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*Contributed equally to the study.

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Corresponding author: Nicolás Lopez-Villalobos; Email: n.lopez-villalobos@massey.ac.nz

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Effects of breed and genomic inbreeding on milk, fat and protein lactation yields and fertility traits in pasture-based dairy cows in Argentina

María José Beribe^{1,2,*}, Hugo Adrián Carignano^{3,4,*} , Mario Andrés Poli^{5,6} and Nicolás Lopez-Villalobos⁷

¹Estación Experimental Agropecuaria Pergamino, Instituto Nacional de Tecnología Agropecuaria, Pergamino, Buenos Aires, Argentina; ²Facultad de Ciencias Bioquímicas y Farmacéuticas, Universidad Nacional de Rosario, Rosario, Argentina; ³Instituto de Virología e Innovaciones Tecnológicas, Instituto Nacional de Tecnología Agropecuaria – Consejo Nacional de Investigaciones Científicas y Técnicas, Hurlingham, Argentina; ⁴Consejo Nacional de Investigaciones Científicas y Técnicas, Ciudad Autónoma de Buenos Aires, Argentina; ⁵Instituto de Genética, Instituto Nacional de Tecnología Agropecuaria, Hurlingham, Buenos Aires, Argentina; ⁶Facultad de Ciencias Agrarias y Veterinaria, Universidad del Salvador, Pilar, Buenos Aires, Argentina and ⁷School of Agriculture and Environment, Massey University, Palmerston North, New Zealand

Abstract

The objective of the current study was to evaluate the effects of breed and genomic inbreeding on 305-day lactation yields of milk, fat and protein; and fertility traits of pasture-based dairy cows in Argentina. The genomic inbreeding and heterozygosity of 890 first-lactation cows and 27 bulls were calculated through methods based on the genomic relationship matrix and run of homozygosity using 44 174 single-nucleotide polymorphisms. Cows were classified into four breed groups: Holstein, Holstein crossbred, Holstein–Jersey crossbred and Jersey crossbred. The effect of genomic inbreeding was not significant on production traits, but inbred cows increased 3.0 days calving to conception interval (CCI) per 1% genomic inbreeding. On average, purebred Holstein cows produced 1119 kg milk, 22 kg fat and 30 kg protein more than Jersey crossbred cows. In the case of the fertility traits, Jersey crossbred cows had 45 days shorter CCI than purebred Holstein cows. A possible reason for the non-significant effects of genomic inbreeding of production and fertility traits is that these effects were evaluated in a crossbred population in which rates of heterozygosity would operate to some extent in the opposite direction to rates of genomic inbreeding.

Introduction

The dominant breed of dairy cattle in Argentina is Holstein (Gastaldi *et al.*, 2020). Traditionally, the cows have been produced using dairy sires from the USA of high-genetic potential for milk production, which are suitable for indoor systems using total mixed rations (Lazzarini *et al.*, 2019). Contradictorily to the intensive indoor systems, in Argentina 149/155 of dairy farms are pastoral (Gastaldi *et al.*, 2020). Feeding is based mainly on grazing alfalfa (*Medicago sativa* L.) throughout the year, in combination with silage and concentrates (Gastaldi *et al.*, 2020). Some dairy farmers have used Jersey sires to incorporate Jersey × Holstein crossbred cows, which produce lower milk yields but higher proportions of fat and protein. In 2021, approximately 3 197 025 of 3 475 028 semen doses from dairy bulls that were marketed in Argentina were from Holstein, 0.05 from Jersey and the rest from other dairy breeds such as Brown Swiss and Guernsey (Cámara Argentina de Biotecnología de la Reproducción e Inseminación Artificial, 2021).

Inbreeding depression is the reduction of an individual's fitness due to fixation of deleterious recessive genes (Falconer and Mackay, 1996). Progeny that results from mating of genetically related animals results in high inbreeding coefficient. A inbreeding coefficient has been originally defined as the probability that an individual inherits two identical alleles at the same locus from the parents (Malécot, 1969) or as the correlation between two homologous alleles in uniting gametes (Wright, 1922). Therefore, inbreeding could also be reflected as an increase in autozygosity (i.e. homozygosity due to the inheritance of identical alleles by offspring) and a loss of genetic variability (deficit in heterozygosity). Inbreeding coefficients can be calculated through either ancestral pedigrees or genomic analysis. Parentage data can be used to construct the relationship matrix (A) among the individuals represented in the pedigree file. This matrix is symmetric, the elements of the diagonal represent the genetic relationship among the individuals of the pedigree and the diagonal elements represent the degree of inbreeding of each animal in the pedigree, where the value is 1 plus the expected inbreeding coefficient (Falconer and Mackay, 1996). Analysis of inbreeding in large populations poses difficulty due to the need for a simple and accurate, but high-throughput method to calculate the inbreeding coefficient. Such calculations of inbreeding predict only the expected proportion of animal genome that is considered identical by descent (Nietlisbach *et al.*, 2017). Inaccurate estimates of inbreeding coefficients can arise if pedigrees are small or incomplete, missing parentage can underestimate actual rates of inbreeding (Nietlisbach *et al.*, 2017; Gutiérrez-Reinoso *et al.*, 2022).

Currently, the availability of molecular markers distributed throughout the whole genome enables the estimation of genomebased inbreeding coefficients without the need to record extensive genealogy or parental relationship among individuals (Leutenegger et al., 2003). Therefore, genomic-based inbreeding coefficients demand less time and effort than pedigree-based measures. Besides, they can be more accurate due to their capacity to account for variation in the random process of meiotic recombination (Mendelian sampling). The simplest estimations of this realized inbreeding coefficient rely on single-nucleotide polymorphism (SNP) analysis of different genomic relationship matrix (GRM) formulations (equivalent to the F calculation when the A matrix is used) (VanRaden, 2008) or analysing multi-SNP consecutive DNA stretches, known as run of homozygosity (ROH) (Keller et al., 2011; Purfield et al., 2012). Depending on the GRMs utilized, the single-SNP inbreeding coefficients could describe (1) deviations from Hardy-Weinberg equilibrium (HWE) (F_{HOM}), (2) correlations between uniting gametes (F_{UNI}) and (3) the variance of additive genetic values (F_{GRM}) (Villanueva *et al.*, 2021). On the other hand, the ROH-based inbreeding coefficient, calculated as the sum of the ROHs identified in an individual divided by the total genome length, ideally represents the proportion of the individual's autosomal genome that is autozygous (Howrigan et al., 2011).

Similarly, the rate of heterozygosity of an individual can be calculated from genotypes and requires no knowledge of ancestry in contrast to pedigree inbreeding (Iversen *et al.*, 2019).

The effect of inbreeding on production and reproduction traits has been well documented in Holstein-Friesian dairy cattle, depicting that as inbreeding increases milk production, fertility is reduced (Cassell et al., 2003; McParland et al., 2007). Although a similar relationship has been reported in the Jersey breed, the effect of inbreeding has a greater influence on milk yield in the Holstein-Friesian breed (Maiwashe et al., 2008; Pryce et al., 2014). Other studies have reported that fat and protein yield follow the same trend as milk yield, but inbreeding has substantially less influence on change of milk yield, resulting in small regression coefficients (Dezetter et al., 2015; Doekes et al., 2019). Also, increases in inbreeding coefficients have been associated with larger concentrations of somatic cell counts (SCCs) in milk, with greater incidence seen in older animals (McParland et al., 2007). Contrary to these findings, some studies have found no significant effect of inbreeding on SCCs (Rokouei et al., 2010; Dezetter et al., 2015).

Crossbreeding is the mating of individuals from different lines, breeds or populations (Lopez-Villalobos *et al.*, 2000). As a breeding strategy, crossbreeding offers the removal of the negative effects associated with inbreeding depression in particular traits associated with fitness and survival (Falconer and Mackay, 1996). Crossbreeding also offers a potentially attractive avenue for farmers to improve economic efficiency by using breed complementary and exploiting heterosis for milk production, fertility and survival (Buckley *et al.*, 2014). The heterozygosity rate has been indicated as a proxy for heterosis (Iversen *et al.*, 2019); in F1 crosses all animals are expected to be heterozygotes (maximum heterozygosity rate), but in backcross and subsequent generations a reduction in heterozygotes frequency would be observed, altogether a heterosis decline. Therefore, given the interplay between crossbreeding, inbreeding and heterozygosity, quantification of these values in crossbreed cattle would be relevant for management decisions in dairy cattle.

Comparisons between Jersey, Holstein and Jersey × Holstein crossbred cows for production and reproduction traits in pasturebased production systems of Argentina have been reported in the literature (Baudracco *et al.*, 2011; Mancuso and Marini, 2012; Biga *et al.*, 2022). To our knowledge, there are no studies in the literature that report the rate of genomic inbreeding and heterozygosity calculated using genetic markers in a crossbred population and the effects of genomic inbreeding in dairy cattle in Argentina. The objective of the current research was to evaluate breed and genomic inbreeding on lactation milk yields, fat and protein content and fertility in pasture-based dairy cows from Argentina.

Materials and methods

Data

Study population

Records of 20 005 Holstein and Holstein × Jersey cows born between 1994 and 2010 were obtained from 37 dairy farms. Dates of birth, services, calving, dairy controls, drying-off, transfers and rejection added up to a total of 755 141 records. The cows (4804/20 005 Holstein and 5201/20 005 Holstein × Jersey) belong to 30 half-sisters families. In particular, the crossbred groups considered in this study were the following: H: purebred cows with 1.00 Holstein (639); HX: backcross to Holstein with 0.75 Holstein and 0.25 Jersey (64); HJ: first crossbred Holstein– Jersey with 0.50 Holstein and 0.50 Jersey (157); JX: backcross to Jersey with 0.25 Holstein and 0.75 Jersey (30).

The dairy farms are located in the central dairy basin of Argentina (central-eastern region). On each farm, two daily milkings were performed within a semi-stabled feeding system with cows outside grazing alfalfa during the hot summer months and oats or barley in the winter. The diet was systematically supplemented with maize silage, soybean meal and concentrates elaborated in house containing 0.16–0.18 of crude protein.

The cow's reproductive management system involves continuous calving, where calving periods are uniformly distributed throughout the year.

Genotyping and quality control

A total of 970 cows (718 purebred Holstein and 252 crossbred Holstein × Jersey) and 29 bulls (24 Holstein and 5 Jersey) were chosen for genotyping as described elsewhere (Carignano *et al.*, 2018). Briefly, quality control of the genotypic data was carried out using a set of tools and routines provided by PLINK v1.9 (Purcell *et al.*, 2007). Individuals with a genotype call rate (CR_{IND}) <90% were excluded from further analysis. SNPs with a call rate (CR_{SNPs}) <90%, a deviation from HWE $P < 1.10^{-8}$ and minor allele frequency <0.01 were removed from the study. After the quality control, the data set for genomic inbreeding calculations comprised of 44 174 SNPs and 917 individuals (890 cows and 27 bulls).

Genomic inbreeding and heterozygosity coefficients

Genomic-based individual's inbreeding coefficients (F) were calculated using different approaches as follows:

 F_{ROH} : The inbreeding coefficient F_{ROH} for each cow was calculated using the - *-homozyg* routine implemented in PLINK version 1.9 (Purcell *et al.*, 2007). The runs of homozygous genotypes for each individual were defined using sliding windows of 50 SNPs across the genome, requiring 25 homozygous SNPs spanning a ≥ 1000 kb distance. A sliding window hit ('homozygous') contained at most one heterozygous SNP and none missing calls. An SNP was included in an ROH segment if the hit rate (proportion of 'homozygous' windows that overlap that position) was >0.05. The final ROH segments were constrained to a maximum interval between two consecutive SNPs of 500 kb, allowing one heterozygous genotype and a minimal density of 1 SNP/100 kb.

The coefficient F_{ROH} was defined as the proportion of the autosomal genome in runs of homozygosity:

$$F_{\rm ROH} = \frac{L_{\rm ROH}}{L_{\rm aut}}$$

where $L_{\rm ROH}$ is the total length of all ROH segments identified in an individual and $L_{\rm aut}$ is the length of the autosomal genome covered by SNPs (i.e. between the first SNP and the last SNP per chromosome for all autosomal chromosomes).

 F_{GRM} : The inbreeding coefficient F_{GRM} for each cow was calculated using the diagonal elements of the GRM according to VanRaden (2008):

$$F_{\rm GRM} = \frac{1}{N} \sum_{i=1}^{N} \frac{(X_i - 2P_i)^2}{h_i} - 1$$

where X_i is the genotype coded as the number of reference allele copies for the *i*th SNP; P_i is the observed frequency of this allele (i.e. the allele whose homozygous genotype was coded as '0'), N is the total number of markers and $h_i = 2P_i(1 - P_i)$ is the expected heterozygosity.

 $F_{\rm HOM}$: This inbreeding coefficient was based on the homozygous excess and is defined as

$$F_{\text{HOM}} = \frac{O(hom) - E(hom)}{N - E(hom)} = 1 - \frac{1}{N} \sum_{i=1}^{N} \frac{X_i(2 - X_i)}{h_i}$$

where O(hom) is the observed number of homozygous markers of the individual and E(hom) is the expected number of homozygous markers under the HWE calculated from the allele frequencies estimated on the sample. The expected number of homozygotes is calculated assuming HWE as $1 - 2P_i(1 - P_i)$.

 F_{UNI} : The inbreeding coefficient F_{UNI} is based on the correlation between uniting gametes following the method proposed by Ritland (1996):

$$F_{\text{UNI}} = \frac{1}{N} \sum_{i=1}^{N} \frac{X_i^2 - (1+2P_i)X_i + 2P_i^2}{h_i}$$

where $2P_i(1 - P_i) = 1$ if X_i is heterozygous and = 0 if it is homozygous.

The inbreeding coefficients F_{GRM} , F_{HOM} and F_{UNI} were calculated using the –ibc routine implemented in GCTA v1.24 software (Yang *et al.*, 2011).

Heterozygosity was measured as the proportion of heterozygous marker genotypes for each individual as (N - O)/N, where

Phenotypic traits

The initial production data set included 316 855 monthly herd tests from 48 367 lactations belonging to 20 005 cows. The lactation curves for daily milk, fat and protein production for each cow-lactation were modelled using a sixth-order Legendre polynomial. Predicted daily yields were then used to calculate 305-day lactation yields of milk (MY305), fat yield (FY305) and protein yield (PY305), for each lactation of each cow (Beribe, 2020).

Two fertility traits were calculated, calving to first service interval (CFSI) and calving to conception interval (CCI). CFSI was calculated as the number of days between calving date and the first service date, and CCI was calculated as the number of days between calving date and conception date.

Finally, for this study the values of MY305, FY305, PY305, CFSI and CCI were merged with the 890 first-lactation cows that had genotypic data. Not all cows that were genotyped and passed quality control for genetic markers had the full set of phenotypic traits.

Statistical analysis

The effects of breed and inbreeding were estimated using the following mixed model:

$$y_{ijklm} = \mu + \alpha_i + \beta_i + \gamma_k + \delta x_l + \tau z_m + \varepsilon_{ijklm}$$

where:

 y_{ijklm} is any of the traits evaluated: MY305, FY305, PY305, CFSI and CCI.

 α_i is the fixed effect of breed group with four classes (H: proportion of Holstein = 1, HX: proportion of Holstein = 0.75, HJ: proportion of Holstein = 0.5 and JX: proportion of Holstein = 0.25).

 β_j is the fixed effect of calving season with five classes: November–January, February–April, May–June, July–August and September–October.

 γ_k is the random effect of the contemporary group, defined as the group of cows that started lactation in the same herd and year.

 δ is the regression coefficient of the dependent trait on age at calving x_l .

 τ is the regression coefficient of the dependent trait on the genomic inbreeding coefficients z_m (expressed as a percentage).

 ε_{ijklm} is the random residual associated with observation y_{ijklm} .

Calving season was defined based on monthly average temperatures and ensuring a representative number of records in each class (>19 records).

Least-squares means and standard errors were obtained for each breed group and calving season and used for multiple mean comparisons using Fisher least significant difference test.

Analyses were performed using the MIXED procedure of SAS v9.4 (SAS* Institute Inc., 2013, Cary, NC, USA).

Comparisons of mean rates of genomic inbreeding and heterozygosity among the different breed groups were performed using individual *t*-tests.

Results

Descriptive statistics for the production and fertility traits and measures of genomic inbreeding coefficients are presented in Table 1. The fertility traits presented greater variability (higher coefficient of variation [CV]) than production traits. For example, CCI was in the range of 18–673 days (CV = 64%), whereas FY305 was in the range of 156–281 kg (CV = 12%).

The distributional properties of $F_{\rm GRM}$, $F_{\rm HOM}$ and $F_{\rm UNI}$ coefficients were similar; median at -2.69 ($F_{\rm GRM}$), -1.31 ($F_{\rm HOM}$) and -2.26 ($F_{\rm UNI}$) with (min to max) values at (-13.90 to 46.11), (-18.72 to 16.04) and (-11.38 to 24.68), respectively. The range of the $F_{\rm ROH}$ values was 0.00–19.00, with positive values for the mean and median. The range of heterozygosity was from 0.29 to 0.39 with median and mean values of 0.35 and low standard deviation (Table 1).

Scatter plots and distributions (histograms) of each of the genomic inbreeding coefficients and pairwise Pearson's correlation coefficients are presented in Fig. 1. All correlation coefficients were statistically significant (P < 0.05). The correlation between $F_{\rm ROH}$ and $F_{\rm UNI}$ was strong positive (0.70), but the correlation between $F_{\rm ROH}$ and $F_{\rm HOM}$ was stronger (0.86). No correlation between $F_{\rm ROH}$ and $F_{\rm GRM}$ (-0.08) was observed. Among the inbreeding coefficients derived from the genomic matrix, a weak negative correlation was found between $F_{\rm GRM}$ and $F_{\rm HOM}$ (-0.36). The correlations between $F_{\rm GRM}$ and $F_{\rm UNI}$, and between $F_{\rm HOM}$ and $F_{\rm UNI}$ were moderately positive.

Mixed models were used to study the effects of breed and genomic inbreeding on production and fertility traits. For milk production traits, F-statistics indicated that breed and calving age explained the majority of the variation (Table 2). Besides, calving season had a significant effect on CFSI. In the case of F coefficients, only $F_{\rm ROH}$ had a significant effect for CCI (P = 0.019). The rest of the genomic F coefficients evaluated did not influence MY305, FY305, PY305 or CFSI in the studied population (data not shown). A simple linear regression analysis of CCI on $F_{\rm ROH}$ showed that CCI increased by 4.87 days per 1% increase of $F_{\rm ROH}$ (P < 0.0001; Fig. 2(a)), but when corrected (full model) by breed, season calving and age at calving the partial regression coefficient was 3.0 days per 1% inbreeding. Similarly, regression analysis of MY305 on $F_{\rm GRM}$ showed that MY305 reduced by 22.7 kg per 1% of F_{GRM} (Fig. 2(b)). However, when the effect of $F_{\rm GRM}$ was evaluated considering the full model the partial regression coefficient was not statistically significant.

Effects of genomic F_{GRM} , F_{HOM} and F_{UNI} on the productive and fertility traits were not significant (data not shown).

The effect of breed group on productivity and fertility was evaluated through least-squares analysis (Table 3). In general, milk yield increased as the proportion of Holstein breed increased in the population. The differences between first-lactation purebred Holstein and JX crossbred cows were 1119 kg milk, 22 kg fat and 30 kg protein (P < 0.001). The breed groups' mean comparisons for CFSI were not significantly different (Table 2). However, a trend (P = 0.099) was observed to reduce CCI as the proportion of Jersey increased; JX cows had 45 days shorter CCI than H purebred.

The relationship between heterozygosity and $F_{\rm ROH}$ across the different breed groups is presented in Fig. 3. First crossbred HJ cows which are close to 0.50 H and 0.50 Jersey had the highest heterozygosity and the lowest $F_{\rm ROH}$ coefficients, whereas purebred H cows tended to have the lowest values of heterozygosity and highest values of $F_{\rm ROH}$ coefficients.

In accordance with Fig. 3, crossbred HJ cows had the lowest (P < 0.05) rate of inbreeding, measured by the $F_{\rm ROH}$ coefficient, and the highest (P < 0.05) rate of heterozygosity.

Discussion

The current study investigated the effect of breed and genomic inbreeding coefficients on production and fertility traits in pasture-based dairy cows from commercial farms of Argentina. The average rate of milk production found in the current study (5687 kg milk, 213 kg fat and 197 kg protein per cow) was similar to that reported for the Holstein breed in Argentina (5760 kg milk, 207 kg fat and 193 kg protein per cow, respectively) (FCA, 2016). Subtle differences in cows' productivity between the current research and national herd could be explained by variation in food supplementation intensity and farm management conditions as shown in Lazzarini *et al.* (2019).

The mean CFSI was 87.5 ± 43.2 days, which was similar to values reported in USA for Holstein cows (81 ± 4 days; Mullen *et al.*, 2015) and UK Holstein and Jersey cows (71 ± 1.9 days; Coffey *et al.*, 2016). The average CCI (155 ± 99.3 days) was in

Table 1. Descriptive statistics for production and fertility traits and different measures of genomic inbreeding of pasture-based dairy cows from commercial herds of Argentina

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Trait ^a	Ν	Mean	Median	SD	CV	Min	Мах
MY305 (kg)	833	5687	5631	84.9	15	3295	8375
FY305 (kg)	816	213	213	26.3	12	145	308
PY305 (kg)	816	197	195	26.8	14	156	281
CFSI (days)	890	87.5	75.5	43.2	49	15	297
CCI (days)	890	155.0	126.0	99.3	64	18	673
F _{GRM}	890	-1.95	-2.69	6.2		-13.9	46.1
F _{HOM}	890	-2.06	-1.31	6.3		-18.7	16.0
F _{UNI}	890	-2.01	-2.26	3.5		-11.4	24.7
F _{ROH}	890	5.22	5.39	3.5		0.00	19.0
Heterozygosity	890	0.35	0.35	0.02		0.29	0.39

N, number of cows; SD, standard deviation; CV, coefficient of variation (%); Min, minimum value; Max, maximum value.

^aMY305, 305-day milk production; FY305, 305-day fat production; PY305, 305-day protein production; CFSI, calving to first service interval; CCI, calving to conception interval; *F*_{ROH}, coefficient of genomic inbreeding calculated based on proportion of the autosomal genome in runs of homozygosity, *F*_{GRM}, coefficient of genomic inbreeding obtained from the diagonal elements of the genomic relationship matrix, *F*_{HOM}, coefficient of genomic inbreeding calculated based in the homozygous excess and *F*_{UNI}, coefficient of genomic inbreeding calculated based on the correlation between uniting gametes. Heterozygosity was calculated as the proportion of heterozygous marker genotypes for each individual.



Figure 1. Pearson correlations and *P*-values (within brackets) (above diagonal), scatter plots (below diagonal) between each pair of the four genomic inbreeding estimators and densities (diagonal) of the inbreeding estimators of dairy cows in commercial herds of Argentina. ${}^{1}F_{ROH}$, coefficient of genomic inbreeding calculated based on proportion of the autosomal genome in runs of homozygosity; ${}^{2}F_{GRM}$, coefficient of genomic inbreeding obtained from the diagonal elements of the genomic relationship matrix; ${}^{3}F_{HOM}$, coefficient of genomic inbreeding calculated based in the homozygous excess; ${}^{4}F_{UNI}$, coefficient of genomic inbreeding calculated based on the correlation between uniting gametes.

line with those reported in the USA (115–177 days) and Argentina (92–157 days) (Dutour and Melucci, 2011). However, much shorter CCI values have been reported in Canada (102–108 days), New Zealand (90–101 days) and UK (100–115 days) (reviewed by Dutour and Melucci, 2011). The differences between our results and other studies are explained by the differences in genetic, management and feeding programmes.

Despite the coefficient of inbreeding being an important variable for breeding mating management in dairy herds, aiming to reduce inbreeding depression, there is currently no consensus on which genomic inbreeding coefficient estimator is appropriate (Dadousis *et al.*, 2022). Furthermore, the evaluation of genomic inbreeding coefficients in crossbred cattle populations has been scarcely reported. In the current study, the range of genomic inbreeding coefficient values for F_{GRM} , F_{HOM} and F_{UNI} were (as percentage) [-13.90 to 46.11], [-18.72 to 16.04] and [-11.38 to 24.68], respectively. These values did not meet Wright's original definitions of inbreeding, since these estimates of genomic

inbreeding are outside of the ranges allowed by probabilities (0-100%). Alternatively, these estimators can be interpreted as measures of genotypic variability (heterozygosity) in the current population compared to a base population (Villanueva et al., 2021). However, it is important to note that the allelic frequencies in the base population are often unknown and must be estimated from the current population assuming HWE (Villanueva et al., 2021; Dadousis et al., 2022). In this regard, in a study by Villanueva et al. (2021) F_{GRM} , F_{UNI} and F_{HOM} were analysed through simulations using different scenarios of allelic frequencies in the base and current populations. They described that F_{GRM} and F_{UNI} could indicate variability increasing (genomic inbreeding <0) in the current population when in fact it has decreased (genomic inbreeding >0), or vice-versa. Also, F_{GRM} and F_{UNI} could indicate that more variability than present in the base population has been lost (genomic inbreeding >100%), which makes no sense. On the other hand, by definition, $F_{\rm ROH}$ has only positive values, ranging from 0 to 19% in the current research. Among

Effect Trait^a Breed Calving season Age at calving FROH 22.52 (<0.001) 0.82 (0.485) 16.63 (<0.001) 0.62 (0.430) MY305 FY305 9.01 (<0.001) 0.84 (0.503) 12.20 (<0.001) 0.05 (0.821) 14.53 (<0.001) 1.03 (0.391) PY305 14.86 (<0.001) 0.58 (0.447) CFSI 1.37 (0.252) 2.52 (0.040) 2.68 (0.102) 1.90 (0.168) CCI 2.10 (0.099) 2.07 (0.083) 0.24 (0.622) 5.44 (0.019)

Table 2. F-values and P-values (within brackets) for factors affecting production and fertility traits of pasture-based dairy cows from commercial herds of Argentina

F_{ROH}, coefficient of genomic inbreeding calculated based on runs of homozygosity.

^aMY305, 305-day milk production; FY305, 305-day fat production; PY305, 305-day protein production; CFSI, Calving to first service interval; CCI, calving to conception interval.

several alternative estimates of genomic inbreeding coefficients, it is accepted that $F_{\rm ROH}$ values are more accurate and/or comparable to classical inbreeding coefficient based on pedigree information to determine inbreeding in several livestock species and humans, as they reflect the shared ancestry of genomic haplotypes (Curik *et al.*, 2014; Purfield *et al.*, 2017; Yengo *et al.*, 2017; Yoshida *et al.*, 2020).

Overall, based on the $F_{\rm ROH}$, the rate of inbreeding present in this sample of purebred and crossbred cows is similar to the rates of inbreeding reported in other studies; 1.93–3.05% in Irish Holstein–Friesian (McParland *et al.*, 2007), 3.6% in US Holstein (Cassell *et al.*, 2003) and 5.00 in French Holstein (Dezetter *et al.*, 2015).

In our study the effects of inbreeding and crossbreeding on milk production and fertility traits were evaluated using a single model. A decrease in milk, fat and protein yields with a decreasing breed proportion of Holstein is in line with previous research in New Zealand (Lembeye *et al.*, 2016), Ireland (Prendiville *et al.*, 2011) and Argentina (Baudracco *et al.*, 2011; Mancuso and Marini, 2012; Biga *et al.*, 2022). The productive performance of the HJ crossbred cows was lower than the productive performance

of the purebred Holstein and higher than the JX crossbred cows. Because there were not purebred Jersey cows it was not possible to estimate heterosis effects. Crossbreeding systems have been adopted widely in the New Zealand dairy industry because of the favourable heterosis for milk production and fertility resulting in improved farm profitability (Lopez-Villalobos *et al.*, 2000). All crossbred cows in the current study tended to have shorter CFSI and CCI than purebred Holstein but differences were not significant. These results can be used for modelling studies to evaluate the effects of these fertility rates on farm profitability under pasture-based conditions of Argentina.

The effects of inbreeding, evaluated as the regression coefficient of the milk production and fertility traits on the estimators of genomic inbreeding (F_{GRM} , F_{HOM} and F_{UNI}) considered in the current research were not significant. These results are in disagreement with previous reports by Cassell *et al.* (2003) in US Holstein cows and McParland *et al.* (2007) in Irish dairy cattle. The only trait that was affected by inbreeding, considering the F_{ROH} estimator, was CCI (P < 0.05), which agrees with the effect of inbreeding on the lengthening of the calving interval by up to 0.31 days per 1% increase in inbreeding (Fuerst and Sölkner,



Figure 2. Pearson correlations and *P*-values (within brackets) (above diagonal), scatter plots (below diagonal) between each pair of the four genomic inbreeding estimators and densities (diagonal) of the inbreeding estimators of dairy cows in commercial herds of Argentina. ${}^{1}F_{ROH}$, coefficient of genomic inbreeding calculated based on proportion of the autosomal genome in runs of homozygosity; ${}^{2}F_{GRM}$, coefficient of genomic inbreeding obtained from the diagonal elements of the genomic relationship matrix; ${}^{3}F_{HOM}$, coefficient of genomic inbreeding calculated based in the homozygous excess; ${}^{4}F_{UNI}$, coefficient of genomic inbreeding calculated based on the correlation between uniting gametes.

Table 3. Least-squares means and standard errors for production and fertility traits, and means and standard errors of inbreeding coefficients of purebred Holstein and crossbred Holstein × Jersey cows from pasture-based commercial herds of Argentina

		Breed ^b							
Trait ^a	H 639	HX 64	HJ 157	JX 30	<i>P</i> -value ^c				
MY305 (kg)	5932 ± 67.0	5566 ± 122.0	5125 ± 117.0	4813 ± 159.0	<0.001				
FY305 (kg)	219 ± 2.2	215 ± 4.1	203 ± 3.9	197 ± 5.3	<0.001				
PY305 (kg)	205 ± 2.3	195 ± 4.1	182 ± 3.9	175 ± 5.3	<0.001				
CFSI (days)	90 ± 3.0	77 ± 6.5	81 ± 5.9	81 ± 8.9	0.252				
CCI (days)	160 ± 6.3	151 ± 14.0	140 ± 12.2	115 ± 19.4	0.099				
F _{ROH}	6.6 ± 0.11	3.7 ± 0.18	0.020 ± 0.0004	5.2 ± 0.76					
F _{GRM}	-3.4 ± 0.16	-0.2 ± 0.51	-0.9 ± 0.35	21 ± 1.8					
F _{HOM}	0.8 ± 0.17	-6.7 ± 0.38	-11.2 ± 0.18	-6 ± 1.2					
F _{UNI}	-1.3 ± 0.09	-3.5 ± 0.19	-6.0 ± 0.14	7 ± 1.4					
Heterozygosity	0.35 ± 0.0004	0.36 ± 0.0004	0.38 ± 0.0003	0.34 ± 0.0004					

^aMY305, 305-day milk production; FY305, 305-day fat production; PY305, 305-day protein production; CFSI, calving to first service interval; CCI, calving to conception interval. *F*_{ROH}, mean of genomic inbreeding coefficient calculated based on proportion of the autosomal genome in runs of homozygosity. Heterozygosity was calculated as the proportion of heterozygous marker genotypes for each individual.

^bH, purebred cows with 1.00 Holstein; HX, crossbred Holstein with 0.75 Holstein and 0.25 Jersey; HJ, crossbred Holstein–Jersey with 0.50 Holstein and 0.50 Jersey; JX, crossbred Jersey with 0.25 Holstein and 0.75 Jersey.

^cP-value for breed group effect.

1994; Smith *et al.*, 1998). In this study, the Holstein sires were imported from the US Holstein population using frozen semen, which is a common practice to produce herd replacements. The apparent constriction of no significant effect of inbreeding on milk and fertility traits in this study compared with the significant effects reported by Cassell *et al.* (2003) in US Holstein cows and McParland *et al.* (2007) in Irish Holstein–Friesian cows can be explained considering that in this study we evaluated the effect of inbreeding using a crossbreed population.

The apparent no effects of inbreeding on MY305, FY305, PY305 and CFSI are likely attributed to the fact that the effect of genomic inbreeding was evaluated across breed groups. Crossbreeding and inbreeding have opposite effects (Falconer and Mackay, 1996); inbreeding is the mating of animals that are

more closely related than the average in a population increasing homozygosity, which increases the frequency of unfavourable genotypes, and crossbreeding is the mating of animals that are less related than the average in a population increasing heterozygosity, which allows the expression of heterosis (interactions between alleles in the same locus). Table 3 shows that the HJ crossbred cows, which have the maximum rate of heterozygosits (0.38) in fact, had the lowest mean rate of inbreeding (0.02%) measured according to the $F_{\rm ROH}$ formula. Furthermore, $F_{\rm ROH}$ would credibly measure autozygosity (homozygosity genomic segments produced by identity by descent) and is nowadays considered the most consistent coefficient and the one that best captures inbreeding depression in livestock (Keller *et al.*, 2011; Peripolli *et al.*, 2017; Caballero *et al.*, 2020, 2022; Lozada-Soto



Figure 3. Relationship between heterozygosity and F_{ROH} in purebred Holstein and crossbred Holstein×Jersey cows from pasture-based commercial herds of Argentina. H, purebred cows with 1.00 Holstein; HX, crossbred Holstein with 0.75 Holstein and 0.25 Jersey; HJ, crossbred Holstein–Jersey with 0.50 Holstein and 0.50 Jersey; JX, crossbred Jersey with 0.25 Holstein and 0.75 Jersey. ¹Heterozygosity was calculated as the proportion of heterozygous marker genotypes for each individual. ² F_{ROH} , coefficient of genomic inbreeding calculated based on proportion of the autosomal genome in runs of homozygosity.

et al., 2022). Based on the available information so far, F_{ROH} shows the highest correlation with the classical *F* pedigree-based (Purfield *et al.*, 2012, 2017; Ferenčaković *et al.*, 2013; Marras *et al.*, 2015; Caballero *et al.*, 2022; Dadousis *et al.*, 2022).

Other studies have reported no significant effects of inbreeding on production and fertility traits (Hodges *et al.*, 1979) using a small sample of cows, but concordantly with the results obtained in the current study. In a much larger study, Hofmannová *et al.* (2019) conclude that inbreeding has a negligible influence on the breeding values for conception.

This study reports significant effects of breed on milk production and fertility using data from a large commercial herd. These results can be used for simulation studies that assist the design of crossbreeding programmes (Lopez-Villalobos *et al.*, 2000; Clasen *et al.*, 2020) that increases farm profitability. This study also reports rates of genomic inbreeding and heterozygosity calculated using genetic markers. It was found that, in general, the effects of genomic inbreeding on milk production and fertility traits were not significant, except on the CCI. A possible reason for these nonsignificant effects is that these effects were evaluated in a crossbred population in which rates of heterozygosity would operate to some extent in the opposite direction to rates of genomic inbreeding.

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