

Genetic analysis using the sommer package

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The sommer package has been developed to provide R users with a powerful mixed model solver for different genetic and non-genetic analysis in diploid and polyploid organisms. This package allows the user to estimate variance components for a mixed model with the advantage of specifying the variance-covariance structure of the random effects and obtain other parameters such as BLUPs, BLUEs, residuals, fitted values, variances for fixed and random effects, etc.

The package is focused on genomic prediction (or genomic selection) and GWAS analysis, although general mixed models can be fitted as well. The package provides kernels to estimate additive (**A.mat**), dominance (**D.mat**), and epistatic (**E.mat**) relationship matrices that have been shown to increase prediction accuracy under certain scenarios. The package provides flexibility to fit other genetic models such as full and half diallel models as well.

Vignettes aim to provide several examples in how to use the sommer package under different scenarios in breeding and genetics. We will spend the rest of the space providing examples for:

- 1) heritability (h^2) calculation
- 2) Half and full diallel designs
- 3) Genome wide association analysis (GWAS) in diploids and tetraploids
- 4) Genomic selection
- 5) Single cross prediction.

Background

The core of the package is the **mmer** function and solves the mixed model equations proposed by Henderson (1975). An user friendly version named **mmer2** has been added as well, using the ASReml syntax. The functions are an interface to call one of the 4 ML/REML methods supported in the package; **EMMA** efficient mixed model association (Kang et al. 2008), **AI** average information (Gilmour et al. 1995; Lee et al. 2015), **EM** expectation maximization (Searle 1993; Bernardo 2010), and **NR** Newton-Raphson (Tunnicliffe 1989). The **EMMA** method can be implemented when only one variance component other than the error variance component (σ_e^2) is being estimated. On the other hand when more than one variance component needs to be estimated the **AI**, **EM** or **NR** methods should be used.

The mixed model solved by the algorithms has the form:

$$y = X\beta + Zu + \epsilon$$

or

$$y = X\beta + Zu_1 + \dots + Zu_i + \epsilon$$

where:

X is an incidence matrix for fixed effects

Z is an incidence matrix for random effects

β is the vector for BLUEs of fixed effects

u is the vector for BLUPs of random effects

ϵ are the residuals

The variance of the response is known to be the random part of the model:

$$\text{Var}(y) = \text{Var}(Zu + \epsilon) = ZGZ + R$$

and with

$$u \sim \text{MVN}(u, G\sigma_u^2)$$

$$\epsilon \sim \text{MVN}(\epsilon, R\sigma_e^2)$$

When multiple random effects are present the Z matrix becomes the column binding of each of the Z_i matrices for the i random effects. And the G matrix becomes the diagonal binding of each of the variance covariance structures (K matrices) for the random effects:

$$Z = [Z_1 \dots Z_i]$$

$$G = \begin{bmatrix} K_1 & 0 & 0 \\ 0 & \dots & 0 \\ 0 & 0 & K_i \end{bmatrix}$$

The program takes the Z and G for each random effect and construct the neccesary structure inside and estimate the variance components by ML/REML using any of the 4 methods available in sommer; Average Information, Expectation-Maximization, Newton-Raphson, and Efficient Mixed Model Association. Please refer to the canonical paper liste in the Literature section to check how the methods work. We have tested widely the methods to make sure they provide the same solution when the likelihood behaves well but for complex problems they might lead to different answers. If you have any concer please contact me at cova_ruber@live.com.mx or covarrubiasp@wisc.edu.

1) Marker and non-marker based heritability calculation

The heritability is one of the most famous parameters in breeding and genetics theory. The heritability is usually estimated as narrow sense (h^2 ; only additive variance in the numerator σ_A^2), and broad sense (H^2 ; all genetic variance in the numerator σ_G^2).

In a classical experiment with no molecular markers, special designs are performed to estimate and dissect the additive (σ_A^2) and dominance (σ_D^2) variance along with environmental variability. Designs such as generation analysis, North Carolina designs ae used to dissect σ_A^2 and σ_D^2 to estimate the narrow sense heritability (h^2). When no special design is available we can still dissect the genetic variance (σ_G^2) and estimate the broad sense. In this example we will show the broad sense estimation.

The dataset has 41 potato lines evaluated in 5 locations across 3 years in an RCBD design. We show how to fit the model and extract the variance components to calculate the h^2 .

```
library(sommer)
data(h2)
head(h2)

##          Name Env Loc Year     Block y
## 1      W8822-3 FL.2012 FL 2012 FL.2012.1 2
## 2      W8867-7 FL.2012 FL 2012 FL.2012.2 2
## 3    MSL007-B MO.2011 MO 2011 MO.2011.1 3
## 4 C000270-7W FL.2012 FL 2012 FL.2012.2 3
## 5 Manistee(MSL292-A) FL.2013 FL 2013 FL.2013.2 3
## 6      MSM246-B FL.2012 FL 2012 FL.2012.2 3
```

```
ans1 <- mmmer2(y~1, random=~Name + Env + Name:Env + Block, data=h2, silent = TRUE)
```

```
## Estimating variance components
```

```
vc <- ans1$var.comp
V_E <- vc[2,1]; V_GE <- vc[3,1]; V_G <- vc[1,1]; Ve <- vc[5,1]

n.env <- length(levels(h2$Env))
h2 <- V_G/(V_G + V_GE/n.env + Ve/(2*n.env)) #the 2 is a reference for block
h2
```

```
## [1] 0.9621128
```

Recently with markers becoming cheaper, thousand of markers can be run in the breeding materials. When markers are available, an special design is not necessary to dissect the additive variance. The estimation of the additive relationship matrix allow us to estimate the σ_A^2 and σ_D^2 .

Assume you have a population and a similar model like the one displayed previously has been fitted. Now we have BLUPs for the genotypes but in addition we have genetic markers. NOTICE WE WILL USE THE mmmer FUNCTION THIS TIME.

```
data(CPdata)
CPpheno <- CPdata$pheno
CPgeno <- CPdata$geno
### look at the data
head(CPpheno)
```

```
##      color Yield FruitAver Firmness
## 3 0.10075269 154.67     41.93  588.917
## 4 0.13891940 186.77     58.79  640.031
## 5 0.08681502  80.21     48.16  671.523
## 6 0.13408561 202.96     48.24  687.172
## 7 0.13519278 174.74     45.83  601.322
## 8 0.17406685 194.16     44.63  656.379
```

```
CPgeno[1:5,1:4]
```

```
##      scaffold_50439_2381 scaffold_39344_153 uneak_3436043 uneak_2632033
## P003                      0                      0                      0                      1
## P004                      0                      0                      0                      1
## P005                      0                     -1                      0                      1
## P006                     -1                     -1                     -1                      0
## P007                      0                      0                      0                      1
```

```
## fit a model including additive and dominance effects
y <- CPpheno$color
Za <- diag(length(y)); Zd <- diag(length(y)); Ze <- diag(length(y))
A <- A.mat(CPgeno) # additive relationship matrix
D <- D.mat(CPgeno) # dominance relationship matrix
E <- E.mat(CPgeno) # epistatic relationship matrix
```

```
ETA.ADE <- list(list(Z=Za,K=A),list(Z=Zd,K=D),list(Z=Ze,K=E))
ans.ADE <- mmmer(y=y, Z=ETA.ADE,silent = TRUE)
```

```

## Estimating variance components
##
## One or more variance components close to zero. Boundary constraint applied.

(H2 <- sum(ans.ADE$var.comp[1:3,1])/sum(ans.ADE$var.comp[,1]))
```

```

## [1] 0.7521069
```

```

(h2 <- sum(ans.ADE$var.comp[1,1])/sum(ans.ADE$var.comp[,1]))
```

```

## [1] 0.6222834
```

In the previous example we showed how to estimate the additive (σ_A^2), dominance (σ_D^2), and epistatic (σ_I^2) variance components based on markers and estimate broad (H^2) and narrow sense heritability (h^2).

2) Half and full diallel designs

When breeders are looking for the best single cross combinations, diallel designs have been by far the most used design in crops like maize. There are 4 types of diallel designs depending if reciprocate and self cross are performed. In this example we will show a full diallel design (reciprocate crosses are performed) and half diallel designs (only one of the directions is performed).

In the first data set we show a full diallel among 40 lines from 2 heterotic groups, 20 in each. Therefore 400 possible hybrids are possible. We have phenotypic data for 100 of them across 4 locations. We use the data available to fit a model of the form:

$$y = X\beta + Zu_1 + Zu_2 + Zu_S + \epsilon$$

We estimate variance components for GCA_1 , GCA_2 and SCA and use them to estimate heritability. Additionally BLUPs for GCA and SCA effects can be used to predict crosses.

```

data(cornHybrid)
hybrid2 <- cornHybrid$hybrid # extract cross data
head(hybrid2)
```

	Location	GCA1	GCA2	SCA	Yield	PlantHeight
## 1	1	A258	AS5707	A258:AS5707	NA	NA
## 2	1	A258	B2	A258:B2	NA	NA
## 3	1	A258	B99	A258:B99	NA	NA
## 4	1	A258	LH51	A258:LH51	NA	NA
## 5	1	A258	Mo44	A258:Mo44	NA	NA
## 6	1	A258	NC320	A258:NC320	NA	NA

```
modFD <- mmmer2(Yield~Location, random=~GCA1+GCA2+SCA, data=hybrid2,silent = TRUE)
```

```

## Estimating variance components
##
## One or more variance components close to zero. Boundary constraint applied.
```

```

summary(modFD)

##
## Information contained in this fitted model:
## * Variance components
## * Residuals and conditional residuals
## * BLUPs and BLUPs
## * Inverse phenotypic variance(V)
## * Variance-covariance matrix for fixed effects
## * Variance-covariance matrix for random effects
## * Predicted error variance (PEV)
## * LogLikelihood
## * AIC and BIC
## * Fitted values
## Use the 'str' function to access such information
##
## =====
## Linear mixed model fit by restricted maximum likelihood
## ***** sommer 1.6 *****
## =====
## Method: [1] "NR"
##
## logLik      AIC      BIC
## -1342     2691     2707
## =====
## Random effects:
##          VarianceComp
## Var(GCA1)      0.000
## Var(GCA2)      7.409
## Var(SCA)       187.563
## Var(Error)    221.142
## Number of obs: 400 Groups: 20 20 400
## =====
## Fixed effects:
##           Value Std.Error t.value
## (Intercept) 1.3794e+02 2.1227e+00 64.9815
## Location2   -1.5987e-13 2.1031e+00  0.0000
## Location3    7.8353e+00 2.1031e+00  3.7257
## Location4   -9.0975e+00 2.1031e+00 -4.3258
## =====
## Var-Cov for Fixed effects:
## (diagonals are variances)
##           (Intercept) Location2 Location3 Location4
## (Intercept)      4.5058   -2.2114   -2.2114   -2.2114
## Location2      -2.2114    4.4228    2.2114    2.2114
## Location3      -2.2114    2.2114    4.4228    2.2114
## Location4      -2.2114    2.2114    2.2114    4.4228
## =====
## Use the 'str' function to access all information

```

```

Vgca <- sum(modFD$var.comp[1:2,1])
Vsca <- modFD$var.comp[3,1]
Ve <- modFD$var.comp[4,1]

```

```
Va = 4*Vgca
Vd = 4*Vsca
Vg <- Va + Vd
(H2 <- Vg / (Vg + (Ve)) )
```

```
## [1] 0.7790852
```

```
(h2 <- Va / (Vg + (Ve)) )
```

```
## [1] 0.02960619
```

In this second data set we show a small half diallel with 7 parents crossed in one direction. $n(n-1)/2$ crosses are possible $7(6)/2 = 21$ unique crosses. Parents appear as males or females indistinctly. Each with two replications in a CRD. For a half diallel design a single GCA variance component can be estimated and an SCA as well ($\sigma_G^2 CA$ and $\sigma_S^2 CA$ respectively). And BLUPs for GCA and SCA of the parents can be extracted. We would create the design matrices in sommer using the `hdm` and `model.matrix` functions for the GCA and SCA matrices respectively.

$$y = X\beta + Zu_g + Zu_s + \epsilon$$

```
data(HDdata)
head(HDdata)
```

```
##   rep geno male female    sugar
## 1   1    12     1      2 13.950509
## 2   2    12     1      2  9.756918
## 3   1    13     1      3 13.906355
## 4   2    13     1      3  9.119455
## 5   1    14     1      4  5.174483
## 6   2    14     1      4  8.452221
```

```
# GCA matrix for half diallel using male and female columns
Z1 <- hdm(HDdata[,c(3:4)])
# SCA matrix
Z2 <- model.matrix(~as.factor(geno)-1, data=HDdata)
# Fit the model
y <- HDdata$sugar
ETA <- list(GCA=list(Z=Z1), SCA=list(Z=Z2)) # Zu component
modHD <- mmer(y=y, Z=ETA, silent = TRUE)
```

```
## Estimating variance components
```

```
summary(modHD)
```

```
##
## Information contained in this fitted model:
## * Variance components
## * Residuals and conditional residuals
## * BLUEs and BLUPs
## * Inverse phenotypic variance(V)
```

```

## * Variance-covariance matrix for fixed effects
## * Variance-covariance matrix for random effects
## * Predicted error variance (PEV)
## * LogLikelihood
## * AIC and BIC
## * Fitted values
## Use the 'str' function to access such information
##
## =====
## Linear mixed model fit by restricted maximum likelihood
## ***** sommer 1.6 *****
## =====
## Method:[1] "AI"
##
## logLik      AIC      BIC
## -7.543 17.086 18.824
## =====
## Random effects:
##          VarianceComp
## Var(GCA)           5.509
## Var(SCA)           1.816
## Var(Error)         3.117
## Number of obs: 42 Groups: 7 21
## =====
## Fixed effects:
##          Value Std.Error t.value
## Intercept 10.3332   1.8189  5.6809
## =====
## Var-Cov for Fixed effects:
## (diagonals are variances)
##          1
## Intercept 3.3086
## =====
## Use the 'str' function to access all information

```

```

Vgca <- modHD$var.comp[1,1]
Vsca <- modHD$var.comp[2,1]
Ve <- modHD$var.comp[3,1]
Va = 4*Vgca
Vd = 4*Vsca
Vg <- Va + Vd
(H2 <- Vg / (Vg + (Ve/2))) # 2 technical reps

```

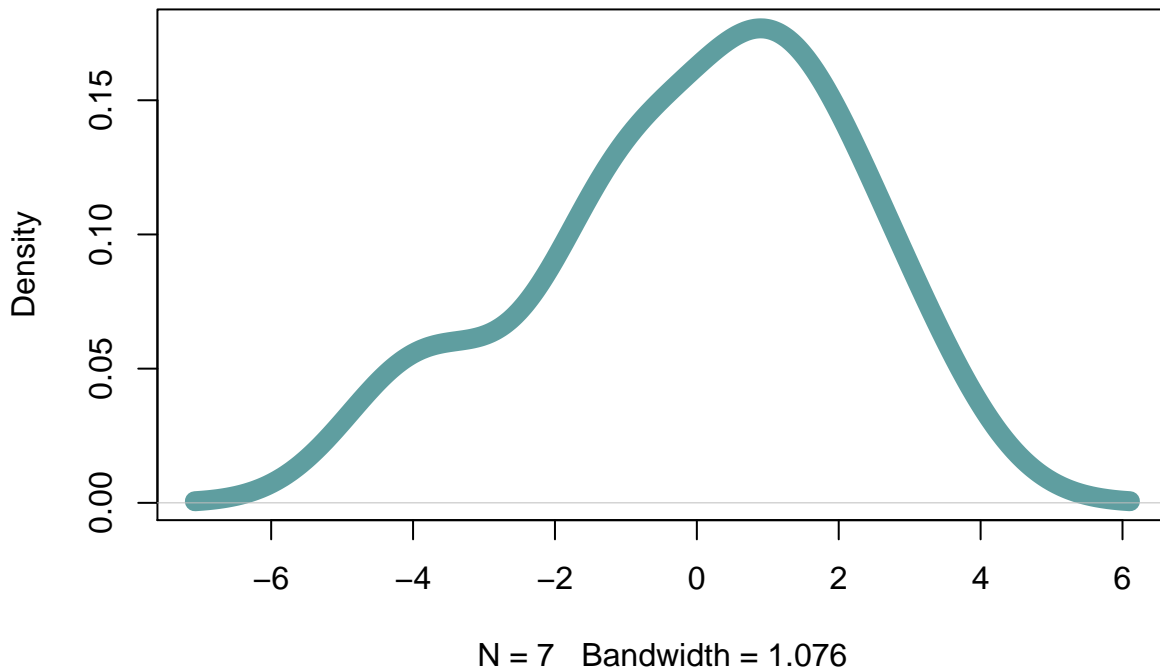
```
## [1] 0.9494882
```

```
(h2 <- Va / (Vg + (Ve/2)))
```

```
## [1] 0.7140884
```

```
plot(density(randef(modHD)$GCA[,1]), col="cadetblue",
lwd=10, main="GCA BLUPs")
```

GCA BLUPs



3) Genome wide association analysis (GWAS) in diploids and tetraploids

With the development of modern statistical machinery the detection of markers associated to phenotypic traits have become quite straight forward. The days of QTL mapping using biparental populations exclusively are in the past. In this section we will show how to perform QTL mapping for diploid and polyploid organisms with complex genetic relationships. In addition we will show QTL mapping in biparental populations to clarify that the fact that is not required anymore doesn't limit the capabilities of modern mixed model machinery.

First we will start doing the GWAS in a biparental population with 363 individuals genotyped with 2889 SNP markers. This is easily done by creating the variance covariance among individuals and using it in the random effect for genotypes. The markers are added in the *W* argument to fit the model of the form:

$$y = X\beta + Zu + Wg + \epsilon$$

In this case $X\beta$ is the fixed part only for the intercept, Zu is the random effect for genotypes with the additive relationship matrix (*A*) as the variance-covariance of the random effect, Wg is the marker matrix and the effects of each marker. This is done in this way:

```
data(CPdata)
CPpheno <- CPdata$pheno
CPgeno <- CPdata$geno
my.map <- CPdata$map
### look at the data
head(CPpheno); CPgeno[1:5,1:4]
```

```

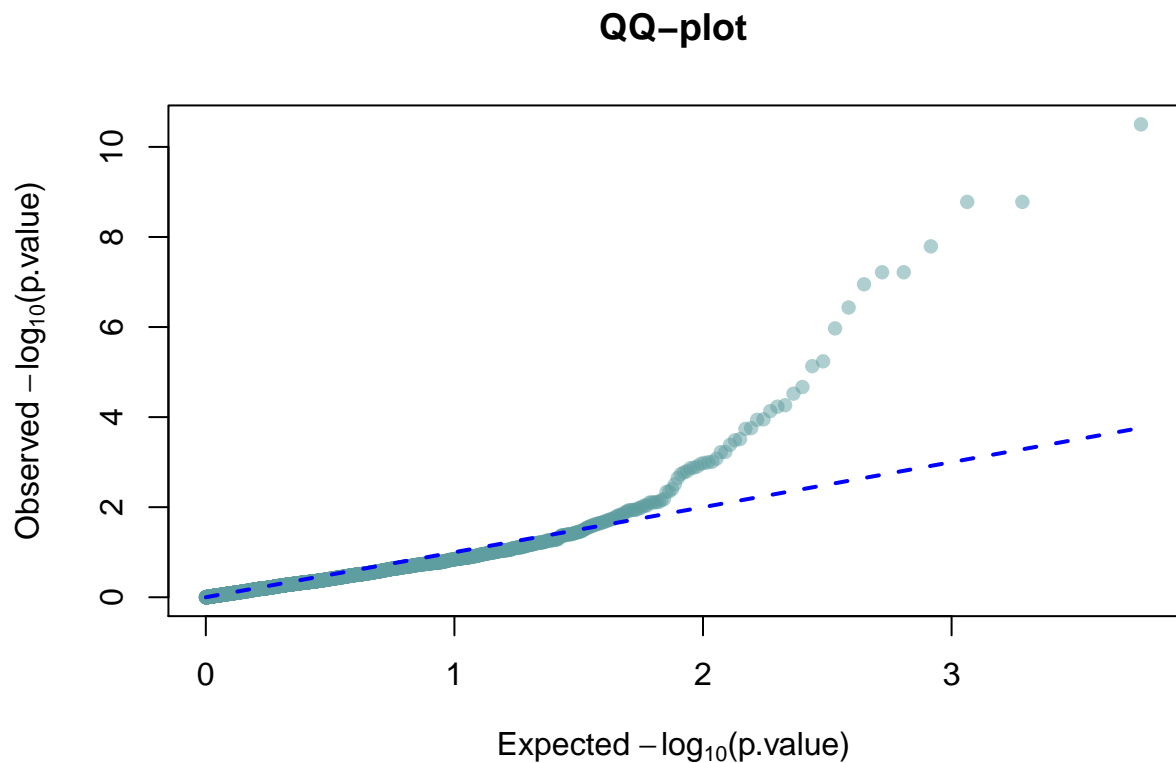
##          color Yield FruitAver Firmness
## 3 0.10075269 154.67    41.93   588.917
## 4 0.13891940 186.77    58.79   640.031
## 5 0.08681502  80.21    48.16   671.523
## 6 0.13408561 202.96    48.24   687.172
## 7 0.13519278 174.74    45.83   601.322
## 8 0.17406685 194.16    44.63   656.379

## scaffold_50439_2381 scaffold_39344_153 uneak_3436043 uneak_2632033
## P003                  0           0           0           1
## P004                  0           0           0           1
## P005                  0           -1          0           1
## P006                 -1          -1          -1          0
## P007                  0           0           0           1

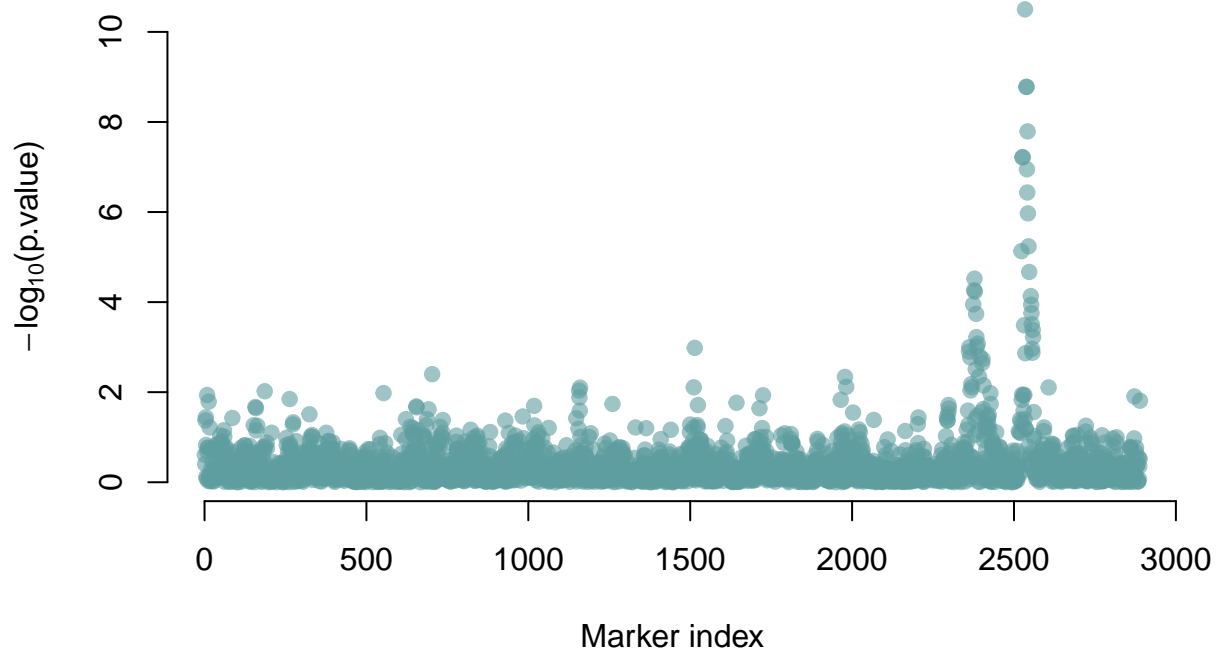
y <- CPpheno$color # response
Za <- diag(length(y)) # incidence matrix for random effect
A <- A.mat(CPgeno) # additive relationship matrix
ETA.A <- list(add=list(Z=Za,K=A)) # create random component
ans.A <- mmmer(y=y, Z=ETA.A, W=CPgeno, silent=TRUE) # fit the model

## Estimating variance components
##
## Performing GWAS
## Running additive model

```



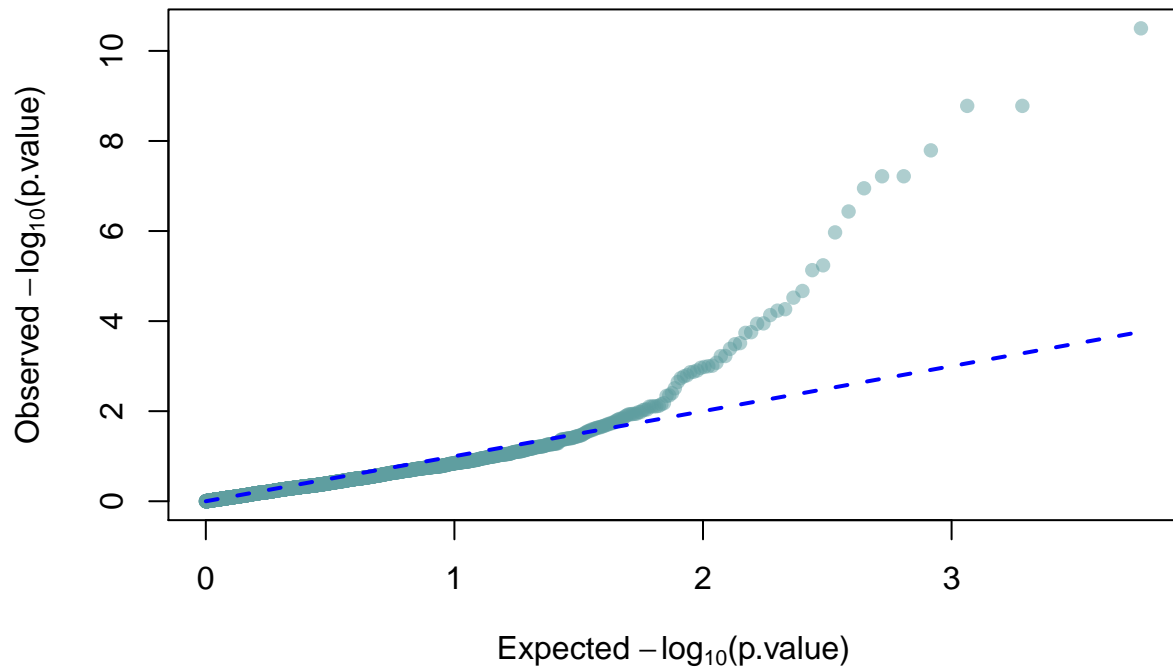
additive model

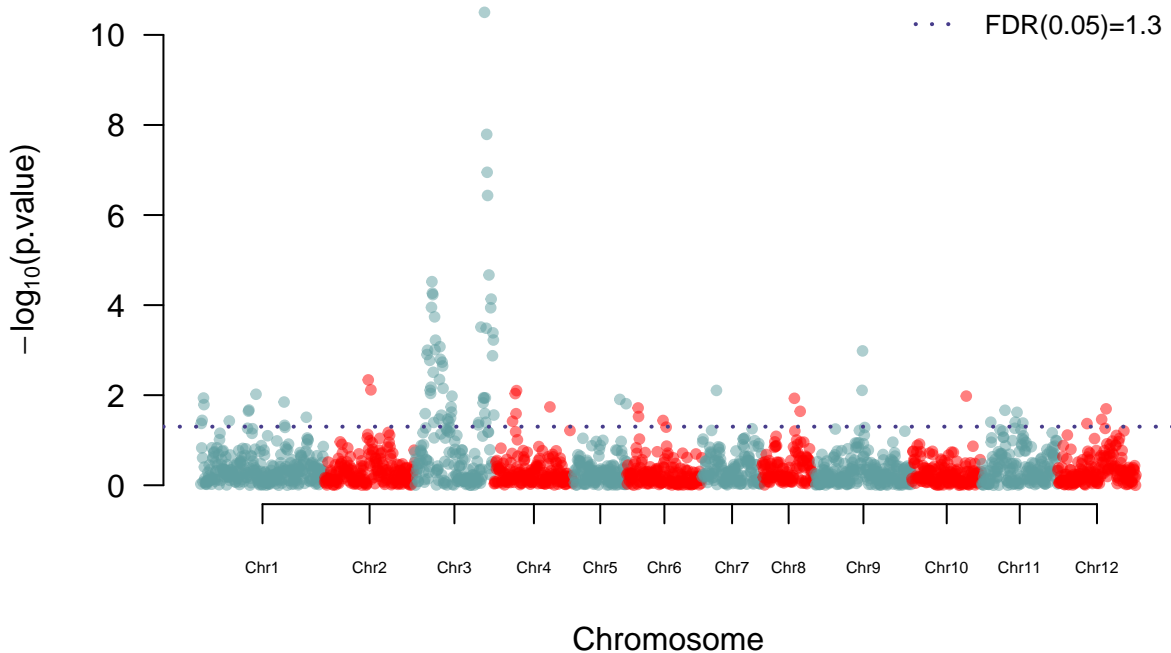


```
### if you have a genetic map you can use it  
ans.B <- mmmer(y=y, Z=ETA.A, W=CPgeno, silent=TRUE, map=my.map) # fit the model
```

```
## Estimating variance components  
##  
## Performing GWAS  
## Running additive model
```

QQ-plot





Now we will show how to do GWAS in a tetraploid using potato data. Is not very different from diploids. We only need to pay attention to the `ploidy` argument in the `atcg1234` and `A.mat` functions. In addition, when running the `mmer` model there is more models that can be implemented according to Rosyara et al. (2016).

```

data(PolyData)
genotypes <- PolyData$PGeno
phenotypes <- PolyData$PPheno
## convert markers to numeric format
numo <- atcg1234(data=genotypes, ploidy=4, silent = TRUE); numo[1:5,1:5]; dim(numo)

## Obtaining reference alleles
## Checking for markers with more than 2 alleles. If found will be removed.
## Converting to numeric format
## Calculating minor allele frequency (MAF)
## Imputing missing data with mode

##          c2_41437 c2_24258 c2_21332 c2_21320 c2_21318
## A96104-2      1        2        2        4        0
## A97066-42      2        3        2        4        1
## ACBrador       2        4        2        4        0
## ACLPI175395    0        4        0        4        0
## ADGPI195204    0        4        0        4        0

## [1] 221 3521

```

```

# get only plants with both genotypes and phenotypes
common <- intersect(phenotypes>Name, rownames(numo))
marks <- numo[common,]; marks[1:5,1:5]

##          c2_41437 c2_24258 c2_21332 c2_21320 c2_21318
## A97066-42           2       3       2       4       1
## ACBrador            2       4       2       4       0
## AdirondackBlue     2       2       2       4       1
## AF2291-10           0       4       2       4       0
## AF2376-5            1       3       2       4       0

phenotypes2 <- phenotypes[match(common,phenotypes>Name),];
phenotypes2[1:5,1:5]

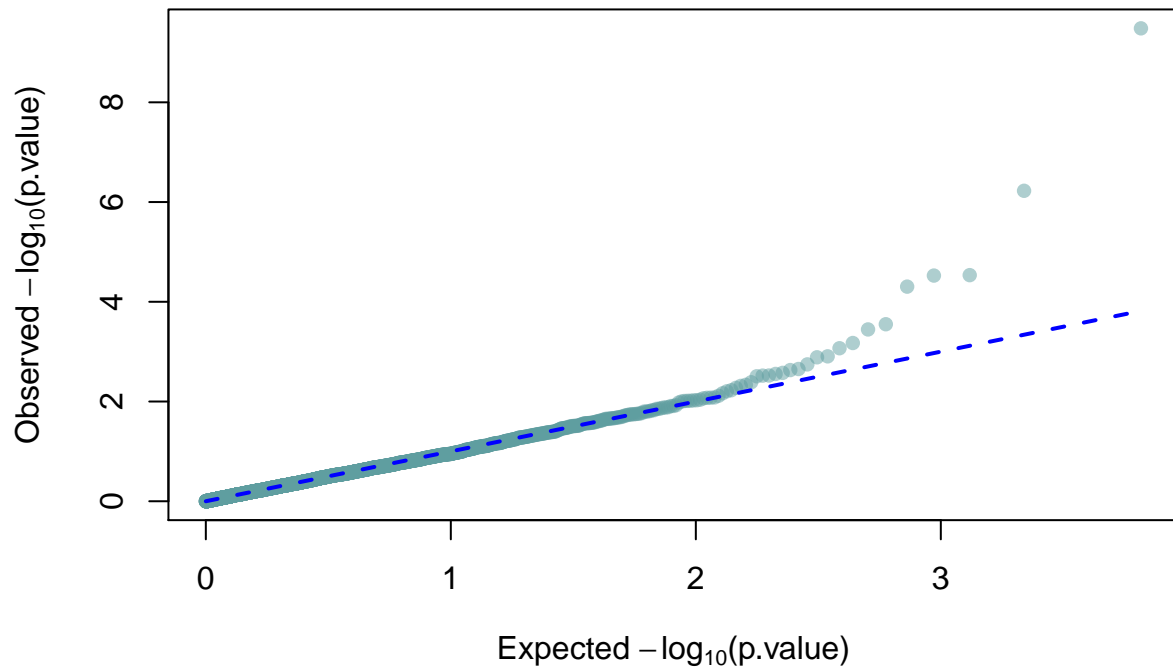
##          Name total_yield chip_color tuber_eye_depth tuber_shape
## 1    A97066-42      13.10      2.35          3.03      4.71
## 2    ACBrador        15.56      2.63          4.37      3.59
## 3 AdirondackBlue    11.77      2.82          3.76      4.07
## 4    AF2291-10      13.43      1.50          4.50      2.81
## 5    AF2376-5       12.58      1.83          4.50      2.81

# Additive relationship matrix, specify ploidy
yy <- phenotypes2$tuber_shape
K1 <- A.mat(marks, ploidy=4)
Z1 <- diag(length(yy))
ETA <- list( list(Z=Z1, K=K1)) # random effects for genotypes
# run the model
models <- c("additive","1-dom-alt","1-dom-ref","2-dom-alt","2-dom-ref")
ans2 <- mmmer(y=yy, Z=ETA, W=marks, method="EMMA",
               ploidy=4, models=models[1], silent = TRUE)

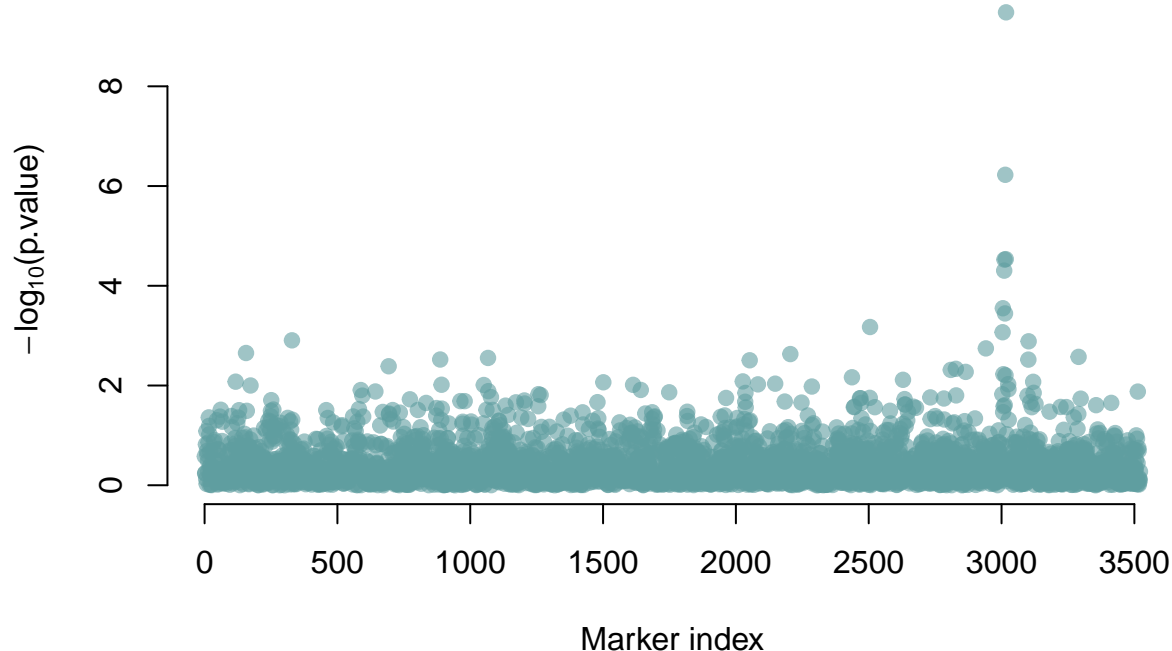
## Estimating variance components
##
## Performing GWAS
## Running additive model

```

QQ-plot



additive model



```
summary(ans2)
```

```
##  
## Information contained in this fitted model:  
## * Variance components  
## * Residuals and conditional residuals  
## * BLUEs and BLUPs  
## * Inverse phenotypic variance(V)  
## * Variance-covariance matrix for fixed effects  
## * Variance-covariance matrix for random effects  
## * Predicted error variance (PEV)  
## * LogLikelihood  
## * AIC and BIC  
## * Fitted values  
## Use the 'str' function to access such information  
##  
## =====  
## Linear mixed model fit by restricted maximum likelihood  
## ***** sommer 1.6 *****  
## =====  
## Method: [1] "EMMA"  
##  
## logLik     AIC      BIC  
## -192.5   387.0   390.3  
## =====
```

```

## Random effects:
##      VarianceComp
## V(u)      0.60778
## V(e)      0.03807
## Number of obs: 187  Groups: 187
## =====
## Fixed effects:
##           Value Std.Error t.value
## Intercept 3.307861  0.014268 231.83
## =====
## Var-Cov for Fixed effects:
## (diagonals are variances)
##          1
## Intercept 2e-04
## =====
## Use the 'str' function to access all information

```

4) Genomic selection

In this section we will use wheat data from CIMMYT to show how is genomic selection performed. This is the case of prediction of specific individuals within a population. It basically uses a similar model of the form:

$$y = X\beta + Zu + \epsilon$$

and takes advantage of the variance covariance matrix for the genotype effect known as the additive relationship matrix (A) and calculated using the `A.mat` function to establish connections among all individuals and predict the BLUPs for individuals that were not measured. The prediction accuracy depends on several factors such as the heritability (h^2), training population used (TP), size of TP, etc.

```

data(wheatLines)
X <- wheatLines$wheatGeno; X[1:5,1:5]; dim(X)

##      wPt.0538 wPt.8463 wPt.6348 wPt.9992 wPt.2838
## [1,]      -1       1       1       1       1
## [2,]       1       1       1       1       1
## [3,]       1       1       1       1       1
## [4,]      -1       1       1       1       1
## [5,]      -1       1       1       1       1

## [1] 599 1279

Y <- wheatLines$wheatPheno
rownames(X) <- rownames(Y)
# select environment 1
y <- Y[,1] # response grain yield
Z1 <- diag(length(y)) # incidence matrix
K <- A.mat(X) # additive relationship matrix
# GBLUP pedigree-based approach
set.seed(12345)
y.trn <- y
vv <- sample(1:length(y), round(length(y)/5))
y.trn[vv] <- NA
ETA <- list(g=list(Z=Z1, K=K))
ans <- mmer(y=y.trn, Z=ETA, method="EMMA", silent = TRUE) # kinship based

```

```

## Estimating variance components

cor(ans$u.hat$g[vv],y[vv])

## [1] 0.4885687

## maximum prediction value that can be achieved
sqrt(ans$var.comp[1,1]/sum(ans$var.comp[,1]))


##          V(u)
## 0.5771923

```

5) Single cross prediction

When doing prediction of single cross performance the phenotype can be dissected in three main components, the general combining abilities (GCA) and specific combining abilities (SCA). This can be expressed with the same model analyzed in the diallel experiment mentioned before:

$$y = X\beta + Zu_1 + Zu_2 + Zu_s + \epsilon$$

with:

$$u_1 \sim N(0, K_1 \sigma_u^2 1)$$

$$u_2 \sim N(0, K_2 \sigma_u^2 2)$$

$$u_s \sim N(0, K_3 \sigma_u^2 s)$$

And we can specify the K matrices. The main difference between this model and the full and half diallel designs is the fact that this model will include variance covariance structures in each of the three random effects (GCA1, GCA2 and GCA3) to be able to predict the crosses that have not occurred yet. We will use the data published by Technow et al. (2015) to show how to do prediction of single crosses.

```

data(Technow_data)

A.flint <- Technow_data$AF # Additive relationship matrix Flint
A.dent <- Technow_data$AD # Additive relationship matrix Dent
M.flint <- Technow_data$MF # Marker matrix Flint
M.dent <- Technow_data$MD # Marker matrix Dent

pheno <- Technow_data$pheno # phenotypes for 1254 single cross hybrids
pheno$hy <- paste(pheno$dent, pheno$flint, sep=":");head(pheno);dim(pheno)

##      hybrid dent flint      GY      GM      hy
## 1 518.298 518    298 -8.04 -0.85 518:298
## 2 518.305 518    305 -11.10  1.70 518:305
## 3 518.306 518    306 -16.85  2.24 518:306
## 4 518.316 518    316   2.08 -1.33 518:316
## 5 518.323 518    323   5.65 -2.71 518:323
## 6 518.327 518    327 -16.95 -0.52 518:327

## [1] 1254     6

```

```

# CREATE A DATA FRAME WITH ALL POSSIBLE HYBRIDS
DD <- kronecker(A.dent,A.flint,make.dimnames=TRUE)

hybs <- data.frame(sca=rownames(DD),yield=NA,matter=NA,gcad=NA, gcaf=NA)
hybs$yield[match(pheno$hy, hybs$sca)] <- pheno$GY
hybs$matter[match(pheno$hy, hybs$sca)] <- pheno$GM
hybs$gcad <- as.factor(gsub(":.*","",hybs$sca))
hybs$gcaf <- as.factor(gsub(".*:","",hybs$sca))
head(hybs)

##          sca yield matter gcad gcaf
## 1 513:316 10.02 -2.05  513   316
## 2 513:323  6.97 -3.78  513   323
## 3 513:330     NA     NA  513   330
## 4 513:336     NA     NA  513   336
## 5 513:340     NA     NA  513   340
## 6 513:341     NA     NA  513   341

# CREATE INCIDENCE MATRICES
Z1 <- model.matrix(~gcad-1, data=hybs)
Z2 <- model.matrix(~gcaf-1, data=hybs)
# SORT INCIDENCE MATRICES ACCORDING TO RELATIONSHIP MATRICES, REAL ORDERS
real1 <- match( colnames(A.dent), gsub("gcad","",colnames(Z1)))
real2 <- match( colnames(A.flint), gsub("gcaf","",colnames(Z2)))
Z1 <- Z1[,real1]
Z2 <- Z2[,real2]
# RUN THE PREDICTION MODEL
y.trn <- hybs$yield
vv1 <- which(!is.na(hybs$yield))
vv2 <- sample(vv1, 100)
y.trn[vv2] <- NA
ETA2 <- list(GCA1=list(Z=Z1, K=A.dent), GCA2=list(Z=Z2, K=A.flint))
anss2 <- mmer(y=y.trn, Z=ETA2, method="EM", silent=TRUE)

## Estimating variance components

summary(anss2)

## 
## Information contained in this fitted model:
## * Variance components
## * Residuals and conditional residuals
## * BLUEs and BLUPs
## * Inverse phenotypic variance(V)
## * Variance-covariance matrix for fixed effects
## * Variance-covariance matrix for random effects
## * Predicted error variance (PEV)
## * LogLikelihood
## * AIC and BIC
## * Fitted values
## Use the 'str' function to access such information
##

```

```

## =====
## Linear mixed model fit by restricted maximum likelihood
## ***** sommer 1.6 *****
## =====
## Method: [1] "EM"
##
## logLik      AIC      BIC
## -4998    9999   10004
## =====
## Random effects:
##          VarianceComp
## Var(GCA1)      16.01
## Var(GCA2)      11.16
## Var(Error)     17.71
## Number of obs: 1154 Groups: 123 86
## =====
## Fixed effects:
##           Value Std.Error t.value
## Intercept 0.24778  0.20360  1.2169
## =====
## Var-Cov for Fixed effects:
## (diagonals are variances)
##          1
## Intercept 0.0415
## =====
## Use the 'str' function to access all information

cor(anss2$fitted.y[vv2], hybs$yield[vv2])

```

```
## [1] 0.8778897
```

In the previous model we only used the GCA effects (GCA1 and GCA2) for practicality, although it's been shown that the SCA effect doesn't actually help that much in increasing prediction accuracy and increase a lot the computation intensity required since the variance covariance matrix for SCA is the kronecker product of the variance covariance matrices for the GCA effects, resulting in a 10578x10578 matrix that increases in a very intensive manner the computation required.

A model without covariance structures would show that the SCA variance component is insignificant compared to the GCA effects. This is why including the third random effect doesn't increase the prediction accuracy.

Literature

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