User manual for the psych package Version 1.0.58

William Revelle Department of Psychology Northwestern University

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psych	A package for personality,	psychometric,	$and \ psychological \ re-$
	search		

Description

The psych package has been developed at Northwestern University to include functions most useful for personality and psychological research. Some of the functions (e.g., read.clipboard, describe, pairs.panels, error.bars) are useful for basic daa entry and descriptive analyses.

Psychometric applications include routines for Very Simple Structure (VSS), Minimum Average Partial correlation (MAP), Item Cluster Analysis (ICLUST) and principal axes factor analysis (factor.pa), as well as functions to do Schmid Leiman transformations (schmid) to transform a hierarchical factor structure into a bifactor solution, and to calculate reliability coefficients alpha (score.items, score.multiple.choice), beta (ICLUST) and McDonald's omega (omega and omega.graph).

The score.items, and score.multiple.choice functions may be used to form single or multiple scales from sets of dichotomous, multilevel, or multiple choice items by specifying scoring keys.

Additional functions make for more convenient descriptions of item characteristics. Functions under development include 1 and 2 parameter Item Response measures.

A number of procedures have been developed as part of the Synthetic Aperture Personality Assessment (SAPA) project. These routines facilitate forming and analyzing composite scales equivalent to using the raw data but doing so by adding within and between cluster/scale item correlations. These functions include extracting clusters from factor loading matrices (factor2cluster), synthetically forming clusters from correlation matrices (cluster.cor), and finding multiple correlation from correlation matrices (mat.regress).

Functions to generate simulate data with particular structures include circ.sim (for circumplex structures), item.sim (for general structures) and congeneric.sim (for a specific demonstration of congeneric measurement).

Functions to apply various standard statistical tests include p.rep and its variants for testing the probability of replication, r.con for the confidence intervals of a correlation, r.test to test single, paired, or sets of correlations.

The most recent development version of the package is always available for download as a *source* file from the repository at http://personality-project.org/r/src/contrib/.

Details

The psych package was originally a combination of multiple source files maintained at the http://personality-project.org/r repository: "useful.r", VSS.r., ICLUST.r, omega.r, etc. "useful.r" is a set of routines for easy data entry (read.clipboard), simple descriptive statistics (describe), and splom plots combined with correlations (pairs.panels, adapted from the help files of pairs). It is now a single package.

The VSS routines allow for testing the number of factors (VSS), showing plots (VSS.plot) of goodness of fit, and basic routines for estimating the number of factors/components to extract by using the MAP's procedure, the examining the scree plot (VSS.scree) or comparing with the scree of an equivalent matrix of random numbers (VSS.parallel).

In addition, there are routines for hierarchical factor analysis using Schmid Leiman tranformations (omega, omega.graph) as well as Item Cluster analysis (ICLUST, ICLUST.graph).

The more important functions in the package are for the analysis of multivariate data, with an emphasis upon those functions useful in scale construction of item composites.

When given a set of items from a personality inventory, one goal is to combine these into higher level item composites. This leads to several questions:

- 1) What are the basic properties of the data? describe reports basic summary statistics (mean, sd, median, mad, range, minimum, maximum, skew, kurtosis, standard error) for vectors, columns of matrices, or data.frames. describe.by provides descriptive statistics, organized by a grouping variable. pairs.panels shows scatter plot matrices (SPLOMs) as well as histograms and the Pearson correlation for scales or items. error.bars will plot variable means with associated confidence intervals.
- 2) What is the most appropriate number of item composites to form? After finding either standard Pearson correlations, or finding tetrachoric or polychoric correlations using a wrapper (poly.mat) for John Fox's hetcor function, the dimensionality of the correlation matrix may be examined. The number of factors/components problem is a standard question of factor analysis, cluster analysis, or principal components analysis. Unfortunately, there is no agreed upon answer. The Very Simple Structure (VSS) set of procedures has been proposed as on answer to the question of the optimal number of factors. Other procedures (VSS.scree, VSS.parallel, fa.parallel, and MAP) also address this question.
- 3) What are the best composites to form? Although this may be answered using principal components (principal) or factor analysis (factor.pa) and to show the results graphically (fa.graph), it is sometimes more useful to address this question using cluster analytic techniques. (Better yet is to use maximum likelihood factor analysis using factanal from the stats package.) Previous versions of ICLUST (e.g., Revelle, 1979) have been shown to be particularly successful at doing this. Graphical output from ICLUST.graph uses the Graphviz dot language and allows one to write files suitable for Graphviz.

Graphical organizations of cluster and factor analysis output can be done using cluster.plot which plots items by cluster/factor loadings and assigns items to that dimension with the highest loading.

- 4) How well does a particular item composite reflect a single construct? This is a question of reliability and general factor saturation. Multiple solutions for this problem result in (Cronbach's) alpha (score.items), (Revelle's) Beta (ICLUST), and (McDonald's) omega. Functions to estimate all three of these are included in psych.
- 5) For some applications, data matrices are synthetically combined from sampling different items for different people. So called Synthetic Aperture Personality Assessement (SAPA) techniques allow the formation of large correlation or covariance matrices even though no one person has taken all of the items. To analyze such data sets, it is easy to form item composites based upon the covariance matrix of the items, rather than original data set. These matrices may then be analyzed using a number of functions (e.g., cluster.cor, factor.pa, ICLUST, principal, mat.regress, and factor2cluster.

6) More typically, one has a raw data set to analyze. score.items will score data sets on multiple scales, reporting the scale scores, item-scale and scale-scale correlations, as well as coefficient alpha and alpha-1. Using a 'keys' matrix (created by make.keys or by hand), scales can have overlapping or independent items. score.multiple.choice scores multiple choice items or converts multiple choice items to dichtomous (0/1) format for other functions.

An additional set of functions generate simulated data to meet certain structural properties. item.sim creates simple structure data, circ.sim will produce circumplex structured data, item.dichot produces circumplex or simple structured data for dichotomous items. These item structures are useful for understanding the effects of skew, differential item endorsement on factor and cluster analytic soutions.

When examining personality items, some people like to discuss them as representing items in a two dimensional space with a circumplex structure. Tests of circumplex fit circ.tests have been developed. When representing items in a circumplex, it is convenient to view them in polar coordinates.

Additional functions for testing the difference between two independent or dependent correlation r.test, to find the phi or Yule coefficients from a two by table, or to find the confidence interval of a correlation coefficient.

Ten data sets are included: bfi represents 25 personality items thought to represent five factors of personality, iqitems has 14 multiple choice iq items. sat.act has data on self reported test scores by age and gender. galton Galton's data set of the heights of parents and their children. peas recreates the original Galton data set of the genetics of sweet peas. heights and cubits provide even more Galton data, vegetables provides the Guilford preference matrix of vegetables. cities provides airline miles between 11 US cities (demo data for multidimensional scaling).

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Type: Package
Version: 1.0-57
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License: GPL version 2 or newer

Index:

psych A package for personality, psychometric, and psychological research.

Useful data entry and descriptive statistics

describe Basic descriptive statistics useful for psychometrics describe.by Find summary statistics by groups headtail combines the head and tail functions for showing data sets read.clipboard shortcut for reading from the clipboard read.clipboard.csv shortcut for reading comma delimited files from clipboard read.clipboard.lower shortcut for reading lower triangular matrices from the clipboard read.clipboard.upper shortcut for reading upper triangular matrices from the clipboard pairs.panels SPLOM and correlations for a data matrix multi.hist Histograms and densities of multiple variables arranged in matrix form

skew Calculate skew for a vector, each column of a matrix, or data.frame kurtosi Calculate kurtosis for a vector, each column of a matrix or dataframe geometric.mean Find the geometric mean of a vector or columns of a data.frame harmonic.mean Find the harmonic mean of a vector or columns of a data.frame error.bars Plot means and error bars error.bars.by Plot means and error bars for separate groups error.crosses Two way error bars interp.median Find the interpolated median, quartiles, or general quantiles. rescale Rescale data to specified mean and standard deviation table2df Convert a two dimensional table of counts to a matrix or data frame

Data reduction through cluster and factor analysis

factor.pa Do a principal Axis factor analysis fa.graph Show the results of a factor analysis or principal components analysis graphically principal Do an eigen value decomposition to find the principal components of a matrix fa.parallel Scree test and Parallel analysis guttman 8 different measures of reliability (6 from Guttman (1945) ICLUST Apply the ICLUST algorithm

ICLUST.graph Graph the output from ICLUST using the dot language ICLUST.rgraph Graph the output from ICLUST using rgraphviz poly.mat Find the polychoric correlations for items (uses J. Fox's hetcor omega Calculate the omega estimate of factor saturation (requires the GPArotation pack-

omega.graph Draw a hierarchical or Schmid Leiman orthogonalized solution schmid Apply the Schmid Leiman transformation to a correlation matrix

score.items Combine items into multiple scales and find alpha

score.multiple.choice Combine items into multiple scales and find alpha and basic scale statistics

smc Find the Squared Multiple Correlation (used for initial communality estimates)

VSS Apply the Very Simple Structure criterion to determine the appropriate number of factors.

VSS.parallel Do a parallel analysis to determine the number of factors for a random matrix VSS.plot Plot VSS output

VSS.scree Show the scree plot of the factor/principal components

VSS.simulate Generate simulated data for the factor model

make.hierarchical Generate simulated correlation matrices with hierarchical structure

MAP Apply the Velicer Minimum Absolute Partial criterion for number of factors

Procedures particularly useful for Synthetic Aperture Personality Assessment

alpha.scale Find coefficient alpha for a scale (see also score.items) make.keys Create the keys file for score.items or cluster.cor correct.cor Correct a correlation matrix for unreliability count.pairwise Count the number of complete cases when doing pair wise correlations cluster.cor find correlations of composite variables from larger matrix

cluster.loadings find correlations of items with composite variables from a larger matrix

eigen.loadings Find the loadings when doing an eigen value decomposition

factor.pa Do a Principal Axis factor analysis and estimate factor scores

factor2cluster extract cluster definitions from factor loadings

factor.congruence Factor congruence coefficient

factor.fit How well does a factor model fit a correlation matrix

factor.model Reproduce a correlation matrix based upon the factor model

factor.residuals Fit = data - model

factor.rotate "hand rotate" factors

guttman 8 different measures of reliability mat.regress standardized multiple regression from raw or correlation matrix input

principal Do an eigen value decomposition to find the principal components of a matrix

Functions for generating simulated data sets

circ.sim Generate a two dimensional circumplex item structure

item.sim Generate a two dimensional simple structrue with particular item characteristics

congeneric.sim Generate a one factor congeneric reliability structure

phi.demo Create artificial data matrices for teaching purposes

make.hierarchical Generate simulated correlation matrices with hierarchical or any structure

Miscellaneous functions

comorbidity Convert base rate and comorbity to phi, Yule and tetrachoric fisherz Apply the Fisher r to z transform

fisherz2r Apply the Fisher z to r transform

ICC Intraclass correlation coefficients

linkjennrich.cortest

mat.cortest Test for equality of two matrices (see also linknormal.cortest, linkjennrich.cortest) linknormal.cortest

paired.r Test for the difference of two paired or two independent correlations

r.con Confidence intervals for correlation coefficients

r.test Test of significance of r, differences between rs.

p.rep The probability of replication given a p, r, t, or F

phi Find the phi coefficient of correlation from a 2 x 2 table

phi.demo Demonstrate the problem of phi coefficients with varying cut points

phi2poly Given a phi coefficient, what is the polychoric correlation

phi2poly.matrix Given a phi coefficient, what is the polychoric correlation (works on matrices)

polar Convert 2 dimensional factor loadings to polar coordinates.

poly.mat Use John Fox's hetcor to create a matrix of correlations from a data.frame or matrix of integer values

polychor.matrix Use John Fox's polycor to create a matrix of polychoric correlations from a matrix of Yule correlations

scaling.fits Compares alternative scaling solutions and gives goodness of fits

thurstone Thurstone Case V scaling

tr Find the trace of a square matrix

wkappa weighted and unweighted versions of Cohen's kappa

Yule Find the Yule Q coefficient of correlation

Yule.inv What is the two by two table that produces a Yule Q with set marginals? Yule2phi What is the phi coefficient corresponding to a Yule Q with set marginals? Yule2phi.matrix Convert a matrix of Yule coefficients to a matrix of phi coefficients. Yule2phi.matrix Convert a matrix of Yule coefficients to a matrix of polychoric coefficients.

Functions that are under development and not recommended for casual use irt.item.diff.rasch IRT estimate of item difficulty with assumption that theta = 0 irt.person.rasch Item Response Theory estimates of theta (ability) using a Rasch like model

Data sets included in the psych package

bfi represents 25 personality items thought to represent five factors of personality cities The airline distances between 11 cities (used to demonstrate MDS) epi.bfi 13 personality scales

igitems 14 multiple choice ig items

sat.act Self reported ACT and SAT Verbal and Quantitative scores by age and gender galton Galton's data set of the heights of parents and their children

heights Galton's data set of the relationship between height and forearm (cubit) length cubits Galton's data table of height and forearm length

peas Galton's data set of the diameters of 700 parent and offspring sweet peas vegetables Guilford's preference matrix of vegetables (used for thurstone)

test.psych Run a test of the major functions on 5 different data sets. Primarily for development purposes. Although the output can be used as a demo of the various functions.

Note

Development versions (source code) of this package are maintained at the repository http://personality-project.org/r along with further documentation. Specify that you are downloading a source package.

Some functions require other packages. Specifically, omega and schmid require the GPArotation package, and poly.mat, phi2poly and polychor.matrix requires John Fox's polychor package. ICLUST.rgraph and fa.graph require Rgraphviz. i.e.:

function requires **GPArotation** omega schmid **GPArotation** poly.mat polychor polychor phi2poly polychor polychor.matrix ICLUST.rgraph Rgraphviz fa.graph Rgraphviz

Author(s)

William Revelle Department of Psychology

```
Northwestern University
Evanston, Illiniois
http://personality-project.org/revelle.html
```

Maintainer: William Revelle < revelle@northwestern.edu>

References

A general guide to personality theory and research may be found at the personality-project.org. See also the short guide to R at http://personality-project.org/r. In addition, see

Revelle, W. (in preparation) An Introduction to Psychometric Theory with applications in R. Springer. at http://personality-project.org/r/book/

Examples

```
#See the separate man pages
test.psych()
```

alpha.scale

Cronbach alpha for a scale

Description

Find Cronbach's coefficient alpha given a scale and a data.frame of the items in the scale. For X, a total score composed of items in the data.frame Y, find Cronbach's alpha.

Usage

```
alpha.scale(x, y)
```

Arguments

x A scale made by summing items

y A data frame of items

Details

Alpha is one of several estimates of the internal consistency reliability of a test. Perhaps because it is so easy to calculate, it is without doubt the most frequently reported measure of internal consistency reliability. Alpha is the mean of all possible spit half reliabilities (corrected for test length). For a unifactorial test, it is a reasonable estimate of the first factor saturation, although if the test has any microstructure (i.e., if it is "lumpy") coefficients beta (see ICLUST and link{omega} are more appropriate estimates of the general factor saturation.

Alpha is a positive function of the number of items in a test as well as the average intercorrelation of the items in the test. When calculated from the item variances and total test variance, as is done here, alpha is sensitive to differences in the item variances. Alternative functions score.items and cluster.cor will also score multiple scales and report more useful statistics. "Standardized" alpha is calculated from the inter-item correlations and will differ from raw alpha. Standardized alpha can be found by using cluster.cor.

Value

Coefficient alpha

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

References

```
http://personality-project.org/revelle/syllabi/405.syllabus.html
```

See Also

```
score.items, cluster.cor, ICLUST, omega, describe, pairs.panels, alpha in psychometrics
```

Examples

```
y <- attitude  #from the datasets package
x <- rowSums(y)  #find the sum of the seven attitudes
alpha.scale(x,y)
#[1] 0.8431428
#compare with standardized alpha:
st.alpha <- cluster.cor(rep(1,7),cor(attitude),digits=4)
st.alpha
#compare with score.items
si <- score.items(rep(1,7), attitude,digits=3)
si$alpha</pre>
```

bartlett.cortest

Bartlett's test that a correlation matrix is an identity matrix

Description

Bartlett (1951) proposed that $-\ln(\det(R)^*(N-1-(2p+5)/6))$ was distributed as chi square if R were an identity matrix. A useful test that residuals correlations are all zero.

Usage

```
bartlett.cortest(R, n = NULL)
```

Arguments

R A correlation matrix. (If R is not square, correlations are found and a

warning is issued.

n Sample size (if not specified, 100 is assumed.

Details

More useful for pedagogical purposes than actual applications. The Bartlett test is asymptotically chi square distributed.

Value

chisq Assymptotically chisquare

p.value Of chi square

df The degrees of freedom

Author(s)

William Revelle

References

Bartlett, M. S., (1951), The Effect of Standardization on a χ^2 Approximation in Factor Analysis, Biometrika, 38, 337-344.

See Also

```
mat.cortest, normal.cortest, jennrich.cortest
```

Examples

```
set.seed(42)
x <- matrix(rnorm(1000),ncol=10)
r <- cor(x)
bartlett.cortest(r)  #random data don't differ from an identity matrix
data(bfi)
bartlett.cortest(bfi)  #not an identity matrix</pre>
```

bfi

25 Personality items representing 5 factors

Description

25 personality self report items taken from the International Personality Item Pool (ipip.ori.org) were included as part of the Synthetic Aperture Personality Assessment (SAPA) web based personality assessment project. The data from 1000 subjects are included here as a demonstration set for scale construction and factor analysis.

Usage

data(bfi)

Format

A data frame with 1000 observations on the following 25 variables.

- A1 Am indifferent to the feelings of others.
- A2 Inquire about others' well-being.
- A3 Know how to comfort others.
- A4 Love children.
- A5 Make people feel at ease.
- C1 Am exacting in my work.
- C2 Continue until everything is perfect.
- C3 Do things according to a plan.
- C4 Do things in a half-way manner.
- C5 Waste my time.
- E1 Don't talk a lot.
- E2 Find it difficult to approach others.
- E3 Know how to captivate people.
- E4 Make friends easily.
- E5 Take charge.
- N1 Get angry easily.
- N2 Get irritated easily.
- N3 Have frequent mood swings.
- N4 Often feel blue.
- N5 Panic easily.
- 01 Am full of ideas.
- 02 Avoid imposing my will on others.
- 03 Carry the conversation to a higher level.
- 04 Spend time reflecting on things.
- 05 Will not probe deeply into a subject.

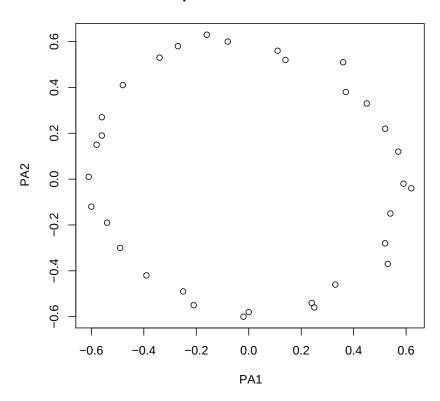
Details

The 25 items are organized by five putative factors: Agreeableness, Conscientiousness, Extraversion, Neuroticism, and Opennness. The scoring key is created using make.keys, the scores are found using score.items

Source

The items are from the ipip. The data are from the SAPA project, collected fall,2006.

A circumplex structure of 32 variables



Examples

```
data(bfi)
describe(bfi)
data(bfi)
keys.list <- list(Agree=c(-1,2:5),Conscientious=c(6:8,-9,-10),Extraversion=c(-11,-12,13:15),Neuroticism=
keys <- make.keys(25,keys.list,item.labels=colnames(bfi))
scores <- score.items(keys,bfi,short=TRUE)
scores</pre>
```

circ.simulation

 $Simulations\ of\ circumplex\ and\ simple\ structure$

Description

Rotations of factor analysis and principal components analysis solutions typically try to represent correlation matrices as simple structured. An alternative structure, appealing to some, is a circumplex structure where the variables are uniformly spaced on the perimeter

of a circle in a two dimensional space. Generating these data is straightforward, and is useful for exploring alternative solutions to affect and personality structure.

Usage

```
circ.simulation(samplesize=c(100,200,400,800), numberofvariables=c(16,32,48,72))
```

Arguments

samplesize a vector of sample sizes to simulate numberofvariables

vector of the number of variables to simulate

Details

"A common model for representing psychological data is simple structure (Thurstone, 1947). According to one common interpretation, data are simple structured when items or scales have non-zero factor loadings on one and only one factor (Revelle & Rocklin, 1979). Despite the commonplace application of simple structure, some psychological models are defined by a lack of simple structure. Circumplexes (Guttman, 1954) are one kind of model in which simple structure is lacking.

"A number of elementary requirements can be teased out of the idea of circumplex structure. First, circumplex structure implies minimally that variables are interrelated; random noise does not a circumplex make. Second, circumplex structure implies that the domain in question is optimally represented by two and only two dimensions. Third, circumplex structure implies that variables do not group or clump along the two axes, as in simple structure, but rather that there are always interstitial variables between any orthogonal pair of axes (Saucier, 1992). In the ideal case, this quality will be reflected in equal spacing of variables along the circumference of the circle (Gurtman, 1994; Wiggins, Steiger, & Gaelick, 1981). Fourth, circumplex structure implies that variables have a constant radius from the center of the circle, which implies that all variables have equal communality on the two circumplex dimensions (Fisher, 1997; Gurtman, 1994). Fifth, circumplex structure implies that all rotations are equally good representations of the domain (Conte & Plutchik, 1981; Larsen & Diener, 1992)." (Acton and Revelle, 2004)

Acton and Revelle reviewed the effectiveness of 10 tests of circumplex structure and found that four did a particularly good job of discriminating circumplex structure from simple structure, or circumplexes from ellipsoidal structures. Unfortunately, their work was done in Pascal and is not easily available. Here we release R code to do the four most useful tests:

The Gap test of equal spacing

Fisher's test of equality of axes

A test of indifference to Rotation

A test of equal Variance of squared factor loadings across arbitrary rotations.

Included in this set of functions are simple procedure to generate circumplex structured or simple structured data, the four test statistics, and a simple simulation showing the effectiveness of the four procedures.

circ.sim.plot compares the four tests for circumplex, ellipsoid and simple structure data as function of the number of variables and the sample size. What one can see from this plot is that although no one test is sufficient to discriminate these alternative structures, the set of four tests does a very good job of doing so. When testing a particular data set for structure, comparing the results of all four tests to the simulated data will give a good indication of the structural properties of the data.

Value

A data frame with simulation results for circumplex, ellipsoid, and simple structure data sets for each of the four tests.

Note

The simulations default values are for sample sizes of 100, 200, 400, and 800 cases, with 16, 32, 48 and 72 items.

Author(s)

William Revelle

References

Acton, G. S. and Revelle, W. (2004) Evaluation of Ten Psychometric Criteria for Circumplex Structure. Methods of Psychological Research Online, Vol. 9, No. 1 http://www.dgps.de/fachgruppen/methoden/mpr-online/issue22/mpr110_10.pdf

See Also

```
See also circ.tests, circ.sim
```

Examples

```
demo <- circ.simulation()
boxplot(demo[3:14])
title("4 tests of Circumplex Structure", sub="Circumplex, Ellipsoid, Simple Structure")
circ.sim.plot(demo[3:14]) #compare these results to real data</pre>
```

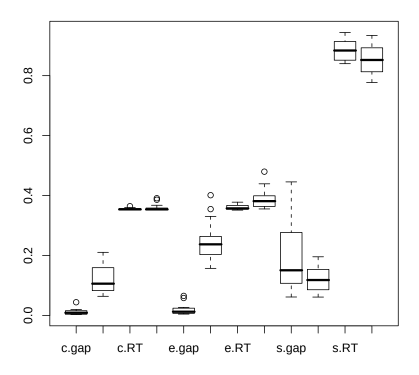
circ.tests

Apply four tests of circumplex versus simple structure

Description

Rotations of factor analysis and principal components analysis solutions typically try to represent correlation matrices as simple structured. An alternative structure, appealing to some, is a circumplex structure where the variables are uniformly spaced on the perimeter of a circle in a two dimensional space. Generating these data is straightforward, and is useful for exploring alternative solutions to affect and personality structure.

4 tests of Circumplex Structure



Circumplex, Ellipsoid, Simple Structure

Figure 1: Tests for circumplex structure can be shown to discriminate circumplex and simple structure. Here we compare four tests of circumplex structure, the Gap Test, the Fisher Test, the Rotation Test, and the Variance Test of Circumplex structure. The data sets vary on whether or not they have a circumplex or simple structure. See Acton and Revelle (2004) for details.

Usage

```
circ.tests(loads, loading = TRUE, sorting = TRUE)
```

Arguments

loads A matrix of loadings loads here

loading Are these loadings or a correlation matrix loading

sorting Should the variables be sorted sorting

Details

"A common model for representing psychological data is simple structure (Thurstone, 1947). According to one common interpretation, data are simple structured when items or scales have non-zero factor loadings on one and only one factor (Revelle & Rocklin, 1979). Despite the commonplace application of simple structure, some psychological models are defined by a lack of simple structure. Circumplexes (Guttman, 1954) are one kind of model in which simple structure is lacking.

"A number of elementary requirements can be teased out of the idea of circumplex structure. First, circumplex structure implies minimally that variables are interrelated; random noise does not a circumplex make. Second, circumplex structure implies that the domain in question is optimally represented by two and only two dimensions. Third, circumplex structure implies that variables do not group or clump along the two axes, as in simple structure, but rather that there are always interstitial variables between any orthogonal pair of axes (Saucier, 1992). In the ideal case, this quality will be reflected in equal spacing of variables along the circumference of the circle (Gurtman, 1994; Wiggins, Steiger, & Gaelick, 1981). Fourth, circumplex structure implies that variables have a constant radius from the center of the circle, which implies that all variables have equal communality on the two circumplex dimensions (Fisher, 1997; Gurtman, 1994). Fifth, circumplex structure implies that all rotations are equally good representations of the domain (Conte & Plutchik, 1981; Larsen & Diener, 1992). (Acton and Revelle, 2004)

Acton and Revelle reviewed the effectiveness of 10 tests of circumplex structure and found that four did a particularly good job of discriminating circumplex structure from simple structure, or circumplexes from ellipsoidal structures. Unfortunately, their work was done in Pascal and is not easily available. Here we release R code to do the four most useful tests:

- 1 The Gap test of equal spacing
- 2 Fisher's test of equality of axes
- 3 A test of indifference to Rotation
- 4 A test of equal Variance of squared factor loadings across arbitrary rotations.

To interpret the values of these various tests, it is useful to compare the particular solution to simulated solutions representing pure cases of circumplex and simple structure. See the example output from circ.simulation and compare these plots with the results of the circ.test.

Value

A list of four items is returned. These are the gap, fisher, rotation and variance test results.

gaps gap.test
fisher fisher.test
RT rotation.test
VT variance.test

Note

Of the 10 criterion discussed in Acton and Revelle (2004), these tests operationalize the four most useful.

Author(s)

William Revelle

References

Acton, G. S. and Revelle, W. (2004) Evaluation of Ten Psychometric Criteria for Circumplex Structure. Methods of Psychological Research Online, Vol. 9, No. 1 http://personality-project.org/revelle/publications/acton.revelle.mpr110_10.pdf

See Also

```
circ.simulation, circ.sim
```

```
circ.data <- circ.sim(24,500)
circ.fa <- factor.pa(circ.data,2)
#plot(circ.fa$loadings)
ct <- circ.tests(circ.fa)
#compare with non-circumplex data
simp.data <- item.sim(24,500)
simp.fa <- factor.pa(simp.data,2)
#plot(simp.fa$loadings)
st <- circ.tests(simp.fa)
print(rbind(ct,st),digits=2)</pre>
```

Description

Airline distances between 11 US cities may be used as an example for multidimensional scaling or cluster analysis.

Usage

data(cities)

Format

A data frame with 11 observations on the following 11 variables.

ATL Atlana, Georgia

BOS Boston, Massachusetts

ORD Chicago, Illinois

DCA Washington, District of Columbia

DEN Denver, Colorado

LAX Los Angeles, California

MIA Miami, Florida

JFK New York, New York

SEA Seattle, Washington

SFO San Francisco, California

MSY New Orleans, Lousianna

Details

An 11 x11 matrix of distances between major US airports. This is a useful demonstration of multiple dimensional scaling.

city.location is a dataframe of longitude and latitude for those cities.

Note that the 2 dimensional MDS solution does not perfectly capture the data from these city distances. Boston, New York and Washington, D.C. are located slightly too far west, and Seattle and LA are slightly too far south.

Source

http://www.timeanddate.com/worldclock/distance.html

Examples

```
data(cities)
city.location[,1] <- -city.location[,1]
if(require(maps)) {map("usa")}
title("MultiDimensional Scaling of US cities")
points(city.location)} else {plot(city.location, xlab="Dimension 1", ylab="Dimension 2",main ="multidimen
city.loc <- cmdscale(cities, k=2) #ask for a 2 dimensional solution round(city.loc,0)
city.loc <- -city.loc
city.loc <- rescale(city.loc,mean(city.location),sd(city.location))
points(city.loc,type="n")
text(city.loc,labels=names(cities))</pre>
```

cluster.cor

Find correlations of composite variables from a larger matrix

Description

Given a n x c cluster definition matrix of -1s, 0s, and 1s (the keys), and a n x n correlation matrix, find the correlations of the composite clusters. The keys matrix can be entered by hand, copied from the clipboard (read.clipboard), or taken as output from the factor2cluster function.

Usage

```
cluster.cor(keys, r.mat, correct = TRUE, digits=2)
```

Arguments

keys A matrix of cluster keys
r.mat A correlation matrix

correct TRUE shows both raw and corrected for attenuation correlations

digits round off answer to digits

Details

This is one of the functions used in the SAPA procedures to form synthetic correlation matrices. Given any correlation matrix of items, it is easy to find the correlation matrix of scales made up of those items. This can also be done from the original data matrix using score.items.

A typical use in the SAPA project is to form item composites by clustering or factoring (see ICLUST, principal), extract the clusters from these results (factor2cluster), and then form the composite correlation matrix using cluster.cor. The variables in this reduced matrix may then be used in multiple R procedures using mat.regress.

The original correlation is pre and post multiplied by the (transpose) of the keys matrix.

If some correlations are missing from the original matrix this will lead to missing values (NA) for scale intercorrelations based upon those lower level correlations.

Because the alpha estimate of reliability is based upon the correlations of the items rather than upon the covariances, this estimate of alpha is sometimes called "standardized alpha". If the raw items are available, it is useful to compare standardized alpha with the raw alpha found using <code>score.items</code>. They will differ substantially only if the items differ a great deal in their variances.

Value

cor the (raw) correlation matrix of the clusters
sd standard deviation of the cluster scores

corrected raw correlations below the diagonal, alphas on diagonal, disattenuated

above diagonal

alpha The (standardized) alpha reliability of each scale.

size How many items are in each cluster?

Note

See SAPA e.g., Revelle, W. (2006) Synthetic Aperture Personality Assessment. Invited paper at the Midwestern Psychological Association Annual Meeting, Chicago, May, 2006. pdf available at http://personality-project.org/revelle/publications/sapa.mpa.key.pdf

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

See Also

```
factor2cluster, mat.regress, alpha.scale, score.items
```

```
## Not run:
data(attitude)
0,0,0,1,1,1,1), ncol=2
colnames(keys) <- c("first", "second")</pre>
r.mat <- cor(attitude)</pre>
cluster.cor(keys,r.mat)
## End(Not run)
#$cor
       first second
#first
         1.0
               0.6
#second 0.6
               1.0
#$sd
# first second
 2.57
         3.01
```

```
# #$corrected
# first second
#first 0.82 0.77
#second 0.60 0.74
#
#$size
# first second
# 3 4
```

cluster.fit

cluster Fit: fit of the cluster model to a correlation matrix

Description

How well does the cluster model found by ICLUST fit the original correlation matrix? A similar algorithm factor.fit is found in VSS. This function is internal to ICLUST but has more general use as well.

In general, the cluster model is a Very Simple Structure model of complexity one. That is, every item is assumed to represent only one factor/cluster. Cluster fit is an analysis of how well this model reproduces a correlation matrix. Two measures of fit are given: cluster fit and factor fit. Cluster fit assumes that variables that define different clusters are orthogonal. Factor fit takes the loadings generated by a cluster model, finds the cluster loadings on all clusters, and measures the degree of fit of this somewhat more complicated model. Because the cluster loadings are similar to, but not identical to factor loadings, the factor fits found here and by factor.fit will be similar.

Usage

```
cluster.fit(original, load, clusters, diagonal = FALSE, digits = 2)
```

Arguments

original The original correlation matrix being fit

load Cluster loadings – that is, the correlation of individual items with the

clusters, corrected for item overlap

clusters The cluster structure

diagonal Should we fit the diagonal as well?

digits Number of digits to be used in the output

Details

The cluster model is similar to the factor model: R is fitted by C'C. Where C <- Cluster definition matrix x the loading matrix. How well does this model approximate the original correlation matrix and how does this compare to a factor model?

The fit statistic is a comparison of the original (squared) correlations to the residual correlations. Fit = $1 - r^2/r^2$ where r^* is the residual correlation of data - model and model = C'C.

Value

clusterfit The cluster model is a reduced form of the factor loading matrix. That

is, it is the product of the elements of the cluster matrix * the loading

matrix

factorfit How well does the complete loading matrix reproduce the correlation ma-

trix?

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

References

```
http://personality-project.org/r/r.ICLUST.html
```

See Also

```
VSS, ICLUST, factor2cluster, cluster.cor, factor.fit
```

Examples

```
r.mat<- Harman74.cor$cov
iq.clus <- ICLUST(r.mat,nclusters =2)
fit <- cluster.fit(r.mat,iq.clus$loadings,iq.clus$clusters)
fit</pre>
```

cluster.loadings

Find item by cluster correlations, corrected for overlap and reliability

Description

Given a n x n correlation matrix and a n x c matrix of -1,0,1 cluster weights for those n items on c clusters, find the correlation of each item with each cluster. If the item is part of the cluster, correct for item overlap. Part of the ICLUST set of functions, but useful for many item analysis problems.

Usage

```
cluster.loadings(keys, r.mat, correct = TRUE, digits = 2)
```

Arguments

keys Cluster keys: a matrix of -1,0,1 cluster weights

r.mat A correlation matrixcorrect Correct for reliabilitydigits Number of digits output

Details

Given a set of items to be scored as (perhaps overlapping) clusters and the intercorrelation matrix of the items, find the clusters and then the correlations of each item with each cluster. Correct for item overlap by replacing the item variance with its average within cluster inter-item correlation.

Although part of ICLUST, this may be used in any SAPA application where we are interested in item- whole correlations of items and composite scales.

These loadings are particularly interpretable when sorted by absolute magnitude for each cluster (see ICLUST.sort).

Value

loadings A matrix of item-cluster correlations (loadings)

cor Correlation matrix of the clusters

corrected Correlation matrix of the clusters, raw correlations below the diagonal,

alpha on diagonal, corrected for reliability above the diagonal

sd Cluster standard deviations

alpha alpha reliabilities of the clusters

count Number of items in the cluster

Note

Although part of ICLUST, this may be used in any SAPA application where we are interested in item- whole correlations of items and composite scales.

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

References

```
ICLUST: http://personality-project.org/r/r.iclust.html
```

See Also

```
ICLUST, factor2cluster, cluster.cor
```

```
## Not run:
    r.mat<- Harman74.cor$cov
    clusters <- matrix(c(1,1,1,rep(0,24),1,1,1,1,rep(0,17)),ncol=2)
    cluster.loadings(clusters,r.mat)
## End(Not run)</pre>
```

cluster.plot	Plot factor/cluster loadings and assign items to clusters by their
	highest loading.

Description

Cluster analysis and factor analysis are procedures for grouping items in terms of a smaller number of (latent) factors or (observed) clusters. Graphical presentations of clusters typically show tree structures, although they can be represented in terms of item by cluster correlations.

Cluster.plot plots items by their cluster loadings (taken, e.g., from ICLUST). Cluster membership may be assigned apriori or may be determined in terms of the highest (absolute) cluster loading for each item.

If the input is an object of class "kmeans", then the cluster centers are plotted.

Usage

```
cluster.plot(ic.results, cluster = NULL, cut = 0, labels=NULL,title = "Cluster plot",...)
```

Arguments

ic.results	A factor analysis or	cluster analysis output	including the loadings, or a
------------	----------------------	-------------------------	------------------------------

matrix of item by cluster correlations. Or the output from a kmeans

cluster analysis.

cluster A vector of cluster membership

cut Assign items to clusters if the absolute loadings are > cut

labels If row.names exist they will be added to the plot, or, if they don't, labels

can be specified. If labels = NULL, and there are no row names, then

variables are labeled by row number.)

title Any title

... Further options to plot

Value

Graphical output is presented

Note

September 5, 2007

Author(s)

William Revelle

See Also

ICLUST, ICLUST. graph

Examples

```
circ.data <- circ.sim(24,500)
circ.fa <- factor.pa(circ.data,2)
cluster.plot(circ.fa,cut=.5)</pre>
```

cluster2keys

Convert a cluster vector (from e.g., kmeans) to a keys matrix suitable for scoring item clusters.

Description

The output of the kmeans clustering function produces a vector of cluster membership. The score.items and cluster.cor functions require a matrix of keys. cluster2keys does this.

Usage

```
cluster2keys(c)
```

Arguments

С

A vector of cluster assignments or an object of class "kmeans" that contains a vector of clusters.

Details

Note that because kmeans will not reverse score items, the clusters defined by kmeans will not necessarily match those of ICLUST with the same number of clusters extracted.

Value

keys

A matrix of keys suitable for score.items or cluster.cor

Author(s)

William Revelle

See Also

```
cluster.cor,score.items
```

```
test.data <- Harman74.cor$cov
kc <- kmeans(test.data,4)
keys <- cluster2keys(kc)
keys #these match those found by ICLUST
cluster.cor(keys,test.data)</pre>
```

comorbidity	Convert base rates of two	diagnoses	$and\ their$	comorbidity into
	ohi Vule and tetrachoric	e		

Description

In medicine and clinical psychology, diagnoses tend to be categorical (someone is depressed or not, someone has an anxiety disorder or not). Cooccurrence of both of these symptoms is called comorbidity. Diagnostic categories vary in their degree of comorbidity with other diagnostic categories. From the point of view of correlation, comorbidity is just a name applied to one cell in a four fold table. It is thus possible to analyze comorbidity rates by considering the probability of the separate diagnoses and the probability of the joint diagnosis. This gives the two by two table needed for a phi, Yule, or tetrachoric correlation.

Usage

```
comorbidity(d1, d2, com, labels = NULL)
```

Arguments

d1	Proportion of diagnostic category 1
d2	Proportion of diganostic category 2

com Proportion of comorbidity (diagnostic category 1 and 2)

labels Names of categories 1 and 2

Value

twobytwo The two by two table implied by the input
phi Phi coefficient of the two by two table
Yule Yule coefficient of the two by two table

tetrachoric Tetrachoric coefficient of the two by two table

Note

Requires the polycor package

Author(s)

William Revelle

See Also

```
phi, Yule
```

```
if(require(polycor)) {comorbidity(.2,.15,.1,c("Anxiety","Depression")) }
```

Simulate a congeneric data set

congeneric.sim

Description

Classical Test Theory (CTT) considers four or more tests to be congenerically equivalent if all tests may be expressed in terms of one factor and a residual error. Parallel tests are the special case where (usually two) tests have equal factor loadings. Tau equivalent tests have equal factor loadings but may have unequal errors. Congeneric tests may differ in both factor loading and error variances.

Usage

```
congeneric.sim(N = 1000, loads = c(0.8, 0.7, 0.6, 0.5), err=NULL, short = TRUE)
```

Arguments

N How many subjects to simulate

loads A vector of factor loadings for the tests

err A vector of error variances – if NULL then error = 1 - loading 2

short short=TRUE: Just give the test scores, short=FALSE, report observed

test scores as well as the implied pattern matrix

Details

When constructing examples for reliability analysis, it is convenient to simulate congeneric data structures. These are the most simple of item structures, having just one factor. Mainly used for a discussion of reliability theory as well as factor score estimates.

The implied covariance matrix is just pattern %*% t(pattern).

Value

model The implied population correlation matrix

pattern The pattern matrix implied by the loadings and error variances

observed a matrix of test scores for n tests

1atent The latent trait and error scores

Author(s)

William Revelle

See Also

item.sim for other simulations, factor.pa for an example of factor scores.

Examples

```
test <- congeneric.sim(short=FALSE)
round(cor(test$observed),2) # show a congeneric correlation matrix
f1=factor.pa(test$observed,1,scores=TRUE)
round(cor(f1$scores,test$latent),2) #factor score estimates are correlated with but not equal to the fac</pre>
```

correct.cor

Find dis-attenuated correlations and give alpha reliabilities

Description

Given a raw correlation matrix and a vector of reliabilities, report the disattenuated correlations above the diagonal.

Usage

```
correct.cor(x, y)
```

Arguments

x A raw correlation matrix

y Vector of reliabilities

Details

Disattenuated correlations may be thought of as correlations between the latent variables measured by a set of observed variables. That is, what would the correlation be between two (unreliable) variables be if both variables were measured perfectly reliably.

This function is mainly used if importing correlations and reliabilities from somewhere else. If the raw data are available, use score.items, or cluster.loadings or cluster.cor.

Examples of the output of this function are seen in cluster.loadings and cluster.cor

Value

Raw correlations below the diagonal, reliabilities on the diagonal, disattenuated above the diagonal.

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

References

http://personality-project.org/revelle/syllabi/405.syllabus.html

See Also

```
cluster.loadings and cluster.cor
```

Examples

```
# attitude from the datasets package
#example 1 is a rather clunky way of doing things
## Not run:
a1 <- attitude[,c(1:3)]
a2 <- attitude[,c(4:7)]
x1 <- rowSums(a1) #find the sum of the first 3 attitudes
x2 <- rowSums(a2) #find the sum of the last 4 attitudes
alpha1 <- alpha.scale(x1,a1)
alpha2 <- alpha.scale(x2,a2)
x <- matrix(c(x1,x2),ncol=2)</pre>
x.cor <- cor(x)</pre>
alpha <- c(alpha1,alpha2)</pre>
round(correct.cor(x.cor,alpha),2)
#much better - although uses standardized alpha
clusters \leftarrow matrix(c(rep(1,3),rep(0,7),rep(1,4)),ncol=2)
cluster.loadings(clusters,cor(attitude))
# or
clusters <- matrix(c(rep(1,3),rep(0,7),rep(1,4)),ncol=2)</pre>
cluster.cor(clusters,cor(attitude))
#best
scores <- score.items(matrix(c(rep(1,3),rep(0,7),rep(1,4)),ncol=2),attitude)</pre>
scores$corrected
## End(Not run)
```

count.pairwise

Count number of pairwise cases for a data set with missing (NA) data.

Description

When doing cor(x, use="pairwise"), it is nice to know the number of cases for each pairwise correlation. This is particularly useful when doing SAPA type analyses.

Usage

```
count.pairwise(x, y = NULL)
```

Arguments

x An input matrix, typically a data matrix ready to be correlated.

y An optional second input matrix

Value

result = matrix of counts of pairwise observations

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

Examples

```
## Not run:
x <- matrix(rnorm(1000),ncol=6)
y <- matrix(rnorm(500),ncol=3)
x[x < 0] <- NA
y[y> 1] <- NA

count.pairwise(x)
count.pairwise(y)
count.pairwise(x,y)
## End(Not run)</pre>
```

cubits

Galton's example of the relationship between height and 'cubit' or forearm length

Description

Francis Galton introduced the 'co-relation' in 1888 with a paper discussing how to measure the relationship between two variables. His primary example was the relationship between height and forearm length. The data table (cubits) is taken from Galton (1888). Unfortunately, there seem to be some errors in the original data table in that the marginal totals do not match the table.

The data frame, heights, is converted from this table.

Usage

```
data(cubits)
```

Format

A data frame with 9 observations on the following 8 variables.

```
16.5 Cubit length of lowest category
16.75 a numeric vector
17.25 a numeric vector
18.25 a numeric vector
18.75 a numeric vector
19.25 a numeric vector
19.75 a numeric vector
19.75 a numeric vector
```

Details

Sir Francis Galton (1888) published the first demonstration of the correlation coefficient. The regression (or reversion to mediocrity) of the height to the length of the left forearm (a cubit) was found to .8. There seem to be some errors in the table as published in that the row sums do not agree with the actual row sums. These data are used to create a matrix using table2matrix for demonstrations of analysis and displays of the data.

Source

```
Galton (1888)
```

References

Galton, Francis (1888) Co-relations and their measurement. Proceedings of the Royal Society. London Series, 45, 135-145,

See Also

```
table2matrix, table2df,heights,ellipses,galton
```

```
data(cubits)
cubits
heights <- table2df(cubits,labs <- c("height","cubit"))
ellipses(heights,n=1,main="Galton's co-relation data set")
ellipses(jitter(heights$cubit,3),jitter(heights$height,3),pch=".",main="Galton's co-relation data set") #a</pre>
```

Basic summary statistics by group

describe.by

Description

Report basic summary statistics by a grouping variable. Useful if the grouping variable is some experimental variable and data are to be aggregated for plotting. Just a wrapper for by and describe.

Usage

```
describe.by(x, group,mat=FALSE,...)
```

Arguments

x a data.frame or matrixgroupa grouping variable

mat provide a matrix output rather than a list
... parameters to be passed to describe

Value

A data frame of the relevant statistics broken down by group:

item name item number

number of valid cases

mean

standard deviation

median

mad: median absolute deviation (from the median)

minimum maximum skew

standard error

Author(s)

William Revelle

See Also

describe

Examples

```
set.seed(42) #to get the same values each time
  x.df <- data.frame(group=sample(2,20,replace=TRUE), matrix(rnorm(100),ncol=5))
  x <- describe.by(x.df,x.df$group)
  x #shows all the results
  x[1] #shows just the first group
  x <- matrix(sample(4,200,replace=TRUE),ncol=5)
  y <- describe.by(x,x[,1])
  v</pre>
```

describe

Basic descriptive statistics useful for psychometrics

Description

There are many summary statistics available in R; this function provides the ones most useful for scale construction and item analysis in classic psychometrics. Range is most useful for the first pass in a data set, to check for coding errors.

Usage

```
describe(x, digits = 2, na.rm = TRUE, interp=FALSE, skew = TRUE, ranges = TRUE, trim=.1)
```

Arguments

x	A data frame or matrix
digits	How many significant digits to report
na.rm	The default is to delete missing data. na.rm=FALSE will delete the case.
interp	Should the median be standard or interpolated
skew	Should the skew and kurtosis be calculated?
ranges	Should the range be calculated?
trim	trim=.1 – trim means by dropping the top and bottom trim fraction

Details

In basic data analysis it is vital to get basic descriptive statistics. Procedures such as summary and hmisc::describe do so. The describe function in the psych package is meant to produce the most frequently requested stats in psychometric and psychology studies, and to produce them in an easy to read data.frame. The results from describe can be used in graphics functions (e.g., error.crosses).

The range statistics (min, max, range) are most useful for data checking to detect coding errors, and should be found in early analyses of the data.

Although describe will work on data frames as well as matrices, it is important to realize that for data frames, descriptive statistics will be reported only for those variables where this makes sense (i.e., not for factors or for alphanumeric data).

In a typical study, one might read the data in from the clipboard (read.clipboard), show the splom plot of the correlations (pairs.panels), and then describe the data.

na.rm=FALSE is equivalent to describe(na.omit(x))

Value

A data.frame of the relevant statistics: item name item number number of valid cases mean standard deviation trimmed mean (with trim defaulting to .1) median (standard or interpolated mad: median absolute deviation (from the median) minimum maximum skew kurtosis standard error

Note

Describe uses either the mean or colMeans functions depending upon whether the data are a data.frame or a matrix. The mean function supplies means for the columns of a data.frame, but the overall mean for a matrix. Mean will throw a warning for non-numeric data, but colMeans stops with non-numeric data. Thus, the describe function uses either mean (for data frames) or colMeans (for matrices). This is true for skew and kurtosi as well.

Author(s)

```
http://personality-project.org/revelle.html
```

Maintainer: William Revelle (revelle@northwestern.edu)

See Also

describe.by, skew, kurtosi interp.median, pairs.panels, read.clipboard, error.crosses

```
describe(attitude)
describe(attitude,skew=FALSE) #attitude is taken from R data sets
```

eigen.loadings

Convert eigen vectors and eigen values to the more normal (for psychologists) component loadings

Description

The default procedures for principal component returns values not immediately equivalent to the loadings from a factor analysis. eigen loadings translates them into the more typical metric of eigen vectors multiplied by the squareroot of the eigenvalues. This lets us find pseudo factor loadings if we have used princomp or eigen.

If we use principal to do our principal components analysis, then we do not need this routine.

Usage

```
eigen.loadings(x)
```

Arguments

x

the output from eigen or a list of class princomp derived from princomp

Value

A matrix of Principal Component loadings more typical for what is expected in psychometrics. That is, they are scaled by the square root of the eigenvalues.

Note

Useful for SAPA analyses

Author(s)

```
\ revelle@northwestern.edu \>
http://personality-project.org/revelle.html
```

Examples

```
x <- eigen(Harman74.cor$cov)
x$vectors[1:8,1:4] #as they appear from eigen
y <- princomp(covmat=Harman74.cor$cov)
y$loadings[1:8,1:4] #as they appear from princomp
eigen.loadings(x)[1:8,1:4] # rescaled by the eigen values</pre>
```

Description

For teaching correlation, it is useful to draw ellipses around the mean to reflect the correlation. This variation of the ellipse function from John Fox's car package does so. Input may be either two vectors or a matrix or data.frame. In the latter cases, if the number of variables >2, then the ellipses are done in the pairs.panels function. Ellipses may be added to existing plots.

Usage

```
ellipses(x, y = NULL, add = FALSE, smooth=TRUE, lm=FALSE, data=TRUE, n = 2, span=2/3, iter=3, col
```

Arguments

x	a vector,matrix, or data.frame
у	Optional second vector
add	Should a new plot be created, or should it be added to?
smooth	smooth = TRUE -> draw a loess fit
lm	$lm=TRUE \rightarrow draw$ the linear fit
data	data=TRUE implies draw the data points
n	Should 1 or 2 ellipses be drawn
span	averaging window parameter for the lowess fit
iter	iteration parameter for lowess
col	color of ellipses (default is red
xlab	label for the x axis
ylab	label for the y axis
	Other parameters for plotting

Details

Ellipse dimensions are calculated from the correlation between the x and y variables and are scaled as sqrt(1+r) and sqrt(1-r).

Value

A single plot (for 2 vectors or data frames with fewer than 3 variables. Otherwise a call is made to pairs.panels.

Note

Adapted from John Fox's ellipse and data.ellipse functions.

Author(s)

William Revelle

References

Galton, Francis (1888), Co-relations and their measurement. Proceedings of the Royal Society. London Series, 45, 135-145.

See Also

```
pairs.panels
```

Examples

```
data(galton)
ellipses(galton,lm=TRUE)
ellipses(galton$parent,galton$child,xlab="Mid Parent Height",ylab="Child Height") #input are two vectors
data(sat.act)
ellipses(sat.act) #shows the pairs.panels ellipses
```

epi.bfi

13 personality scales from the Eysenck Personality Inventory and Big 5 inventory

Description

A small data set of 5 scales from the Eysenck Personality Inventory, 5 from a Big 5 inventory, a Beck Depression Inventory, and State and Trait Anxiety measures. Used for demonstrations of correlations, regressions, graphic displays.

Usage

```
data(epi.bfi)
```

Format

A data frame with 231 observations on the following 13 variables.

```
epiE EPI Extraversion
epiS EPI Sociability (a subset of Extraversion items
epiImp EPI Impulsivity (a subset of Extraversion items
epilie EPI Lie scale
epiNeur EPI neuroticism
bfagree Big 5 inventory (from the IPIP) measure of Agreeableness
bfcon Big 5 Conscientiousness
bfext Big 5 Extraversion
```

```
bfneur Big 5 Neuroticism
bfopen Big 5 Openness
bdi Beck Depression scale
traitanx Trait Anxiety
stateanx State Anxiety
```

Details

Self report personality scales tend to measure the "Giant 2" of Extraversion and Neuroticism or the "Big 5" of Extraversion, Neuroticism, Agreeableness, Conscientiousness, and Openness. Here is a small data set from Northwestern University undergraduates with scores on the Eysenck Personality Inventory (EPI) and a Big 5 inventory taken from the International Personality Item Pool.

Source

Data were collected at the Personality, Motivation, and Cognition Lab (PMCLab) at Northwestern by William Revelle)

References

```
http://personality-project.org/pmc.html
```

Examples

```
data(epi.bfi)
pairs.panels(epi.bfi[,1:5])
describe(epi.bfi)
```

error.bars.by

Plot means and confidence intervals for multiple groups

Description

One of the many functions in R to plot means and confidence intervals. Meant mainly for demonstration purposes for showing the probabilty of replication from multiple samples. Can also be combined with such functions as boxplot to summarize distributions. Means and standard errors for each group are calculated using describe.by.

Usage

```
error.bars.by(x, group,by.var = FALSE,x.cat=TRUE, ylab = "NULL", xlab = "NULL",main="95% Confidence of the confidence of
```

Arguments

х	A data frame or matrix
group	A grouping variable
by.var	A different line for each group (default) or each variable
x.cat	Is the grouping variable categorical (TRUE) or continuous (FALSE
ylab	y label
xlab	x label
main	title for figure
ylim	if specified, the limits for the plot, otherwise based upon the data
ci	What size confidence interval to use
labels	X axis label
pos	where to place text: below, left, above, right
arrow.len	How long should the top of the error bars be?
add	add=FALSE, new plot, add=TRUE, just points and error bars
	other parameters to pass to the plot function, e.g., typ="b" to draw lines,

Details

Drawing the mean +/- a confidence interval is a frequently used function when reporting experimental results. By default, the confidence interval is 1.96 standard errors.

lty="dashed" to draw dashed lines

This function is a wrapper for **error.bars** and allows groups to be organized either as the x axis or as separate lines.

Value

Graphic output showing the means + x% confidence intervals for each group. For ci=1.96, and normal data, this will be the 95% confidence region. For ci=1, the 68% confidence region.

See Also

See Also as error.crosses, error.bars

Examples

```
x <- matrix(rnorm(500),ncol=20)
y <- sample(4,25 ,replace=TRUE)
x <- x+y
error.bars.by(x,y)
error.bars.by(x,y,TRUE)</pre>
```

Plot means and confidence intervals

error.bars

Description

One of the many functions in R to plot means and confidence intervals. Meant mainly for demonstration purposes for showing the probability of replication from multiple samples. Can also be combined with such functions as boxplot to summarize distributions. Means and standard errors are calculated from the raw data using describe.

Usage

error.bars(x, ylab = "Dependent Variable",xlab="Independent Variable", main="95% confidence line

Arguments

x	A data frame or matrix
ylab	y label
xlab	x label
main	title for figure
ylim	if specified, the limits for the plot, otherwise based upon the data
ci	What size confidence interval to use
labels	X axis label
pos	where to place text: below, left, above, right
arrow.len	How long should the top of the error bars be?
add	add=FALSE, new plot, add=TRUE, just points and error bars
	other parameters to pass to the plot function, e.g., typ="b" to draw lines, lty="dashed" to draw dashed lines

Details

Drawing the mean +/- a confidence interval is a frequently used function when reporting experimental results. By default, the confidence interval is 1.96 standard errors.

Value

Graphic output showing the means + x

Author(s)

William Revelle

See Also

See Also as error.crosses, error.bars.by

Examples

```
x <- matrix(rnorm(500),ncol=20)
error.bars(x)
#now do a boxplot and then add error bars
x.df <- as.data.frame(x)
boxplot(x.df)
error.bars(x.df, add=TRUE)
error.bars(attitude) #another example</pre>
```

error.crosses

Plot x and y error bars

Description

Given two vectors of data, plot the means and show standard errors in both X and Y directions.

Usage

```
error.crosses(x, y, labels = NULL, pos = NULL, arrow.len = 0.2, ...)
```

Arguments

x A vector of summary statistics (from Describe)

y A second vector of summary statistics (also from Describe)

labels name the pair

pos Labels are located where with respect to the mean?

arrow.len Arrow length

... Other parameters for plot

Details

For an example of two way error bars describing the effects of mood manipulations upon positive and negative affect, see http://personality-project.org/revelle/publications/happy-sad-appendix/FIG.A-6.pdf

The second example shows how error crosses can be done for multiple variables where the grouping variable is found dynamically.

Author(s)

William Revelle ⟨revelle@northwestern.edu⟩

See Also

```
See Also as error.bars, error.bars.by,describe.by
```

Examples

```
desc <- describe(attitude)
x <- desc[1,]
y <- desc[2,]
plot(x$mean,y$mean,xlab=rownames(x),ylab=rownames(y)) #in graphics window
error.crosses(x,y) #in graphics window
#now for a bit more complicated plotting
desc <- describe.by(attitude,(attitude[,7]>41)) #select a high and low group
g1 <- desc$'FALSE'
g2 <- desc$'TRUE'
plot(g1$mean,g2$mean,xlab = "Low Advance",ylab="High Advance",xlim=c(30,80),ylim=c(50,80))
error.crosses(g1,g2,labels=rownames(g1),pos=rep(1,7))
title("Attitudes grouped by high and low scores on Advance")</pre>
```

fa.graph

Graph factor loading matrices

Description

Factor analysis or principal components analysis results are typically interpreted in terms of the major loadings on each factor. These structures may be represented as a table of loadings or graphically, where all loadings with an absolute value > some cut point are represented as an edge (path).

Usage

```
fa.graph(fa.results, out.file = NULL, labels = NULL, cut = 0.3, simple = TRUE, size = c(8, 6),
```

Arguments

fa.results The output of factor analysis or principal components analysis out.file If it exists, a dot representation of the graph will be stored here

labels Variable labels

cut Loadings with abs(loading) > cut will be shown

simple Only the biggest loading per item is shown

size

node.font
edge.font
rank.direction

A graphNEL graph with directed edges Number of Nodes = 5 Number of Edges = 4

4 congeneric measures

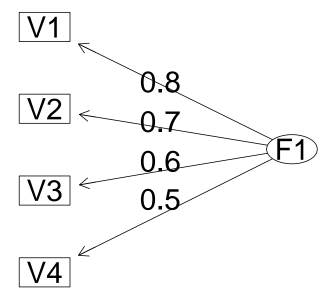


Figure 2: Congeneric tests have one common factor and can differ in their error variances. A demonstration of congeneric.sim, factor.pa, and fa.graph.

digits	Number of digits to show as an edgelable
title	Graphic title
	other parameters

Details

Path diagram representations have become standard in confirmatory factor analysis, but are not yet common in exploratory factor analysis. Representing factor structures graphically helps some people understand the structure.

Value

A graph is drawn using rgraphviz. If an output file is specified, the graph instructions are also saved in the dot language.

Note

Requires Rgraphviz. For the Mac, there are occasional difficulties installing Rgraphviz from Bioconductor in that some libraries are misplaced and need to be relinked using X11.

As of June 1, 2007 there is an occasionally strange result when using the simple=FALSE option in Sweave.

Author(s)

William Revelle

See Also

```
omega.graph, ICLUST.graph
```

Examples

```
test.simple <- factor.pa(item.sim(16),2)
if(require(Rgraphviz)) {fa.graph(test.simple) }</pre>
```

fa.parallel

Scree plots of data or correlation matrix compared to random "parallel" matrices

Description

One way to determine the number of factors or components in a data matrix or a correlation matrix is to examine the "scree" plot of the successive eigenvalues. Sharp breaks in the plot suggest the appropriate number of components or factors to extract. "Parallel" analysis is an alternative technique that compares the scree of the observed data with that of a random data matrix of the same size as the original.

Parallel Analysis Scree Plots

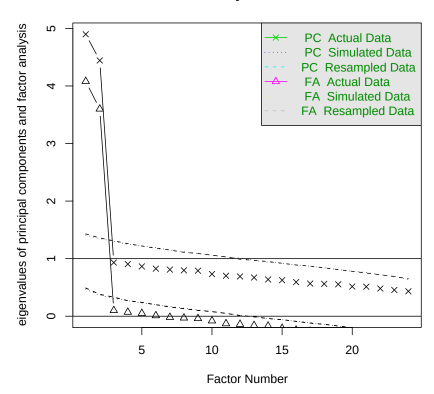


Figure 3: Plots of eigen values for real and simulated "parallel" data give two ways to determine the optimal number of factors: Examine the scree for large breaks or find when the eigen values of random data are greater than for the real data. For this data set, both methods suggest two factors are optimal.

Usage

fa.parallel(x, n.obs = NULL, fa="both", main = "Parallel Analysis Scree Plots",ntrials=20,error

Arguments

х	A data frame or data matrix of scores. If the matrix is square, it is assumed to be a correlation matrix. Otherwise, correlations (with pairwise deletion) will be found
n.obs	n.obs=0 implies a data matrix/data.frame. Otherwise, how many cases were used to find the correlations.
fa	show the eigen values for a principal components (fa="pc") or a principal axis factor analysis (fa="fa") or both principal components and principal factors (fa="both")
main	a title for the analysis
ntrials	Number of simulated analyses to perform
error.bars	Should error.bars be plotted (default = $FALSE$)

Details

Cattell's "scree" test is one of most simple tests for the number of factors problem. Humphreys and Montanelli's "parallel" analysis is an equally compelling procedure. Other procedures for determining the most optimal number of factors include finding the Very Simple Structure (VSS) criterion (VSS) and Velicer's MAP procedure (included in VSS). fa.parallel plots the eigen values for a principal components and principal factor solution and does the same for random matrices of the same size as the original data matrix. For raw data, the random matrices are 1) a matrix of univariate normal data and 2) random samples (randomized across rows) of the original data.

The means of (n.trials) random solutions are shown. Error bars are usually very small and are suppressed by default but can be shown if requested.

Value

A plot of the eigen values for the original data, ntrials of resampling of the original data, and of a equivalent size matrix of random normal deviates. If the data are a correlation matrix, specify the number of observations.

Author(s)

William Revelle

See Also

VSS, VSS. plot, VSS. parallel

Examples

```
test.data <- Harman74.cor$cov
fa.parallel(test.data,n.obs=200)
fa.parallel(attitude)
#</pre>
```

factor.congruence

Coefficient of factor congruence

Description

Given two sets of factor loadings, report their degree of congruence.

Usage

```
factor.congruence(x, y,digits=2)
```

Arguments

x A matrix of factor loadings

y A second matrix of factor loadings

digits Round off to digits

Details

Find the coefficient of factor congruence between two sets of factor loadings.

It is an interesting exercise to compare factor congruences with the correlations of factor loadings. Factor congruences are based upon the raw cross products, while correlations are based upon centered cross products.

Input may either be matrices or factor analysis output (which includes a loadings object), or a mixture of the two.

Factor congruences are the cosines of pairs vectors defined by the loadings matrix and based at the origin. The correlations of factor loadings are cinoses of the vectors based at the mean loading for each factor.

Value

A matrix of factor congruences.

Author(s)

```
\( \text{revelle@northwestern.edu} \)
http://personality-project.org/revelle.html
```

References

Gorsuch, Richard, (1983) Factor Analysis. Lawrence Erlebaum Associates. Revelle, W. (In preparation) An Introduction to Psychometric Theory with applications in R (http://personality-project.org/r/book/)

See Also

```
principal, factor.pa
```

Examples

```
#fa <- factanal(x,4,covmat=Harman74.cor$cov)</pre>
#pc <- principal(Harman74.cor$cov,4)</pre>
#pcv <- varimax(pc$loading)</pre>
#factor.congruence(fa,pcv)
#factor.congruence(pcv,fa)
     Factor1 Factor2 Factor3 Factor4
        1.00
                 0.60
                                  0.55
#PC1
                         0.45
#PC2
        0.44
                 0.49
                          1.00
                                  0.56
#PC3
        0.54
                 0.99
                          0.44
                                  0.55
#PC4
        0.47
                 0.52
                          0.48
                                  0.99
#compare with
#round(cor(fa$loading,pcv$loading),2)
```

factor.fit

How well does the factor model fit a correlation matrix. Part of the VSS package

Description

The basic factor or principal components model is that a correlation or covariance matrix may be reproduced by the product of a factor loading matrix times its transpose: F'F or P'P. One simple index of fit is the 1 - sum squared residuals/sum squared original correlations. This fit index is used by VSS, ICLUST, etc.

Usage

```
factor.fit(r, f)
```

Arguments

r a correlation matrix

f A factor matrix of loadings.

Details

There are probably as many fit indices as there are psychometricians. This fit is a plausible estimate of the amount of reduction in a correlation matrix given a factor model. Note that it is sensitive to the size of the original correlations. That is, if the residuals are small but the original correlations are small, that is a bad fit.

Value

fit

Author(s)

William Revelle

See Also

VSS, ICLUST

Examples

```
## Not run:
#compare the fit of 4 to 3 factors for the Harman 24 variables
fa4 <- factanal(x,4,covmat=Harman74.cor$cov)
round(factor.fit(Harman74.cor$cov,fa4$loading),2)
#[1] 0.9
fa3 <- factanal(x,3,covmat=Harman74.cor$cov)
round(factor.fit(Harman74.cor$cov,fa3$loading),2)
#[1] 0.88
## End(Not run)</pre>
```

factor.model

Find R = F F' + U2 is the basic factor model

Description

The basic factor or principal components model is that a correlation or covariance matrix may be reproduced by the product of a factor loading matrix times its transpose. Find this reproduced matrix. Used by factor.fit, VSS, ICLUST, etc.

Usage

```
factor.model(f)
```

Arguments

f A matrix of loadings.

Value

A correlation or covariance matrix.

Author(s)

```
\( \text{revelle@northwestern.edu } \)
http://personality-project.org/revelle.html
```

References

```
Gorsuch, Richard, (1983) Factor Analysis. Lawrence Erlebaum Associates. Revelle, W. In preparation) An Introduction to Psychometric Theory with applications in R (http://personality-project.org/r/book/)
```

See Also

```
ICLUST.graph,ICLUST.cluster, cluster.fit , VSS, omega
```

Examples

```
f2 <- matrix(c(.9,.8,.7,rep(0,6),.6,.7,.8),ncol=2)
mod <- factor.model(f2)
round(mod,2)</pre>
```

factor.pa

Principal Axis Factor Analysis

Description

Among the many ways to do factor analysis, one of the most conventional is principal axes. An eigen value decomposition of a correlation matrix is done and then the communalities for each variable are estimated by the first n factors. These communalities are entered onto the diagonal and the procedure is repeated until the $\operatorname{sum}(\operatorname{diag}(r))$ does not vary. For well behaved matrices, maximum likelihood factor analysis (factanal) is probably preferred.

Usage

```
factor.pa(r, nfactors=1, residuals = FALSE, rotate = "varimax",n.obs = NULL,
scores = FALSE,SMC=TRUE, missing=FALSE,impute="median",min.err = 0.001, digits = 2, max.iter =
```

A graphNEL graph with directed edges Number of Nodes = 28 Number of Edges = 35

Harman's 24 tests of mental ability

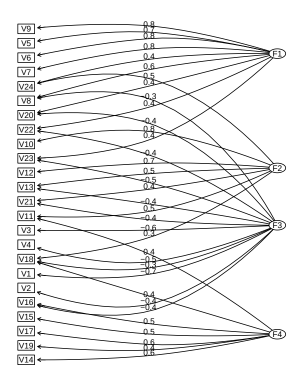


Figure 4: Four factors of the Harman 24 mental tests suggest that a simple structure is difficult to obtain. Note the cross loadings for several items. This figure was created with factor.pa and omega.graph. Compare with a hierarchical solution using the omega function (figure 7) or a cluster solution with ICLUST (figure 5).

Arguments

r	A correlation matrix or a raw data matrix. If raw data, the correlation matrix will be found using pairwise deletion.
nfactors	Number of factors to extract, default is 1
residuals	Should the residual matrix be shown
rotate	"none", "varimax", "promax" or "oblimin" are possible rotations of the solution.
n.obs	Number of observations used to find the correlation matrix if using a correlation matrix. Used for finding the goodness of fit statistics.
scores	If TRUE, estimate factor scores
SMC	Use squared multiple correlations (SMC=TRUE) or use 1 as initial communality estimate. Try using 1 if imaginary eigen values are reported.
missing	if scores are TRUE, and missing=TRUE, then impute missing values using either the median or the mean $$
impute	"median" or "mean" values are used to replace missing values
min.err	Iterate until the change in communalities is less than min.err
digits	How many digits of output should be returned
max.iter	Maximum number of iterations for convergence
symmetric	$\operatorname{symmetric=} \operatorname{TRUE}$ forces symmetry by just looking at the lower off diagonal values
warnings	warnings=TRUE => warn if number of factors is too many

Details

Factor analysis is an attempt to approximate a correlation or covariance matrix with one of lesser rank. The basic model is that ${}_{n}R_{n} \approx_{n} F_{kk}F'_{n} + U^{2}$ where k is much less than n. There are many ways to do factor analysis, and maximum likelihood procedures are probably the most preferred (see factanal). The existence of uniquenesses is what distinguishes factor analysis from principal components analysis (e.g., principal).

Principal axes factor analysis has a long history in exploratory analysis and is a straightforward procedure. Successive eigen value decompositions are done on a correlation matrix with the diagonal replaced with diag (FF') until $\operatorname{sum}(\operatorname{diag}(FF'))$ does not change (very much). The current limit of max.iter =50 seems to work for most problems, but the Holzinger-Harmon 24 variable problem needs about 203 iterations to converge for a 5 factor solution.

Principal axes may be used in cases when maximum likelihood solutions fail to converge.

The algorithm does not attempt to find the best (as defined by a maximum likelihood criterion) solution, but rather one that converges rapidly using successive eigen value decompositions. The maximum likelihood criterion of fit and the associated chi square value are reported, and will be worse than that found using maximum likelihood procedures.

Although for items, it is typical to find factor scores by scoring the salient items (using, e.g., score.items factor scores can be estimated by regression.

Value

values Eigen values of the final solution

communality Communality estimates for each item. These are merely the sum of

squared factor loadings for that item.

rotation which rotation was requested?

n.obs number of observations specified or found

loadings An item by factor loading matrix of class "loadings" Suitable for use in

other programs (e.g., GPA rotation or factor2cluster.

fit How well does the factor model reproduce the correlation matrix. (See

VSS, ICLUST, and principal for this fit statistic.

fit.off how well are the off diagonal elements reproduced?

dof Degrees of Freedom for this model. This is the number of observed corre-

lations minus the number of independent parameters. Let n=Number of

items, nf = number of factors then

dof = n * (n-1)/2 - n * nf + nf * (nf - 1)/2

objective value of the function that is minimized by maximum likelihood proce-

dures. This is reported for comparison purposes and as a way to estimate

chi square goodness of fit. The objective function is

 $f = log(trace((FF' + U2)^{-1}R) - log(|(FF' + U2)^{-1}R|) - n.items.$

STATISTIC If the number of observations is specified or found, this is a chi square

based upon the objective function, f. Using the formula from factanal (which

seems to be Bartlett's test):

 $\chi^2 = (n.obs - 1 - (2 * p + 5)/6 - (2 * factors)/3)) * f$

PVAL If n.obs > 0, then what is the probability of observing a chisquare this

large or larger?

phi If oblique rotations (using oblimin from the GPArotation package) are

requested, what is the interfactor correlation.

communality.iterations

The history of the communality estimates. Probably only useful for teach-

ing what happens in the process of iterative fitting.

residual If residuals are requested, this is the matrix of residual correlations after

the factor model is applied.

Author(s)

William Revelle

References

Gorsuch, Richard, (1983) Factor Analysis. Lawrence Erlebaum Associates.

See Also

principal, VSS, ICLUST

Examples

```
#using the Harman 24 mental tests, compare a principal factor with a principal components solution
pc <- principal(Harman74.cor$cov,4,rotate="varimax")
pa <- factor.pa(Harman74.cor$cov,4,rotate="varimax")
round(factor.congruence(pc,pa),2)

#then compare with a maximum likelihood solution using factanal
mle <- factanal(x,4,covmat=Harman74.cor$cov)
round(factor.congruence(mle,pa),2)
#note that the order of factors and the sign of some of factors differ

#finally, compare the unrotated factor and pca solutions
pc1 <- principal(Harman74.cor$cov,4,rotate="none")
pa1 <- factor.pa(Harman74.cor$cov,4,rotate="none")
round(factor.congruence(pc1,pa1),2)
#note that the order of factors and the sign of some of factors differ</pre>
```

factor.residuals

 $R^* = R - F F'$

Description

The basic factor or principal components model is that a correlation or covariance matrix may be reproduced by the product of a factor loading matrix times its transpose. Find the residuals of the original minus the reproduced matrix. Used by factor.fit, VSS, ICLUST, etc.

Usage

factor.residuals(r, f)

Arguments

r A correlation matrix

f A factor model matrix or a list of class loadings

Details

The basic factor equation is ${}_{n}R_{n} \approx_{n} F_{kk}F'_{n} + U^{2}$. Residuals are just R* = R - F'F. The residuals should be (but in practice probably rarely are) examined to understand the adequacy of the factor analysis. When doing Factor analysis or Principal Components analysis, one usually continues to extract factors/components until the residuals do not differ from those expected from a random matrix.

Value

rstar is the residual correlation matrix.

Author(s)

Maintainer: William Revelle <revelle@northwestern.edu>

See Also

```
factor.pa, principal, VSS, ICLUST
```

Examples

```
fa2 <- factor.pa(Harman74.cor$cov,2,rotate=TRUE)
fa2resid <- factor.residuals(Harman74.cor$cov,fa2)
fa2resid[1:4,1:4] #residuals with two factors extracted
fa4 <- factor.pa(Harman74.cor$cov,4,rotate=TRUE)
fa4resid <- factor.residuals(Harman74.cor$cov,fa4)
fa4resid[1:4,1:4] #residuals with 4 factors extracted</pre>
```

factor.rotate

"Hand" rotate a factor loading matrix

Description

Given a factor or components matrix, it is sometimes useful to do arbitrary rotations of particular pairs of variables. This supplements the much more powerful rotation package GPArotation and is meant for specific requirements to do unusual rotations.

Usage

```
factor.rotate(f, angle, col1=1, col2=2)
```

Arguments

f	original loading matrix or a data frame (can be output from a factor analysis function
angle	angle (in degrees!) to rotate
col1	column in factor matrix defining the first variable
col2	column in factor matrix defining the second variable

Details

Partly meant as a demonstration of how rotation works, factor.rotate is useful for those cases that require specific rotations that are not available in more advanced packages such as GPArotation.

Value

the resulting rotated matrix of loadings.

Note

For a complete rotation package, see GPArotation

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

References

```
http://personality-project.org/revelle/syllabi/405.syllabus.html
```

Examples

```
#using the Harman 24 mental tests, rotate the 2nd and 3rd factors 45 degrees
pc <- principal(Harman74.cor$cov,4,rotate=TRUE)
pcr45 <- factor.rotate(pc,45,2,3)
pcr90 <- factor.rotate(pcr45,45,2,3)
print(factor.congruence(pc,pcr45),digits=3) #poor congruence with original
print(factor.congruence(pc,pcr90),digits=3) #factor 2 and 3 have been exchanged and 3 flipped</pre>
```

factor2cluster

Extract cluster definitions from factor loadings

Description

Given a factor or principal components loading matrix, assign each item to a cluster corresponding to the largest (signed) factor loading for that item. Essentially, this is a Very Simple Structure approach to cluster definition that corresponds to what most people actually do: highlight the largest loading for each item and ignore the rest.

Usage

```
factor2cluster(loads, cut = 0)
```

Arguments

loads either a matrix of loadings, or the result of a factor analysis/principal

components analyis with a loading component

cut Extract items with absolute loadings > cut

Details

A factor/principal components analysis loading matrix is converted to a cluster (-1,0,1) definition matrix where each item is assigned to one and only one cluster. This is a fast way to extract items that will be unit weighted to form cluster composites. Use this function in combination with cluster.cor to find the correlations of these composite scores.

A typical use in the SAPA project is to form item composites by clustering or factoring (see ICLUST, principal), extract the clusters from these results (factor2cluster), and then form the composite correlation matrix using cluster.cor. The variables in this reduced matrix may then be used in multiple R procedures using mat.regress.

The input may be a matrix of item loadings, or the output from a factor analysis which includes a loadings matrix.

Value

a matrix of -1,0,1 cluster definitions for each item.

Author(s)

```
http://personality-project.org/revelle.html
```

Maintainer: William Revelle (revelle@northwestern.edu)

References

```
http://personality-project.org/r/r.vss.html
```

See Also

```
cluster.cor, factor2cluster, factor.pa, principal, ICLUST
```

Examples

```
## Not run:
f <- factanal(x,4,covmat=Harman74.cor$cov)</pre>
factor2cluster(f)
## End(Not run)
                         Factor1 Factor2 Factor3 Factor4
#VisualPerception
                               0
                                        1
                                                0
#Cubes
                                0
                                        1
                                                 0
                                                         0
#PaperFormBoard
                               0
                                        1
                                                0
                                                         0
                               0
                                        1
                                                0
                                                         0
#Flags
                                        0
                                                0
                                                         0
#GeneralInformation
                               1
                                                         0
#PargraphComprehension
                                1
                                        0
                                                0
#SentenceCompletion
                                        0
                                                0
                                                         0
#WordClassification
                                        0
                                                0
                                                         0
#WordMeaning
                               1
                                        0
                                                0
                                                         0
#Addition
                               0
                                        0
                                                         0
                                                1
#Code
                               0
                                        0
                                                1
                                                         0
#CountingDots
                                        0
                                                 1
                                                         0
                                0
```

#StraightCurvedCapitals	0	0	1	0
#WordRecognition	0	0	0	1
#NumberRecognition	0	0	0	1
#FigureRecognition	0	0	0	1
#ObjectNumber	0	0	0	1
#NumberFigure	0	0	0	1
#FigureWord	0	0	0	1
#Deduction	0	1	0	0
#NumericalPuzzles	0	0	1	0
#ProblemReasoning	0	1	0	0
#SeriesCompletion	0	1	0	0
#ArithmeticProblems	0	0	1	0

fisherz

Fisher r to z and z to r and confidence intervals

Description

convert a correlation to a z score or z to r using the Fisher transformation or find the confidence intervals for a specified correlation

Usage

```
fisherz(rho)
fisherz2r(z)
r.con(rho,n,p=.95,twotailed=TRUE)
r2t(rho,n)
```

Arguments

rho	a Pearson r
z	A Fisher z

n Sample size for confidence intervals

p Confidence intervaltwotailed Treat p as twotailed p

Value

z value corresponding to r (fisherz) \setminus r corresponding to z (fisherz2r) \setminus lower and upper p confidence intervals (r.con) \setminus t with n-2 df (r2t)

Author(s)

Examples

```
cors <- seq(-.9,.9,.1)
zs <- fisherz(cors)
rs <- fisherz2r(zs)
round(zs,2)
n <- 30
r <- seq(0,.9,.1)
rc <- matrix(r.con(r,n),ncol=2)
t <- r*sqrt(n-2)/sqrt(1-r^2)
p <- (1-pt(t,n-2))/2
r.rc <- r.rc <- data.frame(r=r,z=fisherz(r),lower=rc[,1],upper=rc[,2],t=t,p=p)
round(r.rc,2)</pre>
```

galton

Galton's Mid parent child height data

Description

Two of the earliest examples of the correlation coefficient were Francis Galton's data sets on the relationship between mid parent and child height and the similarity of parent generation peas with child peas. This is the data set for the Galton height.

Usage

```
data(galton)
```

Format

A data frame with 928 observations on the following 2 variables.

```
parent Mid Parent heights (in inches)
child Child Height
```

Details

Female heights were adjusted by 1.08 to compensate for sex differences. (This was done in the original data set)

Source

This is just the galton data set from UsingR, slightly rearranged.

References

Stigler, S. M. (1999). Statistics on the Table: The History of Statistical Concepts and Methods. Harvard University Press. Galton, F. (1869). Hereditary Genius: An Inquiry into its Laws and Con-sequences. London: Macmillan.

Wachsmuth, A.W., Wilkinson L., Dallal G.E. (2003). Galton's bend: A previously undiscovered nonlinearity in Galton's family stature regression data. The American Statistician, 57, 190-1922.

Examples

```
data(galton)
describe(galton)
pairs.panels(galton,lm=TRUE)
```

geometric.mean

Find the geometric mean of a vector or columns of a data.frame.

Description

The geometric mean is the nth root of n products or e to the mean log of x. Useful for describing non-normal, i.e., geometric distributions.

Usage

```
geometric.mean(x)
```

Arguments

Х

a vector or data.frame

Details

Useful for teaching how to write functions, also useful for showing the different ways of estimating central tendency.

Value

geometric mean(s) of x or x.df.

Note

Not particularly useful if there are elements that are ≤ 0 .

Author(s)

William Revelle

See Also

```
harmonic.mean, mean
```

Examples

```
x <- seq(1,5)
x2 <- x^2
geometric.mean(x)
geometric.mean(x2)</pre>
```

guttman

Alternative estimates of test reliability

Description

Eight alternative estimates of test reliability include the six discussed by Guttman (1945), four discussed by ten Berge and Zergers (1978) ($\mu_0 \dots \mu_3$) as well as beta (the worst split half, Revelle, 1979), the glb (greatest lowest bound) discussed by Bentler and Woodward (1980), and ω_h and ω_t (McDonald, 1999; Zinbarg et al., 2005).

Usage

```
guttman(r,key=NULL,digits=2)
tenberge(r,digits=2)
glb(r,key=NULL,digits=2)
```

Arguments

r A correlation matrix or raw data matrix.

key a vector of -1, 0, 1 to select or reverse items

digits How many digits of accuracy in the output?

Details

Surprisingly, 104 years after Spearman (1904) introduced the concept of reliability to psychologists, there are still multiple approaches for measuring it. Although very popular, Cronbach's α (1951) underestimates the reliability of a test and over estimates the first factor saturation. The guttman function includes the six estimates discussed by Guttman (1945), four of ten Berge and Zergers (1978), as well as Revelle's β (1979) using ICLUST. The companion function, omega calculates omega hierarchical (ω_h) and omega total (ω_t).

$$\lambda_1 = 1 - \frac{tr(\vec{V_x})}{V_x} = \frac{V_x - tr(\vec{V_x})}{V_x}$$
 (1)

The second bound, λ_2 replaces the diagonal with a function of the square root of the sums of squares of the off diagonal elements. Let $C_2 = \vec{1}(\vec{V} - diag(\vec{V}))^2 \vec{1}'$, then

$$\lambda_2 = \lambda_1 + \frac{\sqrt{\frac{n}{n-1}C_2}}{V_x} = \frac{V_x - tr(\vec{V}_x) + \sqrt{\frac{n}{n-1}C_2}}{V_x}.$$
 (2)

Effectively, this is replacing the diagonal with n * the square root of the average squared off diagonal element.

Guttman's 3rd lower bound, λ_3 , also modifies λ_1 and estimates the true variance of each item as the average covariance between items and is, of course, the same as Cronbach's α .

$$\lambda_3 = \lambda_1 + \frac{\frac{V_X - tr(\vec{V}_X)}{n(n-1)}}{V_X} = \frac{n\lambda_1}{n-1} = \frac{n}{n-1} \left(1 - \frac{tr(\vec{V})_x}{V_x} \right) = \frac{n}{n-1} \frac{V_x - tr(\vec{V}_x)}{V_x} = \alpha \quad (3)$$

This is just replacing the diagonal elements with the average off diagonal elements. $\lambda_2 \geq \lambda_3$ with $\lambda_2 > \lambda_3$ if the covariances are not identical.

 λ_3 and λ_2 are both corrections to λ_1 and this correction may be generalized as an infinite set of successive improvements. (Ten Berge and Zegers, 1978)

$$\mu_r = \frac{1}{V_r} \left(p_o + (p_1 + (p_2 + \dots (p_{r-1} + (p_r)^{1/2})^{1/2} \dots)^{1/2} \right), r = 0, 1, 2, \dots$$
 (4)

where

$$p_h = \sum_{i \neq j} \sigma_{ij}^{2h}, h = 0, 1, 2, \dots r - 1$$

and

$$p_h = \frac{n}{n-1}\sigma_{ij}^{2h}, h = r$$

tenberge & Zegers (1978). Clearly $\mu_0 = \lambda_3 = \alpha$ and $\mu_1 = \lambda_2$. $\mu_r \ge \mu_{r-1} \ge \dots \mu_1 \ge \mu_0$, although the series does not improve much after the first two steps.

Guttman's fourth lower bound, λ_4 was originally proposed as any spit half reliability but has been interpreted as the greatest split half reliability. If \vec{X} is split into two parts, \vec{X}_a and \vec{X}_b , with correlation r_{ab} then

$$\lambda_4 = 2\left(1 - \frac{V_{X_a} + V_{X_b}}{V_X}\right) = \frac{4r_{ab}}{V_x} = \frac{4r_{ab}}{V_{X_a} + V_{X_b} + 2r_{ab}V_{X_a}V_{X_b}}$$
(5)

which is just the normal split half reliability, but in this case, of the most similar splits.

 λ_5 , Guttman's fifth lower bound, replaces the diagonal values with twice the square root of the maximum (across items) of the sums of squared interitem covariances

$$\lambda_5 = \lambda_1 + \frac{2\sqrt{\bar{C}_2}}{V_X}.\tag{6}$$

Although superior to λ_1 , λ_5 underestimates the correction to the diagonal. A better estimate would be analogous to the correction used in λ_3 :

$$\lambda_{5+} = \lambda_1 + \frac{n}{n-1} \frac{2\sqrt{\bar{C}_2}}{V_X}. (7)$$

Guttman's final bound considers the amount of variance in each item that can be accounted for the linear regression of all of the other items (the squared multiple correlation or smc), or more precisely, the variance of the errors, e_i^2 , and is

$$\lambda_6 = 1 - \frac{\sum e_j^2}{V_x} = 1 - \frac{\sum (1 - r_{smc}^2)}{V_x} \tag{8}$$

Guttman's λ_4 is the greatest split half reliability. This is found by combining the output from three different approaches, and seems to work for all test cases yet tried. Lambda 4 is reported as the max of these three algorithms.

The algorithms are

- a) Do an ICLUST of the reversed correlation matrix. ICLUST normally forms the most distinct clusters. By reversing the correlations, it will tend to find the most related cluster. Truly a weird approach but tends to work.
- b) Alternatively, a kmeans clustering of the correlations (with the diagonal replaced with 0 to make pseudo distances) can produce 2 similar clusters.
- c) Clusters identified by assigning items to two clusters based upon their order on the first principal factor. (Highest to cluster 1, next 2 to cluster 2, etc.)

Value

beta	The normal beta estimate of cluster similarity from ICLUST. This is an estimate of the general factor saturation.
tenberge\$mu0	tenBerge mu 1 is functionally alpha
tenberge\$mu2	one of the sequence of estimates $\mathrm{mu}1\\ \mathrm{mu}3$
beta.factor	For experimental purposes, what is the split half based upon the two factor solution?
glb.IC	Greatest split half based upon ICLUST of reversed correlations
glb.Km	Greatest split half based upon a kmeans clustering.
glb.Fa	Greatest split half based upon the items assigned by factor analysis.
glb.max	max of the above estimates
keys	scoring keys from each of the alternative methods of forming best splits

Author(s)

William Revelle

References

Cronbach, L.J. (1951) Coefficient alpha and the internal strucuture of tests. Psychometrika, 16, 297-334.

Guttman, L. (1945). A basis for analyzing test-retest reliability. Psychometrika, 10 (4), 255-282.

Revelle, W. (1979). Hierarchical cluster-analysis and the internal structure of tests. Multivariate Behavioral Research, 14 (1), 57-74.

Ten Berge, J. M. F., & Zegers, F. E. (1978). A series of lower bounds to the reliability of a test. Psychometrika, 43 (4), 575-579.

Zinbarg, R. E., Revelle, W., Yovel, I., & Li, W. (2005). Cronbach's α , Revelle's β , and McDonald's ω_h): Their relations with each other and two alternative conceptualizations of reliability. Psychometrika, 70 (1), 123-133.

See Also

```
omega, ICLUST,
```

Examples

```
data(attitude)
glb(attitude)
guttman(attitude)
```

harmonic.mean

Find the harmonic mean of a vector, matrix, or columns of a data.frame

Description

The harmonic mean is merely the reciprocal of the arithmetic mean of the reciprocals.

Usage

```
harmonic.mean(x)
```

Arguments

х

a vector, matrix, or data.frame

Details

Included as an example for teaching about functions. As well as for a discussion of how to estimate central tendencies.

Value

The harmonic mean(s)

Note

Included as a simple demonstration of how to write a function

Examples

```
x <- seq(1,5)
x2 <- x^2
harmonic.mean(x)
harmonic.mean(x2)</pre>
```

headtail

Combine calls to head and tail

Description

A quick way to show the first and last n lines of a data.frame, matrix, or a text object. Just a pretty call to head and tail

Usage

```
headtail(x,hlength=4,tlength=4,digits=2)
```

Arguments

x A matrix or data frame or free text

hlength The number of lines at the beginning to show

tlength The number of lines at the end to show

digits Round off the data to digits

Value

The first hlength and last tlength lines of a matrix or data frame with an ellipsis in between. If the input is neither a matrix nor data frame, the output will be the first hlength and last tlength lines.

See Also

head and tail

Examples

```
x <- matrix(sample(10,1000,TRUE),ncol=5)
headtail(x,4,8)</pre>
```

Description

Francis Galton introduced the 'co-relation' in 1888 with a paper discussing how to measure the relationship between two variables. His primary example was the relationship between height and forearm length. The data table (cubits) is taken from Galton (1888). Unfortunately, there seem to be some errors in the original data table in that the marginal totals do not match the table.

The data frame, heights, is converted from this table using table2df.

Usage

data(heights)

Format

A data frame with 348 observations on the following 2 variables.

```
height Height in inches
cubit Forearm length in inches
```

Details

Sir Francis Galton (1888) published the first demonstration of the correlation coefficient. The regression (or reversion to mediocrity) of the height to the length of the left forearm (a cubit) was found to .8. The original table cubits is taken from Galton (1888). There seem to be some errors in the table as published in that the row sums do not agree with the actual row sums. These data are used to create a matrix using table2matrix for demonstrations of analysis and displays of the data.

Source

Galton (1888)

References

Galton, Francis (1888) Co-relations and their measurement. Proceedings of the Royal Society. London Series, 45, 135-145,

See Also

```
table2matrix, table2df, cubits, ellipses, galton
```

Examples

```
data(heights)
ellipses(heights,n=1,main="Galton's co-relation data set")
```

ICC Intraclass Correlations (ICC1, ICC2, ICC3 from Shrout and Fleiss)

Description

The Intraclass correlation is used as a measure of association when studying the reliability of raters. Shrout and Fleiss (1979) outline 6 different estimates, that depend upon the particular experimental design. All are implemented.

Usage

ICC(x,digits=2)

Arguments

x a matrix or dataframe of ratingsdigits Round the output to digits

Details

Shrout and Fleiss (1979) consider six cases of reliability of ratings done by k raters on n targets.

1) Each target is rated by a different set of k judges. 2) A random sample of k judges rate each target 3) A fixed set of k judges rate each target. Then, for each of these cases, is reliability to be estimated for a single rating or for the average of k ratings? (The 1 rating case is equivalent to the average intercorrelation, the k rating case to the Spearman Brown adjusted reliability.

Value

ICC2Intraclass corelation for the same random judgesICC3Intraclass corelation for the same fixed judgesICC12Intraclass corelation for the average of different judgesICC22Intraclass corelation for the average of random judgesICC3Intraclass corelation for the average of fixed judges	ICC1	Intraclass correlation for different judges
ICC12 Intraclass corelation for the average of different judges ICC22 Intraclass corelation for the average of random judges	ICC2	Intraclass corelation for the same random judges
ICC22 Intraclass corelation for the average of random judges	ICC3	Intraclass corelation for the same fixed judges
· ·	ICC12	Intraclass corelation for the average of different judges
ICC3 Intraclass corelation for the average of fixed judges	ICC22	Intraclass corelation for the average of random judges
	ICC3	Intraclass corelation for the average of fixed judges

Author(s)

William Revelle

References

Shrout, Patrick E. and Fleiss, Joseph L. Intraclass correlations: uses in assessing rater reliability. Psychological Bulletin, 1979, 86, 420-3428.

Revelle, W. (in prep) An introduction to psychometric theory with applications in R. Springer.

Examples

```
ICC(attitude)
```

ICLUST.cluster

Function to form hierarchical cluster analysis of items

Description

The guts of the ICLUST algorithm. Called by ICLUST.

Usage

```
ICLUST.cluster(r.mat, ICLUST.options)
```

Arguments

A list of options (see ICLUST)

Details

See ICLUST

Value

A list of cluster statistics, described more fully in ICLUST

```
comp1 Description of 'comp1'
comp2 Description of 'comp2'
```

. . .

Note

Although the main code for ICLUST is here in ICLUST.cluster, the more extensive documentation is for ICLUST.

Author(s)

```
William Revelle
Department of Psychology
Northwestern University
Evanston, Illinois

( revelle@northwestern.edu )
http://personality-project.org/revelle.html
```

References

Revelle, W. 1979, Hierarchical Cluster Analysis and the Internal Structure of Tests. Multivariate Behavioral Research, 14, 57-74. http://personality-project.org/revelle/publications/iclust.pdf
See also more extensive documentation at http://personality-project.org/r/r.ICLUST.html

See Also

```
ICLUST.graph,ICLUST, cluster.fit , VSS, omega
```

ICLUST.graph

create control code for ICLUST graphical output

Description

Given a cluster structure determined by ICLUST, create dot code to describe the ICLUST output. To use the dot code, use either http://www.graphviz.org/ Graphviz or a commercial viewer (e.g., OmniGraffle).

Usage

```
ICLUST.graph(ic.results, out.file,min.size=1, short = FALSE,labels=NULL,
size = c(8, 6), node.font = c("Helvetica", 14), edge.font = c("Helvetica", 12),
rank.direction=c("RL","TB","LR","BT"), digits = 2, title = "ICLUST", ...)
```

Arguments

ic.results	output list from ICLUST
out.file	name of output file (defaults to console)
min.size	draw a smaller node (without all the information) for clusters $<\min.size$ – useful for large problems
short	if short==TRUE, don't use variable names
labels	vector of text labels (contents) for the variables
size	size of output
node.font	Font to use for nodes in the graph

```
edge.font Font to use for the labels of the arrows (edges)
rank.direction
LR or RL
digits number of digits to show
title any title
... other options to pass
```

Details

Will create (or overwrite) an output file and print out the dot code to show a cluster structure. This dot file may be imported directly into a dot viewer (e.g., http://www.graphviz.org/). The "dot" language is a powerful graphic description language that is particularly appropriate for viewing cluster output. Commercial graphics programs (e.g., OmniGraffle) can also read (and clean up) dot files.

ICLUST.graph takes the output from ICLUST results and processes it to provide a pretty picture of the results. Original variables shown as rectangles and ordered on the left hand side (if rank direction is RL) of the graph. Clusters are drawn as ellipses and include the alpha, beta, and size of the cluster. Edges show the cluster intercorrelations.

It is possible to trim the output to not show all cluster information. Clusters < min.size are shown as small ovals without alpha, beta, and size information.

Value

Output is a set of dot commands written either to console or to the output file. These commands may then be used as input to any "dot" viewer, e.g., Graphviz.

Author(s)

```
\( \text{revelle@northwestern.edu } \)
http://personality-project.org/revelle.html
```

References

```
ICLUST: http://personality-project.org/r/r.iclust.html
```

See Also

```
VSS.plot, ICLUST
```

Examples

```
## Not run:
test.data <- Harman74.cor$cov
ic.out <- ICLUST(test.data)
out.file <- file.choose(new=TRUE)  #create a new file to write the plot commands to
ICLUST.graph(ic.out,out.file)
now go to graphviz (outside of R) and open the out.file you created
print(ic.out,digits=2)
## End(Not run)</pre>
```

```
#test.data <- Harman74.cor$cov
#my.iclust <- ICLUST(test.data)</pre>
#ICLUST.graph(my.iclust)
#digraph ICLUST {
# rankdir=RL;
# size="8,8";
  node [fontname="Helvetica" fontsize=14 shape=box, width=2];
  edge [fontname="Helvetica" fontsize=12];
# label = "ICLUST";
        fontsize=20;
#V1 [label = VisualPerception];
#V2 [label = Cubes];
#V3 [label = PaperFormBoard];
#V4 [label = Flags];
#V5 [label = GeneralInformation];
#V6 [label = PargraphComprehension];
#V7 [label = SentenceCompletion];
#V8 [label = WordClassification];
#V9 [label = WordMeaning];
#V10 [label = Addition];
#V11 [label = Code];
#V12 [label = CountingDots];
#V13 [label = StraightCurvedCapitals];
#V14 [label = WordRecognition];
#V15 [label = NumberRecognition];
#V16 [label = FigureRecognition];
#V17 [label = ObjectNumber];
#V18 [label = NumberFigure];
#V19 [label = FigureWord];
#V20 [label = Deduction];
#V21 [label = NumericalPuzzles];
#V22 [label = ProblemReasoning];
#V23 [label = SeriesCompletion];
     [label = ArithmeticProblems];
#node [shape=ellipse, width ="1"];
#C1-> V9 [ label = 0.78 ];
#C1-> V5 [ label = 0.78 ];
#C2-> V12 [ label = 0.66 ];
\#C2-> V10 [label = 0.66];
#C3-> V18 [ label = 0.53 ];
#C3-> V17 [ label = 0.53 ];
#C4-> V23 [ label = 0.59 ];
#C4-> V20 [ label = 0.59 ];
#C5-> V13 [ label = 0.61 ];
#C5-> V11 [ label = 0.61 ];
\#C6-> V7 [ label = 0.78 ];
\#C6-> V6 [label = 0.78];
\#C7-> V4 [ label = 0.55 ];
#C7-> V1 [ label = 0.55 ];
```

```
#C8-> V16 [ label = 0.5 ];
#C8-> V14 [ label = 0.49 ];
#C9-> C1 [ label = 0.86 ];
#C9-> C6 [ label = 0.86 ];
\#C10-> C4 [ label = 0.71 ];
#C10-> V22 [ label = 0.62 ];
#C11-> V21 [ label = 0.56 ];
#C11-> V24 [ label = 0.58 ];
#C12-> C10 [ label = 0.76 ];
#C12-> C11 [ label = 0.67 ];
#C13-> C8 [ label = 0.61 ];
#C13-> V15 [ label = 0.49 ];
#C14-> C2 [ label = 0.74 ];
#C14-> C5 [ label = 0.72 ];
#C15-> V3 [ label = 0.48 ];
#C15-> C7 [ label = 0.65 ];
#C16-> V19 [ label = 0.48 ];
#C16-> C3 [ label = 0.64 ];
#C17-> V8 [ label = 0.62 ];
#C17-> C12 [ label = 0.8 ];
#C18-> C17 [ label = 0.82 ];
#C18-> C15 [ label = 0.68 ];
#C19-> C16 [ label = 0.66 ];
#C19-> C13 [ label = 0.65 ];
#C20-> C19 [ label = 0.72 ];
#C20-> C18 [ label = 0.83 ];
#C21-> C20 [ label = 0.87 ];
#C21-> C9 [ label = 0.76 ];
#C22-> 0 [ label = 0 ];
#C22-> 0 [ label = 0 ];
#C23-> 0 [ label = 0 ];
#C23-> 0 [ label = 0 ];
#C1 [label =
                "C1\n alpha= 0.84\n beta= 0.84\nN= 2"];
#C2
    [label =
                "C2\n alpha= 0.74\n beta= 0.74\nN= 2"];
#C3
    [label =
                "C3\n alpha= 0.62\n beta= 0.62\nN= 2"];
#C4
    [label =
                "C4\n alpha= 0.67\n beta= 0.67\nN= 2"];
                "C5\n alpha= 0.7\n beta= 0.7\nN= 2"];
#C5
    [label =
#C6
     [label =
                "C6\n alpha= 0.84\n beta= 0.84\nN= 2"];
#C7
     [label =
                "C7\n alpha= 0.64\n beta= 0.64\nN= 2"];
#C8
    [label =
                "C8\n alpha= 0.58\n beta= 0.58\nN= 2"];
#C9
    [label =
                "C9\n alpha= 0.9\n beta= 0.87\nN= 4"];
#C10 [label =
                "C10\n alpha= 0.74\n beta= 0.71\nN= 3"];
#C11
      [label =
                "C11\n alpha= 0.62\n beta= 0.62\n= 2"];
#C12
      [label =
                 "C12\n alpha= 0.79\n beta= 0.74\nN= 5"];
#C13
                 "C13\n alpha= 0.64\n beta= 0.59\nN= 3"];
      [label =
#C14
      [label =
                 "C14\n alpha= 0.79\n beta= 0.74\nN= 4"];
#C15
      [label =
                 "C15\n alpha= 0.66\n beta= 0.58\n= 3"];
#C16
      [label =
                 "C16\n alpha= 0.65\n beta= 0.57\nN= 3"];
#C17
      [label =
                 "C17\n alpha= 0.81\n beta= 0.71\n= 6"] ;
#C18
      [label =
                 "C18\n alpha= 0.84\n beta= 0.75\nN= 9"];
#C19
      [label =
                 "C19\n alpha= 0.74\n beta= 0.65\n= 6"];
#C20
      [label =
                 "C20\n alpha= 0.87\n beta= 0.74\nN= 15"];
#C21
      [label =
                "C21\n alpha= 0.9\n beta= 0.77\nN= 19"];
```

```
#C22 [label = "C22\n alpha= 0\n beta= 0\nN= 0"];
#C23 [label = "C23\n alpha= 0\n beta= 0\nN= 0"];
#{ rank=same;
#V1;V2;V3;V4;V5;V6;V7;V8;V9;V10;V11;V12;V13;V14;V15;V16;V17;V18;V19;V20;V21;V22;V23;V24;}}
#
#copy the above output to Graphviz and draw it
#see \url{http://personality-project.org/r/r.ICLUST.html} for an example.
```

ICLUST.rgraph

Draw an ICLUST graph using the Rgraphviz package

Description

Given a cluster structure determined by ICLUST, create a rgraphic directly using Rgraphviz. To create dot code to describe the ICLUST output with more precision, use ICLUST.graph. As an option, dot code is also generated and saved in a file. To use the dot code, use either http://www.graphviz.org/ Graphviz or a commercial viewer (e.g., OmniGraffle).

Usage

```
ICLUST.rgraph(ic.results, out.file = NULL, min.size = 1, short = FALSE, labels = NULL, size = 0
```

Arguments

output list from ICLUST ic.results out.file File name to save optional dot code. min.size draw a smaller node (without all the information) for clusters < min.size - useful for large problems if short==TRUE, don't use variable names short vector of text labels (contents) for the variables labels size size of output node.font Font to use for nodes in the graph edge.font Font to use for the labels of the arrows (edges) rank.direction LR or TB or RL digits number of digits to show any title title The variable labels can be a different size than the other nodes. This is label.font particularly helpful if the number of variables is large or the labels are long. other options to pass

Details

Will create (or overwrite) an output file and print out the dot code to show a cluster structure. This dot file may be imported directly into a dot viewer (e.g., http://www.graphviz.org/). The "dot" language is a powerful graphic description language that is particularly appropriate for viewing cluster output. Commercial graphics programs (e.g., OmniGraffle) can also read (and clean up) dot files.

ICLUST.rgraph takes the output from ICLUST results and processes it to provide a pretty picture of the results. Original variables shown as rectangles and ordered on the left hand side (if rank direction is RL) of the graph. Clusters are drawn as ellipses and include the alpha, beta, and size of the cluster. Edges show the cluster intercorrelations.

It is possible to trim the output to not show all cluster information. Clusters < min.size are shown as small ovals without alpha, beta, and size information.

Value

Output is a set of dot commands written either to console or to the output file. These commands may then be used as input to any "dot" viewer, e.g., Graphviz.

ICLUST.rgraph is a version of ICLUST.graph that uses Rgraphviz to draw on the screen as well.

Additional output is drawn to main graphics screen.

Note

Requires Rgraphviz

Author(s)

References

```
ICLUST: http://personality-project.org/r/r.iclust.html
```

See Also

```
VSS.plot, ICLUST
```

```
test.data <- Harman74.cor$cov
if(require(Rgraphviz) ) {ic.out <- ICLUST(test.data,labels=colnames(test.data)) }</pre>
```

0 1	7	1 1 ,		r	1 ,	1 1.
Sort items	hni	ahsolute	817P	αt	chieter	Loadinas
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ICLUST.sort

Description

Given a cluster analysis or factor analysis loadings matrix, sort the items by the (absolute) size of each column of loadings. Used as part of ICLUST and SAPA analyses.

Usage

```
ICLUST.sort(ic.load, cut = 0, labels = NULL,keys=FALSE)
```

Arguments

ic.load The output from a factor or principal components analysis, or from ICLUST,

or a matrix of loadings.

cut Do not include items in clusters with absolute loadings less than cut

labels for each item.

keys should cluster keys be returned? Useful if clusters scales are to be scored.

Details

When interpreting cluster or factor analysis outputs, is is useful to group the items in terms of which items have their biggest loading on each factor/cluster and then to sort the items by size of the absolute factor loading.

A stable cluster solution will be one in which the output of these cluster definitions does not vary when clusters are formed from the clusters so defined.

With the keys=TRUE option, the resulting cluster keys may be used to score the original data or the correlation matrix to form clusters from the factors.

Value

sorted A data frame of item numbers, item contents, and item x factor loadings.

cluster A matrix of -1, 0, 1s defining each item by the factor/cluster with the row

wise largest absolute loading.

• • •

Note

Although part of the ICLUST set of programs, this is also more useful for factor or principal components analysis.

Author(s)

William Revelle

ICLUST

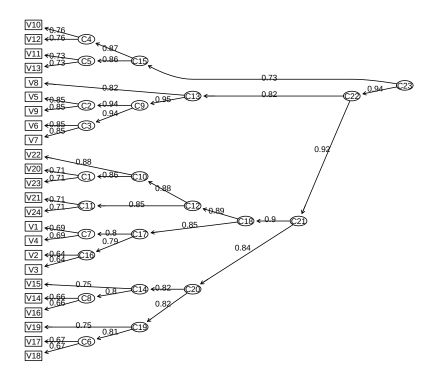


Figure 5: The Harman 24 variables may also be cluster analyzed to highlight the hierarchical structure of abilities. Compare to the hierarchical factor solution using omega (Figure 7) or the orthogonal factor solution (Figure 4)

References

http://personality-project.org/r/r.ICLUST.html

See Also

ICLUST.graph,ICLUST.cluster, cluster.fit , VSS, factor2cluster

Description

A common data reduction technique is to cluster cases (subjects). Less common, but particularly useful in psychological research, is to cluster items (variables). This may be thought of as an alternative to factor analysis, based upon a much simpler model. The cluster model is that the correlations between variables reflect that each item loads on at most one cluster, and that items that load on those clusters correlate as a function of their respective loadings on that cluster and items that define different clusters correlate as a function of their respective cluster loadings and the intercluster correlations. Essentially, the cluster model is a Very Simple Structure factor model of complexity one (see VSS).

This function applies the ICLUST algorithm to hierarchically cluster items to form composite scales. Clusters are combined if coefficients alpha and beta will increase in the new cluster.

Alpha, the mean split half correlation, and beta, the worst split half correlation, are estimates of the reliability and general factor saturation of the test. (See also the omega function to estimate McDonald's coefficient ω_h and ω_t)

Usage

```
ICLUST(r.mat, nclusters=0, alpha=3, beta=1, beta.size=4, alpha.size=3,
correct=TRUE,correct.cluster=TRUE, reverse=TRUE, beta.min=.5, output=1, digits=2,labels=NULL,cu
n.iterations = 0,title="ICLUST",plot=TRUE)

#ICLUST(r.mat)  #use all defaults
#ICLUST(r.mat,nclusters =3)  #use all defaults and if possible stop at 3 clusters
#ICLUST(r.mat, output =3)  #long output shows clustering history
#ICLUST(r.mat, n.iterations =3)  #clean up solution by item reassignment
```

Arguments

r.mat	A correlation matrix or data matrix/data.frame. (If r.mat is not square i.e, a correlation matrix, the data are correlated using pairwise deletion.
nclusters	Extract clusters until nclusters remain (default will extract until the other criteria are met or 1 cluster, whichever happens first). See the discussion below for alternative techniques for specifying the number of clusters.
alpha	Apply the increase in alpha criterion (0) never or for (1) the smaller, (2) the average, or (3) the greater of the separate alphas. (default (2))
beta	Apply the increase in beta criterion (0) never or for (1) the smaller, (2) the average, or (3) the greater of the separate betas. (default (2))
beta.size	Apply the beta criterion after clusters are of beta-size (default $= 4$)
alpha.size	Apply the alpha criterion after clusters are of size alpha. size (default $=$ 3)
correct	Correct correlations for reliability (default $=$ TRUE)
correct.cluste	er
	Correct cluster -sub cluster correlations for reliability of the sub cluster , default is $TRUE)$)
reverse	Reverse negative keyed items (default = $TRUE$

beta.min Stop clustering if the beta is not greater than beta.min (default = .5)

output 1) short, 2) medium, 3) long output (default =1)

labels vector of item content or labels

cut sort cluster loadings > absolute(cut) (default = 0)

n.iterations

digits Precision of digits of output (default = 2)

title Title for this run

plot Should ICLUST.rgraph be called automatically for plotting (requires Rgraphviz

default=TRUE)

Details

Extensive documentation and justification of the algorithm is available in the original MBR 1979 http://personality-project.org/revelle/publications/iclust.pdf paper. Further discussion of the algorithm and sample output is available on the personality-project.org web page: http://personality-project.org/r/r.ICLUST.html

The results are best visualized using ICLUST.graph, the results of which can be saved as a dot file for the Graphviz program. http://www.graphviz.org/. With the installation of Rgraphviz, ICLUST will automatically provide cluster graphs.

A common problem in the social sciences is to construct scales or composites of items to measure constructs of theoretical interest and practical importance. This process frequently involves administering a battery of items from which those that meet certain criteria are selected. These criteria might be rational, empirical, or factorial. A similar problem is to analyze the adequacy of scales that already have been formed and to decide whether the putative constructs are measured properly. Both of these problems have been discussed in numerous texts, as well as in myriad articles. Proponents of various methods have argued for the importance of face validity, discriminant validity, construct validity, factorial homogeneity, and theoretical importance.

Revelle (1979) proposed that hierarchical cluster analysis could be used to estimate a new coefficient (beta) that was an estimate of the general factor saturation of a test. More recently, Zinbarg, Revelle, Yovel and Li (2005) compared McDonald's Omega to Chronbach's alpha and Revelle's beta. They conclude that ω_h is the best estimate. An algorithm for estimating omega is available as part of this package.

ICLUST was completely rewritten for the psych package. Please email me if you want help with this version of ICLUST or if you desire more features.

A requested feature (not yet available) is to specify certain items as forming a cluster. That is, to do confirmatory cluster analysis.

The program currently has three primary functions: cluster, loadings, and graphics.

Although ICLUST will give what it thinks is the best solution in terms of the number of clusters to extract, the user will sometimes disagree. To get more clusters than the default solution, just set the nclusters parameter to the number desired. However, to get fewer than meet the alpha and beta criteria, it is sometimes necessary to set alpha=0 and beta=0 and then set the nclusters to the desired number.

Clustering 24 tests of mental ability

A sample output using the 24 variable problem by Harman can be represented both graphically and in terms of the cluster order. Note that the graphic is created using GraphViz in the dot language. ICLUST.graph produces the dot code for Graphviz. Somewhat lower resolution graphs with fewer options are available in the ICLUST.rgraph function which requires Rgraphviz. Dot code can be viewed directly in Graphviz or can be tweaked using commercial software packages (e.g.,OmniGraffle)

Note that for this problem, with these parameters, the data form one large cluster. (This is consistent with the Very Simple Structure (VSS) output as well, which shows a clear one factor solution for complexity 1 data.) See below for an example with this same data set, but with more stringent parameter settings.

At least for the Harman 24 mental ability measures, it is interesting to compare the cluster pattern matrix with the oblique rotation solution from a factor analysis. The factor congruence of a four factor oblique pattern solution with the four cluster solution is > .99 for three of the four clusters and > .97 for the fourth cluster.

To see the graphic output go to http://personality-project.org/r/r.ICLUST.html or use ICLUST.rgraph (requires Rgraphviz).

Value

title Name of this run
results A list containing

clusters a matrix of -1,0, and 1 values to define cluster membership.

corrected The raw and corrected for alpha reliability cluster intercorrelations.

purified A list of the cluster definitions and cluster loadings of the purified solution.

To show just the most salient items, use the cutoff option in print.psych

cluster.fit, structure.fit, pattern.fit

There are a number of ways to evaluate how well any factor or cluster matrix reproduces the original matrix. Cluster fit considers how well the clusters fit if only correlations with clusters are considered. Structure fit evaluates R=CC while pattern fit evaluate R=C inverse (phi) C where C is the cluster loading matrix, and phi is the intercluster correlation matrix.

Author(s)

William Revelle
Department of Psychology
Northwestern University
Evanston, Illinois

(revelle@northwestern.edu)
http://personality-project.org/revelle.html

References

Revelle, W. Hierarchical Cluster Analysis and the Internal Structure of Tests. Multivariate Behavioral Research, 1979, 14, 57-74. http://personality-project.org/revelle/publications/iclust.pdf

See also more extensive documentation at http://personality-project.org/r/r.ICLUST.html and

Revelle, W. (in prep) An introduction to psychometric theory with applications in R. To be published by Springer. (working draft available at http://personality-project.org/r/book/

See Also

```
ICLUST.graph,ICLUST.cluster, cluster.fit , VSS, omega
```

Examples

```
test.data <- Harman74.cor$cov
ic.out <- ICLUST(test.data)
summary(ic.out)
ic.out <- ICLUST(test.data,nclusters =4)  #use all defaults and stop at 4 clusters
print(ic.out)
plot(ic.out)  #this shows the spatial representation</pre>
```

interp.median

Find the interpolated sample median, quartiles, or specific quantiles for a vector, matrix, or data frame

Description

For data with a limited number of response categories (e.g., attitude items), it is useful treat each response category as range with width, w and linearly interpolate the median, quartiles, or any quantile value within the median response.

Usage

```
interp.median(x, w = 1,na.rm=TRUE)
interp.quantiles(x, q = .5, w = 1,na.rm=TRUE)
interp.quartiles(x,w=1,na.rm=TRUE)
interp.boxplot(x,w=1,na.rm=TRUE)
interp.values(x,w=1,na.rm=TRUE)
interp.qplot.by(y,x,w=1,na.rm=TRUE,xlab="group",ylab="dependent",ylim=NULL,arrow.len=.05,typ="backgroup")
```

Arguments

X	input vector
q	quantile to estimate ($0 < q < 1$
W	category width
у	input vector for interp.qplot.by
na.rm	should missing values be removed

```
xlab x label
ylab Y label
ylim limits for the y axis
arrow.len length of arrow in interp.qplot.by
typ plot type in interp.qplot.by
add add the plot or not
```

... additional parameters to plotting function

Details

If the total number of responses is N, with median, M, and the number of responses at the median value, Nm >1, and Nb= the number of responses less than the median, then with the assumption that the responses are distributed uniformly within the category, the interpolated median is M - .5w + w*(N/2 - Nb)/Nm.

The generalization to 1st, 2nd and 3rd quartiles as well as the general quantiles is straightforward.

A somewhat different generalization allows for graphic presentation of the difference between interpolated and non-interpolated points. This uses the interp.values function.

If the input is a matrix or data frame, quantiles are reported for each variable.

Value

```
im interpolated median(quantile)v interpolated values for all data points
```

See Also

median

```
interp.median(c(1,2,3,3,3)) # compare with median = 3
interp.median(c(1,2,2,5))
interp.quantiles(c(1,2,2,5),.25)
x <- sample(10,100,TRUE)
interp.quartiles(x)
     c(1,1,2,2,2,3,3,3,3,4,5,1,1,1,2,2,3,3,3,3,4,5,1,1,1,2,2,3,3,3,3,4,2)
y \leftarrow c(1,2,3,3,3,3,4,4,4,5,5,1,2,3,3,3,3,4,4,5,5,5,1,5,3,3,3,3,4,4,4,5,5)
x \leftarrow x[order(x)]
                     #sort the data by ascending order to make it clearer
y <- y[order(y)]</pre>
xv <- interp.values(x)</pre>
yv <- interp.values(y)</pre>
barplot(x,space=0,xlab="ordinal position",ylab="value")
lines(1:length(x)-.5,xv)
points(c(length(x)/4, length(x)/2, 3*length(x)/4), interp.quartiles(x))
barplot(y,space=0,xlab="ordinal position",ylab="value")
lines(1:length(y)-.5,yv)
```

```
points(c(length(y)/4,length(y)/2,3*length(y)/4),interp.quartiles(y))
data(galton)
interp.median(galton)
interp.qplot.by(galton$child,galton$parent,ylab="child height"
,xlab="Mid parent height")
```

iqitems

14 multiple choice IQ items

Description

14 multiple choice ability items were included as part of the Synthetic Aperture Personality Assessment (SAPA) web based personality assessment project. The data from 1000 subjects are included here as a demonstration set for scoring multiple choice inventories and doing basic item statistics.

Usage

data(iqitems)

Format

A data frame with 1000 observations on the following 14 variables.

- iq1 In the following number series, what number comes next?
- iq8 Please mark the word that does not match the other words:
- iq10 If you rearrange the letters ATNHIDLA, you will have the name of a:
- iq15 If Jerks are Perks and some Perks are Lerks, then some Jerks are definitely Lerks.
 This statement is:
- iq20 How many total legs do two ducks and three dogs have?
- iq44 Matrix reasoning 2
- iq47 Matrix reasoning 5
- iq2 In the following number series, what number comes next? 1 2 4 7 12
- iq11 The opposite of a 'stubborn' person is a ' ' person.
- iq16 Zach is taller than Matt and Richard is shorter than Zach. Which of the following statements would be most accurate?
- iq32 If the day before yesterday is three days after Saturday then what day is today?
- iq37 In the following alphanumeric series, what letter comes next? Q, S, N, P, L
- iq43 Matrix Reasoning 1
- iq49 Matrix Reasoning 9

Details

14 items were sampled from 54 items given as part of the SAPA project to develop online measures of ability.

Source

```
http://personality-project.org
```

Examples

```
data(iqitems)
iq.keys <- c(4,4,3,1,4,3,2,3,1,4,1,3,4,3)
score.multiple.choice(iq.keys,iqitems)</pre>
```

irt.item.diff.rasch Simple function to estimate item difficulties using IRT concepts

Description

Steps toward a very crude and preliminary IRT program. These two functions estimate item difficulty and discrimination parameters.

Usage

```
irt.item.diff.rasch(items)
irt.discrim(item.diff,theta,items)
```

Arguments

items a matrix of items

item.diff a vector of item difficulties (found by irt.item.diff)

theta ability estimate from irt.person.theta

Details

Item Response Theory (aka "The new psychometrics") models individual responses to items with a logistic function and an individual (theta) and item difficulty (diff) parameter.

irt.item.diff.rasch finds item difficulties with the assumption of theta=0 for all subjects and that all items are equally discriminating.

irt.discrim takes those difficulties and theta estimates from irt.person.rasch to find item discrimination (beta) parameters.

A far better package with these features is the ltm package. The IRT functions in the psych-package are for pedagogical rather than production purposes. They are believed to be accurate, but are not guaranteed. They do seem to be slightly more robust to missing data structures associated with SAPA data sets than the ltm package.

Value

a vector of item difficulties or item discriminations.

Note

Under development. Not recommended for public consumption.

Author(s)

William Revelle

See Also

```
irt.person.rasch
```

irt.1p	Item Response Theory estimate of theta (ability) using a Rasch (like) model

Description

Item Response Theory models individual responses to items by estimating individual ability (theta) and item difficulty (diff) parameters. This is an early and crude attempt to capture this modeling procedure.

Usage

```
irt.person.rasch(diff, items)
irt.0p(items,possible=20)
irt.1p(delta,items)
irt.2p(delta,beta,items)
```

Arguments

diff	A vector of item difficulties –probably taken from irt.item.diff.rasch
items	A matrix of 0.1 items nrows = number of subjects, ncols = number of items
possible	Number of items in the scale – used to determine values of all wrong or all right
delta	delta is the same as diff and is the item difficulty parameter
beta	beta is the item discrimination parameter found in irt.discrim

Details

A very preliminary IRT estimation procedure. Given scores xij for ith individual on jth item

Classical Test Theory ignores item difficulty and defines ability as expected score: ability = theta(i) = x(i.) A zero parameter model rescales these mean scores from 0 to 1 to a quasi logistic scale ranging from - 4 to 4 This is merely a non-linear transform of the raw data to reflect a logistic mapping.

Basic 1 parameter (Rasch) model considers item difficulties (delta j): p(correct on item j for the ith subject | theta i, deltaj) = $1/(1+\exp(\text{deltaj} - \text{thetai}))$ If we have estimates of item difficulty (delta), then we can find theta i by optimization

Two parameter model adds item sensitivity (beta j): p(correct on item j for subject i | thetai, deltaj, betaj) = $1/(1+\exp(\text{betaj *(deltaj-theta i)}))$ Estimate delta, beta, and theta to maximize fit of model to data.

The procedure used here is to first find the item difficulties assuming theta = 0 Then find theta given those deltas Then find beta given delta and theta.

This is not an "official" way to do IRT, but is useful for basic item development.

Value

a data.frame with estimated ability (theta) and quality of fit. (for irt.person.rasch) a data.frame with the raw means, theta0, and the number of items completed

Note

Not recommended for serious use. This code is under development.

Author(s)

William Revelle

See Also

```
irt.item.diff.rasch
```

item.sim

 $Generate\ simulated\ data\ structures\ for\ circumplex\ or\ simple\ structure$

Description

Rotations of factor analysis and principal components analysis solutions typically try to represent correlation matrices as simple structured. An alternative structure, appealing to some, is a circumplex structure where the variables are uniformly spaced on the perimeter of a circle in a two dimensional space. Generating simple structure and circumplex data is straightforward, and is useful for exploring alternative solutions to affect and personality structure.

Simple structure for 32 variables

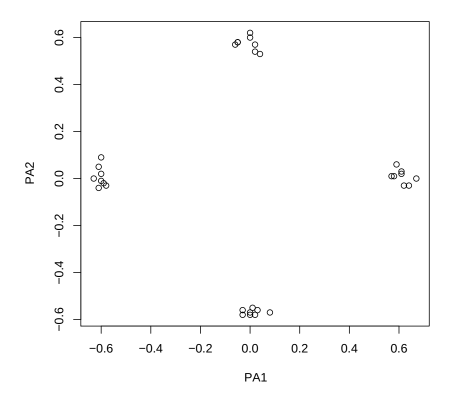


Figure 6: Simple structure is a goal for many factor rotation and extraction procedures. Data may be generated with simple or circumplex structure with varying degrees of skew and correlation. (Compare with Figure)

Usage

```
item.sim(nvar = 72, nsub = 500, circum = FALSE, xloading = 0.6, yloading = 0.6, gloading = 0, zerc.sim(nvar = 72, nsub = 500, circum = TRUE, xloading = 0.6, yloading = 0.6, gloading = 0, zerc.sim(nvar = 72, nsub = 500, circum = FALSE, xloading = 0.6, yloading = 0.6, gloading = 0.6, glo
```

Arguments

nvarNumber of variables to simulatensubNumber of subjects to simulate

circum = TRUE is circumplex structure, FALSE is simple structure

xloading the average loading on the first dimension yloading Average loading on the second dimension

gloading Average loading on a general factor (default=0)

xbias To introduce skew, how far off center is the first dimension

ybias To introduce skew on the second dimension

categorical continuous or categorical variables.

values less than low are forced to low (or 0 in item.dichot)values greater than high are forced to high (or 1 in item.dichot)

truncate Change all values less than cutpoint to cutpoint.

cutpoint What is the cutpoint

Details

This simulation was originally developed to compare the effect of skew on the measurement of affect (see Rafaeli and Revelle, 2005). It has been extended to allow for a general simulation of affect or personality items with either a simple structure or a circumplex structure. Items can be continuous normally distributed, or broken down into n categories (e.g., -2, -1, 0, 1, 2). Items can be distorted by limiting them to these ranges, even though the items have a mean of (e.g., 1).

The addition of item.dichot allows for testing structures with dichotomous items of different difficulty (endorsement) levels. Two factor data with either simple structure or circumplex structure are generated for two sets of items, one giving a score of 1 for all items greater than the low (easy) value, one giving a 1 for all items greater than the high (hard) value. The default values for low and high are 0. That is, all items are assumed to have a 50 percent endorsement rate. To examine the effect of item difficulty, low could be -1, high 1. This will lead to item endorsements of .84 for the easy and .16 for the hard. Within each set of difficulties, the first 1/4 are assigned to the first factor factor, the second to the second factor, the third to the first factor (but with negative loadings) and the fourth to the second factor (but with negative loadings).

Value

A data matrix of (nsub) subjects by (nvar) variables.

Author(s)

William Revelle

References

Variations of a routine used in Rafaeli and Revelle, 2006; Rafaeli, E. & Revelle, W. (2006). A premature consensus: Are happiness and sadness truly opposite affects? Motivation and Emotion.

Acton, G. S. and Revelle, W. (2004) Evaluation of Ten Psychometric Criteria for Circumplex Structure. Methods of Psychological Research Online, Vol. 9, No. 1 http://www.dgps.de/fachgruppen/methoden/mpr-online/issue22/mpr110_10.pdf

See Also

See Also the implementation in this to generate numerous simulations. circ.simulation, circ.tests

Examples

```
round(cor(circ.sim(nvar=8,nsub=200)),2)
plot(factor.pa(circ.sim(16,500),2)$loadings,main="Circumplex Structure") #circumplex structure
#
#
plot(factor.pa(item.sim(16,500),2)$loadings,main="Simple Structure") #simple structure
#
cluster.plot(factor.pa(item.dichot(16,low=0,high=1),2))
```

wkappa

Find Cohen's kappa and weighted kappa coefficients for correlation of two raters

Description

Cohen's kappa (Cohen, 1960) and weighted kappa (Cohen, 1968) may be used to find the agreement of two raters when using nominal scores.

wkappa is (probability of observed matches - probability of expected matches)/(1 - probability of expected matches). Kappa just considers the matches on the main diagonal. Weighted kappa considers off diagonal elements as well

Usage

```
wkappa(x, w = NULL)
```

Arguments

X	Either a two by n data with categorical values from 1 to p or a p x p
	table. If a data array, a table will be found.

w A p x p matrix of weights. If not specified, they are set to be 1 (on the diagonal) and .5(distance from diagonal) off the diagonal.

Details

Some categorical judgments are made using more than two outcomes. For example, two diagnosticians might be asked to categorize patients three ways (e.g., Personality disorder, Neurosis, Psychosis). Just as base rates affect observed cell frequencies in a two by two table, they need to be considered in the n-way table (Cohen, 1960).

Value

```
kappa Unweighted kappa
weighted.kappa
If weights are provided
```

Note

kappa is included in psych more for completeness than necessity. The Kappa function in the vcd package is probably preferred.

To avoid confusion with Kappa (from vcd) or the kappa function from base, the function is named wkappa

Author(s)

William Revelle

References

Cohen, J. (1960). A coefficient of agreement for nominal scales. Educational and Psychological Measurement, 20 37-46

Cohen, J. (1968). Weighted kappa: Nominal scale agreement provision for scaled disagreement or partial credit. Psychological Bulletin, 70, 213-220.

```
cohen <- scan()
0.44 0.05 0.01
0.07 0.20 0.03
0.09 0.05 0.06

cohen <- matrix(cohen,ncol=3)
wkappa(cohen)

fleiss <- scan()
0.53 0.05 0.02</pre>
```

```
0.11 0.14 0.05

0.01 0.06 0.03

fleiss <- matrix(fleiss,ncol=3)

weights <- scan()

1.0000 0.0000 0.4444

0.0000 1.0000 0.6666

0.4444 0.6666 1.0000

weights <- matrix(weights,ncol=3)

wkappa(fleiss,weights)
```

kurtosi

Kurtosis of a vector, matrix, or data frame

Description

Find the kurtosis of a vector, matrix, or dataframe.

Usage

```
kurtosi(x, na.rm = TRUE)
```

Arguments

x vector, matrix, or data frame

na.rm = TRUE removes missing data from the column

Details

Kurtosis in the E1071 package finds the kurtosis for a single vector. This does it for matrices and dataframes. Used in the describe function.

Value

kurtosi a vector of the kurtosis for each column of the matrix

Note

The mean function supplies means for the columns of a data frame, but the overall mean for a matrix. Mean will throw a warning for non-numeric data, but colMeans stops with non-numeric data. Thus, the function uses either mean (for data frames) or colMeans (for matrices). This is true for skew and kurtosi as well.

Author(s)

William Revelle

See Also

skew, describe

Examples

round(kurtosi(attitude),2)

make.hierarchical

Create a population or sample correlation matrix, perhaps with hierarchical structure.

Description

Create a population orthogonal or hierarchical correlation matrix from a set of factor loadings and factor intercorrelations. Samples of size n may be then be drawn from this population. Return either the sample data, sample correlations, or population correlations. This is used to create sample data sets for instruction and demonstration.

Usage

```
make.hierarchical(gload=NULL, fload=NULL, n = 0, raw = FALSE)
```

Arguments

gload	Loadings	of group	factors on	a general factor
-------	----------	----------	------------	------------------

fload Loadings of items on the group factor

n Number of subjects to generate: N=0 => population values

raw =TRUE, report the raw data, raw=FALSE, report the sample corre-

lation matrix.

Details

Many personality and cognitive tests have a hierarchical factor structure. For demonstration purposes, it is useful to be able to create such matrices, either with population values, or sample values.

Given a matrix of item factor loadings (fload) and of loadings of these factors on a general factor (gload), we create a population correlation matrix by using the general factor law (R = F' theta F where theta = g'g).

To create sample values, we use the mvrnorm function from MASS.

The default is to return population correlation matrices. Sample correlation matrices are generated if n > 0. Raw data are returned if raw = TRUE.

The default values for gload and fload create a data matrix discussed by Jensen and Weng, 1994.

Although written to create hierarchical structures, if the gload matrix is all 0, then a non-hierarchical structure will be generated.

Value

a matrix of correlations or a data matrix

Author(s)

William Revelle

References

```
http://personality-project.org/r/r.omega.html
Jensen, A.R., Weng, L.J. (1994) What is a Good g? Intelligence, 18, 231-258.
```

See Also

```
omega, schmid, ICLUST, VSS, mvrnorm
```

gload <- gload<-matrix(c(.9,.8,.7),nrow=3)</pre>

Examples

```
fload <-matrix(c(</pre>
                                                                                                                                                                                                           #a lower order (oblique) factor matrix
                                                             .8,0,0,
                                                              .7,0,.0,
                                                              .6,0,.0,
                                                                 0,.7,.0,
                                                                 0,.6,.0,
                                                                 0,.5,0,
                                                                 0,0,.6,
                                                                 0,0,.5,
                                                                 0,0,.4),
                                                                                                                             ncol=3,byrow=TRUE)
 jensen <- make.hierarchical(gload,fload)</pre>
                                                                                                                                                                                                                                                  #the test set used by omega
round(jensen,2)
fload <- matrix(c(c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,.7,.6),rep(0,20)),c(c(.9,.8,
gload <- matrix(rep(0,5))</pre>
five.factor <- make.hierarchical(gload,fload,500,TRUE) #create sample data set</pre>
```

a higher order factor matrix

make.keys

Create a keys matrix for use by score.items or cluster.cor

Description

When scoring items by forming composite scales either from the raw data using score.items or from the correlatio matrix using cluster.cor, it is necessary to create a keys matrix. This is just a short cut for doing so.

Usage

```
make.keys(nvars, keys.list, key.labels = NULL, item.labels = NULL)
```

Arguments

nvars Number of variables items to be scored

keys.list A list of the scoring keys, one element for each scale

key.labels Labels for the scales can be specified here, or in the key.list item.labels Typically, just the colnames of the items data matrix.

Value

keys a nvars x nkeys matrix of -1, 0, or 1s describing how to score each scale.

nkeys is the length of the keys.list

See Also

```
score.items, cluster.cor,
```

Examples

mat.cortest

Chi square test of whether a single matrix is an identity matrix, or a pair of matrices are equal.

Description

Steiger (1980) pointed out that the sum of the squared elements of a correlation matrix, or the Fisher z score equivalents, is distributed as chi square under the null hypothesis that the values are zero (i.e., elements of the identity matrix). This is particularly useful for examining whether correlations in a single matrix differ from zero or for comparing two matrices. Jennrich (1970 also examined tests of differences between matrices.

Usage

```
normal.cortest(R1, R2 = NULL, n1 = NULL, n2 = NULL, fisher = TRUE)
mat.cortest(R1,R2=NULL,n1=NULL,n2 = NULL)
jennrich.cortest(R1,R2,n1=NULL, n2=NULL)
```

Arguments

R1	A correlation matrix. (If R1 is not rectangular, the correlations are found).
R2	A correlation matrix. If R2 is not rectangular, the correlations are found. If R2 is NULL, then the test is just whether R1 is an identity matrix.
n1	Sample size of R1
n2	Sample size of R2
fisher	Fisher z transform the correlations?

Details

There are several ways to test if a matrix is the identity matrix. The most well known is the chi square test of Bartlett (1951) and Box (1949). A very straightforward test, discussed by Steiger (1980) is to find the sum of the squared correlations or the sum of the squared Fisher transformed correlations. Under the null hypothesis that all the correlations are equal, this sum is distributed as chi square.

Yet another test, is the Jennrich (1970) test of the equality of two matrices.

Value

chi2	The chi square statistic
df	Degrees of freedom for the Chi Square
prob	The probability of observing the Chi Square under the null hypothesis.

Note

Both the jennrich.cortest and normal.cortest are probably overly stringenent. The ChiSquare values for pairs of random samples from the same population are larger than would be expected. This is a good test for rejecting the null of no differences.

Author(s)

William Revelle

References

Steiger, James H. (1980) Testing pattern hypotheses on correlation matrices: alternative statistics and some empirical results. Multivariate Behavioral Research, 15, 335-352.

See Also

bartlett.cortest, jennrich.cortest, mat.cortest

Examples

```
x <- matrix(rnorm(1000),ncol=10)
y <- matrix(rnorm(500),ncol=10)
normal.cortest(x)  #just test if this matrix is an identity
normal.cortest(x,y) #do these two matrices differ?
mat.cortest(x)
mat.cortest(x,y)  #twice the degrees of freedom as the Jennrich
jennrich.cortest(x,y) #</pre>
```

mat.regress

Multiple Regression from matrix input

Description

Extract subsets of variables (x and y) from a correlation or data matrix matrix and find the multiple correlation and beta weights of the (x) set predicting each member of the (y) set.

Usage

```
mat.regress(m, x, y,n.obs=NULL,digits=2)
```

Arguments

m	a matrix of correlations or, if not square of data
x	the column numbers of the x set (e.g., $c(1,3,5)$
У	the column numbers of the y set (e.g., $c(2,4,6)$
n.obs	If specified, then confidence intervals, etc. are calculated, not needed if raw data are given
digits	round the answer to digits

Details

Although it is more common to calculate multiple regression from raw data, it is, of course, possible to do so from a set of correlations. The input to the function is a square covariance or correlation matrix, as well as the column numbers of the x (predictor) and y (criterion) variables. The program will find correlations if given raw data.

The output is a set of multiple correlations, one for each dependent variable in the y set.

A typical use in the SAPA project is to form item composites by clustering or factoring (see ICLUST, principal), extract the clusters from these results (factor2cluster), and then form the composite correlation matrix using cluster.cor. The variables in this reduced matrix may then be used in multiple R procedures using mat.regress.

Although the overall matrix can have missing correlations, the correlations in the subset of the matrix used for prediction must exist. If the number of observations is entered, then the conventional confidence intervals, statistical significance, and shrinkage estimates are reported.

If the input matrix is rectangular, correlations are found from the data.

Value

beta	the beta weights for each variable in X for each variable in Y
R	The multiple R for each equation (the amount of change a unit in the predictor set leads to in the criterion set).
R2	The multiple R2 (% variance acounted for) for each equation

Author(s)

William Revelle

Maintainer: William Revelle <revelle@northwestern.edu>

See Also

```
cluster.cor, factor2cluster,principal,ICLUST
```

```
## Not run:
test.data <- Harman74.cor$cov
                                  #24 mental variables
#choose 3 of them to regress against another 4 -- arbitrary choice of variables
print(mat.regress(test.data,c(1,2,3),c(4,5,10,12)),digits=2)
## End(Not run)
#gives this output
#print(mat.regress(test.data,c(1,2,3),c(4,5,10,12)),digits=2)
#$beta
                  Flags GeneralInformation Addition CountingDots
#VisualPerception 0.40
                                      0.22
                                               0.16
                                                            0.30
#Cubes
                   0.06
                                      0.18
                                               0.06
                                                            0.05
#PaperFormBoard
                   0.12
                                      0.10
                                              -0.16
                                                            0.00
#$R
              Flags GeneralInformation
                                                 Addition
                                                                CountingDots
              0.49
                                 0.38
                                                    0.18
                                                                       0.32
#$R2
              Flags GeneralInformation
                                                                CountingDots
                                                 Addition
#
               0.24
                                 0.15
                                                     0.03
                                                                        0.10
#
data(attitude)
mat.regress(attitude,c(1:3),c(4:7)) #standardized regression from raw data
```

Description

It is sometimes convenient to add two vectors or matrices in an operation analogous to matrix multiplication. For matrices nXm and mYp, the matrix sum of the i,jth element of $nSp = sum(over\ m)$ of iXm + mYj.

Usage

```
x %+% y
```

Arguments

```
x a n by m matrix (or vector if m=1)

y a m by p matrix (or vector if m=1)
```

Details

Used in such problems as Thurstonian scaling. Although not technically matrix addition, as pointed out by Krus, there are many applications where the sum or difference of two vectors or matrices is a useful operation. An alternative operation for vectors is outer(x, y, y, y) but this does not work for matrices.

Value

a n by p matix of sums

Author(s)

William Revelle

References

Krus, D. J. (2001) Matrix addition. Journal of Visual Statistics, 1, (February, 2001).

```
x <- seq(1,4)
z <- x %+% -t(x)
x
z
#compare with outer(x,-x,FUN="+")
x <- matrix(seq(1,6),ncol=2)
y <- matrix(seq(1,10),nrow=2)
z <- x %+% y</pre>
```

```
y z #but compare this with outer(x ,y,FUN="+")
```

multi.hist

Multiple histograms with density and normal fits on one page

Description

Given a matrix or data.frame, produce histograms for each variable in a "matrix" form. Include normal fits and density distributions for each plot.

The number of rows and columns may be specified, or calculated.

May be used for single variables.

Usage

```
multi.hist(x,nrow=NULL, ncol=NULL,density=TRUE,main="Histogram, Density, and Normal Fit")
```

Arguments

matrix or data.frame
 nrow
 number of rows in the plot
 number of columns in the plot

density density=TRUE, show the normal fits and density distributions

main title for each panel

Author(s)

```
William Revelle
Northwestern University
Evanston, Illinois
⟨ revelle@northwestern.edu ⟩
http://personality-project.org/revelle.html
```

```
#multi.hist(attitude[-1])
```

V1 ← V2 **←** V3 **≺** V4 **← 7** V23 **←** V5 . V6 ← V7 **←** V8 **≺ €**10 V9 • V20 V22 V10. V11 ∢ V12 < V13 < V21 < V24 V14 🗸 V15 **←** V16 **←** V17 < V18 **<** V19

Omega

Figure 7: Hierarchical factor solutions are typical in the ability domain where a g factor is thought to reflect the correlations among lower level factors. An alternative transformation is to ortogonalize the g factor from the residual group factors using the Schmid-Leiman transformation (Figure 8)

V1 € V2 **←** V3 **<** \[\V4] \ 7 V23 **⁴** V5 . V6 **∢** V7 **←** V8 **←** V9 **←** V20 V22 → V10 **→** V11 ← V12 **←** V13 < V21 < V24 ' V14 < . V15 ← V16 < V17 < V18 ⁴ V19

Omega

Figure 8: An alternative to the standard hierarchical factor solutions which are typical in the ability domain is to orthgonalize the g factor from the residual group factors using the Schmid-Leiman transformaton. For the hierarchical solution, see Figure 7

Description

Hierarchical factor structures represent the correlations between variables in terms of a smaller set of correlated factors which themselves can be represented by a higher order factor.

Two alternative solutions to such structures are found by the omega function. The correlated factors solutions represents the effect of the higher level, general factor, through its effect on the correlated factors. The other representation makes use of the Schmid Leiman transformation to find the direct effect of the general factor upon the original variables as well as the effect of orthogonal residual group factors upon the items.

Graphic presentations of these two alternatives are helpful in understanding the structure. omega.graph draws both such structures. Graphs are drawn directly onto the graphics window or expressed in "dot" commands for conversion to graphics using implementations of Graphviz.

Using Graphviz allows the user to clean up the Rgraphviz output (e.g., make the arrows go the right direction!

Usage

```
omega.graph(om.results, out.file = NULL, s1 = TRUE, labels = NULL, size = c(8, 6), node.font =
```

Arguments

om.results The output from the omega function

out.file Optional output file for off line analysis using Graphviz

orthogonal clusters using the Schmid-Leiman transform (sl=TRUE) or

oblique clusters

labels variable labels

size size of graphics window

node.font What font to use for the items

edge.font What font to use for the edge labels

rank.direction

Defaults to left to right

digits Precision of labels

title Figure title

... Other options to pass into the graphics packages

Details

Requires the Rgraphviz package. omega requires the GPArotation package.

Value

```
clust.graph A graph object
```

Note

Requires rgraphviz. – omega requires GPArotation

Author(s)

```
http://personality-project.org/revelle.html
Maintainer: William Revelle \( \rmoreal revelle \( \rmoreal revelle \) revelle (northwestern.edu \)
```

References

```
http://personality-project.org/r/r.omega.html
```

Revelle, W. (in preparation) An Introduction to Psychometric Theory with applications in R. http://personality-project.org/r/book

Revelle, W. (1979). Hierarchical cluster analysis and the internal structure of tests. Multivariate Behavioral Research, 14, 57-74. (http://personality-project.org/revelle/publications/iclust.pdf)

Zinbarg, R.E., Revelle, W., Yovel, I., & Li. W. (2005). Cronbach's Alpha, Revelle's Beta, McDonald's Omega: Their relations with each and two alternative conceptualizations of reliability. Psychometrika. 70, 123-133. http://personality-project.org/revelle/publications/zinbarg.revelle.pmet.05.pdf

Zinbarg, R., Yovel, I., Revelle, W. & McDonald, R. (2006). Estimating generalizability to a universe of indicators that all have one attribute in common: A comparison of estimators for omega. Applied Psychological Measurement, 30, 121-144. DOI: 10.1177/0146621605278814 http://apm.sagepub.com/cgi/reprint/30/2/121

See Also

```
omega, make.hierarchical, ICLUST.rgraph
```

```
#24 mental tests from Holzinger-Swineford-Harman
if(require(GPArotation) ) {om24 <- omega(Harman74.cor$cov,4) } #run omega
if(require(Rgraphviz) ){om24pn <- omega.graph(om24,sl=FALSE)} #show the structure
#
#example hierarchical structure from Jensen and Weng
if(require(GPArotation) ) {jen.omega <- omega(make.hierarchical())}
if(require(Rgraphviz) ) {om.jen <- omega.graph(jen.omega,sl=FALSE) }</pre>
```

Description

McDonald has proposed coefficient omega as an estimate of the general factor saturation of a test. One way to find omega is to do a factor analysis of the original data set, rotate the factors obliquely, do a Schmid Leiman transformation, and then find omega. This function estimates omega as suggested by McDonald by using hierarchical factor analysis (following Jensen).

Usage

omega(m, nfactors, pc = "mle",key = NULL, flip=TRUE, digits=2,title="Omega",sl=TRUE,labels=NULI

Arguments

m	A correlation matrix or a data.frame/matrix of data
nfactors	Number of factors believed to be group factors
рс	pc="pa" for principal axes, pc="pc" for principal components, pc="mle" for maximum likelihood.
key	a vector of $+/-$ 1s to specify the direction of scoring of items. The default is to assume all items are positively keyed, but if some items are reversed scored, then key should be specified.
flip	If flip is TRUE, then items are automatically flipped to have positive correlations on the general factor. Items that have been reversed are shown with a - sign.
digits	if specified, round the output to digits
title	Title for this analysis
sl	If plotting the results, should the Schmid Leiman solution be shown or should the hierarchical solution be shown? (default sl=TRUE)
labels	If plotting, what labels should be applied to the variables
plot	plot=TRUE (default) calls omega.graph, plot =FALSE does not
rotate	What rotation to apply? The default is oblimin, the alternative is simplimax.
• • •	Allows additional parameters to be passed through to the factor routines

Details

"Many scales are assumed by their developers and users to be primarily a measure of one latent variable. When it is also assumed that the scale conforms to the effect indicator model of measurement (as is almost always the case in psychological assessment), it is important to support such an interpretation with evidence regarding the internal structure of that scale. In particular, it is important to examine two related properties pertaining to

the internal structure of such a scale. The first property relates to whether all the indicators forming the scale measure a latent variable in common.

The second internal structural property pertains to the proportion of variance in the scale scores (derived from summing or averaging the indicators) accounted for by this latent variable that is common to all the indicators (Cronbach, 1951; McDonald, 1999; Revelle, 1979). That is, if an effect indicator scale is primarily a measure of one latent variable common to all the indicators forming the scale, then that latent variable should account for the majority of the variance in the scale scores. Put differently, this variance ratio provides important information about the sampling fluctuations when estimating individuals' standing on a latent variable common to all the indicators arising from the sampling of indicators (i.e., when dealing with either Type 2 or Type 12 sampling, to use the terminology of Lord, 1956). That is, this variance proportion can be interpreted as the square of the correlation between the scale score and the latent variable common to all the indicators in the infinite universe of indicators of which the scale indicators are a subset. Put yet another way, this variance ratio is important both as reliability and a validity coefficient. This is a reliability issue as the larger this variance ratio is, the more accurately one can predict an individual's relative standing on the latent variable common to all the scale's indicators based on his or her observed scale score. At the same time, this variance ratio also bears on the construct validity of the scale given that construct validity encompasses the internal structure of a scale." (Zinbarg, Yovel, Revelle, and McDonald, 2006).

McDonald has proposed coefficient omega (hierarchical (ω_h) as an estimate of the general factor saturation of a test. Zinbarg, Revelle, Yovel and Li (2005) http://personality-project.org/revelle/publications/zinbarg.revelle.pmet.05.pdf compare McDonald's ω_h to Cronbach's α and Revelle's β They conclude that ω_h is the best estimate. (See also Zinbarg et al., 2006)

One way to find ω_h is to do a factor analysis of the original data set, rotate the factors obliquely, factor that correlation matrix, do a Schmid-Leiman (schmid) transformation to find general factor loadings, and then find ω_h . Here we present code to do that.

 ω_h differs as a function of how the factors are estimated. Three options are available, pc="pa" does a principle axes factor analysis (factor.pa), pc="mle" uses the factanal function, and pc="pc" does a principal components analysis (principal).

For ability items, it is typically the case that all items will have positive loadings on the general factor. However, for non-cognitive items it is frequently the case that some items are to be scored positively, and some negatively. Although probably better to specify which directions the items are to be scored by specifying a key vector, if flip =TRUE (the default), items will be reversed so that they have positive loadings on the general factor. The keys are reported so that scores can be found using the score.items function.

Output from omega will be shown graphically using the omega.graph function. This requires Rgraphviz to be installed. If Rgraphviz is not available, select plot=FALSE.

 β , an alternative to ω , is defined as the worst split half reliability. It can be estimated by using ICLUST (a hierarchical clustering algorithm originally developed for main frames and written in Fortran and that is now available in R. (For a very complimentary review of why the ICLUST algorithm is useful in scale construction, see Cooksey and Soutar, 2005).

The omega function uses exploratory factor analysis to estimate the ω_h coefficient. It is important to remember that "A recommendation that should be heeded, regardless of the method chosen to estimate ω_h , is to always examine the pattern of the estimated general

factor loadings prior to estimating ω_h . Such an examination constitutes an informal test of the assumption that there is a latent variable common to all of the scale's indicators that can be conducted even in the context of EFA. If the loadings were salient for only a relatively small subset of the indicators, this would suggest that there is no true general factor underlying the covariance matrix. Just such an informal assumption test would have afforded a great deal of protection against the possibility of misinterpreting the misleading ω_h estimates occasionally produced in the simulations reported here." (Zinbarg et al., 2006, p 137).

A simple demonstration of the problem of an omega estimate reflecting just one of two group factors can be found in the last example.

Although omega is uniquely defined only for cases where 3 or more subfactors are extracted, it is sometimes desired to have a two factor solution. This is done by forcing the schmid extraction to treat the two subfactors as having equal loadings. See Zinbarg et al., 2007.

In addition to ω_h , another of McDonald's coefficients is ω_t . This is an estimate of the total reliability of a test.

McDonald's ω_t , which is similar to Guttman's λ_6 , guttman but uses the estimates of uniqueness (u^2) from factor analysis to find e_j^2 . This is based on a decomposition of the variance of a test score, V_x , into four parts: that due to a general factor, \vec{g} , that due to a set of group factors, \vec{f} , (factors common to some but not all of the items), specific factors, \vec{s} unique to each item, and \vec{e} , random error. (Because specific variance can not be distinguished from random error unless the test is given at least twice, some combine these both into error).

Letting

$$\vec{x} = \vec{cg} + \vec{Af} + \vec{Ds} + \vec{e} \tag{9}$$

then the communality of item $_{i}$, based upon general as well as group factors,

$$h_j^2 = c_j^2 + \sum f_{ij}^2 \tag{10}$$

and the unique variance for the item

$$u_j^2 = \sigma_j^2 (1 - h_j^2) \tag{11}$$

may be used to estimate the test reliability. That is, if h_j^2 is the communality of item_j, based upon general as well as group factors, then for standardized items, $e_j^2 = 1 - h_j^2$ and

$$\omega_t = \frac{\vec{1}\vec{c}\vec{c}'\vec{1} + \vec{1}\vec{A}\vec{A}'\vec{1}'}{V_x} = 1 - \frac{\sum(1 - h_j^2)}{V_x} = 1 - \frac{\sum u^2}{V_x}$$
(12)

Because $h_j^2 \ge r_{smc}^2$, $\omega_t \ge \lambda_6$.

It is important to distinguish here between the two ω coefficients of McDonald, 1978 and Equation 6.20a of McDonald, 1999, ω_t and ω_h . While the former is based upon the sum of squared loadings on all the factors, the latter is based upon the sum of the squared loadings on the general factor.

$$\omega_h = \frac{\vec{1}c\vec{c}'\vec{1}}{V_x} \tag{13}$$

Value

omega hierarchical

The ω_h coefficient

omega total The $omega_t$ coefficient

alpha Cronbach's α

schmid The Schmid Leiman transformed factor matrix and associated matrices

schmid\$sl The g factor loadings as well as the residualized factors

schmid\$orthog Varimax rotated solution of the original factors

schmid\$oblique

The oblimin transformed factors

schmid\$fcor the correlation matrix of the oblique factors

schmid\$gloading

The loadings on the higher order, g, factor of the oblimin factors

key A vector of -1 or 1 showing which direction the items were scored.

Note

Requires the GPArotation package

Author(s)

```
http://personality-project.org/revelle.html
```

Maintainer: William Revelle (revelle@northwestern.edu)

References

```
http://personality-project.org/r/r.omega.html
```

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See Also

```
omega.graph ICLUST, ICLUST.graph, VSS, schmid , make.hierarchical
```

Examples

```
## Not run:
test.data <- Harman74.cor$cov
my.omega <- omega(test.data,3)</pre>
print(my.omega,digits=2)
## End(Not run)
#create 9 variables with a hierarchical structure
jen.data <- make.hierarchical()</pre>
#with correlations of
jen.data
#find omega
jen.omega <- omega(jen.data,digits=2)</pre>
jen.omega
#create 8 items with a two factor solution, showing the use of the flip option
#sim2 <- item.sim(8)
#omega(sim2) #an example of misidentification-- remember to look at the loadings matrices.
#apply omega to analyze 6 mental ability tests
data(ability.cov) #has a covariance matrix
omega(ability.cov$cov)
```

p.rep

Find the probability of replication for an F, t, or r and estimate effect size

Description

The probability of replication of an experimental or correlational finding as discussed by Peter Killeen (2005) is the probability of finding an effect in the same direction upon an exact replication. For articles submitted to Psychological Science, p.rep needs to be reported.

F, t, p and r are all estimates of the size of an effect. But F, t, and p also are also a function of the sample size. Effect size, d prime, may be expressed as differences between means compared to within cell standard deviations, or as a correlation coefficient. These functions convert p, F, and t to d prime and the r equivalent.

Usage

```
p.rep(p = 0.05, n=NULL,twotailed = FALSE)
p.rep.f(F,df2,twotailed=FALSE)
p.rep.r(r,n,twotailed=TRUE)
p.rep.t(t,df,df2=NULL,twotailed=TRUE)
```

Arguments

- p conventional probability of statistic (e.g., of F, t, or r)
- F The F statistic

df Degrees of freedom of the t-test, or of the first group if unequal sizes

Degrees of freedom of the denominator of F or the second group in an unequal sizes t test

Correlation coefficient

Total sample size if using r

t t-statistic if doing a t-test or testing significance of a regression slope

twotailed Should a one or two tailed test be used?

Details

The conventional Null Hypothesis Significance Test (NHST) is the likelihood of observing the data given the null hypothesis of no effect. But this tells us nothing about the probability of the null hypothesis. Peter Killeen (2005) introduced the probability of replication as a more useful measure. The probability of replication is the probability that an exact replication study will find a result in the *same direction* as the original result.

p.rep is based upon a 1 tailed probability value of the observed statistic.

Other frequently called for statistics are estimates of the effect size, expressed either as Cohen's d, Hedges g, or the equivalent value of the correlation, r.

For p.rep.t, if the cell sizes are unequal, the effect size estimates are adjusted by the ratio of the mean cell size to the harmonic mean cell size (see Rownow et al., 2000).

Value

p.rep	Probability of replication
dprime	Effect size (Cohen's d) if more than just p is specified
prob	Probability of F, t, or r. Note that this can be either the one-tailed or two tailed probability value.
r.equivalent	For t-tests, the r equivalent to the t (see Rosenthal and Rubin (2003), Rosnow, Rosenthal, and Rubin, 2000))

Note

The p.rep value is the one tailed probability value of obtaining a result in the same direction.

References

Cummings, Geoff (2005) Understanding the average probability of replication: comment on Killeen 2005). Psychological Science, 16, 12, 1002-1004).

Killeen, Peter H. (2005) An alternative to Null-Hypothesis Significance Tests. Psychological Science, $16,\,345\text{-}353$

Rosenthal, R. and Rubin, Donald B.(2003), r-sub(equivalent): A Simple Effect Size Indicator. Psychological Methods, 8, 492-496.

Rosnow, Ralph L., Rosenthal, Robert and Rubin, Donald B. (2000) Contrasts and correlations in effect-size estimation, Psychological Science, 11. 446-453.

Examples

```
p.rep(.05) #probability of replicating a result if the original study had a p = .05 p.rep.f(9.0,98) #probability of replicating a result with F = 9.0 with 98 df p.rep.r(.4,50) #probability of replicating a result if r = .4 with n = 50 p.rep.t(3,98) #probability of replicating a result if t = 3 with df =98 p.rep.t(2.14,84,14) #effect of equal sample sizes (see Rosnow et al.)
```

paired.r

Test the difference between (un)paired correlations

Description

Test the difference between two (paired or unpaired) correlations. Given 3 variables, x, y, z, is the correlation between xy different than that between xz? If y and z are independent, this is a simple t-test of the z transformed rs. But, if they are dependent, it is a bit more complicated.

Usage

```
paired.r(xy, xz, yz=NULL, n, n2=NULL,twotailed=TRUE)
```

Arguments

xy	r(xy)
xz	r(xz)
yz	r(yz)
n	Number of subjects for first group
n2	Number of subjects in second group (if not equal to n)
twotailed	Calculate two or one tailed probability values

Details

To find the z of the difference between two independent correlations, first convert them to z scores using the Fisher r-z transform and then find the z of the difference between the two correlations. The default assumption is that the group sizes are the same, but the test can be done for different size groups by specifying n2.

If the correlations are not independent (i.e., they are from the same sample) then the correlation with the third variable r(yz) must be specified. Find a t statistic for the difference of thee two dependent correlations.

Value

a list containing the calculated t or z values and the associated two (or one) tailed probability.

t test of the difference between two dependent correlations

p probability of the t or of the z

z z test of the difference between two independent correlations

Author(s)

```
William Revelle
Northwestern University
Evanston, Illinois
⟨ revelle@northwestern.edu ⟩
http://personality-project.org/revelle.html
```

See Also

```
p.rep.r, cor.test
```

Examples

```
paired.r(.5,.3, .4, 100) #dependent correlations
paired.r(.5,.3,NULL,100) #independent correlations same sample size
paired.r(.5,.3,NULL, 100, 64) # independent correlations, different sample sizes
```

pairs.panels

SPLOM, histograms and correlations for a data matrix

Description

Adapted from the help page for pairs, pairs.panels shows a scatter plot of matrices (SPLOM), with bivariate scatter plots below the diagonal, histograms on the diagonal, and the Pearson correlation above the diagonal. Useful for descriptive statistics of small data sets. If lm=TRUE, linear regression fits are shown for both y by x and x by y. Correlation ellipses are also shown.

Usage

```
pairs.panels(x, smooth = TRUE, scale = FALSE, density=TRUE,ellipses=TRUE,digits = 2, pch = 20,]
```

Arguments

x a data.frame or matrixsmoothTRUE draws loess smooths

scale TRUE scales the correlation font by the size of the absolute correlation.

density TRUE shows the density plots as well as histograms

ellipses TRUE draws correlation ellipses

lm Plot the linear fit rather than the LOESS smoothed fits.

digits the number of digits to show

pch The plot character (defaults to 20 which is a '.').
jiggle Should the points be jittered before plotting?

... other options for pairs

Details

Shamelessly adapted from the pairs help page. Uses panel.cor, panel.cor.scale, and panel.hist, all taken from the help pages for pairs. Also adapts the ellipse function from John Fox's car package.

pairs.panels is most useful when the number of variables to plot is less than about 6-8. It is particularly useful for an initial overview of the data.

Value

A scatter plot matrix (SPLOM) is drawn in the graphic window. The lower off diagonal draws scatter plots, the diagonal histograms, the upper off diagonal reports the Pearson correlation (with pairwise deletion).

If lm=TRUE, then the scatter plots are drawn above and below the diagonal, each with a linear regression fit. Useful to show the difference between regression lines.

See Also

pairs

```
pairs.panels(attitude) #see the graphics window
data(peas)
pairs.panels(peas,lm=TRUE,xlim=c(14,22),ylim=c(14,22))
```

partial.r Find the partial correlations for a set (x) of variables with set (y) removed.

Description

A straightforward application of matrix algebra to remove the effect of the variables in the y set from the x set. Input may be either a data matrix or a correlation matrix. Variables in x and y are specified by location.

Usage

```
partial.r(m, x, y, digits = 2)
```

Arguments

m	A data or correlation matrix
x	The variable numbers associated with the X set.
У	The variable numbers associated with the Y set
digits	Report correlations to digits of accuracy (default =2)

Details

It is sometimes convenient to partial the effect of a number of variables (e.g., sex, age, education) out of the correlations of another set of variables. This could be done laboriously by finding the residuals of various multiple correlations, and then correlating these residuals. The matrix algebra alternative is to do it directly.

Value

The matrix of partial correlations.

Author(s)

William Revelle

References

Revelle, W. (in prep) An introduction to psychometric theory with applications in R. To be published by Springer. (working draft available at http://personality-project.org/r/book/

See Also

mat.regress for a similar application for regression

Examples

```
jen <- make.hierarchical() #make up a correlation matrix
round(jen[1:5,1:5],2)
par.r <- partial.r(jen,c(1,3,5),c(2,4))
par.r</pre>
```

peas

Galton's Peas

Description

Francis Galton introduced the correlation coefficient with an analysis of the similarities of the parent and child generation of 700 sweet peas.

Usage

data(peas)

Format

A data frame with 700 observations on the following 2 variables.

parent The mean diameter of the mother pea for 700 peas child The mean diameter of the daughter pea for 700 sweet peas

Source

Stanton, Jeffrey M. (2001) Galton, Pearson, and the Peas: A brief history of linear regression for statistics intstructors, Journal of Statistics Education, 9. (retrieved from the web from http://www.amstat.org/publications/jse/v9n3/stanton.html) reproduces the table from Galton, 1894, Table 2.

The data were generated from this table.

References

Galton, Francis (1877) Typical laws of heredity. paper presented to the weekly evening meeting of the Royal Institution, London. Volume VIII (66) is the first reference to this data set. The data appear in

Galton, Francis (1894) Natural Inheritance (5th Edition), New York: MacMillan).

```
data(peas)
pairs.panels(peas,lm=TRUE,xlim=c(14,22),ylim=c(14,22))
describe(peas)
```

Create demo data for psychometrics

phi.demo

Description

A not very interesting demo of what happens if bivariate continuous data are dichotomized. Bascially a demo of r, phi, and polychor.

Usage

```
phi.demo(n=1000, r=.6, cuts=c(-2,-1,0,1,2))
```

Arguments

n number of cases to simulate

r correlation between latent and observed

cuts form dichotomized variables at the value of cuts

Details

A demonstration of the problem of different base rates on the phi correlation, and how these are partially solved by using the polychoric correlation. Not one of my more interesting demonstrations. See http://personality-project.org/r/simulating-personality.html and http://personality-project.org/r/r.datageneration.html for better demonstrations of data generation.

Value

a matrix of correlations and a graphic plot)

Author(s)

William Revelle

References

http://personality-project.org/r/simulating-personality.html and http://personality-project.org/r/r.datageneration.html for better demonstrations of data generation.

See Also

```
VSS.simulate, item.sim
```

Examples

round(phi.demo(),2) #compare the phi (lower off diagonal and polychoric correlations (upper off diagonal

Find the phi coefficient of correlation between two dichotomous variables

phi

Description

Given a 1×4 vector or a 2×2 matrix of frequencies, find the phi coefficient of correlation. Typical use is in the case of predicting a dichotomous criterion from a dichotomous predictor.

Usage

```
phi(t, digits = 2)
```

Arguments

t a 1 x 4 vector or a 2 x 2 matrix digits round the result to digits

Details

In many prediction situations, a dichotomous predictor (accept/reject) is validated against a dichotomous criterion (success/failure). Although a polychoric correlation estimates the underlying Pearson correlation as if the predictor and criteria were continuous and bivariate normal variables, the phi coefficient is the Pearson applied to a matrix of 0's and 1s.

Given a two x two table of counts

$$\begin{array}{cccc} a & b & a+b \\ c & d & c+d \\ a+c & b+d & a+b+c+d \end{array}$$

convert all counts to fractions of the total and then $\ Phi = a-(a+b)*(a+c)/sqrt((a+b)(c+d)(a+c)(b+d))$

Value

phi coefficient of correlation

Author(s)

William Revelle with modifications by Leo Gurtler

See Also

```
phi2poly ,Yule, Yule2phi
```

Examples

```
phi(c(30,20,20,30))
phi(c(40,10,10,40))
x <- matrix(c(40,5,20,20),ncol=2)
phi(x)</pre>
```

phi2poly

Convert a phi coefficient to a polychoric correlation

Description

Given a phi coefficient (a Pearson r calculated on two dichotomous variables), and the marginal frequencies (in percentages), what is the corresponding estimate of the polychoric correlation?

Given a two x two table of counts

 $\begin{array}{cc} a & b \\ c & d \end{array}$

The phi coefficient is (a - (a+b)*(a+c))/sqrt((a+b)(a+c)(b+d)(c+c)).

This function reproduces the cell entries for specified marginals and then calls John Fox's polychor function.

Usage

```
phi2poly(ph, cp, cc)
```

Arguments

ph phi

cp probability of the predictor – the so called selection ratio
 cc probability of the criterion – the so called success rate.

Details

Uses John Fox's polycor package, which in turn requires the mytnorm package

Value

a polychoric correlation

Author(s)

William Revelle

See Also

```
polychor.matrix, Yule2phi.matrix, phi2poly.matrix
```

Examples

```
#phi2poly(.3,.5,.5)
#phi2poly(.3,.3,.7)
```

plot.psych

Plotting functions for the psych package of class "psych"

Description

Combines several plotting functions into one for objects of class "psych". This can be used to plot the results of VSS, ICLUST, omega, factor.pa, or principal.

Usage

```
plot.psych(x,labels=NULL,...)
```

Arguments

x The object to plotlabels Variable labels... other calls to plot

Details

Passes the appropriate values to plot

Note

More precise plotting control is available in the separate plot functions.

Author(s)

William Revelle

See Also

```
VSS.plot and cluster.plot
```

```
test.data <- Harman74.cor$cov
f4 <- factor.pa(test.data,4)
plot(f4)</pre>
```

Description

Factor and cluster analysis output typically presents item by factor correlations (loadings). Tables of factor loadings are frequently sorted by the size of loadings. This style of presentation tends to make it difficult to notice the pattern of loadings on other, secondary, dimensions. By converting to polar coordinates, it is easier to see the pattern of the secondary loadings.

Usage

```
polar(f, sort = TRUE)
```

Arguments

f A matrix of loadings or the output from a factor or cluster analysis pro-

gram

sort = TRUE: sort items by the angle of the items on the first pair of

factors.

Details

Although many uses of factor analysis/cluster analysis assume a simple structure where items have one and only one large loading, some domains such as personality or affect items have a more complex structure and some items have high loadings on two factors. (These items are said to have complexity 2, see VSS). By expressing the factor loadings in polar coordinates, this structure is more readily perceived.

For each pair of factors, item loadings are converted to an angle with the first factor, and a vector length corresponding to the amount of variance in the item shared with the two factors

For a two dimensional structure, this will lead to a column of angles and a column of vector lengths. For n factors, this leads to n^* (n-1)/2 columns of angles and an equivalent number of vector lengths.

Value

polar

A data frame of polar coordinates

Author(s)

William Revelle

References

Rafaeli, E. & Revelle, W. (2006). A premature consensus: Are happiness and sadness truly opposite affects? Motivation and Emotion. \

Hofstee, W. K. B., de Raad, B., & Goldberg, L. R. (1992). Integration of the big five and circumplex approaches to trait structure. Journal of Personality and Social Psychology, 63, 146-163.

See Also

```
ICLUST, cluster.plot, circ.tests, factor.pa
```

Examples

```
circ.data <- circ.sim(24,500)
circ.fa <- factor.pa(circ.data,2)
circ.polar <- round(polar(circ.fa),2)
circ.polar
#compare to the graphic
cluster.plot(circ.fa)</pre>
```

poly.mat

Find polychoric correlations of item data

Description

Uses John Fox's hetcor function (from polychor package) to find a matrix of polychoric correlations for integer data. Essentially a wrapper for hetcor to convert integer item data into factor (categorical) data and then use hetcor. Just a useful shortcut for subsequent factor analysis.

Usage

```
poly.mat(x, short = TRUE, std.err = FALSE, ML = FALSE)
```

imum likelihood

Arguments

x	A matrix or data frame of integer data
short	short=TRUE, just show the correlations, short=FALSE give the full hetcor output
std.err	std.err=FALSE does not report the standard errors (faster)
ML	ML=FALSE do a quick two step procedure, ML=TRUE, do longer max-

Details

Typical personality and item data are integer values (0,1 for ability; 1,2, 3, 4 for attitude scales). The normal correlation procedures will find Pearson correlations (cor). The polycor and hetcor functions from John Fox's polychor package will find polychoric correlations for categorical data. This wrapper function converts integer data to categorical data and then calls hetcor.

Value

A matrix of polychoric correlations (if short=TRUE), otherwise a list of various estimates (see hetcor).

Note

requires polycor

Author(s)

William Revelle

Examples

round(phi.demo(),2) #compare the phi (lower off diagonal and polychoric correlations (upper off diagona

polychor.matrix

Phi or Yule coefficient matrix to polychoric coefficient matrix

Description

Given a matrix of phi or Yule correlation coefficients and a vector of marginals, use John Fox's polycor function to convert these to polychoric correlations.

Some older correlation matrices were reported as matrices of Phi or of Yule correlations. That is, correlations were found from the two by two table of counts:

a b c d

Yule Q is (ad - bc)/(ad+bc).

With marginal frequencies of a+b, c+d, a+c, b+d.

Given a square matrix of such correlations, and the proportions for each variable that are in the a + b cells, it is possible to reconvert each correlation into a two by two table and then estimate the corresponding polychoric correlation (using John Fox's polychor function.

Usage

```
Yule2poly.matrix(x, v)
phi2poly.matrix(x, v)
Yule2phi.matrix(x, v)
```

Arguments

x a matrix of phi or Yule coefficientsv A vector of marginal frequencies

Details

These functions call Yule2poly, Yule2phi or phi2poly for each cell of the matrix. See those functions for more details. See phi.demo for an example.

Value

A matrix of correlations

Author(s)

William Revelle

Examples

round(phi.demo() ,2) #compare the phi (lower off diagonal and polychoric correlations (upper off diagonal

principal

Principal components analysis

Description

Does an eigen value decomposition and returns eigen values, loadings, and degree of fit for a specified number of components. Basically it is just doing a principal components for n principal components. Can show the residual correlations as well. The quality of reduction in the squared correlations is reported by comparing residual correlations to original correlations. Unlike princomp, this returns a subset of just the best nfactors. The eigen vectors are rescaled by the sqrt of the eigen values to produce the component loadings more typical in factor analysis.

Usage

principal(r, nfactors = 1, residuals = FALSE,rotate="varimax",n.obs=NULL, scores=FALSE,missing=

Arguments

r a correlation matrix. If a raw data matrix is used, the correlations will

be found using pairwise deletions for missing values.

nfactors Number of components to extract

residuals FALSE, do not show residuals, TRUE, report residuals

rotate

n.obs Number of observations used to find the correlation matrix if using a

correlation matrix. Used for finding the goodness of fit statistics.

scores If TRUE, estimate component scores

missing if scores are TRUE, and missing=TRUE, then impute missing values using

either the median or the mean

impute "median" or "mean" values are used to replace missing values

digits =2. Accuracy of answers as well as display

Details

Useful for those cases where the correlation matrix is improper (perhaps because of SAPA techniques).

There are a number of data reduction techniques including principal components and factor analysis. Both PC and FA attempt to approximate a given correlation or covariance matrix of rank n with matrix of lower rank (p). ${}_{n}R_{n} \approx_{n} F_{kk}F'_{n} + U^{2}$ where k is much less than n. For principal components, the item uniqueness is assumed to be zero and all elements of the correlation matrix are fitted. That is, ${}_{n}R_{n} \approx_{n} F_{kk}F'_{n}$ The primary empirical difference between a components versus a factor model is the treatment of the variances for each item. Philosophically, components are weighted composites of observed variables while in the factor model, variables are weighted composites of the factors.

For a n x n correlation matrix, the n principal components completely reproduce the correlation matrix. However, if just the first k principal components are extracted, this is the best k dimensional approximation of the matrix.

Some of the statistics reported are more appropriate for maximum likelihood factor analysis rather than principal components analysis, and are reported to allow comparisons with these other models.

Although for items, it is typical to find component scores by scoring the salient items (using, e.g., score.items component scores can be estimated by regression. This is done just to be parallel with the principal axis factor analysis function factor.pa

Value

values Eigen Values of all components – useful for a scree plot

rotation which rotation was requested?

n.obs number of observations specified or found

communality Communality estimates for each item. These are merely the sum of

squared factor loadings for that item.

loadings A standard loading matrix of class "loadings"

fit Fit of the model to the correlation matrix

fit.off how well are the off diagonal elements reproduced?

residual Residual matrix – if requested

communality The history of the communality estimates. Probably only useful for teach-

ing what happens in the process of iterative fitting.

dof Degrees of Freedom for this model. This is the number of observed cor-

relations minus the number of independent parameters (number of items * number of factors - $nf^*(nf-1)/2$. That is, dof = niI * (ni-1)/2 - ni * nf

+ nf*(nf-1)/2.

objective value of the function that is minimized by maximum likelihood proce-

dures. This is reported for comparison purposes and as a way to estimate

chi square goodness of fit. The objective function is

 $f = log(trace((FF' + U2)^{-1}R) - log(|(FF' + U2)^{-1}R|) - n.items.$

STATISTIC If the number of observations is specified or found, this is a chi square

based upon the objective function, f. Using the formula from factanal:

 $\chi^2 = (n.obs - 1 - (2 * p + 5)/6 - (2 * factors)/3)) * f$

PVAL If n.obs > 0, then what is the probability of observing a chisquare this

large or larger?

phi If oblique rotations (using oblimin from the GPArotation package) are

requested, what is the interfactor correlation.

scores If scores=TRUE, then estimates of the factor scores are reported

Author(s)

William Revelle

See Also

VSS,factor2cluster,factor.pa, factor.congruence

```
#Four principal components of the Harmon 24 variable problem
#compare to a four factor principal axes solution using factor.congruence
pc <- principal(Harman74.cor$cov,4,rotate=TRUE)
pa <- factor.pa(Harman74.cor$cov,4,rotate=TRUE)
round(factor.congruence(pc,pa),2)</pre>
```

Print and summary functions for the psych class

print.psych

Description

Give limited output (print) or somewhat more detailed (summary) for the score.items, cluster.cor, cluster.loadings, or ICLUST functions

Usage

```
print.psych(x,digits=2,all=FALSE,cutoff=0,sort=FALSE,...)
summary.psych(object,digits=2,items=FALSE,...)
```

Arguments

X	Output from score.items
object	Output from score.items
items	items=TRUE (default) does not print the item whole correlations
digits	Number of digits to use in printing
all	if all=TRUE, then the object is declassed and all output from the function is printed $$
cutoff	Cluster loadings < cutoff will not be printed
sort	Cluster loadings are in sorted order
	More options to pass to summary and print

Details

Most of the psych functions produce too much output. print.psych and summary.psych use generic methods for printing just the highlights.

To get complete output, unclass(theobject) and then print it.

Value

Various psyc functions produce copious output. This is a way to summarize the most important parts of the output of the score.items, cluster.scores, and ICLUST functions. See those (score.items, cluster.cor, cluster.loadings, or ICLUST) for details on what is produced.

Note

See score.items, cluster.cor, cluster.loadings, or ICLUST for details on what is printed.

Author(s)

William Revelle

Examples

```
data(bfi)
keys.list <- list(agree=c(-1,2:5),conscientious=c(6:8,-9,-10),extraversion=c(-11,-12,13:15),neuroticism=
keys <- make.keys(25,keys.list,item.labels=colnames(bfi))
scores <- score.items(keys,bfi,short=TRUE)
scores
summary(scores)</pre>
```

r.test

Tests of significance for correlations

Description

Tests the significance of a single correlation, the difference between two independent correlations, the difference between two dependent correlations sharing one variable, or the difference between two dependent correlations with different variables.

Usage

```
r.test(n, r12, r34 = NULL, r23 = NULL, r13 = NULL, r14 = NULL, r24 = NULL, n2 = NULL, pooled=TRU
```

Arguments

n	Sample size of first group
r12	Correlation to be tested
r34	Test if this correlation is different from r12, if r23 is specified, but r13 is not, then r34 becomes r13
r23	if $ra=r(12)$ and $rb=r(13)$ then test for differences of dependent correlations given $r23$
r13	implies ra =r(12) and rb =r(34) test for difference of dependent correlations
r14	implies ra $=$ r (12) and rb $=$ r (34)
r24	ra = r(12) and rb = r(34)
n2	$\rm n2$ is specified in the case of two independent correlations. $\rm n2$ defaults to $\rm n$ if if not specified
pooled	use pooled estimates of correlations
twotailed	should a twotailed or one tailed test be used

Details

Depending upon the input, one of four different tests of correlations is done. \ 1) For a sample size n, find the t value for a single correlation. \ 2) For sample sizes of n and n2 (n2 = n if not specified) find the z of the difference between the z transformed correlations divided by the standard error of the difference of two z scores. \ 3) For sample size n, and correlations ra= r12, rb= r23 and r13 specified, test for the difference of two dependent correlations. \ 4) For

sample size n, test for the difference between two dependent correlations involving different variables.\ For clarity, correlations may be specified by value. If specified by location and if doing the test of dependent correlations, if three correlations are specified, they are assumed to be in the order r12, r13, r23.

Value

test	Label of test done
z	z value for tests 2 or 4 $$
t	t value for tests 1 and 3 $$
р	probability value of z or t

Note

Steiger specifically rejects using the Hotelling T test to test the difference between correlated correlations. These tests follow Steiger's advice.

Author(s)

William Revelle

References

Olkin, I. and Finn, J. D. (1995). Correlations redux. Psychological Bulletin, 118(1):155-164. \ Steiger, J.H. (1980), Tests for comparing elements of a correlation matrix, Psychological Bulletin, 87, 245-251.

See Also

This extends the tests in paired.r,r.con

```
n <- 30
r <- seq(0,.9,.1)
rc <- matrix(r.con(r,n),ncol=2)
test <- r.test(n,r)
r.rc <- data.frame(r=r,z=fisherz(r),lower=rc[,1],upper=rc[,2],t=test$t,p=test$p)
round(r.rc,2)
r.test(50,r)
r.test(30,.4,.6)  #test the difference between two independent correlations
r.test(103,.4,.5,.1)  #Steiger case A
r.test(103,.5,.6,.7,.5,.8)  #steiger Case B</pre>
```

shortcut for reading from the clipboard

read.clipboard

Description

input from the keyboard is easy but a bit obscure, particularly for Mac users. This is just an easier mnemonic to do so. Also will do some processing to read lower triangular matrices and fill them out to square matrices.

Usage

Arguments

header Does the first row have variable labels

diag for upper or lower triangular matrices, is the diagonal specified or not

names for read.clipboard.lower or upper, what colnames to use

... Other parameters to pass to read

Details

A typical session of R might involve data stored in text files, generated on line, etc. Although it is easy to just read from a file (particularly if using file.locate(), copying from the file to the clipboard and then reading from the clipboard is also very convenient (and somewhat more intuitive to the naive user. This is particularly convenient when copying from a text book or article and just moving a section of text into R.)

Based upon a suggestion by Ken Knoblauch to the R-help listserve.

read.clipboard.lower and read.clipboard.upper are adapted from John Fox's read.moments function in the sem package. They will read a lower (or upper) triangular matrix from the clipboard and return a full, symmetric matrix for use by factanal, factor.pa, ICLUST, etc. If the diagonal is false, it will be replaced by 1.0s. These two function were added to allow easy reading of examples from various texts and manuscripts with just triangular output.

Value

the contents of the clipboard.

Author(s)

William Revelle

Examples

```
#my.data <- read.clipboad()
#my.data <- read.clipboard.csv()
#my.data <- read.clipboad(header=FALSE)</pre>
```

rescale

Function to convert scores to "conventional" metrics

Description

Psychologists frequently report data in terms of transformed scales such as "IQ" (mean=100, sd=15, "SAT/GRE" (mean=500, sd=100), "ACT" (mean=18, sd=6), "T-scores" (mean=50, sd=10), or "Stanines" (mean=5, sd=2). The **rescale** function converts the data to standard scores and then rescales to the specified mean(s) and standard deviation(s).

Usage

```
rescale(x, mean = 100, sd = 15,df=TRUE)
```

Arguments

X	A matrix or data frame
mean	Desired mean of the rescaled scores- may be a vector
sd	Desired standard deviation of the rescaled scores
df	if TRUE, returns a data frame, otherwise a matrix

Value

A data frame (default) or matrix of rescaled scores.

Author(s)

William Revelle

See Also

See Also scale

```
T <- rescale(attitude,50,10) #all put on same scale
describe(T)
T1 <- rescale(attitude,seq(0,300,50),seq(10,70,10)) #different means and sigmas
describe(T1)</pre>
```

Description

Self reported scores on the SAT Verbal, SAT Quantitative and ACT were collected as part of the Synthetic Aperture Personality Assessment (SAPA) web based personality assessment project. Age, gender, and education are also reported. The data from 700 subjects are included here as a demonstration set for correlation and analysis.

Usage

```
data(sat.act)
```

Format

A data frame with 700 observations on the following 6 variables.

```
gender males = 1, females = 2
education self reported education 1 = high school ... 5 = graduate work
age age
ACT ACT composite scores may range from 1 - 36. National norms have a mean of 20.
SATV SAT Verbal scores may range from 200 - 800.
SATQ SAT Quantitative scores may range from 200 - 800
```

Details

hese items were collected as part of the SAPA project to develop online measures of ability. The score means are higher than national norms suggesting both self selection for people taking on line personality and ability tests and a self reporting bias in scores.

See also the ig. items data set.

Source

```
http://personality-project.org
```

```
data(sat.act)
describe(sat.act)
pairs.panels(sat.act)
```

 $\begin{tabular}{lll} {\it Scaling.fits} & {\it Test~the~adequacy~of~simple~choice,~logistic,~or~Thurstonian~scaling.} \\ & {\it ing.} \\ \hline \end{tabular}$

Description

Given a matrix of choices and a vector of scale values, how well do the scale values capture the choices? That is, what is size of the squared residuals given the model versus the size of the squared choice values?

Usage

```
scaling.fits(model, data, test = "logit", digits = 2, rowwise = TRUE)
```

Arguments

model A vector of scale values

data A matrix or dataframe of choice frequencies

test "choice", "logistic", "normal"

digits Precision of answer

rowwise Are the choices ordered by column over row (TRUE) or row over column

False)

Value

GF Goodness of fit of the model
original Sum of squares for original data

resid Sum of squares for residuals given the data and the model

residual Residual matrix

Note

Mainly for demonstration purposes for a course on psychometrics

Author(s)

William Revelle

References

Revelle, W. (in preparation) Introduction to psychometric theory with applications in R, Springer. http://personality-project.org/r/book

See Also

thurstone, vegetables

Description

One way to find omega is to do a factor analysis of the original data set, rotate the factors obliquely, do a Schmid Leiman transformation, and then find omega. Here is the code for Schmid Leiman. The S-L transform takes a factor or PC solution, transforms it to an oblique solution, factors the oblique solution to find a higher order (g) factor, and then residualizes g out of the the group factors.

Usage

```
schmid(model, nfactors = 3, pc = "pa",digits=2,rotate="oblimin",...)
```

Arguments

model	A correlation matrix
nfactors	Number of factors to extract
pc	pc="pa" for principal axes, pc="pc" for principal components, pc="mle" for maximum likelihood
digits	if digits not equal NULL, rounds to digits
rotate	The default, oblimin, produces somewhat more correlated factors than the alternative, simplimax. $$
	Allows additional parameters to be passed to the factoring routines

Details

Schmid Leiman orthogonalizations are typical in the ability domain, but are not seen as often in the non-cognitive personality domain. S-L is one way of finding the loadings of items on the general factor for estimating omega.

A typical example would be in the study of anxiety and depression. A general neuroticism factor (g) accounts for much of the variance, but smaller group factors of tense anxiety, panic disorder, depression, etc. also need to be considerd.

An alternative model is to consider hierarchical cluster analysis techniques such as ICLUST. Requires the GPArotation package.

Although 3 factors are the minimum number necessary to define the solution uniquely, it is occasionally useful to allow for a two factor solution. This is done here by setting the general factor loadings between the two lower order factors as the sqrt(oblique correlations between the factors). A warning message is issued.

Value

sl	loadings on	g + nfactors	group factors.	communalities,	uniqueness
~-	1000111160 011	8 1110000010	group ractors,	COLLING COLUMN	arrique recire

orthog original orthogonal factor loadings

oblique factor loadings

phi correlations among the transformed factors gload loadings of the lower order factors on g

. . .

Author(s)

William Revelle

References

http://personality-project.org/r/r.omega.html gives an example taken from Jensen and Weng, 1994 of a S-L transformation.

See Also

```
omega, omega.graph, fa.graph, ICLUST, VSS
```

Examples

```
s.jen <- schmid(make.hierarchical(),digits=2) #create a hierarchical demo
s.jen</pre>
```

score.alpha	$Score\ scales$	and find	$Cronbach \'s$	alpha	as	well	as	associated
	statistics							

Description

Given a matrix or data frame of k keys for m items (-1, 0, 1), and a matrix or data frame of items scores for m items and n people, find the sum scores or average scores for each person and each scale. In addition, report Cronbach's alpha, the average r, the scale intercorrelations, and the item by scale correlations. (Superseded by score.items).

Usage

```
score.alpha(keys, items, labels = NULL, totals=TRUE, digits = 2)
```

Arguments

keys	A matrix or dataframe of -1, 0, or 1 weights for each item on each scale
items	Data frame or matrix of raw item scores
labels	column names for the resulting scales
totals	Find sum scores (default) or average score
digits	Number of digits for answer (default $=2$)

Details

The process of finding sum or average scores for a set of scales given a larger set of items is a typical problem in psychometric research. Although the structure of scales can be determined from the item intercorrelations, to find scale means, variances, and do further analyses, it is typical to find the sum or the average scale score.

Various estimates of scale reliability include "Cronbach's alpha", and the average interitem correlation. For k = number of items in a scale, and av.r = average correlation between items in the scale, alpha = k * av.r/(1+(k-1)*av.r). Thus, alpha is an increasing function of test length as well as the test homeogeneity.

Alpha is a poor estimate of the general factor saturation of a test (see Zinbarg et al., 2005) for it can seriously overestimate the size of a general factor, and a better but not perfect estimate of total test reliability because it underestimates total reliability. None the less, it is a useful statistic to report.

Value

scores	Sum or average scores for each subject on the k scales
alpha	Cronbach's coefficient alpha. A simple (but non-optimal) measure of the internal consistency of a test. See also beta and omega.
av.r	The average correlation within a scale, also known as alpha 1 is a useful index of the internal consistency of a domain.
n.items	Number of items on each scale
cor	The intercorrelation of all the scales
item.cor	The correlation of each item with each scale. Because this is not corrected for item overlap, it will overestimate the amount that an item correlates with the other items in a scale.

Author(s)

William Revelle

References

```
An introduction to psychometric theory with applications in R (in preparation). http://personality-project.org/r/book
```

See Also

```
score.items, alpha.scale, correct.cor, alpha.scale, cluster.loadings, omega
```

```
y <- attitude  #from the datasets package
keys <- matrix(c(rep(1,7),rep(1,4),rep(0,7),rep(-1,3)),ncol=3)
labels <- c("first","second","third")
x <- score.alpha(keys,y,labels)</pre>
```

score.items	Score item composite scales and find Cronbach's alpha as well as
	$associated\ statistics$

Description

Given a matrix or data.frame of k keys for m items (-1, 0, 1), and a matrix or data.frame of items scores for m items and n people, find the sum scores or average scores for each person and each scale. In addition, report Cronbach's alpha, the average r, the scale intercorrelations, and the item by scale correlations. Replace missing values with the item median or mean if desired. Will adjust scores for reverse scored items. See make.keys for a convenient way to make the keys file.

Usage

```
score.items(keys, items, totals = FALSE, ilabels = NULL, missing = TRUE,impute="median", min
```

Arguments

keys	A matrix or dataframe of -1, 0, or 1 weights for each item on each scale. May be created by hand, or by using make.keys
items	Matrix or dataframe of raw item scores
totals	if TRUE find total scores, if FALSE (default), find average scores
ilabels	a vector of item labels.
missing	TRUE: Replace missing values with the corresponding item median or mean. FALSE: do not score that subject
impute	impute="median" replaces missing values with the item median, impute = "mean" replaces values with the mean response.
min	May be specified as minimum item score allowed, else will be calculated from data
max	May be specified as maximum item score allowed, else will be calculated from data
digits	Number of digits to report
short	if short is TRUE, then just give the item and scale statistics and do not report the scores

Details

The process of finding sum or average scores for a set of scales given a larger set of items is a typical problem in psychometric research. Although the structure of scales can be determined from the item intercorrelations, to find scale means, variances, and do further analyses, it is typical to find scores based upon the sum or the average item score. For some strange reason, personality scale scores are typically given as totals, but attitude scores as averages. The default for score.items is the average.

Various estimates of scale reliability include "Cronbach's alpha", and the average interitem correlation. For k = number of items in a scale, and av.r = average correlation between items in the scale, alpha = k * av.r/(1+(k-1)*av.r). Thus, alpha is an increasing function of test length as well as the test homeogeneity.

Alpha is a poor estimate of the general factor saturation of a test (see Zinbarg et al., 2005) for it can seriously overestimate the size of a general factor, and a better but not perfect estimate of total test reliability because it underestimates total reliability. None the less, it is a useful statistic to report. To estimate the omega cofficient, use the omega function.

Correlations between scales are attenuated by a lack of reliability. Correcting correlations for reliability (by dividing by the square roots of the reliabilities of each scale) sometimes help show structure.

By default, missing values are replaced with the corresponding median value for that item. Means can be used instead (impute="mean"), or subjects with missing data can just be dropped (missing = FALSE).

Value

scores	Sum or average scores for each subject on the k scales
alpha	Cronbach's coefficient alpha. A simple (but non-optimal) measure of the internal consistency of a test. See also beta and omega. Set to 1 for scales of length 1.
av.r	The average correlation within a scale, also known as alpha 1 is a useful index of the internal consistency of a domain. Set to 1 for scales with 1 item.
n.items	Number of items on each scale
item.cor	The correlation of each item with each scale. Because this is not corrected for item overlap, it will overestimate the amount that an item correlates with the other items in a scale.
cor	The intercorrelation of all the scales
corrected	The correlations of all scales (below the diagonal), alpha on the diagonal,

Author(s)

William Revelle

References

An introduction to psychometric theory with applications in R (in preparation). http://personality-project.org/r/book

See Also

make.keys for a convenient way to create the keys file, score.multiple.choice for multiple choice items,

alpha.scale, correct.cor, cluster.cor, cluster.loadings, omega for item/scale analysis

Examples

```
#see the example including the bfi data set
data(bfi)
keys.list <- list(agree=c(-1,2:5),conscientious=c(6:8,-9,-10),extraversion=c(-11,-12,13:15),neuroticism=
keys <- make.keys(25,keys.list,item.labels=colnames(bfi))
scores <- score.items(keys,bfi,short=TRUE)
scores</pre>
```

score.multiple.choice

Score multiple choice items and provide basic test statistics

Description

Ability tests are typically multiple choice with one right answer. score.multiple.choice takes a scoring key and a data matrix (or data.frame) and finds total or average number right for each participant. Basic test statistics (alpha, average r, item means, item-whole correlations) are also reported.

Usage

```
score.multiple.choice(key, data, score = TRUE, totals = FALSE, ilabels = NULL, missing = TRUE,
```

Arguments

key	A vector of the correct item alternatives
data	a matrix or data frame of items to be scored.
score	score=FALSE, just convert to right (1) or wrong (0). score=TRUE, find the totals or average scores and do item analysis
totals	total=FALSE: find the average number correct total=TRUE: find the total number correct
ilabels	item labels
missing	missing=TRUE: missing values are replaced with means or medians mising=FALSE missing values are not scored
impute	impute="median", replace missing items with the median score impute="mean": replace missing values with the item mean
digits	How many digits of output
short	short=TRUE, just report the item statistics, short=FALSE, report item statistics and subject scores as well

Details

Basically combines score.items with a conversion from multiple choice to right/wrong.

The item-whole correlation is inflated because of item overlap.

Value

scores Subject scores on one scale

missing Number of missing items for each subject

item.stats scoring key, response frequencies, item whole correlations, n subjects

scored, mean, sd, skew, kurtosis and se for each item

alpha Cronbach's coefficient alpha av.r Average interitem correlation

Author(s)

William Revelle

See Also

```
score.items, omega
```

Examples

```
data(iqitems)
iq.keys <- c(4,4,3,1,4,3,2,3,1,4,1,3,4,3)
score.multiple.choice(iq.keys,iqitems)
#just convert the items to true or false
iq.tf <- score.multiple.choice(iq.keys,iqitems,score=FALSE)
describe(iq.tf) #compare to previous results</pre>
```

SD

Find the Standard deviation for a vector, matrix, or data.frame - do not return error if there are no cases

Description

Find the standard deviation of a vector, matrix, or data.frame. In the latter two cases, return the sd of each column. Unlike the sd function, return NA if there are no observations rather than throw an error.

Usage

```
SD(x, na.rm = TRUE)
```

Arguments

x a vector, data.frame, or matrixna.rm is assumed to be TRUE

Details

Finds the standard deviation of a vector, matrix, or data.frame. Returns NA if no cases. Just an adaptation of the stats:sd function to return the functionality found in R < 2.7.0

Value

The standard deviation

Note

Until R 2.7.0, sd would return a NA rather than an error if no cases were observed. SD brings back that functionality. Although unusual, this condition will arise when analyzing data with high rates of missing values.

Author(s)

William Revelle

See Also

These functions use SD rather than sd: describe.by, skew, kurtosi

Examples

```
data(attitude)
sd(attitude) #all complete
attitude[,1] <- NA
SD(attitude) #missing a column
describe(attitude)</pre>
```

skew

Calculate skew for a vector, matrix, or data.frame

Description

Find the skew for each variable in a data.frame or matrix. Unlike skew in e1071, this calculates a different skew for each variable or column of a data.frame/matrix.

Usage

```
skew(x, na.rm = TRUE)
```

Arguments

x A data.frame or matrix
na.rm how to treat missing data

Details

given a matrix or data.frame x, find the skew for each column.

Value

if input is a matrix or data.frame, skew is a vector of skews

Note

The mean function supplies means for the columns of a data frame, but the overall mean for a matrix. Mean will throw a warning for non-numeric data, but colMeans stops with non-numeric data. Thus, the function uses either mean (for data frames) or colMeans (for matrices). This is true for skew and kurtosi as well.

Author(s)

William Revelle

See Also

```
describe, describe.by, kurtosi
```

Examples

round(skew(attitude),2)

smc

Find the Squared Multiple Correlation (SMC) of each variable with the remaining variables in a matrix

Description

The squared multiple correlation of a variable with the remaining variables in a matrix is sometimes used as initial estimates of the communality of a variable.

Usage

smc(R)

Arguments

R A correlation matrix or a dataframe. In the latter case, correlations are found.

Value

a vector of squared multiple correlations.

Author(s)

William Revelle

See Also

```
mat.regress, factor.pa
```

Examples

```
R <- make.hierarchical()
round(smc(R),2)</pre>
```

table2matrix

Convert a table with counts to a matrix or data frame representing those counts.

Description

Some historical sets are reported as summary tables of counts in a limited number of bins. Transforming these tables to data.frames representing the original values is useful for pedagogical purposes. (E.g., transforming the original Galton table of height x cubits in order to demonstrate regression.) The column and row names must be able to be converted to numeric values.

Usage

```
table2matrix(x, labs = NULL)
table2df(x, labs = NULL)
```

Arguments

x A two dimensional table of counts with row and column names that can

be converted to numeric values.

labs Labels for the rows and columns. These will be used for the names of the

two columns of the resulting matrix

Details

The original Galton (1888) of heights by cubits (arm length) is in tabular form. To show this as a correlation or as a scatter plot, it is useful to convert the table to a matrix or data frame of two columns.

Value

A matrix (or data.frame) of sum(x) rows and two columns.

Author(s)

William Revelle

See Also

cubits

Examples

```
data(cubits)
cubit <- table2matrix(cubits,labs=c("height","cubit"))
describe(cubit)
ellipses(cubit,n=1)</pre>
```

test.psych

Testing of functions in the psych package

Description

Test to make sure the psych functions run on basic test data sets

Usage

```
test.psych(first=1,last=5,short=TRUE)
```

Arguments

first first=1: start with dataset first
last last=5: test for datasets until last

short short=TRUE - don't return any analyses

Details

When modifying the psych package, it is useful to make sure that adding some code does not break something else. The test psych function tests the major functions on various standard data sets. It also shows off a number of the capabilities of the psych package.

Uses 5 standard data sets:

USArrests Violent Crime Rates by US State (4 variables)

attitude The Chatterjee-Price Attitude Data

Harman23.cor\$cov Harman Example 2.3 8 physical measurements

Harman74.cor\$cov Harman Example 7.4 24 mental measurements

ability.cov\$cov 8 Ability and Intelligence Tests

Value

out if short=FALSE, then list of the output from all functions tested

Warning

Warning messages will be thrown by fa.parallel and sometimes by factor.pa for random datasets.

Note

Although test.psych may be used as a quick demo of the various functions in the psych packge, in general, it is better to try the specific functions themselves. The main purpose of test.psych is to make sure functions throw error messages or correct for weird conditions.

The datasets tested are part of the standard R data sets and represent some of the basic problems encountered.

Author(s)

William Revelle

Examples

```
test <- test.psych()</pre>
```

thurstone

Thurstone Case V scaling

Description

Thurstone Case V scaling allows for a scaling of objects compared to other objects. As one of the cases considered by Thurstone, Case V makes the assumption of equal variances and uncorrelated distributions.

Usage

```
thurstone(x, ranks = FALSE, digits = 2)
```

Arguments

x A square matrix or data frame of preferences, or a rectangular data frame

or matrix rank order choices.

ranks TRUE if rank orders are presented digits number of digits in the goodness of fit

Details

Louis L. Thurstone was a pioneer in psychometric theory and measurement of attitudes, interests, and abilities. Among his many contributions was a systematic analysis of the process of comparative judgment (thurstone, 1927). He considered the case of asking subjects to successively compare pairs of objects. If the same subject does this repeatedly, or if subjects act as random replicates of each other, their judgments can be thought of as sampled from a normal distribution of underlying (latent) scale scores for each object, Thurstone proposed that the comparison between the value of two objects could be represented as representing the differences of the average value for each object compared to the standard deviation of the differences between objects. The basic model is that each item has a normal distribution of response strength and that choice represents the stronger of the two response strengths. A justification for the normality assumption is that each decision represents the sum of many independent inputs and thus, through the central limit theorem, is normally distributed.

Thurstone considered five different sets of assumptions about the equality and independence of the variances for each item (thurston, 1927). Torgerson expanded this analysis slightly by considering three classes of data collection (with individuals, between individuals and mixes of within and between) crossed with three sets of assumptions (equal covariance of decision process, equal correlations and small differences in variance, equal variances).

The data may be either a square matrix of dataframe of preferences (as proportions with the probability of the column variable being chosen over the row variable) or a matrix or dataframe of rank orders (1 being prefered to 2, etc.)

Value

GF Goodness of fit 1 = 1 - sum(squared residuals/squared original) for lower

off diagonal.

Goodness of fit 2 = 1 - sum(squared residuals/squared original) for full

matrix.

residual square matrix of residuals (of class dist)

data The original choice data

...

Author(s)

William Revelle

References

Thurstone, L. L. (1927) A law of comparative judgments. Psychological Review, 34, 273-286.

Revelle, W. An introduction to psychometric theory with applications in R. (in preparation), Springer. http://personality-project.org/r/book

Examples

```
data(vegetables)
thurstone(veg)
```

tr

Find the trace of a square matrix

Description

Hardly worth coding, if it didn't appear in so many formulae in psychometrics, the trace of a (square) matrix is just the sum of the diagonal elements.

Usage

tr(m)

Arguments

m

A square matrix

Details

The tr function is used in various matrix operations and is the sum of the diagonal elements of a matrix.

Value

```
The sum of the diagonal elements of a square matrix. i.e. \operatorname{tr}(m) < -\operatorname{sum}(\operatorname{diag}(m)).
```

Examples

```
m <- matrix(1:16,ncol=4)
m
tr(m)</pre>
```

Description

A classic data set for demonstrating Thurstonian scaling is the preference matrix of 9 vegetables from Guilford (1954). Used by Guiford, Nunnally, and Nunally and Bernstein, this data set allows for examples of basic scaling techniques.

Usage

data(vegetables)

Format

A data frame with 9 choices on the following 9 vegetables. The values reflect the perecentage of times where the column entry was preferred over the row entry.

Turn Turnips

Cab Cabbage

Beet Beets

Asp Asparagus

Car Carrots

Spin Spinach

S.Beans String Beans

Peas Peas

Corn Corn

Details

Louis L. Thurstone was a pioneer in psychometric theory and measurement of attitudes, interests, and abilities. Among his many contributions was a systematic analysis of the process of comparative judgment (thurstone, 1927). He considered the case of asking subjects to successively compare pairs of objects. If the same subject does this repeatedly, or if subjects act as random replicates of each other, their judgments can be thought of as sampled from a normal distribution of underlying (latent) scale scores for each object, Thurstone proposed that the comparison between the value of two objects could be represented as representing the differences of the average value for each object compared to the standard deviation of the differences between objects. The basic model is that each item has a normal distribution of response strength and that choice represents the stronger of the two response strengths. A justification for the normality assumption is that each decision represents the sum of many independent inputs and thus, through the central limit theorem, is normally distributed.

Thurstone considered five different sets of assumptions about the equality and independence of the variances for each item (thurston, 1927). Torgerson expanded this analysis slightly

by considering three classes of data collection (with individuals, between individuals and mixes of within and between) crossed with three sets of assumptions (equal covariance of decision process, equal correlations and small differences in variance, equal variances).

This vegetable data set is used by Guilford and by Nunnally to demonstrate Thurstonian scaling.

Source

Guilford, J.P. (1954) Psychometric Methods. McGraw-Hill, New York.

References

```
Nunnally, J. C. (1967). Psychometric theory., McGraw-Hill, New York.
```

Revelle, W. An introduction to psychometric theory with applications in R. (in preparation), Springer. http://personality-project.org/r/book

See Also

thurstone

Examples

data(vegetables)
thurstone(veg)

VSS.parallel

Compare real and random VSS solutions

Description

Another useful test for the number of factors is when the eigen values of a random matrix are greater than the eigen values of a real matrix. Here we show VSS solutions to random data.

Usage

```
VSS.parallel(ncases, nvariables,scree=FALSE,rotate="none")
```

Arguments

ncases Number of simulated cases nvariables number of simulated variables

scree Show a scree plot for random data – see omega

rotate "none" or rotate="varimax"

Value

VSS like output to be plotted by VSS.plot

Author(s)

William Revelle

References

Very Simple Structure (VSS)

See Also

```
fa.parallel, VSS.plot, ICLUST, omega
```

Examples

```
#VSS.plot(VSS.parallel(200,24))
```

VSS.plot

Plot VSS fits

Description

The Very Simple Structure criterion (VSS) for estimating the optimal number of factors is plotted as a function of the increasing complexity and increasing number of factors.

Usage

```
VSS.plot(x, title = "Very Simple Structure", line = FALSE)
```

Arguments

x output from VSS

title any title

line connect different complexities

Details

Item-factor models differ in their "complexity". Complexity 1 means that all except the greatest (absolute) loading for an item are ignored. Basically a cluster model (e.g., ICLUST). Complexity 2 implies all except the greatest two, etc.

Different complexities can suggest different number of optimal number of factors to extract. For personality items, complexity 1 and 2 are probably the most meaningful.

The Very Simple Structure criterion will tend to peak at the number of factors that are most interpretable for a given level of complexity. Note that some problems, the most interpretable number of factors will differ as a function of complexity. For instance, when

doing the Harman 24 psychological variable problems, an unrotated solution of complexity one suggests one factor (g), while a complexity two solution suggests that a four factor solution is most appropriate. This latter probably reflects a bi-factor structure.

For examples of VSS.plot output, see http://personality-project.org/r/r.vss.html

Value

A plot window showing the VSS criterion varying as the number of factors and the complexity of the items.

Author(s)

Maintainer: William Revelle (revelle@northwestern.edu)

References

```
http://personality-project.org/r/r.vss.html
```

See Also

```
VSS, ICLUST, omega
```

Examples

VSS.scree

Plot a scree test

Description

Cattell's scree test is one of most simple ways of testing the number of components in a correlation matrix. Here we plot the eigen values of a correlation matrix.

Usage

```
VSS.scree(rx, main = "scree plot")
```

Arguments

rx a correlation matrix or a data matrix. If data, then correlations are found

using pairwise deletions.

main Title

Author(s)

William Revelle Department of Psychology Northwestern University Evanston, Illiniois

Maintainer: William Revelle <revelle@northwestern.edu>

References

```
http://personality-project.org/r/vss.html
```

See Also

```
VSS.plot, ICLUST, omega
```

Examples

```
#VSS.scree(attitude)
#VSS.scree(cor(attitude)
```

VSS.sim

create VSS like data

Description

Simulation is one of most useful techniques in statistics and psychometrics. Here we simulate a correlation matrix with a simple structure composed of a specified number of factors. Each item is assumed to have complexity one. See circ.sim and item.sim for alternative simulations.

Usage

```
VSS.sim(ncases=1000, nvariables=16, nfactors=4, meanloading=.5,dichot=FALSE,cut=0)
```

Arguments

ncases number of simulated subjects

nvariables Number of variables

nfactors Number of factors to generate

meanloading with a mean loading

dichot=FALSE give continuous variables, dichot=TRUE gives dichoto-

mous variables

cut if dichotomous = TRUE, then items with values > cut are assigned 1,

otherwise 0.

Value

a neases x nvariables matrix

Author(s)

William Revelle

See Also

```
VSS, ICLUST
```

Examples

```
## Not run:
simulated <- VSS.simulate(1000,20,4,.6)
vss <- VSS(simulated,rotate="varimax")
VSS.plot(vss)
## End(Not run)</pre>
```

VSS

Apply the Very Simple Structure and MAP criteria to determine the appropriate number of factors.

Description

There are multiple ways to determine the appropriate number of factors in exploratory factor analysis. Routines for the Very Simple Structure (VSS) criterion allow one to compare solutions of varying complexity and for different number of factors. Graphic output indicates the "optimal" number of factors for different levels of complexity. The Velicer MAP criterion is another good choice.

Usage

```
VSS(x, n = 8, rotate = "varimax", diagonal = FALSE, pc = "pa", n.obs=NULL,plot=TRUE,title="Very
```

Arguments

x a correlation matrix or a data matrix

n Number of factors to extract – should be more than hypothesized!

rotate what rotation to use c("none", "varimax", "promax")

diagonal Should we fit the diagonal as well

pc pc="pa" Principal Axis Factor Analysis, pc="mle" Maximum Likelihood

FA, pc="pc" Principal Components"

n.obs Number of observations if doing a factor analysis of correlation matrix.

This value is ignored by VSS but is necessary for the ML factor analysis

package.

Figure 9: The Very Simple Structure criterion plots goodness of fit as a function of the number of factors extracted and the factorial complexity for each item/test. Note how a complexity one solution best fits the data if only one factor is extracted, but a complexity three solution is optimized at 4 factors.

Number of Factors

plot=TRUE Automatically call VSS.plot with the VSS output, otherwise plot

don't plot

a title to be passed on to VSS.plot title

parameters to pass to the factor analysis program The most important of

these is if using a correlation matrix is covmat= xx

Details

Determining the most interpretable number of factors from a factor analysis is perhaps one of the greatest challenges in factor analysis. There are many solutions to this problem, none of which is uniformly the best. "Solving the number of factors problem is easy, I do it everyday before breakfast. But knowing the right solution is harder" (Kaiser, 195x).

Techniques most commonly used include

- 1) Extracting factors until the chi square of the residual matrix is not significant.
- 2) Extracting factors until the change in chi square from factor n to factor n+1 is not significant.
- 3) Extracting factors until the eigen values of the real data are less than the corresponding eigen values of a random data set of the same size (parallel analysis) fa.parallel.
- 4) Plotting the magnitude of the successive eigen values and applying the scree test (a sudden drop in eigen values analogous to the change in slope seen when scrambling up the talus slope of a mountain and approaching the rock face.
- 5) Extracting principal components until the eigen value <1.
- 6) Extracting factors as long as they are interpetable.
- 7) Using the Very Structure Criterion (VSS).
- 8) Using Wayne Velicer's Minimum Average Partial (MAP) criterion.

Each of the procedures has its advantages and disadvantages. Using either the chi square test or the change in square test is, of course, sensitive to the number of subjects and leads to the nonsensical condition that if one wants to find many factors, one simlpy runs more subjects. Parallel analysis is partially sensitive to sample size in that for large samples the eigen values of random factors will be very small. The scree test is quite appealling but can lead to differences of interpretation as to when the scree "breaks". The eigen value of 1 rule, although the default for many programs, seems to be a rough way of dividing the number of variables by 3. Extracting interpretable factors means that the number of factors reflects the investigators creativity more than the data. VSS, while very simple to understand, will not work very well if the data are very factorially complex. (Simulations suggests it will work fine if the complexities of some of the items are no more than 2).

Most users of factor analysis tend to interpret factor output by focusing their attention on the largest loadings for every variable and ignoring the smaller ones. Very Simple Structure operationalizes this tendency by comparing the original correlation matrix to that reproduced by a simplified version (S) of the original factor matrix (F). R = SS' +U2. S is composed of just the c greatest (in absolute value) loadings for each variable. C (or complexity) is a parameter of the model and may vary from 1 to the number of factors.

The VSS criterion compares the fit of the simplified model to the original correlations: VSS = 1 -sumsquares(r^*)/sumsquares(r) where R^* is the residual matrix $R^* = R$ - SS' and r^* and r are the elements of R* and R respectively.

VSS for a given complexity will tend to peak at the optimal (most interpretable) number of factors (Revelle and Rocklin, 1979).

Although originally written in Fortran for main frame computers, VSS has been adapted to micro computers (e.g., Macintosh OS 6-9) using Pascal. We now release R code for calculating VSS.

Note that if using a correlation matrix (e.g., my.matrix) and doing a factor analysis, the parameters n.obs should be specified for the factor analysis: e.g., the call is VSS(my.matrix,n.obs=500). Otherwise it defaults to 1000.

Wayne Velicer's MAP criterion has been added as an additional test for the optimal number of components to extract. Note that VSS and MAP will not always agree as to the optimal number.

Value

```
A data.frame with entries: map: Velicer's MAP values (lower values are better) dof: degrees of freedom (if using FA) chisq: chi square (from the factor analysis output (if using FA) prob: probability of residual matrix > 0 (if using FA) sqresid: squared residual correlations fit: factor fit of the complete model cfit.1: VSS fit of complexity 1 cfit.2: VSS fit of complexity 2 ... cfit.8: VSS fit of complexity 8 cresidiual.1: sum squared residual correlations for complexity 1 ...: sum squared residual correlations for complexity 2 ...
```

Author(s)

William Revelle Department of Psychology Northwestern University Evanston, Illiniois

Maintainer: William Revelle < revelle@northwestern.edu>

References

http://personality-project.org/r/vss.html, Revelle, W. An introduction to psychometric theory with applications in R (in prep) Springer. Draft chapters available at http://personality-project.org/r/book/

see also Revelle, W. and Rocklin, T. 1979, Very Simple Structure: an Alternative Procedure for Estimating the Optimal Number of Interpretable Factors, Multivariate Behavioral Research, 14, 403-414. http://personality-project.org/revelle/publications/vss.pdf

Velicer, W. (1976) Determining the number of components from the matrix of partial correlations. Psychometrika, 41, 321-327.

See Also

```
VSS.plot, ICLUST, omega, fa.parallel
```

Examples

```
test.data <- Harman74.cor$cov
my.vss <- VSS(test.data,title="VSS of 24 mental tests")
#print(my.vss[,1:12],digits =2)
#VSS.plot(my.vss, title="VSS of 24 mental tests")
#now, some simulated data with two factors
VSS(circ.sim(nvar=24),pc="mle" ,title="VSS of 24 circumplex variables")
VSS(item.sim(nvar=24),pc="mle" ,title="VSS of 24 simple structure variables")</pre>
```

winsor

Find the Winsorized scores or means for a vector, matrix, or data.frame

Description

Among the robust estimates of central tendency are trimmed means and Winsorized means. This function finds the Winsorized mean. The top and bottom trim values are given values of the trimmed and 1- trimmed quantiles. Then means are found.

Usage

```
winsor(x, trim = 0.2, na.rm = TRUE)
winsor.means(x, trim = 0.2, na.rm = TRUE)
```

Arguments

x A data vector, matrix or data frame

trim Percentage of data to move from the top and bottom of the distributions

na.rm Missing data are removed

Details

Among the many robust estimates of central tendency, some recommend the Winsorized mean. Rather than just dropping the top and bottom trim percent, these extreme values are replaced with values at the trim and 1- trim quantiles.

Value

A scalar or vector of winsorized scores or winsorized means (depending upon the call).

Author(s)

William Revelle with modifications suggested by Joe Paxton

References

Wilcox, Rand R. (2005) Introduction to robust estimation and hypothesis testing. Elsevier/Academic Press. Amsterdam; Boston.

See Also

```
interp.median
```

Examples

```
data(sat.act)
winsor.means(sat.act)
x <- matrix(1:100,ncol=5)
winsor(x)
winsor.means(x)
y <- 1:11
winsor(y,trim=.5)</pre>
```

Yule

From a two by two table, find the Yule coefficients of association, convert to phi, or polychoric, recreate table the table to create the Yule coefficient.

Description

One of the many measures of association is the Yule coefficient. Given a two $\mathbf x$ two table of counts

a b c d

Yule Q is (ad - bc)/(ad+bc).

Conceptually, this is the number of pairs in agreement (ad) - the number in disagreement (bc) over the total number of paired observations.

ad/bc is the odds ratio and Q = (OR-1)/(OR+1)

Yule's coefficient of colligation is Y = (sqrt(OR) - 1)/(sqrt(OR) + 1) Yule.inv finds the cell entries for a particular Q and the marginals $(a+b,c+d,a+c,\ b+d)$. This is useful for converting old tables of correlations into more conventional **phi** or polychoric correlations. Yule2phi and Yule2poly convert the Yule Q with set marginals to the correponding phi or polychoric correlation.

Usage

```
Yule(x,Y=FALSE)
Yule.inv(Q,m)
Yule2phi(Q,m)
```

Yule2poly(Q,m)

Arguments

x	A vector of four elements or a two by two matrix
Y	Y=TRUE return Yule's Y coefficient of colligation
Q	The Yule coefficient
m	A two x two matrix of marginals or a four element vector of marginals

Details

Yule developed two measures of association for two by two tables. Both are functions of the odds ratio

Value

Q The Yule Q coefficient
R A two by two matrix of counts

Note

Currently done by using the optimize function, but presumably could be redone by solving a quadratic equation.

Author(s)

William Revelle

References

Yule, G. Uday (1912) On the methods of measuring association between two attributes. Journal of the Royal Statistical Society, LXXV, 579-652

See Also

```
See Also as phi, Yule2poly.matrix, Yule2phi.matrix
```

Examples

```
Nach <- matrix(c(40,10,20,50),ncol=2,byrow=TRUE)
Yule(Nach)
Yule.inv(.81818,c(50,70,60,60))
Yule2phi(.81818,c(50,70,60,60))
if(require(polycor)) Yule2poly(.81818,c(50,70,60,60))
phi(Nach) #much less</pre>
```