Module 5: Genomic Selection and Dissecting G x E in R

Fundamentals of Genomic Prediction and Data-Drive Crop Breeding

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Background Information

For this module, we will use a multi bi-parental mapping population derived by crossing multiple parents. The population is fixed at F_7 generation. The total number of genotypes in the population are **844**. The whole population has been genotyped with the **GBS SNP data** with total number of ~396511 SNP markers.

This mapping population has been divided into training set with 252 genotypes phenotyped across multiple enviornments for Grain yield and other traits. The rest of the 592 genotypes has only genotyped but not phenotyped. We will predict the performance of the 592 genotypes and estimate breeding values for grian yield for them.

What is Our Goal

• Part 2: Disect the G x E interactions and Estimate the Breeding Value

We will use R packages **Sommer** to fit different G x E models. More on this can be found here Click Link. We will aslo **BGLR** package to demonstrate the G x E dissection

NOTE: Due to IRRI's data policies, the actual names of lines is not given

Load the R Packages

```
> library(AGHmatrix)
> library(BGLR)
> library(lme4)
> library(ggplot2)
> library(sommer)
```

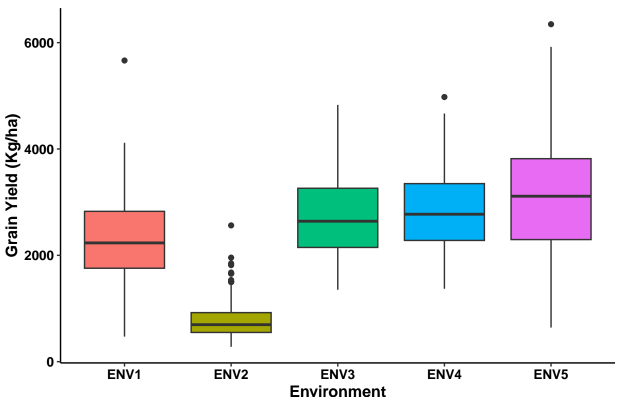
Read the Data Sets

The Data has 5 environments and has yield data. The data comes from the different locations in Bangladesh and India. BLUEs alrady extracted. We will upload the file and use it for analysis.

```
> rm(list=ls()) # remove History
> # Read the phenotypic data
> BLUEs.all<-read.csv(file="./Data/BLUES.ALL.csv")</pre>
```

Visualization of Data

```
> #png(file="~/boxplot.png", width=10, height =6, units = 'in',res=300)
>
          boxplot<-ggplot(BLUEs.all, aes(x=Environment, y=BLUEs))+</pre>
+
            geom_boxplot(aes(fill=Environment))+ # fill by timepoint to give different color
            #scale_fill_manual(values = c("", ""))+
            \#scale\_color\_manual(values = c("", "")) +
            theme_classic()+ #choose the theme for background
            labs(title="",x="Environment", y = "Grain Yield (Kg/ha)")+#Add the labels to the plots
            theme (plot.title = element_text(color="black", size=14, face="bold",hjust=0.5), # add and
                   axis.title.x = element_text(color="black", size=12, face="bold"), # add and modify t
                   axis.title.y = element_text(color="black", size=12, face="bold")) + # add and modify
            theme(axis.text= element_text(face = "bold", color = "black", size = 10))+ # modify the axi
            theme(legend.position="none") # remove the theme from the plot
            \#aes(x = fct\_inorder(timepoint)) + \# order the levels
            #dev.off()
           # ggplotly(p)
      boxplot
```



Read Genotype Data

This marker data as 844 genotypes with 396511 SNP Markers, and the file is saved as .rds. We will subset 252 genotypes and use it estimate the GEBVs.

```
> geno<-readRDS("./Data/GBS_datav2.rds")
> dim(geno)
```

[1] 844 396511

```
> # Match genotype with Phenotype
> Ids<-unique(BLUEs.all$Genotype)
> length(Ids)

[1] 252
> # Now subst the genotype Data based on IDs
> geno<-geno[row.names(geno)%in%Ids,]
> dim(geno)
```

Build the G matrix

252 396511

[1]

[1] 252 252

- Here we will construct the Genomic Relationship Matrix (GRM) using marker data. The GRM will be based on VanRanden (2008).
- The steps used to create this GRM is:
 - Create a center of marker data (X matrix)
 - Create a Cross Product (XX)
 - Divide the (XX) by number of markers

$$GRM = XX^t/m$$

- More on relationship matrix can be found here Source 1, Source 2
- We will use the AGHmatrix package to build G matrix.

```
> GM<- Gmatrix(SNPmatrix=geno, missingValue=NA,
                             maf=0.05, method="VanRaden")
Initial data:
    Number of Individuals: 252
    Number of Markers: 396511
Missing data check:
    Total SNPs: 396511
     O SNPs dropped due to missing data threshold of 0.5
    Total of: 396511 SNPs
MAF check:
     24891 SNPs dropped with MAF below 0.05
    Total: 371620 SNPs
Heterozigosity data check:
    No SNPs with heterozygosity, missing threshold of = 0
Summary check:
    Initial: 396511 SNPs
    Final: 371620 SNPs (24891 SNPs removed)
Completed! Time = 31.512 seconds
    dim(GM)
```

Fit Various G x E models

The more description on G x E models I recommend going over these resources Resource 1 and Resource 2

Main MET Model

Here we will fit model using sommer R package.

The model assumes GxE doesn't exist and that the main genotype effect plus the fixed effect for environment is enough to predict the genotype effect in all locations.

```
> # Fit Model
   gs.model1<- mmes(BLUEs~Environment, # Environment Fixed
                  random= ~ vsm(ism(Genotype), Gu=GM), #usm is covariance function to assign matrices
                  rcov= ~ units, # Residuals have no-covariance
                  data=BLUEs.all, verbose = FALSE)
> # Get summary
   summary(gs.model1)
        Multivariate Linear Mixed Model fit by REML
*************** sommer 4.4 *************
        logLik
                   AIC
                           BIC Method Converge
Value -114.9267 239.8534 265.5438
                                          TRUE
Variance-Covariance components:
                VarComp VarCompSE Zratio Constraint
Genotype:GM:mu:mu
                  20735
                            7139 2.905 Positive
units:mu:mu
                            22767 23.505
                 535135
                                         Positive
_____
Fixed effects:
        Estimate Std.Error t.value
                   46.18 50.146
Intercept 2315.7
ENV2
         -1537.6
                     65.24 -23.570
ENV3
           433.7
                     65.24 6.648
ENV4
           525.7
                     65.24
                           8.058
           780.3
                     65.24 11.960
ENV5
Use the '$' sign to access results and parameters
> # Extract the GEBVs (random effects)
   estimated.all<-data.frame(GEBVs= gs.model1$u) # stored in u
   estimated.all$GEBVs<-estimated.all$GEBVs+ gs.model1$b[1] # adding intercept
   gs.model1$AIC # Check AIC and BIC values
[1] 239.8534
   gs.model1$BIC
[1] 265.5438
> kable(head(estimated.all)) # View as table
                                              GEBVs
```

Genotype_10162 2051.910

	GEBVs
Genotype_10164	2431.821
Genotype_10169	2157.124
Genotype_10173	2057.640
Genotype_10175	2124.616
Genotype_10176	2174.124

MET: diagonal model (DG)

The diagonal model assumes that GxE exists and that the genotype variation is expressed differently at each location, therefore fitting a variance component for the genotype effect at each location. The main drawback is that this model assumes no covariance among locations, as if genotypes were independent.

```
gs.model3<- mmes(BLUEs~Environment,
+
             random= ~ vsm(dsm(Environment),ism(Genotype), Gu=GM),
+
             rcov= ~ units,
             data=BLUEs.all, verbose = FALSE)
   summary(gs.model3)
______
         Multivariate Linear Mixed Model fit by REML
_____
      logLik
               AIC
                      BIC Method Converge
Value -20.67411 51.34823 77.03859
                            ΑI
                                  TRUE
______
Variance-Covariance components:
                         VarComp VarCompSE Zratio Constraint
                          133567
Environment:Genotype:GM:ENV1:ENV1
                                   31523 4.237
                                              Positive
Environment:Genotype:GM:ENV2:ENV2
                                   7265 0.000
                                              Positive
Environment: Genotype: GM: ENV3: ENV3
                                   39035 4.976
                          194228
                                              Positive
Environment:Genotype:GM:ENV4:ENV4
                          127188
                                   30619 4.154
                                              Positive
Environment:Genotype:GM:ENV5:ENV5
                          398478
                                   61539 6.475
                                              Positive
units:mu:mu
                          286935
                                   20286 14.144
                                              Positive
Fixed effects:
       Estimate Std.Error t.value
Intercept
        2317.2
               33.86 68.432
ENV2
        -1539.1
                 47.80 -32.197
ENV3
         432.2
                 47.80
                       9.042
ENV4
         524.2
                 47.80 10.965
ENV5
         778.8
                 47.80 16.291
______
```

Use the '\$' sign to access results and parameters

MET: unstructured model (US)

We assume that that GxE exists and that an environment-specific variance exists in addition to as many covariances for each environment-to-environment combinations. The main drawback is that is difficult to make this models converge because of the large number of variance components, the fact that some of these variance or covariance components are zero, and the difficulty in choosing good starting values. The fixed effect for environment plus the environment specific BLUP (adjusted by covariances) is used to predict the genotype effect in each location of interest.

MET: compund symmetry model (CS)

The compound symmetry model assumes that GxE exists and that a main genotype variance-covariance component is expressed across all location. In addition, it assumes that a main genotype-by-environment variance is expressed across all locations. The main drawback is that the model assumes the same variance and covariance among locations.

```
E <- diag(length(unique(BLUEs.all$Environment)))</pre>
      rownames(E) <- colnames(E) <- unique(BLUEs.all$Environment)</pre>
>
>
      Ei <- solve(E)
      Gi <- solve(GM)</pre>
      EGi <- kronecker(Ei,Gi, make.dimnames = TRUE)
      Ei <- as(as(as( Ei, "dMatrix"), "generalMatrix"), "CsparseMatrix")
      Gi <- as(as(as( Gi, "dMatrix"), "generalMatrix"), "CsparseMatrix")</pre>
      EGi <- as(as(as( EGi, "dMatrix"), "generalMatrix"), "CsparseMatrix")
      attr(Gi, "inverse")=TRUE
>
      attr(EGi, "inverse")=TRUE
      model5<- mmes(BLUEs~Environment,</pre>
                     random= ~ vsm(ism(Genotype), Gu=Gi) + vsm(ism(Environment:Genotype), Gu=EGi),
+
                     rcov= ~ units,
                     data=BLUEs.all, verbose = FALSE)
+
      summary(gs.model5)
```

Fit Same Model in BGLR

Here again we will explain in detail, y: vector of phenotypic values (Yield), BLUES.all is a dataframe with columns Genotype, Environment, and Yield, Z is genotype incidence matrix GM is genomic relationship matrix (e.g., from markers) and X is environment design matrix. Before fitting model is BGLR we will build X and Z matrices, then we will create the Kernal Matrices of Genomic matrix and interaction matrices.

Then we will build ETA list to supply the matrices to function.

```
> # Convert Environment to design matrix
> X <- model.matrix(~ Environment, data = BLUEs.all)
> # Convert Genotype to design matrix
> Z <- model.matrix(~ Genotype - 1, data = BLUEs.all)
> # Genotype kernel
> KG <- tcrossprod(Z %*% GM)
> # GxE interaction kernel
> KGE <- tcrossprod(X) * KG # Element-wise multiplication
>

# ETA list for BGLR Package
> ETA <- list(
+ ENV = list(X = X, model = "FIXED"),
+ G = list(K = KG, model = "RKHS"),
+ GxE = list(K = KGE, model = "RKHS")
+ )
> # Fit the Model
```

```
> modelGxE<- BGLR(y = BLUEs.all$BLUEs,
+ ETA = ETA,
+ nIter = 1000,
+ burnIn = 100,
+ thin = 2,
+ verbose = FALSE)</pre>
```

Extract the Results

```
> # Variance components
> modelGxE$ETA$G$varU  # Genotype variance

[1] 2440.601
> modelGxE$ETA$GxE$varU  # GxE variance

[1] 2721.165
> modelGxE$varE  # Residual variance

[1] 570511.8
> # Breeding values (genotype effects)
> GEBVs <- data.frame(GEBVs_All=modelGxE$ETA$G$u)
> #BGLR directly outputs the genotype predictions as yHat
> GEBVs2<- data.frame(modelGxE$yHat)</pre>
```

 $Note:\ For\ questions\ specific\ to\ data\ analysis\ shown\ here\ contact\ waseem. hussain@irri.org$

If your experiment needs a statistician, you need a better experiment - Ernest Rutherford

For any suggestions or comments, please feel to reach at waseem.hussain@irri.org; and m.anumalla@irri.org