

# Package ‘PROsetta’

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**Type** Package

**Title** Linking Patient-Reported Outcomes Measures

**Version** 0.4.1

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**Description** Perform scale linking to establish relationships between instruments that measure similar constructs according to the PROsetta Stone methodology, as in Choi, Schalet, Cook, & Cella (2014) <[doi:10.1037/a0035768](https://doi.org/10.1037/a0035768)>.

**URL** <https://www.prosettastone.org/> (project description),  
<https://choi-phd.github.io/PROsetta/> (documentation)

**BugReports** <https://github.com/choi-phd/PROsetta/issues>

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**Author** Seung W. Choi [aut, cre] (<<https://orcid.org/0000-0003-4777-5420>>),  
 Sangdon Lim [aut] (<<https://orcid.org/0000-0002-2988-014X>>),  
 Benjamin D. Schalet [ctb],  
 Aaron J. Kaat [ctb],  
 David Cella [ctb]

**Maintainer** Seung W. Choi <[schoi@austin.utexas.edu](mailto:schoi@austin.utexas.edu)>

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checkFrequency	<i>Check frequency table for unobserved response categories</i>
----------------	---

---

## Description

`checkFrequency` is a descriptive function for checking whether all response categories in a frequency table have a frequency of at least 1.

**Usage**

```
checkFrequency(data)
```

**Arguments**

`data` a PROsetta\_data object. See [loadData](#) for loading a dataset.

**Value**

`checkFrequency` returns TRUE if all response categories have a frequency of at least 1, and FALSE if not.

**Examples**

```
checkFrequency(data_asq) # TRUE

## Not run:
data_asq$response$EDANX01[data_asq$response$EDANX01 == 4] <- 3
checkFrequency(data_asq) # FALSE

## End(Not run)
```

---

compareScores

*Compare two sets of scores*

---

**Description**

`compareScores` is a helper function for comparing two sets of scores.

**Usage**

```
compareScores(left, right, type = c("corr", "mean", "sd", "rmsd", "mad"))
```

**Arguments**

`left` scores on the left side of comparison.  
`right` scores on the right side of comparison. This is subtracted from 'left'.  
`type` type of comparisons to include. Accepts corr, mean, sd, rmsd, mad. Defaults to all types.

**Value**

`compareScores` returns a `data.frame` containing the comparison results.

## Examples

```
set.seed(1)
true_theta <- rnorm(100)
theta_est <- true_theta + rnorm(100, 0, 0.3)
compareScores(theta_est, true_theta)
```

dataset\_asq

*ASQ dataset*

## Description

This dataset is associated with the following objects:

### Details

- [response\\_asq](#) a `data.frame` containing raw response data of 751 participants and 41 variables.
  - prosettaid participant IDs.
  - EDANX01 -- MASQ11 response to items.
- [itemmap\\_asq](#) a `data.frame` containing the item map, describing the items in each instrument.
  - item\_order item numeric IDs. This column refers to the column `item_order` in anchor item attributes.
  - instrument the instrument ID that each item belongs to.
  - item\_id item ID strings. This column refers to column names in raw response data, excluding the participant ID column.
  - item\_name new item ID strings to be used in the combined scale.
  - ncat the number of response categories.
- [anchor\\_asq](#) a `data.frame` containing anchor item parameters for 29 items.
  - item\_order item numeric IDs.
  - item\_id item ID strings. This column refers to column names in raw response data, excluding the participant ID column.
  - a the discrimination (slope) parameter for the graded response model.
  - cb1 – cb4 the boundaries between each category-pair for the graded response model.
- [data\\_asq](#) a `PROsetta_data` object containing the datasets above. See `loadData` for creating `PROsetta_data` objects.

## Examples

```
## load datasets into a PROsetta_data object
data_asq <- loadData(
  response = response_asq,
  itemmap  = itemmap_asq,
  anchor   = anchor_asq
```

```
)  
  
## run descriptive statistics  
runDescriptive(data_asq)  
  
## run item parameter calibration on the response data, linking to the anchor item parameters  
runLinking(data_asq, method = "FIXEDPAR")
```

---

dataset\_dep

*DEP dataset*

---

## Description

This dataset is associated with the following objects:

## Details

- [response\\_dep](#) a `data.frame` containing raw response data of 747 participants and 49 variables.
  - prosettaid participant IDs.
  - EDDEP04 -- CESD20 response to items.
- [itemmap\\_dep](#) a `data.frame` containing the item map, describing the items in each instrument.
  - item\_order item numeric IDs. This column refers to the column `item_order` in anchor item parameters.
  - instrument the instrument ID that each item belongs to.
  - item\_id item ID strings. This column refers to column names in raw response data, excluding the participant ID column.
  - item\_name new item ID strings to be used in the combined scale.
  - ncat the number of response categories.
- [anchor\\_dep](#) a `data.frame` containing anchor item parameters for 28 items.
  - item\_order item numeric IDs.
  - item\_id item ID strings. This column refers to column names in raw response data, excluding the participant ID column.
  - a the discrimination (slope) parameter for the graded response model.
  - cb1 – cb4 the boundaries between each category-pair for the graded response model.
- [data\\_dep](#) a `PROsetta_data` object containing the datasets above. See `loadData` for creating `PROsetta_data` objects.

## Examples

```
## load datasets into a PROsetta_data object  
data_dep <- loadData(  
  response = response_dep,  
  itemmap = itemmap_dep,  
  anchor = anchor_dep
```

```
6                                         getEscore  
  
)  
  
## run descriptive statistics  
runDescriptive(data_dep)  
  
## run item parameter calibration on the response data, linking to the anchor item parameters  
runLinking(data_dep, method = "FIXEDPAR")
```

---

**getCompleteData** *Get complete data*

---

### Description

`getCompleteData` is a helper function for performing casewise deletion of missing values.

### Usage

```
getCompleteData(data, scale = NULL, verbose = FALSE)
```

### Arguments

- |         |   |
|---------|---|
| data    | a <code>PROsetta_data</code> object.  |
| scale   | the index of the scale to perform casewise deletion. Leave empty or set to "combined" to perform on all scales. |
| verbose | if TRUE, print status messages. (default = FALSE)   |

### Value

`getCompleteData` returns an updated `PROsetta_data` object.

### Examples

```
d <- getCompleteData(data_asq, verbose = TRUE)
```

---

**getEscore** *Calculate expected scores at theta*

---

### Description

`getEscore` is a helper function for obtaining expected scores at supplied thetas.

### Usage

```
getEscore(ipar, model, theta, is_minscore_0)
```

**Arguments**

ipar	item parameters.
model	item model to use.
theta	theta values.
is_minscore_0	if TRUE the score begins from 0 instead of 1.

**Value**

`getEscore` returns a vector of expected scores.

---

`getItemNames`*Get item names***Description**

`getItemNames` is a helper function for extracting item names for a specified scale from a `PROsetta_data` object.

**Usage**

```
getItemNames(d, scale_id)
```

**Arguments**

d	a <code>PROsetta_data</code> object.
scale_id	scale IDs to extract item names.

**Value**

`getItemNames` returns a vector containing item names.

**Examples**

```
getItemNames(data_asq, 1)
getItemNames(data_asq, 2)
getItemNames(data_asq, c(1, 2))
getItemNames(data_asq, c(2, 1))
```

<code>getResponse</code>	<i>Extract scale-wise response</i>
--------------------------	------------------------------------

## Description

`getResponse` is a helper function for extracting scale-wise response from a `PROsetta_data` object.

## Usage

```
getResponse(d, scale_id = "all", person_id = FALSE)
```

## Arguments

d	a <code>PROsetta_data</code> object.
scale_id	scale IDs to extract response. If all, use all scale IDs. (default = all)
person_id	if TRUE, also return person IDs. (default = FALSE)

## Value

`getResponse` returns a `data.frame` containing scale-wise response.

## Examples

```
getResponse(data_asq)
getResponse(data_asq, 1)
getResponse(data_asq, 2)
getResponse(data_asq, c(1, 2))
getResponse(data_asq, c(2, 1))
getResponse(data_asq, c(1, 2), person_id = TRUE)
```

<code>getRSSS</code>	<i>Compute a Crosswalk Table</i>
----------------------	----------------------------------

## Description

`getRSSS` is a function for generating a raw-score to standard-score crosswalk table.

## Usage

```
getRSSS(ipar, theta_grid, is_minscore_0, prior_mu_sigma)
```

## Arguments

- `ipar` an item parameter matrix for graded response items. Accepts both a/b and a/d format parameters. Accepts multidimensional item parameters.
- `theta_grid` the theta grid to use for numerical integration.
- `is_minscore_0` if TRUE, the score of each item begins from 0. if FALSE, the score of each item begins from 1.
- `prior_mu_sigma` a named list containing prior distribution parameters. All values must be in the theta metric.
- `mu` the prior means
  - `sigma` the covariance matrix
  - `sd` the prior standard deviations
  - `corr` the correlation matrix

## Examples

```
## Free calibration without using anchor

o <- runCalibration(data_asq, technical = list(NCYCLES = 1000))

ipar <- mirt::coef(o, IRTpars = TRUE, simplify = TRUE)$items
items <- getItemNames(data_asq, 2)

getRSSS(
  ipar = ipar[items, ],
  theta_grid = seq(-4, 4, .1),
  is_minscore_0 = TRUE,
  prior_mu_sigma = list(mu = 0, sigma = 1)
)
```

## getScaleSum

*Calculate raw sum scores of a scale*

## Description

`getScaleSum` is a helper function for calculating instrument-wise raw sum scores from response data.

## Usage

```
getScaleSum(data, scale_idx)
```

## Arguments

- `data` a `PROsetta_data` object.
- `scale_idx` the instrument index to obtain the raw sum scores.

## Examples

```
getScaleSum(data_asq, 1)
getScaleSum(data_asq, 2)
```

<code>getTheta</code>	<i>Obtain theta estimates</i>
-----------------------	-------------------------------

## Description

`getTheta` is a helper function for obtaining theta estimates. Estimates are obtained using an *expected a posteriori* (EAP) method.

## Usage

```
getTheta(
  data,
  ipar,
  scale = "combined",
  model = "grm",
  theta_grid = seq(-4, 4, 0.1),
  prior_dist = "normal",
  prior_mean = 0,
  prior_sd = 1
)
```

## Arguments

<code>data</code>	a <code>PROsetta_data</code> object.
<code>ipar</code>	a <code>data.frame</code> containing item parameters.
<code>scale</code>	the index of the scale to use. <code>combined</code> refers to the combined scale. (default = <code>combined</code> )
<code>model</code>	the item model to use. Accepts <code>grm</code> or <code>gpcm</code> . (default = <code>grm</code> )
<code>theta_grid</code>	the theta grid to use for numerical integration. (default = <code>seq(-4, 4, .1)</code> )
<code>prior_dist</code>	the type of prior distribution. Accepts <code>normal</code> or <code>logistic</code> . (default = <code>normal</code> )
<code>prior_mean</code>	mean of the prior distribution. (default = <code>0.0</code> )
<code>prior_sd</code>	SD of the prior distribution. (default = <code>1.0</code> )

## Value

`getTheta` returns a `list` containing EAP estimates.

## Examples

```
x <- runLinking(data_asq, method = "FIXEDPAR")

o <- getTheta(data_asq, x$ipar_linked, scale = 1)
o$theta
o$item_idx

o <- getTheta(data_asq, x$ipar_linked, scale = 2)
o$theta
o$item_idx

o <- getTheta(data_asq, x$ipar_linked, scale = "combined")
o$theta
o$item_idx
```

**loadData**

*Load data from supplied config*

## Description

**loadData** is a data loading function for creating a [PROsetta\\_data](#) object, for performing scale linking/equating in the 'PROsetta' package. **loadData** assumes the response data has been reverse-coded for applicable items.

## Usage

```
loadData(
  response,
  itemmap,
  anchor,
  item_id = NULL,
  person_id = NULL,
  scale_id = NULL,
  input_dir = getwd()
)
```

## Arguments

<b>response</b>	response data containing case IDs and item responses. This can be a .csv filename or a <a href="#">data.frame</a> object.
<b>itemmap</b>	an item map containing item IDs and scale IDs. This can be a .csv filename or a <a href="#">data.frame</a> object.
<b>anchor</b>	anchor data containing item parameters for anchor items. This can be a .csv filename or a <a href="#">data.frame</a> object.
<b>item_id</b>	the column name to look for item IDs. Automatically determined if not specified.

<code>person_id</code>	the column name to look for case IDs. Automatically determined if not specified.
<code>scale_id</code>	the column name to look for scale IDs. Automatically determined if not specified.
<code>input_dir</code>	the directory to look for the files.

**Value**

`loadData` returns a `PROsetta_data` object containing the loaded data.

`plot,PROsetta_data,ANY-method`

*Plot frequency distribution*

**Description**

This is an extension of `plot` to visualize frequency distribution from `PROsetta_data` object.

**Usage**

```
## S4 method for signature 'PROsetta_data,ANY'
plot(
  x,
  y,
  scale_id = "combined",
  filename = NULL,
  title = NULL,
  xlim = NULL,
  color = "blue",
  nbar = 20,
  rug = FALSE,
  filetype = "pdf",
  savefile = FALSE,
  bg = "white",
  width = 6,
  height = 6,
  pointsize = 12
)
```

**Arguments**

<code>x</code>	a <code>PROsetta_data</code> object.
<code>y</code>	unused argument, exists for compatibility with <code>plot</code> in the base R package.
<code>scale_id</code>	scale ID to plot. <code>combined</code> represents the combined scale. (default = <code>combined</code> )
<code>filename</code>	the filename to write if <code>savefile</code> argument is TRUE.

title	the title of the figure.
xlim	the range of scores to plot.
color	the color to fill the histogram. (default = blue)
nbar	the number of histogram bars. (default = 20)
rug	if TRUE, display the actual distribution of scores below each bar. (default = FALSE)
filetype	the type of file to write if savefile argument is TRUE. Accepts pdf, jpeg, png, and tiff. (default = pdf)
savefile	if TRUE, save the figure as a file. (default = FALSE)
bg	the background color to use when saving the figure as a file. (default = white)
width	the width of the plot. (default = 6)
height	the height of the plot. (default = 6)
pointsize	the point size to use when saving the figure as a file. (default = 12)

## Examples

```
plot(data_asq)
plot(data_asq, scale_id = 1)
plot(data_asq, scale_id = 2)
```

**plotInfo**

*Plot scale information*

## Description

`plotInfo` is a plotting function to visualize scale-level information.

## Usage

```
plotInfo(
  object,
  data,
  theta = seq(-4, 4, 0.1),
  t_score = FALSE,
  scale_label = c(1, 2, "Combined"),
  color = c("red", "blue", "black"),
  lty = c(3, 2, 1)
)

## S4 method for signature 'SingleGroupClass'
plotInfo(
  object,
  data,
```

```

theta = seq(-4, 4, 0.1),
t_score = FALSE,
scale_label = c(1, 2, "Combined"),
color = c("red", "blue", "black"),
lty = c(3, 2, 1)
)

```

### Arguments

object	a <a href="#">SingleGroupClass</a> object from <a href="#">runCalibration</a> .
data	a <a href="#">PROsetta_data</a> object.
theta	(optional) theta values to plot on the x-axis. (default = <code>seq(-4, 4, .1)</code> )
t_score	(optional) set to TRUE to plot T-scores on the x-axis instead of thetas. (default = FALSE)
scale_label	(optional) names of each scale. This must have three values. (default = <code>c(1, 2, "Combined")</code> )
color	(optional) line colors to plot. This must have three values. (default = <code>c("red", "blue", "black")</code> )
lty	(optional) line types to plot. This must have three values. (default = <code>c(3, 2, 1)</code> )

### Examples

```

o <- runCalibration(data_asq, technical = list(NCYCLES = 1000))
plotInfo(o, data_asq)

```

### Description

[PROsetta](#) is a caller function for launching a Shiny app locally.

### Usage

```
PROsetta()
```

### Examples

```

if (interactive()) {
  PROsetta()
}

```

---

<code>runCalibration</code>	<i>Run Calibration</i>
-----------------------------	------------------------

---

## Description

`runCalibration` is a function for performing item parameter calibration on the response data.

## Usage

```
runCalibration(
  data,
  dimensions = 1,
  fix_method = "free",
  fixedpar = NULL,
  ignore_nonconv = FALSE,
  verbose = FALSE,
  ...
)
```

## Arguments

<code>data</code>	a PR0setta_data object. See <code>loadData</code> for loading a dataset.
<code>dimensions</code>	the number of dimensions to use. Must be 1 or 2. If 1, use one underlying dimension for all instruments combined. If 2, use each dimension separately for the anchor instrument and the developing instrument. Covariance between dimensions is freely estimated. (default = 1)
<code>fix_method</code>	the type of constraints to impose. (default = <code>free</code> ) <ul style="list-style-type: none"> <li>• <code>item</code> for fixed parameter calibration using anchor item parameters</li> <li>• <code>theta</code> for using the mean and the variance obtained from a unidimensional calibration of anchor items</li> <li>• <code>free</code> for free calibration</li> </ul>
<code>fixedpar</code>	this argument exists for backward compatibility. TRUE is equivalent to <code>fix_method = "item"</code> , and FALSE is equivalent to <code>fix_method = "free"</code> .
<code>ignore_nonconv</code>	if TRUE, return results even when calibration does not converge. If FALSE, raise an error when calibration does not converge. (default = FALSE)
<code>verbose</code>	if TRUE, print status messages. (default = FALSE)
...	additional arguments to pass onto <code>mirt</code> in ' <code>mirt</code> ' package.

## Value

`runCalibration` returns a `SingleGroupClass` object containing item calibration results.

This object can be used in `coef`, `itemfit`, `itemplot` in '`mirt`' package to extract wanted information.

## Examples

```
## Not run:
out_calib <- runCalibration(data_asq) # errors

## End(Not run)

out_calib <- runCalibration(data_asq, technical = list(NCYCLES = 1000))

mirt::coef(out_calib, IRTpars = TRUE, simplify = TRUE)
mirt::itemfit(out_calib, empirical.plot = 1)
mirt::itemplot(out_calib, item = 1, type = "info")
mirt::itemfit(out_calib, "S_X2", na.rm = TRUE)
```

**runCFA**

*Run a confirmatory factor analysis*

## Description

**runCFA** is a function for performing a one-factor confirmatory factor analysis (CFA) to test unidimensionality.

## Usage

```
runCFA(data, estimator = "WLSMV", std.lv = TRUE, scalewise = FALSE, ...)
```

## Arguments

<code>data</code>	a <a href="#">PROsetta_data</a> object. See <a href="#">loadData</a> for loading a dataset.
<code>estimator</code>	the estimator to be used. Passed onto <code>cfa</code> in ' <a href="#">lavaan</a> ' package. (default = WLSMV)
<code>std.lv</code>	if TRUE, the metric of the latent variable is determined by fixing their (residual) variances to 1.0. If FALSE, the metric of each latent variable is determined by fixing the factor loading of the first indicator to 1.0. Passed onto <code>cfa</code> . (default = TRUE)
<code>scalewise</code>	if TRUE, run analysis for each instrument as well as for the combined instrument. If FALSE, run analysis only for the combined instrument. (default = FALSE)
<code>...</code>	additional arguments to pass onto <code>cfa</code> .

## Value

**runCFA** returns a list containing the CFA results.

## Examples

```
out_cfa <- runCFA(data_asq, scalewise = TRUE)
lavaan::summary(out_cfa$`1`      , fit.measures = TRUE, standardized = TRUE, estimates = FALSE)
lavaan::summary(out_cfa$`2`      , fit.measures = TRUE, standardized = TRUE, estimates = FALSE)
lavaan::summary(out_cfa$`combined`, fit.measures = TRUE, standardized = TRUE, estimates = FALSE)
```

---

runClassical	<i>Run CTT-based reliability analysis</i>
--------------	---

---

### Description

`runClassical` is a function for performing a Classical Test Theory (CTT) based reliability analysis.

### Usage

```
runClassical(data, omega = FALSE, scalewise = TRUE, ...)
```

### Arguments

data	a <code>PROsetta_data</code> object. See <code>loadData</code> for loading a dataset.
omega	if TRUE, also obtain McDonald's omega using <code>omega</code> in <code>psych</code> package. (default = FALSE)
scalewise	if TRUE, run analysis for each instrument as well as for the combined instrument. If FALSE, run analysis only for the combined instrument. (default = TRUE)
...	additional arguments to pass onto <code>omega</code> .

### Value

`runClassical` returns a `list` containing reliability analysis results.

### Examples

```
out_alpha <- runClassical(data_asq)
out_omega <- runClassical(data_asq, omega = TRUE) # also obtain omega
```

---

---

runDescriptive	<i>Obtain a descriptive statistics table</i>
----------------	--

---

### Description

`runDescriptive` is a descriptive function for obtaining descriptive statistics for each item in the dataset.

### Usage

```
runDescriptive(data = NULL)
```

### Arguments

data	a <code>PROsetta_data</code> object. See <code>loadData</code> for loading a dataset.
------	---

**Value**

`runDescriptive` returns a `data.frame` containing descriptive statistics (mean, standard deviation, median, ...) of the items in the dataset. These are calculated with `describe` in 'psych' package.

**Examples**

```
out_desc <- runDescriptive(data_asq)
```

<code>runEquateObserved</code>	<i>Run Test Equating</i>
--------------------------------	--------------------------

**Description**

`runEquateObserved` is a function for performing equipercentile equating between two scales. `runEquateObserved` also produces a concordance table, mapping the observed raw scores from one scale to the scores from another scale.

**Usage**

```
runEquateObserved(
  data,
  scale_from = 2,
  scale_to = 1,
  type_to = "raw",
  rsss = NULL,
  eq_type = "equipercentile",
  smooth = "loglinear",
  degrees = list(3, 1),
  boot = TRUE,
  reps = 100,
  verbose = FALSE,
  ...
)
```

**Arguments**

- `data` a `PROsetta_data` object. See `loadData` for loading a dataset.
- `scale_from` the scale ID of the input scale. References to `itemmap` in `data` argument. (default = 2)
- `scale_to` the scale ID of the target scale to equate to. References to `itemmap` in `data` argument. (default = 1)
- `type_to` the type of score to use in the target scale frequency table. Accepts `raw`, `tscore`, and `theta`. `tscore` and `theta` require argument `rsss` to be supplied. (default = `raw`)

<code>rsss</code>	the RSSS table to use to map each raw score level onto a t-score or a theta. See <a href="#">runRSSS</a> .
<code>eq_type</code>	the type of equating to be passed onto <code>equate</code> in ' <code>equate</code> ' package. (default = <code>equipercentile</code> )
<code>smooth</code>	the type of smoothing method to be passed onto <code>presmoothing</code> in ' <code>equate</code> ' package. (default = <code>loglinear</code> )
<code>degrees</code>	the degrees of smoothing to be passed onto <code>presmoothing</code> . (default = <code>list(3, 1)</code> )
<code>boot</code>	performs bootstrapping if TRUE. (default = TRUE)
<code>reps</code>	the number of replications to perform in bootstrapping. (default = 100)
<code>verbose</code>	if TRUE, print status messages. (default = FALSE)
<code>...</code>	other arguments to pass onto <code>equate</code> .

### Value

`runEquateObserved` returns an `equate` object containing the test equating result.

The printed summary statistics indicate the distributional properties of the two supplied scales and the equated scale.

- `x` corresponds to `scale_from`.
- `y` corresponds to `scale_to`.
- `yx` corresponds to `scale_from` after equating to `scale_to`.

See `equate` for details.

The concordance table is stored in `concordance` slot.

### Examples

```
out_eq_raw <- runEquateObserved(data_asq,
  scale_to = 1, scale_from = 2,
  eq_type = "equipercentile", smooth = "loglinear"
)
out_eq_raw$concordance

out_link <- runLinking(data_asq, method = "FIXEDPAR")
out_rsss <- runRSSS(data_asq, out_link)
out_eq_tscore <- runEquateObserved(data_asq,
  scale_to = 1, scale_from = 2,
  type_to = "tscore", rsss = out_rsss,
  eq_type = "equipercentile", smooth = "loglinear"
)
out_eq_tscore$concordance
```

<code>runFrequency</code>	<i>Obtain a frequency table</i>
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## Description

`runFrequency` is a descriptive function for obtaining a frequency table from the dataset.

## Usage

```
runFrequency(data, check_frequency = TRUE)
```

## Arguments

<code>data</code>	a <a href="#">PROsetta_data</a> object. See <a href="#">loadData</a> for loading a dataset.
<code>check_frequency</code>	if TRUE, check the frequency table for missing response categories, and display warning message if any is missing. (default = TRUE)

## Value

`runFrequency` returns a [data.frame](#) containing the frequency table.

## Examples

```
freq_asq <- runFrequency(data_asq)
freq_dep <- runFrequency(data_dep)
```

<code>runLinking</code>	<i>Run Scale Linking</i>
-------------------------	--------------------------

## Description

`runLinking` is a function for obtaining item parameters from the response data in the metric of supplied anchor item parameters.

## Usage

```
runLinking(data, method, verbose = FALSE, ...)
```

## Arguments

data	a <a href="#">PROsetta_data</a> object. See <a href="#">loadData</a> for loading a dataset.
method	the linking method to use. Accepts:
	<ul style="list-style-type: none"> <li>• MM for mean-mean method</li> <li>• MS for mean-sigma method</li> <li>• HB for Haebara method</li> <li>• SL for Stocking-Lord method</li> <li>• FIXEDPAR for fixed parameter calibration</li> <li>• CP for calibrated projection using fixed parameter calibration on the anchor dimension</li> <li>• CPLA for linear approximation of calibrated projection. This is identical to 'CP' but uses approximation in <a href="#">runRSSS</a></li> <li>• CPFIXEDDIM for calibrated projection using mean and variance constraints on the anchor dimension</li> </ul>
	Linear transformation methods (i.e., MM, MS, HB, SL) are performed with <a href="#">plink</a> in ' <a href="#">plink</a> ' package.
verbose	if TRUE, print status messages. (default = FALSE)
...	additional arguments to pass onto <a href="#">mirt</a> in ' <a href="#">mirt</a> ' package.

## Value

`runLinking` returns a [list](#) containing the scale linking results.

- constants linear transformation constants. Only available when linear transformation methods were used (i.e., MM, MS, HB, SL).
- ipar\_linked item parameters calibrated to the response data, and linked to the metric of anchor item parameters.
- ipar\_anchor anchor item parameters used in linking.

## Examples

```
out_link <- runLinking(data_asq, "SL", technical = list(NCYCLES = 1000))
out_link$constants # transformation constants
out_link$ipar_linked # item parameters linked to anchor
out_link <- runLinking(data_asq, "FIXEDPAR")
out_link$ipar_linked # item parameters linked to anchor
```

---

**runRSSS***Compute Crosswalk Tables*

---

**Description**

**runRSSS** is a function for generating raw-score to standard-score crosswalk tables from supplied calibrated item parameters.

**Usage**

```
runRSSS(
  data,
  ipar_linked,
  prior_mean = 0,
  prior_sd = 1,
  min_theta = -4,
  max_theta = 4,
  inc = 0.05,
  min_score = 1
)
```

**Arguments**

<b>data</b>	a <a href="#">PROsetta_data</a> object. See <a href="#">loadData</a> for loading a dataset.
<b>ipar_linked</b>	an object returned from <a href="#">runLinking</a> or <a href="#">runCalibration</a> .
<b>prior_mean</b>	prior mean. (default = 0.0)
<b>prior_sd</b>	prior standard deviation. (default = 1.0)
<b>min_theta</b>	the lower limit of theta quadratures for numerical integration. (default = -4)
<b>max_theta</b>	the upper limit of theta quadratures for numerical integration. (default = 4)
<b>inc</b>	the increment between theta quadratures for numerical integration. (default = 0.05)
<b>min_score</b>	minimum item score (0 or 1) for each scale (1, 2, and combined). If a single value is supplied, the value is applied to all scales. (default = 1)

**Value**

**runRSSS** returns a [list](#) containing crosswalk tables.

**Examples**

```
out_link    <- runLinking(data_asq, method = "FIXEDPAR")
score_table <- runRSSS(data_asq, out_link)
```

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