Application of single-step genomic evaluation using social genetic effect model for growth in pig

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Objective: Social genetic effects (SGE) are an important genetic component for growth, group productivity, and welfare in pigs. The present study was conducted to evaluate i) the feasibility of the single-step genomic best linear unbiased prediction (ssGBLUP) approach with the inclusion of SGE in the model in pigs, and ii) the changes in the contribution of heritable SGE to the phenotypic variance with different scaling ω constants for genomic relationships.

Methods: The dataset included performance tested growth rate records (average daily gain) from 13,166 and 21,762 pigs Landrace (LR) and Yorkshire (YS), respectively. A total of 1,041 (LR) and 964 (YS) pigs were genotyped using the Illumina PorcineSNP60 v2 BeadChip panel. With the BLUPF90 software package, genetic parameters were estimated using a modified animal model for competitive traits. Giving a fixed weight to pedigree relationships (τ: 1), several weights (ωx, 0.1 to 1.0; with a 0.1 interval) were scaled with the genomic relationship for best model fit with Akaike information criterion (AIC).

Results: The genetic variances and total heritability estimates (T²) were mostly higher with ssGBLUP than in the pedigree-based analysis. The model AIC value increased with any level of ω other than 0.6 and 0.5 in LR and YS, respectively, indicating the worse fit of those models. The theoretical accuracies of direct and social breeding value were increased by decreasing ω in both breeds, indicating the better accuracy of ω0.1 models. Therefore, the optimal values of ω to minimize AIC and to increase theoretical accuracy were 0.6 in LR and 0.5 in YS.

Conclusion: In conclusion, single-step ssGBLUP model fitting SGE showed significant improvement in accuracy compared with the pedigree-based analysis method; therefore, it could be implemented in a pig population for genomic selection based on SGE, especially in South Korean populations, with appropriate further adjustment of tuning parameters for relationship matrices.

Keywords: Accuracy; Breeding Value; Landrace; Single-step Genomic Best Linear Unbiased Prediction (BLUP); Social Genetic Effects; Yorkshire

INTRODUCTION

The genetic effect of an individual on the phenotypes of its social partners (i.e., pen mates) is often termed the social genetic effect (SGE) or the indirect genetic effect [1]. The growth rate is a key trait in pig breeding goals because it contributes to economic efficiency. However, negative effects of social interactions, such as tail biting, or excessive aggression can inhibit growth of pen mates, resulting in reduce productivity in pig farming. The report by Bergsma et al [2] on pigs indicated that the heritable social interaction among various group members might play a role in their average daily gain (ADG). In this regard, Bijma et al [1] stated that the total breeding value (TBV), expressed as the combined direct breeding value (DBV) of an individual and social breeding values (SBV) of pen mates, for growth
performance can be used in selection that includes SGE. Although the estimation of SBV in pigs is achievable through the traditional pedigree-based approach directly using phenotypes, this will inevitably result in lower accuracy [3]. Therefore, it is also highly desirable to use a model i.e., genomic best linear unbiased prediction (BLUP) [4], for the selection of pigs which will include SGE as well as improve accuracy in predictions of breeding values.

In recent years, a single-step genomic BLUP (ssGBLUP) which provides better predictions for all animals in a pedigree [5], is a method of choice for genomic evaluation. In pigs, Christensen et al [6] has provided evidence of improved predictions through ssGBLUP, compared with the pedigree-based method. They also suggested an adjustment of the genomic relationship matrix with ssGBLUP. As genomic information is now often used to optimize selection outcomes, it seems reasonable that the inclusion of social interactions alongside genomic information might benefit the selection of in Korean pigs as well. Therefore, this study was established with the following aims: i) to verify the feasibility of the SGE model using ssGBLUP in dam lines and ii) to evaluate the influences of different scales of matrices on the contribution of heritable social effects to the phenotypic variance.

**MATERIALS AND METHODS**

**Animal phenotypes**

The phenotypic dataset on the growth rate of animals was obtained through performance tests of Yorkshire (YS; n = 21,762) and Landrace (LR; n = 13,166) pigs between 2005 and 2015. These pigs were born and raised in a closed nucleus (breeding) farm in South Korea. Animals were reared in pens where each pen housed 4 to 10 YS or 3 to 8 LR pigs of the same sex. Each group of animals comprised 1 to 7 YS or 1 to 6 LR full-sibs. The performance evaluations on the ADG of pigs started soon after each animal reached a live body weight of 30 kg, and were finished when a target weight of 90 kg was attained. On average, fewer than 160 days were required to attain this target weight. The animal phenotypes selected to be studied were based on the size of the group (frequency >10%). The average ADG was recorded to be 787.9±88.3 g/d in LR and 792.5±92.1 g/d in YS.

Complete pedigrees of the studied animals were obtained from the Korean Animal Improvement Association database. The total numbers of animals in the pedigree of the two breeds were 14,013 (LR) and 22,536 (YS). The numbers of individuals for which both parents were known were 13,916 (LR) and 22,324 (YS). In the whole pedigree, about 96% (LR) and 62% (YS) of the animals were inbred. The average inbreeding coefficients were 0.035 and 0.015 in LR and YS, respectively. The ranges of inbreeding coefficients were 0.0001 to 0.263 (YS) and 0.001 to 0.274 (LR). The observed average family sizes were 3.97 and 4.11 in LR and YS, with ranges of 2 to 15 and 2 to 17, respectively. The population structures of these breeds were determined using the CFC v1.0 software package [7].

The experimental protocols describing the management and care of the animals were reviewed and approved according to the guide for the care and use of laboratory animals (National Institute of Animal Science, Animal Care Committee of Korea) on 7 March 2014 (approval number: NIAS 2014-289). The management practices on the studied population stated that each pen was 2.5×3.6 m (~0.9 m²/pig) in size with solid concrete flooring. Pigs were fed ad libitum and water was constantly accessible through nipple drinkers. The feeding program was applied in accordance with pig testing standards of the Korean Animal Improvement Association (http://www.aiak.or.kr/eng/index.jsp).

**Genotyping of animals**

The genomic DNA of pigs was extracted from their blood samples using a standard protocol. A total of 2,005 pigs from both LR (1,041) and YS (964) breeds were genotyped using the Illumina PorcineSNP60 v2 BeadChip panel, which comprised 61,565 SNP (single nucleotide polymorphism) markers [8]. The quality control (QC) procedure for the genotype data included the deletion of individuals with pedigree errors, removal of monomorphic SNP genotypes, SNPs on sex chromosomes or SNPs with minimum allele frequencies (<0.9), genotype call rate of <0.90, animal missing rate of >0.90, Hardy-Weinberg equilibrium of 0.15, and the SNPs with displaced segregation distortion [9,10]. After QC, the final dataset contained genotypes from a total of 1,915 pigs (LR, 1,029; YS, 886). The total number of autosomal SNPs was reduced to 37,779 in LR and 41,503 in YS, respectively. The correlations of off-diagonal elements of G and A matrix were 0.75 and 0.76 for LR and YS, respectively.

**Statistical analysis**

*Estimation using pedigree information: The variances and covariances of the studied traits were estimated by an animal model applying the REML approach. The effects of batch, sex, and group size were fitted as fixed effects. In the model, age at target weight was fitted as a covariate effect. The effect of batch was formed by combining year, month, and week based on each performance test. The models also included two nongenetic random effects, namely, birth litter and group effects [1,11]. To take into account differences in group size and to prevent the overestimation of heritable variances for SGE, an additional covariate term known as dilution, \((\frac{\text{Average group size} - 1}{\text{Group size} - 1})^{0.5}\) [12], was added to the SGE [3,11,13]. Animals were fitted as a random effect in...*
the model. The BLUPF90 software package Misztal et al [14] was used for the estimation of parameters by fitting a classical model with pedigree relationships only (PED\textsubscript{classic}) and a social model pedigree relationships only modified for competitive traits (PED\textsubscript{social}) [11] as follows:

\[
y = Xb + Z_{\text{ped}} + Wl + Vg + e \quad (\text{PED\textsubscript{classic}})
\]

\[
y = Xb + Z_{\text{ped}} + Z_{\text{sub}} + Wl + Vg + e \quad (\text{PED\textsubscript{social}})
\]

where \(y\) is the vector of observations (ADG), \(b\) is the vector of fixed effects, \(a_i\) is the vector of random direct additive genetic effects, \(a_{s,j}\) is the vector of random SGEs, \(l\) is the vector for random birth litter, \(g\) is the vector of random group, and \(e\) is the vector of residuals. \(X, Z_{\text{ped}}, Z_{\text{sub}}, W,\) and \(V\) are the corresponding incidence matrices. Assumptions for the probability distributions were \(g \sim N(0, I\sigma^2_g), l \sim N(0, I\sigma^2_l), c \sim N(0, I\sigma^2_c),\) and \(e \sim N(0, I\sigma^2_e),\) in which \(N(\cdot)\) indicates a normal distribution; \(I\) is an identity matrix of appropriate dimensions; and \(\sigma^2_g, \sigma^2_l, \sigma^2_c,\) and \(\sigma^2_e\) are the variances of the corresponding effects. In Model 1, direct additive genetic effects had the following distribution: \(a_d \sim N(0, A\sigma^2_{ad})\), in which \(A\) is the numerator relationship matrix and \(\sigma^2_{ad}\) is the variance of direct additive genetic effects. In Model 2, direct and indirect additive genetic effects had the following distribution (MVN) normal distribution:

\[
\begin{bmatrix}
a_D \\
a_S
\end{bmatrix} \sim \text{MVN} \left(0, C \otimes A\right),
\]

where \(C\) is defined by the matrix:

\[
\begin{bmatrix}
\sigma^2_D & \sigma_{ad} \\
\sigma_{ad} & \sigma^2_S
\end{bmatrix}
\]

\(\sigma^2_D\) is the variance of direct genetic effects, \(\sigma^2_S\) is the variance of indirect genetic effects, \(\sigma_{ad}\) is the covariance between direct and indirect genetic effects, and \(C \otimes A\) denotes the Kronecker product of two matrices.

According to Bijma et al [1] for traits affected by heritable social effects, the variance of TBV represents the total heritable variation that is exploitable for selection. The TBV of the \(i\)th animal is defined as follows:

\[
TBV_i = a_{D,i} + (n - 1)a_{S,i}
\]

The TBV is the heritable effect of an individual on trait values in the population, the sum of its direct genetic effect \((a_{D,i})\) on its own phenotype and its SGE \((a_{S,i})\) on the phenotypes of its \(n - 1\) group mates. Bijma et al [1] also stated that the total heritable variance determines the population’s potential in response to selection and can be expressed as:

\[
\sigma^2_{TBV} = \sigma^2_D + 2(n - 1)\sigma_{ad} + (n - 1)^2\sigma^2_S
\]

According to Bergsma et al [2], the phenotypic variance for such a model can be calculated as follows:

\[
\sigma^2_P = \sigma^2_D + (n - 1)\sigma^2_{ad} + \sigma^2_G + \sigma^2_I + \sigma^2_e
\]

where \(n\) indicates the average size of social groups. The total heritable variance can be expressed relative to phenotypic variance [2] as follows:

\[
T^2 = \frac{\sigma^2_{TBV}}{\sigma^2_P}
\]

\textbf{Estimation using single-step method:} The relationship matrix \(H\), in single-step evaluation, defines the relationship among genotyped and nongenotyped animals. The inverse of the \(H\) matrix is rather simple in structure [15,16] and can be given as:

\[
H^{-1} = A^{-1} + \begin{bmatrix}
0 & 0 \\
0 & G^{-1} - A_{22}^{-1}
\end{bmatrix}
\]

where \(A_{22}\) is the matrix for genotyped animals only (a submatrix derived from the pedigree-based relationship matrix, \(A\) and \(G\) is the relationship matrix among individuals based on genomic information. The \(G\) matrix was constructed according to VanRaden [4]. Both \(A_{22}\) and \(G\) matrices were subsequently combined. Thus, overall, the two matrices represented similar diagonals. However, the mixed model equations for a single step mainly differed from the pedigree-based model by a matrix block, \(\tau (0.95 + 0.05 A_{22})^{-1} - \omega A_{22}^{-1}\), given to the genotyped animals [17,18]. The constant \(\omega\) represents the proportion of polygenic variances that were unexplained by markers. The parameters \(\tau\) and \(\omega\) scale the size of the genomic and pedigree relationships, respectively. The weights for the \(\omega\) parameter were between 0.1 and 1.0, whereas \(\tau\) was fixed at 1. The models including genomic information are denoted as \(\omega_{1.0}\), \(\omega_{0.8}\), \(\omega_{0.7}\), \(\omega_{0.6}\), \(\omega_{0.5}\), \(\omega_{0.4}\), \(\omega_{0.3}\), \(\omega_{0.2}\), and \(\omega_{0.1}\), in accordance with their values of the constant \(\omega\), whereas a model with pedigree information only is denoted as PED in later sections.

\textbf{Validation process:} Accuracy of breeding value was calculated in two different ways (theoretical accuracy [3] and cross validation [6]). The last 2 years were masked as the validation data set and predictions were made using the first 9 years as the training data set. The validation data set for LR and YS contained 10% and 8% of the observations, respectively. The theoretical accuracy of the estimated breeding value for the \(i\)th individual with the \(n\)th model was calculated as follows:

\[
R_{i,n} = \frac{1 - \frac{PEV_{i,n}}{(1 + F_i) \sigma^2_i}}
\]

where \(PEV\) is the prediction error variance of its breeding value, \(F\) is the inbreeding coefficient of an individual as computed from the pedigree, and \(\sigma^2_i\) is the additive genetic variance of the model. We also calculated correlation between cor-
rected phenotype and the combined breeding value (CBV) for the validation pigs. Accuracy was defined as

$$r = \text{cor}(\text{CBV}, y_i)$$

where CBV is the sum of pig's own direct breeding value and SBVs of pen mates, $y_i$ is corrected ADG for fixed effects.

**RESULTS AND DISCUSSION**

**Model fitness**

The variances, covariances, and various model parameters obtained from the studied models for LR and YSs are presented in Tables 1 and 2, respectively. The Akaike information criterion (AIC) parameter of the pedigree-classical model was higher than the pedigree-social model in both breeds. This result showed that model including SGE fitted the data significantly better than a classical animal model. In addition, AIC parameter of the pedigree-social model was the highest in both breeds compared with those of all ssGBLUP methods. The AIC as an indicator of the goodness fit of the models indicates that the ssGBLUP models performed better in general, which was as expected due to the addition of genomic information alongside the pedigree relationship. This is a feasible approach with a single-step method as it provides more accurate predictions for both genotyped and nongenotyped pigs.

**Table 1.** Estimates of variances, covariances, genetic parameters, and accuracies for different models in Landrace pigs

| Method   | $\sigma_{a_d}^2$ | $\sigma_{d_a}^2$ | $\sigma_{e_d}^2$ | $\sigma_{g}^2$ | $\sigma_{l}^2$ | $\sigma_p^2$ | $r$ | $T^2$ | $\Delta$AIC |
|----------|-----------------|-----------------|-----------------|---------------|--------------|------------|-----|------|------------|
| PED_classic | 2,078           | -               | 433             | 208           | 3,039        | -          | 5,758 | 0.36 | 72.9       |
| PED_social | 2,069           | 3               | 14              | 383           | 208          | 3,037      | 0.02 | 5,825 | 0.41       |
| $\omega_{d}$ | 2,136           | -6              | 9               | 410           | 210          | 2,999      | -0.05 | 5,840 | 0.39       |
| $\omega_{s}$ | 2,392           | -2              | 14              | 395           | 204          | 2,883      | -0.01 | 5,997 | 0.44       |
| $\omega_{l}$ | 2,589           | 2               | 17              | 382           | 201          | 2,796      | 0.01  | 6,122 | 0.48       |
| $\omega_{e}$ | 2,747           | 7               | 20              | 372           | 200          | 2,728      | 0.03  | 6,226 | 0.52       |
| $\omega_{g}$ | 2,878           | 11              | 23              | 362           | 200          | 2,673      | 0.04  | 6,316 | 0.54       |
| $\omega_{l}$ | 2,988           | 15              | 25              | 353           | 200          | 2,628      | 0.05  | 6,395 | 0.57       |
| $\omega_{o}$ | 3,081           | 19              | 27              | 344           | 201          | 2,590      | 0.06  | 6,463 | 0.59       |
| $\omega_{r}$ | 3,162           | 22              | 30              | 336           | 202          | 2,558      | 0.07  | 6,524 | 0.61       |
| $\omega_{p}$ | 3,233           | 26              | 31              | 329           | 203          | 2,532      | 0.08  | 6,580 | 0.62       |
| $\omega_{a}$ | 3,294           | 29              | 33              | 322           | 204          | 2,509      | 0.09  | 6,630 | 0.64       |

$\sigma_{a_d}^2$, direct genetic variance; $\sigma_{d_a}^2$, covariance between direct and social genetic effects; $\sigma_{g}^2$, random group variance; $\sigma_{l}^2$, random litter variance; $\sigma_{r}^2$, random residual variance; $r$, correlation between direct and social genetic effects; $\sigma_p^2$, phenotypic variance; $T^2$, total heritability for model including social genetic effects; $\Delta$AIC, change in Akaike's information criterion from the best (minimum) model; PED_classic, the classic model with pedigree relationships only; PED_social, the social model with pedigree relationships only; $\omega_{w}$, the model with weighted $A^{-1/2}$ matrix by different $\omega$ constants.

**Table 2.** Estimates of variances, covariances, genetic parameters, and accuracies for different models in Yorkshire pigs

| Method   | $\sigma_{a_d}^2$ | $\sigma_{d_a}^2$ | $\sigma_{e_d}^2$ | $\sigma_{g}^2$ | $\sigma_{l}^2$ | $\sigma_p^2$ | $r$ | $T^2$ | $\Delta$AIC |
|----------|-----------------|-----------------|-----------------|---------------|--------------|------------|-----|------|------------|
| PED_classic | 2,255           | -               | 675             | 260           | 3,712        | -          | 6,901 | 0.33 | 102.1      |
| PED_social | 2,320           | 72              | 23              | 479           | 256          | 3,739      | 0.31 | 7,084 | 0.58       |
| $\omega_{d}$ | 2,323           | 69              | 23              | 483           | 268          | 3,732      | 0.30 | 7,092 | 0.57       |
| $\omega_{s}$ | 2,565           | 79              | 29              | 453           | 262          | 3,621      | 0.29 | 7,260 | 0.64       |
| $\omega_{l}$ | 2,763           | 86              | 33              | 430           | 258          | 3,530      | 0.28 | 7,395 | 0.69       |
| $\omega_{e}$ | 2,930           | 93              | 37              | 412           | 256          | 3,454      | 0.28 | 7,511 | 0.73       |
| $\omega_{g}$ | 3,074           | 99              | 40              | 396           | 255          | 3,389      | 0.28 | 7,611 | 0.77       |
| $\omega_{l}$ | 3,201           | 104             | 43              | 383           | 254          | 3,332      | 0.28 | 7,700 | 0.80       |
| $\omega_{o}$ | 3,313           | 109             | 45              | 372           | 253          | 3,282      | 0.28 | 7,778 | 0.82       |
| $\omega_{a}$ | 3,414           | 113             | 47              | 361           | 253          | 3,238      | 0.28 | 7,851 | 0.84       |
| $\omega_{r}$ | 3,504           | 117             | 49              | 352           | 254          | 3,198      | 0.28 | 7,915 | 0.86       |
| $\omega_{p}$ | 3,585           | 121             | 51              | 343           | 254          | 3,163      | 0.28 | 7,975 | 0.88       |

$\sigma_{a_d}^2$, direct genetic variance; $\sigma_{d_a}^2$, covariance between direct and social genetic effects; $\sigma_{g}^2$, random group variance; $\sigma_{l}^2$, random litter variance; $\sigma_{r}^2$, random residual variance; $r$, correlation between direct and social genetic effects; $\sigma_p^2$, phenotypic variance; $T^2$, total heritability for model including social genetic effects; $\Delta$AIC, change in Akaike's information criterion from the best (minimum) model; PED_classic, the classic model with pedigree relationships only; PED_social, the social model with pedigree relationships only; $\omega_{w}$, the model with weighted $A^{-1/2}$ matrix by different $\omega$ constants.
animals [6,11,18,19]. Therefore, a ssGBLUP analysis including SGE in the model would be a better choice for the prediction of traits in pigs. However, differences were observed among the various model fits with different scaling factors in the single-step methods. Among the ssGBLUP models, the model with $\omega$ of 1.0 showed the worst fit, regardless of the breed. The best fitting models in this study were those with $\omega_{\text{gen}}$ and $\omega_{\text{nongen}}$ in LR (Table 1) and YR (Table 2), respectively, as indicated by them having the lowest AIC estimates. The model AIC value increased with any level of $\omega$ other than 0.6 and 0.5 in LR and YS, respectively, indicating the worse fit of those models. Our results obtained through testing different levels of $\omega$ (0.1 to 1.0) indicate that a ssGBLUP method essentially relies on tuning the scales of matrices related to pedigree and genotype relationships, which will lead to less biased model estimates [6,15,17,20-22]. This study strongly coincides with many previous reports in that the choices of appropriate levels of constants ($\tau$ and $\omega$) are rather arbitrary, and are to be determined through fine tuning. For instance, Misztal et al [17] reported the best combination of $\tau = 1.5$ and $\omega = 0.6$ in their study on dairy cattle. Another study in dairy cattle by Harris et al [23] also used both parameters at levels as low as 0.5. Likewise, Koivula et al [20] reported using various combinations of $A$ and $G$ matrices to find the best option in their study. In pig, Christensen et al [6] suggested a single-step method that is adjusted for the genomic relationship matrix. In another study by Misztal et al [24], a model with slower convergence at $\omega$ values greater than 1 was reported, as their $H$ matrix was found to be nonpositive at higher values of this constant. In this context, it is crucial to find appropriate scaling parameters that will ensure better accuracy, lower bias, and easier convergence. It is also important to consider appropriate weights for relationship matrices through scaling factors as any smaller constant for $\omega$ is likely to decrease the emphasis on the genomic relationships and increase the importance of the pedigree relationships [20]. This might explain our estimates obtained with levels of $\omega$ lower than those in best fit models, where model estimates might have been associated with some biases due to the lower weight in genotyped animals through their genomic relationships.

**Genetic parameters**

The genetic variances and total heritability estimates ($T^2$) were mostly higher with ssGBLUP than in the pedigree-based analysis (Tables 1, 2). Among the ssGBLUP models, the genetic variances and $T^2$ were increased by decreasing $\omega$ in both breeds. Therefore, the $T^2$ of $\omega_{\text{gen}}$ model was the highest in both breeds (LR, 0.64; YS, 0.88). The best single-step models ($\Delta$AIC = 0) showed larger estimates of direct and social variances than pedigree-based methods, and thus also larger covariance estimates, resulting in higher total heritability estimates with those models. The $T^2$ estimates with the best fitting models were 0.54 and 0.80 in LR and YS, respectively. They were also greater than those of the pedigree-based analysis method by 0.13 and 0.22 in these two breeds, respectively. However, our $T^2$ estimates for LR with the $\omega_{\text{nongen}}$ model coincided strongly with those of Bergsma et al [25] and Duijvesteijn [3]. Comparing the breeds, both direct and social genetic contributions were higher in YS than in LR, so their $T^2$ estimates also exhibited the same trend. Note that even when the social variance is markedly smaller than direct genetic variance, its contribution to $\sigma^2_{DBV}$ would be substantial due to the factor $(n–1)^2$, especially when group sizes are large, as was the case with YS. The lower $T^2$ estimates in LR could also be due to the larger nongenetic litter effects and negative covariances between direct and social effects. According to Bijma et al [1], the positive covariance between direct and social genetic variances is likely to increase the total heritable variation, which coincides well with the present study. The correlation coefficients between DBV and SBV were somewhat weaker in LR (–0.05 to 0.09) than in YS (0.28 to 0.31). Some earlier reports [2,3,26] also stated somewhat similar correlations, mostly positive but not significant. In this study, the positive correlation in YS could indicate that their pen mates might also have stimulated a greater ADG. Given that SBV is passed on to pen mates, the positive genetic correlation between the direct and associative effects indicates that pigs with a high DBV will also have a high SBV. In other words, the YR pigs in our study may show more positive responses to selection for social interactions than the LR pigs. Nonetheless, breed differences for social interactions are not unlikely. Bergsma et al [2] suggested that the absence of conflict between an individual’s own growth and mate growth might be a consequence of neutral or slightly cooperative social interactions. For the negative or neutral associative effects in LR pigs in this study, it is possible that these pigs in less competition for food and growth as the amount of space that each of them had on average (3 to 8 pigs/9 m² pen) was lower than that of YR (4 to 10 pigs/9 m² pen).

**Validation**

Table 3 illustrates the accuracy for breeding values obtained with different models. The levels of theoretical accuracy obtained for DBV with PED_classic and PED_social models in each breed were same and also the lowest among the different models (LR, 0.52; YS, 0.55). The $\omega_{\text{nongen}}$ models also performed poorly in DBV prediction (LR, 0.55; YS, 0.58). Among the ssGBLUP models, the theoretical accuracy of DBV was increased by decreasing $\omega$ in both breeds (LR, 0.55 to 0.66; YS, 0.58 to 0.64). The best fit models based on AIC exhibited an increase of accuracy by 5% to 8% compared with the $\omega_{\text{nongen}}$ models in both breeds. The ranges of SBV accuracies with the PED_social in LR and in YS were 0.16 and 0.31, respectively. Simi-
lar to DBV, both PED\textsubscript{social} and ω<sub>1</sub> models performed poorly in SBV prediction. However, unlike the DBVs from the single-step methods, the best fitting models exhibited notable increases in SBV accuracies by 39% (LR) and 19% (YS) with ω<sub>1</sub> and ω<sub>2</sub>, respectively, compared with each of the breed’s worst fit (ω<sub>0</sub>) model. In cross validation, the correlations between CBV and corrected phenotype were also mostly higher with ssGBLUP than in the pedigree-based analysis. However, there were little differences among the ssGBLUP models. The ranges of correlations between CBV and corrected phenotype in LR and YS were 0.31 to 0.33 and 0.21 to 0.22, respectively. The correlative prediction methods showed more variability in terms of ranking of models across traits and replicates so care should be taken interpreting these results with small sample sizes [27]. Putz et al [27] also suggested that for within-breed selection, theoretical accuracy using the prediction error variance was consistent and accurate in ssGBLUP. However, selection programmes should be careful which validation method they choose and should inspect multiple methods if possible [27]. Therefore, to minimize AIC and to increase theoretical accuracy in this study, the optimal values of ω in LR and YS were 0.6 and 0.5, respectively. Martini [28] reported that increasing τ or decreasing ω may mainly decrease inflation by decreasing the variance of the estimated breeding values, which indicate the possibility of further adjustment of τ in the H matrix.

**Prospect of social genetic effects**

The phenotypic variability of some traits that are expressed in the social environment could be significantly influenced by SGEs. Earlier reports on such traits, for instance, social dominance or aggressiveness, also suggested that SGEs can substantially influence total phenotypic variability [29-32]. The importance of SGEs can also be recognized from many previous reports [33-35], which showed that the higher SBV and some desirable characteristics in pigs i.e., fearlessness, stress-tolerance are associated to each other. These characteristics in commercial pig production are particularly beneficial for ease of farm management. For this reason, appropriate attention to such socially influenced traits alongside the pig population structure is vital when genomic selection is considered [36]. Certain strategies could also be applied during selection to achieve a high SBV for a desirable trait. One such approach is to select animals with higher TBVs to improve group performance, especially for growth traits [1-3]. Direct selection of pigs for SBV could be another strategy to alter their social behavior. Earlier evidence suggested that high SBV, due to apathy of the animal, could reduce negative social effects on the growth of others [37-39]. Moreover, the inclusion of SNP effects with SGEs in the model could provide better predictions [40]. For successful realization of TBV, it is also important to consider social environments, such as the mixing method of suckling piglets [41].

**CONCLUSION**

For SGEs, our study showed greater improvement in parameter estimates through ssGBLUP over the traditional pedigree-based method. Both breeds differed to some extent for their estimated parameters. The value of ω used for adjusting A<sub>ω</sub> matrix also differed between the best fitting models for the LR and YS breeds. But it was clear that the models with ω of 1.0 in the H matrix were the worst fitting. Our study also indicated the possibility of further adjustment of other model parameters (α, β, τ) in the H matrix to reduce inflation of the estimated breeding values. Our results

### Table 3. The accuracy of estimated breeding values for different models in pigs

| Method       | Landrace | Yorkishire |
|--------------|----------|------------|
|              | DBV<sub>acc</sub> | SBV<sub>acc</sub> | Cor | DBV<sub>acc</sub> | SBV<sub>acc</sub> | Cor |
| PED<sub>classic</sub> | 0.52 (0.06) | - | 0.27 | 0.55 (0.04) | - | 0.20 |
| PED<sub>social</sub> | 0.52 (0.06) | 0.16 (0.05) | 0.28 | 0.55 (0.04) | 0.31 (0.03) | 0.21 |
| ω<sub>0</sub> | 0.55 (0.09) | 0.11 (0.09) | 0.33 | 0.58 (0.05) | 0.33 (0.04) | 0.21 |
| ω<sub>1</sub> | 0.58 (0.07) | 0.33 (0.09) | 0.33 | 0.59 (0.05) | 0.42 (0.04) | 0.21 |
| ω<sub>2</sub> | 0.60 (0.07) | 0.42 (0.07) | 0.33 | 0.60 (0.05) | 0.46 (0.05) | 0.22 |
| ω<sub>3</sub> | 0.61 (0.07) | 0.47 (0.07) | 0.32 | 0.61 (0.05) | 0.49 (0.05) | 0.22 |
| ω<sub>4</sub> | 0.62 (0.07) | 0.50 (0.07) | 0.32 | 0.62 (0.06) | 0.51 (0.06) | 0.22 |
| ω<sub>5</sub> | 0.63 (0.07) | 0.52 (0.07) | 0.32 | 0.63 (0.06) | 0.53 (0.06) | 0.22 |
| ω<sub>6</sub> | 0.64 (0.07) | 0.54 (0.07) | 0.32 | 0.63 (0.06) | 0.54 (0.06) | 0.22 |
| ω<sub>7</sub> | 0.65 (0.07) | 0.56 (0.07) | 0.32 | 0.64 (0.06) | 0.55 (0.06) | 0.22 |
| ω<sub>8</sub> | 0.65 (0.07) | 0.57 (0.08) | 0.31 | 0.64 (0.06) | 0.56 (0.06) | 0.22 |
| ω<sub>9</sub> | 0.66 (0.07) | 0.58 (0.08) | 0.31 | 0.64 (0.06) | 0.57 (0.07) | 0.22 |

DBV<sub>acc</sub>, the theoretical accuracy of direct breeding value; SBV<sub>acc</sub>, the theoretical accuracy of social breeding value; Cor, the correlation between corrected phenotype and the combined breeding value (CBV); PED<sub>classic</sub>, the classic model with pedigree relationships only; PED<sub>social</sub>, the social model with pedigree relationships only; ω<sub>ω</sub>, the model with weighted A<sub>ω</sub> matrix by different ω constants.
also indicated the value of further analysis with a greater sample size to obtain a more robust estimation of breeding values. We believe that our results provide useful insights for future modeling of SGE in the genomic selection of pig breeds, especially in South Korea.

CONFLICT OF INTEREST

We certify that there is no conflict of interest with any financial organization regarding the material discussed in the manuscript.

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REFERENCES

1. Bijma P, Muir WM, Van Arendonk JA. Multilevel selection 1: quantitative genetics of inheritance and response to selection. Genetics 2007;175:277-88. https://doi.org/10.1534/genetics.106.062711
2. Bergsma R, Kanis E, Knol EF, Bijma P. The contribution of social effects to heritable variation in finishing traits of domestic pigs (Sus scrofa). Genetics 2008;178:1559-70. https://doi.org/10.1534/genetics.107.084236
3. Duijvesteijn N. Sociable swine: prospects of indirect genetic effects for the improvement of productivity, welfare and quality [Ph. D. thesis]. Wageningen, NL, USA: Wageningen University; 2014.
4. VanRaden PM. Efficient methods to compute genomic predictions. J Dairy Sci 2008;91:4414-23. https://doi.org/10.3168/jds.2008-0980
5. Misztal I, Legarra A, Aguilar I. Computing procedures for genomic evaluation including phenotypic, full pedigree, and genomic information. J Dairy Sci 2009;92:4648-55. https://doi.org/10.3168/jds.2009-0964
6. Christensen OF, Madsen P, Nielsen B, Ostersen T, Su G. Single-step methods for genomic evaluation in pigs. Animal 2012;6:1565-71. https://doi.org/10.1017/S1751731112000742
7. Sargolzaei M, Iwaisaki H, Colleau J. CFC: A tool for monitoring genetic diversity. 8th World Congress on Genetics Applied to Livestock Production; 2006 Aug 13-18: Belo Horizonte, MG, Brasil. CD-ROM Communication; 2006.
8. Ramos AM, Crooijmans RP, Affara NA, et al. Design of a high density SNP genotyping assay in the pig using SNPs identified and characterized by next generation sequencing technology. PloS One 2009;4:e6524. https://doi.org/10.1371/journal.pone.0006524
9. Wiggans GR, VanRaden PM, Bacheller LR, et al. Selection and management of DNA markers for use in genomic evaluation. J Dairy Sci 2010;93:2287-92. https://doi.org/10.3168/jds.2009-2773
10. Aguilar I, Misztal I, Legarra A, Tsuruta S. Efficient computation of the genomic relationship matrix and other matrices used in single-step evaluation. J Anim Breed Genet 2011;128:422-8. https://doi.org/10.1111/j.1439-0388.2010.00912.x
11. Arango J, Misztal I, Tsuruta S, Culbertson M, Herring W. Estimation of variance components including competitive effects of Large White growing gilts. J Anim Sci 2005;83:1241-6. https://doi.org/10.2527/2005.8361241x
12. Bijma P. Estimating indirect genetic effects: precision of estimates and optimum designs. Genetics 2010;186:1013-28. https://doi.org/10.1534/genetics.110.120493
13. Bijma P. Multilevel selection 4: modeling the relationship of indirect genetic effects and group size. Genetics 2010;186:1029-31. https://doi.org/10.1534/genetics.110.120485
14. Misztal I, Tsuruta S, Strabel T, et al. BLUPF90 and related programs (BGF90). The 7th World Congress on Genetics Applied to Livestock Production; 2002 Aug 19-23: Montpellier, France.
15. Aguilar I, Misztal I, Johnson DL, Legarra A, Tsuruta S, Lawlor TJ. Hot topic: A unified approach to utilize phenotypic, full pedigree, and genomic information for genetic evaluation of Holstein final score. J Dairy Sci 2010;93:743-52. https://doi.org/10.3168/jds.2009-2730
16. Christensen OF, Lund MS. Genomic prediction when some animals are not genotyped. Genet Sel Evol 2010;42:2. https://doi.org/10.1186/1297-9686-42-2
17. Misztal I, Aguilar I, Legarra A, Lawlor T. Choice of parameters for single-step genomic evaluation for type. J Dairy Sci 2010;93(Suppl 1):533.
18. Tsuruta S, Misztal I, Aguilar I, Lawlor TJ. Multiple-trait genomic evaluation of linear type traits using genomic and phenotypic data in US Holsteins. J Dairy Sci 2011;94:4198-204. https://doi.org/10.3168/jds.2011-4256
19. Chen C, Misztal I, Aguilar I, et al. Genome-wide marker-assisted selection combining all pedigree phenotypic information with genotypic data in one step: an example using broiler chickens. J Anim Sci 2011;89:23-8. https://doi.org/10.2527/jas.2010-3071
20. Koivula M, Strandén I, Pööö J, Aamand GP, Mäntysaari EA. Single-step genomic evaluation using multtrait random regression model and test-day data. J Dairy Sci 2015;98:2775-84. https://doi.org/10.3168/jds.2014-8975
21. Tsuruta S, Lourenco DAL, Misztal I. Bias in singlestep genomic evaluations attributable to unknown parent group estimates. J Dairy Sci 2013;96(E-Suppl 1):75 (Abstr.).
22. Onogi A, Komatsu T, Shoji N, et al. Genomic prediction in Japanese Black cattle: application of a single-step approach to beef cattle. J Anim Sci 2014;92:1931-8. https://doi.org/10.2527/jas.2014-7168
23. Harris BL, Winkelman A, Johnson D. Large-scale single-step
genomic evaluation for milk production traits. Proceedings of the 2012 Interbull Meeting; 2012 May 28-31: Cork, Ireland. No. 46.

24. Misztal I, Aggrey SE, Muir WM. Experiences with a single-step genome evaluation. Poult Sci 2013;92:2530-4. https://doi.org/10.3382/ps.2012-02739

25. Bergsma R, Mathur P, Kanis E, Verstegen MW, Knol EF, Van Arendonk JA. Genetic correlations between lactation performance and growing-finishing traits in pigs. J Anim Sci 2013;91:3601-11. https://doi.org/10.2527/jas.2012-6200

26. Chen CY, Kachman SD, Johnson RK, Newman S, Van Vleck LD. Estimation of genetic parameters for average daily gain using models with competition effects. J Anim Sci 2008;86: 2525-30. https://doi.org/10.2527/jas.2007-0660

27. Putz A, Tiezzi F, Maltecca C, Gray KA, Knauer MT. A comparison of accuracy validation methods for genomic and pedigree-based predictions of swine litter size traits using Large White and simulated data. J Anim Breed Genet 2018;135:5-13. https://doi.org/10.1111/jbg.12302

28. Martini JWR, Schrauf MF, Garcia-Baccino CA, et al. The effect of the H-1 scaling factors τ and ω on the structure of H in the single-step procedure. Genet Sel Evol 2017;50:16. https://doi.org/10.1186/s12711-018-0386-x

29. Wilson AJ, Morrissey M, Adams M, et al. Indirect genetics effects and evolutionary constraint: an analysis of social dominance in red deer, *Cervus elaphus*. J Evol Biol 2011;24:772-83. https://doi.org/10.1111/j.1420-9101.2010.02212.x

30. Alemu SW, Bijma P, Møller SH, Janss L, Berg P. Indirect genetic effects contribute substantially to heritable variation in aggression-related traits in group-housed mink (*Neovison vison*). Genet Sel Evol 2014;46:30. https://doi.org/10.1186/1297-9686-46-30

31. Moore AJ, Brodie ED, Wolf JB. Interacting phenotypes and the evolutionary process: I. direct and indirect genetic effects of social interactions. Evolution 1997;51:1352-62. https://doi.org/10.1111/j.1558-5646.1997.tb01458.x

32. Sartori C, Mantovani R. Indirect genetic effects and the genetic bases of social dominance: evidence from cattle. Heredity 2012;110:3-9. https://doi.org/10.1038/hdy.2012.56

33. Camerlink I, Turner SP, Bijma P, Bolhuis JE. Indirect genetic effects and housing conditions in relation to aggressive behaviour in pigs. PloS One 2013;8:e65136. https://doi.org/10.1371/journal.pone.0065136

34. Reimert I, Rodenburg TB, Ursinus WW, et al. Backtest and novelty behavior of female and castrated male piglets, with diverging social breeding values for growth. J Anim Sci 2013;91:4589-97. https://doi.org/10.2527/jas.2013-6673

35. Reimert I, Rodenburg TB, Ursinus WW, Kemp B, Bolhuis JE. Responses to novel situations of female and castrated male pigs with divergent social breeding values and different backtest classifications in barren and straw-enriched housing. Appl Anim Behav Sci 2014;151:24-35. https://doi.org/10.1016/j.applanim.2013.11.015

36. Samorè AB, Fontanesi L. Genomic selection in pigs: state of the art and perspectives. Italian J Anim Sci 2016;15:211-32.

37. D’Eath R, Conington J, Lawrence A, Olsson I, Sandoe P. Breeding for behavioural change in farm animals: practical, economic and ethical considerations. Anim Welf 2010;19:17-27.

38. Rodenburg T, Bijma P, Ellen E, et al. Breeding amiable animals? Improving farm animal welfare by including social effects in breeding programmes. Anim Welf 2010;19 (Suppl 1):77-82.

39. Camerlink I, Ursinus WW, Bijma P, Kemp B, Bolhuis JE. Indirect genetic effects for growth rate in domestic pigs alter aggressive and manipulative biting behaviour. Behav Genet 2015;45:117-26. https://doi.org/10.1007/s10519-014-9671-9

40. Hong JK, Jeong YD, Cho ES, et al. A genome-wide association study of social genetic effects in landrace pigs. Asian-Australas J Anim Sci 2018;31:784-90. https://doi.org/10.5713/ajas.17.0440

41. Hong JK, Kim KH, Hwang HS, Lee JK, Eom TK, Rhim SJ. Behaviors and body weight of suckling piglets in different social environments. Asian-Australas J Anim Sci 2017;30:902-6. https://doi.org/10.5713/ajas.16.0653