

Two co-inherited novel SNPs in the MC4R gene related to live body weight and hormonal assays in Awassi and Arabi sheep breeds of Iraq.

T. M. Al-Thuwaini, M. B. S. Al-Shuhaib, Frédéric Leprêtre, H. H. Dawud

▶ To cite this version:

T. M. Al-Thuwaini, M. B. S. Al-Shuhaib, Frédéric Leprêtre, H. H. Dawud. Two co-inherited novel SNPs in the MC4R gene related to live body weight and hormonal assays in Awassi and Arabi sheep breeds of Iraq.. Veterinary Medicine and Science, 2021, Veterinary Medicine and Science, 7 (3), pp.897-907. 10.1002/vms3.421 . hal-04475882

HAL Id: hal-04475882 https://hal.univ-lille.fr/hal-04475882v1

Submitted on 23 Feb 2024

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.



ORIGINAL ARTICLE

WILEY

Two co-inherited novel SNPs in the *MC4R* gene related to live body weight and hormonal assays in Awassi and Arabi sheep breeds of Iraq

Tahreer M. Al-Thuwaini¹ | Mohammed Baqur S. Al-Shuhaib¹ | Frederic Lepretre² | Halla Hassan Dawud¹

Correspondence

Tahreer M. Al-Thuwaini, Department of Animal Production, College of Agriculture, Al-Qasim Green University, Babil, Iraq.

Abstract

Melanocortin-4 receptor (MC4R) gene plays a key role in the regulation of body weight and energy homeostasis. This study aims to evaluate the association of single nucleotide polymorphisms (SNPs) of the MC4R gene with live body weight and hormonal assays in two breeds of sheep that differ in productive performance, Awassi and Arabi. All known coding sequences of the MC4R gene were covered in this study. DNA samples from 150 animals (Awassi and Arabi breed) were genotyped by PCRsingle-strand conformation polymorphism (PCR-SSCP) to assess their pattern of genetic variation. Concerning exon 1, clear heterogeneity was detected with three different SSCP-banding patterns. The sequencing reactions confirmed these variations by detecting the presence of the two novel SNPs, 107G/C and 138A/C, and three genotypes, GC, AC and AA. The 107G/C SNP was detected in GC genotype, while the 138A/C was detected on both GC and AC genotypes. The other SSCPbanding pattern (AA genotype) did not show any detectable unique variation. Both SNPs were closely and strongly linked in both breeds (D' and r^2 values were 1.00), which signifies that both loci were co-inherited as one unit. Association analysis indicated that both breeds with GC/AC haplotype showed higher live body weight (37.250 ± 0.790) relative to the GG/AA (30.244 ± 0.968) and CC/CC (47.231 ± 1.230) haplotypes (p < .05). Concerning the genotyping of exon 2, only 362 bp showed heterogeneity with a missense mutation, with no significant association (p > .05) with the measured traits. In conclusion, the two novel SNPs (107G/C and 138 A/C) were highly associated with live body weight in both breeds. Haplotype analysis confirmed that these two novel SNPs were in strong linkage disequilibrium (LD) and could be used as genetic markers for sheep phenotypic trait improvement.

KEYWORDS

genetic polymorphism, haplotype, in silico, production trait

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2020 The Authors. Veterinary Medicine and Science Published by John Wiley & Sons Ltd

¹Department of Animal Production, College of Agriculture, Al-Qasim Green University, Babil, Iraq

²University of Lille, Plateau de Genomique Fonctionnelle et Structurale, Lille, France

1 | INTRODUCTION

Body weight in sheep is a pivotal trait for economic breeding and is controlled by multiple genetic loci (Esmailizadeh, 2010). Among these loci, the melanocortin-4 receptor (MC4R) gene plays an important role in the regulation of body weight, energy balance and reproduction in humans and animals (Bakos et al., 2016; Siljee et al., 2013; Zeng et al., 2014). The ovine MC4R gene is mapped on chromosome 23 and consists of two exons (Shishay et al., 2019). The MC4R gene is involved in the regulation of phenotype expressions of several economic traits in ruminant and non-ruminant animals. These traits are largely regulated by the binding of the MC4R gene-encoded receptors to four ligands of melanocyte-stimulating hormone (α -, β - and γ -MSH) and the adrenocorticotropic hormone (ACTH; Switonski et al., 2013).

The MCR family consists of five members (MC1R to MC5R); each of which has seven transmembrane domains belonging to the G protein-coupled receptor superfamily (Li & Li, 2006). Out of these five melanocortin receptors, MC4R has received more attention. It is expressed in the appetite-regulating areas of the brain that involve food intake, which can act as a critical mediator between appetite and reproduction in animals (Zandi et al., 2014). Thus, the MC4R gene can be a major mediator of leptin effects on food intake and body weight (El-Sabrout & Soliman, 2018; Hwa et al., 2001; Li & Li, 2006). The effects of the MC4R gene variation have been recently implicated in the rapid growth selection program of beef cattle (Prihandini & Maharani, 2019). On the other hand, it has been shown that some of the variations detected in the MC4R gene have been associated with economic traits in livestock (Shishay et al., 2019). It has been recently reported that the MC4R gene polymorphism has exhibited a noticeable association with weight, body length and chest circumference gains, in addition to the average daily gain in goats (Latifah et al., 2018). Likewise, several reports have also indicated a remarkable role of MC4R gene in live weight, backfat thickness, carcass traits and marbling score in several breeds of cattle (Lee et al., 2013; Liu et al., 2010; Seong et al., 2012). Additionally, genotyping studies of the MC4R gene have revealed a significant association of this gene with the milk yield and fat percentage in buffalos (Deng et al., 2016). Moreover, the MC4R gene polymorphism has been associated with backfat thickness in sheep (Zuo et al., 2014). The combined effect of multiple mutation sites within MC4R gene has also been included as a valuable factor for assessing its effect on economic traits (El-Sabrout & Aggag, 2018). However, to accommodate such haplotyping studies, it is necessary to assess multiple genetic loci simultaneously (Lu et al., 2003).

Taking into consideration the above-stated data, the combining of the possible interaction between *MC4R* polymorphism in Awassi and Arabi breed with live body weight and hormonal assays may have some potential in the detection of the possible correlation between both patterns.

Awassi, followed by Arabi, is the most predominant one in the middle and southern portions of Iraq (Al-Shuhaib et al., 2019). Though Awassi sheep have well-known living adaptation capacity,

it has recently reported that Arabi sheep showed a higher genetic capability than Awassi sheep to cope up with harsh circumstances (Al-Thuwaini et al., 2020). Furthermore, it has recently been stated that genetic diversity is considerably correlated with the productive performance in both breeds (Aljubouri & Al-Shuhaib, 2020). Taking these genetic differences altogether, it is crucial to discriminate between both breeds to broaden our knowledge in critical production traits. Several biological functions have been associated with such adaptation conditions in sheep, such as live body weight and hormonal secretion (Marai et al., 2007). Thus, it is consequential to measure these values in both breeds taking advantage of the highly dynamic strategy sheep use to enhance their survival chances in harsh conditions (Niyas et al., 2015). Thus, it would be expected that Awassi and Arabi breeds could also exhibit different genetic variations in the MC4R gene in such a way it could affect functions associated with production traits. Therefore, the current study aims to describe the possible association between the MC4R gene polymorphism and the physiological differences between Awassi and Arabi breeds. Thus, single nucleotide polymorphisms (SNPs) of MC4R gene were identified, and association with live body weight was investigated to acquire possible molecular markers related to production traits for marker-assisted selection. According to our knowledge, no report has screened the association of the coding sequences of the MC4R gene with live body weight and hormonal assays in sheep. Therefore, this research is the first one to describe such association in two breeds of sheep.

2 | MATERIALS AND METHODS

2.1 | Sheep population and ethical approval process

The study was conducted according to the international recommendations for the care and use of animals (Federation of Animal Science Societies,), and the animal experimentations were approved by the Al-Qasim Green University (Approval No. 12.10.18). A total of 150 sheep (Ovis aries) were selected randomly from three stations for raising sheep (Babylon, Karbala and Kufa provinces, Iraq) from January 2018 to August 2018. The three stations were approximately situated at a longitude of 32.6027°N, the latitude of 44.0197°E and 32 m a.s.l. Expert veterinarians confirmed the included sheep to be clinically healthy. Two types of breeds, including Awassi (n = 75, 22 rams and 53 ewes) and Arabi (n = 75, 15 rams and 60 ewes), were included in this study. Both included breeds were aged between 2.5 and 3 years. All included ewes were in non-pregnant and non-lactation status, and the parity of ewes was one to two parities. The animals of both breeds had different genetic and geographical backgrounds (Al-Thuwaini et al., 2020). Awassi breed prevails in the Middle Euphrates regions with summer temperatures of <50°C, while Arabi breed prevails in Southern portions of Iraq with summer temperatures exceeding 50°C. Both breeds are fattailed, carpet-wool producers, with some potential to produce milk. However, both breeds had slightly different head appearance and body characteristics (Alkass & Juma, 2005). Animals were fed ad libitum on seasonal grass during summer, while in winter, animals were kept indoors and fed with a concentrated mixture consisting of barley grain (59%), bran (40%) and salt (1%). The live body weight of the sheep was recorded in the morning before the animals were grazing using a suspended spring balance, while blood tests were assessed. The descriptive statistics of estimated traits are shown in Table 1.

2.2 | Hormonal assay

Within 20 min of blood collection from the jugular vein, plasma was separated from peripheral blood by centrifugation at 3,500 g at 4°C for 15 min. Then, the plasma from each sample was collected and stored at -20°C. To assess the possible association between *MC4R* gene and sex hormones in both investigated breeds, testosterone and estradiol were measured using enzyme-linked immunosorbent assay (ELISA) kit based on the sandwich principle (cat. no. E0013Sh for testosterone and cat. no. E0047 Sh for estradiol, Bioassay Technology Laboratory Co.). The absorbance of hormones was measured at 450 nm using a microplate reader (ELx808 Ultraplate Reader, BioTek Instruments, Inc.). Concentrations were presented as pg/ml and ng/ml for estradiol and testosterone, respectively. A standard curve was generated, and samples were interloped according to the manufacturer's instructions (Bioassay Technology Laboratory Co.).

2.3 | Genomic DNA extraction, primer design and PCR

Genomic DNA was isolated from the whole blood using a rapid and efficient salting-out method (Al-Shuhaib, 2017). Four pairs of specific polymerase chain reaction (PCR) oligonucleotides were designed to cover all the coding sequences of the ovine *MC4R* gene using NCBI Primer Blast online server (Ye et al., 2012). For exon 1, only one PCR primer pair was designed, while three pairs of primers were designed to adequately cover the coding sequences in the exon

TABLE 1 Descriptive statistics of estimated traits used in the association analysis

Breed	Phenotypic traits	N	Mean	SD	Minimum	Maximum
A) Awassi						
	Live body weight (Kg)	75	41.909	4.113	25.0	52.0
	Estradiol (pg/ml)	75	28.980	9.272	12.040	67.148
	Testosterone (ng/ ml)	75	0.928	0.035	0.107	4.666
B) Arabi						
	Live body weight (Kg)	75	37.650	2.225	25.0	48.0
	Estradiol (pg/ml)	75	25.915	10.174	10.630	51.869
	Testosterone (ng/ ml)	75	0.711	0.079	0.147	4.185

Abbreviation: N, number of observation; SD, standard deviation.

2 (Figure 1a). The lyophilized oligonucleotides were purchased from Bioneer Company (Bioneer, Korea). The PCR primers were designed based on GenBank accession no. NC_019480.2 and the details of the primer used in this study are shown in Table 2. PCR experiments were conducted using *AccuPower®* PCR PreMix (Bioneer), and initiated by denaturation for 5 min, followed by 30 cycles (annealing at 57.8–59.1°C for 30 s each), and finalized with polymerase extension (72°C) for 5 min. The target amplified PCR products were confirmed by 1.5% agarose gel electrophoresis and then submitted to single-strand conformation polymorphism (SSCP) protocols (Hashim & Al-Shuhaib, 2019).

2.4 | Single-strand conformation polymorphism (SSCP) and sequencing analysis

The initial denaturation of the PCR products, as well as SSCP protocol, was performed according to Al-Shuhaib et al., (2018) with some optimization suitable for the designed amplicons. Briefly, SSCP analyses were conducted in polyacrylamide gels (37.5:1) with TBE (0.5X) buffer at a constant temperature of 10° C using 216×110 mm mini-wide gels with 1 mm gel thickness (JY-CZ-B, Junyi-Dongfang Electrophoresis Equipment). Gel concentrations were adjusted to 10% and 8% for exon 1 and exon 2 amplicons, respectively. Electrophoresis conditions applied were 210 V/105 mA/5 hr and 200 V/100 mA/4 hr for exon 1 and exon 2 amplicons, respectively. Bands were fixed and stained according to the protocol described by Byun et al., (2009). Each detected SSCP-banding pattern of all investigated samples was sent for sequencing from both termini according to the instruction manual described by Macrogen laboratories (Geumcheon). The electropherograms were edited and aligned using EditSeq software, ver. 7.1.0 (DNASTAR, Lasergene). The observed mutations were visualized and annotated using SnapGene Viewer, ver. 4.0.4. (GSL. Biotech. LLC). The novelty of the observed variants was checked by exploring variants of the ovine MC4R gene database in the Ensembl genome browser 96 (https://asia.ensembl.org/index. html).

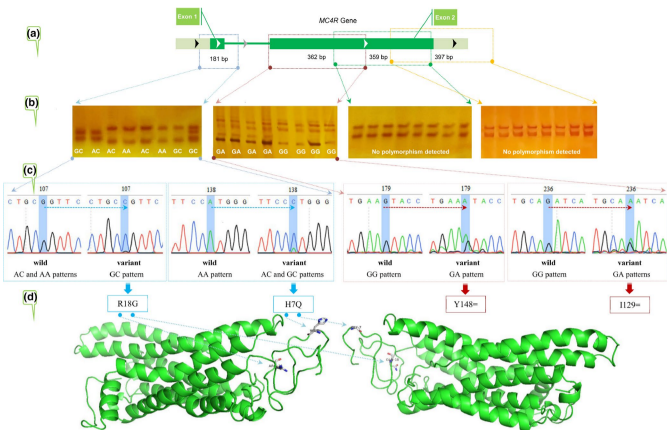


FIGURE 1 A schematic diagram of the present study to assess the *MC4R* gene polymorphism in the Awassi and Arabi sheep. (a) The exact genomic positions of the amplify a portion of the ovine *MC4R* gene were described according to GenBank acc. no. NC_019480.2. (b) PCR-single-strand conformation polymorphism (PCR-SSCP) genotyping of the amplified loci. (c) DNA sequencing chromatogram of the polymorphic fragment. (d) The *in silico* prediction of *MC4R* SNPs

Set	Primer code	Primer sequence (5'→3')	Length	Annealing temp.
1	MC4R,exo1-F	GTCACAAACACCTCGGGAGA	181 bp	57.8°C
	MC4R,exo1-R	TCCAGAGGGGACCTGAATCC		
2	MC4R,exo2,1-F	TGGGGCAGGAGATGTAGAA	359 bp	57.8°C
	MC4R,exo2,1-R	GCGCTCCAGTACCATAGCAT		
3	MC4R,exo2,2-F	TGAGAGCCAGCATGGTGAAG	362 bp	60.4°C
	MC4R,exo2,2-R	TGTGGCTGACATGTTGGTGA		
4	MC4R,exo2,3-F	ACCGCAGTTTGTCCCCATTT	397 bp	59.1°C
	MC4R,exo2,3-R	CATGGCGTCTCTCTACGTCC		

TABLE 2 Oligonucleotide primer sets designed for the amplification of the MC4R in Awassi and Arabi sheep population. The present annotations of this study variants were based on GenBank accession number NC_019480.2

2.5 | In silico prediction

Several computational tools were utilized to assess the consequences of the observed missense variants on the resulting mutant protein structures and functions, namely SIFT (Ng & Henikoff, 2003), PolyPhen-2 (Adzhubei et al., 2010), PROVEAN (Choi et al., 2012), Panther (Tang & Thomas, 2016) and PhD SNP (Capriotti et al., 2006). The 3D structure of *MC4R* was generated by RaptorX server before and after mutation (Källberg et al., 2012),

and validated in verify3D and PROCHECK servers (http://servicesn.mbi.ucla.edu/Verify3D/).

2.6 | Genetic polymorphism and statistical analyses

The allele and genotype frequencies, observed heterozygosity (*Ho*), expected heterozygosity (*He*) and an effective number of alleles were analysed using PopGen32 software, ver. 1.31 (Yeh & Yang, 1999). The polymorphism information content (*PIC*) was calculated by

TABLE 3 In silico prediction of H7Q and R18G on ovine MC4R protein, in terms of structure and function

	H7Q		R18G		
Tool	Score	Prediction	Score	Prediction	
SIFT	0.44	Tolerated	0.24	Tolerated	
PolyPhen-2	0.148	Benign	0.132	Benign	
PROVEAN	-0.042	Neutral	0.24	Neutral	
PANTHER	Unknown preservation time	Invalid substitution	30 (preservation time)	Probably benign	
PhD SNP	89% confidence	Neutral	78% confidence	Neutral	

utilizing the HET software ver. 1.8 (Ott, 2001). Pairwise linkage disequilibrium (LD) between SNPs was calculated by r^2 and D' values using SHEsis software (She & He, 2006).

The significant effects of breed, sex and SNP genotype on the various studied parameters were analysed by SPSS software ver. 23.0 (SPSS Inc), using the general linear model:

$$y_{ijkl} = \mu + B_i + S_j + G_k + e_{ijkl},$$

where y_{ijkl} is phenotypic traits, μ is the overall mean, B_i is the fixed effect of ith breed (i = Awassi, Arabi), S_j is the fixed effect of jth sex (j = ram, ewe), i_k is the fixed effect of kth SNP genotype or combined genotype and e_{ijkl} is the random error associated with y_{ijkl} observation and assumed to be NID (0, σ 2e). Means were compared using the Tukey-Kramer test with a significance level of p < .05. Preliminary statistical analysis indicated that the effects of age, parity, season, station and the interaction between these effects were not included in the final model because they did not have a significant effect on phenotypic traits.

2.7 | SNP genotype effects estimation

For the SNP that showed significant association with the phenotypic traits, differences between the means of each genotype and allelic frequencies were used to estimate additive effects (Hill & Mackay, 2004). The following formula was utilized to find additive genetic variance (Var_{Δ}) imputed to a SNP:

$$Var_A = 2p_i q_i \alpha_i^2$$
,

where q and p were the allelic frequencies for the jth SNP predicted across the entire population, α_i – SNP allele substitution effect obtained from a linear regression model in a statistical program, in which the genotypes recorded as a variate of 0, 1 and 2 copies of a particular allele.

The proportion of the phenotypic variance explained by SNP(s) was calculated by (%) var $P=100\times 2p_iq_i\alpha_i^2/V_P$, where p and q denote the SNP allele frequency; α_i^2 is the SNP allele substitution effect, $2p_iq_i\alpha_i^2$ is the additive genetic variance and V_p is the phenotypic variance.

3 | RESULTS

3.1 | The genetic polymorphism and in silico analysis

Four amplicons were used to scan all the coding regions of the MC4R gene, including only one amplicon (181 bp) for exon 1, and three amplicons (362 bp, 359 bp and 397 bp) for the exon 2. Only exon 1 amplicons showed heterogeneity with a missense effect, while the exon 2 (362 bp) amplicons showed heterogeneity with a silent mutation. However, only monomorphous SSCP-banding patterns were revealed in both 359 and 397 bp amplicons. Thus, the genotyping by the SSCP method was utilized in exon 1 to assess the pattern of genetic variation in both breeds. Three SSCP-banding patterns were detected in exon 1 in the same SSCP gel conditions (Figure 1b). Sequencing reactions confirmed these patterns and allowed the identification of 10 SNPs in exon 1 with missense and silent effects. Eight of these SNPs, 102G > C (S19R), 113A > C (W16G), 117G > A (H14=), 122G > A (L13F), 124G > A (S12F), 126G > A (T11=), 153G > A (N2=) and 168A > C (N2=), were detected in all analysed samples, while two SNPs, 138A > C (H7Q) and 107G > C (R18G), exhibited variable distributions in the exon 1. Both 107G > C and 138A > C SNPs were found to be responsible for the observed heterogeneity of samples having GC and AC genotypes, respectively. The other samples with a homologous SSCP-banding pattern (AA genotype) did not exhibit any detectable unique variation. The 107G > C SNP was found to cause the amino acid R18G (arginine to glycine) substitution. Whereas the (H7Q) histidine to glutamine amino acid change was caused by the 138A > C SNP (Figure 1c). Concerning exon 2, both PCR-SSCP method as well as sequencing reactions revealed a remarkably less polymorphic status than the exon 1. No polymorphism was detected in both 359 bp and 397 bp amplicons, while two different SSCP-banding patterns were identified in the 362 bp amplicons. Sequencing reactions confirmed the presence of two SSCP patterns in the 362 bp amplicons with the detection of two SNPs in analysed samples with GA genotype (179G > A and 236G > A), while no polymorphism was observed in samples with GG genotype.

A series of computational tools were utilized to assess the final consequences of this missense variant on the altered MC4R protein in structure and function, such as SIFT, PolyPhen-2, PROVEAN, Panther and PhD SNP (Table 3). All of these tools lead to neutral/non-deleterious signals for both assessed R18G and H7Q. This

Genotype, allele frequencies and genetic diversity parameters in the MC4R gene for both Awassi and Arabi breeds 4 TABLE

	SNP	Genotypes (N)	Genotype frequencies	Allele	Allele frequencies	2%	Я	Не	Ne	PIC
A) Arabi	179G > A	GG(37) GA(38)	0.49 0.51	ე ∢	0.75 0.25	8.362*	0.506	0.380	1.608	0.304
	236G > A	GG(41) GA(34)	0.55 0.45	ე ∢	0.77	6.223*	0.453	0.352	1.539	0.291
	107G > C 138A > C	GG/AA(7) GC/AC(42) CC/CC(26)	0.09 0.56 0.35	& - CC CC	0.37	55. 733*	906:0	0.629	2.667	0.357
B) Awassi	179G > A	GG(42) GA(33)	0.56 0.44	ს ∢	0.78	5.757*	0.440	0.345	1.522	0.284
	236G > A	GG(39) GA(36)	0.52 0.48	ე ∢	0.76	7.238*	0.480	0.367	1.574	0.298
	107G > C 138A > C	GG/AA(12) GC/AC(45) CC/CC(18)	0.16 0.60 0.24	G - CC	0.46 - 0.54	46.127*	0.840	0.601	2.485	0.373

Abbreviations: He, Expected heterozygosity; Ho, observed heterozygosity; Ne, effective allele number; PIC, polymorphism information content; y², Chi-square. tests have two degrees of freedom and within the significance *All Chi-square non-deleterious effect may be due to their non-critical positions in the generated 3D structure of MC4R protein (Figure 1d).

3.2 | Assessment of MC4R polymorphism and association analysis

Genotype, allele frequencies and genetic diversity parameters for exon 1 amplicon in the analysed populations of Awassi and Arabi sheep are presented in Table 4, A and B. According to the value of Chi-square, the population under study was not in Hardy-Weinberg equilibrium (HWE), which was statistically significant at p < .05. According to the classification of PIC (low polymorphism, if PIC value < 0.25; median polymorphism if 0.25 < PIC value < 0.5 and high polymorphism if PIC value > 0.5; Trakovická et al., 2013), analyses showed that four SNPs (107G > C, 138A > C, 179G > A and 236G > A) were considered as moderate PIC. Pairwise LD between SNPs was calculated and indicated that the D' and r² values were between 0.99 and 1.00, signifying a close connection and strong co-inheritance for both 107 G > C and 138 A > C in both breeds (Figure 2). The haplotypes of the two SNPs were constructed for each breed and in all analysed populations. In relative terms, there was no LD between the two SNPs (179G > A and 236G > A) in exon 2 also with SNP pairs in exon 1.

Awassi breed showed higher live body weight (44.855 \pm 0.683) in comparison to Arabi breed (38.333 \pm 0.790; p < .05). Meanwhile, the conducted sex hormone assays showed no significant (p > .05) differences (0.618 \pm 0.005 ng/ml and 0.987 \pm 0.003 ng/ml for testosterone, and 26.826 \pm 1.966 pg/ml and 29.800 \pm 3.296 ng/ ml for estradiol) between Arabi and Awassi breeds, respectively. Association analyses for live body weight and sex hormone assays according to the breed and sex effect are presented in Table 5. These results indicated that Awassi breed had more body weight gain than Arabi breed. In regards to sex effect, significant differences (p < .05) between rams and ewes were identified, being heavier weight in the ewes (39.833 \pm 1.328) in comparison to rams (34.167 \pm 0.511). Based on the haplotype analysis, a combined analysis of genotypes revealed that the haplotype block was significantly associated with phenotypic traits (p < .05). The sheep with CC/CC haplotype showed higher live body weight (48.567 \pm 2.956; 45.855 \pm 3.626; 47.231 ± 1.230) and lower estradiol level (16.745 \pm 1.460; 14.410 ± 1.883 ; 15.577 ± 1.266) in Awassi, Arabi and both breeds, respectively, than the other combined genotypes. Whereas the testosterone level showed no significant differences among the observed haplotypes (p > .05). Individuals with GC/AC haplotype showed higher live body weight (41.324 \pm 3.961; 33.176 \pm 2.581; 37.250 ± 0.790) than those with GG/AA haplotype (30.800 ± 2.347; 29.696 ± 2.899 ; 30.244 ± 0.968 ; p < .05) in Awassi, Arabi and both breeds, respectively. Whereas no significant differences (p > .05) were observed in the concentrations of estradiol and testosterone between GG/AA and GC/AC haplotypes. These results indicated that the individuals with GC/AC haplotype had a higher body weight and could be used in the improvement of growth traits in Awassi

are governed by the applicable Creative Commons I

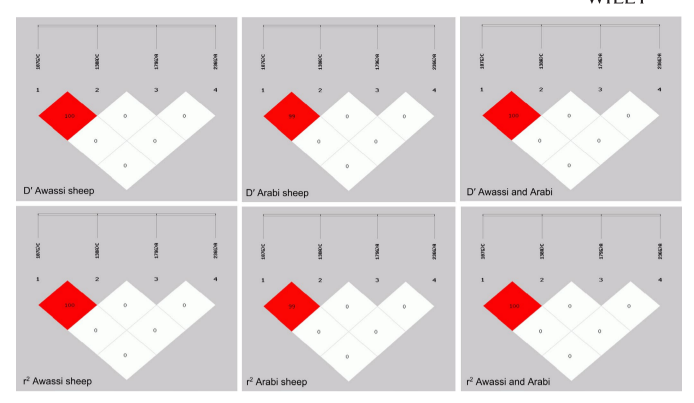


FIGURE 2 Linkage disequilibrium plot of the observed four SNPs within the exon 1 and exon 2 amplicons of the ovine *MC4R* gene in Awassi, Arabi and two breeds. (a) D' value, (b) r^2 value. The red diamonds indicate strong LD between pairs of SNPs

and Arabi breeds, respectively. Furthermore, the genotype effect prediction confirmed that the GC/AC haplotype was associated with higher body weight (p < .05). Greater genetic variance percentage with the phenotypic traits (>1%) is detailed in Table 6A and B. Meanwhile, the individuals with CC/CC haplotype had higher live body weight with a lower level of estradiol indicating that this haplotype should not be extensively selected in sheep breeding for litter size in Awassi breed. However, the association analysis in exon 2 showed no significant difference (p > .05) between 179G > A and 236G > A and phenotypic traits.

4 | DISCUSSION

Association analyses for live body weight revealed that the Awassi breed showed higher live body weight in comparison to Arabi breed, indicating that Awassi breed had more body weight gain than Arabi breed. The breed of sheep was known to have an important impact on body weight (Aktaş et al., 2015). Awassi sheep were shown to respond favourably to body weight than other breeds (Galal et al., 2008). In regards to sex effect, this study found that ewes showed heavier weight compared to rams in both breeds. These differences might be attributed to the physiological and sexual hormones of rams and ewes (Kratochvílová et al., 2002). Indeed, oestrogen hormone is stimulating specific receptors in the pro-opiomelanocortin (POMC) neurons in the hypothalamus by the melanocortin system (Clegg et al., 2006). It has been reported that increased POMC levels result in more binding of α -MSH to MC4R,

promoting increased food intake and reduced energy expenditure in ewes (El-Sabrout & Soliman, 2018; Hewagalamulage et al., 2015). Furthermore, females are more severely affected than males in weight increase (Vaisse et al., 2000).

The MC4R gene/protein variation is a key component in the MC4R pathway, which regulates phenotypic traits. Sequencing analyses of the current study confirmed the identification of two missense SNPs, 107G > C and 138A > C, in exon 1 of the MC4R gene. Both identified SNPs were associated with live body weight and sex hormone levels. El-Sabrout (2017) found that the SNPs of the MC4R gene were associated with body weight and sexual desire behaviour in rabbits. Our results are consistent with numerous studies conducted in livestock species that reported the association of the MC4R genetic polymorphism with phenotypic traits. Individuals with a synonymous 93G > A mutation in the MC4R gene have significantly associated with a backfat thickness in sheep (Zuo et al., 2014). The g.998A/G SNP was significantly associated with weight gain, body length and chest circumference gain in Bligon goats (Latifah et al., 2018). Add to that, individuals with the AG and GG genotypes (-129A > G) in the MC4R gene had higher live body weight than individuals with AA genotype in cattle (Liu et al., 2010). In another study, individuals with heterozygous genotype at g.1104C > T of the MC4R gene showed better performance than individuals with homozygous genotype for milk yield in Chinese buffaloes (Deng et al., 2016). Although the association study between single SNP with phenotypic traits is likely to be easier and more efficient in breeding programs, the haplotype analysis is valuable for assessing the effect of genes on phenotypic traits. In this study, statistical analysis showed that

TABLE 5 Least square means \pm SE for live body weight and sex hormone assay in association with the MC4R polymorphism in Awassi (A), Arabi (B) and both breeds (C)

Breed	Locus	Genotype	Live body weight (Kg)	Testosterone (ng/ml)	Estradiol (pg/ ml)
A) Awassi	Exon 1;107G > C	GG/AA	30.800 ± 2.347°	1.306 ± 0.036 ^a	25.109 ± 3.972 ^a
	138A > C	GC/AC	41.324 ± 3.961^{a}	0.869 ± 0.029^{a}	22.101 ± 2.725 ^a
		CC/CC	48.567 ± 2.956 ^b	0.613 ± 0.057^{a}	16.745 ± 1.460 ^b
	Exon 2;179G > A	GG	29.191 ± 2.725°	0.920 ± 0.036^a	22.203 ± 2.809^a
	236G > A	GA	$31.400 \pm 3.460^{\circ}$	1.120 ± 0.029^a	22.635 ± 3.388^a
B) Arabi	Exon 1;107G > C	GG/AA	29.696 ± 2.899°	1.106 ± 0.027^{a}	24.149 ± 2.010^{a}
	138A > C	GC/AC	33.176 ± 2.581 ^a	0.860 ± 0.010^{a}	20.098 ± 3.250^{a}
		CC/CC	45.855 ± 3.626^{b}	0.534 ± 0.024^{a}	14.410 ± 1.883^{b}
	Exon 2;179G > A	GG	$31.062 \pm 2.167^{\circ}$	0.923 ± 0.057^a	22.218 ± 2.972^{a}
	236G > A	GA	31.040 ± 2.081^{c}	0.922 ± 0.035^{a}	20.408 ± 2.756^{a}
C) Awassi and Arabi	Exon 1;107G > C	GG/AA	30.244 ± 0.968^{c}	1.204 ± 0.004^{a}	24.627 ± 2.870^{a}
	138A > C	GC/AC	37.250 ± 0.790^{a}	0.865 ± 0.014^{a}	21.098 ± 2.870^{a}
		CC/CC	47.231 ± 1.230^{b}	0.574 ± 0.003^a	15.577 ± 1.266^{b}
	Exon 2;179G > A	GG	$30.123 \pm 0.526^{\circ}$	0.921 ± 0.001^a	22.210 ± 1.320^{a}
	236G > A	GA	31.220 ± 0.466^{c}	1.021 ± 0.002^{a}	21.521 ± 4.421^{a}

Note: Different superscript in the same column within each classification indicates significant differences (p < .05).

Different superscript a, b, c indicates the significant difference.

individuals with GC/AC haplotype were more associated with live body weight gain than those with GG/AA and CC/CC haplotypes, respectively. These results confirmed that the priority of genomic and inbreed selection should be held on heterozygous states compared to homozygous ones. The study of Cai et al., (2015) showed that individuals with CGACG and CTCCC haplotypes of the MC4R gene were associated with an increased body weight of animals aged 18 months in yaks. Based on previous information, Awassi sheep is a low prolific breed (Abdullah et al., 2002), and very low incidence of twinning (Al-Sa'aidi et al., 2018). The present study demonstrated that individuals with CC/CC haplotype had higher live body weight and lower levels of estradiol, which indicates that this haplotype has not been extensively selected in sheep breeding to enhance litter size and twinning ratio in Awassi breed. This fact was confirmed by Chen et al., (2017) who reported that the variation of MC4R gene implies a reduction of GnRH-LH secretion and dysfunction of the ovary in mice, resulting in a reduction in the number of developed corpora lutea and decreased litter sizes.

5 | CONCLUSION

This study identified two co-inherited novel SNPs (107G/C and 138 A/C) of MC4R gene in two breeds of sheep. Both of the 107G/C and 138A/C SNPs were found to be highly associated with live body weight and hormonal (testosterone and estradiol) assays. We demonstrated that the GC/AC haplotype affects the sheep live body weight and hormonal assays and it is therefore highly recommended to be selected and fixed for sheep production.

ACKNOWLEDGEMENTS

This study was supported by the college of agriculture, Al-Qasim Green University. The authors are grateful to the staff of stations for raising sheep for their facilities that provided the sheep population.

CONFLICT OF INTEREST

The authors declare that they have no conflict of interest.

AUTHOR CONTRIBUTIONS

Tahreer M. Al-Thuwaini: Conceptualization; supervision; Methodology (lead). Mohammed Baqur S. Al-Shuhaib: Writing-original draft, editing, Data curation and investigation (lead). Frederic Lepretre: Data curation and investigation (equal). Halla Hassan Dawud: Methodology (equal).

FUNDING INFORMATION

The authors have not declared a specific grant for this research from any funding agency in the public, commercial or not-for-profit sectors.

ETHICAL APPROVAL

The study was conducted according to international recommendations for the care and use of animals (Federation of Animal Science Societies, 2010) and the animal experimentations were approved by Al-Qasim Green University (Approval No. 12.10.18).

PEER REVIEW

The peer review history for this article is available at https://publo ns.com/publon/10.1002/vms3.421.

20531095, 2021, 3, Downloaded from https://onlinelibrary.wiley.com/doi/10.1002/vms3.421 by Cochrane France, Wiley Online Library on [23.022024]. See the Terms and Conditions (https://onlinelibrary.wiley .com/terms-and-conditions) on Wiley Online Library for rules of use; OA articles are governed by the applicable Creative Commons License

TABLE 6 Effect of substitution in 107G > C and 138A > C polymorphism of MC4R on the live body weight and estradiol hormone of Awassi and Arabi breed. (A) SNP allele substitution effect (α) is shown. (B) Haplotype (h) substitution effect is shown

A)	Phenotypic traits	SNPs	Substitu	ution allele	α Allele substitution effect	varP (%)
Awassi	Live body weight (Kg)	107G > C	G		1.821	3.315
	Estradiol (pg/ml)	107G > C	G		1.742	1.741
	Live body weight (Kg)	138A > C	Α		1.952	3.808
	Estradiol (pg/ml)	138A > C	Α		1.724	2.971
Arabi	Live body weight (Kg)	107G > C	G		1.573	2.473
	Estradiol (pg/ml)	107G > C	G		1.421	2.018
	Live body weight (Kg)	138A > C	Α		1.601	2.561
	Estradiol (pg/ml)	138A > C	Α		1.324	1.752
В)	Phenotypic traits	SNPs		Haplotype	h substitution effect	varP (%)
Awassi	Live body weight (Kg)	107G > C + 138	A > C	GG/AA	1.930	3.723
	Estradiol (pg/ml)	107G > C + 138	A > C	GG/AA	1.724	2.971
	Live body weight (Kg)	107G > C + 138	A > C	GC/AC	1.952	3.808
	Estradiol (pg/ml)	107G > C + 138	A > C	GC/AC	1.742	3.033
	Live body weight (Kg)	107G > C + 138	A > C	CC/CC	1.992	3.967
	Estradiol (pg/ml)	107G > C + 138	A > C	CC/CC	1.821	3.315
Arabi	Live body weight (Kg)	107G > C + 138	A > C	GG/AA	1.853	3.432
	Estradiol (pg/ml)	107G > C + 138	A > C	GG/AA	1.830	3.348
	Live body weight (Kg)	107G > C + 138	A > C	GC/AC	1.873	3.507
	Estradiol (pg/ml)	107G > C + 138	A > C	GC/AC	1.850	3.421
	Live body weight (Kg)	107G > C + 138	A > C	CC/CC	1.923	3.695
	Estradiol (pg/ml)	107G > C + 138	A > C	CC/CC	1.911	3.650

Note: varP proportion of phenotypic variance explained. The percentage of variance was calculated only for the SNPs that showed a significant association with the traits.

ORCID

Tahreer M. Al-Thuwaini Phttps://orcid.org/ 0000-0003-4534-2963 Mohammed Baqur S. Al-Shuhaib Dhttps://orcid.org/ 0000-0002-6458-2068

REFERENCES

- Abdullah, A. Y., Husein, M. Q., & Kridli, R. T. (2002). Protocols for estrus synchronization in Awassi ewes under arid environmental conditions. Asian-Australasian Journal of Animal Sciences, 15(7), 957-962. https://doi.org/10.5713/ajas.2002.957
- Adzhubei, I. A., Schmidt, S., Peshkin, L., Ramensky, V. E., Gerasimova, A., Bork, P., Kondrashov, A. S., & Sunyaev, S. R. (2010). A method and server for predicting damaging missense mutations. Nature Methods, 7(4), 248. https://doi.org/10.1038/nmeth0410-248
- Aktas, A. H., Dursun, S., Doğan, S., Kiyma, Z., Demirci, U., & Halıcı, I. (2015). Effects of ewe live weight and age on reproductive performance, lamb growth, and survival in Central Anatolian Merino sheep. Archives Animal Breeding, 58(2), 451-459. https://doi.org/10.5194/ aab-58-451-2015
- Aljubouri, T. R. S., & Al-Shuhaib, M. B. S. (2020). Genotyping of mitochondrial D-loop sequences in three breeds of sheep. Biologia. https://doi. org/10.2478/s11756-020-00543-6
- Alkass, J. E., & Juma, K. H. (2005). Chapter 3: Small Ruminant breeds of Iraq. In L. Iniguez (Ed.), Characterization of Small Ruminant Breeds in

- West Asia and North Africa (Vol. 1, pp. 63-101). International Center for Agricultural Research in Dry Areas (ICARDA).
- Al-Sa'aidi, J. A. A., Khudair, K. K., & Khafaji, S. S. (2018). Reproductive fecundity of Iraqi Awassi ewes immunized against synthetic inhibin- α subunit or steroid-free bovine follicular fluid. Asian-Australasian Journal of Animal Sciences, 31(8), 1169. https://doi.org/10.5713/ ajas.17.0660
- Al-Shuhaib, M. B. S. (2017). A universal, rapid, and inexpensive method for genomic DNA isolation from the whole blood of mammals and birds. Journal of Genetics, 96(1), 171-176. https://doi.org/10.1007/ s12041-017-0750-6
- Al-Shuhaib, M. B. S., Al-Kafajy, F. R., Badi, M. A., AbdulAzeez, S., Marimuthu, K., Al-Juhaishi, H. A. I., & Borgio, J. F. (2018). Highly deleterious variations in COX1, CYTB, SCG5, FK2, PRL and PGF genes are the potential adaptation of the immigrated African ostrich population. Computers in Biology and Medicine, 100, 17-26. https://doi. org/10.1016/j.compbiomed.2018.06.019
- Al-Shuhaib, M. B. S., Al-Thuwaini, T. M., Fadhil, I. A., & Aliobouri, T. R. S. (2019). GHRL gene-based genotyping of ovine and caprine breeds reveals highly polymorphic intronic sequences in Awassi sheep with several RNA motifs. Journal of Genetic Engineering and Biotechnology, 17, 3. https://doi.org/10.1186/s43141-019-0004-5
- Al-Thuwaini, T. M., Al-Shuhaib, M. B. S., & Hussein, Z. M. (2020). A highly deleterious novel T177P missense variant in the HSPA8 gene associated with the low tolerance of Awassi sheep to heat stress. Tropical Animal Health and Production. Ahead of print. https://doi. org/10.1007/s11250-020-02267-w

- Bakos, J., Zatkova, M., Bacova, Z., & Ostatnikova, D. (2016). The role of hypothalamic neuropeptides in neurogenesis and neuritogenesis. *Neural Plasticity*, 2016, 3276383. https://doi. org/10.1155/2016/3276383
- Byun, S. O., Fang, Q., Zhou, H., & Hickford, J. G. H. (2009). An effective method for silver-staining DNA in large numbers of polyacrylamide gels. *Analytical Biochemistry*, 385(1), 174–175. https://doi.org/10.1016/j.ab.2008.10.024
- Cai, X., Mipam, T. D., Zhao, F. F., & Sun, L. (2015). SNPs detected in the yak MC4R gene and their association with growth traits. *Animal*, *9*(7), 1097–1103. https://doi.org/10.1017/S1751731115000397
- Capriotti, E., Calabrese, R., & Casadio, R. (2006). Predicting the insurgence of human genetic diseases associated to single point protein mutations with support vector machines and evolutionary information. *Bioinformatics*, 22(22), 2729–2734. https://doi.org/10.1093/bioinformatics/btl423
- Chen, X., Huang, L., Tan, H. Y., Li, H., Wan, Y., Cowley, M., Veldhuis, J. D., & Chen, C. (2017). Deficient melanocortin-4 receptor causes abnormal reproductive neuroendocrine profile in female mice. *Reproduction*, 153(3), 267–276. https://doi.org/10.1530/REP-16-0341
- Choi, Y., Sims, G. E., Murphy, S., Miller, J. R., & Chan, A. P. (2012). Predicting the functional effect of amino acid substitutions and indels. *PLoS One*, 7(10), e46688. https://doi.org/10.1371/journal.pone.0046688
- Clegg, D. J., Brown, L. M., Woods, S. C., & Benoit, S. C. (2006). Gonadal hormones determine sensitivity to central leptin and insulin. *Diabetes*, 55(4), 978–987. https://doi.org/10.2337/diabetes.55.04.06. db05-1339
- Deng, T. X., Pang, C. Y., Liu, M. Q., Zhang, C., & Liang, X. W. (2016). Synonymous single nucleotide polymorphisms in the MC4R gene that are significantly associated with milk production traits in water buffaloes. Genetics and Molecular Research, 15, 1–8. https://doi. org/10.4238/gmr.15028153
- El-Sabrout, K. (2017). Associations between single-nucleotide polymorphisms of melanocortin gene and sexual desire behavior in rabbit (*Oryctolagus cuniculus*). *Journal of Veterinary Behavior*, 19, 69–71. https://doi.org/10.1016/j.jveb.2017.02.005
- El-Sabrout, K., & Aggag, S. (2018). Association of Melanocortin (MC4R) and Myostatin (MSTN) genes with carcass quality in rabbit. *Meat Science*, 137, 67–70. https://doi.org/10.1016/j.meatsci.2017.11.008
- El-Sabrout, K., & Soliman, F. (2018). Association of single-nucleotide polymorphism of melanocortin gene with feed intake in rabbit (Oryctolagus cuniculus). Journal of Animal Physiology and Animal Nutrition, 102(2), 564–567. https://doi.org/10.1111/jpn.12788
- Esmailizadeh, A. K. (2010). A partial genome scan to identify quantitative trait loci affecting birthweight in Kermani sheep. *Small Ruminant Research*, 94(1–3), 73–78. https://doi.org/10.1016/j.small rumres.2010.07.003
- Fan, B., Onteru, S. K., Plastow, G. S., & Rothschild, M. F. (2009). Detailed characterization of the porcine MC4R gene in relation to fatness and growth. *Animal Genetics*, 40(4), 401–409. https://doi. org/10.1111/j.1365-2052.2009.01853.x
- McGlone, J. (2010). Guide for the care and use of agricultural animals in research and teaching. (3rd ed.).Champaign, IL: Federation of Animal Science Societies.
- Galal, S., Gürsoy, O., & Shaat, I. (2008). Awassi sheep as a genetic resource and efforts for their genetic improvement—A review. Small Ruminant Research, 79(2–3), 99–108. https://doi.org/10.1016/j.small rumres.2008.07.018
- Hashim, H. O., & Al-Shuhaib, M. B. S. (2019). Exploring the potential and limitations of PCR-RFLP and PCR-SSCP for SNP detection: A review. *Journal of Applied Biotechnology Reports*, 6(4), 137–144. https://doi. org/10.29252/JABR.06.04.02
- Hewagalamulage, S. D., Clarke, I. J., Young, I. R., Rao, A., & Henry, B. A. (2015). High cortisol response to adrenocorticotrophic hormone identifies ewes with reduced melanocortin signalling and increased

- propensity to obesity. *Journal of Neuroendocrinology*, 27(1), 44–56. https://doi.org/10.1111/jne.12233
- Hill, W. G., & Mackay, T. F. (2004). DS Falconer and Introduction to quantitative genetics. *Genetics*, 167(4), 1529–1536.
- Hwa, J. J., Ghibaudi, L., Gao, J., & Parker, E. M. (2001). Central melanocortin system modulates energy intake and expenditure of obese and lean Zucker rats. American Journal of Physiology-Regulatory, Integrative and Comparative Physiology, 281(2), R444–R451. https://doi.org/10.1152/ajpregu.2001.281.2.R444
- Källberg, M., Wang, H., Wang, S., Peng, J., Wang, Z., Lu, H., & Xu, J. (2012). Template-based protein structure modeling using the RaptorX web server. *Nature Protocols*, 7(8), 1511. https://doi.org/10.1007/978-1-4939-0366-5_2
- Kratochvílová, M., Hyánková, L., Knížetová, H., Fiedler, J., & Urban, F. (2002). Growth curve analysis in cattle from early maturity and mature body size viewpoint. Czech Journal of Animal Science, 47(4), 125–132
- Latifah, L., Maharani, D., Kustantinah, A., & Hartatik, T. (2018). Association of melanocortin 4 receptor gene polymorphism with growth traits in Bligon goat. *Journal of the Indonesian Tropical Animal Agriculture*, 43(4), 343–351. https://doi.org/10.14710/jitaa.43.4.343-351
- Lee, Y.-S., Park, S., Kim, H., Lee, S.-K., Kim, J.-W., Lee, H.-K., Jeong, D.-K., & Lee, S.-J. (2013). A C1069G SNP of the MC4R gene and its association with economic traits in Korean native cattle (brown, brindle, and black). Electronic Journal of Biotechnology, 16(5), 14. https://doi.org/10.2225/vol16-issue5-fulltext-5
- Li, C. Y., & Li, H. (2006). Association of MC4R gene polymorphisms with growth and body composition traits in chicken. Asian-Australasian Journal of Animal Sciences, 19(6), 763–768. https://doi.org/10.2225/vol16-issue5-fulltext-5
- Liu, H., Tian, W., Zan, L., Wang, H., & Cui, H. (2010). Mutations of MC4R gene and its association with economic traits in Qinchuan cattle. Molecular Biology Reports, 37(1), 535–540. https://doi.org/10.1007/s11033-009-9706-0
- Lu, X., Niu, T., & Liu, J. S. (2003). Haplotype information and linkage disequilibrium mapping for single nucleotide polymorphisms. *Genome Research*, 13, 2112–2117. https://doi.org/10.1101/gr.586803
- Marai, I. F. M., El-Darawany, A. A., Fadiel, A., & Abdel-Hafez, M. A. M. (2007). Physiological traits as affected by heat stress in sheep – A review. Small Ruminant Research, 71, 1–12. https://doi.org/10.1016/j. smallrumres.2006.10.003
- Ng, P. C., & Henikoff, S. (2003). SIFT: Predicting amino acid changes that affect protein function. *Nucleic Acids Research*, 31(13), 3812–3814. https://doi.org/10.1093/nar/gkg509
- Niyas, P. A., Chaidanya, K., Shaji, S., Sejian, V., Bhatta, R., Bagtah, M., & Girish, V. (2015). Adaptation of livestock to environmental challenges. *Journal of Veterinary Science & Medical Diagnostics*, 4, 3. https://doi.org/10.4172/2325-9590.1000162
- Ott, J. (2001). Program Het version 1.8. Utility programs for analysis of genetic linkage. Rockefeller University.
- Prihandini, P. W., & Maharani, D. (2019). Melanocortin-4 receptor (MC4R) gene as the main gene for rapid growth selection in beef cattle. Indonesian Bulletin of Animal and Veterinary Sciences, 29(2), 85–95. https://doi.org/10.14334/wartazoa.v29i2.1955
- Seong, J., Suh, D. S., Do Park, K., Lee, H. K., & Kong, H. S. (2012). Identification and analysis of MC4R polymorphisms and their association with economic traits of Korean cattle (Hanwoo). *Molecular Biology Reports*, 39(4), 3597–3601. https://doi.org/10.1007/s1103 3-011-1133-3
- She, Y. Y., & He, S. L. (2006). SHEsis, a powerful software platform for analyses of linkage disequilibrium, haplotype construction, and genetic association at polymorphism loci. *Cell Research*, 16, 851. https:// doi.org/10.1038/sj.cr.7290272
- Shishay, G., Liu, G., Jiang, X., Yu, Y., Teketay, W., Du, D., Jing, H., & Liu, C. (2019). Variation in the promoter region of the MC4R gene

- Siljee, J. E., Unmehopa, U. A., Kalsbeek, A., Swaab, D. F., Fliers, E., & Alkemade, A. (2013). Melanocortin 4 receptor distribution in the human hypothalamus. *European Journal of Endocrinology*, 168(3), 361–369. https://doi.org/10.1530/EJE-12-0750
- Switonski, M., Mankowska, M., & Salamon, S. (2013). Family of melanocortin receptor (*MCR*) genes in mammals—mutations, polymorphisms and phenotypic effects. *Journal of Applied Genetics*, *54*(4), 461–472. https://doi.org/10.1007/s13353-013-0163-z
- Tang, H., & Thomas, P. D. (2016). PANTHER-PSEP: Predicting disease-causing genetic variants using position-specific evolutionary preservation. *Bioinformatics*, 32(14), 2230–2232. https://doi.org/10.1093/bioinformatics/btw222
- Trakovická, A., Moravčíková, N., & Kasarda, R. (2013). Genetic polymorphisms of leptin and leptin receptor genes in relation with production and reproduction traits in cattle. *Acta Biochimica Polonica*, 60(4), 783–787.
- Vaisse, C., Clement, K., Durand, E., Hercberg, S., Guy-Grand, B., & Froguel, P. (2000). Melanocortin-4 receptor mutations are a frequent and heterogeneous cause of morbid obesity. *The Journal of Clinical Investigation*, 106(2), 253–262. https://doi.org/10.1172/JCI9238
- Ye, J., Coulouris, G., Zaretskaya, I., Cutcutache, I., Rozen, S., & Madden, T. L. (2012). Primer-BLAST: A tool to design target-specific primers for polymerase chain reaction. *BMC Bioinformatics*, 13(1), 134. https:// doi.org/10.1186/1471-2105-13-134

- Yeh, F., & Yang, R. C. (1999). POPGENE 32v. 1.31 Microsoft Window-based Freeware for Population Genetic Analysis. University of Alberta and Tim Boyle, Centre for International Forestry Research.
- Zandi, M. R., Shirazi, M. R. J., Tamadon, A., Akhlaghi, A., Salehi, M. S., Niazi, A., & Moghadam, A. (2014). Hypothalamic expression of melanocortin-4 receptor and agouti-related peptide mRNAs during the estrous cycle of rats. International Journal of Molecular and Cellular Medicine, 3(3), 183.
- Zeng, R., Zhang, Y., & Du, P. (2014). The SNPs of melanocortin 4 receptor (MC4R) associated with body weight in beagle dogs. *Experimental Animals*, 63(1), 73–78. https://doi.org/10.1016/j.pepti des.2005.02.029
- Zuo, B., Liu, G., Peng, Y., Qian, H., Liu, J., Jiang, X., & Mara, A. (2014). Melanocortin-4 receptor (MC4R) polymorphisms are associated with growth and meat quality traits in sheep. *Molecular Biology Reports*, 41(10), 6967–6974. https://doi.org/10.1007/s1103 3-014-3583-x

How to cite this article: Al-Thuwaini TM, Al-Shuhaib MBS, Lepretre F, Dawud HH. Two co-inherited novel SNPs in the *MC4R* gene related to live body weight and hormonal assays in Awassi and Arabi sheep breeds of Iraq. *Vet Med Sci.* 2021;7:897–907. https://doi.org/10.1002/vms3.421