

Comparative assessment of methods for estimating genomic relationships and their use in predictions in an admixed population

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Introduction

- Accurate estimation of relationships between animals is an important step in any routine genetic evaluations
- Relationships were previously based on pedigree information only
- Conversely, most current evaluations use both markerderived relationship matrix (G) and pedigree-based relationships (A)
- G estimators are more accurate than A because they have more variation between closely related individuals





Introduction

- The accuracy of **G** estimators may be even higher
 - If founder population allele frequencies were available
- In the absence, current population allele frequencies are used to make G and that defines the founder population
- The use of observed allele frequencies in structured populations however, may lead to biased estimation of G



- To estimate A and G matrices
 - Different G matrices were estimated using either observed allele frequencies across breeds or breed allele means
- To estimate breeding values (EBV) and direct genomic values (DGV) using different G matrices

Estimated coefficients and their respective DGVs were compared

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The population

- The Nordic Red dairy cattle (RDC) is a combined population
 - 3 sub-populations from **DNK**, **SWE & FIN**
 - 2nd largest breeding population, with *Ne* larger than Holsteins
 - Most animals in the data (~98%) are composites of breeds
- Absence of pure breed animals remains a major limiting factor for the estimation of breed-specific allele frequencies



- Data were genotypes of 38194 SNP markers for 4106 bulls
- Breed proportions for bulls were estimated from the full Nordic RDC pedigree (>4m animals)
 - 3 main breeds defined with mean BP>10% were,
 - SRB, FAY & NRF
 - Remaining breeds with mean BP<10% -> breed "OTHER"
- Phenotypes were cow IDDs for milk, protein & fat, from 2010 NAV routine evaluations



- Genomic relationships (G) were estimated following methods 1 and 2 by VanRaden (2008)
- **G** estimated using observed allele frequencies (GOF)

GOF = ZZ'/k

•
$$Z_{i,j} \leftarrow (0-2p_j); (1-2p_j); (2-2p_j)$$

Number of "second" alleles

 p_i is the frequency for the 2nd allele & $k=2\sum_i p_i(1-p_i)$ ERSITY OF HELSINKI



G matrices estimated using breed allele means (GBM and GBM2)

GBM = MM'/k

•
$$M_{i,j} \leftarrow (0 - 2p_{ij}); (1 - 2p_{ij}); (2 - 2p_{ij})$$

 p_{ij} is the expected allele frequency of marker *j* for bull *i* given it's base breed proportions

computed by multiple regression of genotypes on BP



Modification of VanRaden method II

There,

$G_{PvRII} = ZDZ'/m = ZD^{0.5}D^{0.5}Z'/m$

m is the number of markers



Following the same, we define:

GBM2 = **M*****M***´/m

$$M *_{i,j} \leftarrow \frac{-2p_{ij}}{\sqrt{2p_{ij}(1-p_{ij})}}; \frac{1-2p_{ij}}{\sqrt{2p_{ij}(1-p_{ij})}}; \frac{2-2p_{ij}}{\sqrt{2p_{ij}(1-p_{ij})}}$$

• m is the number of markers



- Pedigree relationships (A) were estimated for genotyped bulls only, using *RelaX2* computer program
- GOF and GBM2 were combined with 20% weight on A to yield GAOF and GABM2
 - $G^* = wG + (1-w)A$



 Variance components, EBVs & DGVs were estimated separately for each matrix, using a GBLUP model

y = Xb + Za + e,

- y is a vector of cow IDD
- X and Z are design matrices allocating records to b and a
- **b** is a vector of fixed mean and breed regression effects
- **a** is a vector of breeding values
- **e** is a vector of residuals
- Breed regression effects were used only for predictions with GBM and GABM2

Predicted values included fixed regression solutions



Statistics of (diagonals)-1 by estimator

	Mean	Min	Max	Mean	Min	Max
	Acros	s population	ons	Within Swedish bulls		
Α	0.012	0.000	0.135	0.008	0.000	0.081
GOF	0.019	-0.129	0.379	0.006	-0.129	0.184
GBM	-0.051	-0.254	0.310	-0.043	-0.226	0.234
GBM2	-0.242	-0.387	0.093	-0.238	-0.387	0.029
	Withi	n Danish bi	ulls	Within Finnish bulls		
Α	0.007	0.000	0.109	0.016	0.000	0.135
GOF	0.136	-0.027	0.328	-0.021	-0.123	0.157
GBM	-0.040	-0.173	0.310	-0.062	-0.217	0.283
HIGI BIMI PISTO HELSINGFORS UNIVERSITET	-0.233	-0.339	0.093	-0.250	-0.377	0.077

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Histograms of (diagonals)-1 for A and G matrices



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	Across populations		Within Swedish bulls	
Α	0.702	0.661	0.789	0.781
GOF		0.537		0.784
GBM				
	Within Danish bulls		Within Finnish bulls	
Α	0.644	0.856	0.819	0.759
GOF		0.625		0.876
GBM				

Correlations between EBV & DGV from different estimators for validation bulls

	GOF	GBM	GBM2	GAOF	GABM2
A	0.67	0.67	0.66	0.76	0.76
GOF		1.00	1.00	0.98	0.98
GBM	EBV Paren averag	i.e. it je	1.00	0.98	0.98
GBM2				0.98	0.98
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Conclusions

- The use of simple observed allele frequencies across breeds over-estimate values in G for:
 - Populations with the least number of animals in the combined data and/or,
 - Individuals from distantly related populations
- Estimated breed allele means reduced country differences in coefficients, similarly, but shifted them too much torwards zero or less

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Conclusions

- The prediction of DGV converged to similar solutions regardless of allele frequencies used
 - Inclusion of breed regressions for GBM & GBM2 brought breed means back into the DGV
- The validation accuracy slightly increased when A and G matrices were combined
- A single-step GBM2 and A including non-genotyped animals could increase the prediction of DGV even more

