Multiple Traits

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So Far . . .

- Prediction of Breeding Values for one trait
- → univariate analyses
 - In Livestock Breeding, populations are improved with respect to several traits
- \rightarrow multi-trait or multiple trait
 - Different selection strategies and different approaches of how data is analysed are possible

Multiple Trait Selection

- Selection index theory provides a tool for optimal integration of different sources of information
- But still other strategies are applied
 - ► Tandem selection
 - Selection based on independent thresholds

Tandem Selection

- Improve one trait at the time until they all reach a certain threshold
- Problem: For traits which are not improved
 - only correlated selection responses
 - can be negative
- Populations with long generation intervals, response per year is very small

Independent Selection Thresholds

- Applied before selection index
- ▶ Define selection thresholds in each of the traits
- Select animals as parents which are above thresholds for all traits

Example

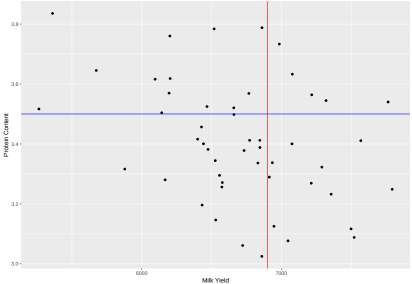


Figure 1: Milk Yield and Protein Content For Dairy Cows

Pros and Cons

- Selection response in all traits
- Thresholds often set to only positive predicted breeding values in all traits
- \rightarrow exclusion of very many animals and reduction in genetic variability
 - Genetic relationships between traits ignored
- ightarrow genetic gain will not be as expected
 - 3. Differences in the economic relevance ignored.
- ightarrow threshold in all traits above positive predicted breeding values emphasizes traits with high heritability

Aggregate Genotype

- ▶ Define the set of important traits for which population should be improved
- ▶ Determine economic values w for these traits
- ► Aggregate genotype *H* follows as

$$H = w^T a$$

Selection Index

Use index I to estimate H where I is a linear combination of information sources

$$I = b^T \hat{a}$$

▶ Index weights *b* are determined using selection index theory as

$$b = P^{-1}Gw$$

- Information sources are predicted breeding values
- ▶ If traits in a and \hat{a} are the same and \hat{a} were estimated using BLUP, then b = w

Implementations

- ▶ First possible implementation
 - Do univariate predictions of breeding values using BLUP animal model
 - ► Combine â with appropriate b-values
- Imrprovement
 - ▶ get â from multivariate analysis

Multivariate Analysis

Given two traits with univariate models

$$y_1 = X_1\beta_1 + Z_1a_1 + e_1$$

 $y_2 = X_2\beta_2 + Z_2a_2 + e_2$

► Combine both univariate models by stacking one on top of the other, resulting in

$$\left[\begin{array}{c} y_1 \\ y_2 \end{array}\right] = \left[\begin{array}{c} X_1 & 0 \\ 0 & X_2 \end{array}\right] \left[\begin{array}{c} \beta_1 \\ \beta_2 \end{array}\right] + \left[\begin{array}{cc} Z_1 & 0 \\ 0 & Z_2 \end{array}\right] \left[\begin{array}{c} a_1 \\ a_2 \end{array}\right] + \left[\begin{array}{c} e_1 \\ e_2 \end{array}\right]$$

Multivariate Model

$$\begin{bmatrix} y_1 \\ y_2 \end{bmatrix} = \begin{bmatrix} X_1 & 0 \\ 0 & X_2 \end{bmatrix} \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix} + \begin{bmatrix} Z_1 & 0 \\ 0 & Z_2 \end{bmatrix} \begin{bmatrix} a_1 \\ a_2 \end{bmatrix} + \begin{bmatrix} e_1 \\ e_2 \end{bmatrix}$$

can be written as

$$y = X\beta + Za + e$$
with $y = \begin{bmatrix} y_1 \\ y_2 \end{bmatrix}$, $\beta = \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix}$, $a = \begin{bmatrix} a_1 \\ a_2 \end{bmatrix}$, $e = \begin{bmatrix} e_1 \\ e_2 \end{bmatrix}$

$$X = \begin{bmatrix} X_1 & 0 \\ 0 & X_2 \end{bmatrix}$$
, $Z = \begin{bmatrix} Z_1 & 0 \\ 0 & Z_2 \end{bmatrix}$

Multivariate Variance-Covariance Matrices

$$G_0 = \begin{bmatrix} \sigma_{g_1}^2 & \sigma_{g1,g2} \\ \sigma_{g1,g2} & \sigma_{g_2}^2 \end{bmatrix} = \begin{bmatrix} g_{11} & g_{12} \\ g_{21} & g_{22} \end{bmatrix}$$

$$var(a) = var \begin{bmatrix} a_1 \\ a_2 \end{bmatrix} = \begin{bmatrix} g_{11}A & g_{12}A \\ g_{21}A & g_{22}A \end{bmatrix} = G_0 \otimes A = G$$

$$R_0 = \begin{bmatrix} r_{11} & r_{12} \\ r_{21} & r_{22} \end{bmatrix}$$

$$R = var(e) = var \begin{bmatrix} e_1 \\ e_2 \end{bmatrix} = \begin{bmatrix} r_{11}I_n & r_{12}I_n \\ r_{21}I_n & r_{22}I_n \end{bmatrix} = R_0 \otimes I_n$$

Solutions

Mixed Model Equations

$$\begin{bmatrix} X^T R^{-1} X & Z^T R^{-1} X \\ Z^T R^{-1} X & Z^T R^{-1} Z + G^{-1} \end{bmatrix} \begin{bmatrix} \hat{\beta} \\ \hat{a} \end{bmatrix} = \begin{bmatrix} X^T R^{-1} y \\ Z^T R^{-1} y \end{bmatrix}$$

Advantages

- some traits have lower heritability than others
- environmental correlations exist between traits measured on the same animal
- some traits are available only a subset of all animals
- some traits were used for a first round of selection
- accuracies are higher in multivariate analyses