Package ‘sitar’

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Type Package

Title Super Imposition by Translation and Rotation Growth Curve Analysis

Version 1.4.0

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Description Functions for fitting and plotting SITAR (Super Imposition by Translation And Rotation) growth curve models. SITAR is a shape-invariant model with a regression B-spline mean curve and subject-specific random effects on both the measurement and age scales. The model was first described by Lindstrom (1995) <doi:10.1002/sim.4780141807> and developed as the SITAR method by Cole et al (2010) <doi:10.1093/ije/dyq115>.

License GPL (>= 2)

URL https://github.com/statist7/sitar

Depends nlme, R (>= 3.0.0)

Imports dplyr, forcats, ggplot2, glue, purrr, rlang, rsample, stats, tibble, tidyr, magrittr

Suggests knitr, rmarkdown

VignetteBuilder knitr

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LazyLoad yes

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Description

SITAR is a method of growth curve analysis, based on nlme, that estimates a single mean growth curve as a regression B-spline, plus a set of up to four fixed and random effects (a, b, c and d) (d was added in version 1.2.0) defining how individual growth curves differ from the mean curve. SITAR stands for SuperImposition by Translation And Rotation.

Details

The package also contains some utility functions for the LMS method, as used to construct growth reference centiles (see gamlss).

Package: sitar
Type: Package
Version: 1.0
Date: 2013-09-23
License: GPL-2

Effect a (or alpha) measures size, and is a random intercept relative to the spline curve intercept. Effect b (or beta) measures tempo, the timing of the growth process, and reflects a shift on the x scale relative to the mean. Effect c (or gamma) is velocity, and indicates how the x scale is stretched or shrunk reflecting the rate at which 'time' passes for individuals. Effect d is a rotation in the plane. The aim is for individual curves, adjusted for abcd to lie on top of (i.e. be superimposed on) the mean curve.

The package creates an object of class sitar, based on nlme, representing the nonlinear mixed-effects model fit. Generic functions such as print, plot and summary have methods to show the results of the fit, along with resid, coef, fitted, fixed.effects and random.effects to extract some of its components. The functions AICadj, BICadj and varexp compare respectively the AIC, BIC and variance explained of a series of models, taking into account any transformations of the y variable. Functions plotclean, velout, codeplot and zapvelout are useful to clean the data file.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

References

The idea of SITAR growth curve analysis arose from the paper by Beath (2007) and was first described in Cole et al (2010). The other references describe applications of SITAR to a variety of data forms.


**anova.sitar**

Compare Likelihoods of Fitted SITAR Objects

**Description**

anova method for sitar objects, based on anova.lme.
Usage

## S3 method for class 'sitar'
anova(
  object,
  ..., 
  test = TRUE,
  type = c("sequential", "marginal"),
  adjustSigma = TRUE,
  Terms,
  L,
  verbose = FALSE
)

Arguments

object
  an object inheriting from class sitar.
...
  other optional fitted model objects.
test
  an optional logical value controlling whether likelihood ratio tests should be used.
type
  an optional character string specifying the type of sum of squares to be used.
adjustSigma
  see anova.lme.
Terms
  see anova.lme.
L
  see anova.lme.
verbose
  an optional logical value.

Value

a data frame inheriting from class "anova.lme".

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

---

**apv_se**

*Bootstrap standard errors for SITAR peak velocity and age at peak velocity*

Description

*apv_se* bootstraps a SITAR model to generate standard errors for age at peak velocity (apv) and peak velocity (pv).

Usage

*apv_se(object, fun = getPeak, nboot = 10, seed = NULL, plot = FALSE, ...)*

Arguments

object     SITAR model.
fun        function to extract apv and pv from velocity curve (default getPeak), alternative
           getTakeoff or getTrough.
nboot      number of bootstrap samples (default 10).
seed       integer to initialize the random number generator (default NULL).
plot       logical to control plotting (default FALSE).
...        optional arguments defining the velocity curve to be bootstrapped (plot.sitar
           arguments xfun, yfun, subset, ns or abc), and graphical par parameters.

Details

If plot is TRUE, the original velocity curve is plotted along with each bootstrap sample’s pv versus
apv.

Value

a 2x2 array giving the mean and standard error of apv and pv, with attribute "bs" a tibble containing
the bootstrap estimates of apv and pv, with NAs removed.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

Examples

data(heights)
## fit sitar model for height
model <- sitar(x = age, y = height, id = id, data = heights, df = 4)

## bootstrap standard errors for age at peak velocity and peak velocity
output <- apv_se(model, nboot=3, seed=111, plot=TRUE)

berkeley

The Berkeley Child Guidance Study

Description

The Berkeley Child Guidance Study dataset contains longitudinal anthropometry data for 136 chi-
dren from birth to 21 years.

Usage

berkeley
Format

A data frame with 4884 observations on the following 10 variables:

- **id** factor with levels 201-278 male and 301-385 female
- **age** years, numeric vector
- **height** cm, numeric vector
- **weight** kg, numeric vector
- **stem.length** cm, numeric vector
- **bi.acromial** cm, numeric vector
- **bi.iliac** cm, numeric vector
- **leg.circ** cm, numeric vector
- **strength** lb, numeric vector
- **sex** factor with level 1 male and level 2 female

Details

The data are for 66 boys and 70 girls from Berkeley, California born in 1928-29 of north European ancestry, and followed from birth to 21 years. Measurements were at ages 0, 0.085, 0.25 to 2 (3-monthly), 2 to 8 (annually), and 8 to 21 (6-monthly) years.

The children were measured for height, weight (undressed), stem length, biacromial diameter, biiliac diameter, leg circumference, and dynamometric strength. The data were provided as an appendix to the book by Tuddenham and Snyder (1954), and a few transcription errors are corrected here. A further 19 errors in height and weight as reported in sitar issue #7 are also now corrected. The growth dataset in the fda package uses heights from the same study.

References


Examples

data(berkeley)

```r
## frequencies of age of measurement for each variable
## weight and length/height from birth, other variables from 6-8 years
## few measurements after 18 years
. <- as.factor(berkeley$age)
plot(levels(.), summary(.), type='s', las=1,
     xlab='age of measurement (years)', ylab='frequency of measurements')
points(levels(.), levels(.) < 0, pch=15)
for (i in 3:9) {
  .. <- .[!is.na(berkeley[, names(berkeley)[i]])]
  lines(levels(..), summary(..), type='s', col=i)
}
legend('topright', names(berkeley)[c(3:9)], text.col=c(3:9), bty='n', inset=0.04)
```
BICadj  

Ways to compare SITAR models for fit

Description

BICadj and AICadj calculate the BIC and AIC for SITAR models, adjusting the likelihood for Box-Cox transformed y variables. varexp calculates the variance explained by SITAR models, compared to the corresponding fixed effect models. getL is used by [AIC]ICadj to find what power the y variable is raised to.

Usage

BICadj(..., pattern = NULL)
AICadj(..., k = 2, pattern = NULL)
varexp(..., pattern = NULL)
getL(expr)

Arguments

... one or more SITAR models.
pattern regular expression defining names of models.
k numeric, the penalty per parameter to be used; the default k = 2 is the classical AIC.
expr quoted or unquoted expression containing a single variable name.

Details

The deviance is adjusted if the y variable is power-transformed, using the formula

\[
adjusted\text{deviance} = \text{deviance} - 2n((\lambda - 1) * \log(gm) + \log(\text{abs}(\lambda)))
\]

where \(\lambda\) is the power transform, and \(n\) and \(gm\) are the length and geometric mean of \(y\).

The variance explained is given by

\[
\%\text{explained} = 100 * (1 - (\sigma_2/\sigma_1)^2)
\]

where \(\sigma_1\) is the fixed effects RSD and \(\sigma_2\) the SITAR random effects RSD.

BICadj and AICadj accept non-sitar models with a logLik class. varexp ignores objects not of class sitar.

getL does not detect if the variable in expr, or its log, contains a multiplying constant, so that the expressions \(\log(x)\) and \(1 + 2 * \log(3 * x)\) both return 0.
Value

For BICadj and AICadj a named vector of deviances in increasing order. For varexp a named vector of percentages in decreasing order. For getL the power the variable in expr is raised to, or NA if expr is not a power of (a multiple of) the variable.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

BIC, AIC

Examples

data(heights)
## fit sitar model for height
m1 <- sitar(x=age, y=height, id=id, data=heights, df=5)

## update it for log(height)
m2 <- update(m1, y=sqrt(height))

## compare variance explained in the two models
varexp(m1, m2)

## compare BIC adjusting for sqrt transform
## the pattern matches names starting with "m" followed by a digit
BICadj(pattern="^m[0-9]")

## find what power height is raised to
getL(quote(sqrt(sqrt(height))))

bupdate

Update the b fixed effect to minimise the b-c random effect correlation

Description

A function to update the value of bstart, the starting value for the b fixed effect, to minimise the correlation between the random effects b and c.

Usage

bupdate(x)

Arguments

x a sitar object.
Value

Returns an updated value of the b fixed effect, based on the random effect covariance matrix.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

Examples

```r
## fit sitar model with b fixed effect starting value defaulting to 'mean'
ml <- sitar(x=age, y=height, id=id, data=heights, df=5)
print(fixef(ml)['b'])

## refit with starting value chosen to minimise b-c correlation and df increased
m2 <- update(ml, bstart=bupdate(ml), df=6)
print(fixef(m2)['b'])
```

cdc2000

The CDC 2000 growth reference

Description

The CDC growth reference (Kuczmarski et al 2000) for height, weight, body mass index and head circumference, fitted by the LMS method and summarised by values of L, M and S by sex from birth to 19 years.

Usage

cdc2000

Format

A tibble with 484 observations on the following 14 variables:

- `years` age from 0 to 19 years
- `L.ht` numeric vector
- `M.ht` numeric vector
- `S.ht` numeric vector
- `L.wt` numeric vector
- `M.wt` numeric vector
- `S.wt` numeric vector
- `L.bmi` numeric vector
- `M.bmi` numeric vector
S.bmi numeric vector
L.hc numeric vector
M.hc numeric vector
S.hc numeric vector
sex two-level factor with level 1 male and level 2 female

Details
BMI starts at 2 years, and head circumference stops at 3 years.
The L, M and S values for each measurement correspond respectively to the Box-Cox power, me-
dian and coefficient of variation of the distribution by age and sex (Cole & Green 1992). The short
names and units for each measurement (see LMS2z) are as follows: height (ht, cm), weight (wt, kg),
body mass index (bmi, kg/m2), head circumference (hc, cm).

References
Kuczmarski RJ, Ogden CL, Guo SS, Grummer-Strawn LM, Flegal KM, Mei Z, Wei R, Curtin LR,
Roche AF, Johnson CL. 2000 CDC growth charts for the United States: methods and development.

Examples
data(cdc2000)
## calculate 98th centile for weight in girls from birth to 19 years
round(setNames(LMS2z(x = 0:19, y = 2, sex = 2, measure = 'wt', ref = 'cdc2000',
toz = FALSE), 0:19), 1)

cLMS LMS conversion to and from z-scores

Description
Routines to handle references constructed with the LMS method. Given a set of LMS values, the
functions convert z-scores to measurement centiles and vice versa.

Usage
cLMS(z, L = 1, M, S)
zLMS(x, L = 1, M, S)
Arguments

- **z**: vector or one-column matrix of z-scores to be converted to measurements.
- **L**: vector of Box-Cox transformation (lambda) values, L in the LMS method.
- **M**: vector of medians (mu), M in the LMS method.
- **S**: vector of coefficients of variation (sigma), S in the LMS method.
- **x**: vector or one-column matrix of measurements to be converted to z-scores.

Details

L, M and S – and if vectors then x and z – should all be the same length, recycled if necessary. The formulae converting \( x \) to \( z \) and vice versa are:

\[
z = \frac{(x/M)^L - 1}{LS}
\]

\[
x = M \left(1 + Lsz\right)^{1/L}
\]

where L is reset to \(10^{-7}\) if it is zero. The LMS method is the same as the BCCG family in the gamlss package, except that lambda in LMS is referred to as nu in BCCG.

Value

If \( x \) and \( z \) are vectors \(zlMS\) and \(clMS\) each return a vector, respectively of z-scores and measurement centiles, with length matching the length of (the longest of) \( x \) or \( z \), L, M and S. If \( x \) or \( z \) are matrices \(zlMS\) and \(clMS\) each return a matrix, the number of rows matching the length of (the longest of) L, M and S, and the number of columns matching the length of \( x \) or \( z \).

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

- \(z2cent\), \(LMS2z\), \(pdLMS\)

Examples

\[
cLMS(z = as.matrix(-2:2), L = 1:-1, M = 5:7, S = rep(0.1, 3))
cLMS(z = 0:2, L = 1:-1, M = 7, S = 0.1)
cLMS(z = as.matrix(0:2), L = 1:-1, M = 7, S = 0.1)
zLMS(x = 6.5, L = 1:-1, M = 5:7, S = rep(0.1, 3))
\]
Description

Handles output from velout function to display growth curves with outlying points, either plotting or zapping the outliers.

Usage

```r
codeplot(outliers, icode = 4, ..., print = TRUE)
zapvelout(outliers, icode)
```

Arguments

- `outliers`: Data frame returned from velout.
- `icode`: The code number(s) defining the subset of curves to be displayed or zapped (between 1 and 6).
- `...`: Optional plot parameters.
- `print`: Option to print as well as plot information on each curve.

Details

The function velout identifies putative outliers for y in data, codeplot plots them, and zapvelout sets missing those confirmed as outliers. Codes range from 0 (normal) to 8, where 4 and 6 are conventional outliers (see velout).

Value

codeplot returns summary information on each curve with an outlier of the relevant code, and optionally plots the curve. zapvelout sets to NA values of y whose code is contained in icode, and returns the modified data frame.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

velout
Examples

```r
## identify outliers
outliers <- velout(age, height, id, heights, limit=2)

## plot outliers with code 4 or 6
codeplot(outliers, icode=c(4,6))

## set the 8 outliers missing
newheights <- zapvelout(outliers, icode=6)
```

### deren

**Deren prevalence data on child thinness, overweight and obesity**

**Description**

Age-sex-specific prevalence rates of thinness, overweight and obesity in Ukraine children based on body mass index and IOTF, WHO and CDC cut-offs.

**Usage**

deren

**Format**

A tibble with 22 observations on the following 11 variables:

- **Age** postnatal age from 7 to 17 completed years
- **Sex** two-level factor - Boys and Girls
- **N** integer - group sample size
- **IOTF18.5** thinness prevalence based on IOTF reference and 18.5 cutoff
- **WHO-2** thinness prevalence based on WHO reference and -2 cutoff
- **CDC5** thinness prevalence based on CDC reference and 5 cutoff
- **IOTF25** overweight prevalence based on IOTF reference and 25 cutoff
- **WHO+1** overweight prevalence based on WHO reference and +1 cutoff
- **CDC85** overweight prevalence based on CDC reference and 85 cutoff
- **IOTF30** obesity prevalence based on IOTF reference and 30 cutoff
- **WHO+2** obesity prevalence based on WHO reference and +2 cutoff
- **CDC95** obesity prevalence based on CDC reference and 95 cutoff

**Details**

Note that the overweight prevalences are for overweight excluding obesity, i.e. the prevalence for BMI between the overweight and obesity cutoffs.
dfpower

Source

The values are obtained from Table 2 of Deren et al (2020), recalculated to full accuracy. https://journals.plos.org/plosone/article?id=10.1371/journal.pone.0244300.

References


Examples

```r
## convert IOTF obesity prevalence to WHO obesity prevalence
## and compare with true WHO obesity prevalence - boys and girls age 7-17
data(deren)
ob_convertr(age = Age, sex = Sex, from = 'IOTF 30', to = 'WHO +2',
pfrom = IOTF30, pto = 'WHO+2', data = deren, plot = 'compare')
```

dfpower

Tabulate BIC of SITAR models by degrees of freedom, fixed effects and xy power transformations

Description

dfpower fits a series of sitar models tabulated by combinations of a) specified degrees of freedom for the spline curve, b) specified fixed effects a, b, c, d, c) specified power transformations of x, and d) specified power transformations of y, returning a four-way array of function values (e.g. BIC) applied to each model. The function provides a convenient way to optimise the model.
Usage

\[ \text{dfpower}( \text{object}, \text{df}, \text{fixed}, \text{xpowers}, \text{ypowers}, \text{FUN} = \text{BICadj}, \text{maxIter} = 50, \text{drop} = \text{TRUE}, \text{verbose} = \text{FALSE}) \]

Arguments

- **object**: fitted sitar model to be updated.
- **df**: vector of integer spline degrees of freedom to be fitted (defaults to df in object).
- **fixed**: character vector of fixed effects to be included (defaults to fixed in object, typically 'a + b + c').
- **xpowers**: vector of powers to apply to x (defaults to the power of x in object).
- **ypowers**: vector of powers to apply to y (defaults to the power of y in object).
- **FUN**: function to be tabulated (default BICadj).
- **maxIter**: maximum number of iterations per fit (default \( \text{nlmeControl()$maxIter} \)).
- **drop**: logical which if TRUE (default) drops redundant dimensions and labels from the returned array.
- **verbose**: logical controlling monitoring, which gives numIter for each model.

Details

xpowers and ypowers treat power 0 as log. The formula for x in object must be of the form \( x^{\text{power}} \) or \( \text{fun}(x) \), e.g. \( x, x^{0.5} \) or \( \log(x) \). More complex formulae e.g. \( \log(x + 1) \) will fail. In this case fit the model with the variable \( x_1 = x + 1 \) instead.

FUN can be any function returning a single numerical value, e.g. BICadj, BIC, AIC, varexp or sigma.

Other fixed effects in object for covariates in a.formula, b.formula, c.formula or d.formula are propagated through all the models. This also applies to the control argument if set in object.

The run-time can be shortened by reducing maxIter, as models often converge quickly or not at all.

Value

Four-way array of returned values, ranked with the largest dimensions first, and by default with single-level dimensions dropped.

Values are returned with changed sign if the model fit generates a warning, or as NA if there is an error.
dfset  

**Author(s)**  
Tim Cole <tim.cole@ucl.ac.uk>  

**See Also**  
aperm transposes the returned array; addmargins adds margins.  

**Examples**  

data(heights)  
m1 <- sitar(x = age, y = height, id = id, data = heights, df = 4)  

dfpower(m1, df = 4:6, fixed = c('a', 'a+b', 'a+c', 'a+b+c'),  
       xpowers = 0:1, ypowers = 0:1, maxIter = 8)  

---  

**dfset**  
Find degrees of freedom for a natural spline curve to minimise BIC or AIC  

**Description**  

dfset fits a natural cubic spline for a range of degrees of freedom, and returns the df minimising the BIC or AIC.  

**Usage**  

dfset(x, y, data = parent.frame(), FUN = BIC, df = 1:15, plot = FALSE, ...)  

**Arguments**  

- **x** vector of x coordinates.  
- **y** vector of y coordinates.  
- **data** environment containing x and y.  
- **FUN** function to be minimised (e.g. BIC or AIC).  
- **df** vector of degrees of freedom to be searched.  
- **plot** logical controlling plotting of FUN versus df.  
- **...** parameters to pass to plot.  

**Value**  

degrees of freedom and value of FUN at minimum.  

**Author(s)**  
Tim Cole <tim.cole@ucl.ac.uk>
Examples

```r
data(heights)
dfset(age, height, heights, FUN=BIC, plot=TRUE)
dfset(age, height, heights, FUN=function(a) AIC(a, k=1))
```

---

**funcall**  
*Function call with optional inverse*

---

**Description**

Applies an expression to vector `v`, optionally inverting the expression first. For example if the expression is `log`, `funcall` returns `log(v)` if `inverse` is `FALSE`, and `exp(v)` if `inverse` is `TRUE`.

**Usage**

```r
funcall(v, vcall, inverse = FALSE)
```

**Arguments**

- `v`  
  vector
- `vcall`  
  expression
- `inverse`  
  logical

**Details**

Inverse covers functions `log, exp, sqrt, ^, *, /, +, -`.

**Value**

Returns a vector of length `v`.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>
**getData.sitar**

*Extract elements of fitted SITAR models*

**Description**

getData, getCovariate and getVarCov methods for sitar objects, based on lme.

**Usage**

```r
## S3 method for class 'sitar'
getData(object)

## S3 method for class 'sitar'
getCovariate(object, ...)

## S3 method for class 'sitar'
getVarCov(obj, ...)
```

**Arguments**

- `object, obj` an object inheriting from class sitar.
- `...` other optional arguments.

**Value**

Respectively the data frame and x variable used in the fit, and the returned variance-covariance matrix.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>

---

**getPeakTrough**

*Identify peak or trough on curve*

**Description**

Given vectors x and y, returns their values at the peak or trough of the smooth (e.g. cubic spline) curve y ~ x.
Usage

getPeakTrough(x, y = NULL, peak = TRUE, takeoff = FALSE)

gPeak(x, y = NULL, peak = TRUE, takeoff = FALSE)

gTrough(x, y = NULL, peak = FALSE, takeoff = FALSE)

gTakeoff(x, y = NULL, peak = FALSE, takeoff = TRUE)

Arguments

x vector.
y vector.
peak logical determining whether peak or trough is returned.
takeoff logical determining whether, if peak is FALSE, the trough is takeoff.

Details

Optionally the trough can be specified as takeoff, which is defined for a growth velocity curve as
the lowest velocity before the pubertal peak, and if there is no peak then there is by definition no
takeoff.

Value

A length-2 vector containing the values of x and y at the peak or trough. If none are identified NA's
are returned.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

Examples

## create mean height velocity curve
data(heights)
m1 <- sitar(age, height, id, heights, 4)
## plot velocity curve
plot(m1, v')
## mark peak, trough and takeoff
xy <- plot_v(m1)
points(t(getPeak(xy)), pch=17)
points(t(getTrough(xy)), pch=25, col=2, bg=2)
points(t(getTakeoff(xy)), pch=25, col=3, bg=3)
### heights

Serial heights measured in 12 girls

**Description**

Heights of 12 girls from the Chard Growth Study measured twice a year between 8 and 16 years of age.

**Usage**

```
heights
```

**Format**

A data frame with 124 observations on the following 4 variables:

- **id**  factor of subject ids (levels 1:12).
- **age**  vector of ages (years).
- **height**  vector of heights (cm).
- **men**  vector of ages at menarche (years), where negative values are right censored.

**Examples**

```
require(graphics)
data(heights)
coplot(height ~ age | id, data = heights, panel=panel.smooth,
      show.given=FALSE, xlab='age (years)', ylab='height (cm)', pch=19)
```

### ifun

Invert an expression defining a data transformation

**Description**

Given a transformed variable and the expression used to transform it, ifun creates a function containing the inverse expression that will back-transform the variable.

**Usage**

```
ifun(expr, verbose = FALSE)
```

**Arguments**

- **expr**  a single-variable call or quoted expression to be inverted. The variable’s name in expr is referred to here as varname.
- **verbose**  a logical controlling printing of the intermediate functions $f(\cdot)$, $g(\cdot)$, $h(\cdot)$ etc (see ‘Details’).
Details

ifun returns the inverting function such that ifun(expr)(eval(expr)) = varname, where expr can include any of the invertible functions in the Math and Ops groups, plus identity and I.

To illustrate its use, consider variants of the sitar model height ~ age where age and/or height are transformed, e.g. height ~ log(age) or log(height) ~ sqrt(age). Each model is of the form y ~ x but the units of x and y vary.

The models are compared by plotting the fitted curves in their original units, by first applying suitable functions to back-transform x and y. For example with log(age), where expr = quote(log(age)), the function ifun = function(x) exp(x) back-transforms eval(expr) to give age. See the first example.

ifun generalises this process for increasingly complex expr, as the next two examples show.

The final example shows ifun in action with plot.sitar, which uses ifun as the default function for arguments xfun and yfun - they are used to back-transform x and y using the values of expr for x and y extracted from the model’s sitar call.

Structuring expr suitably ensures it can be inverted - it should contain a single mention of a single variable (varname here), and possibly functions such as f(.), g(.), h(.) etc such that expr = f(g(h((varname)))). The number of such functions is in principle unlimited. ifun returns function(x) h^{-1}(g^{-1}(f^{-1}(x))), which ensures that expr is invertible so long as the individual functions are invertible.

Value

The required inverting function, with single argument x. Its "varname" attribute contains varname as a character string.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

plot.sitar

Examples

## for best effect run all the code

## define varname variable
(age <- 1:9)

## simple case - age transformed to log(age)
(expr <- quote(log(age)))

## transformed age
eval(expr)

## inverting function, with "varname" attribute set to "age"
ifun(expr)

## inverted transformed age identical to age
all.equal(age, ifun(expr)(eval(expr)))
## more complex case - age transformed to log age since conception

```r
(expr <- quote(log(age + 0.75))
```

## inverting function

```r
ifun(expr)
```

## inverted transformed age identical to age

```r
all.equal(age, ifun(expr)(eval(expr)))
```

## ludicrously complex case involving exp, log10, ^, pi and trigonometry

```r
(expr <- quote((exp(sin(pi * log10(age + 0.75)/2) - 1)^4)))
```

## inverting function, showing intermediate stages

```r
ifun(expr, verbose=TRUE)
```

## identical to original

```r
all.equal(age, ifun(expr)(eval(expr)))
```

## example of plot.sitar back-transforming transformed x and y in sitar models

```r
## fit sitar models
m1 <- sitar(x=age, y=height^2, id=id, data=heights, df=6)
m2 <- update(m1, x=log(age+0.75), y=height)
```

## default plot options for xfun & yfun back-transform x & y to original scales

```r
## xfun=ifun(x$call.sitar$x)
## yfun=ifun(x$call.sitar$y)
```

## compare mean curves for the two models where x & y are on the original scales

```r
plot(m1, 'd', las=1)
lines(m2, 'd', col=2)
```

---

**iotf**

**IOTF international body mass index reference**

### Description

The IOTF (International Obesity TaskForce) BMI growth reference (Cole and Lobstein 2012), fitted by the LMS method and summarised by values of L, M and S by sex and postnatal age from 2 to 18 years.

### Usage

```r
iotf
```

### Format

A tibble with 66 observations on the following 5 variables:

- **years** numeric vector - postnatal age in years
- **L.bmi** numeric vector
- **M.bmi** numeric vector
- **S.bmi** numeric vector
- **sex** two-level factor with level 1 male and level 2 female
Details

The IOTF cutoffs for overweight and obesity (and also thinness) (see Cole et al. 2000, 2007) can be obtained from this BMI reference. See the example for how to convert between cutoffs and z-scores.

The L, M and S values for each measurement correspond respectively to the Box-Cox power, median and coefficient of variation of the distribution by age and sex (Cole & Green 1992). The measurement short name and units for LMS2z are bmi (kg/m2).

Source

The values are tabulated in the Excel spreadsheet IOTF_LMS.xls provided with the Excel add-in LMSgrowth from https://www.healthforallchildren.com/shop-base/software/lmsgrowth/

References


Examples

data(iotf)
## calculate z-scores by sex corresponding to IOTF cutoffs for thinness, ## overweight and obesity
co <- data.frame(cutoff = c(16, 17, 18.5, 25, 30),
   grade = c('thinness 3', 'thinness 2', 'thinness 1',
     'overweight', 'obesity'))
sexes <- c('boys', 'girls')
with(co,
   cbind(co, lapply(setNames(sexes, sexes), function(x)
     LMS2z(x = 18, y = cutoff, sex = x,
       measure = 'bmi', ref = 'iotf')))))

LMS2z

Convert to/from measurement from/to z-score with growth reference

Description

A function to convert between measurements and z-scores using a growth reference previously fitted by the LMS method.
Usage

LMS2z(x, y, sex, measure, ref, toz = TRUE, LMStable = FALSE)

Arguments

x
vector of ages in units of years.

y
vector or one-column matrix of either measurements or z-scores, depending on
the value of toz.

sex
vector where 1/2 = males/females = boys/girls = TRUE/FALSE, based on the
uppercased first character of the string.

measure
unique measurement name, as character string, the choice depending on the
choice of ref (see e.g. references uk90, who06 and ukwhopt).

ref
unique growth reference, either as name or character string, available as a data
object or data frame (e.g. uk90, who06 or ukwhopt).

toz
logical set to TRUE for conversion from measurement to z-score, or FALSE for
the reverse.

LMStable
logical set to TRUE to return the associated LMS table as a data frame in at-
tribute LMStable.

Details

Growth references fitted by the LMS method consist of a table of L, M and S values by age and
sex. Vectors of L, M and S corresponding to x and sex are extracted using cubic interpolation and
passed to either cLMS or zLMS, depending on toz.

Disjunct references are supported, where there is a disjunction in the centiles at a particular age.
This may be because the measurement changes, e.g. from length to height, or because two different
references have been joined together. The disjunction is flagged by including two rows at the
common age, but with different L, M and S values, and measurements at this age are ascribed to the
older reference. For example the who06 reference has a disjunction at 2 years reflecting the switch
from length to height. As a result height at just below and just above 2 years returns a different
z-score.

Value

A vector or matrix containing the transformed values. If y is a vector then a vector of length(x)
is returned, else if y is a one-column matrix then a matrix is returned, with length(x) rows and
length(y) columns. The matrix row names are set to x, and the column names to either y or if toz
is FALSE, z2cent(y). If LMStable is TRUE the associated LMS table is returned as a data frame
in attribute LMStable.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

z2cent. The LMS method can be fitted to data using the package gamlss with the BCCG or BCCGo
family, where nu (originally lambda), mu and sigma correspond to L, M and S respectively.
LMSfit

Examples

## convert girls' heights data to UK 90 z-scores
data(heights)
data(uk90)
with(heights, LMS2z(age, height, sex = 2, measure = 'ht', ref = 'uk90'))

## construct table of boys' weight centiles by age for WHO standard
data(who06)
zs <- -4:4*2/3 # z-scores for 9 centiles
ages <- 0:20/4 # 3-month ages to 5 years
LMS2z(ages, as.matrix(zs), sex = 'm', measure = 'wt', ref = who06,
toz = FALSE, LMStable = TRUE)

LMSfit

Estimate LMS curves from tabulated growth reference centiles

Description

A function to summarise an existing set of growth reference centiles as the L, M and S curves of the LMS method.

Usage

LMSfit(
  x,
  y,
  sex,
  data = parent.frame(),
  centiles = c(3, 10, 25, 50, 75, 90, 97),
  df = c(6, 10, 8),
  L1 = FALSE,
  plot = TRUE,
  ...
)

Arguments

x vector of tabulated ages.
y matrix of corresponding measurement centiles, e.g. of height or weight, with
nrows = length(x) and ncols = length(centiles).
sex two-level factor where level 1 corresponds to male and level 2 to female.
data optional data frame containing x, y and sex.
centiles vector of centiles corresponding to the columns of y, default c(3, 10, 25, 50, 75, 90, 97).
df
length-3 vector with the cubic smoothing spline equivalent degrees of freedom (edf) for the L, M and S curves, default c(6, 10, 8).
L1
logical constraining the L curve to 1, i.e. a Normal distribution, default FALSE.
plot
logical to plot the estimated L, M and S curves, default TRUE.
... optional graphical parameters for the plots.

Details
At each age the optimal Box-Cox power \( L_{opt} \) is estimated to render the centiles closest to Normal, and the corresponding median \( M_{opt} \) and coefficient of variation \( S_{opt} \) are derived. The three sets of values are then smoothed across age to give L, M and S.

Value
A list with the results:

- `list("LMS")` data frame of sex, x, L, M, S, Lopt, Mopt, Sopt.
- `list("ey")` matrix of predicted values of \( y \).
- `list("ez")` matrix of predicted values of \( z \).
- `list("fit")` matrix of summary statistics for \( ey \), giving for each column cmean the mean centile, zmean the mean z-score, zSD the SD of the z-score, and zmin and zmax the minimum and maximum z-scores.

Author(s)
Tim Cole <tim.cole@ucl.ac.uk>

See Also
LMS2z, z2cent. The LMS method can be fitted to data using the package gamlss with the BCCG family, where nu (originally lambda), mu and sigma correspond to L, M and S respectively.

Examples
```r
## first construct table of boys weight centiles by age for WHO standard
data(who06)
zs <- -4:4*2/3 # z-scores for centiles
ages <- 0:12/4 # ages 0-3 years by 3 months
v <- vapply(as.list(zs), function(z)
  LMS2z(ages, z, sex = 1, measure = 'wt', ref = 'who06', toz = FALSE),
  rep(0, length(ages))
  round(v, 2)

## then back-calculate the original LMS curves and display summary statistics
LMSfit(x=ages, y=v, sex=1, centiles=pnorm(zs)*100, plot=FALSE)
```
mplot

*Plot multiple growth curves*

**Description**

Function to plot multiple growth curves indexed by subject id.

**Usage**

```r
mplot(x, y, id, data = parent.frame(), subset = NULL, add = FALSE, ...)
```

**Arguments**

- `x` vector of x coordinates.
- `y` vector of y coordinates.
- `id` factor denoting subject levels.
- `data` optional dataframe containing `x`, `y` and `id`.
- `subset` optional logical defining a subset of rows in `data`.
- `add` optional logical defining whether the plot is pre-existing (TRUE) or new (FALSE).
- `...` Further graphical parameters (see `par`) may also be supplied as arguments, particularly background colour `bg`, character expansion `cex`, colour `col`, line type `lty`, line width `lwd` and character `pch`.

**Details**

The arguments `x`, `y` and `id` can be given as character strings. The `par` parameters can be functions of vector variables in `data`, e.g. to colour curves separately by `id` use: `col = id`.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>

**Examples**

```r
mplot(age, height, id, heights, col=id)
```
ob_convertr

Convert between IOTF, WHO and CDC prevalence rates for child thinness, overweight and obesity

Description
Child thinness, overweight and obesity are defined as the child’s body mass index (BMI) lying beyond a pre-specified reference cutoff. Three references are compared: IOTF (International Obesity Task Force), WHO (World Health Organization) and CDC (US Centers for Disease Control and Prevention), each of which have their own cutoffs. ob_convertr takes age-sex-specific prevalence rates of thinness, overweight or obesity based on one of the cutoffs, and converts them to the corresponding rates based on a different cutoff. ob_convertr2 uses paired prevalence rates of overweight and obesity on one cutoff to estimate those based on another cutoff.

Usage

```r
ob_convertr(
  age,
  sex,
  from,
  to,
  pfrom = NA,
  pto = NA,
  data = parent.frame(),
  report = c("vector", "wider", "longer"),
  plot = c("no", "density", "compare")
)
```

```r
ob_convertr2(
  age,
  sex,
  from,
  to,
  pfrom = NA,
  pto = NA,
  data = parent.frame(),
  report = c("vector", "wider", "longer"),
  plot = c("no", "density", "compare")
)
```

Arguments

- **age**: vector of ages between 2 and 18 years corresponding to prevalence rates pfrom.
- **sex**: vector of sexes corresponding to pfrom, coded as either 'boys/girls' or 'male/female' or '1/2' (upper or lower case, based on the first character).
- **from**: name(s) of the BMI cutoff(s) on which the prevalence pfrom is based (see Details).
to: name(s) of the BMI cutoff(s) on which to base the predicted prevalence (see Details).

pfrom: vector of age-sex-specific percentage prevalence rates based on from (ob_convertr) or the names of two or more such prevalence rates (ob_convertr2).

pto: vector (needed for plot = "compare") of known percentage prevalence rates based on to (ob_convertr) or the names of two or more such prevalence rates (ob_convertr2).

data: optional data frame containing age, sex, pfrom and pto.

report: character controlling the format of the returned data: 'vector' for the estimated prevalence rates, 'wider' for the working tibble in wide format, i.e. the from and to data side by side, or 'longer' for the tibble in long format, i.e. two rows per rate, one for from and one for to. For ob_convertr2 the three settings return progressively more information.

plot: character controlling what if anything is plotted: 'no' for no plot, 'density' to display the BMI density distributions and cutoffs corresponding to from and to, or 'compare' to display the predicted prevalence rates plotted against the observed rates (pto).

Details

The IOTF cutoffs correspond to the value of BMI \((kg/m^2)\) at age 18: IOTF 35 (morbid obesity), IOTF 30 (obesity), IOTF 25 (overweight), IOTF 18.5 (grade 1 thinness), IOTF 17 (grade 2 thinness) and IOTF 16 (grade 3 thinness).

The WHO cutoffs correspond to BMI z_scores. Age 5-19 years, WHO +2 (obesity), WHO +1 (overweight) and WHO -2 (thinness). Age 0-5 years, WHO +3 (obesity), WHO +2 (overweight) and WHO -2 (thinness).

The CDC cutoffs correspond to BMI centiles: CDC 95 (obesity), CDC 85 (overweight) and CDC 5 (thinness).

Note: the overweight category needs to be analysed as overweight prevalence plus obesity prevalence, i.e. the prevalence above the overweight cutoff. To predict overweight prevalence excluding obesity prevalence, first calculate predicted overweight prevalence including obesity then subtract predicted obesity prevalence.

The algorithms for ob_convertr and ob_convertr2 are distinguished by the number of prevalence rates used for the prediction. For ob_convertr (Cole & Lobstein, 2022) just one rate is used – in this case the algorithm is commutative, meaning that converting a prevalence rate from cutoff A to cutoff B and then from B to A returns the original value. from and to are the names of the cutoffs, and pfrom and optionally pto are vectors of percentage prevalence rates.

ob_convertr2 uses two known prevalence rates (Cole & Lobstein, 2023), typically overweight and obesity based on one reference. It is more accurate than ob_convertr though not exactly commutative. from and to are the names of the cutoffs as length-2 character strings, while pfrom and optionally pto are length-2 character strings giving the names of the corresponding vector prevalence rates. For convenience the from or to names 'CDC', 'IOTF' or 'WHO' expand to the corresponding pairs of cutoffs for overweight and obesity, e.g. 'CDC' expands to c("CDC 85", 'CDC 95").
Alternatively `ob_convertr2` can be used to interpolate or extrapolate to one or more specified z-score cutoffs assuming the same reference for all cutoffs. Here the values of from and to are numerical z-score cutoffs, with at least two for from. See the final example.

The algorithms require the prevalences of obesity and overweight net of obesity to be non-zero, and if they are zero they are set to missing.

**Value**

The predicted prevalence rates, optionally with a plot visualizing the findings, depending on the `report` and `plot` settings. Each predicted rate is given the name of the relevant cutoff followed by "pred".

With `report` set to "wider" or "longer", extra information is returned reflecting the internal workings of the algorithms. In particular `ob_convertr2` returns $b$ the regression coefficient of z-score prevalence on z-score cutoff as described in Cole & Lobstein (2023).

If a `plot` is selected, the underlying data and plot are returned invisibly with names `data` and `plot`.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>

**References**


**Examples**

```r
## convert 10% IOTF overweight prevalence (cutoff IOTF 25, including obesity)
## in 8-year-old boys to overweight prevalence for cutoff WHO +1
ob_convertr(age = 8, sex = 'boys', from = 'IOTF 25', to = 'WHO +1', pfrom = 10)
```
## compare the BMI density functions and cutoffs for IOTF and WHO
## in 8-year-old boys
ob_convertr2(age = 8, sex = 'boys', from = 'IOTF', to = 'WHO', plot = 'density')

## convert IOTF overweight prevalence to WHO overweight prevalence
## and compare with true value - boys and girls aged 7-17 (22 groups)
## note the need to first add obesity prevalence to overweight prevalence
data(deren)
deren <- within(deren, {
  CDC85 = CDC85 + CDC95
  IOTF25 = IOTF25 + IOTF30
  \('WHO+1' = \('WHO+1' + \('WHO+2'\')\))
  ob_convertr(age = Age, sex = Sex, from = 'IOTF 25', to = 'WHO +1',
              pfrom = IOTF25, pto = 'WHO+1', data = deren, plot = 'compare')

## convert IOTF overweight and obesity prevalence to WHO using
## ob_convertr2 - which is more accurate than ob_convertr
ob_convertr2(age = Age, sex = Sex, from = 'IOTF', to = 'WHO',
             pfrom = c('IOTF25', 'IOTF30'), pto = c('WHO+1', 'WHO+2'),
             data = deren, plot = 'compare')

## extrapolate WHO overweight and obesity prevalence (cutoffs +1 and +2)
## to severe obesity prevalence based on cutoffs +2.5 or +3
ob_convertr2(Age, Sex, from = 1:2, to = c(2.5, 3),
             pfrom = c('WHO+1', 'WHO+2'), data = deren, report = 'wider')

---

### optimal_design

**Optimal design for growth reference centile studies**

#### Description

Two functions for estimating optimal sample size and sample composition when constructing growth reference centiles.

#### Usage

```r
optimal_design(z = -2, lambda = NA, N = NA, SEz = NA, age = 10)
n_agegp(
  z = -2,
  lambda = NA,
  N = NA,
  SEz = NA,
  minage = 0,
  maxage = 20,
  n_groups = 20
)
```
Arguments

- **z**: z-score on which to base the design, with default -2 which equates to the 2nd centile. If NA, optimal z is calculated from lambda.
- **lambda**: power of age that defines the sample composition. The default NA means calculate optimal lambda from z.
- **N**: total sample size per sex. The default NA means calculate from z or lambda, and SEz if provided.
- **SEz**: target z-score standard error. The default NA means calculate from z or lambda, and N if provided.
- **age**: age at which to calculate SEz. The default 10 returns mean SEz, and if z or lambda are optimal SEz is independent of age.
- **minage**: youngest age (default 0).
- **maxage**: oldest age (default 20).
- **n_groups**: number of age groups (default 20).

Details

Studies to construct growth reference centiles using GAMLSS need to be of optimal size. Cole (SMMR, 2020) has shown that the sample composition, i.e. the age distribution of the measurements, needs to be optimised as well as the sample size. Sample composition is defined in terms of the age power lambda which determines the degree of infant oversampling.

There are two criteria that determine the optimal sample size and sample composition: the centile of interest (as z-score z) and the required level of precision for that centile (as the z-score standard error SEz).

Value

For `optimal_design`, a tibble with columns:

- **z**: as above.
- **lambda**: as above.
- **N**: as above.
- **SEz**: as above.
- **age**: as above.
- **p**: the centile corresponding to z.
- **plo**: lower 95% confidence interval for p.
- **phi**: upper 95% confidence interval for p.

For `n_agegp`, a tibble giving the numbers of measurements to be collected per equal width age group, with columns:

- **n_varying**: numbers for equal width age groups.
- **age**: mean ages for equal width age groups.
- **n**: number for each unequal width age group (only for longitudinal studies).
- **age_varying**: target ages for unequal width age groups (only for longitudinal studies).
Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

gamlss to fit the centiles with the BCCG, BCT or BCPE family.

Examples

## estimate optimal sample composition lambda and precision SEz for 9 centiles
## spaced 2/3 of a z-score apart, based on a sample of 10,000 children
optimal_design(z = -4:4*2/3, N = 10000)

## calculate age group sizes optimised for centiles from the 50th to the 99.6th
## (or equivalently from the 50th to the 0.4th)
## with a sample of 10,000 children from 0 to 20 years in one-year groups
purrr::map_dfc(0:4*2/3, ~{
  n_agegp(z = .x, N = 10000) %>%
  dplyr::select(!l2cent(.x) := n_varying)
}) %>%
  dplyr::bind_cols(tibble::tibble(age = paste(0:19, 1:20, sep='-')), .)

---

pdLMS

Plot frequency distributions(s) for given L, M and S values in LMS method

Description

The LMS method defines frequency distributions in terms of L, M and S parameters. pdLMS plots one or more LMS distributions and optionally returns specified centiles on each distribution.

Usage

pdLMS(
  L = 1,
  M = 1,
  S = 0.2,
  zcent = NULL,
  zlim = 3.5,
  N = 1000,
  plot = TRUE,
  ...
)
Arguments

- **L**: vector of Box-Cox transformation (lambda) values, L in the LMS method (default 1 corresponding to the Normal distribution).
- **M**: vector of medians (mu), M in the LMS method (default 1).
- **S**: vector of coefficients of variation (sigma), S in the LMS method (default 0.2).
- **zcent**: optional vector of z-scores for conversion to the measurement scale under each distribution.
- **zlim**: scalar defining z-score limits underlying x-axis (default 3.5).
- **N**: number of points per distribution curve (default 1000).
- **plot**: logical for plotting (default TRUE).
- **...**: Further graphical parameters (see **par**) may also be supplied as arguments, particularly colour **col**, line type **lty**, line width **lwd** and character **pch**.

Details

L, M and S should all be the same length, recycled if necessary.

Value

An invisible list with the following components:

- **x**: vector of x values for plotting.
- **density**: matrix of densities for each distribution.
- **centile**: matrix of measurement centiles corresponding to **zcent** under each distribution.

The distributions can be plotted with `matplot(x, density, type='l')`.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

z2cent, LMS2z, cLMS

Examples

```r
## plot normal distribution
pdLMS()
## compare variety of distributions
## with centiles corresponding to +3 z-scores
##
pdLMS(L=-2:3, M=2:3, S=1:3/10, zcent=3, lty=1)
```
Description

plot and lines methods for objects of class sitar, providing various flavours of plot of the fitted growth curves. Also helper functions to return the data for plotting, e.g. with ggplot2.

Usage

```r
## S3 method for class 'sitar'
plot(
x,
opt = "dv",
labels = NULL,
apv = FALSE,
xfun = identity,
yfun = identity,
subset = NULL,
ns = 101,
design = NULL,
abc = NULL,
trim = 0,
add = FALSE,
nlme = FALSE,
returndata = FALSE,
...
,xlab = NULL,
 ylab = NULL,
 vlab = NULL,
 xlim = c(NA, NA),
 ylim = c(NA, NA),
 vlim = c(NA, NA),
 legend = list(x = "topleft", inset = 0.04, bty = "o")
)
```

```r
## S3 method for class 'sitar'
lines(x, ...)
```

```r
plot_d(x, ...)
```

```r
plot_v(x, ...)
```

```r
plot_D(x, ...)
```

```r
plot_V(x, ...)
```
plot.sitar

plot_u(x, ...)

plot_a(x, ...)

plot_c(x, ...)

Arguments

x object of class sitar.

opt character string containing a subset of letters corresponding to the options: 'd' for fitted Distance curve, 'v' for fitted Velocity curve, 'c' for fitted Crosssectional distance curve, 'D' for individual fitted Distance curves, 'V' for individual fitted Velocity curves, 'u' for Unadjusted individual growth curves, and 'a' for Adjusted individual growth curves. Options 'dvcDV' give spline curves, while 'ua' give data curves made up as line segments. If both distance and velocity curves are specified, the axis for the velocity curve appears on the right side of the plot (y2), and a legend identifying the distance and velocity curves is provided.

labels optional character vector containing plot labels for x, y and y velocity from the original SITAR model. The three elements can alternatively be provided via parameters xlab, ylab and vlab. The latter take precedence. Default labels are the names of x and y, and "y velocity", suitably adjusted to reflect any back-transformation via xfun and yfun.

apv optional logical specifying whether or not to calculate the age at peak velocity from the velocity curve. If TRUE, age at peak velocity is calculated as the age when the second derivative of the fitted curve changes from positive to negative (after applying xfun and/or yfun). Age at peak velocity is marked in the plot with a vertical dotted line, and its value, along with peak velocity, is printed and returned. NB their standard errors can be obtained using the bootstrap with the function apv.se. Values of apv for individual subjects or groups are also returned invisibly.

xfun optional function to be applied to the x variable prior to plotting (default identity, see Details).

yfun optional function to be applied to the y variable prior to plotting (default identity, see Details).

subset optional logical vector of length x defining a subset of data rows to be plotted, for x and data in the original sitar call.

ns scalar defining the number of points for spline curves (default 101).

design formula defining the variables to use to group data for multiple mean distance and/or velocity curves (opt = 'dv'). By default includes all the categorical variables named in a.formula, b.formula, c.formula and d.formula.

abc vector of named values of random effects a, b, c and d used to define an individual growth curve, e.g. abc = c(a = 1, c = -0.1). Alternatively a single character string defining an id level whose random effect values are used. If abc is set, level is ignored. If abc is NULL (default), or if a, b, c or d values are missing, values of zero are assumed.
trim number (default 0) of long line segments to be excluded from plot with option 'u' or 'a'. See Details.

add optional logical defining if the plot is pre-existing (TRUE) or new (FALSE). TRUE is equivalent to using lines.

nlme optional logical which set TRUE plots the model as an nlme object, using plot.nlme arguments.

returndata logical defining whether to plot the data (default FALSE) or just return the data for plotting (TRUE). See Value.

... Further graphical parameters (see par) may also be supplied as arguments, e.g. line type lty, line width lwd, and colour col. For the velocity (y2) plot y2par can be used (see Details).

xlab optional label for x axis

ylab optional label for y axis

vlab optional label for v axis (velocity)

xlim optional x axis limits

ylim optional y axis limits

vlim optional v axis limits

legend optional list of arguments for legend with distance-velocity plots

Details

For options involving both distance curves (options 'dcDua') and velocity curves (options 'vV') the velocity curve plot (with right axis) can be annotated with par parameters given as a named list called y2par. To suppress the legend that comes with it set legend = NULL.

The transformations xfun and yfun are applied to the x and y variables after back-transforming any transformations in the original SITAR call. So for example if y = log(height) in the SITAR call, then yfun is applied to height. Thus the default yfun = identity has the effect of back-transforming the SITAR call transformation - this is achieved by setting yfun = yfun(ifun(x$call.sitar$y)). For no transformation set yfun = NULL. The same applies to xfun.

For models that include categorical fixed effects (e.g. a.formula = ~sex + region) the options 'dv' plot mean curves for each distinct group. Any continuous (as opposed to grouped) fixed effect variables are set to their mean values in the plots, to ensure that the mean curves are smooth. Setting design allows the grouping variables to be selected, e.g. design = ~sex, and design = ~1 gives a single mean curve. The resulting plots can be formatted with par in the usual way, indexed either by the individual grouping variables (e.g. sex or region in the example) or the subject factor id which indexes all the distinct plots.

The helper functions plot_d, plot_v, plot_D, plot_V, plot_u, plot_a and plot_c correspond to the seven plot options defined by their last letter, and return the data for plotting as a tibble, e.g. for use with ggplot2. Setting returndata = TRUE works similarly but handles multiple options, returning a list of tibbles corresponding to each specified option.

The trim option allows unsightly long line segments to be omitted from plots with options 'a' or 'u'. It ranks the line segments on the basis of the age gap (dx) and the distance of the midpoint of the line from the mean curve (dy) using the formula abs(dx)/mad(dx) + abs(dy)/mad(dy) and omits those with the largest values.
plot.sitar

Value

If `returnData` is `FALSE` returns invisibly a list of (up to) three objects:

- `usr`: value of `par('usr')` for the main plot.
- `usr2`: the value of `par('usr')` for the velocity (y2) plot.
- `apv`: if argument `apv` is `TRUE` a named list giving the age at peak velocity (`apv`) and peak velocity (`pv`) from the fitted velocity curve, either overall or (with options `D` or `V`, invisibly) for all subjects.

If `returnData` is `TRUE` (which it is with the helper functions) returns invisibly either a tibble or named list of tibbles, containing the data to be plotted. The helper functions each return a tibble where the first three variables are `.x`, `.y` and `.id`, plus variable `.groups` for curves grouped by design) and other covariates in the model. Note that `.x` and `.y` are returned after applying `xfun` and `yfun`. Hence if for example `x = log(age)` in the SITAR call then `.x` corresponds by default to age.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

`mplot`, `plotclean`, `ifun`, `apv_se`

Examples

```r
## fit sitar model
m1 <- sitar(x = age, y = height, id = id, data = heights, df = 4)

## draw fitted distance and velocity curves
## with velocity curve in blue
## adding age at peak velocity (apv)
plot(m1, y2par = list(col = 'blue'), apv = TRUE)

## bootstrap standard errors for apv and pv
## Not run:
res <- apv_se(m1, nboot = 20, plot = TRUE)

## End(Not run)
## draw individually coloured growth curves adjusted for random effects
## using same x-axis limits as for previous plot
plot(m1, opt = 'a', col = id, xlim = xaxsd())

## add mean curve in red
lines(m1, opt = 'd', col = 'red', lwd = 2)

## add mean curve for a, b, c = -1 SD
lines(m1, opt = 'd', lwd = 2, abc = -sqrt(diag(getVarCov(m1))))

## use subset to plot mean curves by group
```
## compare curves for early versus late menarche
heights <- within(sitar::heights, {
  men <- abs(men)
  late <- factor(men > median(men))
})
# fit model where size and timing differ by early vs late menarche
m2 <- sitar(log(age), height, id, heights, 5,
  a.formula = ~late, b.formula = ~late)
## early group
plot(m2, subset = late == FALSE, col = 4, lwd = 3,
  y2par = list(col = 4, lwd = 2), ylim = range(heights$height))
## late group
lines(m2, subset = late == TRUE, col = 2, lwd = 3,
  y2par = list(col = 2, lwd = 2))
## add legend
legend("right", paste(c("early", "late"), "menarche"),
  1ty = 1, col = c(4, 2), inset = 0.04)
## alternatively plot both groups together
plot(m2, lwd = 3, col = late, y2par = list(lwd = 3, col = late))
legend("right", paste(c("early", "late"), "menarche"),
  lwd = 3, col = 1:2, inset = 0.04)
## draw fitted height distance curves coloured by subject, using ggplot
## Not run:
require(ggplot2)
ggplot(plot_D(m1), aes(.x, .y, colour = .id)) +
  labs(x = "age", y = "height") +
  geom_line(show.legend = FALSE)
## End(Not run)

plotclean

Plot multiple growth curves to identify outliers

### Description

A version of mplot to plot growth curves and identify outliers. When outliers are clicked on, and if id is specified, the corresponding growth curve is highlighted. If id is not specified the selected point is highlighted. Use right-click to exit.

### Usage

```r
plotclean(
  x,
  y,
  id = NULL,
  data = parent.frame(),
  n = length(x),
  par.out = list(pch = 20),
  ...
)
```
**Arguments**

- **x**: vector of x coordinates.
- **y**: vector of y coordinates.
- **id**: factor of subject levels indexing each growth curve.
- **data**: optional dataframe containing x, y and id.
- **n**: maximum number of points to be identified.
- **par.out**: list of optional graphical parameters to control appearance of selected outlying points and lines.
- **...**: Further graphical parameters (see `par`) may also be supplied as arguments for lines and points, particularly line type, lty, line width, lwd and color, col.

**Value**

`plotclean` returns either a vector `rows` (if data is not specified) or a list:

- **rows**: a vector of row numbers corresponding to the selected points.
- **data**: a subset of `data` consisting of rows `rows`, and columns `id`, x and y.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>

**Examples**

```r
if (interactive()) plotclean(age, height, id, heights)
```

---

**predict.sitar**

*Predict SITAR model*

**Description**

Predict method for sitar objects, based on `predict.lme`.

**Usage**

```r
## S3 method for class 'sitar'
predict(
  object, 
  newdata = getData(object), 
  level = 1L, 
  ..., 
  deriv = 0L, 
  abc = NULL, 
  xfun = identity, 
  yfun = identity
)
```
Arguments

object an object inheriting from class sitar.

newdata an optional data frame to be used for obtaining the predictions, defaulting to the data used to fit object. It requires named columns for x, and for id if level = 1, matching the names in object. Variables with the reserved names x=.x or id=.id take precedence over the model x and id variables. Any covariates in a.formula, b.formula, c.formula or d.formula can also be included. By default their values are set to the mean, so when level = 0 the prediction represents the mean curve.

level an optional integer vector giving the level(s) of grouping to be used in obtaining the predictions, level 0 corresponding to the population predictions. Defaults to level 1, and level = 0:1 fits both levels.

... other optional arguments: asList, na.action and naPattern.

deriv an optional integer specifying predictions corresponding to either the fitted curve or its derivative. deriv = 0 (default) specifies the distance curve, deriv = 1 the velocity curve and deriv = 2 the acceleration curve.

abc an optional named vector containing values of a subset of a, b, c and d, default NULL. Ignored if level = 0. It gives predictions for a single subject with the specified values of a, b, c and d, where missing values are set to 0. Alternatively abc can contain the value for a single id.

xfun an optional function to apply to x to convert it back to the original scale, e.g. if x = log(age) then xfun = exp. Only relevant if deriv > 0 - see Details.

yfun an optional function to apply to y to convert it back to the original scale, e.g. if y = sqrt(height) then yfun = function(z) z^2.

Details

When deriv = 1 the returned velocity is in units of yfun(y) per xfun(x). So if x and/or y are transformed, velocity in units of y per x can be obtained by specifying xfun and/or yfun to back-transform them appropriately.

Value

A vector of the predictions, or a list of vectors if asList = TRUE and level == 1, or a data frame if length(level) > 1.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

ifun for a way to generate the functions xfun and yfun automatically from the sitar model call.
Examples

data(heights)
## fit model
m1 <- sitar(x=age, y=height, id=id, data=heights, df=5)

## predictions at level 0
predict(m1, newdata=data.frame(age=9:16), level=0)

## predictions at level 1 for subject 5
predict(m1, newdata=data.frame(age=9:16, id=5), level=1)

## velocity predictions for subjects with early and late puberty
vel1 <- predict(m1, deriv=1, abc=c(b=-1))
mplot(age, vel1, id, heights, col=id)
vel1 <- predict(m1, deriv=1, abc=c(b=1))
mplot(age, vel1, id, heights, col=id, add=TRUE)

print.sitar  

Print SITAR model

Description

Print method for sitar objects, based on print.lme.

Usage

## S3 method for class 'sitar'
print(x, ...)

Arguments

x  an object inheriting class sitar.

... other optional arguments.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>
print.summary.sitar  
Print summary of SITAR model

Description
A print.summary method for sitar objects.

Usage
## S3 method for class 'summary.sitar'
print(x, verbose = FALSE, ...)

Arguments
x an object inheriting from class summary.sitar.
verbose a logical to control the amount of output.
... to specify extra arguments.

Value
A formatted summary of the object.

Author(s)
Tim Cole <tim.cole@ucl.ac.uk>

recalib  
Recalibrate x, y data using SITAR random effects

Description
A function to recalibrate x,y data using SITAR random effects

Usage
recalib(xc, yc, id = NULL, data, xcnew = NULL, ycnew = NULL, model, from, to)
**Arguments**

- **xc**: character vector defining column name(s) of x data to be recalibrated.
- **yc**: character vector defining column name(s) of y data to be recalibrated.
- **id**: factor defining from and to rows. If NULL then recalibrate all rows.
- **data**: dataframe containing xc, yc and id.
- **xcnew**: column names for replacement columns xc. If default NULL then use names xcnew1...
- **ycnew**: column names for replacement columns yc. If default NULL then use names ycnew1...
- **model**: sitar model defining the random effects to be used for recalibration.
- **from**: level of id defining existing data (must be a single row in coef(model)).
- **to**: level of id defining data to be recalibrated (a single row in coef(model)).

**Details**

recalib recalibrates the values of xc and yc based on model. xc values are changed to:

\[(xc-c(coef[from,"b"]))*exp(coef[from,"c"]-coef[to,"c"])+coef[to,"b"]\].

yc values are changed to: \(yc-coef[from,"a"]+coef[to,"a"]\).

**Value**

Returns the dataframe data with the from rows of xc and yc recalibrated.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>

---

**sitar**

Fit SITAR growth curve model

**Description**

SITAR is a method of growth curve analysis, based on nlme, that summarises a set of growth curves with a mean growth curve as a regression spline, plus a set of up to four fixed and random effects (a, b, c and d) defining how individual growth curves differ from the mean curve.

**Usage**

```r
sitar(
x, y, id, data, df,
```
knots,
  fixed = NULL,
  random = "a + b + c",
  pdDiag = FALSE,
  a.formula = ~1,
  b.formula = ~1,
  c.formula = ~1,
  d.formula = ~1,
  bounds = 0.04,
  start,
  xoffset = "mean",
  bstart = xoffset,
  returndata = FALSE,
  verbose = FALSE,
  correlation = NULL,
  weights = NULL,
  subset = NULL,
  method = "ML",
  na.action = na.fail,
  control = nlmeControl(msMaxIter = 100, returnObject = TRUE),
  keep.data = TRUE
)

## S3 method for class 'sitar'
update(object, ..., evaluate = TRUE)

Arguments

- **x**: vector of ages.
- **y**: vector of measurements.
- **id**: factor of subject identifiers.
- **data**: data frame containing variables x, y and id.
- **df**: degrees of freedom for cubic regression spline (0 or more, see Details).
- **knots**: vector of values for knots (default df quantiles of x distribution).
- **fixed**: character string specifying a, b, c, d fixed effects (default random or the subset of "a + b + c + d" within random).
- **random**: character string specifying a, b, c, d random effects (default "a+b+c"). Alternatively nlme formula e.g. "list(id = pdDiag(a + b + c ~ 1))".
- **pdDiag**: logical which if TRUE fits a diagonal random effects covariance matrix, or if FALSE (default) a general covariance matrix.
- **a.formula**: formula for fixed effect a (default ~ 1).
- **b.formula**: formula for fixed effect b (default ~ 1).
- **c.formula**: formula for fixed effect c (default ~ 1).
- **d.formula**: formula for fixed effect d (default ~ 1).
- **bounds**: span of x for regression spline, or fractional extension of range (default 0.04).
sitar

optional numeric vector of initial estimates for the fixed effects, or list of initial estimates for the fixed and random effects (see \texttt{nlme}).

\texttt{xoffset} optional value of offset for \texttt{x} (either "mean" (default), "apv" or value).

\texttt{bstart} optional starting value for fixed effect \texttt{b} (either "mean", "apv" or value (default \texttt{xoffset})).

\texttt{returndata} logical which if TRUE causes the model matrix to be returned, or if FALSE (default) the fitted model. Setting returndata TRUE is useful in conjunction with subset and \texttt{subsample} for simulation purposes.

\texttt{verbose} optional logical value to print information on the evolution of the iterative algorithm (see \texttt{nlme}).

\texttt{correlation} optional \texttt{corStruct} object describing the within-group correlation structure (see \texttt{nlme}).

\texttt{weights} optional \texttt{varFunc} object or one-sided formula describing the within-group heteroscedasticity structure (see \texttt{nlme}).

\texttt{subset} optional expression indicating the subset of the rows of data that should be used in the fit (see \texttt{nlme}).

\texttt{method} character string, either "REML" or "ML" (default) (see \texttt{nlme}).

\texttt{na.action} function for when the data contain NAs (see \texttt{nlme}).

\texttt{control} list of control values for the estimation algorithm (see \texttt{nlme}) (default \texttt{nlmeControl(returnObject = TRUE)}).

\texttt{keep.data} logical to control saving data as part of the model object (default TRUE).

\texttt{object} object of class \texttt{sitar}.

... further parameters for update consisting of any of the above \texttt{sitar} parameters.

\texttt{evaluate} logical to control evaluation. If TRUE (default) the expanded update call is passed to \texttt{sitar} for evaluation, while if FALSE the expanded call itself is returned.

Details

The SITAR model usually has up to three random effects (a, b and c), termed size, timing and intensity respectively. \texttt{df} sets the degrees of freedom for the mean spline curve, taking values from 1 (i.e. linear) upwards. In addition there is a random effect for the slope, d, which is fitted when \texttt{df} = 0, and combined with a, it provides the classic random intercept random slope model, which is similar to the 1 df spline model. In addition d can be fitted, along with a, b and c, to extend SITAR to model variability in the adult slope of the growth curve.

\texttt{xoffset} allows the origin of \texttt{x} to be varied, while \texttt{bstart} specifies the starting value for \texttt{b}, both of which can affect the model fit and particularly \texttt{b}. The values of \texttt{bstart}, knots and bounds are offset by \texttt{xoffset} for fitting purposes, and similarly for fixed effect \texttt{b}.

The formulae \texttt{a.formula}, \texttt{b.formula}, \texttt{c.formula} and \texttt{d.formula} allow for \texttt{cov.names} and can include functions and interactions. \texttt{make.names} is used to ensure that the names of the corresponding model terms are valid. The modified not the original names need to be specified in \texttt{predict.sitar}.

update updates the model by taking the \texttt{object} call, adding any new parameters and replacing changed ones. Where feasible the fixed and random effects of the model being updated are suitably modified and passed via the \texttt{start} argument.
Value
An object inheriting from class sitar representing the nonlinear mixed-effects model fit, with all the components returned by nlme (see nlmeObject for a full description) plus the following components:

- `fitnlme`: the function returning the predicted value of \( y \).
- `data`: copy of data (if keep.data true).
- `constants`: data frame of mean a-b-c-d values for unique combinations of covariates (excluding x).
- `call.sitar`: the internal sitar call that produced the object.
- `xoffset`: the value of xoffset.
- `ns`: the lm object providing starting values for the B-spline curve.

Generic functions such as print, plot, anova and summary have methods to show the results of the fit. The functions residuals, coef, fitted, fixed.effects, random.effects, predict, getData, getGroups, getCovariate and getVarCov can be used to extract some of its components.

Author(s)
Tim Cole <tim.cole@ucl.ac.uk>

Examples

```r
data(heights)
## fit simple model
(m1 <- sitar(x=age, y=height, id=id, data=heights, df=5))

## relate random effects to age at menarche (with censored values +ve)
## both a (size) and b (timing) are positively associated with age at menarche
(m2 <- update(m1, a.formula = ~abs(men), b.formula = ~abs(men), c.formula = ~abs(men)))
```

Description
A function to sample from a SITAR dataset for experimental design purposes. Two different sampling schemes are offered, based on the values of id and x.

Usage

```r
subsample(x, id, data, prob = 1, xlim = NULL)
```
Arguments

- **x**: vector of age.
- **id**: factor of subject identifiers.
- **data**: dataframe containing x and id.
- **prob**: scalar defining sampling probability. See Details.
- **xlim**: length 2 vector defining range of x to be selected. See Details.

Details

With the first sampling scheme xlim is set to NULL (default), and rows of data are sampled with probability prob without replacement. With the second sampling scheme xlim is set to a range within range(x). Subjects id are then sampled with probability prob without replacement, and all their rows where x is within xlim are selected. The second scheme is useful for testing the power of the model to predict later growth when data only up to a certain age are available. Setting xlim to range(x) allows data to be sampled by subject. The returned value can be used as the subset argument in sitar or update.sitar.

Value

Returns a logical the length of x where TRUE indicates a sampled value.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

- **sitar**

Examples

```r
## draw 50% random sample
s50 <- subsample(age, id, heights, prob=0.5)

## truncate age range to 7-12 for 50% of subjects
t50 <- subsample(age, id, heights, prob=0.5, xlim=c(7, 12))
```
summary.sitar

Create summary of SITAR model

Description

A summary method for sitar objects based on summary.lme.

Usage

## S3 method for class 'sitar'
summary(object, adjustSigma = TRUE, verbose = FALSE, ...)

Arguments

- **object**: object inheriting from class sitar.
- **adjustSigma**: optional logical (see summary.lme).
- **verbose**: optional logical to control the amount of output in print.summary.sitar.
- **...**: some methods for this generic require additional arguments. None are used in this method.

Value

an object inheriting from class summary.sitar with all components included in object (see lmeObject for a full description of the components) plus the components for summary.lme and the following components:

- **x.adj**: vector of length x in object with x values adjusted for subject-specific random effects b and c.
- **y.adj**: vector of length y in object with y values adjusted for subject-specific random effects a.
- **apv**: length 2 vector giving respectively age at peak velocity and peak velocity based on the fitted distance curve (using transformed x and y where specified).

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>
\texttt{timegap} \hspace{1cm} \textit{Select equally spaced ages from a vector of ages}

\textbf{Description}

\texttt{timegap} indexes elements in a vector of ages such that the indexed ages are spaced integer multiples of a time interval apart, to within a given tolerance. \texttt{timegap.id} is a wrapper to apply \texttt{timegap} within levels of factor \texttt{id}. The selected ages can then be split into age groups the specified time interval wide, ensuring that (virtually) every subject has at most one measurement per interval.

\textbf{Usage}

\begin{verbatim}
timegap(age, gap, tol = 0.1 * gap, multiple = FALSE)
timegap.id(
    age,
    id,
    data = parent.frame(),
    gap,
    tol = 0.1 * gap,
    multiple = FALSE
)

diffid(
    age,
    id,
    data = parent.frame(),
    lag = 1,
    differences = 1,
    sort = FALSE,
    keepNA = FALSE
)
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
\item \texttt{age} \hspace{1cm} vector of ages.
\item \texttt{gap} \hspace{1cm} numeric, the required positive time gap between selected ages.
\item \texttt{tol} \hspace{1cm} numeric, the positive tolerance around the gap (default 0.1 * gap).
\item \texttt{multiple} \hspace{1cm} logical, whether or not to return multiple solutions when found (default FALSE).
\item \texttt{id} \hspace{1cm} factor of subject ids.
\item \texttt{data} \hspace{1cm} data frame optionally containing age and id.
\item \texttt{lag} \hspace{1cm} an integer indicating which lag to use.
\item \texttt{differences} \hspace{1cm} an integer indicating the order of the difference.
\item \texttt{sort} \hspace{1cm} a logical indicating whether to first sort by id and age.
\item \texttt{keepNA} \hspace{1cm} a logical indicating whether to keep generated NAs.
\end{itemize}
Details

timegap calculates all possible differences between pairs of ages, expresses them as integer multiples of gap, restricts them to those within tolerance and identifies those providing the longest sequences. For sequences of the same length, those with the smallest standard deviation of successive differences (modulo the time interval) are selected.

Value

With timegap, for unique solutions, or multiple solutions with multiple FALSE, a vector of indices named with age. With timegap.id the subject vectors are returned invisibly, concatenated.

With multiple TRUE, where there are multiple solutions they are returned as a named matrix.

diffid returns diff(age) applied within id. With keepNA TRUE a suitable number of NAs are added at the end, while if FALSE all NAs are omitted.

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

Examples

data(heights)

## bin age into 1-year groups by id
## gives multiple measurements per id per year
with(heights, table(floor(age), id))

## now select heights measured multiples of 1 year apart
(tg1 <- timegap.id(age, id, heights, 1))

## no more than one measurement per id per year
with(heights[tg1, ], table(floor(age), id))

## most time intervals close to 1 year
summary(diffid(age, id, heights[tg1, ], lag=1))

uk90

UK 1990 growth reference

Description

The UK 1990 growth reference (Freeman et al 1995, Cole et al 1998) for height, weight, body mass index, circumferences and percent body fat, fitted by the LMS method and summarised by values of L, M and S by sex from 23 weeks gestation to 23 years.

Usage

uk90
Format

A tibble with 588 observations on the following 26 variables:

- **years** numeric vector
- **L.ht** numeric vector
- **M.ht** numeric vector
- **S.ht** numeric vector
- **L.wt** numeric vector
- **M.wt** numeric vector
- **S.wt** numeric vector
- **L.bmi** numeric vector
- **M.bmi** numeric vector
- **S.bmi** numeric vector
- **L.head** numeric vector
- **M.head** numeric vector
- **S.head** numeric vector
- **L.sitht** numeric vector
- **M.sitht** numeric vector
- **S.sitht** numeric vector
- **L.leglen** numeric vector
- **M.leglen** numeric vector
- **S.leglen** numeric vector
- **L.waist** numeric vector
- **M.waist** numeric vector
- **S.waist** numeric vector
- **L.bfat** numeric vector
- **M.bfat** numeric vector
- **S.bfat** numeric vector
- **sex** two-level factor with level 1 male and level 2 female

Details

The L, M and S values for each measurement correspond respectively to the Box-Cox power, median and coefficient of variation of the distribution by age and sex (Cole & Green 1992). The short names and units for each measurement (see LMS2z) are as follows: height (ht, cm), weight (wt, kg), body mass index (bmi, kg/m2), head circumference (head, cm), sitting height (sitht, cm), leg length (leglen, cm), waist circumference (waist, cm) and percent body fat (fat).

Source

The values are tabulated in the spreadsheet British1990.xls provided with the Excel add-in LMS-growth from: https://www.healthforallchildren.com/shop-base/software/lmsgrowth/
References


Examples

data(uk90)
## calculate median BMI in girls from birth to 10 years
LMS2z(x = 0:10, y = 0, sex = 2, measure = 'bmi', ref = 'uk90', toz = FALSE)

---

ukwhopt

**UK-WHO growth reference including preterm**

Description

The UK-WHO growth reference for height, weight, BMI and head circumference (see Wright et al 2010), fitted by the LMS method and summarised by values of L, M and S by sex from 26 weeks gestation to 20 years.

Usage

ukwhopt

Format

A tibble with 542 observations on the following 17 variables:

- **age_wm** numeric vector - age in weeks or months - see `wm`
- **wm** three-level factor indicating weeks or months: wkga = gestational weeks, wk = postnatal weeks, mth = postnatal months
- **years** numeric vector - age in years
- **L.ht** numeric vector
- **M.ht** numeric vector
- **S.ht** numeric vector
- **L.wt** numeric vector
- **M.wt** numeric vector
- **S.wt** numeric vector
- **L.bmi** numeric vector
- **M.bmi** numeric vector
S.bmi  numeric vector
L.head numeric vector
M.head numeric vector
S.head numeric vector

origin  two-level factor indicating the provenance of the data, with levels British1990 and WHO2006
sex  two-level factor with level 1 male and level 2 female

Details

The growth reference combines the birth section of the British 1990 growth reference (Cole et al 2011) from 26 to 42 weeks gestation, the WHO growth standard from 2 postnatal weeks to 4 years, and the British 1990 reference from 4 to 20 years.

Age is measured in years, where 40 weeks gestation is 0 years. The conversion from weeks gestation to years is: \( \text{years} = (\text{weeks} - 40) \times 7 / 365.25 \).

The L, M and S values for each measurement correspond respectively to the Box-Cox power, median and coefficient of variation of the distribution by age and sex (Cole & Green 1992). The measurement short names and units (see LMS2z) are as follows: height (ht, cm), weight (wt, kg), BMI (bmi, kg/m2) and head circumference (head, cm).

Source

The values are tabulated in the Excel spreadsheet UK_WHO_preterm.xls provided with the Excel add-in LMSgrowth from https://www.healthforallchildren.com/shop-base/software/lmsgrowth/

References


Examples

data(ukwhopt)
## calculate median birth weight (kg) in girls from 26 to 44 weeks gestation
v <- LMS2z(x = (26:44-40) * 7 / 365.25, y = 0, sex = 2, measure = 'wt',
        ref = 'ukwhopt', toz = FALSE)
setNames(v, 26:44)
Description

The UK-WHO growth reference for height, weight, BMI and head circumference (see Wright et al 2010), fitted by the LMS method and summarised by values of L, M and S by sex and postnatal age from term birth (see Details) to 20 years.

Usage

ukwhoterm

Format

A tibble with 512 observations on the following 15 variables:

- **years** numeric vector - postnatal age in years
- **L.ht** numeric vector
- **M.ht** numeric vector
- **S.ht** numeric vector
- **L.wt** numeric vector
- **M.wt** numeric vector
- **S.wt** numeric vector
- **L.bmi** numeric vector
- **M.bmi** numeric vector
- **S.bmi** numeric vector
- **L.head** numeric vector
- **M.head** numeric vector
- **S.head** numeric vector
- **origin** two-level factor indicating the provenance of the data, with levels British1990 and WHO2006
- **sex** two-level factor with level 1 male and level 2 female

Details

The growth reference combines term birth data from the British 1990 growth reference (Cole et al 2011), the WHO growth standard from 2 postnatal weeks to 4 years, and the British 1990 reference from 4 to 20 years.

Age is measured in years, and term birth corresponds to ages between 37 and 42 weeks gestation, where 40 weeks gestation is 0 years. The conversion is: 
\[ \text{years} = (\text{weeks} - 40) \times 7 / 365.25. \]

The L, M and S values for each measurement correspond respectively to the Box-Cox power, median and coefficient of variation of the distribution by age and sex (Cole & Green 1992). The measurement short names and units (see **LMS2z**) are as follows: height (ht, cm), weight (wt, kg), BMI (bmi, kg/m²) and head circumference (head, cm).
Source

The values are tabulated in the Excel spreadsheet UK_WHO_preterm.xls provided with the Excel add-in LMSgrowth from https://www.healthforallchildren.com/shop-base/software/lmsgrowth/

References


Examples

data(ukwhoterm)
## calculate median weight (kg) in girls from 0 to 10 years
v <- LMS2z(x = 0:10, y = 0, sex = 2, measure = 'wt',
  ref = 'ukwhoterm', toz = FALSE)
setNames(v, 0:10)

velout

---

Identify outliers with abnormal velocity in growth curves

Description

Quickly identifies putative outliers in a large number of growth curves.

Usage

velout(x, y, id, data, lag = 1, velpower = 0.5, limit = 5, linearise = FALSE)

Arguments

x
age vector.
y
outcome vector, typically weight or height.
id
factor identifying each subject.
data
data frame containing x, y and id.
lag
lag between measurements for defining growth velocity.
velpower
a value, typically between 0 and 1, defining the power of delta x to use when calculating velocity as delta(y)/delta(x)^velpower. The default of 0.5 is midway between velocity and increment.
limit
the number of standard deviations beyond which a velocity is deemed to be an outlier.
linearise
if TRUE y is converted to a residual about the median curve of y versus x.
Details

The algorithm works by viewing serial measurements in each growth curve as triplets (A-B-C) and comparing the velocities between them. Velocity is calculated as

\[
\text{diff}(y, \text{lag} = \text{lag}) / \text{diff}(x, \text{lag} = \text{lag}) ^ \text{velpower}
\]

Missing values for \(x\) or \(y\) are ignored. If any of the \(AB\), \(BC\) or \(AC\) velocities are abnormal (more than \(\text{limit}\) SDs in absolute value from the median for the dataset) the code for \(B\) is non-zero.

Value

Returns a data frame with columns: \(\text{id}, \text{x}, \text{y}\) (from the call), \(\text{code}\) (as described below), \(\text{vel1}, \text{vel2}\) and \(\text{vel3}\) (corresponding to the velocities \(AB\), \(BC\) and \(AC\) above). The 'data' attribute contains the name of 'data'.

Code is a factor taking values between 0 and 8, with 0 normal (see table below). Values 1-6 depend on the pattern of abnormal velocities, while 7 and 8 indicate a duplicate age (7 for the first in an individual and 8 for later ones). Edge outliers, i.e. first or last for an individual, have just one velocity. Code 4 indicates a conventional outlier, with both \(AB\) and \(BC\) abnormal and \(AC\) normal. Code 6 is an edge outlier. Other codes are not necessarily outliers, e.g. codes 1 or 3 may be adjacent to a code 4. Use \(\text{codeplot}\) to look at individual curves, and \(\text{zapvelout}\) to delete outliers.

<table>
<thead>
<tr>
<th>code</th>
<th>(AB+BC)</th>
<th>(AC)</th>
<th>interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>no outlier</td>
</tr>
<tr>
<td>0</td>
<td>0</td>
<td>NA</td>
<td>no outlier</td>
</tr>
<tr>
<td>1</td>
<td>0</td>
<td>1</td>
<td>rare pattern</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>0</td>
<td>complicated - look at curve</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>1</td>
<td>adjacent to simple outlier</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>0</td>
<td>single outlier</td>
</tr>
<tr>
<td>5</td>
<td>2</td>
<td>1</td>
<td>double outlier</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>NA</td>
<td>edge outlier</td>
</tr>
<tr>
<td>7</td>
<td>-</td>
<td>-</td>
<td>first duplicate age</td>
</tr>
<tr>
<td>8</td>
<td>-</td>
<td>-</td>
<td>later duplicate age</td>
</tr>
</tbody>
</table>

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

See Also

\(\text{codeplot}, \text{zapvelout}\)

Examples

```r
outliers <- velout(age, height, id, heights, limit=3)
```
Description

The WHO growth standard (WHO 2006) for height, weight, body mass index, circumferences and skinfold thicknesses, fitted by the LMS method and summarised by values of L, M and S by sex from birth to 5 years.

Usage

who06

Format

A tibble with 150 observations on the following 23 variables:

- years  age from 0 to 5 years
- L.ht   numeric vector
- M.ht   numeric vector
- S.ht   numeric vector
- L.wt   numeric vector
- M.wt   numeric vector
- S.wt   numeric vector
- L.bmi  numeric vector
- M.bmi  numeric vector
- S.bmi  numeric vector
- L.head numeric vector
- M.head numeric vector
- S.head numeric vector
- L.arm  numeric vector
- M.arm  numeric vector
- S.arm  numeric vector
- L.subscap numeric vector
- M.subscap numeric vector
- S.subscap numeric vector
- L.tricep numeric vector
- M.tricep numeric vector
- S.tricep numeric vector
- sex    two-level factor with level 1 male and level 2 female
Details

The L, M and S values for each measurement correspond respectively to the Box-Cox power, median and coefficient of variation of the distribution by age and sex (Cole & Green 1992). The short names and units for each measurement (see LMS2z) are as follows: height (ht, cm), weight (wt, kg), body mass index (bmi, kg/m2), head circumference (head, cm), arm circumference (arm, cm), subscapular skinfold (subscap, mm), and tricep skinfold (tricep, mm).

Source

https://www.who.int/toolkits/child-growth-standards

References


Examples

data(who06)
## calculate z-score for length 60 cm in boys at age 0:12 months
LMS2z(x = 0:12/12, y = 60, sex = 1, measure = 'ht', ref = 'who06')

who0607  The WHO 2006 growth standard and WHO 2007 growth reference

Description

The WHO growth standard (WHO 2006) and growth reference (2007) for height, weight and body mass index, fitted by the LMS method and summarised by values of L, M and S by sex from birth to 19 years.

Usage

who0607

Format

A tibble with 486 observations on the following 11 variables:

years  age from 0 to 19 years
L.ht  numeric vector
M.ht  numeric vector
S.ht  numeric vector
L.wt numeric vector
M.wt numeric vector
S.wt numeric vector
L.bmi numeric vector
M.bmi numeric vector
S.bmi numeric vector

sex two-level factor with level 1 male and level 2 female

Details

The L, M and S values for each measurement correspond respectively to the Box-Cox power, median and coefficient of variation of the distribution by age and sex (Cole & Green 1992). The short names and units for each measurement (see LMS2z) are as follows: height (ht, cm), weight (wt, kg) and body mass index (bmi, kg/m²).

References


Examples

data(who0607)
## calculate 98th centile for BMI in girls from birth to 19 years
round(
  setNames(
    LMS2z(x = 0:19, y = 2, sex = 2, measure = 'bmi', ref = 'who0607',
             toz = FALSE), 0:19), 1)

xaxsd

Par args xaxs and yaxs option 'd'

Description

Implements par('xaxs') and par('yaxs') option 'd'.

Usage

xaxsd(usr = par()$usr[1:2])
yaxsd(usr = par()$usr[3:4])
Arguments

usr  a length-2 vector defining the length of the x-axis or y-axis.

Details

Implements par('xaxs') and par('yaxs') option 'd', i.e. uses previous axis scales in a new plot.

Value

By default returns xlim/ylim args to match current setting of par()$usr, i.e. previous plot scales. Specifying usr gives scales with the usr args at the extremes. If par('xlog') or par('ylog') are set the returned limits are antilogged (to base 10).

Author(s)

Tim Cole <tim.cole@ucl.ac.uk>

Examples

```r
## generate and plot 100 data points
x <- rnorm(100)
y <- rnorm(100)
plot(x, y, pch=19)

## generate and plot 10 more
## constraining axis scales to be as before
x <- rnorm(10)
y <- rnorm(10)
plot(x, y, pch=19, xlim=xaxsd(), ylim=yaxsd())

## force axis extremes to be -3 and 3
plot(x, y, pch=19, xlim=xaxsd(c(-3,3)), ylim=yaxsd(c(-3,3)))
```

---

**xyadj**  
Adjust x and y variables for SITAR random effects

Description

xyadj Adjusts x and y and optionally v values for subject-specific random effects from a SITAR model.

Usage

```r
xyadj(object, x, y = 0, v = 0, id, abc = NULL, tomean = TRUE)
```
**Arguments**

- `object`: a SITAR model.
- `x`: a vector of x coordinates. If missing, x and y and id are obtained from object.
- `y`: a vector of y coordinates (default 0).
- `v`: a vector of velocity coordinates (default 0).
- `id`: a factor denoting the subject levels corresponding to x and y and v.
- `abc`: a data frame containing random effects for a, b, c and d (default `ranef(object)[id,]`).
- `tomean`: a logical defining the direction of adjustment. TRUE (default) indicates that individual curves are translated and rotated to match the mean curve, while FALSE indicates the reverse, the mean curve being translated and rotated to match individual curves.

**Details**

When `tomean = TRUE` the x and y and v values are adjusted to

\[(x - \text{xoffset} - b < \text{fixed}> - b < \text{random}>) \times \exp(c < \text{random}>) + \text{xoffset} + b < \text{fixed} > \]

\[y - a < \text{random} > - d < \text{random} > \times x\]

\[(v - d < \text{random} >) / \exp(c < \text{random}>)\]

When `tomean = FALSE` they are adjusted to

\[(x - \text{xoffset} - b < \text{fixed} >) / \exp(c < \text{random}>) + \text{xoffset} + b < \text{fixed} > + b < \text{random} > \]

\[y + a < \text{random} > + d < \text{random} > \times x\]

\[v \times \exp(c < \text{random}>) + d < \text{random} >\]

In each case missing values of the fixed or random effects are set to zero.

**Value**

The list of adjusted values:

- `x`: numeric vector.
- `y`: numeric vector the same length as x, or NULL.
- `v`: numeric vector the same length as x, or NULL.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>
Examples

data(heights)
## fit sitar model for height
m1 <- sitar(x = age, y = height, id = id, data = heights, df = 5)

## plot unadjusted data as growth curves
plot(m1, opt='u')

## overplot with adjusted data as points
with(heights, points(xyadj(m1), col='red', pch = 19))

---

**z2cent**

Express z-scores as centile character strings for plotting

---

**Description**

Converts z-scores, typically defining centiles in a growth chart, to character strings that can be used to label the centile curves.

**Usage**

z2cent(z)

**Arguments**

- **z**: a scalar or vector of z-scores.

**Value**

A character string is returned, the same length as `z`. Z-scores between the 1st and 99th centile are converted to centiles with one or two significant figures (lower tail) or to their complement (upper tail). For larger z-scores in absolute value the character consists of "SDS" appended to the z-score rounded to one decimal place.

**Author(s)**

Tim Cole <tim.cole@ucl.ac.uk>

**See Also**

cLMS

**Examples**

z2cent(-4:4)
z2cent(qnorm(0:100/100))
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