Supplementary material for "lme4qtl: linear mixed models with flexible covariance structure for genetic studies of related individuals"

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Supplementary Tables and Figures

Supplementary Table 1

Feature	lme4qtl	SOLAR	ASReml	GEMMA
Covariance for random effects	1	✓	1	1
Covariance for residuals	(1)	✓	1	X
Methods for sparse covariances	✓	X	✓	X
Methods for dense covariances	(2)	✓	✓	✓
More than one covariances	✓	✓	✓	X
Restriction on parameters	✓	X	×	X
Gene-by-environment design	✓	(3)	✓	×
Longitudinal design	✓	(3)	✓	×
Free software	✓	✓	X	✓
Open source	✓	X	×	✓

Table 1: Comparison among lme4qtl and selected stand-alone tools for genetic association analysis: SOLAR [1], ASReml [2] and GEMMA [3]. Notes: (1) The lme4qtl packages does not support any structures of residual variance, as the lme4 package does not has this feature yet. However, we showed two *ad hoc* solutions in Supplementary Note 3. (2) The lme4qtl package is based on sparse matrix methods from the lme4 package. In principle, dense matrix operations are still possible, but that might lead to considerable overhead in computation resources, as presented in Supplementary Figure 4 and discussed in the main text. (3) SOLAR requires specific tcl scripts (not publicly available) to parametrize either gene-by-environment or longitudinal models.

Supplementary Table 2

Feature	lme4qtl	pedigreemm	lmekin	Gaston	regress	rrBLUP
Extension of lme4	✓	✓	Х	Х	Х	Х
Covariance for random effects	✓	(1)	✓	✓	✓	✓
Covariance for residuals	(2)	(2)	✓	X	X	X
Methods for sparse covariances	✓	✓	✓	X	X	X
Methods for dense covariances	(3)	(3)	✓	✓	✓	✓
More than one covariances	✓	✓	✓	✓	✓	X
Restriction on parameters	✓	×	X	×	X	X

Table 2: Comparison among lme4qtl and other selected R packages that implement linear mixed models and can be used in genetic studies: pedigreemm [4] that extends lme4 [5], lmekin function in the R package coxme [6], Gaston [7], regress [8] and rrBLUP [9]. Notes: (1) The pedigreemm package does not support custom covariances, but allows to define relationship matrices based on the pedigree information. (2) Both lme4qtl and pedigreemm packages do not support any structures of residual variance, as the lme4 package does not has this feature yet. However, we showed two *ad hoc* solutions in Supplementary Notes 2 and 3. (3) Both lme4qtl and pedigreemm packages are based on sparse matrix methods from the lme4 package. In principle, dense matrix operations are still possible, but that might lead to considerable overhead in computation resources, as presented in Supplementary Figure 4 and discussed in the main text.

Supplementary Table 3

Model	Fast SOLAR, hours	SOLAR, hours	<i>lme4qtl,</i> hours
aptt \sim age + sex + (1 id)	3.8	16.8	6.6
aptt \sim age + sex + (1 hhid) + (1 id)	_	25.1	7.6
aptt \sim age + sex + (1 hhid) + (1 id) + (1 id7)	_	27.9	23.3

Table 3: We performed several genome-wide screenings of the activated partial thromboplastin time (APTT) phenotype in the GAIT2 data [10]. We considered three types of models and compared the computation time between our software lme4qtl and SOLAR [1]. The three models differed in the number of random effects: a single genetic additive effect (expressed in the model formula as (1|id)); two house-hold and genetic additive effect ((1|hhid) + (1|id)); and three house-hold, genetic additive and dominance effects ((1|hhid) + (1|id) + (1|id7)). The GAIT2 study included 903 individuals (those with measured values of APTT) in 35 extended families. The number of tested genetic markers consisted of 263,764 SNPs and indels, which passed the minimum allele frequency threshold of 1%. The analysis was performed on a desktop computer (2.8GHz quad-core Intel Core i5 processor, 8GB RAM).

Supplementary Figures

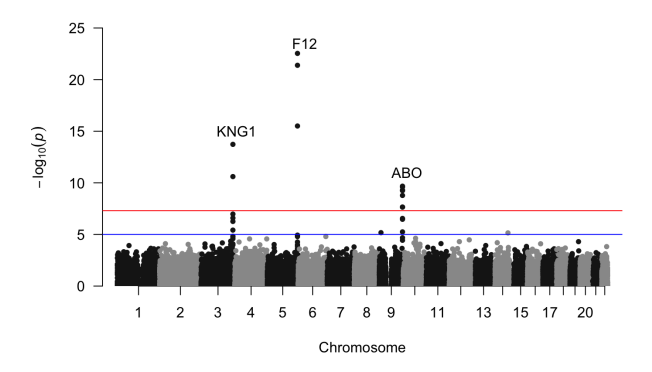


Figure 1: Genome-wide association study of APTT in the GAIT2 data computed by the *lme4qtl* R package partially replicates previously reported loci in a larger cohort of 9,240 individuals [11]. Three loci, in genes *KNG1*, *F12* and *ABO*, passed the genome-wide significant threshold at 5×10^{-8} , depicted as red horizontal line on the plot.

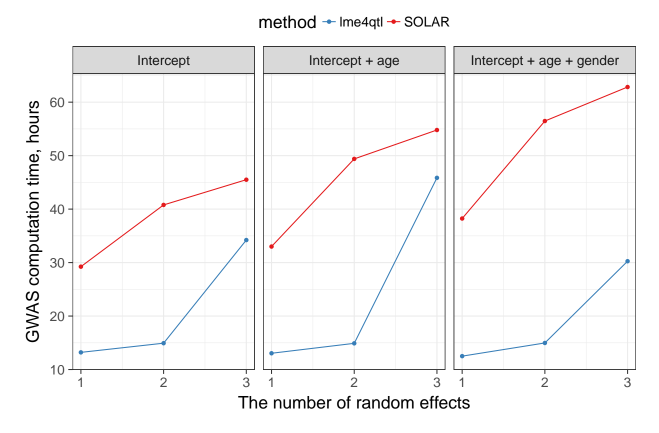


Figure 2: The plot represents the computation times reported in Table 3 on right panel. Other left and central panels show the results for the same experiment as in Table 3, but the list of fixed effects is less, either one (the intercept) or two (the intercept and age).

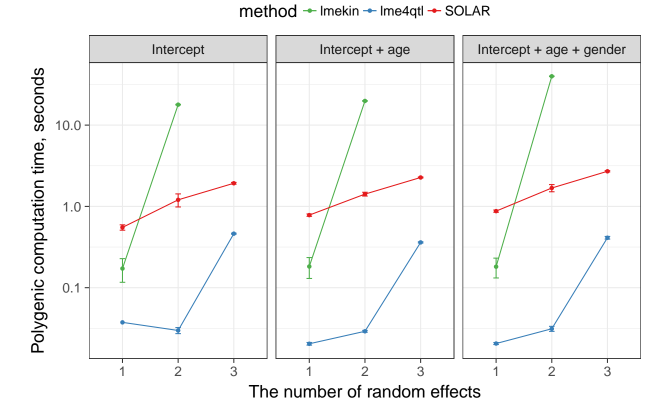


Figure 3: Comparison in computation time among three tools – our software *lme4qtl*, SOLAR [1] and *lmekin* [6] – showed the fastest performance of *lme4qtl*. We fitted a polygenic model of the activated partial thromboplastin time (APTT) phenotype measured in the GAIT2 data [10]. Models were different in the number of random effects: a single genetic additive effect (expressed in the model formula as (1|id)); two house-hold and genetic additive effect ((1|hhid) + (1|id)); and three house-hold, genetic additive and dominance effects ((1|hhid) + (1|id) + (1|id7)). Models also were different in the number of fixed effects (covariates): a single covariate (Intercept), two covariates (Intercept + age) and three covariates (Intercept + age + gender). The GAIT2 study included 903 individuals (those with measured values of APTT) in 35 extended families. The analysis was performed on a desktop computer (2.8GHz quad-core Intel Core i5 processor, 8GB RAM). We repeated each measurement of computational time 10 times and reported the mean value and its standard error in the figure. The numbers on y axis are base-10 log scaled.

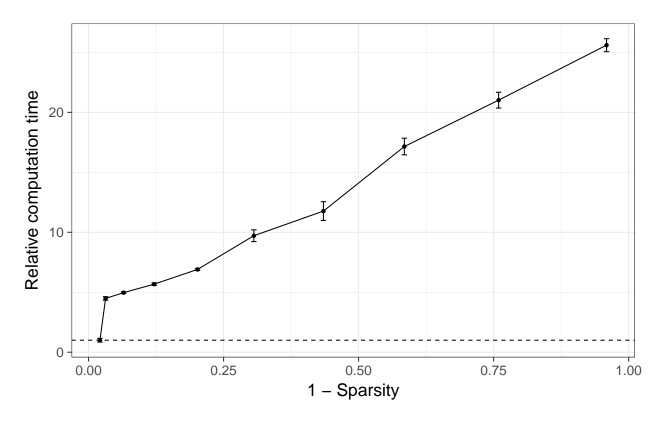


Figure 4: The computation time of polygenic model (devided by the number of interations in the optimization algorithm) fitted by lme4qtl increases as the sparsity of the genetic relatedness matrix is reduced, since the lme4 machinery is optimized for linear algebra operations on sparse matrices. The sparsity is measured as the proportion of zero entries in the relatedness matrix (mat). Point on the plot with the lowest sparsity corresponds to a model fitted with the original GAIT2 genetic additive relatedness matrix. The dashed line marks the reference computation time. Other points come from models fitted with modified matrices varying their sparsity. The polygenic model is estimated for the activated partial thromboplastin time (APTT) measured in 903 individuals from the family-based GAIT2 study. The R code used for computation was relmatLmer(aptt \sim age + gender + (1|id), dat, relmat = list(ID = mat)). The analysis was performed on a desktop computer (2.8GHz quad-core Intel Core i5 processor, 8GB RAM). We repeated each measurement of computational time 10 times and reported the mean value and its standard error in the figure.

References

- [1] L Almasy and J Blangero. Multipoint quantitative-trait linkage analysis in general pedigrees. *American journal of human genetics*, 62(5):1198–211, May 1998.
- [2] Arthur R Gilmour, BJ Gogel, BR Cullis, R Thompson, D Butler, et al. *ASReml user guide release* 3.0, 2009. VSN International Ltd, Hemel Hempstead, UK.
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- [9] Jeffrey B Endelman. Ridge regression and other kernels for genomic selection with r package rrblup. *The Plant Genome*, 4(3):250–255, 2011.
- [10] Laura Martin-Fernandez, Andrey Ziyatdinov, Marina Carrasco, Juan Antonio Millon, Angel Martinez-Perez, Noelia Vilalta, Helena Brunel, Montserrat Font, Anders Hamsten, Juan Carlos Souto, et al. Genetic determinants of thrombin generation and their relation to venous thrombosis: results from the GAIT-2 project". *PloS one*, 11(1):e0146922, 2016.
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Supplementary Note 1: R code to compare lme4qtl and pedigreemm R packages

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About

The R package pedigreemm was first in extending the lme4 R package for particular applications in the animal breeding field (Vazquez et al. 2010). Custom covariance (genetic additive) matrix are defined using the pedigree annotation information (pedigree argument of pedigreemm function). Although the lme4qtl package borrows the same idea of pedigreemm, lme4qtl provides a larger list of genetic models that are not possible with pedigreemm. In particular, these models include:

- models with a single or several custom covariances (not necessary linked to pedigree information);
- models with random slopes and other similar models like gene-by-environment interaction models;
 the restriction on model parameters, e.g. the correlation coefficient is zero, is supported.

Here, we show models that are available with lme4qtl and not with pedigreemm.

Include

First, we load R packages necessary for data analysis.

```
library(Matrix)
library(magrittr)

library(pedigreemm)

library(lme4qtl)
```

Data

We use an example data set milk from the pedigreemm package. See ?milk for description.

Here, we work on a subset of this dataset (milk_subset) to reduce the computation time.

```
data(milk)
milk <- within(milk, {
   id <- as.character(id)
      sdMilk <- milk / sd(milk)
})

ids <- with(milk, id[sire %in% 1:3]) # for more ids: c(1:3, 319-321)
milk_subset <- subset(milk, id %in% ids)

milk_subset <- within(milk_subset, {
   herd <- droplevels(herd)
   herd_id <- paste0("herd", seq(1, length(herd)))
})</pre>
```

Covariance matrices

A mixed model we are going to fit will have two random effects, groupings based on two ID variables:

- id, a numeric identifier of cow (the genetic additive effect);
- herd, a factor indicating the herd (the shared environmental effect).

Further we derive the covariance matrices (among samples) due to these two random effects.

```
A_herd <- with(milk_subset, model.matrix(~ herd - 1)) %>%
    tcrossprod %>% Matrix

rownames(A_herd) <- milk_subset$herd_id

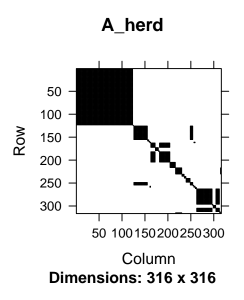
colnames(A_herd) <- milk_subset$herd_id

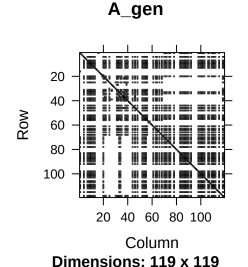
A_gen <- getA(pedCowsR)

stopifnot(all(ids %in% rownames(A_gen)))
ind <- rownames(A_gen) %in% ids

A_gen <- A_gen[ind, ind]

image(A_herd, main = "A_herd")
image(A_gen, main = "A_gen")</pre>
```





Models

id

herd

Residual

A single kinship matrix

Both packages can fit a basic model with a single genetic effect, for which the pedigreemm R package was sought.

```
m1_pmm <- pedigreemm(sdMilk ~ lact + log(dim) + (1|id) + (1|herd),
  data = milk_subset, pedigree = list(id = pedCowsR))
VarCorr(m1_pmm)
Groups
                      Std.Dev.
 id
          (Intercept) 0.55436
herd
          (Intercept) 0.55630
Residual
                      0.59894
m1_relmat <- relmatLmer(sdMilk ~ lact + log(dim) + (1|id) + (1|herd),
  data = milk_subset, relmat = list(id = A_gen))
VarCorr(m1_relmat)
Groups
          Name
                      Std.Dev.
```

We see that the estimation of variance components from both packages are identical.

A single custom covariance matrix

(Intercept) 0.55436

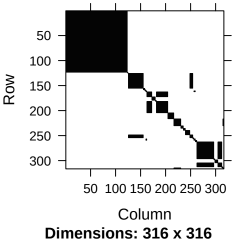
(Intercept) 0.55630

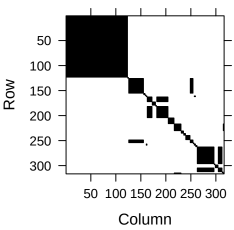
0.59894

lme4qtl packages allows for custom covariance matrices, while pedigreemm does not.

```
m2_lmer <- lmer(sdMilk ~ (1|herd), milk_subset)
VarCorr(m2_lmer)</pre>
```

```
Groups
          Name
                      Std.Dev.
herd
          (Intercept) 0.54833
Residual
                      0.81060
m2_relmat <- relmatLmer(sdMilk ~ (1|herd_id), milk_subset,</pre>
  relmat = list(herd_id = A_herd))
VarCorr(m2_relmat)
Groups
                      Std.Dev.
          Name
herd_id (Intercept) 0.54833
Residual
                      0.81060
(try(m2_pmm <- pedigreemm(sdMilk ~ (1|herd_id), milk_subset,</pre>
  pedigree = list(herd_id = A_herd))))
[1] "Error : all(sapply(pedigree, is, class2 = \"pedigree\")) is not TRUE\n"
attr(,"class")
[1] "try-error"
attr(,"condition")
<simpleError: all(sapply(pedigree, is, class2 = "pedigree")) is not TRUE>
getME(m2 lmer, "Ztlist")[[1]] %>% crossprod %>% image
getME(m2_relmat, "Ztlist")[[1]] %>% crossprod %>% image
```





Dimensions: 316 x 316

Rank deficiency

A_herd is a low-rank matrix, but lme4qtl is able to deal with this rank deficiency situation by replacing the Cholesky decomposition by the EVD operation. The pedigreemm package only uses the Cholesky decomposition.

```
A_herd %>% dim
[1] 316 316
A_herd %>% as.matrix %>% qr %$% rank
```

[1] 21

A single kinship matrix + random slope

```
Complex models are possible with lme4qtl, for example, those with a random slope effect.
```

```
m3_relmat <- relmatLmer(sdMilk ~ lact + log(dim) + (1 + lact | id) + (1 | herd),
  data = subset(milk_subset, relmat = list(id = A_gen)))
VarCorr(m3_relmat)
Groups
          Name
                      Std.Dev. Corr
          (Intercept) 0.400268
 id
                      0.094301 0.489
          (Intercept) 0.593480
herd
Residual
                      0.585815
(try(m3_pmm <- pedigreemm(sdMilk ~ lact + log(dim) + (1 + lact|id) + (1|herd),
 data = milk_subset, pedigree = list(id = pedCowsR))))
[1] "Error in `levels<-`(`*tmp*`, value = if (nl == nL) as.character(labels) else pasteO(labels, : \n
attr(,"class")
[1] "try-error"
attr(,"condition")
<simpleError in `levels<-`(`*tmp*`, value = if (nl == nL) as.character(labels) else pasteO(labels,</pre>
```

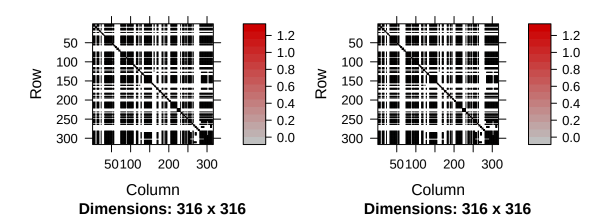
Restriction on model parameters

Two covariance matrices

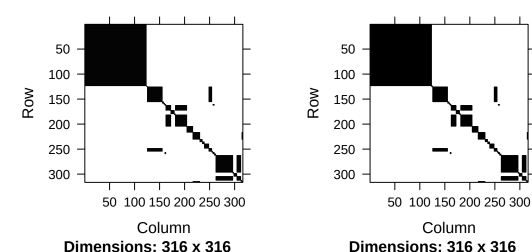
```
m5 <- relmatLmer(sdMilk ~ (1|herd) + (1|id), milk_subset,
 relmat = list(id = A_gen))
VarCorr(m5)
Groups
                      Std.Dev.
id
          (Intercept) 0.54054
herd
          (Intercept) 0.54286
Residual
                      0.64997
m6 <- relmatLmer(sdMilk ~ (1|herd_id) + (1|id), milk_subset,
 relmat = list(herd_id = A_herd, id = A_gen))
VarCorr(m6)
                      Std.Dev.
Groups
          Name
herd_id (Intercept) 0.54286
 id
          (Intercept) 0.54054
```

```
Residual 0.64997

getME(m5, "Ztlist")[[1]] %>% crossprod %>% image
getME(m6, "Ztlist")[[2]] %>% crossprod %>% image
```



getME(m5, "Ztlist")[[2]] %>% crossprod %>% image
getME(m6, "Ztlist")[[1]] %>% crossprod %>% image



```
(try(m3 <- pedigreemm(sdMilk ~ lact + log(dim) + (1|id) + (1|herd_id),
  data = milk_subset, pedigree = list(id = pedCowsR, herd_id = A_herd))))</pre>
```

```
[1] "Error : all(sapply(pedigree, is, class2 = \"pedigree\")) is not TRUE\n"
attr(,"class")
[1] "try-error"
attr(,"condition")
<simpleError: all(sapply(pedigree, is, class2 = "pedigree")) is not TRUE>
```

References

Vazquez, AI, DM Bates, GJM Rosa, D Gianola, and KA Weigel. 2010. "Technical Note: An R Package for Fitting Generalized Linear Mixed Models in Animal Breeding." *Journal of Animal Science* 88 (2). American Society of Animal Science: 497–504.

Supplementary Note 2: Multi-trait and multi-environment linear mixed models

We consider a simple polygenic model with two random effects, the additive genetic effect and the residual errors (also referred to as environment effect).

Single-trait linear mixed model

A linear model describes observations of a trait, measured in n individuals and stored in a vector $y_{n\times 1}$.

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

where $X_{n\times p}$ and $Z_{n\times n}$ are incidence matrices, p is the number of fixed effects, $\beta_{p\times 1}$ is a vector of fixed effects, $u_{n\times 1}$ is a vector of a random polygenic effect, and $e_{n\times 1}$ is a vector of the residuals errors. The random vectors u and e are mutually uncorrelated and multivariate normally distributed, $\mathcal{N}(0,G_{n\times n})$ and $\mathcal{N}(0,R_{n\times n})$. The covariance matrices are parametrized with a few scalar parameters and have the form $G_{n\times n}=\sigma_g^2A_{n\times n}$ and $R_{n\times n}=\sigma_e^2I_{n\times n}$, where A is a genetic additive relationship matrix and I is the identity matrix.

Multi-trait linear mixed model

The model for a single trait can be extended to a more general case of two and more traits by stacking observations from traits together [1].

A linear model describes observations in two traits, measured in n individuals and stored in a vector $y_{2n\times 1}$.

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

where $X_{2n\times p}$ and $Z_{2n\times 2n}$ are incidence matrices, p is the number of fixed effects, $\beta_{p\times 1}$ is a vector of fixed effects, $u_{2n\times 1}$ is a vector of a random polygenic effect, and $e_{2n\times 1}$ is a vector of the residuals errors. The random vectors u and e are mutually uncorrelated and multivariate normally distributed, $\mathcal{N}(0, G_{2n\times 2n})$ and $\mathcal{N}(0, R_{2n\times 2n})$.

The variance-covariance matrices $G_{2n\times 2n}$ and $R_{2n\times 2n}$ have a block structure and can be represented using as the Kronecker operator.

$$G_{2n \times 2n} = C_{2 \times 2} \otimes A_{n \times n} = \begin{pmatrix} c_{11}A & c_{12}A \\ c_{21}A & c_{21}A \end{pmatrix} = \begin{pmatrix} \sigma_{1}^{2}A & \rho\sigma_{1}\sigma_{2}A \\ \rho\sigma_{1}\sigma_{2}A & \sigma_{2}^{2}A \end{pmatrix}_{g}$$

$$R_{2n \times 2n} = E_{2 \times 2} \otimes I_{n \times n} = \begin{pmatrix} e_{11}I & e_{12}I \\ e_{21}I & e_{21}I \end{pmatrix} = \begin{pmatrix} \sigma_{1}^{2}I & \rho\sigma_{1}\sigma_{2}I \\ \rho\sigma_{1}\sigma_{2}I & \sigma_{2}^{2}I \end{pmatrix}_{e}$$

The diagonal entries $\sigma_{g_1}^2$ and $\sigma_{g_2}^2$ in the symmetric matrix C are marginal genetic variances for each trait, and the off-diagonal entries $\rho_g \sigma_{g_1} \sigma_{g_2}$ are covariances between the traits. The environment covariance $R_{2n \times 2n}$ is represented similarly.

Multi-environment linear mixed model

If a trait is measured in two environments, the previous model for two different traits can be applied [2]. Thus, the diagonal entries $\sigma_{g_1}^2$ and $\sigma_{g_2}^2$ in the symmetric matrix C are marginal genetic variances for each of two environment, and the off-diagonal entries $\rho_g \sigma_{g_1} \sigma_{g_2}$ are covariances between the environments. The environment covariance $R_{2n\times 2n}$ has a similar interpretation.

Blangero proposed statistical tests for the null hypothesis of no gene-environment interaction based on the likelihood ratio statistic, when comparing the full model and a reduced model. The first null model assumes that the genetic variances are equal in the null model [2, p. 535]. The second null model assumes that the genetic correlation coefficient is equal to 1.

Following the *lme4* authors' guidelines [3, Section A.1], we implemented three types of restrictions: the correlation is zero ($\rho_g = 0$), the variances are equal ($\sigma_{g_1} = \sigma_{g_2}$), and the correlation is one ($\rho_g = 1$). These types of restrictions can be extended to more general cases with multiple environments.

Multi-environment linear mixed model: a special case of sex-specificity

A sex-specificity model is a special case of gene-environment interactions where individuals are measured in single environments [2, p. 530].

A linear model describes observations in a trait, measured in n individuals and stored in a vector $y_{n\times 1}$.

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

where $X_{n \times p}$ and $Z_{n \times n}$ are incidence matrices, p is the number of fixed effects, $\beta_{p \times 1}$ is a vector of fixed effects, $u_{n \times 1}$ is a vector of a random polygenic effect, and $e_{n \times 1}$ is a vector of the residuals errors. The random vectors u and e are mutually uncorrelated and multivariate normally distributed, $\mathcal{N}(0, G_{n \times n})$ and $\mathcal{N}(0, R_{n \times n})$.

The variance-covariance matrices $G_{n \times n}$ and $R_{n \times n}$ have a block structure stratified by gender.

$$G_{n \times n} = \begin{pmatrix} \sigma_1^2 A_{11} & \rho \sigma_1 \sigma_2 A_{12} \\ \rho \sigma_1 \sigma_2 A_{21} & \sigma_2^2 A_{22} \end{pmatrix}_g$$

$$R_{n \times n} = \begin{pmatrix} \sigma_1^2 I & \rho \sigma_1 \sigma_2 I \\ \rho \sigma_1 \sigma_2 I & \sigma_2^2 I \end{pmatrix}_g$$

Matrices A_{11} , A_{12} , A_{21} and A_{22} are four blocks of the matrix A stratified by gender. For example, A_{11} is the genetic relationship matrix that corresponds to males. As A is symmetric, $A_{12} = A_{21}$.

Parameters $\sigma_{g_1}^2$ and $\sigma_{g_2}^2$ are marginal genetic variances in males and females, and parameter ρ_g is the genetic correlation coefficient between the two genders. The environment covariance parameters have a similar interpretation.

The correlation coefficient ρ_e is restricted to zero, so the sex-specificity model is identifiable [2].

$$R_{n\times n} = \begin{pmatrix} \sigma_1^2 I & 0\\ 0 & \sigma_2^2 I \end{pmatrix}_e$$

Sex-specificity linear mixed model in the GAIT2 data

In the main text of the manuscript, we showed two basic models for the analysis of APTT in the GAIT2 data, polygenic and association. Here, we present an advanced model that assesses the sex-specificity in the APTT phenotype.

Before conducting the analysis, we stored phenotype, age, gender, individual id, house-hold hhid variables and SNPs in a table dat. The additive genetic relatedness matrix was estimated by SOLAR using the pedigree information and stored in a matrix mat. A polygenic model m1 was fitted to the data as follows.

```
m1 <- relmatLmer(aptt ~ age + gender + (1|id), dat,
  relmat = list(id = mat))</pre>
```

To assess the hypothesis of sex-specificity [2] for APTT, our package allows to fit such a polygenic model m3 with multiple levels of relatedness.

```
m3 <- relmatLmer(aptt ~ age + gender + (0 + gender|id) + (0 + gender|rid), dat,
relmat = list(id = mat), vcControl = list(rho0 = list(rid = 5)),
weights = rep(1e10, nrow(dat)))</pre>
```

The first genetic random effect, denoted as (0 + gender|id), has three parameters σ_{g_1} , σ_{g_2} and ρ_g , as described in the previous section. The second residual random effect, denoted as (0 + gender|rid), also has three parameters, but the correlation coefficient is restricted to zero as specified in the vcControl argument. This restriction is necessary because the model is a special case of gene-environment interactions where individuals are measured in single environments [2, p. 530]. The variable rid is a copy of id, and using large values in the last argument weights is an *ad hoc* solution to cancel the independent and identically distributed residual error. We note that the m3 model can be fitted without the *ad hoc*.

```
m3 <- relmatLmer(aptt ~ age + gender + (0 + gender|id) + (0 + dummy(gender)|rid), dat,
relmat = list(id = mat))</pre>
```

Once the evidence of the gene-environment interaction in m3 is confirmed [2], a new association model m4 can be considered for the GWAS, in which a SNP, for example, rs1, has both marginal and interaction terms with the gender variable.

```
m4 <- update(m3, . ~ . + rs1 + rs1:gender)
anova(m3, m4)</pre>
```

Implementation of restriction on model parameters

The R code used in the previous section to fit the sex-specificity model has a special use of the vcControl parameter, that defines the restriction on variance components.

```
m3 <- relmatLmer(aptt ~ age + gender + (0 + gender|id) + (0 + gender|rid), dat,
relmat = list(id = mat), vcControl = list(rho0 = list(rid = 5)),
weights = rep(1e10, nrow(dat)))</pre>
```

To understand how the vcControl argument works, we need to write the covariance structure of random effects ((0 + gender|id) and (0 + gender|rid) using its associated Cholesky decomposition [3, Appendix A.1, p. 44, formula (69)].

$$\begin{pmatrix} \sigma_1^2 & \rho \sigma_1 \sigma_2 \\ \rho \sigma_1 \sigma_2 & \sigma_2^2 \end{pmatrix}_{\mathcal{S}} = \begin{pmatrix} \theta_1 & 0 \\ \theta_2 & \theta_3 \end{pmatrix} \begin{pmatrix} \theta_1 & \theta_2 \\ 0 & \theta_3 \end{pmatrix} = \begin{pmatrix} \theta_1^2 & \theta_1 \theta_2 \\ \theta_1 \theta_2 & \theta_2^2 + \theta_3^2 \end{pmatrix}$$
$$\begin{pmatrix} \sigma_1^2 & \rho \sigma_1 \sigma_2 \\ \rho \sigma_1 \sigma_2 & \sigma_2^2 \end{pmatrix}_{\mathcal{S}} = \begin{pmatrix} \theta_4 & 0 \\ \theta_5 & \theta_6 \end{pmatrix} \begin{pmatrix} \theta_4 & \theta_5 \\ 0 & \theta_6 \end{pmatrix} = \begin{pmatrix} \theta_4^2 & \theta_4 \theta_5 \\ \theta_4 \theta_5 & \theta_5^2 + \theta_6^2 \end{pmatrix}$$

Now it is clear that the environmental correlation can be restricted to zero by setting $\theta_5 = 0$. Consequently, the value of the vcControl argument is list(rho0 = list(rid = 5)).

The following table shows more options of using vcControl.

Condition	Parameter restrictions	vcControl value
$\rho_g = 0$	$\theta_2 = 0$	list(rho0 = list(id = 2))
$\rho_g = 1$		list(rho1 = list(id = 3))
$(\sigma_1)_g = (\sigma_2)_g$	$\theta_1^2 = \theta_2^2 + \theta_3^2$	list(vareq = list(id = c(1, 2, 3)))
$\rho_g = 0, \rho_e = 0$	$\theta_2 = 0, \theta_5 = 0$	list(rho0 = list(id = 2, rid = 5))

The use of names such as rho0, rho1 and vareq is required, as these names are bound to particular implementation (the second column of the table given above) in the body of the relmatLmer function.

References

[1] Michael Lynch, Bruce Walsh, et al. *Genetics and analysis of quantitative traits*, volume 1. Sinauer Sunderland, MA, 1998.

- [2] John Blangero. Statistical genetic approaches to human adaptability. *Human biology*, 81(5):523–546, 2009.
- [3] Douglas Bates, Martin Mächler, Ben Bolker, and Steve Walker. Fitting linear mixed-effects models using lme4. *Journal of Statistical Software*, 67(1):1–48, 2015.

Supplementary Note 3: R code applied to the GAIT2 data

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Introduction

Load packages

We need a list of R package, including our lme4qtl package, to perform the analysis.

```
library(plyr)
library(Matrix)
library(gridExtra)
library(lme4)
library(boot)
```

The next two packages complement the lme4 functionality with additional inference procedures.

```
library(lmerTest)

##
## Attaching package: 'lmerTest'

## The following object is masked from 'package:lme4':
##
## lmer

## The following object is masked from 'package:stats':
##
## step
library(RLRsim)
```

Parameters

The GAIT2 family-based sample consists of 934 individuals. Here, we use a small subset of 10 markers from Chromosome 22 in the association analysis.

```
N <- 934

chr <- 22

M <- 10
```

Load data

We need the following R packages (not publicly available) to load the GAIT2 data.

```
library(gait)
library(solarius)
```

Data variables include

- table of phenotypes phen with such variables as
 - aptt outcome, the activated partial thromboplastin time (APTT)
 - gender and age as covariates or fixed effect
 - id, the individual identifier
 - famid, the family identifier
 - hhid, the house-hold identifier (not the same as famid)
- dkin, the double kinship matrix (additive genetic effect)
- delta7, matrix of dominance genetic effect

Phenotype data

```
dir_phen <- "~/Data/GAIT2/phen/"
dir_snp <- "~/Data/GAIT2/ncdf/"

phen <- gait2.phen(dir_phen, transforms = "tr1", id.alert = TRUE, traits = "tr1_APTT")

phen <- rename(phen,
    aptt = tr1_APTT,</pre>
```

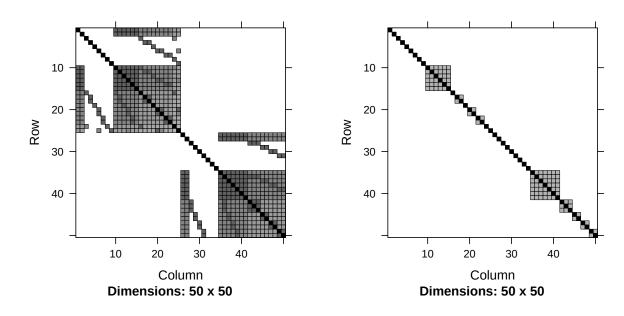
```
gender = SEXf, age = AGEc,
id = ID, famid = FAMID, hhid = HHID)

phen <- mutate(phen, rid = id, id7 = id)</pre>
```

Covariance matrices

```
dkin <- Matrix(solarKinship2(phen))
delta7 <- Matrix(solarKinship2(phen, coef = "d"))</pre>
```

The next plot depicts sub-matrices (first 50 individuals) of the genetic additive (left) and dominance (right) covariance matrices.



Polygenic analysis

The polygenic model of APTT has two random effects (apart from the residual variance), genetic additive and house-hold. In the case of the genetic effect, the covariance matrix dkin is introduced using relmat argument.

```
m1 <- relmatLmer(aptt ~ age + gender + (1|id) + (1|hhid), phen, relmat = list(id = dkin))
m1
## Linear mixed model fit by REML ['lmerMod']
  Formula: aptt ~ age + gender + (1 | id) + (1 | hhid)
##
      Data: phen
## REML criterion at convergence: 2355.299
  Random effects:
    Groups
             Name
                         Std.Dev.
##
##
    id
             (Intercept) 0.7270
##
    hhid
             (Intercept) 0.2582
    Residual
                         0.5926
```

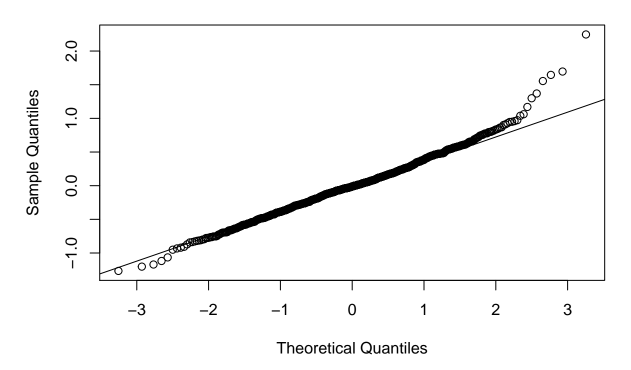
```
## Number of obs: 884, groups: id, 884; hhid, 448
## Fixed Effects:
## (Intercept) age gender2
## 0.17759 -0.01687 -0.07376
```

Diagnostics

The residuals are expected to be normally distributed.

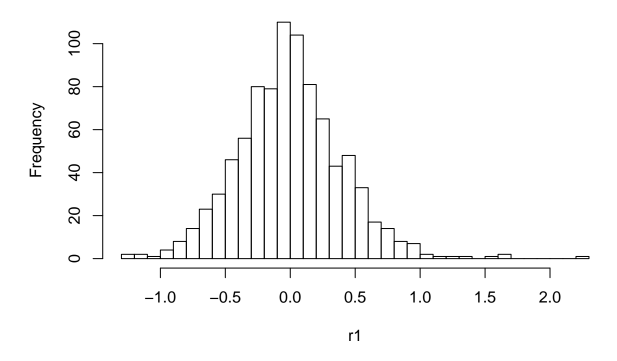
```
r1 <- residuals(m1)
qqnorm(r1)
qqline(r1)</pre>
```

Normal Q-Q Plot



hist(r1, breaks = 30)

Histogram of r1



Inference

We use the step function from the R package lmerTest.

```
# `?lmerTest::step`
# the p-value thr. are set to 1 to disable terms dropping
step(m1, alpha.random = 1, alpha.fixed = 1)
##
  Random effects:
        Chi.sq Chi.DF elim.num p.value
##
## id
         70.44
                    1
                          kept <1e-07
         2.98
## hhid
                    1
                          kept 0.0842
##
## Fixed effects:
##
           Sum Sq Mean Sq NumDF DenDF F.value elim.num Pr(>F)
          53.2671 53.2671
                              1 650.17 151.6804
                                                    kept <1e-07
  gender 0.5382 0.5382
                              1 794.99
                                         1.5326
                                                    kept 0.2161
##
## Least squares means:
             gender Estimate Standard Error DF
                                                 t-value Lower CI Upper CI
## gender 1
                      0.1826
                                     0.0586 459
                                                  3.1200
                                                            0.183
                                                                      0.183
                  1
                      0.1088
## gender
           2
                  2
                                     0.0589 439
                                                  1.8500
                                                            0.109
                                                                      0.109
##
             p-value
## gender 1 0.0019 **
## gender 2 0.0655.
```

Inference for heritability

By definition, heritability is the proportion of explained variance.

```
vf <- as.data.frame(VarCorr(m1))[, c("grp", "vcov")]
vf$prop <- with(vf, vcov / sum(vcov))</pre>
```

grp	vcov	prop
id	0.53	0.56
hhid	0.07	0.07
Residual	0.35	0.37

Confidence interval

```
# `?lme4::profile`
prof <- profile(m1, which = "theta_", prof.scale = "varcov")

# `?lme4qtl::varpropProf`
prof_prop <- varpropProf(prof)

ci <- confint(prof_prop, level = 0.95)
ci

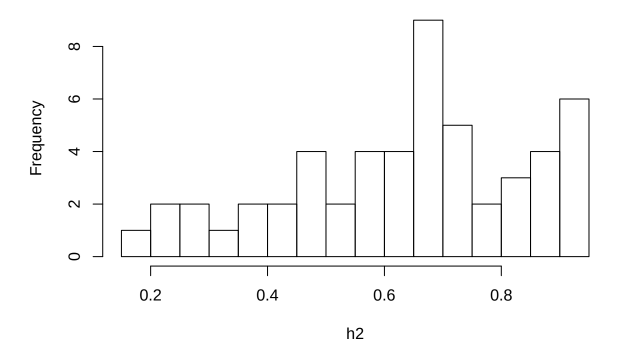
## 2.5 % 97.5 %

## .sigprop01 0.4450731 0.84293219

## .sigprop02 0.0000000 0.06472766

## .sigmaprop 0.1461354 0.50813557</pre>
```

Profiled heritability



Likelihood ratio tests (LRTs)

```
# ?RLRsim::exactRLRT
m1\_reduced \leftarrow update(m1, . \sim . - (1|hhid))
m1_null <- update(m1, . ~ . - (1|id))</pre>
rlrt_h2 <- exactRLRT(</pre>
  m1_reduced, # the reduced model with only the effect to be tested
  mA = m1, # the full model under the alternative
  m0 = m1_null, # the model under the null
  seed = 1
rlrt_h2
##
    simulated finite sample distribution of RLRT.
##
##
##
    (p-value based on 10000 simulated values)
##
## data:
## RLRT = 70.443, p-value < 2.2e-16
lrt_h2 <- anova(m1_null, m1)</pre>
## refitting model(s) with ML (instead of REML)
```

lrt_h2

```
## Data: phen
## Models:
## m1_null: aptt ~ age + gender + (1 | hhid)
## m1: aptt ~ age + gender + (1 | id) + (1 | hhid)
## Df AIC BIC logLik deviance Chisq Chi Df Pr(>Chisq)
## m1_null 5 2416.2 2440.1 -1203.1 2406.2
## m1 6 2348.0 2376.7 -1168.0 2336.0 70.185 1 < 2.2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1</pre>
```

Summary

Heritability estimates / tests	Value
Estimate	0.56
95% CI	[0.45; 0.84]
Exact RLRT p-value	< 2.2e-16
LRT p-value	$<2.2\mathrm{e}\text{-}16$

Polygenic sex-specificity analysis

The the advanced polygenic model of sex-specificity the variance depends on gender.

```
##
    Groups
             Name
                            Std.Dev. Corr
##
    id
             gender1
                            0.82909
##
                            0.69727
             gender2
                                      1.000
             dummy(gender) 0.40729
##
    rid
    Residual
                            0.52552
```

We see from the previous output of variance components that there are some sex-specific differences. To assess these difference quantitively, we will fit two null models and perform LRT:

- the genetic variances are equal;
- the genetic correlation coefficient is 1.

The later model does not make sense, as the alternative model m3 indicates that the genetic correlation coefficient is 1.

```
m3_vareq <- relmatLmer(aptt ~ age + gender + (0 + gender|id) + (0 + dummy(gender)|rid),
    phen, relmat = list(id = dkin), vcControl = list(vareq = list(id = c(1, 2, 3))), REML = FALSE)
VarCorr(m3_vareq)</pre>
```

```
Groups
             Name
                            Std.Dev. Corr
##
    id
             gender1
                            0.76577
##
             gender2
                            0.76577
                                    1.000
##
   rid
             dummy(gender) 0.17475
    Residual
                            0.58803
```

The LRT suggests that we cannot conclude that there is sex-specificity.

```
stat <- 2 * (logLikNum(m3) - logLikNum(m3_vareq))</pre>
pval <- pchisq(stat, df = 1, lower = FALSE)</pre>
pval
## [1] 0.1882005
The following code shows how to fit a model with a restriction that the genetic correlation coefficient is 1.
m3_rho1 <- relmatLmer(aptt ~ age + gender + (0 + gender | id) + (0 + dummy(gender) | rid),
    phen, relmat = list(id = dkin), vcControl = list(rho1 = list(id = 3)), REML = FALSE)
VarCorr(m3_rho1)
   Groups
                             Std.Dev. Corr
             Name
##
              gender1
                             0.82909
    id
                             0.69727 1.000
##
              gender2
## rid
              dummy(gender) 0.40729
## Residual
                             0.52552
```

Summary

Null Model	LRT p-value
$ \frac{\rho_g = 1}{(\sigma_m)_g = (\sigma_f)_g} $	1 0.1882005

Additional analyses

Dominance effect in addition to additive genetic and house-hold effects

A single genetic additive effect:

```
mod1 <- relmatLmer(aptt ~ age + gender + (1|id) + (1|hhid), phen, relmat = list(id = dkin))</pre>
## Linear mixed model fit by REML ['lmerMod']
## Formula: aptt ~ age + gender + (1 | id) + (1 | hhid)
      Data: phen
## REML criterion at convergence: 2355.299
## Random effects:
## Groups
             Name
                         Std.Dev.
## id
             (Intercept) 0.7270
## hhid
             (Intercept) 0.2582
## Residual
                         0.5926
## Number of obs: 884, groups: id, 884; hhid, 448
## Fixed Effects:
## (Intercept)
                                 gender2
                        age
       0.17759
##
                   -0.01687
                                 -0.07376
Two genetic additive and dominance effects:
mod2 <- relmatLmer(aptt ~ age + gender + (1|id) + (1|hhid) + (1|id7), phen, relmat = list(id = dkin, id
## Linear mixed model fit by REML ['lmerMod']
## Formula: aptt ~ age + gender + (1 | id) + (1 | hhid) + (1 | id7)
```

```
Data: phen
## REML criterion at convergence: 2353.095
## Random effects:
## Groups
                        Std.Dev.
            Name
## id7
             (Intercept) 0.7180
## id
             (Intercept) 0.5024
             (Intercept) 0.2687
## hhid
## Residual
                         0.3375
## Number of obs: 884, groups: id7, 884; id, 884; hhid, 448
## Fixed Effects:
## (Intercept)
                                 gender2
                        age
      0.17302
                   -0.01676
                                -0.06852
anova(mod1, mod2)
## refitting model(s) with ML (instead of REML)
## Data: phen
## Models:
## mod1: aptt ~ age + gender + (1 | id) + (1 | hhid)
## mod2: aptt ~ age + gender + (1 | id) + (1 | hhid) + (1 | id7)
             AIC
                    BIC logLik deviance Chisq Chi Df Pr(>Chisq)
## mod1 6 2348.0 2376.7 -1168.0
                                   2336.0
## mod2 7 2347.9 2381.4 -1166.9
                                   2333.9 2.125
                                                           0.1449
Two fitting methods for gene-by-gender
```

```
mod3 <- relmatLmer(aptt ~ age + gender + (0 + gender | id) + (0 + dummy(gender) | rid),
   phen, relmat = list(id = dkin), REML = FALSE)
mod3
## Linear mixed model fit by maximum likelihood ['lmerMod']
## Formula: aptt ~ age + gender + (0 + gender | id) + (0 + dummy(gender) |
##
       rid)
      Data: phen
##
##
         AIC
                   BIC
                          logLik deviance
## 2353.077 2391.352 -1168.538 2337.077
                                                  876
## Random effects:
                           Std.Dev. Corr
## Groups
             Name
## id
                           0.8291
             gender1
##
                           0.6973
                                    1.00
             gender2
## rid
             dummy(gender) 0.4073
## Residual
                           0.5255
## Number of obs: 884, groups: id, 884; rid, 884
## Fixed Effects:
## (Intercept)
                                 gender2
                        age
                                -0.08525
##
       0.18150
                   -0.01685
mod4 <- relmatLmer(aptt ~ age + gender + (0 + gender | id) + (0 + gender | rid),
   phen, relmat = list(id = dkin), vcControl = list(rho0 = list(rid = 5)),
    weights = rep(1e10, nrow(phen)), REML = FALSE)
mod4
## Linear mixed model fit by maximum likelihood ['lmerMod']
## Formula: aptt ~ age + gender + (0 + gender | id) + (0 + gender | rid)
```

```
##
      Data: phen
## Weights: rep(1e+10, nrow(phen))
##
        AIC
                 BIC
                        logLik deviance df.resid
##
         NA
                                              874
                  NΑ
                            NΑ
                                     NΑ
## Random effects:
    Groups
             Name
                      Std.Dev. Corr
##
##
             gender1 0.8292
##
             gender2 0.6969
                               1.00
##
             gender1 0.5249
   rid
##
             gender2 0.6654
                               0.00
## Residual
                      0.8131
## Number of obs: 884, groups: id, 884; rid, 884
## Fixed Effects:
## (Intercept)
                         age
                                  gender2
##
       0.18138
                                 -0.08519
                    -0.01685
```

R session info

```
sessionInfo()
```

```
## R version 3.4.0 (2017-04-21)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: OS X El Capitan 10.11.6
## Matrix products: default
## BLAS: /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLib.framework/Version
## LAPACK: /System/Library/Frameworks/Accelerate.framework/Versions/A/Frameworks/vecLib.framework/Versi
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                               datasets methods
                                                                    base
## other attached packages:
  [1] solarius_0.3.0.2 gait_0.1
                                            data.table_1.10.4
  [4] RLRsim_3.1-3
                          lmerTest_2.0-33
                                            lme4qtl_0.1.9
## [7] boot_1.3-19
                                            gridExtra_2.2.1
                          lme4_1.1-15
## [10] Matrix_1.2-9
                          dplyr_0.7.4
                                            plyr_1.8.4
## [13] rmarkdown_1.5
                          knitr_1.15.1
                                            devtools_1.13.1
##
## loaded via a namespace (and not attached):
   [1] splines_3.4.0
                            lattice_0.20-35
                                                 colorspace_1.3-2
##
  [4] htmltools_0.3.6
                            mgcv_1.8-17
                                                 yaml_2.1.14
## [7] base64enc_0.1-3
                            survival_2.41-3
                                                 rlang_0.1.2
## [10] nloptr_1.0.4
                                                 glue_1.1.1
                            foreign_0.8-67
## [13] withr_1.0.2
                            RColorBrewer_1.1-2
                                                bindrcpp_0.2
## [16] bindr_0.1
                            stringr_1.2.0
                                                 munsell_0.4.3
## [19] gtable_0.2.0
                            htmlwidgets_0.9
                                                 memoise_1.1.0
## [22] evaluate_0.10
                            latticeExtra_0.6-28 highr_0.6
## [25] htmlTable_1.9
                            Rcpp_0.12.13
                                                 acepack_1.4.1
## [28] scales_0.5.0
                            backports_1.0.5
                                                 checkmate_1.8.2
```

##	[31]	$Hmisc_4.0-3$	ggplot2_2.2.1	digest_0.6.12
##	[34]	stringi_1.1.5	grid_3.4.0	rprojroot_1.2
##	[37]	quadprog_1.5-5	kinship2_1.6.4	tools_3.4.0
##	[40]	magrittr_1.5	lazyeval_0.2.0	tibble_1.3.4
##	[43]	Formula_1.2-1	cluster_2.0.6	pkgconfig_2.0.1
##	[46]	MASS_7.3-47	assertthat_0.2.0	$minqa_1.2.4$
##	[49]	R6_2.2.1	rpart_4.1-11	nnet_7.3-12
##	[52]	nlme_3.1-131	compiler_3.4.0	