The RepeatABEL package - a tutorial

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Introduction

This vignette gives an introduction to the RepeatABEL package. The package performs GWAS for data where there are repeated observations on individuals that may be related. Various random effects can be fitted. Random polygenic effects are fitted by default, but also other random effects can be fitted including spatial effects. A model without the fixed SNP effect is fitted using the hglm package for estimating variance components (see Ronnegard, Shen & Alam (2010) hglm: A package for fitting hierarchical generalized linear models. The R Journal 2:20-28). These variance component estimates are subsequently used in the GWAS where each marker is fitted one at a time. Thus, this is similar to using the polygenic_hglm function and subsequently the mmscore function in GenABEL. The package consists of three main functions rGLS, preFitModel and simulate_PhenData.

Theory for the rGLS function

The rGLS function is the main function of the package and by default fits a linear mixed model including permanent environmental effects p and polygenic effects q with correlation matrix given by the genomic relationship matrix (aka kinship matrix) in

$$y = X\beta + Zg + Zp + e \tag{1}$$

where \boldsymbol{y} is the studied trait, $\boldsymbol{\beta}$ are fixed effects, \boldsymbol{e} are residuals with $\boldsymbol{e} \sim N(0, \sigma_e^2)$ having residual variance σ_e^2 . Furthermore, \boldsymbol{X} and \boldsymbol{Z} are model matrices. The random effects are assumed multivariate normal such that $\boldsymbol{p} \sim N(0, \boldsymbol{I}\sigma_p^2)$ and $\boldsymbol{g} \sim N(0, \boldsymbol{G}\sigma_g^2)$ where \boldsymbol{G} is the genomic relationship matrix. Thus the estimated (co)variance matrix for this model is

$$\hat{\mathbf{V}} = \mathbf{Z}\mathbf{G}\mathbf{Z}^T\hat{\sigma}_q^2 + \mathbf{Z}\mathbf{Z}^T\hat{\sigma}_p^2 + \mathbf{I}\hat{\sigma}_e^2.$$
 (2)

Subsequently, a linear model is fitted (using generalized least squares, GLS) for each marker where the covariate x_{SNP} is coded as 0,1,2:

$$y = X\beta + x_{SNP}\beta + \varepsilon \tag{3}$$

with

$$\varepsilon \sim N(0, \sigma_{\epsilon}^2 \hat{V}).$$
 (4)

A Wald test is used to compute the P value for SNP effect β .

The computations are made fast by applying a eigen-decomposition of \hat{V} and using the built-in qr function in R to fit the linear models (3).

Using the rGLS function

The following example illustrates the use of the function. There are two data objects to be included in the input: a GenABEL object including the genotypic information and a data frame including the phenotypic information. The name of the ID variable in the phenotype data frame should be "id" (otherwise specify id.name equal to the ID variable name).

In this example there are 360 observations from 100 individuals, and there are 5792 SNP to be tested.

- > library(RepeatABEL)
- > #GenABEL object including IDs and marker genotypes
- > data(gen.data)
- > #Phenotype data with repeated observations
- > data(Phen.Data)

The data frame Phen.Data includes the trait value y, two covariates (age and sex) and the ID of the individuals. We wish to include age and sex as fixed effects so the function input is

```
> GWAS1 <- rGLS(y ~ age + sex, genabel.data = gen.data,
   phenotype.data = Phen.Data)
[1] "GRM ready"
[1] "Variance component estimation ready"
```

- [1] "Rotation matrix ready"
- [1] "Rotate LMM started"
- [1] "Rotate LMM ready"

The computations in this function consists of four parts: construction of the GRM, variance component estimation using the hglm function, rotating the GLS using eigen-decomposition transforming it to an ordinary least squares (OLS) problem, and finally fitting an OLS for each SNP. How far the computations have come is shown in the output.

The class of the output object GWAS1 is gwaa.scan, so we can apply the generic functions summary() and plot() defined by the GenABEL package.

```
> summary(GWAS1)
Summary for top 10 results, sorted by P1df
          Chromosome Position Strand A1 A2
                                                 effB
rs120315
                 1 3725352 + T C 2.1696316
                                   - C A -2.0412127
rs9670687
                 1 4779800
rs891586
                 2 8024318
                                  + A G -1.6669267
                 2 8022651
                                   - A C 1.6169429
rs1352451
                 2 8167984
                                   - A T -0.9982089
rs1252282
                 1 3721952
                                   + C T 1.8586037
rs7633966
                 1 3729983
rs9922492
                                  + C T -1.8586037
rs4378234
                  1 4800348
                                   - G T -1.9012073
rs7499832
                 3 10242953
                                   - G C -1.0897248
                                   - T G -1.9073305
rs6561272
                  1 3715538
           se_effB chi2.1df
rs120315 0.4258979 NA 3.501204e-07
                                             NΑ
rs9670687 0.5351582
                        NA 1.366120e-04
rs891586 0.4478499 NA 1.975997e-04
rs1352451 0.4564078 NA 3.959643e-04
rs1252282 0.2825403 NA 4.109058e-04
                                             NA
                                             NA
                                             NA
rs7633966 0.5325691
                        NA 4.832322e-04
                                             NΑ
```

```
rs9922492 0.5325691 NA 4.832322e-04 NA rs4378234 0.5509388 NA 5.588239e-04 NA rs7499832 0.3182487 NA 6.167705e-04 NA rs6561272 0.5726170 NA 8.656541e-04 NA
```

Extracting the genotypic variance, a 95% CI and the heritability

From the GWAS1 object, the estimated variance components for the prefitted model, not including SNP effects, can be extracted as follows.

Furthermore, the standard errors for the estimated variance components are computed on a log scale. Thereby, 95% confidence intervals can be computed for the variance components. Here the computations for the genotypic variance are shown.

The heritability for this example is computed as:

Using the preFitModel function

The function preFitModel is used for variance component estimation and increases the flexibility of modeling in the rGLS function.

Fitting the same model as above

To start with we have a look at how the model in the previous section can be fitted in two steps using the preFitModel function and thereafter the rGLS function. Note that the results are exactly the same as in the previous section.

```
> #The same results can be computed using the preFitModel as follows
> fixed=y ~ age + sex
> Mod1 <- preFitModel(fixed, random=~1|id,
     genabel.data = gen.data,
     phenotype.data = Phen.Data,
     corStruc=list( id=list("GRM","Ind") ))
[1] "GRM ready"
> GWAS1b <- rGLS(fixed, genabel.data = gen.data,
    phenotype.data = Phen.Data, V = Mod1$V)
[1] "Rotation matrix ready"
[1] "Rotate LMM started"
[1] "Rotate LMM ready"
> summary(GWAS1b)
Summary for top 10 results, sorted by P1df
         Chromosome Position Strand A1 A2
                                               effB
                                + T C
                 1 3725352
rs120315
                                         2.1696316
rs9670687
                 1 4779800
                                 - C A -2.0412127
rs891586
                 2 8024318
                                 + A G -1.6669267
                                 - A C 1.6169429
                2 8022651
rs1352451
                                 - A T -0.9982089
rs1252282
                2 8167984
rs7633966
                1 3721952
                                 + C T 1.8586037
rs9922492
                 1 3729983
                                 + C T -1.8586037
                 1 4800348
                                 - G T -1.9012073
rs4378234
                                 - G C -1.0897248
rs7499832
                 3 10242953
rs6561272
                 1 3715538
                                   T G -1.9073305
           se_effB chi2.1df
                                   P1df Pc1df
rs120315 0.4258979
                        NA 3.501204e-07
rs9670687 0.5351582
                        NA 1.366120e-04
                                           NA
rs891586 0.4478499
                      NA 1.975997e-04
                                           NA
rs1352451 0.4564078
                       NA 3.959643e-04
                                           NΑ
                       NA 4.109058e-04
rs1252282 0.2825403
                                           NA
rs7633966 0.5325691
                        NA 4.832322e-04
                                           NA
rs9922492 0.5325691
                        NA 4.832322e-04
                                           NΑ
rs4378234 0.5509388
                        NA 5.588239e-04
                                           NΑ
rs7499832 0.3182487
                        NA 6.167705e-04
                                           NA
rs6561272 0.5726170
                        NA 8.656541e-04
                                           NA
```

The only information transferred from the preFitModel function to rGLS is the estimated (co)variance matrix Mod1\$V. The corStruc option specifies the correlation structure to be applied on each random effect. In the example we wish to fit polygenic effects and permanent environmental effects. The former requires a correlation structure given by the GRM whereas the permanent environmental effects are iid. Consequently, corStruc=list(id=list("GRM", "Ind")

)) is specfied.

A model having several different random effects

In this (fake) example there are 60 observations on each nest and there are 6 different nests. We wish to include these as random effect too, and to start with they are modelled as independent random effects.

Spatial modelling

This example shows how random effects having a spatial correlation structure can be included to account for populaton structure. The spatial model used is a Conditional AutoRegressive (CAR) model and the input spatial information is given by a neighborhood matrix. (A neighborhood matrix has a non-zero value for an element (i,j) where the subject (nest in our example) i and j come from neighboring locations. The diagonal elements are zero.) Here the neighborhood matrix (D) is a 6×6 matrix

```
> D= matrix(0,6,6)

> D[1,2] = D[2,1] = 1

> D[5,6] = D[6,5] = 1

> D[2,4] = D[4,2] = 1

> D[3,5] = D[5,3] = 1

> D[1,6] = D[6,1] = 1

> D[3,4] = D[4,3] = 1
```

where for instance nests 1 and 2 are defined as neighbors. The model including polygenic effects, permanent environmental effects and a spatial correlation between nests is then fitted as

Using the simulate_PhenData function

The third function included in the RepeatABEL package is a simulation function where one can simulate phenotypic data having repeated observations. The input genotype information is given by a GenABEL object.

Suppose for instance we want to simulate 4 observations from each individual and the three variance components (polygenic, permanent env., residual) are 1.

```
> VC.poly <- VC.perm <- VC.res <- 1
> n.obs <- rep(4, nids(gen.data))</pre>
```

In this example, the GenABEL object gen.data is used as input and an additive genetic effect of 2.0 is simulated at the location of SNP number 1000. The phenotype data is then simulated as

```
> Phen.Sim <- simulate_PhenData(y ~ 1,
    genabel.data = gen.data,
    n.obs = n.obs, SNP.eff = 2, SNP.nr = 1000,
    VC = c(VC.poly, VC.perm, VC.res))</pre>
```

The simulated data can then be fitted as

```
> GWAS.sim1 <- rGLS(y ~ 1, genabel.data = gen.data,
    phenotype.data = Phen.Sim)</pre>
```

The data set gen.data includes information on sex, so we can also add a fixed sex effect to our simulations. Here a sex effect of 1.0 is simulated

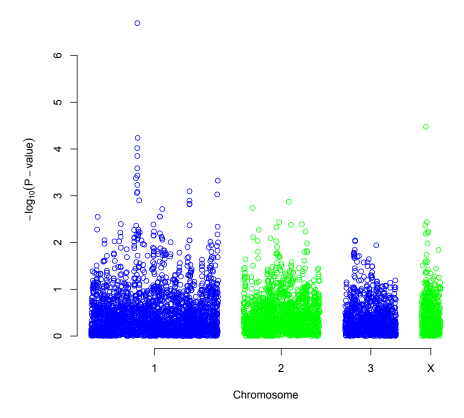
```
> Phen.Sim <- simulate_PhenData(y ~ sex,
    genabel.data = gen.data, n.obs = n.obs, SNP.eff = 2,
    SNP.nr = 1000, VC = c(VC.poly, VC.perm, VC.res), beta = c(0,1))</pre>
```

where beta is a vector specifying the simulated values of the fixed effects and since we fit an intercept and a sex effect the length of beta is two. The data can then be fitted as

```
> GWAS.sim1 <- rGLS(y ~ sex, genabel.data = gen.data,
     phenotype.data = Phen.Sim)
> plot(GWAS.sim1, main="Simulation results")
```

The produced Manhattan plot is given below

Simulation results



Simulating year effects using the simulate_PhenData function

Year effects can be added to the simulated response after running the simulate_PhenData function.

```
> Phen.Sim <- simulate_PhenData(y ~ sex,
    genabel.data = gen.data, n.obs = n.obs, SNP.eff = 2,
    SNP.nr = 1000, VC = c(VC.poly, VC.perm, VC.res), beta = c(0,1))
> ###### PRODUCE YEAR EFFECTS FOR EACH INDIVIDUAL
> sd.year = 1 #Standard Deviation of Year Effects
> beta <- rnorm(max(n.obs), 0, sd.year) #Simulated Year Effects
> year.effects <- years <- NULL
> for (i in 1:length(n.obs)) {
      yr.i <- sort(sample(1:max(n.obs), n.obs[i]))</pre>
      years <- c(years, yr.i)</pre>
            year.effects <- c(year.effects, beta[yr.i])</pre>
    }
> #######################
> #### A FUNCTION TO ADD A VARIABLE TO A LIST
> add.var <- function(x, add.new, new.name) {</pre>
      x[[length(x) + 1]] \leftarrow add.new
            names(x)[length(x)] <- new.name</pre>
            return(x)
    }
> #######################
> #ADDS A NEW PHENOTYPE WITH YEAR EFFECTS ADDED
> Phen.Sim <- add.var(Phen.Sim, Phen.Sim$y + year.effects, "y.yrs")
> #ADD YEAR AS FACTOR
> Phen.Sim <- add.var(Phen.Sim, as.factor(years), "Years")
> ######################
> #RUN THE ANALYSIS
> GWAS.sim1 <- rGLS(y.yrs ~ sex + Years, genabel.data = gen.data,
    phenotype.data = Phen.Sim)
```

Simulating binary data using the simulate_PhenData function

Binary data can be simulated using a threshold model where a Gaussian response is first simulated and all values greater than a specified threshold τ are 1's on the observed scale. The binary data can be simulated and analyzed using the following code.