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Genetic and environmental relationships among milk yield, persistency of milk yield, somatic cell count and calving interval in Holstein cows

Relaciones genéticas y ambientales entre el rendimiento lechero, la persistencia del rendimiento lechero, el recuento de células somáticas y el intervalo de partos en vacas Holstein

Relações genéticas e ambientais entre produção de leite, persistência de produção de leite, contagem de células somáticas e intervalo de partos em vacas Holstein

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Abstract

Background: Fertility and health traits, other than production traits, have a major role in the profitability of dairy cattle. Therefore, it seems necessary to include the afore mentioned traits in breeding programs. Hence, genetic parameters are needed to establish breeding plans. **Objective:** To estimate heritabilities as well as genetic and environmental relationships among total milk yield (TMY), persistency of milk yield (PMY), mean somatic cell count (SCC), mean log_e somatic cell count (LnSCC), standard deviation of somatic cell count (stdSCC), and calving interval (CI) using two-trait and multi-trait analyses in Iranian Holstein. **Methods:** The dataset consisted of 25,883 first lactation records collected from 2002 to 2007 in 97 Holstein dairy herds in Iran. Four criteria of persistency of milk yield (PMY) were calculated using the Wood's gamma function. The WOMBAT1.0 software was used to estimate the (co)variance components using the Average Information Restricted Maximum Likelihood algorithm. **Results:** Total milk yield (TMY) resulted in the highest heritability estimate (0.29). Heritability estimates for different criteria of persistency of milk yield (PMY) ranged from 0.05 to 0.10. The unfavorable genetic correlation between TMY and calving interval (CI) was 0.71, while that of PMY with CI was 0.46. The estimated environmental correlations were lower than the genetic correlations for all traits, but the trends were generally similar. **Conclusion:** The results indicate that including PMY in the breeding goals could increase TMY and CI, and decrease somatic cell count (SCC).

Keywords: *breeding goal, correlation, fertility, heritability, Wood's gamma function.*

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Resumen

Antecedentes: Las características de fertilidad y salud, además de las de producción, tienen un papel importante en la rentabilidad de los hatos lecheros. Por lo tanto, parece necesario incluir los rasgos mencionados en los sistemas de mejoramiento. Por ende, los parámetros genéticos son necesarios para establecer planes de cría. **Objetivo:** Estimar la heredabilidad, así como las relaciones genéticas y ambientales entre el rendimiento total de leche (TMY), la persistencia del rendimiento lechero (PMY), el recuento medio de células somáticas (SCC), el recuento de células somáticas de \log_e promedio (LnSCC), la desviación estándar del recuento de células somáticas (stdSCC) y el intervalo de partos (CI) en vacas Holstein mediante análisis de dos rasgos y análisis múltiples. **Métodos:** Se analizaron registros de primera lactancia de 25.883 vacas Holstein, recolectados entre 2002 y 2007 en 97 rebaños lecheros de Irán. Se calcularon cuatro criterios de persistencia del rendimiento lácteo utilizando la función gamma de Wood. El software Wombat1.0 se utilizó para estimar los componentes de (co)varianza que emplean el algoritmo de máxima verosimilitud restringida de información promedio. **Resultados:** El rendimiento de leche (TMY) obtuvo la mayor heredabilidad estimada (0,29). Las estimaciones de heredabilidad para diferentes criterios de persistencia del rendimiento de la leche variaron de 0,05 a 0,10. La correlación genética desfavorable entre TMY e intervalo entre partos fue 0,71, mientras que la de PMY con el CI fue de 0,46. Las correlaciones ambientales estimadas fueron inferiores a las correlaciones genéticas para todos los rasgos, pero las tendencias fueron, en general, similares. **Conclusión:** Incluir el PMY en la meta de cría podría aumentar el TMY y CI, además de disminuir el recuento de células somáticas (SCC).

Palabras clave: *correlación, fertilidad, función gamma de Wood, heredabilidad, objetivo de cría.*

Resumo

Antecedentes: As características de fertilidade e saúde, além das características de produção, têm um papel importante na rentabilidade das vacas leiteiras. Por isso, parece necessário incluir as características já mencionadas nos sistemas de melhoramento genético. Assim, são necessários parâmetros genéticos para estabelecer planos de melhoramento genético. **Objetivo:** Estimar herdabilidade, bem como as relações genéticas e ambientais entre a produção total de leite (TMY), persistência da produção de leite (PMY), contagem média de células somáticas (SCC), contagem média de células somáticas \log_e (LnSCC), desvio padrão da contagem média de células somáticas (stdSCC) e intervalo de parto (CI) vacas Holstein por duas análises de características e traços múltiplos. **Métodos:** O conjunto de dados consistiu em registros de 25.883 vacas Holstein de primeira lactação coletadas de 2002 a 2007 em 97 rebanhos leiteiros em Irã. Foram calculados quatro critérios de persistência da produção de leite utilizando a função gama de Wood. O software Wombat1.0 foi usado para estimar os componentes de (co)variância que empregam o algoritmo de máxima verossimilhança com informações restritas. **Resultados:** A maior herdabilidade (0,29) foi estimada para produção de leite. As estimativas de herdabilidade para diferentes critérios de persistência da produção de leite variaram 0,05 a 0,10. A correlação genética desfavorável entre o intervalo TMY e o intervalo de parto foi de 0,71 enquanto que a de PMY com CI foi de 0,46. As correlações ambientais estimadas foram inferiores às correlações genéticas para todos os traços, mas as tendências foram geralmente semelhantes. **Conclusão:** Os resultados indicaram que a inclusão de PMY na meta de reprodução poderia aumentar TMY e CI, no entanto, diminuir a contagem de células somáticas.

Palavras chave: *correlação, fertilidade, função gama do Wood, herdabilidade, objetivo de criação.*

Introduction

High milk yield has been the primary selection emphasis in dairy breeding. However, many non-production traits, such as reproduction traits (Lucy, 2007), are important in minimizing cost and maximizing the net return of the dairy enterprise. Some researchers (Dematawewa and Berger, 1998; Haile-Mariam *et al.*, 2003) mentioned that selection for increased milk yield may reduce reproductive efficiency, and increase

susceptibility to certain diseases as well as the risk of culling due to diseases such as milk fever, displaced abomasum, and fatty liver. Milk persistency has been shown to be an economically important trait in dairy cattle (Haile-Mariam *et al.*, 2003). For the same total yield, cows with lower peak yield and greater persistency may experience less energy imbalance and thus less reproductive and health problems than cows with higher peak yield (Haile-Mariam *et al.*, 2003). In addition of these advantages, persistency can minimize

feed costs, because the proportion of roughage in the diet of cows with flat lactation curves can be increased (Solkner and Fuchs, 1987). Hence, including PMY and other functional and reproduction traits in addition to production traits is necessary to improve the overall economic performance of dairy herds.

Several studies have been conducted to estimate genetic parameters for milk yield and composition (Sahebbonar, 2007; Ghavi Hossein-Zadeh and Ardalan, 2011), milk yield persistency (Seyed Sharifi *et al.*, 2006), and reproductive traits (Toghiani Pozveh *et al.*, 2009) in Iranian Holstein cows. The heritability and genetic relationship among economic performance traits are needed for planning efficient breeding systems and develop effective genetic evaluations.

One of the main advantages of multivariate analyses is that it increases the accuracy of evaluations. The gain in accuracy is dependent on the absolute difference between the genetic and residual correlations between traits. The larger the differences in these correlations, the greater the gain in accuracy of evaluations (Schaeffer, 1984; Thompson and Meyer, 1986). Also, there is an additional increase in accuracy with multivariate analysis resulting from better connections in the data due to residual covariance between traits (Thompson and Meyer, 1986). Therefore, the objective of this study was to estimate heritabilities as well as genetic and environmental relationships among total milk yield (TMY), persistency of milk yield (PMY), mean somatic cell count (SCC), mean \log_e somatic cell count (LnSCC), standard deviation of somatic cell count (stdSCC), and calving interval (CI) using two-trait and multi-trait analyses in Iranian Holstein.

Material and methods

Data

Test-day records of milk yield, somatic cell count and calving date of Holstein cows were obtained from the Animal Breeding Center of Iran. Herds with more than 500 test-day records from daughters of at least four different sires were included. Cows that calved for the first time between 20 and 40 months of age,

and those that had at least 6 test-day records were considered in the analysis. Calving intervals between 290 to 600 days were included. Only records of cows with known pedigree were included. The final dataset consisted of records of 25,883 cows with first calving from July 2002 to September 2007 in 97 herds. The average number of daughters per sire was 23.28, and about 6% of the sires had more than 100 daughters. The total number of sires was 1,112. A summary of pedigree information is shown in Table 1.

Table 1. Summary of pedigree information in this study.

Description	Total
Total number of animals	38,291
Animals with record	20,544
Animals without progeny	19,022
Animals with progeny	4,382
Animals with progeny and record	1,522
Base animals	3,026
Sires	837
Dams	3,545
Average inbreeding coefficient	0.0021 (%)
Number of herd-test days	1,324

Statistical methods

Using test-day records of milk yield, the 305-day total milk yield, first 100-day milk yield, and second 100-day milk yield were calculated with Wood's gamma function. The MATLAB 7.7.1 software (Mathworks, 2008) was used for fitting the Wood's gamma function and obtaining the corresponding parameters. The gamma function described by Wood (1967) is one of the most popular mathematical models used to describe the lactation curve as follows:

$$Y_t = at^b e^{-ct}$$

where y_t = milk yield on day t , a = a scaling factor to represent yield at the beginning of lactation, and b and c are factors associated with the inclining and declining slopes of the lactation curve, and e is the

Napier number (~ 2.71828). Typical lactation curves have positive b and c (Tekerli *et al.*, 2000). Atypical lactation curves were excluded from further analysis.

Two types of persistency measures were used in this study. The first type used ratios of different parts of the lactation, namely P2:1, P3:1 and P_{weller} . The P2:1 and P3:1 are the ratios of total milk yield in the second and third 100 days to the first 100 days of lactation, respectively. P_{weller} was introduced by Weller *et al.* (2006) as the estimated milk production at 180 day after peak, divided by the estimated peak production in percent as follows:

$$P_{\text{weller}} = 100\% \times [\text{PROD}(270)] / [\text{PROD}(90)]^{-1}$$

Where, $\text{PROD}(270)$ = is the production at 270 DIM, and calculated by Wood's function. They assumed peak production at first lactation occur at 90 DIM. Higher values of the above mentioned criteria indicate greater persistency.

The second type of measure for persistency used the Wood's function as follows:

$$s = -(b + 1) \ln(c)$$

Estimated genetic and environmental covariance components are sensitive to data amount and structure. In order to control over consistency of these estimates, two types of analysis were performed separately, including two and six-traits analyses. The WOMBAT1.0 software (Meyer, 2007) was used to estimate the variance and covariance components. The fixed part of the model included the effect of region, herd-year-season (four seasons i.e., winter, spring, summer, fall were assumed), age at first calving (months) and percent genetic contribution from the Holstein breed considered as covariate effects. The following linear model was used for data analysis:

$$Y_{ijklmn} = \mu + R_i + \text{HYS}_j + \text{AFC}_k + b\text{BL}_l + A_m + e_{ijklmn}$$

Where: Y_{ijklmn} = is the $ijklmn$ observation on TMY, PMY, mean SCC, mean LnSCC and stdSCC; μ = is the population mean; R_i = is the fixed effect of region; HYS_j is the fixed effect of herd-year-season of calving; AFC_k = is the fixed effect of age at first calving; BL_l = is the

covariate effect of percent genetic contribution from the Holstein breed; b is the linear regression coefficient of traits on percent genetic contribution from the Holstein breed; A_m = is the random genetic effect of the animal, and e_{ijklmn} = is the random residual error.

Results

Summary statistics for studied traits are shown in Table 2 (estimated mean and standard deviations of traits). Mean TMY was 9,065 kg. Averages of P2:1, P3:1, P_{weller} and s were estimated as 1.063, 0.915, 0.803 and 7.709, respectively. Also, means of the test-day SCC and LnSCC were 245.1 and 4.869, respectively.

Table 2. Summary of descriptive statistics for the phenotypic values (based on 20,544 first lactation records).

Trait	Mean (SD)
TMY(kg)	9,065 (1,521.5)
CI (days)	398.6 (66.7)
P2:1	1.063 (0.216)
P3:1	0.915 (0.246)
P_{weller}	0.803 (0.157)
s	7.709 (1.04)
SCC ('000)	245.1 (343.9)
LnSCC	4.869 (1.109)
stdSCC	318.5 (541.2)

SD: standard deviations; TMY: total milk yield; CI: calving interval; P2:1: [(milk yield second 100 days)/(milk yield first 100 days) $^{-1}$] $\times 100$; P3:1: [(milk yield third 100 days)/(milk yield first 100 days) $^{-1}$] $\times 100$; P_{weller} : [(milk yield 270d)/(milk yield 90d) $^{-1}$] $\times 100$; $s = -(b + 1) \ln(c)$; SCC: mean somatic cell count; LnSCC: mean \log_e somatic cell count; stdSCC: standard deviation of somatic cell count.

Estimates of variance components and heritabilities using single-trait analyses are shown in Table 3. Heritability of TMY was 0.29. There was not a large difference between estimates of heritabilities by single-trait and six-trait analyses (Tables 3 and 4). Among different measures of persistency of lactation, P_{weller} had the highest h^2 (0.104 ± 0.015), and P2:1 had the lowest h^2 (0.05 ± 0.011). The highest h^2 among measures of SCC was for LnSCC (0.064 ± 0.012). Also, heritability of CI by single-trait and six-trait analyses were 0.044 and 0.041, respectively.

Table 3. Estimates of variance components and heritabilities of the traits obtained from single-trait analyses.

Trait	Additive variance	Residual variance	Heritability (SE)
TMY	323,561	793,678	0.29 (0.024)
CI	177.508	3,889.45	0.044 (0.01)
P2:1	0.00170814	0.0325266	0.05 (0.011)
P3:1	0.0042058	0.0411223	0.093 (0.015)
P _{weller}	0.00209998	0.0180623	0.104 (0.015)
s	0.0425813	0.792954	0.051 (0.011)
SCC ('000)	3,086.94	98,284.2	0.03 (0.008)
LnSCC	0.0567233	0.832363	0.064 (0.012)
stdSCC	3,837.74	258,296	0.015 (0.007)

TMY: total milk yield; CI: calving interval; P2:1: [(milk yield second 100 days)(milk yield first 100 days)⁻¹] \times 100; P3:1: [(milk yield third 100 days)(milk yield first 100 days)⁻¹] \times 100; P_{weller}: [(milk yield 270d)(milk yield 90d)⁻¹] \times 100; s = -(b + 1)ln(c); SCC: mean somatic cell count; LnSCC: mean log_e somatic cell count; stdSCC: standard deviation of somatic cell count; SE: standard error of estimates.

Table 4. Heritabilities (bold on diagonal), genetic (above) and environmental correlations (below) for total milk yield (MY), calving interval(CI), P3:1, P_{weller}, s and LnSCC by six-trait analyses.

Trait	TMY	CI	P3:1	P _{weller}	s	LnSCC
TMY	0.285 (0.023)	0.732 (0.086)	0.536 (0.07)	0.573 (0.063)	0.436 (0.09)	-0.092 (0.093)
CI	0.019 (0.014)	0.041 (0.009)	0.442 (0.118)	0.417 (0.115)	0.447 (0.14)	-0.076 (0.144)
P3:1	0.049 (0.016)	0.127 (0.01)	0.098 (0.015)	0.979 (0.012)	0.93 (0.031)	-0.348 (0.111)
P _{weller}	0.184 (0.015)	0.182 (0.01)	0.827 (0.004)	0.107 (0.015)	0.843 (0.063)	-0.354 (0.107)
s	-0.081 (0.015)	0.045 (0.01)	0.734 (0.005)	0.38 (0.009)	0.057 (0.011)	-0.162 (0.136)
LnSCC	-0.138 (0.015)	0.048 (0.01)	-0.043 (0.011)	-0.075 (0.011)	-0.01 (0.011)	0.063 (0.012)

TMY: total milk yield; CI: calving interval; P3:1: [(milk yield third 100 days)(milk yield first 100 days)⁻¹] \times 100; P_{weller}: [(milk yield 270d)(milk yield 90d)⁻¹] \times 100; s = -(b + 1)ln(c); LnSCC: mean log_e somatic cell count; SE: standard error of estimates.

Estimates of genetic and environmental correlations (r_g and r_e , respectively) among traits are shown in Table 5. Genetic correlations between TMY and measures of PMY using two-trait and six-trait analyses were ranged from 0.436 (for TMY with s) to 0.573 (for TMY with P_{weller}). Genetic correlation between TMY and CI by two- and six-trait analyses estimated as 0.71 and 0.732, respectively.

Estimates of genetic correlations obtained by six-trait analyses were relatively higher than those of two trait analysis, although none of those correlation coefficients were significantly different from zero. Genetic correlations between measures of PMY and CI estimated by both two- and six-trait analyses ranged from 0.4 to 0.46. Genetic correlations between the measures of PMY and SCC were negative and moderate. Absolute value of r_g between measures of PMY and SCC showed that SCC was more correlated with amount of milk production in late lactation rather than other lactation stages.

Table 5. Genetic (above) and environmental correlations (below) for total milk yield (TMY), calving interval (CI), different criteria of milk persistency and somatic cell count by two-trait analyses.

Trait	TMY	CI	P2:1	P3:1	P _{weller}	s	SCC	LnSCC	stdSCC
TMY		0.71 (0.086)	0.467 (0.089)	0.526 (0.07)	0.566 (0.062)	0.44 (0.09)	-0.023 (0.12)	-0.076 (0.092)	-0.103 (0.156)
CI	0.021 (0.013)		0.46 (0.14)	0.444 (0.12)	0.4 (0.115)	0.43 (0.14)	0.004 (0.176)	-0.051 (0.143)	-0.036 (0.228)
P2:1	-0.06 (0.015)	0.05 (0.01)		0.97 (0.015)	0.906 (0.05)	0.999 (0.01)	-0.35 (0.165)	-0.275 (0.135)	-0.348 (0.217)
P3:1	0.05 (0.015)	0.125 (0.01)	0.89 (0.002)		0.98 (0.012)	0.912 (0.034)	-0.416 (0.14)	-0.336 (0.113)	-0.404 (0.188)
P _{weller}	0.185 (0.014)	0.185 (0.01)	0.494 (0.008)	0.827 (0.003)		0.797 (0.07)	-0.395 (0.14)	-0.328 (0.11)	-0.362 (0.185)
s	-0.086 (0.015)	0.046 (0.01)	0.892 (0.002)	0.752 (0.004)	0.403 (0.01)		-0.156 (0.17)	-0.14 (0.14)	-0.11 (0.224)
SCC	-0.145 (0.013)	0.032 (0.009)	-0.023 (0.01)	-0.051 (0.01)	-0.083 (0.01)	-0.012 (0.01)		0.992 (0.028)	1 (0.04)
LnSCC	-0.14 (0.015)	0.049 (0.01)	-0.016 (0.01)	-0.044± (0.01)	-0.077 (0.01)	-0.013 (0.01)	0.768 (0.004)		0.999 (0.067)
stdSCC	-0.094 (0.013)	0.029 (0.007)	-0.01 (0.009)	-0.029 (0.01)	-0.054 (0.01)	-0.001 (0.01)	0.873 (0.002)	0.73 (0.004)	

TMY: total milk yield; CI: calving interval; P2:1: [(milk yield second 100 days)(milk yield first 100 days)⁻¹] \times 100; P3:1: [(milk yield third 100 days)(milk yield first 100 days)⁻¹] \times 100; P_{weller}: [(milk yield 270d)(milk yield 90d)⁻¹] \times 100; s = -(b + 1)ln(c); SCC: mean somatic cell count; LnSCC: mean log_e somatic cell count; stdSCC: standard deviation of somatic cell count; SE: standard error of estimates.

Discussion

Phenotypic parameters and heritabilities

Mean TMY was (9,065 kg) in the range from 5,558 to 10,881 kg reported by other researchers for Holstein breed (Dematawewa and Berger, 1998; Haile-Mariam *et al.*, 2003; Kadarmideen *et al.*, 2003; Muir *et al.*, 2004; Weller *et al.*, 2006; Bakir and Kaygisiz, 2013). The mean of CI was (398.6 d) slightly longer than that reported (from 382 to 398.3) by Haile-Mariam *et al.* (2003), Kadarmideen *et al.* (2003), Muir *et al.* (2004), and Dong and Van Vleck (1989), but is shorter than that of 406 reported by Ojango and Pollott (2001) for Holstein-Friesian cattle in Kenya. This difference is expected because CI is highly influenced by environment and management factors.

There was not a large difference between heritability estimates of by single-trait and six-trait analyses (Tables 3 and 4). The h^2 of TMY (0.29) was in the range of estimates reported for Australian, Canadian and US Holstein cattle (Dong and Van Vleck, 1989; Dematawewa and Berger, 1998; Haile-Mariam *et al.*, 2003; Muir *et al.*, 2004; Weller *et al.*, 2006; Kaygisiz, 2013).

As can be seen (Table 3), heritability estimates for different measures of persistency of lactation were low, averaging 0.075. The heritability of persistency has been reported to be between 0.01 and 0.3 (Gengler, 1996; Swalve and Gengler, 1999), and varied according to the measure used. The highest h^2 for measures of SCC was for LnSCC (0.064±0.012). Estimates of Kennedy *et al.* (1982) for heritability of LnSCC ranged from 0.05 for ≤ 2 years old to 0.10 for mature (≥ 6 years) cows, and averaged 0.08 over all ages. Other researchers reported the heritability for this trait from 0.04 to 0.44 (Afifi, 1967; Ali and Shook, 1980; Reents *et al.*, 1995; Haile-Mariam *et al.*, 2003).

Heritability of CI in this study by single-trait and six-trait analyses were 0.044 and 0.041, respectively. Kadarmideen *et al.* (2003) obtained h^2 for CI as 0.024 for Holsteins in UK that was lower than our result. However, some other estimates of heritabilities for this trait were between 0.04 and 0.15 (Dong and Van Vleck, 1989; Ojango and Pollott, 2001; Haile-Mariam *et al.*, 2003; Muir *et al.*, 2004).

Correlations

Correlation between TMY and PMY. Genetic correlations between TMY and measures of PMY using two-trait and six-trait analyses were relatively high (ranged from 0.436 to 0.573), being in agreement with Solkner and Fuchs (1987), Swalve (1995) and Rekaya *et al.* (2000), whose estimates of genetic correlations between TMY and measures of s, P2:1 and P3:1 ranged from 0.33 to 0.56. However, the observed values disagreed with Haile-Mariam *et al.* (2003), and Weller *et al.* (2006), who reported lower estimates. Gengler (1996) recommended the use of a measure of persistency that is genetically independent of 305-d milk yield. In this study a moderate positive genetic correlation was estimated between yield and measures of persistency, indicating that selection for increased TMY can cause a correlated response for persistency.

Correlation between TMY and CI. The existence of a positive genetic correlation between TMY and CI (~0.7) indicates that selection for increasing TMY would increase CI, therefore it is unfavorable because of the reduced number of calves produced per cow's lifetime. The r_g obtained in the present study is higher than that reported for dairy cattle in Australia (~0.5) (Haile-Mariam *et al.*, 2003), and was inconsistent with Ojango and pollot (2001), who reported a negative r_g of -0.64 between calving interval and milk yield in the first lactation. Also, Ojango and pollot (2001) stated that genes that positively affect milk production are likely to reduce calving interval. Muir *et al.* (2004) reported a strong genetic correlation between TMY and CI (0.51 ± 0.11), and declared that a longer CI is associated with greater TMY in first lactation. The environmental correlation between TMY and CI was close to zero in the present study (0.02). This implies environmental conditions that increase TMY would not increase CI.

Correlation between TMY and SCC. Genetic correlations of TMY with measures of SCC were low. Environmental correlations between TMY and measures of SCC were negative, indicating that cows with higher production probably are experiencing better health and sanitation conditions. The correlation of TMY and stdSCC was slightly stronger than that of TMY and LnSCC (-0.103 vs. -0.076, respectively). It implies that the higher the consistency of SCC over the

lactation period, the higher the TMY. Although h^2 of stdSCC was lower than that of the mean of SCC and LnSCC, its correlation with other traits was higher; it may be useful and applicable for performing a correlated selection.

Haile-Mariam *et al.* (2003) estimated a negative genetic relationship between slope of LnSCC with mean MY (-0.27). They remarked that increase in LnSCC later in lactation is associated with reduced MY. However, a positive genetic correlation between SCC and milk yield in the first lactation has been reported by several researchers (Kennedy *et al.*, 1982; Weller *et al.*, 2006). Taking into account that those correlations were not significantly different from zero, it could be concluded that selection on TMY would not increase SCC.

In general, absolute environmental correlations were lower in magnitude than genetic correlations (Table 5). The exceptions was the environmental correlation between TMY and SCC measures. Estimates of phenotypic correlations (not shown) between all traits were also similar in direction to environmental correlations.

Correlation between PMY and CI. Measures of PMY had relatively high genetic correlations with CI (0.4 to 0.46). Estimates of the relationship between PMY and fertility are rare in the literature, though the relationship is assumed to be less antagonistic than that with TMY (Dekkers *et al.*, 1998), as we found (~0.4). Lean *et al.* (1989) reported that cows with higher than average peak milk yields and poorer than average persistency were less likely to conceive in 1 or 2 services. In agreement with the result of the present study, Muir *et al.* (2004) reported an unfavorable correlation between PMY and CI (0.17), and expressed that the greater the persistency was in first lactation, the longer the interval was from first to second calving. It is important to know that Weller and Folman (1990) stated that late conception reduced profitability and showed early breeding to be advantageous, especially if the calf has a high value.

The antagonist genetic correlation of CI with P2:1 is higher than that with P3:1 and P_{weller} , which implies that the higher the ratios of milk production early in lactation to milk production late in lactation, the longer the CI. The main reason is that cows are usually in negative energy

balance early in lactation and, as a result, animals with relatively low production are inseminated earlier and will consequently have shorter CI.

Correlation between PMY and SCC. The estimated negative and moderate genetic correlations between PMY and SCC in this study indicate that more persistent cows have lower SCC, healthier udders, and lower incidence of mastitis. Estimates of correlations by six-trait analyses were slightly higher. Absolute value of r_g between measures of PMY and SCC showed that SCC was more correlated with milk production in late lactation rather than other stages of lactation. This agrees with Haile-Mariam *et al.* (2001), who reported that the magnitude of the correlation between milk yield and SCC within parity is relatively higher at the end of lactation. Also, our result was consistent with Haile-Mariam *et al.* (2003) and Weller *et al.* (2006), who reported negative r_g .

Correlation between CI and SCC. Genetic correlations estimated for CI and measures of SCC were between -0.076 and 0.004. These results are in accordance with findings by Österman *et al.* (2005), who showed that the LnSCC throughout the lactation does not differ as a result of an extended CI, compared to the traditional 12 months CI, but disagreed with Haile-Mariam *et al.* (2003), Weller *et al.* (2006) and Rekik *et al.* (2008), who reported positive r_g between CI and SCC. Rekik *et al.* (2008) reported that elevated SCS following parturition of heifers lengthened calving to first service and calving to conception intervals by up to 2 days. Estimates of the environmental relationships between CI and SCC are rare in the literature, though estimated correlations in the present study indicate that the factors decreasing SCC are those making CI longer. However, more research is required to clarify this relationship.

In conclusion, regarding the genetic correlation of PMY with TMY, CI and SCC, selection on PMY would result in some increase in the amount of 305-d milk production, improved health status (reduced number of somatic cells), but causes a depletion of fertility traits (increased CI). Among persistency measures, P_{weller} was easier to compute, had higher h^2 , higher r_g with TMY, lower r_g with CI, and lower standard deviation of estimated genetic correlations with measures of SCC. The genetic correlation

between TMY and CI (0.71 to 0.732) was unfavorable, suggesting that selection for increased TMY increased CI. The absolute values of the correlations estimated with multiple-trait analyses were slightly higher than those of two-trait analyses.

Conflicts of interest

The authors declare they have no conflicts of interest with regard to the work presented in this report.

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