# Package 'random.cdisc.data'

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Type Package

```
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random.cdisc.data-package

random.cdisc.data Package

### **Description**

Package to create random SDTM and ADAM datasets.

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### See Also

Useful links:

- https://insightsengineering.github.io/random.cdisc.data/
- https://github.com/insightsengineering/random.cdisc.data/
- Report bugs at https://github.com/insightsengineering/random.cdisc.data/issues

apply\_metadata

Apply Metadata

#### **Description**

Apply label and variable ordering attributes to domains.

#### Usage

```
apply_metadata(
   df,
   filename,
   add_adsl = TRUE,
   adsl_filename = "metadata/ADSL.yml"
)
```

#### **Arguments**

df (data.frame)

Data frame to which metadata is applied.

filename (yaml)

File containing domain metadata.

add\_adsl (logical)

Should ADSL data be merged to domain.

 ${\tt adsl\_filename} \quad ({\tt yaml})$ 

File containing ADSL metadata.

#### Value

Data frame with metadata applied.

#### **Examples**

```
seed <- 1
adsl <- radsl(seed = seed)
adsub <- radsub(adsl, seed = seed)
yaml_path <- file.path(path.package("random.cdisc.data"), "inst", "metadata")
adsl <- apply_metadata(adsl, file.path(yaml_path, "ADSL.yml"), FALSE)
adsub <- apply_metadata(</pre>
```

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```
adsub, file.path(yaml_path, "ADSUB.yml"), TRUE,
file.path(yaml_path, "ADSL.yml")
)
```

cadab

Cached ADAB

### **Description**

Cached ADAB data generated with seed = 1

### Usage

```
data(cadab)
```

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 6916 rows and 21 columns.

cadae

Cached ADAE

# Description

Cached ADAE data generated with seed = 1

### Usage

data(cadae)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 1934 rows and 92 columns.

cadaette

Cached ADAETTE

# Description

Cached ADAETTE data generated with seed = 1

### Usage

```
data(cadaette)
```

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 3600 rows and 66 columns.

6 cadeg

cadcm

Cached ADCM

### **Description**

Cached ADCM data generated with seed = 1

# Usage

data(cadcm)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 3685 rows and 83 columns.

caddv

Cached ADDV

# Description

Cached ADDV data generated with seed = 1

# Usage

data(caddv)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 119 rows and 66 columns.

cadeg

Cached ADEG

# Description

Cached ADEG data generated with seed = 1

# Usage

data(cadeg)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 13600 rows and 88 columns.

cadex 7

cadex

Cached ADEX

### **Description**

Cached ADEX data generated with seed = 1

# Usage

data(cadex)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 6400 rows and 79 columns.

cadhy

Cached ADHY

# **Description**

Cached ADHY data generated with seed = 1

### Usage

data(cadhy)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 20000 rows and 71 columns.

cadlb

Cached ADLB

### **Description**

Cached ADLB data generated with seed = 1

# Usage

data(cadlb)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 8400 rows and 102 columns.

8 cadpp

cadmh

Cached ADMH

#### **Description**

Cached ADMH data generated with seed = 1

### Usage

data(cadmh)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 1934 rows and 67 columns.

cadpc

Cached ADPC

# Description

Cached ADPC data generated with seed = 1

### Usage

data(cadpc)

### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 6640 rows and 72 columns.

cadpp

Cached ADPP

### **Description**

Cached ADPP data generated with seed = 1

# Usage

data(cadpp)

#### **Format**

An object of class data. frame with 26268 rows and 68 columns.

cadqlqc 9

cadqlqc

Cached ADQLQC

# Description

Cached ADQLQC data generated with seed = 1

### Usage

data(cadqlqc)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 116803 rows and 50 columns.

cadqs

Cached ADQS

# **Description**

Cached ADQS data generated with seed = 1

### Usage

data(cadqs)

### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 14000 rows and 73 columns.

cadrs

Cached ADRS

### **Description**

Cached ADRS data generated with seed = 1

# Usage

data(cadrs)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 3200 rows and 65 columns.

10 cadtr

cadsl

Cached ADSL

# Description

Cached ADSL data generated with seed = 1

# Usage

data(cadsl)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 400 rows and 55 columns.

cadsub

Cached ADSUB

# Description

Cached ADSUB data generated with seed = 1

### Usage

data(cadsub)

### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 2000 rows and 65 columns.

cadtr

Cached ADTR

# Description

Cached ADTR data generated with seed = 1

### Usage

data(cadtr)

#### **Format**

An object of class data. frame with 2800 rows and 76 columns.

cadtte 11

cadtte

Cached ADTTE

# Description

Cached ADTTE data generated with seed = 1

### Usage

data(cadtte)

#### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 2000 rows and 67 columns.

cadvs

Cached ADVS

### **Description**

Cached ADVS data generated with seed = 1

### Usage

data(cadvs)

### **Format**

An object of class tbl\_df (inherits from tbl, data.frame) with 16800 rows and 87 columns.

mutate\_na

Replace Values with NA

# Description

### [Stable]

Replace column values with NAs.

```
mutate_na(ds, na_vars = NULL, na_percentage = 0.05)
```

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#### **Arguments**

ds (data.frame)
Any data set.

na\_vars (list)
A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)
The seed to be used for this element - can be NA.

• percentage (proportion)
Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

na\_percentage (proportion)
Default percentage of values to be replaced by NA.

#### Value

dataframe without NA values.

radab

Anti-Drug Antibody Analysis Dataset (ADAB)

#### **Description**

#### [Stable]

Function for generating a random Anti-Drug Antibody Analysis Dataset for a given Subject-Level Analysis Dataset and Pharmacokinetics Analysis Dataset.

```
radab(
  adsl,
  adpc,
  constants = c(D = 100, ka = 0.8, ke = 1),
 paramcd = c("R1800000", "RESULT1", "R1800001", "RESULT2", "ADASTAT1", "INDUCD1",
  "ENHANC1", "TRUNAFF1", "EMERNEG1", "EMERPOS1", "PERSADA1", "TRANADA1", "BFLAG1",
  "TIMADA1", "ADADUR1", "ADASTAT2", "INDUCD2", "ENHANC2", "EMERNEG2", "EMERPOS2",
    "BFLAG2", "TRUNAFF2"),
  param = c("Antibody titer units", "ADA interpreted per sample result",
    "Neutralizing Antibody titer units", "NAB interpreted per sample result",
   "ADA Status of a patient", "Treatment induced ADA", "Treatment enhanced ADA",
    "Treatment unaffected", "Treatment Emergent - Negative",
  "Treatment Emergent - Positive", "Persistent ADA", "Transient ADA", "Baseline".
    "Time to onset of ADA", "ADA Duration", "NAB Status of a patient",
    "Treatment induced ADA, Neutralizing Antibody",
    "Treatment enhanced ADA, Neutralizing Antibody",
```

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#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

adpc (data.frame)

Pharmacokinetics Analysis Dataset.

constants (character vector)

Constant parameters to be used in formulas for creating analysis values.

paramcd (character vector)

Parameter code values.

param (character vector)

Parameter values.

avalu (character)

Analysis value units.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

cached

boolean whether the cached ADAB data cadab should be returned or new data should be generated. If set to TRUE then the other arguments to radab will be ignored.

#### Details

One record per study per subject per parameter per time point: "R1800000", "RESULT1", "R1800001", "RESULT2".

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#### Value

```
data.frame
```

### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adpc <- radpc(adsl, seed = 2, duration = 9 * 7)

adab <- radab(adsl, adpc, seed = 2)
adab</pre>
```

radae

Adverse Event Analysis Dataset (ADAE)

### Description

### [Stable]

Function for generating random Adverse Event Analysis Dataset for a given Subject-Level Analysis Dataset.

# Usage

```
radae(
  adsl,
  max_n_aes = 10L,
  lookup = NULL,
  lookup_aag = NULL,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(AEBODSYS = c(NA, 0.1), AEDECOD = c(1234, 0.1), AETOXGR = c(1234, 0.1)),
  cached = FALSE
)
```

# Arguments

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

max\_n\_aes (integer)

Maximum number of AEs per patient. Defaults to 10.

lookup (data.frame)

Additional parameters.

lookup\_aag (data.frame)

Additional metadata parameters.

seed (numeric)

Seed to use for reproducible random number generation.

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na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

cached

boolean whether the cached ADAE data cadae should be returned or new data should be generated. If set to TRUE then the other arguments to radae will be ignored.

#### **Details**

One record per each record in the corresponding SDTM domain.

Keys: STUDYID, USUBJID, ASTDTM, AETERM, AESEQ

#### Value

data.frame

#### **Examples**

```
adsl <- radsl(N = 10, study_duration = 2, seed = 1)
adae <- radae(adsl, seed = 2)</pre>
adae
# Add metadata.
aag <- utils::read.table(</pre>
  sep = ",", header = TRUE,
  text = paste(
    "NAMVAR, SRCVAR, GRPTYPE, REFNAME, REFTERM, SCOPE",
    "CQ01NAM, AEDECOD, CUSTOM, D. 2.1.5.3/A.1.1.1.1 AESI, dcd D. 2.1.5.3, ",
    "CQ01NAM, AEDECOD, CUSTOM, D.2.1.5.3/A.1.1.1.1 AESI, dcd A.1.1.1.1,",
    "SMQ01NAM, AEDECOD, SMQ, C.1.1.1.3/B.2.2.3.1 AESI, dcd C.1.1.1.3, BROAD",
    "SMQ01NAM, AEDECOD, SMQ, C.1.1.1.3/B.2.2.3.1 AESI, dcd B.2.2.3.1, BROAD",
    "SMQ02NAM, AEDECOD, SMQ, Y.9.9.9.9/Z.9.9.9.9 AESI, dcd Y.9.9.9.9, NARROW",
    "SMQ02NAM, AEDECOD, SMQ, Y.9.9.9.9/Z.9.9.9.9 AESI, dcd Z.9.9.9.9, NARROW",
    sep = "\n"
  ), stringsAsFactors = FALSE
adae <- radae(adsl, lookup_aag = aag)</pre>
with(
  adae,
  cbind(
```

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```
table(AEDECOD, SMQ01NAM),
  table(AEDECOD, CQ01NAM)
)
```

radaette

*Time to Adverse Event Analysis Dataset (ADAETTE)* 

# Description

#### [Stable]

Function to generate random Time-to-AE Dataset for a given Subject-Level Analysis Dataset.

#### Usage

```
radaette(
  adsl,
  event_descr = NULL,
  censor_descr = NULL,
  lookup = NULL,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(CNSR = c(NA, 0.1), AVAL = c(1234, 0.1)),
  cached = FALSE
)
```

### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

event\_descr (character vector)

Descriptions of events. Defaults to NULL.

censor\_descr (character vector)

Descriptions of censors. Defaults to NULL.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

 ${\tt na\_percentage} \quad ({\tt proportion})$ 

Default percentage of values to be replaced by NA.

na\_vars (list

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

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percentage (proportion)
 Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

cached

boolean whether the cached ADAETTE data cadaette should be returned or new data should be generated. If set to TRUE then the other arguments to radaette will be ignored.

#### **Details**

```
Keys: STUDYID, USUBJID, PARAMCD
```

#### Value

```
data.frame
```

#### Author(s)

Xiuting Mi

#### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adaette <- radaette(adsl, seed = 2)
adaette</pre>
```

radcm

Previous and Concomitant Medications Analysis Dataset (ADCM)

### **Description**

### [Stable]

Function for generating random Concomitant Medication Analysis Dataset for a given Subject-Level Analysis Dataset.

```
radcm(
  adsl,
  max_n_cms = 10L,
  lookup = NULL,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(CMCLAS = c(NA, 0.1), CMDECOD = c(1234, 0.1), ATIREL = c(1234, 0.1)),
  who_coding = FALSE,
  cached = FALSE
)
```

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#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

max\_n\_cms (integer)

Maximum number of concomitant medications per patient. Defaults to 10.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each

element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is  $\frac{1}{2}$ 

used as a default.

who\_coding (flag)

Whether WHO coding (with multiple paths per medication) should be used.

cached boolean whether the cached ADCM data cadcm should be returned or new data

should be generated. If set to TRUE then the other arguments to radcm will be

ignored.

### Details

One record per each record in the corresponding SDTM domain.

```
Keys: STUDYID, USUBJID, ASTDTM, CMSEQ
```

#### Value

```
data.frame
```

### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adcm <- radcm(adsl, seed = 2)
adcm
adcm_who <- radcm(adsl, seed = 2, who_coding = TRUE)
adcm_who</pre>
```

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raddv

Protocol Deviations Analysis Dataset (ADDV)

#### **Description**

#### [Stable]

Function for generating random Protocol Deviations Analysis Dataset for a given Subject-Level Analysis Dataset.

# Usage

```
raddv(
  adsl,
  max_n_dv = 3L,
  p_dv = 0.15,
  lookup = NULL,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(ASTDT = c(seed = 1234, percentage = 0.1), DVCAT = c(seed = 1234, percentage = 0.1)),
  cached = FALSE
)
```

#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

max\_n\_dv (integer)

Maximum number of deviations per patient. Defaults to 3.

p\_dv (proportion)

Probability of a patient having protocol deviations.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

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cached

boolean whether the cached ADDV data caddv should be returned or new data should be generated. If set to TRUE then the other arguments to raddv will be ignored.

#### **Details**

One record per each record in the corresponding SDTM domain.

```
Keys: STUDYID, USUBJID, ASTDT, DVTERM, DVSEQ
```

#### Value

```
data.frame
```

#### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
addv <- raddv(adsl, seed = 2)
addv</pre>
```

radeg

ECG Analysis Dataset (ADEG)

### **Description**

### [Stable]

Function for generating random dataset from ECG Analysis Dataset for a given Subject-Level Analysis Dataset.

```
radeg(
           adsl,
           egcat = c("INTERVAL", "INTERVAL", "MEASUREMENT", "FINDING"),
         param = c("QT Duration", "RR Duration", "Heart Rate", "ECG Interpretation"),
paramcd = c("QT", "RR", "HR", "ECGINTP"),
           paramu = c("msec", "msec", "beats/min", ""),
           visit_format = "WEEK",
           n_{assessments} = 5L,
           n_{days} = 5L
          max_n_eg = 10L,
          lookup = NULL,
           seed = NULL,
         na_percentage = 0,
       na_vars = list(ABLFL = c(1235, 0.1), BASE = c(NA, 0.1), BASEC = c(NA, 0.1), CHG = 
                      c(1234, 0.1), PCHG = c(1234, 0.1)),
           cached = FALSE
)
```

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#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

(character vector) egcat

EG category values.

(character vector) param

Parameter values.

(character vector) paramcd

Parameter code values.

(character vector) paramu

Parameter unit values.

visit\_format (character)

Type of visit. Options are "WEEK" and "CYCLE".

n\_assessments

Number of weeks or cycles.

n\_days (integer)

Number of days in each cycle (only used if visit\_format is "CYCLE").

(integer) max\_n\_eg

Maximum number of EG results per patient. Defaults to 10.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each

element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADEG data cadeg should be returned or new data

should be generated. If set to TRUE then the other arguments to radeg will be

ignored.

#### **Details**

One record per subject per parameter per analysis visit per analysis date.

Keys: STUDYID, USUBJID, PARAMCD, BASETYPE, AVISITN, ATPTN, DTYPE, ADTM, EGSEQ, ASPID

#### Value

data.frame

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#### Author(s)

tomlinsj, npaszty, Xuefeng Hou, dipietrc

#### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adeg <- radeg(adsl, visit_format = "WEEK", n_assessments = 7L, seed = 2)
adeg
adeg <- radeg(adsl, visit_format = "CYCLE", n_assessments = 2L, seed = 2)
adeg</pre>
```

radex

Exposure Analysis Dataset (ADEX)

#### Description

#### [Stable]

Function for generating random Exposure Analysis Dataset for a given Subject-Level Analysis Dataset.

```
radex(
  adsl,
  param = c("Dose administered during constant dosing interval",
    "Number of doses administered during constant dosing interval",
    "Total dose administered", "Total number of doses administered"),
 paramcd = c("DOSE", "NDOSE", "TDOSE", "TNDOSE"),
paramu = c("mg", " ", "mg", " "),
  parcat1 = c("INDIVIDUAL", "OVERALL"),
  parcat2 = c("Drug A", "Drug B"),
  visit_format = "WEEK",
  n_assessments = 5L,
  n_{days} = 5L,
 max_n_{exs} = 6L,
  lookup = NULL,
  seed = NULL,
  na_percentage = 0,
 na_vars = list(AVAL = c(NA, 0.1), AVALU = c(NA), 0.1),
  cached = FALSE
)
```

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#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

param (character vector)

Parameter values.

paramcd (character vector)

Parameter code values.

paramu (character vector)

Parameter unit values.

parcat1 (character vector)

Dose amount categories. Defaults to "Individual" and "Overall".

parcat2 (character vector)

Types of drug received. Defaults to "Drug A" and "Drug B".

visit\_format (character)

Type of visit. Options are "WEEK" and "CYCLE".

n\_assessments (integer)

Number of weeks or cycles.

n\_days (integer)

Number of days in each cycle (only used if visit\_format is "CYCLE").

max\_n\_exs (integer)

Maximum number of exposures per patient. Defaults to 6.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADEX data cadex should be returned or new data

should be generated. If set to TRUE then the other arguments to radex will be

ignored.

### Details

One record per each record in the corresponding SDTM domain.

Keys: STUDYID, USUBJID, EXSEQ, PARAMCD, PARCAT1, ASTDTM, AENDTM, ASTDY, AENDY, AVISITN, EXDOSFRQ, EXROUTE, VISIT, VISITDY, EXSTDTC, EXENDTC, EXSTDY, EXENDY

24 radhy

#### Value

```
data.frame
```

#### **Examples**

```
ads1 <- rads1(N = 10, study_duration = 2, seed = 1)
adex <- radex(ads1, seed = 2)
adex</pre>
```

radhv

Hy's Law Analysis Dataset (ADHY)

#### **Description**

#### [Stable]

Function for generating a random Hy's Law Analysis Dataset for a given Subject-Level Analysis Dataset.

```
radhy(
  adsl,
  param = c("TBILI <= 2 times ULN and ALT value category",
    "TBILI > 2 times ULN and AST value category",
    "TBILI > 2 times ULN and ALT value category"
    "TBILI <= 2 times ULN and AST value category"
    "TBILI > 2 times ULN and ALKPH <= 2 times ULN and ALT value category",
    "TBILI > 2 times ULN and ALKPH <= 2 times ULN and AST value category",
    "TBILI > 2 times ULN and ALKPH <= 5 times ULN and ALT value category"
    "TBILI > 2 times ULN and ALKPH <= 5 times ULN and AST value category",
  "TBILI <= 2 times ULN and two consecutive elevations of ALT in relation to ULN",
   "TBILI > 2 times ULN and two consecutive elevations of AST in relation to ULN",
  "TBILI <= 2 times ULN and two consecutive elevations of AST in relation to ULN"
  "TBILI > 2 times ULN and two consecutive elevations of ALT in relation to ULN",
  "TBILI > 2 times ULN and two consecutive elevations of ALT in relation to Baseline",
  "TBILI <= 2 times ULN and two consecutive elevations of ALT in relation to Baseline".
  "TBILI > 2 times ULN and two consecutive elevations of AST in relation to Baseline",
  "TBILI <= 2 times ULN and two consecutive elevations of AST in relation to Baseline",
    "ALT > 3 times ULN by Period", "AST > 3 times ULN by Period",
   "ALT or AST > 3 times ULN by Period", "ALT > 3 times Baseline by Period",
   "AST > 3 times Baseline by Period", "ALT or AST > 3 times Baseline by Period"),
 paramcd = c("BLAL", "BGAS", "BGAL", "BLAS", "BA2AL", "BA2AS", "BA5AL", "BA5AS",
  "BL2AL2CU", "BG2AS2CU", "BL2AS2CU", "BG2AL2CU", "BG2AL2CB", "BL2AL2CB", "BG2AS2CB",
```

radhy 25

```
"BL2AS2CB", "ALTPULN", "ASTPULN", "ALTASTPU", "ALTPBASE", "ASTPBASE", "ALTASTPB"), seed = NULL, cached = FALSE
)
```

### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

param (character vector)

Parameter values.

paramcd (character vector)

Parameter code values.

seed (numeric)

Seed to use for reproducible random number generation.

cached boolean whether the cached ADHY data cadhy should be returned or new data

should be generated. If set to TRUE then the other arguments to radhy will be

ignored.

#### **Details**

One record per subject per parameter per analysis visit per analysis date.

Keys: STUDYID, USUBJID, PARAMCD, AVISITN, ADTM, SRCSEQ

#### Value

data.frame

### Author(s)

wojciakw

### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adhy <- radhy(adsl, seed = 2)
adhy</pre>
```

26 radlb

radlb

Laboratory Data Analysis Dataset (ADLB)

#### **Description**

### [Stable]

Function for generating a random Laboratory Data Analysis Dataset for a given Subject-Level Analysis Dataset.

### Usage

```
radlb(
 adsl,
 lbcat = c("CHEMISTRY", "CHEMISTRY", "IMMUNOLOGY"),
 param = c("Alanine Aminotransferase Measurement", "C-Reactive Protein Measurement",
    "Immunoglobulin A Measurement"),
 paramcd = c("ALT", "CRP", "IGA"),
 paramu = c("U/L", "mg/L", "g/L"),
 aval_mean = c(18, 9, 2.9),
 visit_format = "WEEK",
  n_assessments = 5L,
  n_{days} = 5L,
 max_n_lbs = 10L,
 lookup = NULL,
  seed = NULL,
 na_percentage = 0,
 na_vars = list(LOQFL = c(NA, 0.1), ABLFL2 = c(1234, 0.1), ABLFL = c(1235, 0.1), BASE2 =
   c(NA, 0.1), BASE = c(NA, 0.1), CHG2 = c(1235, 0.1), PCHG2 = c(1235, 0.1), CHG =
   c(1234, 0.1), PCHG = c(1234, 0.1)),
 cached = FALSE
```

### **Arguments**

adsl	(data.frame) Subject-Level Analysis Dataset (ADSL).
lbcat	(character vector) LB category values.
param	(character vector) Parameter values.
paramcd	(character vector) Parameter code values.
paramu	(character vector) Parameter unit values.

radlb 27

aval\_mean (numeric vector)

Mean values corresponding to each parameter.

visit\_format (character)

Type of visit. Options are "WEEK" and "CYCLE".

n\_assessments (integer)

Number of weeks or cycles.

n\_days (integer)

Number of days in each cycle (only used if visit\_format is "CYCLE").

max\_n\_lbs (integer)

Maximum number of labs per patient. Defaults to 10.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADLB data cadlb should be returned or new data

should be generated. If set to TRUE then the other arguments to radlb will be

ignored.

### **Details**

One record per subject per parameter per analysis visit per analysis date.

Keys: STUDYID, USUBJID, PARAMCD, BASETYPE, AVISITN, ATPTN, DTYPE, ADTM, LBSEQ, ASPID

#### Value

```
data.frame
```

#### Author(s)

```
tomlinsj, npaszty, Xuefeng Hou
```

### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adlb <- radlb(adsl, visit_format = "WEEK", n_assessments = 7L, seed = 2)</pre>
```

28 radmh

```
adlb
adlb <- radlb(adsl, visit_format = "CYCLE", n_assessments = 2L, seed = 2)
adlb</pre>
```

radmh

Medical History Analysis Dataset (ADMH)

# Description

#### [Stable]

Function for generating a random Medical History Analysis Dataset for a given Subject-Level Analysis Dataset.

### Usage

```
radmh(
  adsl,
  max_n_mhs = 10L,
  lookup = NULL,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(MHBODSYS = c(NA, 0.1), MHDECOD = c(1234, 0.1)),
  cached = FALSE
)
```

### Arguments

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

max\_n\_mhs (integer)

Maximum number of MHs per patient. Defaults to 10.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

radpc 29

cached

boolean whether the cached ADMH data cadmh should be returned or new data should be generated. If set to TRUE then the other arguments to radmh will be ignored.

#### **Details**

One record per each record in the corresponding SDTM domain.

```
Keys: STUDYID, USUBJID, ASTDTM, MHSEQ
```

#### Value

```
data.frame
```

#### **Examples**

```
adsl <- radsl(N = 10, study_duration = 2, seed = 1)
admh <- radmh(adsl, seed = 2)
admh</pre>
```

radpc

Pharmacokinetics Analysis Dataset (ADPC)

# Description

### [Stable]

Function for generating a random Pharmacokinetics Analysis Dataset for a given Subject-Level Analysis Dataset.

```
radpc(
  adsl,
  avalu = "ug/mL",
  constants = c(D = 100, ka = 0.8, ke = 1),
  duration = 2,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(AVAL = c(NA, 0.1)),
  cached = FALSE
)
```

30 radpc

#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

avalu (character)

Analysis value units.

constants (character vector)

Constant parameters to be used in formulas for creating analysis values.

duration (numeric)

Duration in number of days.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADPC data cadpc should be returned or new data

should be generated. If set to TRUE then the other arguments to radpc will be

ignored.

#### **Details**

One record per study, subject, parameter, and time point.

### Value

```
data.frame
```

### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adpc <- radpc(adsl, seed = 2)
adpc
adpc <- radpc(adsl, seed = 2, duration = 3)
adpc</pre>
```

radpp 31

radpp

Pharmacokinetics Parameters Dataset (ADPP)

#### **Description**

#### [Stable]

Function for generating a random Pharmacokinetics Parameters Dataset for a given Subject-Level Analysis Dataset.

#### Usage

```
radpp(
 adsl,
 ppcat = c("Plasma Drug X", "Plasma Drug Y", "Metabolite Drug X", "Metabolite Drug Y"),
 ppspec = c("Plasma", "Plasma", "Plasma", "Matrix of PD", "Matrix of PD", "Urine",
    "Urine", "Urine", "Urine"),
 paramcd = c("AUCIFO", "CMAX", "CLO", "RMAX", "TON", "RENALCL", "RENALCLD", "RCAMINT",
    "RCPCINT"),
 param = c("AUC Infinity Obs", "Max Conc", "Total CL Obs", "Time of Maximum Response",
   "Time to Onset", "Renal CL", "Renal CL Norm by Dose", "Amt Rec from T1 to T2",
    "Pct Rec from T1 to T2"),
 paramu = c("day*ug/mL", "ug/mL", "ml/day/kg", "hr", "hr", "L/hr", "L/hr/mg", "mg",
    "%"),
 aval_mean = c(200, 30, 5, 10, 3, 0.05, 0.005, 1.5613, 15.65),
 visit_format = "CYCLE",
 n_{days} = 2L
  seed = NULL,
 na_percentage = 0,
 na\_vars = list(AVAL = c(NA, 0.1)),
 cached = FALSE
)
```

### Arguments

adsl	(data.frame) Subject-Level Analysis Dataset (ADSL).
ppcat	(character vector) Categories of parameters.
ppspec	(character vector) Specimen material types.
paramcd	(character vector) Parameter code values.
param	(character vector) Parameter values.
paramu	(character vector) Parameter unit values.

32 radqlqc

aval\_mean (numeric vector)

Mean values corresponding to each parameter.

 $\verb|visit_format| (\verb|character|)|$ 

Type of visit. Options are "WEEK" and "CYCLE".

n\_days (integer)

Number of days in each cycle (only used if visit\_format is "CYCLE").

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADPP data cadpp should be returned or new data

should be generated. If set to TRUE then the other arguments to radpp will be

ignored.

#### **Details**

One record per study, subject, parameter category, parameter and visit.

#### Value

```
data.frame
```

#### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adpp <- radpp(adsl, seed = 2)
adpp</pre>
```

radqlqc

EORTC QLQ-C30 V3 Analysis Dataset (ADQLQC)

#### **Description**

#### [Stable]

Function for generating a random EORTC QLQ-C30 V3 Analysis Dataset for a given Subject-Level Analysis Dataset.

radqs 33

#### Usage

```
radqlqc(adsl, percent, number, seed = NULL, cached = FALSE)
```

#### Arguments

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

percent (numeric)

Completion - Completed at least y percent of questions, 1 record per visit

number (numeric)

Completion - Completed at least x question(s), 1 record per visit

seed (numeric)

Seed to use for reproducible random number generation.

cached boolean whether the cached ADQLQC data cadqlqc should be returned or new

data should be generated. If set to TRUE then the other arguments to radqlqc

will be ignored.

#### **Details**

Keys: STUDYID, USUBJID, PARCAT1N, PARAMCD, BASETYPE, AVISITN, ATPTN, ADTM, QSSEQ

#### Value

```
data.frame
```

### **Examples**

```
adsl <- radsl(N = 10, study_duration = 2, seed = 1)
adqlqc <- radqlqc(adsl, seed = 1, percent = 80, number = 2)
adqlqc</pre>
```

radqs

Questionnaires Analysis Dataset (ADQS)

# Description

#### [Stable]

Function for generating a random Questionnaires Analysis Dataset for a given Subject-Level Analysis Dataset.

34 radgs

#### **Usage**

```
radqs(
    adsl,
    param = c("BFI All Questions", "Fatigue Interference",
        "Function/Well-Being (GF1,GF3,GF7)", "Treatment Side Effects (GP2,C5,GP5)",
        "FKSI-19 All Questions"),
    paramcd = c("BFIALL", "FATIGI", "FKSI-FWB", "FKSI-TSE", "FKSIALL"),
    visit_format = "WEEK",
    n_assessments = 5L,
    n_days = 5L,
    seed = NULL,
    na_percentage = 0,
    na_vars = list(LOQFL = c(NA, 0.1), ABLFL2 = c(1234, 0.1), ABLFL = c(1235, 0.1), CHG2 =
        c(1235, 0.1), PCHG2 = c(1235, 0.1), CHG = c(1234, 0.1), PCHG = c(1234, 0.1)),
    cached = FALSE
)
```

#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

param (character vector)

Parameter values.

paramcd (character vector)

Parameter code values.

visit\_format (character)

Type of visit. Options are "WEEK" and "CYCLE".

n\_assessments (integer)

Number of weeks or cycles.

n\_days (integer)

Number of days in each cycle (only used if visit\_format is "CYCLE").

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

cached

boolean whether the cached ADQS data cadqs should be returned or new data should be generated. If set to TRUE then the other arguments to radqs will be ignored.

radrs 35

#### **Details**

One record per subject per parameter per analysis visit per analysis date.

```
Keys: STUDYID, USUBJID, PARAMCD, AVISITN
```

#### Value

```
data.frame
```

### Author(s)

npaszty

# **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adqs <- radqs(adsl, visit_format = "WEEK", n_assessments = 7L, seed = 2)
adqs <- radqs(adsl, visit_format = "CYCLE", n_assessments = 3L, seed = 2)
adqs</pre>
```

radrs

Tumor Response Analysis Dataset (ADRS)

# Description

### [Stable]

Function for generating a random Tumor Response Analysis Dataset for a given Subject-Level Analysis Dataset.

```
radrs(
  adsl,
  avalc = NULL,
  lookup = NULL,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(AVISIT = c(NA, 0.1), AVAL = c(1234, 0.1), AVALC = c(1234, 0.1)),
  cached = FALSE
)
```

36 radrs

### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

avalc (character vector)

Analysis value categories.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADRS data cadrs should be returned or new data

should be generated. If set to TRUE then the other arguments to radrs will be

ignored.

### **Details**

One record per subject per parameter per analysis visit per analysis date. SDTM variables are populated on new records coming from other single records. Otherwise, SDTM variables are left blank.

Keys: STUDYID, USUBJID, PARAMCD, AVISITN, ADT, RSSEQ

### Value

```
data.frame
```

### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adrs <- radrs(adsl, seed = 2)
adrs</pre>
```

radsaftte 37

radsaftte

Time to Safety Event Analysis Dataset (ADSAFTTE)

## Description

Function to generate random Time-to-Safety Event Dataset for a given Subject-Level Analysis Dataset.

## Usage

```
radsaftte(adsl, ...)
```

## **Arguments**

```
adsl (data.frame)
```

Subject-Level Analysis Dataset (ADSL).

... Additional arguments to be passed to radaette

#### **Details**

Keys: STUDYID, USUBJID, PARAMCD

#### Value

data.frame

#### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adsaftte <- radsaftte(adsl, seed = 2)
adsaftte</pre>
```

radsl

Subject-Level Analysis Dataset (ADSL)

# Description

## [Stable]

The Subject-Level Analysis Dataset (ADSL) is used to provide the variables that describe attributes of a subject. ADSL is a source for subject-level variables used in other analysis data sets, such as population flags and treatment variables. There is only one ADSL per study. ADSL and its related metadata are required in a CDISC-based submission of data from a clinical trial even if no other analysis data sets are submitted.

38 radsl

#### Usage

```
radsl(
  N = 400,
  study_duration = 2,
  seed = NULL,
  with_trt02 = TRUE,
  na_percentage = 0,
  na_vars = list(AGE = NA, SEX = NA, RACE = NA, STRATA1 = NA, STRATA2 = NA, BMRKR1 =
    c(seed = 1234, percentage = 0.1), BMRKR2 = c(1234, 0.1), BEP01FL = NA),
  ae_withdrawal_prob = 0.05,
  cached = FALSE
)
```

#### **Arguments**

N (numeric)

Number of patients.

study\_duration (numeric)

Duration of study in years.

seed (numeric)

Seed to use for reproducible random number generation.

with\_trt02 (logical)

Should period 2 be added.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

ae\_withdrawal\_prob

(proportion)

Probability that there is at least one Adverse Event leading to the withdrawal of

a study drug.

cached

boolean whether the cached ADSL data cads1 should be returned or new data should be generated. If set to TRUE then the other arguments to rads1 will be

ignored.

#### **Details**

One record per subject.

Keys: STUDYID, USUBJID

radsub 39

#### Value

```
data.frame
```

## **Examples**

```
adsl <- radsl(N = 10, study_duration = 2, seed = 1)
adsl

adsl <- radsl(
   N = 10, seed = 1,
   na_percentage = 0.1,
   na_vars = list(
     DTHDT = c(seed = 1234, percentage = 0.1),
     LSTALVDT = c(seed = 1234, percentage = 0.1)
   )
)
adsl

adsl <- radsl(N = 10, seed = 1, na_percentage = .1)
adsl</pre>
```

radsub

Subcategory Analysis Dataset (ADSUB)

#### **Description**

## [Stable]

Function for generating a random Subcategory Analysis Dataset for a given Subject-Level Analysis Dataset.

## Usage

```
radsub(
  adsl,
  param = c("Baseline Weight", "Baseline Height", "Baseline BMI", "Baseline ECOG",
      "Baseline Biomarker Mutation"),
  paramcd = c("BWGHTSI", "BHGHTSI", "BBMISI", "BECOG", "BBMRKR1"),
  seed = NULL,
  na_percentage = 0,
  na_vars = list(),
  cached = FALSE
)
```

#### **Arguments**

```
adsl (data.frame)
Subject-Level Analysis Dataset (ADSL).
```

40 radsub

param (character vector)

Parameter values.

paramcd (character vector)

Parameter code values.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each

element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADSUB data cadsub should be returned or new

data should be generated. If set to TRUE then the other arguments to radsub will

be ignored.

#### **Details**

One record per subject.

Keys: STUDYID, USUBJID, PARAMCD, AVISITN, ADTM, SRCSEQ

## Value

data.frame

## Author(s)

tomlinsj, npaszty, Xuefeng Hou, dipietrc

## **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adsub <- radsub(adsl, seed = 2)
adsub</pre>
```

radtr 41

radtr

Tumor Response Analysis Dataset (ADTR)

## **Description**

## [Stable]

Function for generating a random Tumor Response Analysis Dataset for a given Subject-Level Analysis Dataset.

## Usage

```
radtr(
  adsl,
  param = c("Sum of Longest Diameter by Investigator"),
  paramcd = c("SLDINV"),
  seed = NULL,
  cached = FALSE,
  ...
)
```

## **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

param (character vector)

Parameter values.

paramcd (character vector)

Parameter code values.

seed (numeric)

Seed to use for reproducible random number generation.

cached boolean whether the cached ADTR data cadtr should be returned or new data

should be generated. If set to TRUE then the other arguments to radtr will be

ignored.

... Additional arguments to be passed to radrs.

#### **Details**

One record per subject per parameter per analysis visit per analysis date.

Keys: STUDYID, USUBJID, PARAMCD, BASETYPE, AVISITN, DTYPE

#### Value

```
data.frame
```

42 radtte

#### Author(s)

tomlinsj, npaszty, Xuefeng Hou, dipietrc

#### **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adtr <- radtr(adsl, seed = 2)
adtr</pre>
```

radtte

Time-to-Event Analysis Dataset (ADTTE)

#### **Description**

#### [Stable]

Function for generating a random Time-to-Event Analysis Dataset for a given Subject-Level Analysis Dataset.

# Usage

```
radtte(
  adsl,
  event_descr = NULL,
  censor_descr = NULL,
  lookup = NULL,
  seed = NULL,
  na_percentage = 0,
  na_vars = list(CNSR = c(NA, 0.1), AVAL = c(1234, 0.1), AVALU = c(1234, 0.1)),
  cached = FALSE
)
```

#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

event\_descr (character vector)

Descriptions of events. Defaults to NULL.

censor\_descr (character vector)

Descriptions of censors. Defaults to NULL.

lookup (data.frame)

Additional parameters.

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

radvs 43

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is used as a default.

cached

boolean whether the cached ADTTE data cadtte should be returned or new data should be generated. If set to TRUE then the other arguments to radtte will be ignored.

#### **Details**

```
Keys: STUDYID, USUBJID, PARAMCD
```

#### Value

```
data.frame
```

## **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
adtte <- radtte(adsl, seed = 2)
adtte</pre>
```

radvs

Vital Signs Analysis Dataset (ADVS)

# Description

# [Stable]

Function for generating a random Vital Signs Analysis Dataset for a given Subject-Level Analysis Dataset.

#### Usage

```
radvs(
  adsl,
  param = c("Diastolic Blood Pressure", "Pulse Rate", "Respiratory Rate",
    "Systolic Blood Pressure", "Temperature", "Weight"),
  paramcd = c("DIABP", "PULSE", "RESP", "SYSBP", "TEMP", "WEIGHT"),
  paramu = c("Pa", "beats/min", "breaths/min", "Pa", "C", "Kg"),
  visit_format = "WEEK",
  n_assessments = 5L,
  n_days = 5L,
```

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```
seed = NULL,
na_percentage = 0,
na_vars = list(CHG2 = c(1235, 0.1), PCHG2 = c(1235, 0.1), CHG = c(1234, 0.1), PCHG =
    c(1234, 0.1), AVAL = c(123, 0.1), AVALU = c(123, 0.1)),
    cached = FALSE
)
```

#### **Arguments**

adsl (data.frame)

Subject-Level Analysis Dataset (ADSL).

param (character vector)

Parameter values.

paramcd (character vector)

Parameter code values.

paramu (character vector)

Parameter unit values.

visit\_format (character)

Type of visit. Options are "WEEK" and "CYCLE".

n\_assessments (integer)

Number of weeks or cycles.

n\_days (integer)

Number of days in each cycle (only used if visit\_format is "CYCLE").

seed (numeric)

Seed to use for reproducible random number generation.

na\_percentage (proportion)

Default percentage of values to be replaced by NA.

na\_vars (list)

A named list where the name of each element is a column name of ds. Each element of this list should be a numeric vector with two elements:

• seed (numeric)

The seed to be used for this element - can be NA.

• percentage (proportion)

Percentage of elements to be replaced with NA. If NA, na\_percentage is

used as a default.

cached boolean whether the cached ADVS data cadvs should be returned or new data

should be generated. If set to TRUE then the other arguments to radvs will be

ignored.

## Details

One record per subject per parameter per analysis visit per analysis date.

Keys: STUDYID, USUBJID, PARAMCD, BASETYPE, AVISITN, ATPTN, DTYPE, ADTM, VSSEQ, ASPID

rel\_var 45

## Value

```
data.frame
```

## Author(s)

npaszty

## **Examples**

```
adsl <- radsl(N = 10, seed = 1, study_duration = 2)
advs <- radvs(adsl, visit_format = "WEEK", n_assessments = 7L, seed = 2)
advs <- radvs(adsl, visit_format = "CYCLE", n_assessments = 3L, seed = 2)
advs</pre>
```

rel\_var

Related Variables: Assign

## **Description**

Assign values to a related variable within a domain.

## Usage

```
rel_var(df, var_name, related_var, var_values = NULL)
```

## **Arguments**

df (data.frame)

Data frame containing the related variables.

var\_name (character)

Name of variable related to rel\_var to add to df.

related\_var (character)

Name of variable within df with values to which values of var\_name must relate.

var\_values (any)

Vector of values related to values of related\_var.

#### Value

df with added factor variable var\_name containing var\_values corresponding to related\_var.

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#### **Examples**

```
# Example with data.frame.
params <- c("Level A", "Level B", "Level C")</pre>
adlb_df <- data.frame(</pre>
 ID = 1:9,
 PARAM = factor(
    rep(c("Level A", "Level B", "Level C"), 3),
    levels = params
 )
)
rel_var(
 df = adlb_df,
 var_name = "PARAMCD",
 var_values = c("A", "B", "C"),
  related_var = "PARAM"
)
# Example with tibble.
adlb_tbl <- tibble::tibble(</pre>
 ID = 1:9,
 PARAM = factor(
    rep(c("Level A", "Level B", "Level C"), 3),
    levels = params
  )
)
rel_var(
 df = adlb_tbl,
 var_name = "PARAMCD",
 var_values = c("A", "B", "C"),
  related_var = "PARAM"
)
```

replace\_na

Replace Values in a Vector by NA

#### **Description**

# [Stable]

Randomized replacement of values by NA.

# Usage

```
replace_na(v, percentage = 0.05, seed = NULL)
```

## **Arguments**

v (any) Vector of any type. rtexp 47

percentage (proportion)

Value between 0 and 1 defining how much of the vector shall be replaced by NA.

This number is randomized by +/- 5% to have full randomization.

seed (numeric)

Seed to use for reproducible random number generation.

#### Value

The input vector v where a certain number of values are replaced by NA.

rtexp

Truncated Exponential Distribution

# **Description**

#### [Stable]

This generates random numbers from a truncated Exponential distribution, i.e. from  $X \mid X > 1$  or  $X \mid X < r$  when  $X \sim Exp(rate)$ . The advantage here is that we guarantee to return exactly n numbers and without using a loop internally. This can be derived from the quantile functions of the left- and right-truncated Exponential distributions.

#### Usage

```
rtexp(n, rate, 1 = NULL, r = NULL)
```

#### **Arguments**

n (numeric)

Number of random numbers.

rate (numeric)

Non-negative rate.

1 (numeric)

Positive left-hand truncation parameter.

r (numeric)

Positive right-hand truncation parameter.

#### Value

The random numbers. If neither 1 nor r are provided then the usual Exponential distribution is used.

48 rtpois

#### **Examples**

```
x <- stats::rexp(1e6, rate = 5)
x <- x[x > 0.5]
hist(x)

y <- rtexp(1e6, rate = 5, 1 = 0.5)
hist(y)

z <- rtexp(1e6, rate = 5, r = 0.5)
hist(z)</pre>
```

rtpois

Zero-Truncated Poisson Distribution

## **Description**

## [Stable]

This generates random numbers from a zero-truncated Poisson distribution, i.e. from  $X \mid X > 0$  when  $X \sim Poisson(lambda)$ . The advantage here is that we guarantee to return exactly n numbers and without using a loop internally. This solution was provided in a post by Peter Dalgaard.

#### Usage

```
rtpois(n, lambda)
```

## **Arguments**

n (numeric)

Number of random numbers.

lambda (numeric)

Non-negative mean(s).

# Value

The random numbers.

# **Examples**

```
x <- rpois(1e6, lambda = 5)
x <- x[x > 0]
hist(x)

y <- rtpois(1e6, lambda = 5)
hist(y)</pre>
```

sample\_fct 49

sample_fct (	Create a Factor with Random Elements of x
--------------	---

# **Description**

Sample elements from x with replacement to build a factor.

# Usage

```
sample_fct(x, N, ...)
```

#### **Arguments**

x (character vector or factor)

If character vector then it is also used as levels of the returned factor. If factor

then the levels are used as the new levels.

N (numeric)

Number of items to choose.

... Additional arguments to be passed to sample.

#### Value

A factor of length N.

## **Examples**

```
sample_fct(letters[1:3], 10)
sample_fct(iris$Species, 10)
```

visit\_schedule

Create Visit Schedule

# Description

Create a visit schedule as a factor.

# Usage

```
visit_schedule(visit_format = "WEEK", n_assessments = 10L, n_days = 5L)
```

## **Arguments**

```
\verb|visit_format| & (\verb|character|) \\
```

Type of visit. Options are "WEEK" and "CYCLE".

n\_assessments (integer)

Number of weeks or cycles.

n\_days (integer)

Number of days in each cycle (only used if visit\_format is "CYCLE").

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# **Details**

X number of visits, or X number of cycles and Y number of days.

# Value

A factor of length  $n\_assessments$ .

# Examples

```
visit_schedule(visit_format = "WEeK", n_assessments = 10L)
visit_schedule(visit_format = "CyCLE", n_assessments = 5L, n_days = 2L)
```

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