Mixed models applied to breeding

Alencar Xavier March 13th - March 15th, 2019

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Schedule

- · Module 1: Intro to mixed models
- Module 2: Fitting mixed models
- Module 3: Advanced topics
- · Module 4: Signal detection
- Module 5: Association analysis

Module 1 - Introduction to mixed models

Outline

Part 1: Concepts

- History of mixed models
- Mixed models in plant breeding
- · Fixed and random terms
- Model notation
- · Variance decomposition

Part 2: Applications

- · Selection models
- Practical examples
- · Variance components
- Ridges and Kernels

Part 1 - Concepts

History of mixed models

- 1886: Regression and heritability
- 1918: Infinitesimal model (P = G + E)
- 1922: Genetic relationship
 - 1968: BLUP using relationship



Mixed models in plant breeding

- of plant breeding (Xavier et al 2017)
- Variance components and heritability (Johnson and Thompson 1994)
- Trait associations (Gianola and Sorensen 2014)
- Estimation of genetic and breedingvalues (Piepho et al 2008)
- Prediction of unphenotyped lines (de los Campos et al 2013)
- Selection index (Wientjes et al. 2016)
- Genome-wide association analysis (Yang et al 2014)
- All sorts of inference (Robinson 1991)

Fixed and random terms

Fixed effect

- Assumed to be invariable (often you cannot recollect the data)
- Inferences are made upon the parameters
- Results can not be extrapolated to other datasets
- Example: Overall mean and environmental effects

Random effects

- You may not have all the levels available
- · Inference are made on variance components
- Prior assumption: coefficients are normally distributed
- Results can not be extrapolated to other datasets
- · Example: Genetic effects

Let's unleash the beast



Model notation

- Linear model: y = Xb + Zu + e
- With variance: $y \sim N(Xb, ZKZ\sigma_u^2 + I\sigma_e^2)$

Assuming: $u \sim N(0, K\sigma_u^2)$ and $e \sim N(0, I\sigma_e^2)$

Henderson equation

$$egin{bmatrix} X'X & Z'X \ X'Z & Z'Z + \lambda K^{-1} \end{bmatrix} egin{bmatrix} b \ u \end{bmatrix} = egin{bmatrix} X'y \ Z'y \end{bmatrix}$$

Summary:

- We know (data): $x = \{y, X, Z, K\}$
- We want (parameters): $\theta = \{b, u, \sigma_a^2, \sigma_e^2\}$
- Estimation based on Gaussian likelihood: $L(x|\theta)$

Model notation

```
y = vector of observations ( )
• X = design matrix of fixed effects (x)
· Z = design (or incidence) matrix of random effects ( x )

    K = random effect correlation matrix ( x )

    u = vector of random effect coefficients ( )

    b = vector of fixed effect coefficients ( )

• e = vector of residuals ( )
• \sigma_a^2 = marker effect variance (1)
• \sigma_u^2 = random effect variance (1)
• \sigma_e^2 = residual variance (1)
• \lambda = \sigma_e^2/\sigma_u^2 (Regularization parameters) (1)
```

Model notation

The mixed model can also be notated as follows

$$y = Wg + e$$

Solved as

$$[W'W + \Sigma]g = [W'y]$$

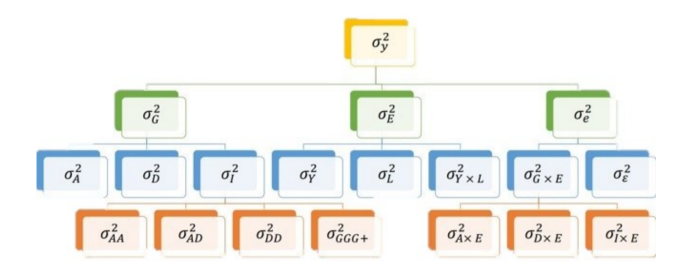
Where

$$W = [X, Z]$$

$$g = [b, u]$$

$$\Sigma = egin{bmatrix} 0 & 0 \ 0 & \lambda K^{-1} \end{bmatrix}$$

Variance decomposition



Part 2 - Applications

Selection

1

- BLUPs or BLUEs from replicated trials
- Captures additive and non-additive genetics together

2

- · Use pedigree information to create K
- Captures additive genetics (heritable)
- Trials not necessarily replicated

3

- Genotypic information replaces pedigree
- · Any signal: additivity, dominance and epistasis

- Example 1: Balanced data, no kinship
- · Example 2: Balanced data, with kinship
- · Example 3: Unbalanced data, with kinship
- · Example 4: Balanced data, missing individual

Data:

```
Env Gen Phe
## 1
      E1
          G1 47
      E1 G2 51
          G3
              46
## 4
      E1 G4
              58
## 5
          G1
              52
## 6
              46
## 7
          G3
              52
## 8
              54
## 9
          G1
              53
## 10
              48
      E3
          G2
## 11
      E3
          G3
              58
## 12 E3 G4 52
```

 $Model: Phenotype = Environment_{(F)} + Genotype_{(R)}$

Design matrix W:

##		EnvE1	EnvE2	EnvE3	GenG1	GenG2	GenG3	GenG4
##	1	1	0	0	1	0	0	0
##	2	1	0	0	0	1	0	0
##	3	1	0	0	0	0	1	0
##	4	1	0	0	0	0	0	1
##	5	0	1	0	1	0	0	0
##	6	0	1	0	0	1	0	0
##	7	0	1	0	0	0	1	0
##	8	0	1	0	0	0	0	1
##	9	0	0	1	1	0	0	0
##	10	0	0	1	0	1	0	0
##	11	0	0	1	0	0	1	0
##	12	0	0	1	0	0	0	1

W'W:

##		EnvE1	EnvE2	EnvE3	GenG1	GenG2	GenG3	GenG4
##	EnvE1	4	0	0	1	1	1	1
##	EnvE2	0	4	0	1	1	1	1
##	EnvE3	0	0	4	1	1	1	1
##	GenG1	1	1	1	3	0	0	0
##	GenG2	1	1	1	0	3	0	0
##	GenG3	1	1	1	0	0	3	0
##	GenG4	1	1	1	0	0	0	3

Left-hand side ($W'W + \Sigma$):

```
EnvE1 EnvE2 EnvE3 GenG1 GenG2 GenG3 GenG4
##
## EnvE1
                               1.00 1.00 1.00
## EnvE2
                          1.00
                               1.00
                                     1.00 1.00
## EnvE3
                       4 1.00
                               1.00
                                     1.00 1.00
## GenG1
                       1 3.17
                               0.00
                                     0.00 0.00
## GenG2
                               3.17
                                     0.00 0.00
## GenG3
                       1 0.00
                               0.00
                                     3.17 0.00
## GenG4
                       1 0.00 0.00 0.00 3.17
```

Assuming independent individuals: K = I

Regularization:
$$\lambda = \sigma_e^2/\sigma_u^2 = 1.64/9.56 = 0.17$$

Right-hand side (W'y):

```
## EnvE1 202
## EnvE2 204
## EnvE3 211
## GenG1 152
## GenG2 145
## GenG3 156
## GenG4 164
```

We can find coefficients through least-square solution

$$g = (LHS)^{-1}(RHS) = (W'W + \Sigma)^{-1}W'y$$

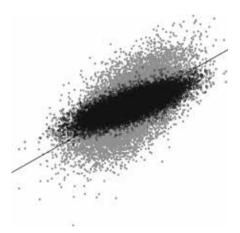
```
## [,1]
## EnvE1 50.50
## EnvE2 51.00
## EnvE3 52.75
## GenG1 -0.71
## GenG2 -2.92
## GenG3 0.55
## GenG4 3.08
```

Shrinkage

$$BLUE = \frac{w'y}{w'w} = \frac{sum}{n} =$$

$$BLUP = rac{w'y}{w'w+\lambda} = rac{sum}{n+\lambda}$$
 =

$$= BLUE \times h^2$$



Note:

- More observations = less shrinkage
- Higher heritability = less shrinkage: $\lambda = \frac{h^2-1}{h^2}$

If we know the relationship among individuals:

```
## GenG1 GenG2 GenG3 GenG4
## GenG1 1.00 0.64 0.23 0.48
## GenG2 0.64 1.00 0.33 0.67
## GenG3 0.23 0.33 1.00 0.31
## GenG4 0.48 0.67 0.31 1.00
```

Then we estimate λK^{-1}

```
## GenG1 GenG2 GenG3 GenG4
## GenG1 0.15 -0.09 0.00 -0.01
## GenG2 -0.09 0.22 -0.02 -0.10
## GenG3 0.00 -0.02 0.10 -0.02
## GenG4 -0.01 -0.10 -0.02 0.17
```

Regularization: $\lambda = \sigma_e^2/\sigma_u^2 = 1.64/17.70 = 0.09$

And the left-hand side becomes

##		EnvE1	EnvE2	EnvE3	GenG1	GenG2	GenG3	GenG4
##	EnvE1	4	0	0	1.00	1.00	1.00	1.00
##	EnvE2	0	4	0	1.00	1.00	1.00	1.00
##	EnvE3	0	0	4	1.00	1.00	1.00	1.00
##	GenG1	1	1	1	3.15	-0.09	0.00	-0.01
##	GenG2	1	1	1	-0.09	3.22	-0.02	-0.10
##	GenG3	1	1	1	0.00	-0.02	3.10	-0.02
##	GenG4	1	1	1	-0.01	-0.10	-0.02	3.17

We can find coefficients through least-square solution

$$g = (LHS)^{-1}(RHS) = (W'W + \Sigma)^{-1}W'y$$

```
## EnvE1 51.05
## EnvE2 51.55
## EnvE3 53.30
## GenG1 -1.32
## GenG2 -3.34
## GenG3 0.03
## GenG4 2.45
```

Genetic coefficients shrink more: Var(A) < Var(G)

What if we have missing data?

```
Env Gen Phe
## 1
      E1
          G1 47
      E1 G2 51
          G3
              NA
## 4
      E1 G4
              58
      E2 G1
              52
## 6
          G2
              46
          G3
              52
## 8
      E2 G4
              NA
## 9
          G1
              53
## 10
      E3
          G2
              48
## 11
      E3
          G3
              58
## 12
      E3 G4 52
```

Rows of missing points are removed

##		EnvE1	EnvE2	EnvE3	GenG1	GenG2	GenG3	GenG4
##	1	1	0	0	1	0	0	0
##	2	1	0	0	0	1	0	0
##	4	1	0	0	0	0	0	1
##	5	0	1	0	1	0	0	0
##	6	0	1	0	0	1	0	0
##	7	0	1	0	0	0	1	0
##	9	0	0	1	1	0	0	0
##	10	0	0	1	0	1	0	0
##	11	0	0	1	0	0	1	0
##	12	0	0	1	0	0	0	1

W'W:

##		EnvE1	EnvE2	EnvE3	GenG1	GenG2	GenG3	GenG4
##	EnvE1	3	0	0	1	1	0	1
##	EnvE2	0	3	0	1	1	1	0
##	EnvE3	0	0	4	1	1	1	1
##	GenG1	1	1	1	3	0	0	0
##	GenG2	1	1	1	0	3	0	0
##	GenG3	0	1	1	0	0	2	0
##	GenG4	1	0	1	0	0	0	2

Left-hand side ($W'W + \Sigma$):

```
EnvE1 EnvE2 EnvE3 GenG1 GenG2 GenG3 GenG4
##
## EnvE1
                          1.00 1.00 0.00 1.00
## EnvE2
                         1.00
                                1.00 1.00 0.00
## EnvE3
                       4 1.00 1.00 1.00 1.00
                       1 3.10 -0.06 0.00 -0.01
## GenG1
## GenG2
                       1 -0.06 3.15 -0.01 -0.07
                       1 0.00 -0.01 2.07 -0.01
## GenG3
## GenG4
                       1 -0.01 -0.07 -0.01 2.11
```

Regularization:
$$\lambda = \sigma_e^2/\sigma_u^2 = 1.21/19.61 = 0.06$$

Right-hand side (W'y):

```
## EnvE1 156
## EnvE2 150
## EnvE3 211
## GenG1 152
## GenG2 145
## GenG3 110
## GenG4 110
```

Find coefficients through least-square solution

$$g = (LHS)^{-1}(RHS) = (W'W + \Sigma)^{-1}W'y$$

```
## [,1]
## EnvE1 54.14
## EnvE2 51.70
## EnvE3 53.82
## GenG1 -2.56
## GenG2 -4.68
## GenG3 2.15
## GenG4 0.81
```

What if we are missing data from a individual?

```
Env Gen Phe
## 1
      E1
          G1 NA
      E1 G2 51
          G3
              46
## 4
      E1 G4 58
         G1
              NA
## 6
              46
## 7
          G3
              52
## 8
      E2 G4 54
## 9
          G1 NA
## 10
              48
      E3
          G2
## 11
      E3
          G3
              58
## 12
      E3 G4 52
```

Rows of missing points are removed

##		EnvE1	EnvE2	EnvE3	GenG1	GenG2	GenG3	GenG4
##	2	1	0	0	0	1	0	0
##	3	1	0	0	0	0	1	0
##	4	1	0	0	0	0	0	1
##	6	0	1	0	0	1	0	0
##	7	0	1	0	0	0	1	0
##	8	0	1	0	0	0	0	1
##	10	0	0	1	0	1	0	0
##	11	0	0	1	0	0	1	0
##	12	0	0	1	0	0	0	1

W'W:

##		EnvE1	EnvE2	EnvE3	GenG1	GenG2	GenG3	GenG4
##	EnvE1	3	0	0	0	1	1	1
##	EnvE2	0	3	0	0	1	1	1
##	EnvE3	0	0	3	0	1	1	1
##	GenG1	0	0	0	0	0	0	0
##	GenG2	1	1	1	0	3	0	0
##	GenG3	1	1	1	0	0	3	0
##	GenG4	1	1	1	0	0	0	3

Left-hand side ($W'W + \Sigma$):

```
EnvE1 EnvE2 EnvE3 GenG1 GenG2 GenG3 GenG4
##
## EnvE1
                          0.00 1.00 1.00 1.00
## EnvE2
                                1.00 1.00 1.00
                          0.00
## EnvE3
                        3 0.00 1.00 1.00 1.00
                       0 0.14 -0.08 0.00 -0.01
## GenG1
## GenG2
                       1 -0.08 3.19 -0.02 -0.09
                       1 0.00 -0.02 3.09 -0.01
## GenG3
## GenG4
                        1 -0.01 -0.09 -0.01 3.15
```

Regularization: $\lambda = \sigma_e^2/\sigma_u^2 = 1.79/22.78 = 0.08$

Right-hand side (W'y):

```
## EnvE1 155
## EnvE2 152
## EnvE3 158
## GenG1 0
## GenG2 145
## GenG3 156
## GenG4 164
```

Find coefficients through least-square solution

$$g = (LHS)^{-1}(RHS) = (W'W + \Sigma)^{-1}W'y$$

```
## [,1]
## EnvE1 52.06
## EnvE2 51.06
## EnvE3 53.06
## GenG1 -1.82
## GenG2 -3.48
## GenG3 -0.07
## GenG4 2.38
```

Variance components

Expectation-Maximization REML (1977)

$$\sigma_u^2=rac{u'K^{-1}u}{q-\lambda tr(K^{-1}C^{22})}$$
 and $\sigma_e^2=rac{e'y}{n-p}$

Bayesian Gibbs Sampling (1993)

$$\sigma_u^2=rac{u'K^{-1}u+S_u
u_u}{\chi^2(q+
u_u)}$$
 and $\sigma_e^2=rac{e'e+S_e
u_e}{\chi^2(n+
u_e)}$

Predicted Residual Error Sum of Squares (PRESS) (2017)

- $\lambda = argmin(\sum e_i^2/(1-h_{ii})^2)$
- Where $H=(I+K\lambda)^{-1}$ and $e=y-\mu-Hy$

Kernel methods:

- Genetic signal is captured by the relationship matrix K
- Random effect coefficients are the breeding values (BV)
- Efficient to compute BV when $markers \gg individuals$
- Easy use and combine pedigree, markers and interactions

Ridge methods:

- ullet Genetic signal is captured by the design matrix M
- · Random effect coefficients are the marker effects
- · Easy way to make predictions of unobserved individuals
- Enables to visualize where the QTLs are in the genome

Kernel

$$y = Xb + Zu + e$$
 , $u \sim N(0, K\sigma_u^2)$

Ridge

$$y = Xb + Ma + e$$
 , $a \sim N(0, I\sigma_a^2)$

Where

- M is the genotypic matrix, $m_{ij} = \{0, 1, 2\}$
- $K = \alpha MM'$
- u = Ma
- $\sigma_a^2 = \alpha \sigma_u^2$

Kernel model

$$egin{bmatrix} X'X & Z'X \ X'Z & Z'Z + K^{-1}(\sigma_e^2/\sigma_u^2) \end{bmatrix} egin{bmatrix} b \ u \end{bmatrix} = egin{bmatrix} X'y \ Z'y \end{bmatrix}$$

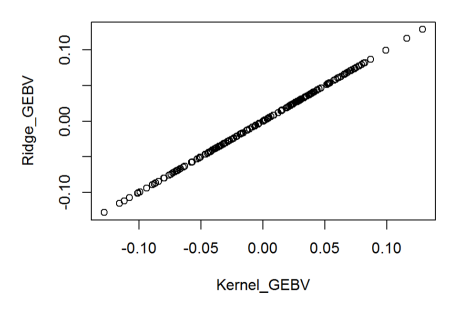
Ridge model

$$egin{bmatrix} X'X & M'X \ X'M & M'M + I^{-1}(\sigma_e^2/\sigma_a^2) \end{bmatrix} egin{bmatrix} b \ a \end{bmatrix} = egin{bmatrix} X'y \ M'y \end{bmatrix}$$

Both models capture same genetic signal (de los Campos 2015)

```
K = tcrossprod(M)/ncol(M)
GBLUP = reml(y=y,K=K); Kernel_GEBV = GBLUP$EBV
RRBLUP = reml(y=y,Z=M); Ridge_GEBV = M%*%RRBLUP$EBV
plot(Kernel_GEBV,Ridge_GEBV, main='Comparing results')
```

Comparing results



Break

Module 2 - Fitting mixed models

Example 1 - Sorghum

Example 1 - load data

```
Example dataset from Kevin's
```

data(adugna.sorghum, package = 'agridat')

package

```
dt = adugna.sorghum
head(dt)
     gen trial env yield year
## 1 G16
           T2 E01
                    590 2001 Mieso
## 2 G17
           T2 E01
                    554 2001 Mieso
## 3 G18
           T2 E01
                    586 2001 Mieso
           T2 E01
                    738 2001 Mieso
## 4 G19
## 5 G20
           T2 E01
                    489 2001 Mieso
## 6 G21
           T2 E01
                    684 2001 Mieso
```

Example 1 - Getting design matrix

- Linear model: Pheotype = Env + Gen
- In algebra notation: y = Xb + Zu + e

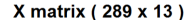
```
y = dt$yield
X = model.matrix(y~env,dt)
Z = model.matrix(y~gen-1,dt) # "-1" means no intercept
```

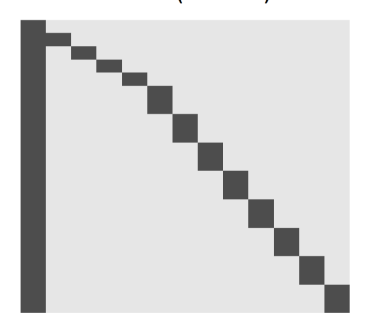
Assuming:

- $u \sim N(0, I\sigma_g^2)$
- · $e \sim N(0, I\sigma_e^2)$

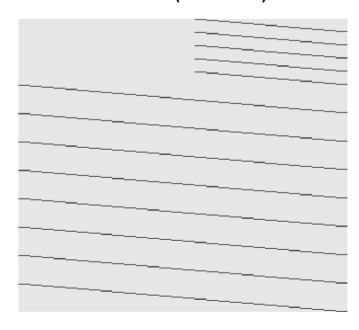
Example 1 - Visualize X and Z matrices

```
SEE=function(A,...)image(t(1-A[nrow(A):1,]),axes=F,col=gray.colors(2),...)
par(mfrow=c(1,2),mar=c(1,1,3,1))
SEE(X, main=paste("X matrix (",paste(dim(X),collapse=' x '),")" ))
SEE(Z, main=paste("Z matrix (",paste(dim(Z),collapse=' x '),")" ))
```





Z matrix (289 x 28)



Example 1 - Fit the model

```
# Using the NAM package (same for rrBLUP, EMMREML, BGLR)
require(NAM, quietly = TRUE)
fit1 = reml(y=y,X=X,Z=Z)
# Alternatively, you can also use formulas with NAM
fit1b = reml(y=dt$yield,X=~dt$env,Z=~dt$gen )
# Using the Lme4 package
require(lme4, quietly = TRUE)
fit2 = lmer(yield ~ env + (1|gen), data=dt)
```

Example 1 - Variance components

fit1\$VC[c(1:2)] # same with fit1b\$VC

data.frame((summary(fit2))\$varcor)\$vcov

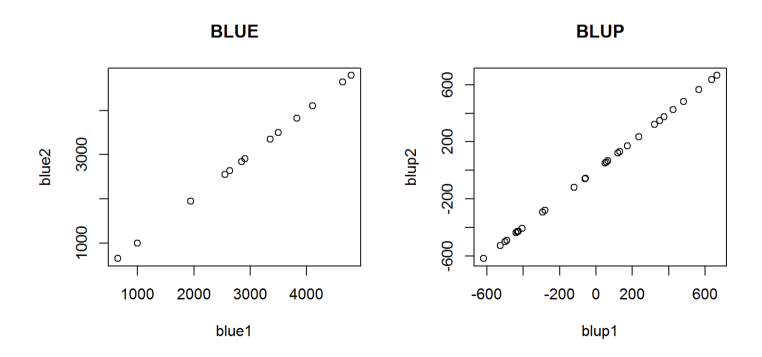
· VC can be used to measure

heritability

$$H=rac{\sigma_g^2}{\sigma_g^2+\sigma_e^2/n}=rac{189680.4}{189680.4+442075.6/10.32}=0.82$$

Example 1 - The coefficients

```
blue1 = fit1$Fixed[,1]; blup1 = fit1$EBV
blue2 = fit2@beta; blup2 = rowMeans(ranef(fit2)$gen)
par(mfrow=c(1,2));
plot(blue1,blue2,main="BLUE"); plot(blup1,blup2,main="BLUP")
```

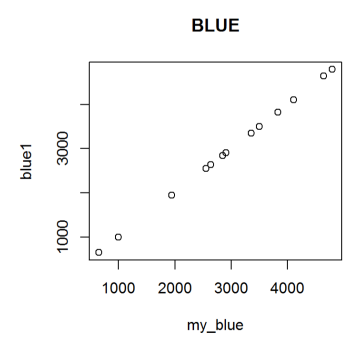


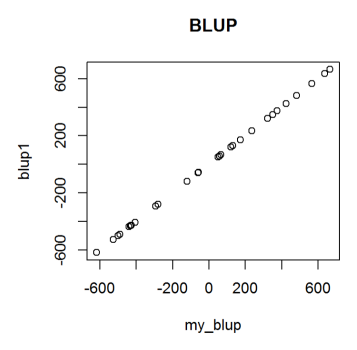
Example 1 - DIY BLUPs

```
iK = diag(ncol(Z))
Lambda = 442075.6/189680.4
W = cbind(X,Z)
Sigma = as.matrix(Matrix::bdiag(diag(0,ncol(X)),iK*Lambda))
LHS = crossprod(W) + Sigma
RHS = crossprod(W,y)
g = solve(LHS,RHS)
my_blue = g[ c(1:ncol(X))]
my_blue = g[-c(1:ncol(X))]
```

Example 1 - DIY BLUPs

```
par(mfrow=c(1,2))
plot(my_blue,blue1,main="BLUE")
plot(my_blup,blup1,main="BLUP")
```





Example 1 - DIY Variance components

```
\sigma_e^2=rac{e'y}{n-p} and \sigma_u^2=rac{u'K^{-1}u+tr(K^{-1}C^{22}\sigma_e^2)}{q}
e = y - X \% my blue - Z %*% my blup
Ve = c(y%*\%e)/(length(y)-ncol(X))
Ve
## [1] 442075.6
trKC22 = sum(diag(iK%*%(solve(LHS)[-c(1:ncol(X)),-c(1:ncol(X))])))
Vg = Vg = c(t(my blup)%*%iK%*%my blup+trKC22*Ve)/ncol(Z)
Vg
## [1] 189680.4
```

Starting from bad variance components

```
Ve = Vg = 1
for(i in 1:25){
 Lambda = Ve/Vg;
  Sigma = as.matrix(Matrix::bdiag(diag(0,ncol(X)),iK*Lambda))
  LHS = crossprod(W) + Sigma; RHS = crossprod(W,y); g = solve(LHS,RHS)
 my blue = g[c(1:ncol(X))]; my blup = g[-c(1:ncol(X))]
  e = v - X%*mv blue - Z%*mv blup; Ve = c(v)*%e)/(length(v)-ncol(X))
 trKC22 = sum(diag(iK%*%(solve(LHS)[(ncol(X)+1):(ncol(W)),(ncol(X)+1):(ncol(W))])))
 Vg = c(t(my blup)%*%iK%*%my blup+trKC22*Ve)/ncol(Z)
  if(!i%5){cat('It',i,'VC: Vg =',Vg,'and Ve =',Ve,'\n')}}
## It 5 VC: Vg = 191751.1 and Ve = 441110.7
## It 10 VC: Vg = 189728.4 and Ve = 442053.2
## It 15 VC: Vg = 189681.5 and Ve = 442075.1
## It 20 VC: Vg = 189680.4 and Ve = 442075.6
## It 25 VC: Vg = 189680.4 and Ve = 442075.6
```

Example 2 - Barley

Example 2 - load data

Another example dataset from Kevin's package

```
data(steptoe.morex.pheno,package='agridat')
dt = steptoe.morex.pheno
head(dt)
```

```
env amylase diapow hddate lodging malt height protein yield
##
        gen
## 1 Steptoe
             MN92
                     22.7
                              46 149.5
                                             NA 73.6
                                                       84.5
                                                              10.5 5.5315
                              72 178.0
## 2 Steptoe MTi92
                     30.1
                                             10 76.5
                                                        NA
                                                              11.2 8.6403
## 3 Steptoe MTd92
                     26.7
                              78 165.0
                                             15 74.5
                                                      75.5
                                                              13.4 5.8990
                     26.2
                                             NA 74.1 111.0
## 4 Steptoe ID91
                              74 179.0
                                                              12.1 8.6290
## 5 Steptoe
             OR91
                     19.6
                              62 191.0
                                             NA 71.5
                                                       90.0
                                                              11.7 5.3440
## 6 Steptoe WA91
                     23.6
                              54 181.0
                                             NA 73.8 112.0
                                                              10.0 6.2700
```

Example 2 - Getting design matrix

- Linear model: Phe = Env + Gen
- In algebra notation: y = Xb + Zu + e

```
X = model.matrix(~env,dt)
Z = model.matrix(~gen-1,dt) # "-1" means no intercept
y = dt$yield
```

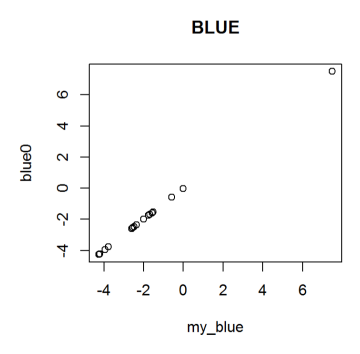
Example 2 - Fit the model

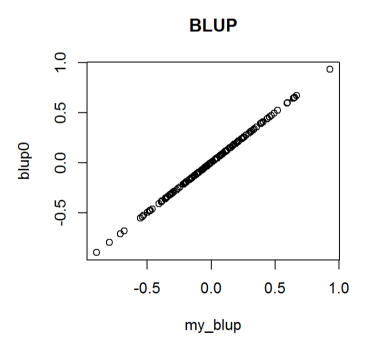
Example 2 - DIY BLUPs

```
iK = diag(ncol(Z))
Lambda = 0.637997/0.132009
W = cbind(X,Z)
Sigma = as.matrix(Matrix::bdiag(diag(0,ncol(X)),iK*Lambda))
LHS = crossprod(W) + Sigma
RHS = crossprod(W,y)
g = solve(LHS,RHS)
my_blue = g[ c(1:ncol(X))]
my_blue = g[-c(1:ncol(X))]
```

Example 2 - DIY BLUPs

```
par(mfrow=c(1,2))
plot(my_blue,blue0,main="BLUE")
plot(my_blup,blup0,main="BLUP")
```





Example 2 - Check variance components

```
\sigma_e^2=rac{e'y}{n-p} and \sigma_u^2=rac{u'K^{-1}u+tr(K^{-1}C^{22}\sigma_e^2)}{q}
e = y - X \% my blue - Z %*% my blup
Ve = c(y%*\%e)/(length(y)-ncol(X))
Ve
## [1] 0.6379967
trKC22 = sum(diag(iK%*%(solve(LHS)[-c(1:ncol(X)),-c(1:ncol(X))])))
Vg = c(t(my blup)%*%iK%*%my blup+trKC22*Ve)/ncol(Z)
Vg
## [1] 0.1320091
```

Starting from bad variance components

```
Ve = Vg = 1
for(i in 1:25){
 Lambda = Ve/Vg;
  Sigma = as.matrix(Matrix::bdiag(diag(0,ncol(X)),iK*Lambda))
  LHS = crossprod(W) + Sigma; RHS = crossprod(W,y); g = solve(LHS,RHS)
 my blue = g[c(1:ncol(X))]; my blup = g[-c(1:ncol(X))]
  e = v - X%*mv blue - Z%*mv blup; Ve = c(v)*%e)/(length(v)-ncol(X))
 trKC22 = sum(diag(iK%*%(solve(LHS)[(ncol(X)+1):(ncol(W)),(ncol(X)+1):(ncol(W))])))
 Vg = c(t(my blup)%*%iK%*%my blup+trKC22*Ve)/ncol(Z)
  if(!i%5){cat('It',i,'VC: Vg =',Vg,'and Ve =',Ve,'\n')}}
## It 5 VC: Vg = 0.1336139 and Ve = 0.6370751
## It 10 VC: Vg = 0.1320386 and Ve = 0.6379797
## It 15 VC: Vg = 0.1320097 and Ve = 0.6379964
## It 20 VC: Vg = 0.1320092 and Ve = 0.6379967
## It 25 VC: Vg = 0.1320092 and Ve = 0.6379967
```

Example 3 - Barley GEBV

Using genomic information!

```
data(steptoe.morex.geno,package='agridat')
gen = do.call("cbind",lapply(steptoe.morex.geno$geno,function(x) x$data))
gen = rbind(0,2,gen)
rownames(gen) = c('Morex','Steptoe',as.character(steptoe.morex.geno$pheno$gen))
rownames(gen)[10] = "SM8"
gen = gen[gsub('gen','',colnames(Z)),]
K = GRM(IMP(gen),T)
```

Example 3 - Fit the model

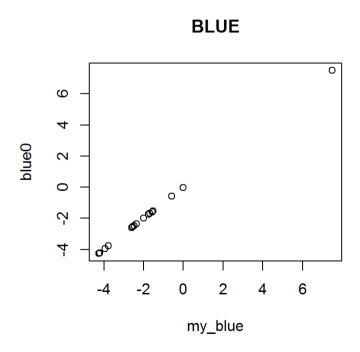
Example 3 - DIY BLUPs

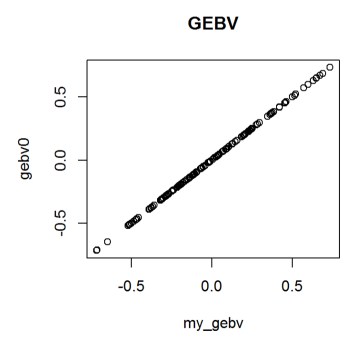
```
diag(K) = diag(K)+1e-08; iK = solve(K)
Lambda = 0.6575/1.0280

W = cbind(X,Z)
Sigma = as.matrix(Matrix::bdiag(diag(0,ncol(X)),iK*Lambda))
LHS = crossprod(W) + Sigma
RHS = crossprod(W,y)
g = solve(LHS,RHS)
my_blue = g[ c(1:ncol(X))]
my_gebv = g[-c(1:ncol(X))]
```

Example 3 - DIY BLUPs

```
par(mfrow=c(1,2))
plot(my_blue,blue0,main="BLUE")
plot(my_gebv,gebv0,main="GEBV")
```





Example 3 - Check variance components

$$\begin{split} \sigma_e^2 &= \frac{e'y}{n-p} \text{ and } \sigma_u^2 = \frac{u'K^{-1}u + tr(K^{-1}C^{22}\sigma_e^2)}{q} \\ \text{e = y - X \% my_blue - Z \% my_blup} \\ \text{Ve = c(y\%\%e)/(length(y)-ncol(X))} \\ \text{Ve} \\ &\# \text{ [1] } 0.6379967 \\ \\ \text{trKC22 = sum(diag(iK\%\%(solve(LHS)[(ncol(X)+1):(ncol(W)),(ncol(X)+1):(ncol(W))])))} \\ \text{Vg = c(t(my_blup)\%\%iK\%\%my_blup+trKC22*Ve)/ncol(Z)} \\ \text{Vg} \\ &\# \text{ [1] } 14.12085 \end{split}$$

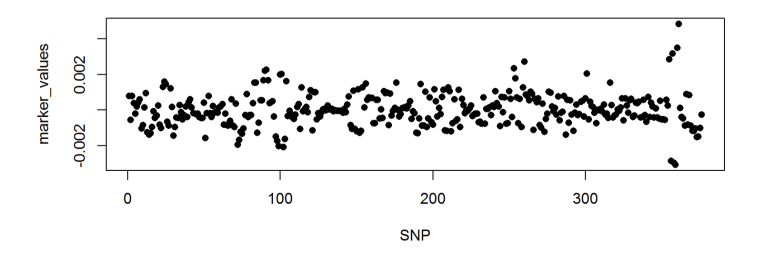
Example 4 - Soybeans

snp-BLUP

```
data(tpod,package='NAM')
X = matrix(1,length(y),1)
Z = gen
dim(Z)
## [1] 196 376
```

Example 4 - Fit the model

```
# Fit using the Lme4 package
fit0 = reml(y=y,X=X,Z=Z) # same as reml(y=y,Z=gen)
marker_values = fit0$EBV
gebv0 = c(gen %*% marker_values)
# Marker effects
plot(marker_values,pch=16, xlab='SNP')
```

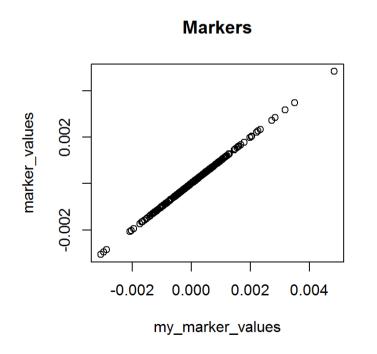


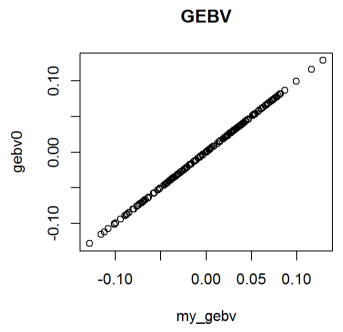
Example 4 - DIY BLUPs

```
iK = diag(ncol(Z))
Lambda = fit0$VC[2] / fit0$VC[1]
W = cbind(X,Z)
Sigma = diag( c(0,rep(Lambda,ncol(Z))) )
LHS = crossprod(W) + Sigma
RHS = crossprod(W,y)
g = solve(LHS,RHS)
my_mu = g[ c(1:ncol(X))]
my_marker_values = g[-c(1:ncol(X))]
my_gebv = c(gen %*% my_marker_values) # GEBVs from RR
```

Example 4 - DIY BLUPs

```
par(mfrow=c(1,2))
plot(my_marker_values, marker_values, main="Markers")
plot(my_gebv, gebv0, main="GEBV")
```





Example 4 - Heritability from RR

```
fit0$VC
                 Ve
              Vg
                                     h2
## 1 1.659819e-05 0.03167014 0.0005238214
Scale=sum(apply(gen,2,var)); Va=fit0$VC[1]*Scale; Ve=fit0$VC[2]
round((Va/(Va+Ve)),2)
      Vg
## 1 0.16
K = tcrossprod(apply(gen,2,function(x) x-mean(X)))
K = K/mean(diag(K)); round(reml(y,K=K)$VC,2)
      Vg
          Ve h2
## 1 0.01 0.03 0.16
```

Estimate VC from bad starters

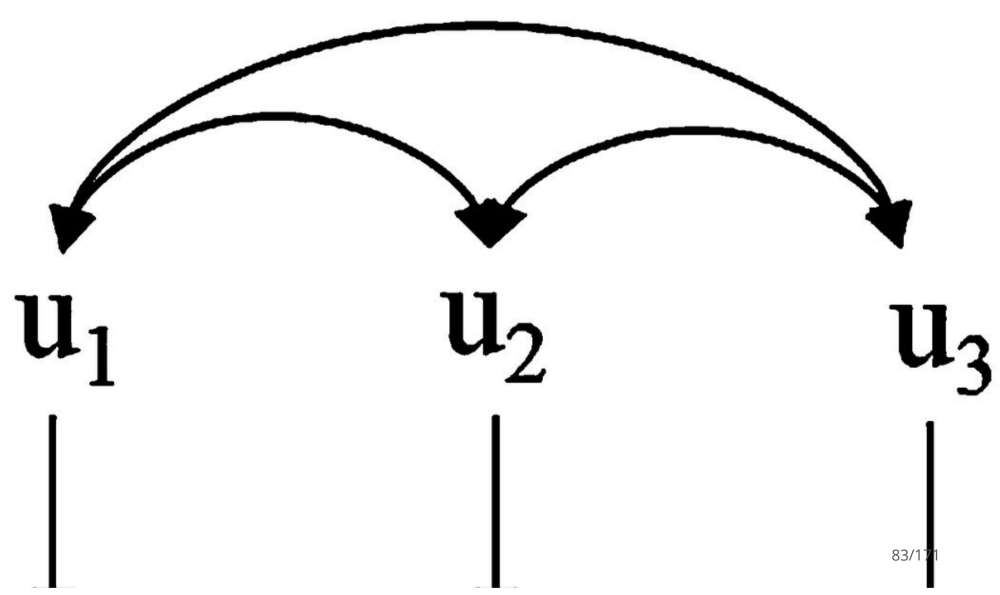
```
W = cbind(X,Z); iK = diag(ncol(Z))
Ve = Vg = 1 # Bad starting values
for(i in 1:100){ # Check the VC convergence after few iterations
 Lambda = Ve/Vg;
  Sigma = as.matrix(Matrix::bdiag(diag(0,ncol(X)),iK*Lambda))
  LHS = crossprod(W) + Sigma; RHS = crossprod(W,y); g = solve(LHS,RHS)
  my blue = g[c(1:ncol(X))]; my blup = g[-c(1:ncol(X))]
  e = y - X%*my blue - Z%*my blup; Ve = c(y)*%e)/(length(y)-ncol(X))
 trKC22 = sum(diag(iK%*%(solve(LHS)[(ncol(X)+1):(ncol(W)),(ncol(X)+1):(ncol(W))])))
 Vg = c(t(my blup)%*%iK%*%my blup+trKC22*Ve)/ncol(Z)
  if(!i%25){cat('It',i,'VC: Vg =',Vg,'and Ve =',Ve,'\n')}}
## It 25 VC: Vg = 0.0002960602 and Ve = 0.01801226
## It 50 VC: Vg = 7.428217e-05 and Ve = 0.02531553
## It 75 VC: Vg = 4.000965e-05 and Ve = 0.02818575
## It 100 VC: Vg = 2.887763e-05 and Ve = 0.02956763
```

Break

Module 3 - Advanced topics

Outline

- · Multivariate models
- Bayesian methods
- Machine learning
- · G x E interactions



Mixed models also enable us to evaluate multiple traits:

- More accurate parameters: BV and variance components
- Information: Inform how traits relate to each other
- Constrains: May increase computation time considerably

It preserves the same formulation

$$y = Xb + Zu + e$$

However, we now stack the traits together:

$$y=\{y_1,y_2,\ldots,y_k\}$$
 , $X=\{X_1,X_2,\ldots,X_k\}'$, $b=\{b_1,b_2,\ldots,b_k\}$, $Z=\{Z_1,Z_2,\ldots,Z_k\}'$, $u=\{u_1,u_2,\ldots,u_k\}$, $e=\{e_1,e_2,\ldots,e_k\}$.

The multivariate variance looks nice at first

$$Var(y) = Var(u) + Var(e)$$

But can get ugly with a closer look:

$$Var(u)=Z(G\otimes \Sigma_a)Z'=egin{bmatrix} Z_1GZ_1'\sigma_{a_1}^2 & Z_1GZ_2'\sigma_{a_{12}}\ Z_2GZ_1'\sigma_{a_{21}} & Z_2GZ_2'\sigma_{a_2}^2 \end{bmatrix}$$

and

$$Var(e) = R \otimes \Sigma_e = egin{bmatrix} R\sigma_{e_1}^2 & R\sigma_{e_1e_2} \ R\sigma_{e_2e_1} & R\sigma_{e_2}^2 \end{bmatrix}$$

You can still think the multivariate mixed model as

$$y = Wg + e$$

Where

$$y=egin{bmatrix} y_1\ y_2 \end{bmatrix}, W=egin{bmatrix} X_1 & 0 & Z_1 & 0\ 0 & X_2 & 0 & Z_2 \end{bmatrix}, g=egin{bmatrix} b_1\ b_2\ u_1\ u_2 \end{bmatrix}, e=egin{bmatrix} e_1\ e_2 \end{bmatrix}$$

Left-hand side $(W'R^{-1}W + \Sigma)$

$$\begin{bmatrix} X_1'X_1\Sigma_{e_{11}}^{-1} & X_1'X_2\Sigma_{e_{12}}^{-1} & X_1'Z_1\Sigma_{e_{11}}^{-1} & X_1'Z_2\Sigma_{e_{12}}^{-1} \\ X_2'X_1\Sigma_{e_{12}}^{-1} & X_2'X_2\Sigma_{e_{22}}^{-1} & X_2'Z_1\Sigma_{e_{12}}^{-1} & X_2'Z_2\Sigma_{e_{22}}^{-1} \\ Z_1'X_1\Sigma_{e_{11}}^{-1} & Z_1'X_2\Sigma_{e_{12}}^{-1} & G^{-1}\Sigma_{a_{11}}^{-1} + Z_1'Z_1\Sigma_{e_{11}}^{-1} & G^{-1}\Sigma_{a_{12}}^{-1} + Z_1'Z_2\Sigma_{e_{12}}^{-1} \\ Z_2'X_1\Sigma_{e_{12}}^{-1} & Z_2'X_2\Sigma_{e_{22}}^{-1} & G^{-1}\Sigma_{a_{11}}^{-1} + Z_2'Z_1\Sigma_{e_{12}}^{-1} & G^{-1}\Sigma_{a_{22}}^{-1} + Z_2'Z_2\Sigma_{e_{22}}^{-1} \end{bmatrix}$$

Right-hand side $(W'R^{-1}y)$

$$egin{bmatrix} X_1'(y_1\Sigma_{e_1}^{-1}+y_2\Sigma_{e_12}^{-1})\ X_2'(y_1\Sigma_{e_22}^{-1}+y_2\Sigma_{e_12}^{-1})\ Z_1'(y_1\Sigma_{e_12}^{-1}+y_2\Sigma_{e_1}^{-1})\ Z_2'(y_1\Sigma_{e_12}^{-1}+y_2\Sigma_{e_12}^{-1}) \end{bmatrix}$$

```
data(wheat, package = 'BGLR')
G = NAM::GRM(wheat.X);
Y = \text{wheat.Y}; \text{colnames}(Y) = c('E1', 'E2', 'E3', 'E4')
mmm = NAM::reml(y = Y, K = G)
knitr::kable( round(mmm$VC$GenCor,2) )
                                   E1
                                                           E2
                                                                                                          E4
                                                                                   E3
 E1
                                  1.00
                                                         -0.25
                                                                                -0.22
                                                                                                        -0.50
 E2
                                 -0.25
                                                         1.00
                                                                                 0.96
                                                                                                         0.55
 E3
                                 -0.22
                                                         0.96
                                                                                                         0.72
                                                                                 1.00
 E4
                                 -0.50
                                                         0.55
                                                                                 0.72
                                                                                                         1.00
```

mmm\$VC\$Vg

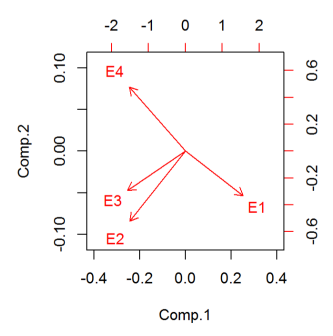
```
## E1 0.6277835 -0.1446924 -0.1102175 -0.2743640
## E2 -0.1446924 0.5440731 0.4419945 0.2822577
## E3 -0.1102175 0.4419945 0.3919626 0.3130735
## E4 -0.2743640 0.2822577 0.3130735 0.4828705
```

mmm\$VC\$Ve

```
## E1 0.53504246 0.08247812 -0.1159118 0.06882868
## E2 0.08247812 0.56214755 0.2973841 0.15795801
## E3 -0.11591175 0.29738408 0.6714234 0.11086214
## E4 0.06882868 0.15795801 0.1108621 0.59405228
```

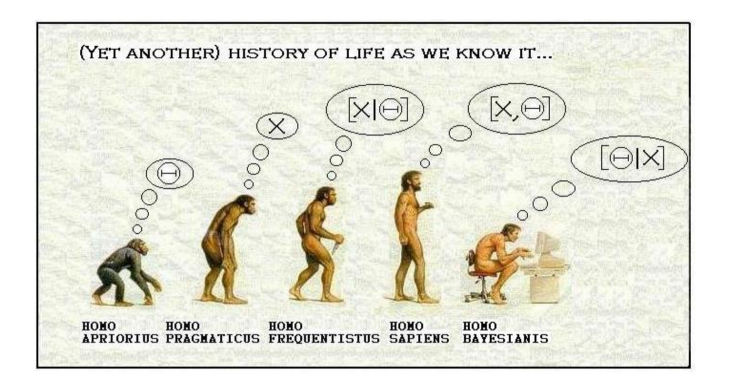
- · Selection indeces, co-heritability, indirect response to selection
- · Study residual and additive genetic association among traits

biplot(princomp(mmm\$VC\$GenCor,cor=T),xlim=c(-.4,.4),ylim=c(-.11,.11))



When do additional traits contribute?

```
# Fit E4, predict E2
fit E4=bWGR::mrr(matrix(Y[,4]),wheat.X); cor(fit E4$hat[,1],Y[,2])
## [1] 0.3988844
# Fit E4 and E1, E4 predict E2
fit E4E1=bWGR::mrr(Y[,c(1,4)],wheat.X); cor(fit E4E1$hat[,2],Y[,2])
## [1] 0.3931193
# Fit E4 and E3, E4 predict E2
fit_E4E3=bWGR::mrr(Y[,3:4],wheat.X); cor(fit_E4E3$hat[,2], Y[,2])
## [1] 0.5279279
```



The general framework on a hierarchical Bayesian model follows:

$$p(\theta|x) \propto p(x|\theta)p(\theta)$$

Where:

- Posterior probability: $p(\theta|x)$
- · Likelihood: $p(x|\theta)$
- Prior probability: $p(\theta)$

For the model:

$$y=Xb+Zu+e,~~u\sim N(0,K\sigma_a^2),~e\sim N(0,I\sigma_e^2)$$

- Data ($x = \{y, X, Z, K\}$)
- Parameters ($\theta = \{b, u, \sigma_a^2, \sigma_e^2\}$)

Probabilistic model:

$$egin{aligned} p(b,u,\sigma_a^2,\sigma_e^2|y,X,Z,K) &\propto N(y,X,Z,K|b,u,\sigma_a^2,\sigma_e^2) imes \ N(b,u|\sigma_a^2,\sigma_e^2) imes \chi^{-2}(\sigma_a^2,\sigma_e^2|S_a,S_e,
u_a,
u_e) \end{aligned}$$

REML: the priors (S_a, S_e, ν_a, ν_e) are estimated from data.

Hierarchical Bayes: You provide priors. Here is how:

$$\sigma_a^2 = rac{u'K^{-1}u + S_a
u_a}{\chi^2(q+
u_a)}$$

sigma2a=(t(u)%*%iK%*%u+Sa*dfa)/rchisq(df=ncol(Z)+dfa,n=1)

$$\sigma_e^2 = rac{e'e + S_e
u_e}{\chi^2 (n +
u_e)}$$

sigma2e=(t(e)%*%e+Se*dfe)/rchisq(df=length(y)+dfe,n=1)

What does it mean for **you**? If your "prior knowledge" tells you that a given trait has approximately $h^2=0.5$ (nothing unreasonable). In which case, half of the phenotypic variance is due to genetics, and the other half is due to error. Your prior shape is:

$$S_a = S_e = \sigma_y^2 imes 0.5$$

We usually assign small a prior degrees of freeds. Samething like four or five prior degrees of freedom. That means that assuming $\nu_0=5$, you are yielding to your model 5 data points that support heritability 0.5

$$\nu_a = \nu_e = 5$$

Example of prior influence: In a dataset with 300 data points, 1.6% of the variance components information comes from prior (5/305), and 98.4% comes from data (300/305).

For whole-genome regression models

$$y=\mu+Ma+e,~~a\sim N(0,I\sigma_b^2),~e\sim N(0,I\sigma_e^2)$$

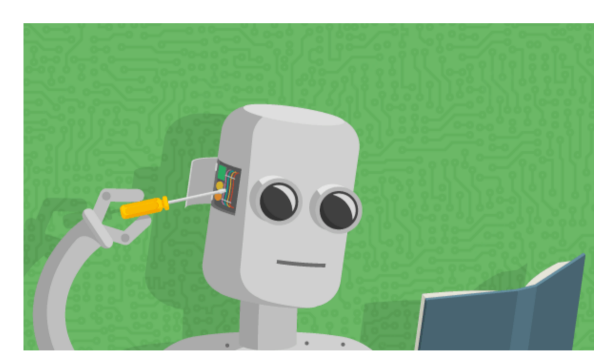
We scale the prior genetic variance based on allele frequencies

$$S_b = rac{\sigma_y^2 imes 0.5}{2 \sum p_j (1-p_j)}$$

Two common settings:

- · All markers, one random effect:
- · Each markers as a random effect:

- Parametric methods for prediction: L1-L2
- · Semi-parametric methods for prediction: Kernels
- · Non-parametric methods for prediction: Trees and nets



L1-L2 machines include all mixed and Bayesian models we have seen so far. The basic framework is driven by a single (random) term model:

$$y = Xb + e$$

The univariate soltion indicates how the model is solved. A model without regularization yields the least square (LS) solution. If we regularize by deflating the nominator, we get the L1 regularization (LASSO). If we regularize by inflating the denominator, we get the L2 regularization (Ridge). For any combination of both, we get a elastic-net (EN). Thus:

$$b_{LS} = rac{x'y}{x'x}, ~~ b_{Lasso} = rac{x'y-\lambda}{x'x}, ~~ b_{Ridge} = rac{x'y}{x'x+\lambda}, ~~ b_{EN} = rac{x'y-\lambda_1}{x'x+\lambda_2}$$

Whereas the Bayesian and mixed model framework resolves the regularization as $\lambda = \sigma_e^2/\sigma_b^2$, ML methods search for λ through (-fold) cross-validation.

Common loss functions in L1-L2 machines

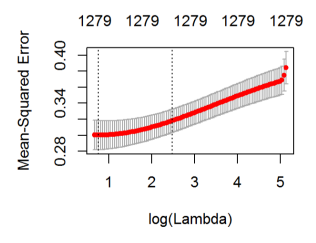
- LS (no prior, no shrinkage): $argmin(\sum e_i^2)$
- L1 (Laplace prior with variable selection): $argmin(\sum e_i^2 + \lambda \sum |b_j|)$
- · L2 (Gaussian prior, unique solution): $argmin(\sum e_i^2 + \lambda \sum b_j^2)$

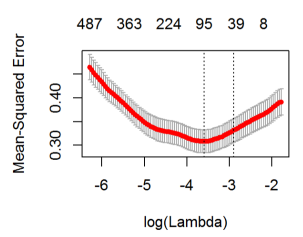
Other losses that are less popular

- Least absolute: $argmin(\sum |e_i|)$ based on $b_{LA} = rac{nMD(x imes y)}{x'x}$
- ϵ -loss: $argmin(\sum e_i^2, |e_i| > \epsilon)$ used in support vector machines

Cross-validations to search for best value of lambda

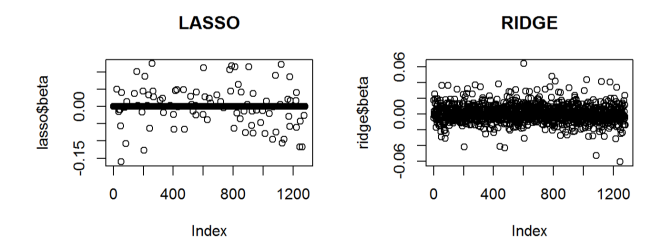
```
lasso = glmnet::cv.glmnet(x=wheat.X,y=rowMeans(Y),alpha=1);
ridge = glmnet::cv.glmnet(x=wheat.X,y=rowMeans(Y),alpha=0);
par(mfrow=c(1,2)); plot(ridge); plot(lasso)
```





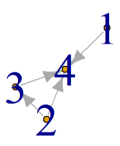
Re-fit the model using this best value

```
lasso = glmnet::glmnet(x=wheat.X,y=rowMeans(Y),lambda=lasso$lambda.min,alpha=1)
ridge = glmnet::glmnet(x=wheat.X,y=rowMeans(Y),lambda=ridge$lambda.min,alpha=0)
par(mfrow=c(1,2)); plot(lasso$beta,main='LASSO'); plot(ridge$beta,main='RIDGE');
```



Of course, the losses presented above are not limited to the application of prediction and classification. Below, we see an example of deploying LASSO for a graphical model (Markov Random Field): How the traits of the multivariate model relate in terms of additive genetics:

```
ADJ=huge::huge(mmm$VC$GenCor,.3,method='glasso',verbose=F)$path[[1]] plot(igraph::graph.adjacency(adjmatrix=ADJ),vertex.label.cex=3)
```



Reproducing kernel Hilbert Spaces (RKHS), is a generalization of a GBLUP... Most commonly instead of using the linear kernel ($ZZ'\alpha$), RKHS commonly uses one or more Gaussian or exponential kernels:

$$K = \exp(-\theta D^2)$$

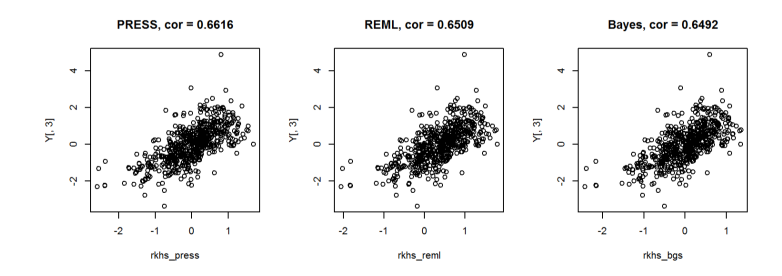
Where D^2 is the squared Euclidean distance, and θ is a bandwidth:

- Single kernel: $1/mean(D^2)$
- Three kernels: θ ={5/q, 1/q, 0.2/q}, where q=quantile(D2,0.05)

We can use REML, PRESS (=cross-validation) or Bayesian approach to solve RKHS

```
# Make the kernel.
D2 = as.matrix(dist(wheat.X)^2)
K = \exp(-D2/mean(D2))
# Below we are going to calibrate models on Env 2 and predict Env 3
rkhs_press = NAM::press(y=Y[,2],K=K)$hat
rkhs_reml = NAM::reml(y=Y[,2],K=K)$EBV
rkhs bgs = NAM::gibbs(y=Y[,2],iK=solve(K))$Fit.mean
##
                                                                          0%
                                                                          1%
                                                                          1%
                                                                          2%
                                                                                              105/171
```

```
par(mfrow=c(1,3))
plot(rkhs_press,Y[,3],main=paste('PRESS, cor =',round(cor(rkhs_press,Y[,3]),4) ))
plot(rkhs_reml,Y[,3],main=paste('REML, cor =',round(cor(rkhs_reml,Y[,3]),4) ))
plot(rkhs_bgs,Y[,3],main=paste('Bayes, cor =',round(cor(rkhs_bgs,Y[,3]),4) ))
```



RKHS for epistasis and variance component analysis

Epistasis



For the same task (E2 predict E3), let's check members of the Bayesian alphabet

```
fit_BRR = bWGR::wgr(Y[,2],wheat.X); cor(c(fit_BRR$hat),Y[,3])

## [1] 0.5842116

fit_BayesB = bWGR::BayesB(Y[,2],wheat.X); cor(fit_BayesB$hat,Y[,3])

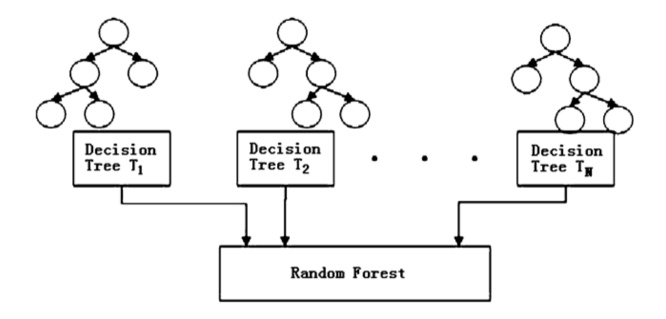
## [1] 0.5355413

fit_emBayesA = bWGR::emBA(Y[,2],wheat.X); cor(fit_emBayesA$hat,Y[,3])

## [1] 0.6388318
```

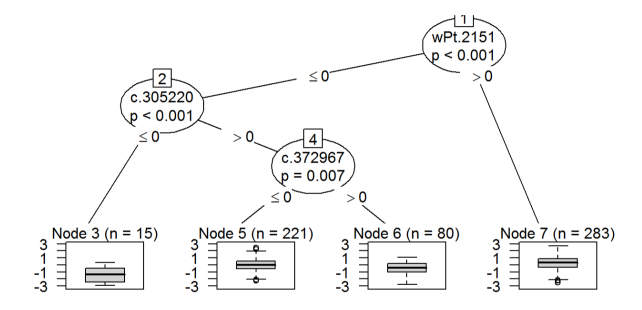
Machine learning methods

Tree regression and classifiers



Machine learning methods

fit_tree = party::ctree(y~.,data.frame(y=Y[,2],wheat.X)); plot(fit_tree)

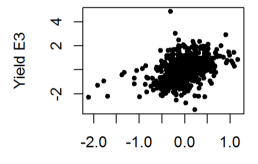


cor(c(fit_tree@predict_response()),Y[,3])

[1] 0.265622

Machine learning methods

```
fit_rf = ranger::ranger(y~.,data.frame(y=Y[,2],wheat.X))
plot(fit_rf$predictions,Y[,3],xlab='RF predictions from E2',ylab='Yield E3',pch=20)
```



RF predictions from E2

```
cor(fit_rf$predictions,Y[,3])
```

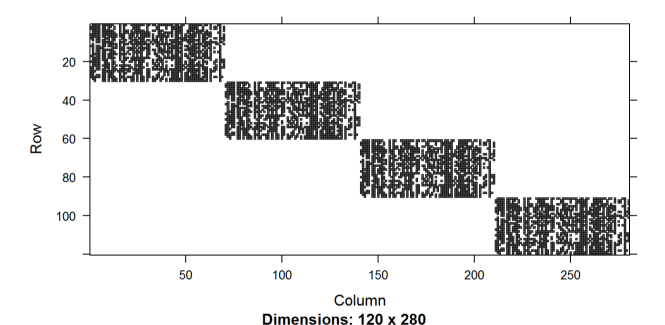
[1] 0.4056496



```
y=as.vector(wheat.Y); Z=wheat.X; Zge=as.matrix(Matrix::bdiag(Z,Z,Z,Z))
fit g = bWGR::BayesRR(rowMeans(wheat.Y),Z)
fit_ge = bWGR::BayesRR(y,Zge)
fit gge = bWGR::BayesRR2(y,rbind(Z,Z,Z,Z),Zge)
#
fit g$h2
## [1] 0.4590341
fit ge$h2
## [1] 0.6762359
fit gge$h2
## [1] 0.6580924
```

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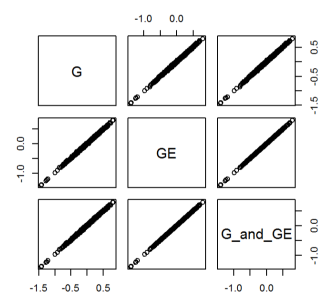
GxE design matrix: Example of 4 environments, 30 individuals, 70 SNPs



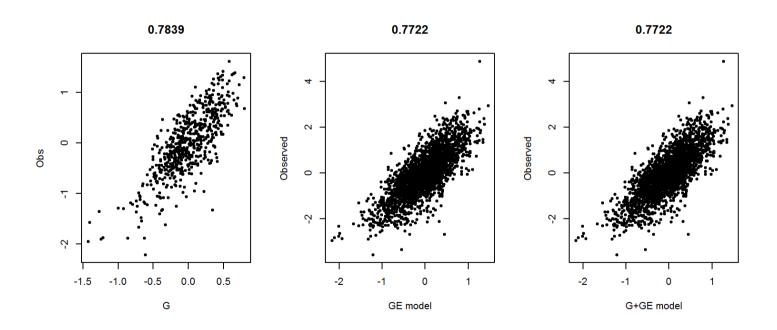
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```
GE1=matrix(fit_ge$hat,ncol=4); GE2=matrix(fit_ge$hat,ncol=4)
plot(data.frame(G=fit_g$hat,GE=rowMeans(GE1),G_and_GE=rowMeans(GE2)),main='GEBV across E')
```

GEBV across E



```
par(mfrow=c(1,3))
plot(fit_g$hat,rowMeans(Y),main=round(cor(fit_g$hat,rowMeans(Y)),4),xlab='G',ylab='Obs',pch=20)
plot(c(GE1),y,main=round(cor(c(GE1),y),4),xlab='GE model',ylab='Observed',pch=20)
plot(c(GE2),y,main=round(cor(c(GE2),y),4),xlab='G+GE model',ylab='Observed',pch=20)
```



Break

Module 4 - Signal detection

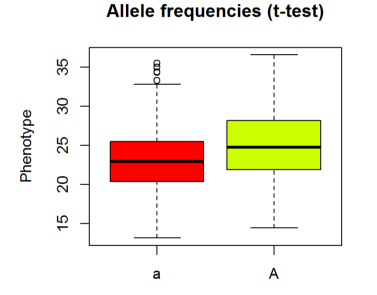
Outline

- Test statistics
- · Allele coding
- · Power & resolution
- Linkage mapping
- · LD mapping
- Structure
- Imputation

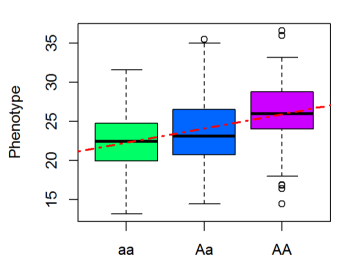
- · GLM
- · MLM
- · WGR
- · Rare-variants
- · Validation studies

Test statistics

Testing associations are as simple as t-test and ANOVA



Genotypic frequencies (ANCOVA)



Test statistics

· A more generalized framework: Likelihood test

$$LRT = L_0/L_1 = -2(logL_1 - logL_0)$$

For the model:

$$y = Xb + Zu + e$$
 $y \sim N(Xb, V)$

REML function is given by:

$$L(\sigma_u^2,\sigma_e^2) = -0.5(ln|V| + ln|X'V^{-1}X| + y'Py)$$

Where $V=ZKZ'\sigma_u^2+I\sigma_e^2$ and y'Py=y'e

Allele coding

Types of allele coding

- 1. Add. (1 df): {-1,0,1} or {0,1,2} **Very popular** (Lines, GCA)
- 2. Dom. (1 df): {0,1,0} Popular (Trees, clonals and Hybrids)
- 3. Jointly A+D (2 df): Popular on QTL mapping in F2s
- 4. Complete dominance (1 df): {0,0,1} or {0,1,1} Very unusual
- 5. Interactions (X df): (epistasis and GxE)

Power and resolution

Power

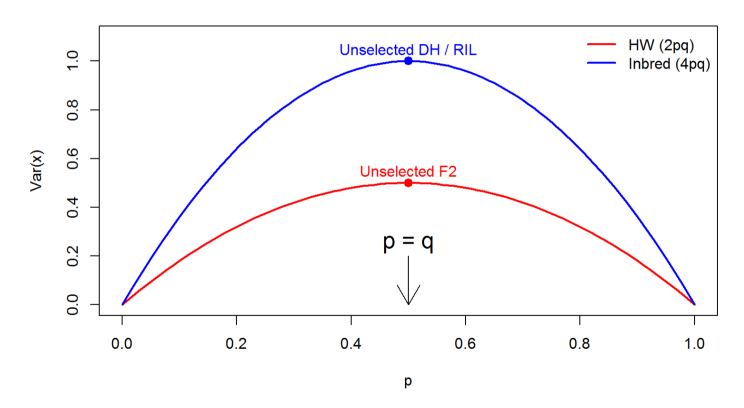
- Key: Number of individuals & allele frequency
- More DF = lower power
- Multiple testing: Bonferroni and FDR
- Tradeoff: Power vs false positives

Resolution

- Genotyping density
- LD blocks
- Recombination

Power: Variance of X

Marker variance



Beavis effect: 1000 is just OK

Xu S. Theoretical basis of the Beavis effect. Genetics. 2003 1;165(4):2259-68.

TABLE 4 Comparisons of predicted and observed (estimated) biases in estimated QTL effects and variances from Beavis F_2 simulation experiments

Simulated conditions ^a	Variance explained			Additive effect			Average estimate
	Simulated	Observed	Predicted ^b	Simulated	Observed	Predicted	Average estimated location
10-30-100	3.00	16.76	16.0537	2.45	4.96	5.6410	11.3
10-30-500	3.00	4.33	4.1890	2.45	2.89	2.8617	10.53
10-30-1000	3.00	3.02	3.1846	2.45	2.56	2.4868	10.8
10-63-100	6.25	12.65	16.5984	3.55	4.68	5.7328	10.51
10-63-500	6.25	7.08	6.5581	3.55	3.73	3.5829	10.96
10-63-1000	6.25	6.34	6.3566	3.55	3.60	3.5500	11.04
10-95-100	9.50	18.68	17.3883	4.36	5.85	5.8466	10.58
10-95-500	9.50	10.1	9.7082	4.36	4.49	4.3607	11.08
10-95-1000	9.50	9.67	9.6028	4.36	4.44	4.3600	11.19
40-30-100	0.75	15.78	15.6270	1.22	4.40	5.5436	10.83
40-30-500	0.75	3.17	3.3332	1.22	2.35	2.5671	10.17
40-30-1000	0.75	1.46	1.7961	1.22	1.85	1.8790	10.17
40-63-100	1.56	16.31	15.7983	1.77	4.71	5.5999	10.45
40-63-500	1.56	3.54	3.5783	1.77	2.59	2.6582	10.13
40-63-1000	1.56	1.96	2.1435	1.77	2.09	2.0494	10.37
40-95-100	2.40	16.55	15.9694	2.18	5.02	5.6236	10.45
40-95-500	2.40	3.97	3.9190	2.18	2.79	2.7641	10.12
40-95-1000	2.40	2.58	2.6970	2.18	2.36	2.2784	10.29

^a Numerical values denote the number of QTL-heritability-number of progeny.

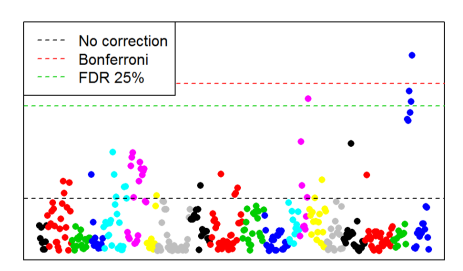
^b Using Equation 17.

^{&#}x27;Using Equation 8.

Multiple testing:

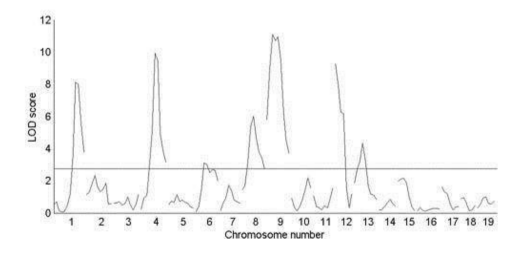
GWAS tests m hypothesis:

- No correction: $\alpha = 0.05$
- Bonferroni: $\alpha = 0.05/m$
- FDR (25%): lpha = 0.05/(m imes 0.75)



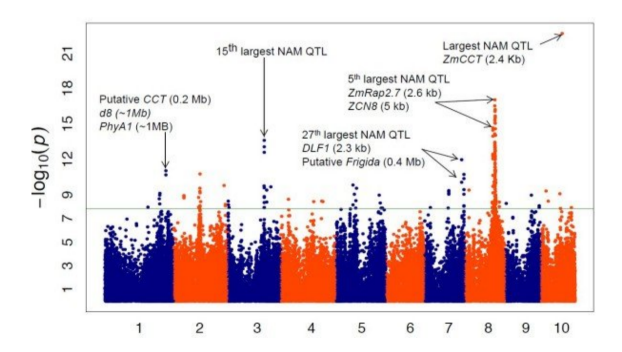
Linkage mapping

- Generally on experimental pops (F2, DH, RIL, BC)
- · Based on single-marker analysis or interval mapping
- · Recombination rates would increase power



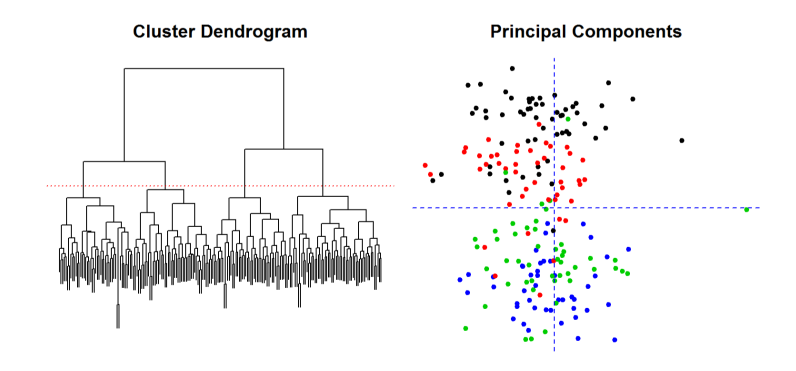
LD mapping (or association mapping)

- · Use of historical LD between marker and QTL
- AM allowed studies on random panels
- Dense SNP panels would increase resolution

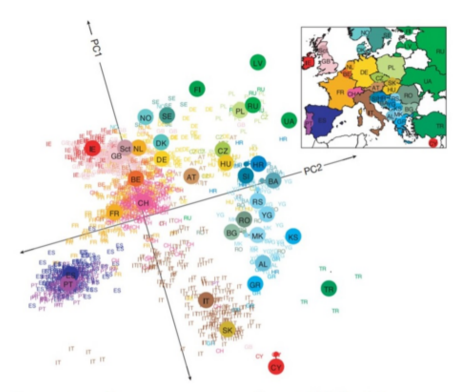


Structure

- 1. Confounding associations with sub-populations
- 2. Major limitation of association mapping
- 3. Structure: , , (eg. race)



Structure



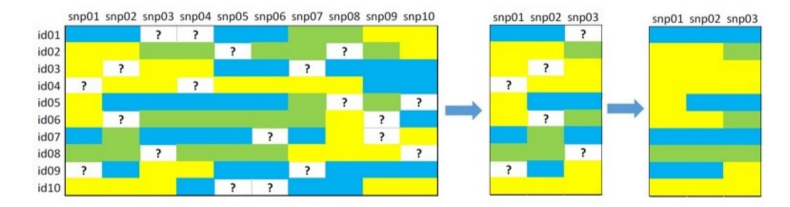
Genes mirror geography within Europe

nature Vol 456 6 November 2008 doi:10.1038/nature07331

Imputation

Less missing values = more obs. = more detection power

- Markov models: Based on flanking markers
- · Random forest: Multiple decision trees capture LD
- kNN & Projections: Fill with similar haplotypes



GLM (generalized linear models)

• Full model (L_1):

$$y = Xb + m_j a + e$$

• Null model (L_0):

$$y = Xb + e$$

- 1. Advantage: Fast, not restricted to Gaussian traits
- 2. Popular methodology on human genetic studies
- 3. Xb includes (1) environment, (2) structure and (3) covariates

MLM (mixed linear models)

• Full model (L_1):

$$y = Xb + Zu + m_j a + e$$

· Null model (L_0):

$$y = Xb + Zu + e$$

- 1. The famous "Q+K model"
- 2. Advantage: Better control of false positives, no need for PCs
- 3. Polygenic effect (u) assumes $u \sim N(0, K\sigma_u^2)$
- 4. Faster if we don't reestimate $\lambda = \sigma_e^2/\sigma_u^2$ for each SNP

cMLM (compressed MLM)

- 1. Uses the same base model as MLM
- 2. Advantage: Faster than MLM
- 3. Based on clustered individuals:
- $\cdot Z$ is indicates the subgroup
- \cdot K is the relationship among subgroup
- * Often needs PCs to complement K

WGR (whole-genome regression)

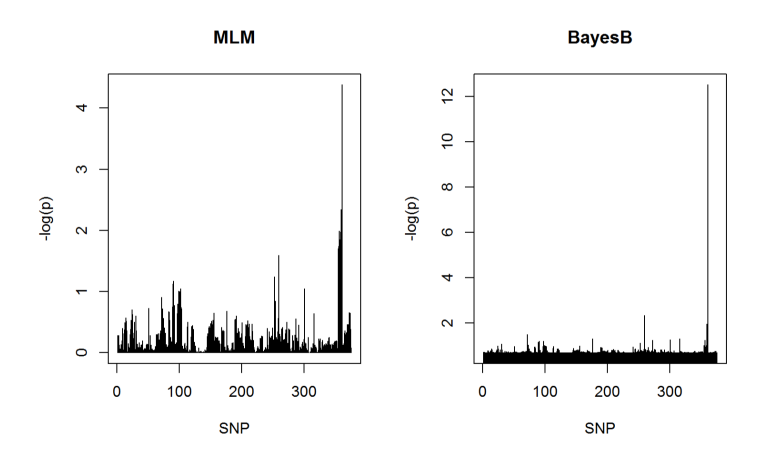
- 1. Tests all markers at once
- 2. Advantage: No double-fitting, no PCs, no Bonferroni
- Model (BayesB, BayesC, SSVS):

$$y = Xb + Ma + e$$

Marker effects are from a mixture of distributions

$$a_j \sim Binomial$$
 with $p(\pi) = 0$ and $p(1-\pi) = a_j$

WGR (whole-genome regression)



Rare variants

- 1. Screen a set (s) of low MAF markers on NGS data
- 2. Advantage: Detect signals from low power SNPs
- 3. Applied to uncommon diseases (not seen in plant breeding)
- 4. Two possible model
- Full model 1 (L_1): $y = Xb + M_sa + e$
- Full model 2 (L_2): $y = Xb + PC_1(M_s) + e$
- Null model (L_0): y = Xb + e

Test either $LR(L_1, L_0)$ or $LR(L_2, L_0)$

Validation studies

- · QTLs detected with 3 methods, across 3 mapping pops
- · Validations made on 3 unrelated populations

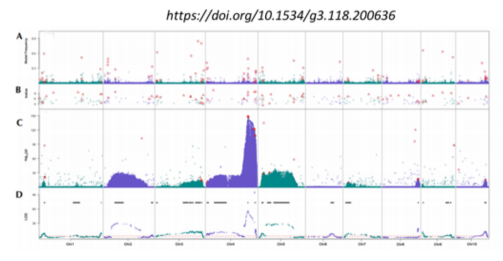


Figure 2 Stacked plots of GWAS and QTL results. From upper to lower panels are results from the Bayesian-based multi-variant (A) stepwise regression (B) and single variant(C) models for GWAS and the joint QTL mapping result (D). The red dashed line in the QTL plot indicates the 1,000 permutation threshold and black lines show the QTL confidence intervals. Red squares in panel (A), triangles in panel (B) and circles in panel (C) indicate the kernel row number associated variants selected for further genetic validation.

Break

Module 5 - Association analysis

Prelude: Data & Structure

Getting some data

Example dataset from the package. We are querying two of the forty biparental families with a shared parental IA3023, grown in 18 environment.

```
Data = SoyNAM::BLUP(trait = 'yield', family = 2:3)

### solving BLUE of checks

### solving BLUP of phenotypes

### No redundant SNPs found

### There are 312 markers with MAF below the threshold

### Removing markers with more than 50% missing values

### Imputing with expectation (based on transition prob)

### removing repeated genotypes

### solving identity matrix

### indiviual 1 had 37 duplicate(s)

### indiviual 169 had 1 duplicate(s)

### indiviual 182 had 1 duplicate(s)
```

Genomic relationship matrix

```
y = Data$Phen
M = Data$Gen
#
Z = apply(M,2,function(snp) snp-mean(snp))
ZZ = tcrossprod(Z)
Sum2pq = sum(apply(M,2,function(snp){p=mean(snp)/2; return(2*p*(1-p))}))
G = ZZ/Sum2pq
```

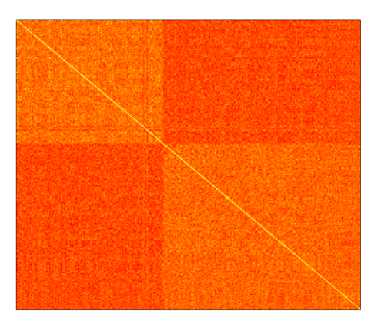
Kernel commonly deployed, referred in VanRaden (2008)

$$G = rac{(M-P)(M-p)'}{2\sum_{j=1}^{J} p_j (1-p_j)}$$

Genomic relationship matrix

image(G[,241:1], main='GRM heatmap',xaxt='n',yaxt='n')

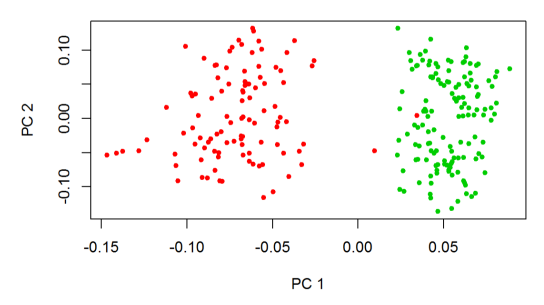
GRM heatmap



Structure parameters (1) PCs

```
Spectral = eigen(G,symmetric = TRUE)
PCs = Spectral$vectors[,1:5]
plot(PCs,xlab='PC 1',ylab='PC 2',main='Population on eigen spaces',col=Data$Fam,pch=20)
```

Population on eigen spaces



Structure parameters (2) Clusters

```
GeneticDistance = Gdist(M,method=6)

## Modified Rogers' distance

Tree = hclust(GeneticDistance,method = 'ward.D2')
Clst = factor(cutree(Tree,k=2))
plot(Tree,labels = FALSE)
```

Single marker analysis

GLM (1) - No structure

```
Marker = M[,117]
fit = lm(y \sim Marker)
anova( fit )
## Analysis of Variance Table
## Response: y
             Df Sum Sq Mean Sq F value Pr(>F)
## Marker 1 476321 476321 20.172 1.102e-05 ***
## Residuals 239 5643504 23613
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
-log(anova(fit)$`Pr(>F)`[1],base = 10)
## [1] 4.957736
```

GLM (2) - Principal Components

```
reduced model = lm(y \sim PCs)
full model = lm( y \sim PCs + Marker )
anova( reduced model, full model )
## Analysis of Variance Table
##
## Model 1: y ~ PCs
## Model 2: y ~ PCs + Marker
    Res.Df RSS Df Sum of Sq F Pr(>F)
## 1 235 4060362
## 2 234 3562067 1 498295 32.734 3.215e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
-log((anova( reduced model, full model ))$`Pr(>F)`[2],base = 10)
## [1] 7.492813
```

GLM (3) - Population Clusters

```
reduced model = lm(y \sim Clst)
full model = lm( y \sim Clst + Marker )
anova( reduced model, full model )
## Analysis of Variance Table
##
## Model 1: y ~ Clst
## Model 2: y ~ Clst + Marker
    Res.Df RSS Df Sum of Sq F Pr(>F)
## 1 239 4275698
## 2 238 3652041 1 623657 40.643 9.4e-10 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
-log( anova(reduced model, full model)$\Pr(>F)\[2], base = 10)
## [1] 9.026884
```

MLM - K+Q model

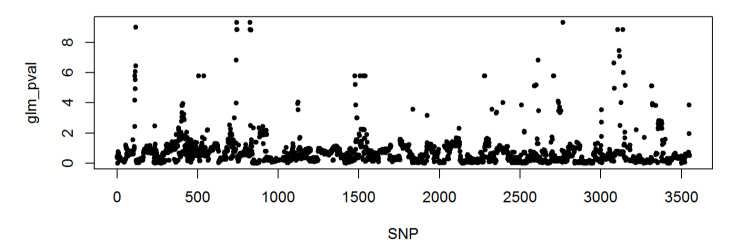
```
Q = model.matrix(~Clst)
reduced_model = reml( y=y, X=Q, K=G)
full_model = reml( y=y, X=cbind(Q, Marker), K=G)
LRT = -2*(full_model$loglik - reduced_model$loglik)
-log(pchisq(LRT,1,lower.tail=FALSE),base=10)
## [1] 10.80903
```

Whole genome screening

DIY (example with GLM)

```
reduced_model = lm( y ~ Clst )
glm_pval = apply(M,2,function(Marker){
  pval = anova(reduced_model, lm(y~Clst+Marker) )$`Pr(>F)`[2]
  return(-log(pval,base = 10))})
plot(glm_pval,pch=20,xlab='SNP',main='My first GLM GWAS')
```

My first GLM GWAS



Using CRAN implementations

```
NAM random model: y = \mu + Marker \times Pop + Zu + e
fit gwa = gwas3(y=y, gen=M, fam=c(Clst), chr=Data$Chrom)
## Calculating G matrix
## Solving polygenic model
## Starting Eigendecomposition
## Starting Marker Analysis
##
                                                                      0%
                                                                      1%
                                                                      1%
                                                                      2%
```

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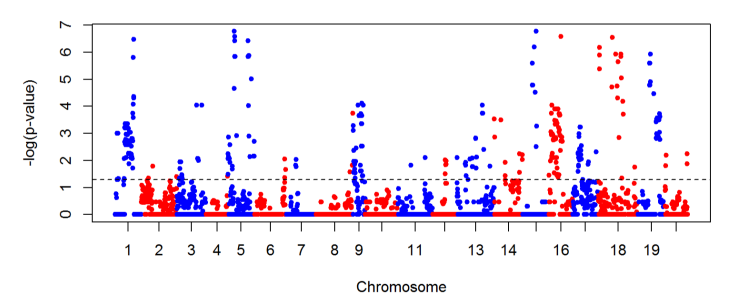
2%

==

Manhattan plot

plot(fit_gwa, pch=20, main = "My first MLM GWAS")

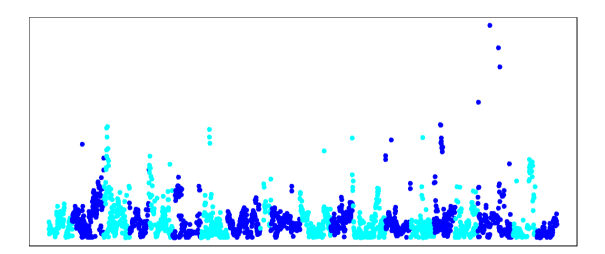
My first MLM GWAS



Yet another R implementations

```
require(rrBLUP,quietly = TRUE); COL = fit_gwa$MAP[,1]%2+1 # Color chromosomes
geno=data.frame(colnames(M),fit_gwa$MAP[,1:2],t(M-1),row.names=NULL)
pheno=data.frame(line=colnames(geno)[-c(1:3)],Pheno=y,Clst,row.names=NULL)
fit_another_gwa=GWAS(pheno,geno,fixed='Clst',plot=FALSE)
```

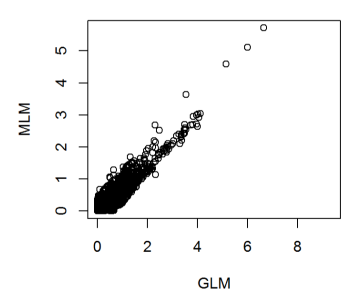
```
## [1] "GWAS for trait: Pheno"
## [1] "Variance components estimated. Testing markers."
```



Comparing results

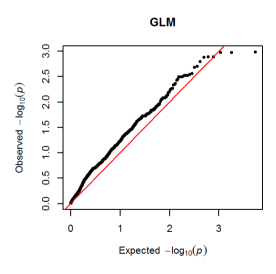
mlm_pval=fit_another_gwa\$Pheno; mlm_pval[mlm_pval==0]=NA
plot(glm_pval,mlm_pval,xlab='GLM',ylab='MLM',main='Compare')

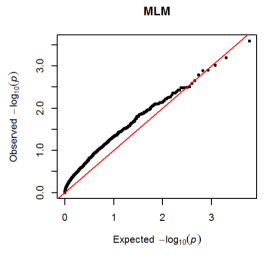
Compare

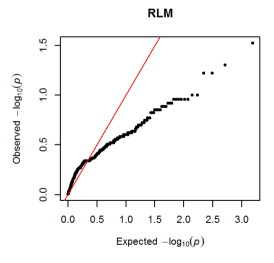


Power analysis - QQ plot

```
nam_pval = fit_gwa$PolyTest$pval
par(mfrow=c(1,3))
qqman::qq(glm_pval,main='GLM')
qqman::qq(mlm_pval,main='MLM')
qqman::qq(nam_pval,main='RLM')
```







Multiple testing

Multiple testing

In statistics, the multiple comparisons, multiplicity or multiple testing problem occurs when one considers a set of statistical inferences simultaneously or infers a subset of parameters selected based on the observed values. In certain fields it is known as the look-elsewhere effect:

Several

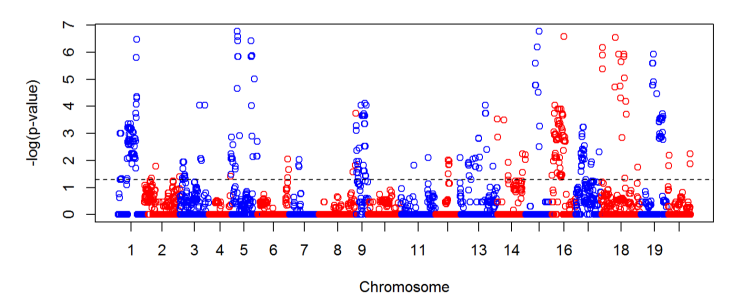
statistical techniques have been developed to prevent this from happening, allowing significance levels for single and multiple comparisons to be directly compared. These techniques generally require a **stricter significance threshold** for individual comparisons, so as to compensate for the number of inferences being made.

Baseline - No correction

Base significance threshold: lpha=0.05

plot(fit_gwa, alpha=0.05, main = "No correction")

No correction

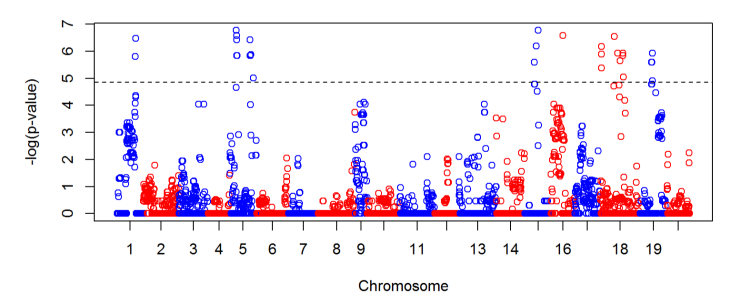


Multiple testing correction

Bonferroni: $\alpha=0.05/m$

plot(fit_gwa, alpha=0.05/ncol(M), main = "Bonferroni correction")

Bonferroni correction

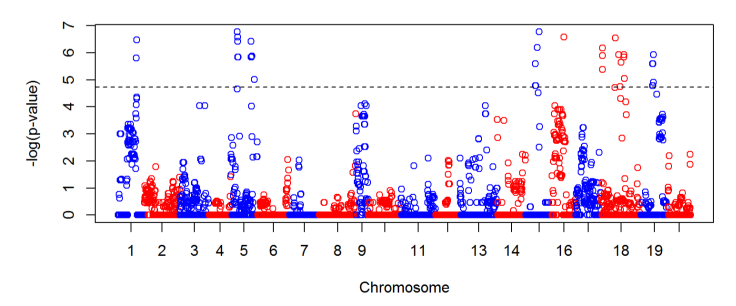


False-Discovery Rate

Benjamini-Hochberg FDR:
$$lpha = rac{0.05}{m imes (1-FDR)}$$

plot(fit_gwa, alpha=0.05/(ncol(M)*.75), main = "FDR 25%")

FDR 25%

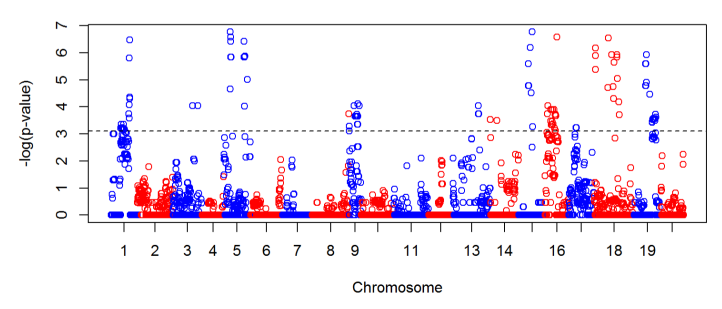


False-Discovery Rate

Unique segments based on Eigenvalues: $m^* = D > 1$

```
m_star = sum(Spectral$values>1)
plot(fit_gwa, alpha=0.05/m_star, main="Bonferroni on unique segments")
```

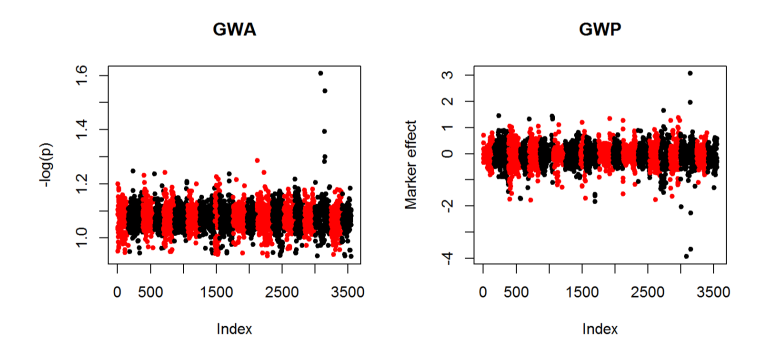
Bonferroni on unique segments



Multi-loci analysis

Whole genome regression

```
fit_wgr = bWGR::BayesDpi(y=y,X=M,it=3000); par(mfrow=c(1,2));
plot(fit_wgr$PVAL,col=COL,pch=20,ylab='-log(p)',main='GWA')
plot(fit_wgr$b,col=COL,pch=20,ylab='Marker effect',main='GWP')
```

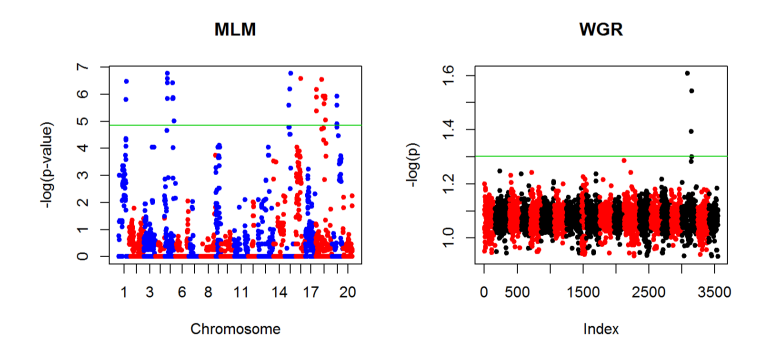


plot(fit_wgr\$hat,y,pch=20)

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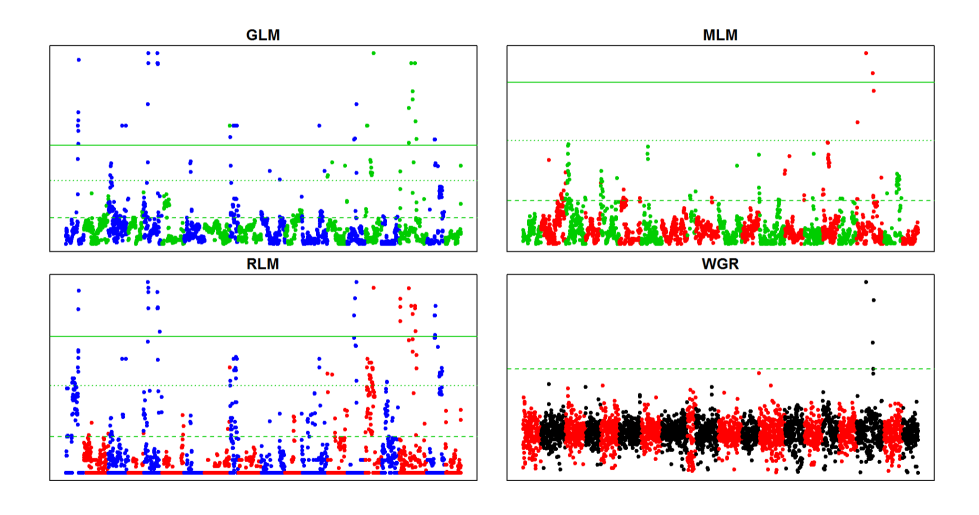
WGR - No need for multiple testing

```
thr_none = -log(pchisq(qchisq(1-0.05/ncol(M),1),1,lower.tail=FALSE),base=10)
thr_bonf = -log(pchisq(qchisq(1-0.05,1),1,lower.tail=FALSE),base=10)
par(mfrow=c(1,2)); plot(fit_gwa,alpha=NULL,main="MLM",pch=20); abline(h=thr_none,col=3)
plot(fit_wgr$PVAL,col=COL,ylab='-log(p)',main="WGR",pch=20); abline(h=thr_bonf,col=3)
```



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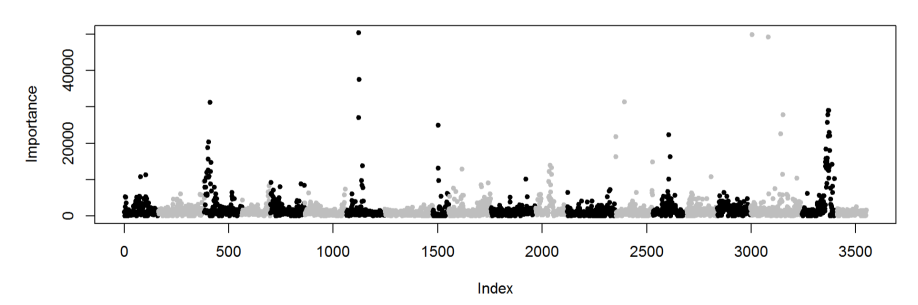
Approaches are complementary



Random forest

fit_rf = ranger::ranger(y~.,data= data.frame(y=y,M),importance='impurity')
plot(fit_rf\$variable.importance,ylab='Importance',main='Random Forest',col=COL+7,pch=20)

Random Forest



Thanks!

Thanks!

- e-mail: xaviera@purdue.edu or alencar.xavier@corteva.com
- other resources: https://alenxav.wixsite.com/home
- material: https://github.com/alenxav/Lectures/tree/master/Purdue_MLM

Break