion, and cell proliferation (Fagan et al., 1998).

Based on our previous experiments, and considering the importance of SSTR5 in controlling the growth hormone axis and the lack of research on the effect of the SSTR5 gene on growth traits in sheep, we conducted a molecular characterization of SSTR5, identified polymorphisms, and analyzed associations between different genotypes and growth traits in Hulun Buir sheep. The current study generated novel information about the genetic resources of Hulun Buir sheep, serving as a foundation for future applications of molecular markers in breeding.

2 MATERIALS AND METHODS

2.1 Experimental Animals and Growth Trait Data Acquisition

The study included 884 Hulun Buir sheep, the progeny of unrelated rams, born in the same pasture in March 2019 at the Hulun Buir Sheep Breeding Farm in Hulun Buir, Inner Mongolia, China. The growth traits of each sheep were recorded from birth to adulthood (16 months of age). During the experiment, all sheep were allowed to graze freely on natural pasture and had free access to water.

At birth, only birth weight (BRW) was recorded. Body weight (BW), body length (BL), body height (BH), chest circumference (ChC), chest depth (ChD), chest width (ChW), hip width (HW), and cannon circumference (CaC) were recorded at 16 months of age. Size and weight were measured using a veterinary measuring tape and a sensitive platform balance, respectively (Zhang et al., 2016).

Based on the growth traits, a total of 233 healthy, disease-free sheep were selected as experimental animals, comprising 119 sheep with the fastest growth rate and 114 sheep with the slowest growth rate (124 females and 109 males). There were significant differences in growth traits between the two extreme populations (Supplementary Table S1). All animal experiments were conducted following the procedures described in the “Guidelines for animal care and use” manually approved by the Animal Care and Use Committee, Northwest Institute of Plateau Biology, Chinese Academy of Sciences (NWIPB2020302, 13 April 2020).

| Primer name | Primer sequences (5’-3’) | Size (bp) | Tm (°C) |
|-------------|-------------------------|----------|---------|
| E1-2        | F: CCGCGCTGCTGTCGCTGTC  | 761      | 60      |
|             | R: TACAGCCAACCAGATGACCAG  |          |         |
| E3-4        | F: TGAAACACTGCAGAACCTCAG  | 759      | 60      |
|             | R: GCTCTCCGTCTTGCTGTCGACG  |          |         |
### 2.2 Primer Design and Sequencing

A 0.5-cm² ear tissue sample was collected from each sheep at birth for DNA extraction and preserved in 75% alcohol. DNA was purified using a DNA extraction kit (Tiangen, Beijing, China), and the quality was evaluated by running samples on a gel. Primers were designed for all exons of the SSTR5 using Primer3 v0.4.0 (1) (Koressaar and Remm, 2007). The SSTR5 transcript has four exons (ENSOARG00000014478.1); one primer pair was designed to cover exons 1 to 2 and another to cover exons 3 and 4. Information about primers is presented in Table 1.

PCR amplifications were performed in a 30 μl reaction volume consisting of 1.0 μl of DNA, 15 μl of 2×Taq PCR Master Mix (Sangon, Shanghai, China), 1.0 μl of each primer, and double-distilled water (dH₂O) to make up the volume. Amplifications were performed using Bio-Rad S1000 thermal cyclers (Bio-Rad, Hercules, CA, United States). The thermal profile was as follows: initial denaturation at 94°C for 2 min, followed by 35 cycles at 94°C for 10 s (denaturation), 60°C for 30 s (annealing), 72°C for 60 s (elongation), with a final extension step at 72°C for 5 min. The PCR products were visualized using 1% agarose gel electrophoresis to determine amplicon quality and quantity. The sequencing was performed using Sanger sequencing (Agilent 3,730, United States). Sequence alignment and SNP identification were conducted via MEGA (version 5.0) (Electronics Ltd., Kuopio, Finland). DNAMAN software (version 5.2.10) (Lynnon BioSoft, Vaudreuil, Canada) was used to conduct sequence analyses.

### 2.3 Bioinformatics Analysis of Non-Synonymous Mutations

Protein analyses were conducted with ExPASy tools (http://expasy.org/tools/), and parameters including molecular weight, isoelectric point, instability index, aliphatic index, and grand average of hydropathicity were computed. SignalP 4.0 (http://www.cbs.dtu.dk/services/SignalP/) was used to predict the presence of signal peptides. NetOGlyc 3.1 (http://www.cbs.dtu.dk/services/NetOGlyc/) and NetNGlyc 1.0 (http://www.cbs.dtu.dk/services/NetNGlyc/) were used to predict potential O– and N–glycosylation sites, respectively. NetPhos2.0 (http://www.cbs.dtu.dk/services/NetPhos/) was used to predict phosphorylation sites.

### 2.4 Population Genetic Analyses

Population genetic indices including allele frequency, heterozygosity (He), observed heterozygosity (Ho), effective allele numbers (Ne), and the polymorphism information content (PIC) were analyzed as previously reported (Nei and Roychoudhury, 1974). Genotypes of SNPs were tested for the Hardy–Weinberg equilibrium (HWE) (Ortega et al., 2016). Linkage disequilibrium (LD) and haplotypes analysis were conducted using Haploview (v.4.2) (Barrett et al., 2005).

### 2.5 Statistical Analysis

Measured traits were tested for normality by using the Shapiro–Wilk test in SPSS Statistics (V.19, IBM, Armonk, NY, United States). Pearson’s correlation coefficients were calculated to determine the correlation among the following measured traits at 4, 9, and 16 months of age: BW, BL, BH, ChC, ChD, ChW, HW, and CaC. SPSS was used to perform all analyses, and values are expressed as mean ± standard error. General linear mixed models were established to examine the associations between the genotypes and individual growth traits, and statistical significance was defined at p < 0.05. In this model, genotype and gender were fixed factors, and their interaction was tested. If an interaction between genotype and gender was identified, the following statistical model was used:

### Table 2 | Correlations between growth traits of Hulun Buir sheep a.

|      | BW       | BL       | BH       | ChW      | ChD      | ChC      | HW       |
|------|----------|----------|----------|----------|----------|----------|----------|
| 4 months of age |          |          |          |          |          |          |          |
| BW   | 0.782**  |          |          | 0.607**  | 0.507**  | 0.507**  |          |
| BL   | 0.303**  | 0.512**  |          | 0.368**  | 0.404**  | 0.404**  |          |
| ChW  | 0.368**  | 0.724**  | 0.698**  |          |          |          |          |
| ChD  | 0.507**  | 0.724**  | 0.756**  |          |          |          |          |
| ChC  | 0.507**  | 0.724**  | 0.756**  | 0.698**  |          |          |          |
| HW   | 0.368**  | 0.724**  | 0.756**  | 0.698**  | 0.671**  |          |          |
| CaC  | 0.447**  | 0.388**  | 0.331**  | 0.411**  | 0.495**  | 0.348**  | 0.456**  |
| 9 months of age |          |          |          |          |          |          |          |
| BW   | 0.631**  |          |          | 0.588**  | 0.431**  |          |          |
| BL   | 0.631**  | 0.588**  |          | 0.431**  | 0.605**  |          |          |
| ChW  | 0.507**  | 0.724**  |          |          |          |          |          |
| ChD  | 0.507**  | 0.724**  |          |          |          |          |          |
| ChC  | 0.507**  | 0.724**  |          |          |          |          |          |
| HW   | 0.561**  | 0.678**  | 0.602**  | 0.468**  |          |          |          |
| CaC  | 0.697**  | 0.545**  | 0.642**  | 0.602**  | 0.526**  | 0.565**  | 0.425**  |
| 16 months of age |          |          |          |          |          |          |          |
| BW   | 0.479**  |          |          |          |          |          |          |
| BL   | 0.184*   | 0.346**  |          |          |          |          |          |
| ChW  | 0.345**  | 0.173*   | 0.363**  |          |          |          |          |
| ChD  | 0.386**  | 0.381**  | 0.581**  | 0.532**  |          |          |          |
| HW   | 0.498**  | 0.046    | −0.424** | 0.310**  |          |          |          |
| CaC  | 0.498**  | 0.046    | −0.424** | 0.310**  |          |          |          |

BW= body weight; BL= body length; BH= body height; ChW = chest width; ChD = chest depth; ChC = chest circumference; HW= hip width; CaC = cannon circumference.

Correlations with |r| > 0.7 are in bold, *p < 0.05, **p < 0.01.

aData represent means ± SEM (n = 233).
\[ Y = \mu + \text{Genotype} + \text{Gender} + \text{Combination} + \epsilon, \]
where \( Y \) is the trait measured for each animal (BW, BL, BH, ChW, ChD, ChW, HW, and CaC), \( \mu \) is the mean value of \( Y \), Genotype is the genotype effect, Gender is the gender effect, Combination is the combined effect of the gender and genotype, and \( \epsilon \) is the random error, assumed to be independent and normally distributed; \( N(0, \sigma^2) \).

If no interaction between genotype and gender was identified, the following statistical model was used:
\[ Y = \mu + \text{Genotype} + \epsilon, \]
where \( Y \) is the trait measured for each animal (BW, BL, BH, ChW, ChD, ChW, HW, and CaC), \( \mu \) is the mean value of \( Y \), Genotype is the genotype effect, and \( \epsilon \) is the random error, assumed to be independent and normally distributed; \( N(0, \sigma^2) \).

### 3 RESULTS

#### 3.1 Correlations Between Growth Traits

All data conformed to a normal distribution, indicating suitability for subsequent analyses. At 4 months of age, BW exhibited strong correlations \((|r| > 0.7)\) with BL, ChW, and ChD; BL exhibited strong correlations with ChW and ChD; BH exhibited a strong correlation with ChD; ChD exhibited a strong correlation with HW. At 9 months, BW exhibited strong correlations with BL, BH, ChC, and HW; BL exhibited a strong correlation with BH; BH exhibited a strong correlation with ChC; and ChC exhibited a strong correlation with HW. At 16 months, BW exhibited strong correlations with BL, BH, ChC, and HW; BL exhibited a strong correlation with BH; BH exhibited a strong correlation with ChC; and ChC exhibited a strong correlation with HW. At 16 months, BW exhibited a negligible correlation \((|r| \leq 0.3)\) with BH, BL exhibited negligible correlations with CaC and ChD, and HW exhibited a negligible correlation with CaC. Moderate correlations \((0.3 < |r| \leq 0.7)\) were observed among the other traits (Table 2).

#### 3.2 Polymorphism in SSTR5

Seven SNPs were identified by sequencing: C186T (rs400944340), C351T (rs404123088), T384C (rs413380618), T537C (rs403055255), C576T (rs593868112), and T989C (rs601836309).
The first four SNPs were located in exon 2, SNP5 and SNP6 in exon 3, and SNP7 in exon 4. The genetic map of the mutated sites in SSTR5 based on the sequencing results is illustrated in Figure 1.

### 3.3 Analysis of Physicochemical Properties of Protein With Non-Synonymous Mutations

SNP1–6 were synonymous mutations. SNP7 was a non-synonymous mutation that caused the amino acid at the 330 position to change from methionine (Met) to threonine (Thr). According to analyses of the physicochemical properties of the mutant and wild-type proteins, the molecular weight of the wild-type was less than that of the mutant, and the wild-type had two O-glycosylation sites, but the mutant had only one (Table 3).

### 3.4 Population Genetics and the Linkage Disequilibrium Analysis

The Ne (effective allele numbers), calculated for each SNP, ranged from 1 to 2. The allele frequency of SNPs was in Hardy–Weinberg equilibrium ($p > 0.05$). Based on the PIC, SNP1–7 were classified as moderate polymorphic loci (Table 4). LD analysis revealed a strong LD ($D' > 0.85$) among SNP1–7 (Figure 2), and five common haplotypes were identified in this LD region.

### 3.5 Association Analysis of Genetic Variants and Haplotypes in SSTR5 With Growth Traits of Hulun Buir Sheep

The results of association analyses of the SSTR5 SNP genotypes and the growth traits at birth, 4, 9, and 16 months of age are shown in Supplementary Tables S2–S4.

#### 3.5.1 Association Analysis of SSTR5 With Growth Traits

The genotypes of SNP1 were significantly associated with BW, ChW, and ChC at 4 months of age and HW at 9 months of age ($p < 0.05$). They also showed extremely significant association with CaC at 4 months of age ($p < 0.01$). CaC at 9 months of age was significantly associated with the genotypes of SNP3, SNP4, and SNP7, and the genotypes of SNP7 were significantly associated with ChW at 4 months of age ($p < 0.05$, Figure 3). No significant differences were observed among the rest of the SNPs with other growth traits ($p > 0.05$).

#### 3.5.2 Haplotype Association Analysis With Growth Traits

The results of association analyses between the haplotypes and growth traits at birth, 4, 9, and 16 months of age are shown in Supplementary Table S5. There were no detectable differences among the five haplotypes of the seven SNPs in strong LD ($p > 0.05$).

### 4 DISCUSSION

This is the first report of associations between SNPs of SSTR5 and growth traits in sheep. Growth is one of the most important

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**Table 4** | Population genetics analyses of SSTR5 of in Hulun Buir sheep

| SNP     | Gene frequency | Ho  | He  | PIC  | Ne  | HW  |
|---------|---------------|-----|-----|------|-----|-----|
| SNP1 (C/T) | 0.2756 0.7244 | 0.3993 | 0.3991 | 0.3196 | 1.6648 | 0.5888 |
| SNP2 (C/T) | 0.6197 0.3803 | 0.4714 | 0.4807 | 0.3603 | 1.8917 | 0.6642 |
| SNP3 (T/C) | 0.2035 0.7965 | 0.3241 | 0.3391 | 0.2716 | 1.4796 | 0.5052 |
| SNP4 (T/C) | 0.2137 0.7863 | 0.3360 | 0.3519 | 0.2796 | 1.5061 | 0.5188 |
| SNP5 (C/T) | 0.2756 0.7244 | 0.3993 | 0.3991 | 0.3196 | 1.6648 | 0.5888 |
| SNP6 (G/A) | 0.6923 0.3077 | 0.4260 | 0.4893 | 0.3353 | 1.7423 | 0.6172 |
| SNP7 (T/C) | 0.2158 0.7842 | 0.3395 | 0.3962 | 0.2812 | 1.5117 | 0.5216 |

He = heterozygosity; Ho = homozygosity; PIC= polymorphism information content; Ne = effective allele numbers; HW= Hardy–Weinberg equilibrium.

*a*Group size of population genetics analyses was n = 233.
economical traits monitored in domestic animals; therefore, it is a significant research topic in most genetic selection programs (Koller et al., 2020). As the main measure of growth traits, body weight and size have important impacts on the production of meat and wool (Luo, et al., 2021). In this study, we recorded these production traits from birth to adulthood in Hulun Buir sheep. We measured body weight at 4, 9, and 16 months of age, using seven body size indicators at 4 and 9 months of age, and using five body size indicators at 16 months of age. The phenotypic data contained the main growth traits of the sheep, providing a comprehensive image of growth trends in the Hulun Buir sheep. Our study was more comprehensive than most other publications and featured a longer period.

Body size indicators are important tools for accurately measuring livestock. These measurements are used to study the appearance, characteristics of the breed, and production performance. The growth rate of body size of ovine used for meat varies with age (Chao et al., 2016). In Hulun Buir sheep, we identified significant positive correlations between body size indicators and body weight, and individual size indicators affected body weight directly or indirectly. At 4 months, ChD, BL, and ChW had the strongest correlations with BW, successively. At 9 months, ChC, BH, and BL had the strongest correlation with BW, successively. At 16 months, CaC, BL, and HW had the strongest correlation with BW, successively. Moreover, with an increase in age, the correlation between body size and body weight weakened.

SNPs, defined as a substitution, insertion, or deletion of a single nucleotide, are important genetic sources for animal breeding. Gene expression and protein functions are related to the location of the SNPs in regulatory sequences or coding regions (Stevenson, 2015). Exons are protein-coding regions consisting of only 1%–2% of the genome, and the mutation rate in exons is approximately one-fifth of that in non-coding regions (Komar, 2009). However, almost 85% of reported disease-causing genes harbor mutations in their exons, which is of great significance for the study of genetic diseases (Li H et al., 2019). Thus, exome sequencing is the most efficient approach for identifying potentially functional mutations related to phenotypes in domestic animals (Li et al., 2013). Exome sequencing is also the most cost-efficient sequencing approach for conducting genome research and animal phenotyping (Guo et al., 2014), and this technology can be used to more effectively identify SNPs affecting growth traits.

Linked loci are a particular concern as there is substantial LD among causal SNPs (Koch, 2019). Studies have shown that body size is affected by the buildup of interpopulation LD among loci, caused by selection (Routtu et al., 2014). In this study, we observed LD among SNP1–7 and identified five common haplotypes in this LD region. However, there was no correlation between these haplotypes and growth traits possibly because of the small sample size or because of interactions between other genes and the SNPs in SSTR5.
SSTR5 reduces the activity of MAPK, which is considered a key molecule in the transduction of growth factor proliferation signals (Cattaneo et al., 1996; Cordelier et al., 1997). Thus, we speculated that SSTR5 plays an important role in reducing GH secretion (Franck et al., 2017). In addition, because these SSTR5-specific agonists are 1,000 times more powerful than other SSTR5-specific agonists in inhibiting GHRH-stimulated GH release in the primary culture of rat pituitary cells, SSTR5 is considered one of the main mediators of SST-induced inhibition of GH release (Shimon et al., 1997). Finally, SSTR5 is involved in the regulation of ACTH release and may mediate SST-induced secretion of insulin expression/secretion and cell proliferation by down-regulating the duodenal homolog box-1 (PDX-1) (Park et al., 2003; Zhou et al., 2012); both ACTH and PDX-1 are key hormones related to animal growth traits. Therefore, SSTR5 may regulate the growth traits of Hulun Buir sheep. However, studies on SSTR5 have mainly focused on tumors; there has been no prior study on the association between SSTR5 and growth traits of livestock.

To discover the potential functional mutation related to the growth traits of Hulun Buir sheep, we conducted exome sequencing of SSTR5 and then performed association analyses of the phenotypic records. Seven SNPs (C186T, C351T, T384C, T537C, C576T, G768A, and T989C) were all in Hardy–Weinberg equilibrium, which indicated that the population size under random mating conditions (without selection) was adequate for the experiment (Liu et al., 2018).

Synonymous mutations can affect translation dynamics and protein folding, leading to phenotypic changes (McCarthy et al., 2017; Chu and Wei, 2019). This was consistent with our findings, while SNP1, SNP3, and SNP4 did not cause synonymous mutations in amino acid sequences, SNP1 genotypes were significantly associated with BW at 4 months of age and with ChC, Chw, CaC, and HW at 9 months of age; SNP3 and SNP4 were significantly associated with CaC at 4 months of age.

These types of mutations are often deleterious (Saavedra-Rodriguez et al., 2021). In this study, SNP7 was identified as a non-synonymous mutation significantly associated with CaC at 9 months of age. The CaC of the sheep with a wild-type genotype (TT) was significantly larger than that of the sheep with two mutants (TC and CC). These results were also observed in other association analyses of SNPs and growth traits, suggesting that the SNPs identified in this study may very likely be deleterious. These findings indicate that Hulun Buir sheep may exhibit a tendency for breed deterioration; therefore, further studies on the effect of SSTR5 on the growth traits of Hulun Buir sheep are necessary.

Analyses of the physicochemical properties of SNP7 indicated that the wild-type SSTR5 protein had two O-glycosylation sites, whereas the mutant had only one. The O-glycosylated stalk domain serves as a functional element for delivering proteins to the apical plasma membrane (Yeaman et al., 1997) and plays diverse, highly specific roles in fine-tuning protein functions (Schjoldager and Clausen, 2012). Therefore, SNP7 mutation may affect the function of the SSTR5 protein.

Association analyses of SNPs and growth traits in different ages revealed that the genotypes of SNP1, SNP3, SNP4, and SNP7 were associated with various growth indexes at 4 and 9 months of age. We inferred that the SSTR5 might affect the early growth and development of Hulun Buir sheep. These four SNPs were significantly correlated with CaC and may serve as molecular markers to determine CaC.

5 CONCLUSION

In the current study, we used exon sequencing technology to screen SSTR5 and discovered seven SNPs in Hulun Buir sheep. SNP1, SNP3, SNP4, and SNP7 were associated with CaC, demonstrating potential as molecular markers for the selection of CaC in Hulun Buir sheep. The genotypes of SNP1 were also associated with BW and ChC at 4 months of age, and HW at 9 months of age, indicating that SNP1 could be used as molecular markers for the selection of growth traits in Hulun Buir sheep. These molecular markers may provide a theoretical basis for improving the growth traits of Hulun Buir sheep.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

ETHICS STATEMENT

The animal study was reviewed and approved by the Ethics Committee of Experimental Animal Welfare, Northwest Institute of Plateau Biology, Chinese Academy of Sciences.

AUTHOR CONTRIBUTIONS

XL, DT, FT and KZ provided the study concept and design. XL wrote and revised the manuscript. ND, ZZ, DL, BH, SL, DF and XS collected and analyzed the data. All authors have read and approved the final manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2022.831599/full#supplementary-material
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**Conflict of Interest:** DF and XS were employed by the Inner Mongolia Daxing ‘Anling Agricultural Reclamation Group Co. LTD. They only help with sample collection and have no business cooperation or conflict of interest with their company.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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