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Pedigree relationships to control inbreeding in optimum-contribution selection realise more genetic gain than genomic relationships

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Abstract

Background: We tested the premise that optimum-contribution selection with pedigree relationships to control inbreeding (POCS) realises at least as much true genetic gain as optimum-contribution selection with genomic relationships (GOCS) at the same rate of true inbreeding.

Methods: We used stochastic simulation to estimate rates of true genetic gain realised by POCS and GOCS at a 0.01 rate of true inbreeding in three breeding schemes with best linear unbiased predictions of breeding values based on pedigree (PBLUP) and genomic (GBLUP) information. The three breeding schemes differed in number of matings and litter size. Selection was for a single trait with a heritability of 0.2. The trait was controlled by 7702 biallelic quantitative-trait loci (QTL) that were distributed across a 30-M genome. The genome contained 54,218 biallelic markers that were used in GOCS and GBLUP. A total of 6012 identity-by-descent loci were placed across the genome in base populations. Unique alleles at these loci were used to calculate rates of true inbreeding. Breeding schemes were run for 10 discrete generations. Selection candidates were genotyped and phenotyped before selection.

Results: POCS realised more true genetic gain than GOCS at a 0.01 rate of true inbreeding in all combinations of breeding scheme and prediction method. POCS realised 14 to 33% more true genetic gain than GOCS with PBLUP in the three breeding schemes. It realised 1.5 to 5.7% more true genetic gain than GOCS with GBLUP.

Conclusions: POCS realised more true genetic gain than GOCS because it managed expected genetic drift without restricting selection at QTL. By contrast, GOCS penalised changes in allele frequencies at markers that were generated by genetic drift and selection. Because these marker alleles were in linkage disequilibrium with QTL alleles, GOCS restricted changes in allele frequencies at QTL. This provides little incentive to use GOCS and highlights that we have more to learn before we can control inbreeding using genomic relationships in selective-breeding schemes. Until we can do so, POCS remains a worthy method of optimum-contribution selection because it realises more true genetic gain than GOCS at the same rate of true inbreeding.

Background

The aim of most animal-breeding schemes is to maximise rates of true genetic gain (ΔG_{true}) at acceptable rates of true inbreeding (ΔF_{true}). ΔG_{true} is calculated as the increase in true breeding value (TBV) averaged across animals in a breeding population. ΔF_{true} is

calculated from the average true inbreeding coefficient of the animals, where the true inbreeding coefficient of an individual is the proportion of loci in its genome with alleles that are identical-by-descent (IBD). Both ΔG_{true} and ΔF_{true} are unobservable in practice. They need to be predicted. The best selection method to use these predictions and fulfil the aim of most animal-breeding schemes is optimum-contribution selection (OCS). OCS maximises rates of predicted genetic gain while controlling inbreeding at given rates of predicted

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inbreeding [1, 2]. It does this by optimising the genetic contribution of each selection candidate to the next generation. One of the benefits of OCS is that it can optimise genetic contributions when different sources of information are used to predict ΔG_{true} and control ΔF_{true} [3]. ΔG_{true} is, more often than not, predicted using best linear unbiased prediction (BLUP) of breeding values based on pedigree or genomic information, hereafter referred to as PBLUP and GBLUP. ΔF_{true} is predicted and controlled using pedigree or genomic relationships, hereafter referred to as OCS with pedigree (POCS) or genomic relationships (GOCS). GOCS became the method-of-choice for OCS with GBLUP when Sonesson et al. [4] used stochastic simulation to recommend that the information used to predict ΔG_{true} should also be used to predict and control ΔF_{true} . Their reasoning was that GOCS predicted and controlled ΔF_{true} more accurately when it was used with GBLUP, while POCS predicted and controlled ΔF_{true} more accurately with PBLUP. However, this reasoning did not consider ΔG_{true} . When we plotted ΔG_{true} realised by Sonesson et al. [4] against ΔF_{true} , we saw that POCS realised more ΔG_{true} than GOCS, even at similar ΔF_{true} (Fig. 1). We are generally supported by Clark et al. [5], who found that, with few exceptions, POCS realised just as much ΔG_{true} as GOCS with both PBLUP and GBLUP, despite being compared at the same rates of genomic inbreeding. Comparing POCS and GOCS at the same rates of genomic inbreeding, rather than ΔF_{true} , would have favoured GOCS, given that GOCS maximises rates of predicted genetic gain while controlling rates of genomic inbreeding. Our interpretation of these studies led us to believe that POCS realises at least as much ΔG_{true} as GOCS at the same ΔF_{true} . We tested this premise by stochastic simulation. We compared ΔG_{true} realised by POCS and GOCS at $\Delta F_{\text{true}} = 0.01$ ($0.01\Delta F_{\text{true}}$) in three breeding schemes with PBLUP and GBLUP. We also simulated OCS with IBD relationships (IOCS) and replaced predictions of breeding values with TBV as points of reference. Results that highlight the mechanisms underlying POCS and GOCS are presented.

Methods

Design

We used stochastic simulation to estimate ΔG_{true} realised by POCS and GOCS at $0.01\Delta F_{\text{true}}$ in three breeding schemes with PBLUP and GBLUP. Put simply, we calibrated POCS and GOCS to realise $0.01\Delta F_{\text{true}}$ and compared their ΔG_{true} . We also simulated IOCS—OCS with IBD relationships—and replaced predictions of breeding values with TBV as points of reference. Selection was for a single trait that had a heritability of 0.2

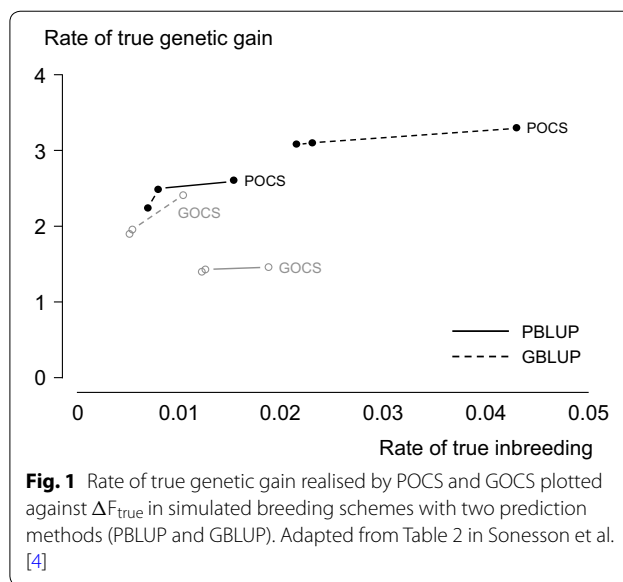


Fig. 1 Rate of true genetic gain realised by POCS and GOCS plotted against ΔF_{true} in simulated breeding schemes with two prediction methods (PBLUP and GBLUP). Adapted from Table 2 in Sonesson et al. [4]

and was controlled by 7702 biallelic quantitative-trait loci (QTL). The QTL were randomly distributed across a 30-M genome that consisted of 18 pairs of autosomal chromosomes. Each chromosome was 167 cM long. The genome contained 54,218 biallelic markers that were used in GOCS and GBLUP. These markers were randomly distributed across the genome and in linkage disequilibrium (LD) with the QTL. A total of 6012 IBD loci were placed evenly across the genomes of animals in base populations. Unique alleles at these loci were used to calculate ΔF_{true} and carry out IOCS. The number of chromosomes and LD between alleles at the markers were simulated to resemble those in three commercial breeds of Danish pigs [6]. Breeding schemes were run for 10 discrete generations ($t = 1 \dots 10$). Animals in the base populations were randomly selected in generation $t = 1$. In generations $t = 2 \dots 10$, selection candidates were allocated matings by OCS. All animals were genotyped before selection; all candidates in generations $t = 2 \dots 10$ were phenotyped for the trait under selection. Each combination of OCS method, breeding scheme, and prediction method was replicated 200 times. We present ΔG_{true} realised at $0.01\Delta F_{\text{true}}$ in generations $t = 4 \dots 10$ and results that highlight the mechanisms underlying POCS and GOCS.

Breeding schemes

The three breeding schemes differed in number of matings and litter size.

M25L5

Twenty-five matings were allocated to 125 selection candidates by OCS in generations $t = 2 \dots 10$. There was no

upper limit for the number of matings that were allocated to each male; males were allocated 0, 1, 2 ... or 25 matings. Twenty-five females were allocated a single mating. The 25 sire and dam matings were paired randomly. Each pair (dam) produced five offspring, resulting in 25 full-sib families and 125 offspring. Offspring were assigned as males or females with a probability of 0.5.

M25L20

This scheme is as for breeding scheme M25L5 with two exceptions. First, 25 matings were allocated to 500 candidates. Second, each dam produced 20 offspring, resulting in 25 full-sib families and 500 offspring.

M100L5

This scheme is as for breeding scheme M25L5 with two exceptions. First, 100 matings were allocated to 500 candidates. Males were allocated 0, 1, 2 ... or 100 matings and 100 females were allocated a single mating. Second, each dam produced five offspring, resulting in 100 full-sib families and 500 offspring.

Simulation procedure

Generations – 1000 to – 1: founder population

LD between the 54,218 markers and 7702 QTL was established in a founder population using a Fisher-Wright inheritance model [7, 8]. The founder population was simulated for 1000 discrete generations ($t = -1000 \dots -1$) with 25 males and 25 females, and an effective-population size of 50, in each generation. In generation $t = -1$, the founder population was in recombination-drift-mutation-selection equilibrium. We considered the founder population to be in equilibrium when the numbers of segregating markers and QTL, the level of heterozygosity averaged over all segregating markers and QTL, and the average LD between segregating markers that were 0.25, 0.5, 1, 2, 5, and 10 cM apart became constant across generations.

The founder population was initiated with 25 males and 25 females in generation $t = -1000$. Their 30-M genomes consisted of 3×10^7 monomorphic loci with wild-type alleles that were placed evenly across the genome at 10^4 loci per cM. Every eighth locus harboured a QTL that controlled the trait under selection. The remaining loci were markers.

The males and females in subsequent generations were simulated by randomly sampling a sire and dam with replacement from the 25 males and 25 females in the previous generation. Bi-allelic polymorphism at each locus was generated with a mutation rate of 4×10^{-6} per locus using an infinite-sites mutation model [9]. An additive-genetic effect for the mutant allele at each QTL was sampled from an exponential distribution. The sign

of each additive-genetic effect was negative with a probability of 0.9. The additive-genetic effects of the wild-type alleles were zero. Selection was introduced by sampling 25 males and 25 females that were above a 5% percentile for TBV. The TBV of the i th animal in the founder population, a_i , was calculated as $a_i = \sum_{j=1}^{n_{\text{QTL}}} x_{ij}g_j$, where $n_{\text{QTL}} = 3.75 \times 10^6$ is the number of QTL across the genome, x_{ij} is the number of copies of the mutant allele that animal i inherited at the j th QTL ($x_{ij} = 0, 1, 2$), and g_j is the additive-genetic effect of the mutant allele at the j th QTL. We introduced selection because animal populations are always under selection, which influences LD between alleles.

The 54,218 markers and 7702 QTL in our three breeding schemes were all segregating in generation $t = -1$ of the founder population. The additive-genetic effects of the mutant alleles at the 7702 segregating QTL were standardised so that the total additive-genetic variance for the trait under selection was equal to 1.0. No new mutations were generated after the founder population was simulated.

Chromosomes from the 50 animals in generation $t = -1$ of the founder population were pooled: 18 pools of 100 chromosomes. Each pool consisted of 50 chromosome pairs of the i th chromosome ($i = 1 \dots 18$) from 50 founder animals. The breeding schemes were initiated by sampling base populations from these chromosome pools.

Generation 0: base populations

Each replicate combination of OCS method, breeding scheme, and prediction method was initiated by sampling a unique base population. Twenty-six males and 25 females were sampled in breeding schemes M25L5 and M25L20. Eleven males and 100 females were sampled in breeding scheme M100L5. The genotype of each base animal was sampled from the 18 pools of chromosomes in generation $t = -1$ of the founder population. For chromosome i ($i = 1 \dots 18$), two chromosomes were randomly sampled without replacement from the i th pool of 100 chromosomes. The sampled chromosomes were replaced before the next base animal was sampled. Base animals were assumed to be unrelated and non-inbred based on pedigree and IBD alleles. They were genotyped, but not phenotyped for the trait under selection.

Generation 1: random selection in base populations

Animals in the base populations were selected in generation $t = 1$ by randomly culling a single male. In breeding schemes M25L5 and M25L20, 25 sires and 25 dams were selected. Each sire was mated with one dam. Each dam produced five offspring in breeding scheme M25L5

and 20 offspring in breeding scheme M25L20. In breeding scheme M100L5, 10 sires and 100 dams were selected. Each sire was mated with 10 dams and each dam produced five offspring. Randomly culling a single male enabled us to construct genomic-relationship matrices that were positive-definite. This is explained in more detail in section ‘Genomic and IBD-relationship matrices.’

Generations 2–10: optimum-contribution selection

Animals were selected and allocated matings by OCS in generations $t=2 \dots 10$. The phenotype of animal i , p_i , was calculated as $p_i = a_i + e_i$, where a_i is the animal’s TBV and e_i is its residual value. The TBV of animal i was calculated as described for the founder population using the standardised additive-genetic effects of the mutant alleles at the 7702 QTL. Its residual value, e_i , was sampled from $e_i \sim N(0, \sigma_e^2 = 4)$.

IBD loci

The 6012 IBD loci used to calculate ΔF_{true} and carry out IOCS were placed evenly across the genome of animals in the base populations at two IBD loci per cM (334 loci per chromosome). Each base animal was assigned two unique alleles at each IBD locus. IBD alleles could be traced back from each descendant to the base animal from which it was derived. A descendant was IBD at an IBD locus when it inherited two copies of a unique allele (i.e., both alleles at the IBD locus descended from the same unique allele in the base population). IBD loci were not involved in prediction.

Optimum-contribution selection

POCS allocated matings to selection candidates in generations $t=2 \dots 10$ conditional on predicted breeding values and pedigree relationships. It did this by maximising a quadratic function, U_t , with respect to \mathbf{c} :

$$U_t(\mathbf{c}) = \mathbf{c}'\mathbf{g} - \omega\mathbf{c}'\mathbf{A}\mathbf{c}, \quad (1)$$

where \mathbf{c} is an n vector of genetic contributions to the next generation and the number of matings allocated to each candidate is a linear function of these contributions, n is the number of animals in the population traced back from the candidates in generation t to the base population, \mathbf{g} is an n vector of PBLUP, GBLUP, or TBV, ω is a penalty applied to the expected average relationship of the next generation, and \mathbf{A} is an $n \times n$ pedigree-relationship matrix. Elements of \mathbf{c} were constrained to $0 \leq c_i \leq 0.5$ ($i = 1 \dots n$) with $c_i = 0$ for animals that were not candidates for selection in generation t . Using these definitions, $\mathbf{c}'\mathbf{g}$ is the expected breeding value and $\mathbf{c}'\mathbf{A}\mathbf{c}$ is the expected average relationship of the next generation. The penalty, ω , was constant across generations. It was calibrated to realise $0.01\Delta F_{\text{true}}$. We calibrated it by simulating 200 replicates of POCS in each combination

of breeding scheme and prediction method with an initial ω and calculating the mean ΔF_{true} across the replicates. This process was repeated using different ω until the mean ΔF_{true} deviated from 0.01 by less than 0.0001. GOCS was carried out by replacing \mathbf{A} with an $n \times n$ genomic-relationship matrix, \mathbf{G} . IOCS was carried out by replacing \mathbf{A} with an $n \times n$ IBD-relationship matrix, \mathbf{B} . The method of POCS is described in full by Henryon et al. [10].

Predicted breeding values

PBLUP for the trait under selection were estimated in generations $t=2 \dots 10$ by fitting an animal model to the phenotypes observed in generations 2 to t . The model was:

$$\mathbf{y} = \mathbf{X}\mathbf{b} + \mathbf{Z}\mathbf{a} + \mathbf{e},$$

where \mathbf{y} is an n vector of phenotypes, \mathbf{b} is an h vector of fixed generation effects, h is the number of generations with phenotypes, \mathbf{a} is an n vector of random animal effects, \mathbf{e} is an n vector of residual errors, and \mathbf{X} and \mathbf{Z} are incidence matrices. The (co)variance structure was:

$$\begin{pmatrix} \mathbf{a} \\ \mathbf{e} \end{pmatrix} \sim N \left(\begin{bmatrix} \mathbf{0} \\ \mathbf{0} \end{bmatrix}, \begin{bmatrix} \mathbf{A}\sigma_a^2 & \mathbf{0} \\ \mathbf{0} & \mathbf{I}\sigma_e^2 \end{bmatrix} \right),$$

where \mathbf{I} is an identity matrix, $\sigma_a^2 = 1$ is the additive-genetic variance in generation $t = -1$ of the founder population, and $\sigma_e^2 = 4$ is the residual variance that was used to sample phenotypes. GBLUP were estimated by replacing \mathbf{A} with the genomic-relationship matrix, \mathbf{G} .

Genomic and IBD-relationship matrices

Genomic-relationship matrices

Genomic-relationship matrices used in GOCS and GBLUP were constructed as $\mathbf{G} = \mathbf{Z}\mathbf{Z}'/s$, where \mathbf{G} is an $n \times n$ matrix of genomic relationships, $\mathbf{Z} = \mathbf{M} - 1(2\mathbf{p})'$, \mathbf{M} is an $n \times m$ matrix of counts of the mutant allele for the n animals at the $m=54,218$ markers with element $M_{ij} = 0, 1$, or 2 for animal i at marker j ($i = 1 \dots n$, $j = 1 \dots m$), $\mathbf{1}$ is an n vector of ones, $\mathbf{p} = (p_1, p_2, \dots, p_m)$ is an m vector with p_j the frequency of the mutant allele at marker j in the base populations, and $s = 2\mathbf{p}'(1 - \mathbf{p})$ transforms \mathbf{G} towards the same scale as a pedigree-relationship matrix (adapted from VanRaden [11]).

We carried out two additional steps to ensure that \mathbf{G} was positive-definite. First, the allele frequencies in \mathbf{p} were calculated using all animals in the base populations, including the single male that was culled in generation $t = 1$ of each breeding scheme. Second, all base animals and selection candidates, except for the culled male, were included in \mathbf{G} . These steps generated linear independence in \mathbf{G} because allele frequencies in \mathbf{p} were calculated using an animal that was not included in \mathbf{G} .

IBD-relationship matrices

The IBD-relationship matrix, \mathbf{B} , used in IOCS was an $n \times n$ matrix constructed as $B_{ij} = \frac{1}{n_{\text{IBD}}} \sum_{k=1}^{n_{\text{IBD}}} \left[\frac{1}{2} \sum_{u=1}^2 \sum_{v=1}^2 \delta_{uv} \right]$, where element B_{ij} is the IBD relationship between animals i and j ($i = 1 \dots n, j = 1 \dots n$), $n_{\text{IBD}} = 6012$ is the number of IBD loci, and δ_{uv} is the allele-sharing status. δ_{uv} was equal to 1 if allele u of animal i was identical to allele v of animal j at IBD locus k , and 0 otherwise.

Rates of true genetic gain and true inbreeding

We present ΔG_{true} realised by POCS, GOCS, and IOCS at $0.01\Delta F_{\text{true}}$ in each combination of breeding scheme and prediction method. ΔG_{true} and ΔF_{true} are presented as means (\pm SD) of the 200 replicates. We also scaled ΔG_{true} by setting ΔG_{true} realised by POCS to 100 in each combination of breeding scheme and prediction method.

ΔG_{true} in each replicate was calculated as a linear regression of G_t on t , where G_t is the average TBV of animals born in generations $t=4 \dots 10$. ΔG_{true} was presented as a linear regression because G_t was linear over t . ΔF_{true} in each replicate was calculated as $1 - \exp(\beta)$, where β is a linear regression of $\ln(1 - F_t)$ on t , and F_t is the average coefficient of true inbreeding for animals born in generations $t=4 \dots 10$ [12, 13]. These transformations were made because $\ln(1 - F_t)$, not F_t , was linear over t . F_t was calculated as $F_t = \frac{1}{n_t n_{\text{IBD}}} \sum_{i=1}^{n_t} \sum_{j=1}^{n_{\text{IBD}}} \delta_{ij}$, where n_t is the number of animals born in generation t , $n_{\text{IBD}} = 6012$ is the number of IBD loci, and δ_{ij} is the IBD status of animal i ($i = 1 \dots n_t$) at IBD locus j ($j = 1 \dots n_{\text{IBD}}$). δ_{ij} was equal to 1 if animal i was homozygous at IBD locus j , and 0 otherwise.

Rates of pedigree and genomic inbreeding

We present rates of pedigree inbreeding realised by POCS and rates of genomic inbreeding realised by GOCS at $0.01\Delta F_{\text{true}}$ in each combination of breeding scheme and prediction method. Rates of pedigree and genomic inbreeding were calculated as for ΔF_{true} with F_t replaced by average coefficients of pedigree and genomic inbreeding for animals born in generations $t=4 \dots 10$. The coefficient of genomic inbreeding for animal i was calculated as $G_{ii} - 1.0$, where G_{ii} is the i th diagonal element of \mathbf{G} used in GOCS.

Mechanisms underlying POCS and GOCS

We present results that highlight the mechanisms underlying POCS and GOCS. These results are only presented for POCS and GOCS with PBLUP and GBLUP in breeding scheme M25L5; results from breeding schemes M100L5 and M25L20 were similar to those from breeding scheme M25L5. Two of the results—response

frontiers and minimum ΔF_{true} —involved additional simulations. All of these results are presented as means (\pm SD) of 200 replicates.

Changes in allele frequencies at markers and QTL

We present the average absolute changes in allele frequencies at markers and QTL at $0.01\Delta F_{\text{true}}$ and the average increase in the frequencies of favourable alleles at the QTL. Changes in allele frequencies were calculated from generations $t=4$ to $t=10$ using animals born in generations $t=4$ and $t=10$. The frequency changes in each replicate were averaged over the 54,218 markers and 7702 QTL.

Variance in rate of identity-by-descent

We present the variance in rate of IBD between the 6012 IBD loci at $0.01\Delta F_{\text{true}}$. Rate of IBD at each locus in each replicate was calculated as $1 - \exp(\beta)$, where β is a linear regression of $\ln(1 - F_{it})$ on t , and F_{it} is the proportion of animals born in generations $t=4 \dots 10$ that were IBD at locus i ($i = 1 \dots 6012$).

Numbers of candidates and families that were allocated matings

We present the number of male candidates that were allocated matings at $0.01\Delta F_{\text{true}}$ and the numbers of half and full-sib families with male or female candidates that were allocated matings. The numbers in each replicate were averaged over generations $t=4 \dots 10$. The number of female candidates that were allocated matings was not presented because 25 females were always allocated a single mating in breeding scheme M25L5.

Rank and rank deviations

We present the average ranks and average-rank deviations of males and females that were allocated matings within full-sib families at $0.01\Delta F_{\text{true}}$ when males and females within each full-sib family were ranked by predicted breeding value. The average rank of males that were allocated matings in each full-sib family was calculated when males were ranked from 1 \dots $n_{\text{male}i}$, where $n_{\text{male}i}$ is the number of males in the i th full-sib family. The average-rank deviation of males in each full-sib family was calculated as the difference between their average rank and their average-minimum rank, where average-minimum rank is the average rank had those males that were allocated matings been the highest-ranked males in their full-sib families. The average ranks and average-rank deviations in each generation were averaged across full-sib families with males that were allocated matings. The generation averages in each replicate were averaged over generations $t=4 \dots 10$. The average rank and

average-rank deviation of females were calculated as for males.

Response frontiers

We present response frontiers for POCS and GOCS with PBLUP and GBLUP: ΔG_{true} plotted against ΔF_{true} . Different ΔF_{true} were realised by applying different penalties, ω , in Eq. (1). Response frontiers tell us if the relative ΔG_{true} realised by POCS and GOCS at $0.01\Delta F_{\text{true}}$ is also realised across a range of ΔF_{true} .

Minimum rates of true inbreeding

We present the minimum ΔF_{true} realised by POCS and GOCS when we relaxed selection for predicted breeding value. POCS was carried out as described previously with the exception that Eq. (1) was reduced to $U_i(c) = -c'Ac$ for POCS. With GOCS, A was replaced by G . Minimum ΔF_{true} provides insight into the effectiveness of POCS and GOCS to control ΔF_{true} .

Software

The simulations were run using the program ADAM [14]. PBLUP and GBLUP were predicted using DMU version 6 [15]. OCS was carried out by EVA [16].

Results

Rates of true genetic gain

POCS realised more ΔG_{true} than GOCS at $0.01\Delta F_{\text{true}}$ in all combinations of breeding scheme and prediction method. POCS realised 14 to 33% more ΔG_{true} than GOCS with PBLUP in our three breeding schemes (Table 1). It realised 1.5 to 5.7% more ΔG_{true} than GOCS with GBLUP and 0.3 to 1.4% more ΔG_{true} than GOCS with our reference prediction, TBV.

POCS also realised more ΔG_{true} than IOCS at $0.01\Delta F_{\text{true}}$ with PBLUP. With PBLUP, POCS realised 3.7 to 6.5% more ΔG_{true} than IOCS in our three breeding schemes (Table 1). In turn, IOCS realised 10 to 27% more ΔG_{true} than GOCS. While POCS realised more ΔG_{true} than IOCS with PBLUP at $0.01\Delta F_{\text{true}}$, IOCS realised a little more ΔG_{true} than POCS with GBLUP and TBV. With GBLUP and TBV, IOCS realised 1.8 to 2.8% and 1.1 to 1.3% more ΔG_{true} than POCS. IOCS realised 3.4 to 8.3% and 1.6 to 2.6% more ΔG_{true} than GOCS with GBLUP and TBV.

Rates of pedigree and genomic inbreeding

Pedigree relationships used by POCS underestimated $0.01\Delta F_{\text{true}}$ in all combinations of breeding scheme and prediction method. POCS underestimated $0.01\Delta F_{\text{true}}$ by 2 to 6% with PBLUP in our three breeding schemes (Table 1). With GBLUP and TBV, it underestimated $0.01\Delta F_{\text{true}}$ by 10 to 26%. By contrast, genomic

relationships used by GOCS underestimated $0.01\Delta F_{\text{true}}$ by 20 to 39% with PBLUP, but overestimated $0.01\Delta F_{\text{true}}$ by as much as 31% with GBLUP and TBV.

The following sections present results that highlight the mechanisms underlying POCS and GOCS. The results are presented for breeding scheme M25L5 with PBLUP and GBLUP.

Changes in allele frequencies at markers and QTL

POCS generated larger changes in allele frequencies at markers and QTL than GOCS at $0.01\Delta F_{\text{true}}$. In breeding scheme M25L5 with PBLUP and GBLUP, the average absolute changes in allele frequencies generated by POCS at markers and QTL were about 4% larger than the changes generated by GOCS (Table 2). By contrast, POCS increased the average frequency of favourable alleles at QTL by 20% more than GOCS with PBLUP and by 4.8% more than GOCS with GBLUP.

Variance in rate of identity-by-descent

POCS and GOCS generated similar variances in rate of IBD between the 6012 IBD loci at $0.01\Delta F_{\text{true}}$. This was highlighted by breeding scheme M25L5 with PBLUP and GBLUP (Table 3).

Numbers of candidates and families that were allocated matings

Males

POCS allocated matings to more male candidates than GOCS at $0.01\Delta F_{\text{true}}$. In breeding scheme M25L5 with PBLUP, POCS allocated matings to 10.1% more male candidates than GOCS (Table 4). With GBLUP, POCS allocated matings to 5.9% more male candidates.

Half and full-sib families

Selection candidates that were allocated matings by POCS were from more half and full-sib families than GOCS at $0.01\Delta F_{\text{true}}$. In breeding scheme M25L5 with PBLUP, POCS allocated matings to candidates from 10.6% more half-sib and 2.3% more full-sib families than GOCS (Table 4). With GBLUP, POCS allocated matings to candidates from 10.5 and 5.0% more half and full-sib families.

Rank and rank deviations

POCS allocated matings to higher-ranked candidates within full-sib families than GOCS at $0.01\Delta F_{\text{true}}$. In breeding scheme M25L5 with PBLUP and GBLUP, the average ranks of males and females that were allocated

Table 1 Rate of true genetic gain realised by POCS, GOCS, and IOCS at $0.01\Delta F_{\text{true}}$ in three breeding schemes with three predictions methods (PBLUP, GBLUP, and TBV)

Prediction	Scheme	OCS	ΔG_{true}	ΔG_{scaled}	$\Delta F_{\text{pedigree}}$	$\Delta F_{\text{genomic}}$	
PBLUP	M25L5	POCS	0.379	100.0	0.0098	0.0071	
		GOCS	0.317	83.6			
		IOCS	0.356	93.9			
	M25L20	POCS	0.570	100.0		0.0094	0.0061
		GOCS	0.429	75.3			
		IOCS	0.544	95.4			
	M100L5	POCS	0.554	100.0		0.0094	0.0080
		GOCS	0.485	87.5			
		IOCS	0.534	96.4			
GBLUP	M25L5	POCS	0.398	100.0	0.0088	0.0099	
		GOCS	0.390	98.0			
		IOCS	0.409	102.8			
	M25L20	POCS	0.703	100.0		0.0074	0.0117
		GOCS	0.665	94.6			
		IOCS	0.720	102.4			
	M100L5	POCS	0.658	100.0		0.0076	0.0127
		GOCS	0.648	98.5			
		IOCS	0.670	101.8			
TBV	M25L5	POCS	0.773	100.0	0.0090	0.0101	
		GOCS	0.762	98.6			
		IOCS	0.782	101.2			
	M25L20	POCS	1.149	100.0		0.0079	0.0131
		GOCS	1.143	99.5			
		IOCS	1.162	101.1			
	M100L5	POCS	0.999	100.0		0.0080	0.0124
		GOCS	0.996	99.7			
		IOCS	1.012	101.3			

Rates of absolute and scaled true genetic gain (ΔG_{true} and ΔG_{scaled}), rates of pedigree inbreeding ($\Delta F_{\text{pedigree}}$) realised by POCS, and rates of genomic inbreeding ($\Delta F_{\text{genomic}}$) realised by GOCS are means of 200 simulation replicates. ΔG_{scaled} was calculated by setting ΔG_{true} realised by POCS to 100 in each combination of breeding scheme and prediction method. SD between the replicates ranged from 0.00114 to 0.00256 (ΔF_{true}), 0.0288 to 0.0559 (ΔG_{true}), 2.51 to 14.45 (ΔG_{scaled}), 0.00743 to 0.00979 ($\Delta F_{\text{pedigree}}$), and 0.00153 to 0.00270 ($\Delta F_{\text{genomic}}$)

Table 2 Average absolute changes in allele frequencies at markers and QTL, and average increase in the frequencies of favourable QTL alleles generated by POCS and GOCS at $0.01\Delta F_{\text{true}}$ in breeding scheme M25L5 with two prediction methods (PBLUP and GBLUP)

Prediction	OCS	Absolute-marker alleles	Absolute-QTL alleles	Favourable-QTL alleles
PBLUP	POCS	0.0475	0.0472	0.00428
	GOCS	0.0456	0.0455	0.00356
GBLUP	POCS	0.0493	0.0487	0.00459
	GOCS	0.0471	0.0468	0.00438

Changes in allele frequencies are means of 200 simulation replicates. SD between the replicates ranged from 0.00132 to 0.00159 (absolute-marker alleles and absolute-QTL alleles) and from 0.00078 to 0.00085 (favourable QTL alleles)

matings by POCS were 8.5 to 10.8% lower than those allocated matings by GOCS (Table 5). Not only did POCS allocate matings to higher-ranked candidates within full-sib families, candidates that were allocated matings by POCS were always the highest-ranked males and females in their full-sib families. The average ranks of the males and females that were allocated matings by POCS did not deviate from their average-minimum ranks—their average-rank deviations were zero. With GOCS, the average ranks of the males and females deviated from their average-minimum ranks by about 10%.

Table 3 Variance in rate of IBD between 6012 IBD loci generated by POCS and GOCS at $0.01\Delta F_{\text{true}}$ in breeding scheme M25L5 with two prediction methods (PBLUP and GBLUP)

Prediction	OCS	Variance
PBLUP	POCS	4.52×10^{-5}
	GOCS	4.56×10^{-5}
GBLUP	POCS	4.48×10^{-5}
	GOCS	4.64×10^{-5}

Variances are means of 200 simulation replicates. SD between the replicates ranged from 5.607×10^{-6} to 7.687×10^{-6}

Table 4 Number of male candidates allocated matings, and numbers of half and full-sib families with candidates allocated matings by POCS and GOCS at $0.01\Delta F_{\text{true}}$ in breeding scheme M25L5 with two prediction methods (PBLUP and GBLUP)

Prediction	OCS	Males	Half-sibs	Full-sibs
PBLUP	POCS	19.7	17.8	21.8
	GOCS	17.9	16.1	21.3
GBLUP	POCS	19.9	18.9	23.0
	GOCS	18.8	17.1	21.9

Numbers are means of 200 simulation replicates. SD between the replicates ranged from 0.72 to 0.87 (males), 0.79 to 0.93 (half-sibs), and 0.50 to 0.60 (full-sibs)

Response frontiers

There were two main features of our response frontiers. First, POCS continued to realise more ΔG_{true} than GOCS across a range of ΔF_{true} . Second, both POCS and GOCS realised less ΔG_{true} as ΔF_{true} decreased, but ΔG_{true} realised by POCS decreased at a slower rate than GOCS. These two features are illustrated by the response frontiers for POCS and GOCS with PBLUP and GBLUP in breeding scheme M25L5 (Fig. 2). With PBLUP, POCS realised about 6% more ΔG_{true} than GOCS at $0.03\Delta F_{\text{true}}$, 20% more ΔG_{true} at $0.01\Delta F_{\text{true}}$, and 40% more ΔG_{true} at $0.006\Delta F_{\text{true}}$. With GBLUP, POCS and GOCS realised

similar ΔG_{true} at ΔF_{true} higher than about 0.015. POCS realised 2% more ΔG_{true} than GOCS at $0.01\Delta F_{\text{true}}$ and almost 10% more ΔG_{true} at about $0.006\Delta F_{\text{true}}$.

Minimum rates of true inbreeding

POCS realised higher minimum ΔF_{true} than GOCS when we relaxed selection for predicted breeding value. In breeding scheme M25L5, the minimum ΔF_{true} realised by POCS was 6.1% higher than the minimum ΔF_{true} realised by GOCS (mean \pm SD of 200 replicates: 0.0050 ± 0.00147 vs 0.0047 ± 0.00144).

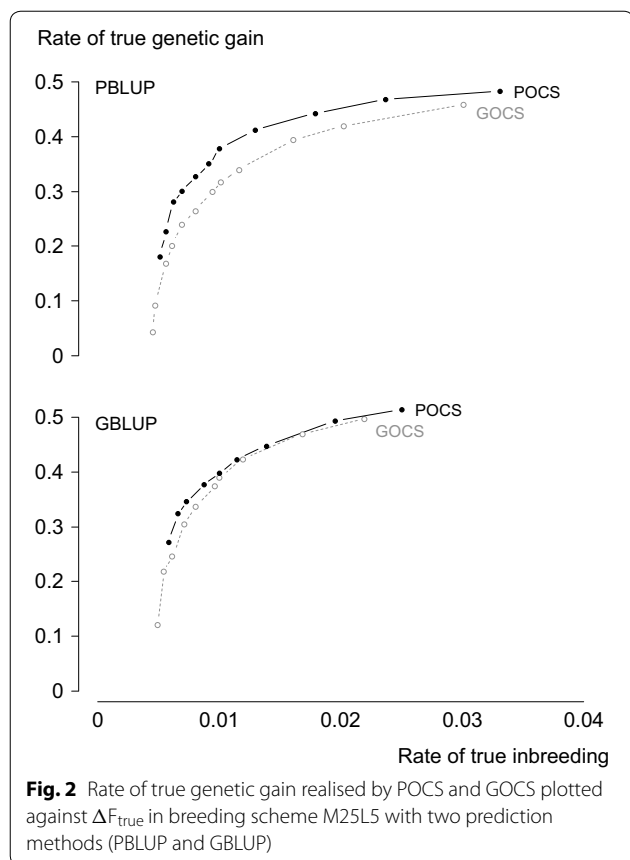
Discussion

Our findings supported our premise that POCS realises at least as much ΔG_{true} as GOCS at the same ΔF_{true} . When we calibrated POCS and GOCS to $0.01\Delta F_{\text{true}}$, we found that POCS always realised more ΔG_{true} than GOCS, regardless of the breeding scheme that we simulated or the information used to predict breeding values. This makes POCS an attractive method of OCS to use in breeding schemes, including schemes that use GBLUP. It also highlights that the potential for GOCS to trace changes in allele frequencies at markers does not guarantee more ΔG_{true} . Our findings are supported by the results of Sonesson et al. [4] and Clark et al. [5], but not the recommendation by Sonesson et al. [4], who reasoned that the same information used to predict ΔG_{true} should also be used to predict and control ΔF_{true} . It would be worthwhile reassessing the recommendation by Sonesson et al. [4], given that their reasoning did not consider ΔG_{true} and their study made GOCS the method-of-choice for OCS with GBLUP. While we predicted that POCS would realise at least as much ΔG_{true} as GOCS, we were surprised to find that our reference OCS, IOCS, realised only marginally more ΔG_{true} than POCS with GBLUP and TBV. This result provides little incentive to use GOCS and highlights that we have more to learn before we can control inbreeding using genomic relationships in selective-breeding schemes. Until we can

Table 5 Average ranks and average-rank deviations of males and females allocated matings by POCS and GOCS at $0.01\Delta F_{\text{true}}$ in breeding scheme M25L5 with two prediction methods (PBLUP and GBLUP)

Prediction	OCS	Rank _{Males}	Deviation _{Males}	Rank _{Females}	Deviation _{Females}
PBLUP	POCS	1.18 ± 0.025	0	1.26 ± 0.026	0
	GOCS	1.29 ± 0.056	0.13 ± 0.044	1.39 ± 0.044	0.12 ± 0.035
GBLUP	POCS	1.16 ± 0.022	0	1.24 ± 0.026	0
	GOCS	1.30 ± 0.052	0.13 ± 0.044	1.39 ± 0.043	0.13 ± 0.038

Average ranks and average-rank deviations are means (\pm SD) of 200 simulation replicates



do so, POCS remains a worthy method of OCS because it realises more ΔG_{true} than GOCS at the same ΔF_{true} .

POCS realised more ΔG_{true} than GOCS because it managed expected genetic drift without restricting selection at QTL. It did this by applying a penalty to $c'Ac$. $c'Ac$ can be rewritten as $c'Ac = c'LL'c$, where L is a normed lower-triangular matrix that describes the expected genetic contribution that an ancestor makes to its descendants, and $L'c$ is a vector of expected genetic contributions from candidates and ancestors to the next generation [17, 18]. Penalising $c'Ac = c'LL'c$ penalised increases in expected genetic contributions quadratically, where the sum of squares of expected contributions is a function of expected genetic drift [19]. Managing expected genetic drift managed the variance in changes in allele frequencies at hypothetical neutral loci. These neutral loci are assumed to be unlinked to QTL alleles [3]. Because they were unlinked, POCS allowed the frequencies of favourable alleles at QTL to be increased by selection. By contrast, GOCS penalised changes in allele frequencies at markers that were generated by genetic drift and selection. It applied a penalty to $c'Gc = c'ZZ'c/s$, where $Z'c/2$ is a vector of changes in allele frequency at each marker [3] and these changes

were measured as deviations from allele frequencies in the base populations. Penalising $c'Gc = c'ZZ'c/s$ penalised changes in allele frequencies at markers quadratically; markers with the largest frequency changes were penalised hardest. Because these marker alleles were in LD with QTL alleles, GOCS restricted changes in allele frequencies at QTL. This explanation highlights the problem with GOCS in its current form: it penalises changes in allele frequencies at all markers when, in fact, we need to change allele frequencies at some markers to increase the frequencies of favourable alleles at QTL. So, by managing expected genetic drift, POCS realises more ΔG_{true} than GOCS at the same ΔF_{true} because it allows selection to increase the frequencies of favourable alleles at QTL more than GOCS.

Deductive reasoning tells us that POCS also generated different IBD profiles across the genome than GOCS at $0.01\Delta F_{true}$. POCS must have generated more IBD than GOCS in regions of the genome that harboured QTL and less IBD in regions that lacked QTL, given that (1) POCS realised more ΔG_{true} than GOCS, (2) POCS generated larger increases in the frequencies of favourable alleles at QTL, (3) QTL alleles were in LD with IBD alleles, (4) areas under IBD profiles increase at the same rate at the same ΔF_{true} , and (5) POCS and GOCS generated similar variances in rate of IBD between IBD loci. While the IBD generated by POCS was associated with QTL location, the IBD generated by GOCS must have been associated with both QTL location and marker density because the markers used by GOCS to predict and control ΔF_{true} were randomly distributed across the genome. GOCS presumably generated most IBD in regions of the genome that harboured QTL with low marker densities, least IBD in regions that lacked QTL with high marker densities, and intermediate IBD in other regions of the genome. We did not present IBD profiles for POCS and GOCS because we simulated many QTL, each with a small change in allele frequency. This resulted in differences in IBD between POC and GOCS at each IBD locus that were small and difficult to detect visually. It would be worthwhile carrying out simulations to test unequivocally that POCS generates more IBD than GOCS in regions of the genome that harbour QTL. Increasing the frequencies of favourable alleles at QTL while restricting most of the IBD to regions of the genome that harbour these QTL is, after all, how we want to realise ΔG_{true} at acceptable ΔF_{true} in animal breeding. Therefore, not only does POCS allow selection to increase the frequencies of favourable alleles at QTL more than GOCS at the same ΔF_{true} , it is probably also more aligned with the objectives of animal breeding by restricting most IBD to regions of the genome that harbour QTL.

A direct consequence of managing expected genetic drift was that POCS allocated matings to different selection candidates than GOCS. There were two major differences. First, POCS allocated matings to more candidates from more half and full-sib families than GOCS to realise $0.01\Delta F_{\text{true}}$. POCS did this because it could neither differentiate between pairwise relationships within full-sib families—they had the same expected relationships—nor could it trace increases in allele frequencies at IBD loci that were in regions of the genome under the influence of genetic drift and selection. These regions were prone to higher ΔF_{true} than predicted by pedigree relationships. Candidates that were allocated matings by POCS tended to share more QTL alleles, more genomic regions flanking the QTL, and more IBD alleles than predicted. POCS compensated for this by allocating matings at rates of pedigree inbreeding that were lower than $0.01\Delta F_{\text{true}}$. It used variation in expected relationships between families to allocate matings to more candidates from more families. By contrast, GOCS allocated matings to fewer candidates from fewer families than POCS because it could differentiate between pairwise relationships and exploit some of the variation in IBD relationships within full-sib families. Realising the same ΔF_{true} with fewer breeding animals made GOCS a more effective control of ΔF_{true} than POCS and shows that GOCS does provide valuable information for inbreeding control—it was just not as effective at realising ΔG_{true} . Further evidence that GOCS controlled ΔF_{true} more effectively was that it realised lower minimum ΔF_{true} than POCS when we relaxed selection for predicted breeding value. With no selection for predicted breeding value, the objective was to restrict increases in the frequencies of IBD alleles, which GOCS did more effectively than POCS. But despite GOCS being a more effective control of ΔF_{true} , POCS still realised a minimum ΔF_{true} of 0.005 with only 25 matings per generation in breeding scheme M25L5. This was well within the 0.005–0.01 range of ΔF_{true} that is considered acceptable for breeding schemes [20]. Therefore, POCS should still be able to realise ΔF_{true} that is considered acceptable in most breeding schemes by allocating matings to more candidates from more full and half-sib families.

The second difference was that POCS allocated matings to higher-ranked candidates within full-sib families than GOCS at $0.01\Delta F_{\text{true}}$. Candidates that were allocated matings by POCS were always the highest-ranked males and females in their full-sib families. Allocating matings to the highest-ranked candidates generated the extra ΔG_{true} realised by POCS as these candidates tended to share favourable alleles at QTL. POCS allowed matings to be allocated to these candidates because all full-sibs have the same pairwise relationships based on pedigree relationships; candidates from the same full-sib

family incurred the same penalty regardless of rank. On the other hand, GOCS could not always allocate matings to the highest-ranked candidates. Just as these candidates tended to share QTL alleles, they also shared marker alleles. Allocating matings to them was penalised by GOCS because it generated larger changes in allele frequencies at markers. So, POCS more than compensated for allocating matings to more candidates from more half and full-sib families to realise the same ΔF_{true} as GOCS. It allocated these matings to higher-ranked candidates within full-sib families, which increased the frequencies of favourable alleles at QTL and realised more ΔG_{true} .

Not only did POCS realise more ΔG_{true} than GOCS at $0.01\Delta F_{\text{true}}$, it also realised more ΔG_{true} across a range of ΔF_{true} . At ΔF_{true} higher than $0.01\Delta F_{\text{true}}$, POCS and GOCS realised similar ΔG_{true} . Most selection emphasis was on ΔG_{true} , and both POCS and GOCS tended to allocate matings to the same highly-ranked candidates that would have been allocated matings by truncation selection. At lower ΔF_{true} , the mechanisms that differentiate POCS from GOCS became more pronounced and POCS realised relatively more ΔG_{true} than GOCS. POCS allocated matings to even more candidates from more full and half-sib families to reduce ΔF_{true} . Candidates that were allocated matings by POCS continued to be the highest-ranked males and females in their full-sib families and the frequencies of favourable alleles at QTL continued to increase, albeit at slower rates. On the other hand, GOCS penalised changes in allele frequencies at markers even harder at lower ΔF_{true} . Candidates that were allocated matings by GOCS differed more for predicted IBD relationships and they were less likely to be the highest-ranked males and females in their full-sib families. This further restricted changes in allele frequencies at QTL. That is, penalising changes in allele frequencies at markers imposes increasingly larger restrictions on changes in allele frequencies at QTL at lower ΔF_{true} than penalising increases in expected genetic contributions. Therefore, the mechanisms that underlie POCS and GOCS apply across a range of ΔF_{true} with POCS realising relatively more ΔG_{true} than GOCS at lower ΔF_{true} because it allows relatively larger changes in the frequencies of favourable alleles at QTL.

Pedigree and genomic relationships used by POCS and GOCS were only predictors of ΔF_{true} . Pedigree relationships used by POCS always underestimated $0.01\Delta F_{\text{true}}$ when we selected for predicted breeding value because they could not trace increases in allele frequencies at IBD loci that were in regions of the genome under the influence of selection. Genomic relationships used by GOCS did not predict ΔF_{true} more accurately than pedigree relationships, even though GOCS controlled ΔF_{true} more effectively than POCS. GOCS underestimated $0.01\Delta F_{\text{true}}$

with PBLUP, but overestimated $0.01\Delta F_{\text{true}}$ with GBLUP and TBV. There are two reasons why GOCS did not predict ΔF_{true} more accurately than POCS. First, the marker alleles used to predict ΔF_{true} were not in complete LD with IBD alleles. Second, the markers were randomly distributed across the genome. Random distribution implies uneven control of IBD across the genome because GOCS could only control inbreeding using marker alleles. GOCS had to increase inbreeding control in regions of the genome with high marker densities to compensate for reduced inbreeding control in regions with low marker densities. Observable and accurate predictors of ΔF_{true} are central to breeding schemes for two reasons. First, they provide a measure of risk; the risk of breeding schemes being adversely impacted by inbreeding depression and loss of genetic variation [3]. Second, they enable OCS to increase selection differentials by allocating matings to selection candidates that realise predicted rates of inbreeding that are close to desired ΔF_{true} . With no obvious relationship between ΔF_{true} and rates of pedigree and genomic inbreeding, we are unable to calibrate POCS and GOCS to realise the desired ΔF_{true} . This makes inbreeding control using POCS or GOCS challenging. Clearly, observable and accurate predictors of ΔF_{true} are needed to better manage risk and increase selection differentials in animal breeding.

Even with accurate predictors of ΔF_{true} , GOCS in its current form is still unlikely to realise more ΔG_{true} than POCS at the same ΔF_{true} . In other words, accurate prediction of ΔF_{true} is not enough to maximise ΔG_{true} at the same ΔF_{true} . The reason is that prediction of ΔF_{true} and inbreeding control are different concepts when maximising ΔG_{true} at the same ΔF_{true} . This was highlighted by our reference OCS, IOCS. IOCS realised more ΔG_{true} than GOCS at $0.01\Delta F_{\text{true}}$ because it had perfect knowledge of ΔF_{true} . It controlled ΔF_{true} with the same IBD alleles that were used to calculate ΔF_{true} . This suggests that GOCS will realise more ΔG_{true} if genomic relationships could be used to predict ΔF_{true} more accurately. However, the amount of extra ΔG_{true} is unlikely to result in GOCS realising more ΔG_{true} than POCS, given that IOCS, at best, only realised marginally more ΔG_{true} than POCS at $0.01\Delta F_{\text{true}}$. IOCS realised only marginally more ΔG_{true} than POCS because it penalised increases in allele frequencies at IBD loci in the same way that GOCS penalised changes in allele frequencies at markers. It applied a penalty to $\mathbf{c}'\mathbf{B}\mathbf{c} = \mathbf{c}'\mathbf{D}\mathbf{D}'\mathbf{c}$, where \mathbf{D} is a matrix of counts of each unique allele at each IBD locus that was inherited by each animal, and $\mathbf{D}'\mathbf{c}$ is a vector of the numbers of each allele at each IBD locus that were expected to be passed on to the next generation. Penalising $\mathbf{c}'\mathbf{B}\mathbf{c} = \mathbf{c}'\mathbf{D}\mathbf{D}'\mathbf{c}$ penalised increases in the expected numbers of IBD

alleles quadratically. This presumably generated flat IBD profiles across the genome and restricted changes in allele frequencies at QTL. Like GOCS, IOCS needed to increase the frequencies of some IBD alleles to increase the frequencies of favourable alleles at QTL. So, GOCS in its current form, where changes in allele frequencies at all markers are penalised, is unlikely to ever realise more ΔG_{true} than POCS at the same ΔF_{true} .

If GOCS is to realise more ΔG_{true} than POCS at the same ΔF_{true} , we will need to change the way that genomic relationships are used to control ΔF_{true} . Rather than penalise changes in allele frequencies at all markers, we should probably allow changes in allele frequencies at some markers by varying the level of inbreeding control and rate of IBD across the genome while controlling ΔF_{true} at acceptable levels. This will involve relaxing inbreeding control in regions of the genome that harbour QTL, allowing selection to increase the frequencies of favourable alleles at QTL. At the same time, we will need to increase inbreeding control to reduce genetic drift in regions of the genome that lack QTL. Varying the level of inbreeding control across the genome could be carried out in GOCS by constructing genomic-relationship matrices that weight markers in regions of the genome that harbour QTL lower than markers in regions that lack QTL. Weighted genomic-relationship matrices have been used in genomic prediction [21, 22]. An alternative approach is to construct genomic-relationship matrices by fixing the frequencies of mutant alleles at markers, \mathbf{p} , to desired frequencies rather than frequencies in base populations. This approach would cause GOCS to penalise deviations from the desired allele frequencies. While these approaches are simple in theory, implementing them in practice requires that we overcome a major hurdle: we do not know where many, if any, of the QTL are located on the genome. We do not know what we want to change and in what direction, nor do we know which regions of the genome can tolerate being IBD. Overcoming this hurdle will require biological information about the QTL that control traits under selection, traits that might be under selection in future, and unobserved fitness traits. Unfortunately, this information is unlikely to become available soon. Without it, there is no guarantee that GOCS will realise more ΔG_{true} or that it will control IBD in regions of the genome that are susceptible to IBD. Therefore, GOCS should realise more ΔG_{true} than POCS at the same ΔF_{true} when we relax inbreeding control in regions of the genome that harbour QTL, but implementing this in practice will require biological information about QTL.

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Authors' contributions

All authors designed the study, interpreted the results, and revised the manuscript. MH ran the simulations, analysed the simulated data, and wrote the manuscript. MH, ACS, and PB co-wrote ADAM. PB wrote EVA. GS wrote a program that calculated genomic-relationship matrices. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

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References

- Wray NR, Goddard ME. Increasing long-term response to selection. *Genet Sel Evol.* 1994;26:431–51.
- Meuwissen THE. Maximizing the response of selection with a predefined rate of inbreeding. *J Anim Sci.* 1997;75:934–40.
- Woolliams JA, Berg P, Dagnachew BS, Meuwissen THE. Genetic contributions and their optimisation. *J Anim Breed Genet.* 2015;132:89–99.
- Sonesson AK, Woolliams JA, Meuwissen THE. Genomic selection requires genomic control of inbreeding. *Genet Sel Evol.* 2012;44:27.
- Clark SA, Kinghorn BP, Hickey JM, van der Werf JH. The effect of genomic information on optimal contribution selection in livestock breeding programs. *Genet Sel Evol.* 2013;45:44.
- Wang L, Sørensen P, Janss L, Ostersen T, Edwards D. Genome-wide and local pattern of linkage disequilibrium and persistence of phase for 3 Danish pig breeds. *BMC Genet.* 2013;14:115.
- Fisher RA. *The genetical theory of natural selection.* Oxford: Clarendon Press; 1930.
- Wright S. Evolution in Mendelian populations. *Genetics.* 1931;16:97–159.
- Kimura M. Number of heterozygous nucleotide sites maintained in a finite population due to steady flux of mutations. *Genetics.* 1969;61:893–903.
- Henryon M, Ostersen T, Ask B, Sørensen AC, Berg P. Most of the long-term genetic gain from optimum-contribution selection can be realised with restrictions imposed during optimisation. *Genet Sel Evol.* 2015;47:21.
- VanRaden PM. Efficient methods to compute genomic predictions. *J Dairy Sci.* 2008;91:4414–23.
- Sonesson AK, Woolliams J, Meuwissen THE. Kinship, relationship and inbreeding. In: Gjedrem T, Doordrecht K, editors. *Selection and breeding programs in aquaculture.* Dordrecht: Springer; 2004. p. 73–87.
- Hinrichs D, Meuwissen THE, Ødegard J, Holt M, Vangen O, Woolliams JA. Analysis of inbreeding depression in the first litter size of mice in a long-term selection experiment with respect to the age of the inbreeding. *Heredity (Edinb).* 2007;99:81–8.
- Pedersen LD, Sørensen AC, Henryon M, Ansari-Mahyari S, Berg P. ADAM: a computer program to simulate selective breeding schemes for animals. *Livest Sci.* 2009;121:343–4.
- Madsen P, Sørensen P, Su G, Damgaard LH, Thomsen, H, Labouriau R. DMU—a package for analyzing multivariate mixed models. In: *Proceedings of the 8th world congress on genetics applied to livestock production: 13–18 August 2006; Belo Horizonte; 2006.* Communication, p. 27–11.
- Berg P, Nielsen J, Sørensen MK. EVA: realized and predicted optimal genetic contributions. In: *Proceedings of the 8th world congress on genetics applied to livestock production: 13–18 August 2006; Belo Horizonte; 2006.* Communication, p. 27–09.
- Henderson CR. A simple method for computing the inverse of a numerator relationship matrix used in prediction of breeding values. *Biometrics.* 1976;32:69–83.
- Quaas RL. Computing the diagonal elements and inverse of a large numerator relationship matrix. *Biometrics.* 1976;32:949–53.
- Wray NR, Thompson R. Prediction of rates of inbreeding in selected populations. *Genet Res.* 1990;55:41–54.
- Bijma P. Long-term genetic contributions: prediction of rates of inbreeding and genetic gain in selected populations. Ph.D. thesis. Wageningen Agricultural University; 2000.
- Zhang Z, Liu J, Ding X, Bijma P, de Koning DJ, Zhang Q. Best linear unbiased prediction of genomic breeding values using a trait-specific marker-derived relationship matrix. *PLoS One.* 2010;5:e12648.
- Su G, Christensen OF, Janss L, Lund MS. Comparison of genomic predictions using genomic relationship matrices built with different weighting factors to account for locus-specific variances. *J Dairy Sci.* 2014;97:6547–59.

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