

# Package ‘CodataGS’

January 20, 2025

**Type** Package

**Title** Genomic Prediction Using SNP Codata

**Version** 1.43

**Date** 2019-05-17

**Author** Lars Ronnegard

**Maintainer** Lars Ronnegard <lrn@du.se>

**Description** Computes genomic breeding values using external information on the markers. The package fits a linear mixed model with heteroscedastic random effects, where the random effect variance is fitted using a linear predictor and a log link. The method is described in Mouresan, Selle and Ronnegard (2019) <[doi:10.1101/636746](https://doi.org/10.1101/636746)>.

**License** GPL

**Depends** Matrix

**NeedsCompilation** no

**Repository** CRAN

**Date/Publication** 2019-05-17 15:40:07 UTC

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CodataGS-package

*Genomic Prediction Using SNP Codata***Description**

Computes genomic breeding values using external information on the markers. The package fits a linear mixed model with heteroscedastic random effects, where the random effect variance is fitted using a linear predictor and a log link. The method is described in Mouresan, Selle and Ronnegard (2019) <doi:10.1101/636746>.

**Details**

The DESCRIPTION file:

```
Package:          CodataGS
Type:            Package
Title:           Genomic Prediction Using SNP Codata
Version:         1.43
Date:            2019-05-17
Author:          Lars Ronnegard
Maintainer:     Lars Ronnegard <lrn@du.se>
Description:    Computes genomic breeding values using external information on the markers. The package fits a linear
License:        GPL
Depends:        Matrix
NeedsCompilation: no
```

Index of help topics:

```
CodataGS-package      Genomic Prediction Using SNP Codata
MME                   Mixed model equations
Transform              Transforms hat values
compute_GL            Computes genomic relationship matrix
compute_phitau        Computes models for the variance components
genomicEBV.w.codata  Performs genomic prediction based on SNP
                      codata.
hat.transf            Transforms hat values
scaleZ                Scales the genotype matrix.
summary.CodataGS     Summary method for CodataGS objects
```

This package performs genomic prediction based on SNP codata. The main function is genomicEBV.w.codata.

**Author(s)**

Lars Ronnegard

Maintainer: Lars Ronnegard <lrn@du.se>

---

 compute\_GL

*Computes genomic relationship matrix*


---

**Description**

This function computes the genomic relationship matrix, G, together with its matrix square root, L.

**Usage**

```
compute_GL(Z, w)
```

**Arguments**

Z	Scaled matrix with genotype information
w	weights

**Value**

L	Square root matrix of G
svdVec	Vectors in the Single Value Decomposition of G
svdD	Diagonal elements in the Single Value Decomposition of G
wZt	weights times the transpose of Z

**Author(s)**

Lars Ronnegard

**Examples**

```
set.seed(1234)
N <- 20 #Number of individuals
k <- 30 #Number of SNPs with all marker positions including a QTL
Z1 <- matrix(0, N, k )
Z2 <- matrix(0, N, k )
Z1[1:N, 1] <- rbinom(N, 1, 0.5) #Simulated phased SNP matrices
Z2[1:N, 1] <- rbinom(N, 1, 0.5)
LD.par <- 0.2 #A parameter to simulate LD. 0 gives full LD, and 0.5 no LD
for (j in 2:k) {
  Z1[1:N, j] <- abs( Z1[1:N, j-1] - rbinom(N, 1, LD.par) )
  Z2[1:N, j] <- abs( Z2[1:N, j-1] - rbinom(N, 1, LD.par) )
}
Z <- Z1 + Z2 #Genotypic SNP matrix
sim.res <- compute_GL(Z, w = rep(1,k))
```

---

compute\_phitau                    *Computes models for the variance components*

---

### Description

This function computes the residual variance, the SNP variances and the linear predictor for the SNP variance model.

### Usage

```
compute_phitau(dev, hv, devu, hvu, X.rand.disp)
```

### Arguments

dev	Deviance values
hv	Hat values for the observed response values
devu	Deviance values computed for the random effects
hvu	Hat values for the random effects
X.rand.disp	Design matrix used in the linear predictor for the SNP variance model.

### Value

var.e	Residual variance
phi	Vector of SNP variances
coef	Fitted coefficients for the linear predictor in the SNP variance model

### Author(s)

Lars Ronnegard

### Examples

```
set.seed(1234)
N <- 20 #Number of individuals
k <- 30 #Number of SNPs with all marker positions including a QTL
#Simulated deviances and hat values
dev <- rnorm(N)^2
hv <- runif(N, 0.1, 0.5)
devu <- rnorm(k)^2
hvu <- runif(k, 0.1, 0.85)
X.rand.disp <- matrix(1, k, 1)
sim.res <- compute_phitau(dev, hv, devu, hvu, X.rand.disp)
```

---

genomicEBV.w.codata *Performs genomic prediction based on SNP codata.*

---

### Description

The main function of the package. The input includes response values, a design matrix for the fixed effects, a matrix with SNP genotype data and a design matrix for the SNP codata.

### Usage

```
genomicEBV.w.codata(y, X, Z, X.SNPcodata, Z.test = NULL, max.iter = 100, conv.crit = 1e-5)
```

### Arguments

y	Response values
X	Design matrix for the fixed effects
Z	Genotype matrix with element values of 0, 1 or 2
X.SNPcodata	Design matrix for the linear predictor of the SNP variances.
Z.test	An optional genotype matrix for a test data set.
max.iter	The maximum number of iterations
conv.crit	The value of the convergence criterion.

### Details

By specifying the matrix `Z.test` in the input, the function computes predicted genomic breeding values for an out-of-sample data set.

### Value

gEBV	Genomic breeding values
predicted.gEBV	Genomic breeding values based on the genotypes in <code>Z.test</code>
w	Computed SNP weights
u	Fitted SNP effects
beta	Fitted fixed effects
disp.beta	Fitted coefficients in the linear predictor for the SNP variance model
Converge	Shows whether the algorithm has converged or not
iter	The number of iterations used

### Author(s)

Lars Ronnegard

## Examples

```
#####
#Simulation part
set.seed(1234)
N <- 200 #Number of individuals
k <- 300 #Number of SNPs with all marker positions including a QTL
Z1 <- matrix(0, N, k )
Z2 <- matrix(0, N, k )
Z1[1:N, 1] <- rbinom(N, 1, 0.5) #Simulated phased SNP matrices
Z2[1:N, 1] <- rbinom(N, 1, 0.5)
LD.par <- 0.2 #A parameter to simulate LD. 0 gives full LD, and 0.5 no LD
for (j in 2:k) {
  Z1[1:N, j] <- abs( Z1[1:N, j-1] - rbinom(N, 1, LD.par) )
  Z2[1:N, j] <- abs( Z2[1:N, j-1] - rbinom(N, 1, LD.par) )
}
Z <- Z1 + Z2 #Genotypic SNP matrix
x1 <- c(rep(1,k/2), rep(0,k/2)) #An indicator for the SNPs.
#The first k/2 SNPs and the last k/2 have different variances
#Simulate linear predictor for the random effect variance
lin.pred <- 0 + 2*x1
X.snp <- model.matrix( ~ x1 ) #Corresponding design matrix
u <- rnorm(k, 0 , sqrt( exp(lin.pred) ))
#Took the square root here because it is the SD that is specified.
#and exp() because we are modelling a log link.
u.scaled <- u/as.numeric( sqrt( var( crossprod(t(Z), u) ) ) )
#Scaled by the variance of the breeding values
e <- rnorm(N) #A residual variance
mu <- 0
y <- mu + crossprod(t(Z),u.scaled) + e
#####
#Estimation part
mod1 <- genomicEBV.w.codata(y = as.numeric(y),
  X = matrix(1, N, 1), Z = Z, X.SNPcodata = X.snp)
#To fit gBLUP just specify X.SNPcodata = matrix(1, k, 1)
cat("Correlation between true and estimated BV for the codata model:")
cat(cor(crossprod(t(Z),u.scaled), mod1$gEBV), "\n")
```

---

hat.transf

*Transforms hat values*


---

## Description

Transforms hat values between the SNP-BLUP model and the gBLUP model.

## Usage

```
hat.transf(C22, transf, vc, k, N, w)
```

**Arguments**

C22	Submatrix of the inverse of the LHS in the MME
transf	A transformation matrix.
vc	Genetic variance
k	Number of SNPs
N	Number of individuals
w	SNP weights

**Value**

Transformed hat values

**Author(s)**

Lars Ronnegard

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MME

*Mixed model equations*

---

**Description**

A fast version of the Henderson's mixed model equations (MME)

**Usage**

MME(y, X, Z, var.e, var.u)

**Arguments**

y	Response
X	Design matrix for fixed effects
Z	Design matrix for the random effects
var.e	Residual variance
var.u	Genetic variance

**Value**

beta	Estimates of fixed effects
v	Fitted random effects
hv	Hat values
dev	Deviances

**Author(s)**

Lars Ronnegard

---

`scaleZ`*Scales the genotype matrix.*

---

**Description**

Scales the genotype matrix so that  $ZZ'$  gives the genomic relationship matrix.

**Usage**

```
scaleZ(Z, freq1)
```

**Arguments**

<code>Z</code>	Genotype matrix with element values 0, 1 and 2
<code>freq1</code>	Optional input parameter with allele frequencies. A vector of length equal to the number of columns in <code>Z</code> .

**Value**

<code>Z</code>	Scaled genotype matrix
----------------	------------------------

**Author(s)**

Lars Ronnegard

**Examples**

```
#####  
#Simulation part  
set.seed(1234)  
N <- 200 #Number of individuals  
k <- 300 #Number of SNPs with all marker positions including a QTL  
Z1 <- matrix(0, N, k )  
Z2 <- matrix(0, N, k )  
Z1[1:N, 1] <- rbinom(N, 1, 0.5) #Simulated phased SNP matrices  
Z2[1:N, 1] <- rbinom(N, 1, 0.5)  
LD.par <- 0.2 #A parameter to simulate LD. 0 gives full LD, and 0.5 no LD  
for (j in 2:k) {  
  Z1[1:N, j] <- abs( Z1[1:N, j-1] - rbinom(N, 1, LD.par) )  
  Z2[1:N, j] <- abs( Z2[1:N, j-1] - rbinom(N, 1, LD.par) )  
}  
Z <- Z1 + Z2 #Genotypic SNP matrix  
sim.res <- scaleZ(Z)
```



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summary.CodataGS	<i>Summary method for CodataGS objects</i>
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---

## Description

A summary method for the object class CodataGS

## Usage

```
## S3 method for class 'CodataGS'
summary(object, ...)
```

## Arguments

object	A CodataGS object
...	arguments not used

## Details

Provides a concise summary of CodataGS objects.

## Examples

```
#####
#Simulation part
set.seed(1234)
N <- 200 #Number of individuals
k <- 300 #Number of SNPs with all marker positions including a QTL
Z1 <- matrix(0, N, k )
Z2 <- matrix(0, N, k )
Z1[1:N, 1] <- rbinom(N, 1, 0.5) #Simulated phased SNP matrices
Z2[1:N, 1] <- rbinom(N, 1, 0.5)
LD.par <- 0.2 #A parameter to simulate LD. 0 gives full LD, and 0.5 no LD
for (j in 2:k) {
  Z1[1:N, j] <- abs( Z1[1:N, j-1] - rbinom(N, 1, LD.par) )
  Z2[1:N, j] <- abs( Z2[1:N, j-1] - rbinom(N, 1, LD.par) )
}
Z <- Z1 + Z2 #Genotypic SNP matrix
x1 <- c(rep(1,k/2), rep(0,k/2)) #An indicator for the SNPs.
#The first k/2 SNPs and the last k/2 have different variances
#Simulate linear predictor for the random effect variance
lin.pred <- 0 + 2*x1
X.snp <- model.matrix( ~ x1 ) #Corresponding design matrix
u <- rnorm(k, 0 , sqrt( exp(lin.pred) ))
#Took the square root here because it is the SD that is specified.
#and exp() because we are modelling a log link.
u.scaled <- u/as.numeric( sqrt( var( crossprod(t(Z), u) ) ) )
#Scaled by the variance of the breeding values
e <- rnorm(N) #A residual variance
```

```

mu <- 0
y <- mu + crossprod(t(Z),u.scaled) + e
#####
#Estimation part
mod1 <- genomicEBV.w.codata(y = as.numeric(y),
                           X = matrix(1, N, 1), Z = Z, X.SNPcodata = X.snp)
summary(mod1)

```

---

Transform

*Transforms hat values*


---

### Description

The function calls the `hat.transf` function.

### Usage

```
Transform(X, L, var.e, var.u, v, svdVec, svdD, wZt, w)
```

### Arguments

X	Design matrix for the fixed effects
L	Square root matrix of the genomic relationship matrix, G
var.e	Residual variance
var.u	Genetic variance
v	Random effects
svdVec	Vector from the Single Value Decomposition of G
svdD	Diagonal elements of the Single Value Decomposition of G
wZt	Weights times the transpose of the scaled genotype matrix
w	Fitted SNP weights

### Value

u	SNP effects
qu	Hat values for the SNP effects

### Author(s)

Lars Ronnegard

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