

# Package ‘cNORM’

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**Title** Continuous Norming

**Version** 3.5.3

**Description** A comprehensive toolkit for generating continuous test norms in psychometrics and biometrics, and analyzing model fit. The package offers both distribution-free modeling using Taylor polynomials and parametric modeling using the beta-binomial and the 'Sinh-Arcsinh' distribution. Originally developed for achievement tests, it is applicable to a wide range of mental, physical, or other test scores dependent on continuous or discrete explanatory variables. The package provides several advantages: It minimizes deviations from representativeness in subsamples, interpolates between discrete levels of explanatory variables, and significantly reduces the required sample size compared to conventional norming per age group. cNORM enables graphical and analytical evaluation of model fit, accommodates a wide range of scales including those with negative and descending values, and even supports conventional norming. It generates norm tables including confidence intervals. It also includes methods for addressing representativeness issues through Iterative Proportional Fitting. Based on Lenhard et al. (2016)  
<doi:10.1177/1073191116656437>, Lenhard et al. (2019)  
<doi:10.1371/journal.pone.0222279>, Lenhard and Lenhard (2021)  
<doi:10.1177/0013164420928457> and Gary et al. (2023)  
<doi:10.1007/s00181-023-02456-0>.

**License** AGPL-3

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bestModel	<i>Determine Regression Model</i>
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---

### Description

Computes Taylor polynomial regression models by evaluating a series of models with increasing predictors. It aims to find a consistent model that effectively captures the variance in the data. It draws on the `regsubsets` function from the `leaps` package and builds up to 20 models for each number of predictors, evaluates these models regarding model consistency and selects consistent model with the highest  $R^2$ . This automatic model selection should usually be accompanied with visual inspection of the percentile plots and assessment of fit statistics. Set  $R^2$  or number of terms manually to retrieve a more parsimonious model, if desired.

### Usage

```
bestModel(
  data,
  raw = NULL,
  R2 = NULL,
  k = NULL,
  t = NULL,
  predictors = NULL,
  terms = 0,
  weights = NULL,
  force.in = NULL,
  plot = TRUE,
  extensive = TRUE,
  subsampling = TRUE
)
```

### Arguments

<code>data</code>	Preprocessed dataset with 'raw' scores, powers, interactions, and usually an explanatory variable (like age).
<code>raw</code>	Name of the raw score variable (default: 'raw').
<code>R2</code>	Adjusted $R^2$ stopping criterion for model building.
<code>k</code>	Power constant influencing model complexity (default: 4, max: 6).
<code>t</code>	Age power parameter. If unset, defaults to 'k'.
<code>predictors</code>	List of predictors or regression formula for model selection. Overrides 'k' and can include additional variables.
<code>terms</code>	Desired number of terms in the model.
<code>weights</code>	Optional case weights. If set to <code>FALSE</code> , default weights (if any) are ignored.
<code>force.in</code>	Variables forcibly included in the regression.

plot	If TRUE (default), displays a percentile plot of the model and information about the regression object. FALSE turns off plotting and report.
extensive	If TRUE (default), screen models for consistency and - if possible, exclude inconsistent ones
subsampling	If TRUE (default), model coefficients are calculated using 10-folds and averaged across the folds. This produces more robust estimates with a slight increase in bias.

### Details

The functions `rankBySlidingWindow`, `rankByGroup`, `bestModel`, `computePowers` and `prepareData` are usually not called directly, but accessed through other functions like `cnorm`.

Additional functions like `plotSubset(model)` and `cnorm.cv` can aid in model evaluation.

### Value

The model. Further exploration can be done using `plotSubset(model)` and `plotPercentiles(data, model)`.

### See Also

`plotSubset`, `plotPercentiles`, `plotPercentileSeries`, `checkConsistency`

Other model: `checkConsistency()`, `cnorm.cv()`, `derive()`, `modelSummary()`, `print.cnorm()`, `printSubset()`, `rangeCheck()`, `regressionFunction()`, `summary.cnorm()`

### Examples

```
# Example with sample data
## Not run:
# It is not recommende to use this function. Rather use 'cnorm' instead.
normData <- prepareData(elfe)
model <- bestModel(normData)
plotSubset(model)
plotPercentiles(buildCnormObject(normData, model))

# Specifying variables explicitly
preselectedModel <- bestModel(normData, predictors = c("L1", "L3", "L1A3", "A2", "A3"))
print(regressionFunction(preselectedModel))

## End(Not run)
```

---

betaCoefficients

*Compute Parameters of a Beta Binomial Distribution*

---

### Description

This function calculates the  $\alpha$  (a) and  $\beta$  (b) parameters of a beta binomial distribution, along with the mean (m), variance (var) based on the input vector 'x' and the maximum number 'n'.

**Usage**

```
betaCoefficients(x, n = NULL)
```

**Arguments**

**x** A numeric vector of non-negative integers representing observed counts.

**n** The maximum number or the maximum possible value of 'x'. If not specified, uses `max(x)` instead.

**Details**

The beta-binomial distribution is a discrete probability distribution that models the number of successes in a fixed number of trials, where the probability of success varies from trial to trial. This variability in success probability is modeled by a beta distribution. Such a calculation is particularly relevant in scenarios where there is heterogeneity in success probabilities across trials, which is common in real-world situations, as for example the number of correct solutions in a psychometric test, where the test has a fixed number of items.

**Value**

A numeric vector containing the calculated parameters in the following order: alpha (a), beta (b), mean (m), standard deviation (sd), and the maximum number (n).

---

<code>buildCnormObject</code>	<i>Build cnorm object from data and bestModel model object</i>
-------------------------------	--

---

**Description**

Helper function to build a cnorm object from a data object and a model object from the `bestModel` function for compatibility reasons.

**Usage**

```
buildCnormObject(data, model)
```

**Arguments**

**data** A data object from 'prepareData', or from 'rankByGroup' and 'computePower'

**model** Object obtained from the `bestModel` function

**Value**

A cnorm object

**Examples**

```
## Not run:
data <- prepareData(elfe)
model <- bestModel(data, k = 4)
model.cnorm <- buildCnormObject(data, model)

## End(Not run)
```

---

buildFunction	<i>Build regression function for bestModel</i>
---------------	--

---

**Description**

Build regression function for bestModel

**Usage**

```
buildFunction(raw, k, t, age)
```

**Arguments**

raw	name of the raw score variable
k	the power degree for location
t	the power degree for age
age	use age

**Value**

regression function

---

calcPolyInL	<i>Internal function for retrieving regression function coefficients at specific age</i>
-------------	--

---

**Description**

The function is an inline for searching zeros in the inverse regression function. It collapses the regression function at a specific age and simplifies the coefficients.

**Usage**

```
calcPolyInL(raw, age, model)
```

**Arguments**

raw	The raw value (subtracted from the intercept)
age	The age
model	The cNORM regression model

**Value**

The coefficients

---

calcPolyInLBase2	<i>Internal function for retrieving regression function coefficients at specific age</i>
------------------	--

---

**Description**

The function is an inline for searching zeros in the inverse regression function. It collapses the regression function at a specific age and simplifies the coefficients. Optimized version of the prior 'calcPolyInLBase'

**Usage**

```
calcPolyInLBase2(raw, age, coeff, k)
```

**Arguments**

raw	The raw value (subtracted from the intercept)
age	The age
coeff	The cNORM regression model coefficients
k	The cNORM regression model power parameter

**Value**

The coefficients

---

CDC

*BMI growth curves from age 2 to 25*

---

## Description

By the courtesy of the Center of Disease Control (CDC), cNORM includes human growth data for children and adolescents age 2 to 25 that can be used to model trajectories of the body mass index and to estimate percentiles for clinical definitions of under- and overweight. The data stems from the NHANES surveys in the US and was published in 2012 as public domain. The data was cleaned by removing missing values and it includes the following variables from or based on the original dataset.

## Usage

CDC

## Format

A data frame with 45053 rows and 7 variables:

**age** continuous age in years, based on the month variable

**group** age group; chronological age in years at the time of examination

**month** chronological age in month at the time of examination

**sex** sex of the participant, 1 = male, 2 = female

**height** height of the participants in cm

**weight** weight of the participants in kg

**bmi** the body mass index, computed by  $(\text{weight in kg})/(\text{height in m})^2$

A data frame with 45035 rows and 7 columns

## References

Center for Disease Control and Prevention (2012). National Health and Nutrition Examination Survey: Questionnaires, datasets and related documentation. U.S. Department of Health and Human Services (original source not available anymore).

---

checkConsistency      *Check the consistency of the norm data model*

---

### Description

While abilities increase and decline over age, within one age group, the norm scores always have to show a monotonic increase or decrease with increasing raw scores. Violations of this assumption are an indication for problems in modeling the relationship between raw and norm scores. There are several reasons, why this might occur:

1. Vertical extrapolation: Choosing extreme norm scores, e. g. values  $-3 \leq x$  and  $x \geq 3$  In order to model these extreme values, a large sample dataset is necessary.
2. Horizontal extrapolation: Taylor polynomials converge in a certain radius. Using the model values outside the original dataset may lead to inconsistent results.
3. The data cannot be modeled with Taylor polynomials, or you need another power parameter (k) or R2 for the model.

### Usage

```
checkConsistency(
  model,
  minAge = NULL,
  maxAge = NULL,
  minNorm = NULL,
  maxNorm = NULL,
  minRaw = NULL,
  maxRaw = NULL,
  stepAge = NULL,
  stepNorm = 1,
  warn = FALSE,
  silent = FALSE
)
```

### Arguments

model	The model from the bestModel function or a cnorm object
minAge	Age to start with checking
maxAge	Upper end of the age check
minNorm	Lower end of the norm value range
maxNorm	Upper end of the norm value range
minRaw	clipping parameter for the lower bound of raw scores
maxRaw	clipping parameter for the upper bound of raw scores
stepAge	Stepping parameter for the age check. values indicate higher precision / closer checks

stepNorm	Stepping parameter for the norm table check within age with lower scores indicating a higher precision. The choice depends of the norm scale used. With T scores a stepping parameter of 1 is suitable
warn	If set to TRUE, already minor violations of the model assumptions are displayed (default = FALSE)
silent	turn off messages

### Details

In general, extrapolation (point 1 and 2) can carefully be done to a certain degree outside the original sample, but it should in general be handled with caution. Please note that at extreme values, the models most likely become independent and it is thus recommended to restrict the norm score range to the relevant range of abilities, e.g. +/- 2.5 SD via the minNorm and maxNorm parameter.

### Value

Boolean, indicating model violations (TRUE) or no problems (FALSE)

### See Also

Other model: [bestModel\(\)](#), [cnorm.cv\(\)](#), [derive\(\)](#), [modelSummary\(\)](#), [print.cnorm\(\)](#), [printSubset\(\)](#), [rangeCheck\(\)](#), [regressionFunction\(\)](#), [summary.cnorm\(\)](#)

### Examples

```
## Not run:
model <- cnorm(raw = elfe$raw, group = elfe$group, plot = FALSE)
modelViolations <- checkConsistency(model, minNorm = 25, maxNorm = 75)
plotDerivative(model, minNorm = 25, maxNorm = 75)

## End(Not run)
```

---

checkWeights	<i>Check, if NA or values &lt;= 0 occur and issue warning</i>
--------------	---

---

### Description

Check, if NA or values <= 0 occur and issue warning

### Usage

```
checkWeights(weights)
```

### Arguments

weights	Raking weights
---------	----------------

---

check_monotonicity	<i>Check Monotonicity of Predicted Values</i>
--------------------	---

---

### Description

This function checks if the predicted values from a linear model are monotonically increasing or decreasing across a range of L values for multiple age points.

### Usage

```
check_monotonicity(lm_model, pred_data, minRaw, maxRaw)
```

### Arguments

lm_model	An object of class 'lm' representing the fitted linear model.
pred_data	Matrix with prediction values
minRaw	lowest raw score in prediction
maxRaw	highest raw score in prediction

### Details

The function creates a prediction data frame using all combinations of the provided L values and age points. It then generates predictions using the provided linear model and checks if these predictions are monotonically increasing or decreasing for each age point across the range of L values.

### Value

A named character vector where each element corresponds to an age point. Possible values for each element are 1 for "Monotonically increasing" -1 for "Monotonically decreasing", or 0 for "Not monotonic".

---

cnorm	<i>Continuous Norming</i>
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---

### Description

Conducts continuous norming in one step and returns an object including ranked raw data and the continuous norming model. Please consult the function description of 'rankByGroup', 'rankBySlidingWindow' and 'bestModel' for specifics of the steps in the data preparation and modeling process. In addition to the raw scores, either provide

- a numeric vector for the grouping information (group)
- a numeric age vector and the width of the sliding window (age, width)

for the ranking of the raw scores. You can adjust the grade of smoothing of the regression model by setting the `k` and `terms` parameter. In general, increasing `k` to more than 4 and the number of terms lead to a higher fit, while lower values lead to more smoothing. The power parameter for the age trajectory can be specified independently by `t`. If both parameters are missing, `cnorm` uses `k = 5` and `t = 3` by default.

### Usage

```
cnorm(
  raw = NULL,
  group = NULL,
  age = NULL,
  width = NA,
  weights = NULL,
  scale = "T",
  method = 4,
  descend = FALSE,
  k = NULL,
  t = NULL,
  terms = 0,
  R2 = NULL,
  plot = TRUE,
  extensive = TRUE,
  subsampling = TRUE
)
```

### Arguments

<code>raw</code>	Numeric vector of raw scores
<code>group</code>	Numeric vector of grouping variable, e.g. grade. If no group or age variable is provided, conventional norming is applied.
<code>age</code>	Numeric vector with chronological age. If used without <code>'group'</code> , please additionally specify <code>'width'</code> .
<code>width</code>	Size of the sliding window in case an age vector is used.
<code>weights</code>	Optional numeric vector of case weights for post-stratification.
<code>scale</code>	Type of norm scale, either "T" (default), "IQ", "z" or "percentile" (= no transformation); a numeric vector with mean and SD can also be provided, e.g. <code>'c(10, 3)'</code> for Wechsler scale index points.
<code>method</code>	Ranking method in case of ties; an integer index from 1 (Blom 1958) through 7 (Yu & Huang 2001). Default is 4 (Rankit).
<code>descend</code>	If TRUE, inverts the ranking order so that higher raw scores receive lower norm scores (e.g. for error scores).
<code>k</code>	Power degree for the location dimension (max 6).
<code>t</code>	Power degree for the age dimension (max 6).
<code>terms</code>	If > 0, fix the model to this number of terms.
<code>R2</code>	Stopping criterion (adjusted R-squared) for model selection.

plot	If TRUE (default), display percentile plot and report.
extensive	If TRUE (default), screen models for monotonic consistency.
subsampling	If TRUE (default), use 10-fold subsampled coefficient averaging in 'bestModel'.

### Value

cnorm object including the ranked raw data and the regression model.

### References

1. Gary, S. & Lenhard, W. (2021). In norming we trust. *Diagnostica*.
2. Gary, S., Lenhard, W. & Lenhard, A. (2021). Modelling Norm Scores with the cNORM Package in R. *Psych*, 3(3), 501-521. <https://doi.org/10.3390/psych3030033>
3. Gary, S., Lenhard, W., Lenhard, A., & Herzberg, D. (2023). A tutorial on automatic post-stratification and weighting in conventional and regression-based norming of psychometric tests. *Behavior Research Methods*. <https://doi.org/10.3758/s13428-023-02207-0>
4. Gary, S., Lenhard, A., Lenhard, W., & Herzberg, D. S. (2023). Reducing the Bias of Norm Scores in Non-Representative Samples: Weighting as an Adjunct to Continuous Norming Methods. *Assessment*, 30(8), 2491–2509. <https://doi.org/10.1177/10731911231153832>
5. Lenhard, A., Lenhard, W., Suggate, S. & Segerer, R. (2016). A continuous solution to the norming problem. *Assessment*, Online first, 1-14. doi:10.1177/1073191116656437
6. Lenhard, A., Lenhard, W., Gary, S. (2018). Continuous Norming (cNORM). The Comprehensive R Network, Package cNORM, available: <https://CRAN.R-project.org/package=cNORM>
7. Lenhard, A., Lenhard, W., Gary, S. (2019). Continuous norming of psychometric tests: A simulation study of parametric and semi-parametric approaches. *PLoS ONE*, 14(9), e0222279. doi:10.1371/journal.pone.0222279
8. Lenhard, W., & Lenhard, A. (2020). Improvement of Norm Score Quality via Regression-Based Continuous Norming. *Educational and Psychological Measurement(Online First)*, 1-33. <https://doi.org/10.1177/0013164420928457>

### See Also

rankByGroup, rankBySlidingWindow, computePowers, bestModel

### Examples

```
## Not run:
# Conventional norming
cnorm(raw = elfe$raw)

# Continuous norming via group
m1 <- cnorm(raw = elfe$raw, group = elfe$group)

# Continuous norming via continuous age + sliding window
m2 <- cnorm(raw = ppvt$raw, age = ppvt$age, width = 1)

# Norm tables with confidence intervals
normTable(c(2.125, 2.375, 2.625), m1, CI = .9, reliability = .95)
```

```
rawTable (c(2.125, 2.375, 2.625), m1, CI = .9, reliability = .95)

## End(Not run)
```

---

cnorm.betabinomial      *Fit a beta-binomial regression model for continuous norming*

---

### Description

This function fits a beta-binomial regression model where both the  $\alpha$  and  $\beta$  parameters of the beta-binomial distribution are modeled as polynomial functions of the predictor variable (typically age). Setting mode to 1 fits a beta-binomial model on the basis of  $\mu$  and  $\sigma$ , setting it to 2 (default) fits a beta-binomial model directly on the basis of  $\alpha$  and  $\beta$ .

### Usage

```
cnorm.betabinomial(
  age,
  score,
  n = NULL,
  weights = NULL,
  mode = 2,
  alpha = 3,
  beta = 3,
  control = NULL,
  scale = "T",
  plot = T
)
```

### Arguments

age	A numeric vector of predictor values (e.g., age).
score	A numeric vector of response values.
n	The maximum score (number of trials in the beta-binomial distribution). If NULL, max(score) is used.
weights	A numeric vector of weights for each observation. Default is NULL (equal weights).
mode	Integer specifying the mode of the model. Default is 2 (direct modelling of $\gamma$ and $\beta$ ). If set to 1, the model is fitted on the basis of $\mu$ and $\sigma$ , the predicted mean and standard deviation over age.
alpha	Integer specifying the degree of the polynomial for the alpha model. Default is 3. If mode is set to 1, this parameter is used to specify the degree of the polynomial for the $\mu$ model.
beta	Integer specifying the degree of the polynomial for the beta model. Default is 3. If mode is set to 1, this parameter is used to specify the degree of the polynomial for the $\sigma$ model.

control	A list of control parameters to be passed to the 'optim' function. If NULL, default values are used, namely control = list(reltol = 1e-8, maxit = 1000) for mode 1 and control = list(factr = 1e-8, maxit = 1000) for mode 2.
scale	Type of norm scale, either "T" (default), "IQ", "z" or a double vector with the mean and standard deviation.
plot	Logical indicating whether to plot the model. Default is TRUE.

### Details

The function standardizes the input variables, fits polynomial models for both the alpha and beta parameters, and uses maximum likelihood estimation to find the optimal parameters. The optimization is performed using the L-BFGS-B method.

### Value

A list of class "cnormBetaBinomial" or "cnormBetaBinomial2". In case of mode 2 containing:

alpha_est	Estimated coefficients for the alpha model
beta_est	Estimated coefficients for the beta model
se	Standard errors of the estimated coefficients
alpha_degree	Degree of the polynomial for the alpha model
beta_degree	Degree of the polynomial for the beta model
result	Full result from the optimization procedure

### Examples

```
## Not run:
# Fit a beta-binomial regression model to the PPVT data
model <- cnorm.betabinomial(ppvt$age, ppvt$raw, n = 228)
summary(model)

# Use weights for post-stratification
marginals <- data.frame(var = c("sex", "sex", "migration", "migration"),
                       level = c(1,2,0,1),
                       prop = c(0.51, 0.49, 0.65, 0.35))
weights <- computeWeights(ppvt, marginals)
model <- cnorm.betabinomial(ppvt$age, ppvt$raw, n = 228, weights = weights)

## End(Not run)
```

**Description**

Assists in determining the optimal number of terms for the regression model using repeated Monte Carlo cross-validation. It leverages an 80-20 split between training and validation data, with stratification by norm group or random sample in case of using sliding window ranking.

**Usage**

```
cnorm.cv(
  data,
  formula = NULL,
  repetitions = 5,
  norms = TRUE,
  min = 1,
  max = 12,
  cv = "full",
  pCutoff = NULL,
  width = NA,
  raw = NULL,
  group = NULL,
  age = NULL,
  weights = NULL
)
```

**Arguments**

data	Data frame of norm sample or a cnorm object. Should have ranking, powers, and interaction of L and A.
formula	Formula from an existing regression model; min/max functions ignored. If using a cnorm object, this is automatically fetched.
repetitions	Number of repetitions for cross-validation.
norms	If TRUE, computes norm score crossfit and R <sup>2</sup> . Note: Computationally intensive.
min	Start with a minimum number of terms (default = 1).
max	Maximum terms in model, up to $(k + 1) * (t + 1) - 1$ .
cv	"full" (default) splits data into training/validation, then ranks. Otherwise, expects a pre-ranked dataset.
pCutoff	Checks stratification for unbalanced data. Performs a t-test per group. Default set to 0.2 to minimize beta error.
width	If provided, ranking done via 'rankBySlidingWindow'. Otherwise, by group.
raw	Name of the raw score variable.

group	Name of the grouping variable.
age	Name of the age variable.
weights	Name of the weighting parameter.

## Details

Successive models, with an increasing number of terms, are evaluated, and the RMSE for raw scores plotted. This encompasses the training, validation, and entire dataset. If 'norms' is set to TRUE (default), the function will also calculate the mean norm score reliability and crossfit measures. Note that due to the computational requirements of norm score calculations, execution can be slow, especially with numerous repetitions or terms.

When 'cv' is set to "full" (default), both test and validation datasets are ranked separately, providing comprehensive cross-validation. For a more streamlined validation process focused only on modeling, a pre-ranked dataset can be used. The output comprises RMSE for raw score models, norm score  $R^2$ , delta  $R^2$ , crossfit, and the norm score SE according to Oosterhuis, van der Ark, & Sijtsma (2016).

This function is not yet prepared for the 'extensive' search strategy, introduced in version 3.3, but instead relies on the first model per number of terms, without consistency check.

For assessing overfitting:

$$CROSSFIT = R(Training; Model)^2 / R(Validation; Model)^2$$

A CROSSFIT > 1 suggests overfitting, < 1 suggests potential underfitting, and values around 1 are optimal, given a low raw score RMSE and high norm score validation  $R^2$ .

Suggestions for ideal model selection:

- Visual inspection of percentiles with 'plotPercentiles' or 'plotPercentileSeries'.
- Pair visual inspection with repeated cross-validation (e.g., 10 repetitions).
- Aim for low raw score RMSE and high norm score  $R^2$ , avoiding terms with significant overfit (e.g., crossfit > 1.1).

## Value

Table with results per term number: RMSE for raw scores,  $R^2$  for norm scores, and crossfit measure.

## References

Oosterhuis, H. E. M., van der Ark, L. A., & Sijtsma, K. (2016). Sample Size Requirements for Traditional and Regression-Based Norms. *Assessment*, 23(2), 191–202. <https://doi.org/10.1177/1073191115580638>

## See Also

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [derive\(\)](#), [modelSummary\(\)](#), [print.cnorm\(\)](#), [printSubset\(\)](#), [rangeCheck\(\)](#), [regressionFunction\(\)](#), [summary.cnorm\(\)](#)

**Examples**

```
## Not run:
# Example: Plot cross-validation RMSE by number of terms (up to 9) with three repetitions.
result <- cnorm(raw = elfe$raw, group = elfe$group)
cnorm.cv(result$data, min = 2, max = 9, repetitions = 3)

# Using a cnorm object examines the predefined formula.
cnorm.cv(result, repetitions = 1)

## End(Not run)
```

---

cNORM.GUI	<i>Launcher for the graphical user interface of cNORM for distribution free continuous norming</i>
-----------	--

---

**Description**

Launcher for the graphical user interface of cNORM for distribution free continuous norming

**Usage**

```
cNORM.GUI(launch.browser = TRUE)
```

**Arguments**

launch.browser Default TRUE; automatically open browser for GUI

**Examples**

```
## Not run:
# Launch graphical user interface
cNORM.GUI()

## End(Not run)
```

---

cNORM.GUI2	<i>Launch the cNORM Parametric Modeling Shiny Application</i>
------------	---

---

**Description**

Launch the cNORM Parametric Modeling Shiny Application

**Usage**

```
cNORM.GUI2(launch.browser = TRUE)
```

**Arguments**

launch.browser Logical, whether to launch browser automatically (default: TRUE)

---

cnorm.shash *Fit a Sinh-Arcsinh (shash) Regression Model for Continuous Norming*

---

**Description**

This function fits a Sinh-Arcsinh (shash; Jones & Pewsey, 2009) regression model for continuous norm score modeling, where the distribution parameters vary smoothly as polynomial functions of age or other predictors. The shash distribution is well-suited for psychometric data as it can flexibly model skewness and tail weight independently, making it ideal for handling floor effects, ceiling effects, and varying degrees of individual differences across age groups. In a simulation study (Lenhard et al, 2019), the shash model demonstrated superior performance compared to other parametric approaches from the Box Cox family of functions. In contrast to Box Cox, Sinh-Arcsinh can model distributions including zero and negativ values.

**Usage**

```
cnorm.shash(
  age,
  score,
  weights = NULL,
  mu_degree = 3,
  sigma_degree = 2,
  epsilon_degree = 2,
  delta_degree = 1,
  delta = 1,
  control = NULL,
  scale = "T",
  plot = TRUE
)
```

**Arguments**

age	A numeric vector of predictor values (typically age, but can be any continuous predictor).
score	A numeric vector of response values (raw test scores). Must be the same length as age. The value range is unrestricted and it can include zeros and negative values.
weights	An optional numeric vector of weights for each observation. Useful for incorporating sampling weights. If NULL (default), all observations are weighted equally.
mu_degree	Integer specifying the degree of the polynomial for modeling the location parameter $\mu(\text{age})$ . Default is 3. Higher degrees allow more flexible modeling of how the central tendency changes with age, but may lead to overfitting with small samples. Common choices:

	<ul style="list-style-type: none"> <li>• 1: Linear change with age</li> <li>• 2: Quadratic change (allows one inflection point)</li> <li>• 3: Cubic change (allows two inflection points, suitable for most developmental curves)</li> <li>• 4+: Higher-order changes (use cautiously, mainly for large samples)</li> </ul>
sigma_degree	Integer specifying the degree of the polynomial for modeling the scale parameter $\sigma(\text{age})$ . Default is 2. This controls how the variability (spread) of scores changes with age. Lower degrees are often sufficient as variability typically changes more smoothly than location.
epsilon_degree	Integer specifying the degree of the polynomial for modeling the skewness parameter $\epsilon(\text{age})$ . Default is 2. This controls how the asymmetry of the distribution changes with age.
delta_degree	Integer specifying the polynomial for modelling the tail weight parameter $\delta(\text{age})$ . Default is 1. The tail weight can be fixed as well in case of numerical instability. In that case, set 'delta_degree' to NULL and specify a value for delta instead. Recommendation: Keep delta_degree low to avoid overfitting.
delta	Fixed tail weight parameter (must be $> 0$ ). Default is 1. This parameter controls the heaviness of the distribution tails and is kept constant across all ages in this implementation. It is only used, if 'delta_degree' is set to NULL. Common values: <ul style="list-style-type: none"> <li>• <math>\delta = 1</math>: Normal-like tail behavior (baseline)</li> <li>• <math>\delta &gt; 1</math>: Heavier tails, higher kurtosis (more extreme scores than normal distribution)</li> <li>• <math>\delta &lt; 1</math>: Lighter tails, lower kurtosis (fewer extreme scores than normal distribution)</li> </ul>
control	An optional list of control parameters passed to the <code>optim</code> function for maximum likelihood estimation. If NULL, sensible defaults are chosen automatically based on the model complexity. Common parameters to adjust: <ul style="list-style-type: none"> <li>• <code>factr</code>: Controls precision of optimization (default: <math>1e-8</math>)</li> <li>• <code>maxit</code>: Maximum number of iterations (default: <math>n\_parameters * 200</math>)</li> <li>• <code>lmm</code>: Memory limit for L-BFGS-B (default: <math>\min(n\_parameters, 20)</math>)</li> </ul> <p>Increase <code>maxit</code> or decrease <code>factr</code> if optimization fails to converge.</p>
scale	Character string or numeric vector specifying the type of norm scale for output. This affects the scaling of derived norm scores but does not influence model fitting: <ul style="list-style-type: none"> <li>• "T": T-scores (mean = 50, SD = 10) - default</li> <li>• "IQ": IQ-like scores (mean = 100, SD = 15)</li> <li>• "z": z-scores (mean = 0, SD = 1)</li> <li>• <code>c(M, SD)</code>: Custom scale with specified mean M and standard deviation SD</li> </ul>
plot	Logical indicating whether to automatically display a diagnostic plot of the fitted model. Default is TRUE.

## Details

This implementation uses the Jones & Pewsey (2009) parameterization of the Sinh-Arcsinh distribution. Parameters are estimated using maximum likelihood via the L-BFGS-B algorithm. In case, optimization fails, try reducing model complexity by reducing polynomial degrees or fixing the delta parameter.

**The Sinh-Arcsinh Distribution:** The shash distribution is defined by the transformation:

$$X = \mu + \sigma \cdot \sinh\left(\frac{\operatorname{arcsinh}(Y) - \epsilon}{\delta}\right)$$

where Y is a standard normal variable,  $Y \sim N(0,1)$ .

This transformation provides:

- mu: Location parameter (similar to mean)
- sigma: Scale parameter (similar to standard deviation)
- epsilon: Skewness parameter (epsilon = 0 for symmetry)
- delta: Tail weight parameter (delta = 1 for normal-like tails)

**Model Selection:** Choose polynomial degrees based on:

- Sample size (higher degrees need more data)
- Theoretical expectations about developmental trajectories
- Model comparison criteria (AIC, BIC)
- Visual inspection of fitted curves

For most applications, mu\_degree = 3, sigma\_degree = 2, epsilon\_degree = 2, delta\_degree = 1 provides a good balance of flexibility and parsimony.

## Value

An object of class "cnormShash" containing the fitted model results. This is a list with components:

mu_est	Numeric vector of estimated coefficients for the location parameter mu(age). The first coefficient is the intercept, subsequent coefficients correspond to polynomial terms.
sigma_est	Numeric vector of estimated coefficients for the scale parameter log(sigma(age)). Note: These are coefficients for log(sigma) to ensure sigma > 0.
epsilon_est	Numeric vector of estimated coefficients for the skewness parameter epsilon(age).
delta	The fixed tail weight parameter value used in fitting.
delta_est	Numeric vector of estimated coefficients for the tail weight parameter delta(age) - in case, a degree has been set.
se	Numeric vector of standard errors for all estimated coefficients (if Hessian computation succeeds).
mu_degree, sigma_degree, epsilon_degree	The polynomial degrees used for each parameter.
result	Complete output from the optim function, including convergence information, log-likelihood value, and other optimization details.

**Note**

- The function requires the input data to have sufficient variability. Very small datasets or datasets with little age spread may cause convergence problems.
- Polynomial models can exhibit edge effects at the boundaries of the age range. Predictions outside the observed age range should be made cautiously.
- If convergence fails, try: (1) reducing polynomial degrees, (2) adjusting the delta parameter, (3) providing custom control parameters, or (4) checking for data quality issues.
- The tail weight parameter delta is fixed across ages by default. For applications where tail behavior changes substantially with age, consider setting the delta\_degree parameter to 1 or 2.

**Author(s)**

Wolfgang Lenhard

**References**

Jones, M. C., & Pewsey, A. (2009). Sinh-arcsinh distributions. *Biometrika*, 96(4), 761-780.

Lenhard, A., Lenhard, W., Gary, S. (2019). Continuous norming of psychometric tests: A simulation study of parametric and semi-parametric approaches. *PLoS ONE*, 14(9), e0222279. <https://doi.org/10.1371/journal.pone.0222279>

**See Also**

[plot](#) for plotting fitted models, [predict](#) for generating predictions, [cnorm.betabinomial2](#) for discrete beta-binomial alternative

**Examples**

```
## Not run:
# Basic usage with default settings
model <- cnorm.shash(age = children$age, score = children$raw_score)

# Custom polynomial degrees for complex developmental pattern
model_complex <- cnorm.shash(
  age = adolescents$age,
  score = adolescents$vocabulary_score,
  mu_degree = 4,          # Complex mean trajectory
  sigma_degree = 3,      # Changing variability pattern
  epsilon_degree = 2,    # Skewness shifts
  delta_degree = NULL,   # set to NULL to activate fixed delta
  delta = 1.3            # Slightly heavy tails
)

# With sampling weights
model_weighted <- cnorm.shash(
  age = survey$age,
  score = survey$score,
  weights = survey$sample_weight
```

```

)

# Custom optimization control for difficult convergence
model_robust <- cnorm.shash(
  age = mixed$age,
  score = mixed$score,
  control = list(factr = 1e-6, maxit = 2000),
  delta = 1.5
)

# Compare model fit
compare(model, model_complex)

## End(Not run)

```

---

compare

*Compare Two Norm Models Visually*


---

## Description

This function creates a visualization comparing two norm models by displaying their percentile curves. The first model is shown with solid lines, the second with dashed lines. If age and score vectors are provided, manifest percentiles are displayed as dots. The function works with regular cnorm models, beta-binomial models, and shash models, allowing comparison between different model types.

## Usage

```

compare(
  model1,
  model2,
  percentiles = c(0.025, 0.1, 0.25, 0.5, 0.75, 0.9, 0.975),
  age = NULL,
  score = NULL,
  weights = NULL,
  title = NULL,
  subtitle = NULL
)

```

## Arguments

model1	First model object (distribution free, beta-binomial, or shash)
model2	Second model object (distribution free, beta-binomial, or shash)
percentiles	Vector with percentile scores, ranging from 0 to 1 (exclusive)
age	Optional vector with manifest age or group values
score	Optional vector with manifest raw score values

weights	Optional vector with manifest weights
title	Custom title for plot (optional)
subtitle	Custom subtitle for plot (optional)

**Value**

A ggplot object showing the comparison of both models

**See Also**

Other plot: [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

**Examples**

```
## Not run:
# Compare different types of models
model1 <- cnorm(group = elfe$group, raw = elfe$raw)
model2 <- cnorm.betabinomial(elfe$group, elfe$raw)
model3 <- cnorm.shash(elfe$group, elfe$raw)

# Compare traditional cnorm with shash
compare(model1, model3, age = elfe$group, score = elfe$raw)

# Compare beta-binomial with shash
compare(model2, model3, age = elfe$group, score = elfe$raw)

## End(Not run)
```

---

computePowers	<i>Compute powers of the explanatory variable a as well as of the person location l (data preparation)</i>
---------------	--

---

**Description**

The function computes powers of the norm variable e. g. T scores (location, L), an explanatory variable, e. g. age or grade of a data frame (age, A) and the interactions of both (L X A). The k variable indicates the degree up to which powers and interactions are build. These predictors can be used later on in the [bestModel](#) function to model the norm sample. Higher values of k allow for modeling the norm sample closer, but might lead to over-fit. In general k = 3 or k = 4 (default) is sufficient to model human performance data. For example, k = 2 results in the variables L1, L2, A1, A2, and their interactions L1A1, L2A1, L1A2 and L2A2 (but k = 2 is usually not sufficient for the modeling). Please note, that you do not need to use a normal rank transformed scale like T or IQ; you can use the percentiles for the 'normValue' as well.

**Usage**

```
computePowers(data, k = 5, norm = NULL, age = NULL, t = 3, silent = FALSE)
```

**Arguments**

data	data.frame with the norm data
k	degree of the location polynomial (1..6)
norm	name of the norm variable in the data.frame (T scores, IQ, percentiles, ...). If 'NULL', the "normValue" attribute of 'data' is used.
age	explanatory variable (e.g. age or grade). May be a column name, a numeric vector of 'nrow(data)', 'FALSE' to disable age handling, or 'NULL' to fall back to the "age" attribute of 'data'.
t	age power parameter (1..6). If 'NULL', falls back to 'k'.
silent	set to TRUE to suppress messages

**Details**

The functions `rankBySlidingWindow`, `rankByGroup`, `bestModel`, `computePowers` and `prepareData` are usually not called directly, but accessed through other functions like `cnorm`.

**Value**

data.frame with the powers and interactions of location and explanatory variable / age

**See Also**

`bestModel`

Other prepare: [prepareData\(\)](#), [rankByGroup\(\)](#), [rankBySlidingWindow\(\)](#)

**Examples**

```
## Not run:  
# Dataset with grade levels as grouping  
data.elfe <- rankByGroup(elfe)  
data.elfe <- computePowers(data.elfe)  
  
# Dataset with continuous age variable and k = 5  
data.ppvt <- rankByGroup(ppvt)  
data.ppvt <- computePowers(data.ppvt, age = "age", k = 5)  
  
## End(Not run)
```

---

computeWeights	<i>Weighting of cases through iterative proportional fitting (Raking)</i>
----------------	---

---

### Description

Computes and standardizes weights via raking to compensate for non-stratified samples. It is based on the implementation in the survey R package. It reduces data collection #' biases in the norm data by the means of post stratification, thus reducing the effect of unbalanced data in percentile estimation and norm data modeling.

### Usage

```
computeWeights(data, population.margins, standardized = TRUE)
```

### Arguments

data	data.frame with norm sample data.
population.margins	A data.frame including three columns, specifying the variable name in the original dataset used for data stratification, the factor level of the variable and the according population share. Please ensure, the original data does not include factor levels, not present in the population.margins. Additionally, summing up the shares of the different levels of a variable should result in a value near 1.0. The first column must specify the name of the stratification variable, the second the level and the third the proportion
standardized	If TRUE (default), the raking weights are scaled to weights/min(weights)

### Details

This function computes standardized raking weights to overcome biases in norm samples. It generates weights, by drawing on the information of population shares (e. g. for sex, ethnic group, region ...) and subsequently reduces the influence of over-represented groups or increases under-represented cases. The returned weights are either raw or standardized and scaled to be larger than 0.

Raking in general has a number of advantages over post stratification and it additionally allows cNORM to draw on larger datasets, since less cases have to be removed during stratification. To use this function, additionally to the data, a data frame with stratification variables has to be specified. The data frame should include a row with (a) the variable name, (b) the level of the variable and (c) the according population proportion.

### Value

a vector with the standardized weights

**Examples**

```
## Not run:
# cNORM features a dataset on vocabulary development (ppvt)
# that includes variables like sex or migration. In order
# to weight the data, we have to specify the population shares.
# According to census, the population includes 52% boys
# (factor level 1 in the ppvt dataset) and 70% / 30% of persons
# without / with a a history of migration (= 0 / 1 in the dataset).
# First we set up the population margins with all shares of the
# different levels:

margins <- data.frame(variables = c("sex", "sex",
                                   "migration", "migration"),
                      levels = c(1, 2, 0, 1),
                      share = c(.52, .48, .7, .3))

head(margins)

# Now we use the population margins to generate weights
# through raking

weights <- computeWeights(ppvt, margins)

# There are as many different weights as combinations of
# factor levels, thus only four in this specific case

unique(weights)

# To include the weights in the cNORM modelling, we have
# to pass them as weights. They are then used to set up
# weighted quantiles and as weights in the regression.

model <- cnorm(raw = ppvt$raw,
               group=ppvt$group,
               weights = weights)

## End(Not run)
```

---

derivationTable	<i>Create a table based on first order derivative of the regression model for specific age</i>
-----------------	--

---

**Description**

In order to check model assumptions, a table of the first order derivative of the model coefficients is created.

**Usage**

```
derivationTable(A, model, minNorm = NULL, maxNorm = NULL, step = 0.1)
```

**Arguments**

A	the age
model	The regression model or a cnorm object
minNorm	The lower bound of the norm value range
maxNorm	The upper bound of the norm value range
step	Stepping parameter with lower values indicating higher precision

**Value**

data.frame with norm scores and the predicted scores based on the derived regression function

**See Also**

plotDerivative, derive

Other predict: [getNormCurve\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predict.cnormShash\(\)](#), [predictNorm\(\)](#), [predictRaw\(\)](#), [rawTable\(\)](#)

**Examples**

```
## Not run:
# Generate cnorm object from example data
cnorm.elfe <- cnorm(raw = elfe$raw, group = elfe$group)

# retrieve function for time point 6
d <- derivationTable(6, cnorm.elfe, step = 0.5)

## End(Not run)
```

---

derive	<i>Derivative of regression model</i>
--------	---------------------------------------

---

**Description**

Calculates the derivative of the location / norm value from the regression model with the first derivative as the default. This is useful for finding violations of model assumptions and problematic distribution features as f. e. bottom and ceiling effects, non-progressive norm scores within an age group or in general #' intersecting percentile curves.

**Usage**

```
derive(model, order = 1)
```

**Arguments**

model	The regression model or a cnorm object
order	The degree of the derivate, default: 1

**Value**

The derived coefficients

**See Also**

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [cnorm.cv\(\)](#), [modelSummary\(\)](#), [print.cnorm\(\)](#), [printSubset\(\)](#), [rangeCheck\(\)](#), [regressionFunction\(\)](#), [summary.cnorm\(\)](#)

**Examples**

```
## Not run:
m <- cnorm(raw = elfe$raw, group = elfe$group)
derivedCoefficients <- derive(m)

## End(Not run)
```

---

diagnostics.betabinomial

*Diagnostic Information for Beta-Binomial Model*

---

**Description**

This function provides diagnostic information for a fitted beta-binomial model from the `cnorm.betabinomial` function. It returns various metrics related to model convergence, fit, and complexity. In case, age and raw scores are provided, the function as well computes R2, rmse and bias for the norm scores based on the manifest and predicted norm scores.

**Usage**

```
diagnostics.betabinomial(model, age = NULL, score = NULL, weights = NULL)
```

**Arguments**

<code>model</code>	An object of class "cnormBetaBinomial", typically the result of a call to <code>cnorm.betabinomial()</code> .
<code>age</code>	An optional vector with age values
<code>score</code>	An optional vector with raw values
<code>weights</code>	An optional vector with weights

**Details**

The AIC and BIC are calculated as:  $AIC = 2k - 2\ln(L)$   $BIC = \ln(n)k - 2\ln(L)$  where  $k$  is the number of parameters,  $L$  is the maximum likelihood, and  $n$  is the number of observations.

**Value**

A list containing the following diagnostic information:

- `converged`: Logical indicating whether the optimization algorithm converged.
- `n_evaluations`: Number of function evaluations performed during optimization.
- `n_gradient`: Number of gradient evaluations performed during optimization.
- `final_value`: Final value of the objective function (negative log-likelihood).
- `message`: Any message returned by the optimization algorithm.
- `AIC`: Akaike Information Criterion.
- `BIC`: Bayesian Information Criterion.
- `max_gradient`: Maximum absolute gradient at the solution (if available).

**Examples**

```
## Not run:
# Fit a beta-binomial model
model <- cnorm.betabinomial(ppvt$sage, ppvt$raw)

# Get diagnostic information
diag_info <- diagnostics.betabinomial(model)

# Print the diagnostic information
print(diag_info)

# Summary the diagnostic information
summary(diag_info)

# Check if the model converged
if(diag_info$converged) {
  cat("Model converged successfully.\n")
} else {
  cat("Warning: Model did not converge.\n")
}

# Compare AIC and BIC
cat("AIC:", diag_info$AIC, "\n")
cat("BIC:", diag_info$BIC, "\n")

## End(Not run)
```

**Description**

A dataset containing the raw data of 1400 students from grade 2 to 5 in the sentence comprehension test from ELFE 1-6 (Lenhard & Schneider, 2006). In this test, students are presented lists of sentences with one gap. The student has to fill in the correct solution by selecting from a list of 5 alternatives per sentence. The alternatives include verbs, adjectives, nouns, pronouns and conjunctives. Each item stems from the same word type. The text is speeded, with a time cutoff of 180 seconds. The variables are as follows:

**Usage**

elfe

**Format**

A data frame with 1400 rows and 3 variables:

**personID** ID of the student

**group** grade level, with x.5 indicating the end of the school year and x.0 indicating the middle of the school year

**raw** the raw score of the student, spanning values from 0 to 28

A data frame with 1400 rows and 3 columns

**Source**

<https://www.psychometrica.de/elfe2.html>

**References**

Lenhard, W. & Schneider, W.(2006). Ein Leseverstaendnistest fuer Erst- bis Sechstklaesser. Goettingen/Germany: Hogrefe.

**Examples**

```
## Not run:  
# prepare data, retrieve model and plot percentiles  
model <- cnorm(elfe$group, elfe$raw)  
  
## End(Not run)
```

---

`getGroups`*Determine groups and group means*

---

**Description**

Helps to split the continuous explanatory variable into groups and assigns the group mean. The groups can be split either into groups of equal size (default) or equal number of observations.

**Usage**

```
getGroups(x, n = NULL, equidistant = FALSE)
```

**Arguments**

<code>x</code>	The continuous variable to be split
<code>n</code>	The number of groups; if NULL then the function determines a number of groups with usually 100 cases or $3 \leq n \leq 20$ .
<code>equidistant</code>	If set to TRUE, builds equidistant interval, otherwise (default) with equal number of observations

**Value**

vector with group means for each observation

**Examples**

```
## Not run:  
x <- rnorm(1000, m = 50, sd = 10)  
m <- getGroups(x, n = 10)  
  
## End(Not run)
```

---

`getNormCurve`*Computes the curve for a specific T value*

---

**Description**

As with this continuous norming regression approach, raw scores are modeled as a function of age and norm score (location), `getNormCurve` is a straightforward approach to show the raw score development over age, while keeping the norm value constant. This way, e. g. academic performance or intelligence development of a specific ability is shown.

**Usage**

```
getNormCurve(  
  norm,  
  model,  
  minAge = NULL,  
  maxAge = NULL,  
  step = 0.1,  
  minRaw = NULL,  
  maxRaw = NULL  
)
```

**Arguments**

norm	The specific norm score, e. g. T value
model	The model from the regression modeling obtained with the cnorm function
minAge	Age to start from
maxAge	Age to stop at
step	Stepping parameter for the precision when retrieving of the values, lower values indicate higher precision (default 0.1).
minRaw	lower bound of the range of raw scores (default = 0)
maxRaw	upper bound of raw scores

**Value**

data.frame of the variables raw, age and norm

**See Also**

Other predict: [derivationTable\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predict.cnormShash\(\)](#), [predictNorm\(\)](#), [predictRaw\(\)](#), [rawTable\(\)](#)

**Examples**

```
## Not run:  
# Generate cnorm object from example data  
cnorm.elfe <- cnorm(raw = elfe$raw, group = elfe$group)  
getNormCurve(35, cnorm.elfe)  
  
## End(Not run)
```

---

getNormScoreSE	<i>Calculates the standard error (SE) or root mean square error (RMSE) of the norm scores In case of large datasets, both results should be almost identical</i>
----------------	--

---

**Description**

Calculates the standard error (SE) or root mean square error (RMSE) of the norm scores In case of large datasets, both results should be almost identical

**Usage**

```
getNormScoreSE(model, type = 2)
```

**Arguments**

model	a cnorm object
type	either '1' for the standard error senso Oosterhuis et al. (2016) or '2' for the RMSE (default)

**Value**

The standard error (SE) of the norm scores sensu Oosterhuis et al. (2016) or the RMSE

**References**

Oosterhuis, H. E. M., van der Ark, L. A., & Sijtsma, K. (2016). Sample Size Requirements for Traditional and Regression-Based Norms. *Assessment*, 23(2), 191–202. <https://doi.org/10.1177/1073191115580638>

---

modelSummary	<i>Prints the results and regression function of a cnorm model</i>
--------------	--

---

**Description**

Prints the results and regression function of a cnorm model

**Usage**

```
modelSummary(object, ...)
```

**Arguments**

object	A regression model or cnorm object
...	additional parameters

**Value**

A report on the regression function, weights, R2 and RMSE

**See Also**

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [cnorm.cv\(\)](#), [derive\(\)](#), [print.cnorm\(\)](#), [printSubset\(\)](#), [rangeCheck\(\)](#), [regressionFunction\(\)](#), [summary.cnorm\(\)](#)

---

normTable

*Create a norm table based on model for specific age*

---

**Description**

This function generates a norm table for a specific age based on the regression model by assigning raw scores to norm scores. Please specify the range of norm scores, you want to cover. A T value of 25 corresponds to a percentile of .6. As a consequence, specifying a range of T = 25 to T = 75 would cover 98.4 the population. Please be careful when extrapolating vertically (at the lower and upper end of the age specific distribution). Depending on the size of your standardization sample, extreme values with T < 20 or T > 80 might lead to inconsistent results. In case a confidence coefficient (CI, default .9) and the reliability is specified, confidence intervals are computed for the true score estimates, including a correction for regression to the mean (Eid & Schmidt, 2012, p. 272).

**Usage**

```
normTable(
  A,
  model,
  minNorm = NULL,
  maxNorm = NULL,
  minRaw = NULL,
  maxRaw = NULL,
  step = NULL,
  monotonuous = TRUE,
  CI = 0.9,
  reliability = NULL,
  pretty = TRUE
)
```

**Arguments**

A	the age as single value or a vector of age values
model	The regression model from the cnorm function
minNorm	The lower bound of the norm score range
maxNorm	The upper bound of the norm score range
minRaw	clipping parameter for the lower bound of raw scores

maxRaw	clipping parameter for the upper bound of raw scores
step	Stepping parameter with lower values indicating higher precision
monotonuous	corrects for decreasing norm scores in case of model inconsistencies (default)
CI	confidence coefficient, ranging from 0 to 1, default .9
reliability	coefficient, ranging between 0 to 1
pretty	Format table by collapsing intervals and rounding to meaningful precision

### Value

either data.frame with norm scores, predicted raw scores and percentiles in case of simple A value or a list of norm tables if vector of A values was provided

### References

Eid, M. & Schmidt, K. (2012). Testtheorie und Testkonstruktion. Hogrefe.

### See Also

rawTable

Other predict: [derivationTable\(\)](#), [getNormCurve\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predict.cnormShash\(\)](#), [predictNorm\(\)](#), [predictRaw\(\)](#), [rawTable\(\)](#)

### Examples

```
## Not run:
# Generate cnorm object from example data
cnorm.elfe <- cnorm(raw = elfe$raw, group = elfe$group)

# create single norm table
norms <- normTable(3.5, cnorm.elfe, minNorm = 25, maxNorm = 75, step = 0.5)

# create list of norm tables
norms <- normTable(c(2.5, 3.5, 4.5), cnorm.elfe,
  minNorm = 25, maxNorm = 75,
  step = 1, minRaw = 0, maxRaw = 26
)

# conventional norming, set age to arbitrary value
model <- cnorm(raw = elfe$raw)
normTable(0, model)

## End(Not run)
```

---

 normTable.betabinomial

*Calculate Cumulative Probabilities, Density, Percentiles, and Z-Scores for Beta-Binomial Distribution*

---

### Description

This function generates a norm table for a specific ages based on the beta binomial regression model. In case a confidence coefficient (CI, default .9) and the reliability is specified, confidence intervals are computed for the true score estimates, including a correction for regression to the mean (Eid & Schmidt, 2012, p. 272).

### Usage

```
normTable.betabinomial(
  model,
  ages,
  n = NULL,
  m = NULL,
  range = 3,
  CI = 0.9,
  reliability = NULL
)
```

### Arguments

model	The model, which was fitted using the ‘optimized.model’ function.
ages	A numeric vector of age points at which to make predictions.
n	The number of items resp. the maximum score.
m	An optional stop criterion in table generation. Positive integer lower than n.
range	The range of the norm scores in standard deviations. Default is 3. Thus, scores in the range of +/- 3 standard deviations are considered.
CI	confidence coefficient, ranging from 0 to 1, default .9
reliability	coefficient, ranging between 0 to 1

### Value

A list of data frames with columns: x, Px, Pcum, Percentile, z, norm score and possibly confidence interval

---

normTable.shash	<i>Calculate Norm Tables for Sinh-Arcsinh Distribution</i>
-----------------	--

---

**Description**

Generates norm tables for specific ages based on a fitted Sinh-ArcSinh (shash) regression model. Computes probabilities, percentiles, z-scores, and norm scores for a specified range of raw scores. Optionally includes confidence intervals when reliability is provided.

**Usage**

```
normTable.shash(
  model,
  ages,
  start = NULL,
  end = NULL,
  step = 1,
  CI = 0.9,
  reliability = NULL
)
```

**Arguments**

model	Fitted shash model object of class "cnormShash"
ages	Numeric vector of age points for norm table generation
start	Minimum raw score value for the norm table
end	Maximum raw score value for the norm table
step	Step size between consecutive raw scores (default: 1)
CI	Confidence coefficient (0-1, default: 0.9) for confidence intervals
reliability	Reliability coefficient (0-1) for true score confidence intervals

**Details**

For continuous shash distributions, probability densities are computed and converted to cumulative probabilities and percentiles. When reliability is specified, confidence intervals include correction for regression to the mean.

**Value**

List of data frames (one per age) containing:

x	Raw scores
Px	Probability density values
Pcum	Cumulative probabilities
Percentile	Percentile ranks (0-100)

z	Standardized z-scores
norm	Norm scores in specified scale
lowerCI, upperCI	Confidence intervals (if reliability provided)
lowerCI_PR, upperCI_PR	CI as percentile ranks (if reliability provided)

### Examples

```
## Not run:
# Basic norm table
model <- cnorm.shash(age, score)
tables <- normTable.shash(model, ages = c(7, 8, 9), start = 0, end = 50)

# With confidence intervals and finer granularity
tables_ci <- normTable.shash(model, ages = c(8, 9), start = 10, end = 40,
                             step = 0.5, CI = 0.95, reliability = 0.85)

## End(Not run)
```

---

plot.cnorm

*S3 function for plotting cnorm objects*

---

### Description

S3 function for plotting cnorm objects

### Usage

```
## S3 method for class 'cnorm'
plot(x, y, ...)
```

### Arguments

x	the cnorm object
y	the type of plot as a string, can be one of 'raw' (1), 'norm' (2), 'curves' (3), 'percentiles' (4), 'series' (5), 'subset' (6), or 'derivative' (7), either as a string or the according index
...	additional parameters for the specific plotting function

### See Also

Other plot: [compare\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

---

`plot.cnormBetaBinomial`*Plot cnormBetaBinomial Model with Data and Percentile Lines*

---

## Description

This function creates a visualization of a fitted `cnormBetaBinomial` model, including the original data points manifest percentiles and specified percentile lines.

## Usage

```
## S3 method for class 'cnormBetaBinomial'  
plot(x, ...)
```

## Arguments

<code>x</code>	A fitted model object of class "cnormBetaBinomial" or "cnormBetaBinomial2".
<code>...</code>	Additional arguments passed to the plot method. <ul style="list-style-type: none"><li>• <code>age</code> A vector the age data.</li><li>• A vector of the score data.</li><li>• <code>weights</code> An optional numeric vector of weights for each observation.</li><li>• <code>percentiles</code> An optional vector with the percentiles to plot.</li><li>• <code>points</code> Logical indicating whether to plot the data points. Default is TRUE.</li></ul>

## Value

A ggplot object.

## See Also

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

## Examples

```
## Not run:  
# Computing beta binomial models already displays plot  
model.bb <- cnorm.betabinomial(elfe$group, elfe$raw)  
  
# Without data points  
plot(model.bb, age = elfe$group, score = elfe$raw, weights=NULL, points=FALSE)  
  
## End(Not run)
```

---

```
plot.cnormBetaBinomial2
```

*Plot cnormBetaBinomial Model with Data and Percentile Lines*

---

### Description

This function creates a visualization of a fitted cnormBetaBinomial model, including the original data points manifest percentiles and specified percentile lines.

### Usage

```
## S3 method for class 'cnormBetaBinomial2'
plot(x, ...)
```

### Arguments

x	A fitted model object of class "cnormBetaBinomial" or "cnormBetaBinomial2".
...	Additional arguments passed to the plot method. <ul style="list-style-type: none"> <li>• age A vector the age data.</li> <li>• A vector of the score data.</li> <li>• weights An optional numeric vector of weights for each observation.</li> <li>• percentiles An optional vector with the percentiles to plot.</li> <li>• points Logical indicating whether to plot the data points. Default is TRUE.</li> </ul>

### Value

A ggplot object.

### See Also

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

---

```
plot.cnormShash
```

*Plot SinH-ArcSinH Model with Data and Percentile Lines*

---

### Description

Plot SinH-ArcSinH Model with Data and Percentile Lines

### Usage

```
## S3 method for class 'cnormShash'
plot(x, ...)
```

**Arguments**

x                    A fitted model object of class "cnormShash"  
 ...                  Additional arguments including age, score, weights, percentiles, points

**Value**

A ggplot object

---

plotCnorm                    *General convenience plotting function*

---

**Description**

General convenience plotting function

**Usage**

```
plotCnorm(x, y, ...)
```

**Arguments**

x                    a cnorm object  
 y                    the type of plot as a string, can be one of 'raw' (1), 'norm' (2), 'curves' (3),  
 'percentiles' (4), 'series' (5), 'subset' (6), or 'derivative' (7), either as a string or  
 the according index  
 ...                  additional parameters for the specific plotting function

---

plotDensity                    *Plot the density function per group by raw score*

---

**Description**

This function plots density curves based on the regression model against the raw scores. It supports both traditional continuous norming models and beta-binomial models. The function allows for customization of the plot range and groups to be displayed.

**Usage**

```
plotDensity(  
  model,  
  minRaw = NULL,  
  maxRaw = NULL,  
  minNorm = NULL,  
  maxNorm = NULL,  
  group = NULL  
)
```

**Arguments**

model	The model from the bestModel function, a cnorm object, a cnormBetaBinomial, a cnormBetaBinomial2 or cnormShash object.
minRaw	Lower bound of the raw score. If NULL, it's automatically determined based on the model type.
maxRaw	Upper bound of the raw score. If NULL, it's automatically determined based on the model type.
minNorm	Lower bound of the norm score. If NULL, it's automatically determined based on the model type.
maxNorm	Upper bound of the norm score. If NULL, it's automatically determined based on the model type.
group	Numeric vector specifying the age groups to plot. If NULL, groups are automatically selected.

**Details**

The function generates density curves for specified age groups, allowing for easy comparison of score distributions across different ages.

For beta-binomial models, the density is based on the probability mass function, while for traditional models, it uses a normal distribution based on the norm scores.

**Value**

A ggplot object representing the density functions.

**Note**

Please check for inconsistent curves, especially those showing implausible shapes such as violations of biuniqueness in the cnorm models.

**See Also**

[plotNormCurves](#), [plotPercentiles](#)

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

**Examples**

```
## Not run:
# For traditional continuous norming model
result <- cnorm(raw = elfe$raw, group = elfe$group)
plotDensity(result, group = c(2, 4, 6))

# For beta-binomial model
bb_model <- cnorm.betabinomial(age = ppvt$age, score = ppvt$raw, n = 228)
plotDensity(bb_model)
```

```
## End(Not run)
```

---

plotDerivative	<i>Plot first order derivative of regression model</i>
----------------	--

---

### Description

This function plots the scores obtained via the first order derivative of the regression model in dependence of the norm score.

### Usage

```
plotDerivative(  
  model,  
  minAge = NULL,  
  maxAge = NULL,  
  minNorm = NULL,  
  maxNorm = NULL,  
  stepAge = NULL,  
  stepNorm = NULL,  
  order = 1  
)
```

### Arguments

model	The model from the bestModel function, a cnorm object.
minAge	Minimum age to start checking. If NULL, it's automatically determined from the model.
maxAge	Maximum age for checking. If NULL, it's automatically determined from the model.
minNorm	Lower end of the norm score range. If NULL, it's automatically determined from the model.
maxNorm	Upper end of the norm score range. If NULL, it's automatically determined from the model.
stepAge	Stepping parameter for the age check, usually 1 or 0.1; lower values indicate higher precision.
stepNorm	Stepping parameter for norm scores.
order	Degree of the derivative (default = 1).

## Details

The results indicate the progression of the norm scores within each age group. The regression-based modeling approach relies on the assumption of a linear progression of the norm scores. Negative scores in the first order derivative indicate a violation of this assumption. Scores near zero are typical for bottom and ceiling effects in the raw data.

The regression models usually converge within the range of the original values. In case of vertical and horizontal extrapolation, with increasing distance to the original data, the risk of assumption violation increases as well.

## Value

A ggplot object representing the derivative of the regression function.

## Note

This function is currently incompatible with reversed raw score scales ('descent' option).

## See Also

[checkConsistency](#), [bestModel](#), [derive](#)

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

## Examples

```
## Not run:  
# For traditional continuous norming model  
result <- cnorm(raw = elfe$raw, group = elfe$group)  
plotDerivative(result, minAge=2, maxAge=5, stepAge=.2, minNorm=25, maxNorm=75, stepNorm=1)  
  
## End(Not run)
```

---

plotNorm

*Plot manifest and fitted norm scores*

---

## Description

This function plots the manifest norm score against the fitted norm score from the inverse regression model per group. This helps to inspect the precision of the modeling process. The scores should not deviate too far from the regression line. Applicable for Taylor polynomial models.

**Usage**

```
plotNorm(
  model,
  age = NULL,
  score = NULL,
  width = NULL,
  weights = NULL,
  group = FALSE,
  minNorm = NULL,
  maxNorm = NULL,
  type = 0
)
```

**Arguments**

model	The regression model, usually from the 'cnorm' or 'cnorm.betabinomial' function
age	In case of beta binomial model, please provide the age vector
score	In case of beta binomial model, please provide the score vector
width	In case of beta binomial model, please provide the width for the sliding window. If null, the function tries to determine a sensible setting.
weights	Vector or variable name in the dataset with weights for each individual case. If NULL, no weights are used.
group	On optional grouping variable, use empty string for no group, the variable name for Taylor polynomial models or a vector with the groups for beta binomial models
minNorm	lower bound of fitted norm scores
maxNorm	upper bound of fitted norm scores
type	Type of display: 0 = plot manifest against fitted values, 1 = plot manifest against difference values

**Value**

A ggplot object representing the norm scores plot.

**See Also**

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

**Examples**

```
## Not run:
# Load example data set, compute model and plot results

# Taylor polynomial model
```

```

model <- cnorm(raw = elfe$raw, group = elfe$group)
plot(model, "norm")

# Beta binomial models; maximum number of items in elfe is n = 28
model.bb <- cnorm.betabinomial(elfe$group, elfe$raw, n = 28)
plotNorm(model.bb, age = elfe$group, score = elfe$raw)

## End(Not run)

```

---

plotNormCurves

*Plot norm curves*


---

### Description

This function plots the norm curves based on the regression model. It supports both Taylor polynomial models and beta-binomial models.

### Usage

```

plotNormCurves(
  model,
  normList = NULL,
  minAge = NULL,
  maxAge = NULL,
  step = 0.1,
  minRaw = NULL,
  maxRaw = NULL
)

```

### Arguments

model	The model from the bestModel function, a cnorm object, or a cnormBetaBinomial / cnormBetaBinomial2 object.
normList	Vector with norm scores to display. If NULL, default values are used.
minAge	Age to start with checking. If NULL, it's automatically determined from the model.
maxAge	Upper end of the age check. If NULL, it's automatically determined from the model.
step	Stepping parameter for the age check, usually 1 or 0.1; lower scores indicate higher precision.
minRaw	Lower end of the raw score range, used for clipping implausible results. If NULL, it's automatically determined from the model.
maxRaw	Upper end of the raw score range, used for clipping implausible results. If NULL, it's automatically determined from the model.

## Details

Please check the function for inconsistent curves: The different curves should not intersect. Violations of this assumption are a strong indication of violations of model assumptions in modeling the relationship between raw and norm scores.

Common reasons for inconsistencies include: 1. Vertical extrapolation: Choosing extreme norm scores (e.g., scores  $\leq -3$  or  $\geq 3$ ). 2. Horizontal extrapolation: Using the model scores outside the original dataset. 3. The data cannot be modeled with the current approach, or you need another power parameter (k) or R2 for the model.

## Value

A ggplot object representing the norm curves.

## See Also

[checkConsistency](#), [plotDerivative](#), [plotPercentiles](#)

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

## Examples

```
## Not run:
# For Taylor continuous norming model
m <- cnorm(raw = ppvt$raw, group = ppvt$group)
plotNormCurves(m, minAge=2, maxAge=5)

# For beta-binomial model
bb_model <- cnorm.betabinomial(age = ppvt$age, score = ppvt$raw, n = 228)
plotNormCurves(bb_model)

## End(Not run)
```

---

plotPercentiles

*Plot norm curves against actual percentiles*

---

## Description

The function plots the norm curves based on the regression model against the actual percentiles from the raw data. As in 'plotNormCurves', please check for inconsistent curves, especially intersections. Violations of this assumption are a strong indication for problems in modeling the relationship between raw and norm scores. In general, extrapolation (point 1 and 2) can carefully be done to a certain degree outside the original sample, but it should in general be handled with caution. The original percentiles are displayed as distinct points in the according color, the model based projection of percentiles are drawn as lines. Please note, that the estimation of the percentiles of the raw data is done with the quantile function with the default settings. In case, you get 'jagged' or disorganized percentile curve, try to reduce the 'k' and/or 't' parameter in modeling.

**Usage**

```
plotPercentiles(
  model,
  minRaw = NULL,
  maxRaw = NULL,
  minAge = NULL,
  maxAge = NULL,
  raw = NULL,
  group = NULL,
  percentiles = c(0.025, 0.1, 0.25, 0.5, 0.75, 0.9, 0.975),
  scale = NULL,
  title = NULL,
  subtitle = NULL,
  points = FALSE
)
```

**Arguments**

model	The Taylor polynomial regression model object from the cNORM
minRaw	Lower bound of the raw score (default = 0)
maxRaw	Upper bound of the raw score
minAge	Variable to restrict the lower bound of the plot to a specific age
maxAge	Variable to restrict the upper bound of the plot to a specific age
raw	The name of the raw variable
group	The name of the grouping variable; the distinct groups are automatically determined
percentiles	Vector with percentile scores, ranging from 0 to 1 (exclusive)
scale	The norm scale, either 'T', 'IQ', 'z', 'percentile' or self defined with a double vector with the mean and standard deviation, f. e. c(10, 3) for Wechsler scale index points; if NULL, scale information from the data preparation is used (default)
title	custom title for plot
subtitle	custom title for plot
points	Logical indicating whether to plot the data points. Default is TRUE.

**See Also**

plotNormCurves, plotPercentileSeries

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

**Examples**

```
## Not run:
# Load example data set, compute model and plot results
result <- cnorm(raw = elfe$raw, group = elfe$group)
plotPercentiles(result)

## End(Not run)
```

---

plotPercentileSeries *Generates a series of plots with number curves by percentile for different models*

---

**Description**

This functions makes use of 'plotPercentiles' to generate a series of plots with different number of predictors. It draws on the information provided by the model object to determine the bounds of the modeling (age and standard score range). It can be used as an additional model check to determine the best fitting model. Please have a look at the 'plotPercentiles' function for further information.

**Usage**

```
plotPercentileSeries(
  model,
  start = 1,
  end = NULL,
  group = NULL,
  percentiles = c(0.025, 0.1, 0.25, 0.5, 0.75, 0.9, 0.975),
  filename = NULL
)
```

**Arguments**

model	The Taylor polynomial regression model object from the cNORM
start	Number of predictors to start with
end	Number of predictors to end with
group	The name of the grouping variable; the distinct groups are automatically determined
percentiles	Vector with percentile scores, ranging from 0 to 1 (exclusive)
filename	Prefix of the filename. If specified, the plots are saves as png files in the directory of the workspace, instead of displaying them

**Value**

the complete list of plots

**See Also**

plotPercentiles

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#), [plotSubset\(\)](#)

**Examples**

```
## Not run:
# Load example data set, compute model and plot results
result <- cnorm(raw = elfe$raw, group = elfe$group)
plotPercentileSeries(result, start=4, end=6)

## End(Not run)
```

---

plotRaw

*Plot manifest and fitted raw scores*

---

**Description**

The function plots the raw data against the fitted scores from the regression model per group. This helps to inspect the precision of the modeling process. The scores should not deviate too far from regression line.

**Usage**

```
plotRaw(model, group = FALSE, type = 0)
```

**Arguments**

model	The regression model from the 'cnorm' function
group	Should the fit be displayed by group?
type	Type of display: 0 = plot manifest against fitted values, 1 = plot manifest against difference values

**See Also**

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotSubset\(\)](#)

**Examples**

```
## Not run:
# Compute model with example dataset and plot results
result <- cnorm(raw = elfe$raw, group = elfe$group)
plotRaw(result)

## End(Not run)
```

---

plotSubset

*Evaluate information criteria for regression model*


---

**Description**

This function plots various information criteria and model fit statistics against the number of predictors or adjusted R-squared, depending on the type of plot selected. It helps in model selection by visualizing different aspects of model performance. Models, which did not pass the initial consistency check are depicted with an empty circle.

**Usage**

```
plotSubset(model, type = 0)
```

**Arguments**

model	The regression model from the bestModel function or a cnorm object.
type	Integer specifying the type of plot to generate: <ul style="list-style-type: none"> <li>• 0: Adjusted R2 by number of predictors (default)</li> <li>• 1: Log-transformed Mallows' Cp by adjusted R2</li> <li>• 2: Bayesian Information Criterion (BIC) by adjusted R2</li> <li>• 3: Root Mean Square Error (RMSE) by number of predictors</li> <li>• 4: Residual Sum of Squares (RSS) by number of predictors</li> <li>• 5: F-test statistic for consecutive models by number of predictors</li> <li>• 6: p-value for model tests by number of predictors</li> </ul>

**Details**

The function generates different plots to help in model selection:

- For types 1 and 2 (Mallows' Cp and BIC), look for the "elbow" in the curve where the information criterion begins to drop. This often indicates a good balance between model fit and complexity.
- For type 0 (Adjusted R2), higher values indicate better fit, but be cautious of overfitting with values approaching 1.
- For types 3 and 4 (RMSE and RSS), lower values indicate better fit.
- For type 5 (F-test), higher values suggest significant improvement with added predictors.
- For type 6 (p-values), values below the significance level (typically 0.05) suggest significant improvement with added predictors.

**Value**

A ggplot object representing the selected information criterion plot.

**Note**

It's important to balance statistical measures with practical considerations and to visually inspect the model fit using functions like `plotPercentiles`.

**See Also**

[bestModel](#), [plotPercentiles](#), [printSubset](#)

Other plot: [compare\(\)](#), [plot.cnorm\(\)](#), [plot.cnormBetaBinomial\(\)](#), [plot.cnormBetaBinomial2\(\)](#), [plotDensity\(\)](#), [plotDerivative\(\)](#), [plotNorm\(\)](#), [plotNormCurves\(\)](#), [plotPercentileSeries\(\)](#), [plotPercentiles\(\)](#), [plotRaw\(\)](#)

**Examples**

```
## Not run:
# Compute model with example data and plot information function
cnorm.model <- cnorm(raw = elfe$raw, group = elfe$group)
plotSubset(cnorm.model)

# Plot BIC against adjusted R-squared
plotSubset(cnorm.model, type = 2)

# Plot RMSE against number of predictors
plotSubset(cnorm.model, type = 3)

## End(Not run)
```

---

ppvt

*Vocabulary development from 2.5 to 17*

---

**Description**

A dataset based on an unstratified sample of PPVT4 data (German adaption). The PPVT4 consists of blocks of items with 12 items each. Each item consists of 4 pictures. The test taker is given a word orally and he or she has to point out the picture matching the oral word. Bottom and ceiling blocks of items are determined according to age and performance. For instance, when a student knows less than 4 word from a block of 12 items, the testing stops. The sample is not identical with the norm sample and includes doublets of cases in order to align the sample size per age group. It is primarily intended for running the cNORM analyses with regard to modeling and stratification.

**Usage**

```
ppvt
```

## Format

A data frame with 4542 rows and 6 variables:

**age** the chronological age of the child

**sex** the sex of the test taker, 1=male, 2=female

**migration** migration status of the family, 0=no, 1=yes

**region** factor specifying the region, the data were collected; grouped into south, north, east and west

**raw** the raw score of the student, spanning values from 0 to 228

**group** age group of the child, determined by the `getGroups()`-function with 12 equidistant age groups

A data frame with 5600 rows and 9 columns

## Source

<https://www.psychometrica.de/ppvt4.html>

## References

Lenhard, A., Lenhard, W., Segerer, R. & Suggate, S. (2015). Peabody Picture Vocabulary Test - Revision IV (Deutsche Adaption). Frankfurt a. M./Germany: Pearson Assessment.

## Examples

```
## Not run:
# Example with continuous age variable, ranked with sliding window
model.ppvt.sliding <- cnorm(age=ppvt$age, raw=ppvt$raw, width=1)

# Example with age groups; you might first want to experiment with
# the granularity of the groups via the 'getGroups()' function
model.ppvt.group <- cnorm(group=ppvt$group, raw=ppvt$raw) # with predefined groups
model.ppvt.group <- cnorm(group=getGroups(ppvt$age, n=15, equidistant = T),
                          raw=ppvt$raw) # groups built 'on the fly'

# plot information function
plot(model.ppvt.group, "subset")

# check model consistency
checkConsistency(model.ppvt.group)

# plot percentiles
plot(model.ppvt.group, "percentiles")

## End(Not run)
```

---

predict.cnormBetaBinomial

*Predict Norm Scores from Raw Scores*

---

### Description

This function calculates norm scores based on raw scores, age, and a fitted cnormBetaBinomial model.

### Usage

```
## S3 method for class 'cnormBetaBinomial'
predict(object, ...)
```

### Arguments

object	A fitted model object of class 'cnormBetaBinomial' or 'cnormBetaBinomial2'.
...	Additional arguments passed to the prediction method: <ul style="list-style-type: none"> <li>• age A numeric vector of ages, same length as raw.</li> <li>• score A numeric vector of raw scores.</li> <li>• range The range of the norm scores in standard deviations. Default is 3. Thus, scores in the range of +/- 3 standard deviations are considered.</li> </ul>

### Details

The function first predicts the alpha and beta parameters of the beta-binomial distribution for each age using the provided model. It then calculates the cumulative probability for each raw score given these parameters. Finally, it converts these probabilities to the norm scale specified in the model.

### Value

A numeric vector of norm scores.

### See Also

Other predict: [derivationTable\(\)](#), [getNormCurve\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predict.cnormShash\(\)](#), [predictNorm\(\)](#), [predictRaw\(\)](#), [rawTable\(\)](#)

### Examples

```
## Not run:
# Assuming you have a fitted model named 'bb_model':
model <- cnorm.betabinomial(ppvt$age, ppvt$raw)
raw <- c(100, 121, 97, 180)
ages <- c(7, 8, 9, 10)
norm_scores <- predict(model, ages, raw)

## End(Not run)
```

---

`predict.cnormBetaBinomial2`*Predict Norm Scores from Raw Scores*

---

## Description

This function calculates norm scores based on raw scores, age, and a fitted `cnormBetaBinomial` model.

## Usage

```
## S3 method for class 'cnormBetaBinomial2'  
predict(object, ...)
```

## Arguments

<code>object</code>	A fitted model object of class <code>'cnormBetaBinomial'</code> or <code>'cnormBetaBinomial2'</code> .
<code>...</code>	Additional arguments passed to the prediction method: <ul style="list-style-type: none"><li>• <code>age</code> A numeric vector of ages, same length as <code>raw</code>.</li><li>• <code>score</code> A numeric vector of raw scores.</li><li>• <code>range</code> The range of the norm scores in standard deviations. Default is 3. Thus, scores in the range of +/- 3 standard deviations are considered.</li></ul>

## Details

The function first predicts the alpha and beta parameters of the beta-binomial distribution for each age using the provided model. It then calculates the cumulative probability for each raw score given these parameters. Finally, it converts these probabilities to the norm scale specified in the model.

## Value

A numeric vector of norm scores.

## See Also

Other predict: [derivationTable\(\)](#), [getNormCurve\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormShash\(\)](#), [predictNorm\(\)](#), [predictRaw\(\)](#), [rawTable\(\)](#)

## Examples

```
## Not run:  
# Assuming you have a fitted model named 'bb_model':  
model <- cnorm.betabinomial(ppvt$age, ppvt$raw)  
raw <- c(100, 121, 97, 180)  
ages <- c(7, 8, 9, 10)  
norm_scores <- predict(model, ages, raw)  
  
## End(Not run)
```

---

predict.cnormShash      *Predict Norm Scores from Raw Scores*

---

### Description

This function calculates norm scores based on raw scores, age, and a fitted cnormShash model.

### Usage

```
## S3 method for class 'cnormShash'
predict(object, ...)
```

### Arguments

object	A fitted model object of class 'cnormShash'.
...	Additional arguments passed to the prediction method: <ul style="list-style-type: none"> <li>• age A numeric vector of ages, same length as score.</li> <li>• score A numeric vector of raw scores.</li> <li>• range The range of the norm scores in standard deviations. Default is 3. Thus, scores in the range of +/- 3 standard deviations are considered.</li> </ul>

### Details

The function predicts the SinH-ArcSinH (shash) distribution parameters (mu, sigma, epsilon, delta) for each age using the provided model. It then calculates the cumulative probability for each raw score given these parameters using the continuous shash distribution. Finally, it converts these probabilities to the norm scale specified in the model.

### Value

A numeric vector of norm scores.

### See Also

Other predict: [derivationTable\(\)](#), [getNormCurve\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predictNorm\(\)](#), [predictRaw\(\)](#), [rawTable\(\)](#)

### Examples

```
## Not run:
# Assuming you have a fitted model named 'shash_model':
model <- cnorm.shash(children$age, children$score)
raw_scores <- c(25.5, 30.2, 18.7, 45.3)
ages <- c(7, 8, 9, 10)
norm_scores <- predict(model, ages, raw_scores)

## End(Not run)
```

---

predictNorm	<i>Retrieve norm value for raw score at a specific age</i>
-------------	--

---

### Description

This functions numerically determines the norm score for raw scores depending on the level of the explanatory variable A, e. g. norm scores for raw scores at given ages.

### Usage

```
predictNorm(
  raw,
  A,
  model,
  minNorm = NULL,
  maxNorm = NULL,
  force = TRUE,
  silent = FALSE
)
```

### Arguments

raw	The raw value, either single numeric or numeric vector
A	the explanatory variable (e. g. age), either single numeric or numeric vector
model	The regression model or a cnorm object
minNorm	The lower bound of the norm score range
maxNorm	The upper bound of the norm score range
force	Try to resolve missing norm scores in case of inconsistent models
silent	set to TRUE to suppress messages

### Value

The predicted norm score for a raw score, either single value or vector

### See Also

Other predict: [derivationTable\(\)](#), [getNormCurve\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predict.cnormShash\(\)](#), [predictRaw\(\)](#), [rawTable\(\)](#)

### Examples

```
## Not run:
# Generate cnorm object from example data
cnorm.elfe <- cnorm(raw = elfe$raw, group = elfe$group)

# return norm value for raw value 21 for grade 2, month 9
```

```
specificNormValue <- predictNorm(row = 21, A = 2.75, cnorm.elfe)

## End(Not run)
```

---

predictRaw                      *Predict raw values*

---

### Description

Most elementary function to predict raw score based on Location (L, T score), Age (grouping variable) and the coefficients from a regression model.

### Usage

```
predictRaw(norm, age, coefficients, minRaw = -Inf, maxRaw = Inf)
```

### Arguments

norm	The norm score, e. g. a specific T score or a vector of scores
age	The age value or a vector of scores
coefficients	The a cnorm object or the coefficients from the regression model
minRaw	Minimum score for the results; can be used for clipping unrealistic outcomes, usually set to the lower bound of the range of values of the test (default: 0)
maxRaw	Maximum score for the results; can be used for clipping unrealistic outcomes usually set to the upper bound of the range of values of the test

### Value

the predicted raw score or a data.frame of scores in case, lists of norm scores or age is used

### See Also

Other predict: [derivationTable\(\)](#), [getNormCurve\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predict.cnormShash\(\)](#), [predictNorm\(\)](#), [rawTable\(\)](#)

### Examples

```
## Not run:
# Prediction of single scores
model <- cnorm(row = elfe$raw, group = elfe$group)
predictRaw(35, 3.5, model)

## End(Not run)
```

---

prepareData	<i>Prepare data for modeling in one step (convenience method)</i>
-------------	---

---

### Description

This is a convenience method to either load the inbuilt sample dataset, or to provide a data frame with the variables "raw" (for the raw scores) and "group". The function ranks the data within groups, computes norm values, powers of the norm scores and interactions. Afterwards, you can use these preprocessed data to determine the best fitting model.

### Usage

```
prepareData(
  data = NULL,
  group = "group",
  raw = "raw",
  age = "group",
  k = 4,
  t = NULL,
  width = NA,
  weights = NULL,
  scale = "T",
  descend = FALSE,
  silent = FALSE
)
```

### Arguments

data	data.frame with a grouping variable named 'group' and a raw score variable named 'raw'.
group	grouping variable in the data, e. g. age groups, grades ... Setting group = FALSE deactivates modeling in dependence of age. Use this in case you do want conventional norm tables.
raw	the raw scores
age	the continuous explanatory variable; by default set to "group"
k	The power parameter, default = 4
t	the age power parameter (default NULL). If not set, cNORM automatically uses k. The age power parameter can be used to specify the k to produce rectangular matrices and specify the course of scores per independently from k
width	if a width is provided, the function switches to rankBySlidingWindow to determine the observed raw scores, otherwise, ranking is done by group (default)
weights	Vector or variable name in the dataset with weights for each individual case. It can be used to compensate for moderate imbalances due to insufficient norm data stratification. Weights should be numerical and positive. Please use the 'computeWeights' function for this purpose.

scale	type of norm scale, either T (default), IQ, z or percentile (= no transformation); a double vector with the mean and standard deviation can as well, be provided f. e. c(10, 3) for Wechsler scale index point
descend	ranking order (default descent = FALSE): inverses the ranking order with higher raw scores getting lower norm scores; relevant for example when norming error scores, where lower scores mean higher performance
silent	set to TRUE to suppress messages

### Details

The functions `rankBySlidingWindow`, `rankByGroup`, `bestModel`, `computePowers` and `prepareData` are usually not called directly, but accessed through other functions like `cnorm`.

### Value

data frame including the norm scores, powers and interactions of the norm score and grouping variable

### See Also

Other prepare: [computePowers\(\)](#), [rankByGroup\(\)](#), [rankBySlidingWindow\(\)](#)

### Examples

```
## Not run:
# conducts ranking and computation of powers and interactions with the 'elfe' dataset
data.elfe <- prepareData(elfe)

# use vectors instead of data frame
data.elfe <- prepareData(raw=elfe$raw, group=elfe$group)

# variable names can be specified as well, here with the BMI data included in the package
data.bmi <- prepareData(CDC, group = "group", raw = "bmi", age = "age")

## End(Not run)

# modeling with only one group with the 'elfe' dataset as an example
# this results in conventional norming
data.elfe2 <- prepareData(data = elfe, group = FALSE)
m <- bestModel(data.elfe2)
```

---

print.cnorm

*S3 method for printing model selection information*

---

### Description

After conducting the model fitting procedure on the data set, the best fitting model has to be chosen. The `print` function shows the  $R^2$  and other information on the different best fitting models with increasing number of predictors.

**Usage**

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

**Arguments**

x                    The model from the 'bestModel' function or a cnorm object  
...                   additional parameters

**Value**

A table with information criteria

**See Also**

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [cnorm.cv\(\)](#), [derive\(\)](#), [modelSummary\(\)](#),  
[printSubset\(\)](#), [rangeCheck\(\)](#), [regressionFunction\(\)](#), [summary.cnorm\(\)](#)

---

print.cnormShash            *Print method for SinH-ArcSinH objects*

---

**Description**

Print method for SinH-ArcSinH objects

**Usage**

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

**Arguments**

x                    A cnormShash object  
...                   Additional arguments

---

printSubset	<i>Print Model Selection Information</i>
-------------	--

---

**Description**

Displays  $R^2$  and other metrics for models with varying predictors, aiding in choosing the best-fitting model after model fitting.

**Usage**

```
printSubset(x, ...)
```

**Arguments**

x	Model output from 'bestModel' or a cnorm object.
...	Additional parameters.

**Value**

Table with model information criteria.

**See Also**

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [cnorm.cv\(\)](#), [derive\(\)](#), [modelSummary\(\)](#), [print.cnorm\(\)](#), [rangeCheck\(\)](#), [regressionFunction\(\)](#), [summary.cnorm\(\)](#)

**Examples**

```
## Not run:
# Using cnorm object from sample data
result <- cnorm(raw = elfe$raw, group = elfe$group)
printSubset(result)

## End(Not run)
```

---

rangeCheck	<i>Check for horizontal and vertical extrapolation</i>
------------	--

---

**Description**

Regression model only work in a specific range and extrapolation horizontally (outside the original range) or vertically (extreme norm scores) might lead to inconsistent results. The function generates a message, indicating extrapolation and the range of the original data.

**Usage**

```
rangeCheck(  
  object,  
  minAge = NULL,  
  maxAge = NULL,  
  minNorm = NULL,  
  maxNorm = NULL,  
  digits = 3,  
  ...  
)
```

**Arguments**

object	The regression model or a cnorm object
minAge	The lower age bound
maxAge	The upper age bound
minNorm	The lower norm value bound
maxNorm	The upper norm value bound
digits	The precision for rounding the norm and age data
...	additional parameters

**Value**

the report

**See Also**

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [cnorm.cv\(\)](#), [derive\(\)](#), [modelSummary\(\)](#), [print.cnorm\(\)](#), [printSubset\(\)](#), [regressionFunction\(\)](#), [summary.cnorm\(\)](#)

**Examples**

```
## Not run:  
  m <- cnorm(raw = elfe$raw, group = elfe$group)  
  rangeCheck(m)  
  
## End(Not run)
```

rankByGroup

*Determine the norm scores of the participants in each subsample***Description**

This is the initial step, usually done in all kinds of test norming projects, after the scale is constructed and the norm sample is established. First, the data is grouped according to a grouping variable and afterwards, the percentile for each raw value is retrieved. The percentile can be used for the modeling procedure, but in case, the samples do not deviate too much from normality, T, IQ or z scores can be computed via a normal rank procedure based on the inverse cumulative normal distribution. In case of bindings, we use the medium rank and there are different methods for estimating the percentiles (default RankIt).

**Usage**

```
rankByGroup(
  data = NULL,
  group = "group",
  raw = "raw",
  weights = NULL,
  method = 4,
  scale = "T",
  descend = FALSE,
  descriptives = TRUE,
  na.rm = TRUE,
  silent = FALSE
)
```

**Arguments**

data	data.frame with norm sample data. If no data.frame is provided, the raw score and group vectors are directly used
group	name of the grouping variable (default 'group') or numeric vector, e. g. grade, setting group to FALSE cancels grouping (data is treated as one group)
raw	name of the raw value variable (default 'raw') or numeric vector
weights	Vector or variable name in the dataset with weights for each individual case. It can be used to compensate for moderate imbalances due to insufficient norm data stratification. Weights should be numerical and positive. Please use the 'computeWeights' function for this purpose.
method	Ranking method in case of bindings, please provide an index, choosing from the following methods: 1 = Blom (1958), 2 = Tukey (1949), 3 = Van der Warden (1952), 4 = Rankit (default), 5 = Levenbach (1953), 6 = Filliben (1975), 7 = Yu & Huang (2001)
scale	type of norm scale, either T (default), IQ, z or percentile (= no transformation); a double vector with the mean and standard deviation can as well, be provided f. e. c(10, 3) for Wechsler scale index points

descend	ranking order (default descent = FALSE): inverses the ranking order with higher raw scores getting lower norm scores; relevant for example when norming error scores, where lower scores mean higher performance
descriptives	If set to TRUE (default), information in n, mean, median and standard deviation per group is added to each observation
na.rm	remove values, where the percentiles could not be estimated, most likely happens in the context of weighting
silent	set to TRUE to suppress messages

**Value**

the dataset with the percentiles and norm scales per group

**Remarks on using covariates**

So far the inclusion of a binary covariate is experimental and far from optimized. The according variable name has to be specified in the ranking procedure and the modeling includes this in the further process. At the moment, during ranking the data are split into the according cells group x covariate, which leads to small sample sizes. Please take care to have enough cases in each combination. Additionally, covariates can lead to unstable modeling solutions. The question, if it is really reasonable to include covariates when norming a test is a decision beyond the pure data modeling. Please use with care or alternatively split the dataset into the two groups beforehand and model them separately.

The functions rankBySlidingWindow, rankByGroup, bestModel, computePowers and prepareData are usually not called directly, but accessed through other functions like cnorm.

**See Also**

rankBySlidingWindow, computePowers, computeWeights, weighted.rank

Other prepare: [computePowers\(\)](#), [prepareData\(\)](#), [rankBySlidingWindow\(\)](#)

**Examples**

```
## Not run:
# Transformation with default parameters: RankIt and converting to T scores
data.elfe <- rankByGroup(elfe, group = "group") # using a data frame with vector names
data.elfe2 <- rankByGroup(raw=elfe$raw, group=elfe$group) # use vectors for raw score and group

# Transformation into Wechsler scores with Yu & Huang (2001) ranking procedure
data.elfe <- rankByGroup(raw = elfe$raw, group = elfe$group, method = 7, scale = c(10, 3))

# cNORM can as well be used for conventional norming, in case no group is given
d <- rankByGroup(raw = elfe$raw)
d <- computePowers(d)
m <- bestModel(d)
rawTable(0, m) # please use an arbitrary value for age when generating the tables

## End(Not run)
```

---

rankBySlidingWindow *Determine the norm scores of the participants by sliding window*

---

### Description

The function retrieves all individuals in the predefined age range ( $x \pm \text{width}/2$ ) around each case and ranks that individual based on this individually drawn sample. This function can be directly used with a continuous age variable in order to avoid grouping. When collecting data on the basis of a continuous age variable, cases located far from the mean age of the group receive distorted percentiles when building discrete groups and generating percentiles with the traditional approach. The distortion increases with distance from the group mean and this effect can be avoided by the sliding window. Nonetheless, please ensure, that the optional grouping variable in fact represents the correct mean age of the respective age groups, as this variable is later on used for displaying the manifest data in the percentile plots.

### Usage

```
rankBySlidingWindow(
  data = NULL,
  age = "age",
  raw = "raw",
  weights = NULL,
  width,
  method = 4,
  scale = "T",
  descend = FALSE,
  descriptives = TRUE,
  nGroup = 0,
  group = NA,
  na.rm = TRUE,
  silent = FALSE
)
```

### Arguments

data	data.frame with norm sample data
age	the continuous age variable. Setting 'age' to FALSE inhibits computation of powers of age and the interactions
raw	name of the raw value variable (default 'raw')
weights	Vector or variable name in the dataset with weights for each individual case. It can be used to compensate for moderate imbalances due to insufficient norm data stratification. Weights should be numerical and positive. It can be resource intense when applied to the sliding window. Please use the 'computeWeights' function for this purpose.
width	the width of the sliding window

method	Ranking method in case of bindings, please provide an index, choosing from the following methods: 1 = Blom (1958), 2 = Tukey (1949), 3 = Van der Warden (1952), 4 = Rankit (default), 5 = Levenbach (1953), 6 = Filliben (1975), 7 = Yu & Huang (2001)
scale	type of norm scale, either T (default), IQ, z or percentile (= no transformation); a double vector with the mean and standard deviation can as well, be provided f. e. c(10, 3) for Wechsler scale index points
descend	ranking order (default descent = FALSE): inverses the ranking order with higher raw scores getting lower norm scores; relevant for example when norming error scores, where lower scores mean higher performance
descriptives	If set to TRUE (default), information in n, mean, median and standard deviation per group is added to each observation
nGroup	If set to a positive value, a grouping variable is created with the desired number of equi distant groups, named by the group mean age of each group. It creates the column 'group' in the data.frame and in case, there is already one with that name, overwrites it.
group	Optional parameter for providing the name of the grouping variable (if present; overwritten if ngroups is used)
na.rm	remove values, where the percentiles could not be estimated, most likely happens in the context of weighting
silent	set to TRUE to suppress messages

### Details

In case of bindings, the function uses the medium rank and applies the algorithms already described in the [rankByGroup](#) function. At the upper and lower end of the data sample, the sliding stops and the sample is drawn from the interval  $\text{min} + \text{width}$  and  $\text{max} - \text{width}$ , respectively.

### Value

the dataset with the individual percentiles and norm scores

### Remarks on using covariates

So far the inclusion of a binary covariate is experimental and far from optimized. The according variable name has to be specified in the ranking procedure and the modeling includes this in the further process. At the moment, during ranking the data are split into the according degrees of the covariate and the ranking is done separately. This may lead to small sample sizes. Please take care to have enough cases in each combination. Additionally, covariates can lead to unstable modeling solutions. The question, if it is really reasonable to include covariates when norming a test is a decision beyond the pure data modeling. Please use with care or alternatively split the dataset into the two groups beforehand and model them separately.

The functions `rankBySlidingWindow`, `rankByGroup`, `bestModel`, `computePowers` and `prepareData` are usually not called directly, but accessed through other functions like `cnorm`.

**See Also**

rankByGroup, computePowers, computeWeights, weighted.rank, weighted.quantile  
 Other prepare: `computePowers()`, `prepareData()`, `rankByGroup()`

**Examples**

```
## Not run:
# Transformation using a sliding window
data.elfe2 <- rankBySlidingWindow(relfe, raw = "raw", age = "group", width = 0.5)

# Comparing this to the traditional approach should give us exactly the same
# values, since the sample dataset only has a grouping variable for age
data.elfe <- rankByGroup(elfe, group = "group")
mean(data.elfe$normValue - data.elfe2$normValue)

## End(Not run)
```

---

rawTable

*Create a table with norm scores assigned to raw scores for a specific age based on the regression model*

---

**Description**

This function is comparable to 'normTable', despite it reverses the assignment: A table with raw scores and the according norm scores for a specific age based on the regression model is generated. This way, the inverse function of the regression model is solved numerically with brute force. Please specify the range of raw values, you want to cover. With higher precision and smaller stepping, this function becomes computational intensive. In case a confidence coefficient (CI, default .9) and the reliability is specified, confidence intervals are computed for the true score estimates, including a correction for regression to the mean (Eid & Schmidt, 2012, p. 272).

**Usage**

```
rawTable(
  A,
  model,
  minRaw = NULL,
  maxRaw = NULL,
  minNorm = NULL,
  maxNorm = NULL,
  step = 1,
  monotonuous = TRUE,
  CI = 0.9,
  reliability = NULL,
  pretty = TRUE
)
```

**Arguments**

A	the age, either single value or vector with age values
model	The regression model or a cnorm object
minRaw	The lower bound of the raw score range
maxRaw	The upper bound of the raw score range
minNorm	Clipping parameter for the lower bound of norm scores (default 25)
maxNorm	Clipping parameter for the upper bound of norm scores (default 25)
step	Stepping parameter for the raw scores (default 1)
monotonuous	corrects for decreasing norm scores in case of model inconsistencies (default)
CI	confidence coefficient, ranging from 0 to 1, default .9
reliability	coefficient, ranging between 0 to 1
pretty	Format table by collapsing intervals and rounding to meaningful precision

**Value**

either data.frame with raw scores and the predicted norm scores in case of simple A value or a list of norm tables if vector of A values was provided

**References**

Eid, M. & Schmidt, K. (2012). Testtheorie und Testkonstruktion. Hogrefe.

**See Also**

normTable

Other predict: [derivationTable\(\)](#), [getNormCurve\(\)](#), [normTable\(\)](#), [predict.cnormBetaBinomial\(\)](#), [predict.cnormBetaBinomial2\(\)](#), [predict.cnormShash\(\)](#), [predictNorm\(\)](#), [predictRaw\(\)](#)

**Examples**

```
## Not run:
# Generate cnorm object from example data
cnorm.elfe <- cnorm(raw = elfe$raw, group = elfe$group)
# generate a norm table for the raw value range from 0 to 28 for the time point month 7 of grade 3
table <- rawTable(3 + 7 / 12, cnorm.elfe, minRaw = 0, maxRaw = 28)

# generate several raw tables
table <- rawTable(c(2.5, 3.5, 4.5), cnorm.elfe, minRaw = 0, maxRaw = 28)

# additionally compute confidence intervals
table <- rawTable(c(2.5, 3.5, 4.5), cnorm.elfe, minRaw = 0, maxRaw = 28, CI = .9, reliability = .94)

# conventional norming, set age to arbitrary value
model <- cnorm(raw = elfe$raw)
rawTable(0, model)

## End(Not run)
```

---

regressionFunction      *Regression function*

---

### Description

The method builds the regression function for the regression model, including the beta weights. It can be used to predict the raw scores based on age and location.

### Usage

```
regressionFunction(model, raw = NULL, digits = NULL)
```

### Arguments

model	The regression model from the bestModel function or a cnorm object
raw	The name of the raw value variable (default 'raw')
digits	Number of digits for formatting the coefficients

### Value

The regression formula as a string

### See Also

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [cnorm.cv\(\)](#), [derive\(\)](#), [modelSummary\(\)](#), [print.cnorm\(\)](#), [printSubset\(\)](#), [rangeCheck\(\)](#), [summary.cnorm\(\)](#)

### Examples

```
## Not run:
  result <- cnorm(raw = elfe$raw, group = elfe$group)
  regressionFunction(result)

## End(Not run)
```

---

shash      *Sinh-Arcsinh (shash) Distribution*

---

### Description

Density, distribution function, quantile function and random generation for the Sinh-Arcsinh distribution with location parameter  $\mu$ , scale parameter  $\sigma$ , skewness parameter  $\epsilon$ , and tail weight parameter  $\delta$ .

**Usage**

```
dshash(x, mu = 0, sigma = 1, epsilon = 0, delta = 1, log = FALSE)
```

```
pshash(
  q,
  mu = 0,
  sigma = 1,
  epsilon = 0,
  delta = 1,
  lower.tail = TRUE,
  log.p = FALSE
)
```

```
qshash(
  p,
  mu = 0,
  sigma = 1,
  epsilon = 0,
  delta = 1,
  lower.tail = TRUE,
  log.p = FALSE
)
```

```
rshash(n, mu = 0, sigma = 1, epsilon = 0, delta = 1)
```

**Arguments**

<code>x, q</code>	vector of quantiles
<code>mu</code>	location parameter (default: 0)
<code>sigma</code>	scale parameter (must be > 0, default: 1)
<code>epsilon</code>	skewness parameter (default: 0, symmetric distribution)
<code>delta</code>	tail weight parameter (must be > 0, default: 1 for normal-like tails)
<code>log, log.p</code>	logical; if TRUE, probabilities <code>p</code> are given as $\log(p)$
<code>lower.tail</code>	logical; if TRUE (default), probabilities are $P[X \leq x]$ otherwise, $P[X > x]$
<code>p</code>	vector of probabilities
<code>n</code>	number of observations. If $\text{length}(n) > 1$ , the length is taken to be the number required.

**Details**

The Sinh-Arcsinh distribution (Jones & Pewsey, 2009) is defined by the transformation:

$$X = \mu + \sigma \cdot \sinh\left(\frac{\text{asinh}(Z) - \epsilon}{\delta}\right)$$

where  $Z \sim N(0, 1)$  is a standard normal variable.

The four parameters control:

- mu: Location (similar to mean)
- sigma: Scale (similar to standard deviation)
- epsilon: Skewness (epsilon = 0 gives symmetry)
- delta: Tail weight (delta = 1 gives normal-like tails, delta > 1 gives heavier tails, delta < 1 gives lighter tails)

### Value

dshash gives the density, pshash gives the distribution function, qshash gives the quantile function, and rshash generates random deviates.

The length of the result is determined by n for rshash, and is the maximum of the lengths of the numerical arguments for the other functions.

### References

Jones, M. C., & Pewsey, A. (2009). Sinh-arcsinh distributions. *Biometrika*, 96(4), 761-780. doi:[10.1093/biomet/asp053](https://doi.org/10.1093/biomet/asp053)

### See Also

[Normal](#) for the normal distribution.

### Examples

```
## Not run:
# Generate random samples
x <- rshash(1000, mu = 0, sigma = 1, epsilon = 0.5, delta = 1.2)

# Density
plot(density(x))
curve(dshash(x, mu = 0, sigma = 1, epsilon = 0.5, delta = 1.2),
      add = TRUE, col = "red")

# Cumulative probability
pshash(0, mu = 0, sigma = 1, epsilon = 0.5, delta = 1.2)

# Quantiles
qshash(c(0.025, 0.5, 0.975), mu = 0, sigma = 1, epsilon = 0.5, delta = 1.2)

# Compare with normal distribution (epsilon = 0, delta = 1)
par(mfrow = c(2, 2))
x_vals <- seq(-4, 4, length.out = 200)
plot(x_vals, dshash(x_vals), type = "l", main = "Symmetric (like normal)")
plot(x_vals, dshash(x_vals, epsilon = 1), type = "l", main = "Right skewed")
plot(x_vals, dshash(x_vals, delta = 2), type = "l", main = "Heavy tails")
plot(x_vals, dshash(x_vals, delta = 0.5), type = "l", main = "Light tails")

## End(Not run)
```

---

simMean	<i>Simulate mean per age</i>
---------	------------------------------

---

**Description**

Simulate mean per age

**Usage**

```
simMean(age)
```

**Arguments**

age                    the age variable

**Value**

return predicted means

**Examples**

```
## Not run:  
x <- simMean(a)  
  
## End(Not run)
```

---

simSD	<i>Simulate sd per age</i>
-------	----------------------------

---

**Description**

Simulate sd per age

**Usage**

```
simSD(age)
```

**Arguments**

age                    the age variable

**Value**

return predicted sd

**Examples**

```
## Not run:
x <- simSD(a)

## End(Not run)
```

---

simulateRasch

*Simulate raw test scores based on Rasch model*


---

**Description**

For testing purposes only: The function simulates raw test scores based on a virtual Rasch based test with  $n$  results per age group, an evenly distributed age variable,  $\text{items.n}$  test items with a simulated difficulty and standard deviation. The development trajectories over age group are modeled by a curve linear function of age, with at first fast progression, which slows down over age, and a slightly increasing standard deviation in order to model a scissor effects. The item difficulties can be accessed via  $\$theta$  and the raw data via  $\$data$  of the returned object.

**Usage**

```
simulateRasch(
  data = NULL,
  n = 100,
  minAge = 1,
  maxAge = 7,
  items.n = 21,
  items.m = 0,
  items.sd = 1,
  Theta = "random",
  width = 1
)
```

**Arguments**

<code>data</code>	data.frame from previous simulations for recomputation (overrides <code>n</code> , <code>minAge</code> , <code>maxAge</code> )
<code>n</code>	The sample size per age group
<code>minAge</code>	The minimum age (default 1)
<code>maxAge</code>	The maximum age (default 7)
<code>items.n</code>	The number of items of the test
<code>items.m</code>	The mean difficulty of the items
<code>items.sd</code>	The standard deviation of the item difficulty
<code>Theta</code>	irt scales difficulty parameters, either "random" for drawing a random sample, "even" for evenly distributed or a set of predefined values, which then overrides the <code>items.n</code> parameters

**width** The width of the window size for the continuous age per group; +- 1/2 width around group center on items.m and item.sd; if set to FALSE, the distribution is not drawn randomly but normally nonetheless

### Value

a list containing the simulated data and thetas

**data** the data.frame with only age, group and raw

**sim** the complete simulated data with item level results

**theta** the difficulty of the items

### Examples

```
## Not run:
# simulate data for a rather easy test (m = -1.0)
sim <- simulateRasch(n=150, minAge=1,
                    maxAge=7, items.n = 30, items.m = -1.0,
                    items.sd = 1, Theta = "random", width = 1.0)

# Show item difficulties
mean(sim$theta)
sd(sim$theta)
hist(sim$theta)

# Plot raw scores
boxplot(raw~group, data=sim$data)

# Model data
data <- prepareData(sim$data, age="age")
model <- bestModel(data, k = 4)
printSubset(model)
plotSubset(model, type=0)

## End(Not run)
```

---

standardize

*Standardize a numeric vector*

---

### Description

This function standardizes a numeric vector by subtracting the mean and dividing by the standard deviation. The resulting vector will have a mean of 0 and a standard deviation of 1.

### Usage

```
standardize(x)
```

**Arguments**

x                    A numeric vector to be standardized.

**Value**

A numeric vector of the same length as x, containing the standardized values.

**Examples**

```
## Not run:  
data <- c(1, 2, 3, 4, 5)  
standardized_data <- standardize(data)  
print(standardized_data)  
  
## End(Not run)
```

---

standardizeRakingWeights

*Function for standardizing raking weights Raking weights get divided by the smallest weight. Thereby, all weights become larger or equal to 1 without changing the ratio of the weights to each other.*

---

**Description**

Function for standardizing raking weights Raking weights get divided by the smallest weight. Thereby, all weights become larger or equal to 1 without changing the ratio of the weights to each other.

**Usage**

```
standardizeRakingWeights(weights)
```

**Arguments**

weights            Raking weights computed by computeWeights()

**Value**

the standardized weights

---

subsample_lm	<i>K-fold Resampled Coefficient Estimation for Linear Regression</i>
--------------	--

---

### Description

Performs k-fold resampling to estimate averaged coefficients for linear regression. The coefficients are averaged across k different subsets of the data to provide more stable estimates. For small samples ( $n < 100$ ), returns a standard linear model instead.

### Usage

```
subsample_lm(text, data, weights, k = 10)
```

### Arguments

text	A character string or formula specifying the model to be fitted
data	A data frame containing the variables in the model
weights	Optional numeric vector of weights. If NULL, unweighted regression is performed
k	Integer specifying the number of resampling folds (default = 10)

### Details

The function splits the data into k subsets, fits a linear model on k-1 subsets, and stores the coefficients. This process is repeated k times, and the final coefficients are averaged across all iterations to provide more stable estimates.

### Value

An object of class 'lm' with averaged coefficients from k-fold resampling. For small samples, returns a standard lm object.

---

summary.cnorm	<i>S3 method for printing the results and regression function of a cnorm model</i>
---------------	--

---

### Description

S3 method for printing the results and regression function of a cnorm model

### Usage

```
## S3 method for class 'cnorm'
summary(object, ...)
```

**Arguments**

object            A regression model or cnorm object  
 ...                additional parameters

**Value**

A report on the regression function, weights, R2 and RMSE

**See Also**

Other model: [bestModel\(\)](#), [checkConsistency\(\)](#), [cnorm.cv\(\)](#), [derive\(\)](#), [modelSummary\(\)](#), [print.cnorm\(\)](#), [printSubset\(\)](#), [rangeCheck\(\)](#), [regressionFunction\(\)](#)

---

summary.cnormBetaBinomial

*Summarize a Beta-Binomial Continuous Norming Model*

---

**Description**

This function provides a summary of a fitted beta-binomial continuous norming model, including model fit statistics, convergence information, and parameter estimates.

**Usage**

```
## S3 method for class 'cnormBetaBinomial'
summary(object, ...)
```

**Arguments**

object            An object of class "cnormBetaBinomial" or "cnormBetaBinomial2", typically the result of a call to [cnorm.betabinomial](#).

...                Additional arguments passed to the summary method:

- age An optional numeric vector of age values corresponding to the raw scores. If provided along with raw, additional fit statistics (R-squared, RMSE, bias) will be calculated.
- score An optional numeric vector of raw scores. Must be provided if age is given.
- weights An optional numeric vector of weights for each observation.

**Details**

The summary includes:

- Basic model information (type, number of observations, number of parameters)
- Model fit statistics (log-likelihood, AIC, BIC)

- R-squared, RMSE, and bias (if age and raw scores are provided) in comparison to manifest norm scores
- Convergence information
- Parameter estimates with standard errors, z-values, and p-values

### Value

Invisibly returns a list containing detailed diagnostic information about the model. The function primarily produces printed output summarizing the model.

### See Also

[cnorm.betabinomial](#), [diagnostics.betabinomial](#)

### Examples

```
## Not run:
model <- cnorm.betabinomial(ppvt$age, ppvt$raw, n = 228)
summary(model)

# Including R-squared, RMSE, and bias in the summary:
summary(model, age = ppvt$age, score = ppvt$raw)

## End(Not run)
```

---

```
summary.cnormBetaBinomial2
```

*Summarize a Beta-Binomial Continuous Norming Model*

---

### Description

This function provides a summary of a fitted beta-binomial continuous norming model, including model fit statistics, convergence information, and parameter estimates.

### Usage

```
## S3 method for class 'cnormBetaBinomial2'
summary(object, ...)
```

### Arguments

- |        |  |
|--------|--|
| object | An object of class "cnormBetaBinomial" or "cnormBetaBinomial2", typically the result of a call to <a href="#">cnorm.betabinomial</a> .   |
| ...    | Additional arguments passed to the summary method: <ul style="list-style-type: none"> <li>• age An optional numeric vector of age values corresponding to the raw scores. If provided along with raw, additional fit statistics (R-squared, RMSE, bias) will be calculated.</li> </ul> |

- score An optional numeric vector of raw scores. Must be provided if age is given.
- weights An optional numeric vector of weights for each observation.

### Details

The summary includes:

- Basic model information (type, number of observations, number of parameters)
- Model fit statistics (log-likelihood, AIC, BIC)
- R-squared, RMSE, and bias (if age and raw scores are provided) in comparison to manifest norm scores
- Convergence information
- Parameter estimates with standard errors, z-values, and p-values

### Value

Invisibly returns a list containing detailed diagnostic information about the model. The function primarily produces printed output summarizing the model.

### See Also

[cnorm.betabinomial](#), [diagnostics.betabinomial](#)

### Examples

```
## Not run:
model <- cnorm.betabinomial(ppvt$age, ppvt$raw, n = 228)
summary(model)

# Including R-squared, RMSE, and bias in the summary:
summary(model, age = ppvt$age, raw = ppvt$raw)

## End(Not run)
```

---

summary.cnormShash      *Summarize a SinH-ArcSinH Continuous Norming Model*

---

### Description

This function provides a summary of a fitted SinH-ArcSinH (shash) continuous norming model, including model fit statistics, convergence information, and parameter estimates.

### Usage

```
## S3 method for class 'cnormShash'
summary(object, ...)
```

## Arguments

- object      An object of class "cnormShash", typically the result of a call to [cnorm.shash](#).
- ...      Additional arguments passed to the summary method:
- age An optional numeric vector of age values corresponding to the raw scores. If provided along with score, additional fit statistics (R-squared, RMSE, bias) will be calculated.
  - score An optional numeric vector of raw scores. Must be provided if age is given.
  - weights An optional numeric vector of weights for each observation.

## Details

The summary includes:

- Basic model information (polynomial degrees, delta parameter, number of observations)
- Model fit statistics (log-likelihood, AIC, BIC)
- R-squared, RMSE, and bias (if age and raw scores are provided)
- Convergence information
- Parameter estimates with standard errors, z-values, and p-values

## Value

Invisibly returns a list containing detailed diagnostic information about the model. The function primarily produces printed output summarizing the model.

## See Also

[cnorm.shash](#), [diagnostics.shash](#)

## Examples

```
## Not run:
model <- cnorm.shash(children$age, children$score)
summary(model)

# Including R-squared, RMSE, and bias in the summary:
summary(model, age = children$age, score = children$score)

## End(Not run)
```

---

taylorSwift	<i>Swiftly compute Taylor regression models for distribution free continuous norming</i>
-------------	--

---

### Description

Conducts distribution free continuous norming and aims to find a fitting model. Raw data are modelled as a Taylor polynomial of powers of age and location and their interactions. In addition to the raw scores, either provide a numeric vector for the grouping information (`group`) for the ranking of the raw scores. You can adjust the grade of smoothing of the regression model by setting the `k`, `t` and `terms` parameter. In general, increasing `k` and `t` leads to a higher fit, while lower values lead to more smoothing. If both parameters are missing, `taylorSwift` uses `k = 5` and `t = 3` by default.

### Usage

```
taylorSwift(
  raw = NULL,
  group = NULL,
  age = NULL,
  width = NA,
  weights = NULL,
  scale = "T",
  method = 4,
  descend = FALSE,
  k = NULL,
  t = NULL,
  terms = 0,
  R2 = NULL,
  plot = TRUE,
  extensive = TRUE,
  subsampling = TRUE
)
```

### Arguments

<code>raw</code>	Numeric vector of raw scores
<code>group</code>	Numeric vector of grouping variable, e. g. grade. If no group or age variable is provided, conventional norming is applied
<code>age</code>	Numeric vector with chronological age, please additionally specify width of window
<code>width</code>	Size of the sliding window in case an age vector is used
<code>weights</code>	Vector or variable name in the dataset with weights for each individual case. It can be used to compensate for moderate imbalances due to insufficient norm data stratification. Weights should be numerical and positive.

scale	type of norm scale, either T (default), IQ, z or percentile (= no transformation); a double vector with the mean and standard deviation can as well, be provided f. e. <code>c(10, 3)</code> for Wechsler scale index points
method	Ranking method in case of bindings, please provide an index, choosing from the following methods: 1 = Blom (1958), 2 = Tukey (1949), 3 = Van der Warden (1952), 4 = Rankit (default), 5 = Levenbach (1953), 6 = Filliben (1975), 7 = Yu & Huang (2001)
descend	ranking order (default <code>descend = FALSE</code> ): inverses the ranking order with higher raw scores getting lower norm scores; relevant for example when norming error scores, where lower scores mean higher performance
k	The power constant. Higher values result in more detailed approximations but have the danger of over-fit ( <code>max = 6</code> ). If not set, it uses <code>t</code> and if both parameters are <code>NULL</code> , <code>k</code> is set to 5.
t	The age power parameter ( <code>max = 6</code> ). If not set, it uses <code>k</code> and if both parameters are <code>NULL</code> , <code>k</code> is set to 3, since age trajectories are most often well captured by cubic polynomials.
terms	Selection criterion for model building. The best fitting model with this number of terms is used
R2	Adjusted R square as a stopping criterion for the model building (default <code>R2 = 0.99</code> )
plot	Default <code>TRUE</code> ; plots the regression model and prints report
extensive	If <code>TRUE</code> , screen models for consistency and - if possible, exclude inconsistent ones
subsampling	If <code>TRUE</code> (default), model coefficients are calculated using 10-folds and averaged across the folds. This produces more robust estimates with a slight increase in bias.

### Value

`cnorm` object including the ranked raw data and the regression model

### References

1. Gary, S. & Lenhard, W. (2021). In norming we trust. *Diagnostica*.
2. Gary, S., Lenhard, W. & Lenhard, A. (2021). Modelling Norm Scores with the `cNORM` Package in R. *Psych*, 3(3), 501-521. <https://doi.org/10.3390/psych3030033>
3. Lenhard, A., Lenhard, W., Suggate, S. & Segerer, R. (2016). A continuous solution to the norming problem. *Assessment*, Online first, 1-14. doi:10.1177/1073191116656437
4. Lenhard, A., Lenhard, W., Gary, S. (2018). Continuous Norming (`cNORM`). The Comprehensive R Network, Package `cNORM`, available: <https://CRAN.R-project.org/package=cNORM>
5. Lenhard, A., Lenhard, W., Gary, S. (2019). Continuous norming of psychometric tests: A simulation study of parametric and semi-parametric approaches. *PLoS ONE*, 14(9), e0222279. doi:10.1371/journal.pone.0222279
6. Lenhard, W., & Lenhard, A. (2020). Improvement of Norm Score Quality via Regression-Based Continuous Norming. *Educational and Psychological Measurement(Online First)*, 1-33. <https://doi.org/10.1177/0013164420928457>

**See Also**

rankByGroup, rankBySlidingWindow, computePowers, bestModel

**Examples**

```
## Not run:
# Using this function with the example dataset 'ppvt'
# You can use the 'getGroups()' function to set up grouping variable in case,
# you have a continuous age variable.
model <- taylorSwift(raw = ppvt$raw, group = ppvt$group)

# return norm tables including 90% confidence intervals for a
# test with a reliability of r = .85; table are set to mean of quartal
# in grade 3 (children completed 2 years of schooling)
normTable(c(5, 15), model, CI = .90, reliability = .95)

# ... or instead of raw scores for norm scores, the other way round
rawTable(c(8, 12), model, CI = .90, reliability = .95)

## End(Not run)
```

---

weighted.quantile	<i>Weighted quantile estimator</i>
-------------------	------------------------------------

---

**Description**

Computes weighted quantiles (code from Andrey Akinshin (2023) "Weighted quantile estimators" arXiv:2304.07265 [stat.ME] Code made available via the CC BY-NC-SA 4.0 license) on the basis of either the weighted Harrell-Davis quantile estimator or an adaption of the type 7 quantile estimator of the generic quantile function in the base package. Please provide a vector with raw values, the probabilities for the quantiles and an additional vector with the weight of each observation. In case the weight vector is NULL, a normal quantile estimation is done. The vectors may not include NAs and the weights should be positive non-zero values. Please draw on the computeWeights() function for retrieving weights in post stratification.

**Usage**

```
weighted.quantile(x, probs, weights = NULL, type = "Harrell-Davis")
```

**Arguments**

x	A numerical vector
probs	Numerical vector of quantiles
weights	A numerical vector with weights; should have the same length as x

`type` Type of estimator, can either be "inflation", "Harrell-Davis" using a beta function to approximate the weighted percentiles (Harrell & Davis, 1982) or "Type7" (default; Hyndman & Fan, 1996), an adaption of the generic quantile function in R, including weighting. The inflation procedure is essentially a numerical, non-parametric solution that gives the same results as Harrel-Davis. It requires less resources with small datasets and always finds a solution (e. g. 1000 cases with weights between 1 and 10). If it becomes too resource intense, it switches to Harrell-Davis automatically. Harrel-Davis and Type7 code is based on the work of Akinshin (2023).

### Value

the weighted quantiles

### References

1. Harrell, F.E. & Davis, C.E. (1982). A new distribution-free quantile estimator. *Biometrika*, 69(3), 635-640.
2. Hyndman, R. J. & Fan, Y. (1996). Sample quantiles in statistical packages, *American Statistician* 50, 361–365.
3. Akinshin, A. (2023). Weighted quantile estimators arXiv:2304.07265 [stat.ME]

### See Also

weighted.quantile.inflation, weighted.quantile.harrell.davis, weighted.quantile.type7

---

weighted.quantile.harrell.davis

*Weighted Harrell-Davis quantile estimator*

---

### Description

Computes weighted quantiles; code from Andrey Akinshin (2023) "Weighted quantile estimators" arXiv:2304.07265 [stat.ME] Code made available via the CC BY-NC-SA 4.0 license

### Usage

```
weighted.quantile.harrell.davis(x, probs, weights = NULL)
```

### Arguments

<code>x</code>	A numerical vector
<code>probs</code>	Numerical vector of quantiles
<code>weights</code>	A numerical vector with weights; should have the same length as <code>x</code> . If no weights are provided (NULL), it falls back to the base quantile function, type 7

**Value**

the quantiles

---

weighted.quantile.inflation

*Weighted quantile estimator through case inflation*

---

**Description**

Applies weighted ranking numerically by inflating cases according to weight. This function will be resource intensive, if inflated cases get too high and in this cases, it switches to the parametric Harrell-Davis estimator.

**Usage**

```
weighted.quantile.inflation(
  x,
  probs,
  weights = NULL,
  degree = 3,
  cutoff = 1e+07
)
```

**Arguments**

x	A numerical vector
probs	Numerical vector of quantiles
weights	A numerical vector with weights; should have the same length as x.
degree	power parameter for case inflation (default = 3, equaling factor 1000) If no weights are provided (NULL), it falls back to the base quantile function, type 7
cutoff	stop criterion for the sum of standardized weights to switch to Harrell-Davis, default = 1000000

**Value**

the quantiles

---

 weighted.quantile.type7

*Weighted type7 quantile estimator*


---

**Description**

Computes weighted quantiles; code from Andrey Akinshin (2023) "Weighted quantile estimators" arXiv:2304.07265 [stat.ME] Code made available via the CC BY-NC-SA 4.0 license

**Usage**

```
weighted.quantile.type7(x, probs, weights = NULL)
```

**Arguments**

x	A numerical vector
probs	Numerical vector of quantiles
weights	A numerical vector with weights; should have the same length as x. If no weights are provided (NULL), it falls back to the base quantile function, type 7

**Value**

the quantiles

---

weighted.rank

*Weighted rank estimation*


---

**Description**

Conducts weighted ranking on the basis of sums of weights per unique raw score. Please provide a vector with raw values and an additional vector with the weight of each observation. In case the weight vector is NULL, a normal ranking is done. The vectors may not include NAs and the weights should be positive non-zero values.

**Usage**

```
weighted.rank(x, weights = NULL)
```

**Arguments**

x	A numerical vector
weights	A numerical vector with weights; should have the same length as x

**Value**

the weighted absolute ranks

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