# Package 'cdashQC'

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<b>Description</b> This package creates table and listings for QC of cdash format data.
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 $assign\_ae\_trt$ 

Assing AE to treatment

### Description

Assign ae to a treatment.

### Usage

```
assign_ae_trt(ae, ex)
```

### Arguments

ae the dataset ae read from sas ex the dataset ex read from sas

### Value

a data frame with one more column specifying which treatment the AE should be in.

clean\_lab 3

clean\_lab

Clean the lab test data set.

#### **Description**

clean the lab data, handling rechecks/unscheduled included.

#### Usage

```
clean_lab(lab, included, return_prob = FALSE)
```

#### **Arguments**

lab the dataset returned by create\_lab

return\_prob should the problematic observations be shown? FALSE by default.

ex the ex data set.

#### Value

lab data with necessary variables kept for further summarization.

#### See Also

```
create_lab and create_included
```

codes

the data set codes.

### Description

A dataset containing the lab test code and its corresponding full name

#### Usage

codes

### **Format**

a data frame containing all possible LB\_TESTC and the corresponding full names.

**TESTNUM** Maybe the number code?

LB\_TESTC the code for laboratory test

TTL the full name

**COLWIDE** The column width? ...

#### **Source**

https://github.com/statswork/cdashQC/tree/master/data-raw/

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count\_percent

add percentage to a frequency table

#### **Description**

count to percent

#### Usage

```
count_percent(data, var1, var2, digit_keep = 3)
```

### Arguments

data the data should contain at least 2 columns

var1 which column should be the group

var2 which columns are used to calculate percentage (could be a vector)

digit\_keep how many digits should be kept.

#### Value

a data frame

### **Examples**

```
trait = rep(c("A", "B", "C"), 3); subtype = paste(trait, rep(1:3, each=3), sep = "")
data <- data.frame(trait, subtype, count1 = rpois(9, 5), count2 = rpois(9, 10))
count_percent(data, var1= 1, var2 = 3:4)

r1 <- get_summary_stats(eg, group = "PERIOD", var = "EG_TEST")
count_percent(r1, var1 = 1, var2 = 3:5)</pre>
```

create\_aet

do what createaet does.

#### **Description**

createaet.

### Usage

```
create_aet(ae, ex, improv = 99)
```

#### **Arguments**

ae the dataset ae read from sas ex the dataset ex read from sas

improv the same argument as the sas macro createaet

included the dataset included from sas, can be created using create\_included()

#### Value

aet the data

create\_baseline 5

create\_baseline

create baseline indicator.

### Description

create an extra column inidcating whether the current row should be considered as baseline.

### Usage

```
create_baseline(data, var_identifier = "_TEST")
```

### **Arguments**

```
data currently support vs, eg and 1b_cq.
var_identifier a string that can be used to identify the variable name e.g.(VS_TEST, EG_TEST, LB_TEST)
```

#### Value

a data frame with an extra column status whose value could be one of BASELINE, POSTDOSE and PREDOSE (NOT BASELINE)

#### See Also

```
guess_base_phour
```

create\_dem

create the demographics data.

#### Usage

```
create_dem(dm, vs, included)
```

#### **Arguments**

dm the dm data set vs the vs data set

inlcuded the included data set created by create\_included

#### Value

a data frame

#### See Also

```
create_included
```

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create\_eg

create variable columns for the ECGs parameters.

### Description

dcast the ecg variables

### Usage

```
create_eg(eg, included)
```

#### **Arguments**

eg the dataset eg read from sas

included the included data set created by create\_included

#### Value

eg1 data with necessary variables kept for further summarization.

create\_included an integrated data set.

### Description

create\_included.

### Usage

```
create_included(ex, cr, ds)
```

### Arguments

ex the dataset ex read from sas
cr the dataset cr read from sas
ds the dataset ds read from sas

### Value

the included data set which contains the following information

CLIENTID the patient ID

SCRID the corresponding screening ID

SEQ the treatment sequence, from ex data

PERIOD CRF study period, from ex data

EX\_ROUTE Route

EX\_FORM Formulation

EX\_STDAT Date of first dosing

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EX\_STTIM Time of first dosing EX\_DOSE Dose Amount EX\_LASTDOSEDAT Date of last dosing (Created in this function) EX\_LASTDOSETIM Time of last dosing CHK\_IN\_DAT date of check-in for each period time of check in for each period CHK\_IN\_TIM RETURN\_DAT date of return for each period RETURN\_TIM time of return for each period Primary reason for discontinuation DS\_TRM\_D

DS\_SPEC specify

Date of completion or discontinuation per subject DS\_STDAT

DS\_LSDAT Date of last contact per subject

create\_lab create the lab test data set.

### **Description**

create the lab data

#### Usage

```
create_lab(lb_cq, included)
```

### **Arguments**

lb\_cq the dataset lb\_cq read from sas

included the included data set created by create\_included

### Value

lab data with necessary variables kept for further summarization.

Create protocol hour create\_phour

### **Description**

Create protocol hour variable if it's not already in the data set.

### Usage

```
create_phour(data, digit = 2)
```

8 create\_vs

### **Arguments**

data either eg, vs or lb\_cq

digit round the hour by digit, default set to be 2.

#### Value

the same data set with one more column if Phour is created, otherwise returns the input data.

create\_seq Create SEQ

### Description

Create SEQ variable (treatment) if it's not already in the data set.

### Usage

```
create_seq(data, included)
```

### Arguments

data either eg, vs or lb\_cq

included the included data set created by create\_included

#### Value

the same data set with one more column if SEQ is created, otherwise returns the input data.

create\_vs create the vital sign data set.

### Description

create the vs data

#### Usage

```
create_vs(vs, included)
```

### **Arguments**

vs the dataset dm read from sas

included the included data set created by create\_included

### Value

vs data with necessary variables kept for further summarization.

ex\_start 9

ex\_start

Get the start time of each treatment

### Description

Get the start time of each treatment

### Usage

```
ex_start(ex)
```

#### **Arguments**

ex

the dataset ex read from sas

#### Value

a data frame specifying the start time of each treatment

format\_time

Format the time variable in a data set

### **Description**

Format the time variable in a data set

#### Usage

```
format_time(data, date = "AE_STDT", time = "AE_STTM")
```

### **Arguments**

data the data which contains the time variables

date the date variable time the time (in seconds)

### Value

the same data set but time has been changed to "hh:mm:ss" format.

10 guess\_base\_phour

<pre>get_summary_stats</pre>	get summary statistics
------------------------------	------------------------

### Usage

```
get_summary_stats(data, group = "EX_TRT_C", var = "race", na.rm = TRUE)
```

#### **Arguments**

data the data

group which column should be the group var which columns are used to summarize

na.rm should missing value be removed? TRUE by default.

#### Value

a data frame

#### **Examples**

```
 \begin{split} &\text{SEQ} = \text{rep}(\text{c}(\text{"A", "B", "C"}), 3); \text{ subtype} = \text{sample}(\text{c}(\text{"ONE", "TWO", "THREE"}), 9, \text{ replace} = \text{TRUE}) \\ &\text{data} <- \text{ data.frame}(\text{SEQ, subtype, BMI} = \text{rnorm}(9, 25, 4), \text{ HEIGHT} = \text{rnorm}(9, 175, 3)) \\ &\text{get\_summary\_stats}(\text{data, group} = \text{"SEQ", var} = \text{"subtype"}) \\ &\text{get\_summary\_stats}(\text{data, group} = \text{"SEQ", var} = \text{"BMI"}) \\ \end{aligned}
```

guess\_base\_phour

find the baseline hour for each test

#### **Description**

find the baseline hour for each test category

### Usage

```
guess_base_phour(data, var_identifier = "_TEST")
```

### **Arguments**

```
data currently support vs, eg and 1b_cq.

var_identifier a string that can be used to identify the variable name e.g.(VS_TEST, EG_TEST, LB_TEST)
```

### Value

a data frame where by-subject baseline hour is determined.

guess\_race\_age 11

#### **Examples**

```
# the following two will give you exactly the same result
r1 <- guess_base_phour(vs)
r2 <- guess_base_phour(vs, var_identifier = "VS_TEST")

# If you want to find baseline hours for eg, the following two methods are equivalent
r3 <- guess_base_phour(eg)
r4 <- guess_base_phour(eg, var_identifier = "EG_TEST")</pre>
```

guess\_race\_age

race and ethnicity indicator, age.

#### **Description**

create necessary variable for the demographic summary tables and listing.

### Usage

```
guess_race_age(dm, included)
```

### Arguments

dm the dataset dm read from sas
ex the dataset ex read from sas

#### Value

a data frame with additional columns listed as follows

race the race of the subject

ethnic has two levels, "NOT HISPANIC OR LATINO" and "HISPANIC OR LATINO".

EX\_TRT\_C the treatment groups

ptno convert CLIENTID to numerical values of subject number

age Age calculated from start date of treatment

guess\_reps

guess the number of replicates

#### **Description**

guess the number of replicates for each SEQ/PERIOD/PHOUR/TESTCODE

### Usage

```
guess_reps(data, var_identifier = "_TEST")
```

12 guess\_test

#### **Arguments**

```
data data of eg (created by create_eg) or vs (created by create_vs)
var_identifier,
which columns is the variable for test code names.
```

#### Value

a data frame containing number of reps per PERIOD/PHOUR/TESTCODE

#### **Examples**

```
eg0 <- create(eg, included)
d1 <- guess_reps(eg0)</pre>
```

guess\_test

guess the name of the test

### Description

guess the name of the test code: it should end with var\_identifier.

#### Usage

```
guess_test(data, var_identifier = "_TEST")
```

### Arguments

#### Value

the test code

### **Examples**

```
guess_test(eg)
```

lab\_oor

lab\_oor

Find OOR

### Description

Find the out-of-range values

### Usage

```
lab_oor(lab)
```

### **Arguments**

lab

the data returned by create\_lab.

### Value

a list containg OOR.

listing\_ae

list Adverse Envent

### Description

list the ae

### Usage

```
listing_ae(aet, type = 1)
```

### Arguments

aet

created by create\_aet

type

an indicator. Should I list ae1 (type = 1), ae2 (type = 2) or ae3 (type = 3)?

### Value

a data frame

### See Also

```
create_aet
```

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listing\_concom

list concomitant.

### Description

list concomitant medication data

### Usage

```
listing_concom(cm)
```

### Arguments

cm

the dataset cm read from sas

### Value

the data with related columns

listing\_dem

demographics listing.

### Description

Summarize the demographic data

### Usage

```
listing_dem(dmt)
```

### **Arguments**

dmt

the data set created by  ${\tt create\_dem}.$ 

#### Value

a data frame

### See Also

```
create_included and create_dem
```

normal\_range 15

normal\_range

Find normal range for lab test

### Description

Find the normal range of each test code

### Usage

```
normal_range(lab)
```

### Arguments

lab

the dataset generated by create\_lab

#### Value

a data frame containing test codes and their corresponding normal range

replicate\_average

 $get \ the \ averanges \ of \ the \ replicates for \ vs \ or \ eg \ data.$ 

### **Description**

Get the averages of the replicates

### Usage

```
replicate_average(data_clean, digits = NULL, na_rm = TRUE)
```

### Arguments

data\_clean an object returned from replicate\_clean digits should the averages be rounded? Default NO.

na\_rm should missing values be excluede? Default TRUE.

### Value

the averages

### See Also

```
replicate_clean
```

### **Examples**

```
eg2 \leftarrow replicate\_data(eg) \# find the triplicates
```

16 replicate\_clean

replicate\_check

check the replicates and issue a message if there's issue

### Description

check data qualities, reporting missing and RECHECKS

#### Usage

```
replicate_check(data, reps = NULL, printed = TRUE)
```

#### **Arguments**

data of eg (created by create\_eg) or vs (created by create\_vs)

reps the number of replicates planned. (should be tabled by TESTCODE, SEQ, PE-

RIOD and PHOUR). If NULL, guess\_reps will be invoked to guess the number

of replicates.

printed whether to print the observations having issues at the console.

#### Value

a data frame containing subject and issues at given period and protocol hours

#### See Also

```
guess_reps.
```

### Examples

```
eg0 <- create_eg(eg, incldued)
d1 <- replicate_check(eg1)</pre>
```

replicate\_clean

get the cleaned replicates by combining the "clean" data and "dirty"

data

### Description

get cleaned replicates

### Usage

```
replicate_clean(data, rm_row = NULL)
```

### **Arguments**

data an object returned from replicate\_data (works for vs and eg)

rm\_row a vector of integers specifying which rows should be removed from the dirty

data.

replicate\_data 17

#### Value

the cleaned replicates

#### See Also

```
replicate_data
```

#### **Examples**

```
eg2 <- replicate_data(eg) # step 1: find the triplicates
eg_prob <- eg2$data_dirty  # need manual check
# the following rows should be removed
rows_removed <- c(2, 4, 7, 13, 14, 15, 19, 25, 28, 32, 37, 40, 44, 52, 57, 59, 64, 65)
eg3 <- replicate_clean(eg2, rows_removed)</pre>
```

replicate\_data

get the replicates for eg or vs data.

### Description

get the replicates for each protocol hour (PHOUR)

#### Usage

```
replicate_data(data, reps = NULL)
```

#### **Arguments**

data the data set created by create\_eg (for ecg data) or by create\_vs (for vital signs

data).

reps specifies the structures of reps. If not specified, guess\_resp will be invoked.

Default set to be NULL.

#### Value

a list

data\_clean the subjects containing correct number of resplicates

data\_dirty the subjects that have different number of replicates than desired

### See Also

```
create_eg, guess_resp and replicate_check
```

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round df	
----------	--

round data frame

#### **Description**

round a data frame if the column is numerical

#### Usage

```
round_df(df, digits = 3)
```

### **Arguments**

df the data frame

digits how many digits you want to keep

#### Value

a data frame with numerical columns rounded.

summary\_ae\_rel

Adverse event by relationship to study drug

#### **Description**

summarize adverse events by relationship to study drug

#### Usage

```
summary_ae_rel(data, group = "STUDYNO", var = "EX_TRT_C", by = "subject")
```

#### **Arguments**

data the data set created by create\_aet

group choose a variable that has all values identical (e.g., "STUDYNO").

var the treatment variable name, "EX\_TRT\_C" for example

by has two options, either "event" or "subject". If by = "subject", that means

When a subject experienced the same AE at more than one level of drug rela-

tionship during a treatment period, each AE was counted separately

#### Value

a list

pt frequency table by preferred term total frequency table by treatment

summary\_ae\_sev 19

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Sullillai y	_ac_s	C V

Adverse event by severity

#### **Description**

summarize adverse events by severity

### Usage

```
summary_ae_sev(data, group = "STUDYNO", var = "EX_TRT_C", by = "subject")
```

### **Arguments**

data the data set created by create\_aet

group choose a variable that has all values identical (e.g., "STUDYNO").

var the treatment variable name, "EX\_TRT\_C" for example

by has two options, either "event" or "subject". If by = "subject", the subject

will be counted under the highest severity level.

### Value

a list

pt frequency table by preferred term total frequency table by treatment

summary\_dem

demographic summary.

### **Description**

Summarize the demographic data

#### Usage

```
summary_dem(dmt, group = "SEQ", na.rm = TRUE)
```

### Arguments

dmt the data set created by create\_dem.

group which variable name would be used to calculate summary statistics?

na.rm should missing values be included? TRUE by default.

### Value

a data frame

20 summary\_lab

#### See Also

```
create_included and create_dem
```

### **Examples**

```
included <- create_included(ex, dm, cr, ds)
dmt <- create_dem(dm, ex, vs, included)
summary_dem(dmt, group = "SEQ")  # the summary by group
summary_dem(dmt, group = "SPONSOR")  # to get the overall summary</pre>
```

summary\_lab

summary statistics for lab shift.

### **Description**

Summarize the lab statistics

### Usage

```
summary_lab(lb_clean, digits = 3)
```

### **Arguments**

lb\_clean the dataset returned by clean\_lab.

digits how many digits should be kept for the final data.

### Value

the summary statistics by test code, time point and by treatment.

### See Also

```
clean_lab
```

### **Examples**

```
included <- create_included(ex, dm, cr, ds)
lab <- create_lab(lb_cq)
lb_clean <- clean_lab(lab, included= included, ex= ex)
s1 <- summary_lab(clean_lab) #not urinalysis</pre>
```

summary\_labshift 21

summary\_labshift

create the lab shift table.

### **Description**

Create the lab shift table.

#### Usage

```
summary_labshift(lb_clean)
```

#### **Arguments**

lb\_clean the dataset returned by clean\_lab.

#### Value

the shift table

#### See Also

clean\_lab

#### **Examples**

```
included <- create_included(ex, dm, cr, ds)
lab <- create_lab(lb_cq)
lb_clean <- clean_lab(lab, included, ex)
lbshift <- summary_labshift(lb_clean)</pre>
```

summary\_teae

teae by number of subjects or by number of events

### Description

summarize treatment-emergent adverse event

### Usage

```
summary_teae(data, group = "STUDYNO", var = "EX_TRT_C", by = "subject")
```

### Arguments

data the data set created by create\_aet

group choose a variable that has all values identical (e.g., "STUDYNO").

var the treatment variable name, "EX\_TRT\_C" for example

by has two options, either "event" or "subject"

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#### Value

a list

ae frequency table for teae

soc summarization by system organ class and by preferred term

summary\_vs\_eg

summary statistics

#### **Description**

Summary statistics for vital signs or eg

#### Usage

```
summary_vs_eg(data_clean, inter_digit = NULL, final_digits = 3,
    na_rm = TRUE, ischangefrombase = FALSE)
```

#### **Arguments**

data\_clean the clean data set returned by replicate\_clean.

inter\_digit if rounding happens for the intermediate averages, what digits should be kept.

na\_rm should missing values be excluede? Default set to be TRUE.

ischangefrombase

Is this summary for change from baseline? Default set to be FALSE

final\_digit what is the digit for final summary?

#### Value

a data frame with summary statistics by test, time point and treatment.

weight\_height\_bmi

read BMI Weight and Height info from Screening stage.

### Description

extract BMI, weight and height from vs.sas7bat.

### Usage

```
weight_height_bmi(vs, period = "SCREEN")
```

#### **Arguments**

٧S

the vs sas data

### Value

a data frame containing BMI, WEIGHT and HEIGHT from admission stage

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