# Package 'jlmerclusterperm'

March 4, 2024

**Title** Cluster-Based Permutation Analysis for Densely Sampled Time Data **Version** 1.1.3

Description An implementation of fast cluster-based permutation analysis (CPA) for densely-sampled time data developed in Maris & Oostenveld, 2007 <doi:10.1016/j.jneumeth.2007.03.024>. Supports (generalized, mixed-effects) regression models for the calculation of timewise statistics. Provides both a wholesale and a piecemeal interface to the CPA procedure with an emphasis on interpretability and diagnostics. Integrates 'Julia' libraries 'MixedModels.jl' and 'GLM.jl' for performance improvements, with additional functionalities for interfacing with 'Julia' from 'R' powered by the 'JuliaConnectoR' package.

License MIT + file LICENSE

URL https://github.com/yjunechoe/jlmerclusterperm,
https://yjunechoe.github.io/jlmerclusterperm/

BugReports https://github.com/yjunechoe/jlmerclusterperm/issues

**Depends** R (>= 3.5)

**Imports** backports (>= 1.1.7), cli, generics, JuliaConnectoR, lme4, stats, tools, utils

**Suggests** broom, broom.mixed, covr, dplyr, eyetrackingR, forcats, future, ggplot2, knitr, MASS, patchwork, readr, rmarkdown, scales, testthat (>= 3.0.0), tibble

VignetteBuilder knitr

Config/testthat/edition 3

**Encoding UTF-8** 

RoxygenNote 7.2.3

**SystemRequirements** Julia (>= 1.8)

Collate 'jlmerclusterperm-package.R' 'aaa.R' 'utils.R' 'interop-utils.R' 'interop-utils-unexported.R' 'julia\_rng.R' 'jlmer\_spec.R' 'jlmer.R' 'compute\_timewise\_statistics.R' 'permute.R' 'permute\_timewise\_statistics.R' 'clusters\_methods.R' 'extract\_clusters.R' 'calculate\_pvalue.R' 'clusterpermute.R' 'threshold\_search.R' 'tidy.R' 'zzz.R' 'srr-stats-standards.R'

### NeedsCompilation no

**Author** June Choe [aut, cre, cph] (<a href="https://orcid.org/0000-0002-0701-921X">https://orcid.org/0000-0002-0701-921X</a>)

Maintainer June Choe <jchoe001@gmail.com>

**Repository** CRAN

**Date/Publication** 2024-03-04 19:40:02 UTC

# **R** topics documented:

Index		25
	walk_threshold_steps	23
	to_jlmer	
	permute_timewise_statistics	20
	permute_by_predictor	18
	make_jlmer_spec	17
	julia_setup_ok	17
	julia_rng	16
	julia_progress	15
	julia_model_tidiers	14
	jlmerclusterperm_setup	13
	jlmer	12
	extract_null_cluster_dists	10
	extract_empirical_clusters	9
	compute_timewise_statistics	7
	cluster_permutation_tidiers	6
	clusterpermute	4
	calculate_clusters_pvalues	2

calculate\_clusters\_pvalues

Calculate bootstrapped p-values of cluster-mass statistics

### Description

Calculate bootstrapped p-values of cluster-mass statistics

#### Usage

```
calculate_clusters_pvalues(
  empirical_clusters,
  null_cluster_dists,
  add1 = FALSE
)
clusters_are_comparable(empirical_clusters, null_cluster_dists, error = FALSE)
```

#### **Arguments**

```
empirical_clusters

A empirical_clusters object

null_cluster_dists

A null_cluster_dists object

add1 Whether to add 1 to the numerator and denominator when calculating the p-
value. Use TRUE to effectively count the observed statistic as part of the per-
muted null distribution (recommended with larger nsim prior to publishing re-
sults).

error Whether to throw an error if incompatible
```

#### Value

An empirical\_clusters object augmented with p-values.

#### See Also

```
extract_empirical_clusters(), extract_null_cluster_dists()
```

```
library(dplyr, warn.conflicts = FALSE)

# Specification object
spec <- make_jlmer_spec(
    weight ~ 1 + Diet, filter(ChickWeight, Time <= 20),
    subject = "Chick", time = "Time"
)
spec

# Make empirical clusters
empirical_statistics <- compute_timewise_statistics(spec)
empirical_clusters <- extract_empirical_clusters(empirical_statistics, threshold = 2)
empirical_clusters</pre>
# Make null cluster-mass distribution
```

4 clusterpermute

```
reset_rng_state()
null_statistics <- permute_timewise_statistics(spec, nsim = 100)</pre>
null_cluster_dists <- extract_null_cluster_dists(null_statistics, threshold = 2)</pre>
# Significance test the empirical cluster(s) from each predictor against the simulated null
calculate_clusters_pvalues(empirical_clusters, null_cluster_dists)
# Set `add1 = TRUE` to normalize by adding 1 to numerator and denominator
calculate_clusters_pvalues(empirical_clusters, null_cluster_dists, add1 = TRUE)
# This sequence of procedures is equivalent to `clusterpermute()`
reset_rng_state()
clusterpermute(spec, threshold = 2, nsim = 100, progress = FALSE)
# The empirical clusters and the null cluster-mass distribution must be comparable
empirical_clusters2 <- extract_empirical_clusters(empirical_statistics, threshold = 3)</pre>
# For example, below code errors because thresholds are different (2 vs. 3)
try( calculate_clusters_pvalues(empirical_clusters2, null_cluster_dists) )
# Check for compatibility with `clusters_are_comparable()`
clusters_are_comparable(empirical_clusters, null_cluster_dists)
clusters_are_comparable(empirical_clusters2, null_cluster_dists)
```

clusterpermute

Conduct a cluster-based permutation test

#### **Description**

Conduct a cluster-based permutation test

### Usage

```
clusterpermute(
   jlmer_spec,
   family = c("gaussian", "binomial"),
   statistic = c("t", "chisq"),
   threshold,
   nsim = 100L,
   predictors = NULL,
   binned = FALSE,
   top_n = Inf,
   add1 = TRUE,
   ...,
   progress = TRUE
)
```

clusterpermute 5

### Arguments

jlmer_spec	Data prepped for jlmer from make_jlmer_spec()
family	A GLM family. Currently supports "gaussian" and "binomial".
statistic	Test statistic for calculating cluster mass. Can be one of "t" (default) from the regression model output or "chisq" from a likelihood ratio test (takes about twice as long to calculate).
threshold	The threshold value that the statistic must pass to contribute to cluster mass. Interpretation differs on the choice of statistic (more below):
	• If statistic = "t", the threshold for t-value (beta/std.err) from the regression model.
	• If statistic = "chisq", the threshold for the p-value of chi-squared statistics from likelihood ratio tests.
nsim	Number of simulations description
predictors	(Optional) a subset of predictors to test. Defaults to NULL which tests all predictors.
binned	Whether the data has been aggregated/collapsed into time bins. Defaults to FALSE, which requires a cluster to span at least two time points. If TRUE, allows length-1 clusters to exist.
top_n	How many clusters to return, in the order of the size of the cluster-mass statistic. Defaults to Inf which return all detected clusters.
add1	Whether to add 1 to the numerator and denominator when calculating the p-value. Use TRUE to effectively count the observed statistic as part of the permuted null distribution (recommended with larger nsim prior to publishing results).
•••	Optional arguments passed to Julia for model fitting. Defaults to fast = TRUE (when family = "binomial") and progress = FALSE.
progress	Defaults to TRUE, which prints progress on each step of the cluster permutation test.

#### Value

A list of null\_cluster\_dists and empirical\_clusters with p-values

### See Also

```
compute\_timewise\_statistics(), permute\_timewise\_statistics(), extract\_empirical\_clusters(), extract\_null\_cluster\_dists(), calculate\_clusters\_pvalues()
```

```
library(dplyr, warn.conflicts = FALSE)
```

```
# Specification object
spec <- make_jlmer_spec(
    weight ~ 1 + Diet, filter(ChickWeight, Time <= 20),
    subject = "Chick", time = "Time"
)
spec

# Should minimally provide `threshold` and `nsim`, in addition to the spec object
reset_rng_state()
CPA <- clusterpermute(spec, threshold = 2, nsim = 100, progress = FALSE)
CPA

# CPA is a list of `<null_cluster_dists>` and `<empirical_clusters>` objects
sapply(CPA, class)

# You can extract the individual components for further inspection
CPA$null_cluster_dists
CPA$empirical_clusters
```

cluster\_permutation\_tidiers

Tidiers for cluster permutation test objects

### **Description**

Tidiers for cluster permutation test objects

#### Usage

```
## S3 method for class 'timewise_statistics'
tidy(x, ...)

## S3 method for class 'empirical_clusters'
tidy(x, ...)

## S3 method for class 'null_cluster_dists'
tidy(x, ...)
```

#### **Arguments**

x An object of class <timewise\_statistics>, <empirical\_clusters>, or <null\_cluster\_dists>
... Unused

#### Value

A data frame

#### **Examples**

```
library(dplyr, warn.conflicts = FALSE)
# Specification object
spec <- make_jlmer_spec(</pre>
  weight \sim 1 + Diet, filter(ChickWeight, Time <= 20),
  subject = "Chick", time = "Time"
spec
# Method for `<timewise_statistics>`
empirical_statistics <- compute_timewise_statistics(spec)</pre>
class(empirical_statistics)
tidy(empirical_statistics)
reset_rng_state()
null_statistics <- permute_timewise_statistics(spec, nsim = 100)</pre>
class(null_statistics)
tidy(null_statistics)
# Method for `<empirical_clusters>`
empirical_clusters <- extract_empirical_clusters(empirical_statistics, threshold = 2)</pre>
class(empirical_clusters)
tidy(empirical_clusters)
# Method for `<null_cluster_dists>`
null_cluster_dists <- extract_null_cluster_dists(null_statistics, threshold = 2)</pre>
class(null_cluster_dists)
tidy(null_cluster_dists)
```

compute\_timewise\_statistics

Fit Julia regression models to each time point of a time series data

### **Description**

Fit Julia regression models to each time point of a time series data

#### Usage

```
compute_timewise_statistics(
  jlmer_spec,
  family = c("gaussian", "binomial"),
 statistic = c("t", "chisq"),
)
```

#### **Arguments**

jlmer\_spec Data prepped for ilmer from make\_jlmer\_spec() A GLM family. Currently supports "gaussian" and "binomial". family Test statistic for calculating cluster mass. Can be one of "t" (default) from the statistic regression model output or "chisq" from a likelihood ratio test (takes about twice as long to calculate). Optional arguments passed to Julia for model fitting. Defaults to fast = TRUE (when family = "binomial") and progress = FALSE.

#### Value

A predictor-by-time matrix of cluster statistics, of class timewise\_statistics.

#### See Also

```
jlmer(), make_jlmer_spec()
```

```
library(dplyr, warn.conflicts = FALSE)
# Specification object
spec <- make_jlmer_spec(</pre>
  weight ~ 1 + Diet, filter(ChickWeight, Time <= 20),</pre>
  subject = "Chick", time = "Time"
)
spec
# Predictor x Time matrix of t-statistics from regression output
empirical_statistics <- compute_timewise_statistics(spec)</pre>
round(empirical_statistics, 2)
# Collect as dataframe with `tidy()`
empirical_statistics_df <- tidy(empirical_statistics)</pre>
empirical_statistics_df
# Timewise statistics are from regression models fitted to each time point
```

```
# - Note the identical statistics at `Time == 0`
empirical_statistics_df %>%
  filter(time == 0)
to_jlmer(weight ~ 1 + Diet, filter(ChickWeight, Time == 0)) %>%
  tidy() %>%
  select(term, statistic)
```

extract\_empirical\_clusters

Detect largest clusters from a time sequence of predictor statistics

#### **Description**

Detect largest clusters from a time sequence of predictor statistics

#### Usage

```
extract_empirical_clusters(
  empirical_statistics,
  threshold,
  binned = FALSE,
  top_n = Inf
)
```

### Arguments

empirical\_statistics

A predictor-by-time matrix of empirical timewise statistics.

threshold

The threshold value that the statistic must pass to contribute to cluster mass. Interpretation differs on the choice of statistic (more below):

- If statistic = "t", the threshold for t-value (beta/std.err) from the regression model.
- If statistic = "chisq", the threshold for the p-value of chi-squared statistics from likelihood ratio tests.

binned

Whether the data has been aggregated/collapsed into time bins. Defaults to FALSE, which requires a cluster to span at least two time points. If TRUE, allows length-1 clusters to exist.

top\_n

How many clusters to return, in the order of the size of the cluster-mass statistic. Defaults to Inf which return all detected clusters.

### Value

An empirical\_clusters object.

#### See Also

```
compute_timewise_statistics()
```

#### **Examples**

```
library(dplyr, warn.conflicts = FALSE)
# Specification object
spec <- make_jlmer_spec(</pre>
  weight ~ 1 + Diet, filter(ChickWeight, Time <= 20),</pre>
  subject = "Chick", time = "Time"
)
spec
# Empirical clusters are derived from the timewise statistics
empirical_statistics <- compute_timewise_statistics(spec)</pre>
empirical_clusters <- extract_empirical_clusters(empirical_statistics, threshold = 2)</pre>
empirical_clusters
# Collect as dataframe with `tidy()`
empirical_clusters_df <- tidy(empirical_clusters)</pre>
empirical_clusters_df
# Changing the `threshold` value identifies different clusters
extract_empirical_clusters(empirical_statistics, threshold = 1)
# A predictor can have zero or multiple clusters associated with it
extract_empirical_clusters(empirical_statistics, threshold = 3)
```

 ${\tt extract\_null\_cluster\_dists}$ 

Construct a null distribution of cluster-mass statistics

#### **Description**

Construct a null distribution of cluster-mass statistics

### Usage

```
extract_null_cluster_dists(null_statistics, threshold, binned = FALSE)
```

#### **Arguments**

null\_statistics

A simulation-by-time-by-predictor 3D array of null (permuted) timewise statistics.

threshold

The threshold value that the statistic must pass to contribute to cluster mass. Interpretation differs on the choice of statistic (more below):

- If statistic = "t", the threshold for t-value (beta/std.err) from the regression model.
- If statistic = "chisq", the threshold for the p-value of chi-squared statistics from likelihood ratio tests.

binned

Whether the data has been aggregated/collapsed into time bins. Defaults to FALSE, which requires a cluster to span at least two time points. If TRUE, allows length-1 clusters to exist.

#### Value

A null\_cluster\_dists object.

#### See Also

```
permute_timewise_statistics()
```

```
library(dplyr, warn.conflicts = FALSE)
# Specification object
spec <- make_jlmer_spec(</pre>
  weight ~ 1 + Diet, filter(ChickWeight, Time <= 20),</pre>
  subject = "Chick", time = "Time"
)
spec
# Null cluster-mass distributions are derived from the permuted timewise statistics
reset_rng_state()
null_statistics <- permute_timewise_statistics(spec, nsim = 100)</pre>
null_cluster_dists <- extract_null_cluster_dists(null_statistics, threshold = 2)</pre>
null_cluster_dists
# Collect as dataframe with `tidy()`
# - Each simulation contributes one (largest) cluster-mass statistic to the null
# - When no clusters are found, the `sum_statistic` value is zero
null_cluster_dists_df <- tidy(null_cluster_dists)</pre>
null_cluster_dists_df
# Changing the `threshold` value changes the shape of the null
```

jlmer

```
extract_null_cluster_dists(null_statistics, threshold = 1)
extract_null_cluster_dists(null_statistics, threshold = 3)
```

jlmer

Fit a Julia regression model using ilmer specifications

#### **Description**

Fit a Julia regression model using jlmer specifications

#### Usage

```
jlmer(jlmer_spec, family = c("gaussian", "binomial"), ..., progress = FALSE)
```

### Arguments

family A GLM family. Currently supports "gaussian" and "binomial".

... Optional arguments passed to Julia for model fitting.

progress If TRUE, prints the timing of iterations.

#### Value

A jlmer\_mod object.

#### See Also

```
make_jlmer_spec()
```

```
# Fitting a regression model with a specification object
spec <- make_jlmer_spec(weight ~ 1 + Diet, ChickWeight)
jlmer(spec)

# `lm()` equivalent
summary(lm(weight ~ 1 + Diet, ChickWeight))$coef</pre>
```

jlmerclusterperm\_setup 13

```
jlmerclusterperm_setup
```

Initial setup for the ilmerclusterperm package

### **Description**

Initial setup for the jlmerclusterperm package

### Usage

```
jlmerclusterperm_setup(..., cache_dir = NULL, restart = TRUE, verbose = TRUE)
```

### Arguments

	Ignored

cache\_dir The location to write out package cache files (namely, Manifest.toml). If NULL

(default), attempts to write to the package's cache directory discovered via R\_user\_dir()

and falls back to tempdir().

restart Whether to set up a fresh Julia session, given that one is already running. If

FALSE and jlmerclusterperm\_setup() has already been called, nothing hap-

pens.

verbose Whether to print progress and messages from Julia in the console

### Value

TRUE

```
options("jlmerclusterperm.nthreads" = 2)
jlmerclusterperm_setup(cache_dir = tempdir(), verbose = FALSE)
```

14 julia\_model\_tidiers

```
julia_model_tidiers Tidier methods for Julia regression models
```

#### **Description**

Tidier methods for Julia regression models

### Usage

```
## S3 method for class 'jlmer_mod'
tidy(x, effects = c("var_model", "ran_pars", "fixed"), ...)
## S3 method for class 'jlmer_mod'
glance(x, ...)
```

#### **Arguments**

```
x An object of class jlmer_mod
effects One of "var_model", "ran_pars", or "fixed"
... Unused
```

#### Value

A data frame

```
# Fixed-effects only model
mod1 <- to_jlmer(weight ~ 1 + Diet, ChickWeight)
tidy(mod1)
glance(mod1)

# Mixed model
mod2 <- to_jlmer(weight ~ 1 + Diet + (1 | Chick), ChickWeight)
tidy(mod2)
glance(mod2)

# Select which of fixed/random effects to return
tidy(mod2, effects = "fixed")
tidy(mod2, effects = "ran_pars")</pre>
```

julia\_progress 15

### Description

Set/get options for Julia progress bar

#### Usage

```
julia_progress(show, width)
```

#### **Arguments**

show Whether to show the progress bar. You may also pass in a list of "show" and

"width".

width Width of the progress bar. If "auto", adjusts the progress bar width to fit the

console.

#### Value

Previous values for show and width

```
# Show current progress options
julia_progress()

# Set options and save previous options
old_progress_opts <- julia_progress(show = FALSE, width = 100)
julia_progress()

# Restoring progress settings by passing a list of old options
old_progress_opts
julia_progress(old_progress_opts)
identical(julia_progress(), old_progress_opts)

# Alternatively, reset to default settings using this syntax:
julia_progress(show = TRUE, width = "auto")</pre>
```

julia\_rng

julia\_rng

Interface to the Julia RNG

### Description

Interface to the Julia RNG

### Usage

```
set_rng_state(i)
reset_rng_state()
get_rng_state()
set_rng_seed(seed)
get_rng_seed()
```

### Arguments

i Counter number

seed Seed

#### Value

The current seed or counter

```
# RNG initializes to seed=1 counter=0
get_rng_seed()
get_rng_state()

# setter/getter for RNG counter
set_rng_state(123)
get_rng_state()

# setter/getter for RNG seed
set_rng_seed(2)
get_rng_seed(2)
get_rng_seed()

# restore to initial setting (seed=1, counter=0)
set_rng_seed(1)
set_rng_state(0)
```

julia\_setup\_ok 17

julia\_setup\_ok

Check Julia requirements for jlmerclusterperm

### Description

Check Julia requirements for jlmerclusterperm

### Usage

```
julia_setup_ok()
```

#### Value

Boolean

#### **Examples**

```
julia_setup_ok()
```

make\_jlmer\_spec

Create a specifications object for fitting regression models in Julia

### Description

Create a specifications object for fitting regression models in Julia

## Usage

```
make_jlmer_spec(
  formula,
  data,
  subject = NULL,
  trial = NULL,
  time = NULL,
  drop_terms = NULL,
  ...
)
```

#### **Arguments**

formula	Model formula in R syntax
data	A data frame
subject	Column for subjects in the data.
trial	Column for trials in the data. Must uniquely identify a time series within subject (for example, the column for items in a counterbalanced design where each subject sees exactly one item).
time	Column for time in the data.
drop_terms	(Optional) any terms to drop from the reconstructed model formula
	Unused, for extensibility.

#### Value

An object of class jlmer\_spec.

### Examples

```
# Bare specification object (minimal spec for fitting a global model)
spec <- make_jlmer_spec(weight ~ 1 + Diet, ChickWeight)
spec

# Constraints on specification for CPA:
# 1) The combination of `subject`, `trial`, and `time` must uniquely identify rows in the data
# 2) `time` must have constant sampling rate (i.e., evenly spaced values)
spec_wrong <- make_jlmer_spec(
    weight ~ 1 + Diet, ChickWeight,
    time = "Time"
)
unique(ChickWeight$Time)

# Corrected specification for the above
spec_correct <- make_jlmer_spec(
    weight ~ 1 + Diet, subset(ChickWeight, Time <= 20),
    subject = "Chick", time = "Time"
)
spec_correct</pre>
```

permute\_by\_predictor Permute data while respecting grouping structure(s) of observations

### **Description**

Permute data while respecting grouping structure(s) of observations

permute\_by\_predictor 19

### Usage

```
permute_by_predictor(
    jlmer_spec,
    predictors,
    predictor_type = c("guess", "between_participant", "within_participant"),
    n = 1L
)
```

#### **Arguments**

#### Value

A long dataframe of permuted re-samples with .id column representing replication IDs.

```
# Example data setup
chickweights_df <- ChickWeight</pre>
chickweights_df <- chickweights_df[chickweights_df$Time <= 20, ]</pre>
chickweights_df$DietInt <- as.integer(chickweights_df$Diet)</pre>
head(chickweights_df)
# Example 1: Spec object using the continuous `DietInt` predictor
chickweights_spec1 <- make_jlmer_spec(</pre>
 formula = weight ~ 1 + DietInt,
 data = chickweights_df,
 subject = "Chick", time = "Time"
)
chickweights_spec1
# Shuffling `DietInt` values guesses `predictor_type = "between_participant"`
reset_rng_state()
spec1_perm1 <- permute_by_predictor(chickweights_spec1, predictors = "DietInt")</pre>
# This calls the same shuffling algorithm for CPA in Julia, so counter is incremented
get_rng_state()
# Shuffling under shared RNG state reproduces the same permutation of the data
reset_rng_state()
spec1_perm2 <- permute_by_predictor(chickweights_spec1, predictors = "DietInt")</pre>
```

```
identical(spec1_perm1, spec1_perm2)

# Example 2: Spec object using the multilevel `Diet` predictor
chickweights_spec2 <- make_jlmer_spec(
    formula = weight ~ 1 + Diet,
    data = chickweights_df,
    subject = "Chick", time = "Time"
)
chickweights_spec2

# Levels of a category are automatically shuffled together
reset_rng_state()
spec2_perm1 <- permute_by_predictor(chickweights_spec2, predictors = "Diet2")
reset_rng_state()
spec2_perm2 <- permute_by_predictor(chickweights_spec2, predictors = c("Diet2", "Diet3", "Diet4"))
identical(spec2_perm1, spec2_perm2)</pre>
```

permute\_timewise\_statistics

Simulate cluster-mass statistics via bootstrapped permutations

### Description

Simulate cluster-mass statistics via bootstrapped permutations

#### Usage

```
permute_timewise_statistics(
    jlmer_spec,
    family = c("gaussian", "binomial"),
    statistic = c("t", "chisq"),
    nsim = 100L,
    predictors = NULL,
    ...
)
```

#### **Arguments**

```
    (Optional) a subset of predictors to test. Defaults to NULL which tests all predictors.
    Optional arguments passed to Julia for model fitting. Defaults to fast = TRUE (when family = "binomial") and progress = FALSE.
```

#### Value

A simulation-by-time-by-predictor 3D array of cluster statistics, of class timewise\_statistics.

#### See Also

```
make_jlmer_spec()
```

```
library(dplyr, warn.conflicts = FALSE)
# Specification object
spec <- make_jlmer_spec(</pre>
 weight ~ 1 + Diet, filter(ChickWeight, Time <= 20),</pre>
  subject = "Chick", time = "Time"
)
spec
# Simulation x Time x Predictor array of t-statistics from regression output
reset_rng_state()
null_statistics <- permute_timewise_statistics(spec, nsim = 3)</pre>
round(null_statistics, 2)
# Collect as dataframe with `tidy()`
permuted_timewise_stats_df <- tidy(null_statistics)</pre>
permuted_timewise_stats_df
# Permutations ran under the same RNG state are identical
reset_rng_state()
null_statistics2 <- permute_timewise_statistics(spec, nsim = 3)</pre>
identical(null_statistics, null_statistics2)
get_rng_state()
null_statistics3 <- permute_timewise_statistics(spec, nsim = 3)</pre>
identical(null_statistics, null_statistics3)
```

to\_jlmer

to\_jlmer

Fit a Julia regression model using lme4 syntax

### **Description**

Fit a Julia regression model using lme4 syntax

#### Usage

```
to_jlmer(
  formula,
  data,
  family = c("gaussian", "binomial"),
  jlmer_spec_opts = list(),
    ...,
  progress = FALSE
)
```

### **Arguments**

```
formula Model formula in R syntax

data A data frame

family A GLM family. Currently supports "gaussian" and "binomial".

jlmer_spec_opts

List of options passed to make_jlmer_spec()

Optional arguments passed to Julia for model fitting.

progress If TRUE, prints the timing of iterations.
```

## Value

```
A jlmer_mod object.
```

#### See Also

```
jlmer(), make_jlmer_spec()
```

```
# Fitting a regression model with R formula syntax
to_jlmer(weight ~ 1 + Diet, ChickWeight)

# `lm()` equivalent
summary(lm(weight ~ 1 + Diet, ChickWeight))$coef
```

walk\_threshold\_steps 23

```
# Fitting a mixed model with {lme4} syntax
to_jlmer(weight ~ 1 + Diet + (1 | Chick), ChickWeight)

# Passing MixedModels.jl fit options to the `...`
to_jlmer(weight ~ 1 + Diet + (1 | Chick), ChickWeight, REML = TRUE)
```

walk\_threshold\_steps

Test the probability of cluster-mass statistics over a range of threshold values

#### **Description**

Test the probability of cluster-mass statistics over a range of threshold values

#### Usage

```
walk_threshold_steps(
  empirical_statistics,
  null_statistics,
  steps,
  top_n = Inf,
  binned = FALSE,
  add1 = TRUE,
  progress = TRUE
)
```

#### Arguments

empirical\_statistics

A predictor-by-time matrix of empirical timewise statistics.

null\_statistics

A simulation-by-time-by-predictor 3D array of null (permuted) timewise statis-

tics.

steps A vector of threshold values to test

top\_n How many clusters to return, in the order of the size of the cluster-mass statistic.

Defaults to Inf which return all detected clusters.

binned Whether the data has been aggregated/collapsed into time bins. Defaults to

FALSE, which requires a cluster to span at least two time points. If TRUE, al-

lows length-1 clusters to exist.

add1 Whether to add 1 to the numerator and denominator when calculating the p-

value. Use TRUE to effectively count the observed statistic as part of the permuted null distribution (recommended with larger nsim prior to publishing re-

sults).

progress Whether to display a progress bar

#### Value

A data frame of predictor clusters-mass statistics by threshold.

# **Index**

```
calculate_clusters_pvalues, 2
                                               tidy.jlmer_mod(julia_model_tidiers), 14
calculate_clusters_pvalues(), 5
                                                tidy.null_cluster_dists
\verb|cluster_permutation_tidiers|, 6
                                                        (cluster_permutation_tidiers),
clusterpermute, 4
clusters_are_comparable
                                                tidy.timewise_statistics
        (calculate_clusters_pvalues), 2
                                                        (cluster_permutation_tidiers),
compute_timewise_statistics, 7
compute_timewise_statistics(), 5, 10
                                                to_jlmer, 22
extract_empirical_clusters, 9
                                               walk_threshold_steps, 23
extract_empirical_clusters(), 3, 5
extract_null_cluster_dists, 10
extract_null_cluster_dists(), 3, 5
get_rng_seed(julia_rng), 16
get_rng_state(julia_rng), 16
glance.jlmer_mod(julia_model_tidiers),
jlmer, 12
jlmer(), 8, 22
jlmerclusterperm_setup, 13
julia_model_tidiers, 14
julia_progress, 15
julia_rng, 16
julia_setup_ok, 17
make_jlmer_spec, 17
make_jlmer_spec(), 8, 12, 21, 22
permute_by_predictor, 18
permute_timewise_statistics, 20
permute_timewise_statistics(), 5, 11
reset_rng_state (julia_rng), 16
set_rng_seed(julia_rng), 16
set_rng_state(julia_rng), 16
tidy.empirical_clusters
        (cluster_permutation_tidiers),
        6
```