

# Package ‘trajmsm’

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**Type** Package

**Title** Marginal Structural Models with Latent Class Growth Analysis of Treatment Trajectories

**Version** 0.1.4

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**Description** Implements marginal structural models combined with a latent class growth analysis framework for assessing the causal effect of treatment trajectories. Based on the approach described in “Marginal Structural Models with Latent Class Growth Analysis of Treatment Trajectories” Diop, A., Sirois, C., Guertin, J.R., Schnitzer, M.E., Candas, B., Cossette, B., Poirier, P., Brophy, J., Mésidor, M., Blais, C. and Hamel, D., (2023) <[doi:10.1177/09622802231202384](https://doi.org/10.1177/09622802231202384)>.

**License** GPL (>= 3)

**Encoding** UTF-8

**Imports** stats, e1071, flexmix, ggplot2, survival, sandwich, utils

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**URL** <https://github.com/awamaeva/R-package-trajmsm>

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**NeedsCompilation** no

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|            |                            |
|------------|----------------------------|
| build_traj | <i>Wrapper for flexmix</i> |
|------------|----------------------------|

---

**Description**

Call the package flexmix to build trajectory groups

**Usage**

```
build_traj(  
  obsdata,  
  formula,  
  number_traj,  
  identifier,  
  family = "binomial",  
  seed = 945,  
  control = list(iter.max = 1000, minprior = 0),  
  ...  
)
```

**Arguments**

|             |                                                                                  |
|-------------|----------------------------------------------------------------------------------|
| obsdata     | Data to build trajectory groups in long format.                                  |
| formula     | Designate the formula to model the longitudinal variable of interest.            |
| number_traj | An integer to fix the number of trajectory groups.                               |
| identifier  | A string to designate the column name for the unique identifier.                 |
| family      | Designate the type of distribution ("gaussian", "binomial", "poisson", "gamma"). |
| seed        | Set a seed for replicability.                                                    |
| control     | Object of class FLXcontrol.                                                      |
| ...         | Additional arguments passed to the flexmix function.                             |

**Value**

A list containing the posterior probability matrix and the fitted trajectory model.

**Examples**

```
obsdata_long = gendata(n = 1000, format = "long", total_followup = 6, seed = 945)
formula = as.formula(cbind(statins, 1 - statins) ~ time)
restraj = build_traj(obsdata = obsdata_long, number_traj = 3, formula = formula, identifier = "id")
```

gendata

*Generate data trajectories for MSM***Description**

Provides datasets for running examples for LCGA-MSM and LCGA-HRMSM.

**Usage**

```
gendata(
  n,
  include_censor = FALSE,
  format = c("long", "wide"),
  start_year = 2011,
  total_followup,
  timedep_outcome = FALSE,
  seed
)
```

**Arguments**

|                 |                                                                             |
|-----------------|-----------------------------------------------------------------------------|
| n               | Number of observations to generate.                                         |
| include_censor  | Logical, if TRUE, includes censoring.                                       |
| format          | Character, either "long" or "wide" for the format of the output data frame. |
| start_year      | Baseline year.                                                              |
| total_followup  | Number of measuring times.                                                  |
| timedep_outcome | Logical, if TRUE, includes a time-dependent outcome.                        |
| seed            | Use a specific seed value to ensure the simulated data is replicable.       |

**Value**

A data frame with generated data trajectories.

**Examples**

```
gendata(n = 100, include_censor = FALSE, format = "wide", total_followup = 3, seed = 945)
```

gformula

*Counterfactual means via G-Formula***Description**

Calculates counterfactual means using the g-formula approach.

**Usage**

```
gformula(
  formula,
  baseline,
  covariates,
  treatment,
  outcome,
  ntimes_interval,
  obsdata
)
```

**Arguments**

|                 |                                                          |
|-----------------|----------------------------------------------------------|
| formula         | Specification of the model for the outcome to be fitted. |
| baseline        | Names of the baseline covariates.                        |
| covariates      | Names of the time-varying covariates (should be a list). |
| treatment       | Names of the time-varying treatment.                     |
| outcome         | Name of the outcome variable.                            |
| ntimes_interval | Length of a time-interval (s).                           |
| obsdata         | Observed data in wide format.                            |

**Value**

list\_gform\_countermeans  
List of counterfactual means obtained with g-formula.

**Author(s)**

Awa Diop, Denis Talbot

**Examples**

```
obsdata = gendata(n = 1000, format = "wide", total_followup = 6, seed = 945)
years <- 2011:2016
baseline_var <- c("age", "sex")
variables <- c("hyper", "bmi")
var_cov <- c("statins", "hyper", "bmi")
covariates <- lapply(years, function(year) {
```

```

paste0(variables, year)))
treatment_var <- paste0("statins", 2011:2016)
formula = paste0("y ~", paste0(treatment_var, collapse = "+"), "+",
                  paste0(unlist(covariates), collapse = "+"), "+",
                  paste0(baseline_var, collapse = "+"))
res_gform <- gformula(formula = formula, baseline = baseline_var, covariates = covariates,
                      treatment = treatment_var, outcome = "y", ntimes_interval = 6, obsdata = obsdata )

```

ggtraj

*ggplot Trajectory***Description**

Use "ggplot2" to plot trajectory groups produced by the function "build\_traj" using the observed treatment.

**Usage**

```
ggtraj(traj_data, treatment, time, identifier, class, FUN = mean, ...)
```

**Arguments**

|            |                                                                                |
|------------|--------------------------------------------------------------------------------|
| traj_data  | Merged datasets containing observed data in long format and trajectory groups. |
| treatment  | Name of the time-varying treatment.                                            |
| time       | Name of the time variable.                                                     |
| identifier | Name of the identifier variable.                                               |
| class      | Name of the trajectory groups.                                                 |
| FUN        | Specify which statistics to display, by default calculate the mean.            |
| ...        | Additional arguments to be passed to ggplot functions.                         |

**Value**

A ggplot object representing the trajectory groups using the observed treatment.

**Examples**

```

obsdata_long = gendata(n = 1000, format = "long", total_followup = 12, seed = 945)
restraj = build_traj(obsdata = obsdata_long, number_traj = 3,
formula = as.formula(cbind(statins, 1 - statins) ~ time), identifier = "id")
datapost = restraj$data_post
head(datapost)
traj_data_long <- merge(obsdata_long, datapost, by = "id")
  AggFormula <- as.formula(paste("statins", "~", "time", "+", "class"))
  Aggtraj_data <- aggregate(AggFormula, data = traj_data_long, FUN = mean)
  Aggtraj_data
#Aggtraj_data with labels

```

```

traj_data_long[, "traj_group"] <- factor(iffelse(traj_data_long[, "class"] == "3" ,"Group1" ,
iffelse (traj_data_long[, "class"]== "1" , "Group2" ,"Group3"))
AggFormula <- as.formula(paste("statins", "~", "time", "+", "traj_group"))
Aggtraj_data <- aggregate(AggFormula, data = traj_data_long, FUN = mean)
ggtraj(traj_data = Aggtraj_data,
treatment = "statins",time= "time",identifier="id",class = "traj_group", FUN = mean)

```

---

inverse\_probability\_weighting

*Inverse Probability Weighting*

---

## Description

Compute stabilized and unstabilized weights, with or without censoring.

## Usage

```

inverse_probability_weighting(
  numerator = c("stabilized", "unstabilized"),
  identifier,
  baseline,
  covariates,
  treatment,
  include_censor = FALSE,
  censor,
  obsdata
)

```

## Arguments

|                |                                                                  |
|----------------|------------------------------------------------------------------|
| numerator      | To choose between stabilized and unstabilized weights.           |
| identifier     | Name of the column of the unique identifier.                     |
| baseline       | Name of the baseline covariates.                                 |
| covariates     | Name of the time-varying covariates.                             |
| treatment      | Name of the time-varying treatment.                              |
| include_censor | Logical value TRUE/FALSE to include or not a censoring variable. |
| censor         | Name of the censoring variable.                                  |
| obsdata        | Observed data in wide format.                                    |

## Value

Inverse Probability Weights (Stabilized and Unstabilized) with and without censoring.

## Author(s)

Awa Diop, Denis Talbot

## Examples

```
obsdata = gendata(n = 1000, format = "wide", total_followup = 3, seed = 945)
baseline_var <- c("age", "sex")
covariates <- list(c("hyper2011", "bmi2011"),
  c("hyper2012", "bmi2012"), c("hyper2013", "bmi2013"))
treatment_var <- c("statins2011", "statins2012", "statins2013")
stabilized_weights = inverse_probability_weighting(numerator = "stabilized",
  identifier = "id", covariates = covariates, treatment = treatment_var,
  baseline = baseline_var, obsdata = obsdata)
```

---

pltmle

---

*Counterfactual means for a Pooled LTMLE*


---

## Description

Function to estimate counterfactual means for a pooled LTMLE.

## Usage

```
pltmle(
  formula,
  outcome,
  treatment,
  covariates,
  baseline,
  ntimes_interval,
  number_traj,
  time,
  time_values,
  identifier,
  obsdata,
  traj,
  total_followup,
  treshold = treshold,
  class_var,
  class_pred
)
```

## Arguments

|            |                                                          |
|------------|----------------------------------------------------------|
| formula    | Specification of the model for the outcome to be fitted. |
| outcome    | Name of the outcome variable.                            |
| treatment  | Time-varying treatment.                                  |
| covariates | Covariates.                                              |
| baseline   | Name of baseline covariates.                             |

|                              |                                                       |
|------------------------------|-------------------------------------------------------|
| <code>ntimes_interval</code> | Length of a time-interval (s).                        |
| <code>number_traj</code>     | An integer to choose the number of trajectory groups. |
| <code>time</code>            | Name of the time variable.                            |
| <code>time_values</code>     | Measuring times.                                      |
| <code>identifier</code>      | Name of the column of the unique identifier.          |
| <code>obsdata</code>         | Observed data in wide format.                         |
| <code>traj</code>            | Matrix of indicators for the trajectory groups.       |
| <code>total_followup</code>  | Number of measuring times per interval.               |
| <code>treshold</code>        | For weight truncation.                                |
| <code>class_var</code>       | Name of the trajectory group variable.                |
| <code>class_pred</code>      | Vector of predicted trajectory groups.                |

### Value

|                                       |                                                                     |
|---------------------------------------|---------------------------------------------------------------------|
| <code>list_pltmle_countermeans</code> | Counterfactual means and influence functions with the pooled ltmle. |
| <code>D</code>                        | Influence functions                                                 |

### Author(s)

Awa Diop, Denis Talbot

### Examples

```
obsdata_long = gendata(n = 2000, format = "long", total_followup = 3, seed = 945)
baseline_var <- c("age", "sex")
covariates <- list(c("hyper2011", "bmi2011"),
c("hyper2012", "bmi2012"), c("hyper2013", "bmi2013"))
treatment_var <- c("statins2011", "statins2012", "statins2013")
time_values <- c(2011, 2012, 2013)
formulaA = as.formula(cbind(statins, 1 - statins) ~ time)
restraj = build_traj(obsdata = obsdata_long, number_traj = 3,
formula = formulaA, identifier = "id")
datapost = restraj$data_post
trajmsm_long <- merge(obsdata_long, datapost, by = "id")
  AggFormula <- as.formula(paste("statins", "~", "time", "+", "class"))
  AggTrajData <- aggregate(AggFormula, data = trajmsm_long, FUN = mean)
  AggTrajData
trajmsm_long[, "traj_group"] <- trajmsm_long[, "class"]
obsdata = reshape(trajmsm_long, direction = "wide", idvar = "id",
v.names = c("statins", "bmi", "hyper"), timevar = "time", sep = "")
formula = as.formula(" y ~ statins2011 + statins2012 + statins2013 +
hyper2011 + bmi2011 + hyper2012 + bmi2012 +
hyper2013 + bmi2013 + age + sex ")
class = factor(predict_traj(identifier = "id", total_followup = 3,
treatment = "statins", time = "time", time_values = time_values,
trajmodel = restraj$traj_model)$post_class);
```



```

traj=t(sapply(1:8,function(x)sapply(1:3,function(i)ifelse(class[x]==i,1,0))))
traj[,1]=1
res_pltmle = pltmle(formula = formula, outcome = "y",treatment = treatment_var,
covariates = covariates, baseline = baseline_var, ntimes_interval = 3, number_traj = 3,
  time = "time",time_values = time_values,identifier = "id",obsdata = obsdata,
traj=traj, threshold = 0.99, class_pred= class, class_var = "class")
res_pltmle$counter_means

```

---

|              |                                                                      |
|--------------|----------------------------------------------------------------------|
| predict_traj | <i>Predict trajectory groups for deterministic treatment regimes</i> |
|--------------|----------------------------------------------------------------------|

---

### Description

Function to predict trajectory groups for deterministic treatment regimes used with gformula and pooled LTMLE.

### Usage

```

predict_traj(
  identifier,
  total_followup,
  treatment,
  time,
  time_values,
  trajmodel
)

```

### Arguments

|                |                                                     |
|----------------|-----------------------------------------------------|
| identifier     | Name of the column of the unique identifier.        |
| total_followup | Number of measuring times.                          |
| treatment      | Name of the time-varying treatment.                 |
| time           | Name of the variable time.                          |
| time_values    | Values of the time variable.                        |
| trajmodel      | Trajectory model built with the observed treatment. |

### Value

A data.frame with the posterior probabilities.

### Author(s)

Awa Diop, Denis Talbot

---

|            |                                                  |
|------------|--------------------------------------------------|
| split_data | <i>Split observed data into multiple subsets</i> |
|------------|--------------------------------------------------|

---

**Description**

Function to split the data into multiple subsets of size `s` each one subset corresponding to one time-interval.

**Usage**

```
split_data(
  obsdata,
  total_followup,
  ntimes_interval,
  time,
  time_values,
  identifier
)
```

**Arguments**

|                              |                                         |
|------------------------------|-----------------------------------------|
| <code>obsdata</code>         | Observed data in wide format.           |
| <code>total_followup</code>  | Total length of follow-up.              |
| <code>ntimes_interval</code> | Number of measuring times per interval. |
| <code>time</code>            | Name of the time variable.              |
| <code>time_values</code>     | Measuring times.                        |
| <code>identifier</code>      | Identifier of individuals.              |

**Value**

|                     |                                      |
|---------------------|--------------------------------------|
| <code>all_df</code> | All subsets, list of time intervals. |
|---------------------|--------------------------------------|

**Author(s)**

Awa Diop Denis Talbot

**Examples**

```
## Not run:
obsdata = gendata(n = 1000, format = "long", total_followup = 8, seed = 945)
years <- 2011:2018
res = split_data(obsdata = obsdata, total_followup = 8,
  ntimes_interval = 6, time = "time", time_values = years, identifier = "id")

## End(Not run)
```

---

|                 |                                                                                             |
|-----------------|---------------------------------------------------------------------------------------------|
| trajhrmsm_gform | <i>History Restricted MSM and Latent Class of Growth Analysis estimated with G-formula.</i> |
|-----------------|---------------------------------------------------------------------------------------------|

---

### Description

Estimate parameters of LCGA-HRMSM using g-formula. and bootstrap to get standard errors.

### Usage

```
trajhrmsm_gform(
  degree_traj = c("linear", "quadratic", "cubic"),
  rep = 50,
  treatment,
  covariates,
  baseline,
  outcome,
  ntimes_interval,
  total_followup,
  time,
  time_values,
  identifier,
  var_cov,
  number_traj = 3,
  family = "poisson",
  obsdata
)
```

### Arguments

|                 |                                                                            |
|-----------------|----------------------------------------------------------------------------|
| degree_traj     | To specify the polynomial degree for modelling the time-varying treatment. |
| rep             | Number of repetition for the bootstrap.                                    |
| treatment       | Name of the time-varying treatment.                                        |
| covariates      | Names of the time-varying covariates (should be a list).                   |
| baseline        | Name of baseline covariates.                                               |
| outcome         | Name of the outcome variable.                                              |
| ntimes_interval | Length of a time-interval (s).                                             |
| total_followup  | Total length of follow-up.                                                 |
| time            | Name of the time variable.                                                 |
| time_values     | Measuring times.                                                           |
| identifier      | Name of the column of the unique identifier.                               |
| var_cov         | Names of the time-varying variables.                                       |
| number_traj     | Number of trajectory groups.                                               |

|         |                                                                                    |
|---------|------------------------------------------------------------------------------------|
| family  | Specification of the error distribution and link function to be used in the model. |
| obsdata | Data in a long format.                                                             |

### Value

A list containing the following components:

**results\_hrmsm\_gform** Matrix of estimates for LCGA-MSM, obtained using the g-formula method.

**result\_coef\_boot** Matrix of estimates obtained with bootstrap.

**restraj** Fitted trajectory model.

**mean\_adh** Matrix of mean adherence per trajectory group.

### Author(s)

Awa Diop Denis Talbot

### Examples

```
obsdata_long = gendata(n = 5000, format = "long", total_followup = 8,
  timedep_outcome = TRUE, seed = 845)
baseline_var <- c("age", "sex")
years <- 2011:2018
variables <- c("hyper", "bmi")
covariates <- lapply(years, function(year) {
  paste0(variables, year)})
treatment_var <- paste0("statins", 2011:2018)
var_cov <- c("statins", "hyper", "bmi")
reshrmsm_gform = trajhrmsm_gform(degree_traj = "linear", rep=50 ,
  treatment = treatment_var, covariates = covariates, baseline = baseline_var,
  outcome = "y", var_cov = var_cov, ntimes_interval = 6, total_followup = 8,
  time = "time", time_values = years, identifier = "id",
  number_traj = 3, family = "poisson", obsdata = obsdata_long)
reshrmsm_gform$results_hrmsm_gform
```

---

trajhrmsm\_ipw

---

*History Restricted MSM and Latent Class of Growth Analysis estimated with IPW.*


---

### Description

Estimate parameters of LCGA-HRMSM using IPW.

**Usage**

```
trajhrmsm_ipw(
  degree_traj = c("linear", "quadratic", "cubic"),
  numerator = c("stabilized", "unstabilized"),
  identifier,
  baseline,
  covariates,
  treatment,
  outcome,
  var_cov,
  include_censor = FALSE,
  ntimes_interval,
  total_followup,
  time,
  time_values,
  family = "poisson",
  censor = censor,
  number_traj,
  obsdata,
  weights = NULL,
  threshold = 0.999
)
```

**Arguments**

|                 |                                                                                    |
|-----------------|------------------------------------------------------------------------------------|
| degree_traj     | To specify the polynomial degree for modelling the time-varying treatment.         |
| numerator       | To choose between stabilized and unstabilized weights.                             |
| identifier      | Name of the column of the unique identifier.                                       |
| baseline        | Names of the baseline covariates.                                                  |
| covariates      | Names of the time-varying covariates (should be a list).                           |
| treatment       | Name of the time-varying treatment.                                                |
| outcome         | Name of the outcome variable.                                                      |
| var_cov         | Names of the time-varying variables.                                               |
| include_censor  | Logical, if TRUE, includes censoring.                                              |
| ntimes_interval | Length of a time-interval (s).                                                     |
| total_followup  | Total length of follow-up.                                                         |
| time            | Name of the time variable.                                                         |
| time_values     | Values of the time variable.                                                       |
| family          | specification of the error distribution and link function to be used in the model. |
| censor          | Name of the censoring variable.                                                    |
| number_traj     | Number of trajectory groups.                                                       |
| obsdata         | Data in a long format.                                                             |

|          |                                                                                   |
|----------|-----------------------------------------------------------------------------------|
| weights  | A vector of estimated weights. If NULL, the weights are computed by the function. |
| treshold | For weight truncation.                                                            |

**Value**

Provides a matrix of estimates for LCGA-HRMSM, obtained using IPW.

**Author(s)**

Awa Diop, Denis Talbot

**Examples**

```
obsdata_long = gendata(n = 5000, format = "long", total_followup = 8,
  timedep_outcome = TRUE, seed = 845)
baseline_var <- c("age", "sex")
years <- 2011:2018
variables <- c("hyper", "bmi")
covariates <- lapply(years, function(year) {
  paste0(variables, year)})
treatment_var <- paste0("statins", 2011:2018)
var_cov <- c("statins", "hyper", "bmi", "y")
reshrmsm_ipw <- trajhrmsm_ipw(degree_traj = "linear", numerator = "stabilized",
  identifier = "id", baseline = baseline_var,
  covariates = covariates, treatment = treatment_var,
  outcome = "y", var_cov= var_cov, include_censor = FALSE,
  ntimes_interval = 6, total_followup = 8, time = "time", time_values = 2011:2018,
  family = "poisson", number_traj = 3, obsdata = obsdata_long, treshold = 1)
reshrmsm_ipw$res_trajhrmsm_ipw
```

---

|                  |                                                                                                  |
|------------------|--------------------------------------------------------------------------------------------------|
| trajhrmsm_pltmle | <i>History Restricted MSM and Latent Class of Growth Analysis estimated with a Pooled LTMLE.</i> |
|------------------|--------------------------------------------------------------------------------------------------|

---

**Description**

Estimate parameters of LCGA-HRMSM using a Pooled LTMLE.

**Usage**

```
trajhrmsm_pltmle(
  degree_traj = c("linear", "quadratic", "cubic"),
  treatment,
  covariates,
  baseline,
  outcome,
  ntimes_interval,
```

```

    total_followup,
    time,
    time_values,
    identifier,
    var_cov,
    number_traj = 3,
    family = "poisson",
    obsdata,
    treshold = 0.99
  )

```

### Arguments

|                 |                                                                                    |
|-----------------|------------------------------------------------------------------------------------|
| degree_traj     | To specify the polynomial degree for modelling the time-varying treatment.         |
| treatment       | Name of time-varying treatment.                                                    |
| covariates      | Names of time-varying covariates (should be a list).                               |
| baseline        | Names of baseline covariates.                                                      |
| outcome         | Name of the outcome variable.                                                      |
| ntimes_interval | Length of a time-interval (s).                                                     |
| total_followup  | Total length of follow-up.                                                         |
| time            | Name of the time variable.                                                         |
| time_values     | Measuring times.                                                                   |
| identifier      | Name of the column for unique identifiant.                                         |
| var_cov         | Names of the time-varying variables.                                               |
| number_traj     | Number of trajectory groups.                                                       |
| family          | Specification of the error distribution and link function to be used in the model. |
| obsdata         | Data in a long format.                                                             |
| treshold        | For weight truncation.                                                             |

### Value

A list containing the following components:

**results\_hrmsm\_pltmle** Matrix of estimates for LCGA-HRMSM, obtained using the pooled ltmle method.

**restraj** Fitted trajectory model.

**mean\_adh** Matrix of the mean adherence per trajectory group.

### Author(s)

Awa Diop Denis Talbot

### Examples

```
obsdata_long = gendata(n = 5000, format = "long",
total_followup = 8, timedep_outcome = TRUE, seed = 845)
baseline_var <- c("age", "sex")
years <- 2011:2018
variables <- c("hyper", "bmi")
covariates <- lapply(years, function(year) {
  paste0(variables, year)})
treatment_var <- paste0("statins", 2011:2018)
var_cov <- c("statins", "hyper", "bmi", "y")
respltmle = trajhrmsm_pltmle(degree_traj = "linear", treatment = treatment_var,
covariates = covariates, baseline = baseline_var,
outcome = paste0("y", 2016:2018), var_cov = var_cov, ntimes_interval = 6,
total_followup = 8, time = "time", time_values = years, identifier = "id",
number_traj = 3, family = "poisson", obsdata = obsdata_long, treshold = 1)
respltmle$results_hrmsm_pltmle
```

trajmsm\_gform

*Parametric g-formula*

### Description

Estimate parameters of LCGA-MSM using g-formula and bootstrap to get standard errors.

### Usage

```
trajmsm_gform(
  formula = formula,
  rep = 50,
  identifier,
  baseline,
  covariates,
  treatment,
  outcome,
  total_followup,
  time = time,
  time_values,
  var_cov,
  trajmodel,
  ref,
  obsdata
)
```

### Arguments

|         |                                                          |
|---------|----------------------------------------------------------|
| formula | Specification of the model for the outcome to be fitted. |
| rep     | Number of repetitions for the bootstrap.                 |



|                |                                                     |
|----------------|-----------------------------------------------------|
| identifier     | Name of the column of the unique identifier.        |
| baseline       | Vector of names of the baseline covariates.         |
| covariates     | List of names of the time-varying covariates.       |
| treatment      | Vector of names of the time-varying treatment.      |
| outcome        | Name of the outcome of interest.                    |
| total_followup | Total length of follow-up.                          |
| time           | Name of the time variable.                          |
| time_values    | Measuring times.                                    |
| var_cov        | Names of the time-varying covariates.               |
| trajmodel      | Trajectory model built with the observed treatment. |
| ref            | The reference trajectory group.                     |
| obsdata        | Observed data in wide format.                       |

### Value

Provides a matrix of estimates for LCGA-MSM, obtained using the g-formula method.

### Author(s)

Awa Diop Denis Talbot

### Examples

```
obsdata_long = gendata(n = 1000, format = "long", total_followup = 6, seed = 845)
years <- 2011:2016
baseline_var <- c("age", "sex")
variables <- c("hyper", "bmi")
var_cov <- c("statins", "hyper", "bmi")
covariates <- lapply(years, function(year) {
  paste0(variables, year)})
treatment_var <- paste0("statins", 2011:2016)
formula_treatment = as.formula(cbind(statins, 1 - statins) ~ time)
restraj = build_traj(obsdata = obsdata_long, number_traj = 3,
  formula = formula_treatment, identifier = "id")
datapost = restraj$data_post
trajmsm_long <- merge(obsdata_long, datapost, by = "id")
  AggFormula <- as.formula(paste("statins", "~", "time", "+", "class"))
  AggTrajData <- aggregate(AggFormula, data = trajmsm_long, FUN = mean)
  AggTrajData
obsdata = reshape(data = trajmsm_long, direction = "wide", idvar = "id",
  v.names = c("statins", "bmi", "hyper"), timevar = "time", sep = "")
formula = paste0("y ~", paste0(treatment_var, collapse = "+"), "+",
  paste0(unlist(covariates), collapse = "+"), "+",
  paste0(baseline_var, collapse = "+"))
resmsm_gform <- trajmsm_gform(formula = formula, identifier = "id", rep = 5,
  baseline = baseline_var, covariates = covariates, var_cov = var_cov,
  treatment = treatment_var, outcome = "y", total_followup = 6, time = "time",
  time_values = years, trajmodel = restraj$traj_model, ref = "1", obsdata = obsdata )
```

resmsm\_gform

---

|             |                                                                                         |
|-------------|-----------------------------------------------------------------------------------------|
| trajmsm_ipw | <i>Marginal Structural Model and Latent Class of Growth Analysis estimated with IPW</i> |
|-------------|-----------------------------------------------------------------------------------------|

---

## Description

Estimate parameters of LCGA-MSM using IPW.

## Usage

```
trajmsm_ipw(
  formula1,
  formula2,
  family,
  identifier,
  treatment,
  covariates,
  baseline,
  obsdata,
  numerator = "stabilized",
  include_censor = FALSE,
  censor,
  weights = NULL,
  threshold = 0.99
)
```

## Arguments

|                |                                                                                                  |
|----------------|--------------------------------------------------------------------------------------------------|
| formula1       | Specification of the model for the outcome to be fitted for a binomial or gaussian distribution. |
| formula2       | Specification of the model for the outcome to be fitted for a survival outcome.                  |
| family         | Specification of the error distribution and link function to be used in the model.               |
| identifier     | Name of the column of the unique identifier.                                                     |
| treatment      | Time-varying treatment.                                                                          |
| covariates     | Names of the time-varying covariates (should be a list).                                         |
| baseline       | Name of the baseline covariates.                                                                 |
| obsdata        | Dataset to be used in the analysis.                                                              |
| numerator      | Type of weighting ("stabilized" or "unstabilized").                                              |
| include_censor | Logical, if TRUE, includes censoring.                                                            |
| censor         | Name of the censoring variable.                                                                  |
| weights        | A vector of estimated weights. If NULL, the weights are computed by the function IPW.            |
| threshold      | For weight truncation.                                                                           |

**Value**

Provides a matrix of estimates for LCGA-MSM, obtained using IPW.

Provides a matrix of estimates for LCGA-MSM, obtained using IPW.

**Examples**

```
obsdata_long = gendata(n = 1000, format = "long", total_followup = 6, seed = 845)
years <- 2011:2016
baseline_var <- c("age", "sex")
variables <- c("hyper", "bmi")
covariates <- lapply(years, function(year) {
  paste0(variables, year)})
treatment_var <- paste0("statins", 2011:2016)
formula_treatment = as.formula(cbind(statins, 1 - statins) ~ time)
restraj = build_traj(obsdata = obsdata_long, number_traj = 3,
  formula = formula_treatment, identifier = "id")
datapost = restraj$data_post
trajmsm_long <- merge(obsdata_long, datapost, by = "id")
  AggFormula <- as.formula(paste("statins", "~", "time", "+", "class"))
  AggTrajData <- aggregate(AggFormula, data = trajmsm_long, FUN = mean)
  AggTrajData
trajmsm_long$ipw_group <- releval(trajmsm_long$class, ref = "1")
obsdata = reshape(data = trajmsm_long, direction = "wide", idvar = "id",
  v.names = c("statins", "bmi", "hyper"), timevar = "time", sep = "")
formula = paste0("y ~", paste0(treatment_var, collapse = "+"), "+",
  paste0(unlist(covariates), collapse = "+"), "+",
  paste0(baseline_var, collapse = "+"))

resmsm_ipw = trajmsm_ipw(formula1 = as.formula("y ~ ipw_group"),
  identifier = "id", baseline = baseline_var, covariates = covariates,
  treatment = treatment_var, family = "binomial",
  obsdata = obsdata, numerator = "stabilized", include_censor = FALSE, threshold = 0.99)
resmsm_ipw
```

---

|                |                     |
|----------------|---------------------|
| trajmsm_pltmle | <i>Pooled LTMLE</i> |
|----------------|---------------------|

---

**Description**

Estimate parameters of LCGA-MSM using pooled LTMLE with influence functions to estimate standard errors.

**Usage**

```
trajmsm_pltmle(
  formula = formula,
  identifier,
  baseline,
```

```

    covariates,
    treatment,
    outcome,
    number_traj,
    total_followup,
    time,
    time_values,
    trajmodel,
    ref,
    treshold = 0.99,
    obsdata,
    class_var
  )

```

### Arguments

|                |                                                          |
|----------------|----------------------------------------------------------|
| formula        | Specification of the model for the outcome to be fitted. |
| identifier     | Name of the column for unique identifiant.               |
| baseline       | Names of the baseline covariates.                        |
| covariates     | Names of the time-varying covariates (should be a list). |
| treatment      | Name of the time-varying treatment.                      |
| outcome        | Name of the outcome variable.                            |
| number_traj    | An integer to choose the number of trajectory groups.    |
| total_followup | Total length of follow-up.                               |
| time           | Name of the time variable.                               |
| time_values    | Measuring times.                                         |
| trajmodel      | Trajectory model built with the observed treatment.      |
| ref            | The reference group.                                     |
| treshold       | For weight truncation.                                   |
| obsdata        | Observed data in wide format.                            |
| class_var      | Name of the trajectory group variable.                   |

### Value

Provides a matrix of estimates for LCGA-MSM, obtained using the pooled ltmle method.

```
results_msm_pooledltmle
```

Estimates of a LCGA-MSM with pooled LTMLE.

### Author(s)

Awa Diop, Denis Talbot

**Examples**

```

obsdata_long = gendata(n = 1000, format = "long", total_followup = 6, seed = 845)
years <- 2011:2016
baseline_var <- c("age", "sex")
variables <- c("hyper", "bmi")
covariates <- lapply(years, function(year) {
  paste0(variables, year)})
treatment_var <- paste0("statins", 2011:2016)
formula_treatment = as.formula(cbind(statins, 1 - statins) ~ time)
restraj = build_traj(obsdata = obsdata_long, number_traj = 3,
  formula = formula_treatment, identifier = "id")
datapost = restraj$data_post
trajmsm_long <- merge(obsdata_long, datapost, by = "id")
  AggFormula <- as.formula(paste("statins", "~", "time", "+", "class"))
  AggTrajData <- aggregate(AggFormula, data = trajmsm_long, FUN = mean)
trajmsm_wide = reshape(data = trajmsm_long, direction = "wide", idvar = "id",
  v.names = c("statins", "bmi", "hyper"), timevar = "time", sep = "")
formula = paste0("y ~", paste0(treatment_var, collapse = "+"), "+",
  paste0(unlist(covariates), collapse = "+"), "+",
  paste0(baseline_var, collapse = "+"))
resmsm_pltmle <- trajmsm_pltmle(formula = formula, identifier = "id",
  baseline = baseline_var,
  covariates = covariates, treatment = treatment_var,
  outcome = "y", time = "time", time_values = years,
  number_traj = 3, total_followup = 6,
  trajmodel = restraj$traj_model, ref = "1", obsdata = trajmsm_wide,
  treshold = 1, class_var = "class")
resmsm_pltmle

```

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