

# Package ‘pencal’

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**Title** Penalized Regression Calibration (PRC)

**Version** 0.3.2

**Description** Computes penalized regression calibration (PRC), a statistical method that allows to predict survival from high-dimensional longitudinal predictors. PRC is described in Signorelli et al. (in review, arXiv preprint: <arXiv:2101.04426>).

**License** GPL-3

**URL** <https://mirkosignorelli.github.io/r>

**Depends** R (>= 4.0.0)

**VignetteBuilder** knitr

**Encoding** UTF-8

**LazyData** true

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**Suggests** ptmixed, survminer, knitr, rmarkdown

**NeedsCompilation** no

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fitted_prclmm	<i>A fitted PRC LMM</i>
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### Description

This list contains a fitted PRC LMM, where the CBOCP is computed using 50 cluster bootstrap samples. It is used to reduce the computing time in the example of the function `performance_prc`

### Usage

```
data(fitted_prclmm)
```

### Format

A list comprising step 2 and step 3 as obtained during the estimation of the PRC LMM

### Author(s)

Mirko Signorelli

### References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

### See Also

[performance\\_prc](#)

### Examples

```
data(fitted_prclmm)
ls(fitted_prclmm)
```

---

fit\_lmms *Step 1 of PRC-LMM (estimation of the linear mixed models)*

---

### Description

This function performs the first step for the estimation of the PRC-LMM model proposed in Signorelli et al. (2021, in review)

### Usage

```
fit_lmms(y.names, fixefs, ranefs, long.data, surv.data, t.from.base,
         n.boots = 0, n.cores = 1, verbose = TRUE)
```

### Arguments

y.names	character vector with the names of the response variables which the LMMs have to be fitted to
fixefs	fixed effects formula for the model, example: <code>~ time</code>
ranefs	random effects formula for the model, specified using the representation of random effect structures of the R package <code>nlme</code>
long.data	a data frame with the longitudinal predictors, comprehensive of a variable called <code>id</code> with the subject ids
surv.data	a data frame with the survival data and (if relevant) additional baseline covariates. <code>surv.data</code> should at least contain a subject id (called <code>id</code> ), the time to event outcome ( <code>time</code> ), and binary event variable ( <code>event</code> )
t.from.base	name of the variable containing time from baseline in <code>long.data</code>
n.boots	number of bootstrap samples to be used in the cluster bootstrap optimism correction procedure (CBOCP). If 0, no bootstrapping is performed
n.cores	number of cores to use to parallelize the computation of the CBOCP procedure. If <code>ncores = 1</code> (default), no parallelization is done. Pro tip: you can use <code>parallel::detectCores()</code> to check how many cores are available on your computer
verbose	if TRUE (default and recommended value), information on the ongoing computations is printed in the console

### Value

A list containing the following objects:

- `call.info`: a list containing the following function call information: `call`, `y.names`, `fixefs`, `ranefs`;
- `lmm.fits.orig`: a list with the LMMs fitted on the original dataset (it should comprise as many LMMs as the elements of `y.names` are);
- `df.sanitized`: a sanitized version of the supplied `long.data` dataframe, without the longitudinal measurements that are taken after the event or after censoring;



fit\_mlpms

*Step 1 of PRC-MLPMM (estimation of the linear mixed models)***Description**

This function performs the first step for the estimation of the PRC-MLPMM model proposed in Signorelli et al. (2021, in review)

**Usage**

```
fit_mlpms(y.names, fixefs, ranef.time, randint.items = TRUE, long.data,
  surv.data, t.from.base, n.boots = 0, n.cores = 1, verbose = TRUE,
  maxiter = 1000, conv = rep(0.001, 3))
```

**Arguments**

y.names	a list with the names of the response variables which the MLPMMs have to be fitted to. Each element in the list contains all the items used to reconstruct a latent biological process of interest
fixefs	a fixed effects formula for the model, where the time variable (specified also in ranef.time) is included as first element and within the function contrast(). Examples: ~ contrast(age), ~ contrast(age) + group + treatment
ranef.time	a character with the name of the time variable for which to include a shared random slope
randint.items	logical: should item-specific random intercepts be included in the MLCMMs? Default is TRUE. It can also be a vector, with different values for different elements of y.names
long.data	a data frame with the longitudinal predictors, comprehensive of a variable called id with the subject ids
surv.data	a data frame with the survival data and (if relevant) additional baseline covariates. surv.data should at least contain a subject id (called id), the time to event outcome (time), and binary event variable (event)
t.from.base	name of the variable containing time from baseline in long.data
n.boots	number of bootstrap samples to be used in the cluster bootstrap optimism correction procedure (CBOCP). If 0, no bootstrapping is performed
n.cores	number of cores to use to parallelize the computation of the CBOCP procedure. If ncores = 1 (default), no parallelization is done. Pro tip: you can use parallel::detectCores() to check how many cores are available on your computer
verbose	if TRUE (default and recommended value), information on the ongoing computations is printed in the console
maxiter	maximum number of iterations to use when calling the function multlcm. Default is 1e3
conv	a vector containing the three convergence criteria (convB, convL and convG) to use when calling the function multlcm. Default is c(1e-3, 1e-3, 1e-3)



```

# specify options for cluster bootstrap optimism correction
# procedure and for parallel computing
do.bootstrap = FALSE
# IMPORTANT: set do.bootstrap = TRUE to compute the optimism correction!
n.boots = ifelse(do.bootstrap, 100, 0)
parallelize = TRUE
# IMPORTANT: set parallelize = TRUE to speed computations up!
if (!parallelize) n.cores = 1
if (parallelize) {
  # identify number of available cores on your machine
  n.cores = parallel::detectCores()
  if (is.na(n.cores)) n.cores = 1
}

# step 1 of PRC-MLPMM: estimate the MLPMMs
y.names = vector('list', length(n.items))
for (i in 1:length(n.items)) {
  y.names[[i]] = paste('marker', i, '_', 1:n.items[i], sep = '')
}

step1 = fit_mlpmm(y.names, fixefs = ~ contrast(age),
                 ranef.time = age, randint.items = TRUE,
                 long.data = simdata$long.data,
                 surv.data = simdata$surv.data,
                 t.from.base = t.from.base,
                 n.boots = n.boots, n.cores = n.cores)

```

---

fit\_prclmm

*Step 3 of PRC-LMM (estimation of the penalized Cox model(s))*


---

### Description

This function performs the third step for the estimation of the PRC-LMM model proposed in Signorelli et al. (2021, in review)

### Usage

```

fit_prclmm(object, surv.data, baseline.covs = NULL, penalty = "ridge",
           standardize = TRUE, pfac.base.covs = 0, n.alpha.elnet = 11,
           n.folds.elnet = 5, n.cores = 1, verbose = TRUE)

```

### Arguments

object	the output of step 2 of the PRC-LMM procedure, as produced by the <a href="#">summarize_lmms</a> function
surv.data	a data frame with the survival data and (if relevant) additional baseline covariates. <code>surv.data</code> should at least contain a subject id (called <code>id</code> ), the time to event outcome ( <code>time</code> ), and binary event variable ( <code>event</code> )

<code>baseline.covs</code>	a formula specifying the variables (e.g., baseline age) in <code>surv.data</code> that should be included as baseline covariates in the penalized Cox model. Example: <code>baseline.covs = '~ baseline.age'</code> . Default is NULL
<code>penalty</code>	the type of penalty function used for regularization. Default is 'ridge', other possible values are 'elasticnet' and 'lasso'
<code>standardize</code>	logical argument: should the predicted random effects be standardized when included in the penalized Cox model? Default is TRUE
<code>pfac.base.covs</code>	a single value, or a vector of values, indicating whether the baseline covariates (if any) should be penalized (1) or not (0). Default is <code>pfac.base.covs = 0</code> (no penalization of all baseline covariates)
<code>n.alpha.elnet</code>	number of alpha values for the two-dimensional grid of tuning parameters in elasticnet. Only relevant if <code>penalty = 'elasticnet'</code> . Default is 11, so that the resulting alpha grid is <code>c(1, 0.95, 0.90, ..., 0.05, 0)</code>
<code>n.folds.elnet</code>	number of folds to be used for the selection of the tuning parameter in elasticnet. Only relevant if <code>penalty = 'elasticnet'</code> . Default is 5
<code>n.cores</code>	number of cores to use to parallelize the computation of the cluster bootstrap optimism correction procedure. If <code>ncores = 1</code> (default), no parallelization is done. Pro tip: you can use <code>parallel::detectCores()</code> to check how many cores are available on your computer
<code>verbose</code>	if TRUE (default and recommended value), information on the ongoing computations is printed in the console

### Value

A list containing the following objects:

- `call`: the function call
- `pcox.orig`: the penalized Cox model fitted on the original dataset;
- `surv.data`: the supplied survival data (ordered by subject id)
- `n.boots`: number of bootstrap samples;
- `boot.ids`: a list with the ids of bootstrapped subjects (when `n.boots > 0`);
- `pcox.boot`: a list where each element is a fitted penalized Cox model for a given bootstrap sample (when `n.boots > 0`).

### Author(s)

Mirko Signorelli

### References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

### See Also

[fit\\_lmms](#) (step 1), [summarize\\_lmms](#) (step 2), [performance\\_prc](#)



**Examples**

```

# generate example data
set.seed(1234)
p = 4 # number of longitudinal predictors
simdata = simulate_prclmm_data(n = 100, p = p, p.relev = 2,
                              seed = 123, t.values = c(0, 0.2, 0.5, 1, 1.5, 2))

# specify options for cluster bootstrap optimism correction
# procedure and for parallel computing
do.bootstrap = FALSE
# IMPORTANT: set do.bootstrap = TRUE to compute the optimism correction!
n.boots = ifelse(do.bootstrap, 100, 0)
parallelize = FALSE
# IMPORTANT: set parallelize = TRUE to speed computations up!
if (!parallelize) n.cores = 1
if (parallelize) {
  # identify number of available cores on your machine
  n.cores = parallel::detectCores()
  if (is.na(n.cores)) n.cores = 1
}

# step 1 of PRC-LMM: estimate the LMMs
y.names = paste('marker', 1:p, sep = '')
step1 = fit_lmms(y.names = y.names,
                fixeFs = ~ age, ranefs = ~ age | id,
                long.data = simdata$long.data,
                surv.data = simdata$surv.data,
                t.from.base = t.from.base,
                n.boots = n.boots, n.cores = n.cores)

# step 2 of PRC-LMM: compute the summaries
# of the longitudinal outcomes
step2 = summarize_lmms(object = step1, n.cores = n.cores)

# step 3 of PRC-LMM: fit the penalized Cox models
step3 = fit_prclmm(object = step2, surv.data = simdata$surv.data,
                  baseline.covs = ~ baseline.age,
                  penalty = 'ridge', n.cores = n.cores)

```

---

fit\_prclpmm

*Step 3 of PRC-MLPMM (estimation of the penalized Cox model(s))*


---

**Description**

This function performs the third step for the estimation of the PRC-MLPMM model proposed in Signorelli et al. (2021, in review)

**Usage**

```
fit_prcmlpmm(object, surv.data, baseline.covs = NULL, include.b0s = TRUE,
  penalty = "ridge", standardize = TRUE, pfac.base.covs = 0,
  n.alpha.elnet = 11, n.folds.elnet = 5, n.cores = 1, verbose = TRUE)
```

**Arguments**

<code>object</code>	the output of step 2 of the PRC-MLPMM procedure, as produced by the <a href="#">summarize_mlpmms</a> function
<code>surv.data</code>	a data frame with the survival data and (if relevant) additional baseline covariates. <code>surv.data</code> should at least contain a subject id (called <code>id</code> ), the time to event outcome ( <code>time</code> ), and binary event variable ( <code>event</code> )
<code>baseline.covs</code>	a formula specifying the variables (e.g., baseline age) in <code>surv.data</code> that should be included as baseline covariates in the penalized Cox model. Example: <code>baseline.covs = '~ baseline.age'</code> . Default is <code>NULL</code>
<code>include.b0s</code>	logical. If <code>TRUE</code> , the PRC-MLPMM(U+B) model is estimated; if <code>FALSE</code> , the PRC-MLPMM(U) model is estimated. See Signorelli et al. (2021) for details
<code>penalty</code>	the type of penalty function used for regularization. Default is <code>'ridge'</code> , other possible values are <code>'elasticnet'</code> and <code>'lasso'</code>
<code>standardize</code>	logical argument: should the predicted random effects be standardized when included in the penalized Cox model? Default is <code>TRUE</code>
<code>pfac.base.covs</code>	a single value, or a vector of values, indicating whether the baseline covariates (if any) should be penalized (1) or not (0). Default is <code>pfac.base.covs = 0</code> (no penalization of all baseline covariates)
<code>n.alpha.elnet</code>	number of alpha values for the two-dimensional grid of tuning parameters in elasticnet. Only relevant if <code>penalty = 'elasticnet'</code> . Default is 11, so that the resulting alpha grid is <code>c(1, 0.95, 0.90, ..., 0.05, 0)</code>
<code>n.folds.elnet</code>	number of folds to be used for the selection of the tuning parameter in elasticnet. Only relevant if <code>penalty = 'elasticnet'</code> . Default is 5
<code>n.cores</code>	number of cores to use to parallelize the computation of the cluster bootstrap optimism correction procedure. If <code>ncores = 1</code> (default), no parallelization is done. Pro tip: you can use <code>parallel::detectCores()</code> to check how many cores are available on your computer
<code>verbose</code>	if <code>TRUE</code> (default and recommended value), information on the ongoing computations is printed in the console

**Value**

A list containing the following objects:

- `call`: the function call
- `pcox.orig`: the penalized Cox model fitted on the original dataset;
- `surv.data`: the supplied survival data (ordered by subject id)
- `n.boots`: number of bootstrap samples;
- `boot.ids`: a list with the ids of bootstrapped subjects (when `n.boots > 0`);

- `pcox.boot`: a list where each element is a fitted penalized Cox model for a given bootstrap sample (when `n.boots > 0`).

### Author(s)

Mirko Signorelli

### References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

### See Also

[fit\\_mlpmm](#) (step 1), [summarize\\_mlpmm](#) (step 2), [performance\\_prc](#)

### Examples

```
# generate example data
set.seed(123)
n.items = c(4,2,2,3,4,2)
simdata = simulate_prcmlpmm_data(n = 100, p = length(n.items),
                                p.relev = 3, n.items = n.items,
                                type = 'u+b', seed = 1)

# specify options for cluster bootstrap optimism correction
# procedure and for parallel computing
do.bootstrap = FALSE
# IMPORTANT: set do.bootstrap = TRUE to compute the optimism correction!
n.boots = ifelse(do.bootstrap, 100, 0)
parallelize = TRUE
# IMPORTANT: set parallelize = TRUE to speed computations up!
if (!parallelize) n.cores = 1
if (parallelize) {
  # identify number of available cores on your machine
  n.cores = parallel::detectCores()
  if (is.na(n.cores)) n.cores = 1
}

# step 1 of PRC-MLPMM: estimate the MLPMMs
y.names = vector('list', length(n.items))
for (i in 1:length(n.items)) {
  y.names[[i]] = paste('marker', i, '_', 1:n.items[i], sep = '')
}

step1 = fit_mlpmm(y.names, fixefs = ~ contrast(age),
                 ranef.time = age, randint.items = TRUE,
                 long.data = simdata$long.data,
                 surv.data = simdata$surv.data,
                 t.from.base = t.from.base,
```

```

n.boots = n.boots, n.cores = n.cores)

# step 2 of PRC-MLPMM: compute the summaries
step2 = summarize_mlpmm(object = step1, n.cores = n.cores)

# step 3 of PRC-LMM: fit the penalized Cox models
step3 = fit_prcmlmm(object = step2, surv.data = simdata$surv.data,
                    baseline.covs = ~ baseline.age,
                    include.b0s = TRUE,
                    penalty = 'ridge', n.cores = n.cores)

```

---

performance\_prc      *Predictive performance of the PRC-LMM and PRC-MLPMM models*

---

### Description

This function computes the naive and optimism-corrected measures of performance (C index and time-dependent AUC) for the PRC models proposed in Signorelli et al. (2021, in review). The optimism correction is computed based on a cluster bootstrap optimism correction procedure (CBOCP)

### Usage

```
performance_prc(step2, step3, times = 1, n.cores = 1, verbose = TRUE)
```

### Arguments

step2	the output of either <code>summarize_lmms</code> or <code>summarize_mlpmm</code> (step 2 of the estimation of PRC)
step3	the output of <code>fit_prcmlmm</code> or <code>fit_prcmlpm</code> (step 3 of PRC)
times	numeric vector with the time points at which to estimate the time-dependent AUC
n.cores	number of cores to use to parallelize the computation of the CBOCP procedure. If <code>ncores = 1</code> (default), no parallelization is done. Pro tip: you can use <code>parallel::detectCores()</code> to check how many cores are available on your computer
verbose	if TRUE (default and recommended value), information on the ongoing computations is printed in the console

### Value

A list containing the following objects:

- `call`: the function call;
- `concordance`: a data frame with the naive and optimism-corrected estimates of the concordance (C) index;
- `tdAUC`: a data frame with the naive and optimism-corrected estimates of the time-dependent AUC at the desired time points.

**Author(s)**

Mirko Signorelli

**References**

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

**See Also**

for the PRC-LMM model: [fit\\_lmms](#) (step 1), [summarize\\_lmms](#) (step 2) and [fit\\_prclmm](#) (step 3);  
for the PRC-MLPMM model: [fit\\_mlpms](#) (step 1), [summarize\\_mlpms](#) (step 2) and [fit\\_prclpmm](#) (step 3).

**Examples**

```
data(fitted_prclmm)

parallelize = FALSE
# IMPORTANT: set parallelize = TRUE to speed computations up!
if (!parallelize) n.cores = 1
if (parallelize) {
  # identify number of available cores on your machine
  n.cores = parallel::detectCores()
  if (is.na(n.cores)) n.cores = 1
}

# compute the performance measures
perf = performance_prc(fitted_prclmm$step2, fitted_prclmm$step3,
  times = c(0.5, 1, 1.5, 2), n.cores = n.cores)

# concordance index:
perf$concordance
# time-dependent AUC:
perf$tdAUC
```

---

simulate\_prclmm\_data *Simulate data that can be used to fit the PRC-LMM model*

---

**Description**

This function allows to simulate a survival outcome from longitudinal predictors. Specifically, the longitudinal predictors are simulated from linear mixed models (LMMs), and the survival outcome from a Weibull model where the time to event depends on the random effects from the LMMs. It is an implementation of the simulation method used in Signorelli et al. (2021, in review)

## Usage

```
simulate_prclmm_data(n = 100, p = 10, p.relev = 4, lambda = 0.2,  
  nu = 2, seed = 1, base.age.range = c(3, 5), cens.range = c(0.5, 10),  
  t.values = c(0, 0.5, 1, 2))
```

## Arguments

n	sample size
p	number of longitudinal outcomes
p.relev	number of longitudinal outcomes that are associated with the survival outcome (min: 1, max: p)
lambda	Weibull location parameter, positive
nu	Weibull scale parameter, positive
seed	random seed (defaults to 1)
base.age.range	range for age at baseline (set it equal to c(0, 0) if you want all subjects to enter the study at the same age)
cens.range	range for censoring times
t.values	vector specifying the time points at which longitudinal measurements are collected (NB: for simplicity, this function assumes a balanced designed; however, pencial is designed to work both with balanced and with unbalanced designs!)

## Value

A list containing the following elements:

- a dataframe `long.data` with data on the longitudinal predictors, comprehensive of a subject id (`id`), baseline age (`base.age`), time from baseline (`t.from.base`) and the longitudinal biomarkers;
- a dataframe `surv.data` with the survival data: a subject id (`id`), baseline age (`baseline.age`), the time to event outcome (`time`) and a binary vector (`event`) that is 1 if the event is observed, and 0 in case of right-censoring;
- `perc.cens` the proportion of censored individuals in the simulated dataset.

## Author(s)

Mirko Signorelli

## References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

**Examples**

```
# generate example data
simdata = simulate_prcmlmm_data(n = 20, p = 10,
                               p.relev = 4, seed = 1)
# view the longitudinal markers:
library(ptmixed)
make.spaghetti(x = age, y = marker1,
              id = id, group = id,
              data = simdata$long.data,
              legend.inset = - 1)
# proportion of censored subjects
simdata$censoring.prop
# visualize KM estimate of survival
library(survival)
surv.obj = Surv(time = simdata$surv.data$time,
                event = simdata$surv.data$event)
kaplan <- survfit(surv.obj ~ 1,
                 type="kaplan-meier")
plot(kaplan)
```

---

simulate\_prcmlpmm\_data

*Simulate data that can be used to fit the PRC-LMM model*

---

**Description**

This function allows to simulate a survival outcome from longitudinal predictors. Specifically, the longitudinal predictors are simulated from multivariate latent process mixed models (MLPMMs), and the survival outcome from a Weibull model where the time to event depends on the random effects from the MLPMMs. It is an implementation of the simulation method used in Signorelli et al. (2021, in review)

**Usage**

```
simulate_prcmlpmm_data(n = 100, p = 5, p.relev = 2, n.items = c(3, 2,
  3, 4, 1), type = "u", lambda = 0.2, nu = 2, seed = 1,
  base.age.range = c(3, 5), cens.range = c(0.5, 10), t.values = c(0, 0.5,
  1, 2))
```

**Arguments**

n	sample size
p	number of longitudinal latent processes
p.relev	number of latent processes that are associated with the survival outcome (min: 1, max: p)
n.items	number of items that are observed for each latent process of interest. It must be either a scalar, or a vector of length p

type	the type of relation between the longitudinal outcomes and survival time. Two values can be used: 'u' refers to the PRC-MLPMM(U) model, and 'u+b' to the PRC-MLPMM(U+B) model presented in Section 2.3 of Signorelli et al. (2021). See the article for the mathematical details
lambda	Weibull location parameter, positive
nu	Weibull scale parameter, positive
seed	random seed (defaults to 1)
base.age.range	range for age at baseline (set it equal to c(0, 0) if you want all subjects to enter the study at the same age)
cens.range	range for censoring times
t.values	vector specifying the time points at which longitudinal measurements are collected (NB: for simplicity, this function assumes a balanced designed; however, pncal is designed to work both with balanced and with unbalanced designs!)

### Value

A list containing the following elements:

- a dataframe `long.data` with data on the longitudinal predictors, comprehensive of a subject id (`id`), baseline age (`base.age`), time from baseline (`t.from.base`) and the longitudinal biomarkers;
- a dataframe `surv.data` with the survival data: a subject id (`id`), baseline age (`baseline.age`), the time to event outcome (`time`) and a binary vector (`event`) that is 1 if the event is observed, and 0 in case of right-censoring;
- `perc.cens` the proportion of censored individuals in the simulated dataset.

### Author(s)

Mirko Signorelli

### References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

### Examples

```
# generate example data
simdata = simulate_prcmlpmm_data(n = 40, p = 6,
                                p.relev = 3, n.items = c(3,4,2,5,4,2),
                                type = 'u+b', seed = 1)

# names of the longitudinal outcomes:
names(simdata$long.data)
# markerx_y is the y-th item for latent process (LP) x
# we have 6 latent processes of interest, and for LP1
# we measure 3 items, for LP2 4, for LP3 2 items, and so on
```



```
# visualize trajectories of marker1_1
library(ptmixed)
make.spaghetti(x = age, y = marker1_1,
              id = id, group = id,
              data = simdata$long.data,
              legend.inset = - 1)

# proportion of censored subjects
simdata$censoring.prop
# visualize KM estimate of survival
library(survival)
surv.obj = Surv(time = simdata$urv.data$time,
               event = simdata$urv.data$event)
kaplan <- survfit(surv.obj ~ 1,
                 type="kaplan-meier")
plot(kaplan)
```

---

simulate\_t\_weibull      *Generate survival data from a Weibull model*

---

## Description

This function implements the algorithm proposed by Bender et al. (2005) to simulate survival times from a Weibull model

## Usage

```
simulate_t_weibull(n, lambda, nu, X, beta, seed = 1)
```

## Arguments

n	sample size
lambda	Weibull location parameter, positive
nu	Weibull scale parameter, positive
X	design matrix (n rows, p columns)
beta	p-dimensional vector of regression coefficients associated to X
seed	random seed (defaults to 1)

## Value

A vector of survival times

## Author(s)

Mirko Signorelli

## References

Bender, R., Augustin, T., & Blettner, M. (2005). Generating survival times to simulate Cox proportional hazards models. *Statistics in medicine*, 24(11), 1713-1723.

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

## Examples

```
# generate example data
set.seed(1)
n = 50
X = cbind(matrix(1, n, 1),
           matrix(rnorm(n*9, sd = 0.7), n, 9))
beta = rnorm(10, sd = 0.7)
times = simulate_t_weibull(n = n, lambda = 1, nu = 2,
                          X = X, beta = beta)
hist(times, 20)
```

---

summarize\_lmms

*Step 2 of PRC-LMM (computation of the predicted random effects)*

---

## Description

This function performs the second step for the estimation of the PRC-LMM model proposed in Signorelli et al. (2021, in review)

## Usage

```
summarize_lmms(object, n.cores = 1, verbose = TRUE)
```

## Arguments

object	a list of objects as produced by <code>fit_lmms</code>
n.cores	number of cores to use to parallelize the computation of the cluster bootstrap optimism correction procedure. If <code>ncores = 1</code> (default), no parallelization is done. Pro tip: you can use <code>parallel::detectCores()</code> to check how many cores are available on your computer
verbose	if TRUE (default and recommended value), information on the ongoing computations is printed in the console

## Value

A list containing the following objects:

- `call`: the function call
- `ranef.orig`: a matrix with the predicted random effects computed for the original data;

- `n.boots`: number of bootstrap samples;
- `boot.ids`: a list with the ids of bootstrapped subjects (when `n.boots > 0`);
- `ranef.boot.train`: a list where each element is a matrix that contains the predicted random effects for each bootstrap sample (when `n.boots > 0`);
- `ranef.boot.valid`: a list where each element is a matrix that contains the predicted random effects on the original data, based on the lmms fitted on the cluster bootstrap samples (when `n.boots > 0`);

### Author(s)

Mirko Signorelli

### References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

### See Also

[fit\\_lmms](#) (step 1), [fit\\_prclmm](#) (step 3), [performance\\_prc](#)

### Examples

```
# generate example data
set.seed(1234)
p = 4 # number of longitudinal predictors
simdata = simulate_prclmm_data(n = 100, p = p, p.relev = 2,
                              seed = 123, t.values = c(0, 0.2, 0.5, 1, 1.5, 2))

# specify options for cluster bootstrap optimism correction
# procedure and for parallel computing
do.bootstrap = FALSE
# IMPORTANT: set do.bootstrap = TRUE to compute the optimism correction!
n.boots = ifelse(do.bootstrap, 100, 0)
parallelize = FALSE
# IMPORTANT: set parallelize = TRUE to speed computations up!
if (!parallelize) n.cores = 1
if (parallelize) {
  # identify number of available cores on your machine
  n.cores = parallel::detectCores()
  if (is.na(n.cores)) n.cores = 1
}

# step 1 of PRC-LMM: estimate the LMMs
y.names = paste('marker', 1:p, sep = '')
step1 = fit_lmms(y.names = y.names,
                 fixeFs = ~ age, ranefs = ~ age | id,
                 long.data = simdata$long.data,
                 surv.data = simdata$surv.data,
                 t.from.base = t.from.base,
```

```

n.boots = n.boots, n.cores = n.cores)

# step 2 of PRC-LMM: compute the summaries
# of the longitudinal outcomes
step2 = summarize_lmms(object = step1, n.cores = n.cores)

```

---

summarize_mlpmms	<i>Step 2 of PRC-MLPMM (computation of the predicted random effects)</i>
------------------	--

---

### Description

This function performs the second step for the estimation of the PRC-MLPMM model proposed in Signorelli et al. (2021, in review)

### Usage

```
summarize_mlpmms(object, n.cores = 1, verbose = TRUE)
```

### Arguments

object	a list of objects as produced by <code>fit_mlpmms</code>
n.cores	number of cores to use to parallelize the computation of the cluster bootstrap optimism correction procedure. If <code>ncores = 1</code> (default), no parallelization is done. Pro tip: you can use <code>parallel::detectCores()</code> to check how many cores are available on your computer
verbose	if TRUE (default and recommended value), information on the ongoing computations is printed in the console

### Value

A list containing the following objects:

- `call`: the function call
- `ranef.orig`: a matrix with the predicted random effects computed for the original data;
- `n.boots`: number of bootstrap samples;
- `boot.ids`: a list with the ids of bootstrapped subjects (when `n.boots > 0`);
- `ranef.boot.train`: a list where each element is a matrix that contains the predicted random effects for each bootstrap sample (when `n.boots > 0`);
- `ranef.boot.valid`: a list where each element is a matrix that contains the predicted random effects on the original data, based on the `mlpmms` fitted on the cluster bootstrap samples (when `n.boots > 0`);

### Author(s)

Mirko Signorelli

## References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

## See Also

[fit\\_mlpmm](#) (step 1), [fit\\_prcmlpmm](#) (step 3), [performance\\_prc](#)

## Examples

```
# generate example data
set.seed(123)
n.items = c(4,2,2,3,4,2)
simdata = simulate_prcmlpmm_data(n = 100, p = length(n.items),
                                p.relev = 3, n.items = n.items,
                                type = 'u+b', seed = 1)

# specify options for cluster bootstrap optimism correction
# procedure and for parallel computing
do.bootstrap = FALSE
# IMPORTANT: set do.bootstrap = TRUE to compute the optimism correction!
n.boots = ifelse(do.bootstrap, 100, 0)
parallelize = TRUE
# IMPORTANT: set parallelize = TRUE to speed computations up!
if (!parallelize) n.cores = 1
if (parallelize) {
  # identify number of available cores on your machine
  n.cores = parallel::detectCores()
  if (is.na(n.cores)) n.cores = 1
}

# step 1 of PRC-MLPMM: estimate the MLPMMs
y.names = vector('list', length(n.items))
for (i in 1:length(n.items)) {
  y.names[[i]] = paste('marker', i, '_', 1:n.items[i], sep = '')
}

step1 = fit_mlpmm(y.names, fixefs = ~ contrast(age),
                 ranef.time = age, randint.items = TRUE,
                 long.data = simdata$long.data,
                 surv.data = simdata$surv.data,
                 t.from.base = t.from.base,
                 n.boots = n.boots, n.cores = n.cores)

# step 2 of PRC-MLPMM: compute the summaries
step2 = summarize_mlpmm(object = step1, n.cores = n.cores)
```

---

survpred_prc	<i>Compute the predicted survival probabilities obtained from the PRC models</i>
--------------	--

---

### Description

This function computes the predictive survival probabilities for the for the PRC models proposed in Signorelli et al. (2021, in review)

### Usage

```
survpred_prc(step2, step3, times = 1)
```

### Arguments

step2	the output of either <code>summarize_lmms</code> or <code>summarize_mlpmms</code> (step 2 of the estimation of PRC)
step3	the output of <code>fit_prclmm</code> or <code>fit_prcmlpmm</code> (step 3 of PRC)
times	numeric vector with the time points at which to estimate the time-dependent AUC

### Value

A data frame with the predicted survival probabilities computed at the supplied time points

### Author(s)

Mirko Signorelli

### References

Signorelli, M., Spitali, P., Al-Khalili Szigyarto, C, The MARK-MD Consortium, Tsonaka, R. (2021). Penalized regression calibration: a method for the prediction of survival outcomes using complex longitudinal and high-dimensional data. arXiv preprint: arXiv:2101.04426.

### See Also

for the PRC-LMM model: `fit_lmms` (step 1), `summarize_lmms` (step 2) and `fit_prclmm` (step 3);  
for the PRC-MLPMM model: `fit_mlpmms` (step 1), `summarize_mlpmms` (step 2) and `fit_prcmlpmm` (step 3).

### Examples

```
# generate example data
set.seed(1234)
p = 4 # number of longitudinal predictors
simdata = simulate_prclmm_data(n = 100, p = p, p.relev = 2,
                              seed = 123, t.values = c(0, 0.2, 0.5, 1, 1.5, 2))
```

```
# step 1 of PRC-LMM: estimate the LMMs
y.names = paste('marker', 1:p, sep = '')
step1 = fit_lmms(y.names = y.names,
                 fixeFs = ~ age, ranefs = ~ age | id,
                 long.data = simdata$long.data,
                 surv.data = simdata$surv.data,
                 t.from.base = t.from.base,
                 n.boots = 0)

# step 2 of PRC-LMM: compute the summaries
# of the longitudinal outcomes
step2 = summarize_lmms(object = step1)

# step 3 of PRC-LMM: fit the penalized Cox models
step3 = fit_prclmm(object = step2, surv.data = simdata$surv.data,
                  baseline.covs = ~ baseline.age,
                  penalty = 'ridge')

# predict survival probabilities at times 1, 2, 3
surv.probs = survpred_prc(step2, step3, times = 1:3)
head(surv.probs)
```

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