ENLARGING A TRAINING SET FOR GENOMIC SELECTION
BY IMPUTATION OF UN-GENOTYPED ANIMALS IN
POPULATIONS OF VARYING GENETIC ARCHITECTURE

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Introduction

The most common application of imputation is to infer genotypes (GT) of a high-density panel of markers on animals that are genotyped for a low-density panel.

Another application of imputation is to increase the size of the training set with un-genotyped animals. This strategy can be particularly successful when a set of closely related individuals are genotyped.

The posterior probabilities of the dam’s three possible GT can be calculated following Bayes’ theorem and the allele frequency (use the given GT for MGS (M=G) and Sire (S=S0) - \( \rightarrow P(D=G|M=S_0, P=G) = P(D=G|P=G) \), \( P(P=G|S=S_0, D=D_0) = P(P=G | D=D_0) \), \( P(D=G|M=S) = P(D=G) \))

Data

The impact of enlarging a training set (TS) with imputed dams (TSA) on the accuracy of genomic predictions was evaluated for:

- different populations
- varying levels of heritability
- different sizes of genotyped females (TS)

Variants of differing heritability were generated by adding a residual term to the simulated true breeding values (\( h^2 = 0.05, 0.10, ..., 0.50 \)).

Results

Accuracies of genomic prediction for different values of \( h^2 \), number of female progeny in the last generation, and population structure, red/green surfaces: with TS/TSA (90% of the progeny in TS/TS+imputed dams) Single_Step, AlphaImpute

(allele substitution effects of every locus on the simulated phenotypes were fitted in a multiple random regression model similar to the GBLUP method of Meuwissen et al. (2001), accuracies were evaluated as correlation between simulated TBV and estimated GEBV)

Conclusion

With the underlying family structure (typical for Holsteins) imputation can be used to achieve an extra increase in accuracy of genomic predictions by enlarging the training set with completely un-genotyped dams. This strategy was shown to be particularly useful for populations with lower levels of linkage disequilibrium, for genomic selection on traits with low heritability, and for species or breeds for which the size of the reference population is limited.