

# Package ‘CodataGS’

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**Type** Package

**Title** Genomic Prediction Using SNP Codata

**Version** 1.43

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**Author** Lars Ronnegard

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**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

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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 <lrm@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 <lrm@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

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|        |                                    |
|--------|------------------------------------|
| scaleZ | <i>Scales the genotype matrix.</i> |
|--------|------------------------------------|

---

## Description

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

## Usage

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

## Arguments

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

## Value

|   |                        |
|---|------------------------|
| Z | 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)
```

---

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

---

## 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 | <i>Transforms hat values</i> |
|-----------|------------------------------|

---

**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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