# Predicted response of genomic selection for new traits using combined cow and bull reference populations

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#### Introduction

Genomic selection particularly is beneficial for dairy cattle breeding because it allows programs, to significantly reduce generation interval, and cheaply increase selection intensity, while the accuracy of selection is only marginally lower compared to progeny testing schemes. It relaxes the requirement of traditional dairy cattle breeding schemes to measure phenotypes from progeny groups for each male selection candidate. Therefore, genomic selection holds the promise to allow selection for new traits, that are difficult or expensive to measure.

For traits that have been part of the breeding goal, currently, reference populations are composed of up to 16,000 daughter proven sires (Lund, et al., 2010). For new more expensive traits reference populations may only be feasible for perhaps a few thousand cows with a single phenotypic measurement. Such reference population may be formed by combining data from different research herds, such as realized within the RobustMilk project.

Examples of new traits are methane emission, energy balance, progesterone profiles as a proxy for fertility, and susceptibility to paratuberculosis (Johne's disease).

Small reference populations comprising only up to a few thousand cows with records, are expected to yield direct genomic values (DGV) with relatively low reliability. Since the generation interval is decreased with genomic selection, the selection response might still be noteworthy. Also, it might be possible to increase reliability by including information from correlated indicator traits a bivariate analysis (Calus and in Veerkamp, 2011), using a reference population with daughter proven bulls. Important questions are whether such approach would help to significantly increase accuracy of genomic selection for new traits, and what the response to genomic selection for those new traits is, even if their DGV reliability is low.

The objectives of this paper, therefore, are 1) to predict DGV reliability for small cow reference populations, that may be supplemented with sire reference populations using an indicator trait, and 2) to predict response to genomic selection for a new trait across the range of the DGV reliabilities.

#### Material and methods

#### Evaluated traits

The evaluated simplified breeding program was based on an index modelled as a single trait with a heritability of 30%. The breeding program was executed using genomic selection, assuming that DGV of selection candidates have a reliability of 0.64 for the index. In this breeding program, a new trait was introduced. We considered a heritability of 30% for the new trait, and genetic correlations with the index ( $r_g$ (index, new trait)) of -0.5, 0, or 0.5. Both the index and the new trait had a phenotypic variance of 1.

#### Predicted DGV reliability

Expected reliabilities of genomic predictions were derived as explained below. First, reliabilities are predicted considering either only cow or bull data. Those reliabilities are then blended afterwards using selection index theory. Reliabilities using cow or bull data are predicted using the formula (Daetwyler, et al., 2008, Daetwyler, 2009):

$$r_{g\hat{g}}^2 = q^2 \frac{\frac{n_p h^2}{n_G}}{\frac{n_p h^2 + 1}{n_G} h^2 + 1}$$
[1]

where  $q^2$  is the proportion of the total genetic variance captured by the markers,  $n_p$  is the number of phenotypes used,  $n_G$  is the number of effective QTL or chromosome segments, and  $h^2$  is the reliability of the trait. For  $q^2$  we used a value of 0.8, which is reported for the commonly used 50k SNP chip (Daetwyler, 2009). Values for  $n_G$  were varied from 0 to 4,000 for cows, and was considered to be 0, 200, 500, 2000, 5000, 20,000 for bulls.

The reliability using cow data only  $(R_{cow})$ was predicted using 0.3 for the heritability. To predict the reliability using bull data (R<sub>bull</sub>), the heritability used reflects the reliability of average offspring performances such as daughter yield deviations (VanRaden and Wiggans, 1991) or de-regressed proofs (Sigurdsson and Banos, 1995). We considered that bulls in the reference population had an estimated breeding value based on 100 daughters with records. Combined with the heritability of 0.3 using the formula  $r_{IH}^2 = \frac{\frac{1}{4}nh^2}{1 + \frac{1}{4}(n-1)h^2}$  (Mrode, 2005), this

yields a reliability  $(r_{1H}^2)$  of 0.89, that was used as  $h^2$  in the Daetwyler formula.

Since the cow and bull data partly explain the same variance, we used the 'information source method' (<u>Harris and</u> <u>Johnson, 1998</u>) to blend  $R_{cow}$  and  $R_{bull}$ . Considering the two pieces of information ( $R_{cow}$  and  $R_{bull}$ ), the combined reliability (R) for the new trait is

$$R = \frac{R_{cow} + R_{bull} - 2R_{cow}R_{bull}}{1 - R_{cow}R_{bull}}$$

where

$$R_{bull}(new trait) =$$

 $r_g^2(index, new trait)R_{bull}(index)$ 

#### Predicted response of genomic selection

In the evaluated breeding program, the new trait was considered to have the same

economic value as the index. Note that results for a negative economic value for the new trait, while the genetic correlation between the index and the new trait is positive (e.g. 0.5), are the same as those for a genetic correlation of -0.5.

Numbers of cows and bulls included, were similar to the study of Schrooten et al. (2005). Each generation, 2,000 male and 2,000 female selection candidates were available for selection. For the males, 2% were selected, while for the females 20% were selected for use in the breeding program. In our study, we adopted single stage genomic selection. For the DGV of the new trait, the accuracy of selection was varied from 0.0 to 1.0 with steps of 0.1. Generation intervals of sires to breed bulls and cows, were considered to be 2 years. Generation intervals of dams to breed bulls and cows, were considered to be 3.33 and 5 years, respectively. Those values were adopted from Pryce et al., (2010). All analyses were performed using SelAction (Rutten, et al., 2002), following the guidelines from Dekkers (2007).

## Results

## Predicted DGV reliability

Predicted DGV reliabilities for the new trait are given in Figure 1, when an  $r_g(index, new trait)$  of 0.5 was assumed. Having 1,000 cows in the reference population, yielded a DGV reliability for the new trait of 0.18. Increasing the number of cows to 2,000, yielded a DGV reliability of 0.3. The same was achieved by adding 5,000 bulls with a DGV for the index to a cow reference population of 1,000 animals. The marginal contribution of additional bulls decreases quickly, when the number of already included bulls becomes larger. Note that a genetic correlation between the index and the new trait of -0.5 gives the same results as those in Figure 1. When  $r_g(index, new trait) = 0.0$ , the obtained reliability follows the same curve as for '0 bulls' in Figure 1.

## Predicted response of genomic selection

Predicted responses for genomic selection of the new trait are shown in Figure 2, in genetic standard deviations per year, considering simultaneous selection with the overall index. The vertical line in Figure 2 at a reliability of 0.3 indicates a reference population of 2,000 cows, or one of 1,000 cows supplemented with 5,000 bulls considering a genetic correlation of 0.5 (see Figure 1). Figure 2 indicates that a DGV reliability of 0.3 can already have an important impact on the genetic response of a new trait. For instance, with  $r_g(index,$ new trait) = -0.5, a DGV reliability of 0.3 is sufficient to prohibit a negative genetic trend in the new trait. When  $r_g(index, new)$ trait) = 0.0, a DGV reliability of  $\sim 0.3$ yields a genetic response that is half the genetic response for the index (results for the index are not shown).

## Conclusion

Our results indicate that response to genomic selection for new traits can be important, even when the DGV reliability is much lower than commonly accepted in dairy cattle breeding programs. This relatively low DGV reliability is offset by a decrease in generation intervals. The response for the new trait depends heavily on the genetic correlation with the new trait.

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Figure 1. DGV reliability for a new trait, depending on the number of cows and bulls included in the reference population



Figure 2. Response to genomic selection for the index and the new trait, depending on the genetic correlation with the overall index  $(r_g)$ .