Package 'sitar'

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Type Package

Title Super Imposition by Translation and Rotation Growth Curve Analysis

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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

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sitar-package

SITAR (SuperImposition by Translation And Rotation) growth curve analysis

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 three fixed and random effects (a, b and c) 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**).

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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. The aim is for individual curves, adjusted for a, b and c, 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 mixedeffects 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.

Beath KJ. Infant growth modelling using a shape invariant model with random effects. Stat Med 2007;26:2547-64.

Cole TJ, Cortina Borja M, Sandhu J, et al. Nonlinear growth generates age changes in the moments of the frequency distribution: the example of height in puberty. Biostatistics 2008;9:159-71.

Cole TJ, Donaldson MD, Ben-Shlomo Y. SITAR–a useful instrument for growth curve analysis. Int J Epidemiol 2010;39:1558-66.

Gault EJ, Perry RJ, Cole TJ, et al. Effect of oxandrolone and timing of pubertal induction on final height in Turner's syndrome: randomised, double blind, placebo controlled trial. BMJ 2011;\-342:d1980.

Johnson L, Llewellyn CH, van Jaarsveld CHM, et al. Genetic and environmental influences on infant growth: prospective analysis of the Gemini twin birth cohort. PLoS ONE 2011;6:e19918.

Prentice A, Dibba B, Sawo Y, et al. The effect of prepubertal calcium carbonate supplementation on the age of peak height velocity in Gambian adolescents. Am J Clin Nutr 2012;96:1042-50.

Dean MC, Cole TJ. Human life history evolution explains dissociation between the timing of tooth eruption and peak rates of root growth. PLoS ONE 2013;8:e54534.

Cole TJ, Statnikov Y, Santhakumaran S, et al. Birth weight and longitudinal growth in infants born below 32 weeks gestation: a UK population study. Arch Dis Child Fetal Neonatal Ed 2014;99:F34-F40.

Cole TJ, Rousham EK, Hawley NL, et al. Ethnic and sex differences in skeletal maturation among the Birth to Twenty cohort in South Africa. Arch Dis Child 2014;100:138-43.

Cole TJ, Pan H, Butler GE. A mixed effects model to estimate timing and intensity of pubertal growth from height and secondary sexual characteristics. Ann Hum Biol 2014;41:7683.

Johnson L, van Jaarsveld CHM, Llewellyn CH, et al. Associations between infant feeding and the size, tempo and velocity of infant weight gain: SITAR analysis of the Gemini twin birth cohort. Int J Obes 2014;38:980-7.

Pizzi C, Cole TJ, Richiardi L, et al. Prenatal influences on size, velocity and tempo of iInfant growth: findings from three contemporary cohorts. PLoS ONE 2014;9:e90291.

Ward KA, Cole TJ, Laskey MA, et al. The Effect of Prepubertal Calcium Carbonate Supplementation on Skeletal Development in Gambian Boys-A 12-Year Follow-Up Study. J C E M 2014;99:3169-76.

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,
```

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apv_se

```
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)</pre>
```

berkeley

The Berkeley Child Guidance Study

Description

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

Usage

berkeley

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. The growth dataset in the fda package uses heights from the same study.

References

Tuddenham RD, Snyder MM. Physical growth of California boys and girls from birth to eighteen years. University of California Publications in Child Development 1954;1:183-364.

Examples

```
data(berkeley)
## 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)</pre>
```

BICadj

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 [AB]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

 $adjusted deviance = deviance - 2n((\lambda - 1) * log(gm) + log(abs(\lambda)))$

where λ is the power transform, and n and gm are the length and geometric mean of y. The variance explained is given by

$$\% explained = 100 * (1 - (\sigma_2/\sigma_1)^2)$$

where σ_1 is the fixed effects RSD and σ_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.

bupdate

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))))</pre>
```

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

```
## fit sitar model with b fixed effect starting value defaulting to 'mean'
m1 <- sitar(x=age, y=height, id=id, data=heights, df=5)
print(fixef(m1)['b'])
## refit with starting value chosen to minimise b-c correlation and df increased
m2 <- update(m1, bstart=bupdate(m1), df=6)</pre>
```

```
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

cLMS

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, 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 (hc, cm).

References

Cole TJ, Green PJ. Smoothing reference centile curves: the LMS method and penalized likelihood. Stat Med 1992;11:1305-19.

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. Vital Health Stat, 2002, 11, 246, 1-190.

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.
Μ	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:

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

$$x = M(1 + LSz)^{1/L})$$

where L is reset to 10⁻⁷ if it is zero. The LMS method is the same as the BCCG family in the gam1ss 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))
```

codeplot

Description

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

Usage

```
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

```
## 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)</pre>
```

dfpower

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

Description

dfpower fits a series of SITAR models tabulated by specified degrees of freedom and power transformations of x and y, returning a table of function values (e.g. BIC) applied to each model.

Usage

```
dfpower(
  model,
  df,
  xpowers,
  ypowers,
  FUN = BICadj,
  maxIter = nlmeControl()$maxIter,
  verbose = FALSE
)
```

Arguments

model	fitted sitar model to be updated.
df	vector of degrees of freedom to be fitted (defaults to df in model).
xpowers	vector of powers to apply to x (defaults to x power in model).
ypowers	vector of powers to apply to y (defaults to y power in model).
FUN	function to be tabulated (default BICadj, or e.g. AICadj or varexp).
maxIter	maximum number of iterations per fit.
verbose	logical controlling monitoring.

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dfset

Details

The function provides a convenient way to optimise the model's degrees of freedom and explore transformations of x and y, based by default on adjusted BIC. The function value is returned with changed sign if there is a warning, or as NA if there is an error. The run-time can be shortened by reducing maxIter, as the models that converge do so in relatively few iterations, and much of the run-time is spent on models that fail to converge.

FUN can be any function returning a single numerical value.

The returned table can be rearranged using aperm.

Value

Table or vector of returned values.

Author(s)

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

Examples

```
data(heights)
m1 <- sitar(log(age), height, id, heights, 4)
dfpower(m1, df=4:5, xpowers=0:1, maxIter=4)</pre>
```

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

х	vector of x coordinates.
У	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.

funcall

Value

degrees of freedom and value of FUN at minimum.

Author(s)

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

Examples

```
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

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>

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getData.sitar

Description

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

Usage

```
## 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 $y \sim x$.

Usage

```
getPeakTrough(x, y = NULL, peak = TRUE, takeoff = FALSE)
getPeak(x, y = NULL, peak = TRUE, takeoff = FALSE)
getTrough(x, y = NULL, peak = FALSE, takeoff = FALSE)
getTakeoff(x, y = NULL, peak = FALSE, takeoff = TRUE)
```

Arguments

х	vector.
У	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

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, if un 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 $f(.)$, $g(.)$, $h(.)$ etc
	(see 'Details').

Details

if un returns the inverting function such that if un(expr)(eval(expr)) = varname, where expr can include any of the invertible functions in the 'Math' and 'Ops' groups.

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 $y \sim x$ 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.

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

The final example shows if un in action with plot.sitar, which uses if un 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. if un 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)</pre>

```
## 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)))</pre>
```

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```
## more complex case - age transformed to log age since conception
(expr <- quote(log(age + 0.75)))
## inverting function
ifun(expr)
## inverted transformed age identical to age
all.equal(age, ifun(expr)(eval(expr)))
## ludicrously complex case involving exp, log10, ^, pi and trigonometry
(expr <- quote((exp(sin(pi * log10(age + 0.75)/2) - 1)^4)))</pre>
## inverting function, showing intermediate stages
ifun(expr, verbose=TRUE)
## identical to original
all.equal(age, ifun(expr)(eval(expr)))
## example of plot.sitar back-transforming transformed x and y in sitar models
## fit sitar models
m1 <- sitar(x=age, y=height^2, id=id, data=heights, df=6)</pre>
m2 <- update(m1, x=log(age+0.75), y=height)</pre>
## default plot options for xfun & yfun back-transform x & y to original scales
## 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
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

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 http://www.healthforallchildren.com/?product_cat=software.

References

Cole TJ, Green PJ. Smoothing reference centile curves: the LMS method and penalized likelihood. Stat Med 1992;11:1305-19.

Cole TJ, Bellizzi MC, Flegal KM, Dietz WH. Establishing a standard definition for child overweight and obesity worldwide: international survey. BMJ 2000;320:1240-3.

Cole TJ, Flegal KM, Nicholls D, Jackson AA. Body mass index cut offs to define thinness in children and adolescents: international survey. BMJ 2007;335:194-7.

Cole TJ, Lobstein T. Extended international (IOTF) body mass index cut-offs for thinness, overweight and obesity. Ped Obes 2012;7:284-94.

Examples

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.

LMS2z

Usage

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

Arguments

x	vector of ages in units of years.
У	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	measurement name, as character string, the choice depending on the choice of ref (see e.g. references uk90, who06 and ukwhopt).
ref	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.

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)</pre>
```

```
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.
У	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).

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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 Lopt is estimated to render the centiles closest to Normal, and the corresponding median Mopt and coefficient of variation Sopt 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

```
## 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

Description

Function to plot multiple growth curves indexed by subject id.

Usage

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

Arguments

x	vector of x coordinates.
У	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, par- ticularly 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

mplot(age, height, id, heights, col=id)

pdLMS

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 (de- fault 1 corresponding to the Normal distribution).
Μ	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).
Ν	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, par- ticularly 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

```
## 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)
```

plot.sitar Plot SITAR model

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

```
## S3 method for class 'sitar'
plot(
    x,
    opt = "dv",
    labels,
    apv = FALSE,
    xfun = I,
    yfun = I,
    subset = NULL,
    ns = 101,
    abc = NULL,
    trim = 0,
    add = FALSE,
    nlme = FALSE,
    returndata = FALSE,
```

plot.sitar

```
...,
  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")
)
## S3 method for class 'sitar'
lines(x, ...)
plot_d(x, ...)
plot_v(x, ...)
plot_D(x, ...)
plot_V(x, ...)
plot_u(x, ...)
plot_a(x, ...)
plot_c(x, ...)
```

Arguments

х	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 Ad- justed 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.
ару	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 sign (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.
xfun	optional function to be applied to the x variable prior to plotting (default I, see Details).
yfun	optional function to be applied to the y variable prior to plotting (default I, 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).
abc	vector of named values of random effects a, b and c 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 or c 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 1ty, line width 1wd, 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 inverting any transformations applied 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 = I has the effect of inverting the transformation. This is achieved by setting yfun = yfun(ifun(x\$call.sitar\$y)). For no transformation set yfun = NULL.

plot.sitar

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.

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.

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.
арv	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. The variable names are '.x', '.y' and (for curves grouped by subject) '.id'. 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

```
## 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))))
## 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

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

Arguments

х	vector of x coordinates.
У	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.

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predict.sitar

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, \boldsymbol{x} and $\boldsymbol{y}.$

Author(s)

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

Examples

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

predict.sitar Predict SITAR model

Description

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

Usage

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

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 or c.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 and c, default NULL. Ignored if $level = 0$. It gives predictions for a single subject with the specified values of a, b and c, 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

if un 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</pre>
```

print.sitar

```
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)</pre>
```

print.sitar

Print SITAR model

Description

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

Usage

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

Arguments

х	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

Х	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

хс	character vector defining column name(s) of x data to be recalibrated.
ус	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

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 three fixed and random effects (a, b and c) defining how individual growth curves differ from the mean curve.

Usage

```
sitar(
 х,
 у,
 id,
 data,
 df,
 knots,
  fixed = random,
  random = "a+b+c",
 pdDiag = FALSE,
 a.formula = \sim 1,
 b.formula = ~1,
  c.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)
)
```

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

Arguments

x	vector of ages.
У	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 (1 or more).
knots	vector of values for knots (default df quantiles of x distribution).
fixed	character string specifying a, b, c fixed effects (default random or the subset of "a+b+c" within random).
random	character string specifying a, b, c random effects (default "a+b+c"). Alterna- tively 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).
bounds	span of x for regression spline, or fractional extension of range (default 0.04).
start	optional numeric vector of initial estimates for the fixed effects, or list of initial estimates for the fixed and random effects (see nlme).
xoffset	optional value of offset for x (either "mean" (default), "apv" or value).
bstart	optional starting value for fixed effect b (either "mean", "apv" or value (default xoffset)).
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 subsample for simulation purposes.
verbose	optional logical value to print information on the evolution of the iterative algorithm (see nlme).
correlation	optional corStruct object describing the within-group correlation structure (see nlme).
weights	optional varFunc object or one-sided formula describing the within-group heteroscedasticity structure (see nlme).
subset	optional expression indicating the subset of the rows of data that should be used in the fit (see nlme).
method	character string, either "REML" or "ML" (default) (see nlme).
na.action	function for when the data contain NAs (see nlme).
control	list of control values for the estimation algorithm (see nlme) (default nlmeCon- trol(returnObject=TRUE)).
object	object of class sitar.
	further parameters for update consisting of any of the above sitar parameters.
evaluate	logical to control evaluation. If TRUE (default) the expanded update call is passed to sitar for evaluation, while if FALSE the expanded call itself is returned.

Details

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

The formulae a.formula, b.formula and c.formula can include functions and interactions, but 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 predict.sitar.

update updates the model by taking the 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 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.
call.sitar	the internal sitar call that produced the object.
xoffset	the value of xoffset.
ns	the 1m 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 resid, coef, fitted, fixed.effects, random.effects, predict, getData, getGroups, getCovariate and getVarCov can be used to extract some of its components.

Note that versions of sitar prior to 1.0.4 did not return fitnlme. Both plot and predict may require it, in which case they update the SITAR object on the fly, with a message. Also version 1.0.5 altered the defaults for xoffset and bstart. Models fitted with versions prior to 1.0.5 need refitting.

Author(s)

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

Examples

```
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 (tempo) are positively associated with age at menarche
amen <- abs(heights$men)
(m2 <- update(m1, a.form=~amen, b.form=~amen, c.form=~amen))</pre>
```

sitar

subsample

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

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

Arguments

х	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

summary.sitar

Examples

```
## 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))</pre>
```

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>

timegap

Description

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. timegap.id is a wrapper to apply timegap within levels of factor 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.

Usage

```
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
)
```

Arguments

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

uk90

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 LMSgrowth from:

http://www.healthforallchildren.com/shop-base/software/lmsgrowth/.

ukwhopt

References

Cole TJ, Green PJ. Smoothing reference centile curves: the LMS method and penalized likelihood. Stat Med 1992;11:1305-19.

Cole TJ, Freeman JV, Preece MA. British 1990 growth reference centiles for weight, height, body mass index and head circumference fitted by maximum penalized likelihood. Stat Med 1998;17:407-29.

Freeman JV, Cole TJ, Chinn S, et al. Cross sectional stature and weight reference curves for the UK, 1990. Arch Dis Child 1995;73:17-24.

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

ukwhopt

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: years = (weeks -40) * 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 http://www.healthforallchildren.com/?product_cat=software.

References

Cole TJ, Green PJ. Smoothing reference centile curves: the LMS method and penalized likelihood. Stat Med 1992;11:1305-19.

Cole TJ, Williams AF, Wright CM, et al. Revised birth centiles for weight, length and head circumference in the UK-WHO growth charts. Ann Hum Biol 2011;38:7-11.

Wright CM, Williams AF, Elliman D, et al. Using the new UK-WHO growth charts. BMJ 2010;340:c1140.

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)</pre>
```

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ukwhoterm

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: years = (weeks -40) * 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 http://www.healthforallchildren.com/?product_cat=software.

References

Cole TJ, Green PJ. Smoothing reference centile curves: the LMS method and penalized likelihood. Stat Med 1992;11:1305-19.

Cole TJ, Williams AF, Wright CM, et al. Revised birth centiles for weight, length and head circumference in the UK-WHO growth charts. Ann Hum Biol 2011;38:7-11.

Wright CM, Williams AF, Elliman D, et al. Using the new UK-WHO growth charts. BMJ 2010;340:c1140.

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)</pre>
```

```
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

х	age vector.
У	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.

velout

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

 $diff(y, lag = lag) / diff(x, lag = lag) ^ velpower$

Missing values for x or y are ignored. If any of the AB, BC or AC velocities are abnormal (more than 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: id, x, y (from the call), code (as described below), vel1, vel2 and 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 codeplot to look at individual curves, and zapvelout to delete outliers.

AB+BC	AC	interpretation
0	0	no outlier
0	NA	no outlier
0	1	rare pattern
1	0	complicated - look at curve
1	1	adjacent to simple outlier
2	0	single outlier
2	1	double outlier
1	NA	edge outlier
-	-	first duplicate age
-	-	later duplicate age
	0 0 0 1 1	$\begin{array}{cccc} 0 & 0 \\ 0 & NA \\ 0 & 1 \\ 1 & 0 \\ 1 & 1 \\ 2 & 0 \\ 2 & 1 \end{array}$

Author(s)

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

See Also

codeplot, zapvelout

Examples

outliers <- velout(age, height, id, heights, limit=3)</pre>

who06

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

who0607

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

http://www.who.int/childgrowth/en/

References

World Health Organization. WHO Child Growth Standards: Methods and development: Length/heightfor-age, weight-for-age, weight-for-length, weight-for-height and body mass index-for-age. Geneva: WHO; 2006.

Cole TJ, Green PJ. Smoothing reference centile curves: the LMS method and penalized likelihood. Stat Med 1992;11:1305-19.

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')
```

who	0607
-----	------

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

who0607

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).

Source

http://www.who.int/childgrowth/en/

References

Cole TJ, Green PJ. Smoothing reference centile curves: the LMS method and penalized likelihood. Stat Med 1992;11:1305-19.

World Health Organization. WHO Child Growth Standards: Methods and development: Length/heightfor-age, weight-for-age, weight-for-length, weight-for-height and body mass index-for-age. Geneva: WHO; 2006.

de Onis M, Onyango AW, Borghi E, Siyam A, Nishida C, Siekmann J. Development of a WHO growth reference for school-aged children and adolescents. Bull WHO 2007;85:660-7.

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)
```

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xaxsd

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

```
## 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())</pre>
```

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

xyadj

Description

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

Usage

xyadj(object, x, y = NULL, 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.
У	a vector of y coordinates (default NULL).
id	a factor denoting the subject levels corresponding to x and y.
abc	a data frame containing random effects for a, b and c (default ranef(object)[id,]).
tomean	a logical defining the direction of adjustment. TRUE (default) indicates that in- dividual curves are translated and rotated to match the mean curve, while FALSE indicates the reverse, the mean curve being translated and rotated to match indi- vidual curves.

Details

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

(x-xoffset-b < fixed > -b < random >) * exp(c < random >) + xoffset + b < fixed > fi

y - a < random >

When tomean = FALSE they are adjusted to

$$(x-xoffset-b < fixed >)/exp(c < random >) + xoffset + b < fixed > +b < random >) + xoffset + b < random > +b < r$$

y + a < random >

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

Value

The list of adjusted values:

х	numeric vector.
У	numeric vector the same length as x, or NULL.

Author(s)

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

z2cent

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))</pre>
```

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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