Package 'bayesammi'

October 15, 2024

Type Package Title Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model Version 0.2.0 Maintainer Muhammad Yaseen <myaseen208@gmail.com> **Description** Performs Bayesian estimation of the additive main effects and multiplicative interaction (AMMI) model. The method is explained in Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G. and Cornelius, P.L. (2011) (<doi:10.2135/cropsci2010.06.0343>). **Depends** R (>= 3.1)Imports dplyr, ggplot2, lme4, magrittr, MASS, mvtnorm, rlang, rstiefel, scales, stats, tibble, tidyr, tmvtnorm License GPL-2 **Encoding UTF-8** LazyData true RoxygenNote 7.3.2 Note 1. School of Mathematical & Statistical Sciences, Clemson University, Clemson, South Carolina, USA 2. Department of Mathematics & Statistics, University of Agriculture Faisalabad, Faisalabad, Pakistan NeedsCompilation no Author Muhammad Yaseen [aut, cre], Jose Crossa [aut, ctb], Sergio Perez-Elizalde [aut, ctb], Diego Jarquin [aut, ctb], Jose Miguel Cotes [aut, ctb], Kert Viele [aut, ctb], Genzhou Liu [aut, ctb], Paul L. Cornelius [aut, ctb]

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bayes_ammi Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model

Description

Performs Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model

Usage

```
bayes_ammi(.data, .y, .gen, .env, .rep, .nIter)
## Default S3 method:
bayes_ammi(.data, .y, .gen, .env, .rep, .nIter)
```

Arguments

.data	data.frame
. y	Response Variable
.gen	Genotypes Factor
.env	Environment Factor
.rep	Replication Factor
.nIter	Number of Iterations

Value

Genotype by Environment Interaction Model

Author(s)

```
    Muhammad Yaseen (<myaseen208@gmail.com>)
```

- 2. Jose Crossa (< j.crossa@cgiar.org>)
- 3. Sergio Perez-Elizalde (<sergiop@colpos.mx>)
- 4. Diego Jarquin (<diego.jarquin@gmail.com>)
- 5. Jose Miguel Cotes
- 6. Kert Viele
- 7. Genzhou Liu
- 8. Paul L. Cornelius

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
fm1 <-
 bayes_ammi(
    .data = Maiz
   , .у
            = y
   , .gen = entry
   , .env
          = site
   , .rep = rep
  , .nIter = 20
  )
names(fm1)
fm1$mu1
fm1$tau1
fm1$tao1
fm1$delta1
fm1$lambdas1
fm1$alphas1
fm1$gammas1
library(ggplot2)
Plot1Mu <-
  ggplot(data = fm1$mu1, mapping = aes(x = 1:nrow(fm1$mu1), y = mu)) +
  geom_line(color = "blue") +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = expression(mu), x = "Iterations") +
  theme_bw()
print(Plot1Mu)
```

```
Plot2Mu <-
  ggplot(data = fm1$mu1, mapping = aes(mu)) +
  geom_histogram() +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = "Frequency", x = expression(mu)) +
  theme_bw()
print(Plot2Mu)
Plot1Sigma2 <-
  ggplot(data = fm1\$tau1, mapping = aes(x = 1:nrow(fm1\$tau1), y = tau)) +
  geom_line(color = "blue") +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = expression(sigma^2), x = "Iterations") +
  theme_bw()
print(Plot1Sigma2)
Plot2Sigma2 <-
  ggplot(data = fm1$tau1, mapping = aes(tau)) +
  geom_histogram() +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = "Frequency", x = expression(sigma^2)) +
  theme_bw()
print(Plot2Sigma2)
# Plot of Alphas
Plot1Alpha1 <-
  ggplot(data = fm1$tao1, mapping = aes(x = 1:nrow(fm1$tao1), y = tao1)) +
  geom_line(color = "blue") +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = expression(alpha[1]), x = "Iterations") +
  theme_bw()
print(Plot1Alpha1)
Plot2Alpha1 <-
  ggplot(data = fm1$tao1, mapping = aes(tao1)) +
  geom_histogram() +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = "Frequency", x = expression(alpha[1])) +
  theme_bw()
print(Plot2Alpha1)
Plot1Alpha2 <-
  ggplot(data = fm1\$tao1, mapping = aes(x = 1:nrow(fm1\$tao1), y = tao2)) +
  geom_line(color = "blue") +
  scale_x_continuous(labels = scales::comma) +
```

```
scale_y_continuous(labels = scales::comma) +
  labs(y = expression(alpha[2]), x = "Iterations") +
  theme_bw()
print(Plot1Alpha2)
Plot2Alpha2 <-
  ggplot(data = fm1$tao1, mapping = aes(tao2)) +
  geom_histogram() +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = "Frequency", x = expression(alpha[2])) +
  theme_bw()
print(Plot2Alpha2)
# Plot of Betas
Plot1Beta1 <-
  ggplot(data = fm1\$delta1, mapping = aes(x = 1:nrow(fm1\$delta1), y = delta1)) +
  geom_line(color = "blue") +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = expression(beta[1]), x = "Iterations") +
  theme_bw()
print(Plot1Beta1)
Plot2Beta1 <-
  ggplot(data = fm1$delta1, mapping = aes(delta1)) +
  geom_histogram() +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = "Frequency", x = expression(beta[1])) +
  theme_bw()
print(Plot2Beta1)
Plot1Beta2 <-
  ggplot(data = fm1\$delta1, mapping = aes(x = 1:nrow(fm1\$delta1), y = delta2)) +
  geom_line(color = "blue") +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = expression(beta[2]), x = "Iterations") +
  theme_bw()
print(Plot1Beta2)
Plot2Beta2 <-
  ggplot(data = fm1$delta1, mapping = aes(delta2)) +
  geom_histogram() +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = "Frequency", x = expression(beta[2])) +
  theme_bw()
print(Plot2Beta2)
```

```
Plot1Beta3 <-
  ggplot(data = fm1$delta1, mapping = aes(x = 1:nrow(fm1$delta1), y = delta3)) +
  geom_line(color = "blue") +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = expression(beta[3]), x = "Iterations") +
  theme_bw()
print(Plot1Beta3)
Plot2Beta3 <-
  ggplot(data = fm1$delta1, mapping = aes(delta3)) +
  geom_histogram() +
  scale_x_continuous(labels = scales::comma) +
  scale_y_continuous(labels = scales::comma) +
  labs(y = "Frequency", x = expression(beta[3])) +
  theme_bw()
print(Plot2Beta3)
BiplotAMMI <-
  ggplot(data = fm1$alphas0, mapping = aes(x = alphas1, y = alphas2)) +
  geom_point() +
  geom_hline(yintercept = 0) +
  geom_vline(xintercept = 0) +
  geom_text(aes(label = 1:nrow(fm1$alphas0)),
            vjust = "inward", hjust = "inward") +
  geom\_point(data = fm1\$gammas0, mapping = aes(x = gammas1, y = gammas2)) +
  geom_segment(data = fm1$gammas0,
               aes(x = 0, y = 0, xend = gammas1, yend = gammas2),
               arrow = arrow(length = unit(0.2, "cm"))
               , alpha = 0.75, color = "red") +
  geom_text(data = fm1$gammas0,
            aes(x = gammas1, y = gammas2,
                label = paste0("E", 1:nrow(fm1$gammas0))),
            vjust = "inward", hjust = "inward") +
  scale_x_continuous(
    limits = c(-max(abs(c(range(fm1$alphas0[, 1:2], fm1$gammas0[, 1:2]))))
               , max(abs(c(range(fm1$alphas0[, 1:2], fm1$gammas0[, 1:2]))))) +
  scale_y_continuous(
    limits = c(-max(abs(c(range(fm1\$alphas0[, 1:2], fm1\$gammas0[, 1:2]))))
               , max(abs(c(range(fm1$alphas0[, 1:2], fm1$gammas0[, 1:2]))))) +
  labs(title = "MCO Method", x = expression(PC[1]), y = expression(PC[2])) +
  theme(plot.title = element_text(hjust = 0.5))
print(BiplotAMMI)
BiplotBayesAMMI <-
  ggplot(data = fm1$alphas1, mapping = aes(x = alphas1, y = alphas2)) +
  geom_point() +
  geom_hline(yintercept = 0) +
  geom_vline(xintercept = 0) +
```

e_eff

```
geom_text(aes(label = 1:nrow(fm1$alphas1)),
            vjust = "inward", hjust = "inward") +
 geom\_point(data = fm1\$gammas1, mapping = aes(x = gammas1, y = gammas2)) +
 geom_segment(data = fm1$gammas1,
              aes(x = 0, y = 0, xend = gammas1, yend = gammas2),
              arrow = arrow(length = unit(0.2, "cm"))
               , alpha = 0.75, color = "red") +
 geom_text(data = fm1$gammas1,
            aes(x = gammas1, y = gammas2,
               label = paste0("E", 1:nrow(fm1$gammas1))),
            vjust = "inward", hjust = "inward") +
 scale_x_continuous(
   limits = c(-max(abs(c(range(fm1$alphas1[, 1:2], fm1$gammas1[, 1:2]))))
               , \max(abs(c(range(fm1\$alphas1[, 1:2], fm1\$gammas1[, 1:2])))))) +
 scale_y_continuous(
   limits = c(-max(abs(c(range(fm1$alphas1[, 1:2], fm1$gammas1[, 1:2]))))
               , \max(abs(c(range(fm1\$alphas1[, 1:2], fm1\$gammas1[, 1:2])))))) +
 labs(title = "Bayesian Method", x = expression(PC[1]), y = expression(PC[2])) +
 theme_bw() +
 theme(plot.title = element_text(hjust = 0.5))
print(BiplotBayesAMMI)
```

e_eff

Environment Effects

Description

Calcuates Environment Effects

Usage

```
## Default S3 method:
e_eff(.data, .y, .gen, .env)
```

Arguments

.data	data.frame
. y	Response Variable
.gen	Genotypes Factor
.env	Environment Factor

Value

Environment Effects

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Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
e_eff(
    .data = Maiz
    , .y = y
    , .gen = entry
    , .env = site
    )
```

ge_ammi

AMMI of Genotype by Environment Interaction Model

Description

Performs Additive Main Effects and Multiplication Interaction Analysis of Genotype by Environment Interaction Model

Usage

```
ge_ammi(.data, .y, .gen, .env, .rep)
## Default S3 method:
ge_ammi(.data, .y, .gen, .env, .rep)
```

Arguments

.data	data.frame
. y	Response Variable
.gen	Genotypes Factor
.env	Environment Factor
.rep	Replication Factor

Value

Genotype by Environment Interaction Model

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Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
fm1 <-
   ge_ammi(
        .data = Maiz
   , .y = y
   , .gen = entry
   , .env = site
   , .rep = rep
   )</pre>
```

ge_eff

Genotype by Environment Interaction Effects

Description

Calcuates Genotype by Environment Interaction Effects

Usage

```
## Default S3 method:
ge_eff(.data, .y, .gen, .env)
```

Arguments

.data	data.frame
. У	Response Variable
.gen	Genotypes Factor
.env	Environment Factor

Value

Genotype by Environment Interaction Effects

10 ge_mean

Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
ge_eff(
    .data = Maiz
    , .y = y
    , .gen = entry
    , .env = site
    )
```

ge_mean

Genotype by Environment Interaction Means

Description

Calcuates Genotype by Environment Interaction Means

Usage

```
## Default S3 method:
ge_mean(.data, .y, .gen, .env)
```

Arguments

.data	data.frame
. У	Response Variable
.gen	Genotypes Factor
.env	Environment Factor

Value

Genotype by Environment Interaction Means

Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

ge_model 11

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
ge_mean(
    .data = Maiz
, .y = y
, .gen = entry
, .env = site
)
```

ge_model

Genotype by Environment Interaction Model

Description

Calcuates Genotype by Environment Interaction Model

Usage

```
ge_model(.data, .y, .gen, .env, .rep)
## Default S3 method:
ge_model(.data, .y, .gen, .env, .rep)
```

Arguments

.data	data.frame
. y	Response Variable
.gen	Genotypes Factor
.env	Environment Factor
.rep	Replication Factor

Value

Genotype by Environment Interaction Model

Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

12 ge_var

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
fm1 <-
   ge_model(
        .data = Maiz
        , .y = y
        , .gen = entry
        , .env = site
        , .rep = rep
    )</pre>
```

ge_var

Genotype by Environment Interaction Variances

Description

Calcuates Genotype by Environment Interaction Variances

Usage

```
## Default S3 method:
ge_var(.data, .y, .gen, .env)
```

Arguments

.data	data.frame
. У	Response Variable
.gen	Genotypes Factor
.env	Environment Factor

Value

Genotype by Environment Interaction Variances

Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

g_eff

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
ge_var(
    .data = Maiz
    , .y = y
    , .gen = entry
    , .env = site
    )
```

g_eff

Genotype Effects

Description

Calcuates Genotype Effects

Usage

```
## Default S3 method:
g_eff(.data, .y, .gen, .env)
```

Arguments

.data	data.frame
. y	Response Variable
.gen	Genotypes Factor
.env	Environment Factor

Value

Genotype Effects

Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

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Examples

```
data(Maiz)
g_eff(
    .data = Maiz
    , .y = y
    , .gen = entry
    , .env = site
    )
```

Maiz

Data for Genotypes by Environment Interaction (GEI)

Description

Maiz is used for performing Genotypes by Environment Interaction (GEI) Analysis.

Usage

```
data(Maiz)
```

Format

A data.frame 1320 obs. of 6 variables.

Details

- Gen Genotype
- Institute Institute
- Rep Replicate
- Block Block
- Env Environment
- Yield Yield Response

Author(s)

- Muhammad Yaseen (<myaseen208@gmail.com>)
- 2. Jose Crossa (<j.crossa@cgiar.org>)
- 3. Sergio Perez-Elizalde (<sergiop@colpos.mx>)
- 4. Diego Jarquin (<diego.jarquin@gmail.com>)
- 5. Jose Miguel Cotes
- 6. Kert Viele
- 7. Genzhou Liu
- 8. Paul L. Cornelius

matrix_k

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

Examples

```
data(Maiz)
```

matrix_k

k Matrix

Description

Gives k matrix

Usage

```
matrix_k(n)
## Default S3 method:
matrix_k(n)
```

Arguments

n

Number of columns

Value

Matrix

Author(s)

Muhammad Yaseen (<myaseen208@gmail.com>)

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

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orthnorm

Orthogonal Normalization

Description

Perform Orthogonal Normalization of a matrix

Usage

```
orthnorm(u = NULL, basis = TRUE, norm = TRUE)
## Default S3 method:
orthnorm(u = NULL, basis = TRUE, norm = TRUE)
```

Arguments

u Matrix

basis Logical argument by default TRUE norm Logical argument by default TRUE

Value

Matrix

Author(s)

1. Muhammad Yaseen (<myaseen208@gmail.com>)

References

Crossa, J., Perez-Elizalde, S., Jarquin, D., Cotes, J.M., Viele, K., Liu, G., and Cornelius, P.L. (2011) Bayesian Estimation of the Additive Main Effects and Multiplicative Interaction Model *Crop Science*, **51**, 1458–1469. (doi: 10.2135/cropsci2010.06.0343)

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