Package 'growthcleanr'

February 26, 2024

Type Package Title Data Cleaner for Anthropometric Measurements Version 2.2.0 Description Identifies implausible anthropometric (e.g., height, weight) measurements in irregularly spaced longitudinal datasets, such as those from electronic health records. URL https://carriedaymont.github.io/growthcleanr/index.html, https://github.com/carriedaymont/growthcleanr BugReports https://github.com/carriedaymont/growthcleanr/issues Imports R.utils (>= 2.11.0), data.table (>= 1.13.0), tidyr (>= 1.1.0), plyr (>= 1.8.6), dplyr (>= 1.0.1), foreach (>= 1.5.0), doParallel (>= 1.0.15), labelled (>= 2.5.0), magrittr (>= 1.5) **Depends** R (>= 2.10) License MIT + file LICENSE **Encoding** UTF-8 LazyData true **Suggests** argparser (≥ 0.6), bit64 ($\geq 4.0.2$), knitr (≥ 1.29), rmarkdown (>= 2.3), testthat (>= 2.3.2) RoxygenNote 7.2.3 NeedsCompilation no Author Carrie Daymont [ctb, cre], Robert Grundmeier [aut], Jeffrey Miller [aut], Diego Campos [aut], Dan Chudnov [ctb], Hannah De los Santos [ctb], Lusha Cao [ctb], Steffani Silva [ctb], Hanzhe Zhang [ctb], Matt Boyas [ctb], David Freedman [ctb],

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acf_answers

Index

acf_answers

Description

Determines what should absolutely be reincluded or definitely excluded for a given dataset, already run through cleangrowth.

Usage

```
acf_answers(
   subjid,
   param,
   agedays,
   sex,
   measurement,
   orig.exclude,
   sd.recenter = NA,
   ewma.exp = -1.5,
   ref.data.path = "",
   quietly = TRUE
)
```

Arguments

subjid	Vector of unique identifiers for each subject in the database.
param	Vector identifying each measurement, may be 'WEIGHTKG', 'HEIGHTCM', or 'LENGTHCM' 'HEIGHTCM' vs. 'LENGTHCM' only affects z-score calculations between ages 24 to 35 months (730 to 1095 days). All linear measurements below 731 days of life (age 0-23 months) are interpreted as supine length, and all linear measurements above 1095 days of life (age 36+ months) are interpreted as standing height. Note: at the moment, all LENGTHCM will be converted to HEIGHTCM. In the future, the algorithm will be updated to consider this difference.
agedays	Numeric vector containing the age in days at each measurement.
sex	Vector identifying the gender of the subject, may be 'M', 'm', or 0 for males, vs. 'F', 'f' or 1 for females.
measurement	Numeric vector containing the actual measurement data. Weight must be in kilograms (kg), and linear measurements (height vs. length) in centimeters (cm).
orig.exclude	Vector of exclusion assessment results from cleangrowth()
sd.recenter	Data frame or table with median SD-scores per day of life

ewma.exp	Exponent to use for weighting measurements in the exponentially weighted moving average calculations. Defaults to -1.5. This exponent should be negative in order to weight growth measurements closer to the measurement being evaluated more strongly. Exponents that are further from zero (e.g3) will increase the relative influence of measurements close in time to the measurement being evaluated compared to using the default exponent.
ref.data.path	Path to reference data. If not supplied, the year 2000 Centers for Disease Control (CDC) reference data will be used.
quietly	Determines if function messages are to be displayed and if log files (parallel only) are to be generated. Defaults to TRUE.

Value

A data frame, containing an index "n" of rows, corresponding to the original order of the input vectors, and "acf_answers", containing the answers on whether a height value should be kept or excluded (returns "Definitely Exclude", "Definitely Include", or "Unknown" for height values, NA for weight values).

adjustcarryforward	adjustcarryforward adjustcarryforward Uses absolute height ve-
	locity to identify values excluded as carried forward values for rein-
	clusion.

Description

adjustcarryforward adjustcarryforward Uses absolute height velocity to identify values excluded as carried forward values for reinclusion.

Usage

```
adjustcarryforward(
  subjid,
  param,
  agedays,
  sex,
 measurement,
  orig.exclude,
  exclude_opt = 0,
  sd.recenter = NA,
  ewma.exp = -1.5,
  ref.data.path = "",
  quietly = TRUE,
  minfactor = 0.5,
  maxfactor = 2,
  banddiff = 3,
  banddiff_plus = 5.5,
  min_ht.exp_under = 2,
```

```
min_ht.exp_over = 0,
max_ht.exp_under = 0.33,
max_ht.exp_over = 1.5
)
```

Arguments

subjid	Vector of unique identifiers for each subject in the database.
param	Vector identifying each measurement, may be 'WEIGHTKG', 'HEIGHTCM', or 'LENGTHCM' 'HEIGHTCM' vs. 'LENGTHCM' only affects z-score calculations between ages 24 to 35 months (730 to 1095 days). All linear measurements below 731 days of life (age 0-23 months) are interpreted as supine length, and all linear measurements above 1095 days of life (age 36+ months) are interpreted as standing height. Note: at the moment, all LENGTHCM will be converted to HEIGHTCM. In the future, the algorithm will be updated to consider this difference.
agedays	Numeric vector containing the age in days at each measurement.
sex	Vector identifying the gender of the subject, may be 'M', 'm', or 0 for males, vs. 'F', 'f' or 1 for females.
measurement	Numeric vector containing the actual measurement data. Weight must be in kilograms (kg), and linear measurements (height vs. length) in centimeters (cm).
orig.exclude	Vector of exclusion assessment results from cleangrowth()
exclude_opt	Number from 0 to 3 indicating which option to use to handle strings of carried-forwards: 0. no change.
	 when deciding to exclude values, if we have a string of carried forwards, drop the most deviant value, and all CFs in the same string, and move on as normal. when deciding to exclude values, if the most deviant in a string of carried forwards is flagged, check all the CFs in that string from 1:N. Exclude all after the first that is flagged for exclusion when comparing to the Include before and after. Do not remove things designated as include. when deciding to exclude values, if the most deviant in a string of carried forwards is flagged, check all the CFs in that string from 1:N. Exclude all after the first that is flagged for exclusion when comparing to the Include before and after. Do not remove things designated as include.
	after the first that is flagged for exclusion when comparing to the Include before and after. Make sure remove things designated as include.
sd.recenter	Data frame or table with median SD-scores per day of life
ewma.exp	Exponent to use for weighting measurements in the exponentially weighted moving average calculations. Defaults to -1.5. This exponent should be negative in order to weight growth measurements closer to the measurement being evaluated more strongly. Exponents that are further from zero (e.g3) will increase the relative influence of measurements close in time to the measurement being evaluated compared to using the default exponent.
ref.data.path	Path to reference data. If not supplied, the year 2000 Centers for Disease Control (CDC) reference data will be used.
quietly	Determines if function messages are to be displayed and if log files (parallel only) are to be generated. Defaults to TRUE.

minfactor	Sweep variable for computing mindiff.next.ht in 15f, default 0.5	
maxfactor	Sweep variable for computing maxdiff.next.ht in 15f, default 2	
banddiff	Sweep variable for computing mindiff.next.ht in 15f, default 3	
banddiff_plus	Sweep variable for computing maxdiff.next.ht in 15, default 5.5	
<pre>min_ht.exp_under</pre>		
	Sweep variable for computing ht.exp in 15f, default 2	
min_ht.exp_over		
	Sweep variable for computing ht.exp in 15f, default 0	
<pre>max_ht.exp_under</pre>		
	Sweep variable for computing ht.exp in 15f, default 0.33	
max_ht.exp_over		
	Sweep variable for computing ht.exp in 15f, default 1.5	

Value

Re-evaluated exclusion assessments based on height velocity.

Examples

```
# Run on a small subset of given data
df <- as.data.frame(syngrowth)</pre>
df <- df[df$subjid %in% unique(df[, "subjid"])[1:2], ]</pre>
clean_df <- cbind(df,</pre>
                   "gcr_result" = cleangrowth(df$subjid,
                                                df$param,
                                                df$agedays,
                                                df$sex,
                                                df$measurement))
# Adjust carry forward values in cleaned data
adj_clean <- adjustcarryforward(subjid = clean_df$subjid,</pre>
                                  param = clean_df$param,
                                  agedays = clean_df$agedays,
                                  sex = clean_df$sex,
                                  measurement = clean_df$measurement,
                                  orig.exclude = clean_df$gcr_result)
```

bmianthro

BMI Anthro

Description

Part of default CDC-derived tables

CDCref_d

Details

Contains BMI data for calculating BMI

bmianthro.txt.gz

Used in function cleangrowth()

CDCref_d

CDC BMI reference data

Description

Used for extended BMIz computation

CDCref_d.csv.gz

Used for extended BMI computation

cleangrowth

Clean growth measurements

Description

Clean growth measurements

Usage

```
cleangrowth(
  subjid,
  param,
  agedays,
  sex,
 measurement,
  recover.unit.error = FALSE,
  sd.extreme = 25,
  z.extreme = 25,
  lt3.exclude.mode = "default",
  height.tolerance.cm = 2.5,
  error.load.mincount = 2,
  error.load.threshold = 0.5,
  sd.recenter = NA,
  sdmedian.filename = "",
  sdrecentered.filename = "",
  include.carryforward = FALSE,
  ewma.exp = -1.5,
```

```
ref.data.path = "",
log.path = NA,
parallel = FALSE,
num.batches = NA,
quietly = TRUE,
adult_cutpoint = 20,
weight_cap = Inf,
adult_columns_filename = "",
prelim_infants = FALSE
)
```

Arguments

subjid	Vector of unique identifiers for each subject in the database.
param	Vector identifying each measurement, may be 'WEIGHTKG', 'WEIGHTLBS', 'HEIGHTCM', 'HEIGHTIN', 'LENGTHCM', or 'HEADCM'. 'HEIGHTCM'/'HEIGHTIN' vs. 'LENGTHCM' only affects z-score calculations between ages 24 to 35 months (730 to 1095 days). All linear measurements below 731 days of life (age 0-23 months) are interpreted as supine length, and all linear measurements above 1095 days of life (age 36+ months) are interpreted as standing height. Note: at the moment, all LENGTHCM will be converted to HEIGHTCM. In the future, the algorithm will be updated to consider this difference. Addition- ally, imperial 'HEIGHTIN' and 'WEIGHTLBS' measurements are converted to metric during algorithm calculations.
agedays	Numeric vector containing the age in days at each measurement.
sex	Vector identifying the gender of the subject, may be 'M', 'm', or 0 for males, vs. 'F', 'f' or 1 for females.
measurement	Numeric vector containing the actual measurement data. Weight must be in kilograms (kg), and linear measurements (height vs. length) in centimeters (cm).
recover.unit.e	rror
	Indicates whether the cleaning algorithm should attempt to identify unit errors (I.e. inches vs. cm, lbs vs. kg). If unit errors are identified, the value will be corrected and retained within the cleaning algorithm as a valid measurement. Defaults to FALSE.
sd.extreme	Measurements more than sd.extreme standard deviations from the mean (either above or below) will be flagged as invalid. Defaults to 25.
z.extreme	Measurements with an absolute z-score greater than z.extreme will be flagged as invalid. Defaults to 25.
lt3.exclude.mo	de
	Determines type of exclusion procedure to use for 1 or 2 measurements of one type without matching same ageday measurements for the other parameter. Op- tions include "default" (standard growthcleanr approach), and "flag.both" (in case of two measurements of one type without matching values for the other parameter, flag both for exclusion if beyond threshold)
height.toleran	

maximum decrease in height tolerated for sequential measurements

error.load.mincount

minimum count of exclusions on parameter before considering excluding all measurements. Defaults to 2.

error.load.threshold

threshold of percentage of excluded measurement count to included measurement count that must be exceeded before excluding all measurements of either parameter. Defaults to 0.5.

- sd.recenter specifies how to recenter medians. May be a data frame or table w/median SDscores per day of life by gender and parameter, or "NHANES" or "derive" as a character vector.
 - If sd.recenter is specified as a data set, use the data set
 - If sd. recenter is specified as "nhanes", use NHANES reference medians
 - If sd.recenter is specified as "derive", derive from input
 - If sd. recenter is not specified or NA:
 - If the input set has at least 5,000 observations, derive medians from input
 - If the input set has fewer than 5,000 observations, use NHANES

If specifying a data set, columns must include param, sex, agedays, and sd.median (referred to elsewhere as "modified Z-score"), and those medians will be used for recentering. A summary of how the NHANES reference medians were derived is available in README.md. Defaults to NA.

sdmedian.filename

Name of file to save sd.median data calculated on the input dataset to as CSV. Defaults to "", for which this data will not be saved. Use for extracting medians for parallel processing scenarios other than the built-in parallel option.

sdrecentered.filename

Name of file to save re-centered data to as CSV. Defaults to "", for which this data will not be saved. Useful for post-processing and debugging.

include.carryforward

Determines whether Carry-Forward values are kept in the output. Defaults to False.

- ewma.exp Exponent to use for weighting measurements in the exponentially weighted moving average calculations. Defaults to -1.5. This exponent should be negative in order to weight growth measurements closer to the measurement being evaluated more strongly. Exponents that are further from zero (e.g. -3) will increase the relative influence of measurements close in time to the measurement being evaluated compared to using the default exponent.
- ref.data.path Path to reference data. If not supplied, the year 2000 Centers for Disease Control (CDC) reference data will be used.
- log.pathPath to log file output when running in parallel (non-quiet mode). Default is
NA. A new directory will be created if necessary. Set to NA to disable log files.
- parallel Determines if function runs in parallel. Defaults to FALSE.
- num.batches Specify the number of batches to run in parallel. Only applies if parallel is set to TRUE. Defaults to the number of workers returned by the getDoParWorkers function in the foreach package.

quietly	Determines if function messages are to be displayed and if log files (parallel only) are to be generated. Defaults to TRUE	
adult_cutpoint	Number between 18 and 20, describing ages when the pediatric algorithm should not be applied (< adult_cutpoint), and the adult algorithm should apply (>= adult_cutpoint). Numbers outside this range will be changed to the closest number within the range. Defaults to 20.	
weight_cap	Positive number, describing a weight cap in kg (rounded to the nearest .1, +/1) within the adult dataset. If there is no weight cap, set to Inf. Defaults to Inf.	
adult_columns_filename		
	Name of file to save original adult data, with additional output columns to as CSV. Defaults to "", for which this data will not be saved. Useful for post-analysis. For more information on this output, please see README.	
prelim_infants	TRUE/FALSE. Run the in-development release of the infants algorithm (expands pediatric algorithm to improve performance for children $0-2$ years). Not recommended for use in research. For more information regarding the logic of the algorithm, see the vignette 'Preliminary Infants Algorithm.' Defaults to FALSE.	

Value

Vector of exclusion codes for each of the input measurements.

Possible values for each code are:

- 'Include', 'Unit-Error-High', 'Unit-Error-Low', 'Swapped-Measurements', 'Missing',
- 'Exclude-Carried-Forward', 'Exclude-SD-Cutoff', 'Exclude-EWMA-Extreme', 'Exclude-EWMA-Extreme-Pair',
- 'Exclude-Extraneous-Same-Day',
- 'Exclude-EWMA-8', 'Exclude-EWMA-9', 'Exclude-EWMA-10', 'Exclude-EWMA-11', 'Exclude-EWMA-12', 'Exclude-EWMA-13', 'Exclude-EWMA-14',
- 'Exclude-Min-Height-Change', 'Exclude-Max-Height-Change',
- 'Exclude-Pair-Delta-17', 'Exclude-Pair-Delta-18', 'Exclude-Pair-Delta-19',
- · 'Exclude-Single-Outlier', 'Exclude-Too-Many-Errors', 'Exclude-Too-Many-Errors-Other-Parameter'

Examples

Once processed you can filter data based on result value

ewma

ewma

Exponentially Weighted Moving Average (EWMA)

Description

ewma calculates the exponentially weighted moving average (EWMA) for a set of numeric observations over time.

Usage

ewma(agedays, z, ewma.exp, ewma.adjacent = TRUE)

Arguments

agedays	Vector of age in days for each z score (potentially transformed to adjust weight- ing).
z	Input vector of numeric z-score data.
ewma.exp	Exponent to use for weighting.
ewma.adjacent	Specify whether EWMA values excluding adjacent measurements should be cal- culated. Defaults to TRUE.

Value

Data frame with 3 variables:

- The first variable (ewma.all) contains the EWMA at observation time excluding only the actual observation for that time point.
- The second variable (ewma.before) contains the EWMA for each observation excluding both the actual observation and the immediate prior observation.
- The third variable (ewma.after) contains the EWMA for each observation excluding both the actual observation and the subsequent observation.

Examples

ext_bmiz

Calculate extended BMI measures

Description

ext_bmiz Calculates the sigma (scale parameter for the half-normal distribution), extended BMI percentile, extended BMIz, and the CDC LMS Z-scores for weight, height, and BMI for children between 2 and 19.9 years of age. Note that for BMIs <= 95th percentile of the CDC growth charts, the extended values for BMI are equal to the LMS values. The extended values differ only for children who have a BMI > 95th percentile.

Usage

```
ext_bmiz(
  data,
  age = "agem",
  wt = "wt",
  ht = "ht",
  bmi = "bmi",
  adjust.integer.age = TRUE,
  ref.data.path = ""
)
```

Arguments

data	Input data frame or data table
age	Name of input column containing subject age in months in quotes, default "agem"
wt	Name of input column containing weight (kg) value in quotes, default "wt"
ht	Name of input column containing height (cm) value in quotes, default "ht"
bmi	Name of input column containing calculated BMI in quotes, default "bmi"

ext_bmiz

adjust.integer.age If age inputs are all integer, add 0.5 if TRUE; default TRUE ref.data.path Path to directory containing reference data

Details

This function should produce output equivalent to the SAS macro provided at https://www.cdc.gov/nccdphp/dnpao/growthcha The macro was updated in December, 2022, according to the findings of the NCHS report available at https://dx.doi.org/10.15620/cdc:121711. This function has been updated to match it as of growthcleanr v2.1.0.

See the README.md file for descriptions of the output columns generated by this function.

data must have columns for at least age, sex, weight, height, and bmi.

age should be coded in months, using the most precise values available. To convert to months from age in years, multiply by 12. To convert to months from age in days, divide by 30.4375 (365.25 / 12).

sex is coded as 1, boys, Boys, b, B, males, Males, m, or M for male subjects or 2, girls, Girls, g, G, females, Females, f, or F for female subjects. Note that this is different from cleangrowth, which uses 0 (Male) and 1 (Female).

wt should be in kilograms.

ht should be in centimeters.

Specify the input data parameter names for age, wt, ht, bmi using quotation marks. See example below.

If the parameter adjust.integer.age is TRUE (the default), 0.5 will be added to all age if all input values are integers. Set to FALSE to disable.

By default, the reference data file CDCref_d.csv, made available at https://www.cdc.gov/nccdphp/dnpao/growthcharts/resour is included in this package for convenience. If you are developing this package, use ref.data.path to adjust the path to this file from your working directory if necessary.

Value

Expanded data frame containing computed BMI values

```
# Run on a small subset of given data
df <- as.data.frame(syngrowth)
df <- df[df$subjid %in% unique(df[, "subjid"])[1:5], ]
df <- cbind(df,
                             "gcr_result" = cleangrowth(df$subjid,
```

```
df$param,
                                          df$agedays,
                                          df$sex,
                                          df$measurement))
df_wide <- longwide(df) # convert to wide format for ext_bmiz</pre>
df_wide_bmi <- simple_bmi(df_wide) # compute simple BMI</pre>
# Calling the function with default column names
df_bmiz <- ext_bmiz(df_wide_bmi)</pre>
# Specifying different column names; note that quotes are used
dfc <- simple_bmi(df_wide)</pre>
colnames(dfc)[colnames(dfc) %in% c("agem", "wt", "ht")] <-</pre>
  c("agemos", "weightkg", "heightcm")
df_bmiz <- ext_bmiz(dfc, age="agemos", wt="weightkg", ht="heightcm")</pre>
# Disabling conversion of all-integer age in months to (age + 0.5)
dfc <- simple_bmi(df_wide)</pre>
df_bmiz <- ext_bmiz(dfc, adjust.integer.age=FALSE)</pre>
```

fentlms_foraga Fenton Growth Curves

Description

Fenton growth curves with premature infant data with sex, age, and integer weight

fentlms_foraga.csv.gz

Used in function cleangrowth()

fentlms_forz Fenton Growth Curve Z-Scores

Description

Fenton growth curves with premature infant z-scores for height and head circumference

fentlms_forz.csv.gz

growth_cdc_ext CDC Growth Percentile Table

Description

Part of default CDC-derived tables

Details

Contains percentiles for various ages, gender, and weights, pre-calculated by CDC

growthfile_cdc_ext.csv.gz

Used in function cleangrowth()

growth_cdc_ext_infants

CDC Growth Percentile Table for Infants

Description

Part of default CDC-derived tables

Details

Contains percentiles for various ages, gender, and weights, pre-calculated by CDC for infants algorithm

$growthfile_cdc_ext_infants.csv.gz$

Used in function cleangrowth()

growth_who_ext WHO Growth Percentile Table

Description

Part of default WHO-derived tables

Details

Contains percentiles for various ages, gender, and weights, pre-calculated by WHO

growthfile_who.csv.gz

lenanthro

Description

Part of default CDC-derived tables

Details

Contains percentiles for various ages, gender, and weights, pre-calculated by CDC

lenanthro.txt.gz

Used in function cleangrowth()

longwide

Transform data in growthcleanr format into wide structure for BMI calculation

Description

longwide transforms data from long to wide format. Ideal for transforming output from growthcleanr::cleangrowth() into a format suitable for growthcleanr::ext_bmiz().

Usage

```
longwide(
   long_df,
   id = "id",
   subjid = "subjid",
   sex = "sex",
   agedays = "agedays",
   param = "param",
   measurement = "measurement",
   gcr_result = "gcr_result",
   include_all = FALSE,
   inclusion_types = c("Include"),
   extra_cols = NULL,
   keep_unmatched_data = FALSE
)
```

longwide

Arguments

long_df	A data frame to be transformed. Expects columns: id, subjid, sex, agedays, param, measurement, and gcr_result.
id	name of observation ID column
subjid	name of subject ID column
sex	name of sex descriptor column
agedays	name of age (in days) descriptor column
param	name of parameter column to identify each type of measurement
measurement	name of measurement column containing the actual measurement data
gcr_result	name of column of results from growthcleanr::cleangrowth()
include_all	Determines whether the function keeps all exclusion codes. If TRUE, all exclusion types are kept and the inclusion_types argument is ignored. Defaults to FALSE.
inclusion_types	
	Vector indicating which exclusion codes from the cleaning algorithm should be included in the data, given that include_all is FALSE. For all options, see growthcleanr::cleangrowth(). Defaults to c("Include").
extra_cols	Vector of additional columns to include in the output. If a column C1 differs on agedays matched height and weight values, then include separate ht_C1 and wt_C1 columns as well as a match_C1 column that gives booleans indicating where ht_C1 and wt_C1 are the same. If the agedays matched height and weight columns are identical, then only include a single version of C1. Defaults to empty vector (not keeping any additional columns).
keep_unmatched_data	
	boolean indicating whether to keep height/weight observations that do not have a matching weight/height on that day

Value

Returns a data frame transformed from long to wide. Includes only values flagged with indicated inclusion types. Potentially includes additional columns if arguments are passed to extra_cols. For each subject, heights without corresponding weights for a given age (and vice versa) will be dropped unless keep_unmatched_data is set to TRUE.

```
# Convert to wide format
wide_df <- longwide(df)
# Include all inclusion types
wide_df <- longwide(df, include_all = TRUE)
# Specify all inclusion codes
wide_df <- longwide(df, inclusion_types = c("Include", "Exclude-Carried-Forward"))</pre>
```

nhanes-reference-medians *NHANES reference medians*

Description

Contains reference median values for default recentering, derived from NHANES years 2009-2018

nhanes-reference-medians.csv.gz

Used in function cleangrowth()

rc-reference-medians Infants reference medians

Description

Contains reference median values for default recentering in the infants algorithm

rcfile-2023-08-15_format.csv.gz

Used in function cleangrowth()

read_anthro

Description

Function to calculate z-scores and csd-scores based on anthro tables.

Usage

```
read_anthro(path = "", cdc.only = FALSE, prelim_infants = FALSE)
```

Arguments

path	Path to supplied reference anthro data. Defaults to package anthro tables.
cdc.only	Whether or not only CDC data should be used. Defaults to false.
prelim_infants	TRUE/FALSE. Run the in-development release of the infants algorithm (expands pediatric algorithm to improve performance for children $0 - 2$ years). Not recommended for use in research. For more information regarding the logic of the algorithm, see the vignette 'Preliminary Infants Algorithm.' Defaults to FALSE.

Value

Function for calculating BMI based on measurement, age in days, sex, and measurement value.

Examples

recode_sex

Recode binary sex variable for compatibility

Description

recode_sex recodes a binary sex variable for a given source column in a data frame or data table. Useful in transforming output from growthcleanr::cleangrowth() into a format suitable for growthcleanr::ext_bmiz().

Usage

```
recode_sex(
    input_data,
    sourcecol = "sex",
    sourcem = "0",
    sourcef = "1",
    targetcol = "sex_recoded",
    targetm = 1L,
    targetf = 2L
)
```

Arguments

input_data	a data frame or data table to be transformed. Expects a source column containing a binary sex variable.
sourcecol	name of sex descriptor column. Defaults to "sex"
sourcem	variable indicating "male" sex in input data. Defaults to "0"
sourcef	variable indicating "female" sex in input data. Defaults to "1"
targetcol	desired name of recoded sex descriptor column. Defaults to "sex_recoded"
targetm	desired name of recoded sex variable indicating "male" sex in output data. Defaults to 1
targetf	desired name of recoded sex variable indicating "female" sex in output data. Defaults to 2

Value

Returns a data table with recoded sex variables.

Examples

```
# Run on given data
df <- as.data.frame(syngrowth)
# Run with all defaults
df_r <- recode_sex(df)
# Specify different targets
df_rt <- recode_sex(df, targetcol = "sexr", targetm = "Male", targetf = "Female")
# Specify different inputs
df_ri <- recode_sex(df_rt, sourcecol = "sexr", sourcem = "Male", sourcef = "Female")</pre>
```

sd_median

Description

Calculate median SD score by age for each parameter.

Usage

sd_median(param, sex, agedays, sd.orig)

Arguments

param	Vector identifying each measurement, may be 'WEIGHTKG', or 'HEIGHTCM'.
sex	Vector identifying the gender of the subject, may be 'M', 'm', or 0 for males, vs. 'F', 'f' or 1 for females.
agedays	Numeric vector containing the age in days at each measurement.
sd.orig	Vector of previously calculated standard deviation (SD) scores for each mea- surement before re-centering.

Value

Table of data with median SD-scores per day of life by gender and parameter.

```
simple_bmi
```

Description

simple_bmi Computes BMI using standard formula. Assumes input compatible with output from longwide().

Usage

```
simple_bmi(wide_df, wtcol = "wt", htcol = "ht")
```

Arguments

wide_df	A data frame or data table containing heights and weights in wide format, e.g., after transformation with longwide()
wtcol	name of observation height value column, default 'wt'
htcol	name of subject weight value column, default 'ht'

Value

Returns a data table with the added column "bmi"

```
# Simple usage
# Run on a small subset of given data
df <- as.data.frame(syngrowth)</pre>
df <- df[df$subjid %in% unique(df[, "subjid"])[1:2], ]</pre>
df <- cbind(df,</pre>
             "gcr_result" = cleangrowth(df$subjid,
                                          df$param,
                                          df$agedays,
                                          df$sex,
                                          df$measurement))
# Convert to wide format
wide_df <- longwide(df)</pre>
wide_df_with_bmi <- simple_bmi(wide_df)</pre>
# Specifying different column names; note that quotes are used
colnames(wide_df)[colnames(wide_df) %in% c("wt", "ht")] <-</pre>
  c("weight", "height")
wide_df_with_bmi <- simple_bmi(wide_df, wtcol = "weight", htcol = "height")</pre>
```

splitinput

Description

splitinput Splits input based on keepcol specified, yielding csv files each with at least the mininum number of rows that are written and saved separately (except for the last split file written, which may be smaller). Allows splitting input data while ensuring all records for each individual subject will stay together in one file. Pads split filenames with zeros out to five digits for consistency, assuming < 100,000 file count result.

Usage

```
splitinput(
   df,
   fname = deparse(substitute(df)),
   fdir = NA,
   min_nrow = 10000,
   keepcol = "subjid"
)
```

Arguments

df	data frame to split
fname	new name for each of the split files to start with
fdir	directory to put each of the split files (use "." for working directory). Must be changed from default (NA), which will trigger error.
min_nrow	minimum number of rows for each split file (default 10000)
keepcol	the column name (default "subjid") to use to keep records with the same values together in the same single split file

Value

the count number referring to the last split file written

```
# Run on given data
df <- as.data.frame(syngrowth)
# Run with all defaults (specifying directory)
splitinput(df, fdir = tempdir())
# Specifying the name, directory and minimum row size
splitinput(df, fname = "syngrowth", fdir = tempdir(), min_nrow = 5000)
```

```
# Specifying a different subject ID column
colnames(df)[colnames(df) == "subjid"] <- "sub_id"
splitinput(df, fdir = tempdir(), keepcol = "sub_id")
```

syngrowth

syngrowth

Description

A synthetic set of measurements from ~3,500 subjects generated using Synthea, with measurement errors for testing with growthcleanr. Contains both pediatric and adult data.

Usage

syngrowth

Format

A data frame with six variables: id, subjid, sex, agedays, param, and measurement

Details

Example electronic health record (heightcm, weightkg) data.

tanner_ht_vel Tanner Growth Velocity Table

Description

Part of default CDC-derived tables

Details

Contains velocities for growth pre-calculated by CDC

tanner_ht_vel.csv.gz

Used in function cleangrowth()

tanner_ht_vel_rev Tanner Growth Velocity Table for Infants

Description

Part of default CDC-derived tables

Details

Contains velocities for growth pre-calculated by CDC, used for the infants algorithm

tanner_ht_vel_rev.csv.gz

Used in function cleangrowth()

tanner_ht_vel_with_2sd

Tanner Growth Velocity Table with (2σ)

Description

Part of default CDC-derived tables

Details

Contains velocities for growth pre-calculated by CDC, including those 2 standard deviations away.

tanner_ht_vel_with_2sd.csv.gz

Used in function acf_answers()

testacf

Function to test adjust carried forward

Description

The goal of this script is to consider the height values that growthclean excludes as "carried forward" for potential re-inclusion by using a reverse absolute height velocity check based on step 15 of the Daymont et al. algorithm

testacf

Usage

```
testacf(
  infile,
  seed = 7,
  searchtype = "random",
  grid.length = 9,
  writeout = FALSE,
  outfile = paste0("test_adjustcarryforward_", format(Sys.time(),
        "%m-%d-%Y_%H-%M-%S")),
  quietly = FALSE,
  param = "none",
  debug = FALSE,
  maxrecs = 0,
  exclude_opt = 0,
  add_answers = TRUE
)
```

Arguments

infile	Input data frame/data table, cleaned by cleangrowth(), with columns as described in main README.md
seed	Numeric random seed, used only when performing random search
searchtype	Type of search to perform: random (default), line-grid, full-grid
grid.length	Number of steps in grid to search
writeout	Write output to file? Default FALSE.
outfile	"Output file name, default 'test_adjustcarrforward_DATE_TIME', where DATE is the current system date and time"
quietly	Verbose progress info
param	"none", or data frame to specify which parameters to run full search on, and values to use if not, used only when performing full-grid search
debug	Produce extra data files for debugging
maxrecs	Limit to specified # subjects, default 0 (no limit)
exclude_opt	Type of exclusion method for carried forward strings, 0 to 3. See adjustcarry- forward documentation for more information
add_answers	TRUE or FALSE, indicating whether or not to add answers (definely include/exclude) for the given dataset. Defaults to TRUE

Value

A list containing: testacf_res: data frame with adjustcarryforward results for each run, params: a data frame containing parameter values for each run. debug_filtered_data: data frame with original data, returned if debug TRUE

Description

Contains results of CDC SAS macro for calculating BMI values.

test_syngrowth_sas_output_compare.csv.gz

Used to test function ext_bmiz()

test_syngrowth_wide CDC SAS BMI Input

Description

Contains input data for CDC SAS macro for calculating BMI values.

test_syngrowth_wide.csv.gz

Used to test function ext_bmiz()

weianthro

Weight Anthro Table

Description

Part of default CDC-derived tables

Details

Contains median and standard deviation for weight by age and gender

weianthro.csv.gz

who_hc_maxvel

Description

Part of default WHO-derived tables

Details

Contains three standard deviations for the World Health Organization values of maximum head circumference velocities.

who_hc_maxvel_3sd_infants.csv.gz

Used in function cleangrowth()

who_hc_vel_3sd WHO Head Circumference Velocity for (3σ)

Description

Part of default WHO-derived tables

Details

Contains three standard deviations for the World Health Organization values of head circumference velocities.

who_hc_vel_3sd_infants.csv.gz

Used in function cleangrowth()

who_ht_maxvel WHO Maximum Height Velocity for (3σ)

Description

Part of default WHO-derived tables

Details

Contains three standard deviations for the World Health Organization values of maximum height velocities.

who_ht_maxvel_3sd.csv.gz

who_ht_maxvel_2sd WHO Maximum Height Velocity for (2σ)

Description

Part of default WHO-derived tables

Details

Contains two standard deviations for the World Health Organization values of maximum height velocities.

who_ht_maxvel_2sd.csv.gz

Used in function acf_answers()

who_ht_vel_2sd WHO Height Velocity for (2σ)

Description

Part of default WHO-derived tables

Details

Contains two standard deviations for the World Health Organization values of height velocities.

who_ht_vel_2sd.csv.gz

Used in function acf_answers()

who_ht_vel_3sd WHO Height Velocity for (3σ)

Description

Part of default WHO-derived tables

Details

Contains three standard deviations for the World Health Organization values of height velocities.

who_ht_vel_3sd.csv.gz

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