Changes and FAQs for the sommer package

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2020-06-25

The sommer package was developed to provide R users a powerful and reliable multivariate mixed model solver. The package is focused in problems of the type p > n (more effects to estimate than observations) and its core algorithm is coded in C++ using the Armadillo library. This package allows the user to fit mixed models with the advantage of specifying the variance-covariance structure for the random effects, and specify heterogeneous variances, and obtain other parameters such as BLUPs, BLUEs, residuals, fitted values, variances for fixed and random effects, etc.

Recently, I decided to code the main algorithm (Newton-Raphson & Average-Information) in C++ which encouraged me to refactor all the machinery including special functions and specification of the models. For a more in depth explanation of how the machinery works please read the "Quick start for the sommer package" vignette [i.e. typing vignette ('sommer.start')]. Here I will focus in just making a translation of the old specification to the new specification.

The purpose of this vignette is to first show the changes in sintax for sommer and frequently asked question.

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SECTION 1: The new sintax of sommer

1) The specification of multiresponse model

In past versions (depending how old your version) there was an argument called MVM which had to be set to TRUE if the user wanted to run a true multi-trait model since the specification

fixed= cbind(y1,y2)~x

would by default fit 2 univariate models in parallel. That is no longer the case, the MVM argument doesn't exist and if a model like the one above is specified it will run a tru multi-trait model.

2) The specification of multivariate unknown covariance structures

In the previous versions when I introduced the multivariate solver I decided to follow the asreml sintax to specify the unknown covariance structured that needed to be estimated. For example, a diagonal model for the multitrait model, assuming a random effect called 're' looked something like this:

fixed= cbind(y1,y2)~x

random= ~ diag(trait):re

and an unstructured multitrait model was:

random= ~ us(trait):re

Although this was easier for users adapted to asreml a good thing, it put a lot of limitations on the way constraints for variance components where specified. The same model in the new versions looks like this:

where the Gtc argument helps us to indicate what type of structure this random effect represents. Here I specified an unstructured model with the function unsm() with a number 2 for 2 traits. The user can use either diag() or uncm(), or any customized matrix with dimensions t x t (t being the number of traits) containing the number 0,1,2,3 that specify the constraint:

0: not to be estimated 1: estimated and constrained to be positive (i.e. variance component) 2: estimated and unconstrained (can be negative or positive, i.e. covariance component) 3: not to be estimated but fixed (value has to be provided in the Gti argument)

All this models fit a model with the following variance for re:

$$var(u) = T \otimes A$$

where:

$$\mathbf{var}(\mathbf{u}) = \left[\begin{array}{cc} \sigma_{g_{t1,t1}}^2 & \sigma_{g_{t1,t2}} \\ \sigma_{g_{t2,t1}} & \sigma_{g_{t2,t2}}^2 \end{array} \right] \otimes A$$

By doing this change now the user has full control on the constraints applied to the estimation of variance components and can provide initial values easily using the Gti argument.

3) The specification of additional unknown covariance structures

If we focus for a moment in an univariate mixed model we can also have other unknown covariance structures specified.

$$var(u) = E \otimes ... \otimes F \otimes A$$

where:

$$\mathbf{var}(\mathbf{u}) = \left[\begin{array}{cccc} \sigma_{g_{e1,e1}}^2 & \sigma_{g_{e1,e2}} & \sigma_{g_{e1,e3}} \\ \sigma_{g_{e2,e1}} & \sigma_{g_{e2,e2}}^2 & \sigma_{g_{e2,e3}} \\ \sigma_{g_{e3,e1}} & \sigma_{g_{e3,e2}} & \sigma_{g_{e3,e3}}^2 \end{array} \right] \otimes ... \otimes \left[\begin{array}{ccc} \sigma_{g_{f1,f1}}^2 & \sigma_{g_{f1,f2}} \\ \sigma_{g_{f2,f1}} & \sigma_{g_{f2,f2}}^2 \end{array} \right] \otimes A$$

If we think in the multi trait model this is very similar but with an additional kroneker product for the multivariate version:

$$var(u) = T \otimes E \otimes ... \otimes F \otimes A$$

where:

$$\mathbf{var}(\mathbf{u}) = \left[\begin{array}{ccc} \sigma_{g_{t1,t1}}^2 & \sigma_{g_{t1,t2}} \\ \sigma_{g_{t2,t1}} & \sigma_{g_{t2,t2}}^2 \end{array} \right] \otimes \left[\begin{array}{cccc} \sigma_{g_{e1,e1}}^2 & \sigma_{g_{e1,e2}} & \sigma_{g_{e1,e3}} \\ \sigma_{g_{e2,e1}} & \sigma_{g_{e2,e2}}^2 & \sigma_{g_{e2,e3}} \\ \sigma_{g_{e3,e1}} & \sigma_{g_{e3,e2}} & \sigma_{g_{e3,e3}}^2 \end{array} \right] \otimes \ldots \otimes \left[\begin{array}{cccc} \sigma_{g_{f1,f1}}^2 & \sigma_{g_{f1,f2}} \\ \sigma_{g_{f2,f1}} & \sigma_{g_{f2,f2}}^2 \end{array} \right] \otimes A$$

Going back to the point. The additional unknown covariance structures besides the multi-trait (T) before where specified with asreml sintax. For example an univariate diagonal and unstructured model, assumed a random effect called 'id' representing the treatments planted in different environments coded in a second random effect called 'env'. The model used to look like:

fixed= y1~x

random= ~ diag(env):id or random= ~ us(env):id

and now it would be specified as:

fixed= y1~x

random= ~ vs(ds(env),id) or random= ~ vs(us(env),id)

where the ds() and us() functions specifie a diagonal and unstructured models respectively. Now in addition cs() for a customized structure is available. The main gain from having changed the formulation is that the new specification trhough the vs() function allows to contruct more complex moels. For example, assume individuals specified in a column called 'id' tested in three environments in a column called 'env' measured at two different time points specified in a column called 'time'. We may want something more flexible than:

fixed= y1~x

random= ~ id

We could actually assume that individuals are correlated within the environments for the different time points but want to consider environments indepedent. The variance for such random effects is the following:

$$\mathbf{var}(\mathbf{u}) = \left[\begin{array}{ccc} \sigma_{g_{e1,e1}}^2 & \sigma_{g_{e1,e2}} & \sigma_{g_{e1,e3}} \\ \sigma_{g_{e2,e1}} & \sigma_{g_{e2,e2}}^2 & \sigma_{g_{e2,e3}} \\ \sigma_{g_{e3,e1}} & \sigma_{g_{e3,e2}} & \sigma_{g_{e3,e3}}^2 \end{array} \right] \otimes \left[\begin{array}{ccc} \sigma_{g_{t1,t1}}^2 & \sigma_{g_{t1,t2}} \\ \sigma_{g_{t2,t1}} & \sigma_{g_{t2,t2}}^2 \end{array} \right] \otimes A$$

which was not possible in previous versions of sommer and now can be specified as:

random= ~ vs(us(env),ds(time),id)

and the same logic can be extended to as many factors interacting as desired.

4) The specification of unknown covariance structures in the residuals

Before sommer was limited to only diagonal models in the residuals (unstructured available only for multi-trait before). Now all the same applications discussed for the random term also applies for the residual term. Just keep in mind that the residual term is always called units.

Previous versions:

random= ~ diag(trait):diag(env):units

random= ~ us(trait):diag(env):units # limit

New versions (>3.7):

random= ~ vs(ds(env),units, Gtc=mm) ## can be extended to more interacting factors

random= ~ vs(us(env),units, Gtc=mm) ## can be extended to more interacting factors

random= ~ vs(at(env), units, Gtc=mm) ## can be extended to more interacting factors

random= ~ vs(cs(env), units, Gtc=mm) ## can be extended to more interacting factors

where mm can be any matrix specifying the type of multi-trait model (constraints). For example we could use unsm() diag(), uncm() or any other customized matrix.

5) Special models

In previous version the use of asreml formulation really limited the expansion of sommer to more sophistiated models. Now there's many more possible models

Previous versions:

Overlay models Previous version: limited to 2 columns and only random and no covariance structures.

```
random = x + and(y)
```

New versions (>3.7): in theory no limits. Can be extended to more interacting factors in the unknown covariance structures and can overlay as many columns as needed. Plus is fully functional with the multivariate models.

```
random=~ vs(..., overlay(x1,...,xn), Gtc=mm)
```

Random regression models Previous version: Not available before

New versions (>3.7): in theory no limits. Can be extended to more interacting factors in the unknown covariance structures and only requires the use of the leg() function. Plus is fully functional with the multivariate models.

```
random=~ vs(us(leg(v,1)),x)
random=~ vs(ds(leg(v,1)),x)
random=~ vs(leg(v,1),x)
```

GWAS models Previous version: Only univariate models available

New versions (>3.7): all the power of the mmer function is available plus you can fit multivariate GWAS models. See details in the sommer start vignettes.

Spatial models Previous version: It was called directly in the formula

```
random=~ spl2D(Row,Col,at=?)
```

New versions (>3.7): It has to be called within the vs() function but now it can be combined with all the unknown covariance structures available.

random=~ vs(...,spl2D(Row,Col,at=?), Gtc=mm) # being mm any multi-trait constraint-structure.

Customized random effects Previous version: It was provided in the grouping argument

```
random=~ grp(x),
grouping=list(x=Z)
```

New versions (>3.7): It has to be called within the vs() function but now it can be combined with all the unknown covariance structures available.

random=~vs(,..., Z, Gtc=mm) # being mm any multi-trait constraint-structure.

SECTION 2: Frequently asked questions

1) I got an error similar to:

```
# iteration LogLik wall cpu(sec) restrained
# 1 -224.676 18:11:23 3 0
# Sistem is singular. Aborting the job. You could try a bigger tolparino value.
```

This error indicates that your model is singular (phenotypic variance V matrix is not invertible) and therefore the model is stopped throwing the error message and returning an empty list. Whether you can try a simpler model or just modify the argument tolparinv in the mmer function. The default is 1e-3, which means that it will try to invert V and if it fails it will try to add a small value to the diagonal of V of 1e-3 to make it invertible and try bigger and biger numbers. If this fails then the program will return the empty list.

Sometimes the model becomes singular when you use variance covariance matrices (i.e. genomic relationship matrices) that are not full-rank. You can try to make it full-rank and try again.

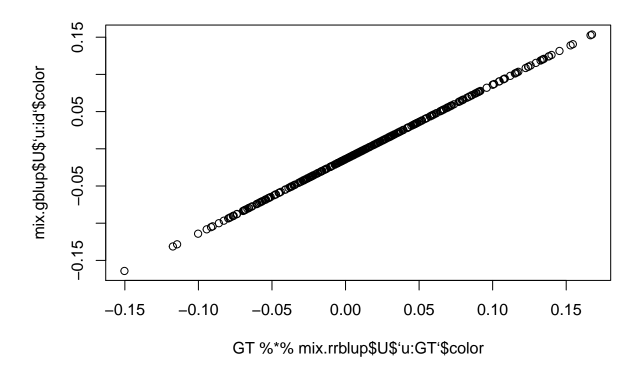
2) My model runs very slow

Keep in mind that sommer uses direct inversion (DI) algorithm which can be very slow for large datasets. The package is focused in problems of the type p > n (more random effect levels than observations) and models with dense covariance structures. For example, for experiment with dense covariance structures with low-replication (i.e. 2000 records from 1000 individuals replicated twice with a covariance structure of 1000×1000) sommer will be faster than MME-based software. Also for genomic problems with large number of random effect levels, i.e. 300 individuals (n) with 100,000 genetic markers (p). For highly replicated trials with small covariance structures or n > p (i.e. 2000 records from 200 individuals replicated 10 times with covariance structure of 200×200) asreml or other MME-based algorithms will be much faster and we recommend you to opt for those software.

3) Can I run both; rrBLUP for markers and GBLUP for individuals in sommer?

Both types of models can be fitted in sommer. The only thing that it changes is what is the random effect of interest; the marker matrix or the identifier for the individual. Here there is a complex example using multi-trait models but can be used with only one trait.

```
## -----
## Variance-Covariance components:
                VarComp VarCompSE Zratio Constraint
               4.213e-06 8.581e-07 4.909
## u:GT.color-color
## u:Rowf.color-color 1.963e-04 1.355e-04 1.449
                                   Positive
## u:units.color-color 2.612e-03 2.926e-04 8.928 Positive
## Fixed effects:
   Trait
          Effect Estimate Std.Error t.value
## 1 color (Intercept) 0.1692 0.03908 4.329
## Groups and observations:
      color
       2889
## u:GT
## u:Rowf
       13
## -----
## Use the '$' sign to access results and parameters
## GBLUP for individuals
A <- A.mat(GT)
mix.gblup <- mmer(fixed=color~1,</pre>
            random=~vs(id,Gu=A, Gtc=unsm(1)) + vs(Rowf,Gtc=diag(1)),
            rcov=~vs(units,Gtc=unsm(1)),
            data=DT, verbose = FALSE)
summary(mix.gblup)
Multivariate Linear Mixed Model fit by REML
## ***************** sommer 4.1 ************
logLik AIC BIC Method Converge
##
## Value -108.1201 218.2403 222.1319 NR TRUE
## Variance-Covariance components:
                VarComp VarCompSE Zratio Constraint
## u:id.color-color 0.0049524 0.0010082 4.912 Positive
## u:Rowf.color-color 0.0001965 0.0001359 1.446 Positive
## u:units.color-color 0.0026123 0.0002926 8.928 Positive
## Fixed effects:
         Effect Estimate Std.Error t.value
## 1 color (Intercept) 0.1831 0.004732 38.7
## Groups and observations:
##
      color
## u:id
        363
## u:Rowf
       13
## -----
## Use the '$' sign to access results and parameters
## Equivalence
plot(GT%*%mix.rrblup$U$`u:GT`$color, mix.gblup$U$`u:id`$color)
```



4) I am missing BLUPs for individuals even when I provided them in the relationship matrix

I got this good question in the past: "when I want to fit an animal model with sommer package using additive relationship matrix(A), this A matrix would contain parents. But the random effects only contains animals in the random effect but not including parents in the A matrix. How can I get the random effects for parents?"

Answer: The easy way to do it is to make sure that even if the parents don't show up in the dataset, you need to make sure that they are present in the levels of the column that contains the individuals (i.e. animal IDs), in addition they have to be provided in the relationship matrix and that's it. They should be returned in the blups.

```
library(sommer)
data(DT_cpdata)
DT <- DT_cpdata
GT <- GT cpdata
MP <- MP_cpdata
#### create the variance-covariance matrix
A <- A.mat(GT) # additive relationship matrix
#### look at the data and fit the model
set.seed(12)
DT2 <- droplevels(DT[sample(1:nrow(DT),100),]) # we simulate a dataset with only 100 animals
nrow(DT2); length(levels(DT2$id))
## [1] 100
## [1] 100
# we fit a model with the reduced datatset where only 100 blups will be returned since only
# 100 levels exist in the "id" column
mix1 <- mmer(Yield~1,
              random=~vs(id,Gu=A)
                      + Rowf + Colf,
```

```
rcov=~units,
         data=DT2, verbose = FALSE)
summary(mix1)
Multivariate Linear Mixed Model fit by REML
## ************** sommer 4.1 ***********
logLik AIC
##
                      BIC Method Converge
## Value -47.00674 96.01348 98.61865 NR TRUE
## -----
## Variance-Covariance components:
              VarComp VarCompSE Zratio Constraint
                    1000.9 1.530
## u:id.Yield-Yield 1531.7
                                 Positive
              157.1
                      297.5 0.528
## Rowf.Yield-Yield
                                 Positive
## Colf.Yield-Yield
                      396.4 0.000 Positive
                 0.0
## units.Yield-Yield 3358.4
                      883.6 3.801 Positive
## Fixed effects:
## Trait
          Effect Estimate Std.Error t.value
## 1 Yield (Intercept) 127.4 7.214 17.66
## Groups and observations:
     Yield
## u:id 100
## Rowf
       13
## Colf
       35
## -----
## Use the '$' sign to access results and parameters
length(mix1$U$`u:id`$Yield) # only 100 levels
## [1] 100
# we add additional levels to the "id" column and also provide them in the relationship matrix
levels(DT2$id) <- c(levels(DT2$id), setdiff(levels(DT$id), levels(DT2$id)))</pre>
mix2 <- mmer(Yield~1,
        random=~vs(id,Gu=A)
        + Rowf + Colf,
        rcov=~units,
        data=DT2, verbose = FALSE)
summary(mix2)
Multivariate Linear Mixed Model fit by REML
## ************** sommer 4.1 ***********
AIC BIC Method Converge
        logLik
## Value -47.00674 96.01348 98.61865 NR
## -----
## Variance-Covariance components:
##
              VarComp VarCompSE Zratio Constraint
## u:id.Yield-Yield 1531.7 1000.9 1.530 Positive
                      297.5 0.528 Positive
## Rowf.Yield-Yield 157.1
## Colf.Yield-Yield
               0.0
                      396.4 0.000 Positive
```

```
## units.Yield-Yield 3358.4
                       883.6 3.801
## Fixed effects:
##
   Trait
           Effect Estimate Std.Error t.value
## 1 Yield (Intercept)
                  127.4
                          7.214
## Groups and observations:
##
     Yield
## u:id
       363
## Rowf
       13
## Colf
       35
## Use the '$' sign to access results and parameters
length(mix2$U$`u:id`$Yield) # now 363 levels
## [1] 363
```

5) How can I use the AR1(), CS() and ARMA() functions

Sommer doesn't support the estimation of additional correlation components like AR1 in the way asreml does. Still, if the user know the correlation value or can do an iterative approach to find the best value then these functions can be used to specify the variance covariance structure for a given random effect.

For example, in the DT_cpdata dataset we have a field with row and column coordinates. This allows to fit row and column as random effects:

```
## VarComp VarCompSE Zratio Constraint
## Rowf.Yield-Yield 832.2879 393.8951 2.112968 Positive
## Colf.Yield-Yield 153.9201 126.7582 1.214281 Positive
## units.Yield-Yield 3647.3486 290.4910 12.555804 Positive
```

If the user wants to relax the independence between rows and define an AR1 covariance structure among rows then the model could be fitted as:

```
## VarComp VarCompSE Zratio Constraint
## u:Rowf.Yield-Yield 791.8219 387.8695 2.041465 Positive
## Colf.Yield-Yield 154.5660 126.8094 1.218885 Positive
```

```
## units.Yield-Yield 3643.6027 290.1689 12.556834 Positive
```

Same could be done for the column random effect:

```
## VarComp VarCompSE Zratio Constraint
## Rowf.Yield-Yield 830.3623 392.8264 2.113815 Positive
## u:Colf.Yield-Yield 178.7490 134.2703 1.331262 Positive
## units.Yield-Yield 3624.6074 287.6072 12.602629 Positive
```

If on the other hand, you would like to stablish the presence of correlation in row and columns at the same time the model would look like this:

```
## VarComp VarCompSE Zratio Constraint
## u:Rowf:Colf.Yield-Yield 2474.339 730.1474 3.388821 Positive
## units.Yield-Yield 2025.584 622.1023 3.256030 Positive
```

Notice that if you specify a random effect that is the interaction between 2 random effects the covariance structure to be specified in the Gu argument has to be built using the kronecker() function. The same applies to the ARMA() and CS() functions. Please keep in mind that the correlation values (rho argument) is a fixed value not estimated by REML like asreml does but you can always follow an iterative approach.

6) Can I run GWAS in MET experiments with replicates?

Although the direct-inversion algorithm that sommer uses in the background is not the best choice to solve GWAS models for the n > p scenario it is still possible to perform GWAS in MET models.

For example, assume a MET model that has 41 lines with 1000 SNP markers, tested in 3 environments. The genetic term in a MET can be modeled as CS, DIAG or US covariance. The whole point of the GWAS function in sommer is to provide the name of the gTerm (random effect) that will match the marker data provided.

We first make up marker data for the MET data for example purposes

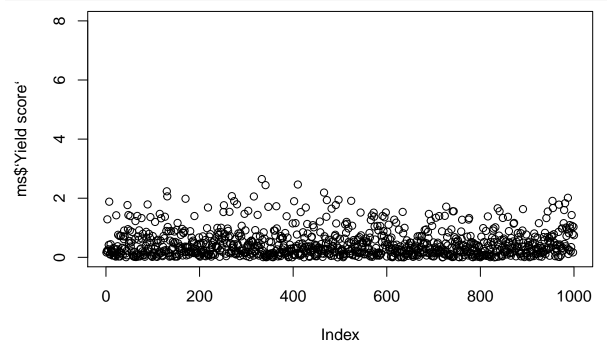
```
library(sommer)
data(DT_example)
DT <- DT_example
A <- A_example
M <- matrix(rep(0,41*1000),1000,41)</pre>
```

```
for (i in 1:41) {
   M[,i] <- ifelse(runif(1000)<0.5,-1,1)
}
tM <- t(M)</pre>
```

We then fit the MET model of the type compound simmetry (CS) and evaluate the GWAS for the main genetic term "Name":

Performing GWAS evaluation

```
ms <- as.data.frame(t(ansx$scores))
plot(ms$`Yield score`, ylim=c(0,8))</pre>
```



We could do the same but evaluate the GWAS considering the genetic term as the interaction term "Env:Name"

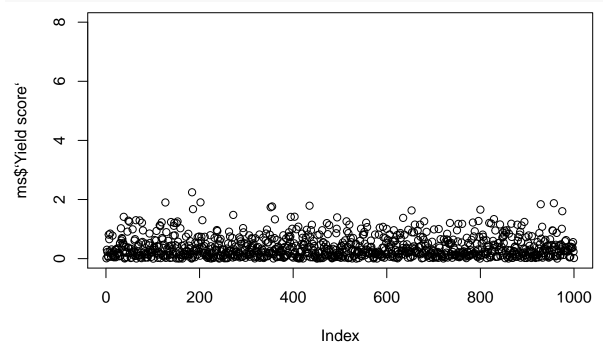
```
## GWAS for the interaction term in CS model
E <- matrix(1,nrow = length(unique(DT$Env)));E</pre>
```

```
data=DT,
M=EtM,
gTerm = "Env:Name", verbose = FALSE)
```

Performing GWAS evaluation

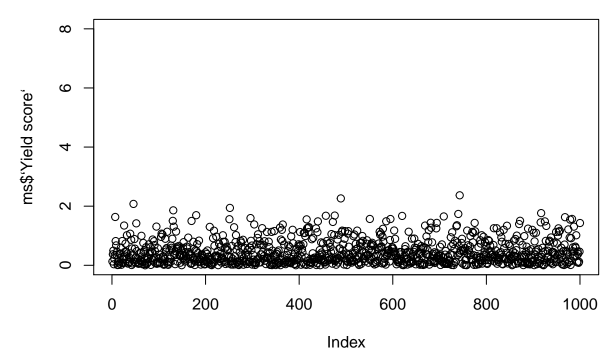
ms <- as.data.frame(t(ansx\$scores))
plot(ms\$'Yield score', ylim=c(0,8))</pre>

```
ms <- as.data.frame(t(ansx$scores))
plot(ms$`Yield score`, ylim=c(0,8))</pre>
```



If the MET is a diagonal model, there's a variance components and BLUPs for each environment. Therefore we can evaluate the GWAS at any environment, here for example we evaluate the GWAS at the genetic term in the environment 'CA.2011'.

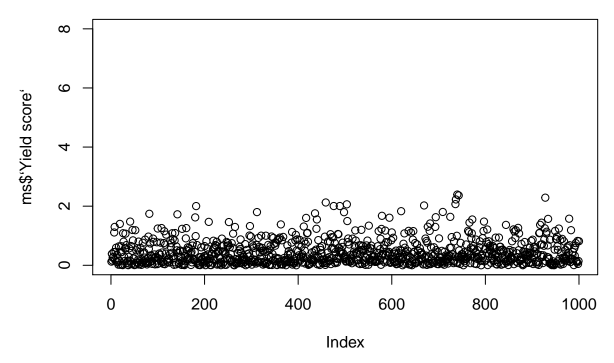
```
## GWAS for the interaction term in DIAG model
E <- matrix(1,nrow = length(unique(DT$Env)));E</pre>
##
         [,1]
## [1,]
            1
## [2,]
## [3,]
EtM <- kronecker(E,tM)</pre>
ansx <- GWAS(Yield~Env,</pre>
              random= ~Name + vs(ds(Env), Name),
              rcov= ~ vs(ds(Env),units),
              data=DT,
              M=tM,
              gTerm = "CA.2011:Name", verbose = FALSE )
## Performing GWAS evaluation
```



If the MET is an unstructured model, there's a variance components and BLUPs for each environment and a covariance component among the different combinations of environments. Therefore we can evaluate the GWAS at any environment as before, here for example we evaluate the GWAS at the genetic term in the environment 'CA.2011'. The difference with the previous model is that here we expect a greater accuracy in the environment CA.2011 since it has borrowed information from the other environments given the covariance fitted among environments.

```
## Performing GWAS evaluation
```

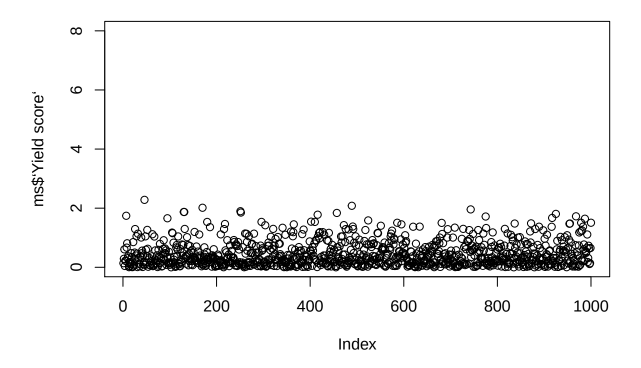
```
ms <- as.data.frame(t(ansx$scores))
plot(ms$`Yield score`, ylim=c(0,8))</pre>
```



Same theory applies for the multitrait model:

Performing GWAS evaluation

```
ms <- as.data.frame(t(ansx$scores))
plot(ms$`Yield score`, ylim=c(0,8))</pre>
```



7) How can I constrain the value of specific random effects?

When using the vs function three additional arguments help to control the following:

Gu: matrix for covariances among levels for the u.th random effect Gti: matrix of initial values for the variance-covariance components Gtc: matrix of constraints for the variance-covariance components

Since each random effect can be seen as a multi-trait variance covariance structure, the univariate models are just an extension where the multi-trait variance covariance structure is a 1 x 1 matrix. When inspecting the results for the mixed models fitted by the mmer() function corresponding to the variance components stored in the sigma element, you will notice that each random effect contains a t x t matrix which corresponds to the multi-trait structure we referred to. For example:

```
## Yield 650.4145
```

Here the sigma element contains the random effects for id and units (error). Each of these contains a matrix of 1 by 1, but in a multi trait model would look like this:

```
data(DT_cpdata)
DT <- DT_cpdata</pre>
```

```
## Yield color
## Yield 634.6295932 0.471715518
## color 0.4717155 0.005126228
```

Notice that for 2 traits this becomes a 2 by 2 matrix. In order to put constraints in some of these random effect matrices you can use the Gtc argument as show above. For the id random effect we have specified that variance components should be estimated and be positive (diagonals with a 1), whereas covariance components should be estimated and unsconstrained to be positive or negative (off-diagonals with a 2).

```
unsm(2)
```

```
## [,1] [,2]
## [1,] 1 2
## [2,] 2 1
mix2$sigma$`u:id`
## Yield color
```

```
## Yield 634.6295932 0.471715518
## color 0.4717155 0.005126228
```

On the other hand, for the units (error) random effect we have specified in the Gtc argument that variance components should be estimated and be positive (diagonals), whereas covariance components should not be estimated (off-diagonals with a 0)

```
diag(2)
```

```
## [,1] [,2]

## [1,] 1 0

## [2,] 0 1

mix2$sigma$`u:units`
```

```
## Yield color
## Yield 4009.336 0.000000000
## color 0.000 0.002563711
```

If the user would like to constraint a value to be fixed and don't change through the estimation process of other random effects the user needs to provide the initial value (scaled with respect to the error variance) of those variance-covariance components in the <code>Gti</code> argument and use a matrix with the value 3 in the <code>Gtc</code> constraint matrix.

```
data=DT, verbose = FALSE)

# analyze variance components
summary(mix1)$varcomp

## VarComp VarCompSE Zratio Constraint
## u:id.Yield-Yield 650.4145 325.5562 1.997856 Positive
## units.Yield-Yield 4031.0153 344.6051 11.697493 Positive
summary(mix3)$varcomp
## VarComp VarCompSE Zratio Constraint
```

```
## u:id.Yield-Yield 325.2072 259.0889 1.255196 Fixed ## u:units.Yield-Yield 4051.3786 352.4804 11.493912 Positive
```

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