HIBLUP User Manual

January 14, 2019 v1.1.0 **Disclaimer**: While extensive tests have been performed by Zhao lab at Huazhong Agricultural University and Yuan lab at Wuhan University of Technology. Results are, in general, reliable, correct, and appropriate. However, results are not guaranteed for any specific data set. We strongly recommend that users validate the HIBLUP results with other software packages, such as GCTA, LDAK, and DMU.

Support documents: Extensive support documents, including the user manual, demo script, demo data and demo results, are available at the zip file.

Questions and comments: Users and developers are recommended to send questions to Lilin Yin (ylilin@163.com), Haohao Zhang (haohaozhang@whut.edu.cn), and Xiaolei Liu (xiaoleiliu@mail.hzau.edu.cn).

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1 Installing HIBLUP and a quick start

1.1 Installation

It is highly recommended to install Microsoft R Open (https://mran.microsoft.com/download/) to speed up the mathematical calculation of HIBLUP, but this is not required. HIBLUP can also work fine with base R. The latest software packages and user manuals are available on the HIBLUP homepage (https://hiblup.github.io). It can be installed with the following code:

```
install.packages("Rcpp")
install.packages("RcppParallel")
install.packages("RcppArmadillo")
install.packages("bigmemory")
install.packages("hiblup_1.1.0_R_x86_64-pc-linux-gnu.tar.gz", repos = NULL)
```

1.2 Quick start

The data embedded in HIBLUP was derived from an animal breeding farm, it includes a total of 2934 genetic related individuals and 573 of them were genotyped with 50K SNP Chip. The genotype was coded as 0, 1, 2 for AA, AB, BB, respectively, and two traits(t1, t2) were recorded for 800 individuals. Sire information and sex information can be treated as random effect and fixed effect, respectively. A quick start of HIBLUP to fit above model is shown below:

```
suppressMessages(library("hiblup"))
data("hidata")
X <- model.matrix(~Sex, data = pheno) # fixed effects</pre>
R <- as.matrix(pheno$Sire) # random effects</pre>
gebv <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,
   pedigree = pedigree, vc.method = c("HI"), mode = "A", CV = X, R = R,
   back.solution = TRUE)
    -----#
  He-aI BLUP
             | | | | | _ | _ | _ \| | | | | | | | _ _ \
             | |__| | | | | |_) | | | | | | | |_) |
             | __ | | | _ <| | | | | | ___/
             |_| |_|___/|__ Version: 1.1.0 #
SSBLUP model is selected based on the provided data
 Analyzed trait: t2
 Number of fixed effects: 2 (intercept included)
 Number of random effects: 1
 Number of individuals with phenotypic observations: 800
 Deriving GA matrix from genotype...Done within 2s
 Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Deriving A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
```

```
Realign index of X matrix...Done!
Realign index of R matrix...Done!
Extracting A11 matrix...Done!
Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
Adjusting GA matrix: GA* = 0.98 * GA + 0.03
Weighting of A11 and GA matrix: 0.05
Calculating the inverse of A11 matrix...Done within Os
Constructing HA matrix...Done within 1s
HE Prior derived: A:0.4034 e:3.166; Done within Os
HE adopted: TRUE
Variance components estimation:
[Iter] Var_R1(SE)
                    Var_K1(SE)
                                                          h2_R1(SE)
                                                                          h2_K1(SE)
                                          Var e(SE)
[AI] 0.471187(0.7591) 0.375320(0.4361) 2.765134(0.4017) 0.1305(0.1833) 0.1039(0.1206)
[AI] 0.325688(0.1923) 0.250717(0.3599) 1.919587(0.3290) 0.1305(0.0688) 0.1004(0.1421)
[AI] 0.356566(0.1103) 0.278890(0.2048) 2.098667(0.1883) 0.1304(0.0360) 0.1020(0.0739)
[AI] 0.359874(0.1265) 0.282026(0.2358) 2.117141(0.2163) 0.1304(0.0409) 0.1022(0.0843)
[AI] 0.359916(0.1283) 0.282024(0.2392) 2.117333(0.2193) 0.1304(0.0415) 0.1022(0.0855)
[AI] 0.359914(0.1283) 0.282027(0.2392) 2.117331(0.2193) 0.1304(0.0415) 0.1022(0.0855)
[AI] 0.359915(0.1283) 0.282026(0.2392) 2.117331(0.2193) 0.1304(0.0415) 0.1022(0.0855)
[Convergence] YES
Done within 2s
Estimating SNP effect...Done within Os
Estimated beta: 11.53 0.8425
Estimated Vg and Ve: 0.282 2.117
HIBLUP IS DONE WITHIN: 7s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

You can also load your own data with the following codes:

```
pheno <- read.table("phenotype.txt", header = F)
geno <- bigmemory::attach.big.matrix("genotype.desc")
geno.id <- read.table("geno.id", header = F)
pedigree <- read.table("pedigree.txt", header = T)
map <- read.table("map.txt", header = F)</pre>
```

Note that two result files will be generated. One is used for storing the estimated genetic values, and the other is used for storing marker effects if "back.solution" is TRUE. The contents of these two files are displayed as follows:

Ind	hiblup.A.ebv	hiblup.D.ebv	hiblup.AD.ebv
P0322	-0.0805	-3.21e-04	-0.0808
P0323	-0.1635	-3.14e-04	-0.1638
P0324	-0.0830	1.04e-06	-0.0830
P0325	-0.0637	1.31e-06	-0.0636

Table 1: Estimated genetic values.

Marker	Chr	Pos	P.Freq	SNP.A.effect	SNP.D.effect
ASGA0000014	1	342481	0.355	-0.0304	0.01134
ASGA0000021	1	489855	0.407	-0.0473	-0.00152
H3GA0000026	1	509928	0.286	0.2445	0.09484
ALGA0000009	1	538161	0.139	-0.2514	-0.07195
ALGA0000014	1	565627	0.390	-0.1583	-0.00737

Table 2: Genetic marker effects.

2 Input of HIBLUP

The data requirements of three BLUP methods in HIBLUP:

ABLUP: Phenotypic observations, Pedigree records GBLUP: Phenotypic observations, Genotype data

SSBLUP: Phenotypic observations, Genotype data, and Pedigree records

2.1 Pedigree Data

The pedigree data file includes 3 columns (sample id, paternal id, and maternal id). Note that the individuals in the pedigree data file do not need to be sorted by the date of birth, and the missing value can be replaced by NA or 0.

2.2 Genotype data

HIBLUP accepts both "big.matrix" format, which is from R bigmemory package and R standard "matrix" format. Each Column represents an individual and each row represents a marker. Here is an example that contains 573 individuals and each individual has 48,353 markers from the demo data. Genotype data in multiple popular formats such as vcf, hapmap, and plink binary format can be converted to "big.matrix" using "MVP.Data" function in the rMVP package (https://github.com/XiaoleiLiuBio/rMVP). Genotype ID list is a one-column matrix that includes the id list of genotyped individuals. The order of individuals in genotype id list should match the order of individuals in Genotype data file.

```
geno.id[1:3, ]

[1] ind799 ind800 ind801

573 Levels: ind1061 ind1063 ind1066 ind1067 ind1068 ... ind842

dim(geno)
```

```
[1] 48353 573

geno[1:3, 1:10]

V1 V2 V3 V4 V5 V6 V7 V8 V9 V10

1 1 1 2 0 1 2 1 1 0 1

2 1 1 2 2 2 0 2 1 2 1

3 1 1 1 1 0 0 1 2 0
```

2.3 Genotypic map data

Genotypic map data includes three columns, which are marker id, Chromosome ID, and physical position. This information is only used for the output.

```
dim(map)

[1] 48353 3

map[1:3, ]

SNP Chrom BP

1 10000345 0 0
2 10007117 0 0
3 12784072 0 0
```

2.4 Phenotype, Fixed effects, and Random effects

Individuals in Phenotype, fixed effects, and random effects must have the same order and the individual ID is only added in the first column of phenotype data. Above three information are always incorporated in a single file. Missing phenotype value should be marked with "NA".

2.5 Variance components

```
For single trait and K model, vc=c(V_g, V_e); for pairs of correlated traits, vc=c(V_g^{(1)}, V_g^{(2)}, COV_g^{(12)}, V_e^{(1)}, V_e^{(2)}, COV_e^{(12)}); for multiple K model, vc=c(V_g^{(1)}, V_g^{(2)}, \ldots, V_g^{(n)}, V_e); if R (Random effects) is added in the model, V_r should be added in the beginning of vc vector.
```

3 Gallery of HIBLUP input parameters

Parameter	Default	Options	Description
Pheno	NULL	Users	Phenotypic observations
CV	NULL	Users	Fixed effects
R	NULL	Users	Random effects
geno	NULL	Users	Genotype data
pedigree	NULL	Users	Pedigree records
map	NULL	Users	Genotypic map
geno.id	NULL	Users	Genotype id list
val.id	NULL	Users	sample id list for prediction on individual genetic value
K	NULL	Users	a list of variance-covariance matrices for random effects
G	NULL	Users	Relationship matrix that derived from geno- type data
A	NULL	Users	Relationship matrix that derived from pedigree records
alpha	0.05	$0 \sim 1$	the weight of A matrix when merging A and G matrices
cpu	NULL	Positive integer	number of threads used for parallel computation, default is NULL and automatically assign the computational task to appropriate number of threads
vc	NULL	Users	A vector includes known variance components. See section 2.5
mode	"A"	"A" or "AD"	"A" and "AD" repsent Additive model and Additive plus Dominant model, respectively; it doesn't work when "K" is not NULL
vc.method	HI	"AI", "EM", "AIEM", "EMAI", "HE", and "HI"	methods for variance components estimation
nAIiter	20	Positive integer	Maximum iteration number for "AI"
nEMiter	1	Positive integer	Maximum iteration number for "EM"
mme.method	"sor"	"solve" and "sor"	methods for solving mixed model equation when vc is known
reliability	FALSE	TRUE or FALSE	if TRUE, the reliability of individual genetic value will be calculated
back.solution	FALSE	TRUE or FALSE	if TRUE, the marker effects will be calculated
file.output	TRUE	TRUE or FALSE	if TRUE, gebv and marker effect will be written out
het.add	FALSE	TRUE or FALSE	if TRUE, the individual heterozygosity will be added as covariates in AD model

Table 3: Gallery of HIBLUP input parameters $\,$

4 Functions and scripts

In this section, we will provide some code snippets to show the HIBLUP functions. For the sake of brevity, output has been hidden. All code has been verified under the built-in data set.

```
suppressMessages(library("hiblup"))
data("hidata")
```

4.1 Load data

You can also load your own data with the following codes:

```
pheno <- read.table("phenotype.txt", header = F)
geno <- bigmemory::attach.big.matrix("genotype.desc")
geno.id <- read.table("geno.id", header = F)
pedigree <- read.table("pedigree.txt", header = T)
map <- read.table("map.txt", header = F)</pre>
```

4.2 Construct relationship matrix

4.2.1 Pedigree based relationship matrix(A matrix)

Construct pedigree based Additive relationship matrix:

```
Acal <- hiblup.AD(pedigree = pedigree, mode = "A")

Deriving A matrix from pedigree...Done within Os

A_PA <- Acal$PA
id <- Acal$order.id
```

Construct pedigree based Additive and Dominant relationship matrix:

```
ADcal <- hiblup.AD(pedigree = pedigree, mode = "AD")

Deriving A and D matrix from pedigree...Done within 1s

AD_PA <- ADcal$PA

AD_PD <- ADcal$PD

id <- ADcal$order.id
```

4.2.2 Genome based relationship matrix(G matrix)

Construct genome based Additive relationship matrix:

```
Acal <- hiblup.K(M = geno, mode = "A")
A_GA <- Acal$GA
```

Construct genome based Additive and Dominant relationship matrix:

```
ADcal <- hiblup.K(M = geno, mode = "AD")

AD_GA <- ADcal$GA

AD_GD <- ADcal$GD
```

4.2.3 Pedigree and genome based relationship matrix(H matrix)

Construct pedigree and genome based Additive relationship matrix:

```
G_ind <- as.character(as.matrix(geno.id)[, 1])</pre>
phe_ind <- as.character(as.matrix(pheno)[, 1])</pre>
Acal <- hiblup.AD(pedigree = pedigree, mode = "A")
Deriving A matrix from pedigree...Done within Os
A_PA <- Acal$PA
A_ind <- Acal$order.id
Acal <- hiblup.AD(pedigree, mode = "A", inverse = TRUE)
Deriving the inverse of A matrix from pedigree...Done within Os
A_PAinv <- Acal$PA
Acal <- hiblup.K(M = geno, mode = "A")
A_GA <- Acal$GA
H <- hiblup.H(A_ind = A_ind, G_ind = G_ind, phe_ind = phe_ind, A = A_PA,
    G = A_GA, Ainv = A_PAinv, alpha = 0.05, tag = "a")
 Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Constructing HA matrix...Done within 1s
```

Construct pedigree and genome based Additive and Dominant relationship matrix:

```
G_ind <- as.character(as.matrix(geno.id)[, 1])
phe_ind <- as.character(as.matrix(pheno)[, 1])

ADcal <- hiblup.AD(pedigree = pedigree, mode = "AD")

Deriving A and D matrix from pedigree...Done within 1s

AD_PA <- ADcal$PA
AD_PD <- ADcal$PD
A_ind <- ADcal$poder.id</pre>
ADcal <- hiblup.AD(pedigree, mode = "AD", inverse = TRUE)
```

```
Deriving A and D matrix from pedigree...Done within 1s
 Deriving the inverse of both A and D matrix from pedigree...Done within 2s
AD_PAinv <- ADcal$PA
AD_PDinv <- ADcal$PD
ADcal <- hiblup.K(M = geno, mode = "AD")
AD_GA <- ADcal$GA
AD_GD <- ADcal$GD
HA <- hiblup.H(A_ind = A_ind, G_ind = G_ind, phe_ind = phe_ind, A = AD_PA,
   G = AD_GA, Ainv = AD_PAinv, alpha = 0.05, tag = "a")
Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Constructing HA matrix...Done within 1s
HD <- hiblup.H(A_ind = A_ind, G_ind = G_ind, phe_ind = phe_ind, A = AD_PD,
   G = AD_GD, Ainv = AD_PDinv, alpha = 0.05, tag = "d")
 Extracting D11 matrix...Done!
 Mean of diagonal and Off-diagonal of PD: 1 0.001
 Mean of diagonal and Off-diagonal of GD: 0.5374 -9e-04
 Adjusting GD matrix: GD* = 1.86 * GD + 0
 Weighting of D11 and GD matrix: 0.05
 Calculating the inverse of D11 matrix...Done within Os
 Constructing HD matrix...Done within 1s
```

4.3 Variance components estimation

Six variance components estimation methods were implemented in HIBLUP, including AI, EM, EMAI, AIEM, HE Regression, and HI. Since the information required for HE Regression is different from other methods, it is implemented separately in a function named hiblup.he. Other methods can be called by setting the method parameter of the hiblup.vc function. nAIiter and nEMiter are the maximum iteration number of "AI" and "EM", valid only in the variance components estimation with "AI" or "EM" method.

4.3.1 Singe K model

```
index <- match(geno.id[, 1], pheno[, 1])

# AI
vc <- hiblup.vc(y = pheno$t2[index], K = A_GA, nAIiter = 20, method = "AI")

# EM
vc <- hiblup.vc(y = pheno$t2[index], K = A_GA, nEMiter = 20, method = "EM")</pre>
```

```
# HI
vc <- hiblup.vc(y = pheno$t2[index], K = A_GA, nAIiter = 20, method = "HI")

# EMAI
vc <- hiblup.vc(y = pheno$t2[index], K = A_GA, nAIiter = 20, nEMiter = 1,
    method = "EMAI")

# AIEM
vc <- hiblup.vc(y = pheno$t2[index], K = A_GA, nAIiter = 5, nEMiter = 20,
    method = "AIEM")

# HE Regression
vc <- hiblup.he(y = pheno$t2[index], K = A_GA)</pre>
```

Fixed effects and random effects can be added by parameters X and R:

```
index <- match(geno.id[, 1], pheno[, 1])</pre>
X <- model.matrix(~Sex, data = pheno) # fixed effects</pre>
R <- as.matrix(pheno$Sire) # random effects</pre>
vc <- hiblup.vc(y = pheno$t2[index], X = X[index,], R = R[index,], K = A_GA,
    nAliter = 20, method = "Al")
 Variance components estimation:
                           Var_K1(SE)
 [Iter] Var_R1(SE)
                                                Var_e(SE)
                                                                 h2_R1(SE)
                                                                                    h2 K1(SE)
  \texttt{[AI]} \ \ 1.146199 (0.6755) \ \ 0.018965 (0.8592) \ \ 3.277526 (0.5790) \ \ 0.2580 (0.1181) \ \ 0.0043 (0.1931) 
  \texttt{[AI]} \ \ 1.262022 (0.4568) \ \ 0.036802 (0.3420) \ \ 3.578946 (0.4950) \ \ 0.2587 (0.0743) \ \ 0.0075 (0.0701) 
  \texttt{[AI]} \ \ 1.274311(0.5270) \ \ 0.040030(0.4036) \ \ 3.609913(0.5720) \ \ 0.2588(0.0849) \ \ 0.0081(0.0820) 
 [AI] 1.274342(0.5347) 0.039942(0.4113) 3.610357(0.5808) 0.2588(0.0861) 0.0081(0.0835)
 [AI] 1.274347(0.5348) 0.039948(0.4113) 3.610349(0.5808) 0.2588(0.0861) 0.0081(0.0835)
 [AI] 1.274347(0.5348) 0.039948(0.4113) 3.610349(0.5808) 0.2588(0.0861) 0.0081(0.0835)
 [Convergence] YES
 Done within Os
```

4.3.2 Multiple K model

Parameter K accepts a list of Ks and execute a multiple random effects model:

```
index <- match(geno.id[, 1], pheno[, 1])</pre>
ADcal <- hiblup.K(M = geno, mode = "AD")
AD_GA <- ADcal$GA
AD_GD <- ADcal$GD
# AI, EM, EMAI, AIEM, and HI algorithm
vc <- hiblup.vc(y = pheno$t2[index], K = list(AD_GA, AD_GD), method = "AI")
 Variance components estimation:
 [Iter] Var_K1(SE)
                                                    Var_e(SE)
                               Var_K2(SE)
                                                                       h2_K1(SE)
                                                                                           h2 K2(SE)
  \texttt{[AI]} \ \ 1.693366(1.3788) \ \ 0.791422(2.4007) \ \ 2.403310(1.6515) \ \ 0.3464(0.2689) \ \ 0.1619(0.4863) 
  \texttt{[AI]} \ \ 1.744592(1.3761) \ \ 0.030725(2.4384) \ \ 3.099038(1.7194) \ \ 0.3579(0.2688) \ \ 0.0063(0.5001) 
  \texttt{[AI]} \ \ 1.746125(1.3661) \ \ 0.009845(2.4571) \ \ 3.118258(1.7663) \ \ 0.3582(0.2669) \ \ 0.0020(0.5041) 
  \hbox{\tt [AI]} \ \ 1.746508 (1.3658) \ \ 0.004647 (2.4575) \ \ 3.123044 (1.7674) \ \ 0.3583 (0.2668) \ \ 0.0010 (0.5042)
```

```
[AI] 1.746699(1.3657) 0.002050(2.4576) 3.125435(1.7677) 0.3584(0.2668) 0.0004(0.5042)
 [AI] 1.746795(1.3657) 0.000753(2.4576) 3.126630(1.7678) 0.3584(0.2668) 0.0002(0.5042)
  \hbox{\tt [AI]} \ \ 1.746842 (1.3657) \ \ 0.000104 (2.4576) \ \ 3.127227 (1.7679) \ \ 0.3584 (0.2668) \ \ 0.0000 (0.5042) 
  \texttt{[AI]} \ \ 1.746848(1.3657) \ \ 0.000023(2.4577) \ \ 3.127302(1.7680) \ \ 0.3584(0.2668) \ \ 0.0000(0.5042) 
  \hbox{\tt [AI]} \ \ 1.746850 (1.3657) \ \ 0.000003 (2.4577) \ \ 3.127320 (1.7680) \ \ 0.3584 (0.2668) \ \ 0.0000 (0.5042) \\
 [AI] 1.942710(1.3657) 0.000000(2.4577) 5.573222(1.7680) 0.2585(0.1751) 0.0000(0.3270)
 [AI] 2.381842(2.4885) 0.000000(4.8410) 7.838853(3.4667) 0.2330(0.2386) 0.0000(0.4736)
 [AI] 2.991915(3.8848) 0.000000(7.7095) 8.706900(5.5106) 0.2557(0.3262) 0.0000(0.6590)
 [AI] 2.483826(4.8247) 0.000000(9.4059) 9.281411(6.7344) 0.2111(0.4026) 0.0000(0.7995)
  \hbox{\tt [AI]} \ \ 3.754822(4.7280) \ \ 0.000000(9.5458) \ \ 8.324399(6.8106) \ \ 0.3108(0.3854) \ \ 0.0000(0.7903) 
 [AI] 0.679796(5.2155) 0.000000(9.7390) 10.455150(6.9943) 0.0611(0.4646) 0.0000(0.8746)
 [AI] 6.435215(3.7816) 0.000000(8.6006) 5.023880(6.0105) 0.5616(0.3462) 0.0000(0.7506)
 [AI] 0.370329(5.1742) 0.000000(7.9743) 8.915135(5.7280) 0.0399(0.5516) 0.0000(0.8588)
  \texttt{[AI]} \ \ 4.729876 \\ (2.7959) \ \ 0.000000 \\ (6.4739) \ \ 6.070746 \\ (4.5053) \ \ 0.4379 \\ (0.2694) \ \ 0.0000 \\ (0.5994) 
  \texttt{[AI]} \ \ 1.498777 (4.6290) \ \ 0.000000 (7.8358) \ \ 8.671031 (5.6422) \ \ 0.1474 (0.4424) \ \ 0.0000 (0.7705) 
  \texttt{[AI]} \ \ 4.455326(3.6159) \ \ 0.000000(7.6755) \ \ 7.414271(5.4385) \ \ 0.3754(0.3047) \ \ 0.0000(0.6467) 
 [Convergence] NO(More iteration number is needed!)
 Done within 1s
# HE Regression
vc <- hiblup.he(y = pheno$t2[index], K = list(AD_GA, AD_GD))</pre>
```

With fixed effects and random effects:

```
index <- match(geno.id[, 1], pheno[, 1])

ADcal <- hiblup.K(M = geno, mode = "AD")
AD_GA <- ADcal$GA
AD_GD <- ADcal$GD

X <- model.matrix(~Sex, data = pheno)  # fixed effects
R <- as.matrix(pheno$Sire)  # random effects

# AI, EM, EMAI, AIEM, and HI algorithm
vc <- hiblup.vc(y = pheno$t2[index], X = X[index, ], R = R[index, ], K = list(AD_GA, AD_GD), method = "AI")

# HE algorithm
vc <- hiblup.he(y = pheno$t2[index], X = X[index, ], R = R[index, ], K = list(AD_GA, AD_GD))</pre>
```

4.3.3 Pairs of correlated traits

Variance and co-variance for pairs of correlated traits can be estimated using following codes. X1 and X2 are the fixed effects of trait1 and trait2, respectively.

4.3.4 With user-provided variance components

The start parameter is used to accept the initial value in the variance component calculation method containing AI or EM. The length of the start vector is equal to the number of K plus one. It should be noted that if R is specified, the value of V_R needs to be given in the first position in start. For pairs of correlated traits, the elements in start are $V_g^{(1)}$, $V_g^{(2)}$, $COV_g^{(12)}$, $V_e^{(1)}$, $V_e^{(2)}$, and $COV_e^{(12)}$;

```
print(start1)
[1] 0.755 4.180
print(start2)
[1] 1.27435 0.03995 3.61035
print(start3)
[1] 4.455 0.000 7.414
print(start4)
[1] 2.115 9.890 0.000 0.755
print(start5)
[1] 0.08812 0.01323 0.02905 3.58966 2.68928 2.66573
```

4.4 BLUP

4.4.1 Additive effect based model

if variance components are unknown:

```
# AI, EM, EMAI, AIEM, and HI algorithm
X <- model.matrix(~Sex, data = pheno) # fixed effects</pre>
R <- as.matrix(pheno$Sire) # random effects
# using HI algorithm if variance components are unknown
gebv.a.hi <- hiblup(pheno = pheno[, c(1, 5)], pedigree = pedigree, CV = X,</pre>
    R = R, vc.method = c("HI"), nAIiter = 5, mode = "A")
  -----Welcome to HIBLUP-----
  He-aI BLUP
              | | | | | _ | _ | _ \| | | | | | | | _ _ \
              | __ | | | | _ <| | | | | | | | ___/
| | | | | | | | | | | | | | |
              |_| |_|___|__/|___\___/|_| Version: 1.1.0 #
ABLUP model is selected based on the provided data
 Analyzed trait: t2
 Number of fixed effects: 2 (intercept included)
Number of random effects: 1
 Number of individuals with phenotypic observations: 800
 Deriving A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Realign index of R matrix...Done!
 HE Prior derived: A:1.007 e:1.66; Done within Os
 HE adopted: TRUE
 Variance components estimation:
 [Iter] Var_R1(SE) Var_K1(SE)
                                        Var_e(SE)
                                                        h2_R1(SE)
                                                                       h2_K1(SE)
 [AI] 0.224987(0.1201) 0.533193(0.4132) 1.948811(0.2899) 0.0831(0.0441) 0.1970(0.1477)
 [AI] 0.242646(0.1355) 0.584247(0.3266) 1.918416(0.2514) 0.0884(0.0472) 0.2128(0.1169)
 [AI] 0.245346(0.1459) 0.582230(0.3415) 1.919892(0.2591) 0.0893(0.0506) 0.2119(0.1219)
 [AI] 0.245362(0.1467) 0.582361(0.3412) 1.919794(0.2590) 0.0893(0.0508) 0.2120(0.1218)
 [AI] 0.245364(0.1467) 0.582351(0.3412) 1.919801(0.2590) 0.0893(0.0508) 0.2120(0.1218)
 [Convergence] NO(More iteration number is needed!)
 Done within 1s
 Estimated beta: 11.52 0.8442
 Estimated Vg and Ve: 0.5824 1.92
 HIBLUP IS DONE WITHIN: 1s
 HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.4.2 Additive and Dominant effect based model

```
gebv.ad <- hiblup(pheno = pheno[, c(1, 5)], pedigree = pedigree, mode = "AD")
#-----#</pre>
```

```
# He-aI BLUP
              #
                                                                #
              #
                                                                #
#
              | __ | | | _ <| | | | | ___/
              |_| |_|___| Version: 1.1.0 #
 ABLUP model is selected based on the provided data
 Analyzed trait: t2
 Number of fixed effects: 1 (intercept included)
 Number of random effects: 0
 Number of individuals with phenotypic observations: 800
 Deriving A and D matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 HE Prior derived: A:1.216 D:0.8789 e:0.7516; Done within Os
 HE adopted: TRUE
 Variance components estimation:
 [Iter] Var_K1(SE) Var_K2(SE)
                                      Var_e(SE)
                                                     h2_K1(SE)
 [AI] 1.210780(0.3632) 0.421778(0.6900) 1.180488(0.5894) 0.4304(0.1140) 0.1499(0.2452)
 [AI] 1.196838(0.3523) 0.032851(0.6563) 1.560590(0.5740) 0.4289(0.1113) 0.0118(0.2352)
 [AI] 1.192947(0.3406) 0.067969(0.6261) 1.532128(0.5599) 0.4271(0.1075) 0.0243(0.2244)
 [AI] 1.194088(0.3405) 0.065163(0.6295) 1.534041(0.5621) 0.4275(0.1075) 0.0233(0.2255)
  \texttt{[AI]} \ \ 1.193903(0.3407) \ \ 0.065467(0.6294) \ \ 1.533882(0.5620) \ \ 0.4274(0.1076) \ \ 0.0234(0.2255) 
 [AI] 1.193931(0.3407) 0.065428(0.6294) 1.533898(0.5620) 0.4274(0.1076) 0.0234(0.2255)
 [AI] 1.193927(0.3407) 0.065434(0.6294) 1.533896(0.5620) 0.4274(0.1076) 0.0234(0.2255)
 [AI] 1.193928(0.3407) 0.065433(0.6294) 1.533897(0.5620) 0.4274(0.1076) 0.0234(0.2255)
 [Convergence] YES
 Done within 1s
 Estimated beta: 13.09
 Estimated additive genetic variacne: 1.194
 Estimated Dominance genetic variacne: 0.06543
 Estimated Ve: 1.534
 HIBLUP IS DONE WITHIN: 2s
 HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.4.3 With user-provided variance components

If the variance components are known and provided by the users, the methods for solving mixed model equation can be controlled by the mme.method parameter. The options are "solve" and "sor".

```
ABLUP model is selected based on the provided data
 Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 1
 Number of individuals with phenotypic observations: 800
 Deriving the inverse of A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
Realign index of R matrix...Done!
 Solving MME...
Done within 2s
 Estimated beta: 10.94 1.085
Estimated Vg and Ve: 0.03995 3.61
HIBLUP IS DONE WITHIN: 2s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
# Solve mixed model equation using SOR method
gebv.a <- hiblup(pheno = pheno[, c(1, 5)], pedigree = pedigree, CV = X,</pre>
   R = R, vc = start2, mme.method = "sor", mode = "A")
          ------Welcome to HIBLUP-----
# He-aI BLUP
             \| |
              | |__| | | | | |_) | | | | | | | |_) |
              | __ | | | _ <| | | | | ___/
              |_| |_|___| Version: 1.1.0 #
ABLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 1
Number of individuals with phenotypic observations: 800
Deriving the inverse of A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Realign index of R matrix...Done!
 Solving MME...
 using SOR method...
Final solution achieved after the 398 th iteration Done within 1m1s
Estimated beta: 10.93 1.117
Estimated Vg and Ve: 0.03995 3.61
HIBLUP IS DONE WITHIN: 1m1s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.4.4 Pairs of correlated traits

HIBLUP supports the estimation of individual genetic values for pairs of correlated traits. Users can specify the columns of the trait1 and trait2 in phenotype file by setting the bivar.pos parameter, for example:

```
gebv <- hiblup(pheno = pheno, bivar.pos = c(4, 5), X1 = X, X2 = X, pedigree = pedigree)
            -----Welcome to HIBLUP-----
# He-aI BLUP
              __ | | | | | < | |
                                     Version: 1.1.0 #
              |_| |_|___/|___/|_
Bivariate GREML analysis started...
ABLUP model is selected based on the provided data
Analyzed trait: t1 & t2
Number of phenotypic observations for trait1: 800
Number of phenotypic observations for trait2: 800
 Deriving A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 HE prior derived... Done within Os
 Variance components estimation:
 Dimension of V: 5048 * 5048
 Bivariate GREML analysis: V(G)_tr1 V(G)_tr2 C(G)_tr12 V(e)_tr1 V(e)_tr2 C(e)_tr12
 [AI] Iter 1 of Max Iter 20: 0.246632 0.287186 0.266138 2.341842 1.437926 0.912211
 [AI] Iter 2 of Max Iter 20: 0.440754 0.586023 0.469265 2.178774 1.436027 1.768835
 [AI] Iter 3 of Max Iter 20: 0.696731 0.899445 0.717611 2.500768 1.583474 1.983699
 [AI] Iter 4 of Max Iter 20: 0.873017 1.149052 0.888256 2.536012 1.508778 1.923990
 [AI] Iter 5 of Max Iter 20: 0.895045 1.240929 0.941279 2.606986 1.469086 1.844294
 [AI] Iter 6 of Max Iter 20: 0.888643 1.184977 0.990134 2.707947 1.522181 1.757735
 [AI] Iter 7 of Max Iter 20: 0.891684 1.092909 0.979085 2.768846 1.600671 1.725523
 [AI] Iter 8 of Max Iter 20: 0.885224 1.094455 0.976973 2.790742 1.606972 1.717332
 [AI] Iter 9 of Max Iter 20: 0.885279 1.094424 0.977076 2.791618 1.607371 1.716731
 [AI] Iter 10 of Max Iter 20: 0.885246 1.094405 0.977054 2.791649 1.607387 1.716748
 [AI] Iter 11 of Max Iter 20: 0.885249 1.094407 0.977057 2.791647 1.607385 1.716746
 [AI] Iter 12 of Max Iter 20: 0.885249 1.094407 0.977057 2.791648 1.607385 1.716746
 [Convergence] YES
Done within 19s
Estimated beta1: 161.3 0.9368
Estimated beta2: 11.45 0.8637
Estimated Vg1 and Ve1: 0.8852 2.792
Estimated Vg2 and Ve2: 1.094 1.607
 Estimated COVg1g2 and COVe1e2: 0.9771 1.717
Estimated genetic correlation: 0.9927
HIBLUP IS DONE WITHIN: 20s
 HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.4.5 Reliability of individual genetic value

The boolean parameter reliability is used to specify whether to calculate the reliability of each individual's genetic value.

```
X <- model.matrix(~Sex, data = pheno) # fixed effects
R <- as.matrix(pheno$Sire) # random effects</pre>
```

```
# get the reliability of individual genetic value
 gebv.a.hi <- hiblup(pheno = pheno[, c(1, 5)], pedigree = pedigree, CV = X,</pre>
         R = R, vc.method = c("HI"), nAliter = 5, mode = "A", reliability = TRUE)
 #-----#
 # He-aI BLUP
                                       | |__| | | | | |_) | |
                                                                                                    | | | | |__) |
                                                 _ | | | _ <| |
                                                                                                  ____/|_|
                                                                                                                                   Version: 1.1.0 #
  ABLUP model is selected based on the provided data
   Analyzed trait: t2
   Number of fixed effects: 2 (intercept included)
   Number of random effects: 1
   Number of individuals with phenotypic observations: 800
   Deriving A matrix from pedigree...Done within Os
   Number of total predicted individuals: 2524
   Realign index of y...Done!
   Realign index of X matrix...Done!
   Realign index of R matrix...Done!
    HE Prior derived: A:1.007 e:1.66; Done within Os
    HE adopted: TRUE
    Variance components estimation:
    [Iter] Var_R1(SE)
                                                      Var_K1(SE)
                                                                                                        Var_e(SE) h2_R1(SE)
    [AI] 0.224987(0.1201) 0.533193(0.4132) 1.948811(0.2899) 0.0831(0.0441) 0.1970(0.1477)
     \texttt{[AI]} \quad 0.242646(0.1355) \quad 0.584247(0.3266) \quad 1.918416(0.2514) \quad 0.0884(0.0472) \quad 0.2128(0.1169) 
     \texttt{[AI]} \quad 0.245346 \\ (0.1459) \quad 0.582230 \\ (0.3415) \quad 1.919892 \\ (0.2591) \quad 0.0893 \\ (0.0506) \quad 0.2119 \\ (0.1219) \quad 0.0893 \\ (0.0506) \quad 0.0893 
     \texttt{[AI]} \quad 0.245362(0.1467) \quad 0.582361(0.3412) \quad 1.919794(0.2590) \quad 0.0893(0.0508) \quad 0.2120(0.1218) 
     \texttt{[AI]} \quad 0.245364(0.1467) \quad 0.582351(0.3412) \quad 1.919801(0.2590) \quad 0.0893(0.0508) \quad 0.2120(0.1218) 
    [Convergence] NO(More iteration number is needed!)
   Done within 1s
    Estimated beta: 11.52 0.8442
   Estimated Vg and Ve: 0.5824 1.92
    Calculating SEP and reliability... Done!
   HIBLUP IS DONE WITHIN: 5s
   HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.5 GBLUP

4.5.1 Additive effect based model

```
| | | | _ <| | | | | | | .
              #
#
              |_| |_|___| Version: 1.1.0 #
GBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 0
 Number of individuals with phenotypic observations: 800
 Deriving GA matrix from genotype...Done within 1s
 Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Number of total predicted individuals: 573
 Realign index of y...Done!
Realign index of X matrix...Done!
 Variance components estimation:
 [Iter] Var_K1(SE) Var_e(SE)
                                      h2_K1(SE)
 [AI] 1.265440(1.1820) 3.659904(0.8541) 0.2569(0.2182)
 [AI] 0.908219(0.9788) 4.057931(0.8841) 0.1829(0.1889)
 [AI] 0.807857(0.8896) 4.156277(0.8818) 0.1627(0.1743)
 [AI] 0.778040(0.8578) 4.183374(0.8754) 0.1568(0.1688)
 [AI] 0.768762(0.8476) 4.191671(0.8728) 0.1550(0.1671)
 [AI] 0.765820(0.8443) 4.194291(0.8719) 0.1544(0.1665)
 [AI] 0.764881(0.8433) 4.195126(0.8716) 0.1542(0.1663)
 [AI] 0.764581(0.8430) 4.195393(0.8715) 0.1542(0.1662)
 [AI] 0.764485(0.8429) 4.195478(0.8715) 0.1541(0.1662)
 [AI] 0.764454(0.8428) 4.195506(0.8715) 0.1541(0.1662)
 [AI] 0.764444(0.8428) 4.195515(0.8715) 0.1541(0.1662)
 [AI] 0.764441(0.8428) 4.195517(0.8715) 0.1541(0.1662)
 [Convergence] YES
 Done within 1s
 Estimated beta: 12.4 0.4291
 Estimated Vg and Ve: 0.7644 4.196
HIBLUP IS DONE WITHIN: 2s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.5.2 Additive and Dominant effect based model

```
gebv.ad <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,
  mode = "AD")
       ------Welcome to HIBLUP-----
# He-aI BLUP
          | | | | __ \
#
          #
#
          | __ | | | _ <| | | | | ___/
          #
          |_| |_|___| Version: 1.1.0 #
GBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 1 (intercept included)
```

```
Number of random effects: 0
Number of individuals with phenotypic observations: 800
Deriving GA and GD matrix from genotype...Done within 3s
Number of genotyped individuals: 573
Number of genotyped individuals with phenotypic observations: 175
Number of genotyped individuals without phenotypic observations: 398
Number of total predicted individuals: 573
Realign index of y...Done!
Realign index of X matrix...Done!
HE Prior derived: A:0.9797 D:0.1 e:5.809; Done within Os
HE adopted: TRUE
Variance components estimation:
[Iter] Var_K1(SE)
                     Var_K2(SE)
                                                              h2_K1(SE)
                                           Var_e(SE)
                                                                               h2 K2(SE)
[AI] 1.006840(2.0141) 0.005793(4.2850) 5.831226(3.0315) 0.1471(0.2935) 0.0008(0.6261)
[AI] 1.008473(1.9961) 0.000119(4.2377) 5.832540(3.0023) 0.1474(0.2908) 0.0000(0.6194)
[AI] 1.008498(1.9950) 0.000030(4.2348) 5.832560(3.0006) 0.1474(0.2907) 0.0000(0.6190)
[AI] 1.008504(1.9950) 0.000008(4.2348) 5.832565(3.0005) 0.1474(0.2907) 0.0000(0.6190)
 \texttt{[AI]} \ \ 1.008506(1.9950) \ \ 0.000003(4.2347) \ \ 5.832567(3.0005) \ \ 0.1474(0.2907) \ \ 0.0000(0.6190) 
[AI] 2.675951(1.9950) 0.000000(4.2347) 7.173500(3.0005) 0.2717(0.2020) 0.0000(0.4299)
[AI] 2.278828(3.7618) 0.000000(7.2427) 9.069668(5.1909) 0.2008(0.3250) 0.0000(0.6382)
[AI] 3.919482(4.4465) 0.000000(9.0505) 8.167389(6.4510) 0.3243(0.3629) 0.0000(0.7488)
 \texttt{[AI]} \ \ 0.238759 \\ (5.2541) \ \ 0.000000 \\ (9.7090) \ \ 10.713283 \\ (6.9766) \ \ 0.0218 \\ (0.4783) \ \ 0.0000 \\ (0.8865) 
[AI] 6.499274(3.4791) 0.000000(8.1872) 4.207605(5.6764) 0.6070(0.3503) 0.0000(0.7647)
 \texttt{[AI]} \quad 0.751425(4.6846) \quad 0.000000(6.9762) \quad 8.034977(5.0013) \quad 0.0855(0.5209) \quad 0.0000(0.7940) 
[AI] 4.129675(2.7308) 0.000000(6.0881) 6.901682(4.2738) 0.3744(0.2519) 0.0000(0.5519)
[AI] 1.845106(4.6792) 0.000000(8.3173) 8.877667(5.9854) 0.1721(0.4246) 0.0000(0.7757)
[AI] 4.316535(3.9961) 0.000000(8.3194) 7.719906(5.9122) 0.3586(0.3298) 0.0000(0.6912)
[AI] 1.739195(5.3001) 0.000000(9.5363) 9.506922(6.8601) 0.1546(0.4604) 0.0000(0.8480)
[AI] 4.943089(4.2312) 0.000000(8.9304) 7.200726(6.3334) 0.4070(0.3482) 0.0000(0.7354)
[AI] 1.212187(5.4675) 0.000000(9.4768) 9.663354(6.8230) 0.1115(0.4928) 0.0000(0.8714)
[AI] 5.502895(3.8664) 0.000000(8.4434) 6.428994(5.9524) 0.4612(0.3294) 0.0000(0.7076)
 \texttt{[AI]} \quad 0.797314(5.4076) \quad 0.000000(8.9926) \quad 9.523833(6.4745) \quad 0.0773(0.5156) \quad 0.0000(0.8713) 
 \texttt{[AI]} \ \ 5.476338(3.4434) \ \ 0.000000(7.7282) \ \ 6.061720(5.4172) \ \ 0.4746(0.3076) \ \ 0.0000(0.6698) 
[Convergence] NO(More iteration number is needed!)
Done within 1s
Estimated beta: 13.24
Estimated additive genetic variacne: 5.476
Estimated Dominance genetic variacne: 0
Estimated Ve: 6.062
HIBLUP IS DONE WITHIN: 4s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.5.3 With user-provided variance components

If the variance components are known and provided by the users, the methods for solving mixed model equation can be controlled by the mme.method parameter. The options are "solve" and "sor".

```
| |__| | | | | | _) | | | | | | | | | __) |
              | __ | | | _ <| | | | | | ___/
#
              | | | | | | | | | | |
                                                               #
                                  ____\__/|_| Version: 1.1.0 #
GBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 1
 Number of individuals with phenotypic observations: 800
 Deriving GA matrix from genotype...Done within 1s
 Number of genotyped individuals: 573
Number of genotyped individuals with phenotypic observations: 175
Number of genotyped individuals without phenotypic observations: 398
Number of total predicted individuals: 573
Realign index of y...Done!
Realign index of X matrix...Done!
Realign index of R matrix...Done!
 Solving MME...
Done within Os
 Estimated Vg and Ve: 0.03995 3.61
HIBLUP IS DONE WITHIN: 2s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
# Solve mixed model equation using SOR method
gebv.a <- hiblup(pheno = pheno[, c(1, 5)], mme.method = "sor", CV = X,
   R = R, vc = start2, geno = geno, map = map, geno.id = geno.id)
           -----Welcome to HIBLUP-----
# He-aI BLUP
             | | | | | _ | _ | _ | | | | | | | _ _ \
#
             | |__| | | | | |_) | | | | | | | |__) |
             | __ | | | _ <| |
                                    #
             |_| |_|__/|___/|__
                                                Version: 1.1.0 #
\ensuremath{\mathtt{GBLUP}} model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 1
Number of individuals with phenotypic observations: 800
Deriving GA matrix from genotype...Done within 1s
Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
Number of total predicted individuals: 573
Realign index of y...Done!
Realign index of X matrix...Done!
 Realign index of R matrix...Done!
 Solving MME...
 using SOR method...
 Final solution achieved after the 2 th iteration Done within Os
 Estimated beta: 12.41 13.23
Estimated Vg and Ve: 0.03995 3.61
```

```
HIBLUP IS DONE WITHIN: 1s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.5.4 Estimate the marker effects

HIBLUP will output the marker effects if back.solution is TRUE. it only works with GBLUP model or SSBLUP model.

```
gebv.a.ai <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,
   CV = X, vc.method = c("AI"), mode = "A", back.solution = TRUE)
#-----Welcome to HIBLUP-----
# He-aI BLUP
              | | | | |_
                                      | | | | __ \
              | |__| | | | | |_) | | | | | | | | |_) |
              | __ | | | _ <| |
                                     | | | | ___/
#
              Version: 1.1.0 #
{\tt GBLUP} model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 0
Number of individuals with phenotypic observations: 800
Deriving GA matrix from genotype...Done within 1s
Number of genotyped individuals: 573
Number of genotyped individuals with phenotypic observations: 175
Number of genotyped individuals without phenotypic observations: 398
Number of total predicted individuals: 573
Realign index of y...Done!
Realign index of X matrix...Done!
Variance components estimation:
[Iter] Var_K1(SE)
                     Var_e(SE)
                                       h2_K1(SE)
[AI] 1.265440(1.1820) 3.659904(0.8541) 0.2569(0.2182)
[AI] 0.908219(0.9788) 4.057931(0.8841) 0.1829(0.1889)
[AI] 0.807857(0.8896) 4.156277(0.8818) 0.1627(0.1743)
[AI] 0.778040(0.8578) 4.183374(0.8754) 0.1568(0.1688)
[AI] 0.768762(0.8476) 4.191671(0.8728) 0.1550(0.1671)
[AI] 0.765820(0.8443) 4.194291(0.8719) 0.1544(0.1665)
[AI] 0.764881(0.8433) 4.195126(0.8716) 0.1542(0.1663)
[AI] 0.764581(0.8430) 4.195393(0.8715) 0.1542(0.1662)
[AI] 0.764485(0.8429) 4.195478(0.8715) 0.1541(0.1662)
[AI] 0.764454(0.8428) 4.195506(0.8715) 0.1541(0.1662)
[AI] 0.764444(0.8428) 4.195515(0.8715) 0.1541(0.1662)
[AI] 0.764441(0.8428) 4.195517(0.8715) 0.1541(0.1662)
[Convergence] YES
Done within 1s
Estimating SNP effect...Done within Os
Estimated beta: 12.4 0.4291
Estimated Vg and Ve: 0.7644 4.196
HIBLUP IS DONE WITHIN: 2s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.5.5 Pairs of correlated traits

HIBLUP supports the estimation of individual genetic values for pairs of correlated traits. Users can specify the position of the trait1 and trait2 in pheno by setting the bivar.pos parameter, for example:

```
gebv <- hiblup(pheno = pheno, bivar.pos = c(4, 5), X1 = X, X2 = X, map = map,
   geno = geno, geno.id = geno.id)
#-----#
# He-aI BLUP
             | __ | | | _ <| | | | | ___/
             |_| |_|___| Version: 1.1.0 #
Bivariate GREML analysis started...
{\tt GBLUP} model is selected based on the provided data
Analyzed trait: t1 & t2
Number of phenotypic observations for trait1: 800
Number of phenotypic observations for trait2: 800
Deriving GA matrix from genotype...Done within 1s
Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
Number of genotyped individuals without phenotypic observations: 398
Number of total predicted individuals: 573
Realign index of y...Done!
Realign index of X matrix...Done!
 HE prior derived... Done within Os
 Variance components estimation:
 Dimension of V: 1146 * 1146
 Bivariate GREML analysis: V(G)_tr1 V(G)_tr2 C(G)_tr12 V(e)_tr1 V(e)_tr2 C(e)_tr12
 [AI] Iter 1 of Max Iter 20: 0.611658 0.901322 0.288246 1.948774 1.707617 1.824215
 [AI] Iter 2 of Max Iter 20: 0.743415 0.869007 0.474235 3.193945 2.820537 2.838791
 [AI] Iter 3 of Max Iter 20: 0.503254 0.777144 0.568722 4.559059 3.791561 3.772494
 [AI] Iter 4 of Max Iter 20: 0.284674 0.738731 0.458582 5.292587 4.160313 4.216107
 [AI] Iter 5 of Max Iter 20: 0.294574 0.732244 0.464436 5.368836 4.195966 4.252565
 [AI] Iter 6 of Max Iter 20: 0.292856 0.731214 0.462753 5.367915 4.196675 4.252062
 [AI] Iter 7 of Max Iter 20: 0.292640 0.730558 0.462375 5.368194 4.197077 4.252404
 [AI] Iter 8 of Max Iter 20: 0.292521 0.730428 0.462240 5.368281 4.197194 4.252506
 [AI] Iter 9 of Max Iter 20: 0.292492 0.730385 0.462203 5.368309 4.197226 4.252536
 [AI] Iter 10 of Max Iter 20: 0.292482 0.730373 0.462192 5.368317 4.197236 4.252545
 [AI] Iter 11 of Max Iter 20: 0.292480 0.730370 0.462189 5.368320 4.197239 4.252547
 [AI] Iter 12 of Max Iter 20: 0.292479 0.730369 0.462188 5.368320 4.197239 4.252548
 [AI] Iter 13 of Max Iter 20: 0.292479 0.730368 0.462188 5.368320 4.197240 4.252548
 [Convergence] YES
 Done within 1s
Estimated beta1: 163 0.05853
Estimated beta2: 13.63 -0.2017
 Estimated Vg1 and Ve1: 0.2925 5.368
 Estimated Vg2 and Ve2: 0.7304 4.197
 Estimated COVg1g2 and COVe1e2: 0.4622 4.253
 Estimated genetic correlation: 1
 HIBLUP IS DONE WITHIN: 2s
 HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.5.6 Reliability of individual genetic value

The boolean parameter reliability is used to specify whether to calculate the reliability of each individual's genetic value.

```
X <- model.matrix(~Sex, data = pheno) # fixed effects</pre>
gebv.a.ai <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,
   CV = X, vc.method = c("AI"), mode = "A", reliability = TRUE)
#-----#
# He-aI BLUP
             #
              | |__| | | | | |_) | | | | | | | |_) |
              | __ | | | | _ <| | | | | | ___/
                  | |_| |_| |_) | |___| |_
              |_| |_|___/|___\___/|_| Version: 1.1.0 #
GBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
 Number of random effects: 0
 Number of individuals with phenotypic observations: 800
 Deriving GA matrix from genotype...Done within 1s
 Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Number of total predicted individuals: 573
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Variance components estimation:
                                      h2_K1(SE)
 [Iter] Var_K1(SE) Var_e(SE)
 [AI] 1.265440(1.1820) 3.659904(0.8541) 0.2569(0.2182)
 [AI] 0.908219(0.9788) 4.057931(0.8841) 0.1829(0.1889)
 [AI] 0.807857(0.8896) 4.156277(0.8818) 0.1627(0.1743)
 [AI] 0.778040(0.8578) 4.183374(0.8754) 0.1568(0.1688)
 [AI] 0.768762(0.8476) 4.191671(0.8728) 0.1550(0.1671)
 [AI] 0.765820(0.8443) 4.194291(0.8719) 0.1544(0.1665)
 [AI] 0.764881(0.8433) 4.195126(0.8716) 0.1542(0.1663)
 [AI] 0.764581(0.8430) 4.195393(0.8715) 0.1542(0.1662)
 [AI] 0.764485(0.8429) 4.195478(0.8715) 0.1541(0.1662)
 [AI] 0.764454(0.8428) 4.195506(0.8715) 0.1541(0.1662)
 [AI] 0.764444(0.8428) 4.195515(0.8715) 0.1541(0.1662)
 [AI] 0.764441(0.8428) 4.195517(0.8715) 0.1541(0.1662)
 [Convergence] YES
 Done within 1s
 Estimated beta: 12.4 0.4291
 Estimated Vg and Ve: 0.7644 4.196
 Calculating SEP and reliability... Done!
 HIBLUP IS DONE WITHIN: 2s
 HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.6 SSBLUP

4.6.1 Additive effect based model

```
gebv.a.ai <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,
   pedigree = pedigree, vc.method = c("AI"), mode = "A")
       -----Welcome to HIBLUP-----
  He-aI BLUP
                                    | | | | __ \
              | | | | _ | _ \| |
              | __ | | | | _ <| | | | | | _ -
                                                 Version: 1.1.0 #
              |_| |_|___/|___\___/|_|
SSBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 1 (intercept included)
Number of random effects: 0
Number of individuals with phenotypic observations: 800
Deriving GA matrix from genotype...Done within 1s
Number of genotyped individuals: 573
Number of genotyped individuals with phenotypic observations: 175
Number of genotyped individuals without phenotypic observations: 398
Deriving A matrix from pedigree...Done within Os
Number of total predicted individuals: 2524
Realign index of y...Done!
Realign index of X matrix...Done!
 Extracting A11 matrix...Done!
Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Constructing HA matrix...Done within 1s
 Variance components estimation:
 [Iter] Var_K1(SE)
                     Var_e(SE)
                                       h2_K1(SE)
 [AI] 1.219009(0.3209) 1.648960(0.2440) 0.4250(0.0976)
 [AI] 1.267629(0.3044) 1.619052(0.2463) 0.4391(0.0933)
 [AI] 1.258991(0.3125) 1.625889(0.2497) 0.4364(0.0955)
 [AI] 1.260852(0.3112) 1.624455(0.2493) 0.4370(0.0952)
 [AI] 1.260460(0.3115) 1.624759(0.2494) 0.4369(0.0952)
 [AI] 1.260543(0.3114) 1.624695(0.2494) 0.4369(0.0952)
 [AI] 1.260525(0.3114) 1.624708(0.2494) 0.4369(0.0952)
 [AI] 1.260529(0.3114) 1.624705(0.2494) 0.4369(0.0952)
 [AI] 1.260528(0.3114) 1.624706(0.2494) 0.4369(0.0952)
 [Convergence] YES
 Done within 1s
 Estimated beta: 13.09
 Estimated Vg and Ve: 1.261 1.625
 HIBLUP IS DONE WITHIN: 4s
 HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.6.2 Additive and Dominant effect based model

```
gebv.ad <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,</pre>
    pedigree = pedigree, mode = "AD")
                ------Welcome to HIBLUP-----
# He-aI BLUP
               | |__| | | | | |_) | | | | | | | |_) |
                | __ | | | _ <| | | | | ___/
                |_| |_|___| Version: 1.1.0 #
 SSBLUP model is selected based on the provided data
 Analyzed trait: t2
 Number of fixed effects: 1 (intercept included)
 Number of random effects: 0
 Number of individuals with phenotypic observations: 800
 Deriving GA and GD matrix from genotype...Done within 3s
 Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Deriving A and D matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Constructing HA matrix...Done within 1s
 Extracting D11 matrix...Done!
 Mean of diagonal and Off-diagonal of PD: 1 0.001
 Mean of diagonal and Off-diagonal of GD: 0.5374 -0.0009
 Adjusting GD matrix: GD* = 1.86 * GD + 0
 Weighting of D11 and GD matrix: 0.05
 Calculating the inverse of D11 matrix...Done within Os
 Constructing HD matrix...Done within 1s
 HE Prior derived: A:0.5493 D:0.1 e:5.179; Done within Os
 HE adopted: TRUE
 Variance components estimation:
 [Iter] Var_K1(SE)
                      Var_K2(SE)
                                             Var_e(SE)
                                                              h2_K1(SE)
                                                                               h2_K2(SE)
 [AI] 0.576766(0.5657) 0.045559(1.8438) 5.153651(1.5875) 0.0999(0.0984) 0.0079(0.3192)
  \texttt{[AI]} \ \ 0.590231 \\ (0.5659) \ \ 0.019153 \\ (1.8193) \ \ 5.140993 \\ (1.5658) \ \ 0.1026 \\ (0.0988) \ \ 0.0033 \\ (0.3164) 
  \texttt{[AI]} \ \ 0.596895 \\ (0.5660) \ \ 0.006147 \\ (1.8073) \ \ 5.134717 \\ (1.5552) \ \ 0.1040 \\ (0.0990) \ \ 0.0011 \\ (0.3150) 
  \texttt{[AI]} \ \ 0.598553 \\ (0.5660) \ \ 0.002919 \\ (1.8015) \ \ 5.133154 \\ (1.5500) \ \ 0.1044 \\ (0.0991) \ \ 0.0005 \\ (0.3141) 
 [AI] 0.599381(0.5660) 0.001309(1.8000) 5.132373(1.5487) 0.1045(0.0991) 0.0002(0.3140)
 [AI] 0.599794(0.5660) 0.000504(1.7993) 5.131983(1.5481) 0.1046(0.0991) 0.0001(0.3139)
 [AI] 0.600001(0.5660) 0.000102(1.7989) 5.131788(1.5477) 0.1047(0.0991) 0.0000(0.3138)
  \texttt{[AI]} \ \ 0.600053(0.5660) \ \ 0.000002(1.7987) \ \ 5.131740(1.5476) \ \ 0.1047(0.0991) \ \ 0.0000(0.3138) 
  \texttt{[AI]} \ \ 3.986695(0.5660) \ \ 0.000000(1.7987) \ \ 1.937711(1.5475) \ \ 0.6729(0.1070) \ \ 0.0000(0.3036) 
  \texttt{[AI]} \ \ 1.556933(1.3225) \ \ 0.000000(1.8229) \ \ 5.053256(1.3631) \ \ 0.2355(0.1871) \ \ 0.0000(0.2758) 
 [AI] 5.146880(0.9594) 0.000000(2.3560) 3.841720(1.9466) 0.5726(0.1048) 0.0000(0.2621)
  \texttt{[AI]} \quad 0.841125(2.3223) \quad 0.000000(3.5623) \quad 8.719804(2.7201) \quad 0.0880(0.2374) \quad 0.0000(0.3726) 
 [AI] 5.233039(1.1566) 0.000000(3.8189) 0.783908(3.3068) 0.8697(0.2360) 0.0000(0.6347)
```

```
[AI] 0.502300(1.4217) 0.000000(1.6389) 4.708608(1.1791) 0.0964(0.2642) 0.0000(0.3145)
[AI] 3.338258(0.4782) 0.000000(1.5484) 2.327104(1.3364) 0.5892(0.0932) 0.0000(0.2733)
[AI] 0.782384(1.1761) 0.000000(1.7709) 7.281153(1.3475) 0.0970(0.1422) 0.0000(0.2196)
[AI] 3.965032(0.9223) 0.000000(2.9823) 2.452116(2.5733) 0.6179(0.1600) 0.0000(0.4647)
[AI] 1.688102(1.4452) 0.000000(2.1093) 5.660310(1.5954) 0.2297(0.1849) 0.00000(0.2870)
[AI] 6.018878(1.1120) 0.000000(2.7586) 3.335304(2.2828) 0.6434(0.1178) 0.00000(0.2949)
[AI] 2.540541(2.5828) 0.000000(3.6694) 6.545733(2.7606) 0.2796(0.2631) 0.0000(0.4038)
[Convergence] NO(More iteration number is needed!)
Done within 3s
Estimated beta: 13.12
Estimated additive genetic variacne: 2.541
Estimated Dominance genetic variacne: 0
Estimated Ve: 6.546
HIBLUP IS DONE WITHIN: 9s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.6.3 With user-provided variance components

If the variance components are known and provided by the users, the methods for solving mixed model equation can be controlled by the mme.method parameter. The options are "solve" and "sor".

```
# Solve mixed model equation directly
gebv.a <- hiblup(pheno = pheno[, c(1, 5)], mme.method = "solve", CV = X,</pre>
   R = R, vc = c(57.893, 0.0686, 0.0008), geno = geno, map = map, geno.id = geno.id,
   pedigree = pedigree)
#-----#
# He-aI BLUP
            #
            SSBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 1
Number of individuals with phenotypic observations: 800
Deriving GA matrix from genotype...Done within 1s
Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Deriving the inverse of A matrix from pedigree...Done within Os
 Deriving A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Realign index of R matrix...Done!
 Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
```

```
Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Calculating the inverse of Gw.A...Done within Os
 Constructing the inverse of HA matrix...Done within Os
 Solving MME...
 Done within 2s
 Estimated beta: 11.5 0.8719
 Estimated Vg and Ve: 0.0686 0.0008
 HIBLUP IS DONE WITHIN: 4s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
# Solve mixed model equation using SOR method
gebv.a <- hiblup(pheno = pheno[, c(1, 5)], mme.method = "sor", CV = X,
   R = R, vc = c(57.893, 0.0686, 0.0008), geno = geno, map = map, geno.id = geno.id,
   pedigree = pedigree)
           -----#
# He-aI BLUP
              | | | | __ \
                                    | | | | | | ___/
              | |__| | | | | |_) | |
                 __ | | | | | <| |
              |_| |_|___|___/|_________/|_| Version: 1.1.0 #
SSBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 2 (intercept included)
Number of random effects: 1
Number of individuals with phenotypic observations: 800
 Deriving GA matrix from genotype...Done within 1s
 Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Deriving the inverse of A matrix from pedigree...Done within Os
 Deriving A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Realign index of R matrix...Done!
 Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
Calculating the inverse of Gw.A...Done within Os
 Constructing the inverse of HA matrix...Done within Os
 Solving MME...
 using SOR method...
 Final solution achieved after the 26 th iteration Done within 4s
 Estimated beta: 10.21 12.86
Estimated Vg and Ve: 0.0686 0.0008
HIBLUP IS DONE WITHIN: 6s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.6.4 Estimate the marker effects

```
gebv.a.ai <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,
   pedigree = pedigree, vc.method = c("AI"), mode = "A", back.solution = TRUE)
#-----#
# He-aI BLUP
             | | | | __ \
             | | | | ___/
             | __ | | | < | |
             |_| |_|___| Version: 1.1.0 #
SSBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 1 (intercept included)
Number of random effects: 0
Number of individuals with phenotypic observations: 800
Deriving GA matrix from genotype...Done within 1s
Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
Number of genotyped individuals without phenotypic observations: 398
Deriving A matrix from pedigree...Done within Os
Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Constructing HA matrix...Done within 1s
 Variance components estimation:
 [Iter] Var_K1(SE) Var_e(SE)
                                    h2 K1(SE)
 [AI] 1.219009(0.3209) 1.648960(0.2440) 0.4250(0.0976)
 [AI] 1.267629(0.3044) 1.619052(0.2463) 0.4391(0.0933)
 [AI] 1.258991(0.3125) 1.625889(0.2497) 0.4364(0.0955)
 [AI] 1.260852(0.3112) 1.624455(0.2493) 0.4370(0.0952)
 [AI] 1.260460(0.3115) 1.624759(0.2494) 0.4369(0.0952)
 [AI] 1.260543(0.3114) 1.624695(0.2494) 0.4369(0.0952)
 [AI] 1.260525(0.3114) 1.624708(0.2494) 0.4369(0.0952)
 [AI] 1.260529(0.3114) 1.624705(0.2494) 0.4369(0.0952)
 [AI] 1.260528(0.3114) 1.624706(0.2494) 0.4369(0.0952)
 [Convergence] YES
 Done within 1s
 Estimating SNP effect...Done within Os
 Estimated beta: 13.09
 Estimated Vg and Ve: 1.261 1.625
HIBLUP IS DONE WITHIN: 4s
 HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.6.5 Pairs of correlated traits

HIBLUP also supports the estimation of individual genetic values for pairs of correlated traits. Users can specify the position of the trait1 and trait2 in pheno by setting the bivar.pos parameter, for example:

```
gebv <- hiblup(pheno = pheno, bivar.pos = c(4, 5), X1 = X, X2 = X, pedigree = pedigree,
   map = map, geno = geno, geno.id = geno.id)
           -----#
# He-aI BLUP
              #
#
              | |__| | | | | |_) | | | | | | | |_) |
              #
 Bivariate GREML analysis started...
 SSBLUP model is selected based on the provided data
 Analyzed trait: t1 & t2
 Number of phenotypic observations for trait1: 800
 Number of phenotypic observations for trait2: 800
 Deriving GA matrix from genotype...Done within 1s
 Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Deriving A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Constructing HA matrix...Done within 1s
 HE prior derived... Done within Os
 Variance components estimation:
 Dimension of V: 5048 * 5048
 Bivariate GREML analysis: V(G)_tr1 V(G)_tr2 C(G)_tr12 V(e)_tr1 V(e)_tr2 C(e)_tr12
 [AI] Iter 1 of Max Iter 20: 0.238334 0.269981 0.253664 1.155193 0.910799 1.025743
 [AI] Iter 2 of Max Iter 20: 0.348750 0.411680 0.378797 1.928973 1.518909 1.711557
 [AI] Iter 3 of Max Iter 20: 0.367117 0.482653 0.420587 2.737928 2.150112 2.425515
 [AI] Iter 4 of Max Iter 20: 0.241355 0.413038 0.314937 3.088985 2.410107 2.725830
 [AI] Iter 5 of Max Iter 20: 0.262136 0.455499 0.329550 2.958427 2.269084 2.583601
 [AI] Iter 6 of Max Iter 20: 0.558726 0.723745 0.581862 2.651230 1.969817 2.266805
 [AI] Iter 7 of Max Iter 20: 0.848500 0.954179 0.800006 2.467214 1.767960 2.041913
 [AI] Iter 8 of Max Iter 20: 0.968468 1.036340 0.872640 2.490140 1.706635 1.947982
 [AI] Iter 9 of Max Iter 20: 0.930618 1.060095 0.906128 2.661935 1.694131 1.883434
 [AI] Iter 10 of Max Iter 20: 0.812805 1.063151 0.926385 2.867885 1.697317 1.837136
 [AI] Iter 11 of Max Iter 20: 0.796443 1.062906 0.918749 2.923089 1.699652 1.829465
 [AI] Iter 12 of Max Iter 20: 0.796688 1.065252 0.920164 2.926795 1.698147 1.827330
 [AI] Iter 13 of Max Iter 20: 0.796408 1.064933 0.919859 2.927031 1.698396 1.827557
 [AI] Iter 14 of Max Iter 20: 0.796453 1.064996 0.919914 2.926997 1.698348 1.827515
 [AI] Iter 15 of Max Iter 20: 0.796444 1.064983 0.919903 2.927004 1.698358 1.827524
 [AI] Iter 16 of Max Iter 20: 0.796446 1.064985 0.919905 2.927003 1.698355 1.827522
```

```
[AI] Iter 17 of Max Iter 20: 0.796446 1.064985 0.919904 2.927003 1.698356 1.827522 [Convergence] YES
Done within 27s
Estimated beta1: 161.1 1.023
Estimated beta2: 11.29 0.9509
Estimated Vg1 and Ve1: 0.7964 2.927
Estimated Vg2 and Ve2: 1.065 1.698
Estimated COVg1g2 and COVe1e2: 0.9199 1.828
Estimated genetic correlation: 0.9988
HIBLUP IS DONE WITHIN: 30s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

4.6.6 Reliability of individual genetic value

The boolean parameter reliability is used to specify whether to calculate the reliability of each individual's genetic value.

```
gebv.a.ai <- hiblup(pheno = pheno[, c(1, 5)], geno = geno, map = map, geno.id = geno.id,
   pedigree = pedigree, vc.method = c("AI"), mode = "A", reliability = TRUE)
               -----#elcome to HIBLUP-----#
# He-aI BLUP
             #
             | __ | | | _ <| | | | | ___/
             | | | | | | | | |
             |_| |_|___| Version: 1.1.0 #
SSBLUP model is selected based on the provided data
Analyzed trait: t2
Number of fixed effects: 1 (intercept included)
Number of random effects: 0
 Number of individuals with phenotypic observations: 800
 Deriving GA matrix from genotype...Done within 1s
 Number of genotyped individuals: 573
 Number of genotyped individuals with phenotypic observations: 175
 Number of genotyped individuals without phenotypic observations: 398
 Deriving A matrix from pedigree...Done within Os
 Number of total predicted individuals: 2524
 Realign index of y...Done!
 Realign index of X matrix...Done!
 Extracting A11 matrix...Done!
 Mean of diagonal and Off-diagonal of PA: 1.001 0.0285
 Mean of diagonal and Off-diagonal of GA: 0.9886 -0.0017
 Adjusting GA matrix: GA* = 0.98 * GA + 0.03
 Weighting of A11 and GA matrix: 0.05
 Calculating the inverse of A11 matrix...Done within Os
 Constructing HA matrix...Done within 1s
 Variance components estimation:
 [Iter] Var_K1(SE) Var_e(SE)
                                     h2_K1(SE)
 [AI] 1.219009(0.3209) 1.648960(0.2440) 0.4250(0.0976)
 [AI] 1.267629(0.3044) 1.619052(0.2463) 0.4391(0.0933)
 [AI] 1.258991(0.3125) 1.625889(0.2497) 0.4364(0.0955)
 \hbox{[AI] 1.260852} (0.3112) \ 1.624455 (0.2493) \ 0.4370 (0.0952)
```

```
[AI] 1.260460(0.3115) 1.624759(0.2494) 0.4369(0.0952)
[AI] 1.260543(0.3114) 1.624695(0.2494) 0.4369(0.0952)
[AI] 1.260525(0.3114) 1.624708(0.2494) 0.4369(0.0952)
[AI] 1.260529(0.3114) 1.624705(0.2494) 0.4369(0.0952)
[AI] 1.260528(0.3114) 1.624706(0.2494) 0.4369(0.0952)
[Convergence] YES
Done within 1s
Estimated beta: 13.09
Estimated Vg and Ve: 1.261 1.625
Calculating SEP and reliability... Done!
HIBLUP IS DONE WITHIN: 7s
HIBLUP ACCOMPLISHED SUCCESSFULLY!
```

5 Function support list of HIBLUP

	HIBLUP
Genotype	
Pedigree	$\sqrt{}$
Phenotype	$\sqrt{}$
Fixed effects	$\sqrt{}$
Random effects	$\sqrt{}$
Relationship matrix	
AI	
EM	$\sqrt{}$
EMAI	$\sqrt{}$
HE Regression	$\sqrt{}$
HI	$\sqrt{}$
Fixed effects	
Random effects	$\sqrt{}$
Repeated records	
ABLUP	$\sqrt{}$
GBLUP	$\sqrt{}$
SSBLUP	$\sqrt{}$
GEBV	
Effect	$\sqrt{}$
Reliability	$\sqrt{}$
	Pedigree Phenotype Fixed effects Random effects Relationship matrix AI EM EMAI HE Regression HI Fixed effects Random effects Random effects RabLUP GBLUP SSBLUP GEBV Effect

Table 4: Function support list of HIBLUP.

6 HIBLUP Biography

Date	Version	Event
Aug-2018	1.0	BLUP/GBLUP/SSBLUP
Sep-2018	1.01	Add function of calculating reliability
Oct-2018	1.02	Fixed some bugs
Jan-2019	1.1	Add Repeated Models