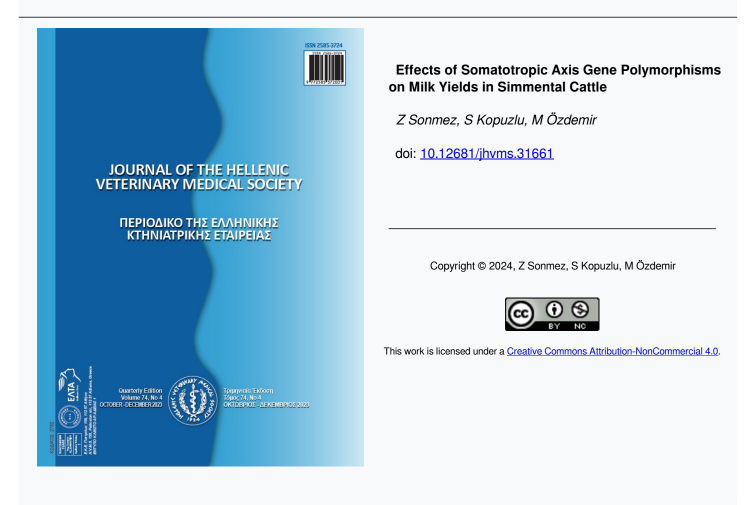




Journal of the Hellenic Veterinary Medical Society

Vol 74, No 4 (2023)



To cite this article:

Sonmez, Z., Kopuzlu, S., & Özdemir, M. (2024). Effects of Somatotropic Axis Gene Polymorphisms on Milk Yields in Simmental Cattle. *Journal of the Hellenic Veterinary Medical Society*, *74*(4), 6589–6598. https://doi.org/10.12681/jhvms.31661

The Effects of Somatotropic Axis Gene Polymorphisms on Milk Yields in Simmental Cattle

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ABSTRACT: Due to the increasing world population, scientists aim to obtain high-yield products using new techniques and methods to meet needs in the fields of food, agriculture, and livestock. It is very important to select the candidate genes and markers correctly, especially in the QTL and MAS techniques in livestock. Somatotropic axis genes affect yield traits, growth, reproduction, and milk production in cattle. In the study, we determined GH/AluI and IGFI/SnaBI gene polymorphisms using the PCR-RFLP technique in DNA samples obtained from 70 Simmental cattle. The allele frequencies of the genes were in the same proportion as 0. 62 and 0. 38, L and V for the GH gene and T and C for the IGFI gene. There was no significant relationship between the polymorphic genotypes of the genes we examined and lactation milk yields, 305-day milk yields, daily milk yields, and lactation periods of Simmental cattle. The probability of using bGH/AluI and IGFI/SnaBI genetic variations as markers in association studies with milk yield in cattle is extremely low.

Keywords: Axis genes; RFLP; MAS; Simmental; milk yield

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Date of initial submission: 27-10-2022 Date of acceptance: 17-02-2023

INTRODUCTION

Towadays, genome sequencing of organisms and the development of comprehensive single-nucleotide polymorphism (SNP) data, along with advancements in genotype identification technologies, have made genetic association maps in the entire genome possible. Somatotropic axis genes, consisting mainly of growth hormone (GH), insulin-like growth factors (IGF I and II), carrier proteins and receptors associated with them (GHR, IGFR, IGFBP), take part in metabolism and physiological processes. They also play a role in many physiological processes, including the regulation of ionic and osmotic balance, the regulation of lipid, carbohydrate and protein metabolisms, growth, reproduction, immune function and behaviour (Renaville et al., 2002; Gabillard et al., 2003; Besseau et al., 2013; Hax et al., 2017).

The IGF I hormone consisting of 70-amino-acid with a weight of 7649 kilodaltons connected by disulfide bridges in the form of A and B chains. The cleavage sites of the SnaBI restriction enzyme, recognized by Ge et al. (2001), the C>T cleavage site at base 512 in the regulatory region of the IGFI gene, have been studied extensively. The IGF-I hormone is functionally separated since it is released by two different tissues. First, it is released by the liver and acts as an extension of the GH axis through the tonic pituitary stimulation of hepatic synthesis (Oster et al., 1995). Second, the GH hormone determines the level of IGF-I in circulation along with nutritional factors (Laron, 2001; Delafontaine et al., 2004).

The bovine growth hormone (bGH) represents a single-chain polypeptide hormone weighing 22 kDa produced in the pre-pituitary gland and comprising 191 amino acids. Studies on bGH polymorphismshave intensely determined L (leucine at codon 127) and V (valine at codon 127) allelic polymorphisms located on the 5th exon and coinciding with the recognition region of the AluI (GH-AluI) restriction enzyme (Zhang et al., 1993). The hormone bGH is known to play an essential role in biological processes, including breast development, lactation, ageing, growth and regulation of metabolism, ovulation in female individuals, and control of sexual behaviours (Ge et al., 2003). Polymorphisms in the GH gene are known to correlate with different production yields, such as milk yield and quality, growth (Pal et al., 2005; Kovacs et al., 2006; Zhou et al., 2006), carcass composition and quality (Grochowska et al., 2001). Furthermore, this genealso playsan important role in postpartum growth and overall metabolism, as well as lactationin cattle (Fedota et al., 2016).

Studies on various cattle breeds have generally reported the L allele in the GH/Alu gene polymorphism and the B (T) allele in the IGFI/SnaBI gene polymorphism at higher frequencies. Tables 1 and 2 present some studies on the GH/AluI and IGFI/SnaBIgenes.

Breeds	Ν	Allele fr	equencies	References
IGF-I/SnaBI		Α	В	
South Anatolian Red (SAR)	50	0.23*	0.77*	Akis et al., 2010
East Anatolian Red (EAR)	50	0.38*	0.62*	Akis et al., 2010
Charolais (Coahuila)	68	0.26	0.74	Reyna et al., 2010
Charolais (Nuevo León)	68	0.46*	0.54*	Reyna et al., 2010
Beefmaster (Tamaulipas)	25	0.03	0.97	Reyna et al., 2010
Angus	275	0.40	0.60	Rogberg-Muñoz et al., 2013
Charolais	238	0.47	0.53	Jeanmaset al., 2014
Black Hereford	36	0.48	0.52	Dașet al., 2019
Limousin	34	0.42	0.58	Dașet al., 2019
Brahman	24	0.34	0.66	Daș et al., 2019
Holstein	250	0.46*	0.54 *	Bonakdar et al., 2010
IGF-1/SnaBI (Eco105I)		Т	С	
Montbeliarde	163	0.67*	0.33*	Szewczuk, 2016
Necdi	84	0.10	0.90	Yazdanpanah et al., 2013
Pesisir cattle	183	0.02	0.98	Putra et al., 2017
Jersey	227	0.52*	0.48 *	Piątkowska et al., 2021
Polish Holstein	147	0.56*	0,44 *	Piątkowska et al., 2021a
Polish Holstein	181	0.52*	0.48 *	Piątkowska et al., 2021b

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Holstein–Friesian bulls	296	0.44 *	0.56 *	Ardicli et al., 2018
Kazakhstan Holstein	100	0.58*	0.42*	Beishova et al., 2019
Holstein	95	0.59*	0.41 *	Silveira et al., 2019
Native Sudanese Baggaracattle (Nyalawi)	64	0.93	0.07	Omer et al., 2018
Native Sudanese Baggaracattle (Mesairi)	63	0.92	0.08	Omer et al., 2018
Holstein-Friesian	70	0.59*	0.41 *	Nicolini et al., 2013
Iranian Holstein	60	0.40*	0.60*	Ararouti et al., 2013
Polish Holstein	191	0.54*	0.46 *	Szewczuk et al., 2013
Holstein	381	0.4(-/-)	0.6(+/+)	Hax et al., 2017
Kazakhstan black and white tawny	99	0.55*	0.45*	Ulyanov et al., 2021

Breeds	N	Allele f	requencies	References	
bGH		L	V		
Native Antioquia (Colombia)	178	0.87	0.13	Echeverri et al., 2015	
Italian Holstein	695	0.86	0.14	Fontanesi et al., 2015	
Grey steppe	76	0.79	0.21	Özdemir, 2011	
Holstein	177	0.90	0.10	Özdemir, 2011	
Brown Swiss	177	0.90	0.10	Özdemir, 2011	
East Anatolian Red	51	0.95	0.05	Özdemir, 2011	
East Anatolian Red	76	0.84	0.16	Özkan et al., 2009	
South Eastern Red	180	0.44*	0.56*	Yardibiet al., 2009	
East Anatolian Red	76	0.84	0.16	Özkan et al., 2009	
East Anatolian Red	50	0.43*	0.57*	Yardibiet al., 2009	
Holstein	410	0.87	0.13	Amiri et al., 2018	
Holstein Friesian	578	0.87	0.13	Pal et al., 2020	
Kazakhstan Holstein	100	0.82	0.18	Beishova et al., 2019	
Pesisir cattle	175	0.79	0.21	Afriani, 2018	
Holstein	158	0.90	0.10	Gilmanov et al., 2021	
Holstein's calves	59	0.96	0.04	Çınar et al., 2018	
Slovak spotted ows	110	0.68*	0.32*	Hazuchova et al., 2013	
Egyptian water buffaloes	200	0.80	0.20	El-Komy et al., 2021	
Slovak spotted black-and-white cows	58	0.64*	0.36*	Moravcikova et al., 2012	
Limousin	114	0.71,	0.29	Sedykh et al., 2020	
Hereford	115	0.69*	0.31*	Sedykh et al., 2020	
Kazakhstan black and white tawny	99	0.7	0.3	Ulyanov et al., 2021	
Holstein	186	0.82	0.18	Özdemir et al., 2015	
Holstein	115	0.65*	0.35*	Sönmez et al., 2018	

Our study aimed to identify the genotypes of somatotropic axis genes in 70 heads of Simmental cattle by the RFLP method and correlate the genotype polymorphisms of the relevant gene regions with milk yield traits

MATERIALS AND METHODS

Animal care, Animal DNA isolation and Primer Sets

In the present study, we collected blood samples from 70heads of Simmental cattle with low degrees of relatedness for analysis. Animals were raisedon a private Simmental cattle farm in Erzurum province.

DNA was isolated from whole blood using the Purgene DNA kit following the manufacturer's instruction (Gentra Systems Minnesota USA). In the PCR step, for growth hormone (bGH)-AluI, the 245-bp DNA region, the primers F: 5'-GTA GGG GAG GGT GGAAAATG -3 and R:5' TGA CCCTCA GGT ACG TCT CC -3 (Özdemir, 2011), and for the 249-bp IGF1/ SnaBI gene, F: 5'-ATT-ACA-AAG-CTG-CCT-GCC-CC-3'and R: 5'-ACC-TTA-CCC-GTA-TGA-AAG-GAA-TAT-ACG Primers -T-3' (Ge et al., 2001) were used.

PCR conditions

Ten pmol/ μ L of each primer and 2. 5 μ L of dNTP-

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mix (D7595: Sigma, St. Louis, MO, USA, 0. 25mM), 0. 5 units of Taq DNA polymerase (D1806: Sigma), 100 ng of template DNA, 5 μ L of 10x PCR Buffer (100 mM Tris-HCl, pH 8. 3, 500 mM KCl, 15 mM MgCl₂ and 0. 01% gelatin), 1. 5 μ L of 0. 25 mM MgCl₂, andddH₂O making the total volume of 30 μ L were used for PCR amplification.

PCR amplification conditions for GH were as follows: 5 min at 94°C, 50 s at 94°C, and 50 s at 58 °C;PCR amplification conditions for the IGFI gene were as follows: 5 min at 94°C, 30 s at 94°C, and 50 s at 61°C, and 30 cycles for both genes, with the final elongation temperature set at 72°C for 5 min and 1 cycle.

For the amplified IGFI and bGH genes, 2-5 U of the AluI and SnaBI restriction enzymes were added to each 7μ I PCR product and incubated at 37°C for 12 hours. After incubation, the samples were run on 3. 0% agarose gel at 45 Volts for 2. 5 hours, and bands were observed under UV light. The bGH and IGFI allele gene frequencies of each Simmental breed were calculated by counting. POPGENE software v1. 32 (Yeh et al., 1999) was utilized to reveal whether the genotype frequencies were in Hardy-Weinberg equilibrium. SPSS 21. 0 software program was used to analyze the association of genotypes with the population's milk data.

To research the impacts of IGFI and GH gene polymorphisms on lactation and some other milk yield traits in Simmental cattle raised on a private enterprise in Erzurum, a correlation analysis was conducted between the relevant polymorphic regions and the milk yield records of animals in different lactations, whose yield records were systematically kept between 2017 and 2020.

The statistical association analysis used milk yield traits, including real milk yield, 305-day milk yield, lactation period, and daily milk yield of the Simmental cattle breed. Data were analyzed by the SPSS 25. 0 statistics package program (IBM SPSS 25. 0 Corp. Inc.) based on the general linear model (Harvey, 1994). The analysis evaluated the effects of environmental factors, e. g., genotype, lactation order, and calving seasons, thought to affect the relevant yield trait.

According to the yield traits in the research, the following statistical model was employed.

Where

 Y_{ijk} is any of the milk yield traits (real milk yield, 305-day milk yield, lactation period, and daily milk yield),

 μ is the population average,

a, is the ith genotype effect,

b_j is the effect of the jth lactation order (j: 3; 1st Lactation: 1, 2nd Lactation: 2, 3rd Lactation: 3, 4rd Lactation,5 rd Lactation, 6rd Lactation,7rd Lactation),

 c_k is the effect of the kth calving season (k:2; 1: winter-spring, 2: summer-autumn),

e_{iii}is the margin of error.

RESULTS

The figures show the PCR amplification results and base sizes of PCR products (Figure 1).

The 249 bp PCR products were digested by SnaBI for the IGFI gene, and the result of digestion was one band (249 bp) for the CC genotype, three bands (249, 223, and 26 bp) for the TC genotype, and two bands (223 and 26 bp) for the TT genotype (Figure 2).

The 246 bp products were acquired after the PCR process, and PCR products were digested by the AluI restriction enzyme for the GH gene. At the end of digestion, one band (246 bp) for the VV genotype, three bands (246, 171, and 52 bp) for the LV genotype, and two bands (171 and 52 bp) for the LL genotype were expected. The band of 52 bp could not be observed on agarose gel electrophoresis (Figure 3).

As a result of determining allele and genotype frequencies in Simmental cattle, it was found that the LL, LV, and VV genotype frequencies for the GH gene were 0. 33, 0. 57, and 0. 10, respectively,while the TT, CT, and CC genotype frequencies for the IGFI gene were 0. 35, 0. 55, and 0. 10, respectively. The LL and VV allele frequencies for the GH gene and the TT and CC allele frequencies for the IGFI gene were measured similarly as 0. 62 and 0. 38 (Table 3).

Hardy-Weinberg genetic equilibrium test showed that the distributions of these genotype frequencies were not in equilibrium in the analysed breed (P > 0. 05).

 $Y_{ijk}: \mu + a_i + b_j + c_k + e_{ijk}$

The overall averages for milk yield, 305-day milk yield, daily milk yield,real milk yield,and lactation

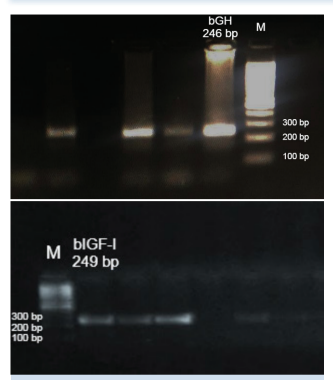


Figure 1. PCR results of the bGHand IGFI gene regions (1st primer pair 246, 2nd primer pair 249, with Standard GENESTA 100 bp DNA marker images, respectively)



Figure 2. IGFI-SnaBI RFLP results (CC, CT, and TT genotypes with Standard GENESTA100 bp DNA marker images, respectively)

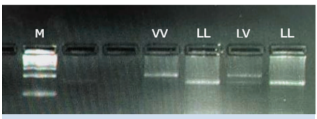


Figure 3. GH-AluI RFLP results (VV, LV, and LL genotypes with Standard GENESTA 100 bp DNA marker images)

IGFI allele and genotype frequencies GH allele and genotype frequencies											
Genotype	N	Т	С	% Frequency	H-W ^a	Genotype	Ν	L	V	% Frequency	H-W ^a
TT	23	46	0	34.85		LL	23	46	0	33.33	
CT	36	36	36	54.55	0.20	LV	39	39	39	56.52	0.11
CC	7	0	14	10.61		VV	7	0	14	10.14	
Total	66	82	50	100.00		Total	69	85	53	100.00	
р	132	0.62	0.38			р	138	0.62	0.38		

a. P-value for the test for Hardy-Weinberg equilibrium, in which P>0.05 indicates that the sampled population is not in Hardy-Weinberg equilibrium.

Table 4. Effects of IGFI/Snal	BI genotypes on mil	k yields in Simmental cattle
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Variation Sources		Ν	Daily milk yield $\overline{X} \pm S_{\overline{x}}$	Lactation milk yield $\overline{X} \pm S_{\overline{x}}$	305-day milk yield $\overline{X} \pm S_{\overline{x}}$	Lactation period $\overline{X} \pm S_{\overline{z}}$
IGF1/SnaBI	GF1/SnaBI CC		18,279±0,702	5468,285±283,960	5571,231±218,310	301,822±13,997
	CT	39	18,265±0,690	5085,252±279,220	5515,907±214,666	282,320±13,763
	TT	8	17,044±1,483	4916,054±600,087	5286,936±461,350	304,416±29,579
		Р	0,732	0,536	0,857	0,545
Calving season	1	48	17,742±0,786	4783,274±318,281	5228,148±244,696	278,548±15,688
-	2	35	17,984±0,809	5529,786±327,522	5687,901±251,801	313,824±16,144
		Р	0,817	0,081	0,160	0,094
Lactation order	2	10	17,026±1,439	4994,060±582,260	5272,492±447,645	297,624±28,700
	3	17	18,423±1,067	5300,083±432,062	5677,206±332,172	290,881±21,297
	4	18	17,422±1,085	5577,652±438,982	5493,044±337,492	332,631±21,638
	5	18	18,687±0,983	5294,563±397,867	5674,684±305,882	280,223±19,611
	6	14	17,440±1,165	5044,346±471,381	5327,560±362,400	298,212±23,235
	7	6	18,177±1,745	4728,477±706,436	5303,162±543,112	277,545±34,821
		Р	0,876	0,893	0,924	0,502
Total		83	17,863±0,605	5156,530±244,727	5458,025±188,148	296,186±12,063

 \overline{X} : General Means, $S_{\overline{x}}$: Standard deviation

period were 5458,025kg, 17,863 kg, 5156,530 kg, and 296,186days, respectively. Among the polymorphic regions, the highest average for all milk yield values was obtained with the CC genotype and the lowest value was obtained with the TT genotype for the IGFI/SnaBI polymorphic region, while the difference between the highest and lowest average milk yield values was not insignificant.

The differences between the genotype according to the GH/AluI fragment differences and the milk

yield averages of 91 Simmental cattle in 6 lactations and the genotype, calving season, and lactation order factors were statistically insignificant (p>0.05). The overall averages for milk yield, 305-day milk yield, daily milk yield,real milk yield,and lactation period were 5443,264 kg, 18,355 kg, 5619,733 kg, and 301,545days, respectively. Among the yield records, the highest genotype average was observed in the LL genotype, while the lowest yield average was observed in the LV genotype.

Table 5. Effects of GH/AluI genotypes on milk yields in Simmental cattle									
Variation Sources		Ν	Daily milk yield $\overline{X} \pm S_{\overline{x}}$	Lactation milk yield $\overline{X} \pm S_{\overline{x}}$	305-day milk yield $\overline{X} \pm S_{\overline{x}}$	Lactation period $\overline{\mathbf{X}} \pm \mathbf{S}_{\overline{\mathbf{x}}}$			
GH/AluI	LL	33	18,349±0,728	5541,241±292,225	5624,453±228,070	304,904±14,130			
	LV	47	18,167±0,623	5059,354±250,305	5499,343±195,353	282,277±12,103			
	VV	11	18,549±1,215	5729,198±487,804	5735,402±380,712	317,454±23,587			
		Р	0,954	0,290	0,824	0,273			
Calving period	1	54	18,447±0,672	5262,943±269,631	5508,863±210,436	289,717±13,037			
	2	37	18,262±0,742	5623,586±297,773	5730,602±232,400	313,374±14,398			
		Р	0,847	0,350	0,461	0,206			
Lactation order	2	10	17,306±1,332	5093,744±534,652	5297,029±417,275	295,945±25,852			
	3	17	18,875±1,040	5584,882±417,492	5817,002±325,836	297,047±20,187			
	4	20	$18,008 \pm 0,970$	$5689,066 \pm 389,565$	5602,114±304,040	326,819±18,836			
	5	21	18,983±0,965	5408,306±387,470	5738,397±302,405	283,487±18,735			
	6	16	17,388±±1,046	5304,829±419,796	5345,561±327,634	312,241±20,298			
	7	7	19,570±1,552	5578,758±623,270	5918,292±486,438	293,732±30,137			
		Р	0,681	0,940	0,794	0,600			
Total		91	18,355±0,522	5443,264±209,579	5619,733±163,568	301,545±10,134			

 \overline{X} : General Means, $S_{\overline{x}}$: Standard deviation

DISCUSSION

The study carried out in the Simmental breed because of LL and CC genotypes are seen at the same rate distributions so allele frequencies are similar for GH/ AluI and IGFI/SnaBI genes. Tables 1 and 2 present the allele and genotype frequencies obtained in previous polymorphic studies on the GH and IGFI genes in cattle. Among these studies, those that show similarities and differences with our studies are marked. The distributions of allele and genotype frequencies for both genes are generally similar to previous studies, and it is seen that the L allele is dominant for the GH hormone, while the T allele is dominant for the IGFI gene. According to the Hardy-Weinberg equilibrium test, the distributions of genotype frequencies in the herd were estimated to be in not balanced because the herd was raised in a closed barn environment, external migration was absent, and there was high mortality due to uncontrolled breeding.

yields of GH/AluI gene polymorphisms in 118 and 186 numbers of Holstein cattle breeds from different farms, no significant relationship was found between genotypes and milk yields. A similar relationship could not be found between GH-AluI and IGFI-Sna-BI gene polymorphisms and milk yield in Simmental cattle (Sönmez et al., 2018; Özdemir et al., 2018). Therefore, considering our three association studies, we examined previous studies to determine the potential for these gene regions to be used as markers.

In our study of associating the IGFI/SnaBI gene polymorphism with milk yield in 70 cattle breeds, we thought that the relevant polymorphic gene marker was not associated with milk yields may be due to a small number of samples. However, similar results were obtained in previous milk yield association studies on different breeds and with a large number of samples. Studies on milk yield generally report that the IGFI/SnaBI gene polymorphism does not have a significant effect. Similar to our study, in the study

In two different studies in which we correlated the

that examined the impact of the IGFI gene polymorphism on milk yield in three breeds (227 Jersey, 147 Polish Holstein-Friesian black-and-white (HO) and 181 Polish Holstein-Friesian red-and-white (RW)) in different lactations, the average milk yield was high in individuals with the CC genotype, but there was no significant difference between this average and other genotype averages (Piatkowska et al., 2021). The study in which the serum IGF-I concentration and reproductive performance of Holstein dairy cows were associated with IGF-I gene polymorphisms reported no significant relationship between milk production and body condition score and IGF-I genotypes (Silveira et al., 2019). In different studies conducted on a number of which is over 100Holstein cattle showed that the IGFI/SnaBI polymorphism did not significantly affect milk yield, but the fat and protein ratios in milk were high, especially in individuals with the AB (CT) genotype (Bonakdar et al., 2010; Zhang et al., 2018; Beishova et al., 2017; Beishova et al., 2018; Beishova et al., 2019). For the IGFI gene, Ge et al. (2001) reported that the c. 512 C>T polymorphic marker defined by studies on milk yield in cattle breeds was generally not associated with milk yield. However, it was reported that different SNPs identified for the IGFI gene were associated with milk yield (Mullen et al., 2011;Lynch et al., 2010;Gui et al., 2018). The IGFI-SnaBI polymorphism studies conducted in different cattle breeds reported that the CC genotype was positively associated with the meat and fat weight of the carcass (Ge et al., 2001; Reyna et al., 2010; Siadkowska et al., 2006). Especially among studies on meat yield, studies are showing that the IGFI gene affects meat quality parameters and meat colour change, positively associated with live body weight at slaughter, cold carcass weight, meat and fat weight in valuable cuts, body weight, subcutaneous back fat, and waist circumference musculature (Curi et al., 2005; Siadkowska et al., 2006; Ardicli, 2018; Szewczuk et al., 2013). In the study determining the effects of the IGFI gene polymorphism on the reproductive data of different cattle breeds, it was revealed that the genotypes defined with IGF-SnaBI were effective on reproductive data but did not affect milk yield. Studies on reproductive data associated the SNP identified in the IGF-1 gene with the resumption of the postpartum lying phase and affected reproductive parameters in Holstein cows (Nicolini et al., 2013; Ararouti et al., 2013). In another study, the SNP within the GH/IGF-1 pathways was associated with the reproductive performance of Holstein cows under

heat stress conditions and associated with services per conception (SPC) (Leyva-Corona et al., 2018). While most studies have reported that the IGFI gene affects reproductive data, some studies have reported no significant relationship between IGF-I SnaBI genotypes in Holstein cattle kept under different breeding conditions and the fertility of Holstein cows raised in semi-intensive or intensive regimens (Hax et al., 2017).

Studies have demonstrated that the GH/AluI gene polymorphism, like the IGFI/SnaBI gene polymorphism is more effective on growth data and body weight than milk yield, especially in cattle. Similar to our study among the association studies on the GH hormone, a study conducted on black and white calico heifers reported that the amount of milk and the protein and fat ratios in milk were high in LL genotype heifers (Gilmanov et al., 2021) but the difference between the highest and lowest average milk yield values was not significant. In the studies GH-AluI gene polymorphism was determined in different breeds and the relationships between the obtained genotype data and body weight gain and body measurements were investigated genotype data showed that the GH-AluI polymorphism was related to the weaning body weights and the length of the chest circumference (Çınar et al., 2018; Afriani and Putra, 2018). In the association analyses conducted milk production length according to GH-AluI genotypes was not found significant, however, other SNPs identified in the GH gene were found to be associated with milk yields. (Zhou et al., 2006; Maylinda, 2011; Moravcikova et al., 2012;Hazuchova et al., 2013). Studies conducted on different cattle types and buffaloes have reported that GH-AluI gene polymorphisms affect meat yield, pre-slaughter body weight, carcass meat weight, slaughter weight and growth data (Pereira et al., 2005; Curi et al., 2006; Sedykh et al., 2020; El-Komy et al., 2021)

Parallel to our research, polymorphic association studies analyzing somatotropic axis genes together have generally associated these genes with growth and reproduction data but not with milk yield, especially GH/AluI and IGFI/SnaBI (Balogh et al., 2008; Ulyanov et al., 2021). The two different META analysis studies show that the common L allele was dominant and no significant difference was found in the results of the meta-analysis and subgroup analysis for the milk yields in the LL, LV, and VV genotypes Akcay et al., 2020; Bangar et al., 2021). The results of the meta-analysis on bGH/AluI and IGFI/SnaBI genetic variations in cattle confirm the results of our study.

In association analysis studies published to determine the potential of the gene polymorphisms to be used as markers in cattle breeding, the above-mentioned genes appear to have significant implications for body weight, meat yield, and reproductive performance in general. Studies to be conducted in the field of animal breeding estimate the effects of the determined bGH/AluI and IGFI/SnaBI genetic variations on different breeds with different performance traits and their potential to be used as markers in breeding programs. However, previous studies and our work have shown that bGH/AluI and IGFI/SnaBI genetic variations in cattle are less likely to be used as markers in milk data analyses.

ACKNOWLEDGMENTS

This study was not supported by any foundation. The study was carried out in the Zootechnie Genetic Laboratories of Atatürk University, therefore, we thank the Zootechnie Genetic Laboratory manager.

CONFLICTS OF INTEREST

The Authors have no conflict of interest to declare

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