### Prediction of Breeding Values

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# Recap Statistical Modelling

- Capture uncertainty due to stochastic relationshipComponents:
  - response variable y
  - predictor variables  $x_1, x_2, \ldots, x_k$
  - error term e
  - ▶ function m(x)

# Model Selection

▶ For fixed effects, select relevant predictors via model selection

- Recommended approach: backwards-elimination
  - 1. start with full model
  - 2. discard predictor variable that increases residual sums of squares the least and get current reduced model
  - 3. repeat step 2 until all predictors are eliminated
  - 4. from above resulting sequence of models, select the one with minimal criterion
- Criterion can be Mallow  $C_p$ , AIC or BIC

### Variation

- Any new trait can only be used for selection, if variation is found in population
- Since change in trait via selection happens between generations, variation must also be at genetic level
- Use mixed linear effects model to estimate genetic variance, often reported as heritability

$$h^2 = \frac{\sigma_u^2}{\sigma_p^2}$$

# Selection Criterion

- If heritability is confirmed
- Selection criterion is needed
- Animals have to be ranked according to the criterion
- Best animals selected as parents of future generation

### What are breeding values

Definition: two times difference between offspring of a given parent from population mean



## Practical Considerations

- Definition of breeding value is based on biological fact that parent passes half of its alleles to offspring
- In practice, definition cannot be used
  - breeding values depend on population (allele frequencies)
  - most parents do not have enough offspring
  - breeding values are needed before animals have offspring
  - different environmental factors not considered

### Solution

- Use genetic model to predict breeding values based on phenotypic observations
- Genetic model decomposes phenotypic observation (y<sub>i</sub>) in different components

$$y_i = \mu + u_i + d_i + i_i + e_i$$

where  $\mu$  is the general mean,  $u_i$  the breeding value,  $d_i$  the dominance deviation,  $i_i$  the epistasis effect and  $e_i$  the random error term.

# Solution II

For predicting breeding values d<sub>i</sub> and i<sub>i</sub> are often ignored, leading to a simplified version of the genetic model

$$y_i = \mu + u_i + e_i$$

Expected values and variance-covariance matrix

$$E\begin{bmatrix} y_i\\ u_i\\ e_i \end{bmatrix} = \begin{bmatrix} \mu\\ 0\\ 0 \end{bmatrix}$$
$$var\begin{bmatrix} y_i\\ u_i\\ e_i \end{bmatrix} = \begin{bmatrix} \sigma_y^2 & \sigma_u^2 & \sigma_e^2\\ \sigma_u^2 & \sigma_u^2 & 0\\ \sigma_e^2 & 0 & \sigma_e^2 \end{bmatrix}$$

## How to Predict Breeding Values

- Predicted breeding values (û) are a function of the observed phenotypic data (y)
- $\rightarrow \hat{u} = f(y)$ 
  - ▶ What should *f*() look like?
  - Goal: Maximize improvement of offspring generation over parents

 $\rightarrow \hat{u}$  should be conditional expected value of true breeding value u given y:

$$\hat{u} = E(u|y)$$

### Derivation

Assume: multivariate normality of u and y and E(u) = 0, then

$$\hat{u} = E(u|y) = E(u) + cov(u, y^{T}) * var(y)^{-1} * (y - E(y)) = E(u|y) = cov(u, y^{T}) * var(y)^{-1} * (y - E(y))$$

- $\hat{u}$  consists of two parts
- 1. (y E(y)): phenotypic observations corrected for environmental effects
- 2.  $cov(u, y^T) * var(y)^{-1}$ : weighting factor of corrected observation

### Unbiasedness

Expected value  $(E(\hat{u}))$ 

$$E(\hat{u}) = E(cov(u, y^{T}) * var(y)^{-1} * (y - E(y)))$$
  
= cov(u, y^{T}) \* var(y)^{-1} \* E(y - E(y))  
= cov(u, y^{T}) \* var(y)^{-1} \* (E(y) - E(y)) = 0

• With E(u) = 0, it follows  $E(\hat{u}) = E(u) = 0$ 

## Variance

•  $var(\hat{u})$  and  $cov(u, \hat{u})$  important for quality of prediction

$$var(\hat{u}) = var(cov(u, y^{T}) * var(y)^{-1} * (y - E(y)))$$
  
=  $cov(u, y^{T}) * var(y)^{-1} * var(y - E(y))$   
\*  $var(y)^{-1} * cov(y, u^{T})$   
=  $cov(u, y^{T}) * var(y)^{-1} * cov(y, u^{T})$   
 $cov(u, \hat{u}) = cov(u, (cov(u, y^{T}) * var(y)^{-1} * (y - E(y)))^{T})$   
=  $cov(u, (y - E(y))^{T}) * var(y)^{-1} * cov(y, u^{T})$   
=  $cov(u, y^{T}) * var(y)^{-1} * cov(y, u^{T}) = var(\hat{u})$ 

### Accuracy

$$r_{u,\hat{u}} = \frac{cov(u,\hat{u})}{\sqrt{var(u) * var(\hat{u})}}$$
$$= \sqrt{\frac{var(\hat{u})}{var(u)}}$$

• Reliability ("Bestimmtheitsmass"):  $B = r_{u,\hat{u}}^2$ 

# Prediction Error Variance (PEV)

• Variability of prediction error:  $u - \hat{u}$ 

$$var(u - \hat{u}) = var(u) - 2cov(u, \hat{u}) + var(\hat{u}) = var(u) - var(\hat{u})$$
$$= var(u) * \left[1 - \frac{var(\hat{u})}{var(u)}\right]$$
$$= var(u) * \left[1 - r_{u,\hat{u}}^{2}\right]$$

Obtained from coefficient matrix of mixed model equations
 Used to compute reliability

### Conditional Density

- Assessment of risk when using animals with predicted breeding values with different reliabilities quantified by f(u|û)
- Multivariate normal density with mean E(u|û) and variance var(u|û)

$$E(u|\hat{u}) = E(u) + cov(u, \hat{u}^{T}) * var(\hat{u})^{-1} * (\hat{u} - E(\hat{u})) = \hat{u}$$
  

$$var(u|\hat{u}) = var(u) - cov(u, \hat{u}^{T}) * var(\hat{u})^{-1} * cov(\hat{u}, u^{T})$$
  

$$= var(u) * \left[1 - \frac{cov(u, \hat{u}^{T})^{2}}{var(u) * var(\hat{u})}\right]$$
  

$$= var(u) * \left[1 - r_{u,\hat{u}}^{2}\right]$$

# Confidence Intervals (CI)

- Assume an error level  $\alpha$ , this results in 100 \*  $(1 \alpha)$ %-Cl
- Typical values of α 0.05 or 0.01
- With α = 0.05, the 95%-CI gives interval around mean which covers a surface of 0.95

CI-Plot



# **CI** Limits

Iower limit I and upper limit m are given by

$$l = \hat{u} - z * SEP$$
  

$$m = \hat{u} + z * SEP$$
(1)

z corresponds to quantile value to cover a surface of (1 - α)
 Use R-function qnorm() to get value of z

## Linear Mixed Effects Model

Use more realistic model for prediction of breeding values

$$y = Xb + Zu + e$$

where

- y vector of length *n* with observations
- *b* vector of length *p* with fixed effects
- *u* vector of length *q* with random breeding values
- *e* vector of length *n* with random error terms
- $X \quad n \times p$  incidence matrix
- Z  $n \times q$  incidence matrix

Expected Values and Variances

$$E\begin{bmatrix} y\\ u\\ e \end{bmatrix} = \begin{bmatrix} Xb\\ 0\\ 0 \end{bmatrix}$$
$$var\begin{bmatrix} y\\ u\\ e \end{bmatrix} = \begin{bmatrix} ZGZ^T + R & ZG & 0\\ GZ^T & G & 0\\ 0 & 0 & R \end{bmatrix}$$

### Solutions

#### Same as for simple model

$$\hat{u} = E(u|y) = GZ^T V^{-1}(y - X\hat{b})$$

with

$$\hat{b} = (X^T V^{-1} X)^- X^T V^{-1} y$$

corresponding to the general least squares solution of b

### Problem

- Solution for û contains V<sup>-1</sup> which is large and difficult to compute
- Use mixed model equations

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

### Sire Model

$$y = Xb + Zs + e$$

where *s* is a vector of length  $q_s$  with all sire effects.

$$var(s) = A_s * \sigma_s^2$$

where  $A_s$ : numerator relationship considering only sires

### Animal Model

$$y = Xb + Za + e$$

where a is a vector of length  $q_a$  containing the breeding values

$$\mathit{var}(\mathit{a}) = \mathit{A}\sigma^2_{\mathit{a}}$$

where A is the numerator relationship matrix

# Genomic BLUP (GBLUP)

- 1. marker-effect models: SNP-loci as random effects (MEM)
- 2. breeding value based models: genomic breeding values as random effects (BVM)

# $\mathsf{MEM} \text{ and } \mathsf{BVM}$



# Marker Effect Model (MEM)

Marker effects (a) as random in a linear mixed effects model

$$y = X\beta + Ma + e$$

Solution of marker effects via mixed model equations

$$\begin{bmatrix} X^{T}R^{-1}X & X^{T}R^{-1}M \\ M^{T}R^{-1}X & M^{T}R^{-1}M + I * \sigma_{a}^{-2} \end{bmatrix} \begin{bmatrix} \hat{\beta} \\ \hat{a} \end{bmatrix} = \begin{bmatrix} X^{T}R^{-1}y \\ M^{T}R^{-1}y \end{bmatrix}$$

where  $\sigma_a^2$  is the SNP-effect variance component.

Genomic breeding value for animal *i* is computed as sum over appropriate values of *â* given by genotype of animal *i* 

## Breeding Value Based Model

 Genomic breeding values (u) as random effects in linear mixed effects model

$$y = X\beta + Wu + e$$



$$\begin{bmatrix} X^{\mathsf{T}}R^{-1}X & X^{\mathsf{T}}R^{-1}W \\ W^{\mathsf{T}}R^{-1}X & W^{\mathsf{T}}R^{-1}W + G^{-1}*\sigma_u^{-2} \end{bmatrix} \begin{bmatrix} \hat{\beta} \\ \hat{u} \end{bmatrix} = \begin{bmatrix} X^{\mathsf{T}}R^{-1}y \\ W^{\mathsf{T}}R^{-1}y \end{bmatrix}$$

• Genomic breeding values correspond to solutions for  $\hat{u}$ 

### How Does GBLUP Work

$$\begin{bmatrix} X^{\mathsf{T}}X & X^{\mathsf{T}}Z & 0\\ Z^{\mathsf{T}}X & Z^{\mathsf{T}}Z + G^{(11)} & G^{(12)}\\ 0 & G^{(21)} & G^{(22)} \end{bmatrix} \begin{bmatrix} \hat{b}\\ \hat{g}_1\\ \hat{g}_2 \end{bmatrix} = \begin{bmatrix} X^{\mathsf{T}}y\\ Z^{\mathsf{T}}y\\ 0 \end{bmatrix}$$

G<sup>(11)</sup>: animals with phenotypic observations
 G<sup>(22)</sup>: animals without phenotypic observations

$$\hat{g}_2 = -\left(G^{22}
ight)^{-1}G^{21}\hat{g}_1$$

# Summary for One Trait

- Predicted breeding values with or without genomic information
- Animals can be ranked according to predicted breeding values
- Problems:
  - Not only one trait should be improved
  - Selection for one trait changes also other traits via correlated selection response

 $\rightarrow$  see genetic trends at: https://1-htz.quagzws.com/shiny/users/ zws/genTrendHolstein\_DE/index.Rmd

### Example Fertility and Production



### Multi-Trait Selection



# Types of Multi-Trait Selection

#### Tandem selection

- select for one trait at the time
- after goal has been reached change to different trait
- Independent selection limits
  - select only animals which fullfill criteria in all traits
- Selection according to aggregate genotype
  - combine traits into aggregate genotype H
  - define H as weighted sum of true breeding values and economic values
  - use selection index I to estimate H

# Aggregate Genotype

Definition in vector notation:  $H = v^T \cdot u$ 

where

- u: vector of true breeding values
- v vector of economic values which are marginal changes in profit for a small change in the population mean of the trait

Estimate *H* via index *I*, hence  $\hat{H} = I = b^T x$ 

with

- x: a vector of information sources
- b: a vector of unknown weights.

Determine b such that var(I - H) is minimal.

### Find *b* . . .

... such that var(I - H)

$$var(I - H) = var(I) - 2 * cov(I, H) + var(H)$$
  
=  $var(b^Tx) - 2 * cov(b^Tx, v^Tu) + var(v^Tu)$   
=  $b^T var(x)b - 2 * b^T cov(x, u^T)v + v^T var(u)v$   
=  $b^T Pb - 2 * b^T Cv + v^T Gv$ 

Setting  $\frac{\partial \operatorname{var}(I-H)}{\partial b} = 0$  leads to

$$Pb = Cv$$

Hence

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

## Special Case

- Same traits in *H* and in *I*
- Use predicted breeding values û from multivariate BLUP animal model as information source x
- Then it follows

$$b = P^{-1}Cv = var(\hat{u})^{-1} \cdot cov(\hat{u}, u^{T}) \cdot v = v$$