Determinant of the genetic and non-genetic variations in growth curve of Zandi lambs by random regression models

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Abstract The aim of this study was to model the variances and covariances of body weight in Zandi sheep from 60 to 365 days of age using random regression models (RRM). Legendre polynomials of different orders were used to model the direct and maternal covariances. Mean trends were also modeled through a quadratic regression on orthogonal polynomials of age. Homogeneity and heterogeneity of the residual variance were considered along the growth trajectory. Different models were compared by log-likelihood ratio test (LRT) and Akaike’s information criterion (AIC). Results showed that simple repeatability model in which orders of 1 were used for all random effects could not adequately model variations in growth curve of Zandi lambs. A RRM with Legendre polynomials of orders 3, 3, 3, and 3 for direct additive genetic, individual permanent environment, maternal additive genetic and maternal permanent environmental effects was selected as the most parsimonious model. The power of the parsimonious model decreased when maternal effects were excluded from the analysis, indicating the necessity of including maternal effects in the model for genetic evaluation of Zandi lambs. Considering the heterogeneity of residual variance along with the growth trajectory improved the overall properties of the model. Direct heritability (h^2) decreased from 0.3 at 60 days of age to 0.15 at about 120 days and then increased with age gradually and reached 0.39 at 365 days of age. The individual permanent environmental effect (p^2) decreased from 0.43 at 60 days of age to 0.23 at 180 days of age and fixed between 0.25 and 0.30 thereafter. Maternal heritability (m^2) was 0.03 at 60 days of age, increased to a peak around 240 days of age (0.22) and decreased with age thereafter. The ratio of maternal permanent environmental variance to phenotypic variance (c^2) was below 0.03 throughout the trajectory. Estimates of coefficients of variation (CV) revealed the presence of considerable genetic and environmental variability in growth curve of Zandi sheep which can be exploited for breeding purposes. Both direct and maternal correlations were positively high between adjacent weights but decreased as the distance between ages increased.

Keywords: sheep, body weight, random regression models, heritability, genetic correlation

Received: 22 Aug. 2017, accepted: 13 Sep. 2018, published online: 08 Dec. 2018

Introduction

Small ruminants, especially native breed types, play an important role in the livelihoods of a considerable part of human population in the tropics from socio-economic aspects (Zamani et al., 2015). Therefore, integrated attempts in terms of management and genetic improvement to enhance production is of crucial importance (Mohammadabadi and Sattayimokhtari, 2013). There are approximately 25 million heads of sheep in Iran (Ghafouri-Kesbi and Gholizadeh, 2017) consisting of 27 breeds and ecotypes which are mainly raised for meat production (Khodabakhshizadeh et al., 2016).

Recently in Iran demand for meat and mutton has increased at a significant rate mainly because of population growth and improved living standards. This means that the meat production should be increased either by increasing the number of animals or increasing the productivity per animal. Statistics show that in the last three decades, the sheep population has experienced a sharp decrease by roughly 30% (Sefidbakht, 2011). Therefore, the only way to increase meat production would be increasing the productivity per animal. This is possible by applying efficient selection programs. This strategy has been followed in recent years and some results have been published (Mokhtari and Rashidi, 2010; Mohammad et al., 2012; Gholizadeh and Ghafouri-Kesbi, 2015). In some cases, the results are somewhat lower than expected. One approach to increase the efficiency of selection programs is to increase the accuracy of genetic evaluation. This can be achieved by replacing traditional evaluation methods with random regression
models (RRM; Henderson Jr., 1984). Originally, the RRM was suggested to model the biological human data but its advantages encouraged the animal breeders to apply it in the genetic evaluation of livestock when repeated measurements or longitudinal data (measurements obtained sequentially from the same animal over time) are available. Milk yield records and body weights measured along the time are typical examples of repeated measurements. Dealing with repeated measurements, the RRM can model covariances through regression, a feature that has been used in studying body weight in sheep (Lewis and Brotherstone, 2002; Ghafouri-Kesbi et al., 2008; Safaei et al., 2010), beef cattle (Albuquerque and Meyer, 2001; Arango et al., 2004) and poultry (Rafat et al., 2011). In contrast to the traditional multi-trait models (MTM) that provide point predictions, RRM allows the prediction of breeding values for the growth curve as a whole, for any point or interval within the growth trajectory (Lewis and Brotherstone, 2002). Meyer (2004) stated that using RRM instead of traditional MTM increase the accuracy of genetic evaluation. Any increase in the accuracy of genetic evaluation is desired as it increases the genetic gain.

The Zandi sheep is a fat-tailed small-size breed. The coat-color of the new-born lamb is black, but gradually changes with age, in such a way that black, light-brown and gray adult animals are found. Historically, Zandi breed had its origin in the southern province of Iran known as Fars (Ghafouri-Kesbi et al., 2011). Today, they are reared in central areas of Iran, in Tehran and Qom provinces (Ghafouri-Kesbi et al., 2011). Due to its large population (about 2 million heads), there is considerable interest in genetically improving Zandi sheep in order to enhance their growth performance. Although, genetic information for growth-related traits has been reported for some of Iranian sheep breeds (Eskandarinasab et al., 2010; Rashidi et al., 2013; Gholizadeh and Ghafouri-Kesbi, 2015; Ghafouri-Kesbi and Gholizadeh, 2017), RRM has hardly been applied to growth data. Since increase in accuracy of breeding values is the direct consequence of applying RRM, use of RRM seems to be necessary. Therefore, the aim of this study was to analysis the growth curve of Zandi sheep by RRM. The findings might provide new insights regarding the genetic and non-genetic variations in growth curve and can be exploited by the breeders of Zandi sheep.

Materials and methods

Data and flock management

Data and pedigree information on Zandi lambs were obtained from the Breeding Station of Zandi sheep. This station was established in 1981 in the Khojir National Park between Tehran and Abali at 35°45’E and 51°40’N, 1547 m above mean sea-level, with temperate summers and cool winters, and an average rainfall of approximately 300 mm/yr. The founder animals of unknown pedigree were purchased from various sheep producers in the region. The first lambs were born in 1982 and the flock has been maintained as a close flock. In general, the flock is reared according to the conventional husbandry procedures. The mating season commences in August. Ewes on heat undergo artificial insemination (AI) and mating between very closely related animals is avoided. Usually, 20–25 ewes are inseminated with semen of one ram. Animals that do not conceive by AI are allowed to mate naturally at a ratio of 10-15 ewes per ram. Lambing commences in December. At birth, lambs are weighed, ear tagged, sexed, and their pedigree information recorded. Birth date is also recorded. The mean suckling period is 90 days. Animals are grazed on natural pasture during the spring, summer and autumn seasons and hand-fed indoors during winter.

No live weight records were available from birth to 60 days of age, thus only 60–365 day BW records were included in the data set. As recommended by Fischer et al. (2004), birth weight was excluded from the analysis because of the low genetic correlation of the birth weight with body weights measured later in life and seems to be a different trait (Fischer et al., 2004; Ghafouri-Kesbi et al. 2008). In addition, there is evidence that inclusion of birth weight may cause some problems such as increased order of polynomial fit, occurrence of “end effects of polynomials” and difficult fitting of data (Fischer et al., 2004). There were few records after 365 days that were omitted to prevent computational problems due to small number of records. Furthermore, records with doubtful dates or weights were also removed from the data file. This yielded a total of 9699 records on 4187 animals that were the progeny of 170 sires and 1450 dams.

Statistical analysis

The least squares analysis, using the GLM procedure of SAS (2004), was used to determine the significance of the fixed effects. The model included the effects of birth year, sex, type of birth and age of dam at lambing. All fixed effects were significant (P<0.05) and therefore included in the final analyses.

Four sets of RR coefficients were fitted to the data. Direct and maternal genetic effects were assumed to be proportionate to the numerator relationship matrix. Further analyses considering different orders of fit for the
four random effects were carried out subsequently to determine the most parsimonious model describing the data. The general model is represented as:

\[
y_{ij} = F_{ij} + \sum_{m=0}^{3} \beta_m x_m(t_{ij}) + \sum_{m=0}^{k_{1}} \alpha_m \phi_m(t_{ij}^*) + \\
\sum_{m=0}^{k_{2}} \gamma_{im} \phi_m(t_{ij}^*) + \sum_{m=0}^{k_{3}} \delta_{im} \phi_m(t_{ij}^*) + \sum_{m=0}^{k_{4}} \rho_{im} \phi_m(t_{ij}^*) + \epsilon_{ij}
\]

(1)

where, \(y_{ij}\) is the \(j_{th}\) record from \(i_{th}\) animal; \(t_{ij}\) is the standardized (-1 < \(t\) < 1) age at recording; \(\phi_m(t_{ij})\) is the \(m_{th}\) legendre polynomials of age; \(F_{ij}\) is a set of fixed effects; \(\beta_m\) are the fixed regression coefficients for modeling the population mean; \(\alpha_m, \gamma_{im}, \delta_{im}\) and \(\rho_{im}\) are the random regression coefficients for direct genetic, maternal genetic, and direct and maternal permanent environmental effects, respectively; \(k_{1}, k_{2}, k_{3}, k_{4}\) and \(k_{C:1}\) are the corresponding order of (polynomial) fit for each effect and \(\epsilon_{ij}\) denotes the residual (measurement error) effect. Log-likelihood ratio test (LRT) and Akaike’s Information Criterion (Akaike, 1973) were used to compare models with different orders of fit and random effects. The WOMBAT program of Meyer (2007) which uses the average information (AI) algorithm was used to analyze the data.

After determining the most parsimonious model including the direct and maternal effects, the maternal genetic and maternal permanent environmental effects were excluded from the model and differences in likelihoods were monitored to determine whether including the maternal effects in genetic evaluation procedure is necessary or not.

It has been argued that heritabilities do not provide a good means of comparing variation at different ages, when different levels of environmental variation exist (Houle, 1992; Kruuk et al., 2000). Therefore, the estimates of variance components were used to calculate the coefficients of variation. The additive coefficient of variation (\(CV_A\)) was calculated as:

\[
CV_A = 100 \times \frac{\sqrt{V_A}}{\bar{X}}
\]

(2)

where, \(V_A\) is the additive genetic variance and \(\bar{X}\) is the sample mean. The “\(\sqrt{V_A/\bar{X}}\)” ratio measures “variability”, while the ratio “\(V_A/V_P\)” measures “heritability” of the trait. Coefficients for other random effects can be expressed in a similar fashion. Compared to the raw variances, coefficients of variation are expected to be less sensitive to the scale effects arising from an increasing mean weight with age (Houle, 1992).

Two strategies were considered for modeling the residual variance (\(\sigma\)). In the first strategy, homogeneity of the residual variance for the whole period was considered. In the second strategy, heterogeneity of residual variance was considered along growth trajectory by dividing the growth trajectory to 5 growth phases, namely 60 to 120, 121 to 180, 181 to 240, 241 to 300, and 301 to 365 days of age and allocating a residual variance to each phase.

Results and discussion

Figure 1 shows the number of records and mean weights at different points of growth trajectory. Two growth phases with different growth rates are detectable, one from 60 to 180 days of age with approximately linear increase in body weight, and another one from 180 to 365 days of age with decreased growth rate. The mean weight and standard deviation for the whole period were 27.46 kg and 7.72 kg, respectively. Data characteristics and pedigree structures are shown in Table 1. The pedigree consisted of 5930 animals distributed in 9 generations; the pedigree was deep enough to allow accurate estimation of the genetic parameters.

Several models with different orders of fit were fitted to the data to find a parsimonious model that described the data adequately. Of all models fitted, the description of five models is listed in Table 2. According to LogL and AIC values, the simple repeatability model (the intercept model; 1-1-1-1) was not statistically suitable to model direct and maternal random effects. Arango et al. (2004), analyzing the growth data of beef cattle, reported that a simple repeatability model could not describe the longitudinal data adequately. In addition, greater estimates of error variances for 5 growth phases obtained with simple repeatability model compared to other models indicated that those variances tended to

Table 1. Summary of pedigree and data structure of the Zandi sheep

| N | No. of Generations (including base generation) | 9 |
|   | No. of Animals in the pedigree file | 5930 |
|   | No. of Animals with progeny | 1620 |
|   | No. of Animals without progeny | 4370 |
|   | No. of Sires with progeny | 170 |
|   | No. of Sires with progeny and record | 168 |
|   | No. of Dams with progeny | 1450 |
|   | No. of Dams with progeny and record | 965 |
|   | No. of Grand sire | 133 |
|   | No. of Grand dam | 752 |
|   | No. of Animals with record | 4187 |
Figure 1. Number of records (gray columns) and mean weights (black line) at different ages

Table 2. Order of fit for direct ($K_A$) and maternal ($K_M$) genetic, individual permanent ($K_P$) and maternal permanent ($K_C$) environmental effects (best model in bold)

<table>
<thead>
<tr>
<th>Model</th>
<th>$K_A$</th>
<th>$K_P$</th>
<th>$K_M$</th>
<th>$K_C$</th>
<th>Np</th>
<th>LogL</th>
<th>AIC</th>
<th>E1</th>
<th>E2</th>
<th>E3</th>
<th>E4</th>
<th>E5</th>
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<td>1</td>
<td>1</td>
<td>1</td>
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<td>36194.26</td>
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<td>14.20</td>
<td>10.99</td>
<td>16.54</td>
<td>26.27</td>
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<td>2</td>
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<td>-18207.40</td>
<td>36448.81</td>
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<td>13.07</td>
<td>10.00</td>
<td>8.48</td>
<td>12.36</td>
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<td>3</td>
<td>29</td>
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<td>36267.52</td>
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<td>14.40</td>
<td>12.25</td>
<td>13.68</td>
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<td>3</td>
<td>3</td>
<td>3</td>
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<td>9.54</td>
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<td>9.54</td>
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<td>3</td>
<td>-</td>
<td>-</td>
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<td>36773.50</td>
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<td>14.01</td>
<td>11.86</td>
<td>17.00</td>
<td>17.96</td>
</tr>
</tbody>
</table>

Np: Number of parameters, LogL: Log likelihood function, AIC: Akaike’s information criterion; E1-E5: estimates for residual variances for 5 growth phases (60 to 120, 121 to 180, 181 to 240, 241 to 300, and 301 to 365 days of age).

pick up variation due to insufficient order of fit (Meyer, 1999). The simple repeatability model is limited with its assumptions. It assumes that all genetic variances and covariances are of the same magnitude, all phenotypic variances are considered identical and phenotypic correlations and covariances among all measurements are equal (Meyer and Hill, 1997). These assumptions contradict most of the published reports (Albuquerque and Meyer, 2001; Lewis and Brotherstone, 2002). Increasing the order of fit from 1-1-1-1 to 3-3-3-3 improved the LRT and AIC values, a finding which has been previously reported by Lewis and Brotherstone, (2002), Arango et al. (2004) and Ghafouri-Kesbi et al. (2008). Therefore, model 3 which had 25 parameters was chosen as the most parsimonious model. Higher-order polynomials are more flexible and consequently are able to follow the changes in covariances more closely. However, the performance of higher-order polynomials is limited by the problem of “oscillation” or Runge’s phenomenon (Kirkpatrick et al., 1994; Meyer, 1998b).

Therefore, we did not increase the orders of fit beyond 3 to prevent computational problems related to high order polynomials (Meyer, 2005). In analyzing sheep growth data with RRM, usually orders equal or lower than 3 have been used (Fischer et al. 2004, Safaei et al. 2010).

The assumption that the residual term is constant over time (Model 4) had undesirable effects on the overall properties of the model, as also reported in other studies (Albuquerque and Meyer, 2001; Meyer, 2001; Huisman et al., 2005). To accommodate the fluctuation in residual variance with the age, considering heterogeneity of the residual variance over time is necessary (Albuquerque and Meyer, 2001).

By excluding the maternal effects from the most parsimonious model (Model 3) a significant decrease in Log L was observed (Model 5). Similar results were reported for growth data in sheep using traditional univariate models (Ghafouri-kesbi and Eskandarinasab, 2008; Ghavi Hossein-Zadeh, 2012; Gholizadeh and
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Ghafouri- Kesbi, 2015) and Random Regression Models (Ghafouri-Kesbi et al., 2008), indicating the necessity of including the maternal effects where lambs are genetically evaluated for growth traits.

Table 3 presents the estimates of covariances and correlations between RR coefficients for model 3 with K= 3, 3, 3, 3. As observed, the most variable coefficient was the intercept. Correlations between the intercept and linear coefficient ranged from -0.173 (maternal permanent environmental effect) to 0.626 (maternal genetic effect). Correlations between the intercept and quadratic regression coefficient ranged from -0.985 (maternal permanent environmental effect) to 0.626 (maternal genetic effect). Correlations between RR coefficients (Albuquerque and Meyer, 2001). Large eigenvalues indicate considerable genetic variation in growth curve and the potential for changing growth curve genetically. The first eigenvalue of the K matrix for A, M, P and C shows that 80-84% of the total variance can be explained by the first eigenfunction of each covariance function. Fischer et al. (2004) reported that 89-94% of total variation in Poll Dorset sheep was be explained by the first eigenfunction of each covariance function, which is close to our findings; this was 95-98% in Mehraban sheep (Ghafouri-Kesbi et al., 2008).

Estimates of variance components, genetic parameters and coefficients of variation for weights from 60 to 365 days of age, extracted from the most parsimonious model (Model 3) are shown in Figures 2, 3, and 4, respectively. The patterns for direct genetic and permanent environmental variances was similar in a way that they decreased after birth to a smallest value around 120 days of age, increased with age thereafter and reached the maximum value at 365 days of age, which is in agreement with Ghafouri-Kesbi et al. (2008). Maternal genetic variance increased after 60 days of age to a peak around 180 days of age and decreased gradually thereafter. The maternal permanent environmental variance decreased from 60 to around 120 days of age and then increased until 365 days of age.

Direct heritability estimates decreased after 60 days of age (0.30) until animals were about 120 days old (0.15) and increased sharply until 365 days of age (0.39). In particular, the direct heritability increased faster at ages at which maternal heritability tended to be the lowest. The increased pattern for heritability is in agreement with most studies (Albuquerque and Meyer, 2001; Ghafouri-Kesbi et al., 2008), and may be due to increase in the expression of genes with additive effects on body development and decrease in variances due to maternal effects at later ages. Likewise, CVₐ which measures the additive genetic variability in body weight of Zandi lambs had a similar pattern with h² in general; with a minimum value found between 150 and 160 days of age. The CVₐ can be high in traits with low heritabilities if there is high residual error variance in the trait (Houle 1992; Wilson et al., 2005). The CVₐ scales the components of the additive genetic variance by the trait mean instead of total variance and therefore is not confounded by the magnitude of other variance components. Under selection, the levels of CVₐ directly determine the magnitude of response to selection. Regarding CVₐ, two peaks were detectable: one between 220 to 250 days and another between 320 to 340 days at which body weights can be the target of selection. The individual permanent environmental effect (p²) decreased rapidly from 0.43 at 60 days to a value of 0.35 at about 120 days of age, slowly decreased until 180 days of age (0.23) and then fluctuated between 0.25 and 0.30 until the end of the trajectory. The pattern for CVₚ differed from p² in such a way that it decreased slowly until 180 days of age and gradually increased thereafter. Maternal heritability estimates (m²) increased from 60 days of age (0.03) to 240 days of age (0.22) and decreased with age thereafter. At weaning (~90-100 days), the estimates of m² were between 0.11 and 0.14. Persistency of m² after weaning may be due to carry-over of maternal effects from weaning weight. Such carry-over of maternal effects beyond weaning are biologically sensible; particularly when lamb growth to weaning is restricted by low ewe milk

Table 3. Estimated coefficient matrices of the covariance functions and the corresponding eigenvalues of the covariance matrices and the correlation (above diagonal) between random regression coefficients for direct genetic (A), maternal genetic (M), individual permanent environmental (P) and maternal permanent environmental (C) effects

<table>
<thead>
<tr>
<th></th>
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<th>Eigenvalue</th>
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<td>17.941</td>
<td>0.516</td>
<td>-0.268</td>
<td>3.39 (14%)</td>
</tr>
<tr>
<td>M</td>
<td>1.5327</td>
<td>0.627</td>
<td>-0.616</td>
<td>3.16 (82%)</td>
</tr>
<tr>
<td>P</td>
<td>3.3739</td>
<td>0.556</td>
<td>0.985</td>
<td>3.87 (84%)</td>
</tr>
<tr>
<td>C</td>
<td>1.435</td>
<td>-0.171</td>
<td>-0.985</td>
<td>1.49 (80%)</td>
</tr>
</tbody>
</table>

61
Figure 2. Estimates of variance components

Figure 3. Estimates of direct and maternal heritability and individual and maternal permanent environmental effects
Genetic and non-genetic variations of growth curve in Zandi lambs

Figure 4. Estimates of direct genetic (CVA), individual permanent (CVP), maternal genetic (CVM) and maternal permanent environmental coefficient of variation (CVC)

Production either due to large litter size or seasonal constraints on feed (Bradford, 1972; Snyman et al., 1995). It is more evidenced by estimates of CVM with a peak between 220 and 240 days of age as well as estimates of \( c^2 \) and CVC with sharp increases after weaning. However, estimates of \( c^2 \) and CVC were below 0.03 and 5% throughout the trajectory, which indicated that maternal permanent environmental effect has little contribution in phenotypic variation of body weight in Zandi lambs; therefore this effect can be excluded from the model of genetic evaluation. By including only one of the maternal effects (additive genetic or permanent environment-

<table>
<thead>
<tr>
<th>Age1</th>
<th>Age2</th>
<th>( r_s )</th>
<th>( r_p )</th>
<th>( r_m )</th>
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<tr>
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</tr>
<tr>
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<td>365</td>
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<td>1</td>
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<tr>
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\( r_c \): Direct genetic correlation, \( r_p \): individual permanent environmental correlation, \( r_m \): Maternal genetic correlation, \( r_c \): Maternal permanent environmental correlation.
Correlations between selected body weights are listed in Table 4. The pattern of direct and maternal genetic and individual and maternal permanent environmental correlations is the same; in that correlations decreased as the distance between the tests increased. The genetic correlations between body weight at 60 days and around 9 and 12 months of age were smaller than 0.1, indicating that genes controlling 60-day body weight has little contribution to development of body weight at higher ages and vice versa. Therefore, small correlated response in body weight at higher ages is expected following selection for 60-day body weight. This is in agreement with previous reports (Lewis and Brotherstone, 2002; Arango et al., 2004; Ghafouri-Kesbi et al., 2008). The direct genetic correlations between body weights taken at higher ages were highly positive; suggesting that they may be under the same genetic control. Maternal genetic correlations between 60-day body weight and body weights at higher ages were negative which indicates that the genes of dams which contribute in milk production have some unfavorable effects on post-weaning body weights.

In general, results showed the inadequacy of simple repeatability model to deal with repeated body weight records as evidenced by LRT and AIC values. An RRM with orders of fit of 3, 3, 3 and 3 for direct genetic, maternal genetic, individual permanent and maternal permanent environmental effects was selected as the most parsimonious model for analysis. Results from this model revealed presence of genetic variation in the growthcurve of Zandi lambs and that this genetic variation was not constant throughout the growth trajectory and fluctuated with advancing age. Body weights beyond 200 days of age had greater genetic variability and heritability and therefore greater genetic response would be expected through genetic selection. While maternal genetic effects had a considerable contribution to the phenotypic variation in body weight of Zandi lambs, the contribution of maternal permanent environmental effect was negligible and therefore can be excluded in genetic evaluation models. Genetic and permanent environmental correlations between body weights decreased as the distance between the pairs of ages increased.

Acknowledgements
The staffs of the Zandi Sheep Breeding Station, in particular Mr Sharif, are gratefully acknowledged for collecting the data used in this research.

References


**Communicating editor:** Ali K. Esmailizadeh
بررسی تنوع زنده و غیرزنده در منحنی رشد گوسفندهای زندی با استفاده از مدل‌های رگرسیون تصادفی

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چکیده
هدف این پژوهش، مدل ساختاری واریانس‌ها و کواریانس‌های وزن بدن گوسفندهای زندی بین سن‌های ۶۰ تا ۳۶۵ روزگری با به کارگیری مدل‌های رگرسیون تصادفی بود. چندجمله‌ای‌های لجبری برای مدل‌سازی اثرهای مستقیم و مادی استفاده شدند. میانگین منحنی رشد جمعیت نیز با روش رگرسیون درجه ۲ از سن روزگری اول در آیینه مدل‌های برآوردگری برای شدن. در ضمن، یکنواختی و عدم یکنواختی واریانس باقی‌مانده در طول منحنی رشد در نظر گرفته شد. مدل‌های مختلف با استفاده از آزمون لگاریتم نسبت درستنمایی (LRT) و شاخص اطلاعات آکائیک (AIC) مورد مقایسه قرار گرفتند. نتایج نشان داد که مدل تکرارپذیری ساده که در آن از درجه برآورد ۱ برای مدل سازی اثرات مستقیم و مادی استفاده کرده و فرضیه رفت. به نظر نشان‌دهنده ضرورت در نظر گرفتن اثرات مادی در مدل ارایش زنده‌ای در طول دوره رشد در طول دوره رشد منجر به بهبود خصوصیات کلی مدل می‌گردد. ورالت‌پذیری مستقیم (۳) از مقدار ۰.۳ در سن ۶۰ روزگری به ۰.۱۵ در سن ۱۲۰ روزگری کاهش یافت و سپس به صورت تدریجی افزایش یافت و در سن ۳۶۵ روزگری به مقدار ۰.۳۹/۰. سی رفت. اثر میانگین دانی منحنی در سن ۶۰ روزگری کاهش یافت و سپس به صورت تدریجی افزایش یافت. در سن ۶۰ روزگری به ۰.۲۳ در سن ۱۸۰ روزگری کاهش یافت و پس از سن ۲۲۰ به ۰.۲۳/۰. سی رسید و پس از آن روند کاهشی به خود گرفت. اثر میانگین دانی مادی (۳) در کل دوره مورد بررسی در حدود ۰.۳/۰. قرار داشت. برآورد‌هاي ضریب تغییرات (CV) مقدار مطلق توجه و تغییرات زنده‌ای و میانگین در منحنی رشد گوسفندهای زندی را نشان داد که می‌توانند جهت اصلاحی مورد بهره‌برداری قرار گیرد. همچنین مدل‌های مستقیم و مادی بین وزن بدن در سن‌های مختلف به طور قابل توجه ملاحظه‌ای مشاهده شده و با بودن ایفای نقش در آینده ابتدا روزهای زنده‌گیری کاهش یافت.