ABSTRACT
Bayes Cp methodology due to the simultaneous use of all SNPs in a random regression model is an alternative to overcome the overestimation of SNPs effects and excess of false positives. Hence, the present study was performed to identify causal single nucleotide polymorphisms (SNPs) associated with body weight (BW) and their pattern of significance over time from 2 to 8 weeks of age, in an F2 crossbred chicken population using Bayes Cp methodology. Illumina 60K SNP bead chip was used to genotype 312 F2 chicks. Sixteen SNPs distributed over 8 chromosomes had a Bayes Factor (BF) greater than 150 for BW in different ages. 14 different genes harbouring these SNPs (±250 kb), of which 12 were protein-encoding (DKK2, LEF1, RASGEF1C, FOXP1, LPCAT1, AKR1D1, GPR137C, EIF2AK3, ROR1, AKR1D1, LRFN5 and XPO7) and two were noncoding RNAs (LOC101751953 and LOC101751953). According to the obtained results, GGA1 and GGA4 were the most important chromosomes containing QTLs associated with observed traits. Different genetic regions were responsible for body weight in early and late ages. The various number of significant Bayes factors across different weeks of ages may indicate the artificial selection for market weight in a given age which has driven larger SNPs effects in intended weeks for desired market weight for the two groups.

HIGHLIGHTS
My research outputs resulted in discovering associated genes and pathways related to gain growth and body weight in chicken. The identified genes in this study can be used in poultry breeding programs. Therefore, this leads to reduced feed costs and increase revenue.

Introduction
Body weight is a complex quantitative trait that is influenced by a large number of genes, each with different sizes and types (Goddard and Hayes 2009). As body weight is the most important trait in broilers especially for a specific stage of life, therefore identification of causative genetic loci for this trait can be essential to enhance genetic improvement by marker-assisted selection (Goddard and Hayes 2009).

The combination of natural and artificial selection for production traits over time in chicken can leave selection signatures in the genome (Walugembe et al. 2018; Weng et al. 2020). Artificial selection has been proved to influence significantly on the genomic loci associated with specific traits of an organism. Also, the relationship among the artificial selection, genomic regions, and interesting traits represented that recent artificial selection can increase runs of homozygosities at certain genetic markers and provide insight into the process of artificial selection (Kim et al. 2018).

The availability of dense marker genotypes for farm animals in recent years has made it possible to associate genetic markers such as single nucleotide polymorphisms (SNPs) with phenotypes of quantitative traits using genome-wide association study (GWAS). Generally, GWAS intends to detect significant SNPs to identify QTLs or causal genes for an interesting trait and alternatively enables to estimate the SNP-based heritability of the trait (Maher 2008). GWASs have firstly conducted based on the least-squares methodology (Costa et al. 2015), which has several drawbacks. Currently, GWAS studies sufficiently improved by using variance component model (Kang et al. 2010) and mixed linear models controlling simultaneously the population structure and identical by descent issues.
between genotyped individuals for achievement of more accurate signals (Zhang et al. 2010).

First of all, SNPs enter the analysis individually so that it may exceed the false positive and overestimation of SNP effects due to multiple testing (Peters et al. 2012). Choosing the right correction to avoid problems due to multiple testing is still a challenge (Wakefield 2009). An alternative to overcome these problems is to apply all SNPs simultaneously in a random regression model (Costa et al. 2015). Since the quantitative traits are influenced by a large number of SNPs, the model that analyzes all markers simultaneously might be more accurate in comparison with models that analyzes markers individually (van den Berg et al. 2013).

Additionally, the F2 population designed for GWAS is more advantageous than a random population in reducing false positives and improving the accuracy of mapping (Ledur et al. 2010). Given that in most animal breeding studies the importance of statistical methods was considered and Bayesian methods based on Gibbs sampling techniques have been concerned. Although there is a published study on the present data using a generalised linear model (GLM) and compressed mixed linear model (cMLM) (Emrani et al. 2017), we hypothesised that there would be differences in the size and locations of important regions if using other statistical methods such as Bayesian methodologies. Alongside, SNP-based GWAS provides a useful framework with which to carry out genomic analyses. So, the goal of this study was conducted to test the above hypothesis to identify SNPs surrounding the genes associated with body weight traits through SNP-based GWAS using the Bayes Cpi method.

Material and methods

Phenotypic data

Phenotypic data used in this study was collected from an F2 generation descended from reciprocal crosses between a commercial broiler line named Arian (AA) and an Iranian indigenous chicken (NN) named Urmia. These birds are bred as a dual-purpose breed (eggs and meat production). Arian line chickens are a commercial fast-growing broiler strain and Urmia population is a slow-growing indigenous strain in Iran. Arian chickens have been under strong selection for body weight and meat quality traits. Otherwise, the Urmia chicken is an indigenous slow growth population under natural selection for natural conditions and more resistant to diseases and pathogens than Arian chickens. The probable genomic distance between these two led us to use the benefits of the F2 generation for association studies based on LD patterns by generating an F2 population through reciprocal crosses between Arian and Urmia chickens. Each of F1 males was crossed to 4–8 females and finally, 450 F2 birds produced in six different hatches, 52 full-sib and 8 half-sib families. Birds were housed individually in battery cages and body weight of all birds recorded weekly from 2 to 8 weeks of age. On day 8, birds were moved to individual cages with a temperature of 30°C and gradually decreased to reach a final temperature of 22°C. Chickens were fed starter diet (22.8% CP and 3025 kcal of ME/kg) for the first two weeks, a crumbled grower diet (20% CP and 2960 kcal of ME/kg) between 3 and 7 weeks; and a crumbled finisher diet (18% CP and 3070 kcal of ME/kg) on the week 8 of age. Chickens were not vaccinated against any diseases. Feed and water were provided ad libitum (Begli et al. 2017).

To increase the accuracy of measurements, 8 hours of fasting were applied to birds before each weighing.

Genotyping and quality control

A random collection of 312 birds were Genotyped using blood samples using Illumina 60 k SNP chip (Illumina Inc., San Diego, CA) (Begli et al. 2017). Each sample was genotyped for 54,340 SNP markers. Data quality controls performed by PLINK 1.9 software. SNPs with missing genotype greater than 10% and minor allele frequency (MAF) smaller than 5% as well as SNPs with \( p \)-value less than \( 1 \times 10^{-6} \) in Hardy-Weinberg test was removed from the analysis (Begli et al. 2017). The total number of SNPs and population used for analysis in each age are shown in Table 1.

Statistical analysis

SNP effects estimated using the Bayes \( C_p \) methodology. This method splits the SNPs into two categories. One category consists of the SNPs having effects and the other category for non-effective SNPs on the phenotypic trait (Habier et al. 2011). Starting values for genetic variance

Table 1. Summary of the phenotypic and genotypic data used for the analysis in different weeks of ages.

| Birds age (week) | Number of birds | Number of SNPs |
|------------------|-----------------|----------------|
| 2                | 294             | 43588          |
| 3                | 292             | 43569          |
| 4                | 305             | 43605          |
| 5                | 304             | 43588          |
| 6                | 299             | 43577          |
| 7                | 307             | 43622          |
| 8                | 301             | 43577          |
and the variance of residual were calculated by DMU software (Madsen and Jensen, 2002) and Gibbs sampling was conducted with Markov Chain Monte Carlo (MCMC) technique. The number of iterations to reach the convergence set to be 100,000 iterations. A number of 20,000 burn-in and a sampling interval of 100 used to calculate the average posterior probability of the parameters. To reach a stable posterior distribution in the Bayesian analysis, we start from a starting point which is called prior parameter which is not informative enough based on the data in MCMC chain. Therefore, there is some fluctuation at the beginning of the chain to estimate the posterior parameter and the posterior distribution. Usually, this fluctuation period discarded from final calculation of the parameters (genetic variance) which called burn-in phase. After this phase the fluctuation becomes more stable around a value that we call it convergence of the parameter. The convergence diagnostic of Markov chain posterior samples (results) was conducted using trace plot approach by plotting the posterior density function of additive and residual variances which reflect chains mixed satisfactorily.

A mixed regression model was used for data analysis. This model contains sex and hatch effects as fixed systematic factors and animal polygenic and SNP effects as random effects. The following model was applied:

\[ y = Xb + Zu + Wg + e \]

In the above model, \( y \) is a vector of phenotypes; \( X \) is an incidence matrix related to systematic effects; \( b \) is a vector of systematic effects; \( Z \) is an incidence matrix related to random polygenic effects; \( u \) is a random vector of polygenic effects of all individuals available in the pedigree; \( W \) is a matrix \((n \times s)\) including genotype codes of SNPs for each animal; \( g \) is a vector of SNP effects and \( e \) is a vector of residual effects. In this approach Bayes Factors were retrieved as an alternative threshold instead of \( p \)-values to assess the significance of SNP effects (Habier et al. 2011). Also, Bayes Cpi enhances all drawbacks of Bayes A and B, treating the prior probability \( \pi \) that SNPs with zero effects are unknown (Habier et al. 2011).

Significant Bayes factors corresponded to SNPs effects under Bayesian regression compare the posterior odds to their prior odds in any two hypotheses (Wilson et al. 2010), which calculated by the following formula (Kass and Raftery 1995; Habier et al. 2011):

\[ BF = \frac{\frac{\pi}{1-\pi}}{\frac{1-\pi}{\pi}}, \]

where \( \pi \) is the posterior probability of the SNP effects with non-zero effect and \( \pi \) is the a priori probability of an SNP to be placed in the analysis (Habier et al. 2011). A Bayes factor is a statistical index that quantifies the evidence for a hypothesis, compared to an alternative hypothesis. Possible values for BF are any value between 0 and infinity; and the interpretation of different magnitudes can be a matter of convention since different studies have been used slightly different classification methods.

The following classification schemes have been used to detect significant SNPs and interpretation of Bayes factors (Kass and Raftery 1995):

\[ BF = 3–20, \text{ they provide suggestive evidence; } \]
\[ BF = 20–150, \text{ they provide strong evidence; } \]
\[ BF > 150, \text{ they provide very strong evidence.} \]

When BF is used, there is no place to Bonferroni correction, because all SNPs can be analysed simultaneously in the model (Costa et al. 2015). In this study, SNPs with BF greater than 150 were considered as significant. Then the log10 of retrieved Bayes factors were provided since log10(150) = 2.17 so the amounts more than 2.17 considered as significant values in Manhattan plots.

The analyses were executed by GS3 software developed by Legarra et al. (2011; https://snp.toulouse.inra.fr/~alegarra). For mapping significant phenotype-associated SNPs to the closest genes making molecular sense of association studies, we considered 0.5 Mbp up and downstream of the significant SNPs. Because, Fu et al. (2015) reported that in the crossbred chicken populations, the average \( r^2 \)LD between adjacent SNPs across the genome measured as only 0.24 (Fu et al. 2015).

Owing to multi-stage data available from a continuous life span of the bird in this study it was worthwhile to look for selection signature for different ages. Therefore, the density plot overall Bayes factors were considered to look for such signals.

**Results**

Using the Bayes factor of 150 as significant thresholds, 16 significant SNPs were detected for body weights of 2–8 weeks of age (Figures 1–7). Two SNPs for week3 (Gga_rs14453946 and Gga_rs312288120), three SNPs for week 4 (Gga_rs13934188, Gga_rs14998801 and Gga_rs313892216), five SNPs for week 5 (Gga_rs314085832, Gga_rs315478364, Gga_rs314695269, Gga_rs14502415 and Gga_rs13934188), three SNPs for week 6 (Gga_rs14657039, Gga_rs314695269 and Gga_rs315478364), two SNPs for week 7 (Gga_rs315478364 and Gga_rs14558098), and only one SNP (Gga_rs15998756)
for body weight of week 8 were significantly associated to their phenotypes. These SNPs were located close to, or inside the 14 genes and they have distributed over 8 different chromosomes, of which 12 were protein-encoding genes and the rest were noncoding regions. Non-coding RNA regions are small and do not carry information for protein synthesis, but they have particular functional roles in the cell (Costa et al. 2015). These sequences are involved in post-transcriptional regulations of gene expression (Anderegg et al. 2013).

To identify genes surrounding each significant SNP we considered 0.5 Mbp up and downstream of the significant SNPs (SNP position ±0.25 Mb). In some cases, the significant SNPs were inside Genes and in some other cases, we presented the nearest gene to the identified SNP (Table 2).

Beside significant SNPs that passed the significant threshold, an overall view of the distribution of Bayes factors across different weeks showed that the density of SNP effects shifts from around zero in early ages of weeks 2 and 3, and reaches its most distance from zero in weeks 4, 5 and 6 (Figure 8). Surprisingly these ages are the market goal ages in commercial broilers nowadays. After these ages, the distribution of Bayes factors shifts slightly back towards zero in weeks 7 and 8. Results show that the selection for market

Figure 1. Manhattan plot of resulted from Bayes factors using BayesC\textsuperscript{\textregistered} methodology for body weight at week 2 in the chicken F2 population. The \( y \)-axes indicate the logarithm (base 10) of Bayes factors and the \( x \)-axes indicate the position of SNPs along the genome in different chromosome numbers. A significant threshold is the base 10 logarithm of Bayes factor equal to 150.

Figure 2. Manhattan plot of resulted from Bayes factors using BayesC\textsuperscript{\textregistered} methodology for body weight at week 3 in the chicken F2 population. The \( y \)-axes indicate the logarithm (base 10) of Bayes factors and the \( x \)-axes indicate the position of SNPs along the genome in different chromosome numbers. A significant threshold is the base 10 logarithm of Bayes factor equal to 150.
weight may be the main driver for the size of SNPs effects of different ages. So, the presence of markers with a larger effect sizes (more deviation from zero) was considered as evidence of the selection signature.

**Discussion**

No significant marker found for body weight at week 2 of age. According to Carlborg et al. (2004) non-additive effect contribution at an early age can be considerable, therefore with models accounting for only additive effects failing to detect significant markers are expected.

Since the present study is only looking for additive effects for the SNPs then the results are comparable with additive model in Carlborg et al. (2004). Frequency distribution of the significant SNPs over the weeks under study showed that the number and the effect sizes of SNPs increasing from early age to the week 5, which reaches its maximum of five significant SNPs and decreasing again weekly to reach its minimum of only one significant SNPs in week 8. This pattern may show the effect of artificial selection that pressures on these ages (weeks 5 and 6) for commercial and market purposes. Selection process may lead to skewed distributions or some significant deviation.

Figure 3. Manhattan plot of resulted from Bayes factors using BayesC\(p\) methodology for body weight at week 4 in the chicken F2 population. The \(y\)-axes indicate the logarithm (base 10) of Bayes factors and the \(x\)-axes indicate the position of SNPs along the genome in different chromosome numbers. A significant threshold is the base 10 logarithm of Bayes factor equal to 150.

Figure 4. Manhattan plot of resulted from Bayes factors using BayesC\(p\) methodology for body weight at week 5 in the chicken F2 population. The \(y\)-axes indicate the logarithm (base 10) of Bayes factors and the \(x\)-axes indicate the position of SNPs along the genome in different chromosome numbers. A significant threshold is the base 10 logarithm of Bayes factor equal to 150.
Signatures of selection (SOS) have the potential to elucidate the characteristics of genes and mutations associated with phenotypic traits. Additionally, genetic diversity, the essential material for breeding programs can be investigated by SOS analysis. Therefore, SOS has practical implications for the implementation of genomic prediction and genomic selection. As Bayesian models by introducing the global and local parameters, which can shrink the effects of some markers to zero and allow large effect markers to escape from the shrinkage it can be considered as an efficient and alternative model for genomic prediction (Habier et al. 2011).

Results showed that chromosome 4 is the second important chromosome in terms of the number of significant SNPs. Chromosome 4 explains a relatively high proportion of the growth variation observed in different ages (Podisi et al. 2013), which is in agreement with our results. Therefore, there is likely to be the key genes related to growth traits on this chromosome.

In this study, most significant markers (9 SNPs) were located on macro chromosomes (GGA1, GGA2, GGA4, GGA5) which is an indication of the higher
contributions of these chromosomes to body weight traits (Xie et al. 2012).

Chromosome 1 reported as the main chromosome hosting several QTLs affecting body, breast muscle, thigh, and wings weights as well as overall growth (Rao et al. 2007). Among the significant markers, Gga_rs315478364 located on chromosome 1, was significant for body weights at three different ages (weeks 5, 6, and 7). This marker is located 83.8 Kb downstream of the AKR1D1 gene. Besides, Gga_rs139341188 located on the same chromosome was significant for two different ages (weeks 4 and 5). This marker is located within the LOC101751953 gene, which is one of the non-coding genes. These two markers are located in a relatively small distance from each other (~10 Mb) that could be an indication of QTL presence in this region as declared by previous studies (Nadaf et al. 2009).

The genomic region between 38.29 Mbp and 46.9 Mbp on chromosome 4 is reported as an effective region on body weight for early ages (Sewalem et al. 2002). Although there was no significant SNP having Bayes factor greater than 150 in this region there were 7 SNPs with Bayes factor of 20–150 for week 2 which provides strong evidence of significance for this trait. It seems the interference of non-additive causes has the main responsibility to prevent SNSs located in this region passing the Bayes factor of 150.

As shown in Table 2, different genes of various metabolic pathways were identified, indicating a polygenic background of body weight. These genes were classified into five categories according to their metabolic functions. These categories are involved in signalling Wnt genes (DKK2, LEF1, ROR-1), biosynthesis of various compounds, and metabolism (LPCAT1, AKR1D1, EIF2AK3), development of the nervous system (FOXp1, LRFN5), and transferring nuclear materials (XPO7). The last category including RASGEF1C and GPR137C genes play roles in internal and external cell transfer (protein transport through the membrane and

| Birds | SNP ID | Chr | Position (bp) | Nearest Gene | Distance |
|-------|--------|-----|---------------|--------------|----------|
| BW3   | Gga_rs14453946 | 4   | 38536266      | DKK2         | 2 Kb D   |
| BW3   | Gga_rs312288120 | 4   | 38295506      | LEF1         | 87.8 Kb D|
| BW4   | Gga_rs139341188 | 1   | 123148233     | LOC101751953 | 0        |
| BW4   | Gga_rs149988013 | 1   | 14041722      | RASGEF1C     | 9 Kb U   |
| BW4   | Gga_rs13892216  | 12  | 15998199      | FOXp1        | 0        |
| BW5   | Gga_rs314085832 | 2   | 86159751      | LPCAT1       | 2 Kb D   |
| BW5   | Gga_rs315478364 | 1   | 56522937      | AKR1D1       | 83.8 Kb D|
| BW5   | Gga_rs314695269 | 5   | 58320627      | GPR137C      | 0        |
| BW5   | Gga_rs14502415  | 4   | 68652277      | EIF2AK3      | 134.8 Kb D|
| BW5   | Gga_rs139341188 | 1   | 123148233     | LOC101751953 | 0        |
| BW6   | Gga_rs14657039  | 8   | 28000752      | ROR1         | 0        |
| BW6   | Gga_rs31495269  | 5   | 58320627      | GPR137C      | 0        |
| BW6   | Gga_rs315478364 | 1   | 56522937      | AKR1D1       | 83.8 Kb D|
| BW7   | Gga_rs314558098 | 5   | 59440955      | LRFN5        | 178.5 Kb U|
| BW8   | Gga_rs15998756  | 22  | 1397579       | XPO7         | 0        |

SNP ID: Single nucleotide polymorphism id; Chr: Chromosome; BW: Body Weight. DKK2: Dickkopf WNT Signalling Pathway Inhibitor 2; LEF1: Lymphoid Enhancer Binding Factor 1; RASGEF1C = RasGEF Domain Family Member 1C; FOXp1: Forkhead Box P1; LPCAT1: Lysophosphatidylcholine Acyltransferase 1; AKR1D1: Aldo-Keto Reductase Family 1 Member D1; GPR137C = G Protein-Coupled Receptor 137 C; EIF2AK3: Eukaryotic Translation Initiation Factor 2 Alpha Kinase 3; ROR1: Receptor Tyrosine Kinase Like Orphan Receptor 1; LRFN5: Leucine Rich Repeat And Fibronectin Type III Domain Containing 5; XPO7: Exportin-7.

1\[\text{U}\] and 2\[\text{D}\] refers to upstream and downstream of gene.
intracellular vesicle transportation, and controls the cell vital processes including protein biosynthesis, differentiation and growth, cellular signalling, cell proliferation, stimulation cell growth, and tissue regeneration.

The FOXP1 gene plays solely in the development of the nervous system and cooperates with other family members of FOXP genes in developing other organs (Cheng et al. 2007). Also XPO7 plays a key role to facilitate the nuclear transfer of the thyroid hormone receptor (TR) gene (Stern 2015). TR is the main receptor of the T₃ hormone which regulates the metabolism and growth rate. Additionally, it plays critical roles in neurodevelopment, the development of lung, skeletal, and bone mass.

Wnt signalling genes play role in embryonic development. Currently, Wnt proteins are known as a major family of growth signalling molecules (Cadigan and Liu 2006). These genes encode diverse families of secreted lipid-modified signalling proteins that are required during the body development (Clevers 2006), osteogenesis (Li et al. 2005), the regulation of cartilage development and foetal growth (Surmann-Schmitt et al. 2009). Also, the activation of canonical Wnt signalling is recognised as an important mechanism for vascular development (Goodwin et al. 2006). Wnt receptors through cooperating with TGF-β, regulate cell fate and proliferation during development and tissue homeostasis (Barker et al. 1999). Wnt signalling extracellular activity is regulated by Dkk family factors (Surmann-Schmitt et al. 2009). Binding DKK2 gene to LRPS/LRP6 through forming a ternary complex with trans-membrane proteins regulates the canonical Wnt pathway. Canonical Wnt signalling stimulates osteogenesis, and Dkk2 gene is involved in late stages of osteoblast differentiation into mineralised matrices. During this time Wnt7b expression reaches its peak level. Dkk2 deficiency in vitro affected osteocalcin and osteopontin expression considerably (Li et al. 2005). Osteocalcin is secreted by osteoblasts and it is

Figure 8. The probability density function of the logarithm of Bayes factors for SNPs effects in different ages. Dotted lines are referred to the weeks 2 and 3. Solid lines referred to the weeks 4, 5, and 6, and the dashed line referred to the weeks 7 and 8.
implicated in bone mineralisation and calcium ion homeostasis. Osteopontin helps to myogenic differentiation and finally muscle regeneration (Uaesoontrachoon et al. 2008). Following the consecutive transition of growth signals, Wnt can induce the transcriptional activity of numerous developmental genes through the activation of β-catenin/LEF-1 complexes in the nucleus (Filali et al. 2002). Members of the LEF1/TCF family of transcriptional factors can activate the transcription of genes by interacting with β-catenin (a downstream component of the Wnt signalling pathway) (Barker et al. 1999). The importance of LEF performance in Wnt signalling has long been proven in multiple developments and morphogenesis models (Filali et al. 2002). RORs genes are members of the nuclear receptor family of intracellular transcription factors that can be considered as the main regulators of Wnt signalling non-canonical pathways. These pathways are activated via the binding of Wnt to the frizzled domain and RORs as co-receptor. ROR-1 is a member of the ROR family of tyrosine kinase-like receptors that are highly conserved between various species (Rodriguez-Niedenfuehr et al. 2004). Receptor tyrosine kinases (RTKs) play an important role in many cellular processes, including differentiation, proliferation, and cell migration, angiogenesis, and survival (Green et al. 2008). In humans, skeletal development is known as the fundamental performance of ROR proteins (Green et al. 2008). Some studies have shown ROR1 signalling is involved in late limb development compared to early limb development (Rodriguez-Niedenfuehr et al. 2004).

The second group including LPCAT1, AKR1D1, EIF2AK3 genes have an important role in growth by participating in the biosynthesis and metabolism of various substances. Steroid 5β-reductase (AKR1D1) is a member of the Aldo keto reductase (AKR) (Mmindnich et al. 2011) enzyme. This enzyme is required for the metabolism of hormones such as androgen, oestrogen, progesterone, and also prostaglandins (Seery et al. 1998). Furthermore, the main physiological function of AKR1D1 is participation in the biosynthesis of bile acids that is known as an important metabolic pathway in all vertebrates (Mmindnich et al. 2011). Steroid hormones such as oestrogens and androgens have been raised as the main growth factor simulators. Oestrogens regulate metabolic activity of growth hormone (GH) either in the secretion or in the action levels (Leung et al. 2004). Oestrogens also can affect the pituitary secretion of GH. Moreover, it specifically can increase the expression of the suppressor cytokine signalling 2 that is a negative regulator of the GHR-JAK2-STAT5 signalling pathways (Rico-Bautista et al. 2006). On the other hand, Bile acids have a critical function to digestion and absorption of dietary fat and fat-soluble vitamins, due to forming mixed micelles in the small intestine.

EIF2AK3 membrane protein acts as a sensor to regulate the total protein synthesis. It is also essential for the normal turnover of insulin secretion from beta cells and bone growth (Liu et al. 2012). Insulin is a hormone that plays a key role in metabolism. It also involves in the storage of glycogen in muscles and amino acid transferring of the skeletal muscle system. This hormone has a GH-like act through increasing the consumption of amino acids and exchanging them to proteins and preventing degradation of available proteins in cells. Insulin is involved to transfer the received carbohydrate by the muscles. In this way, it plays an important role to build a muscular body (Ghorbani and Hashtrodi 2014).

Finally, comparing the compressed mixed linear model used by Emrani et al., in 2017 and a recent study represented the complete differences in terms of potential candidate genes resulted from these different methodologies. As expected, no commonness can be seen in this comparison. 10 candidate genes reported by Emrani et al., in 2017 included DIS3, BORA, UBE2H, CNOT10, SGOL1, ADGRB3, DTNB, SETD3, EFNA5, and SPINZ (Emrani et al. 2017) however, the recent study detected 16 candidate genes including DKK2, LEF1, RASGEF1C, FOXP1, LPCAT1, AKR1D1, GPR137C, EIF2AK3, ROR1, AKR1D1, LRFN5, XPO7, LOC101751953 and LOC101751953.

**Conclusions**

Our results have provided a list of candidate genes for body weight at ages 2–8 weeks. Results can be used in genomic selection and marker or gene assisted selection to improve growth rate in chicken. Macro chromosomes showed a remarkable role in growth control. According to the obtained results, GGA1 and GGA4 are the most important chromosomes that contain QTLs associated with the growth traits in the chicken genome. On the other hand, there were different genetic regions responsible for early and late ages for the observed traits. Ages of 4, 5, and 6 had the highest number of significant SNPs, and weeks 2 and 3 had the lowest number of significant SNPs. It may be due to the effect of non-additive drivers or due to selection goals for broilers which are weeks 4, 5, and 6 in modern commercial broiler types. This conclusion confirmed by drawing the overall density of all SNPs.
effects of different ages. The difference in the number of significant SNPs in different ages indicates the effect of artificial selection, which admit the breeding has been working properly in the chicken groups for the ages of importance. This is important as such that different regions in the genome may cause the trait of interest in different stages of life time.

It shows that the density of SNPs effects for the intended market ages is more segregating than other ages.

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Ethical approval

All experimental procedures have been approved by the Animal Care Committee of Tarbiat Modares University with a letter number 1229734.

Disclosure statement

No potential conflict of interest was reported by the author(s).

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Data and model availability statement

phenotypes:  https://figshare.com/articles/Phenotypes.rar/11369358/1

genotypes:  https://figshare.com/articles/f2_chicken_data/11369352/2

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