

# Package: packDAMipd (via r-universe)

August 19, 2024

**Type** Package

**Title** Decision Analysis Modelling Package with Parameters Estimation  
Ability from Individual Patient Level Data

**Version** 1.1.0

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**Description** A collection of functions to construct Markov model for model-based cost-effectiveness analysis. This includes creating Markov model (both time homogenous and time dependent models), decision analysis, sensitivity analysis (deterministic and probabilistic). The package allows estimation of parameters for the Markov model from a given individual patient level data, provided the data file follows some standard data entry rules.

**License** GPL-3

**Depends** R (>= 4.3.0)

**Imports** readxl, stringr, data.table, reshape2, rlang, stats, lme4, survminer, SurvRegCensCov, survival, MASS, systemfit, IPDFFileCheck, valueEQ5D, car, ggplot2, grDevices, lmtest, broom, effects, gvlma, methods, relaimpo, dplyr, tidyr, hash, haven, flextable, ggpubr, labelled, purrr, ISLR

**Suggests** knitr, rmarkdown, covr, testthat

**VignetteBuilder** knitr

**Encoding** UTF-8

**LazyData** true

**RoxygenNote** 7.3.1

**URL** <https://github.com/sheejamk/packDAMipd>

**BugReports** <https://github.com/sheejamk/packDAMipd/issues>

**NeedsCompilation** no

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**Date/Publication** 2024-03-24 21:10:02 UTC

**Additional\_repositories** <https://cranhaven.r-universe.dev>

**Repository** <https://cranhaven.r-universe.dev>

**RemoteUrl** <https://github.com/cranhaven/cranhaven.r-universe.dev>

**RemoteRef** package/packDAMipd

**RemoteSha** 3e3dfebe8fcf677e6ef2a3f7a4ee24c8436cbc83

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---

 add\_entries\_sameuse\_timepoint

*Function to get sum of entries of resource per individual at diff timepoints if same category has listed multiple time for same id of participant , this method comes in handy to get the sum*

---

**Description**

Function to get sum of entries of resource per individual at diff timepoints if same category has listed multiple time for same id of participant , this method comes in handy to get the sum

**Usage**

```
add_entries_sameuse_timepoint(
    use_data,
    timepointcol,
    timepointval,
    idcolumn,
    result_col
)
```

**Arguments**

use_data	the data where the observations are held
timepointcol	columnname in the data where the timepoints are noted
timepointval	which time point is considered now at which the descriptive analysis is done
idcolumn	id for each participant
result_col	name of the column where the sum of entries to be saved

**Value**

the data with added sum of resource use

**Examples**

```
eg_data <- as.data.frame(list(no = c(1, 2, 3, 4),
  mark_at_1 = c(12, 7, 23, 45), gender = c("M", "F", "M", "F"),
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2"),
  time = c(1,1,2,2), id = c(1, 1, 1, 2)))
add_entries_sameuse_timepoint(eg_data, "time", 1, "id",
("mark_at_2"))
```

---

adl_scoring.df	<i>adl_scoring table</i>
----------------	--------------------------

---

**Description**

adl\_scoring table

**Usage**

adl\_scoring.df

**Format**

A 41 by 3 dataframe

**Source**

created on Jan 15, 2020

---

assign_parameters	<i>Function to assign the values of nested parameters from the parameter list</i>
-------------------	---

---

**Description**

Function to assign the values of nested parameters from the parameter list

**Usage**

```
assign_parameters(param_list)
```

**Arguments**

param_list	list of parameters, can be nested, or can be used the list returned from define_parameters()
------------	--

**Details**

The parameter list should be a list of parameters in the form name value pairs. If the name value pairs is given as a string it throws error as in `assign_parameters(c("cost_A = 100", "a = 10"))` even if you use `assign_parameters(define_parameters(c("cost_A = 100","a = 10")))` but this will be ok if you use the below forms `assign_list2 <- c(a = 10, cost_A = "a + 100", cost_B = 10)` `assign_parameters(assign_list2)` OR `param_list <- define_parameters(a = 10, cost_A = "a + 100", cost_B = 10)` `assign_list <- assign_parameters(param_list)` Also for nested parameters, remember to give the parameters in order so that at run time, the parameters can be evaluated for example, `assign_list = define_parameters(cost_A="a+100", a=10)` `assign_parameters(assign_list)` will throw an error, while `assign_list = define_parameters( a = 10, cost_A = "a + 100")` `assign_parameters(assign_list)` will successfully assign parameters as the parameters 'a' is visible before the calculation of 'cost\_A' Another thing to note is that while using `define_parameters`, just enumerate them, no need to create as a list by using `c()` or `list` function

**Value**

list of assigned parameters

**Examples**

```
param_list <- define_parameters(
  cost_direct_med_A = 1701, cost_comm_care_A = 1055,
  cost_direct_med_B = 1774, cost_comm_care_B = 1278,
  cost_direct_med_C = 6948,
  cost_comm_care_C = 2059, cost_zido = 2456, cost_health_A =
  "cost_direct_med_A + cost_comm_care_A",
  cost_health_B = "cost_direct_med_B + cost_comm_care_B",
  cost_health_C = "cost_direct_med_C + cost_comm_care_C",
  cost_drug = "cost_zido"
)
assign_parameters(param_list)
```

---

 blank.df

---

*Parameter table created*


---

**Description**

Parameter table created

**Usage**

blank.df

**Format**

A 2 column 1 observation

**Source**

created on September 5, 2020

---

calculate\_icer\_nmb      *Estimation of ICER and NMB*

---

**Description**

Estimation of ICER and NMB

**Usage**

```
calculate_icer_nmb(list_markov, threshold, comparator = NULL)
```

**Arguments**

list_markov	list of Markov model objects with their Markov trace, cost matrix and utility matrix
threshold	threshold value of WTP
comparator	the strategy to be compared with

**Value**

ICER and NMB for all the strategies compared to comparator

**Examples**

```
well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 100, utility = 1)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.6, 0.2, 0.2, 0.6, 0.4, 1),
colnames(tmat))
health_states <- combine_state(well, disabled, dead)
this.strategy <- strategy(tm, health_states, "control")
this_markov <- markov_model(this.strategy, 24, c(1000, 0, 0), c(0,0))
well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 10, utility = 0.5)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.4, 0.4, 0.2, 0.6, 0.4, 1),
colnames(tmat))
health_states <- combine_state(well, disabled, dead)
this.strategy <- strategy(tm, health_states, "intervention")
sec_markov <- markov_model(this.strategy, 24, c(1000, 0, 0), c(0,0))
list_markov <- combine_markov(this_markov, sec_markov)
calculate_icer_nmb(list_markov, 20000, comparator = "control")
```



---

 checks\_markov\_pick\_method

*Checks the input to run the Markov cycles and picks correct method*


---

**Description**

Checks the input to run the Markov cycles and picks correct method

**Usage**

```
checks_markov_pick_method(
  current_strategy,
  initial_state,
  discount,
  method,
  half_cycle_correction,
  startup_cost,
  startup_util,
  state_cost_only_prevalent,
  state_util_only_prevalent
)
```

**Arguments**

current_strategy	strategy object
initial_state	value of states initially
discount	rate of discount for costs and qalys
method	what type of half cycle correction needed
half_cycle_correction	boolean to indicate half cycle correction
startup_cost	cost of states initially
startup_util	utility of states initially if any
state_cost_only_prevalent	boolean parameter to indicate if the costs for state occupancy is only for those in the state excluding those that transitioned new. This is relevant when the transition cost is provided for eg. in a state with dialysis the cost of previous dialysis is different from the newly dialysis cases. Then the state_cost_only_prevalent should be TRUE
state_util_only_prevalent	boolean parameter to indicate if the utilities for state occupancy is only for those in the state excluding those that transitioned new.

**Value**

changed method name

**Examples**

```
tmat <- rbind(c(1, 2), c(3, 4))
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")
tm <- populate_transition_matrix(2, tmat, c(0.5, 0.5, 0, 1))
a <- health_state("Healthy", 1, 1, 0, FALSE)
b <- health_state("Dead", 1, 0, 0, TRUE)
health_states <- combine_state(a, b)
this.strategy <- strategy(tm, health_states, "intervention")
checks_markov_pick_method(this.strategy, c(1, 0), c(0, 0),
"half cycle correction", TRUE, NULL, NULL)
```

---

checks\_plot\_dsa

*Function to do some checks before plotting sensitivity analysis results*


---

**Description**

Function to do some checks before plotting sensitivity analysis results

**Usage**

```
checks_plot_dsa(
  result_dsa_control,
  plotfor,
  type,
  result_dsa_treat,
  threshold,
  comparator
)
```

**Arguments**

result_dsa_control	result from deterministic sensitivity analysis for first or control model
plotfor	the variable to plotfor e.g. cost, utility NMB etc
type	type of analysis, range or difference
result_dsa_treat	result from deterministic sensitivity analysis for the comparative Markov model
threshold	threshold value of WTP
comparator	the strategy to be compared with

**Value**

the plot variable

---

`check_equal_columncontents_NAomitted`

*Function to check the equality of column contents between two data sets with omitting NA*

---

## Description

Function to check the equality of column contents between two data sets with omitting NA

## Usage

```
check_equal_columncontents_NAomitted(  
  data1,  
  data2,  
  samecol,  
  column_data1,  
  column_data2  
)
```

## Arguments

<code>data1</code>	first data set
<code>data2</code>	second data set
<code>samecol</code>	a unique col in both datasets, like tno or id
<code>column_data1</code>	column name in data 1 to be compared
<code>column_data2</code>	column name in data 2 to be compared

## Value

0 if they are equal else error message

## Examples

```
eg_data <- as.data.frame(list(no = c(1, 2, 3, 4),  
  mark_at_1 = c(12, 7, 23, 45), gender = c("M", "F", "M", "F"),  
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2"),  
  time = c(1,1,2,2), id = c(1, 1, 1, 2)))  
eg_data2 <- as.data.frame(list(no = c(1, 2, 3, 4),  
  mark_at_1 = c(12, 27, 23, 45), gender = c("M", "F", "M", "F"),  
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2"),  
  time = c(1,1,2,2), id = c(1, 1, 1, 2)))  
check_equal_columncontents_NAomitted(eg_data, eg_data2, "no",  
  "mark_at_1", "mark_at_2")
```

---

check\_equal\_sumcolumncontents\_NAomitted

*Function to check the sum of column contents between two data sets with omitting NA*

---

## Description

Function to check the sum of column contents between two data sets with omitting NA

## Usage

```
check_equal_sumcolumncontents_NAomitted(  
  data1,  
  data2,  
  samecol,  
  column_data1,  
  column_data2  
)
```

## Arguments

data1	first data set
data2	second data set
samecol	a unique col in both datasets, like tno or id
column_data1	column name in data 1 to be compared
column_data2	column name in data 2 to be compared

## Value

0 if they are equal else error message

## Examples

```
eg_data <- as.data.frame(list(no = c(1, 2, 3, 4),  
  mark_at_1 = c(12, 7, 23, 45), gender = c("M", "F", "M", "F"),  
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2"),  
  time = c(1,1,2,2), id = c(1, 1, 1, 2)))  
eg_data2 <- as.data.frame(list(no = c(1, 2, 3, 4),  
  mark_at_1 = c(12, 27, 23, 45), gender = c("M", "F", "M", "F"),  
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2"),  
  time = c(1,1,2,2), id = c(1, 1, 1, 2)))  
check_equal_sumcolumncontents_NAomitted(eg_data, eg_data2, "no",  
  "mark_at_1", "mark_at_2")
```

---

check_link_glm	<i>Function to find the keyword for family of distribution in glm</i>
----------------	---

---

**Description**

Function to find the keyword for family of distribution in glm

**Usage**

```
check_link_glm(family, link)
```

**Arguments**

family	family of distribution
link	function to be used

**Details**

Check and get the link function for the method glm

**Value**

the link if they can be accepted else error

**Examples**

```
check_link_glm("gaussian", "identity")
```

---

check_list_markov_models	<i>check the list of Markov models</i>
--------------------------	--

---

**Description**

check the list of Markov models

**Usage**

```
check_list_markov_models(list_markov)
```

**Arguments**

list_markov	list of Markov model objects with their Markov trace, cost matrix and utility matrix
-------------	--

**Value**

0 if success else error

**Examples**

```

well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 100, utility = 1)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.6, 0.2, 0.2, 0.6, 0.4, 1),
  colnames(tmat))
health_states <- combine_state(well, disabled, dead)
this.strategy <- strategy(tm, health_states, "example")
this_markov <- markov_model(this.strategy, 24, c(1000, 0, 0), c(0,0))
well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 10, utility = 0.5)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.4, 0.4, 0.2, 0.6, 0.4, 1),
  colnames(tmat))
health_states <- combine_state(well, disabled, dead)
this.strategy <- strategy(tm, health_states, "example_two")
sec_markov <- markov_model(this.strategy, 24, c(1000, 0, 0), c(0, 0))
list_markov <- combine_markov(this_markov, sec_markov)
check_list_markov_models(list_markov)

```

---

check\_null\_na

*Function to check the variable null or NA*

---

**Description**

Function to check the variable null or NA

**Usage**

```
check_null_na(variable)
```

**Arguments**

variable            name of variable or list of variable to check

**Value**

-1 or -2 as error, else return 0 as success

**Examples**

```
var = c("a")
check_null_na(var)
```

---

check_trans_prob	<i>Check the transition probabilities for numeric values and unity row sum</i>
------------------	--

---

**Description**

Check the transition probabilities for numeric values and unity row sum

**Usage**

```
check_trans_prob(trans_mat)
```

**Arguments**

trans\_mat      transition matrix

**Details**

checking for rowsum - checks for the class of transition matrix, value of rowsum (to be 1) and numeric values

**Value**

0 if they add to 1 else error

**Examples**

```
tmat <- rbind(c(1, 2), c(3, 4))
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")
tm <- populate_transition_matrix(2, tmat, list_prob = c(0.5, 0.5, 0, 1))
check_trans_prob(tm)
```

check\_treatment\_arm     *Function to return treatment arm*

---

**Description**

Function to return treatment arm

**Usage**

```
check_treatment_arm(arm)
```

**Arguments**

arm                    the arm of the trial

**Value**

0, if success -1, if failure

**Examples**

```
check_treatment_arm("control")
```

---

check\_values\_states     *Check if the values of health states are provided*

---

**Description**

Check if the values of health states are provided

**Usage**

```
check_values_states(health_states)
```

**Arguments**

health\_states     list of health\_state objects

**Details**

This is to check if the values are numeric during the run time, else to throw an error

**Value**

true or false



**Examples**

```

well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 100, utility = 1)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.6, 0.2, 0.2, 0.6, 0.4, 1))
health_states <- combine_state(well, disabled, dead)
check_values_states(health_states)

```

---

combine\_markov

*Join Markov model objects*


---

**Description**

Join Markov model objects

**Usage**

```
combine_markov(markov1, ...)
```

**Arguments**

markov1	object 1 of class markov_model
...	any additional objects

**Details**

Combining Markov models for easiness of comparison

**Value**

joined objects of type markov\_model

**Examples**

```

well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 100, utility = 1)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.6, 0.2, 0.2, 0.6, 0.4, 1))
health_states <- combine_state(well, disabled, dead)
this_strategy <- strategy(tm, health_states, "example")
this_markov <- markov_model(this_strategy, 24, c(1000, 0, 0))
well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 10, utility = 0.5)
dead <- health_state("dead", cost = 0, utility = 0)

```

```
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.4, 0.4, 0.2, 0.6, 0.4, 1))
health_states <- combine_state(well, disabled, dead)
this_strategy <- strategy(tm, health_states, "example")
sec_markov <- markov_model(this_strategy, 24, c(1000, 0, 0))
list_markov <- combine_markov(this_markov, sec_markov)
```

---

combine_state	<i>Join health states</i>
---------------	---------------------------

---

### Description

Join health states

### Usage

```
combine_state(...)
```

### Arguments

... any additional objects

### Details

checking each state is a health state and join them

### Value

joined health states

### Examples

```
a <- health_state("IT", 100, 0.4, 0, FALSE)
b <- health_state("PT", 100, 0.4, 0, FALSE)
combine_state(a, b)
```

---

`convert_freq_diff_basis`*Convert frequency medication to given basis*

---

**Description**

Convert frequency medication to given basis

**Usage**

```
convert_freq_diff_basis(freq_given, basis = "day")
```

**Arguments**

freq_given	given frequency
basis	given basis, default is daily

**Value**

converted frequency

**Examples**

```
convert_freq_diff_basis("once daily")
convert_freq_diff_basis("bd", "week")
convert_freq_diff_basis("Every 4 days", "day")
```

---

`convert_to_given_timeperiod`*Convert period to given basis*

---

**Description**

Convert period to given basis

**Usage**

```
convert_to_given_timeperiod(given_time, basis_time = "day")
```

**Arguments**

given_time	given time
basis_time	given basis, default is "day"

**Value**

converted unit

**Examples**

```
convert_to_given_timeperiod("4 weeks")
convert_to_given_timeperiod("a month")
convert_to_given_timeperiod("1 week")
```

---

convert\_volume\_basis    *Convert volume to given basis*

---

**Description**

Convert volume to given basis

**Usage**

```
convert_volume_basis(given_unit, basis = "ml")
```

**Arguments**

given_unit	given unit
basis	given basis, default is "ml"

**Value**

converted unit

**Examples**

```
convert_volume_basis("ml", "liter")
```

---

convert\_weight\_diff\_basis  
*Convert unit strength to given basis*

---

**Description**

Convert unit strength to given basis

**Usage**

```
convert_weight_diff_basis(given_unit, basis = "mg")
```

**Arguments**

given\_unit      given unit  
basis            given basis, default is "mg"

**Value**

converted unit

**Examples**

```
convert_weight_diff_basis("mg")  
convert_weight_diff_basis("kilogram", "micro gram")
```

---

convert\_wtpertimediff\_basis  
*Convert weight per time to given basis*

---

**Description**

Convert weight per time to given basis

**Usage**

```
convert_wtpertimediff_basis(given_unit, basis = "mcg/hour")
```

**Arguments**

given\_unit      given unit  
basis            given basis, default is "mg"

**Value**

converted unit

**Examples**

```
convert_wtpertimediff_basis("mg/day")  
convert_wtpertimediff_basis("mcg/day")  
convert_wtpertimediff_basis("mg/hour")
```

convert\_wtpervoldiff\_basis

*Convert wt per unit volume to given basis*

---

### **Description**

Convert wt per unit volume to given basis

### **Usage**

```
convert_wtpervoldiff_basis(given_unit, basis = "mg/ml")
```

### **Arguments**

given_unit	given unit
basis	given basis, default is "mg/ml"

### **Value**

converted unit

### **Examples**

```
convert_wtpervoldiff_basis("g/ml")
```

---

costing\_AandE\_admission

*Function to estimate the cost of inpatient admission but taken from GP records where code or description known*

---

### **Description**

Function to estimate the cost of inpatient admission but taken from GP records where code or description known

### **Usage**

```
costing_AandE_admission(  
  ind_part_data,  
  code_ae,  
  descrip_ae,  
  number_use_ae,  
  type_admit_ae,  
  unit_cost_data,  
  code_col,
```

```

    type_admit_col,
    description_col,
    unit_cost_col,
    cost_calculated_in = "attendance",
    sheet = NULL
  )

```

### Arguments

ind_part_data	IPD
code_ae	column name of code (for inpatient admission)
descrip_ae	column name of description for inpatient admission
number_use_ae	the number of days spent in each admission if that is a criteria to be included. Otherwise each admission will be costed
type_admit_ae	term indicating admission and type of attendance
unit_cost_data	unit cost data file with code/descriptions and unit costs are listed for inpatient admission
code_col	code column name in unit cost data
type_admit_col	colname that describes type of the attendance and
description_col	column name of description of inpatient admission in the unit cost data
unit_cost_col	column name of unit cost in unit_cost_data
cost_calculated_in	name of unit where the cost is calculated assumed to be per admission
sheet	sheet where the unit costs are listed in the unit costs data file

### Value

the calculated cost of inpatient admission long with original data

### Examples

```

costs_file <- system.file("extdata",
  "National_schedule_of_NHS_costs_2019_AandE.csv", package = "packDAMipd")
datafile <- system.file("extdata", "resource_use_ae_ip.csv",
  package = "packDAMipd")
ind_part_data <- packDAMipd::load_trial_data(datafile)
unit_cost_data <- packDAMipd::load_trial_data(costs_file)
result <- costing_AandE_admission(ind_part_data = ind_part_data,
  code_ae = "code", descrip_ae = NULL, number_use_ae = "number_use",
  type_admit_ae = "type_admit", unit_cost_data = unit_cost_data,
  code_col = "Currency_Code", type_admit_col = "Service_Code",
  description_col = NULL, unit_cost_col = "National_Average_Unit_Cost",
  cost_calculated_in = "attendance")

```

---

costing\_inpatient\_daycase\_admission

*Function to estimate the cost of inpatient admission but taken from GP records where HRG code or description known*

---

### Description

Function to estimate the cost of inpatient admission but taken from GP records where HRG code or description known

### Usage

```
costing_inpatient_daycase_admission(
  ind_part_data,
  hrg_code_ip_admi,
  descrip_ip_admi,
  number_use_ip_admi,
  elective_col,
  unit_cost_data,
  hrg_code_col,
  description_col,
  unit_cost_col,
  cost_calculated_in = "admission"
)
```

### Arguments

ind_part_data	IPD
hrg_code_ip_admi	column name of hrg code (for inpatient admission)
descrip_ip_admi	column name of description for inpatient admission
number_use_ip_admi	the number of days spent in each admission if that is a criteria to be included. Otherwise each admission will be costed
elective_col	colname to say whether it is an elective admission or non elective admission
unit_cost_data	unit cost data file with hrg code/descriptions and unit costs are listed for inpatient admission
hrg_code_col	hrg code column name in unit cost data
description_col	column name of description of inpatient admission in the unit cost data
unit_cost_col	column name of unit cost in unit_cost_data
cost_calculated_in	name of unit where the cost is calculated assumed to be per admission



**Value**

the calculated cost of inpatient admission long with original data

**Examples**

```
costs_file <- system.file("extdata",
  "National_schedule_of_NHS_costs_2019.csv",
  package = "packDAMipd")
datafile <- system.file("extdata", "resource_use_hc_ip.csv",
  package = "packDAMipd")
ind_part_data <- packDAMipd::load_trial_data(datafile)
unit_cost_data <- packDAMipd::load_trial_data(costs_file)
result <- costing_inpatient_daycase_admission(ind_part_data,
  hrg_code_ip_admi = "HRGcode", descrip_ip_admi = NULL,
  number_use_ip_admi = "number_use", elective_col = "EL",
  unit_cost_data, hrg_code_col = "Currency_Code", description_col = NULL,
  unit_cost_col = "National_Average_Unit_Cost",
  cost_calculated_in = "admission")
```

---

costing\_opioid\_liquids\_averageMED\_long

*Function to estimate the cost of liquids when IPD is in long format*

---

**Description**

Function to estimate the cost of liquids when IPD is in long format

**Usage**

```
costing_opioid_liquids_averageMED_long(
  the_columns,
  ind_part_data_long,
  name_med,
  brand_med = NULL,
  bottle_size,
  bottle_size_unit = NULL,
  bottle_last,
  bottle_last_unit = NULL,
  preparation_dose,
  preparation_unit = NULL,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_brand = NULL,
  list_of_code_bottle_size_unit = NULL,
```

```

list_of_code_bottle_last_unit = NULL,
list_preparation_dose_unit = NULL,
eqdose_covtab = NULL,
basis_strength_unit = NULL
)

```

### Arguments

```

the_columns      columns that are to be used to convert the data from long to wide
ind_part_data_long
                  IPD
name_med         name of medication
brand_med        brand name of medication if revealed
bottle_size      size of the bottle used
bottle_size_unit
                  unit of bottle volume
bottle_last     how long the bottle lasted
bottle_last_unit
                  time unit of how long the bottle lasted
preparation_dose
                  dose if preparation is given
preparation_unit
                  unit of preparatio dose
timeperiod       time period for cost calculation
unit_cost_data   unit costs data
unit_cost_column
                  column name of unit cost in unit_cost_data
cost_calculated_per
                  column name of unit where the cost is calculated
strength_column
                  column column name that has strength of medication
list_of_code_names
                  if names is coded, give the code:name pairs, optional
list_of_code_brand
                  if brand names are coded, give the code:brand pairs, optional
list_of_code_bottle_size_unit
                  list of bottle size units and codes
list_of_code_bottle_last_unit
                  list of time of bottle lasts and codes
list_preparation_dose_unit
                  list of preparation dose units and codes
eqdose_covtab    table to get the conversion factor for equivalent doses, optional
basis_strength_unit
                  strength unit to be taken as basis required for total medication calculations

```

**Value**

the calculated cost of tablets along with original data

**Examples**

```
med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
  package = "packDAMipd")
data_file <- system.file("extdata", "medication_liq_brand_empty.xlsx",
  package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx",
  package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- costing_opioid_liquids_averageMED_wide(
  ind_part_data = ind_part_data, name_med = "liq_name",
  brand_med = "liq_brand", bottle_size = "liq_bottle_size",
  bottle_size_unit = NULL, bottle_last = "liq_last",
  bottle_last_unit = NULL, preparation_dose = "liq_strength",
  preparation_unit = NULL, timeperiod = "1 day",
  unit_cost_data = med_costs, unit_cost_column = "UnitCost",
  cost_calculated_per = "Basis", strength_column = "Strength",
  list_of_code_names = NULL, list_of_code_brand = NULL,
  list_of_code_bottle_size_unit = NULL,
  list_of_code_bottle_last_unit = NULL,
  list_preparation_dose_unit = NULL, eqdose_covtab = table,
  basis_strength_unit = NULL)
```

---

costing\_opioid\_liquids\_averageMED\_wide

*Function to estimate the cost of liquids taken (from IPD)*

---

**Description**

Function to estimate the cost of liquids taken (from IPD)

**Usage**

```
costing_opioid_liquids_averageMED_wide(
  ind_part_data,
  name_med,
  brand_med = NULL,
  bottle_size,
  bottle_size_unit = NULL,
  bottle_last,
  bottle_last_unit = NULL,
  preparation_dose,
  preparation_unit = NULL,
```

```

timeperiod,
unit_cost_data,
unit_cost_column,
cost_calculated_per,
strength_column,
list_of_code_names = NULL,
list_of_code_brand = NULL,
list_of_code_bottle_size_unit = NULL,
list_of_code_bottle_last_unit = NULL,
list_preparation_dose_unit = NULL,
eqdose_covtab = NULL,
basis_strength_unit = NULL
)

```

### Arguments

```

ind_part_data   IPD
name_med        name of medication
brand_med       brand name of medication if revealed
bottle_size     size of the bottle used
bottle_size_unit
                unit of bottle volume
bottle_last    how long the bottle lasted
bottle_last_unit
                time unit of how long the bottle lasted
preparation_dose
                dose if preparation is given
preparation_unit
                unit of preparation dose
timeperiod      time period for cost calculation
unit_cost_data  unit costs data
unit_cost_column
                column name of unit cost in unit_cost_data
cost_calculated_per
                column name of unit where the cost is calculated
strength_column
                column column name that has strength of medication
list_of_code_names
                if names is coded, give the code:name pairs, optional
list_of_code_brand
                if brand names are coded, give the code:brand pairs, optional
list_of_code_bottle_size_unit
                list of bottle size units and codes
list_of_code_bottle_last_unit
                list of time of bottle lasts and codes

```

```

list_preparation_dose_unit
    list of preparation dose units and codes

eqdose_covtab    table to get the conversion factor for equivalent doses, optional, but the column
                  names have to unique Similar to c("Drug", "form", "unit", "factor") or c("Drug",
                  "form", "unit", "conversion")

basis_strength_unit
    strength unit to be taken as basis required for total medication calculations

```

**Value**

the calculated cost of tablets along with original data

**Examples**

```

med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
package = "packDAMipd")
data_file <- system.file("extdata", "medication_liq.xlsx",
package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx",package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- microcosting_liquids_wide(
ind_part_data = ind_part_data, name_med = "liq_name", brand_med = NULL,
dose_med = "liq_strength", unit_med = NULL, bottle_size = "liq_bottle_size",
bottle_size_unit = NULL, bottle_last = "liq_last",
bottle_last_unit = NULL, preparation_dose = NULL, preparation_unit = NULL,
timeperiod = "4 months", unit_cost_data = med_costs,
unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
strength_column = "Strength", list_of_code_names = NULL,
list_of_code_brand = NULL, list_of_code_dose_unit = NULL,
list_of_code_bottle_size_unit = NULL, list_of_code_bottle_last_unit = NULL,
list_preparation_dose_unit = NULL, eqdose_covtab = table,
basis_strength_unit = NULL)

```

---

```

costing_opioid_patches_averageMED_long

```

```

#' #####
Function to estimate the cost of patches when IPD is in long format
using a IPD data of long format

```

---

**Description**

```

#' ##### Function
to estimate the cost of patches when IPD is in long format using a IPD data of long format

```

**Usage**

```

costing_opioid_patches_averageMED_long(
  the_columns,
  ind_part_data_long,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

**Arguments**

the_columns	columns that are to be used to convert the data from long to wide
ind_part_data_long	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken
freq_taken	frequency of medication
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data
cost_calculated_per	column name of unit in the cost is calculated
strength_column	column column name that contain strength of medication
list_of_code_names	if names is coded, give the code:name pairs, optional

list\_of\_code\_freq  
if frequency is coded, give the code:frequency pairs, optional

list\_of\_code\_dose\_unit  
if unit is coded, give the code:unit pairs, optional

list\_of\_code\_brand  
if brand names are coded, give the code:brand pairs, optional

eqdose\_cov\_tab table to get the conversion factor for equivalent doses, optional

basis\_strength\_unit  
strength unit to be taken as basis required for total medication calculations

**Value**

the calculated cost of tablets along with original data

**Examples**

```
med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
  package = "packDAMipd")
data_file <- system.file("extdata", "medication.xlsx",
  package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx", package = "packDAMipd")
table <- load_trial_data(conv_file)
names <- colnames(ind_part_data)
ending <- length(names)
ind_part_data_long <- tidyr::gather(ind_part_data, measurement, value,
  names[2]:names[ending], factor_key = TRUE)
the_columns <- c("measurement", "value")
res <- costing_opioid_patches_averageMED_long(the_columns,
  ind_part_data_long = ind_part_data_long, name_med = "patch_name",
  brand_med = "patch_brand", dose_med = "patch_strength", unit_med = NULL,
  no_taken = "patch_no_taken", freq_taken = "patch_frequency",
  timeperiod = "4 months", unit_cost_data = med_costs,
  unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
  strength_column = "Strength", list_of_code_names = NULL,
  list_of_code_freq = NULL, list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL, eqdose_cov_tab = table,
  basis_strength_unit = "mcg/hr")
```

---

costing\_opioid\_patches\_averageMED\_wide

*Function to estimate the cost of patches taken (from IPD)*

---

**Description**

Function to estimate the cost of patches taken (from IPD)

**Usage**

```

costing_opioid_patches_averageMED_wide(
  ind_part_data,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

**Arguments**

ind_part_data	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken
freq_taken	frequency of medication
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data
cost_calculated_per	column name of unit where the cost is calculated
strength_column	column column name that contain strength of medication
list_of_code_names	if names is coded, give the code:name pairs, optional
list_of_code_freq	if frequency is coded, give the code:frequency pairs, optional
list_of_code_dose_unit	if unit is coded, give the code:unit pairs, optional



list\_of\_code\_brand  
                   if brand names are coded, give the code:brand pairs, optional

eqdose\_cov\_tab table to get the conversion factor for equivalent doses, optional, but the column names have to be unique Similar to c("Drug", "form", "unit", "factor") or c("Drug", "form", "unit", "conversion")

basis\_strength\_unit  
                   strength unit to be taken as basis required for total medication calculations

### Details

Assumes individual level data has name of medication, dose, dose unit, number taken, frequency taken, and basis time Assumes unit cost data contains the name of medication, form/type, strength, unit of strength (or the unit in which the cost calculated), preparation, unit cost, size and size unit (in which name, forms, size, size unit, and preparation are not passed on) @importFrom dplyr %>%

### Value

the calculated cost of tablets along with original data

### Examples

```
med_costs_file <- system.file("extdata", "medication_costs_all.xlsx",
  package = "packDAMipd")
data_file <- system.file("extdata", "medication.xlsx",
  package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx", package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- costing_opioid_patches_averageMED_wide(
  ind_part_data = ind_part_data, name_med = "patch_name",
  brand_med = "patch_brand", dose_med = "patch_strength", unit_med = NULL,
  no_taken = "patch_no_taken", freq_taken = "patch_frequency",
  timeperiod = "4 months", unit_cost_data = med_costs,
  unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
  strength_column = "Strength", list_of_code_names = NULL,
  list_of_code_freq = NULL, list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL, eqdose_cov_tab = table,
  basis_strength_unit = "mcg/hr")
```

---

costing\_opioid\_tablets\_averageMED\_long

*Function to estimate the cost of tablets when IPD is in long format*

---

### Description

Function to estimate the cost of tablets when IPD is in long format

**Usage**

```

costing_opioid_tablets_averageMED_long(
  the_columns,
  ind_part_data_long,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

**Arguments**

the_columns	columns that are to be used to convert the data from long to wide
ind_part_data_long	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken
freq_taken	frequency of medication
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data
cost_calculated_per	column name of unit where the cost is calculated
strength_column	column column name that contain strength of medication
list_of_code_names	if names is coded, give the code:name pairs, optional

```

list_of_code_freq
    if frequency is coded, give the code:frequency pairs, optional
list_of_code_dose_unit
    if unit is coded, give the code:unit pairs, optional
list_of_code_brand
    if brand names are coded, give the code:brand pairs, optional
eqdose_cov_tab  table to get the conversion factor for equivalent doses, optional
basis_strength_unit
    strength unit to be taken as basis required for total medication calculations

```

**Value**

the calculated cost of tablets along with original data

**Examples**

```

med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
package = "packDAMipd")
data_file <- system.file("extdata", "medication_all.xlsx",
package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx", package = "packDAMipd")
table <- load_trial_data(conv_file)
names <- colnames(ind_part_data)
ending <- length(names)
ind_part_data_long <- tidyr::gather(ind_part_data, measurement, value,
names[2]:names[ending], factor_key = TRUE)
the_columns <- c("measurement", "value")
res <- microcosting_tablets_long(the_columns,
ind_part_data_long = ind_part_data_long, name_med = "tab_name",
brand_med = "tab_brand", dose_med = "tab_strength",
unit_med = "tab_str_unit",
no_taken = "tab_no_taken", freq_taken = "tab_frequency",
timeperiod = "2 months", unit_cost_data = med_costs,
unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
strength_column = "Strength", list_of_code_names = NULL,
list_of_code_freq = NULL, list_of_code_dose_unit = NULL,
eqdose_cov_tab = table, basis_strength_unit = "mg")

```

---

costing\_opioid\_tablets\_averageMED\_wide

*Function to estimate the cost of tablets taken as an average cost per equivalent dose in the opioid scenario is the morphine equivalent dose (from IPD)*

---

**Description**

Function to estimate the cost of tablets taken as an average cost per equivalent dose in the opioid scenario is the morphine equivalent dose (from IPD)

**Usage**

```

costing_opioid_tablets_averageMED_wide(
  ind_part_data,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

**Arguments**

ind_part_data	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken
freq_taken	frequency of medication
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data
cost_calculated_per	column name of unit where the cost is calculated
strength_column	column column name that contain strength of medication
list_of_code_names	if names is coded, give the code:name pairs, optional
list_of_code_freq	if frequency is coded, give the code:frequency pairs, optional
list_of_code_dose_unit	if unit is coded, give the code:unit pairs, optional

list\_of\_code\_brand  
                   if brand names are coded, give the code:brand pairs, optional

eqdose\_cov\_tab table to get the conversion factor for equivalent doses, optional, but the column names have to be unique Similar to c("Drug", "form", "unit", "factor") or c("Drug", "form", "unit", "conversion")

basis\_strength\_unit  
                   strength unit to be taken as basis required for total medication calculations

### Details

Assumes individual level data has name of medication, dose, dose unit, number taken, frequency taken, and basis time Assumes unit cost data contains the name of medication, form/type, strength, unit of strength (or the unit in which the cost calculated), preparation, unit cost, size and size unit (in which name, forms, size, size unit, and preparation are not passed on) @importFrom dplyr %>%

### Value

the calculated cost of tablets along with original data

### Examples

```
med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
  package = "packDAMipd")
data_file <- system.file("extdata", "medication_all_brandNull.xlsx",
  package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx",
  package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- costing_opioid_tablets_MED_wide(ind_part_data = ind_part_data,
  name_med = "tab_name", brand_med = "tab_brand", dose_med = "tab_str",
  unit_med = "tab_unit", no_taken = "tab_no_taken",
  freq_taken = "tab_frequency", timeperiod = "one day",
  unit_cost_data = med_costs, unit_cost_column = "UnitCost",
  cost_calculated_per = "Basis", strength_column = "Strength",
  list_of_code_names = NULL, list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL, eqdose_cov_tab = table,
  basis_strength_unit = "mg")
```

---

costing\_opioid\_tablets\_MED\_wide

*Function to estimate the cost of tablets taken (from IPD)*

---

### Description

Function to estimate the cost of tablets taken (from IPD)

**Usage**

```

costing_opioid_tablets_MED_wide(
  ind_part_data,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

**Arguments**

ind_part_data	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken
freq_taken	frequency of medication
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data
cost_calculated_per	column name of unit where the cost is calculated
strength_column	column column name that contain strength of medication
list_of_code_names	if names is coded, give the code:name pairs, optional
list_of_code_freq	if frequency is coded, give the code:frequency pairs, optional
list_of_code_dose_unit	if unit is coded, give the code:unit pairs, optional

list\_of\_code\_brand  
if brand names are coded, give the code:brand pairs, optional

eqdose\_cov\_tab table to get the conversion factor for equivalent doses, optional, but the column names have to be unique Similar to c("Drug", "form", "unit", "factor") or c("Drug", "form", "unit", "conversion")

basis\_strength\_unit  
strength unit to be taken as basis required for total medication calculations

### Details

Assumes individual level data has name of medication, dose, dose unit, number taken, frequency taken, and basis time Assumes unit cost data contains the name of medication, form/type, strength, unit of strength (or the unit in which the cost calculated), preparation, unit cost, size and size unit (in which name, forms, size, size unit, and preparation are not passed on) @importFrom dplyr %>%

### Value

the calculated cost of tablets along with original data

### Examples

```
med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
  package = "packDAMipd")
data_file <- system.file("extdata", "medication_all_brandNull.xlsx",
  package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx",
  package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- costing_opioid_tablets_MED_wide(ind_part_data = ind_part_data,
  name_med = "tab_name", brand_med = "tab_brand", dose_med = "tab_str",
  unit_med = "tab_unit", no_taken = "tab_no_taken",
  freq_taken = "tab_frequency", timeperiod = "one day",
  unit_cost_data = med_costs, unit_cost_column = "UnitCost",
  cost_calculated_per = "Basis", strength_column = "Strength",
  list_of_code_names = NULL, list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL, eqdose_cov_tab = table,
  basis_strength_unit = "mg")
```

---

costing\_resource\_use *Function to estimate the cost of resource use taken (from IPD)*

---

### Description

Function to estimate the cost of resource use taken (from IPD)

**Usage**

```

costing_resource_use(
  ind_part_data,
  name_use_col,
  each_length_num_use = NULL,
  each_use_provider_indicator = NULL,
  unit_length_use = "day",
  unit_cost_data,
  name_use_unit_cost,
  unit_cost_column,
  cost_calculated_in,
  list_code_use_indicator = NULL,
  list_code_provider_indicator = NULL
)

```

**Arguments**

`ind_part_data` IPD

`name_use_col` name of the column containing resource use

`each_length_num_use`  
list of column names that shows length/number of repeated use eg. hospital admission

`each_use_provider_indicator`  
list of column names that shows the bool indicators for the use of resource if this is to be included for the particular provider, say an nhs hospital use

`unit_length_use`  
the column name that contains how many or how long used

`unit_cost_data` unit costs data where the assumption is that the unit cost for resources such as hospital use, gp visit are listed in column resource/resource use with unit costs in another column and the units calculated as in another column

`name_use_unit_cost`  
name of resource use (the column name in the unit cost data is assumed to be name/resource/type etc) in unit cost data

`unit_cost_column`  
column name of unit cost in `unit_cost_data`

`cost_calculated_in`  
column name of unit where the cost is calculated

`list_code_use_indicator`  
if the column `name_use_col` shows codes to indicate the resource use provide the list of codes and resource use for eg., `list(c("yes", "no", c(1,2)))`

`list_code_provider_indicator`  
column `each_use_provider_indicator` shows codes to indicate the resource use provide the list of codes and resource use for eg., `list(c("yes", "no", c(1,2)))`

**Value**

the calculated cost of resource uses along with original data



**Examples**

```

costs_file <- system.file("extdata", "costs_resource_use.csv",
  package = "packDAMipd")
datafile <- system.file("extdata", "resource_use_hc_2.csv",
  package = "packDAMipd")
ind_part_data <- load_trial_data(datafile)
unit_cost_data <- load_trial_data(costs_file)
res <- costing_resource_use(
  ind_part_data[1, ],
  "hospital_admission_1",
  list("length_1", "length_2"),
  list("nhs_1", "nhs_2"),
  "day",
  unit_cost_data, "Inpatient hospital admissions", "UnitCost",
  "UnitUsed",
  NULL, NULL
)

```

---

cost\_data.df

*cost matrix*


---

**Description**

cost matrix

**Usage**

```
cost_data.df
```

**Format**

A 11 by 2 dataframe

**Source**

created on Nov 26, 2019 from `tmat <- rbind(c(1, 2), c(3, 4)) colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead") tm <- transition_matrix(2, tmat, c(0.5, 0.5, 0, 1)) a <- health_state("Healthy", 1, 1, FALSE) b <- health_state("Dead", 1, 0, TRUE) health_states <- combine_state(a, b) this.strategy <- strategy(tm, health_states, "intervention")`

---

create_new_dataset	<i>create new dataset while keeping cox regression results and returned coefficients</i>
--------------------	--

---

**Description**

create new dataset while keeping cox regression results and returned coefficients

**Usage**

```
create_new_dataset(var, covar, dataset, categorical)
```

**Arguments**

var	variable for which the levels have to be identified usually indep variable
covar	the other covariates
dataset	the dataset where these variables contain
categorical	are these variables categorical? True of false

**Value**

new data frame

**Examples**

```
dataset <- survival::lung
new = create_new_dataset("status", c("age"), dataset, c(FALSE))
```

---

create_shorttable_from_gtsummary_compare_twogroups_timepoints	<i>Create a table to compare the descriptive analysis (short) from gtsummary of two groups, but at different timepoints</i>
---	---

---

**Description**

Create a table to compare the descriptive analysis (short) from gtsummary of two groups, but at different timepoints

**Usage**

```
create_shorttable_from_gtsummary_compare_twogroups_timepoints(
  variables,
  gtsummary,
  name_use,
  timepoints
)
```

**Arguments**

variables	variables that interested
gtsummary	a gtsummary object that contains summary parameters
name_use	name of the variable or category
timepoints	the timepoints at which the descriptive analysis is done

**Value**

the table

**Examples**

```
eg_data <- as.data.frame(list(no = c(1, 2, 3, 4),
  mark_at_1 = c(12, 34, 23, 45), gender = c("M", "F", "M", "F"),
  mark_at_2 = c(12, 34, 23, 45)))
outcome_summary <- IPDFileCheck::get_summary_gtsummary(eg_data,
  c("gender", "mark_at_1", "mark_at_2"), byvar = "gender")
variables <- "Mark"
k <- create_shorttable_from_gtsummary_compare_twogroups_timepoints(variables,
  outcome_summary, "Category", c(1, 2))
```

---

```
create_table_from_gtsummary_compare_twogroups
```

*Create a table to compare the descriptive analysis from gtsummary of two groups*

---

**Description**

Create a table to compare the descriptive analysis from gtsummary of two groups

**Usage**

```
create_table_from_gtsummary_compare_twogroups(variables, gtsummary, name_use)
```

**Arguments**

variables	variables that interested
gtsummary	a gtsummary object that contains summary parameters
name_use	name of the variable or category

**Value**

the table

## Examples

```
eg_data <- as.data.frame(list(no = c(1,2,3,4), mark = c(12,34,23,45),
gender = c("M", "F", "M", "F")))
outcome_summary <- IPDFileCheck::get_summary_gtsummary(eg_data,
c("gender", "mark"), byvar = "gender")
variables <- "Mark"
create_table_from_gtsummary_compare_twogroups(variables,
outcome_summary, "Category")
```

---

define\_parameters      *Function to return a list of parameters given*

---

## Description

Function to return a list of parameters given

## Usage

```
define_parameters(...)
```

## Arguments

...                    any parameters set of name value pairs expected

## Details

To return a list of parameters For using with assign\_parameters() just list or enumerate the parameters, do not use c() or list() to create a data type list

## Value

a list of parameters

## Examples

```
define_parameters(rr = 1)
```

---

define\_parameters\_psa *Define parameter lists for deterministic sensitivity analysis*

---

## Description

Define parameter lists for deterministic sensitivity analysis

## Usage

```
define_parameters_psa(base_param_list, sample_list)
```

## Arguments

`base_param_list` list of parameters that used to define Markov model  
`sample_list` list of parameter values with their sampling distributions

## Value

table for probability sensitivity analysis

## Examples

```
param_list <- define_parameters(  
  cost_zido = 2278, cost_direct_med_A = 1701,  
  cost_comm_care_A = 1055, cost_direct_med_B = 1774,  
  cost_comm_care_B = 1278,  
  cost_direct_med_C = 6948, cost_comm_care_C = 2059,  
  tpAtoA = 1251 / (1251 + 483),  
  tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),  
  tpAtoD = 17 / (17 + 1717),  
  tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),  
  tpBtoD = 15 / (15 + 1243),  
  tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312), tpDtoD = 1,  
  cost_health_A = "cost_direct_med_A+ cost_comm_care_A",  
  cost_health_B = "cost_direct_med_B+ cost_comm_care_B",  
  cost_health_C = "cost_direct_med_C+ cost_comm_care_C",  
  cost_drug = "cost_zido"  
)  
sample_list <- define_parameters(cost_zido = "gamma(mean = 2756,  
  sd = sqrt(2756))")  
param_table <- define_parameters_psa(param_list, sample_list)
```

---

```
define_parameters_sens_anal
  ISLR flextable huxtable nlme tm Define parameter lists for deterministic sensitivity analysis
```

---

## Description

ISLR flextable huxtable nlme tm Define parameter lists for deterministic sensitivity analysis

## Usage

```
define_parameters_sens_anal(param_list, low_values, upp_values)
```

## Arguments

param_list	list of parameters that used to define Markov model
low_values	list of lower values of those parameters for whom the sensitivity is to be estimated
upp_values	list of upper values of those parameters for whom the sensitivity is to be estimated

## Details

Get the parameter list, min and maximum values of the parameters. The min and max values should have same entries, but they should be contained in param\_list too. Copy the exact values of parameters that are in param list but not in min and max values

## Value

table for sensitivity analysis

## Examples

```
param_list <- define_parameters(
  cost_zido = 2278, cost_direct_med_A = 1701,
  cost_comm_care_A = 1055, cost_direct_med_B = 1774,
  cost_comm_care_B = 1278,
  cost_direct_med_C = 6948, cost_comm_care_C = 2059,
  tpAtoA = 1251 / (1251 + 483),
  tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),
  tpAtoD = 17 / (17 + 1717),
  tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),
  tpBtoD = 15 / (15 + 1243),
  tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312), tpDtoD = 1,
  cost_health_A = "cost_direct_med_A + cost_comm_care_A",
  cost_health_B = "cost_direct_med_B + cost_comm_care_B",
  cost_health_C = "cost_direct_med_C + cost_comm_care_C",
  cost_drug = "cost_zido"
```

```
)  
low_values <- define_parameters(cost_direct_med_B = 177.4,  
cost_comm_care_C = 205.9)  
upp_values <- define_parameters(cost_direct_med_B = 17740,  
cost_comm_care_C = 20590)  
param_table <- define_parameters_sens_anal(param_list, low_values,  
upp_values)
```

---

define\_transition\_table

*Define the table for transition*

---

## Description

Define the table for transition

## Usage

```
define_transition_table(tmat)
```

## Arguments

tmat                    transition matrix in the format as in package 'mstate'

## Details

Generating a table for transition matrix for efficient understanding and checking The transition matrix in the format as per 'mstate' package is transformed to a table. if tmat is not a square matrix, it gives error else it spells out the transition number, probability name and from state to state

## Value

the transition table with the probabilities

## Examples

```
tmat <- rbind(c(1, 2), c(3, 4))  
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")  
define_transition_table(tmat)
```

do\_psa

*Function to do probabilistic sensitivity analysis***Description**

Function to do probabilistic sensitivity analysis

**Usage**

```
do_psa(this_markov, psa_table, num_rep)
```

**Arguments**

<code>this_markov</code>	Markov model object
<code>psa_table</code>	table object from <code>define_parameters_psa</code>
<code>num_rep</code>	number of repetitions

**Value**

result after sensitivity analysis

**Examples**

```
param_list <- define_parameters(
  cost_zido = 2278, cost_direct_med_A = 1701,
  cost_comm_care_A = 1055, cost_direct_med_B = 1774,
  cost_comm_care_B = 1278,
  cost_direct_med_C = 6948, cost_comm_care_C = 2059,
  tpAtoA = 1251 / (1251 + 483),
  tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),
  tpAtoD = 17 / (17 + 1717),
  tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),
  tpBtoD = 15 / (15 + 1243),
  tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312), tpDtoD = 1,
  cost_health_A = "cost_direct_med_A+ cost_comm_care_A",
  cost_health_B = "cost_direct_med_B+ cost_comm_care_B",
  cost_health_C = "cost_direct_med_C+ cost_comm_care_C",
  cost_drug = "cost_zido"
)
A <- health_state("A", cost = "cost_health_A+ cost_drug ", utility = 1)
B <- health_state("B", cost = "cost_health_B + cost_drug", utility = 1)
C <- health_state("C", cost = "cost_health_C + cost_drug", utility = 1)
D <- health_state("D", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3, 4), c(NA, 5, 6, 7), c(NA, NA, 8, 9),
c(NA, NA, NA, 10))
colnames(tmat) <- rownames(tmat) <- c("A", "B", "C", "D")
tm <- populate_transition_matrix(4, tmat, c(
  "tpAtoA", "tpAtoB", "tpAtoC", "tpAtoD",
  "tpBtoB", "tpBtoC", "tpBtoD", "tpCtoC", "tpCtoD", "tpDtoD"
```



```

), colnames(tmat))
health_states <- combine_state(A, B, C, D)
mono_strategy <- strategy(tm, health_states, "mono")
mono_markov <- markov_model(mono_strategy, 20, discount = c(0.06, 0),
initial_state =c(1,0,0,0),param_list)
sample_list <- define_parameters(cost_zido = "gamma(mean = 2756,
sd = sqrt(2756))")
param_table <- define_parameters_psa(param_list, sample_list)
result <- do_psa(mono_markov, param_table, 10)

```

---

do\_sensitivity\_analysis

*Function to do deterministic sensitivity analysis*

---

### Description

Function to do deterministic sensitivity analysis

### Usage

```
do_sensitivity_analysis(this_markov, param_table)
```

### Arguments

this_markov	Markov model object
param_table	table object from define_parameters_sens_anal() with parameters (base case value, lower and upper)

### Value

result after sensitivity analysis

### Examples

```

param_list <- define_parameters(
cost_zido = 2278, cost_direct_med_A = 1701,
cost_comm_care_A = 1055, cost_direct_med_B = 1774,
cost_comm_care_B = 1278,
cost_direct_med_C = 6948, cost_comm_care_C = 2059,
tpAtoA = 1251 / (1251 + 483),
tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),
tpAtoD = 17 / (17 + 1717),
tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),
tpBtoD = 15 / (15 + 1243),
tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312),
tpDtoD = 1,
cost_health_A = "cost_direct_med_A + cost_comm_care_A",
cost_health_B = "cost_direct_med_B + cost_comm_care_B",

```

```

cost_health_C = "cost_direct_med_C + cost_comm_care_C",
cost_drug = "cost_zido")
low_values <- define_parameters(cost_direct_med_B = 177.4,
cost_comm_care_C = 205.9)
upp_values <- define_parameters(cost_direct_med_B = 17740,
cost_comm_care_C = 20590)
A <- health_state("A", cost = "cost_health_A + cost_drug ", utility = 1)
B <- health_state("B", cost = "cost_health_B + cost_drug", utility = 1)
C <- health_state("C", cost = "cost_health_C + cost_drug", utility = 1)
D <- health_state("D", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3, 4), c(NA, 5, 6, 7), c(NA, NA, 8, 9),
c(NA, NA, NA, 10))
colnames(tmat) <- rownames(tmat) <- c("A", "B", "C", "D")
tm <- populate_transition_matrix(4, tmat, c("tpAtoA", "tpAtoB", "tpAtoC",
"tpAtoD", "tpBtoB", "tpBtoC", "tpBtoD", "tpCtoC", "tpCtoD", "tpDtoD"),
colnames(tmat))
health_states <- combine_state(A, B, C, D)
mono_strategy <- strategy(tm, health_states, "mono")
mono_markov <- markov_model(mono_strategy, 20, c(1, 0, 0, 0),
discount = c(0.06, 0), param_list)
param_table <- define_parameters_sens_anal(param_list, low_values,
upp_values)
result <- do_sensitivity_analysis(mono_markov, param_table)

```

---

encode\_codes\_data

*Function to get the codes and the corresponding entries*

---

## Description

Function to get the codes and the corresponding entries

## Usage

```
encode_codes_data(list_code_values, data_column_nos, the_data)
```

## Arguments

`list_code_values`  
list of codes and values, given as list of lists

`data_column_nos`  
the column numbers of data to look for the entries

`the_data`  
the data where to look for

## Value

weight and vol units

**Examples**

```

data_file <- system.file("extdata", "medication_liq_codes.xlsx",
package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
data_column_nos = c(2,12)
list_of_code_names = list(c(1, 2),c("Morphine", "Oxycodone"))
encode_codes_data(list_of_code_names, data_column_nos, ind_part_data)

```

---

eval\_assign\_trans\_prob

*Attribute parameters to probabilities of transition matrix*

---

**Description**

Attribute parameters to probabilities of transition matrix

**Usage**

```
eval_assign_trans_prob(tm, parameter_values)
```

**Arguments**

**tm**                    A transition matrix in the format from the package 'mstate'

**parameter\_values**       name value pairs of parameter values in the probability matrix

**Details**

Once the transition matrix is populated, the probabilities in transition matrix gets evaluated and assigned in this function call. If the entry in transition matrix is NA, replaces it with zero. Similarly to evaluate and assign health states, the parameter values is expected to be a list from `assign_parameter()` and `define_parameter()`. The exception is that if the parameters are defined directly and no nested calculation is required. For eg. `assign_list = c(p1 = 0.2, p2 = 0.3, p3 = 0.4, p4 = 0.5)` `prob <- eval_assign_trans_prob(tmat, assign_list)` will work. For those with nested calculations, this has to be defined as below `assign_list<-assign_parameters(define_parameters(p1 = 0.2, p2 = 0.3, p3 = 0.4, p4 = 0.5))` `prob <- eval_assign_trans_prob(tmat, assign_list)`. The below will give error `assign_list <- c(p1=0.1, p2 = "p1 + 0.2", p3=0, p4=0.3)` `prob <- eval_assign_trans_prob(tmat, assign_list)`

**Value**

the transition table with the probabilities

**Examples**

```
tmat <- rbind(c(1, 2), c(3, 4))
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")
tmat <- populate_transition_matrix(2, tmat,
list_prob = c("p1", "p2", "p3", "p4"))
tmat_assigned <- eval_assign_trans_prob(tmat,
c(p1 = 0.2, p2 = 0.3, p3 = 0.4, p4 = 0.5))
```

---

eval\_assign\_values\_states

*Attribute values in health states*

---

**Description**

Attribute values in health states

**Usage**

```
eval_assign_values_states(health_states, assigned_param)
```

**Arguments**

health\_states list of health\_state objects

assigned\_param name value pairs of parameter values in the probability matrix expected created using function assign\_parameters()

**Details**

Assigning the param is done for the cost and utility if the param is not numeric, check if it can be evaluated at the run time if yes, assign the evaluated numeric value if not get the parameters between operators, and assign the values to each individual parameters and then evaluate. only works for two levels. For the example shown the cost is sum of cost\_A and cost\_B which will only get added in the call eval\_assign\_values\_states While initialising the state "well" it will be only saved as expression(cost\_A + cost\_B) assigned\_param (a list) can be expected to be created using assign\_parameters() the exception is if parameter is directly assigned with no nested calculation and no missing parameters. For example assigned\_param = c(cost\_a = 10, cost\_b=10) will be ok but not assigned\_param = c(a=10, cost\_A = "a+100", cost\_B =10) as it requires a nested calculation then use define\_parameters() with assign\_parameters() as in param\_list <- define\_parameters(a = 10, cost\_A = "a + 100", cost\_B = 10) assign\_list <- assign\_parameters(param\_list)

**Value**

health states with assigned values

**Examples**

```
well <- health_state("well", cost = "cost_A + cost_B", utility = 1)
disabled <- health_state("disabled", cost = 100, utility = 1)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.6, 0.2, 0.2, 0.6, 0.4, 1))
health_states <- combine_state(well, disabled, dead)
eval_assign_values_states(health_states, c(cost_A = 100, cost_B = 11))
```

---

find\_glm\_distribution *Function to find the keyword for family of distribution in glm*

---

**Description**

Function to find the keyword for family of distribution in glm

**Usage**

```
find_glm_distribution(text)
```

**Arguments**

text	distribution
------	--------------

**Details**

Find the family for glm method

**Value**

the keyword - the name of distribution

**Examples**

```
find_glm_distribution("gamma")
```

---

find\_keyword\_rand\_generation

*Function to find the keyword for generating random numbers the distribution*

---

**Description**

Function to find the keyword for generating random numbers the distribution

**Usage**

```
find_keyword_rand_generation(text)
```

**Arguments**

text                    name of the probability distribution

**Details**

This function returns the keyword for generating random number using a keyword provided that is generally used for prob distribution (but R might require a different keyword)

**Value**

the keyword that should be used in R for generating random numbers

**Examples**

```
find_keyword_rand_generation("gamma")
```

---

find\_keyword\_regression\_method

*Function to find the keyword for regression methods*

---

**Description**

Function to find the keyword for regression methods

**Usage**

```
find_keyword_regression_method(text, additional_info = NA)
```

**Arguments**

text                    regression method  
additional\_info        additional information required

**Details**

This function returns the keyword to use in regression methods. For example linear regression requires lm in R some regression methods require additional info and it has to be provided

**Value**

the keyword that should be used in R for regression analysis

**Examples**

```
find_keyword_regression_method("linear")
```

---

```
find_parameters_btn_operators
```

*Function to return parameters with in a expression containing operators*

---

**Description**

Function to return parameters with in a expression containing operators

**Usage**

```
find_parameters_btn_operators(expr)
```

**Arguments**

expr            an expression

**Details**

This function returns the parameters between the operators if the state value or probabilities are defined as expressions, we need to extract the parameters and then assign First the position of all operators are found and then return the parameters separated by those operators This happens only for one level find\_parameters\_btn\_operators("a+b") provides a and b but for find\_parameters\_btn\_operators("mean(a,b)+b") provides mean(a,b) and b

**Value**

parameters in the expression expr

**Examples**

```
find_parameters_btn_operators("a+b")
```

---

find\_required\_parameter\_combs

*Function to find the parameters that determine the probability distribution*

---

### **Description**

Function to find the parameters that determine the probability distribution

### **Usage**

```
find_required_parameter_combs(name_distri)
```

### **Arguments**

name\_distri      name of the probability distribution

### **Details**

For each of the probability distribution we require certain parameters and this function provides that required list of parameters.

### **Value**

the parameters that determine the distribution

### **Examples**

```
find_required_parameter_combs("gamma")
```

---

find\_rowwise\_sum\_multiplecol

*Function to get sum of multiple columns of observations*

---

### **Description**

Function to get sum of multiple columns of observations

### **Usage**

```
find_rowwise_sum_multiplecol(the_data, colnames, sumcolname)
```

### **Arguments**

the\_data            the data where the observations are held  
colnames            columnname in the data whose sum to be obtained  
sumcolname          name of the new column where sum to be saved



**Value**

the data with added sum

**Examples**

```
eg_data <- as.data.frame(list(no = c(1, 2, 3, 4),
  mark_at_1 = c(12, 7, 23, 45), gender = c("M", "F", "M", "F"),
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2"),
  time = c(1,1,2,2), id = c(1, 1, 1, 2)))
find_rowwise_sum_multiplecol(eg_data, c("mark_at_1", "mark_at_2"),
  "totalmarks")
```

---

find\_survreg\_distribution

*Function to find the keyword for survreg distribution*

---

**Description**

Function to find the keyword for survreg distribution

**Usage**

```
find_survreg_distribution(text)
```

**Arguments**

text	distribution
------	--------------

**Details**

For survreg method, find the distribution

**Value**

the keyword - the name of distribution

**Examples**

```
find_survreg_distribution("weibull")
```

---

form\_expression\_glm *Form expression to use with glm()*

---

### Description

Form expression to use with glm()

### Usage

```
form_expression_glm(  
  param_to_be_estimated,  
  indep_var,  
  family,  
  covariates,  
  interaction,  
  naaction,  
  link  
)
```

### Arguments

param_to_be_estimated	parameter of interest
indep_var	the independent variable (column name in data file)
family	distribution name eg. for logistic regression -binomial
covariates	list of covariates
interaction	boolean value to indicate interaction in the case of generalised linear models, false by default
naaction	action to be taken with the missing values
link	link function if not the default for each family

### Details

Form expression for the method glm

### Value

the formula for glm

### Examples

```
formula <- form_expression_glm("admit",  
  indep_var = "gre", family = "binomial",  
  covariates = c("gpa", "rank"), interaction = FALSE, naaction = "na.omit",  
  link = NA)
```

---

form\_expression\_lm     *Form expression to use with lm()*

---

**Description**

Form expression to use with lm()

**Usage**

```
form_expression_lm(param_to_be_estimated, indep_var, covariates, interaction)
```

**Arguments**

param_to_be_estimated	parameter of interest
indep_var	the independent variable (column name in data file)
covariates	list of covariates
interaction	boolean value to indicate interaction in the case of linear regression, false by default

**Details**

This function helps to create the expression for liner regression model it takes care of covariates and interaction

**Value**

the formula for lm

**Examples**

```
formula <- form_expression_lm("gre", indep_var = "gpa", covariates = NA,  
interaction = FALSE)
```

---

form\_expression\_mixed\_model\_lme4  
*Form expression to use with mixed models*

---

**Description**

Form expression to use with mixed models

**Usage**

```
form_expression_mixed_model_lme4(
  param_to_be_estimated,
  dataset,
  fix_eff,
  fix_eff_interact_vars,
  random_intercept_vars,
  nested_intercept_vars_pairs,
  cross_intercept_vars_pairs,
  uncorrel_slope_intercept_pairs,
  random_slope_intercept_pairs,
  family,
  link
)
```

**Arguments**

param_to_be_estimated	column name of dependent variable
dataset	a dataframe
fix_eff	names of variables as fixed effect predictors
fix_eff_interact_vars	if interaction -true
random_intercept_vars	names of variables for random intercept
nested_intercept_vars_pairs	those of the random intercept variables with nested effect
cross_intercept_vars_pairs	those of the random intercept variables with crossed effect
uncorrel_slope_intercept_pairs	variables with correlated intercepts
random_slope_intercept_pairs	random slopes intercept pairs - this is a list of paired variables
family	family of distribution for non gaussian distribution of predicted variable
link	link function for the variance

**Details**

Form the expression for mixed model

**Value**

result regression result with plot if success and -1, if failure

**Examples**

```
datafile <- system.file("extdata", "data_linear_mixed_model.csv",
  package = "packDAMipd")
dt = utils::read.csv(datafile, header = TRUE)
formula <- form_expression_mixed_model_lme4("extro",
  dataset = dt,
  fix_eff = c("open", "agree", "social"),
  fix_eff_interact_vars = NULL,
  random_intercept_vars = c("school", "class"),
  nested_intercept_vars_pairs = list(c("school", "class")),
  cross_intercept_vars_pairs = NULL,
  uncorrel_slope_intercept_pairs = NULL,
  random_slope_intercept_pairs = NULL, family = "binomial", link = NA
)
```

---

generate\_wt\_time\_units

*Function to get the weight and time units*

---

**Description**

Function to get the weight and time units

**Usage**

```
generate_wt_time_units()
```

**Value**

weight and time units

**Examples**

```
ans <- generate_wt_time_units()
```

---

generate\_wt\_vol\_units *Function to get the weight and volume units*

---

**Description**

Function to get the weight and volume units

**Usage**

```
generate_wt_vol_units()
```

**Value**

weight and vol units

**Examples**

```
ans <- generate_wt_vol_units()
```

---

get\_age\_details      *Function to get the details of the age column*

---

**Description**

Function to get the details of the age column

**Usage**

```
get_age_details(trialdata)
```

**Arguments**

trialdata      data containing individual level trial data

**Details**

expecting the data contains the information on age preferably column names "age", "dob" or "yob" or "date of birth". "year of birth", "birth year" If multiple column names match these, then first match will be chosen.

**Value**

the name of the variable related to age and the unique contents if success, else error

**Examples**

```
get_age_details(data.frame("Age" = c(21, 15),  
"arm" = c("control", "intervention")))
```

---

`get_colnames_codedvalues`

*Function to keep the column name, coded values and non response code into a dataframe*

---

**Description**

Function to keep the column name, coded values and non response code into a dataframe

**Usage**

```
get_colnames_codedvalues(variable, name, code, nrcode = NA)
```

**Arguments**

variable	name of the variable in the column
name	column name
code	coded values
nrcode	code for non response

**Value**

data frame with all the above information

**Examples**

```
get_colnames_codedvalues("arm", "pat_trial_arm", c("Y", "N"))
```

---

`get_cost_AandE_code` *Function to extract the unit hospital inpatient admission by matching code*

---

**Description**

Function to extract the unit hospital inpatient admission by matching code

**Usage**

```
get_cost_AandE_code(  
  code,  
  type_admit,  
  ref_cost_data_file,  
  col_name_code,  
  unit_cost_col,  
  type_admit_col,  
  sheet = NULL  
)
```

**Arguments**

code                   code for AE attendance  
 type\_admit            term indicating admission and type of attendance  
 ref\_cost\_data\_file    file that has unit cost  
 col\_name\_code         name of the column that has the code  
 unit\_cost\_col         name of the column with the unit cost  
 type\_admit\_col        colname that describes type of the attendance and that indicates admitted or not  
 sheet                 sheet if excel file is given

**Value**

unit cost the unit cost matching the code

**Examples**

```
ref_cost_data_file <- system.file("extdata",
  "National_schedule_of_NHS_costs_2019_AandE.csv", package = "packDAMipd")
re = get_cost_AandE_code("VB02Z", "T01A", ref_cost_data_file,
  "Currency_Code", "National_Average_Unit_Cost", "Service_Code")
```

---

get\_cost\_AandE\_description

*Function to extract the unit cost by description of AandE att matching description*

---

**Description**

Function to extract the unit cost by description of AandE att matching description

**Usage**

```
get_cost_AandE_description(
  description,
  type_admit,
  ref_cost_data_file,
  col_name_description,
  unit_cost_col,
  type_admit_col,
  sheet = NULL
)
```



**Arguments**

description      description of the AE attendance  
 type\_admit        term indicating admission and type of attendance  
 ref\_cost\_data\_file  
                   file that has unit cost  
 col\_name\_description  
                   name of the column that has the description  
 unit\_cost\_col    name of the column with the unit cost  
 type\_admit\_col   colname that describes type of the attendance and  
 sheet             sheet if excel file is given

**Value**

unit cost the unit cost matching the hrg code

**Examples**

```

ref_cost_data_file <- system.file("extdata",
  "National_schedule_of_NHS_costs_2019_AandE.csv", package = "packDAMipd")
re = get_cost_AandE_description("Emergency Medicine", "T01A",
  ref_cost_data_file, "Currency_Description", "National_Average_Unit_Cost",
  "Service_Code")

```

---

```
get_cost_ip_dc_description
```

*Function to extract the unit hospital inpatient admission by matching description*

---

**Description**

Function to extract the unit hospital inpatient admission by matching description

**Usage**

```

get_cost_ip_dc_description(
  description,
  ref_cost_data_file,
  col_name_description,
  unit_cost_col,
  sheet = NULL
)

```

**Arguments**

description      description corresponding to the inpatient admission  
 ref\_cost\_data\_file  
                   file that has unit cost  
 col\_name\_description  
                   name of the column that has the description  
 unit\_cost\_col    name of the column with the unit cost  
 sheet            sheet if excel file is given

**Value**

unit cost the unit cost matching the hrg code

**Examples**

```
ref_cost_data_file <- system.file("extdata",
  "National_schedule_of_NHS_costs_2019.csv", package = "packDAMipd")
result <- get_cost_ip_dc_description("Cerebrovascular Accident",
  ref_cost_data_file, "Currency_Description",
  "National_Average_Unit_Cost")
```

---

get\_cost\_ip\_dc\_hrg      *Function to extract the unit hospital inpatient admission by matching HRG code*

---

**Description**

Function to extract the unit hospital inpatient admission by matching HRG code

**Usage**

```
get_cost_ip_dc_hrg(  
  hrg,  
  ref_cost_data_file,  
  col_name_hrg_code,  
  unit_cost_col,  
  sheet = NULL  
)
```

**Arguments**

hrg                    hrg code corresponding to the inpatient admission  
 ref\_cost\_data\_file  
                   file that has unit cost  
 col\_name\_hrg\_code  
                   name of the column that has the hrg code  
 unit\_cost\_col    name of the colum with the unit cost  
 sheet            sheet if excel file is given

**Value**

unit cost the unit cost matching the hrg code

**Examples**

```
ref_cost_data_file <- system.file("extdata",  
  "National_schedule_of_NHS_costs_2019.csv", package = "packDAMipd")  
get_cost_ip_dc_hrg("AA22C", ref_cost_data_file, "Currency_Code",  
  "National_Average_Unit_Cost")
```

---

get\_doses\_combination *Convert the combined dose to its individual component numerical value or can be unit/unit*

---

**Description**

Convert the combined dose to its individual component numerical value or can be unit/unit

**Usage**

```
get_doses_combination(the_string, separator = "/")
```

**Arguments**

the\_string      given combined unit  
separator      given character for separation , default is "/"

**Value**

separated texts

**Examples**

```
get_doses_combination("g/ml")
```

---

```
get_doses_combination_units
```

*Convert the combined dose to its individual component numerical value and units*

---

**Description**

Convert the combined dose to its individual component numerical value and units

**Usage**

```
get_doses_combination_units(the_string, separator = "/")
```

**Arguments**

the\_string      given combined unit  
separator      given character for separation , default is "/"

**Value**

separated numerical value and its units

**Examples**

```
get_doses_combination_units("10g/2ml")
```

---

```
get_eq5d_details
```

*Function to get the details of the EQ5D column*

---

**Description**

Function to get the details of the EQ5D column

**Usage**

```
get_eq5d_details(trialdata)
```

**Arguments**

trialdata      data containing individual level trial data

**Details**

Specific to the EQ5D data - the column names are given as certain sets, Tried to give 15 sets as the column names

**Value**

the name of the variable related to EQ5D and the unique contents if success, else error

**Examples**

```
get_eq5d_details(data.frame(
  "MO" = c(1, 2), "SC" = c(1, 2), "UA" = c(1, 2),
  "PD" = c(1, 2), "AD" = c(1, 2)
))
```

---

`get_extension_file`      *Function to get extension of a file name*

---

**Description**

Function to get extension of a file name

**Usage**

```
get_extension_file(filename)
```

**Arguments**

filename      name of a file

**Details**

if there is no "." character returns error else returns last characters those after string split using "."

**Value**

the extension

**Examples**

```
get_extension_file("data.txt")
```

---

`get_gender_details`      *Function to get the details of the gender column*

---

**Description**

Function to get the details of the gender column

**Usage**

```
get_gender_details(trialdata)
```

**Arguments**

`trialdata`      data containing individual level trial data

**Details**

expecting the data contains the information on gender preferably column names "gender", "sex" or "male" or "female". If multiple column names match these, then first match will be chosen.

**Value**

the name of the variable related to gender and the unique contents if success, else error

**Examples**

```
get_gender_details(data.frame("Age" = c(21, 15), "sex" = c("m", "f")))
```

---

`get_mean_sd_age`      *Function to return mean age from a data frame*

---

**Description**

Function to return mean age from a data frame

**Usage**

```
get_mean_sd_age(this_data, age_nr_code)
```

**Arguments**

`this_data`      the data containing column with age  
`age_nr_code`      non response code

**Details**

Age data is complete with the nr code given and get the mean and sd

**Value**

mean and sd, if success -1, if failure

**Examples**

```
this_data <- as.data.frame(cbind(num = c(1, 2, 3, 4),
age = c(14, 25, 26, 30)))
get_mean_sd_age(this_data, NA)
```

---

```
get_mortality_from_file
```

*Get the mortality rate values from reading a file*

---

**Description**

Get the mortality rate values from reading a file

**Usage**

```
get_mortality_from_file(paramfile, age, mortality_colname, gender = NULL)
```

**Arguments**

paramfile	parameter file to get the mortality eg.national life table data
age	age to get the age specific data
mortality_colname	column name with the mortality rates if it is not gender specific
gender	gender details to get the gender specific mortality data

**Details**

Provides the mortality rates as age and gender dependent Assumes the data contains mortality rate for single year and once it extracted per gender will retrieve single value Age column can consists of range of values, or a particular value also assumes that the mortality rate for each gender is listed under the gender column for gender specific values. if the mortality is not gender specific, the column name should be passed on to the function if gender is not null, mortality\_name will be ignored

**Value**

the paramvalue

**Examples**

```
paramfile <- system.file("extdata", "LifeTable_USA_Mx_2015.csv",
package = "packDAMipd"
)
a <- get_mortality_from_file(paramfile, age = 10, mortality_colname =
"total", gender = NULL)
```

---

```
get_name_value_probdistrb_def
```

*Function to return the two parameters from a given expression separated by comma,*

---

### Description

Function to return the two parameters from a given expression separated by comma,

### Usage

```
get_name_value_probdistrb_def(expr)
```

### Arguments

expr                    an expression

### Details

It will return the parameters of the distribution separated by commas and given in usual notation as brackets. It will identify those in between first occurrence of "(" and last occurrence of ")" and from the characters in between search for comma to indicate different parameters then it will extract (from those extracted parameters separated by commas) that on the left side of "equal" sign `get_name_value_probdistrb_def("gamma(mean = sqrt(2), b =17)")` will be ok but `get_name_value_probdistrb_def("gamma(shape, scale)")` and `get_name_value_probdistrb_def("gamma(shape =1 & scale =1)")` will show error

### Value

parameters in the expression expr

### Examples

```
get_name_value_probdistrb_def("gamma(mean = 10, sd =1)")
```

---

```
get_outcome_details    Function to get the details of the outcome column
```

---

### Description

Function to get the details of the outcome column

### Usage

```
get_outcome_details(trialdata, name, related_words, multiple = FALSE)
```



**Arguments**

trialdata	data containing individual level trial data
name	name of the variable
related_words	probable column names
multiple	indicates true if there are multiple columns

**Details**

if the words related to outcome is given, the function will get the columns and the codes used for the outcome, the difference here is that certain outcomes can be distributed in multiple columns

**Value**

the name of the variable related to health outcome (any) and the unique contents if success, else error

**Examples**

```
get_outcome_details(  
  data.frame("qo1.M0" = c(1, 2), "qo1.PD" = c(1, 2), "qo1.AD" = c(1, 2)),  
  "eq5d", "qo1", TRUE  
)
```

---

get\_parameter\_def\_distribution

*Get the definition of given parameter distribution defined in a file*

---

**Description**

Get the definition of given parameter distribution defined in a file

**Usage**

```
get_parameter_def_distribution(  
  parameter,  
  paramfile,  
  colnames_paramdistr,  
  strategycol = NA,  
  strategyname = NA  
)
```

**Arguments**

parameter	parameter of interest
paramfile	data file to be provided
colnames_paramdistr	list of column names for the parameters that define the distribution
strategycol	treatment strategy column name
strategyname	treatment strategy name in the column strategycol

**Details**

This function reads the parameter distribution from a file and return the parameter obtained This assumes that the file contains parameter, distribution colnames for parameter values for the distribution are passed on to the function assumes the name of each parameter and value are given in the consecutive columns. Once the expression is created using the parameters given in the file, it gets checked for correctness of specifying the distribution in R context using the function `check_estimate_substitute_proper_params` and then evaluated.

**Value**

the definition of parameter from the given distribution

**Examples**

```
paramfile <- system.file("extdata", "table_param.csv",
  package = "packDAMipd")
a <- get_parameter_def_distribution("rr", paramfile, c("Param1_name",
  "Param1_value"))
```

---

get\_parameter\_direct *Get the parameter values from reading a file*

---

**Description**

Get the parameter values from reading a file

**Usage**

```
get_parameter_direct(parameter, paramvalue)
```

**Arguments**

parameter	parameter of interest
paramvalue	parameter value to be assigned

**Details**

Basic function to assign a parameter directly

**Value**

the paramvalue

**Examples**

```
a <- get_parameter_direct("cost_IT", paramvalue = 100)
```

---

```
get_parameter_estimated_regression
```

*Get the parameter values using the provided statistical regression methods*

---

**Description**

Get the parameter values using the provided statistical regression methods

**Usage**

```
get_parameter_estimated_regression(  
  param_to_be_estimated,  
  data,  
  method,  
  indep_var,  
  info_get_method = NA,  
  info_distribution = NA,  
  covariates = NA,  
  timevar_survival = NA,  
  interaction = FALSE,  
  fix_eff = NA,  
  fix_eff_interact_vars = NA,  
  random_intercept_vars = NA,  
  nested_intercept_vars_pairs = NA,  
  cross_intercept_vars_pairs = NA,  
  uncorrel_slope_intercept_pairs = NA,  
  random_slope_intercept_pairs = NA,  
  naaction = "stats::na.omit",  
  param2_to_be_estimated = NA,  
  covariates2 = NA,  
  interaction2 = FALSE,  
  link = NA,  
  cluster_var = NA,  
  package_mixed_model = NA  
)
```

**Arguments**

<code>param_to_be_estimated</code>	parameter of interest
<code>data</code>	data to be provided or the data file containing dataset
<code>method</code>	method of estimation (for example, linear, logistic regression etc)
<code>indep_var</code>	the independent variable (column name in data file)
<code>info_get_method</code>	additional information on methods e.g Kaplan-Meier or hazard
<code>info_distribution</code>	distribution name eg. for logistic regression -binomial
<code>covariates</code>	list of covariates-calculations to be done before passing
<code>timevar_survival</code>	time variable for survival analysis
<code>interaction</code>	boolean value to indicate interaction in the case of linear regression
<code>fix_eff</code>	boolean value to indicate interaction in the case of linear regression
<code>fix_eff_interact_vars</code>	boolean value to indicate interaction in the case of linear regression
<code>random_intercept_vars</code>	boolean value to indicate interaction in the case of linear regression
<code>nested_intercept_vars_pairs</code>	boolean value to indicate interaction in the case of linear regression
<code>cross_intercept_vars_pairs</code>	boolean value to indicate interaction in the case of linear regression
<code>uncorrel_slope_intercept_pairs</code>	boolean value to indicate interaction in the case of linear regression
<code>random_slope_intercept_pairs</code>	boolean value to indicate interaction in the case of linear regression
<code>naaction</code>	what action to be taken for the missing values, default is a missing value.
<code>param2_to_be_estimated</code>	parameter of interest for equation 2 in bivariate regression
<code>covariates2</code>	list of covariates - for equation 2 in bivariate regression
<code>interaction2</code>	boolean value to indicate interaction for equation 2 in bivariate regression
<code>link</code>	link function to be provided if not using the default link for each of the <code>info_distribution</code>
<code>cluster_var</code>	cluster variable if any
<code>package_mixed_model</code>	package to be used for mixed model ie nlme or lme4

**Details**

This function is the top in the layer of functions used for regression analysis Thus it contains many parameters to be passed on The required ones are parameter to be estimated, data that contains the observation, the method of regression to be used, the independent variable and the information for the distribution and method. if the data is given as a file name. it will load the data in that file Then it calls the appropriate functions depending on the regression method that specified. The methods that are considered : Survival analysis, linear regression, logistic regression, generalised linear model, linear multilevel or mixed model, and seemingly unrelated regression

**Value**

results the results of the regression analysis

**Examples**

```
result <- get_parameter_estimated_regression(
  param_to_be_estimated = "Direction",
  data = ISLR::Smarket, method = "logistic", indep_var = "Lag1",
  info_get_method = NA, info_distribution = "binomial",
  covariates = c("Lag2", "Lag3"), interaction = FALSE,
  naaction = "na.omit", link = NA)
```

---

get\_parameter\_read      *Get the parameter values from reading a file*

---

**Description**

Get the parameter values from reading a file

**Usage**

```
get_parameter_read(parameter, paramfile, strategycol = NA, strategyname = NA)
```

**Arguments**

parameter	parameter of interest
paramfile	parameter file to be provided
strategycol	treatment strategy
strategyname	treatment strategy name in the column strategycol

**Details**

This function read the parameter from a file given that the file has these column names (at least) Parameter and Value Strategy col and name are optional. Check if the data file contains column names parameter and value and then get the results.

**Value**

the paramvalue

**Examples**

```
a <- get_parameter_read("cost_IT", paramfile = system.file("extdata",
"table_param.csv", package = "packDAMipd"))
```

---

```
get_single_col_multiple_pattern
```

*Function to get cols for the pattern given*

---

### Description

Function to get cols for the pattern given

### Usage

```
get_single_col_multiple_pattern(pattern, the_data)
```

### Arguments

pattern	the pattern to look for
the_data	data where to look at

### Value

zero or -1

### Examples

```
the_data <- as.data.frame(cbind(c("one", "two"), c("a", "b"), c("aa", "bb")))
colnames(the_data) <- c("name", "brand_one", "two")
get_single_col_multiple_pattern(c("brand", "trade"), the_data)
```

---

```
get_slope_intercept help function to keep slope and intercept portion ready in mixed model expression
```

---

### Description

help function to keep slope and intercept portion ready in mixed model expression

### Usage

```
get_slope_intercept(
  expression,
  random_intercept_vars,
  random_slope_intercept_pairs,
  uncorrel_slope_intercept_pairs
)
```

**Arguments**

expression      expression created so far  
 random\_intercept\_vars  
                  names of variables for random intercept  
 random\_slope\_intercept\_pairs  
                  random slopes intercept pairs this is a list of paired variables  
 uncorrel\_slope\_intercept\_pairs  
                  variables with correlated intercepts

**Value**

expression expression created

---

get\_slope\_intercept\_cross  
                  *help function to keep slope and intercept portion ready in mixed model  
 expression*

---

**Description**

help function to keep slope and intercept portion ready in mixed model expression

**Usage**

```
get_slope_intercept_cross(  
  expression,  
  random_intercept_vars,  
  intercept_vars_pairs,  
  random_slope_intercept_pairs,  
  uncorrel_slope_intercept_pairs  
)
```

**Arguments**

expression      expression created so far  
 random\_intercept\_vars  
                  names of variables for random intercept  
 intercept\_vars\_pairs  
                  those of the random intercept variables with nested effect  
 random\_slope\_intercept\_pairs  
                  random slopes intercept pairs this is a list of paired variables  
 uncorrel\_slope\_intercept\_pairs  
                  variables with correlated intercepts

**Value**

expression expression created

---

```
get_slope_intercept_nested
```

*help function to keep slope and intercept portion ready in mixed model expression*

---

### Description

help function to keep slope and intercept portion ready in mixed model expression

### Usage

```
get_slope_intercept_nested(
  expression,
  random_intercept_vars,
  intercept_vars_pairs,
  random_slope_intercept_pairs,
  uncorrel_slope_intercept_pairs
)
```

### Arguments

expression      expression created so far  
 random\_intercept\_vars  
                   names of variables for random intercept  
 intercept\_vars\_pairs  
                   those of the random intercept variables with nested effect  
 random\_slope\_intercept\_pairs  
                   random slopes intercept pairs this is a list of paired variables  
 uncorrel\_slope\_intercept\_pairs  
                   variables with correlated intercepts

### Value

expression expression created

---

```
get_timepoint_details Function to get the details of the time point column
```

---

### Description

Function to get the details of the time point column

### Usage

```
get_timepoint_details(trialdata)
```



**Arguments**

trialdata      data containing individual level trial data

**Details**

expecting the data contains the information on timepoints preferably column names "time point", "times" or "time" or "timepoint". If multiple column names match these, then first match will be chosen.

**Value**

the name of the variable related to time point and the unique contents if success, else error

**Examples**

```
get_timepoint_details(data.frame("time" = c(21, 15),  
"arm" = c("control", "intervention")))
```

---

get\_trial\_arm\_details    *Function to get the details of the trial arm*

---

**Description**

Function to get the details of the trial arm

**Usage**

```
get_trial_arm_details(trialdata)
```

**Arguments**

trialdata      data containing individual level trial data

**Details**

expecting the data contains the information on trial arm preferably column names "arm", "trial" or "trial arm". If multiple column names match these, then first match will be chosen.

**Value**

the name of the variable related to trial arm and the unique contents if success, else error

**Examples**

```
get_trial_arm_details(data.frame(  
  "Age" = c(21, 15),  
  "arm" = c("control", "intervention")  
)
```

---

get_var_state	<i>Get the attribute for the health state</i>
---------------	---

---

**Description**

Get the attribute for the health state

**Usage**

```
get_var_state(state, var)
```

**Arguments**

state	object of class health state
var	attribute of the health state

**Details**

After checking the given state is a health state and given variable is defined in the health state, the value of the variable is returned

**Value**

modified health state

**Examples**

```
get_var_state(health_state("IT", 100, 0.4, 0, FALSE), "cost")
```

---

health_state	<i>Definition of health state class or health state constructor</i>
--------------	---

---

**Description**

Definition of health state class or health state constructor

**Usage**

```
health_state(name, cost, utility, state_time = 0, absorb = FALSE)
```

**Arguments**

name	name of the health state
cost	value or expression that represents cost of the health state
utility	value or expression that represents utility of the health state
state_time	time denoting how long in the state
absorb	boolean indicating health state absorbing or not

**Details**

Initialising the name, cost, utility and time spent for the health state name is the name of the health state cost/utility can be defined as characters e.g. "cost\_A" if they are characters, the value is assigned after parsing the text. state\_time is integer and absorb is boolean

**Value**

value of the state

**Examples**

```
st <- health_state("IT", 100, 0.4, 0, FALSE)
st <- health_state("IT", "cost_A", 0.4, 0, FALSE)
```

---

init_trace	<i>Define an all zero trace matrix</i>
------------	--

---

**Description**

Define an all zero trace matrix

**Usage**

```
init_trace(health_states, cycles)
```

**Arguments**

health_states	health states
cycles	no of cycles

**Details**

Initialise the trace matrix with all zeros trace matrix will be with no\_cycles+1 by no\_states matrix

**Value**

trace matrix -all zero

**Examples**

```
a <- health_state("Healthy", 1, 1, 0, FALSE)
b <- health_state("Dead", 1, 0.5, 0, FALSE)
health_states <- combine_state(a, b)
init_trace(health_states, 10)
```

---

init\_trace\_sjtime      *Define an all zero trace matrix*

---

**Description**

Define an all zero trace matrix

**Usage**

```
init_trace_sjtime(health_states, cycles)
```

**Arguments**

health\_states    health states  
cycles            no of cycles

**Details**

Initialise the trace matrix with all zeros trace matrix will be with no\_cycles+1 by no\_states matrix

**Value**

trace matrix -all zero

**Examples**

```
a <- health_state("Healthy", 1, 1, 0, FALSE)
b <- health_state("Dead", 1, 0.5, 0, FALSE)
health_states <- combine_state(a, b)
init_trace(health_states, 10)
```

---

keep\_results\_plot\_dsa    *Function to do some checks before plotting sensitivity analysis results*

---

**Description**

Function to do some checks before plotting sensitivity analysis results

**Usage**

```
keep_results_plot_dsa(
  result_dsa_control,
  plotfor,
  result_dsa_treat,
  plot_variable,
  threshold,
  comparator
)
```

**Arguments**

result_dsa_control	result from deterministic sensitivity analysis for first or control model
plotfor	the variable to plotfor e.g. cost, utility NMB etc
result_dsa_treat	result from deterministic sensitivity analysis for the comparative Markov model
plot_variable	variable for plotting
threshold	threshold value of WTP
comparator	the strategy to be compared with

**Value**

results to plot dsa

---

list\_paramwise\_psa\_result

*Function to list probabilistic sensitivity analysis results parameterwise*

---

**Description**

Function to list probabilistic sensitivity analysis results parameterwise

**Usage**

```
list_paramwise_psa_result(
  result_psa_params_control,
  result_psa_params_treat,
  threshold,
  comparator
)
```

**Arguments**

result_psa_params_control	result from probabilistic sensitivity analysis for first or control model
result_psa_params_treat	result from probabilistic sensitivity analysis for the comparative Markov model
threshold	threshold value of WTP
comparator	the strategy to be compared with

**Value**

plot of sensitivity analysis

**Examples**

```

param_list <- define_parameters(
  cost_zido = 2278, cost_direct_med_A = 1701,
  cost_comm_care_A = 1055, cost_direct_med_B = 1774,
  cost_comm_care_B = 1278,
  cost_direct_med_C = 6948, cost_comm_care_C = 2059,
  tpAtoA = 1251 / (1251 + 483),
  tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),
  tpAtoD = 17 / (17 + 1717),
  tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),
  tpBtoD = 15 / (15 + 1243),
  tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312), tpDtoD = 1,
  cost_health_A = "cost_direct_med_A + cost_comm_care_A",
  cost_health_B = "cost_direct_med_B + cost_comm_care_B",
  cost_health_C = "cost_direct_med_C + cost_comm_care_C",
  cost_drug = "cost_zido"
)
A <- health_state("A", cost = "cost_health_A + cost_drug ", utility = 1)
B <- health_state("B", cost = "cost_health_B + cost_drug", utility = 1)
C <- health_state("C", cost = "cost_health_C + cost_drug", utility = 1)
D <- health_state("D", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3, 4), c(NA, 5, 6, 7), c(NA, NA, 8, 9),
c(NA, NA, NA, 10))
colnames(tmat) <- rownames(tmat) <- c("A", "B", "C", "D")
tm <- populate_transition_matrix(4, tmat, c(
  "tpAtoA", "tpAtoB", "tpAtoC", "tpAtoD",
  "tpBtoB", "tpBtoC", "tpBtoD", "tpCtoC", "tpCtoD", "tpDtoD"
), colnames(tmat))
health_states <- combine_state(A, B, C, D)
mono_strategy <- strategy(tm, health_states, "mono")
mono_markov <- markov_model(mono_strategy, 20, initial_state = c(1,0,0,0),
discount = c(0.06, 0),param_list)
sample_list <- define_parameters(cost_zido = "gamma(mean = 2756,
sd = sqrt(2756))")
param_table <- define_parameters_psa(param_list, sample_list)
result <- do_psa(mono_markov, param_table, 10)
list_paramwise_psa_result(result, NULL, NULL, NULL)

```

---

load\_trial\_data

*Function to load the file containing trial data and return it*


---

**Description**

Function to load the file containing trial data and return it

**Usage**

```
load_trial_data(file = NULL, sheet = NULL)
```

**Arguments**

file                    name of the file in full  
sheet                   name of the sheet if excel work book is given

**Value**

trial data if success, else -1

**Examples**

```
load_trial_data(system.file("extdata", "trial_data.csv",  
  package = "packDAMipd"  
))
```

---

map\_eq5d5Lto3L\_VanHout

*Function to map EQ5D5L scores to EQ5D3L scores and then add to IPD data*

---

**Description**

Function to map EQ5D5L scores to EQ5D3L scores and then add to IPD data

**Usage**

```
map_eq5d5Lto3L_VanHout(ind_part_data, eq5d_nrcode)
```

**Arguments**

ind\_part\_data    a data frame  
eq5d\_nrcode      non response code for EQ5D5L, default is NA

**Value**

qaly included modified data, if success -1, if failure

**Source**

[http://eprints.whiterose.ac.uk/121473/1/Devlin\\_et\\_al-2017-Health\\_Economics.pdf](http://eprints.whiterose.ac.uk/121473/1/Devlin_et_al-2017-Health_Economics.pdf)

**Examples**

```
library(valueEQ5D)  
datafile <- system.file("extdata", "trial_data.csv",  
  package = "packDAMipd")  
trial_data <- load_trial_data(datafile)  
map_eq5d5Lto3L_VanHout(trial_data, NA)
```

---

map\_resource\_use\_categories

*Function to to read the text form of resource use and replace it with standard texts of resoure use ie. some one can describe GP visit as GP surgery visit, surgery visit or general practioners visit etc. Here all these texts should be given in a excel or csv file and then corresponidng standard form will be read from the file and will be replaced.*

---

### Description

Function to to read the text form of resource use and replace it with standard texts of resoure use ie. some one can describe GP visit as GP surgery visit, surgery visit or general practioners visit etc. Here all these texts should be given in a excel or csv file and then corresponidng standard form will be read from the file and will be replaced.

### Usage

```
map_resource_use_categories(
  the_data,
  service_actual,
  new_column,
  mapped_data,
  mapped_use,
  analysis,
  replace_only,
  relevant_column = NULL,
  check_value_relevant = NULL,
  nhs_use_column = NULL,
  check_value_nhs_use = NULL
)
```

### Arguments

the_data	the data where the observations are held
service_actual	columna name of the actual service use
new_column	the name of the column where the mapped resource use to be
mapped_data	data where the service name and mapped service name has been stored
mapped_use	columnan name of mapped resource use in mapped_data
analysis	base case or secondary
replace_only	if we want to replace only certain resource use
relevant_column	the name of the column where the mapped resource use is indicated as relevant or not
check_value_relevant	how is the mapped resource is indicated as relevant by a value



nhs\_use\_column the name of the column where the mapped resource use comes under NHS or not  
 check\_value\_nhs\_use value that is used to indicated the nhs use

### Value

the data with added sum

---

markov_model	<i>Definition of Markov model and trace</i>
--------------	---

---

### Description

Definition of Markov model and trace

### Usage

```
markov_model(  
  current_strategy,  
  cycles,  
  initial_state,  
  discount = c(0, 0),  
  parameter_values = NULL,  
  half_cycle_correction = TRUE,  
  state_cost_only_prevalent = FALSE,  
  state_util_only_prevalent = FALSE,  
  method = "half cycle correction",  
  startup_cost = NULL,  
  startup_util = NULL  
)
```

### Arguments

current\_strategy strategy object  
 cycles no of cycles  
 initial\_state value of states initially  
 discount rate of discount for costs and qalys  
 parameter\_values parameters for assigning health states and probabilities  
 half\_cycle\_correction boolean to indicate half cycle correction

state_cost_only_prevalent	boolean parameter to indicate if the costs for state occupancy is only for those in the state excluding those that transitioned new. This is relevant when the transition cost is provided for eg. in a state with dialysis the cost of previous dialysis is different from the newly dialysis cases. Then the state_cost_only_prevalent should be TRUE
state_util_only_prevalent	boolean parameter to indicate if the utilities for state occupancy is only for those in the state excluding those that transitioned new.
method	what type of half cycle correction needed
startup_cost	cost of states initially
startup_util	utility of states initially if any

### Details

Use the strategy, cycles, initial state values creating the markov model and trace. As many probabilities /cost/utility value depend on age/time the evaluation and assignment happens during each cycle. At the heart it does a matrix multiplication using the previous row of the trace matrix and the columns of the transition matrix. Also checks for population loss, calculates cumulative costs and qalys (accounts for discounting and half cycle correction)

### Value

Markov trace

### Examples

```
tmat <- rbind(c(1, 2), c(3, 4))
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")
tm <- populate_transition_matrix(2, tmat, c(0.5, 0.5, 0, 1))
a <- health_state("Healthy", 1, 1, 0, FALSE)
b <- health_state("Dead", 1, 0, 0, TRUE)
health_states <- combine_state(a, b)
this.strategy <- strategy(tm, health_states, "intervention")
markov_model(this.strategy, 10, c(1, 0))
```

---

markov\_model\_sojourntime

*Definition of Markov model and trace*

---

### Description

Definition of Markov model and trace

**Usage**

```

markov_model_sojourntime(
  current_strategy,
  nCycles,
  initial_state,
  discount = c(0, 0),
  parameter_values = NULL,
  half_cycle_correction = TRUE,
  state_cost_only_prevalent = FALSE,
  state_util_only_prevalent = FALSE,
  method = "half cycle correction",
  startup_cost = NULL,
  startup_util = NULL
)

```

**Arguments**

current_strategy	strategy object
nCycles	no of cycles
initial_state	value of states initially
discount	rate of discount for costs and qalys
parameter_values	parameters for assigning health states and probabilities
half_cycle_correction	boolean to indicate half cycle correction
state_cost_only_prevalent	boolean parameter to indicate if the costs for state occupancy is only for those in the state excluding those that transitioned new. This is relevant when the transition cost is provided for eg. in a state with dialysis the cost of previous dialysis is different from the newly dialysis cases. Then the state_cost_only_prevalent should be TRUE
state_util_only_prevalent	boolean parameter to indicate if the utilities for state occupancy is only for those in the state excluding those that transitioned new.
method	what type of half cycle correction needed
startup_cost	cost of states initially
startup_util	utility of states initially if any

**Details**

Use the strategy, cycles, initial state values creating the markov model and trace. As many probabilities /cost/utility value depend on age/time the evaluation and assignment happens during each cycle. At the heart it does a matrix multiplication using the previous row of the trace matrix and the columns of the transition matrix. Also checks for population loss, calculates cumulative costs and qalys (accounts for discounting and half cycle correction)

**Value**

Markov trace

**Examples**

```
tmat <- rbind(c(1, 2), c(3, 4))
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")
tm <- populate_transition_matrix(2, tmat, c(0.5, 0.5, 0, 1))
a <- health_state("Healthy", 1, 1, 0, FALSE)
b <- health_state("Dead", 1, 0, 0, TRUE)
health_states <- combine_state(a, b)
this.strategy <- strategy(tm, health_states, "intervention")
markov_model(this.strategy, 10, c(1, 0))
```

---

microcosting\_liquids\_long

*Function to estimate the cost of liquids when IPD is in long format*

---

**Description**

Function to estimate the cost of liquids when IPD is in long format

**Usage**

```
microcosting_liquids_long(
  the_columns,
  ind_part_data_long,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  bottle_size,
  bottle_size_unit = NULL,
  bottle_last,
  bottle_last_unit = NULL,
  preparation_dose = NULL,
  preparation_unit = NULL,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_brand = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_bottle_size_unit = NULL,
  list_of_code_bottle_last_unit = NULL,
```

```

    list_preparation_dose_unit = NULL,
    eqdose_covtab = NULL,
    basis_strength_unit = NULL
)

```

### Arguments

```

the_columns      columns that are to be used to convert the data from long to wide
ind_part_data_long
                  IPD
name_med         name of medication
brand_med        brand name of medication if revealed
dose_med         dose of medication used
unit_med         unit of medication ; use null if its along with the dose
bottle_size     size of the bottle used
bottle_size_unit
                  unit of bottle volume
bottle_last     how long the bottle lasted
bottle_last_unit
                  time unit of how long the bottle lasted
preparation_dose
                  dose if preparation is given
preparation_unit
                  unit of preparatio dose
timeperiod       time period for cost calculation
unit_cost_data  unit costs data
unit_cost_column
                  column name of unit cost in unit_cost_data
cost_calculated_per
                  column name of unit where the cost is calculated
strength_column
                  column column name that has strength of medication
list_of_code_names
                  if names is coded, give the code:name pairs, optional
list_of_code_brand
                  if brand names are coded, give the code:brand pairs, optional
list_of_code_dose_unit
                  if unit is coded, give the code:unit pairs, optional
list_of_code_bottle_size_unit
                  list of bottle size units and codes
list_of_code_bottle_last_unit
                  list of time of bottle lasts and codes
list_preparation_dose_unit
                  list of preparation dose units and codes
eqdose_covtab   table to get the conversion factor for equivalent doses, optional
basis_strength_unit
                  strength unit to be taken as basis required for total medication calculations

```

**Value**

the calculated cost of tablets along with original data

**Examples**

```
med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
  package = "packDAMipd")
data_file <- system.file("extdata", "medication_liq.xlsx",
  package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx",
  package = "packDAMipd")
table <- load_trial_data(conv_file)
names <- colnames(ind_part_data)
ending <- length(names)
ind_part_data_long <- tidyr::gather(ind_part_data, measurement, value,
  names[2]:names[ending], factor_key = TRUE)
the_columns <- c("measurement", "value")
res <- microcosting_liquids_long(the_columns,
  ind_part_data_long = ind_part_data_long,
  name_med = "liq_name", brand_med = NULL, dose_med = "liq_strength",
  unit_med = NULL, bottle_size = "liq_bottle_size",bottle_size_unit = NULL,
  bottle_last = "liq_last",bottle_last_unit = NULL,preparation_dose = NULL,
  preparation_unit = NULL,timeperiod = "4 months",unit_cost_data = med_costs,
  unit_cost_column = "UnitCost",cost_calculated_per = "Basis",
  strength_column = "Strength",list_of_code_names = NULL,
  list_of_code_brand = NULL,list_of_code_dose_unit = NULL,
  list_of_code_bottle_size_unit = NULL,list_of_code_bottle_last_unit = NULL,
  list_preparation_dose_unit = NULL,eqdose_covtab = table,
  basis_strength_unit = NULL)
```

---

microcosting\_liquids\_wide

*Function to estimate the cost of liquids taken (from IPD)*

---

**Description**

Function to estimate the cost of liquids taken (from IPD)

**Usage**

```
microcosting_liquids_wide(
  ind_part_data,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  bottle_size,
```

```
bottle_size_unit = NULL,  
bottle_last,  
bottle_last_unit = NULL,  
preparation_dose = NULL,  
preparation_unit = NULL,  
timeperiod,  
unit_cost_data,  
unit_cost_column,  
cost_calculated_per,  
strength_column,  
list_of_code_names = NULL,  
list_of_code_brand = NULL,  
list_of_code_dose_unit = NULL,  
list_of_code_bottle_size_unit = NULL,  
list_of_code_bottle_last_unit = NULL,  
list_preparation_dose_unit = NULL,  
eqdose_covtab = NULL,  
basis_strength_unit = NULL  
)
```

### Arguments

ind_part_data	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
bottle_size	size of the bottle used
bottle_size_unit	unit of bottle volume
bottle_last	how long the bottle lasted
bottle_last_unit	time unit of how long the bottle lasted
preparation_dose	dose if preparation is given
preparation_unit	unit of preparatio dose
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data
cost_calculated_per	column name of unit where the cost is calculated
strength_column	column column name that has strength of medication

list\_of\_code\_names  
if names is coded, give the code:name pairs, optional

list\_of\_code\_brand  
if brand names are coded, give the code:brand pairs, optional

list\_of\_code\_dose\_unit  
if unit is coded, give the code:unit pairs, optional

list\_of\_code\_bottle\_size\_unit  
list of bottle size units and codes

list\_of\_code\_bottle\_last\_unit  
list of time of bottle lasts and codes

list\_preparation\_dose\_unit  
list of preparation dose units and codes

eqdose\_covtab table to get the conversion factor for equivalent doses, optional, but the column names have to be unique Similar to c("Drug", "form", "unit", "factor") or c("Drug", "form", "unit", "conversion")

basis\_strength\_unit  
strength unit to be taken as basis required for total medication calculations

**Value**

the calculated cost of tablets along with original data

**Examples**

```
med_costs_file <- system.file("extdata", "medication_costs_all.xlsx",
package = "packDAMipd")
data_file <- system.file("extdata", "medication_liq.xlsx",
package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx", package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- microcosting_liquids_wide(
ind_part_data = ind_part_data, name_med = "liq_name", brand_med = NULL,
dose_med = "liq_strength", unit_med = NULL, bottle_size = "liq_bottle_size",
bottle_size_unit = NULL, bottle_last_unit = "liq_last_unit",
bottle_last_unit = NULL, preparation_dose = NULL, preparation_unit = NULL,
timeperiod = "4 months", unit_cost_data = med_costs,
unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
strength_column = "Strength", list_of_code_names = NULL,
list_of_code_brand = NULL, list_of_code_dose_unit = NULL,
list_of_code_bottle_size_unit = NULL, list_of_code_bottle_last_unit = NULL,
list_preparation_dose_unit = NULL, eqdose_covtab = table,
basis_strength_unit = NULL)
```



---

```

microcosting_patches_long
#'#####
  Function to estimate the cost of patches when IPD is in long format
  using a IPD data of long format

```

---

## Description

```

#'##### Function
  to estimate the cost of patches when IPD is in long format using a IPD data of long format

```

## Usage

```

microcosting_patches_long(
  the_columns,
  ind_part_data_long,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

## Arguments

the_columns	columns that are to be used to convert the data from long to wide
ind_part_data_long	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken

freq\_taken        frequency of medication  
 timeperiod       time period for cost calculation  
 unit\_cost\_data   unit costs data  
 unit\_cost\_column        column name of unit cost in unit\_cost\_data  
 cost\_calculated\_per        column name of unit in the cost is calculated  
 strength\_column        column column name that contain strength of medication  
 list\_of\_code\_names        if names is coded, give the code:name pairs, optional  
 list\_of\_code\_freq        if frequency is coded, give the code:frequency pairs, optional  
 list\_of\_code\_dose\_unit        if unit is coded, give the code:unit pairs, optional  
 list\_of\_code\_brand        if brand names are coded, give the code:brand pairs, optional  
 eqdose\_cov\_tab    table to get the conversion factor for equivalent doses, optional  
 basis\_strength\_unit        strength unit to be taken as basis required for total medication calculations

### Value

the calculated cost of tablets along with original data

### Examples

```

med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
package = "packDAMipd")
data_file <- system.file("extdata", "medication.xlsx",
package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx",package = "packDAMipd")
table <- load_trial_data(conv_file)
names <- colnames(ind_part_data)
ending <- length(names)
ind_part_data_long <- tidyr::gather(ind_part_data, measurement, value,
names[2]:names[ending], factor_key = TRUE)
the_columns <- c("measurement", "value")
res <- microcosting_patches_long(the_columns,
ind_part_data_long = ind_part_data_long, name_med = "patch_name",
brand_med = "patch_brand", dose_med = "patch_strength",unit_med = NULL,
no_taken = "patch_no_taken", freq_taken = "patch_frequency",
timeperiod = "4 months", unit_cost_data = med_costs,
unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
strength_column = "Strength", list_of_code_names = NULL,
list_of_code_freq = NULL, list_of_code_dose_unit = NULL,
list_of_code_brand = NULL, eqdose_cov_tab = table,
basis_strength_unit = "mcg/hr")

```

---

 microcosting\_patches\_wide

*Function to estimate the cost of patches taken (from IPD)*


---

### Description

Function to estimate the cost of patches taken (from IPD)

### Usage

```

microcosting_patches_wide(
  ind_part_data,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

### Arguments

ind_part_data	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken
freq_taken	frequency of medication
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data

`cost_calculated_per`  
                   column name of unit where the cost is calculated  
`strength_column`  
                   column column name that contain strength of medication  
`list_of_code_names`  
                   if names is coded, give the code:name pairs, optional  
`list_of_code_freq`  
                   if frequency is coded, give the code:frequency pairs, optional  
`list_of_code_dose_unit`  
                   if unit is coded, give the code:unit pairs, optional  
`list_of_code_brand`  
                   if brand names are coded, give the code:brand pairs, optional  
`eqdose_cov_tab` table to get the conversion factor for equivalent doses, optional, but the column  
                   names have to be unique Similar to `c("Drug", "form", "unit", "factor")` or `c("Drug",  
                   "form", "unit", "conversion")`  
`basis_strength_unit`  
                   strength unit to be taken as basis required for total medication calculations

### Details

Assumes individual level data has name of medication, dose, dose unit, number taken, frequency taken, and basis time Assumes unit cost data contains the name of medication, form/type, strength, unit of strength (or the unit in which the cost calculated), preparation, unit cost, size and size unit (in which name, forms, size, size unit, and preparation are not passed on) @importFrom dplyr %>%  
 a patient use 1 mg/hr patches 5 patches once a week that patch comes in a pack of 4 with cost £2.50  
 we want to estimate the cost for 3 months that means amount of medication 3 months = 21 weeks  
 number of patches taken =  $21 \times 5 = 105$  patches packs =  $(105/4)$  almost 27 packs cost = 272.50

### Value

the calculated cost of tablets along with original data

### Examples

```

med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
  package = "packDAMipd")
data_file <- system.file("extdata", "medication.xlsx",
  package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx", package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- microcosting_patches_wide(
  ind_part_data = ind_part_data, name_med = "patch_name",
  brand_med = "patch_brand", dose_med = "patch_strength", unit_med = NULL,
  no_taken = "patch_no_taken", freq_taken = "patch_frequency",
  timeperiod = "4 months", unit_cost_data = med_costs,
  unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
  strength_column = "Strength", list_of_code_names = NULL,

```

```
list_of_code_freq = NULL, list_of_code_dose_unit = NULL,
list_of_code_brand = NULL, eqdose_cov_tab = table,
basis_strength_unit = "mcg/hr")
```

---

```
microcosting_tablets_long
```

*Function to estimate the cost of tablets when IPD is in long format*

---

## Description

Function to estimate the cost of tablets when IPD is in long format

## Usage

```
microcosting_tablets_long(
  the_columns,
  ind_part_data_long,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)
```

## Arguments

the_columns	columns that are to be used to convert the data from long to wide
ind_part_data_long	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken

freq\_taken      frequency of medication  
 timeperiod      time period for cost calculation  
 unit\_cost\_data   unit costs data  
 unit\_cost\_column  
                  column name of unit cost in unit\_cost\_data  
 cost\_calculated\_per  
                  column name of unit where the cost is calculated  
 strength\_column  
                  column column name that contain strength of medication  
 list\_of\_code\_names  
                  if names is coded, give the code:name pairs, optional  
 list\_of\_code\_freq  
                  if frequency is coded, give the code:frequency pairs, optional  
 list\_of\_code\_dose\_unit  
                  if unit is coded, give the code:unit pairs, optional  
 list\_of\_code\_brand  
                  if brand names are coded, give the code:brand pairs, optional  
 eqdose\_cov\_tab   table to get the conversion factor for equivalent doses, optional  
 basis\_strength\_unit  
                  strength unit to be taken as basis required for total medication calculations

### Value

the calculated cost of tablets along with original data

### Examples

```

med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
package = "packDAMipd")
data_file <- system.file("extdata", "medication_all.xlsx",
package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx", package = "packDAMipd")
table <- load_trial_data(conv_file)
names <- colnames(ind_part_data)
ending <- length(names)
ind_part_data_long <- tidyr::gather(ind_part_data, measurement, value,
names[2]:names[ending], factor_key = TRUE)
the_columns <- c("measurement", "value")
res <- microcosting_tablets_long(the_columns,
ind_part_data_long = ind_part_data_long, name_med = "tab_name",
brand_med = "tab_brand", dose_med = "tab_strength",
unit_med = "tab_str_unit",
no_taken = "tab_no_taken", freq_taken = "tab_frequency",
timeperiod = "2 months", unit_cost_data = med_costs,
unit_cost_column = "UnitCost", cost_calculated_per = "Basis",
strength_column = "Strength", list_of_code_names = NULL,
list_of_code_freq = NULL, list_of_code_dose_unit = NULL,
eqdose_cov_tab = table, basis_strength_unit = "mg")

```

---

 microcosting\_tablets\_wide

*Function to estimate the cost of tablets taken (from IPD)*


---

### Description

Function to estimate the cost of tablets taken (from IPD)

### Usage

```

microcosting_tablets_wide(
  ind_part_data,
  name_med,
  brand_med = NULL,
  dose_med,
  unit_med = NULL,
  no_taken,
  freq_taken,
  timeperiod,
  unit_cost_data,
  unit_cost_column,
  cost_calculated_per,
  strength_column,
  list_of_code_names = NULL,
  list_of_code_freq = NULL,
  list_of_code_dose_unit = NULL,
  list_of_code_brand = NULL,
  eqdose_cov_tab = NULL,
  basis_strength_unit = NULL
)

```

### Arguments

ind_part_data	IPD
name_med	name of medication
brand_med	brand name of medication if revealed
dose_med	dose of medication used
unit_med	unit of medication ; use null if its along with the dose
no_taken	how many taken
freq_taken	frequency of medication
timeperiod	time period for cost calculation
unit_cost_data	unit costs data
unit_cost_column	column name of unit cost in unit_cost_data

`cost_calculated_per`  
                   column name of unit where the cost is calculated  
`strength_column`  
                   column column name that contain strength of medication  
`list_of_code_names`  
                   if names is coded, give the code:name pairs, optional  
`list_of_code_freq`  
                   if frequency is coded, give the code:frequency pairs, optional  
`list_of_code_dose_unit`  
                   if unit is coded, give the code:unit pairs, optional  
`list_of_code_brand`  
                   if brand names are coded, give the code:brand pairs, optional  
`eqdose_cov_tab` table to get the conversion factor for equivalent doses, optional, but the column  
                   names have to be unique Similar to `c("Drug", "form", "unit", "factor")` or `c("Drug",  
                   "form", "unit", "conversion")`  
`basis_strength_unit`  
                   strength unit to be taken as basis required for total medication calculations

### Details

Assumes individual level data has name of medication, dose, dose unit, number taken, frequency taken, and basis time Assumes unit cost data contains the name of medication, form/type, strength, unit of strength (or the unit in which the cost calculated), preparation, unit cost, size and size unit (in which name, forms, size, size unit, and preparation are not passed on) `@importFrom dplyr %>%`

### Value

the calculated cost of tablets along with original data

### Examples

```

med_costs_file <- system.file("extdata", "medicaton_costs_all.xlsx",
package = "packDAMipd")
data_file <- system.file("extdata", "medication_all.xlsx",
package = "packDAMipd")
ind_part_data <- load_trial_data(data_file)
med_costs <- load_trial_data(med_costs_file)
conv_file <- system.file("extdata", "Med_calc.xlsx",package = "packDAMipd")
table <- load_trial_data(conv_file)
res <- microcosting_tablets_wide(ind_part_data = ind_part_data,
name_med = "tab_name", brand_med = "tab_brand", dose_med = "tab_strength",
unit_med = "tab_str_unit", no_taken = "tab_no_taken",
freq_taken = "tab_frequency",timeperiod = "2 months",
unit_cost_data = med_costs,unit_cost_column = "UnitCost",
cost_calculated_per = "Basis", strength_column = "Strength",
list_of_code_names = NULL, list_of_code_freq = NULL,
list_of_code_dose_unit = NULL, eqdose_cov_tab = table,
basis_strength_unit = "mg")
  
```



---

plot_ceac_psa	<i>Function to plot CEAC</i>
---------------	------------------------------

---

**Description**

Function to plot CEAC

**Usage**

```
plot_ceac_psa(  
  control_markov,  
  trt_markov,  
  psa_table_ctrl,  
  psa_table_trt,  
  thresholds,  
  num_rep,  
  comparator  
)
```

**Arguments**

control_markov	markov model for control
trt_markov	markov model for treatment
psa_table_ctrl	param table for psa for control
psa_table_trt	param table for psa for treatment
thresholds	threshold values of WTP
num_rep	number of repetitions
comparator	the strategy to be compared with

**Value**

plot of ceac

---

plot_dsa	<i>Function to plot results of sensitivity analysis do_sensitivity_analysis()</i>
----------	---

---

**Description**

Function to plot results of sensitivity analysis do\_sensitivity\_analysis()

**Usage**

```
plot_dsa(
  result_dsa_control,
  plotfor,
  type = "range",
  result_dsa_treat = NULL,
  threshold = NULL,
  comparator = NULL
)
```

**Arguments**

result_dsa_control	result from deterministic sensitivity analysis for first or control model
plotfor	the variable to plotfor e.g. cost, utility NMB etc
type	type of analysis, range or difference
result_dsa_treat	result from deterministic sensitivity analysis for the comparative Markov model
threshold	threshold value of WTP
comparator	the strategy to be compared with

**Value**

plot of sensitivity analysis

**Examples**

```
param_list <- define_parameters(
  cost_zido = 2278, cost_direct_med_A = 1701,
  cost_comm_care_A = 1055, cost_direct_med_B = 1774,
  cost_comm_care_B = 1278,
  cost_direct_med_C = 6948, cost_comm_care_C = 2059,
  tpAtoA = 1251 / (1251 + 483),
  tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),
  tpAtoD = 17 / (17 + 1717),
  tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),
  tpBtoD = 15 / (15 + 1243),
  tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312),
  tpDtoD = 1,
  cost_health_A = "cost_direct_med_A + cost_comm_care_A",
  cost_health_B = "cost_direct_med_B + cost_comm_care_B",
  cost_health_C = "cost_direct_med_C + cost_comm_care_C",
  cost_drug = "cost_zido")
low_values <- define_parameters(cost_direct_med_B = 177.4,
  cost_comm_care_C = 205.9)
upp_values <- define_parameters(cost_direct_med_B = 17740,
  cost_comm_care_C = 20590)
A <- health_state("A", cost = "cost_health_A + cost_drug ",
  utility = 1)
```

```

B <- health_state("B", cost = "cost_health_B + cost_drug",
utility = 1)
C <- health_state("C", cost = "cost_health_C + cost_drug",
utility = 1)
D <- health_state("D", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3, 4), c(NA, 5, 6, 7), c(NA, NA, 8, 9),
c(NA, NA, NA, 10))
colnames(tmat) <- rownames(tmat) <- c("A", "B", "C", "D")
tm <- populate_transition_matrix(4, tmat, c("tpAtoA", "tpAtoB", "tpAtoC",
"tpAtoD", "tpBtoB", "tpBtoC", "tpBtoD", "tpCtoC", "tpCtoD", "tpDtoD"),
colnames(tmat))
health_states <- combine_state(A, B, C, D)
mono_strategy <- strategy(tm, health_states, "mono")
mono_markov <- markov_model(mono_strategy, 20, c(1, 0, 0, 0),
discount = c(0.06, 0), param_list)
param_table <- define_parameters_sens_anal(param_list, low_values,
upp_values)
result <- do_sensitivity_analysis(mono_markov, param_table)
param_list_treat <- define_parameters(
cost_zido = 3000, cost_direct_med_A = 890,
cost_comm_care_A = 8976, cost_direct_med_B = 2345,
cost_comm_care_B = 1278,
cost_direct_med_C = 6948, cost_comm_care_C = 2059,
tpAtoA = 1251 / (1251 + 483),
tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),
tpAtoD = 17 / (17 + 1717),
tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),
tpBtoD = 15 / (15 + 1243),
tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312),
tpDtoD = 1,
cost_health_A = "cost_direct_med_A + cost_comm_care_A",
cost_health_B = "cost_direct_med_B + cost_comm_care_B",
cost_health_C = "cost_direct_med_C + cost_comm_care_C",
cost_drug = "cost_zido")
treat_strategy <- strategy(tm, health_states, "treat")
treat_markov <- markov_model(treat_strategy, 20, c(1, 0, 0, 0),
discount = c(0.06, 0), param_list_treat)
treat_low_values <- define_parameters(cost_direct_med_B = 234.5,
cost_comm_care_C = 694.8)
treat_upp_values <- define_parameters(cost_direct_med_B = 23450,
cost_comm_care_C = 69480)
param_table_treat <- define_parameters_sens_anal(param_list_treat,
treat_low_values, treat_upp_values)
result_treat <- do_sensitivity_analysis(treat_markov, param_table)
plot_dsa(result, "NMB", "range", result_treat, 20000, "treat")

```

**Description**

Function to do some checks before plotting sensitivity analysis results

**Usage**

```
plot_dsa_difference(ob_results, plotfor, plot_var)
```

**Arguments**

<code>ob_results</code>	results from deterministic sensitivity analysis
<code>plotfor</code>	the quantity plotting
<code>plot_var</code>	the variable

**Value**

plot

---

`plot_dsa_icer_range`    *Function to do some checks before plotting sensitivity analysis results*

---

**Description**

Function to do some checks before plotting sensitivity analysis results

**Usage**

```
plot_dsa_icer_range(ob_results, plot_var)
```

**Arguments**

<code>ob_results</code>	results from deterministic sensitivity analysis
<code>plot_var</code>	the variable

**Value**

plot

---

plot\_dsa\_nmb\_range     *Function to do some checks before plotting sensitivity analysis results*

---

**Description**

Function to do some checks before plotting sensitivity analysis results

**Usage**

```
plot_dsa_nmb_range(ob_results, plot_var)
```

**Arguments**

ob_results	results from deterministic sensitivity analysis
plot_var	the variable

**Value**

plot

---

plot\_dsa\_others\_range     *Function to do some checks before plotting sensitivity analysis results*

---

**Description**

Function to do some checks before plotting sensitivity analysis results

**Usage**

```
plot_dsa_others_range(ob_results, plot_var)
```

**Arguments**

ob_results	results from deterministic sensitivity analysis
plot_var	the variable

**Value**

plot

---

```
plot_efficiency_frontier
      Plot efficiency frontier
```

---

**Description**

Plot efficiency frontier

**Usage**

```
plot_efficiency_frontier(results_calculate_icer_nmb, threshold)
```

**Arguments**

```
results_calculate_icer_nmb
      results from cea (from calculate_icer_nmb method)
threshold
      threshold value
```

**Value**

```
plot well <- health_state("well", cost = 0, utility = 1) disabled <- health_state("disabled", cost =
100, utility = 1) dead <- health_state("dead", cost = 0, utility = 0) tmat <- rbind(c(1, 2, 3), c(NA, 4,
5), c(NA, NA, 6)) colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead") tm <- popu-
late_transition_matrix(3, tmat, c(0.6, 0.2, 0.2, 0.6, 0.4, 1), colnames(tmat)) health_states <- com-
bine_state(well, disabled, dead) this.strategy <- strategy(tm, health_states, "control") this_markov
<- markov_model(this.strategy, 24, c(1000, 0, 0), c(0,0)) well <- health_state("well", cost = 0, utility
= 1) disabled <- health_state("disabled", cost = 10, utility = 0.5) dead <- health_state("dead", cost =
0, utility = 0) tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6)) colnames(tmat) <- rownames(tmat)
<- c("well", "disabled", "dead") tm <- populate_transition_matrix(3, tmat, c(0.4, 0.4, 0.2, 0.6,
0.4, 1), colnames(tmat)) health_states <- combine_state(well, disabled, dead) this.strategy <- strat-
egy(tm, health_states, "intervention") sec_markov <- markov_model(this.strategy, 24, c(1000, 0, 0),
c(0,0)) list_markov <- combine_markov(this_markov, sec_markov) results_cea <- calculate_icer_nmb(list_markov,
20000, comparator = "control") plot_efficiency_frontier(results_cea, c(1000, 2000))
```

---

```
plot_histogram_onetimepoint_twogroups
      Function to plot mean and SE for longitudinal observations for
      twogroups compared
```

---

**Description**

Function to plot mean and SE for longitudinal observations for twogroups compared

**Usage**

```
plot_histogram_onetimepoint_twogroups(
  thedata,
  colname,
  timepointstring,
  xstring,
  ylimits = NULL,
  nbins = NULL
)
```

**Arguments**

thedata	the data where the observations are held
colname	columnname in the data where the intersted observations
timepointstring	the text that correspond to timepoints at which the descriptive analysis is done
xstring	xlable text
ylimits	on hsitogram plot
nbins	nbins the number of bins to plot histogram

**Value**

the historgram plot at a timepoint for two groups

**Examples**

```
eg_data <- as.data.frame(list(no = c(1, 2, 3, 4),
  mark_at_1 = c(12, 7, 23, 45), gender = c("M", "F", "M", "F"),
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2")))
plot_histogram_onetimepoint_twogroups(eg_data, c("mark_at_2"), c("1", "2"),
  "mark")
```

---

plot\_meanSE\_longitudinal\_twogroups

*Function to plot mean and SE for longitudinal observations for twogroups compared*

---

**Description**

Function to plot mean and SE for longitudinal observations for twogroups compared

**Usage**

```
plot_meanSE_longitudinal_twogroups(
  thedata,
  columnnames,
  timepoints,
  observation
)
```

**Arguments**

thedata	the data where the observations are held
columnnames	columnnames in the data where the interested observations
timepoints	the timepoints at which the descriptive analysis is done
observation	name of the observations

**Value**

the plot that shows mean and SE

**Examples**

```
eg_data <- as.data.frame(list(no = c(1, 2, 3, 4),
  mark_at_1 = c(12, 7, 23, 45), gender = c("M", "F", "M", "F"),
  mark_at_2 = c(12, 34, 89, 45), trialarm = c("1", "1", "2", "2")))
plot_meanSE_longitudinal_twogroups(eg_data, c("mark_at_1", "mark_at_2"),
  c("1", "2"), "mark")
```

---

plot\_model

*E1. Plot a Markov model*

---

**Description**

E1. Plot a Markov model

**Usage**

```
plot_model(markov)
```

**Arguments**

markov	markov_model object
--------	---------------------

**Value**

plots



**Examples**

```
tmat <- rbind(c(1, 2), c(3, 4))
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")
tm <- populate_transition_matrix(2, tmat, c(0.5, 0.5, 0, 1))
a <- health_state("Healthy", 1, 1, 0, FALSE)
b <- health_state("Dead", 1, 0, 0, TRUE)
health_states <- combine_state(a, b)
this_strategy <- strategy(tm, health_states, "intervention")
this_markov <- markov_model(this_strategy, 10, c(1, 0), c(0, 0))
p <- plot_model(this_markov)
```

plot\_nmb\_lambda

*Plot cost effectiveness acceptability curve***Description**

Plot cost effectiveness acceptability curve

**Usage**

```
plot_nmb_lambda(list_markov, threshold_values, comparator, currency = "GBP")
```

**Arguments**

list_markov	markov_model objects
threshold_values	list of threshold values
comparator	the comparator
currency	currency

**Value**

plots

**Examples**

```
well <- health_state("well", cost = 0, utility = 1)
disabled <- health_state("disabled", cost = 100, utility = 1)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.6, 0.2, 0.2, 0.6, 0.4, 1),
colnames(tmat))
health_states <- combine_state(well, disabled, dead)
this_strategy <- strategy(tm, health_states, "control")
this_markov <- markov_model(this_strategy, 24, c(1000, 0, 0), c(0, 0))
well <- health_state("well", cost = 0, utility = 1)
```

```

disabled <- health_state("disabled", cost = 10, utility = 0.5)
dead <- health_state("dead", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("well", "disabled", "dead")
tm <- populate_transition_matrix(3, tmat, c(0.4, 0.4, 0.2, 0.6, 0.4, 1),
colnames(tmat))
health_states <- combine_state(well, disabled, dead)
this.strategy <- strategy(tm, health_states, "intervention")
sec_markov <- markov_model(this.strategy, 24, c(1000, 0, 0), c(0, 0))
list_markov <- combine_markov(this_markov, sec_markov)
plot_nmb_lambda(list_markov, c(1000, 2000, 3000), comparator = "control")

```

---

plot\_prediction\_parametric\_survival

*Plot the predicted survival curves for covariates keeping the others fixed*

---

### Description

Plot the predicted survival curves for covariates keeping the others fixed

### Usage

```

plot_prediction_parametric_survival(
  param_to_be_estimated,
  indep_var,
  covariates,
  dataset,
  fit,
  timevar_survival
)

```

### Arguments

param_to_be_estimated	parameter to be estimated
indep_var	variable for which the levels have to be identified
covariates	the covariates
dataset	the dataset where these variables contain
fit	the fit result survreg
timevar_survival	time variable from the dataset

### Value

plot

**Examples**

```

data_for_survival <- survival::lung
surv_estimated <- use_parametric_survival("status", data_for_survival,
"sex",
info_distribution = "weibull", covariates = c("ph.ecog"), "time")
plot_prediction_parametric_survival("status", "sex",
covariates = c("ph.ecog"), data_for_survival, surv_estimated$fit, "time")

```

---

plot\_return\_residual\_cox

*Plotting and return the residuals after cox proportional hazard model*


---

**Description**

Plotting and return the residuals after cox proportional hazard model

**Usage**

```

plot_return_residual_cox(
  param_to_be_estimated,
  indep_var,
  covariates,
  fit,
  dataset
)

```

**Arguments**

param_to_be_estimated	parameter to be estimated
indep_var	independent variable
covariates	covariates
fit	fit object from coxph method
dataset	data used for cox ph model

**Value**

plot and the residuals

**Examples**

```

data_for_survival <- survival::lung
surv_estimated <- use_coxph_survival("status", data_for_survival, "sex",
covariates = c("ph.ecog"), "time")
plot_return_residual_cox("status", "sex", covariates = c("ph.ecog"),
surv_estimated$fit, data_for_survival )

```

---

`plot_return_residual_survival`*Plotting and return the residuals after survival model*

---

**Description**

Plotting and return the residuals after survival model

**Usage**

```
plot_return_residual_survival(  
  param_to_be_estimated,  
  indep_var,  
  covariates,  
  fit  
)
```

**Arguments**

<code>param_to_be_estimated</code>	parameter to be estimated
<code>indep_var</code>	independent variable
<code>covariates</code>	covariates
<code>fit</code>	fit object from survreg method

**Value**

plot and the residuals

**Examples**

```
data_for_survival <- survival::lung  
surv_estimated <- use_parametric_survival("status", data_for_survival,  
  "sex",  
  info_distribution = "weibull", covariates = c("ph.ecog"), "time")  
plot_return_residual_survival("status", "sex",  
  covariates = c("ph.ecog"), surv_estimated$fit)
```

---

`plot_return_survival_curve`*Plotting survival function for all covariates using survfit*

---

**Description**

Plotting survival function for all covariates using survfit

**Usage**

```
plot_return_survival_curve(  
  param_to_be_estimated,  
  dataset,  
  indep_var,  
  covariates,  
  timevar_survival  
)
```

**Arguments**

<code>param_to_be_estimated</code>	parameter to be estimated
<code>dataset</code>	param describing the methods
<code>indep_var</code>	independent variable
<code>covariates</code>	covariates
<code>timevar_survival</code>	time variable for survival analysis

**Value**

plot and the survival function values

**Examples**

```
data_for_survival <- survival::lung  
plot_return_survival_curve(param_to_be_estimated = "status",  
  dataset = data_for_survival, indep_var = "sex", covariates = c("ph.ecog"),  
  timevar_survival = "time")
```

---

`plot_survival_cox_covariates`

*Plotting survival function for all covariates calculated from cox regression results and returned coefficients*

---

### Description

Plotting survival function for all covariates calculated from cox regression results and returned coefficients

### Usage

```
plot_survival_cox_covariates(  
  coxfit,  
  dataset,  
  param_to_be_estimated,  
  covariates,  
  indep_var  
)
```

### Arguments

<code>coxfit</code>	cox regression fit result
<code>dataset</code>	param describing the methods
<code>param_to_be_estimated</code>	parameter to be estimated
<code>covariates</code>	covariates
<code>indep_var</code>	independent variable

### Value

plot and the survival function values

### Examples

```
data_for_survival <- survival::lung  
surv_estimated <- use_coxph_survival("status", data_for_survival, "sex",  
  covariates = c("ph.ecog"), "time")  
plot_survival_cox_covariates(surv_estimated$fit, data_for_survival,  
  "status", covariates = c("ph.ecog"), "sex")
```

---

populate\_transition\_matrix  
*Populate transition matrix*

---

### Description

Populate transition matrix

### Usage

```
populate_transition_matrix(no_states, tmat, list_prob, name_states = NULL)
```

### Arguments

no_states	number of the health states
tmat	A transition matrix in the format from the package 'mstate'
list_prob	list of probabilities as in the order of transitions (row wise)
name_states	names of the health states

### Details

If the state names are null, they are replaced with numbers starting from 1 First find those missing probabilities, and fill a list from the given list of probabilities and fill those are not NA in the matrix Note that the probabilities need not be numeric here and no checks are needed for sum

### Value

value of the transition matrix

### Examples

```
tmat <- rbind(c(1, 2), c(3, 4))  
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")  
populate_transition_matrix(2, tmat, list_prob = c(0.2, 0.5, 0, 0.3))
```

---

predict\_coxph *Predict risk/hazard function for cox ph regression*

---

### Description

Predict risk/hazard function for cox ph regression

**Usage**

```
predict_coxph(
  coxfit,
  dataset,
  param_to_be_estimated,
  covariates,
  indep_var,
  timevar_survival
)
```

**Arguments**

coxfit	cox regression fit result
dataset	param describing the methods
param_to_be_estimated	parameter to be estimated
covariates	covariates
indep_var	independent variable
timevar_survival	time variable

**Details**

"risk" option for "type" returns the hazard ratio relative to mean e.g given below For lung data with `data_for_survival <- survival::lung` `fit <- use_coxph_survival("status", data_for_survival, "sex", covariates = c("ph.ecog"), "time")` `coeffit = fit$coefficients` `r1234 <- exp(coeffit("sex"))` `lung$sex + coeffit("ph.ecog")` `lung$ph.ecog` `rMean <- exp(sum(coef(fit) * fit$means, na.rm=TRUE))` `rr <- r1234/rMean`

**Value**

plot and the survival function values

**Examples**

```
data_for_survival <- survival::lung
surv_estimated <- use_coxph_survival("status", data_for_survival,
  "sex", covariates = c("ph.ecog"), "time")
predict_coxph(surv_estimated$fit, data_for_survival, "status", "sex",
  covariates = c("ph.ecog"), "time")
```



---

promis3a\_scoring.df    *promis 3a scoring table*

---

**Description**

promis 3a scoring table

**Usage**

promis3a\_scoring.df

**Format**

A 14 by 3 dataframe

**Source**

created on April 08, 2021

---

report\_sensitivity\_analysis  
*Function to report deterministic sensitivity analysis*

---

**Description**

Function to report deterministic sensitivity analysis

**Usage**

```
report_sensitivity_analysis(  
  result_dsa_control,  
  result_dsa_treat = NULL,  
  threshold = NULL,  
  comparator = NULL  
)
```

**Arguments**

result_dsa_control	result from deterministic sensitivity analysis for first or control model
result_dsa_treat	result from deterministic sensitivity analysis for the comparative Markov model
threshold	threshold value of WTP
comparator	the strategy to be compared with

**Value**

report in the form of a table

**Examples**

```

param_list <- define_parameters(
  cost_zido = 2278, cost_direct_med_A = 1701,
  cost_comm_care_A = 1055, cost_direct_med_B = 1774,
  cost_comm_care_B = 1278,
  cost_direct_med_C = 6948, cost_comm_care_C = 2059,
  tpAtoA = 1251 / (1251 + 483),
  tpAtoB = 350 / (350 + 1384), tpAtoC = 116 / (116 + 1618),
  tpAtoD = 17 / (17 + 1717),
  tpBtoB = 731 / (731 + 527), tpBtoC = 512 / (512 + 746),
  tpBtoD = 15 / (15 + 1243),
  tpCtoC = 1312 / (1312 + 437), tpCtoD = 437 / (437 + 1312),
  tpDtoD = 1,
  cost_health_A = "cost_direct_med_A + cost_comm_care_A",
  cost_health_B = "cost_direct_med_B + cost_comm_care_B",
  cost_health_C = "cost_direct_med_C + cost_comm_care_C",
  cost_drug = "cost_zido")
low_values <- define_parameters(cost_direct_med_B = 177.4,
  cost_comm_care_C = 205.9)
upp_values <- define_parameters(cost_direct_med_B = 17740,
  cost_comm_care_C = 20590)
A <- health_state("A", cost = "cost_health_A + cost_drug ", utility = 1)
B <- health_state("B", cost = "cost_health_B + cost_drug", utility = 1)
C <- health_state("C", cost = "cost_health_C + cost_drug", utility = 1)
D <- health_state("D", cost = 0, utility = 0)
tmat <- rbind(c(1, 2, 3, 4), c(NA, 5, 6, 7), c(NA, NA, 8, 9),
  c(NA, NA, NA, 10))
colnames(tmat) <- rownames(tmat) <- c("A", "B", "C", "D")
tm <- populate_transition_matrix(4, tmat, c("tpAtoA", "tpAtoB", "tpAtoC",
  "tpAtoD", "tpBtoB", "tpBtoC", "tpBtoD", "tpCtoC", "tpCtoD", "tpDtoD"),
  colnames(tmat))
health_states <- combine_state(A, B, C, D)
mono_strategy <- strategy(tm, health_states, "mono")
mono_markov <- markov_model(mono_strategy, 20, c(1, 0, 0, 0),
  discount = c(0.06, 0), param_list)
param_table <- define_parameters_sens_anal(param_list, low_values,
  upp_values)
result <- do_sensitivity_analysis(mono_markov, param_table)
reporting <- report_sensitivity_analysis(result)

```

---

return0\_if\_not\_null\_na

*Function to return 0 if the param is not null or NA trimming the white spaces*

---

**Description**

Function to return 0 if the param is not null or NA trimming the white spaces

**Usage**

```
return0_if_not_null_na(param)
```

**Arguments**

param                    the form of medication either tablet or patch

**Value**

zero or -1

**Examples**

```
parame = NULL
ans <- return0_if_not_null_na(parame)
parame = 1
ans <- return0_if_not_null_na(parame)
```

---

return\_equal\_liststring\_col

*Function to get the subset of data compared to a string after trimming the white spaces*

---

**Description**

Function to get the subset of data compared to a string after trimming the white spaces

**Usage**

```
return_equal_liststring_col(col, the_data, list_str)
```

**Arguments**

col                    the form of medication either tablet or patch  
the\_data               the data to be get the subset from  
list\_str               list of strings to be compared

**Value**

the subset data

**Examples**

```
the_data <- as.data.frame(cbind(c("one", "two"), c("a", "b")))
colnames(the_data) <- c("name", "brand")
ans <- return_equal_liststring_col(2, the_data, c("a", "cc"))
```

---

return\_equal\_liststring\_listcol

*Function to get the subset of data compared to a string after trimming the white spaces*

---

**Description**

Function to get the subset of data compared to a string after trimming the white spaces

**Usage**

```
return_equal_liststring_listcol(col, the_data, list_str)
```

**Arguments**

col	the form of medication either tablet or patch
the_data	the data to be get the subset from
list_str	list of strings to be compared

**Value**

the subset data

**Examples**

```
the_data <- as.data.frame(cbind(c("one", "two"), c("tablet", "tablets"),
c("aa", "bb")))
colnames(the_data) <- c("name", "brand_a", "xx")
ans <- return_equal_liststring_listcol(2, the_data, c("tablet", "tablets"))
```

---

return\_equal\_str\_col    *Function to get the subset of data compared to a string after trimming the white spaces*

---

**Description**

Function to get the subset of data compared to a string after trimming the white spaces

**Usage**

```
return_equal_str_col(col, the_data, the_str)
```

**Arguments**

col	the form of medication either tablet or patch
the_data	the data to be get the subset from
the_str	the string to be compared

**Value**

the subset data

**Examples**

```
the_data <- as.data.frame(cbind(c("one", "two"), c("a", "b")))
colnames(the_data) <- c("name", "brand")
ans <- return_equal_str_col(2, the_data, "a")
```

---

set\_var\_state            *Set the attribute for the health state*

---

**Description**

Set the attribute for the health state

**Usage**

```
set_var_state(state, var, new_value)
```

**Arguments**

state	object of class health state
var	attribute of the health state
new_value	new value to be assigned

**Details**

After checking the given state is a health state the value of the variable is set if the value is not numeric, it is being parsed to form an expression

**Value**

modified health state

**Examples**

```
set_var_state(health_state("IT", 100, 0.4, 0, FALSE), "cost", 1)
```

---

strategy	<i>Definition of strategy - or arm</i>
----------	--

---

**Description**

Definition of strategy - or arm

**Usage**

```
strategy(trans_mat, states, name, trans_cost = NULL, trans_util = NULL)
```

**Arguments**

trans_mat	transition matrix
states	health states
name	name of the strategy
trans_cost	values of costs if these are attached to transitions
trans_util	values of utility if these are attached to transitions

**Details**

Defining strategy keeping all transition matrix, states and names together to use in defining Markov model

**Value**

object strategy

**Examples**

```
tmat <- rbind(c(1, 2), c(3, 4))
colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead")
tm <- populate_transition_matrix(2, tmat, c(0.5, 0.5, 0, 1))
a <- health_state("Healthy", 1, 1, 0, FALSE)
b <- health_state("Dead", 1, 0.5, 0, FALSE)
states <- combine_state(a, b)
strategy(tm, states, "intervention")
```

---

summary_plot_psa	<i>Function to summarise and plot probabilistic sensitivity analysis</i>
------------------	--

---

**Description**

Function to summarise and plot probabilistic sensitivity analysis

**Usage**

```
summary_plot_psa(
  result_psa_params_control,
  result_psa_params_treat = NULL,
  threshold = NULL,
  comparator = NULL
)
```

**Arguments**

result_psa_params_control	result from probabilistic sensitivity analysis for first or control model
result_psa_params_treat	result from probabilistic sensitivity analysis for the comparative Markov model
threshold	threshold value of WTP
comparator	the strategy to be compared with

**Value**

plot of sensitivity analysis

**Examples**

```
param_list <- define_parameters(
  cost_direct_med_A = 1701,
  cost_direct_med_B = 1774, tpAtoA = 0.2,
  tpAtoB = 0.5, tpAtoC = 0.3,
  tpBtoB = 0.3, tpBtoC = 0.7,
  tpCtoC = 1, cost_health_A = "cost_direct_med_A",
  cost_health_B = "cost_direct_med_B")
sample_list <- define_parameters(cost_direct_med_A = "gamma(mean = 1701,
  sd = sqrt(1701))")
A <- health_state("A", cost = "cost_health_A ", utility = 1)
B <- health_state("B", cost = "cost_health_B", utility = 1)
C <- health_state("C", cost = 0, utility = 0, absorb = "TRUE")
tmat <- rbind(c(1, 2, 3), c(NA, 4, 5), c(NA, NA, 6))
colnames(tmat) <- rownames(tmat) <- c("A", "B", "C")
tm <- populate_transition_matrix(3, tmat, c(
  "tpAtoA", "tpAtoB", "tpAtoC", "tpBtoB", "tpBtoC", "tpCtoC"),
  colnames(tmat))
```

```

health_states <- combine_state(A, B, C)
mono_strategy <- strategy(tm, health_states, "mono")
mono_markov <- markov_model(mono_strategy, 20, initial_state =c(1,0,0),
discount = c(0.06, 0),param_list)
param_table <- define_parameters_psa(param_list, sample_list)
result <- do_psa(mono_markov, param_table, 3)
result_plot <- summary_plot_psa(result, NULL, NULL, NULL)
param_list_comb <- define_parameters(
cost_direct_med_A = 1800, cost_direct_med_B = 1774, tpAtoA = 0.6,
tpAtoB = 0.1, tpAtoC = 0.3, tpBtoB = 0.3, tpBtoC = 0.7, tpCtoC = 1,
cost_health_A = "cost_direct_med_A", cost_health_B = "cost_direct_med_B")
comb_strategy <- strategy(tm, health_states, "comb")
comb_markov <- markov_model(comb_strategy, 20, c(1, 0, 0),
discount = c(0.06, 0), param_list)
param_table_comb <- define_parameters_psa(param_list_comb, sample_list)
result_comb <- do_psa(comb_markov, param_table_comb, 3)
summary_plot_psa(result, result_comb, 2000, "mono")

```

---

table_param.df	<i>Parameter table created</i>
----------------	--------------------------------

---

### Description

Parameter table created

### Usage

table\_param.df

### Format

A 11 by 2 dataframe

### Source

created on Jan 15, 2020

---

trace_data.df	<i>Trace matrix</i>
---------------	---------------------

---

### Description

Trace matrix

### Usage

trace\_data.df



**Format**

A 11 by 2 dataframe

**Source**

created on Nov 26, 2019 from `tmat <- rbind(c(1, 2), c(3, 4)) colnames(tmat) <- rownames(tmat) <- c("Healthy", "Dead") tm <- transition_matrix(2, tmat, c(0.5, 0.5, 0, 1)) a <- health_state("Healthy", 1, 1, FALSE) b <- health_state("Dead", 1, 0, TRUE) health_states <- combine_state(a, b) this.strategy <- strategy(tm, health_states, "intervention")`

---

`transition_cost_util` *Create the the values of cost and utility while transition*

---

**Description**

Create the the values of cost and utility while transition

**Usage**

```
transition_cost_util(
  no_states,
  tmat_cost_util,
  list_values,
  name_states = NULL
)
```

**Arguments**

`no_states` number of the health states

`tmat_cost_util` A transition matrix for the cost/utility values in the format from the package 'mstate' use NA to indicate if the value is zero

`list_values` list of probabilities as in the order of transitions (row wise)

`name_states` names of the health states

**Details**

Similar to transition matrix but for denoting one time change during transitions

**Value**

value of the transition matrix

**Examples**

```
tmat_cost <- rbind(c(NA, 1), c(NA, NA))
colnames(tmat_cost) <- rownames(tmat_cost) <- c("Healthy", "Dead")
transition_cost_util(2, tmat_cost, list_values = c(500))
```

---

trial_data.df	<i>Example trial data</i>
---------------	---------------------------

---

**Description**

Example trial data

**Usage**

trial\_data.df

**Format**

A 31 by 33 dataframe

**Source**

created on Jan 15, 2020

---

use_coxph_survival	##### <i>Get the parameter values using the survival analysis using cox proportional hazard</i>
--------------------	--

---

**Description**

##### Get the parameter values using the survival analysis using cox proportional hazard

**Usage**

```
use_coxph_survival(
  param_to_be_estimated,
  dataset,
  indep_var,
  covariates,
  timevar_survival
)
```

**Arguments**

param\_to\_be\_estimated  
parameter of interest

dataset  
data set to be provided

indep\_var  
the independent variable (column name in data file)

covariates  
list of covariates - calculations to be done before passing

timevar\_survival  
time variable for survival analysis, default is NA false by default

**Details**

plots baseline cumulative hazard function, survival function for each covariate while keeping the other fixed at the mean value (using plot\_survival\_cox\_covariates), survival function for each combination of covariate using survfit (using plot\_return\_survival\_curve) and test for cox regression results It also returns risk relative to mean (predicted at mean value of each covariate) along with the fit results coefficients, SE of coefficients, summary, and analysis of deviance

**Value**

the results of the regression analysis

**Examples**

```
data_for_survival <- survival::aml
surv_estimated <- use_coxph_survival("status", data_for_survival, "x",
  covariates = NA, "time")
```

```
data_for_survival <- survival::lung
surv_estimated <- use_coxph_survival("status", data_for_survival, "sex",
  covariates = c("ph.ecog"), "time")
```

---

```
use_fh2_survival #####
                  Get the parameter values using the survival analysis using FH2
                  method
```

---

**Description**

```
##### Get
the parameter values using the survival analysis using FH2 method
```

**Usage**

```
use_fh2_survival(
  param_to_be_estimated,
  dataset,
  indep_var,
  covariates,
  timevar_survival
)
```

**Arguments**

```
param_to_be_estimated      parameter of interest
dataset                    data set to be provided
indep_var                  the independent variable (column name in data file)
covariates                 list of covariates
timevar_survival          time variable for survival analysis, default is NA false by default
```

**Details**

This function is for survival analysis using FH2. This plots the cumulative survival function for each combination of covariate. If the covariate is numeric, R takes it as different levels. The plot uses the returned list of survfit and extracts the time and the strata from summary of the fit (implemented in plot\_return\_survival\_curve function)

**Value**

the results of the regression analysis

**Examples**

```
data_for_survival <- survival::aml
surv_estimated <- use_fh2_survival("status", data_for_survival, "x",
  covariates = NA, "time")
```

---

```
use_fh_survival      #####
                    Get the parameter values using the survival analysis method FH
```

---

**Description**

```
##### Get the
parameter values using the survival analysis method FH
```

**Usage**

```
use_fh_survival(  
  param_to_be_estimated,  
  dataset,  
  indep_var,  
  covariates,  
  timevar_survival  
)
```

**Arguments**

param_to_be_estimated	parameter of interest
dataset	data set to be provided
indep_var	the independent variable (column name in data file)
covariates	list of covariates
timevar_survival	time variable for survival analysis, default is NA

**Details**

This function is for survival analysis using FH. This plots the cumulative survival function for each combination of covariate. If the covariate is numeric, R takes it as different levels. The plot uses the returned list of survfit and extracts the time and the strata from summary of the fit (implemented in plot\_return\_survival\_curve function)

**Value**

the results of the regression analysis

**Examples**

```
data_for_survival <- survival::aml  
surv_estimated <- use_fh_survival("status", data_for_survival, "x",  
  covariates = NA, "time"  
)
```

---

use\_generalised\_linear\_mixed\_model

*Function for generalised linear mixed model*

---

**Description**

Function for generalised linear mixed model

**Usage**

```

use_generalised_linear_mixed_model(
  param_to_be_estimated,
  dataset,
  fix_eff,
  fix_eff_interact_vars,
  random_intercept_vars,
  nested_intercept_vars_pairs,
  cross_intercept_vars_pairs,
  uncorrel_slope_intercept_pairs,
  random_slope_intercept_pairs,
  family,
  link,
  package_mixed_model
)

```

**Arguments**

<code>param_to_be_estimated</code>	column name of dependent variable
<code>dataset</code>	a dataframe
<code>fix_eff</code>	names of variables as fixed effect predictors
<code>fix_eff_interact_vars</code>	those of the fixed effect predictors that show interaction
<code>random_intercept_vars</code>	names of variables for random intercept
<code>nested_intercept_vars_pairs</code>	those of the random intercept variables with nested effect
<code>cross_intercept_vars_pairs</code>	those of the random intercept variables with crossed effect
<code>uncorrel_slope_intercept_pairs</code>	variables with no correlated intercepts
<code>random_slope_intercept_pairs</code>	random slopes intercept pairs - this is a list of paired variables
<code>family</code>	family of distributions for the response variable
<code>link</code>	link function for the variances
<code>package_mixed_model</code>	package to be used for mixed model

**Value**

result regression result with plot if success and -1, if failure

**Examples**

```

datafile <- system.file("extdata", "culcita_data.csv",
package = "packDAMipd")
dataset <- read.csv(datafile)
results1 = use_generalised_linear_mixed_model("predation",
dataset = datafile,fix_eff = c("ttt"), family = "binomial",
fix_eff_interact_vars = NULL, random_intercept_vars = c("block"),
nested_intercept_vars_pairs = NULL, cross_intercept_vars_pairs = NULL,
uncorrel_slope_intercept_pairs = NULL, random_slope_intercept_pairs = NULL,
link = NA, package_mixed_model = NA)

```

---

```

use_generalised_linear_model

```

```

#####
Get the parameter values using logistic regression

```

---

**Description**

```

##### Get
the parameter values using logistic regression

```

**Usage**

```

use_generalised_linear_model(
  param_to_be_estimated,
  dataset,
  indep_var,
  family,
  covariates,
  interaction,
  naaction,
  link = NA
)

```

**Arguments**

param_to_be_estimated	parameter of interest
dataset	data set to be provided
indep_var	the independent variable (column name in data file)
family	distribution name eg. for logistic regression -binomial
covariates	list of covariates-calculations to be done before passing
interaction	boolean value to indicate interaction in the case of linear regression
naaction	action to be taken with the missing values
link	link function if not the default for each family

**Details**

This function returns the results and plots after doing linear regression Requires param to be estimated, dataset, independent variables and information on covariates, and interaction variables if there are Uses form\_expression\_glm to create the expression as per R standard for e.g glm(y ~ x ). Returns the fit result,s summary results as returned by summary(), confidence interval for fit coefficients (ci\_coeff), variance covariance matrix, cholesky decomposition matrix, results from correlation test, plot of diagnostic tests and model fit assumptions, plot of model prediction diagnostic include AIC, R2, and BIC. The results of the prediction ie predicted values for fixed other variables will be returned in prediction matrix

**Value**

the results of the regression analysis

**Examples**

```
gm_result <- use_generalised_linear_model(
  param_to_be_estimated = "Direction",
  dataset = ISLR::Smarket, indep_var = "Lag1", family = "binomial",
  covariates = c("Lag2", "Lag3"),
  interaction = FALSE, naaction = "na.omit", link = NA)
```

---

```
use_km_survival #####
                  Get the parameter values using the Kaplan-Meier survival analysis
```

---

**Description**

```
##### Get the
parameter values using the Kaplan-Meier survival analysis
```

**Usage**

```
use_km_survival(
  param_to_be_estimated,
  dataset,
  indep_var,
  covariates,
  timevar_survival
)
```

**Arguments**

```
param_to_be_estimated
                    parameter of interest
dataset             data set to be provided
```



indep\_var        the independent variable (column name in data file)  
 covariates       list of covariates  
 timevar\_survival  
                  time variable for survival analysis, default is NA

### Details

This function is for survival analysis using Kaplan Meier. This plots the cumulative survival function for each combination of covariate. If the covariate is numeric, R takes it as different levels. The plot uses the returned list of survfit and extracts the time and the strata from summary of the fit (implemented in plot\_return\_survival\_curve function)

### Value

the results of the regression analysis, fit results, summary and plot

### Examples

```
data_for_survival <- survival::aml
surv_estimated <- use_km_survival("status", data_for_survival, "x",
  covariates = NA, "time")
```

```
data_for_survival <- survival::lung
surv_estimated <- use_km_survival("status", data_for_survival, "sex",
  covariates = c("ph.ecog"), "time")
```

---

use\_linear\_mixed\_model

*Function for mixed effect regression*

---

### Description

Function for mixed effect regression

### Usage

```
use_linear_mixed_model(
  param_to_be_estimated,
  dataset,
  fix_eff,
  fix_eff_interact_vars,
  random_intercept_vars,
  nested_intercept_vars_pairs,
  cross_intercept_vars_pairs,
  uncorrel_slope_intercept_pairs,
  random_slope_intercept_pairs,
  package_mixed_model
)
```

**Arguments**

param_to_be_estimated	column name of dependent variable
dataset	a dataframe
fix_eff	names of variables as fixed effect predictors
fix_eff_interact_vars	those of the fixed effect predictors that show interaction
random_intercept_vars	names of variables for random intercept
nested_intercept_vars_pairs	those of the random intercept variables with nested effect
cross_intercept_vars_pairs	those of the random intercept variables with crossed effect
uncorrel_slope_intercept_pairs	variables with no correlated intercepts
random_slope_intercept_pairs	random slopes intercept pairs - this is a list of paired variables
package_mixed_model	package to be used for mixed model

**Value**

result regression result with plot if success and -1, if failure

**Examples**

```
datafile <- system.file("extdata", "data_linear_mixed_model.csv",
package = "packDAMipd")
dataset = utils::read.table(datafile, header = TRUE, sep = ",",
na.strings = "NA",
dec = ".", strip.white = TRUE)
result <- use_linear_mixed_model("extro",
dataset = dataset,
fix_eff = c("open", "agree", "social"), fix_eff_interact_vars = NULL,
random_intercept_vars = c("school", "class"),
nested_intercept_vars_pairs = list(c("school", "class")),
cross_intercept_vars_pairs = NULL, uncorrel_slope_intercept_pairs = NULL,
random_slope_intercept_pairs = NULL, package_mixed_model = NA)
```

---

```
use_linear_regression #####
                        Get the parameter values using the linear regression
```

---

## Description

```
##### Get the
parameter values using the linear regression
```

## Usage

```
use_linear_regression(
  param_to_be_estimated,
  dataset,
  indep_var,
  covariates,
  interaction
)
```

## Arguments

param_to_be_estimated	parameter of interest
dataset	data set to be provided
indep_var	the independent variable (column name in data file)
covariates	list of covariates-calculations to be done before passing
interaction	boolean value to indicate interaction in the case of linear regression, false by default

## Details

This function returns the results and plots after doing linear regression Requires param to be estimated, dataset, independent variables and information on covariates, and interaction variables if there are Uses form\_expression\_lm to create the expression as per R standard for e.g lm(y ~ x ). Returns the fit result,s summary results as returned by summary(), confidence interval for fit coefficients (ci\_coeff), variance covariance matrix, cholesky decomposition matrix, Results from correlation test, plot of diagnostic tests and model fit assumptions, plot of model prediction diagnostic include AIC, R2, and BIC. The results of the prediction ie predicted values when each of covariate is fixed will be returned in prediction matrix predicted values will provide the mean value of param\_to\_to\_estimated as calculated by the linear regression formula. ref:<https://www.statmethods.net/stats/regression.html>

## Value

the results of the regression analysis

**Examples**

```
results_lm <- use_linear_regression("dist",
  dataset = cars,
  indep_var = "speed", covariates = NA, interaction = FALSE)
```

```
library(car)
results_lm <- use_linear_regression("mpg",
  dataset = mtcars,
  indep_var = "disp", covariates = c("hp", "wt", "drat"),
  interaction = FALSE)
```

---

```
use_parametric_survival
```

```
#####
Get the parameter values using the survival analysis parametric survival
```

---

**Description**

```
##### Get the
parameter values using the survival analysis parametric survival
```

**Usage**

```
use_parametric_survival(
  param_to_be_estimated,
  dataset,
  indep_var,
  info_distribution,
  covariates,
  timevar_survival,
  cluster_var = NA
)
```

**Arguments**

```
param_to_be_estimated    parameter of interest
dataset                   data set to be provided
indep_var                 the independent variable (column name in data file)
info_distribution         distribution name eg. for logistic regression -binomial
covariates                list of covariates
timevar_survival         time variable for survival analysis, default is NA
cluster_var               cluster variable for survival analysis
```

**Details**

This function is the last in the layer of function for parametric survival analysis. This then returns the parameters of interest, plots the results etc if the distribution is weibull it uses the package SurvRegCensCov for easy interpretation of results Returns the fit result, summary of regression, variance-covariance matrix of coeff, cholesky decomposition, the parameters that define the assumed distribution and the plot of model prediction Using survfit from survival package to plot the survival curve R's weibull distribution is defined as  $\text{std weibull}$  in terms of a and b as  $(a/b) (x/b)^{(a-1)} \exp(-x/b)^a$  where a is the shape and b is the scale In HE the weibull distribution is parameterised as bit different it is like  $\text{gamma.lambda. } t^{(\text{gamma}-1)} \cdot \exp(-\text{lambda} \cdot t^{\text{gamma}})$  where gamma is the shape and lambda is the scale. The relationship is as below.  $\text{HE\_shape} = \text{rweibull\_shape}$   $\text{HE\_scale} = \text{rweibull\_scale}^{(-\text{rweibull\_shape})}$  The survreg shape and scale are again bit different and they are rweibull's shape and scale as below.  $\text{rweibull\_shape} = 1/\text{fit}\$scale$   $\text{rweibull\_scale} = \exp(\text{fit intercept}) = \exp(\text{fit}\$coefficients)$  remember to use 1st of coefficients This has been utilised in SurvRegCensCov::ConvertWeibull predict() for survreg object with type =quantile will provide the failure times as survival function is 1-CDF of failure time.

**Value**

the results of the regression analysis

**Examples**

```
data_for_survival <- survival::lung
surv_estimated <- use_parametric_survival("status",
data_for_survival, "sex", info_distribution = "weibull",
covariates = c("ph.ecog"), "time")
```

---

use\_seemingly\_unrelated\_regression

*Bivariate regression for correlated observations*

---

**Description**

Bivariate regression for correlated observations

**Usage**

```
use_seemingly_unrelated_regression(
  param1_to_be_estimated,
  param2_to_be_estimated,
  dataset,
  indep_var,
  covariates1,
  covariates2,
  interaction1,
  interaction2
)
```



```

    timevar_survival,
    cluster_var = NA
  )

```

### Arguments

```

param_to_be_estimated
                    parameter of interest
dataset             data set to be provided
indep_var          the independent variable (column name in data file)
info_get_method    additional information on methods e.g Kaplan-Meier or hazard
info_distribution  distribution name eg. for logistic regression -binomial
covariates         list of covariates - calculations to be done before passing
timevar_survival  time variable for survival analysis, default is NA
cluster_var       cluster variable for survival analysis

```

### Details

This function helps to get the parameter values after the survival analysis Takes into account many different methods like KM.FH, Cox proportional etc. and then calls appropriate functions to do the survival analysis

### Value

the results of the regression analysis

### Examples

```

data_for_survival <- survival::aml
surv_estimated_aml <- use_survival_analysis("status", data_for_survival,
  "x",
  info_get_method = "parametric", info_distribution = "weibull",
  covariates = NA, "time")

```

---

utility\_data.df

*utility matrix*

---

### Description

utility matrix

**Usage**

```
utility_data.df
```

**Format**

A 11 by 2 dataframe

**Source**

```
created on Nov 26, 2019 from tmat <- rbind(c(1, 2), c(3, 4)) colnames(tmat) <- rownames(tmat) <-
c("Healthy", "Dead") tm <- transition_matrix(2, tmat, c(0.5, 0.5, 0, 1)) a <- health_state("Healthy",
1, 1, FALSE) b <- health_state("Dead", 1, 0, TRUE) health_states <- combine_state(a, b) this.strategy
<- strategy(tm, health_states, "intervention")
```

---

value\_ADL\_scores\_IPD *Function to convert ADL scores to a T score*

---

**Description**

Function to convert ADL scores to a T score

**Usage**

```
value_ADL_scores_IPD(
  ind_part_data,
  adl_related_words,
  adl_nrcode,
  adl_scoring_table = NULL
)
```

**Arguments**

ind\_part\_data a data frame containing IPD data  
 adl\_related\_words related words to find out which columns contain adl data  
 adl\_nrcode non response code for ADL  
 adl\_scoring\_table ADL scoring table, if given as NULL use the default one

**Value**

ADL scores converted to T score included modified data, if success -1, if failure

**Examples**

```
datafile <- system.file("extdata", "trial_data.csv", package = "packDAMipd")
trial_data <- load_trial_data(datafile)
value_ADL_scores_IPD(trial_data, c("tpi"), NA, adl_scoring_table = NULL)
```



---

value\_eq5d3L\_IPD      *Function to add EQ5D3L scores to IPD data*

---

**Description**

Function to add EQ5D3L scores to IPD data

**Usage**

```
value_eq5d3L_IPD(ind_part_data, eq5d_nrcode)
```

**Arguments**

ind\_part\_data    a dataframe  
eq5d\_nrcode      non response code for EQ5D3L, default is NA

**Value**

qaly included modified data, if success -1, if failure

**Source**

[http://eprints.whiterose.ac.uk/121473/1/Devlin\\_et\\_al-2017-Health\\_Economics.pdf](http://eprints.whiterose.ac.uk/121473/1/Devlin_et_al-2017-Health_Economics.pdf)

**Examples**

```
datafile <- system.file("extdata", "trial_data.csv", package = "packDAMipd")  
trial_data <- load_trial_data(datafile)  
value_eq5d5L_IPD(trial_data, NA)
```

---

value\_eq5d5L\_IPD      *Function to add EQ5D5L scores to IPD data*

---

**Description**

Function to add EQ5D5L scores to IPD data

**Usage**

```
value_eq5d5L_IPD(ind_part_data, eq5d_nrcode)
```

**Arguments**

ind\_part\_data    a dataframe  
eq5d\_nrcode      non response code for EQ5D5L, default is NA

**Value**

qaly included modified data, if success -1, if failure

**Source**

[http://eprints.whiterose.ac.uk/121473/1/Devlin\\_et\\_al-2017-Health\\_Economics.pdf](http://eprints.whiterose.ac.uk/121473/1/Devlin_et_al-2017-Health_Economics.pdf)

**Examples**

```
datafile <- system.file("extdata", "trial_data.csv", package = "packDAMipd")
trial_data <- load_trial_data(datafile)
value_eq5d5L_IPD(trial_data, NA)
```

---

value\_promis3a\_scores\_IPD

*Function to convert promis3a scores to a T score*

---

**Description**

Function to convert promis3a scores to a T score

**Usage**

```
value_promis3a_scores_IPD(
  ind_part_data,
  promis3a_related_words,
  promis3a_nrancode,
  promis3a_scoring_table = NULL
)
```

**Arguments**

ind\_part\_data a data frame containing IPD data  
 promis3a\_related\_words related words to find out which columns contain promis3a data  
 promis3a\_nrancode non response code for promis3a  
 promis3a\_scoring\_table promis3a scoring table, if given as NULL use the default one

**Value**

promis3a scores converted to T score included modified data, if success -1, if failure

**Examples**

```
trial_data <- data.frame("tpi.q1" = c(1, 2),
  "tpi.q2" = c(1, 2), "tpi.q3" = c(1, 2))
results <- value_promis3a_scores_IPD(trial_data, c("tpi"), NA, NULL)
```

---

value_Shows_IPD	<i>Function to estimate the cost of tablets taken (from IPD)</i>
-----------------	--

---

**Description**

Function to estimate the cost of tablets taken (from IPD)

**Usage**

```
value_Shows_IPD(ind_part_data, shows_related_words, shows_nrcode)
```

**Arguments**

ind\_part\_data a data frame containing IPD  
shows\_related\_words  
a data frame containing IPD  
shows\_nrcode non response code for ADL, default is NA

**Value**

sum of scores, if success -1, if failure

**Examples**

```
datafile <- system.file("extdata", "trial_data.csv", package = "packDAMipd")  
trial_data <- load_trial_data(datafile)  
value_Shows_IPD(trial_data, "qsy", NA)
```

---

word2num	<i>Function to check the variable null or NA</i>
----------	--

---

**Description**

Function to check the variable null or NA

**Usage**

```
word2num(word)
```

**Arguments**

word word for the number

**Details**

<https://stackoverflow.com/questions/18332463/convert-written-number-to-number-in-r>

**Value**

return the number

**Examples**

```
answer <- word2num("one forty one")
answer <- word2num("forty one and five hundred")
answer <- word2num("five thousand two hundred and eight")
```

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