Estimation of nonadditive genetic impacts on lifetime performance through a grading-up breeding program with Holstein-Friesian
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Abstract

The aim of this study was to estimate the total lifetime milk production and non additive genetic effects (recombination and heterosis) of cows with different proportions of Holstein-Friesian genes, obtained from the Serbian Fleckvieh (SF) and the Holstein-Friesian (HF) crossbreeding program in Vojvodina. Upgrading of local breeds with the Holstein-Friesian breed in Vojvodina started in 1971 and continued 2008. Six genotypes of cows (F₁, R₁, R₂, R₃, R₄, R₅) were obtained with increasing percentage of Holstein genes, in order to attain purebred Holstein cows. Of all obtained genotypes, cows of genotype R₅ with a proportion of Holstein genes from 96.87 % had the highest lifetime milk production (20000 kg), followed by cows R₃ with 19950 kg (93.75 % HF genes) and cows R₅ with 19850 kg (98.44 % HF genes). Finally the process of upgrading resulted in pure Holsteins with 19780 kg of milk. The total lifetime production of milk fat did not show statistically significant difference (P>0.05) among the genotypes R₁ - R₅ which ranged from 675 to 690 kg. The pure Holstein obtained after sixth intermediate generations had the average lifetime milk fat production of 690 kg. With the increase in the proportion of Holstein-Friesian genes percentage of milk fat was decreased, so that the cows of genotypes R₅, R₄, R₃ and pure Holsteins, had less than 3.5 % milk fat. In relation to the total milk yield, the highest realized heterosis effect was observed in the cows of F₁ generation (h²F₁=594 kg), while the lowest was observed in generation R₅ (h²R₅=72 kg), where negative effect of recombination was also found (r²R₅=-77 kg). Positive values of the actual and relative of heterosis effect of the milk fat yield was observed in all genotypes, whereas the negative heterosis effect of the milk fat percentage was observed also in all genotypes, with the exception of R₁ and R₂ cows, in which the typical consequence of the positive recombination in the early crossed Holstein-generations was manifested.

Key words: upgrading, heterosis, recombination, milk, cattle

Introduction

Production of beef and milk is increasingly becoming an important economic sector because of the greater interest and need for these products. Therefore the selection is aimed to create early maturing and productive animals. Since the middle of the nineteenth century to the middle of the twentieth century, the predominant genotype of cattle in Vojvodina was Serbian Fleckvieh Cattle. In the late sixties and early seventies, both in the world and in our
country there has been an interest for the continental European breeds of cattle, such as Black-and-white Lowland Cattle, Danish Red and the Rotbunt cattle of Dutch origin and also the Jersey and Simmental (Fleckvieh) breeds. In the mid-seventies, serious shift in the development of dairy-type cattle in Vojvodina has occurred. But nowadays, we are witnessing that Holstein crosses are required for improvement of other local breeds (McAllister, 2002; Heins et al., 2006; Dechow et al., 2007; Swalve et al., 2008; Wall et al., 2003), and because of that reason there were exemplary initiatives in Serbia to improve the local Fleckvieh with Holstein-Friesian (Perišić, 2008). With organized importation of 660 high quality pregnant Holstein-Friesian heifers at PIK-Bečej and organized work on the genetic improvement of cattle in the population in Vojvodina was planned. There was an introduction of Black-and-white Holstein genes into the Serbian Fleckvieh breed with the aim being achieving 75 % Holstein genetics. The program was implemented in practice, as in some European countries (Germany, Netherlands, France, etc.), by crossing the pure Holstein-Friesian breed into the existing breeds. As a result of specific work in an organized breeding program, maintaining the same number of cows, relatively satisfactory genetic and phenotypic progress was achieved. In addition there was a complete change achieved in genetic make-up in the requested type of cows with the minimal amount of Holstein-Friesian females being imported. A grading-up program includes a series of back crossing in order to transmit new or specific genes into a population from another gene pool in order to replace the genes of the original population with different genes to achieve better performances and more profitable farms. One of the aims of the crossing is to exploit a very mysterious phenomenon, the heterosis. The heterosis is an advantageous deviation of the F1 progeny performance from the theoretical parental mean. From the second progeny generation, the level of this advantage will be influenced by a recombination effect due to the new reassortment of the parental genes in the offspring.

The aim of this paper was to present a long-standing work on the improvement of Serbian Fleckvieh cattle with the Holstein-Friesian breed in the controlled environment of the farm.

Data for the upgrading program were obtained from two cattle farms in Vojvodina, which started in year 1971 and 1975 by crossing the Serbian Fleckvieh (SF) cows with bulls of the Holstein-Friesian (HF) breed imported from the USA and Canada. The whole process lasted until 2008, and six cow genotypes (F1, R1, R2, R3, R4, R5) with different proportions of the Holstein genetics were obtained, in order to finally reach pure Holstein cows. Cows of F1 generation were created by mating SF cows to Holstein bulls, which had 50 % Holstein genes. Each new generation was produced by consequent use of semen of Holstein-Friesian sires. The grading-up was conducted until we obtained a cow with Holstein-Friesian blood with a proportion of over 99 %. As a result of many years of crossing, cows of the following genotypes and proportion of Holstein genes was obtained: R1 with 75 % HF genes, R2 with 87.5 % HF genes, R3 with 93.75 % HF genes, R4 with 96.88 % HF genes, R5 with 98.44 % HF genes, in order to obtain the last generation of HF cows with 100 % (>99 %) proportion of HF genes.

After examining the system records of each cow, the proportion of Holstein genes in the overall genotype, the crossing levels and the number of generations, then the total lifetime quantity of milk and the amount and percentage of milk fat production was determined.

For the investigated traits the adjusted mean value (LSM - Least Square Means) was computed by the multiple trait hierarchical linear model and Duncan’s post-hoc test for statistically significant differences among genotypes was applied (StatSoft, Inc., 2011). For the estimation of the nonadditive crossing effects the Dickerson-model (Dickerson, 1969) was applied. The realised heterosis ($h^2$) is the deviation of the real performance from a theoretical value which is estimated as the sum of the results of the blood proportion $x$ the performance of the pure breeds being present at a given genotype (genetic composition: $p_s p_d + p_d p_s$ where, $p_s = \text{Holstein-Friesian sire’s gene proportion, } p_d = \text{Serbian Fleckvieh sire’s gene proportion, } p_{sd} = \text{Holstein-Friesian dam’s gene proportion, } p_{ds} = \text{Serbian Fleckvieh dam’s gene proportion}$). The relative heterosis ($h^e$) represents, as a percentual value, the level of the previous realised heterosis. The additive component of the realised heterosis
heterosis (defined as individual heterosis, $h'$) has linear association with the degree of heterozygosity. It is made by the consecutive bisection from the heterosis appeared in the $F_1$ genotype. The recombination ($r^2$) is intended to characterize the distance of the realized heterosis from its additive component.

In the case of upgrading a breeding scheme there cannot be crossed sires with paternal heterosis. However, in the dairy production the crossbred stage of the dam contributes, as an environment, hardly to the individual performance. Therefore, the maternal heterosis is negligible. This is the reason that our investigation operates with the $h'$ only like it was in the processing reported by Egger-Danner (2005).

Results and discussion

Table 1 shows the adjusted mean (LSM) and the adjusted standard mean errors ($\text{SE}_{\text{LSM}}$) for the total lifetime milk production of cows with different genotypes. It can be seen that the purebred Serbian Fleckvieh (SF) cows produced 13,980 kg milk, while the improved cow generation $R_4$ with a share in Holstein genes of 96.87 % had the highest life milk production (20,000 kg), followed by genotype $R_3$ with 19,950 kg (93.75 % HF genes), then by the cows of genotype $R_2$ with 19,850 kg (98.44 % HF genes), and finally by the pure Holsteins (HF) gained 19,780 kg of milk.

Among the mentioned genotypes of the cows ($R_4$, $R_3$ and $R_2$) there was no statistically significant difference (P>0.05) as compared to both purebreds (SF and HF) where a significant statistical difference (P<0.05) was observed. The largest heterosis ($h'$) and realized heterosis ($h^R$) was recorded in cows of the $F_1$ generation (594 kg), while the increase in the proportion of genes of Holstein-Friesian cattle led to the decrease of additive component of heterosis from generation to generation by $\frac{1}{2}$. The highest level of recombination ($r^2 = 464$ kg) was recorded in cows with $R_3$ genotype which contained the proportion of Holstein genes from 93.75 %, while the cows of genotype $R_2$ contained a proportion of 87.50 % of HF genes had a negative recombination (-77 kg) and the least realized heterosis ($h^R$) of 72 kg. The smallest relative heterosis ($h'$) for lifetime milk yield - expressed as the percentual difference between the realized and expected production - appeared in the $R_2$ genotype (0.4 %), while cows of $F_1$ generation had the highest relative heterosis (3.5 %). Similar results have been published by other authors (Lopez-Villalobos et al., 2000; Sørensen et al., 2008; Freyer et al., 2008), in their research which had the aim to blend their local cattle breeds with purebred Holsteins.

In the research of Gáspár (1995) the average total lifetime milk production of the purebred Holstein-Friesians was 18,270 kg (with 1774 days long herd life, and with 2.12 lactations) in a Hungarian crossing process. Kräußlich (2002) published data about lifetime performance (15,238 kg milk) of Simmental in Switzerland and he pointed out the superiority of the Simmental×Holstein crossed cows

Table 1. The lifetime milk production and non additive genetic effect in different cow genotypes

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Proportion of Holstein-gene, %</th>
<th>No. of cows</th>
<th>Total life time milk yield, kg LSM ± SE$_{\text{LSM}}$</th>
<th>Additive component of heterosis, kg ($h'$)</th>
<th>Recombination, kg($r^2$)</th>
<th>Realised heterosis, kg($h^R$)</th>
<th>Relative heterosis, % ($h'$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SF</td>
<td>0.00</td>
<td>277</td>
<td>13980*± 353</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.0</td>
</tr>
<tr>
<td>$F_1$</td>
<td>50.00</td>
<td>254</td>
<td>17470*± 354</td>
<td>594</td>
<td>-</td>
<td>594</td>
<td>3.5**</td>
</tr>
<tr>
<td>$R_1$</td>
<td>75.00</td>
<td>243</td>
<td>18860*± 347</td>
<td>297</td>
<td>240</td>
<td>537</td>
<td>2.9**</td>
</tr>
<tr>
<td>$R_2$</td>
<td>87.50</td>
<td>223</td>
<td>19120*± 332</td>
<td>149</td>
<td>-77</td>
<td>72</td>
<td>0.4</td>
</tr>
<tr>
<td>$R_3$</td>
<td>93.75</td>
<td>265</td>
<td>19950*± 308</td>
<td>74</td>
<td>464</td>
<td>538</td>
<td>2.8**</td>
</tr>
<tr>
<td>$R_4$</td>
<td>96.87</td>
<td>967</td>
<td>20000*± 199</td>
<td>37</td>
<td>371</td>
<td>408</td>
<td>2.1*</td>
</tr>
<tr>
<td>$R_5$</td>
<td>98.45</td>
<td>2053</td>
<td>19850*± 175</td>
<td>19</td>
<td>149</td>
<td>168</td>
<td>0.9</td>
</tr>
<tr>
<td>HF</td>
<td>100</td>
<td>6996</td>
<td>19780*± 143</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

*$abcd$ - different subscribed letters show significant differences (P<0.05)

* , ** , *** - number of subscribed stars show a significant difference from the Serbian Fleckvieh (*P<0.05; **P<0.01; *** P<0.001)
in the total amount of produced milk (24,793 kg milk). This yield was between 20,140 and 24,491 kg reported for 59,7560 Holstein and holsteinized cows in Germany (Swalve, 2007). Whereas the average recorded value of lifetime performance of Holsteins in the Netherland was 30,543 kg of milk and 2,406 kg of milk fat and protein (Veeipro Magazine, 2009). In their investigation Fürst and Sölkner (1994) come to a conclusion that in crossed populations the non additive genetic impacts result especially in an increase of milk yield.

The total lifetime fat production (Table 2) among the genotypes from R₁ to R₅ did not show statistically significant difference (P>0.05) which ranged from 675 to 690 kg, while in comparison with the SF breed and F₁ generation they showed statistically proven differences (P<0.05). The purebred Holsteins had an average life production of milk fat of 690 kg, without a significant statistical difference (P>0.05) when compared to cows of genotype from R₁ to R₅. With the increase in the proportion of genes of the Holstein-Friesian, the additive component of heterosis (h₁) was decreased. Positive values of the realised and relative heterosis effect on milk fat yield were observed in all genotypes. In cows of genotype R₁, with proportion of 75 % Holstein genes recorded the highest realised (h² = 32.4 kg) and relative (h' = 5.1 %) heterosis, while the cows of the genotype R₅ with the proportion of 98.45 % Holstein genes had the least individual heterosis. Therefore the lowest realized heterosis, negative recombination, and relative heterosis was not able to be determined, which can be explained by the fact that it is already the sixth improved generation of cattle, where there is an advanced gene fixation and a reduced chance for recombination.

McAllister et al. (1994) recorded high relative heterosis in the total lifetime milk and milk fat production (from 16.5 to 20 %, respectively) in their work about the Holstein and Ayrshire crossing, while Van Raden and Sanders (2003) studied the length of a productive life in the different dairy crossed progenies sired by the Holstein and estimated heterosis which was low (1.2 %), which can be the result of the environmental factors (Bryant et al., 2007).

The Table 3 shows the milk fat content by the genotypes, where we observed that SF cows had the highest percentage of milk fat (3.68 %). The milk fat percentage decreased with the process of grading-up, so that the cow genotypes R₀, R₄, R₅, and HF had less than 3.5 % milk fat. With the increase of the total lifetime milk production from generation to generation the milk fat content is decreasing, as it was expected based on knowledge from single lactation performances, given that these traits are in negative genetic correlation. Negative values of realised and expected heterosis effect on milk fat was observed in almost all genotypes, whereas the positive heterosis effect was only observed in the R₁ and R₂ genotype which is a typical consequence of the positive recombination in the early crossed generations.

Table 2. The lifetime milk fat production and non additive genetic effect in different cow genotypes

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Proportion of Holstein-gene, %</th>
<th>No. of cows</th>
<th>Total life time milk fat yield, kg LSM ± SE₉₅</th>
<th>Additive component of heterosis, kg (h')</th>
<th>Recombination, kg (r')</th>
<th>Realised heterosis, kg (h)</th>
<th>Relative heterosis, % (h')</th>
</tr>
</thead>
<tbody>
<tr>
<td>SF</td>
<td>0.00</td>
<td>277</td>
<td>514±11.4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.0</td>
</tr>
<tr>
<td>F₁</td>
<td>50.00</td>
<td>254</td>
<td>617±11.4</td>
<td>20.0</td>
<td>-</td>
<td>20.0</td>
<td>3.4**</td>
</tr>
<tr>
<td>R₁</td>
<td>75.00</td>
<td>243</td>
<td>675±11.2</td>
<td>10.0</td>
<td>22.4</td>
<td>32.4</td>
<td>5.1***</td>
</tr>
<tr>
<td>R₂</td>
<td>87.50</td>
<td>223</td>
<td>675±10.7</td>
<td>5.0</td>
<td>8.5</td>
<td>13.5</td>
<td>2.1*</td>
</tr>
<tr>
<td>R₃</td>
<td>93.75</td>
<td>265</td>
<td>690±10.0</td>
<td>2.5</td>
<td>11.3</td>
<td>13.8</td>
<td>2.1*</td>
</tr>
<tr>
<td>R₄</td>
<td>96.87</td>
<td>967</td>
<td>686±6.4</td>
<td>1.3</td>
<td>3.4</td>
<td>4.6</td>
<td>0.7</td>
</tr>
<tr>
<td>R₅</td>
<td>98.45</td>
<td>2053</td>
<td>683±5.6</td>
<td>0.6</td>
<td>-5.7</td>
<td>0.1</td>
<td>0.0</td>
</tr>
<tr>
<td>HF</td>
<td>100</td>
<td>6996</td>
<td>690±4.6</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

abc - different subscribed letters show significant differences (P<0.05)
*; **; *** - number of subscribed stars show a significant difference from the Serbian Fleckvieh (*P<0.05; **P<0.01; *** P<0.001)
Lopez-Villalobos et al. (2000) suggested that modern breeding methods, such as artificial insemination, embryo transfer, breeding programs such as TEAM or MOET, use sexed semen, reducing the number of bull lines leading to a reduction in genetic variability, resulting in an increase in the coefficient of inbreeding, while the rotational crossing of two or more breeds contributes to the manifestation of the effects of crossing and maintenance of genetic variability, which is important in maintaining the balance between production costs and future market requirements. When the crossed animals will be bred further (e.g. F₁ x F₁ mating, or grading-up is carried out), the breeders have to realise the loss in traits caused by non-additive genetic impacts, namely by recombination, which could contradict the valued heterosis that is expected. This genetic effect may sometimes be greater than the expected heterosis effect, although in each case the effect is to be expected in reverse. The success of the crossing requires a consistent and strict selection from the breeder.

Dickerson (1969) has shown that the maternal heterosis cancel out when the difference is taken between the crossbred and straight bred females mated to the sires of a different breed. Estimate of maternal heterosis in the beef production is applicable to any system of mating involving purebred bulls and either F₁ dams or crossbred females produced in terminal or rotational crossbreeding systems because its effects expressed in the calf are always present in approximately the same proportion to effects of individual heterosis expected. Alternative designs employing F₁ bulls may provide estimates of maternal heterosis without any effect of recombination but the results for reproduction are confounded with any effect of heterosis on the fertility of crossbred bulls.

Conclusions

With the use of crossing Serbian Fleckvieh cows with Holstein bulls, there is a possibility that after 38 years with a good working organization and breeding work, to get pure Holsteins after six generations. There was a simultaneous increase in the number of cows, and satisfactory genetic and phenotypic improvement was carried out as well as a total change in genetic composition. As result, we came to the desired type of Holstein-Friesian. With the increase in the proportion of Holstein-Friesian genes bred from generation to generation with optimal zootechnical requirements and care. There has been an increase in the total amount of milk, previously from 13,980 kg in SF to 19,780 kg in HF. However, the milk fat decreased from 3.68 % to 3.49 % but there has been an increase in the total milk fat yield from 514 kg to 690 kg.

Research indicates that a planned crossing leads to the optimal gene recombination, which results in the increase in production and maintenance of the necessary variability for the purpose of continual positive success of selection on the observed properties.
Procjena neaditivnog genetskog učinka na životne performanse krava kroz “grading-up” uzgojni program i korištenje holštajn-frizijske pasmine

Sažetak

Cilj ovog istraživanja bio je prikazati procjenu ukupne životne proizvodnje mlijeka i neaditivan genetski učinak (rekonbinacija i heterozis) krava s različitim udjelima gena holštajn-frizijske pasmine, dobivenih pretapanjem domačeg šarenog goveda (SF) u holštajn-frizijsku pasminu goveda (HF) u Vojvodini. Pretapanje domaće pasmine u holštajn-frizijsku pasminu obuhvatio je razdoblje od 1971. do 2008. godine, a dobiveno je šest genotipova krava (F1, RF, RR, R1, R2, R3) s različitim udjelom holštajn gena, do dobivanja krava holštajn pasmina. Od svih dobivenih genotipova, krave R3 s udjelom holštajn gena od 96,87 % imale su najveću životnu proizvodnju mlijeka (20.000 kg), zatim slijede krave R1 sa 19.950 kg (93,75 % HF gena) i krave R2 sa 19.850 kg (98,44 % HF gena), da bi na kraju procesa pretapanja dobiveni čisti holštajn imao 19.780 kg mliječne masti. U životnoj proizvodnji mliječne masti između dobivenih kriza (R1-R3) nije utvrđena statistički značajna razlika (P>0.05) i kretala se od 675 do 690 kg, dok je u usporedbi s roditeljskim čistim pasminama utvrđena statistički značajna razlika (P<0.05). Dobiveni čisti holštajn nakon šeste generacije u prosjeku imao je životnu proizvodnju mliječne masti od 690 kg. S porastom udjela gena holštajn-frizijske pasmine, postotak mliječne masti je opadao, tako da su krave R1, R3, R5, i čisti holštajn imale manje od 35,0 % mliječne masti. U odnosu na ukupnu količinu mliječne masti, najstaknutiliji realizirani heterozis učinak zapažen je kod F1 krava (h21 =594 kg), dok je najniži zapažen kod R3 krava (h21 =72 kg), kod kojih se pojavio negativan efekt rekonbinacije (h21 =-77 kg). Pozitivne vrijednosti ostvarenog i relativnog heterozis efekta na svojstvo prinosa mliječne masti uočene su u svim genotipovima, dok je negativan heterozis efekt za postotak mliječne masti, osim u krava R1 i R2, uočen u svim genotipovima krava, što je tipična posljedica utjecaja holštajn-frizijskog genotipa.

Ključne riječi: pretapanje pasmina, heterozis, rekonbinacija, mlijeko, goveda

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