# Package 'spFW'

# October 14, 2022

Type Package

Title Hierarchical Spatial Finlay-Wilkinson Model
Version 0.1.0
Author Xingche Guo <xguo@iastate.edu></xguo@iastate.edu>
-
Maintainer Xingche Guo <xguo@iastate.edu></xguo@iastate.edu>
<b>Description</b> Estimation and Prediction Functions Using Bayesian Hierarchical Spatial Finlay-Wilkinson Model for Analysis of Multi-Environment Field Trials.
License GPL-3
Encoding UTF-8
LazyData true
RoxygenNote 6.1.1
<b>Depends</b> R (>= $2.10$ )
LinkingTo Rcpp, RcppArmadillo
Imports Rcpp
NeedsCompilation yes
Repository CRAN
<b>Date/Publication</b> 2022-03-29 18:40:02 UTC
R topics documented:
HFWM_est
HFWM_pred
HSFWM_est
HSFWM_pred
spFW_example_data
Index 9

2 HFWM\_est

Н	F١	W	М	est
- 1 1		71	ı.ı	COL

Estimation Function for Hierarchical Finlay-Wilkinson Model

## Description

This function ignores spatial effects.

## Usage

```
HFWM_est(Y, VAR, ENV, kin_info = FALSE, A = NULL, env_info = FALSE,
Z = NULL, inits = NULL, hyper_para = NULL, M_iter = 5000,
burn_in = 3000, thin = 5, save_chain = FALSE, seed = NULL)
```

## Arguments

Υ	A length-N numerical response vector
VAR	A length-N factor/character vector indicating the genotype information of Y
ENV	A length-N factor/character vector indicating the field information of Y
kin_info	A logical parameter controling if to use kinship matrix
Α	kinship matrix, give value only if kin_info = TRUE
env_info	A logical parameter controling whether to use environmental covariates
Z	environmental covariates matrix with rownames = field names, give value only if env_info = TRUE
inits	initial values, default is given
hyper_para	hyper-parameter values, default is given
M_iter	Total iteration number
burn_in	Burn in number
thin	Thinning value
save_chain	A logical parameter controling whether to save MCMC chain: 'Chains.rds' in current working directory
seed	Random seed value

# Value

Mean estimates and RMSE value

# **Examples**

```
library(spFW)

# load data
data(spFW_example_data)
Y <- spFW_example_data$yield
VAR <- spFW_example_data$geno</pre>
```

HFWM\_pred 3

```
ENV <- spFW_example_data$loc
COOR <- spFW_example_data[,c(4,5)]
# run model
fit0 <- HFWM_est(Y, VAR, ENV, M_iter = 1000, burn_in = 500, thin = 5)
# plot estimated Y
plot(Y, fit0$yhat)</pre>
```

HFWM\_pred

Prediction Function for Hierarchical Finlay-Wilkinson Model

## Description

This function ignores spatial effects.

# Usage

```
HFWM_pred(Y, VAR, ENV, VAR2, ENV2, save_int = FALSE, kin_info = FALSE,
  A = NULL, inits = NULL, hyper_para = NULL, M_iter = 5000,
  burn_in = 3000, thin = 5, seed = NULL)
```

## Arguments

A length-N1 factor/character vector indicating the field information of Y  VAR2 A length-N2 factor/character vector indicating the genotype information of testing set  ENV2 A length-N2 factor/character vector indicating the field information of of testing set  save_int A logical parameter controling whether to save prediction credible intervals kin_info A logical parameter controling if to use kinship matrix  A kinship matrix, give value only if kin_info = TRUE  inits initial values, default is given  hyper_para hyper-parameter values, default is given  M_iter Total iteration number	Υ	A length-N1 numerical response vector from training set
A length-N2 factor/character vector indicating the genotype information of testing set  ENV2 A length-N2 factor/character vector indicating the field information of of testing set  save_int A logical parameter controling whether to save prediction credible intervals kin_info A logical parameter controling if to use kinship matrix  A kinship matrix, give value only if kin_info = TRUE  inits initial values, default is given  hyper_para  hyper-parameter values, default is given  M_iter Total iteration number  Burn in number	VAR	A length-N1 factor/character vector indicating the genotype information of Y
ing set  ENV2 A length-N2 factor/character vector indicating the field information of of testing set  save_int A logical parameter controling whether to save prediction credible intervals kin_info A logical parameter controling if to use kinship matrix  A kinship matrix, give value only if kin_info = TRUE  inits initial values, default is given  hyper_para  hyper-parameter values, default is given  M_iter Total iteration number  burn_in Burn in number	ENV	A length-N1 factor/character vector indicating the field information of Y
set  save_int A logical parameter controling whether to save prediction credible intervals  kin_info A logical parameter controling if to use kinship matrix  A kinship matrix, give value only if kin_info = TRUE  inits initial values, default is given  hyper_para  hyper-parameter values, default is given  M_iter Total iteration number  burn_in Burn in number	VAR2	
kin_info A logical parameter controling if to use kinship matrix  A kinship matrix, give value only if kin_info = TRUE  inits initial values, default is given  hyper_para  hyper-parameter values, default is given  M_iter Total iteration number  burn_in Burn in number	ENV2	
A kinship matrix, give value only if kin_info = TRUE  inits initial values, default is given  hyper_para hyper-parameter values, default is given  M_iter Total iteration number  burn_in Burn in number	save_int	A logical parameter controling whether to save prediction credible intervals
inits initial values, default is given  hyper_para hyper-parameter values, default is given  M_iter Total iteration number  burn_in Burn in number	kin_info	A logical parameter controling if to use kinship matrix
hyper_para hyper-parameter values, default is given  M_iter Total iteration number  burn_in Burn in number	Α	kinship matrix, give value only if kin_info = TRUE
M_iter Total iteration number burn_in Burn in number	inits	initial values, default is given
burn_in Burn in number	hyper_para	hyper-parameter values, default is given
	M_iter	Total iteration number
thin Thinning value	burn_in	Burn in number
	thin	Thinning value
seed Random seed value	seed	Random seed value

### Value

Mean prediction values and/or prediction intervals

4 HSFWM\_est

### **Examples**

```
library(spFW)
# load and split data
data(spFW_example_data)
idx_pred <- sample(125, 25)</pre>
Y0 <- spFW_example_data$yield
VAR0 <- spFW_example_data$geno
ENV0 <- spFW_example_data$loc</pre>
Y1 <- Y0[-idx_pred]
Y2 <- Y0[idx_pred]
VAR1 <- VAR0[-idx_pred]</pre>
VAR2 <- VAR0[idx_pred]</pre>
ENV1 <- ENV0[-idx_pred]</pre>
ENV2 <- ENV0[idx_pred]</pre>
order_y <- order(Y2)</pre>
# run model
pred0 <- HFWM_pred(Y1, VAR1, ENV1, VAR2, ENV2, save_int = TRUE,</pre>
                    M_{iter} = 1000, burn_{in} = 500, thin = 5)
# visualize prediction results
plot(1:25, pred0$PY[order_y], ylim = c(50, 250), pch = 15, col = "red",
     xlab = "Plant ID for Prediction", ylab = "Yield",
    main = "95% Prediction Intervals with Predicted Mean (Red) Versus True Yield (Blue)")
points(1:25, Y2[order_y], col = "blue")
for (i in 1:25){
  lines(x = c(i,i), y = c(pred0$PY_CI[,order_y][1,i], pred0$PY_CI[,order_y][4,i]))
```

HSFWM\_est

Estimation Function for Hierarchical Spatial Finlay-Wilkinson Model

### **Description**

This function considers spatial adjustments.

### Usage

```
HSFWM_est(Y, VAR, ENV, COOR, kin_info = FALSE, A = NULL,
  env_info = FALSE, Z = NULL, inits = NULL, hyper_para = NULL,
  M_iter = 5000, burn_in = 3000, thin = 5, save_chain = FALSE,
  seed = NULL)
```

HSFWM\_est 5

# Arguments Y

VAR	A length-N factor/character vector indicating the genotype information of Y
ENV	A length-N factor/character vector indicating the field information of Y
COOR	A N by 2 numerical matrix indicating the spatial locations of Y

kin\_info A logical parameter controling if to use kinship matrix

A kinship matrix, give value only if kin\_info = TRUE

A length-N numerical response vector

env\_info A logical parameter controling whether to use environmental covariates

Z environmental covariates matrix with rownames = field names, give value only

if env\_info = TRUE

inits initial values, default is given

hyper\_para hyper-parameter values, default is given

M\_iter Total iteration number

burn\_in Burn in number thin Thinning value

save\_chain A logical parameter controlling whether to save MCMC chain: 'Chains.rds' in

current working directory

seed Random seed value

### Value

Mean estimates and RMSE value

# **Examples**

6 HSFWM\_pred

HSFWM_pred	Prediction Function for Hierarchical Spatial Finlay-Wilkinson Model

## Description

This function considers spatial adjustments.

## Usage

```
HSFWM_pred(Y, VAR, ENV, COOR, VAR2, ENV2, COOR2, save_int = FALSE,
  kin_info = FALSE, A = NULL, inits = NULL, hyper_para = NULL,
  M_iter = 5000, burn_in = 3000, thin = 5, seed = NULL)
```

# Arguments

Υ	A length-N1 numerical response vector from training set
VAR	A length-N1 factor/character vector indicating the genotype information of Y
ENV	A length-N1 factor/character vector indicating the field information of Y
COOR	A N1 by 2 numerical matrix indicating the spatial locations of Y
VAR2	A length-N2 factor/character vector indicating the genotype information of testing set
ENV2	A length-N2 factor/character vector indicating the field information of of testing set
COOR2	A N2 by 2 numerical matrix indicating the spatial locations of testing set
save_int	A logical parameter controling whether to save prediction credible intervals
kin_info	A logical parameter controling if to use kinship matrix
Α	kinship matrix, give value only if kin_info = TRUE
inits	initial values, default is given
hyper_para	hyper-parameter values, default is given
M_iter	Total iteration number
burn_in	Burn in number
thin	Thinning value
seed	Random seed value

### Value

Mean prediction values and/or prediction intervals

spFW\_example\_data 7

### **Examples**

```
library(spFW)
# load and split data
data(spFW_example_data)
idx_pred <- sample(125, 25)</pre>
Y0 <- spFW_example_data$yield
VAR0 <- spFW_example_data$geno
ENV0 <- spFW_example_data$loc</pre>
COOR0 <- spFW_example_data[,c(4,5)]</pre>
Y1 <- Y0[-idx_pred]
Y2 <- Y0[idx_pred]
VAR1 <- VAR0[-idx_pred]</pre>
VAR2 <- VAR0[idx_pred]</pre>
ENV1 <- ENV0[-idx_pred]</pre>
ENV2 <- ENV0[idx_pred]</pre>
COOR1 <- COOR0[-idx_pred,]</pre>
COOR2 <- COOR0[idx_pred,]</pre>
order_y <- order(Y2)</pre>
# run model
pred1 <- HSFWM_pred(Y1, VAR1, ENV1, COOR1, VAR2, ENV2, COOR2, save_int = TRUE,</pre>
                     M_iter = 1000, burn_in = 500, thin = 5)
# visualize prediction results
plot(1:25, pred1$PY[order_y], ylim = c(50, 250), pch = 15, col = "red",
     xlab = "Plant ID for Prediction", ylab = "Yield",
    main = "95% Prediction Intervals with Predicted Mean (Red) Versus True Yield (Blue)")
points(1:25, Y2[order_y], col = "blue")
for (i in 1:25){
  lines(x = c(i,i), y = c(pred1$PY_CI[,order_y][1,i], pred1$PY_CI[,order_y][4,i]))
}
```

spFW\_example\_data

Example Data for Analysis

### **Description**

A data frame containing the plant yield, genotypes, environments, and within-field positions.

#### Usage

```
spFW_example_data
```

## **Format**

A data frame with 5 elements, which are:

yield Plant yield

geno Plant genotype ID

loc Plant environment ID

range Plant x-coordinate at a given environment

pass Plant y-coordinate at a given environment

# **Index**