Package ‘compositions’

March 4, 2020

Version 1.40-4
Date 2020-03-24
Title Compositional Data Analysis
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Maintainer K. Gerald van den Boogaart <support@boogaart.de>
Depends R (>= 2.2.0), tensorA, robustbase, bayesm
Suggests rgl,combinat, energy
Description Provides functions for the consistent analysis of compositional
data (e.g. portions of substances) and positive numbers (e.g. concentrations)
in the way proposed by J. Aitchison and V. Pawlowsky-Glahn.
License GPL (>= 2)
URL http://www.stat.boogaart.de/compositions
NeedsCompilation yes
Repository CRAN
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R topics documented:

compositions-package .............................................. 5
Aar ................................................................. 12
acom .............................................................. 13
acomparith ....................................................... 15
acompmargin ..................................................... 17
acompscalarproduct .............................................. 18
Activity10 .......................................................... 19
Activity31 .......................................................... 20
alr ................................................................. 22
AnimalVegetation .................................................. 24
aplus ............................................................... 24
aplusarithm ....................................................... 26
<table>
<thead>
<tr>
<th>R topics documented:</th>
</tr>
</thead>
<tbody>
<tr>
<td>apt</td>
</tr>
<tr>
<td>ArcticLake</td>
</tr>
<tr>
<td>arrows3D</td>
</tr>
<tr>
<td>as.data.frame</td>
</tr>
<tr>
<td>axis3D</td>
</tr>
<tr>
<td>balance</td>
</tr>
<tr>
<td>barplot.acomp</td>
</tr>
<tr>
<td>Bayesite</td>
</tr>
<tr>
<td>binary</td>
</tr>
<tr>
<td>biplot3D</td>
</tr>
<tr>
<td>Blood23</td>
</tr>
<tr>
<td>Boxite</td>
</tr>
<tr>
<td>boxplot</td>
</tr>
<tr>
<td>ccomp</td>
</tr>
<tr>
<td>ccompgof</td>
</tr>
<tr>
<td>cdt</td>
</tr>
<tr>
<td>ClamEast</td>
</tr>
<tr>
<td>ClamWest</td>
</tr>
<tr>
<td>clo</td>
</tr>
<tr>
<td>clr</td>
</tr>
<tr>
<td>clr2ilr</td>
</tr>
<tr>
<td>ClusterFinder1</td>
</tr>
<tr>
<td>CoDaDendrogram</td>
</tr>
<tr>
<td>coloredBiplot</td>
</tr>
<tr>
<td>colorsForOutliers</td>
</tr>
<tr>
<td>CompLinModCoReg</td>
</tr>
<tr>
<td>compOKriging</td>
</tr>
<tr>
<td>ConfRadius</td>
</tr>
<tr>
<td>cor.acomp</td>
</tr>
<tr>
<td>Coxite</td>
</tr>
<tr>
<td>cpt</td>
</tr>
<tr>
<td>DiagnosticProb</td>
</tr>
<tr>
<td>dist</td>
</tr>
<tr>
<td>ellipses</td>
</tr>
<tr>
<td>endmemberCoordinates</td>
</tr>
<tr>
<td>Firework</td>
</tr>
<tr>
<td>fitdirichlet</td>
</tr>
<tr>
<td>fitSameMeanDifferentVarianceModel</td>
</tr>
<tr>
<td>gausstest</td>
</tr>
<tr>
<td>geometricmean</td>
</tr>
<tr>
<td>getdetectionlimit</td>
</tr>
<tr>
<td>Glacial</td>
</tr>
<tr>
<td>gof</td>
</tr>
<tr>
<td>groupparts</td>
</tr>
<tr>
<td>Hongite</td>
</tr>
<tr>
<td>HotellingsTsq</td>
</tr>
<tr>
<td>HouseholdExp</td>
</tr>
<tr>
<td>Hydrochem</td>
</tr>
</tbody>
</table>
R topics documented:

idt .............................................................. 94
iit ............................................................... 96
ilr ............................................................... 97
ilrBase .......................................................... 99
ilt ............................................................... 100
ipt ............................................................... 101
is.acomp ........................................................ 103
IsMahalanobisOutlier ......................................... 104
isoPortionLines ................................................ 105
jura ............................................................... 107
kingTetrahedron ............................................... 108
Kongite .......................................................... 110
lines .............................................................. 111
logratioVariogram ............................................ 112
lrvgram .......................................................... 114
MahalanobisDist ............................................... 115
matmult .......................................................... 117
mean.acomp ..................................................... 118
meanrow ........................................................ 119
Metabolites ..................................................... 120
missing.compositions ........................................ 121
missingProjector ............................................... 124
missingsummary ............................................... 126
mix.Read ........................................................ 127
mvar .............................................................. 129
names ............................................................ 131
norm .............................................................. 132
normalize ....................................................... 134
NormalTests ..................................................... 135
oneOrDataset ................................................... 136
outlierclassifier ............................................... 137
outlierplot ....................................................... 139
outliersInCompositions ....................................... 143
pairwiseplot .................................................... 146
parametricMat .................................................. 148
perturbe ......................................................... 149
plot.acomp ...................................................... 151
plot.aplus ....................................................... 154
plot3D ............................................................ 156
plot3Dacomp ................................................... 157
plot3Daplus ..................................................... 159
plot3Drmult .................................................... 160
plot3Drplus ...................................................... 161
plot.logratioVariogram ....................................... 163
plotmissingsummary .......................................... 164
PogoJump ....................................................... 165
powerofpsdmatrix ............................................. 166
princomp.acomp ............................................... 167
R topics documented:

princomp.aplus ................................................. 170
princomp.rcomp ................................................. 173
princomp.rmult ................................................. 175
princomp.rplus ................................................. 177
print.acomp .................................................. 179
pwlrPlot ...................................................... 181
rcomp .......................................................... 183
ratioLoadings .................................................. 184
rAitchison ...................................................... 185
replot .......................................................... 186
rlnorm .......................................................... 187
rMahalanobis ................................................... 188
rmult ........................................................... 189
rmultarithm ..................................................... 190
rmultmatmult ................................................... 191
rnorm ........................................................... 192
robustnessInCompositions ................................. 193
rplus ............................................................ 194
rplusarithm ..................................................... 195
rpois ............................................................ 196
runif ............................................................ 197
ternaryAxis ...................................................... 198
totals ............................................................ 199
transformations from 'mixtures' to 'compositions' classes ......................... 200

Sediments ....................................................... 201
SerumProtein ..................................................... 202
SimulatedAmounts ............................................. 203
Skull ............................................................. 204
SkyeAFM ........................................................ 205
split ............................................................. 206
summary.acomp ............................................... 207
summary.aplus ............................................... 208
summary.rcomp ............................................... 209
sumprojector ................................................... 210
Supervisor ...................................................... 211
ternaryAxis ..................................................... 212
totals ............................................................ 213
transformations from 'mixtures' to 'compositions' classes ......................... 214

Read standard data files ....................................... 215
repplot .......................................................... 216
scale ........................................................... 217
segments ......................................................... 218
Transformations from 'mixtures' to 'compositions' classes ......................... 219
Segments ......................................................... 220

Scalar .......................................................... 221
ShiftOperators .................................................. 222
summary.rcomp ............................................... 223
simulatemissings .............................................. 224

Sediments ....................................................... 225

ternaryAxis ..................................................... 226
totals ............................................................ 227
transformations from 'mixtures' to 'compositions' classes ......................... 228

Read standard data files ....................................... 229
repplot .......................................................... 230
scale ........................................................... 231
segments ......................................................... 232
Transformations from 'mixtures' to 'compositions' classes ......................... 233
SimulatedAmounts ............................................. 234
Skull ............................................................. 235
SkyeAFM ........................................................ 236
split ............................................................. 237
summary.acomp ............................................... 238
summary.aplus ............................................... 239
summary.rcomp ............................................... 240
sumprojector ................................................... 241
Supervisor ...................................................... 242
ternaryAxis ..................................................... 243
totals ............................................................ 244
transformations from 'mixtures' to 'compositions' classes ......................... 245
compositions-package

Description

"compositions" is a package for the analysis of compositional and multivariate positive data (generally called "amounts"), based on several alternative approaches.

Details

The DESCRIPTION file:

Package: compositions
Version: 1.40-4
Date: 2020-03-24
Title: Compositional Data Analysis
Author: K. Gerald van den Boogaart <boogaart@hzdr.de>, Raimon Tolosana-Delgado, Matevz Bren
Maintainer: K. Gerald van den Boogaart <support@boogaart.de>
Depends: R (>= 2.2.0), tensorA, robustbase, bayesm
Suggests: rgl,combinat, energy
Description: Provides functions for the consistent analysis of compositional data (e.g. portions of substances) and positive numbers (e.g. concentrations), based on several alternative approaches.
License: GPL (>= 2)
URL: http://www.stat.boogaart.de/compositions

Index of help topics:

structure
Aar  Composition of glacial sediments from the Aar massif (Switzerland)
acommpmargin  Marginal compositions in Aitchison Compositions
<table>
<thead>
<tr>
<th>Activity10</th>
<th>Activity patterns of a statistician for 20 days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Activity31</td>
<td>Activity patterns of a statistician for 20 days</td>
</tr>
<tr>
<td>alr</td>
<td>Additive log ratio transform</td>
</tr>
<tr>
<td>AnimalVegetation</td>
<td>Animal and vegetation measurement</td>
</tr>
<tr>
<td>+.aplus</td>
<td>Vectorial arithmetic for data sets with aplus class</td>
</tr>
<tr>
<td>aplus</td>
<td>Amounts analysed in log-scale</td>
</tr>
<tr>
<td>apt</td>
<td>Additive planar transform</td>
</tr>
<tr>
<td>ArcticLake</td>
<td>Arctic lake sediment samples of different water depth</td>
</tr>
<tr>
<td>arrows3D</td>
<td>Arrows in 3D, based on package rgl</td>
</tr>
<tr>
<td>as.data.frame.acomp</td>
<td>Convert &quot;compositions&quot; classes to data frames</td>
</tr>
<tr>
<td>axis3D</td>
<td>Drawing a 3D coordinate system to a plot, based on package rgl</td>
</tr>
<tr>
<td>balance</td>
<td>Compute balances for a compositional dataset.</td>
</tr>
<tr>
<td>barplot.acomp</td>
<td>Bar charts of amounts</td>
</tr>
<tr>
<td>Bayesite</td>
<td>Permeabilities of bayesite</td>
</tr>
<tr>
<td>binary</td>
<td>Treating binary and g-adic numbers</td>
</tr>
<tr>
<td>biplot3D</td>
<td>Three-dimensional biplots, based on package rgl</td>
</tr>
<tr>
<td>Blood23</td>
<td>Blood samples</td>
</tr>
<tr>
<td>Boxite</td>
<td>Compositions and depth of 25 specimens of boxite</td>
</tr>
<tr>
<td>boxplot.acomp</td>
<td>Displaying compositions and amounts with box-plots</td>
</tr>
<tr>
<td>cdt</td>
<td>Centered default transform</td>
</tr>
<tr>
<td>ClamEast</td>
<td>Color-size compositions of 20 clam colonies from East Bay</td>
</tr>
<tr>
<td>ClamWest</td>
<td>Color-size compositions of 20 clam colonies from West Bay</td>
</tr>
<tr>
<td>clo</td>
<td>Closure of a composition</td>
</tr>
<tr>
<td>clr</td>
<td>Centered log ratio transform</td>
</tr>
<tr>
<td>clr2ilr</td>
<td>Convert between clr and ilr, and between cpt and ipt.</td>
</tr>
<tr>
<td>ClusterFinder1</td>
<td>Heuristics to find subpopulations of outliers</td>
</tr>
<tr>
<td>CoDaDendrogram</td>
<td>Dendrogram representation of acomp or rcomp objects</td>
</tr>
<tr>
<td>coloredBiplot</td>
<td>A biplot providing somewhat easier access to details of the plot.</td>
</tr>
<tr>
<td>colorsForOutliers1</td>
<td>Create a color/char palette or for groups of outliers</td>
</tr>
<tr>
<td>CompLinModCoReg</td>
<td>Compositional Linear Model of Coregionalisation</td>
</tr>
<tr>
<td>compOKriging</td>
<td>Compositional Ordinary Kriging</td>
</tr>
<tr>
<td>compositions-package</td>
<td>library(compositions)</td>
</tr>
<tr>
<td>ConfRadius</td>
<td>Helper to compute confidence ellipsoids</td>
</tr>
<tr>
<td>cor.acomp</td>
<td>Correlations of amounts and compositions</td>
</tr>
<tr>
<td>Coxite</td>
<td>Compositions, depths and porosities of 25 specimens of coxite</td>
</tr>
<tr>
<td>cpt</td>
<td>Centered planar transform</td>
</tr>
</tbody>
</table>
DiagnosticProb  Diagnostic probabilities
dist            Distances in various approaches
ellipses        Draw ellipses
dendmemberCoordinates Recast amounts as mixtures of end-members
Firework        Firework mixtures
gemetricmean    The geometric mean
getDetectionlimit Gets the detection limit stored in the dataset
Glacial         Compositions and total pebble counts of 92 glacial tills
groupparts     Group amounts of parts
Hongite         Compositions of 25 specimens of hongite
HouseholdExp   Household Expenditures
Hydrochem       Hydrochemical composition data set of Llobregat river basin water (NE Spain)
idt             Isometric default transform
iit             Isometric identity transform
ilr             Isometric log ratio transform
ilrBase         The canonical basis in the clr plane used for ilr and ipt transforms.
ilt             Isometric log transform
ipt             Isometric planar transform
is.acomp       Check for compositional data type
IsMahalanobisOutlier Checking for outliers
isoPortionLines Isoportion- and Isoproportion-lines
juraset         The jura dataset
kingTetrahedron Ploting composition into rotatable tetrahedron
Kongite         Compositions of 25 specimens of kongite
lines.rmult    Draws connected lines from point to point.
logratioVariogram Empirical variograms for compositions
MahalanobisDist Compute Mahalanobis distances based von robust Estimations
mean.acomp     Mean amounts and mean compositions
meanRow        The arithmetic mean of rows or columns
Metabolites    Steroid metabolite patterns in adults and children
missingProjector Returns a projector the the observed space in case of missings.
missingsInCompositions The policy of treatment of missing values in the "compositions" package
missingSummary Classify and summarize missing values in a dataset
mix.2aplus     Transformations from 'mixtures' to 'compositions' classes
mix.Read       Reads a data file in a mixR format
mvar           Metric summary statistics of real, amount or compositional data
names.acomp    The names of the parts
normalize Normalize vectors to norm 1
norm.default Vector space norm
oneOrDataset Treating single compositions as one-row datasets
OutlierClassifier1 Detect and classify compositional outliers.
outlierplot Plot various graphics to analyse outliers.
outliersInCompositions Analysing outliers in compositions.
pairwisePlot Creates a paneled plot like pairs for two different datasets.
parametricPosdefMat Unique parametrisations for matrices.
perturbe Perturbation of compositions
plot3D plot in 3D based on rgl
plot3D.acomp 3D-plot of compositional data
plot3D.aplus 3D-plot of positive data
plot3D.rmult plot in 3D based on rgl
plot3D.rplus plot in 3D based on rgl
plot.acomp Ternary diagrams
plot.aplus Displaying amounts in scatterplots
plot.logratioVariogram Empirical variograms for compositions
plot.missingSummary Plot a Missing Summary
pMaxMahalanobis Compute distributions of empirical Mahalanobis distances based on simulations
PogoJump Honk Kong Pogo-Jumps Championship
power.acomp Power transform in the simplex
powerofpsdmatrix power transform of a matrix
princomp.acomp Principal component analysis for Aitchison compositions
princomp.aplus Principal component analysis for amounts in log geometry
princomp.rcomp Principal component analysis for real compositions
princomp.rmult Principal component analysis for real data
princomp.rplus Principal component analysis for real amounts
print.acomp Printing compositional data.
qHotellingsTsq Hotellings T square distribution
qqnorm.acomp Normal quantile plots for compositions and amounts
R2 R square
rAitchison Aitchison Distribution
+.rcomp Arithmetic operations for compositions in a real geometry
rcomp Compositions as elements of the simplex embedded in the D-dimensional real space
rcompmargin Marginal compositions in real geometry
rDirichlet Dirichlet distribution
read.geoeas Reads a data file in a geoeas format
relativeLoadings: Loadings of relations of two amounts
replot: Modify parameters of compositional plots.
rlnorm.rplus: The multivariate lognormal distribution
+.rmult: Vectorial arithmetic for datasets in a classical vector scale
rmult: Simple treatment of real vectors
rlnorm.acomp: Normal distributions on special spaces
robustnessInCompositions: Handling robustness issues and outliers in compositions.
+.rplus: Vectorial arithmetic for data sets with rplus class
rplus: Amounts i.e. positive numbers analysed as objects of the real vector space
runif.acomp: The uniform distribution on the simplex
scalar: Parallel scalar products
scale: Normalizing datasets by centering and scaling
Sediments: Proportions of sand, silt and clay in sediments specimens
segments.rmult: Draws straight lines from point to point.
SerumProtein: Serum Protein compositions of blood samples
ShiftOperators: Shifts of machine operators
simpleMissingSubplot: Ternary diagrams
SimulatedAmounts: Simulated amount datasets
simulateMissings: Artificial simulation of various kinds of missings
Skulls: Measurement of skulls
SkyeAFM: AFM compositions of 23 aphyric Skye lavas
split.acomp: Splitting datasets in groups given by factors
straight: Draws straight lines.
summary.acomp: Summarizing a compositional dataset in terms of ratios
summary.aplus: Summaries of amounts
summary.rcomp: Summary of compositions in real geometry
sumMissingProjector: Compute the global projector to the observed subspace.
Supervisor: Proportions of supervisor's statements assigned to different categories
ternaryAxis: Axis for ternary diagrams
totals: Total sum of amounts
tryDebugger: Empirical variograms for compositions
ult: Uncentered log transform
var.acomp: Variances and covariances of amounts and compositions
variation: Variation matrices of amounts and compositions
var.lm: Residual variance of a model
vcovAcomp: Variance covariance matrix of parameters in
compositional regression

vgnFit  Compositional variogram model fitting
vgram2rlvgram vgram2rlvgram
vgram.sph Variogram functions
WhiteCells White-cell composition of 30 blood samples by
two different methods
Yatquat Yatquat fruit evaluation
zeroreplace Zero-replacement routine

Further information is available in the following vignettes:

UsingCompositions Using Animal (source, pdf)

To get detailed "getting started" introduction use help.start() or help.start(browser="myfavouritebrowser")
Go to "Packages" then "compositions" and then "overview" and then launch the file "UsingCompositions.pdf" from there. Please also check the web-site: http://www.stat.boogaart.de/compositions for improved material and our new book expected to appear spring 2009.

The package is devoted to the analysis of multiple amounts. Amounts have typically non-negative values, and often sum up to 100% or one. These constraints lead to spurious effects on the covariance structure, as pointed out by Chayes (1960). The problem is treated rigorously in the monograph by Aitchison (1986), who characterizes compositions as vectors having a relative scale, and identifies its sample space with the D-part simplex. However still (i.e. 2005) most statistical packages do not provided any support for this scale.

The grounding idea of the package exploits the class concept: the analyst gives the data a compositional or amount class, and then all further analysis are (should be) automatically done in a consistent way, e.g. x <- acomp(X); plot(x) should plot the data as a composition (in a ternary diagram) directly without any further interaction of the user.

The package provides four different approaches to analyse amounts. These approaches are associated to four R-classes, representing four different geometries of the sampling space of amounts. These geometries depend on two questions: whether the total sum of the amounts is a relevant information, and which is the meaningful measure of difference of the data.

rplus : (Real Plus) The total amount matters, and amounts should be compared on an absolute basis. i.e. the difference between 1g and 2g is the same as the difference between 1kg and 1001g, one gram. 
aplus : (Aitchison Plus) The total amount matters, but amounts should be compared relatively, i.e. the difference between 1mg and 2mg is the same as that of 1g and 2g: the double. 
acomp : (Aitchison composition) the total amount is constant (or an artifact of the sampling/measurement procedure), and the meaningful difference is a relative one. This class follows the original proposals of Aitchison.

rcomp : (Real composition) the sum is a constant, and the difference in amount from 0% to 1% and from 10% to 11% is regarded as equal. This class represents the raw/naive treatment of compositions as elements of the real simplex based on an absolute geometry. This treatment is implicitly used in most amalgamation problems. However the whole approach suffers from the drawbacks and problems discussed in Chayes (1960) and Aitchison (1986).
The aim of the package is to provide all the functionality to do a consistent analysis in all of these 
approaches and to make the results obtained with different geometries as easy to compare as possi-
bile.

Note

The package compositions has grown a lot in the last year: missings, robust estimations, outlier de-
tection and classification, codadendrogram. This makes everything much more complex especially 
from the side of program testing. Thus we would like to urge our users to report all errors and 
problems of the lastest version (please check first) to support@boogaart.de.

Author(s)

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Pawlowsky-Glahn, V. and J.J. Egozcue (2001) Geometric approach to statistical analysis on the 
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Mateu-Figueras, G. and Barcel\’o-Vidal, C. (Eds.) *Proceedings of the 2nd International Workshop 
on Compositional Data Analysis*, Universitat de Girona, ISBN 84-8458-222-1, [http://ima.udg.es/Activitats/CoDaWork05](http://ima.udg.es/Activitats/CoDaWork05)

van den Boogaart, K.G. and R. Tolosana-Delgado (2008) "compositions": a unified R package to an-
See Also
compositions-package, missingsInCompositions, robustnessInCompositions, outliersInCompositions,

Examples
library(compositions) # load library
data(SimulatedAmounts) # load data sa.lognormals
x <- acomp(sa.lognormals) # Declare the dataset to be compositional
# and use relative geometry
plot(x) # plot.acomp : ternary diagram
ellipses(mean(x),var(x),r=2,col="red") # Simplex 2sigma predictive region
pr <- princomp(x)
straight(mean(x),pr$Loadings)

x <- rcomp(sa.lognormals) # Declare the dataset to be compositional
# and use absolute geometry
plot(x) # plot.acomp : ternary diagram
ellipses(mean(x),var(x),r=2,col="red") # Real 2sigma predictive region
pr <- princomp(x)
straight(mean(x),pr$Loadings)

Aar

Composition of glacial sediments from the Aar massif (Switzerland)

Description
Geochemical composition of glacial sediments from the Aar massif region (Switzerland), major
oxides and trace elements.

Usage
data(Aar)

Details
Composition of recent sediments of several moraines and streams from glaciers around the Aar
massif, including both major oxides and trace elements. The major oxides are expressed in weight
percent (total sum reported in column SumOxides), from Silica (SiO2, column 3) to total Iron 3
Oxide (Fe2O3t, column12, incorporating FeO recasted to Fe2O3). The trace elements are reported
in parts per million (ppm, mg/Kg) between columns 14 (Ba) and 29 (Nd). Partial sum of the trace
elements (in ppm) and of all traces and major oxides (in %) are also reported.
Apart of the compositional information, two covariables are included: Sample and GS. The variable
Sample reports the ID of the sample material. This material was sieved in 11 grain size fractions,
and each fraction was analysed separately after drying. The grain size fraction of each subsample
is reported in variable GS, representing the upper limit of the size fraction reported in \( \phi \) scale, e.g.,
the binary log transformation of the average diameter \( \bar{d} \)

\[
\phi = -\log_2(\bar{d})
\]
The Aar is a granitic-granodioritic-gneissic massif of the Alps, in Switzerland, comprised of several intrusions with different compositions within the range of granitoid lithologies. Details of the region, mineralogy, procedures and study questions behind the data can be found in von Eynatten et al (2012) and references thereon.

Note

Courtesy of H. von Eynatten

Source


References


acomp

Aitchison compositions

Description

A class providing the means to analyse compositions in the philosophical framework of the Aitchison Simplex.

Usage

acomp(X,parts=1:NCOL(oneOrDataset(X)),total=1,warn.na=FALSE,
detectionlimit=NULL,BDL=NULL,MAR=NULL,MNAR=NULL,SZ=NULL)

Arguments

X composition or dataset of compositions

parts vector containing the indices xor names of the columns to be used

total the total amount to be used, typically 1 or 100

warn.na should the user be warned in case of NA,NaN or 0 coding different types of missing values?

detectionlimit a number, vector or matrix of positive numbers giving the detection limit of all values, all columns or each value, respectively

BDL the code for ’Below Detection Limit’ in X

SZ the code for ’Structural Zero’ in X

MAR the code for ’Missing At Random’ in X

MNAR the code for ’Missing Not At Random’ in X
Details

Many multivariate datasets essentially describe amounts of D different parts in a whole. This has some important implications justifying to regard them as a scale for its own, called a composition. This scale was in-depth analysed by Aitchison (1986) and the functions around the class "acomp" follow his approach.

Compositions have some important properties: Amounts are always positive. The amount of every part is limited to the whole. The absolute amount of the whole is noninformative since it is typically due to artifacts on the measurement procedure. Thus only relative changes are relevant. If the relative amount of one part increases, the amounts of other parts must decrease, introducing spurious anticorrelation (Chayes 1960), when analysed directly. Often parts (e.g H2O, Si) are missing in the dataset leaving the total amount unreported and longing for analysis procedures avoiding spurious effects when applied to such subcompositions. Furthermore, the result of an analysis should be independent of the units (ppm, g/l, vol.%, mass.%, molar fraction) of the dataset.

From these properties Aitchison showed that the analysis should be based on ratios or log-ratios only. He introduced several transformations (e.g. clr, alr), operations (e.g. perturbe, power.acomp), and a distance (dist) which are compatible with these properties. Later it was found that the set of compositions equipped with perturbation as addition and power-transform as scalar multiplication and the dist as distance form a D-1 dimensional euclidean vector space (Billheimer, Fagan and Guttorp, 2001), which can be mapped isometrically to a usual real vector space by ilr (Pawlowsky-Glahn and Egozcue, 2001).

The general approach in analysing acomp objects is thus to perform classical multivariate analysis on clr/alr/ilr-transformed coordinates and to backtransform or display the results in such a way that they can be interpreted in terms of the original compositional parts.

A side effect of the procedure is to force the compositions to sum up to a total, which is done by the closure operation clo.

Value

a vector of class "acomp" representing one closed composition or a matrix of class "acomp" representing multiple closed compositions each in one row.

Missing Policy

The policy of treatment of zeroes, missing values and values below detection limit is explained in depth in compositions.missing.

Author(s)


References


Aitchison, J, C. Barcel’o-Vidal, J.J. Egozcue, V. Pawlowsky-Glahn (2002) A concise guide to the algebraic geometric structure of the simplex, the sample space for compositional data analysis, Terra Nostra, Schriften der Alfred Wegener-Stiftung, 03/2003


See Also

`clr`, `rcomp`, `aplus`, `princomp`, `acomp`, `plot.acomp`, `boxplot.acomp`, `barplot.acomp`, `mean.acomp`, `var.acomp`, `variation.acomp`, `cov.acomp`, `msd`

Examples

```r
data(SimulatedAmounts)
plot(acomp(sa.lognormals))
```

---

**acomp**

*Power transform in the simplex*

**Description**

The Aitchison Simplex with its two operations perturbation as + and power transform as * is a vector space. This vector space is represented by these operations.

**Usage**

```r
power.acomp(x, s)
## Methods for class "acomp"
## x*y
## x/y
```
Arguments

- **x**: an acomp composition or dataset of compositions (or a number or a numeric vector)
- **y**: a numeric vector of size 1 or nrow(x)
- **s**: a numeric vector of size 1 or nrow(x)

Details

The power transform is the basic multiplication operation of the Aitchison simplex seen as a vector space. It is defined as:

\[(x * y)_i := clo((x^y)_i)_{\alpha i}\]

The division operation is just the multiplication with \(1/y\).

Value

An "acomp" vector or matrix.

Note

For \(*\) the arguments x and y can be exchanged. Note that this definition generalizes the power by a scalar, since y or s may be given as a scalar, or as a vector with as many components as the composition in acomp x. The result is then a matrix where each row corresponds to the composition powered by one of the scalars in the vector.

Author(s)


References


[http://ima.udg.es/Activitats/CoDaWork03](http://ima.udg.es/Activitats/CoDaWork03)

[http://ima.udg.es/Activitats/CoDaWork05](http://ima.udg.es/Activitats/CoDaWork05)

See Also

ilr, clr, alr.
Examples

```r
acomp(1:5)* -1 + accomp(1:5)
data(SimulatedAmounts)
cdata <- accomp(sa.lognormals)
plot(tmp <- (cdata-mean(cdata))/msd(cdata))
class(tmp)
mean(tmp)
msd(tmp)
var(tmp)
```

### acompmargin

**Marginal compositions in Aitchison Compositions**

**Description**

Compute marginal compositions of selected parts, by computing the rest as the geometric mean of the non-selected parts.

**Usage**

```r
acompmargin(X,d=c(1,2),name="*",pos=length(d)+1,what="data")
```

**Arguments**

- `X` composition or dataset of compositions
- `d` vector containing the indices xor names of the columns selected
- `name` The new name of the amalgamation column
- `pos` The position where the new amalgamation column should be stored. This defaults to the last column.
- `what` The role of X either "data" for data (or means) to be transformed or "var" for (acomp-clr)-variances to be transformed.

**Details**

The amalgamation column is simply computed by taking the geometric mean of the non-selected components. This is consistent with the `acomp` approach and gives clear ternary diagrams. However, this geometric mean is difficult to interpret.

**Value**

A closed compositions with class "acomp" containing the variables given by d and the the amalgamation column.

**Missing Policy**

MNAR has the highest priority, MAR afterwards, and WZERO (BDL,SZ) values are considered as 0 and finally reported as BDL.
Author(s)

References
Vera Pawlowsky-Glahn (2003) personal communication. Universitat de Girona. vera.pawlowsky@udg.es


See Also
rcompmargin, acomp

Examples

data(SimulatedAmounts)
plot.acomp(sa.lognormals5,margin="acomp")
plot.acomp(acompmargin(sa.lognormals5,c("Pb","Zn")))
plot.acomp(acompmargin(sa.lognormals5,c(1,2)))

acomp scalar product

inner product for datasets with a vector space structure

Description
acomp and aplus objects are considered as (sets of) vectors. The %*% is considered as the inner multiplication. An inner multiplication with another vector is the scalar product. An inner multiplication with a matrix is a matrix multiplication, where the vectors are either considered as row or as column vector.

Usage

## S3 method for class 'acomp'
x %*% y

## S3 method for class 'aplus'
x %*% y

Arguments

x a acomp or aplus object or a matrix interpreted in clr, ilr or ilt coordinates
y a acomp or aplus object or a matrix interpreted in clr, ilr or ilt coordinates
Details

The operators try to mimic the behavior of \%\% on c()-vectors as inner product, applied in parallel to all row-vectors of the dataset. Thus the product of a vector with a vector of the same type results in the scalar product of both. For the multiplication with a matrix each vector is considered as a row or column, whatever is more appropriate. The matrix itself is considered as representing a linear mapping (endomorphism) of the vector space to a space of the same type. The mapping is represented in clr, ilr or ilt coordinates. Which of the aforementioned coordinate systems is used is judged from the type of x and from the dimensions of the A.

Value

Either a numeric vector containing the scalar products, or an object of type acomp or aplus containing the vectors transformed with the given matrix.

Author(s)


See Also

%\%\%.rmult

Examples

\begin{verbatim}
x <- acomp(matrix( sqrt(1:12), ncol= 3 ))
x%*%x
A <- matrix( 1:9,nrow=3)
x %*% A %*% x
x %*% A
A %*% x
A <- matrix( 1:4,nrow=2)
x %*% A %*% x
x %*% A
A %*% x
x <- aplus(matrix( sqrt(1:12), ncol= 3 ))
x%*%x
A <- matrix( 1:9,nrow=3)
x %*% A %*% x
x %*% A
A %*% x
\end{verbatim}

Description

Proportion of a day in activity teaching, consulting, administrating, research, other wakeful activities and sleep for 20 days are given.
Details

The activity of an academic statistician were divided into following six categories

<table>
<thead>
<tr>
<th>teac</th>
<th>teaching</th>
</tr>
</thead>
<tbody>
<tr>
<td>cons</td>
<td>consultation</td>
</tr>
<tr>
<td>admi</td>
<td>administration</td>
</tr>
<tr>
<td>rese</td>
<td>research</td>
</tr>
<tr>
<td>wake</td>
<td>other wakeful activities</td>
</tr>
<tr>
<td>slee</td>
<td>sleep</td>
</tr>
</tbody>
</table>

Data show the proportions of the 24 hours devoted to each activity, recorded on each of 20 days, selected randomly from working days in alternate weeks, so as to avoid any possible carry-over effects, such as short-sleep day being compensated by make-up sleep on a succeeding day.

The six activity may be divided into two categories 'work' comprising activities 1,2,3,4: and 'leisure' comprising activities 5 and 6.

All rows sum to one.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name STATDAY.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


<table>
<thead>
<tr>
<th>Activity31</th>
<th>Activity patterns of a statistician for 20 days</th>
</tr>
</thead>
</table>

Description

Proportion of a day in activity teaching, consulting, administrating, research, other wakeful activities and sleep for 20 days are given.

Usage

data(Activity31)
Details

The activity of an academic statistician were divided into following six categories
Data shows the proportions of the 24 hours devoted to each activity, recorded on each of 20 days, selected randomly from working days in alternate weeks, so as to avoid any possible carry-over effects, such as short-sleep day being compensated by make-up sleep on a succeeding day.

The six activities may be divided into two categories 'work' comprising activities 1, 2, 3, 4: and 'leisure' comprising activities 5 and 6.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name ACTIVITY.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


---

### alr

**Additive log ratio transform**

#### Description

Compute the additive log ratio transform of a (dataset of) composition(s), and its inverse.

#### Usage

```r
alr( x ,ivar=ncol(x), ... )
alrInv( z, ...,orig=NULL)
```

#### Arguments

- **x**
  - a composition, not necessarily closed
- **z**
  - the alr-transform of a composition, thus a (D-1)-dimensional real vector
- **...**
  - generic arguments, not used.
- **orig**
  - a compositional object which should be mimicked by the inverse transformation. It is especially used to reconstruct the names of the parts.
- **ivar**
  - The column to be used as denominator variable. Unfortunately not yet supported in alrInv. The default works even if x is a vector.
Details

The alr-transform maps a composition in the D-part Aitchison-simplex non-isometrically to a D-1 dimensional euclidian vector, treating the last part as common denominator of the others. The data can then be analysed in this transformation by all classical multivariate analysis tools not relying on a distance. The interpretation of the results is relatively simple, since the relation to the original D-1 first parts is preserved. However distance is an extremely relevant concept in most types of analysis, where a clr or ilr transformation should be preferred.

The additive logratio transform is given by

$$alr(x)_i := \ln \frac{x_i}{x_D}$$

Value

alr gives the additive log ratio transform; accepts a compositional dataset alrInv gives a closed composition with the given alr-transform; accepts a dataset

Author(s)


References


See Also

clr.alr.apt, http://ima.udg.es/Activitats/CoDaWork03

Examples

```r
(tmp <- alr(c(1,2,3)))
alrInv(tmp)
unclass(alrInv(tmp)) - clo(c(1,2,3)) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(alr(cdata),pch=".")
```
AnimalVegetation  Animal and vegetation measurement

Description

Areal compositions by abundance of vegetation and animals for 50 plots in each of regions A and B.

Usage

data(AnimalVegetation)

Details

In a regional ecology study, plots of land of equal area were inspected and the parts of each plot which were thick or thin in vegetation and dense or sparse in animals were identified. From this field work the areal proportions of each plot were calculated for the four mutually exclusive and exhaustive categories: thick-dense, thick-sparse, thin-dense, thin-sparse. These sets of proportions are recorded for 50 plots from each of two different regions A and B. All rows sum to 1, except for some rounding errors.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name ANIVEG.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


aplus  Amounts analysed in log-scale

Description

A class to analyse positive amounts in a logistic framework.

Usage

aplus(X, parts=1:NCOL(oneOrDataset(X)), total=NA, warn.na=FALSE, detectionlimit=NULL, BDL=NULL, MAR=NULL, MNAR=NULL, SZ=NULL)
**Arguments**

- **X**: vector or dataset of positive numbers
- **parts**: vector containing the indices xor names of the columns to be used
- **total**: a numeric vector(s) giving the total amounts of each dataset.
- **warn.na**: should the user be warned in case of NA,NaN or 0 coding different types of missing values?
- **detectionlimit**: a number, vector or matrix of positive numbers giving the detection limit of all values, all columns or each value, respectively
- **BDL**: the code for ‘Below Detection Limit’ in X
- **SZ**: the code for ‘Structural Zero’ in X
- **MAR**: the code for ‘Missing At Random’ in X
- **MNAR**: the code for ‘Missing Not At Random’ in X

**Details**

Many multivariate datasets essentially describe amounts of D different parts in a whole. When the whole is large in relation to the considered parts, such that they do not exclude each other, or when the total amount of each component is indeed determined by the phenomenon under investigation and not by sampling artifacts (such as dilution or sample preparation), then the parts can be treated as amounts rather than as a composition (cf. acomp, rcomp).

Like compositions, amounts have some important properties. Amounts are always positive. An amount of exactly zero essentially means that we have a substance of another quality. Different amounts - spanning different orders of magnitude - are often given in different units (ppm, ppb, g/l, vol.%, mass %, molar fraction). Often, these amounts are also taken as indicators of other non-measured components (e.g. K as indicator for potassium feldspar), which might be proportional to the measured amount. However, in contrast to compositions, amounts themselves do matter. Amounts are typically heavily skewed and in many practical cases a log-transform makes their distribution roughly symmetric, even normal.

In full analogy to Aitchison’s compositions, vector space operations are introduced for amounts: the perturbation perturbe.aplus as a vector space addition (corresponding to change of units), the power transformation power.aplus as scalar multiplication describing the law of mass action, and a distance dist which is independent of the chosen units. The induced vector space is mapped isometrically to a classical $R^D$ by a simple log-transformation called ilt, resembling classical log transform approaches.

The general approach in analysing aplus objects is thus to perform classical multivariate analysis on ilt-transformed coordinates (i.e., logs) and to backtransform or display the results in such a way that they can be interpreted in terms of the original amounts.

The class aplus is complemented by the rplus, allowing to analyse amounts directly as real numbers, and by the classes acomp and rcomp to analyse the same data as compositions disregarding the total amounts, focusing on relative weights only.

The classes rcomp, acomp, aplus, and rplus are designed as similar as possible in order to allow direct comparison between results achieved by the different approaches. Especially the acomp simplex transforms clr, alr, ilr are mirrored in the aplus class by the single bijective isometric transform ilt.
Value

A vector of class "aplus" representing a vector of amounts or a matrix of class "aplus" representing multiple vectors of amounts, each vector in one row.

Missing Policy

The policy of treatment of zeroes, missing values and values below detection limit is explained in depth in `compositions.missing`.

Author(s)


References


See Also

`ilt`, `acomp`, `rplus`, `princomp.aplus`, `plot.aplus`, `boxplot.aplus`, `barplot.aplus`, `mean.aplus`, `var.aplus`, `variation.aplus`, `cov.aplus`, `msd`

Examples

```r
data(SimulatedAmounts)
plot(aplus(sa.lognormals))
```

aplusarithm

Vectorial arithmetic for data sets with aplus class

Description

The positive vectors equipped with the perturbation (defined as the element-wise product) as Abelian sum, and powertransform (defined as the element-wise powering with a scalar) as scalar multiplication forms a real vector space. These vector space operations are defined here in a similar way to `+.rmult`.

Usage

```r
perturbe.aplus(x, y)
## S3 method for class 'aplus'
x + y
## S3 method for class 'aplus'
x - y
## S3 method for class 'aplus'
x * y
```

## S3 method for class 'aplus'
### Methods for `aplus`

- `x+y`
- `x-y`
- `-x`
- `x*r`
- `r*x`
- `x/r`

```r
power.aplus(x, r)
```

### Arguments

- `x` an aplus vector or dataset of vectors
- `y` an aplus vector or dataset of vectors
- `r` a numeric vector of size 1 or `nrow(x)`

### Details

The operators try to mimic the parallel operation of R for vectors of real numbers to vectors of amounts, represented as matrices containing the vectors as rows and works like the operators for `{rmult}`

### Value

an object of class "aplus" containing the result of the corresponding operation on the vectors.

### Author(s)


### See Also

`rmult`, `%*%`.

### Examples

```r
x <- aplus(matrix(sqrt(1:12), ncol = 3))
x
x+x
x + aplus(1:3)
x * 1:4
1:4 * x
x / 1:4
x / 10
power.aplus(x, 1:4)
```
apt \hspace{0.5cm} \textit{Additive planar transform}

\textbf{Description}

Compute the additive planar transform of a (dataset of) compositions or its inverse.

\textbf{Usage}

\begin{verbatim}
apt( x ,...)  
aptInv( z ,..., orig=NULL)
\end{verbatim}

\textbf{Arguments}

\begin{itemize}
  \item \texttt{x} \hspace{0.5cm} a composition or a matrix of compositions, not necessarily closed
  \item \texttt{z} \hspace{0.5cm} the apt-transform of a composition or a matrix of alr-transforms of compositions
  \item \texttt{...} \hspace{0.5cm} generic arguments, not used.
  \item \texttt{orig} \hspace{0.5cm} a compositional object which should be mimicked by the inverse transformation. It is especially used to reconstruct the names of the parts.
\end{itemize}

\textbf{Details}

The apt-transform maps a composition in the D-part real-simplex linearly to a D-1 dimensional euclidian vector. Although the transformation does not reach the whole $\mathbb{R}^{D-1}$, resulting covariance matrices are typically of full rank.

The data can then be analysed in this transformation by all classical multivariate analysis tools not relying on distances. See \texttt{cpt} and \texttt{ipt} for alternatives. The interpretation of the results is easy since the relation to the first D-1 original variables is preserved.

The additive planar transform is given by

$$apt(x)_i := clo(x)_i, i = 1, \ldots, D - 1$$

\textbf{Value}

\begin{itemize}
  \item apt gives the centered planar transform, aptInv gives closed compositions with the given apt-transforms
\end{itemize}

\textbf{Author(s)}

K.Gerald v.d. Boogaart \texttt{http://www.stat.boogaart.de}
References


See Also

alr, cpt, ipt

Examples

```r
(tmp <- apt(c(1,2,3)))
aptInv(tmp)
aptInv(tmp) - clo(c(1,2,3)) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(apt(cdata),pch=".")
```

---

**ArcticLake**

*Artic lake sediment samples of different water depth*

Description

Sand, silt and clay compositions of 39 sediment samples of different water depth in an Arctic lake.

Usage

data(ArcticLake)

Details

Sand, silt and clay compositions of 39 sediment samples at different water depth (in meters) in an Arctic lake. The additional feature is a concomitant variable or *covariate*, water depth, which may account for some of the variation in the compositions. In statistical terminology we have a multivariate regression problem with sediment composition as regressand and water depth as regressor.

All row percentage sums to 100, except for rounding errors.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name ARCTIC.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References

Description

adds 3-dimensional arrows to an rgl plot.

Usage

arrows3D(...)

## Default S3 method:
arrows3D(x0, x1, ..., length = 0.25,
angle = 30, code = 2, col = "black",
lty = NULL, lwd = 2, orth = c(1, 0.0001, 0.0000001),
labs = NULL, size = lwd)

Arguments

x0  a matrix or vector giving the starting points of the arrows
x1  a matrix or vector giving the end points of the arrows
... additional plotting parameters as described in \texttt{rgl.material}
length a number giving the length of the arrowhead
angle numeric giving the angle of the arrowhead
code 0= no arrowhead, 1= arrowhead at x0, 2= arrowhead at x1, 3= double headed
col the color of the arrow
lty Not implemented, here for compatibility reasons with \texttt{arrows}
lwd line width in pixels
orth the flat side of the arrow is not unique by x0 and x1. This ambiguity is solved in a way that the arrow seams as wide as possible from the viewing direction orth.
labs labels to be plotted to the endpoints of the arrows
size size of the plotting symbol

details

The function is called to plot arrows into an rgl plot. The size of the arrow head is given in a absolute way. Therefore it is important to give the right scale for the length, to see the arrow head and that it does not fill the whole window.

Value

the 3D plotting coordinates of the tips of the arrows displayed, returned invisibly

Author(s)

K.Gerald v.d. Boogaart \url{http://www.stat.boogaart.de}
as.data.frame

See Also

points3d, plot, plot3D,

Examples

x <- cbind(rnorm(10), rnorm(10), rnorm(10))
plot3D(x)
x0 <- x*0
arrows3D(x0, x)

as.data.frame( Convert "compositions" classes to data frames

Description

Convert a compositional object to a dataframe

Usage

## S3 method for class 'acompl'
as.data.frame(x, ...)
## S3 method for class 'rcomp'
as.data.frame(x, ...)
## S3 method for class 'aplus'
as.data.frame(x, ...)
## S3 method for class 'rplus'
as.data.frame(x, ...)
## S3 method for class 'rmult'
as.data.frame(x, ...)

Arguments

x an object to be converted to a dataframe

... additional arguments are not used

Value

a data frame containing the given data.

Author(s)

Examples

```r
data(SimulatedAmounts)
as.data.frame(acomp(sa.groups))
# The central purpose of providing this command is that the following
# works properly:
data.frame(acomp(sa.groups), groups=sa.groups.area)
```

Description

Adds a coordinate system to a 3D rgl graphic. In future releases, functionality to add tickmarks will be (hopefully) provided. Now, it is just a system of arrows giving the directions of the three axes.

Usage

```r
axis3D(axis.origin=c(0, 0, 0), axis.scale=1, axis.col="gray", vlabs=c("x", "y", "z"),
vlabs.col=axis.col, bbox=FALSE, axis.lwd=2, axis.len=mean(axis.scale)/10,
axis.angle=30, orth=c(1, 0.0001, 0.000001), axes=TRUE, ...)
```

Arguments

- `axis.origin`: The location where to put the origin of the coordinate arrows typically either 0, the minimum or the mean of the dataset.
- `axis.scale`: either a number or a 3D vector giving the length of the arrows for the axis in the coordinates of the plot.
- `axis.col`: Color to plot the coordinate system.
- `vlabs`: The names of the axes, plotted at the end.
- `vlabs.col`: color for the axes labels.
- `bbox`: boolean, whether to plot a bounding box.
- `axis.lwd`: line width of the axes.
- `axis.angle`: angle of the arrow heads.
- `axis.len`: length of the arrow heads.
- `orth`: the orth argument of `arrows3D`.
- `axes`: a boolean, whether to plot the axes.
- `...`: these arguments are passed to `arrows3D` as `rgl.material` arguments.

Details

The function is called to plot a coordinate system consisting of arrows into an rgl plot.

Value

Nothing
### balance

**Compute balances for a compositional dataset.**

#### Description

Compute balances in a compositional dataset.

#### Usage

```r
balance(X,...)
## S3 method for class 'acomp'
balance(X,expr,...)
## S3 method for class 'rcomp'
balance(X,expr,...)
## S3 method for class 'aplus'
balance(X,expr,...)
## S3 method for class 'rplus'
balance(X,expr,...)
balance01(X,...)
## S3 method for class 'acomp'
balance01(X,expr,...)
## S3 method for class 'rcomp'
balance01(X,expr,...)
balanceBase(X,...)
## S3 method for class 'acomp'
balanceBase(X,expr,...)
## S3 method for class 'rcomp'
balanceBase(X,expr,...)
```

#### Examples

```r
x <- cbind(rnorm(10),rnorm(10),rnorm(10))
plot3D(x)
x0 <- x*0
arrows3D(x0,x)
```
Arguments

- **X**: compositional dataset (or optionally just its column names for balanceBase)
- **expr**: a ~ formula using the column names of X as variables and separating them by '/' and organize by parenthesis ( ). : and * can be used instead of '/' when the corresponding balance should not be created. - can be used as synonym to '/' in the real geometries. 1 can be used in the unclosed geometries to level against a constant.

Details

For acomp-compositions balances are defined as orthogonal projections representing the log ratio of the geometric means of subsets of elements. Based on a recursive subdivision (provided by the expr=) this projections provide a (complete or incomplete) basis of the clr-plane. The basis is given by the balanceBase functions. The transform is given by the balance functions. The balance01 functions are a backtransform of the balances to the amount of the first portion if this was the only balance in a 2 element composition, providing an "interpretation" for the values of the balances.

The package tries to give similar concepts for the other scales. For rcomp objects the concept is mainly unchanges but augmented by a virtual component 1, which always has portion 1.

For rcomp objects, we choose not a "orthogonal" transformation since such a concept anyway does not really exist in the given space, but merely use the difference of one subset to the other. The balance01 is than not really a transform of the balance but simply the portion of the first group of parts in all contrasted parts.

For rplus objects we just used an analog to generalisation from the rcomp definition as aplus is generalized from acomp. However at this time we have no idea wheter this has any usefull interpretation.

Value

- **balance**: a matrix (or vector) with the corresponding balances of the dataset.
- **balance01**: a matrix (or vector) with the corresponding balances in the dataset transformed in the given geometry to a value between 0 and 1.
- **balanceBase**: a matrix (or vector) with column vectors giving the transform in the cdt-transform used to achieve the corresponding balances.

References

http://ima.udg.es/Activitats/CoDaWork08  Papers of Boogaart and Tolosana
http://ima.udg.es/Activitats/CoDaWork05  Paper of Egozcue

See Also

clr, ilr, ipt, ilrBase
### Examples

```r
X <- rnorm(100)
Y <- rnorm.acomp(100, acomp(c(A=1, B=1, C=1)), 0.1*diag(3)) + acomp(t(outer(c(0.2, 0.3, 0.4), X, "^")))
colnames(Y) <- c("A", "B", "C")

subComps <- function(X, ...){
  X <- oneOrDataset(X)
  nams <- sapply(all, function(x) paste(x[[2]], x[[3]], sep = ","))
  val <- sapply(all, function(x){
    a = X[, match(as.character(x[[2]]), colnames(X))]
    b = X[, match(as.character(x[[2]]), colnames(X))]
    c = X[, match(as.character(x[[3]]), colnames(X))]
    return(a/(b+c))
  })
  colnames(val)<-nams
  val
}
pairs(cbind(ilr(Y), X), panel=function(x,y,...) {points(x,y,...); abline(lm(y~x))})
pairs(cbind(balance(Y, ~A/B/C), X), panel=function(x,y,...) {points(x,y,...); abline(lm(y~x))})

pairwisePlot(balance(Y, ~A/B/C), X)
pairwisePlot(X, balance(Y, ~A/B/C), panel=function(x,y,...) {plot(x,y,...); abline(lm(y~x))})
pairwisePlot(X, balance01(Y, ~A/B/C))
pairwisePlot(X, subComps(Y, A~B, A~C, B~C))

balance(rcomp(Y), ~A/B/C)
balance(aplus(Y), ~A/B/C)
balance(rplus(Y), ~A/B/C)
```

### barplot.acomp

**Bar charts of amounts**

### Description

Compositions and amounts displayed as bar plots.

### Usage

```r
## S3 method for class 'acomp'
barplot(height, ..., legend.text=TRUE, beside=FALSE, total=1,
plotMissings=TRUE, missingColor="red", missingPortion=0.01)
## S3 method for class 'rcomp'
barplot(height, ..., legend.text=TRUE, beside=FALSE, total=1,
plotMissings=TRUE, missingColor="red", missingPortion=0.01)
```
```r
## S3 method for class 'aplus'
barplot(height, ..., legend.text=TRUE, beside=TRUE, total=NULL,
plotMissings=TRUE, missingColor="red", missingPortion=0.01)
## S3 method for class 'rplus'
barplot(height, ..., legend.text=TRUE, beside=TRUE, total=NULL,
plotMissings=TRUE, missingColor="red", missingPortion=0.01)
## S3 method for class 'ccomp'
barplot(height, ..., legend.text=TRUE, beside=FALSE, total=1,
plotMissings=TRUE, missingColor="red", missingPortion=0.01)
```

### Arguments

- **height**: an acomp, rcomp, aplus, or rplus object giving amounts to be displayed
- **...**: further graphical parameters as in `barplot`
- **legend.text**: same as legend.text in `barplot`
- **beside**: same as beside in `barplot`
- **total**: The total to be used in displaying the composition, typically 1, 100 or the number of parts. If NULL no normalisation takes place.
- **plotMissings**: logical: shall missings be annotate in the plot
- **missingColor**: color to draw missings
- **missingPortion**: The space portion to be reserved for missings

### Details

These functions are essentially light-weighted wrappers for `barplot`, just adding an adequate default behavior for each of the scales. The missingplot functionality will work well with the default settings.

If `plotMissings` is true, there will be an additional portion introduced, which is not counted in the total. This might make the plots looking less nice, however they make clear to the viewer that it is by no means clear how the rest of the plot should be interpreted and that the missing value really casts some unsureness on the rest of the data.

### Value

A numeric vector (or matrix, when beside = TRUE) giving the coordinates of all the bar midpoints drawn, as in `barplot`

### Author(s)


### See Also

- `acomp`, `rcomp`, `rplus`, `aplus`, `plot.acomp`, `boxplot.acomp`
**Examples**

```r
data(SimulatedAmounts)
barplot(mean(acomp(sa.lognormals[1:10,])))
barplot(mean(rcomp(sa.lognormals[1:10,])))
barplot(mean(aplus(sa.lognormals[1:10,])))
barplot(mean(rplus(sa.lognormals[1:10,])))

barplot(acomp(sa.lognormals[1:10,]))
barplot(rcomp(sa.lognormals[1:10,]))
barplot(aplus(sa.lognormals[1:10,]))
barplot(rplus(sa.lognormals[1:10,]))

barplot(acomp(sa.tnormals))
```
References

binary
Treating binary and g-adic numbers

Description
Allows the access to individual digits in binary (and general g-adic) numbers.

Usage

```r
binary(x, mb=max(maxBit(x, g)), g=2)
unbinary(x, g=2)
bit(x, b, g=2)
## S3 method for class 'numeric'
bit(x, b=0:maxBit(x, g), g=2)
## S3 method for class 'character'
bit(x, b=0:maxBit(x, g), g=2)
bit(x, b, g=2) <- value
## S3 replacement method for class 'numeric'
bit(x, b=0:maxBit(x, g), g=2) <- value
## S3 replacement method for class 'character'
bit(x, b=0:maxBit(x, g), g=2) <- value
maxBit(x, g=2)
## S3 method for class 'numeric'
maxBit(x, g=2)
## S3 method for class 'character'
maxBit(x, g=2)
bitCount(x, mb=max(maxBit(x, g)), g=2)
gsi.orSum(..., g=2)
whichBits(x, mb=max(maxBit(x, g)), g=2, values=c(TRUE))
binary2logical(x, mb=max(maxBit(x, g)), g=2, values=c(TRUE))
```

Arguments

- `x` a number either represented a g-adic character string or as a integral numeric value
- `b` the indices of the bits to be processed. The least significant bit has index 0.
- `mb` maximal bit. The index of the most significant bit to be treated
- `g` the base of the g-adic representation. 2 corresponds to binary numbers, 8 to octal numbers, 16 to hexadecimal numbers. g is limited by 36.
- `value` a vector of bit values to be selected or setted.
- `values` a vector of bit values that should be considered as TRUE.
- `...` some binary numbers
Details

These routines are primarily intended to manipulate g-adic numbers for user interface purposes and condensed representation of information. They are not intended for a long number arithmetic.

Value

- **binary** returns a standard binary (or g-adic) character representation of the number
- **unbinary** returns a binary (or g-adic) representation of the number
- **bit** returns the values of the requested bits. The values are returned as a logical vector for binary numbers and as numeric digit values for other g-adic numbers.
- **maxBit** returns the most significant bit represented in the number. This is the highest bit set in numeric numbers and the highest actually given character in a character representation.
- **bitCount** returns the g-adic crossfoot of the number. For a binary number this is the number of bits set
- **gsi.orSum** Only works for binary numbers and does a parallel or on each of the bits for a list of binary numbers.
- **whichBits** returns the indices of the bits actually set (or more precisely of the bits with value in `values`)
- **binary2logical** returns the a true false vector of the bits actually set (or more precisely of the bits with value in `values`)

Author(s)


See Also

outlierplot

Examples

```r
(x<-unbinary("10101010"))
y<-binary(x)
bit(x,1:3)
bit(y,0:3)
maxBit(x)
maxBit(y)
whichBits(x)
whichBits(y)
binary2logical(y)
bit(x)
bit(y)
gsi.orSum(y,1)
bitCount(x)
bitCount(y)
bit(x,2)<-1
x
```
biplot3D

Three-dimensional biplots, based on package rgl

Description

Plots variables and cases in the same plot, based on a principal component analysis.

Usage

biplot3D(x,...)

## Default S3 method:
biplot3D(x,y,var.axes=TRUE,col=c("green","red"),cex=c(2,2),
  xlabs = NULL, ylabs = NULL, expand = 1, arrow.len = 0.1,
  ... , add=FALSE)

## S3 method for class 'princomp'
biplot3D(x,choices=1:3,scale=1,...,
  comp.col=1,comp.labs=paste("Comp.",1:3),
  scale.scores=lambda[choices]^(1-scale),
  scale.var=scale.comp, scale.comp=sqrt(lambda[choices]),
  scale.disp=1/scale.comp)

Arguments

x          princomp object or matrix of point locations to be drawn (typically, cases)
choices    Which principal components should be used?
scale      a scaling parameter like in biplot
scale.scores a vector giving the scaling applied to the scores
scale.var   a vector giving the scaling applied to the variables
scale.comp  a vector giving the scaling applied to the unit length of each component
scale.disp  a vector giving the scaling of the display in the directions of the components
comp.col    color to draw the axes of the components, defaults to black
comp.labs   labels for the components
...         further plotting parameters as defined in rgl.material
y          matrix of second point/arrow-head locations (typically, variables)
var.axes   logical, TRUE draws arrows and FALSE points for y
col         vector/list of two elements the first giving the color/colors for the first data set
            and the second giving color/colors for the second data set.
cex         vector/list of two elements the first giving the size for the first data set and the
            second giving size for the second data set.
**Blood23**

xlabs  labels to be plotted at x-locations
ylabs  labels to be plotted at y-locations
expand the relative expansion of the y data set with respect to x
arrow.len The length of the arrows as defined in arrows3D
add logical, adding to existing plot or making a new one?

**Details**

This "biplot" is a triplot, relating data, variables and principal components. The relative scaling of
the components is still experimental, meant to mimic the behavior of classical biplots.

**Value**

the 3D plotting coordinates of the tips of the arrows of the variables displayed, returned invisibly

**Author(s)**


**See Also**

gsi

**Examples**

data(SimulatedAmounts)
pc <- princomp(acomp(sa.lognormals5))
pc
summary(pc)
plot(pc)       #plot(pc,type="screeplot")
biplot3D(pc)


---

**Blood23  
**Blood samples

**Description**

Percentage of different leukocytes in blood samples of ten patients, determined by four different
methods.

**Usage**

data(Blood23)

**Details**

In a comparative study of four different methods of assessing leukocytes composition of a blood
sample, aliquots of blood samples from ten patients were assessed by the four methods. Data show
the percentage in the 40 analyses of:
Boxite

P: polymorphonuclear leukocytes,
S: small lymphocytes,
L: large mononuclears.

All rows sum to 100.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name BLOOD.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


Boxite

<table>
<thead>
<tr>
<th>Description</th>
<th>Compositions and depth of 25 specimens of boxite</th>
</tr>
</thead>
</table>

Description

A mineral compositions of 25 rock specimens of boxite type. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite, as well as the depth of location.

Usage

data(Boxite)

Details

A mineral compositions of 25 rock specimens of boxite type are given. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite, and the recorded depth of location of each specimen. We abbreviate the minerals names to A, B, C, D, E.

All row percentage sums to 100, except the 6-th 102.3 and the 10-th 99.9.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name BOXITE.DAT, here included under the GNU Public Library Licence Version 2 or newer.
Displaying compositions and amounts with box-plots

Description

For the different interpretations of amounts or compositional data, a different type of boxplot is feasible. Thus different boxplots are drawn.

Usage

```r
## S3 method for class 'acomp'
boxplot(x,fak=NULL,...,
    xlim=NULL,ylim=NULL,log=TRUE,
    panel=vp.logboxplot,dots=!boxes,boxes=TRUE,
    notch=FALSE,
    plotMissings=TRUE,
    mp=~simpleMissingSubplot(missingPlotRect,
                            missingInfo,c("NM","TM",cn))
)

## S3 method for class 'rcomp'
boxplot(x,fak=NULL,...,
    xlim=NULL,ylim=NULL,log=FALSE,
    panel=vp.boxplot,dots=!boxes,boxes=TRUE,
    notch=FALSE,
    plotMissings=TRUE,
    mp=~simpleMissingSubplot(missingPlotRect,
                            missingInfo,c("NM","TM",cn)))

## S3 method for class 'aplus'
boxplot(x,fak=NULL,...,log=TRUE,
    plotMissings=TRUE,
    mp=~simpleMissingSubplot(missingPlotRect,
                            missingInfo,names(missingInfo)))

## S3 method for class 'rplus'
boxplot(x,fak=NULL,...,ylim=NULL,log=FALSE,
    plotMissings=TRUE,
    mp=~simpleMissingSubplot(missingPlotRect,
                            missingInfo,names(missingInfo)))

vp.boxplot(x,y,...,dots=FALSE,boxes=TRUE,xlim=NULL,ylim=NULL,log=FALSE,
           notch=FALSE,plotMissings=TRUE,
           mp=~simpleMissingSubplot(missingPlotRect,
                                    missingInfo,c("NM","TM",cn)),
           ...)```

References

missingness=attr(y,"missingness")
)
vp.logboxplot(x,y,...,dots=FALSE,boxes=TRUE,xlim,ylim,log=TRUE,notch=FALSE,
plotMissings=TRUE,
mp=simpleMissingSubplot(missingPlotRect,
missingInfo,c("NM","TM",cn)),
missingness=attr(y,"missingness"))

Arguments

x    a data set
fak   a factor to split the data set, not yet implemented in aplus and rplus
xlim  x-limits of the plot.
ylim  y-limits of the plot.
log   logical indicating whether ploting should be done on log scale
panel the panel function to be used or a list of multiple panel functions
...  further graphical parameters
dots  a logical indicating whether the points should be drawn
boxes a logical indicating whether the boxes should be drawn
y     used by pairs
notch logical, should the boxes be notched?
plotMissings Logical indicating that missings should be displayed.
mp    A formula providing a function call, which will be evaluated within each panel
       with missings to plot the missingness situation. The call can use the variables
       missingPlotRect, which provides a rectangle to plot the information to in a
       par("usr") like specification. In the rX is the current data
missingness The missingness information as a result from missingType of the full data
       information the panels could base there missing plots on.

Details

boxplot.aplus and boxplot.rplus are wrappers of bxp, which just take into account the possible
logarithmic scale of the data.

boxplot.acomp and boxplot.rcomp generate a matrix of box-plots, where each cell represents the
difference between the row and column variables. Such difference is respectively computed as a
log-ratio and a rest.

vp.boxplot and vp.logboxplot are only used as panel functions. They should not be directly
called.

Author(s)

**ccomp**

See Also

plot.acomp, qqnorm.acomp

Examples

data(SimulatedAmounts)
boxplot(acomp(sa.lognormals))
boxplot(rcomp(sa.lognormals))
boxplot(aplus(sa.lognormals))
boxplot(rplus(sa.lognormals))
# And now with missing!!!
boxplot(acomp(sa.tnormals))

---

**ccomp**  

**Count compositions**

Description

A class providing the means to analyse count compositions understood as Poisson or multinomial realisation, where the portions are given by an unknown Aitchison compositions.

Usage

```r
ccomp(X, parts = 1:NCOL(oneOrDataset(X)), total = NA, warn.na = FALSE,
      detectionlimit = NULL, BDL = NULL, MAR = NULL, MNAR = NULL, SZ = NULL)
```

Arguments

- **X**  
  composition or dataset of compositions
- **parts**  
  vector containing the indices xor names of the columns to be used
- **total**  
  the total amount to be used, typically 1 or 100
- **warn.na**  
  should the user be warned in case of NA,NaN or 0 coding different types of missing values?
- **detectionlimit**  
  a number, vector or matrix of positive numbers giving the detection limit of all values, all columns or each value, respectively
- **BDL**  
  the code for ‘Below Detection Limit’ in X
- **SZ**  
  the code for ‘Structural Zero’ in X
- **MAR**  
  the code for ‘Missing At Random’ in X
- **MNAR**  
  the code for ‘Missing Not At Random’ in X
Details

A count composition contains an indirect observation of an Aitchison composition by a Poisson or
multinomial variable. A count composition can only contain integer counts. It is assumed that the
total sum is an artefact and does not contain information on the actual composition. But it does
contain information on the precision of the relative observation.

Value

a vector of class "ccomp" representing count composition or a matrix of class "ccomp" representing
multiple count compositions each in one row.

Missing Policy

The policy of treatment of zeroes, missing values and values below detection limit is explained in
depth in compositions.missing.

Author(s)


See Also

barplot.ccomp ccompMultinomialGOF.test ccompPoissonGOF.test cdt.ccomp cdtInv.ccomp
fitSameMeanDifferentVarianceModel groupparts.ccomp idt.ccomp idtInv.ccomp
mean.ccomp names<-.ccomp names.ccomp plot.ccomp PoissonGOF.test rmultinom.ccomp
rnorm.ccomp rpois.ccomp split.ccomp totals.ccomp

Examples

data(SimulatedAmounts)
plot(acomp(sa.lognormals))

---

ccompgoftest

Compositional Goodness of fit test

Description

Goodness of fit tests for count compositional data.

Usage

PoissonGOF.test(x, lambda=mean(x), R=999, estimated=missing(lambda))
ccompPoissonGOF.test(x, simulate.p.value=TRUE, R=1999)
ccompMultinomialGOF.test(x, simulate.p.value=TRUE, R=1999)
Arguments

- **x**: a dataset of integer numbers (PoissonGOF) or count compositions (compPoissonGOF)
- **lambda**: the expected value to check against
- **R**: The number of replicates to compute the distribution of the test statistic
- **estimated**: states whether the lambda parameter should be considered as estimated for the computation of the p-value.
- **simulate.p.value**: should all p-values be inferred by simulation.

Details

The compositional goodness of fit testing problem is essentially a multivariate goodness of fit test. However there is a lack of standardized multivariate goodness of fit tests in R. Some can be found in the energy-package.

In principle there is only one test behind the Goodness of fit tests provided here, a two sample test with test statistic.

$$\frac{\sum_{ij} k(x_i, y_i)}{\sqrt{\sum_{ij} k(x_i, x_i) \sum_{ij} k(y_i, y_i)}}$$

The idea behind that statistic is to measure the cos of an angle between the distributions in a scalar product given by

$$(X, Y) = E[k(X, Y)] = E[\int K(x - X)K(x - Y)dx]$$

where k and K are Gaussian kernels with different spread. The bandwith is actually the standarddeviation of k.

The other goodness of fit tests against a specific distribution are based on estimating the parameters of the distribution, simulating a large dataset of that distribution and apply the two sample goodness of fit test.

Value

A classical "htest" object

- **data.name**: The name of the dataset as specified
- **method**: a name for the test used
- **alternative**: an empty string
- **replicates**: a dataset of p-value distributions under the Null-Hypothesis got from nonparametric bootstrap
- **p.value**: The p.value computed for this test

Missing Policy

Up to now the tests can not handle missings.
Author(s)


References


See Also

fitDirichlet, rDirichlet, runif.acomp, rnorm.acomp.

Examples

## Not run:
x <- runif.acomp(100,4)
y <- runif.acomp(100,4)

erg <- acompGOF.test(x,y)
#continue
erg
unclass(erg)

## Examples:
x <- runif.acomp(100,4)
y <- runif.acomp(100,4)

dd <- replicate(1000, acompGOF.test(runif.acomp(100,4), runif.acomp(100,4))$p.value)
hist(dd)

dd <- replicate(1000, acompGOF.test(runif.acomp(20,4), runif.acomp(100,4))$p.value)
hist(dd)

dd <- replicate(1000, acompGOF.test(runif.acomp(10,4), runif.acomp(100,4))$p.value)

hist(dd)

dd <- replicate(1000, acompGOF.test(runif.acomp(10,4), runif.acomp(400,4))$p.value)

hist(dd)

dd <- replicate(1000, acompGOF.test(runif.acomp(400,4), runif.acomp(10,4), bandwidth=4)$p.value)

hist(dd)

dd <- replicate(1000, acompGOF.test(runif.acomp(20,4), runif.acomp(100,4)+acomp(c(1,2,3,1)))$p.value)

hist(dd)

x <- runif.acomp(100,4)

acompUniformityGOF.test(x)

dd <- replicate(1000, acompUniformityGOF.test(runif.acomp(10,4))$p.value)
cdt

hist(dd)

## End(Not run)

### cdt

**Centered default transform**

**Description**

Compute the centered default transform of a (data set of) compositions or amounts (or its inverse).

**Usage**

```
cdt(x,...)
## Default S3 method:
cdt( x ,...)
## S3 method for class 'acomp'
cdt( x ,...)
## S3 method for class 'rcomp'
cdt( x ,...)
## S3 method for class 'aplus'
cdt( x ,...)
## S3 method for class 'rplus'
cdt( x ,...)
## S3 method for class 'rmult'
cdt( x ,...)
## S3 method for class 'ccomp'
cdt( x ,...)
## S3 method for class 'factor'
cdt( x ,...)
cdtInv(x,orig,...)
## Default S3 method:
cdtInv( x ,orig,...)
## S3 method for class 'acomp'
cdtInv( x ,orig,...)
## S3 method for class 'rcomp'
cdtInv( x ,orig,...)
## S3 method for class 'aplus'
cdtInv( x ,orig,...)
## S3 method for class 'rplus'
cdtInv( x ,orig,...)
## S3 method for class 'rmult'
cdtInv( x ,orig,...)
## S3 method for class 'ccomp'
cdtInv( x ,orig,...)
## S3 method for class 'factor'
```
cdtInv( x ,orig,...)

Arguments

x  a classed (matrix of) amount or composition, to be transformed with its centered
default transform, or its inverse
...
... generic arguments past to underlying functions.
orig  a compositional object which should be mimicked by the inverse transformation.
It is used to determine the backtransform to be used and eventually to reconstruct
the names of the parts. It is the generic argument. Typically this argument is the
data set that has be transformed in the first place.

Details

The general idea of this package is to analyse the same data with different geometric concepts,
in a fashion as similar as possible. For each of the four concepts there exists a unique transform
expressing the geometry in a linear subspace, keeping the relation to the variables. This unique
transformation is computed by cdt. For acomp the transform is clr, for rcomp it is cpt, for aplus
it is ilt, and for rplus it is iit. Each component of the result is identified with a unit vector in the
direction of the corresponding component of the original composition or amount. Keep in mind that
the transform is not necessarily surjective and thus variances in the image space might be singular.

Value

A corresponding matrix or vector containing the transforms.

Author(s)


References

van den Boogaart, K.G. and R. Tolosana-Delgado (2008) "compositions": a unified R package to an-

See Also

idt, clr, cpt, ilt, iit

Examples

## Not run:
# the cdt is defined by
cdt <- function(x) UseMethod("cdt",x)
cdt.default <- function(x) x
cdt.acomp <- clr
cdt.rcomp <- cpt
cdt.aplus <- ilt
cdt.rplus <- iit
```r
## End(Not run)
x <- acomp(1:5)
(ds <- cdt(x))
cdtInv(ds,x)
(ds <- cdt(rcomp(1:5)))
cdtInv(ds,rcomp(x))
data(Hydrochem)
x = Hydrochem[,c("Na","K","Mg","Ca")]
y = acomp(x)
z = cdt(y)
y2 = cdtInv(z,y)
par(mfrow=c(2,2))
for(i in 1:4){plot(y[,i],y2[,i])}
```

### ClamEast

**Color-size compositions of 20 clam colonies from East Bay**

#### Description

From East Bay, 20 clam colonies were randomly selected and from each a sample of clams were taken. Each sample was sieved into three size ranges, large, medium, and small. Then each size range was sorted by the shale colour, dark or light.

#### Usage

```r
data(ClamEast)
```

#### Details

The data consist of 20 cases and $2 \times 3$ variables denoted:

- dl: portion of dark large clams,
- dm: portion of dark medium clams,
- ds: portion of dark small clams,
- ll: portion of light large clams,
- lm: portion of light medium clams,
- ls: portion of light small clams.

All 6-part compositions sum to one, except for rounding errors.

#### Note

Courtesy of J. Aitchison
**Source**

Aitchison: CODA microcomputer statistical package, 1986, the file name CLAMEAST.DAT, here included under the GNU Public Library Licence Version 2 or newer.

**References**


---

**ClamWest**

*Color-size compositions of 20 clam colonies from West Bay*

**Description**

From West Bay, 20 clam colonies were randomly selected, and from each a sample of clams were taken. Each sample was sieved into three size ranges, large, medium, and small. Then each size range was sorted by the shale colour, dark or light.

**Usage**

data(ClamWest)

**Details**

The data consist of 20 cases and $2 \times 3$ variables denoted:

- dl portion of dark large clams,
- dm portion of dark medium clams,
- ds portion of dark small clams,
- ll portion of light large clams,
- lm portion of light medium clams,
- ls portion of light small clams.

All 6-part compositions sum up to one.

**Note**

Courtesy of J. Aitchison

**Source**

Aitchison: CODA microcomputer statistical package, 1986, the file name CLAMWEST.DAT, here included under the GNU Public Library Licence Version 2 or newer.

**References**

Closure of a composition

Description

Closes compositions to sum up to one (or an optional total), by dividing each part by the sum.

Usage

clo( X, parts=1:NCOL(oneOrDataset(X)), total=1,
    detectionlimit=attr(X,"detectionlimit"),
    BDL=NULL, MAR=NULL, MNAR=NULL, SZ=NULL,
    storelimit=!is.null(attr(X,"detectionlimit"))
)

Arguments

X composition or dataset of compositions
parts vector containing the indices xor names of the columns to be used
total the total amount to which the compositions should be closed; either a single number, or a numeric vector of length gsi.getN(X) specifying a different total for each compositional vector in the dataset.
detectionlimit a number, vector or matrix of positive numbers giving the detection limit of all values, all variables, or each value
BDL the code for values below detection limit in X
SZ the code for structural zeroes in X
MAR the code for values missed at random in X
MNAR the code for values missed not at random in X
storelimit a boolean indicating whether to store the detection limit as an attribute in the data. It defaults to FALSE if the detection limit is not already stored in the dataset. The attribute is only needed for very advanced analysis. Most times, this will not be used.

Details

The closure operation is given by

\[ clo(x) := \left( x_i/\sum_{j=1}^{D} x_j \right) \]

clo generates a composition without assigning one of the compositional classes acomp or rcomp. Note that after computing the closed-to-one version, obtaining a version closed to any other value is done by simple multiplication.
Value

A composition or a data matrix of compositions, maybe without compositional class. The individual compositions are forced to sum to 1 (or to the optionally-specified total). The result should have the same shape as the input (vector, row, matrix).

Missing Policy

How missing values are coded in the output always follows the general rules described in *compositions.missing*. The BDL values are accordingly scaled during the scaling operations but not taken into account for the calculation of the total sum.

Note

clo can be used to unclass compositions.

Author(s)


References


See Also

clo, acomp, rcomp

Examples

```r
(tmp <- clo(c(1,2,3)))
clo(tmp, total=100)
data(Hydrochem)
plot( clo(Hydrochem, 8:9) ) # Giving points on a line
```

---

**clr**

*Centered log ratio transform*

**Description**

Compute the centered log ratio transform of a (dataset of) composition(s) and its inverse.

**Usage**

```r
clr( x, ... )
clrInv( z, ... )
```
**clr**

**Arguments**
- **x**: a composition or a data matrix of compositions, not necessarily closed
- **z**: the clr-transform of a composition or a data matrix of clr-transforms of compositions, not necessarily centered (i.e. summing up to zero)
- **...**: for generic use only

**Details**

The clr-transform maps a composition in the D-part Aitchison-simplex isometrically to a D-dimensional euclidian vector subspace: consequently, the transformation is not injective. Thus resulting covariance matrices are always singular.

The data can then be analysed in this transformation by all classical multivariate analysis tools not relying on a full rank of the covariance. See **ilr** and **alr** for alternatives. The interpretation of the results is relatively easy since the relation between each original part and a transformed variable is preserved.

The centered logratio transform is given by

\[ clr(x) := \left( \ln x_i - \frac{1}{D} \sum_{j=1}^{D} \ln x_j \right)_i \]

The image of the clr is a vector with entries summing to 0. This hyperplane is also called the clr-plane.

**Value**

- **clr** gives the centered log ratio transform, **clrInv** gives closed compositions with the given clr-transform

**Author(s)**


**References**


**See Also**

- **ilr**, **alr**, **apt**

**Examples**

```r
tmp <- clr(c(1,2,3))
clrInv(tmp)
clrInv(tmp) - clo(c(1,2,3)) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(clr(cdata),pch=".")
```
**clr2ilr**

*Convert between clr and ilr, and between cpt and ipt.*

**Description**

Compute the centered log ratio transform of a (dataset of) isometric log-ratio transform(s) and its inverse. Equivalently, compute centered and isometric planar transforms from each other. Acts in vectors and in bilinear forms.

**Usage**

```r
clr2ilr(x, V = ilrBase(x))
ilr2clr(z, V = ilrBase(z = z), x = NULL)
clrvar2ilr(varx, V = ilrBase(D = ncol(varx)))
ilrvar2clr(varz, V = ilrBase(D = ncol(varz) + 1), x = NULL)
```

**Arguments**

- `x` the clr/cpt-transform of composition(s) (in the ilr2-routines provided only to give column names.)
- `z` the ilr/ipt-transform of composition(s)
- `varx` variance or covariance matrix of clr/cpt-transformed compositions
- `varz` variance or covariance matrix of ilr/ipt-transformed compositions
- `V` a matrix with columns giving the chosen basis of the clr-plane

**Details**

These functions perform a matrix multiplication with `V` in an appropriate way.

**Value**

- `clr2ilr` gives the ilr/ipt transform of the same composition(s).
- `ilr2clr` gives the clr/cpt transform of the same composition(s).
- `clrvar2ilr` gives the variance-/covariance-matrix of the ilr/ipt transform of the same compositional data set.
- `ilrvar2clr` gives the variance-/covariance-matrix of the clr/cpt transform of the same compositional data set.

**Author(s)**

ClusterFinder1

Heuristics to find subpopulations of outliers

Description

The ClusterFinder is a heuristic to find subpopulations of outliers essentially by looking for secondary modes in a density estimate.

Usage

ClusterFinder1(X,...)
## S3 method for class 'acomp'
ClusterFinder1(X,...,sigma=0.3,radius=1,asig=1,minGrp=3, robust=TRUE)

Arguments

X the dataset to be clustered
...
Further arguments to MahalanobisDist(X,...,robust=robust, pairwise=TRUE)
sigma numeric: The Bandwidth of the density estimation kernel in a robustly Mahalanobis transformed space. (i.e. in the transform, where the main group has unit variance)
radius The minimum size of a cluster in a robustly Mahalanobis transformed space. (i.e. in the transform, where the main group has unit variance)

References

Aitchison, J, C. Barcel'o-Vidal, J.J. Egozcue, V. Pawlowsky-Glahn (2002) A concise guide to the algebraic geometric structure of the simplex, the sample space for compositional data analysis, Terra Nostra, Schriften der Alfred Wegener-Stiftung, 03/2003

See Also

ilr, ipt

Examples

data(SimulatedAmounts)
ilrInv(clr2ilr(clr(sa.lognormals)))-clo(sa.lognormals)
clrInv(ilr2clr(ilr(sa.lognormals)))-clo(sa.lognormals)
ilrvar2clr(var(ilr(sa.lognormals)))-var(clr(sa.lognormals))
clrvar2ilr(var(cpt(sa.lognormals)))-var(ipt(sa.lognormals))
asig a scaling factor for the geometry of the robustly Mahalanobis transformed space when computing the likelihood of an observation to belong to group (under a Gaussian assumption). Higher values

minGrp the minimum size of group to be used. Smaller groups are treated as single outliers

robust A robustness description for estimating the variance of the main group. FALSE is probably not a usefull value. However later other robustness techniques than mcd might be usefull. TRUE just picks the default method of the package.

Details

See outliersInCompositions for a comprehensive introduction into the outlier treatment in compositions.

The ClusterFinder is labeled with a number to make clear that this is just an implementation of some heuristic and not based on some eternal truth. Other might give better Clusterfinders.

Unlike other Clustering Algorithms the basic model of this algorithm assumes that there is one dominating subpopulation and an unkown number of smaller subpopulations with a similar covariance structure but a different mean. The algorithm thus first estimates the covariance structure of the main population by a robust location scale estimator. Then it uses a simplified (Gaussian) kernel density estimator to find nonrandom secondary modes. The it tries to assign the different observations according to discrimination analysis model to the different modes. Groups under a given size are considered as single outliers forming a seperate group. In this way the number of clusters is kept low even if there are many erratic measurements in the dataset.

The main use of the clusters is descriptive plotting. The advantage of these cluster against other cluster techniques like k-mean or hclust is that it does not tear appart the central mass of the data, as these methods do to make the clusters as compact as possible.

Value

A list

types a factor representing the group assignments, when the small groups are ignored

typesTbl a table giving the number of members in each of these groups

groups a factor representing the found group assignments

isMax a logical vector indicating for each observation,whether it represent a local maximum in the density estimate.

prob the inferred probability to belong to the different groups given as an acomp composition.

rmembers a tabel giving the number of members of each group

density the density estimated in each observation location

likeli The inferred likelihood see this observation, for each of the groups

Author(s)

CoDaDendrogram

See Also

hclust, kmeans

Examples

data(SimulatedAmounts)
cl <- ClusterFinder1(sa.outliers5,sigma=0.4,radius=1)
plot(sa.outliers5,col=as.numeric(cl$types),pch=as.numeric(cl$types))
legend(1,1,legend=levels(cl$types),xjust=1,col=1:length(levels(cl$types)),
pch=1:length(levels(cl$types)))

CoDaDendrogram

Dendrogram representation of acomp or rcomp objects

Description

Function for plotting CoDa-dendrograms of acomp or rcomp objects.

Usage

CoDaDendrogram(X, V = NULL, expr=NULL, mergetree = NULL, signary = NULL,
    range = c(-4,4), ..., xlim = NULL, ylim = NULL, yaxt = NULL, box.pos = 0,
    box.space = 0.25, col.tree = "black", lty.tree = 1, lwd.tree = 1,
    col.leaf = "black", lty.leaf = 1, lwd.leaf = 1, add = FALSE,border=NULL,
    type = "boxplot")

Arguments

X          data set to plot (an rcomp or acomp object)
V          basis to use, described as an ilr matrix
expr       a formula describing the partition basis, as with balanceBase
mergetree  basis to use, described as a merging tree (as in hclust)
signary    basis to use, described as a sign matrix (as in the example below)
range      minimum and maximum value for all coordinates (horizontal axes)
...        further parameters to pass to any function, be it a plotting function or one related
to the "type" parameter below; likely to produce lots of warnings
xlim       minimum and maximum values for the horizontal direction of the plot (related
to number of parts)
ylim       minimum and maximum values for the vertical direction of the plot (related to
variance of coordinates)
yaxt       axis type for the vertical direction of the plot (see par)
box.pos  if type="boxplot", this is the relative position of the box in the vertical direction: 0 means centered on the axis, -1 aligned below the axis and +1 aligned above the axis

box.space  if type="boxplot", size of the box in the vertical direction as a portion of the minimal variance of the coordinates

col.tree  color for the horizontal axes

lty.tree  line type for the horizontal axes

lwd.tree  line width for the horizontal axes

col.leaf  color for the vertical connections between an axis and a part (leaf)

lty.leaf  line type for the leaves

lwd.leaf  line width for the leaves

add  should a new plot be triggered, or is the material to be added to an existing CoDa-dendrogram?

border  the color for drawing the rectangles

type  what to represent? one of "boxplot", "density", "histogram", "lines", "nothing" or "points", or an univocal abbreviation

Details

The object and an isometric basis are represented in a CoDa-dendrogram, as defined by Egozcue and Pawlowsky-Glahn (2005). This is a representation of the following elements:

- An hierarchical partition (which can be specified either through an ilrBase matrix (see ilrBase), a merging tree structure (see hclust) or a signary matrix (see gsi.merge2signary))
- The sample mean of each coordinate of the ilr basis associated to that partition
- The sample variance of each coordinate of the ilr basis associated to that partition
- Optionally (potentially!), any graphical representation of each coordinate, as long as this representation is suitable for a univariate data set (box-plot, histogram, dispersion and kernel density are programmed or intended to, but any other may be added with little work).

Each coordinate is represented in a horizontal axis, which limits correspond to the values given in the parameter range. The vertical bar going up from each one of these coordinate axes represent the variance of that specific coordinate, and the contact point the coordinate mean. Note that to be able to represent an initial dendrogram, the first call to this function must be given a full data set, as means and variances must be computed. This information is afterwards stored in a global list, to add any sort of new material to all coordinates.

The default option is type="boxplot", which produces a box-plot for each coordinate, customizable using box.pos and box.space, as well as typical par parameters (col, border, lty, lwd, etc.). To obtain only the first three aspects, the function must be called with type="lines". As extensions, one might represent a single datum/few data (e.g., a mean or a random subsample of the data set) calling the function with add=TRUE and type="points". Other options (calling functions histogram or density, and admitting their parameters) will be also soon available.

Note that the original coda-dendrogram as defined by Egozcue and Pawlowsky-Glahn (2005) works with acomp objects and ilr bases. Functionality is extended to rcomp objects using calls to idt.
Author(s)

References


See Also
ilrBase, balanceBase, rcomp, acomp.

Examples

# first example: take the data set from the example, select only
# compositional parts
data(Hydrochem)
x = acomp(Hydrochem[, -c(1:5)])
gr = Hydrochem[, 4] # river groups (useful afterwards)
# use an ilr basis coming from a clustering of parts
dd = dist(t(clr(x)))
hcl = hclust(dd, method = "ward")
plot(hcl)
mergetree = hcl$merge
CoDaDendrogram(X = acomp(x), mergetree = mergetree, col = "red", range = c(-8, 8), box.space = 1)
# add the mean of each river
color = c("green3", "red", "blue", "darkviolet")
aux = clrInv(t(sapply(split(clr(x), gr), mean)))
CoDaDendrogram(X = aux, add = TRUE, col = color, type = "points",pch = 4)

# second example: box-plots by rivers (filled)
CoDaDendrogram(X = acomp(x), mergetree = mergetree, col = "black", range = c(-8, 8), type = "l")
xsplit = split(x, gr)
for(i in 1:4){
  CoDaDendrogram(X = acomp(xsplits[[i]]), col = color[i], type = "box", box.pos = i - 2.5, box.space = 0.5, add = TRUE)
}

# third example: fewer parts, partition defined by a signary, and empty box-plots
x = acomp(Hydrochem[, c("Na", "K", "Mg", "Ca", "Sr", "Ba", "NH4")])
signary = t(matrix(  
c(1, 1, 1, 1, 1, -1, 1, 1, -1, 1, 1, 1),
1, -1, -1, -1, -1, 1, 0, 0, 0, 0, 0, 0),
0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0), ncol = 7, nrow = 6, byrow = TRUE))
coloredBiplot

A biplot providing somewhat easier access to details of the plot.

Description

This function generates a simple biplot out of various sources and allows to give color and symbol to the x-objects individually.

Usage

```r
## Default S3 method:
coloredBiplot(x, y, var.axes = TRUE, col,
               cex = rep(par("cex"), 2), xlabs = NULL, ylabs = NULL, expand=1,
               xlim = NULL, ylim = NULL, arrow.len = 0.1, main = NULL, sub = NULL,
               xlab = NULL, ylab = NULL, xlabs.col = NULL, xlabs.bg = NULL,
               xlabs.pc=NULL, ...)
## S3 method for class 'princomp'
coloredBiplot(x, choices = 1:2, scale = 1,
               pc.biplot=FALSE, ...)
## S3 method for class 'prcomp'
coloredBiplot(x, choices = 1:2, scale = 1,
               pc.biplot=FALSE, ...)
```

Arguments

- `x`: a representation of the co-information to be plotted, given by a result of princomp or prcomp; or the first set of coordinates to be plotted
- `y`: optional, the second set of coordinates to be plotted
- `var.axes`: if 'TRUE' the second set of points have arrows representing them as (unscaled) axes
- `col`: one color (to be used for the y set) or a vector of two colors (to be used for x and y sets respectively, if xlabs.col is NULL)
- `cex`: the usual cex parameter for plotting; can be a length-2 vector to format differently x and y labels/symbols
coloredBiplot

xbibs  names to write for the points of the first set
ylbs  names to write for the points of the second set
expand  expansion factor to apply when plotting the second set of points relative to the first. This can be used to tweak the scaling of the two sets to a physically comparable scale
xlim  horizontal axis limits
ylim  vertical axis limits
arrow.len  length of the arrow heads on the axes plotted if ‘var.axes’ is true. The arrow head can be suppressed by ‘arrow.len=0’
main  main title
sub  subtitle
xlab  horizontal axis title
ylab  vertical axis title
xbibs.col  the color(s) to draw the points of the first set, if xbibs is null
xbibs.bg  the filling color(s) to draw the points of the first set, if xbibs is null and xbibs.pc is between 21 and 25.
xbibs.pc  the plotting character(s) for the first set, if xbibs is null
scale  the way to distribute the singular values on the right or left singular vectors for princomp and prcomp objects (see biplot)
choices  the components to be plotted (see biplot)
pc.biplot  should be scaled by sqrt(nrow(X))? (see biplot)
...  further parameters for plot

Details
The functions is provided for convenience.

Value
The function is called only for the side effect of plotting. It is a modification of the standard R routine ‘biplot’.

Author(s)

See Also
biplot, plot.acomp

Examples

data(SimulatedAmounts)
coloredBiplot(x=princomp(acomp(sa.outliers5)),pc.biplot=FALSE,
    xlbs.pc=c(1,2,3), xlbs.col=2:4, col="black")
colorsForOutliers  
Create a color/char palette or for groups of outliers

Description
Convenience Functions to generate meaningful color palettes for factors representing different types of outliers.

Usage

```r
colorsForOutliers1(outfac, family=rainbow,
                   extreme="cyan", outlier="red", ok="gray40", unknown="blue")
colorsForOutliers2(outfac, use=whichBits(gsi.orSum(levels(outfac))[, 1:7], "=")
codes=c(2^outer(rep(2, 7), c(24, 16, 8)), ok="yellow")
pchForOutliers1(outfac, ok=\'.', outlier='\004', extreme='\003', unknown='\004',
                  other=c('\001', '\002', '\026', '\027', '\010', '\011', '\012', '\013', '\014', '\015',
                      '\016', strsplit("abcdefghijklmnopqrstuvwxyzABCDEFGHIJKLMNOPQRSTUVWXYZ", ","))[[1]])
```

Arguments

- `outfac` a factor given by an OutlierClassifier (e.g. `OutlierClassifier1`). `colorsForOutliers1` is used for the types "best","type","outlier","grade". `colorsForOutliers2` is used for type all.
- `family` a function generating a color palette from a number of colors requested.
- `extreme` The color/char for extreme but not definitively outlying observations.
- `outlier` The color/char for detected outliers.
- `unknown` The color/char for observation with unclear classification.
- `other` The character codes for other outlier classes.
- `ok` The color/char for nonoutlying usual observations.
- `use` a numerical vector giving the indices of the bits of the output to be represented. The sequence of the bits determines how each bit is represented.
- `codes` The color influences to be used for each bit.
- `...` further codings for other factor levels

Details
This functions are provided for convenience to quickly generate a palette of reasonable colors or plotting chars for groups of outliers classified by OutlierClassifier1.

Value

a character vector of colors or a numeric vector of plot chars.
CompLinModCoReg

Description

Creates a Variogram model according to the linear model of spatial corregionalisation for a compositional geostatistical analysis.

Usage

CompLinModCoReg(formula, comp, D=ncol(comp), envir=environment(formula))

Arguments

formula       A formula without left side providing a formal description of a variogram model.
comp          a compositional dataset, needed to provide the frame size
D             The dimension of the multivariate dataset
envir         The environment in which formula should be interpreted.

Details

The linear model of coregionalisation says uses the fact, that sums of valid variogram models are valid variograms and that scalar variograms multiplied with a positive definite matrix are valid vector variograms. The function computes such a variogram function from a formal description. The formula is a sum. Each summand is either a product of a matrix description and a scalar variogram description or only a scalar variogram description. Scalar variogram descriptions are either formal function calls to

Examples

```r
## Not run:
data(SimulatedAmounts)
data5 <- acomp(sa.outliers5)
olc <- OutlierClassifier1(data5)
plot(data5,col=colorsForOutliers1(olc)[olc])
olc <- OutlierClassifier1(data5,type="all")
plot(data5,col=colorsForOutliers2(olc)[olc])
## End(Not run)
```
• sph(range) for spherical variogram with range range
• exp(range) for an exponential variogram with range range
• gauss(range) for a Gaussian variogram with range range
• gauss(range) for a cardinal sine variogram with (non-effective) range range
• pow(range) for an power variogram with range range
• lin(unit) linear variogram 1 at unit.
• nugget() for adding a nugget effect.

alternatively it can be any expression, which will be evaluated in envir and should depende on a
dataset of distance vectors.
The matrix description always comes first. It can be R1 for a rank 1 matrix PSD for a Positive
Semidefinite matrix or \( S \) for a scalar Sill factor to be multiplied with identity. Or any other con-
struct evaluating to a matrix or a function of some parameters with default values, that if called is
evaluated to a positive semidefinite matrix. R1 and PSD can also be written as calls with providing a
vector or respectively a matrix providing the parameter.
The variogram is created with default parameter values. The parameters can later be modified by
modifying the default parameter with assignments like formal)(vg)$sPSD1 = parameterPosdefMat(4*diag(5)).
We would anyway expect you to fit the model to the data by a command like
vgmFit(logratioVariogram(...),CompLinModCoReg(...)).

Value
A variogram function.

Author(s)

References
What to cite??

See Also
vgram2lrvgram, CompLinModCoReg, vgmFit

Examples
## Not run:
data(juraset)
X <- with(juraset,cbind(X,Y))
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
CompLinModCoReg(~nugget()+sph(0.5)+R1*exp(0.7),comp)
CompLinModCoReg(~nugget()+R1*sph(0.5)+R1*exp(0.7)+(0.3*diag(5))*gauss(0.3),comp)
CompLinModCoReg(~nugget()+R1*sph(0.5)+R1(c(1,2,3,4,5))*exp(0.7),comp)

## End(Not run)
compOKriging

Compositional Ordinary Kriging

Description

Geostatistical prediction for compositional data with missing values.

Usage

compOKriging(comp, X, Xnew, vg, err = FALSE)

Arguments

- `comp`: an acomp compositional dataset
- `X`: A dataset of locations
- `Xnew`: The locations, where a geostatistical prediction should be computed.
- `vg`: A compositional variogram function.
- `err`: boolean: If true kriging errors are computed additionally.

Details

The function performs multivariate ordinary kriging of compositions based on transformes adapted to the missings in every case. The variogram is assumed to be a clr variogram.

Value

A list of class "logratioVariogram"

- `X`: The new locations as given by Xnew
- `Z`: The predicted values as acomp compositions.
- `err`: An ncol(Z)xDxD array with the clr kriging errors.

Author(s)


References


Tolosana (2008) ...

Tolosana, van den Boogaart, Pawlowsky-Glahn (2009) Estimating and modeling variograms of compositional data with occasional missing variables in R, StatGis09
### See Also

vgram2lrvgram, CompLinModCoReg, vgmFit

### Examples

```r
## Not run:
# Load data
data(juraset)
X <- with(juraset,cbind(X,Y))
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
lrv <- logratioVariogram(comp,X,maxdist=1,nbins=10)
plot(lrv)

# Fit a variogram model
vgModel <- CompLinModCoReg(~nugget()+sph(0.5)+R1*exp(0.7),comp)
fit <- vgmFit2lrv(lrv,vgModel)
fit
plot(lrv,lrvg=vgram2lrvgram(fit$vg))

# Define A grid
x <- (0:30/30)*6
y <- (0:30/30)*6
Xnew <- cbind(rep(x,length(y)),rep(y,each=length(x)))

# Kriging
erg <- compOKriging(comp,X,Xnew,fit$vg,err=FALSE)
par(mar=c(0,0,1,0))
pairwisePlot(erg$Z,panel=function(a,b,xlab,ylab) {image(x,y,
structure(log(a/b),dim=c(length(x),length(y))),
main=paste("log(",xlab,"/",ylab,")",sep=""));points(X,pch="."))})

# Check interpolation properties
ergR <- compOKriging(comp,X,X,fit$vg,err=FALSE)
pairwisePlot(ilr(comp),ilr(ergR$Z))
 ergR <- compOKriging(comp,X,X*1E-7,fit$vg,err=FALSE)
pairwisePlot(ilr(comp),ilr(ergR$Z))
 ergR <- compOKriging(comp,X,X[rev(1:31),],fit$vg,err=FALSE)
pairwisePlot(ilr(comp)[rev(1:31),],ilr(ergR$Z))

## End(Not run)
```

---

**ConfRadius**

*Helper to compute confidence ellipsoids*

### Description

Computes the quantile of the Mahalanobis distance needed to draw confidence ellipsoids.
cor.acomp

Usage

ConfRadius(model, prob=1-alpha, alpha)

Arguments

model A multivariate linear model
prob The confidence probability
alpha The alpha error allowed, i.e. the complement of the confidence probability

Details

Calculates the radius to be used in confidence ellipses for the parameters based on the Hottelings $T^2$ distribution.

Value

a scalar

Author(s)


See Also

lm, mvar, AIC

Examples

data(SimulatedAmounts)
model <- lm(ilr(sa.groups)~sa.groups.area)
cf = coef(model)
plot(ilrInv(cf, x=sa.groups))
for(i in 1:nrow(cf)){
  vr = vcovAcomp(model)[,,i,i]
  vr = ilrvar2clr(vr)
  ellipses(ilrInv(cf[i,]), vr, r=ConfRadius(model, alpha=0.05) )
}

---

cor.acomp Correlations of amounts and compositions

Description

Computes the correlation matrix in the various approaches of compositional and amount data analysis.
Usage

```r
cor(x,y=NULL,...)
## Default S3 method:
cor(x, y=NULL, use="everything",
    method=c("pearson", "kendall", "spearman"),...)
## S3 method for class 'acomp'
cor(x,y=NULL,...,robust=getOption("robust"))
## S3 method for class 'rcomp'
cor(x,y=NULL,...,robust=getOption("robust"))
## S3 method for class 'aplus'
cor(x,y=NULL,...,robust=getOption("robust"))
## S3 method for class 'rplus'
cor(x,y=NULL,...,robust=getOption("robust"))
## S3 method for class 'rmult'
cor(x,y=NULL,...,robust=getOption("robust"))
```

Arguments

- **x**: a data set, eventually of amounts or compositions
- **y**: a second data set, eventually of amounts or compositions
- **use**: see `cor`
- **method**: see `cor`
- **...**: further arguments to `cor` e.g. `use`
- **robust**: A description of a robust estimator. FALSE for the classical estimators. See `mean.acomp` for further details.

Details

The correlation matrix does not make much sense for compositions.

In R versions older than v2.0.0, `cor` was defined in package “base” instead of in “stats”. This might produce some misfunction.

Value

The correlation matrix.

Author(s)


See Also

`var.acomp`
Examples

```r
data(SimulatedAmounts)
meanCol(sa.lognormals)
cor(acomp(sa.lognormals5[,1:3]), acomp(sa.lognormals5[,4:5]))
cor(rcomp(sa.lognormals5[,1:3]), rcomp(sa.lognormals5[,4:5]))
cor(aplus(sa.lognormals5[,1:3]), aplus(sa.lognormals5[,4:5]))
cor(rplus(sa.lognormals5[,1:3]), rplus(sa.lognormals5[,4:5]))
cor(acomp(sa.lognormals5[,1:3]), aplus(sa.lognormals5[,4:5]))
```

---

**Coxite**

*Compositions, depths and porosities of 25 specimens of coxite*

---

**Description**

A mineral compositions of 25 rock specimens of coxite type. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite, the depth of location, and porosity.

**Usage**

```r
data(Coxite)
```

**Details**

A mineral compositions of 25 rock specimens of coxite type. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite, the recorded depth of location of each specimen, and porosity. Porosity is the percentage of void space that the specimen contains. We abbreviate the minerals names to A, B, C, D, E.

All row percentage sums to 100.

**Note**

Courtesy of J. Aitchison

**Source**

Aitchison: CODA microcomputer statistical package, 1986, the file name BOXITE.DAT, here included under the GNU Public Library Licence Version 2 or newer.

**References**

**cpt**  
*Centered planar transform*

**Description**

Compute the centered planar transform of a (dataset of) compositions and its inverse.

**Usage**

```r
cpt( x,... )
cptInv( z,... )
```

**Arguments**

- `x`: a composition or a data.matrix of compositions, not necessarily closed
- `z`: the cpt-transform of a composition or a data matrix of cpt-transforms of compositions. It is checked that the `z` sum up to 0.
- `...`: generic arguments. not used.

**Details**

The cpt-transform maps a composition in the D-part real-simplex isometrically to a D-1 dimensional euclidian vector space, identified with a plane parallel to the simplex but passing through the origin. However the transformation is not injective and does not even reach the whole plane. Thus resulting covariance matrices are always singular.

The data can then be analysed in this transformed space by all classical multivariate analysis tools not relying on a full rank of the covariance matrix. See `ipt` and `apt` for alternatives. The interpretation of the results is relatively easy since the relation of each transformed component to the original parts is preserved.

The centered planar transform is given by

\[
\text{cpt}(x)_i := \text{clo}(x)_i - \frac{1}{D}
\]

**Value**

- `cpt` gives the centered planar transform, `cptInv` gives closed compositions with the given cpt-transforms.

**Author(s)**


**References**


**See Also**

clr, apt, ipt

**Examples**

```r
(tmp <- cpt(c(1,2,3)))
cptInv(tmp)
cptInv(tmp) - clo(c(1,2,3)) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(cpt(cdata),pch=".")
```

---

**DiagnosticProb**

*Diagnostic probabilities*

**Description**

Data record the probabilities assigned by subjective diagnostic of 15 clinicians and 15 statisticians.

**Usage**

```r
data(DiagnosticProb)
```

**Details**

The data consist of 30 cases: 15 diagnostics probabilities assigned by clinicians, 15 diagnostics probabilities assigned by statisticians, and 4 variables: probabilities A, B, and C, and type i.e. 1 for clinicians, 2 for statisticians.

In the study of subjective performance in inferential task the subject is faced with the finite set of mutually exclusive and exhaustive hypothesis, and the basis of specific information presented to him/her is required to divide the available unit of probability among these probabilities. In this study the task is presented as a problem of differential diagnosis of three mutually exclusive and exhaustive diseases of students, known under the generic title of 'newmath syndrome',

- A - algebritis,
- B - bilateral paralexia,
- C - calculus deficiency.

The subject, playing the role of diagnostician, is informed that the three diseases types are equally common and is shown the results of 10 diagnostic tests on 60 previous cases of known diagnosis, 20 of each type. The subject is then shown the results of the 10 tests for a new undiagnosed cases and asked to assign diagnostic probabilities to the three possible disease types.
Data record the subjective assessments of 15 clinicians and 15 statisticians for the same case. For this case the objective diagnosis probabilities are known to be $(.08, .05, .87).$ All row probabilities sum to 1, except for some rounding errors.

**Note**

Courtesy of J. Aitchison

**Source**

Aitchison: CODA microcomputer statistical package, 1986, the file name DIAGPROB.DAT, here included under the GNU Public Library Licence Version 2 or newer.

**References**


---

**dist**

*Distances in variouse approaches*

**Description**

Calculates a distance matrix from a data set.

**Usage**

```
dist(x,...)
```

```
## Default S3 method:
dist(x,...)
```

**Arguments**

- `x` a dataset
- `...` further arguments to `dist`

**Details**

The distance is computed based on `cdt`

**Value**

a distance matrix

**Author(s)**

ellipses

See Also
aplus

Examples

```r
data(SimulatedAmounts)
phc <- function(d) { plot(hclust(d))
  phc(dist(iris[,1:4]))
  phc(dist(acomp(sa.lognormals),method="manhattan"))
  phc(dist(rcomp(sa.lognormals)))
  phc(dist(aplus(sa.lognormals)))
  phc(dist(rplus(sa.lognormals)))
```

## S3 method for class 'acomp'
ellipses(mean,...)

## S3 method for class 'rcomp'
ellipses(mean,var,r=1,...,steps=72,thinRatio=NULL,aspanel=FALSE)

## S3 method for class 'aplus'
ellipses(mean,var,r=1,...,steps=72,thinRatio=NULL)

## S3 method for class 'rplus'
ellipses(mean,var,r=1,...,steps=72,thinRatio=NULL)

## S3 method for class 'rmult'
ellipses(mean,var,r=1,...,steps=72,thinRatio=NULL)

Arguments

- mean: a compositional dataset or value of means or midpoints of the ellipses
- var: a variance matrix or a set of variance matrices given by var[i,] (multiple covariance matrices are not consistently implemented as of today). The principal axis of the variance give the axis of the ellipses, whereas the square-root of the eigenvalues times r give the half-diameters of the ellipse.
- r: a scaling of the half-diameters
- ...: further graphical parameters

Description

Draws ellipses from a mean and a variance into a plot.

Usage

```r
ellipses(mean,...)
```
steps: the number of discretisation points to draw the ellipses.

thinRatio: The ellipse function now be default plots the whole ellipsoid by giving its principle circumferences. However this is not reasonable for the thinner directions. If a direction other than the first two eigendirections has an eigenvalue not bigger than thinRatio*rmax it is not plotted. Thus thinRatio=1 reinstatiates the old behavior of the function. Later thinRatio(NULL will become the default, in which case the projection of the ellipse is plotted. However this is not implemented yet.

aspanel: Is the function called as slave to draw in a panel of a gsi.pairs plot, or as a user function setting up the plots.

Details

The ellipsoid/ellipse drawn is given by the solutions of

\[(x - mean)^t var^{-1} (x - mean) = r^2\]

in the respective geometry of the parameter space. Note that these ellipses can be added to panel plots (by means of orthogonal projections in the corresponding geometry).

There are actually three possibilities of drawing a a hyperdimensional ellipsoid or ellipse and non of them is perfect.

- thinRatio=1.1 This works like, what was implemented in the older versions of compositons, but never correctly documented. It draws an ellipse with main axes given by the two largest Eigendirections of the var-Matrix given.
- thinRatio=0 Draws all the ellipses given by every pair of eigendirections. In this way we get a visual impression of the high dimensional ellipsoid represend by the variance matrix. However the plots gets fastly cluttered in dimensions, when D>4. A 0<thinRatio<1 can avoid using eigendirection with small extend (i.e. smaller than thinRatio*largest Eigenvalue.
- thinRatio=NULL Draws in each Panel a two dimensional ellipse representing the marginal variance in the projection of the plot, if var was to be interpreted as a variance matrix. This can be seen as some sort of projection of the high dimensional ellipsoid, but is not necessarily its visual outline.

Author(s)


See Also

plot.acomp.

Examples

data(SimulatedAmounts)
plot(acomp(sa.lognormals))
tt<-acomp(sa.lognormals); ellipses(mean(tt),var(tt),r=2,col="red")
tt<-rcomp(sa.lognormals); ellipses(mean(tt),var(tt),r=2,col="blue")
endmemberCoordinates

Recast amounts as mixtures of end-members

Description

Computes the convex combination of amounts as mixtures of endmembers to explain X as well as possible.

Usage

endmemberCoordinates(X,...)
endmemberCoordinatesInv(K,endmembers,...)

## Default S3 method:
endmemberCoordinates(X,

endmembers=diag(gsi.getD(X)), ...)

## S3 method for class 'acomp'
endmemberCoordinates(X,

endmembers=clrInv(diag(gsi.getD(X))),...)

## S3 method for class 'aplus'
endmemberCoordinates(X,endmembers,...)

## S3 method for class 'rplus'
endmemberCoordinates(X,endmembers,...)

## S3 method for class 'rmult'
endmemberCoordinatesInv(K,endmembers,...)

## S3 method for class 'acomp'
endmemberCoordinatesInv(K,endmembers,...)

## S3 method for class 'rcomp'
endmemberCoordinatesInv(K,endmembers,...)

## S3 method for class 'aplus'
endmemberCoordinatesInv(K,endmembers,...)

## S3 method for class 'rplus'
endmemberCoordinatesInv(K,endmembers,...)
Arguments

- **X**: a data set of amounts or compositions, to be represented in as convex combination of the endmembers in the given geometry
- **K**: weights of the endmembers in the convex combination
- **endmembers**: a dataset of compositions of the same class as X. The number of endmembers given must not exceed the dimension of the space plus one.
- **...**: currently unused

Details

The convex combination is performed in the respective geometry. This means that, for rcomp objects, positivity of the result is only guaranteed with endmembers corresponding to extremal individuals of the sample, or completely outside its hull. Note also that, in acomp geometry, the endmembers must necessarily be outside the hull.

The main idea behind this functions is that the composition actually observed came from a convex combination of some extremal compositions, specified by endmembers. Up to now, this is considered as meaningful only in rplus geometry, and under some special circumstances, in rcomp geometry. It is not meaningful in terms of mass conservation in acomp and aplus geometries, because these geometries do not preserve mass: whether such an operation has an interpretation is still a matter of debate. In rcomp geometry, the convex combination is dependent on the units of measurements, and will be completely different for volume and mass %. Even more, it is valid only if the whole composition is observed (!).

Value

The `endmemberCoordinates` functions give a `rmult` data set with the weights (a.k.a. barycentric coordinates) allowing to build X as good as possible as a convex combination (a mixture) from endmembers. The result is of class rmult because there is no guarantee that the resulting weights are positive (although they sum up to one).

The `endmemberCoordinatesInv` functions reconstruct the convex combination from the weights K and the given endmembers. The class of endmembers determines the geometry chosen and the class of the result.

Author(s)


References


Examples

```r
data(SimulatedAmounts)
ep <- aplus(rbind(c(2,1,2),c(2,2,1),c(1,2,2)))
# mix the endmembers in "ep" with weights given by "sa.lognormals"
dat <- endmemberCoordinatesInv(acomp(sa.lognormals),acomp(ep))
par(mfrow=c(1,2))
```
Firework

plot(dat)
  plot(acomp(ep),add=TRUE,col="red",pch=19)
# compute the barycentric coordinates of the mixture in the "end-member simplex"
plot( acomp(endmemberCoordinates(dat,acomp(ep))))

dat <- endmemberCoordinatesInv(rcomp(sa.lognormals),rcomp(ep))
plot(dat)
plot( rcomp(endmemberCoordinates(dat,rcomp(ep)))))

dat <- endmemberCoordinatesInv(aplus(sa.lognormals),aplus(ep))
plot(dat)
plot( endmemberCoordinates(dat,aplus(ep))))

dat <- endmemberCoordinatesInv(rplus(sa.lognormals),rplus(ep))
plot(dat)
plot(endmemberCoordinates(rplus(dat),rplus(ep))))

Firework

Firework mixtures

Description

Data show two measured properties, brilliance and vorticity, of 81 girandoles composed of different mixtures of five ingredients: a – e. Of these ingredients, a and b are the primary light-producting, c is principal propellant, and d and e are binding agents for c.

Usage

data(Firework)

Details

The data consist of 81 cases and 7 variables: ingredients a, b, c, d, and e, and the two measured properties brilliance and vorticity. The 81 different mixtures form a special experiment design. First the 81 possible quadruples formed from the three values -1, 0, 1 were arranged in ascending order. Then for each such quadruple z, the corresponding mixture x(z)=(a,b,c,d,e)=alrInv(z) is computed. Thus the No. 4 girandole corresponds to z=(-1,-1,0,-1) and so is composed of a mixture x=(.12,.12,.32,.12,.32) of the five ingredients. All 5-part mixtures sum up to one.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name YATQUAD.DAT, here included under the GNU Public Library Licence Version 2 or newer.
References


**fitdirichlet**

**Fitting a Dirichlet distribution**

**Description**

Fits a Dirichtlet Distribution to a dataset by maximum likelihood.

**Usage**

`fitDirichlet(x, elog=mean(ult(x)), alpha0=rep(1, length(elog)), maxIter=20, n=nrow(x))`

**Arguments**

- `x`: a dataset of compositions (`acomph`)
- `elog`: the expected log can provided instead of the dataset itself.
- `alpha0`: the start value for alpha parameter in the iteration
- `maxIter`: The maximum number of iterations in the Fischer scoring method.
- `n`: the number of datapoints used to estimate elog

**Details**

The fitting is done using a modified version of the Fisher-Scoring method using analytiscal expressions for log mean and log variance. The modification is introducted to prevent the algorithm from leaving the admissible parameter set. It reduced the stepsize to at most have of distance to the limit of the admissible parameter set.

**Value**

- `alpha`: the estimated parameter
- `loglikelihood`: the likelihood
- `df`: The dimension of the dataset minus the dimension of the parameter

**Missing Policy**

Up to now the fitting can not handle missings.

**Author(s)**

References


See Also

`rDirichlet`, `acompDirichletGOF.test`, `runif.acomp`, `rnorm.acomp`

Examples

```r
x <- rDirichlet.acomp(100,c(1,2,3,4))
fitDirichlet(x)
```

**fitSameMeanDifferentVarianceModel**

_Fit Same Mean Different Variance Model_

Description

Fits a model of the same mean, but different variances model to a set of several multivariate normal groups by maximum likelihood.

Usage

`fitSameMeanDifferentVarianceModel(x)`

Arguments

- `x` list of rmult type datasets

Details

The function tries to fit a normal model with different variances but the same mean between different groups.

Value

- `mean` the estimated mean
- `vars` a list of estimated variance-covariance matrices
- `N` a vector containing the sizes of the groups

Author(s)

References


See Also

acompNormalLocation.test

Examples

fitSameMeanDifferentVarianceModel

gausstest  *Classical Gauss Test*

Description

One and two sample Gauss test for equal mean of normal random variates with known variance.

Usage

Gauss.test(x,y=NULL,mean=0,sd=1,alternative = c("two.sided", "less", "greater"))

Arguments

x  
a numeric vector providing the first dataset

y  
optional second dataset

mean  
the mean to compare with

sd  
the known standard deviation

alternative  
the alternative to be used in the test

Details

The Gauss test is in every Text-Book, but not in R, because it is nearly never used. However it is included here for educational purposes.

Value

A classical "htest" object

data.name  
The name of the dataset as specified

method  
a name for the test used

parameter  
the mean and variance provided to the test

alternative  
an empty string

p.value  
The p.value computed for this test
Author(s)


See Also
t.test

Examples

```r
x <- rnorm(100)
y <- rnorm(100)
Gauss.test(x,y)
```

---

**geometricmean**

*The geometric mean*

**Description**

Computes the geometric mean.

**Usage**

```r
geometricmean(x,...)
geometricmeanRow(x,...)
geometricmeanCol(x,...)
gsi.geometricmean(x,...)
gsi.geometricmeanRow(x,...)
gsi.geometricmeanCol(x,...)
```

**Arguments**

- `x` a numeric vector or matrix of data
- `...` further arguments to compute the mean

**Details**

The geometric mean is defined as:

\[
\text{geometricmean}(x) := \left( \prod_{i=1}^{n} x_i \right)^{1/n}
\]

The geometric mean is actually computed by \(\exp(\text{mean}(\log(c(\text{unclass}(x))),\ldots))\).
getDetectionLimit

Description

The detection limit of those values below-detection-limit are stored as negative values in compositional dataset. This function extracts that information.

Usage

getDetectionLimit(x, dl=attr(x,"detectionlimit"))

Arguments

x a data set

dl a default to replace the information in the dataset

Value

The geometric means of x as a whole (geometricmean), its rows (geometricmeanRow) or its columns (geometricmeanCol).

Missing Policy

The first three functions take the geometric mean of all non-missing values. This is because they should yield a result in term of data analysis.
Contrarily, the gsi.* functions inherit the arithmetic IEEE policy of R through \( \exp(\text{mean}(\log(c(\text{unclass}(x))),\ldots)) \).
Thus, NA codes a not available i.e. not measured, NaN codes a below detection limit, and 0.0 codes a structural zero. If any of the elements involved is 0, NA or NaN the result is of the same type.
Here 0 takes precedence over NA, and NA takes precedence over NaN. For example, if a structural 0 appears, the geometric mean is 0 regardless of the presence of NaN’s or NA’s in the rest. Values below detection limit become NaN’s if they are coded as negative values.

Author(s)


See Also

mean.rplus

Examples

geometricmean(1:10)
geometricmean(c(1,0,NA,NaN)) # 0
X <- matrix(c(1,NA,NaN,0,1,2,3,4),nrow=4)
X
geometricmeanRow(X)
geometricmeanCol(X)

getDetectionLimit x Gets the detection limit stored in the data set
**Details**

For a proper treatment of truncated data it would be necessary to know the detection limit even for observed data. Unfortunately, there is no clear way to encode this information without annoying the user.

**Value**

A matrix in the same shape as x, with a positive value (the detection limit) where available, and NA in the other cells.

**Author(s)**

K.Gerald van den Boogaart

**References**


**See Also**

`compositions.missings,zeroreplace`

**Examples**

```r
x <- c(2,-0.5,4,3,-0.5,5,BDLvalue,MARvalue,MNARvalue)
getDetectionlimit(x)
```

---

**Description**

In a pebble analysis of glacial tills, the total number of pebbles in each of 92 samples was counted and the pebbles were sorted into four categories red sandstone, gray sandstone, crystalline and miscellaneous. The percentages of these four categories and the total pebble counts are recorded.

**Usage**

data(Glacial)
Details
Percentages by weight in 92 samples of pebbles of glacial tills sorted into four categories, red sandstone, gray sandstone, crystalline and miscellaneous. The percentages of these four categories and the total pebbles counts are recorded. The glaciologist is interested in describing the pattern of variability of his data and whether the compositions are in any way related to abundance. All rows sum to 100, except for some rounding errors.

Note
Courtesy of J. Aitchison

Source
Aitchison: CODA microcomputer statistical package, 1986, the file name GLACIAL.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References

---

gof Compositional Goodness of fit test

Description
Goodness of fit tests for compositional data.

Usage
acompGOF.test(x,...)
acompNormalGOF.test(x,...,method="etest")
## S3 method for class 'formula'
acompGOF.test(formula, data,...,method="etest")
## S3 method for class 'list'
acompGOF.test(x,...,method="etest")
gsi.acompUniformityGOF.test(x,samplesize=nrow(x)*20,R=999)
acompTwoSampleGOF.test(x,y,...,method="etest",data=NULL)

Arguments
- x a dataset of compositions (acomp)
- y a dataset of compositions (acomp)
- samplesize number of observations in a reference sample specifying the distribution to compare with. Typically substantially larger than the sample under investigation
- R The number of replicates to compute the distribution of the test statistic
method  Selecting a method to be used. Currently only "etest" for using an energy test is supported.

... further arguments to the methods

formula an anova model formula defining groups in the dataset
data unused

Details

The compositional goodness of fit testing problem is essentially a multivariate goodness of fit test. However there is a lack of standardized multivariate goodness of fit tests in R. Some can be found in the energy-package.

In principle there is only one test behind the Goodness of fit tests provided here, a two sample test with test statistic.

\[ \frac{\sum_{ij} k(x_i, y_i)}{\sqrt{\sum_{ij} k(x_i, x_i) \sum_{ij} k(y_i, y_i)}} \]

The idea behind that statistic is to measure the cos of an angle between the distributions in a scalar product given by

\[ (X, Y) = E[k(X, Y)] = E[\int K(x - X)K(x - Y)dx] \]

where \( k \) and \( K \) are Gaussian kernels with different spread. The bandwidth is actually the standard deviation of \( k \).

The other goodness of fit tests against a specific distribution are based on estimating the parameters of the distribution, simulating a large dataset of that distribution and apply the two sample goodness of fit test.

For the moment, this function covers: two-sample tests, uniformity tests and additive logistic normality tests. Dirichlet distribution tests will be included soon.

Value

A classical "htest" object

data.name The name of the dataset as specified
method a name for the test used
alternative an empty string
replicates a dataset of p-value distributions under the Null-Hypothesis got from nonparametric bootstrap
p.value The p.value computed for this test

Missing Policy

Up to now the tests can not handle missings.

Author(s)

References


See Also

fitDirichlet, rDirichlet, runif.acomp, rnorm.acomp

Examples

```r
## Not run:
x <- runif.acomp(100,4)
y <- runif.acomp(100,4)

erg <- acompTwoSampleGOF.test(x,y)
#continue
erg
unclass(erg)

# Not run:
x <- runif.acomp(100,4)
y <- runif.acomp(100,4)

dd <- replicate(1000,acompGOF.test(runif.acomp(100,4),runif.acomp(100,4))$p.value)
hist(dd)

dd <- replicate(1000,acompGOF.test(runif.acomp(10,4),runif.acomp(100,4))$p.value)
hist(dd)

dd <- replicate(1000,acompGOF.test(runif.acomp(400,4),runif.acomp(10,4),bandwidth=4)$p.value)
hist(dd)

dd <- replicate(1000,acompGOF.test(runif.acomp(20,4),runif.acomp(100,4)+acomp(c(1,2,3,1)))$p.value)
hist(dd)

# test uniformity
attach("gsi") # the uniformity test is only available as an internal function
x <- runif.acomp(100,4)
gsi.acompUniformityGOF.test.test(x)

# Not run:
```

groupparts

## End(Not run)

groupparts

---

**groupparts**

*Group amounts of parts*

### Description

Groups parts by amalgamation or balancing of their amounts or proportions.

### Usage

```r
groupparts(x,...)
```

#### S3 method for class 'acomp'
```r
groupparts(x,...,groups=list(...))
```

#### S3 method for class 'rcomp'
```r
groupparts(x,...,groups=list(...))
```

#### S3 method for class 'aplus'
```r
groupparts(x,...,groups=list(...))
```

#### S3 method for class 'rplus'
```r
groupparts(x,...,groups=list(...))
```

#### S3 method for class 'ccomp'
```r
groupparts(x,...,groups=list(...))
```

### Arguments

- `x` an amount/compositional dataset
- `...` further parameters to use (actually ignored)
- `groups` a list of numeric xor character vectors, each giving a group of parts

### Details

In the real geometry grouping is done by amalgamation (i.e. adding the parts). In the Aitchison-geometry grouping is done by taking geometric means. The new parts are named by named formal arguments. Not-mentioned parts remain ungrouped.

### Value

A new dataset of the same type with each group represented by a single column

### Missing Policy

For the real geometries, SZ and BDL are considered as 0, and MAR and MNAR are kept as missing of the same type. For the relative geometries, a BDL is a special kind of MNAR, whereas a SZ is qualitatively different (thus a balance with a SZ has no sense). MAR values transfer their MAR property to the resulting new variable.
Author(s)


References


See Also

aplus

Examples

data(SimulatedAmounts)
plot(groupparts(acomp(sa.lognormals5),A=c(1,2),B=c(3,4),C=5))
plot(groupparts(aplus(sa.lognormals5),B=c(3,4),C=5))
plot(groupparts(rcomp(sa.lognormals5),A=c("Cu","Pb"),B=c(2,5)))
hist(groupparts(rplus(sa.lognormals5),1:5))

Hongite

Compositions of 25 specimens of hongite

Description

A mineral compositions of 25 rock specimens of hongite type. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite.

Usage

data(Hongite)

Details

A mineral compositions of 25 rock specimens of hongite type. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite, which we conveniently abbreviate to A, B, C, D, E. All row sums are equal to 100, except for rounding errors.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name HONGITE.DAT, here included under the GNU Public Library Licence Version 2 or newer.
HotellingsTsq

References

---

**Description**

The Hotellings T square distribution is the distribution of the squared Mahalanobis distances with respect to estimated variance covariance matrices.

**Usage**

```
qHotellingsTsq(p,n,m)
pHotellingsTsq(q,n,m)
```

**Arguments**

- `p` a (vector of) probabilities
- `q` a vector of quantiles
- `n` number of parameters, the p parameter of Hotellings $T^2$ distribution
- `m` number of dimensions, the m parameter of the Hotellings $T^2$ distribution

**Details**

The Hotellings $T^2$ with parameter $p$ and $m$ is the distribution empirical squared Mahalanobis distances of a $m$ dimensional vector with respect to a variance covariance matrix estimated based on $np$ degrees of freedom.

**Value**

- `qHotellingsT2` a vector of quantiles
- `pHotellingsT2` a vector of probabilities

**Author(s)**


**See Also**

`ellipses`, `ConfRadius`, `pf`

**Examples**

```
(q <- qHotellingsTsq(seq(0,0.9,by=0.1),3,25))
pHotellingsTsq(q,3,25)
```
**HouseholdExp**  
*Household Expenditures*

**Description**

Household budget survey data on month expenditures of twenty men and twenty women for four commodity groups: housing, foodstuffs, other, and services. Amounts in HK$ are given. There are 40 cases and 5 variables for 4 commodity groups and sex.

**Usage**

data(HouseholdExp)

**Details**

In a sample survey of people living alone in a rented accommodation, twenty men and twenty women were randomly selected and asked to record over a period of one month their expenditures on the following four mutually exclusive and exhaustive commodity groups: Housing, including fuel and lights, Foodstuffs, including alcohol and tobacco, Other goods, including clothing, footwear and durable goods, and Services, including transport and vehicles. Amounts in HK$ are given. There are 40 cases, 20 men and 20 women and 5 variables: 4 for commodity groups Housing, Food, Other, Services and the fifth sex, 1 for men, $-1$ for women. Note that the data has no sum constraint.

**Source**

Aitchison: CODA microcomputer statistical package, 1986, the file name HEMF.DAT, here included under the GNU Public Library Licence Version 2 or newer.

**References**


---

**Hydrochem**  
*Hydrochemical composition data set of Llobregat river basin water (NE Spain)*

**Description**

Contains a hydrochemical amount/compositional data set obtained from several rivers in the Llobregat river basin, in northeastern Spain.

**Usage**

data(Hydrochem)
Format

Data matrix with 485 cases and 19 variables.

Details

This hydrochemical data set contains measurements of 14 components, H, Na, K, Ca, Mg, Sr, Ba, NH\textsubscript{4}, Cl, HCO\textsubscript{3}, NO\textsubscript{3}, SO\textsubscript{4}, PO\textsubscript{4}, TOC. From them, hydrogen was derived by inverting the relationship between its stable form in water, H\textsubscript{3}O\textsuperscript{+}, and pH. Details can be found in Otero et al. (2005). Each of these parameters is measured approximately once each month during 2 years in 31 stations, placed along the rivers and main tributaries of the Llobregat river, one of the medium rivers in northeastern Spain.

The Llobregat river drains an area of 4948.2 km\textsuperscript{2}, and it is 156.6 km long, with two main tributaries, Cardener and Anoia. The headwaters of Llobregat and Cardener are in a rather unpolluted area of the Eastern Pyrenees. Mid-waters these rivers flow through a densely populated and industrialized area, where potash mining activity occurs and there are large salt mine tailings stored with no water proofing. There, the main land use is agriculture and stockbreeding. The lower course flows through one of the most densely populated areas of the Mediterranean region (around the city of Barcelona) and the river receives large inputs from industry and urban origin, while intensive agriculture activity is again present in the Llobregat delta. Anoia is quite different. Its headwaters are in an agricultural area, downwaters it flows through an industrialized zone (paper mills, tannery and textile industries), and near the confluence with Llobregat the main land use is agriculture again, mainly vineyards, with a decrease in industry and urban contribution. Given this variety in geological background and human activities, the sample has been splitted in four groups (higher Llobregat course, Cardener, Anoia and lower Llobregat course), which in turn are splitted into main river and tributaries (Otero et al, 2005). Information on these groupings, the sampling locations and sampling time is included in 5 complementary variables.

Author(s)

Raimon Tolosana-Delgado

Source

The dataset is also accessible in Otero et al. (2005), and are here included under the GNU Public Library Licence Version 2 or newer.

References


Examples

data(Hydrochem)
biplot(princomp(rplus(Hydrochem)))
biplot(princomp(rcomp(Hydrochem)))
biplot(princomp(aplus(Hydrochem)))
biplot(princomp(acomp(Hydrochem)))

idt

**Isometric default transform**

**Description**

Compute the isometric default transform of a vector (or dataset) of compositions or amounts in the selected class.

**Usage**

```r
idt(x,...)
## Default S3 method:
idt( x,... )
## S3 method for class 'acomp'
idt( x ,...)
## S3 method for class 'rcomp'
idt( x ,...)
## S3 method for class 'aplus'
idt( x ,...)
## S3 method for class 'rplus'
idt( x ,...)
## S3 method for class 'rmult'
idt( x ,...)
## S3 method for class 'ccomp'
idt( x ,...)
## S3 method for class 'factor'
```

```r
idtInv(x,orig,...)
## Default S3 method:
idtInv( x ,orig,...)
## S3 method for class 'acomp'
idtInv( x ,orig,...)
## S3 method for class 'rcomp'
idtInv( x ,orig,...)
## S3 method for class 'aplus'
idtInv( x ,orig,...)
## S3 method for class 'rplus'
idtInv( x ,orig,...)
## S3 method for class 'ccomp'
idtInv( x ,orig,...)
## S3 method for class 'rmult'
idtInv( x ,orig,...)
```
Arguments

x a classed amount or composition, to be transformed with its isometric default transform, or its inverse
...
generic arguments past to underlying functions
orig a compositional object which should be mimicked by the inverse transformation. It is used to determine the backtransform to be used, and eventually to reconstruct the names of the parts. It is the generic argument. Typically the orig-argument is the dataset that has been transformed in the first place.

Details

The general idea of this package is to analyse the same data with different geometric concepts, in a fashion as similar as possible. For each of the four concepts there exists an isometric transform expressing the geometry in a full-rank euclidean vector space. Such a transformation is computed by idt. For acomp the transform is ilr, for rcomp it is ipt, for aplus it is ilt, and for rplus it is iit. Keep in mind that the transform does not keep the variable names, since there is no guaranteed one-to-one relation between the original parts and each transformed variable.

The inverse idtInv is intended to allow for an "easy" and automatic back-transformation, without intervention of the user. The argument orig (the one determining the behaviour of idtInv as a generic function) tells the function which back-transformation should be applied, and gives the column names of orig to the back-transformed values of x. Therefore, it is very convenient to give the original classed data set used in the analysis as orig.

Value

A corresponding matrix of row-vectors containing the transforms.

Author(s)


References


See Also

cdt, ilr, ipt, ilt, cdtInv, ilrInv, iptInv, iltInv, iitInv

Examples

## Not run:
# the idt is defined by
idt <- function(x) UseMethod("idt",x)
idt.default <- function(x) x
idt.acomp <- function(x) ilr(x)
idt.rcomp <- function(x) ipt(x)
idt.aplus <- iit
idt.rplus <- iit

## End(Not run)
idt(acomp(1:5))
idt(rcomp(1:5))
data(Hydrochem)
x = Hydrochem[,c("Na","K","Mg","Ca")]
y = acomp(x)
z = idt(y)
y2 = idtInv(z,y)
par(mfrow=c(2,2))
for(i in 1:4){plot(y[,i],y2[,i])}

---

iit

**Isometric identity transform**

**Description**

Compute the isometric identity transform of a vector (dataset) of amounts and its inverse.

**Usage**

```r
iit( x ,...)
```

```r
iitInv( z ,... )
```

**Arguments**

- `x` a vector or data matrix of amounts
- `z` the iit-transform of a vector or data.matrix of iit-transforms of amounts
- `...` generic arguments, to pass to other functions.

**Details**

The iit-transform maps D amounts (considered in a real geometry) isometrically to a D dimensional euclidian vector. The iit is part of the rplus framework. Despite its trivial operation, it is present to achieve maximal analogy between the aplus and the rplus framework.

The data can then be analysed in this transformed space by all classical multivariate analysis tools. The interpretation of the results is easy since the relation to the original variables is preserved. However results may be inconsistent, since the multivariate analysis tools disregard the positivity condition and the inner laws of amounts.

The isometric identity transform is a simple identity given by

\[
iit(x)_i := x_i
\]
ilr

Value

ilt gives the isometric identity transform, i.e. simply the input stripped of the "rplus" class attribute, iptInv gives amounts with class "rplus" with the given iit, i.e. simply the argument checked to be a valid "rplus" object, and with this class attribute.

Note

iit can be used to unclass amounts.

Author(s)


References


See Also

ilt, ilr, rplus

Examples

(tmp <- iit(c(1,2,3)))
iitInv(tmp)
iitInv(tmp) - c(1,2,3) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(iit(cdata))
Arguments

- \( x \): a composition, not necessarily closed
- \( z \): the ilr-transform of a composition
- \( V \): a matrix, with columns giving the chosen basis of the clr-plane
- \( \ldots \): generic arguments, not used.
- \( \text{orig} \): a compositional object which should be mimicked by the inverse transformation. It is especially used to reconstruct the names of the parts.

Details

The ilr-transform maps a composition in the D-part Aitchison-simplex isometrically to a D-1 dimensional euclidian vector. The data can then be analysed in this transformation by all classical multivariate analysis tools. However, the interpretation of the results may be difficult, since there is no one-to-one relation between the original parts and the transformed variables.

The isometric logratio transform is given by

\[ ilr(x) := V^T clr(x) \]

with \( clr(x) \) the centred log ratio transform and \( V \in \mathbb{R}^{d \times (d-1)} \) a matrix which columns form an orthonormal basis of the clr-plane. A default matrix \( V \) is given by \( \text{ilrBase(D)} \).

Value

- \( ilr \) gives the isometric log ratio transform, \( ilrInv \) gives closed compositions with the given ilr-transforms.

Author(s)


References


See Also

- \( clr, alr, apt, ilrBase \)
**Examples**

```r
(tmp <- ilr(c(1,2,3)))
ilrInv(tmp)
ilrInv(tmp) - clo(c(1,2,3)) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(ilr(cdata))
ilrBase(D=3)
```

**Description**

Compute the basis of a clr-plane, to use with isometric log-ratio or planar transform of a (dataset of) compositions.

**Usage**

```r
ilrBase( x=NULL , z=NULL , D = NULL, method = "basic" )
```

**Arguments**

- **x**: optional dataset or vector of compositions
- **z**: optional dataset or vector containing ilr or ipt coordinates
- **D**: number of parts of the simplex
- **method**: method to build the basis, one of "basic", "balanced", "optimal" "PBhclust", "PBmaxvar" or "PBangprox"

**Details**

Method "basic" computes a triangular Helmert matrix (corresponding to the original ilr transformation defined by Egozcue et al, 2003). In this case, ilrBase is a wrapper catching the answers of gsi.ilrBase and is to be used as the more convenient function.

Method "balanced" returns an ilr matrix associated with a balanced partition, splitting the parts in groups as equal as possible. Transforms ilr and ipt computed with this basis are less affected by any component (as happens with "basic").

The following methods are all data-driven and will fail if x is not given. Some of these methods are extended to non-acomp datasets via the cpt general functionality. Use with care with non-acomp objects!

Method "optimal" is a wrapper to gsi.optimalilrBase, providing the ilr basis with less influence of missing values. It is computed as a hierarchical cluster of variables, with parts previously transformed to 1 (if the value is lost) or 0 (if it is recorded).
Methods "PBhclust", "PBmaxvar" and "PBangprox" are principal balance methods (i.e., balances approximating principal components in different ways). These are all resolved by calls to gsi.PrinBal. Principal balances functionality should be considered beta!

Value

All methods give a matrix containing by columns the basis elements for the canonical basis of the clr-plane used for the ilr and ipt transform. Only one of the arguments x, z or D is needed to determine the dimension of the simplex.

References


http://ima.udg.es/Activitats/CoDaWork03

See Also

clr, ilr, ipt

Examples

ilr(c(1,2,3))
ilrBase(D=2)
ilrBase(c(1,2,3))
ilrBase(z= ilr(c(1,2,3)) )
round(ilrBase(D=7),digits= 3)
ilrBase(D=7,method="basic")
ilrBase(D=7,method="balanced")

ilt

Isometric log transform

Description

Compute the isometric log transform of a vector (dataset) of amounts and its inverse.

Usage

ilt( x ,... )
iltInv( z ,... )

Arguments

x          a vector or data matrix of amounts
z          the ilt-transform of a vector or data matrix of ilt-transforms of amounts
...        generic arguments, not used.
Details

The ilt-transform maps D amounts (considered in log geometry) isometrically to a D dimensional euclidean vector. The ilt is part of the aplus framework.

The data can then be analysed in this transformation by all classical multivariate analysis tools. The interpretation of the results is easy since the relation to the original variables is preserved.

The isometric log transform is given by

\[ ilt(x)_i := \ln x_i \]

Value

ilt gives the isometric log transform, i.e. simply the log of the argument, whereas iltInv gives amounts with the given ilt, i.e. simply the exp of the argument.

Author(s)


References


See Also

ilm, iit, aplus

Examples

```r
(tmp <- ilt(c(1,2,3)))
iltInv(tmp)
iltInv(tmp) - c(1,2,3) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(ilt(cdata))
```

Description

Compute the isometric planar transform of a (dataset of) composition(s) and its inverse.
ipt( x , V = ilrBase(x),... )
iptInv( z , V = ilrBase(z=z),...,orig=NULL)
uciptInv( z , V = ilrBase(z=z),...,orig=NULL )

Arguments

x       a composition or a data matrix of compositions, not necessarily closed
z       the ipt-transform of a composition or a data matrix of ipt-transforms of compositions
V       a matrix with columns giving the chosen basis of the clr-plane
...     generic arguments. not used.
orig    a compositional object which should be mimicked by the inverse transformation. It is especially used to reconstruct the names of the parts.

details

The ipt-transform maps a composition in the D-part real-simplex isometrically to a D-1 dimensional euclidian vector. Although the transformation does not reach the whole $R^{D-1}$, resulting covariance matrices are typically of full rank.

The data can then be analysed in this transformation by all classical multivariate analysis tools. However, interpretation of results may be difficult, since the transform does not keep the variable names, given that there is no one-to-one relation between the original parts and each transformed variables. See cpt and apt for alternatives.

The isometric planar transform is given by

$$ipt(x) := V^t cpt(x)$$

with $cpt(x)$ the centred planar transform and $V \in R^{d \times (d-1)}$ a matrix which columns form an orthonormal basis of the clr-plane. A default matrix $V$ is given by $ilrBase(D)$

Value

ipt gives the centered planar transform, iptInv gives closed compositions with with the given ipt-transforms, uciptInv unconstrained iptInv does the same as iptInv but sets illegal values to NA rather than giving an error. This is a workaround to allow procedures not honoring the constraints of the space.

Author(s)


References

is.acomp

See Also

ilr, ilrBase, cpt

Examples

(tmp <- ipt(c(1,2,3)))
iptInv(tmp)
iptInv(tmp) - clo(c(1,2,3)) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(ipt(cdata))

is.acomp

Check for compositional data type

Description

is.XXX returns TRUE if and only if its argument is of type XXX

Usage

is.acomp(x)
is.rcomp(x)
is.aplus(x)
is.rplus(x)
is.rmult(x)
is.ccomp(x)

Arguments

x any object to be checked

Details

These functions only check for the class of the object.

Value

TRUE or FALSE

Author(s)


See Also

acomp, rcomp aplus, rplus
Examples

is.acomp(1:3)
is.acomp(acomp(1:3))
is.rcomp(acomp(1:3))
is.acomp(acomp(1:3)+acomp(1:3))

Description

Detect outliers with respect to a normal distribution model.

Usage

IsMahalanobisOutlier(X,...,alpha=0.05,goodOnly=NULL,
replicates=1000,corrected=TRUE,robust=TRUE,crit=NULL)

Arguments

X        a dataset (e.g. given as acomp, rcomp, aplus, rplus or rmult) object to which \textit{idt} and MahalanobisDist apply.
...      further arguments to MahalanobisDist/gsi.mahOutlier
alpha    The confidence level for identifying outliers.
goodOnly an integer vector. Only the specified index of the dataset should be used for estimation of the outlier criteria. This parameter if only a small portion of the dataset is reliable.
replicates The number of replicates to be used in the Monte Carlo simulations for determination of the quantiles. The replicates not given a minimum is computed from the alpha level to ensure reasonable precision.
corrected logical. Literature often proposed to compare the Mahalanobis distances with Chisq-Approximations of there distributions. However this does not correct for multiple testing. If corrected is true a correction for multiple testing is used. In any case we do not use the chisq-approximation, but a simulation based procedure to compute confidence bounds.
robust    A robustness description as define in \textit{robustnessInCompositions}
crit      The critical value to be used. Typically the routine is called mainly for the purpose of finding this value, which it does, when crit is NULL, however sometimes we might want to specify a value used by someone else to reproduce the results.

Details

See \textit{outliersInCompositions} and \textit{robustnessInCompositions} for a comprehensive introduction into the outlier treatment in compositions.

See \textit{OutlierClassifier1} for a highlevel method to classify observations in the context of outliers.
isoPortionLines

Value
A logical vector giving for each element the result of the alpha-level test for being an outlier. TRUE corresponds to a significant result.

Note
For some unknown reasons the computation sometimes produces NaN's. In this case a warning is issued and a recomputation is tried.

The package robustbase is required for using the robust estimations.

Author(s)

See Also
OutlierClassifier1, outlierplot, ClusterFinder1

Examples
## Not run:
data(SimulatedAmounts)
datas <- list(data1=sa.outliers1,data2=sa.outliers2,data3=sa.outliers3,
data4=sa.outliers4,data5=sa.outliers5,data6=sa.outliers6)

opar<-par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
tmp<-mapply(function(x,y){
  plot(x,col=ifelse(IsMahalanobisOutlier(x),"red","gray"))
  title(y)
},datas,names(datas))

## End(Not run)

isoPortionLines Isoportion- and Isoproportion-lines

Description
Add lines of equal portion and proportion to ternary diagrams, to serve as reference axis.

Usage
isoPortionLines(...)
## S3 method for class 'acomp'
isoPortionLines(by=0.2,at=seq(0,1,by=by),..., parts=1:3,total=1,labs=TRUE,lines=TRUE,unit="")
## S3 method for class 'rcomp'
isoPortionLines(by=0.2, at=seq(0,1,by=by),..., 
    parts=1:3, total=1, labs=TRUE, lines=TRUE, unit="")
isoProportionLines(...)
## S3 method for class 'acomp'
isoProportionLines(by=0.2, at=seq(0,1,by=by),..., 
    parts=1:3, labs=TRUE, lines=TRUE)
## S3 method for class 'rcomp'
isoProportionLines(by=0.2, at=seq(0,1,by=by),..., 
    parts=1:3, labs=TRUE, lines=TRUE)

Arguments

... graphical arguments
at numeric in [0,1]: which portions/proportions should be marked?
by numeric in (0,1]: steps between portions/proportions
parts numeric vector subset of {1,2,3}: the variables to be marked
total the total amount to be used in labeling
labs logical: plot the labels?
lines logical: plot the lines?
unit mark of the units e.g. "%"

Details

Isoportion lines give lines of the same portion of one of the parts, while isoproportion line gives lines of the same ratio between two parts. The isoproportion lines are straight lines in both the Aitchison and the real geometries of the simplex, while the isoportion lines are not straight in an Aitchison sense (only in the real one). However, note that both types of lines remain straight in the real sense when perturbed (von Eynatten et al., 2002).

Note

Currently IsoLines only works with individual plots. This is mainly due to the fact that I have no idea, what the user interface of this function should look like for multipanel plots. This includes philosophical problems with the meaning of isoportions in case of marginal plots.

Author(s)


References

See Also

plot.acomp

Examples

data(SimulatedAmounts)
plot(acomp(sa.lognormals))
isoPortionLines()
plot(acomp(sa.lognormals),center=TRUE)
isoPortionLines()
plot(rcomp(sa.lognormals))
isoPortionLines()
plot(acomp(sa.lognormals))
isoProportionLines()
plot(acomp(sa.lognormals),center=TRUE)
isoProportionLines()
plot(rcomp(sa.lognormals))
isoProportionLines()

---

jura

The jura dataset

Description

A geochemical dataset from the Swiss Jura.

Usage

data(juraset)
data(jura259)

Format

A 359x11 or 259x11 dataframe

Details

The JURA data set provided by J.-P. Dubois, IATE-Paedologie, Ecole Polytechnique Federale de Lausanne, 1015 Lausanne, Switzerland. Spatial coordinates and values of categorial and continuous attributes at the 359 sampled sites. The 100 test locations are denoted with a star. Rock Types: 1: Argovian, 2: Kimmeridgian, 3: Sequanian, 4: Portlandian, 5: Quaternary. Land uses: 1: Forest, 2: Pasture, 3: Meadow , 4: Tillage

X      X location coordinate
Y      Y location coordinate
Rock   Categorical: rocktype,
Land   Categorical: land usage
Cd     element amount,
Cu  element amount,
Pb  element amount,
Co  element amount,
Cr  element amount,
Ni  element amount,

All 3-part compositions sum to one.

Source
AI-Geostats

References
Atteia, O., Dubois, J.-P., Webster, R., 1994, Geostatistical analysis of soil contamination in the Swiss Jura: Environmental Pollution 86, 315-327
Webster, R., Atteia, O., Dubois, J.-P., 1994, Coregionalization of trace metals in the soil in the Swiss Jura: European Journal of Soil Science 45, 205-218

Examples
```r
## Not run:
data(juraset)
X <- with(juraset,cbind(X,Y))
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
lrv <- logratioVariogram(comp,X,maxdist=1,nbins=10)
plot(lrv)
## End(Not run)
```

kingTetrahedron    Ploting composition into rotatable tetrahedron

Description
Plots acomp/rcomp objects into tetrahedron exported in kinemage format.

Usage
```
kingTetrahedron(X, parts=1:4, file="tmptetrahedron.kin",
clu=NULL, vec=NULL, king=TRUE, scale=0.2, col=1,
title="Compositional Tetrahedron")
```
kingTetrahedron

Arguments

x
  a compositional acomp or rcomp object of 4 or more parts

parts
  a numeric or character vector specifying the 4 parts to be used.

file
  file.kin for 3D display with the KiNG (Kinemage, Next Generation) interactive
  system for three-dimensional vector graphics.

clu
  partition determining the colors of points

vec
  vector of values determining points sizes

king
  FALSE for Mage; TRUE for King (described below)

scale
  relative size of points

col
  color of points if clu=NULL

title
  The title of the plot

Details

The routine transforms a 4 parts mixture m quadrays into 3-dimensional XYZ coordinates and
writes them as file.kin. For this transformation we apply K. Urner: Quadrays and XYZ at http://www.grunch.net/synergetics/quadxyz.html. The kin file we display as 3-D animation with
KiNG viewer a free software available at http://kinemage.biochem.duke.edu. A kinemage is
a dynamic, 3-D illustration. The best way to take advantage of that is by rotating it and twisting
it around with the mouse click near the center of the graphics window and slowly dragging right or
left, up or down. Furthermore by clicking on points with the mouse (left button again), the label
associated with each point will appear in the bottom left of the graphics area and also the distance
from this point to the last will be displayed. With the right button drag we can zoom in and out of
the picture. This animation supports coloring and different sizing of points.

We can display the kin file as 3-D animation also with MAGE viewer a previous version of KiNG,
also a free software available at http://kinemage.biochem.duke.edu. For this one has to put king=FALSE
as a parameter.

Value

The function is called for its side effect of generating a file for 3D display with the KiNG (Kinemage,
Next Generation) interactive system for three-dimensional vector graphics. Works only with KiNG
viewer available at http://kinemage.biochem.duke.edu

Note

This routine and the documentation is based on mix.Quad2net from the MixeR-package of Vladimir
Batagelj and Matevz Bren, and has been contributed by Matevz Bren to this package. Only slight
modifications have been applied to make function compatible with the philosophy and objects of
the compositions package.

Author(s)

Vladimir Batagelj and Matevz Bren, with slight modifications of K.Gerald van den Boogaart
Kongite

Compositions of 25 specimens of kongite

Description
A mineral compositions of 25 rock specimens of kongite type. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite.

Usage
data(Kongite)

Details
A mineral compositions of 25 rock specimens of hongite type. Each composition consists of the percentage by weight of five minerals, albite, blandite, cornite, daubite, endite, which we conveniently abbreviate to A, B, C, D, E. All row percentage sums to 100.

Note
Courtesy of J. Aitchison
Source

Aitchison: CODA microcomputer statistical package, 1986, the file name HONGITE.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


Description

Functions taking coordinates given in various ways and joining the corresponding points with line segments.

Usage

```r
## S3 method for class 'a-comp'
lines(x, ..., steps=30, aspanel=FALSE)
## S3 method for class 'r-comp'
lines(x, ..., steps=30, aspanel=FALSE)
## S3 method for class 'aplus'
lines(x, ..., steps=30, aspanel=FALSE)
## S3 method for class 'r-plus'
lines(x, ..., steps=30, aspanel=FALSE)
## S3 method for class 'rmult'
lines(x, ..., steps=30, aspanel=FALSE)
```

Arguments

- `x` a dataset of the given type
- `...` further graphical parameters
- `steps` the number of discretisation points to draw the segments, which might be not visually straight.
- `aspanel` Logical, indicates use as slave to do actual drawing only.

Details

The functions add lines to the graphics generated with the corresponding plot functions. Adding to multipaneled plots, redraws the plot completely and is only possible, when the plot has been created with the plotting routines from this library. For the rcomp/rplus geometries the main problem is providing a function that reasonably works with lines leaving the area. We tried to use a policy of cutting the line at the actual borders of the (high dimensional) simplex. That can lead to very strange visual impression showing lines ending...
somewhere in the middle of the plot. However these lines actually hit some border of the simplex that is not shown in the plot. A hyper dimensional tetrahedron is even more difficult to imagine than a hyperdimensional cube.

Author(s)


See Also

plot.acomp, straight

Examples

data(SimulatedAmounts)

plot(acomp(sa.lognormals))
lines(acomp(sa.lognormals),col="red")
lines(rcomp(sa.lognormals),col="blue")

plot(aplus(sa.lognormals[,1:2]))
lines(aplus(sa.lognormals[,1:2]),col="red")
lines(rplus(sa.lognormals[,1:2],col="blue")

plot(rplus(sa.lognormals[,1:2]))
tt<-aplus(sa.lognormals[,1:2]); ellipses(mean(tt),var(tt),r=2,col="red")
tt<-rplus(sa.lognormals[,1:2]); ellipses(mean(tt),var(tt),r=2,col="blue")
tt<-rmult(sa.lognormals[,1:2]); ellipses(mean(tt),var(tt),r=2,col="green")

logratioVariogram  Empirical variograms for compositions

Description

Computes the matrix of logratio variograms.

Usage

logratioVariogram(comp, loc, maxdist=max(dist(loc))/2, nbins=20, dists=seq(0,maxdist,length.out=nbins+1), bins=cbind(dists[-length(dists)],dists[-1]), azimuth=0, azimuth.tol=180)
Arguments

comp  an acomp compositional dataset
loc   a matrix or dataframe providing the observation locations of the compositions. Any number of dimension >= 2 is supported.
maxdist  the maximum distance to compute the variogram for.
nbins   The number of distance bins to compute the variogram for
dists   The distances seperating the bins
bins    a matrix with lower and upper limit for the distances of each bin. A pair is counted if min<h<=max. min and max are provided as columns. bins is computed from maxdist,nbins and dists. If it is provided, it is used directly.
azimuth  For directional variograms the direction, either as an azimuth angle (i.e. a single real number) for 2D datasets or a unit vector pointing of the same dimension as the locations. The angle is clockwise from North in degree.
azimuth.tol  The angular tolerance it should be below 90 if a directional variogram is intended.

Details

The logratio-variogram is the set of variograms of each of the pairwise logratios. It can be proven that it carries the same information as a usual multivariate variogram. The great advantage is that all the functions have a direct interpretation and can be estimated even with (MAR) missings in the dataset.

Value

A list of class "logratioVariogram".

vg  A nbins x D x D array containing the logratio variograms
h  A nbins x D x D array containing the mean distance the value is computed on.
n  A nbins x D x D array containing the number of nonmissing pairs used for the corresponding value.

Author(s)


References

Tolosana, van den Boogaart, Pawlowsky-Glahn (2009) Estimating and modeling variograms of compositional data with occasional missing variables in R, StatGis09

See Also

vgram2lrvgram, CompLinModCoReg, vgmFit
Examples

```r
## Not run:
data(juraset)
X <- with(juraset,cbind(X,Y))
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
lrv <- logratioVariogram(comp,X,maxdist=1,nbins=10)
plot(lrv)

## End(Not run)
```

Description

Transforms model functions for different types of compositional (logratio)(co)variograms.

Usage

```r
cgram2vgram(cgram)
vgram2lrvgram(vgram)
```

Arguments

- `cgram`: A (matrix valued) covariance function.
- `vgram`: A (matrix valued) variogram functions.

Details

The variogram is given by `cgram(0)-cgram(h)` and `lrvgram(h)[,i,j]=vgram(h)[,i,i]+vgram(h)[,i,j]-2*vgram(h)[,i,j]`.

The logratio-variogram is the set of variograms of each of the pairwise logratios. It can be proven that it carries the same information as a usual multivariate variogram. The great advantage is that all the functions have a direct interpretation and can be estimated even with (MAR) missings in the dataset.

Value

A function that takes the same parameters as the input function (through a...parameterlist), but provides the corresponding variogram values (cgram2vgram) or logratio Variogram (vgram2lrvgram) values.

Author(s)

MahalanobisDist

References

Tolosana, van den Boogaart, Pawlowsky-Glahn (2009) Estimating and modeling variograms of compositional data with occasional missing variables in R, StatGis09

See Also

logratioVariogram, CompLinModCoReg, vgmFit

Examples

data(juraset)
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
vg <- CompLinModCoReg(~nugget()+sph(0.5)+R1*exp(0.7),comp)
vg(1:3)
vgram2lrgram(vg)(1:3)

Description

MahalanobisDist computes the Mahalanobis distances to the center or to other observations.

Usage

MahalanobisDist(x,center=NULL,cov=NULL,inverted=FALSE,...)
## S3 method for class 'rmult'
MahalanobisDist(x,center=NULL,cov=NULL,inverted=FALSE,...,
goodOnly=NULL,pairwise=FALSE,pow=1,
robust=FALSE,giveGeometry=FALSE)
## S3 method for class 'acomp'
MahalanobisDist(x,center=NULL,cov=NULL,inverted=FALSE,...,
goodOnly=NULL, pairwise=FALSE,pow=1,robust=FALSE,giveGeometry=FALSE)

Arguments

x the dataset
robust logical or a robust method description (see robustnessInCompositions) specifying how the center and covariance matrix are estimated, if not given.
... Further arguments to solve.
center An estimated for the center (mean) of the dataset. If center is NULL it will be estimated based using the given robust option.
cov An estimated for the spread (covariance matrix) of the dataset. If cov is NULL it will be estimated based using the given robust option.
inverted TRUE if the inverse of the covariance matrix is given.
goodOnly
An vector of indices to the columns of x that should be used for estimation of center and spread.

pairwise
If FALSE the distances to the center are returned as a vector. If TRUE the distances between the cases are returned as a distance matrix.

pow
The power of the Mahalanobis distance to be used. 1 correponds to the square root of the squared distance in transformed space, like it is defined in most books. The choice 2 corresponds to what is implemented in many software package including the mahalanobis function in R.

giveGeometry
If true an attributes "center" and "cov" given the center and the idt-variance used for the calculations.

Details
The Mahalanobis distance is the distance in a linearly transformed space, where the linear transformation is selected in such a way, that the variance is the unit matrix. Thus the distances are given in multiples of standard deviation.

Value
Either a vector of Mahalanobis distances to the center, or a distance matrix (like from dist) giving the pairwise Mahalanobis distances of the data.

Note
Unlike the mahalanobis function this function does not be default compute the square of the mahalanobis distance. The pow option is provided if the square is needed.
The package robustbase is required for using the robust estimations.

Author(s)

See Also
dist, OutlierClassifier1

Examples

data(SimulatedAmounts)
data5 <- acomp(sa.outliers5)

cl <- ClusterFinder1(data5,sigma=0.4, radius=1)
plot(data5,col=as.numeric(cl$types),pch=as.numeric(cl$types))
legend(1,1,legend=levels(cl$types),xjust=1,col=1:length(levels(cl$types)),
pch=1:length(levels(cl$types)))
matmult

inner product for matrices and vectors

Description

Multiplies two matrices, if they are conformable. If one argument is a vector, it will be coerced to either a row or a column matrix to make the two arguments conformable. If both are vectors it will return the inner product.

Usage

x %*% y

## Default S3 method:
x %*% y

Arguments

x, y
umeric or complex matrices or vectors

Details

This is a copy of the %*% function. The function is made generic to allow the definition of specific methods.

Value

The matrix product. Uses 'drop' to get rid of dimensions which have only one level.

Author(s)


See Also

%*% rmult

Examples

M <- matrix(c(0.2,0.1,0.0,0.1,0.2,0.0,0.0,0.0,0.2),byrow=TRUE,nrow=3)
x <- c(1,1,2)
M %*% x
x %*% M
x %*% x
M %*% M
t(x) %*% M
Mean amounts and mean compositions

Description
Compute the mean in the several approaches of compositional and amount data analysis.

Usage
## S3 method for class 'acomp'
mean(x,...,robust=getOption("robust"))

## S3 method for class 'rcomp'
mean(x,...,robust=getOption("robust"))

## S3 method for class 'aplus'
mean(x,...,robust=getOption("robust"))

## S3 method for class 'rplus'
mean(x,...,robust=getOption("robust"))

## S3 method for class 'ccomp'
mean(x,...,robust=getOption("robust"))

## S3 method for class 'rmult'
mean(x,...,na.action=NULL,robust=getOption("robust"))

Arguments
- x: a classed dataset of amounts or compositions
- ...: further arguments to mean e.g. trim
- na.action: na.action
- robust: A description of a robust estimator. Possible values are FALSE or "pearson" for no robustness, or TRUE or "mcd" for a covMcd based robust location scale estimation. Additional control parameters such as list(trim=0.2) or an rrcov.control object can be given as an attribute "control".

Details
The different compositional approaches acomp, rcomp, aplus, rplus correpond to different geometries. The mean is calculated in the respective canonical geometry by applying a canonical transform (see cdt), taking ordinary meanCol and backtransforming.

The Aitchison geometries imply that mean.acomp and mean.aplus are geometric means, the first one closed. The real geometry implies that mean.rcomp and mean.rplus are arithmetic means, the first one resulting in a closed composition.

In all cases the mean is again an object of the same class.
meanrow

Value
The mean is given as a composition or amount vector of the same class as the original dataset.

Missing Policy
For the additive scales (rcomp,rplus) the SZ and BDT are treated as zeros and MAR and MNAR as missing information. This is not strictly correct for MNAR.
For relative scales (acomp,aplus), all four types of missings are treated as missing information. This corresponds to the idea that BDT are truncated values (and have the corresponding effect in taking means). For SZ and MAR, only the components in the observed subcomposition are fully relevant. Finally, for MNAR the problem is again that nothing could be done without knowing the MNAR mechanism, so the analysis is limited to taking them as MAR, and being careful with the interpretation. Missing and Below Detection Limit Policy is explained in more detail in compositions.missing.

Author(s)

See Also
clo, meanCol, geometricmean, acomp, rcomp, aplus, rplus

Examples

data(SimulatedAmounts)
meanCol(sa.lognormals)
mean(acomp(sa.lognormals))
mean(rcomp(sa.lognormals))
mean(aplus(sa.lognormals))
mean(rplus(sa.lognormals))
mean(rmult(sa.lognormals))

---

meanrow

The arithmetic mean of rows or columns

Description
Computes the arithmetic mean.

Usage

meanRow(x,..., na.action=get(getOption("na.action")))
meanCol(x,..., na.action=get(getOption("na.action")))
Arguments

- **x**: a numeric vector or matrix of data
- **...**: arguments to `mean`
- **na.action**: The na.action to be used: one of `na.omit`, `na.fail`, `na.pass`

Details

Computes the arithmetic means of the rows (meanRow) or columns (meanCol) of x.

Value

The arithmetic means of the rows (meanRow) or columns (meanCol) of x.

Author(s)


See Also

`mean.rplus`

Examples

```r
data(SimulatedAmounts)
meanCol(sa.tnormals)
meanRow(sa.tnormals)
```

---

**Metabolites**

*Steroid metabolite patterns in adults and children*

Description

Data shows the urinary excretion (mg/24 hours) of 37 normal adults and 30 normal children of total cortisol metabolites, total corticosterone metabolites, total pregnanetriol and Δ-5-pregnentriol.

Usage

```r
data(Metabolites)
```

Details

There are 67 cases for 37 adults and 30 children, and 5 columns: Case no., met1, met2, met3 and Type, 1 for adults, $-1$ for children. No sum constraint is placed on this data set: since the urinary excretion in mg for 24 hours are given.
missing.compositions

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name METABOL.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


missing.compositions The policy of treatment of missing values in the "compositions" package

Description

This help section discusses some general strategies of working with missing values in a compositional, relative or vectorial context and shows how the various types of missing values are represented and treated in the "compositions" package, according to each strategy/class of analysis of compositions or amounts.

Usage

is.BDL(x,mc=attr(x,"missingClassifier"))
is.SZ(x,mc=attr(x,"missingClassifier"))
is.MAR(x,mc=attr(x,"missingClassifier"))
is.MNAR(x,mc=attr(x,"missingClassifier"))
is.NMV(x,mc=attr(x,"missingClassifier"))
is.WMNAR(x,mc=attr(x,"missingClassifier"))
is.WZERO(x,mc=attr(x,"missingClassifier"))
has.missings(x,...)
## Default S3 method:
has.missings(x,mc=attr(x,"missingClassifier"),...)
## S3 method for class 'rmult'
has.missings(x,mc=attr(x,"missingClassifier"),...)
SZvalue
MARvalue
MNARvalue
BDLvalue

Arguments

x A vector, matrix, acomp, rcomp, aplus, rplus object for which we would like to know the missing status of the entries
A missing classifier function, giving for each value one of the values BDL (Below Detection Limit), SZ (Structural Zero), MAR (Missing at random), MNAR (Missing not at random), NMV (Not missing value). These functions are introduced to allow a different coding of the missings.

Details

In the context of compositional data we have to consider at least four types of missing and zero values:

- **MAR** (Missing at random) coded by NaN, the amount was not observed or is otherwise missing, in a way unrelated to its actual value. This is the "nice" type of missing.
- **MNAR** (Missing not at random) coded by NA, the amount was not observed or is otherwise missing, but it was missed in a way stochastically dependent on its actual value.
- **BDL** (Below detection limit) coded by 0.0 or a negative number giving the detection limit; the amount was observed but turned out to be below the detection limit and was thus rounded to zero. This is an informative version of MNAR.
- **SZ** (Structural zero) coded by -Inf, the amount is absolutely zero due to structural reasons. E.g. a soil sample was dried before the analysis, or the sample was preprocessed so that the fraction is removed. Structural zeroes are mainly treated as MAR even though they are a kind of MNAR.

Based on these basic missing types, the following extended types are defined:

- **NMV** (Not Missing Value) coded by a real number, it is just an actually-observed value.
- **WMNAR** (Wider MNAR) includes BDL and MNAR.
- **WZERO** (Wider Zero) includes BDL and SZ

Each function of type `is.XXX` checks the status of its argument according to the XXX type of value from those above.

Different steps of a statistical analysis and different understanding of the data will lead to different approaches with respect to missings and zeros. In the first exploratory step, the problem is to keep the methods working and to make the missing structure visible in the analysis. The user should need as less as possible extra thinking about missings, an get nevertheless a true picture of the data. To achieve this we tried to make the basic layer of computational functions working consistently with missings and propagating the missingness character seamlessly. However some of this only works with `acomp`, where a closed form missing theories are available (e.g. proportional imputation [e.g. Martín-Fernández, J.A. et al.(2003)] or estimation with missings [Boogaart&Tolosana 2006]). The main graphics should hint towards missing and try to add missings to the plot by marking the remaining information on the axes. However one again should be clear that this is only reasonably justified in the relative geometries. Unfortunately the missing subsystem is currently not fully compatible with the robustness subsystem.

As a second step, the analyst might want to analyze the missing structure for itself. This is preliminarily provided by these functions, since their result can be treated as a boolean data set in any other R function. Additionally a `missingSummary` provides some a convenience function to provide a fast overview over the different types of missings in the dataset.
In the later inferential steps, the problem is to get results valid with respect to a model. One needs to be able to look through the data on the true processes behind, without being distracted by artifacts stemming from missing values. For the moment, how analyses react to the presence of missings depend on the value of the na.action option. If this is set to na.omit (the default), then cases with missing values on any variable are completely ignored by the analysis. If this is set to na.pass, then some of the following applies.

The policy on how a missing value is to be introduced into the analysis depends on the purpose of the analysis, the type of analysis and the model behind. With respect to this issue this package and probably the whole science of compositional data analysis is still very preliminary.

The four philosophies work with different approaches to these problems:

- **rplus** For positive real vectors, one can either identify BDL with a true 0 or impute a value relative to the detection limit, with a function like `zeroreplace`. A structural zero can either be seen as a true zero or as a MAR value.

- **rcomp** and **acomp** For these relative geometries, a true zero is an alien. Thus a BDL is nothing else but a small unkown value. We could either decide to replace the value by an imputation, or go through the whole analysis keeping this lack of information in mind. The main problem of imputation is that by closing to 1, the absolute value of the detection limit is lost, and the detection limit can correspond to very different portions. Raw differences between all, observed or missed, components (the ground of the rcomp geometry) are completely distorted by the replacement. Contrarily, log-ratios between observed components do not change but ratios between missed components dramatically depend on the replacement. e.g. typically the content of gold is some orders of magnitude smaller than the contend of silver even around a gold deposit, but far away from the deposit they both might be far under detection limit, leading to a ratio of 1, just because nothing was observed. SZ in compositions might be either seen as defining two sub-populations, one fully defined and one where only a subcomposition is defined. But SZ can also very much be like an MAR, if only a subcomposition is measured. Thus, in general we can simply understand that only a subcomposition is available, i.e. a projection of the true value onto a sub-space: for each observation, this sub-space might be different. For MAR values, this approach is stricly valid, and yields unbiased estimations (because these projections are stochastically independent of the observed phenomenon). For MNAR values, the projections depend on the actual value, which strictly speaking yields biased estimations.

- **aplus** Imputation takes place by simple replacement of the value. However this can lead to a dramatic change of ratios and should thus be used only with extra care, by the same reasons explained before.

More information on how missings are actually processed can be found in the help files of each individual functions.

**Value**

A logical vector or matrix with the same shape as x stating wether or not the value is of the given type of missing.

**Author(s)**

References


Aitchison, J., C. Barcel'o-Vidal, J.J. Egozcue, V. Pawlowsky-Glahn (2002) A concise guide to the algebraic geometric structure of the simplex, the sample space for compositional data analysis, Terra Nostra, Schriften der Alfred Wegener-Stiftung, 03/2003


See Also

compositions-package, missingsInCompositions, robustnessInCompositions, outliersInCompositions, zeroreplace, rmult, ilr, mean.acomp, acomp, plot.acomp

Examples

```r
require(compositions)  # load library
data(SimulatedAmounts) # load data sa.lognormals
dat <- acomp(sa.missings)  # load data sa.missings
dat
var(dat)
mean(dat)
plot(dat)
boxplot(dat)
barplot(dat)
```

missingProjector

Returns a projector the observed space in case of missings.

Description

Returns projectors on the observed subspace in the presence of missings.
Usage

missingProjector(x,...,by="s")
## S3 method for class 'acomp'
missingProjector(x,has=is.NMV(x),...,by="s")
## S3 method for class 'aplus'
missingProjector(x,has=is.NMV(x),...,by="s")
## S3 method for class 'rcomp'
missingProjector(x,has=!is.MAR(x) & is.MNAR(x),...,by="s")
## S3 method for class 'rplus'
missingProjector(x,has=!is.MAR(x) & is.MNAR(x),...,by="s")

Arguments

x a dataset or object of the given class
has a boolean matrix of the same size indicating nonmissing values
... additional arguments for generic purpose only
by the name of the dataset dimension on has for tensorial computation with tensorA package

Details

See the references for details on that function.

Value

A dataset of N square matrices of dimension DxD (with N and D respectively equal to the number of rows and columns in x). Each of these matrices gives the projection of a data row onto its observed sub-space.

The function `sumMissingProjector` takes all these matrices and sums them, generating a "summary" of observed sub-spaces. This matrix is useful to obtain estimates of the mean (and variance, in the future) still unbiased in the presence of lost values (only of type MAR, strictly-speaking, but anyway useful for any type of missing value, when used with care).

Author(s)

K.G.van den Boogaart

References


See Also

missingsInCompositions
Examples

```r
data(SimulatedAmounts)
x <- acomp(sa.lognormals)
xnew <- simulateMissings(x,dl=0.05,MAR=0.05,MNAR=0.05,SZ=0.05)
xnew
plot(missingSummary(xnew))

missingProjector(acomp(xnew))
missingProjector(rcomp(xnew))
missingProjector(aplus(xnew))
missingProjector(rplus(xnew))
```

---

missingsummary

Classify and summarize missing values in a dataset

**Description**

Routines classifies codes of missing values as numbers in objects of the compositions package.

**Usage**

```r
missingSummary(x,..., vlabs = colnames(x),
mc=attr(x,"missingClassifier"),
values=eval(formals(missingType)$values))
```

**Arguments**

- `x` a dataset which might contain missings
- `...` additional arguments for `mc`
- `mc` optionally in `missingSummary`, an alternate routine to be used instead of `missingType`
- `vlabs` labels for the variables
- `values` the names of the different types of missings. "Err" is a value that can not be classified e.g. Inf.

**Details**

The function mainly counts the various types of missing values.

**Value**

`missingType` returns a character vector/matrix with the same dimension and dimnames as `x` giving the type of every value.

`missingSummary` returns a table giving the number of missings of each type for each variable.
**mix.Read**

**Reads a data file in a mixR format**

**Description**

Reads a data file, which is formatted in a simple compositional file including the first row with title, the second with data labels and afterwards the matrix with the data itself. In the first column of the matrix are cases labels. This is the format used in the mixR package.

**Usage**

```r
mix.Read(file, eps=1e-6)
```

**Arguments**

- `file` a file name
- `eps` the epsilon to be used for checking null values.

**Details**

The data files must have the adequate structure:

- 1. the first row with a title of the data set,
- 2. the second row with variables names
- 3. the data set in a matrix, rows as cases, variables in columns with the first column comprising cases labels.

**Author(s)**

K. Gerald van den Boogaart

**References**


**See Also**

- `compositions.missing`

**Examples**

```r
data(SimulatedAmounts)
x <- acompsa.lognormals)
xnew <- simulateMissings(x, d1=0.05, MAR=0.05, MNAR=0.05, S2=0.05)
xnew
missingSummary(xnew)
```
A mixture object 'm' consists of m$tit the title, m$mat the matrix with the data, m$sum the value of the rows total, if constant and m$sta the status of the mixture object with values:
mvar

-2 - matrix contains negative elements,
-1 - zero row sum exists,
0  - matrix contains zero elements,
1  - matrix contains positive elements, rows with different row sum(s),
2  - matrix contains constant row sum and
3  - closed mixture, the row sums are all equal to 1.

Value

A mixture object as a data frame with a title, row total, if constant, status (-2, -1, 0, 1, 2 or 3 – see above) and class attributes and the data matrix.

See Also

read.geoeas read.geoEAS read.table

Examples

## Not run:
mix.Read("GLACIAL.DAT")
mix.Read("ACTIVITY.DAT")

## End(Not run)

mvar

Metric summary statistics of real, amount or compositional data

Description

Compute the metric variance, covariance, correlation or standard deviation.

Usage

mvar(x,...)
mcov(x,...)
mcor(x,...)
msd(x,...)

## Default S3 method:
mvar(x,y=NULL,...)
## Default S3 method:
mcov(x,y=x,...)
## Default S3 method:
mcor(x,y,...)
## Default S3 method:
msd(x,y=NULL,...)
Arguments

x  a dataset, eventually of amounts or compositions
y  a second dataset, eventually of amounts or compositions
... further arguments to var or cov. Typically a robust=TRUE argument. e.g. use

Details

The metric variance (mvar) is defined by the trace of the variance in the natural geometry of the data, or also by the generalized variance in natural geometry. The natural geometry is equivalently given by the cdt or idt transforms.

The metric standard deviation (msd) is not the square root of the metric variance, but the square root of the mean of the eigenvalues of the variance matrix. In this way it can be interpreted in units of the original natural geometry, as the radius of a spherical ball around the mean with the same volume as the 1-sigma ellipsoid of the data set.

The metric covariance (mvar) is the sum over the absolute singular values of the covariance of two datasets in their respective geometries. It is always positive. The metric covariance of a dataset with itself is its metric variance. The interpretation of a metric covariance is quite difficult, but useful in regression problems.

The metric correlation (mcor) is the metric covariance of the datasets in their natural geometry normalized to unit variance matrix. It is a number between 0 and the smaller dimension of both natural spaces. A number of 1 means perfect correlation in 1 dimension, but only partial correlations in higher dimensions.

Value

a scalar number, informing of the degree of variation/covariation of one/two datasets.

Author(s)


References


See Also

var, cov, mean.aomp, acomp, rcomp, aplus, rplus
Examples

data(SimulatedAmounts)
mvar(acomp(sa.lognormals))
mvar(rcomp(sa.lognormals))
mvar(aplus(sa.lognormals))
mvar(rplus(sa.lognormals))

msd(acomp(sa.lognormals))
msd(rcomp(sa.lognormals))
msd(aplus(sa.lognormals))
msd(rplus(sa.lognormals))

mcov(acomp(sa.lognormals5[,1:3]),acomp(sa.lognormals5[,4:5]))
mcor(acomp(sa.lognormals5[,1:3]),acomp(sa.lognormals5[,4:5]))
mcov(rcomp(sa.lognormals5[,1:3]),rcomp(sa.lognormals5[,4:5]))
mcor(rcomp(sa.lognormals5[,1:3]),rcomp(sa.lognormals5[,4:5]))
mcov(aplus(sa.lognormals5[,1:3]),aplus(sa.lognormals5[,4:5]))
mcor(aplus(sa.lognormals5[,1:3]),aplus(sa.lognormals5[,4:5]))
mcov(rplus(sa.lognormals5[,1:3]),rplus(sa.lognormals5[,4:5]))
mcor(rplus(sa.lognormals5[,1:3]),rplus(sa.lognormals5[,4:5]))
mcov(acomp(sa.lognormals5[,1:3]),aplus(sa.lognormals5[,4:5]))
mcor(acomp(sa.lognormals5[,1:3]),aplus(sa.lognormals5[,4:5]))

names

The names of the parts

Description

The names function provide a transparent way to access the names of the parts regardless of the shape of the dataset or data item.

Usage

## S3 method for class 'acomp'
names(x)
## S3 method for class 'rcomp'
names(x)
## S3 method for class 'aplus'
names(x)
## S3 method for class 'rplus'
names(x)
## S3 method for class 'rmult'
names(x)
## S3 method for class 'ccomp'
names(x)
## S3 replacement method for class 'acomp'
Arguments

x an amount/amount dataset

value the new names of the parts

Value

da character vector giving the names of the parts

Author(s)


See Also

aplus

Examples

data(SimulatedAmounts)
tmp <- acomp(sa.lognormals)
names(tmp)

names(tmp) <- c("x","y","z")
tmp

norm

Vector space norm

Description

Each of the considered space structures has an associated norm, which is computed for each element by these functions.
Usage

## Default S3 method:
```
norm(X,...)
```

## S3 method for class 'acomp'
```
norm(X,...)
```

## S3 method for class 'rcomp'
```
norm(X,...)
```

## S3 method for class 'aplus'
```
norm(X,...)
```

## S3 method for class 'rplus'
```
norm(X,...)
```

## S3 method for class 'rmult'
```
norm(X,...)
```

Arguments

- **X**
  - a dataset or a single vector of some type
- **...**
  - currently not used, intended to select a different norm rule in the future

Value

The norms of the given vectors.

Author(s)


See Also

- `normalize`

Examples

```r
data(SimulatedAmounts)
tmp <- acomp(sa.lognormals)
mvar(tmp)
sum(norm(tmp - mean(tmp))^2)/(nrow(tmp)-1)
```
normalize  Normalize vectors to norm 1

Description

Normalize vectors to norm 1.

Usage

normalize(x,...)
## Default S3 method:
normalize(x,...)

Arguments

x  a dataset or a single vector of some type
...
 currently not used, intended to select a different norm in the future

Value

The vectors given, but normalized to norm 1.

Author(s)


See Also

norm.rmult

Examples

data(SimulatedAmounts)
normalize(c(1,2,3))
normalize(acomp(c(1,2,3)))
norm(normalize(acomp(sa.groups))))
Description
Tests for several groups of additive lognormally distributed compositions.

Usage
acompNormalLocation.test(x, g=NULL, var.equal=FALSE, paired=FALSE,
R=ifelse(var.equal,999,0))

Arguments
x a dataset of compositions (acomp) or a list of such

g a factor grouping the data, not used if x is a list already. Alternatively, g can be
a second compositional data set.

var.equal a boolean telling whether the variance of the groups should be considered equal

paired true if a paired test should be performed

R number of replicates that should be used to compute p-values. 0 means comparing
the likelihood statistic with the corresponding asymptotic chisq-distribution.

Details
The tests are based on likelihood ratio statistics.

Value
A classical "htest" object

data.name The name of the dataset as specified

method a name for the test used

alternative an empty string

replicates a dataset of p-value distributions under the Null-Hypothesis got from nonpara-
metric bootstrap

p.value The p.value computed for this test

Missing Policy
Up to now the tests cannot handle missings.

Note
Do not trust the p-values obtained forcing var.equal=TRUE and R=0. This will include soon equiv-
alent spread tests.
Author(s)


References


See Also

`fitDirichlet`, `rDirichlet`, `runif.acomp`, `rnorm.acomp`.

Examples

```r
x <- runif.acomp(100,4)
y <- runif.acomp(100,4)
acompNormalLocation.test(list(x,y))
```

---

**oneOrDataset**

*Treating single compositions as one-row datasets*

Description

A dataset is converted to a data matrix. A single data item (i.e. a simple vector) is converted to a one-row data matrix.

Usage

```r
oneOrDataset(W,B=NULL)
```

Arguments

- `W` a vector, matrix or dataframe
- `B` an optional second vector, matrix or data frame having the intended number of rows.

Value

A data matrix containing the same data as `W`. If `W` is a vector it is interpreted as a single row. If `B` is given and `length(dim(B)) != 2` and `W` is a vector, then `W` is repeated `nrow(B)` times.

Author(s)

outlierclassifier

Detect and classify compositional outliers.

Description
Detects outliers and classifies them according to different possible explanations.

Usage

```r
OutlierClassifier1(X,...)
```

## S3 method for class 'acomp'

```r
OutlierClassifier1(X,...,alpha=0.05,
    type=c("best","all","type","outlier","grade"),goodOnly=NULL,
    corrected=TRUE,RedCorrected=FALSE,robust=TRUE)
```

Arguments

- **X**
  the dataset as an acomp object

- **...**
  further arguments to MahalanobisDist/gsi.mahOutlier

- **alpha**
  The confidence level for identifying outliers.

- **type**
  What type of classification should be used: best: Which single component would best explain the outlier. all: Give a binary coding specifying all components, which could explain the outlier. type: Is it a a normal observation "ok", a single component outlier "1" or can it not be explained by a single wrong component "?". outlier: All outliers are marked as "outlier". others are marked as "ok". grade: Proven Outliers are marked as "outlier". suspected outliers, detected without correction of the p-value are reported as "extreme", the rest is reported as "ok".

- **goodOnly**
  an integer vector. Only the specified index of the dataset should be used for estimation of the outlier criteria. This parameter if only a small portion of the dataset is reliable.

- **corrected**
  logical. Literature often proposed to compare the Mahalanobis distances with Chisq-Approximations of there distributions. However this does not correct for multiple testing. If corrected is true a correction for multiple testing is used. In any case we do not use the chisq-approximation, but a simulation based procedure to compute confidence bounds.

- **RedCorrected**
  logical. If an outlier is detected we can try to find out wether a single component would be sufficient to drop the outlier under the outlier detection limit. Since in this second case we only check a few outliers no second correction step applies as long as the number of outliers is not very high.

- **robust**
  A robustness description as define in var.acomp

Examples

oneOrDataset(c(1,2,3))
oneOrDataset(c(1,2,3),matrix(1:12,nrow=4))
oneOrDataset(data.frame(matrix(1:12,nrow=4)))
Details

See `outliersInCompositions` for a comprehensive introduction into the outlier treatment in compositions.

See `ClusterFinder1` for an alternative method to classify observations in the context of outliers.

Value

A factor classifying the observations in the dataset as "ok" or some type of outlier.

Note

The package `robustbase` is required for using the robust estimations.

Author(s)


See Also

`outlierplot`, `ClusterFinder1`

Examples

```r
## Not run:
tmp<-set.seed(1400)
A <- matrix(c(0.1,0.2,0.3,0.1),nrow=2)
Mvar <- 0.1*ilrvar2clr(A%*%t(A))
Mcenter <- acomp(c(1,2,1))
data(SimulatedAmounts)
datas <- list(data1=sa.outliers1,data2=sa.outliers2,data3=sa.outliers3,
data4=sa.outliers4,data5=sa.outliers5,data6=sa.outliers6)
opar<-par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
tmp<-mapply(function(x,y) {
  outlierplot(x,type="scatter",class.type="grade");
title(y)
},datas,names(datas))

par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
tmp<-mapply(function(x,y) {
  myCls2 <- OutlierClassifier1(x,alpha=0.05,type="all",corrected=TRUE)
  outlierplot(x,type="scatter",classifier=OutlierClassifier1,class.type="best",
             Legend=legend(1,1,levels(myCls),xjust=1,col=colcode,pch=pchcode),
pch=as.numeric(myCls2));
  legend(0.1,1,legend=levels(myCls2),pch=1:length(levels(myCls2)))
title(y)
},datas,names(datas))

par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="ecdf",main=names(datas)[i])
```
outlierplot

Plot various graphics to analyse outliers.

Description

A collection of plots emphasising different aspects of possible outliers.

Usage

outlierplot(X,...)
## S3 method for class 'acomp'
outlierplot(X,colcode=colorsForOutliers1,
pchcode=pchForOutliers1,
   type=c("scatter","biplot","dendrogram","ecdf","portion","nout","distdist"),
   legend.position,pch=19,...,clusterMethod="ward",
   myCls=classifier(X,alpha=alpha,type=class.type,corrected=corrected),
   classifier=OutlierClassifier1,
   alpha=0.05,
   class.type="best",
   Legend,pow=1,
   main=paste(deparse(substitute(X))),
   corrected=TRUE,robust=TRUE,princomp.robust=FALSE,
   mahRange=exp(c(-5,5))^pow,
   flagColor="red",
   meanColor="blue",
   grayColor="gray40",
   goodColor="green",
   mahalanobisLabel="Mahalanobis Distance"
)

Arguments

X The dataset as an acomp object

colcode A color palette for factor given by the myCls, or function to create it from the factor. Use colorForOutliers2 if class.method="all" is used.

pchcode A function to create a plot character palette for the factor returned by the myCls call
**type**

The type of plot to be produced. See details for more precise definitions.

**legend.position**

The location of the legend. Must!!! be given to draw a classical legend.

**pch**

A default plotting char

... Further arguments to the used plotting function

**clusterMethod**

The clustering method for `hclust` based outlier grouping.

**myCls**

A factor presenting the groups of outliers

**classifier**

The routine to create a factor presenting the groups of outliers heuristically. It is only used in the default argument to `myCls`.

**alpha**

The confidence level to be used for outlier classification tests

**class.type**

The type of classification that should be generated by `classifier`

**Legend**

The content will be substituted and stored as list entry `legend` in the result of the function. It can than be evaluated to actually create a separate legend on another device (e.g. for publications).

**pow**

The power of Mahalanobis distances to be used.

**main**

The title of the graphic

**corrected**

Literature typically proposes to compare the Mahalanobis distances with the distribution of a random Mahalanobis distance. However it would be needed to correct this for (dependent) multiple testing, since we always test the whole dataset, which means comparing against the distribution of the maximum Mahalanobis distance. This argument switches to this second behavior, giving less outliers.

**robust**

A robustness description as define in `robustnessInCompositions`

**princomp.robust**

Either a logical determining wether or not the principal component analysis should be done robustly or a principal component object for the dataset.

**mahRange**

The range of Mahalanobis distances displayed. This is fixed to make views comparable among datasets. However if the preset default is not enough a warning is issued and a red mark is drawn in the plot

**flagColor**

The color to draw critical situations.

**meanColor**

The color to draw typical curves.

**goodColor**

The color to draw confidence bounds.

**grayColor**

The color to draw less important things.

**mahalanobisLabel**

The axis label to be used for axes displaying Mahalanobis distances.

---

**Details**

See `outliersInCompositions` for a comprehensive introduction into the outlier treatment in compositions.

- **type**="scatter" Produces an appropriate standard plot such as a ternary diagram with the outliers marked by their codes according to the given classifier and colorcoding and pch coding. This shows the actual values of the identified outliers.
• **type**="biplot" Creates a biplot based on a nonrobust principal component analysis showing the outliers classified through outliers in the given color scheme. We use the nonrobust principal component analysis since it rotates according to a good visibility of the extreme values. This shows the position of the outliers in the usual principal components analysis. However note that a **coloredBiplot** is used rather than the usual one.

• **type**="dendrogram" Shows a dendrogram based on robust Mahalanobis distance based hierarchical clustering, where the observations are labeled with the identified outlier classes. This plot can be used to see how good different categories of outliers cluster.

• **type**="ecdf" This plot provides a cummulated distribution function of the Mahalanobis distances along with an expected curve and a lower confidence limit. The empirical cdf is plotted in the default color. The expected cdf is displayed in **meanColor**. The alpha-quantile – i.e. a lower prediction bound – for the cdf is given in **goodColor**. A line in **grayColor** shows the minimum portion of observations above some limit to be outliers, based on the portion of observations necessary to move down to make the empirical distribution function get above its lower prediction limit under the assumption of normality. This plot shows the basic construction for the minimal number of outlier computation done in **type**="portion".

• **type**="portion" This plot focusses on numbers of outliers. The horizontal axis give Mahalanobis distances and the vertical axis number of observations. In **meanColor** we see a curve of an estimated number of outliers above some limit, generated by estimating the portion of outliers with a Mahalanobis distance over the given limit by max(0,1-ecdf/cdf). The minimum number of outliers is computed by replacing cdf by its lower confidence limit and displayed in **goodColor**. The Mahalanobis distances of the individual data points are added as a stacked **stripchart**, such that the influence of individual observations can be seen.

The true problem of outlier detection is to detect "near" outliers. Near outliers are outliers so near to the dataset that they could well be extrem observation. These near outliers would provide no problem unless they are not many showing up in groups. Graphic allows at least to count them and to show there probable Mahalanobis distance such, however it still does not allow to conclude that an individual observation is an outlier. However still the outlier candidates can be identified comparing their mahalanobis distance (returned by the plot as $mahalanobis$) with a cutoff inferred from this graphic.

• **type**="nout" This is a simplification of the previous plot simply providing the number of outliers over a given limit.

?? MORE DOCUMENTATION NEEDED ??

• **type**="distdist" Plots a scatterplot of the the classical and robust Mahalanobis distance with the given classification for colors and plot symbols. Furthermore it plots a horizontal line giving the 0.95-Quantil of the distribution of the maximum robust Mahalanobis distance of normally distributed dataset.

Value

a list representing the criteria computed to create the plots. The content of the list depends on the plotting type selected.

Note

The package **robustbase** is required for using the robust estimations.
Author(s)


See Also

OutlierClassifier1, ClusterFinder1

Examples

## Not run:
data(SimulatedAmounts)
outlierplot(acomp(sa.outliers5))

datas <- list(data1=sa.outliers1, data2=sa.outliers2, data3=sa.outliers3, 
data4=sa.outliers4, data5=sa.outliers5, data6=sa.outliers6)

opar<-par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
tmp<-mapply(function(x,y) {
  outlierplot(x,type="scatter",class.type="grade");
  title(y)
},datas,names(datas))

par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
tmp<-mapply(function(x,y) {
  myCls2 <- OutlierClassifier1(x,alpha=0.05,type="all",corrected=TRUE)
  outlierplot(x,type="scatter",classifier=OutlierClassifier1,class.type="best",
    Legend=legend(1,1,levels(myCls),xjust=1,col=colcode,pch=pchcode),
    pch=as.numeric(myCls2));
  legend(0,1,legend=levels(myCls2),pch=1:length(levels(myCls2)))
  title(y)
},datas,names(datas))

# To slow
par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="ecdf",main=names(datas)[i])
par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="portion",main=names(datas)[i])
par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="nout",main=names(datas)[i])
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="distdist",main=names(datas)[i])
par(opar)

## End(Not run)
Description

The Philosophy behind outlier treatment in library(compositions).

Details

Outliers are omnipresent in all kinds of data analysis. To avoid catastrophic misinterpretations robust statistics has developed some methods to avoid the distracting influence of the outliers. The introduction of robust methods into the compositions package is described in robustnessInCompositions.

However sometimes we are interested directly in the analysis of outliers. The central philosophy of the the outlier classification subsystem in compositions is that outlier are in most cases not simply erroneous observations, but rather products of some systematic anomaly. This can e.g. be an error in an individual component, a secondary process or a minor undetected but different subpopulation. The package provides various concepts to investigate possible reasons for outliers in compositional datasets.

- Proven Outliers The package relies on an additive–lognormal reference distribution in the simplex (and the corresponding normal distribution in each other scale). The central tool for the detection of outliers is the Mahalanobis distance of the observation from a robustly estimated center based on a robustly estimated covariance. The robust estimation can be influenced by the given robust attributes. An outlier is considered as proven if its Mahalanobis distance is larger that the (1-alpha) quantile of the distribution of the maximum Mahalanobis distance of a dataset of the same size with a corresponding (additive)(log)normal distribution. This relies heavily on the presumption that the robust estimation is invariant under linear transformation, but make no assumptions about the actually used robust estimation method. The corresponding distributions are thus only defined with respect to a specific implementation of the robust estimation algorithm. See OutlierClassifier1(...,type="outlier"), outlierplot(...,type=c("scatter","biplot"),class.type="outlier"), qMaxMahalanobis(...).

- Extrem Values / Possible outliers Some cases of the dataset might have unusually high Mahalanobis distances, e.g. such that we would expect the probility of a random case to have such a value or higher might be below alpha. In Literature these cases are often rendered as outliers, because this level is approximated by the corresponding chisq-based criterion proposed. However we consider these only as extrem values, but however provide tools to detect and plot them. See OutlierClassifier1(...,type="grade"), outlierplot(...,type=c("scatter","biplot"),class.type="grade"), qEmpiricalMahalanobis(...).

- Single Component Outliers Some Outliers can be explained by a single component, e.g. because this single measurement error was wrong. These sort of outliers is detected when we reduce the dataset to a subcomposition with one component less and realise that our former outlier is now a fairly normal member of the dataset, maybe not even extrem. Thus a outlier is considered as as single component outlier, when it does not appear extrem in any of the subcompositions with one component less. For other outliers we can prove that they are
still extrem for all subcomposition with one component removed. Thus these have to be as multicomponent outliers, that can not be explained by a single measurement error. For remaining single component outliers, we can ask which component is able to explain the outlying character. See `OutlierClassifier1(...,type=c("best","type","all"))`.

- Counting hidden outliers If outliers are not outlying far enough to be detected by the test for outlyingness are only at first sight harmless. One outlier is within the reasonable bounds of what a normal distribution could have delivered should not harm the analysis and might not even detectable in any way. However if there is more than one they could act together to disrupt our analysis and more interestingly there might be some joint reason, which than might make them an interesting object of investigation in themselves. Thus the package provides methods (e.g. `outlierplot(...,type="portions"))`, to prove the existence of such outliers, to give a lower bound for there number and to provide us with suspects, with an associated outlyingness probability. See `outlierplot(...,type="portions"), outlierplot(...,type="nout"), pQuantileMahalanobis(...)`

- Finding atypical subpopulations When we assume smaller subpopulation we need a tool finding these clusters. However usual cluster analysis tends to ignore the subgroups, split the main mass and then associate the subgroups prematurely to the next part of the main mass. For this task we have developed special tools to find clusters of atypical populations clearly inducing secondary modes, without ripping apart the central nonoutlying mass. See `ClusterFinder1`.

- Identifying multiple distracting processes Outliers that are not due to a separate subpopulation or due to a single component error, might still belong together for being influenced by the same secondary process distorting the composition to a different degrees. Our proposal is to cluster the direction of the outliers from the center, e.g. by a command like: `take<-OutlierClassifier1(data,type="grade")!="ok" hc<-hclust(dist(normalize(acomp(scale(data))[take,])))` and to plot by a command like: `plot(hc)` and `plot(acomp(data[take,]),col=cutree(hc,1.5))`

With these tools we hope to provide a systematic approach to identify various types of outliers in an exploratory analysis.

### Note

The package `robustbase` is required for using the robust estimations and the outlier subsystem of compositions. To simplify installation it is not listed as required, but it will be loaded, whenever any sort of outlier detection or robust estimation is used.

### Author(s)


### References


### See Also

`compositions-package, missingsInCompositions, robustnessInCompositions, outliersInCompositions, outlierplot, OutlierClassifier1, ClusterFinder1`
Examples

## Not run:
# To slow
tmp<-set.seed(1400)
A <- matrix(c(0.1,0.2,0.3,0.1),nrow=2)
Mvar <- 0.1*ilrvar2clr(A%*%t(A))
Mcenter <- acomp(c(1,2,1))
typicalData <- rnorm.acomp(100,Mcenter,Mvar) # main population
colnames(typicalData)<-c("A","B","C")
data1 <- acomp(rnorm.acomp(100,Mcenter,Mvar))
data2 <- acomp(rbind(typicalData+rbinom(100,1,p=0.1)*rnorm(100)*acomp(c(4,1,1)))),
data3 <- acomp(rbind(typicalData,acomp(c(0.5,1.5,2))))
colnames(data3)<colnames(typicalData)
tmp<-set.seed(30)
rcauchy.acomp <- function (n, mean, var){
  D <- gsi.getD(mean)-1
  perturbe(ilrInv(matrix(rnorm(n*D)/rep(rnorm(n),D), ncol = D) %*% chol(clrvar2ilr(var))), mean)
}
data4 <- acomp(rcauchy.acomp(100,acomp(c(1,2,1)),Mvar/4))
colnames(data4)<colnames(typicalData)
data5 <- acomp(rbind(unclass(typicalData)+outer(rbinom(100,1,p=0.1)*runif(100),c(0.1,1,2))))
data6 <- acomp(rbind(unclass(typicalData),rnrom.acomp(20,acomp(c(4,4,1)),Mvar)))
datas <- list(data1=data1,data2=data2,data3=data3,data4=data4,data5=data5,data6=data6)
tmp <-c()
par(opar<-par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
outlierplot(x,type="scatter",class.type="grade";
title(y)
),datas,names(datas))

par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
tmp<-mapply(function(x,y) {
  myCls2 <- OutlierClassifier1(x,alpha=0.05,type="all",corrected=TRUE)
  outlierplot(x,type="scatter",classifier=OutlierClassifier1,class.type="best",
  Legend=legend(1,1,levels(myCls),xjust=1,col=colcode,pch=pchcode),
  pch=as.numeric(myCls2));
  legend(0,1,legend=levels(myCls2),pch=1:length(levels(myCls2)))
  title(y)
},datas,names(datas))

par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="ecdf",main=names(datas)[i])
par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="portion",main=names(datas)[i])
par(mfrow=c(2,3),pch=19,mar=c(3,2,2,1))
for( i in 1:length(datas) )
  outlierplot(datas[[i]],type="nout",main=names(datas)[i])
par(opar)
pairwiseplot

Creates a paneled plot like pairs for two different datasets.

Description

Creates a plot for each element of two lists or each column of each dataset against each of the second.

Usage

```r
pairwisePlot(X, Y, ...)  
## Default S3 method: 
pairwisePlot(X, Y = X, ...,  
  xlab = deparse(substitute(X)), ylab = deparse(substitute(Y)),  
  nm = c(length(Y), length(X)), panel = plot,  
  add.line = FALSE, line.col = 2, add.robust = FALSE, rob.col = 4)
```

Arguments

- `X`: a list, a data.frame, or a matrix representing the first set of things to be displayed.
- `Y`: a list, a data.frame, or a matrix representing the second set of things to be displayed.
- `...`: further parameters to the panel function
- `xlab`: The sequence of labels for the elements of `X`. Alternatively the labels can be given as colnames or names of `X`. This option takes precedence if specified.
- `ylab`: The sequence of labels for the elements of `Y`. Alternatively the labels can be given as colnames or names of `Y`. This option takes precedence if specified.
- `nm`: the parameter to be used in the call `par(mfrow = nm)`. If NULL no parameter is setted and a sequence of plots can be generated.
- `panel`: The panel function to plot the individual panels. If the panel function admits a formula interface, it is called as `panel(y ~ x, xlab = xlab, ylab = ylab, ...)`, otherwise as `panel(x, y, xlab = xlab, ylab = ylab, ...)`. Thus the panel function must be capable of taking these arguments. It must also set up its own plot. There is no negotiation on coordinate system.
- `add.line`: logical, to control the addition of a regression line in each panel
- `line.col`: in case the regression line is added, which color should be used? defaults to red.

### Example

```r
moreData <- acomp(rbind(data3, data5, data6))
take <- OutlierClassifier1(moreData, type = "grade") != "ok"
hc <- hclust(dist(normalize(acomp(scale(moreData)[take, ]))), method = "complete")
plot(hc)
plot(acomp(moreData[take, ]), col = cutree(hc, 1.5))
```

## End(Not run)
pairwiseplot

add.robust logical, to control the addition of a robust regression line in each panel. Ignored if covariable is a factor. This is nowadays based on `lmrob`, but this can change in the future.

rob.col in case the robust regression line is added, which color should be used? Defaults to blue.

Details

This is a light-weight convenience function to plot several aspects of one dataset against several aspects of another dataset. It is far more straight-forward than e.g. the `pairs` function and does not do any internal computation rather than organizing the names. Of course, the rows of the two data sets must be the same.

The current implementation may display a warning about the function `panel` dispatching methods for generic `plot`. It can be ignored without harm.

Optionally, classical and/or robust regression lines can be drawn, though only for non-factor covariables.

It may be convenient to use `par` capabilities to fit the device characteristics to the plot, in particular arguments `mar` and `oma`.

Author(s)


References


http://ima.udg.es/Activitats/CoDaWork03

http://ima.udg.es/Activitats/CoDaWork05

http://ima.udg.es/Activitats/CoDaWork08

See Also

`plot.aplus`, `balance`, `pwlrPlot`

Examples

```r
X <- rnorm(100)
Y <- rnorm.acomp(100, acomp(c(A=1, B=1, C=1))*0.1*diag(3))+acomp(t(outer(c(0.2, 0.3, 0.4), X,"^"))

pairs(cbind(ilr(Y), X), panel=function(x, y,...) {points(x, y,...); abline(lm(y~x))})
pairs(cbind(balance(Y,~A/B/C), X),
     panel=function(x, y,...) {points(x, y,...); abline(lm(y~x))})
pairwisePlot(balance(Y,~A/B/C), X)
pairwisePlot(X, balance(Y, ~A/B/C),
             panel=function(x, y,...) {plot(x, y,...); abline(lm(y~x))})
```
parametricMat

Unique parametrisations for matrices.

Description

Helper functions to parametrize positive semidefinite matrices in multivariate variogram models.

Usage

parametricRank1Mat(p)
parametricPosdefMat(p)
parameterRank1Mat(A)
parameterPosdefMat(A)
parametricRank1ClrMat(p)
parametricPosdefClrMat(p)
parameterRank1ClrMat(A)
parametricPosdefClrMat(A)
Arguments

A    a positiv definit matrix of the given type
p    a vector of parameters describing the matrix, as returned by the parameter functions.

Details

The rank 1 matrix is parametrised by the first eigenvector scaled by the square root of the eigenvalue. The positiv semidefinit matrix the entries of a upper right triangular matrix R with $t(R)\%\%R=A$. The clr matrices are work with the parameters of the corresponding ilr matrix.

Value

A or p, depending on what is not given.

Author(s)


See Also

vgram2lrgram, CompLinModCoReg, vgmFit

Examples

parametricRank1Mat(c(0,0,2))
parametricPosdefMat(c(0,0,1,0,0,0))
parameterRank1Mat(matrix(1,nr=3,nc=3))
parameterPosdefMat(diag(5))

perturbe  Perturbation of compositions

Description

The perturbation is the addition operation in the Aitchison geometry of the simplex.

Usage

perturbe(x,y)
## Methods for class "acomp"
## x + y
## x - y
## - x
Arguments

x compositions of class acomp
y compositions of class acomp

Details

The perturbation is the basic addition operation of the Aitchison simplex as a vector space. It is defined by:

\[(x + y)_i = clo((x_i y_i)_i)\]

and as unary operation respectively as:

\[(-x)_i = clo((1/y_i)_i)\]

Value

An acomp vector or matrix.

Author(s)


References


See Also

acomp, *.aplus, *.rplus

Examples

tmp <- -acomp(1:3)
tmp + acomp(1:3)
plot.acomp

Ternary diagrams

Description

Displaying compositions in ternary diagrams

Usage

## S3 method for class 'acomp'
plot(x,...,labels=names(x),
    aspanel=FALSE,id=FALSE,idlabs=NULL,idcol=2,center=FALSE,
    scale=FALSE,pca=FALSE,col.pca=par("col"),margin="acomp",
    add=FALSE,triangle=!add,col=par("col"),axes=FALSE,
    plotMissings=TRUE,
    lenMissingTck=0.05,colMissingTck="red",
    mp=simpleMissingSubplot(c(0,1,0.95,1),
        missingInfo,c("NM","TM",cn)),
    robust=getOption("robust"))

## S3 method for class 'rcomp'
plot(x,...,labels=names(x),
    aspanel=FALSE,id=FALSE,idlabs=NULL,idcol=2,center=FALSE,
    scale=FALSE,pca=FALSE,col.pca=par("col"),margin="rcomp",
    add=FALSE,triangle=!add,col=par("col"),axes=FALSE,
    plotMissings=TRUE,
    lenMissingTck=0.05,colMissingTck="red",
    mp=simpleMissingSubplot(c(0,1,0.95,1),
        missingInfo,c("NM","TM",cn)),
    robust=getOption("robust"))

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

Arguments

x        a dataset of a compositional class
...

margin    the type of marginalisation to be computed, when displaying the individual panels. Possible values are: "acomp", "rcomp" and any of the variable names/column numbers in the composition. If one of the columns is selected each panel displays a subcomposition given by the row part, the column part and the given part. If one of the classes is given the corresponding margin acompmargin or rcompmargin is used.

add        a logical indicating whether the information should just be added to an existing plot. If FALSE a new plot is created

triangle    a logical indicating whether the triangle should be drawn
col
labels
aspanel
id
idlabs
idcol
center
scale
pca
col.pca
axes
plotMissings
lenMissingTck
colMissingTck
mp
robust

the color to plot the data
the names of the parts
logical indicating that only a single panel should be drawn and not the whole
plot. Internal use only
logical, if TRUE one can identify the points like with the identify command.
a character vector providing the labels to be used with the identification, when
id=TRUE
color of the idlabs labels
a logical indicating whether a the data should be centered prior to the plot. Cen-
tering is done in the choosen geometry. See scale
a logical indicating whether a the data should be scaled prior to the plot. Scaling
is done in the choosen geometry. See scale
a logical indicating whether the first principal component should be displayed
in the plot. Currently, the direction of the principal component of the displayed
subcomposition is displayed as a line. In a future, the projected principal com-
ponent of the whole dataset should be displayed.
The color to draw the principal component.
Either a logical wether to plot the axes, or numerical enumerating the axes sides
to be used e.g. 1 for only plotting the lower axes, or a list of parameters to
ternaryAxis.
logical indicating that missingness should be represented graphically. Compo-
nentes with one missing subcomponent in the plot are represented by tickmarks
at the three axis. Components with two or three missing components are only
represented in a special panel drawn according to the mp parameter if missings
are present. Missings of type BDL (below detection limit) are always plotted,
even if plotMissings is false, but in this case this fact is not specially marked.
In rcomp geometry an actuall 0 in the data is never treated as missing.
length of the tick-marks to be plotted for missing values. If 0 no tickmarks
are plotted. Negative lengths point outside. length 1 draws right through to
the opposit corner. Missing ticks in acomp geometry are inclined showing the
line of possible values in acomp geometry. Missingticks in rcomp-geometry
are vertical to the axis representing the fact that only the other component is
unknown. That these lines can leave the plot is one of the odd consequences
of rcomp geometry.
colors to draw the missing tick-marks. NULL means to take the colors specified
for the observations.
A formula providing a call to a function plotting informations on the missings.
The call is evaluated in the environment of the panel plotting function and has
access (among others) to: cn the names of the components in the current plot, x
the dataset of the current plot, y the transformed dataset, (c60,s60) coordinates
of the upper vertex of the triangle. missingInfo is a table giving the number
of observations of the types NM=Non Missing, TM= Totally missing (i.e. at
least two components of the subcomposition are missing), and the three single
component missing possibilities for the three components.
A robustness description. See robustnessInCompositions for details. The option
is used for centering, scaling and principle components.
Details

The data is displayed in ternary diagrams. Thus, it does not work for two-part compositions. Compositions of three parts are displayed in a single ternary diagram. For compositions of more than three components, the data is arranged in a scatterplot matrix through the command `pairs`. In this case, the third component in each of the panels is chosen according to setting of `margin=`. Possible values of `margin=` are: "acompx", "rcomp" and any of the variable names/column numbers in the composition. If one of the columns is selected each panel displays a subcomposition given by the row part, the column part and the given part. If one of the classes is given the corresponding margin `acompmargin` or `rcompmargin` is used.

Ternary diagrams can be read in multiple ways. Each corner of the triangle corresponds to an extreme composition containing only the part displayed in that corner. Points on the edges correspond to compositions containing only the parts in the adjacent corners. The relative amounts are displayed by the distance to the opposite corner (so-called barycentric coordinates). The individual portions of any point can be inferred by drawing a line through the investigated point, and parallel to the edge opposite to the corner of the part of interest. The portion of this part is constant along the line. Thus we can read it on the sides of the ternary diagram, where the line crosses its borders. Note that these `isoPortionLines` remain straight under an arbitrary perturbation.

ccomp ternary diagrams are always jittered to avoid overplotting.

Author(s)


References


http://ima.udg.es/Activitats/CoDaWork03

http://ima.udg.es/Activitats/CoDaWork05

See Also

`plot.aplus`, `plot3D` (for 3D plot), `kingTetrahedron` (for 3D-plot model export), `qqnorm.acomp`, `boxplot.acomp`
Examples

```r
data(SimulatedAmounts)
plot(acomp(sa.lognormals))
plot(acomp(sa.lognormals),axes=TRUE)
plot(rcomp(sa.lognormals))
plot(rcomp(sa.lognormals5))
plot(acomp(sa.lognormals5),pca=TRUE,col.pca="red")
plot(rcomp(sa.lognormals5),pca=TRUE,col.pca="red",axes=TRUE)
```

plot.aplus Displaying amounts in scatterplots

Description

This function displays multivariate unclosed amount datasets classes "aplus" and "rplus" in a way respecting the chosen geometry eventually in log scale.

Usage

```r
# S3 method for class 'aplus'
plot(x,...,labels=colnames(X),cn=colnames(X),
     aspanel=FALSE,id=FALSE,idlabs=NULL,idcol=2,
     center=FALSE,scale=FALSE,pca=FALSE,col.pca=par("col"),
     add=FALSE,logscale=TRUE,xlim=NULL,ylim=xlim,
     col=par("col"),plotMissings=TRUE,
     lenMissingTck=0.05,colMissingTck="red",
     mp=~simpleMissingSubplot(missingPlotRect,missingInfo,
                              c("NM","TM",cn)),
     robust=getOption("robust"))

# S3 method for class 'rplus'
plot(x,...,labels=colnames(X),cn=colnames(X),
     aspanel=FALSE,id=FALSE,idlabs=NULL,idcol=2,
     center=FALSE,scale=FALSE,pca=FALSE,col.pca=par("col"),
     add=FALSE,logscale=FALSE,
     xlim=NULL,ylim=xlim,col=par("col"),plotMissings=TRUE,
     lenMissingTck=0.05,colMissingTck="red",
     mp=~simpleMissingSubplot(missingPlotRect,missingInfo,
                              c("NM","TM",cn)),
     robust=getOption("robust"))

# S3 method for class 'rmult'
plot(x,...,labels=colnames(X),cn=colnames(X),
     aspanel=FALSE,id=FALSE,idlabs=NULL,idcol=2,
     center=FALSE,scale=FALSE,pca=FALSE,col.pca=par("col"),
     add=FALSE,logscale=FALSE,col=par("col"),
     robust=getOption("robust"))
```
Arguments

- **x**
  - a dataset with class aplus, rplus or rmult

- **...**
  - further graphical parameters passed (see `par`)

- **add**
  - a logical indicating whether the information should just be added to an existing plot. If FALSE, a new plot is created

- **col**
  - the color to plot the data

- **plotMissings**
  - logical indicating that missingness should be represented graphically. Components with one missing subcomponent in the plot are represented by tickmarks at the two axes. Cases with two missing components are only represented in a special panel drawn according to the `mp` parameter if missings are present. Missings of type BDL (below detection limit) are always plotted in nonlogarithmic plots, even if `plotMissings` is false, but in this case this fact is not specially marked.

- **lenMissingTck**
  - length of the tick-marks (in portion of the plotting region) to be plotted for missing values. If 0 no tickmarks are plotted. Negative lengths point outside of the plot. A length of 1 runs right through the whole plot.

- **colMissingTck**
  - colors to draw the missing tick-marks. NULL means to take the colors specified for the observations.

- **mp**
  - A formula providing a call to a function plotting informations on the missings. The call is evaluated in the environment of the panel plotting function and has access (among others) to: `cn` the names of the components in the current plot, `x` the dataset of the current plot, `missingInfo` is a table giving the number of observations of the types NM=Non Missing, TM=Totally missing (i.e. two components of the subcomposition are missing), and the two single component missing possibilities.

- **labels**
  - the labels for names of the parts

- **cn**
  - the names of the parts to be used in a single panel. Internal use only

- **aspanel**
  - logical indicating that only a single panel should be drawn and not the whole plot. Internal use only

- **id**
  - a logical. If TRUE one can identify the points like with the `identify` command

- **idlabs**
  - A character vector providing the labels to be used with the identification, when `id=TRUE`

- **idcol**
  - color of the `idlabs` labels

- **center**
  - a logical indicating whether the data should be centered prior to the plot. Centering is done in the chosen geometry. See `scale`

- **scale**
  - a logical indicating whether the data should be scaled prior to the plot. Scaling is done in the chosen geometry. See `scale`

- **pca**
  - a logical indicating whether the first principal component should be displayed in the plot. Currently, the direction of the principal component of the displayed subcomposition is displayed as a line. In a future, the projected principal component of the whole dataset should be displayed.

- **col.pca**
  - the color to draw the principal component.

- **logscale**
  - logical indicating whether a log scale should be used
xlim
2xncol(x)-matrix giving the xlims for the columns of x

ylim
2xncol(x)-matrix giving the ylims for the columns of x

robust
A robustness description. See `robustnessInCompositions` for details. The option is used for centering, scaling and principle components.

Details
TO DO: fix pca bug

Author(s)

See Also
plot.aplus, qqnorm.acomp.boxplot.acomp

Examples

data(SimulatedAmounts)
plot(aplus(sa.lognormals))
plot(rplus(sa.lognormals))
plot(aplus(sa.lognormals5))
plot(rplus(sa.lognormals5))

plot3D

3-dimensional plots, which can be rotated and zoomed in/out

Description
plot in 3D based on rgl

Usage

plot3D(x,...)

Arguments

x
an object to be plotted, e.g. a data frame or a data matrix

... additional plotting parameters as described in `rgl.material`

add
logical, adding or new plot

bbox
logical, whether to add a bounding box

axes
logical, whether to plot an axes of coordinates

cex
size of the plotting symbol

size
size of the plotting symbol, only size or cex should be used

col
the color used for dots, defaults to black.
Details

The function provides a generic interface for 3-dimensional plotting in analogy to the 2d-plotting interface of plot, using rgl package.

Value

the 3D plotting coordinates of the objects displayed, returned invisibly

Author(s)


See Also

points3d, plot, plot3D.rmult, plot3D.acomp, plot3D.rcomp, plot3D.aplus, plot3D.rplus

Examples

```r
x <- cbind(rnorm(10), rnorm(10), rnorm(10))
plot3D(x)
data(SimulatedAmounts)
plot3D(sa.lognormals, cex=4, col=1:nrow(sa.lognormals))
```

plot3Dacomp

3D-plot of compositional data

Description

3D-plot of compositional data. The plot is mainly an exploratory tool, not intended for exact display of data.

Usage

```r
## S3 method for class 'acomp'
plot3D(x, parts=1:min(ncol(X),4), ..., lwd=2, axis.col="gray", add=FALSE, cex=2, vlabs=colnames(x), vlabs.col=axis.col, center=FALSE, scale=FALSE, log=FALSE, bbox=FALSE, axes=TRUE, size=cex, col=1)

## S3 method for class 'rcomp'
plot3D(x, parts=1:min(ncol(X),4), ..., lwd=2, axis.col="gray", add=FALSE, cex=2, vlabs=colnames(x), vlabs.col=axis.col, center=FALSE, scale=FALSE, log=FALSE, bbox=FALSE, axes=TRUE, size=cex, col=1)
```
Arguments

- **x**: an aplus object to be plotted
- **parts**: a numeric xor character vector of length 3 coding the columns to be plotted
- **...**: additional plotting parameters as described in `rgl.material`
- **add**: logical, adding or new plot
- **cex**: size of the plotting symbols
- **lwd**: line width
- **axis.col**: color of the axis
- **vlabs**: the column names to be plotted, if missing defaults to the column names of the selected columns of X
- **vlabs.col**: color of the labels
- **center**: logical, should the data be centered
- **scale**: logical, should the data be scaled
- **log**: logical, indicating whether to plot in log scale
- **bbox**: logical, whether to add a bounding box
- **axes**: logical, whether plot a coordinate cross
- **size**: size of the plotting symbols
- **col**: the color used for dots, defaults to black.

Details

The routine behaves different when 3 or four components should be plotted. In case of four components:
- If log is TRUE the data is plotted in `ilr` coordinates. This is the isometric view of the data.
- If log is FALSE the data is plotted in `ipt` coordinates and a tetrahedron is plotted around it if `coors` == TRUE. This can be used to do a tetrahedron plot.

In case of three components:
- If log is TRUE the data is plotted in `clr` coordinates. This can be used to visualize the clr plane.
- If log is FALSE the data is plotted as is, showing the embedding of the three-part simplex in the three-dimensional space.

In all cases: If `coors` is true, coordinate arrows are plotted of length 1 in the origin of the space, except in the tetrahedron case.

Value

Called for its side effect of a 3D plot of an acomp object in an rgl plot. It invisibly returns the 3D plotting coordinates of the objects displayed

Note

The function `kingTetrahedron` provides an alternate way of tetrahedron plots, based on a more advanced viewer, which must be downloaded separately.
plot3Daplus

Author(s)

See Also
kingTetrahedron points3d.plot3D.plot.plot3D.rmult.
plot3D.acomp.plot3D.rcomp., plot3D.aplus.plot3D.rplus

Examples

data(SimulatedAmounts)
plot3D(acomp(sa.lognormals5),1:3,col="green")
plot3D(acomp(sa.lognormals5),1:3,log=TRUE,col="green")
plot3D(acomp(sa.lognormals5),1:4,col="green")
plot3D(acomp(sa.lognormals5),1:4,log=TRUE,col="green")

plot3Daplus 3D-plot of positive data

Description

3D-plot of positive data typically in log-log-log scale. The plot is mainly an exploratory tool, and not intended for exact display of data.

Usage

## S3 method for class 'aplus'
plot3D(x,parts=1:3,...,
      vlabs=NULL,add=FALSE,log=TRUE,bbox=FALSE,axes=TRUE,col=1)

Arguments

x an aplus object to be plotted
parts a numeric xor character vector of length 3 coding the columns to be plotted
... additional plotting parameters as described in rgl.material
add logical, adding or new plot
vlabs the column names to be plotted, if missing defaults to the column names of the selected columns of X
log logical, indicating wether to plot in log scale
bbox logical, whether to add a bounding box
axes logical, plot a coordinate system
col the color used for dots, defaults to black.
Details

If `log` is TRUE the data is plotted in `ilt` coordinates. If `coors` is true, coordinate arrows are plotted of length 1 and in the (aplus-)mean of the dataset. If `log` is FALSE the data is plotted with `plot.rplus`

Value

Called for its side effect of a 3D plot of an aplus object in an rgl plot. It invisibly returns the 3D plotting coordinates of the objects displayed

Author(s)


See Also

`points3d`, `plot3D`, `plot`, `plot3D.rmult`, `plot3D.acomp`, `plot3D.rcomp`, `plot3D.aplus`, `plot3D.rplus`

Examples

```r
data(SimulatedAmounts)
plot3D(aplus(sa.lognormals),size=2)
```

plot3Drmult  plot in 3D based on rgl

Description

3-dimensional plots, which can be rotated and zoomed in/out

Usage

```r
## S3 method for class 'rmult'
plot3D(x,parts=1:3,..., 
       center=FALSE,scale=FALSE,add=FALSE,axes=!add, 
       cex=2,vlabs=colnames(x),size=cex,bbox=FALSE,col=1)
```

Arguments

- `x`  an object to be plotted, e.g. a data frame or a data matrix
- `parts`  the variables in the rmult object to be plotted
- `...`  additional plotting parameters as described in `rgl.material`
- `center`  logical, center the data? This might be necessary to stay within the OpenGL-arithmetic used in rgl.
scale logical, scale the data? This might be necessary to stay within the openGL-arithmetic used in rgl.
add logical, adding or new plot
bbox logical, whether to add a bounding box
axes logical, whether to plot a coordinate cross
cex size of the plotting symbol (as expanding factor)
vlabs labels for the variables
size size of the plotting symbol, only size or cex should be used
col the color used for dots, defaults to black.

Details
The function provides a generic interface for 3-dimensional plotting in analogy to the 2d-plotting interface of plot, using rgl package.

Value
the 3D plotting coordinates of the objects displayed, returned invisibly

Author(s)

See Also
points3d, plot, plot3D.rmult,
plot3D.acomp, plot3D.rcomp, plot3D.aplus, plot3D.rplus

Examples
x <- cbind(rnorm(10), rnorm(10), rnorm(10))
plot3D(x)
data(SimulatedAmounts)
plot3D(rmult(sa.lognormals), cex=4, col=1:nrow(sa.lognormals))
Arguments

- **x**: an rplus object to be plotted
- **parts**: the variables in the rplus object to be plotted
- **...**: additional plotting parameters as described in `rgl.material`
- **vlabs**: the labels used for the variable axes
- **add**: logical, adding or new plot
- **bbox**: logical, whether to add a bounding box
- **cex**: size of the plotting symbol (as character expansion factor)
- **size**: size of the plotting symbol, only size or cex should be used
- **axes**: logical, whether to plot a coordinate cross
- **col**: the color used for dots, defaults to black.

Details

The function plots rplus objects in a 3D coordinate system, in an rgl plot.

Value

the 3D plotting coordinates of the objects displayed, returned invisibly

Author(s)


See Also

- `points3d`, `plot3D.rmult`
- `plot3D.acomp`, `plot3D.rcomp`, `plot3D.aplus`, `plot3D`

Examples

```r
x <- cbind(rnorm(10), rnorm(10), rnorm(10))
plot3D(rplus(exp(x)))
data(SimulatedAmounts)
plot3D(rplus(sa.lognormals), cex=4, col=1:nrow(sa.lognormals))
```
**plotlogratioVariogram**  
*Empirical variograms for compositions*

**Description**

Plots a logratioVariogram.

**Usage**

```r
## S3 method for class 'logratioVariogram'
plot(x, ..., type="l", lrvg=NULL, 
fcols=2:length(lrvg), oma=c(4, 4, 4, 4), gap=0, ylim=NULL)
```

**Arguments**

- `x`  
The logratioVariogram created by `logratioVariogram`
- `...`  
further parameters for `plot.default`
- `type`  
as in `plot.default`
- `lrvg`  
a model function for a logratiovariogram or a list of several, to be added to the plot.
- `fcols`  
the colors for the different lrvg variograms
- `oma`  
The outer margin of the paneled plot
- `gap`  
The distance of the plot panals used to determin mar
- `ylim`  
The limits of the Y-axis. If zero it is automatically computed.

**Details**

see `logratioVariogram`

**Value**

Nothing.

**Author(s)**


**See Also**

`vgram2lrvg`, `CompLinModCoReg`
Examples

```r
## Not run:
data(juraset)
X <- with(juraset,cbind(X,Y))
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
lrv <- logratioVariogram(comp,X,maxdist=1,nbins=10)
fff <- CompLinModCoReg(~nugget()+sph(0.5)+R1*exp(0.7),comp)
fit <- vgmFit(lrv,fff)
fit
fff(1:3)
plot(lrv,lrvg=vgm2lrvgram(fit$vg))

## End(Not run)
```

plotmissingsummary  Plot a Missing Summary

Description

Plots a missing summary as a barplot

Usage

```r
## S3 method for class 'missingSummary'
plot(x,...,main="Missings",legend.text=TRUE,
col=c("gray","lightgray","yellow","red","white","magenta"))
as.missingSummary(x,...)
```

Arguments

- `x` a missingSummary table with columns representing different types of missing
- `...` further graphical parameters to barplot
- `main` as in `barplot`
- `legend.text` as in `barplot`
- `col` as in `barplot`

Details

The different types of missings are drawn in quasi-self-understandable colors: normal gray for NMV, and lightgray as for BDT (since they contain semi-numeric information), yellow (slight warning) for MAR, red (serious warning) for MNAR, white (because they are non-existing) for SZ, and magenta for the strange case of errors.

Value

called for its side effect. The return value is not defined.
Description

Yat, yee, sam measurements (in meters) for the final jumps of the 1985 Honk Kong Pogo-Jumps Championship, 4 jumps of the 7 finalists.

Usage

data(PogoJump)

Details

The data consist of 28 cases: 4 jumps of the 7 finalists, and 4 variables: Yat, Yee, Sam measurements in meters, and finalist – 1 to 7.

Pogo-Jumps is similar to the triple jump except that the competitor is mounted on a pogo-stik. After a pogo-up towards the starting board the total jump distance achieved in three consecutive bounces, known as the yat, yee and sam, is recorded.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name HKPOGO.DAT, here included under the GNU Public Library Licence Version 2 or newer.
powerofpsdmatrix

power transform of a matrix

Description

Computes the power of a positive semi-definite symmetric matrix.

Usage

powerofpsdmatrix( M, p,...)

Arguments

M  
  a matrix, preferably symmetric

p  
  a single number giving the power

...  
  further arguments to the singular value decomposition

Details

for a symmetric matrix the computed result can actually be considered as a version of the given power of the matrix fullfilling the relation:

\[ M^p M^q = M^{p+q} \]

The symmetry of the matrix is not checked.

Value

\( \% \% D^p \% \% t(P) \) where the UDP is the singular value decomposition of M.

Author(s)


Examples

data(SimulatedAmounts)
d <- ilr(sa.lognormals)
var( d %*% powerofpsdmatrix(var(d),-1/2)) # Unit matrix
Description

A principal component analysis is done in the Aitchison geometry (i.e. clr-transform) of the simplex. Some gimmicks simplify the interpretation of the computed components as compositional perturbations.

Usage

```r
## S3 method for class 'acomp'
princomp(x,...,scores=TRUE,center=attr(covmat,"center"),
  covmat=var(x,robust=robust,giveCenter=TRUE),
  robust=getOption("robust"))
## S3 method for class 'princomp.acomp'
print(x,...)
## S3 method for class 'princomp.acomp'
plot(x,y=NULL,...,npcs=min(10,length(x$sdev)),
  type=c("screeplot","variance","biplot","loadings","relative"),
  main=NULL,scale.sdev=1)
## S3 method for class 'princomp.acomp'
predict(object,newdata,...)
```

Arguments

- **x**: a acomp-dataset (in princomp) or a result from princomp.acomp
- **y**: not used
- **scores**: a logical indicating whether scores should be computed or not
- **npcs**: the number of components to be drawn in the scree plot
- **type**: type of the plot: "screeplot" is a lined screeplot, "variance" is a boxplot-like screeplot, "biplot" is a biplot, "loadings" displays the loadings as a `barplot.acomp`
- **scale.sdev**: the multiple of sigma to use plotting the loadings
- **main**: title of the plot
- **object**: a fitted princomp.acomp object
- **newdata**: another compositional dataset of class acomp
- **...**: further arguments to pass to internally-called functions
- **covmat**: provides the covariance matrix to be used for the principle component analysis
- **center**: provides the be used for the computation of scores
- **robust**: Gives the robustness type for the calculation of the covariance matrix. See `robustnessInCompositions` for details.
Details

As a metric euclidean space the Aitchison simplex has its own principal component analysis, that should be performed in terms of the covariance matrix and not in terms of the meaningless correlation matrix.

To aid the interpretation we added some extra functionality to a normal princomp(clr(x)). First of all the result contains as additional information the compositional representation of the returned vectors in the space of the data: the center as a composition Center, and the loadings in terms of a composition to perturbe with, either positively (Loadings) or negatively (DownLoadings). The Up- and DownLoadings are normalized to the number of parts in the simplex and not to one to simplify the interpretation. A value of about one means no change in the specific component. To avoid confusion the meaningless last principal component is removed.

The plot routine provides screeplots (type = "s", type = "v"), biplots (type = "b"), plots of the effect of loadings (type = "b") in scale.sdev*sdev-spread, and loadings of pairwise (log-)ratios (type = "r").

The interpretation of a screeplot does not differ from ordinary screeplots. It shows the eigenvalues of the covariance matrix, which represent the portions of variance explained by the principal components.

The interpretation of the biplot strongly differs from a classical one. The relevant variables are not the arrows drawn (one for each component), but rather the links (i.e., the differences) between two arrow heads, which represents the log-ratio between the two components represented by the arrows. The compositional loading plot is introduced with this package. The loadings of all component can be seen as an orthogonal basis in the space of clr-transformed data. These vectors are displayed by a barplot with their corresponding composition. For a better interpretation the total of these compositions is set to the number of parts in the composition, such that a portion of one means no effect. This is similar to (but not exactly the same as) a zero loading in a real principal component analysis.

The loadings plot can work in two different modes: if scale.sdev is set to NA it displays the composition being represented by the unit vector of loadings in the clr-transformed space. If scale.sdev is numeric we use this composition scaled by the standard deviation of the respective component. The relative plot displays the relativeLoadings as a barplot. The deviation from a unit bar shows the effect of each principal component on the respective ratio.

Value

princomp gives an object of type c("princomp.acomp", "princomp") with the following content:

- **sdev**: the standard deviation of the principal components
- **loadings**: the matrix of variable loadings (i.e., a matrix which columns contain the eigenvectors). This is of class "loadings". The last eigenvector is removed since it should contain the irrelevant scaling.
- **center**: the clr-transformed vector of means used to center the dataset
- **Center**: the acomp vector of means used to center the dataset
- **scale**: the scaling applied to each variable
- **n.obs**: number of observations
- **scores**: if scores = TRUE, the scores of the supplied data on the principal components. Scores are coordinates in a basis given by the principal components and thus not compositions
- **call**: the matched call
na.action not clearly understood
Loadings compositions that represent a perturbation with the vectors represented by the loadings of each of the factors
DownLoadings compositions that represent a perturbation with the inverse of the vectors represented by the loadings of each of the factors

predict returns a matrix of scores of the observations in the newdata dataset. The other routines are mainly called for their side effect of plotting or printing and return the object x.

Author(s)

References

Aitchison, J. Barcel’o-Vidal, J.J. Egozcue, V. Pawlowsky-Glahn (2002) A concise guide to the algebraic geometric structure of the simplex, the sample space for compositional data analysis, Terra Nostra, Schriften der Alfred Wegener-Stiftung, 03/2003


http://ima.udg.es/Activitats/CoDaWork03

http://ima.udg.es/Activitats/CoDaWork05

See Also
clr.acomp, relativeLoadings princomp.aplus, princomp.rcomp, barplot.acomp, mean.acomp, var.acomp

Examples

data(SimulatedAmounts)
pc <- princomp(acomp(sa.lognormals5))
pc
summary(pc)
plot(pc) #plot(pc,type="screeplot")
plot(pc, type="v")
plot(pc, type="biplot")
plot(pc, choice=c(1,3), type="biplot")
plot(pc, type="loadings")
plot(pc, type="loadings", scale.sdev=-1) # Downward
plot(pc, type="relative", scale.sdev=NA) # The directions
plot(pc, type="relative", scale.sdev=1) # one sigma Upward
plot(pc, type="relative", scale.sdev=-1) # one sigma Downward
biplot(pc)
screepplot(pc)
loadings(pc)
relativeLoadings(pc,mult=FALSE)
relativeLoadings(pc)
relativeLoadings(pc,scale.sdev=1)
relativeLoadings(pc,scale.sdev=2)

pc$Loadings
pc$DownLoadings
barplot(pc$Loadings)
pc$sdev^2
cov(predict(pc,sa.lognormals5))

princomp.aplus  Principal component analysis for amounts in log geometry

Description
A principal component analysis is done in the Aitchison geometry (i.e. ilt-transform). Some gimpics simplify the interpretation of the computed components as perturbations of amounts.

Usage
## S3 method for class 'plus'
princomp(x,...,scores=TRUE,center=attr(covmat,"center"),
covmat=var(x,robust=robust,giveCenter=TRUE),
        robust=getOption("robust")
## S3 method for class 'princomp.aplus'
print(x,...)
## S3 method for class 'princomp.aplus'
plot(x,y=NULL,..., npcs=min(10,length(x$sdev)),
       type=c("screeplot","variance","biplot","loadings","relative"),
       main=NULL,scale.sdev=1)
## S3 method for class 'princomp.aplus'
predict(object,newdata,...)

Arguments
x an aplus dataset (for princomp) or a result from princomp.aplus
y not used
scores a logical indicating whether scores should be computed or not
npcs the number of components to be drawn in the scree plot
type type of the plot: "screeplot" is a lined screeplot, "variance" is a boxplot-like screeplot, "biplot" is a biplot, "loadings" displays the loadings as a barplot.acomp
scale.sdev the multiple of sigma to use when plotting the loadings
main title of the plot
princomp.aplus

Object

princomp.aplus is a fitted princomp.aplus object.

Arguments

- newdata: another amount dataset of class aplus
- ...: further arguments to pass to internally-called functions
- covmat: provides the covariance matrix to be used for the principle component analysis
- center: provides the be used for the computation of scores
- robust: gives the robustness type for the calculation of the covariance matrix. See var.rmult for details.

Details

As a metric euclidean space, the positive real space described in aplus has its own principal component analysis, that can be performed either in terms of the covariance matrix or the correlation matrix. However, since all parts in a composition or in an amount vector share a natural scaling, they do not need the standardization (which in fact would produce a loss of important information). For this reason, princomp.aplus works on the covariance matrix.

To aid the interpretation we added some extra functionality to a normal princomp(ilt(x)). First of all the result contains as additional information the amount representation of returned vectors in the space of the data: the center as an amount Center, and the loadings in terms of amounts to perturbe with, either positively (Loadings) or negatively (DownLoadings). The Up- and DownLoadings are normalized to the number of parts and not to one to simplify the interpretation. A value of about one means no change in the specific component.

The plot routine provides screeplots (type = "s", type = "v"), biplots (type = "b"), plots of the effect of loadings (type = "b") in scale.sdev*sdev-sdev-spread, and loadings of pairwise (log-)ratios (type = "r").

The interpretation of a screeplot does not differ from ordinary screeplots. It shows the eigenvalues of the covariance matrix, which represent the portions of variance explained by the principal components.

The interpretation of the biplot uses, additionally to the classical one, a compositional concept: The differences between two arrowheads can be interpreted as log-ratios between the two components represented by the arrows.

The amount loading plot is introduced with this package. The loadings of all component can be seen as an orthogonal basis in the space of ilt-transformed data. These vectors are displayed by a barplot with their corresponding amounts. A portion of one means no change of this part. This is equivalent to a zero loading in a real principal component analysis.

The loadings plot can work in two different modes. If scale.sdev is set to NA it displays the amount vector being represented by the unit vector of loadings in the ilt-transformed space. If scale.sdev is numeric we use this amount vector scaled by the standard deviation of the respective component.

The relative plot displays the relativeLoadings as a barplot. The deviation from a unit bar shows the effect of each principal component on the respective ratio. The interpretation of the ratios plot may only be done in an Aitchison-compositional framework (see princomp.acomp).

Value

princomp gives an object of type c("princomp.acomp", "princomp") with the following content:

- sdev: the standard deviation of the principal components
- loadings: the matrix of variable loadings (i.e., a matrix which columns contain the eigenvectors). This is of class "loadings".
center  the ilt-transformed vector of means used to center the dataset
Center  the aplus vector of means used to center the dataset
scale  the scaling applied to each variable
n.obs  number of observations
scores if scores = TRUE, the scores of the supplied data on the principal components.
        Scores are coordinates in a basis given by the principal components and thus not
        compositions
call  the matched call
na.action  not clearly understood
Loadings  vectors of amounts that represent a perturbation with the vectors represented by
          the loadings of each of the factors
DownLoadings  vectors of amounts that represent a perturbation with the inverses of the vectors
               represented by the loadings of each of the factors
predict returns a matrix of scores of the observations in the newdata dataset.
          The other routines are mainly called for their side effect of plotting or printing and return the
          object x.

Author(s)

See Also
ilt.aplus, relativeLoadings princomp.acomp, princomp.rplus, barplot.aplus, mean.aplus,

Examples

data(SimulatedAmounts)
pc <- princomp(aplus(sa.lognormals5))
pc
summary(pc)
plot(pc)  #plot(pc,type="screeplot")
plot(pc,type="v")
plot(pc,type="biplot")
plot(pc,choice=c(1,3),type="biplot")
plot(pc,type="loadings")
plot(pc,type="loadings",scale.sdev=-1)  # Downward
plot(pc,type="relative",scale.sdev=NA)  # The directions
plot(pc,type="relative",scale.sdev=1)  # one sigma Upward
plot(pc,type="relative",scale.sdev=-1)  # one sigma Downward
biplot(pc)
screepplot(pc)
loadings(pc)
relativeLoadings(pc,mult=FALSE)
relativeLoadings(pc)
relativeLoadings(pc,scale.sdev=1)
relativeLoadings(pc,scale.sdev=2)
princomp.rcomp

Principal component analysis for real compositions

Description

A principal component analysis is done in real geometry (i.e. cpt-transform) of the simplex. Some gimmicks simplify the interpretation of the obtained components.

Usage

## S3 method for class 'rcomp'
princomp(x,...,scores=TRUE,center=attr(covmat,"center"),
  covmat=var(x,robust=robust,giveCenter=TRUE),
  robust=getOption("robust"))

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

## S3 method for class 'princomp.rcomp'
plot(x,y=NULL,...,npcs=min(10,length(x$sdev)),
  type=c("screeplot","variance","biplot","loadings","relative"),
  main=NULL,scale.sdev=1)

## S3 method for class 'princomp.rcomp'
predict(object,newdata,...)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>x</td>
<td>an rcomp dataset (for princomp) or a result from princomp.rcomp</td>
</tr>
<tr>
<td>y</td>
<td>not used</td>
</tr>
<tr>
<td>scores</td>
<td>a logical indicating whether scores should be computed or not</td>
</tr>
<tr>
<td>npcs</td>
<td>the number of components to be drawn in the scree plot</td>
</tr>
<tr>
<td>type</td>
<td>type of the plot: &quot;screeplot&quot; is a lined screeplot, &quot;variance&quot; is a boxplot-like screeplot, &quot;biplot&quot; is a biplot, &quot;loadings&quot; displays the loadings as a barplot</td>
</tr>
<tr>
<td>scale.sdev</td>
<td>the multiple of sigma to use when plotting the loadings</td>
</tr>
<tr>
<td>main</td>
<td>title of the plot</td>
</tr>
<tr>
<td>object</td>
<td>a fitted princomp.rcomp object</td>
</tr>
<tr>
<td>newdata</td>
<td>another compositional dataset of class rcomp</td>
</tr>
<tr>
<td>...</td>
<td>further arguments to pass to internally-called functions</td>
</tr>
<tr>
<td>covmat</td>
<td>provides the covariance matrix to be used for the principle component analysis</td>
</tr>
<tr>
<td>center</td>
<td>provides the be used for the computation of scores</td>
</tr>
<tr>
<td>robust</td>
<td>Gives the robustness type for the calculation of the covariance matrix. See var.rmult for details.</td>
</tr>
</tbody>
</table>
Details

Mainly a princomp(cpt(x)) is performed. To avoid confusion, the meaningless last principal component is removed.

The plot routine provides screeplots (type = "s", type = "v"). biplots (type = "b"), plots of the effect of loadings (type = "b") in scale.sdev*sdev-spread, and loadings of pairwise differences (type = "r").

The interpretation of a screeplot does not differ from ordinary screeplots. It shows the eigenvalues of the covariance matrix, which represent the portions of variance explained by the principal components.

The interpretation of the biplot strongly differs from a classical one. The relevant variables are not the arrows drawn (one for each component), but rather the links (i.e., the differences) between two arrow heads, which represents the difference between the two components represented by the arrows, or the transfer of mass from one to the other.

The compositional loading plot is more or less a standard one. The loadings are displayed by a barplot as positive and negative changes of amounts.

The loading plot can work in two different modes: If scale.sdev is set to NA it displays the composition being represented by the unit vector of loadings in cpt-transformed space. If scale.sdev is numeric we use this composition scaled by the standard deviation of the respective component.

The relative plot displays the relativeLoadings as a barplot. The deviation from a unit bar shows the effect of each principal component on the respective differences.

Value

princomp gives an object of type c("princomp.rcomp","princomp") with the following content:

- sdev: the standard deviation of the principal components.
- loadings: the matrix of variable loadings (i.e., a matrix which columns contain the eigenvectors). This is of class "loadings". The last eigenvalue is removed since it should contain the irrelevant scaling.
- Loadings: the loadings as an rmult-object
- center: the cpt-transformed vector of means used to center the dataset
- Center: the rcomp vector of means used to center the dataset
- scale: the scaling applied to each variable
- n.obs: number of observations
- scores: if scores = TRUE, the scores of the supplied data on the principal components. Scores are coordinates in a basis given by the principal components and thus not compositions
- call: the matched call
- na.action: not clearly understood

predict returns a matrix of scores of the observations in the newdata dataset. The other routines are mainly called for their side effect of plotting or printing and return the object x.

Author(s)

princomp.rmult

**See Also**

cpt.rcomp, relativeLoadings, princomp.acomp, princomp.rplus.

**Examples**

data(SimulatedAmounts)
pc <- princomp(rcomp(sa.lognormals5))
summary(pc)
plot(pc)
plot(pc, type="v")
plot(pc, type="biplot")
plot(pc, choice=c(1,3), type="biplot")
plot(pc, type="loadings")
plot(pc, type="relative", scale.sdev=-1) # Downward
plot(pc, type="relative", scale.sdev=NA) # The directions
plot(pc, type="relative", scale.sdev=1) # one sigma Upward
plot(pc, type="relative", scale.sdev=-1) # one sigma Downward
biplot(pc)
screepplot(pc)
loadings(pc)
relativeLoadings(pc, mult=FALSE)
relativeLoadings(pc)
relativeLoadings(pc, scale.sdev=1)
relativeLoadings(pc, scale.sdev=2)

pc$sdev^2
cov(predict(pc, sa.lognormals5))

---

**princomp.rmult**

Principal component analysis for real data

**Description**

Performs a principal component analysis for datasets of type rmult.

**Usage**

```r
## S3 method for class 'rmult'
princomp(x, cor=FALSE, scores=TRUE,
        covmat=var(rmult(x[subset,]), robust=robust, giveCenter=TRUE),
        center=attr(covmat,"center"), subset = rep(TRUE, nrow(x)),
        ..., robust=getOption("robust"))
```

**Arguments**

- `x` a rmult-dataset
- `...` Further arguments to call `princomp.default`
cor logical: shall the computation be based on correlations rather than covariances?

scores logical: shall scores be computed?

covmat provides the covariance matrix to be used for the principle component analysis

center provides the be used for the computation of scores

subset A rowindex to x giving the columns that should be used to estimate the variance.

robust Gives the robustness type for the calculation of the covariance matrix. See var.rmult for details.

Details

The function just does princomp(unclass(x),...,scale=scale) and is only here for convenience.

Value

An object of type princomp with the following fields

sdev the standard deviation of the principal components.

loadings the matrix of variable loadings (i.e., a matrix whose columns contain the eigen-vectors). This is of class "loadings".

center the mean that was substracted from the data set

scale the scaling applied to each variable

n.obs number of observations

scores if scores = TRUE, the scores of the supplied data on the principal components. Scores are coordinates in a basis given by the principal components.

call the matched call

na.action Not clearly understood

Author(s)


See Also

princomp.rplus

Examples

data(SimulatedAmounts)
pc <- princomp(rmult(sa.lognormals5))
pc
summary(pc)
plot(pc)
screeplot(pc)
screeplot(pc,type="l")
biplot(pc)
biplot(pc,choice=c(1,3))
princomp.rplus  Principal component analysis for real amounts

Description

A principal component analysis is done in real geometry (i.e. using iit-transform).

Usage

```
princomp(x,...,scores=TRUE,center=attr(covmat,"center"),
      covmat=var(x,robust=robust,giveCenter=TRUE),
      robust=getOption("robust"))
```

Arguments

- `x` an rplus-dataset (for princomp) or a result from princomp.rplus
- `y` not used
- `scores` a logical indicating whether scores should be computed or not
- `npcs` the number of components to be drawn in the scree plot
- `type` type of the plot: "screeplot" is a lined screeplot, "variance" is a boxplot-like screeplot, "biplot" is a biplot, "loadings" displays the loadings as a barplot
- `scale.sdev` the multiple of sigma to use when plotting the loadings
- `main` title of the plot
- `object` a fitted princomp.rplus object
- `newdata` another amount dataset of class rcomp
- `...` further arguments to pass to internally-called functions
- `covmat` provides the covariance matrix to be used for the principle component analysis
- `center` provides the be used for the computation of scores
- `robust` Gives the robustness type for the calculation of the covariance matrix. See `var.rmult` for details.
Details

Mainly a `princomp(iit(x))` is performed. Note all parts in a composition or in an amount vector share a natural scaling. Therefore, they do not need any preliminary standardization (which in fact would produce a loss of important information). For this reason, `princomp.rplus` works on the covariance matrix.

The plot routine provides screeplots (type = "s", type = "v"), biplots (type = "b"), plots of the effect of loadings (type = "b") in `scale.sdev*sdev-spread`, and loadings of pairwise differences (type = "r").

The interpretation of a screeplot does not differ from ordinary screeplots. It shows the eigenvalues of the covariance matrix, which represent the portions of variance explained by the principal components.

The interpretation of the biplot uses, additionally to the classical interpretation, a compositional concept: the differences between two arrowheads can be interpreted as the shift of mass between the two components represented by the arrows.

The amount loading plot is more or less a standard loadings plot. The loadings are displayed by a barplot as positive and negative changes of amounts.

The loadings plot can work in two different modes: If `scale.sdev` is set to `NA` it displays the amount vector being represented by the unit vector of loadings in the iit-transformed space. If `scale.sdev` is numeric we use this amount vector scaled by the standard deviation of the respective component.

The relative plot displays the `relativeLoadings` as a barplot. The deviation from a unit bar shows the effect of each principal component on the respective differences.

Value

`princomp` gives an object of type c("princomp.rcomp","princomp") with the following content:

- `sdev` the standard deviation of the principal components
- `loadings` the matrix of variable loadings (i.e., a matrix which columns contain the eigenvectors). This is of class "loadings"
- `Loadings` the loadings as an "rmult"-object
- `center` the iit-transformed vector of means used to center the dataset
- `Center` the `rplus` vector of means used to center the dataset (center and `Center` have no difference, except that the second has a class)
- `scale` the scaling applied to each variable
- `n.obs` number of observations
- `scores` if `scores = TRUE`, the scores of the supplied data on the principal components. Scores are coordinates in a basis given by the principal components and thus not compositions
- `call` the matched call
- `na.action` not clearly understood

`predict` returns a matrix of scores of the observations in the `newdata` dataset.

The other routines are mainly called for their side effect of plotting or printing and return the object `x`. 
See Also

iit.rplus, relativeLoadings princomp.rcomp, princomp.aplus.

Examples

data(SimulatedAmounts)
pc <- princomp(rplus(sa.lognormals5))
pc
summary(pc)
plot(pc)  #plot(pc,type="screeplot")
plot(pc,type="v")
plot(pc,type="biplot")
plot(pc,choice=c(1,3),type="biplot")
plot(pc,type="loadings")
plot(pc,type="relative",scale.sdev=-1)  # Downward
plot(pc,type="relative",scale.sdev=NA)  # The directions
plot(pc,type="relative",scale.sdev=1)  # one sigma Upward
plot(pc,type="relative",scale.sdev=-1)  # one sigma Downward
biplot(pc)
screenplot(pc)
loadings(pc)
relativeLoadings(pc,mult=FALSE)
relativeLoadings(pc)
relativeLoadings(pc,scale.sdev=1)
relativeLoadings(pc,scale.sdev=2)

pc$sdev^2
cov(predict(pc,sa.lognormals5))

print.acomp

Printing compositional data.

Description

Prints compositional objects with appropriate missing encodings.

Usage

## S3 method for class 'acomp'
print(x,...,replace0=TRUE)
## S3 method for class 'aplus'
print(x,...,replace0=TRUE)
## S3 method for class 'rcomp'
print(x,...,replace0=FALSE)
## S3 method for class 'rplus'
print(x,...,replace0=FALSE)
Arguments

x  a compositional object
...
replace0 logical: Shall 0 be treated as "Below detection Limit" with unknown limit.

Details

Missings are displayed with an appropriate encoding:

- MARMissing at random: The value is missing independently of its true value.
- MNARMissing NOT at random: The value is missing dependently of its true value, but without a known systematic. Maybe a better name would be: Value dependent missingness.
- BDTbelow detection limit (with unspecified detection limit): The value is missing because it was below an unknown detection limit.
- <Detectionlimitbelow detection limit (with specified detection limit): The value is below the displayed detection limit.
- SZStructural Zero: A true value is either bound to be zero or does not exist for structural nonrandom reasons. E.g. the portion of pregnant girls at a boys school.
- ERRError: An illegal encoding value was found in the object.

Value

An invisible version of x.

Missing Policy

The policy of treatment of zeroes, missing values and values below detection limit is explained in depth in compositions.missings.

Author(s)


References


See Also

clr.acomp, plot.acomp, boxplot.acomp, barplot.acomp, mean.acomp, var.acomp, variation.acomp, zeroreplace
Examples

```r
data(SimulatedAmounts)
mydata <- simulateMissings(sa.groups5, dl=0.01, knownlimit=TRUE, MAR=0.05, MNARprob=0.05, SZprob=0.05)
mydata[1,1]<-BDLvalue
print(aplus(mydata))
print(aplus(mydata), digits=3)
print(acomp(mydata))
print(rplus(mydata))
print(rcomp(mydata))
```

**pwlPlot**  
*Plots of pairwise logratio against a covariable.*

**Description**

Creates a matrix of plots, with each pairwise logratio against a covariable. The covariable can be numeric or factor, and play the role of X or Y axis.

**Usage**

```r
pwlPlot(x,y,...,add.line=FALSE,line.col=2,add.robust=FALSE,rob.col=4)
```

**Arguments**

- `x` 
  a vector, a column of a data.frame, or an acomp representing the first set of things to be displayed. Either `x` or `y` must be an acomp object, and the other must be a covariable. Both factors and continuous covariables allowed here.

- `y` 
  a vector, a column of a data.frame, or an acomp representing the first set of things to be displayed. Either `x` or `y` must be an acomp object, and the other must be a covariable. Factors to be used here with caution.

- `...` 
  further parameters to the panel function

- `add.line` 
  logical, to control the addition of a regression line in each panel. Ignored if covariable is a factor.

- `line.col` 
  in case the regression line is added, which color should be used? Defaults to red.

- `add.robust` 
  logical, to control the addition of a robust regression line in each panel. Ignored if covariable is a factor. This is nowadays based on `lmrob`, but this can change in the future.

- `rob.col` 
  in case the robust regression line is added, which color should be used? Defaults to blue.
Details

This function generates a matrix of plots of all possible pairwise logratios of the `acomp` argument, plotted against a covariable. The covariable can be a factor or a numeric vector, or a column of a matrix or data.frame. Covariable and composition can both be represented in X or Y axis: a factor on X axis generates a `boxplot`; a factor on Y axis generates a `spineplot`; if the covariable is numeric, a default scatterplot is generated. All dot arguments are passed to these plotting functions. In any of these cases, the diagram shows the logratio of the component in the row divided by the component in the column. In the case of a numeric covariable, both classical and robust regression lines can be added.

Author(s)


References

http://ima.udg.es/Activitats/CoDaWork03

http://ima.udg.es/Activitats/CoDaWork05
http://ima.udg.es/Activitats/CoDaWork08

See Also

`plot.aplus.pairwisePlot, boxplot, spineplot, plot.default`

Examples

data(Hydrochem)
xc = acomp(Hydrochem[,c("Ca","Mg","Na","K")])
fk = Hydrochem$River
pH = -log10(Hydrochem$H)
## x=acomp, y=factor
pwlrPlot(xc, fk, border=2:5)
## x=factor, y=acomp
pwlrPlot(fk, xc, col=2:5)
## x=acomp, y=numeric, with colors by river
pwlrPlot(xc, pH, col=as.integer(fk)+1)
## x=numeric, y=acomp, with line
pwlrPlot(pH, xc, add.robust=TRUE)
Normal quantile plots for compositions and amounts

Description

The plots allow to check the normal distribution of multiple univariate marginals by normal quantile-quantile plots. For the different interpretations of amount data a different type of normality is assumed and checked. When an alpha-level is given the marginal displayed in each panel is checked for normality.

Usage

```r
## S3 method for class 'acomp'
qqnorm(y, fak=NULL, ..., panel=vp.qqnorm, alpha=NULL)

## S3 method for class 'rcomp'
qqnorm(y, fak=NULL, ..., panel=vp.qqnorm, alpha=NULL)

## S3 method for class 'aplus'
qqnorm(y, fak=NULL, ..., panel=vp.qqnorm, alpha=NULL)

## S3 method for class 'rplus'
qqnorm(y, fak=NULL, ..., panel=vp.qqnorm, alpha=NULL)

vp.qqnorm(x, y, ..., alpha=NULL)
```

Arguments

- `y`: a dataset
- `fak`: a factor to split the dataset, not yet implemented in aplus and rplus
- `panel`: the panel function to be used or a list of multiple panel functions
- `alpha`: the alpha level of a test for normality to be performed for each of the displayed marginals. The levels are adjusted for multiple testing with a Bonferroni-correction (i.e. dividing each of the alpha-level by the number of test performed)
- `...`: further graphical parameters
- `x`: used by pairs only. Internal use

Details

`qqnorm.rplus` and `qqnorm.rcomp` display `qqnorm` plots of individual amounts (on the diagonal), of pairwise differences of amounts (above the diagonal) and of pairwise sums of amounts (below the diagonal).

`qqnorm.aplus` displays `qqnorm`-plots of individual log-amounts (on the diagonal), of pairwise log-ratios of amounts (above the diagonal) and of pairwise sums of log amount (below the diagonal).

Nothing is displayed on the diagonal.

In all cases a joint normality of the original data in the selected framework would imply normality in all displayed marginal distributions (although the reciprocal is in general not true!).

The marginal normality can be checked in each of the plots using a `shapiro.test` by specifying an
alpha level. The alpha level is corrected for multiple testing. Plots displaying a marginal distribution significantly deviating from a normal distribution at that alpha level are marked by a red exclamation mark.

vp.qqnorm is internally used as a panel function to make high dimensional plots.

Author(s)


See Also

plot.acomp, boxplot.acomp, rnorm.acomp, rnorm.rcomp, rnorm.rnorm.aplus, rnorm.rplus

Examples

data(SimulatedAmounts)
qqnorm(acomp(sa.lognormals),alpha=0.05)
qqnorm(rcomp(sa.lognormals),alpha=0.05)
qqnorm(aplus(sa.lognormals),alpha=0.05)
qqnorm(rplus(sa.lognormals),alpha=0.05)

Description

The R2 measure of determination for linear models

Usage

R2(object,...)
## S3 method for class 'lm'
R2(object,...,adjust=TRUE,ref=0)
## Default S3 method:
R2(object,...,ref=0)

Arguments

object a statistical model
... further not yet used parameters
adjust Logical, whether the estimate of R2 should be adjusted for the degrees of freedom of the model.
ref A reference model for computation of a relative $R^2$. 
Details

The $R^2$ measure of determination is defined as:

$$R^2 = 1 - \frac{\text{var(residuals)}}{\text{var(data)}}$$

and provides the portion of variance explained by the model. It is a number between 0 and 1, where 1 means the model perfectly explains the data and 0 means that the model has no better explanation of the data than a constant mean. In case of multivariate models metric variances are used.

If a reference model is given by ref, the variance of the residuals of that models rather than the variance of the data is used. The value of such a relative $R^2$ estimates how much of the residual variance is explained.

If adjust=TRUE the unbiased estimators for the variances are used, to avoid the automatisme that a more parameters automatically lead to a higher $R^2$.

Value

The R2 measure of determination.

Author(s)


See Also

lm, mvar, AIC

Examples

data(Orange)
R2(lm(circumference~age,data=Orange))
R2(lm(log(circumference)~log(age),data=Orange))

---

**rAitchison**  
*Aitchison Distribution*

Description

The Aitchison distribution is a class of distributions the simplex, containing the normal and the Dirichlet as subfamilies.
Usage

dAitchison(x,
  theta=alpha+sigma %% clr(mu),
  beta=-1/2*gsi.svdinverse(sigma),
  alpha=mean(theta),
  mu=clrInv(c(sigma%%%(theta-alpha))),
  sigma=-1/2*gsi.svdinverse(beta),
  grid=30,
  realdensity=FALSE,
  expKappa=AitchisonDistributionIntegrals(theta,beta,
    grid=grid,mode=1)$expKappa)

rAitchison(n,
  theta=alpha+sigma %% clr(mu),
  beta=-1/2*gsi.svdinverse(sigma),
  alpha=mean(theta),
  mu=clrInv(c(sigma%%%(theta-alpha))),
  sigma=-1/2*gsi.svdinverse(beta), withfit=FALSE)

AitchisonDistributionIntegrals(
  theta=alpha+sigma %% clr(mu),
  beta=-1/2*gsi.svdinverse(sigma),
  alpha=mean(theta),
  mu=clrInv(c(sigma%%%(theta-alpha))),
  sigma=-1/2*gsi.svdinverse(beta),
  grid=30,
  mode=3)

Arguments

x          acomp-compositions the density should be computed for.
n          integer: number of datasets to be simulated
theta      numeric vector: Location parameter vector
beta       matrix: Spread parameter matrix (clr or ilr)
alpha      positiv scalar: departure from normality parameter (positive scalar)
mu         acomp-composition, normal reference mean parameter composition
sigma      matrix: normal reference variance matrix (clr or ilr)
grid       integer: number of discretisation points along each side of the simplex
realdensity logical: if true the density is given with respect to the Haar measure of the real simplex, if false the density is given with respect to the Aitchison measure of the simplex.
mode       integer: desired output: -1: Compute nothing, only transform parameters, 0: Compute only oneIntegral and kappaIntegral, 1: compute also the clrMean, 2: compute also the clrSqExpectation, 3: same as 2, but compute clrVar instead of clrSqExpectation
should a pre-splitting of the Aitchison density be used for simulation?

Details

The Aitchison Distribution is a joint generalisation of the Dirichlet Distribution and the additive log-normal distribution (or normal on the simplex). It can be parametrized by Ait(\theta, \beta) or by Ait(\alpha, \mu, \Sigma). Actually, beta and Sigma can be easily transformed into each other, such that only one of them is necessary. Parameter \theta is a vector in \mathbb{R}^D, \alpha is its sum, \mu is a composition in \mathbb{S}^D, and beta and sigma are symmetric matrices, which can either be expressed in ilr or clr space. The parameters are transformed as

\[
\beta = -1/2\Sigma^{-1} \\
\theta = clr(\mu)\Sigma + \alpha(1, \ldots, 1)^T
\]

The distribution exists, if either, \alpha \geq 0 and Sigma is positive definite (or beta negative definite) in ilr-coordinates, or if each theta is strictly positive and Sigma has at least one positive eigenvalue (or beta has at least one negative eigenvalue). The simulation procedure currently only works with the first case.

AitchisonDistributionIntegral is a convenience function to compute the parameter transformation and several functions of these parameters. This is done by numerical integration over a multinomial simplex lattice of D parts with grid many elements (see xsimplex).

The density of the Aitchison distribution is given by:

\[
f(x, \theta, \beta) = \exp((\theta - 1)^T \log(x) + ilr(x)^T \beta ilr(x)) / \exp(\kappa_{Ait}(\theta, \beta))
\]

with respect to the classical Haar measure on the simplex, and as

\[
f(x, \theta, \beta) = \exp(\theta^T \log(x) + ilr(x)^T \beta ilr(x)) / \exp(\kappa_{Ait}(\theta, \beta))
\]

with respect to the Aitchison measure of the simplex. The closure constant expKappa is computed numerically, in AitchisonDistributionIntegrals.

The random composition generation is done by rejection sampling based on an optimally fitted additive logistic normal distribution. Thus, it only works if the corresponding Sigma in ilr would be positive definite.

Value

dAitchison Returns the density of the Aitchison distribution evaluated at x as a numeric vector.

rAitchison Returns a sample of size n of simulated compositions as an acomp object.

AitchisonDistributionIntegrals Returns a list with

- \theta: \theta parameter given or computed
- \beta: \beta parameter given or computed
- \alpha: \alpha parameter given or computed
- \mu: \mu parameter given or computed
- \Sigma: \Sigma parameter given or computed
• expKappa: the integral over the density without closing constant. I.e. the inverse of the closing constant and the exp of $\kappa_{\text{Ait}}(\theta,\beta)$
• kappaIntegral: The expected value of the mean of the logs of the components as numerically computed
• clrMean: The mean of the clr transformed random variable, computed numerically
• clrSqExpectation: The expectation of $\text{clr}(X)^t \text{clr}(X)$ computed numerically.
• clrVar: The variance covariance matrix of clr(X), computed numerically.

Note

The simulation procedure currently only works with a positive definite Sigma. You need a relatively high grid constant for precise values in the numerical integration.

Author(s)


References


See Also

runif.acomp, rnorm.acomp, rDirichlet.acomp

Examples

```r
(erg <- AitchisonDistributionIntegrals(c(-1,3,-2), ilrvar2clr(-diag(c(1,2))), grid=20))

(myvar <- with(erg, -1/2*ilrvar2clr(solve(clrvar2ilr(beta)))))

(mymean <- with(erg, myvar%*%theta))

with(erg, myvar - clrVar)
with(erg, mymean - clrMean)
```

ratioLoadings  Loadings of relations of two amounts

Description

In a compositional dataset the relation of two objects can be interpreted safer than a single amount. These functions compute, display and plot the corresponding pair-information for the various principal component analysis results.
Usage

relativeLoadings(x,...)
## S3 method for class 'princomp.acomp'
relativeLoadings(x,...,log=FALSE,scale.sdev=NA,
               cutoff=0.1)
## S3 method for class 'princomp.aplus'
relativeLoadings(x,...,log=FALSE,scale.sdev=NA,
               cutoff=0.1)
## S3 method for class 'princomp.rcomp'
relativeLoadings(x,...,scale.sdev=NA,
               cutoff=0.1)
## S3 method for class 'princomp.rplus'
relativeLoadings(x,...,scale.sdev=NA,
               cutoff=0.1)
## S3 method for class 'relativeLoadings.princomp.acomp'
print(x,...,cutoff=attr(x,"cutoff"),
       digits=2)
## S3 method for class 'relativeLoadings.princomp.aplus'
print(x,...,cutoff=attr(x,"cutoff"),
       digits=2)
## S3 method for class 'relativeLoadings.princomp.rcomp'
print(x,...,cutoff=attr(x,"cutoff"),
       digits=2)
## S3 method for class 'relativeLoadings.princomp.rplus'
print(x,...,cutoff=attr(x,"cutoff"),
       digits=2)
## S3 method for class 'relativeLoadings.princomp.acomp'
plot(x,...)
## S3 method for class 'relativeLoadings.princomp.aplus'
plot(x,...)
## S3 method for class 'relativeLoadings.princomp.rcomp'
plot(x,...)
## S3 method for class 'relativeLoadings.princomp.rplus'
plot(x,...)

Arguments

x a result from an amount PCA princomp.acomp/princomp.aplus/princomp.rcomp/princomp.rplus
log a logical indicating to use log-ratios instead of ratios
scale.sdev if not NA, a number specifying the multiple of a standard deviation, used to scale the components
cutoff a single number. Changes under that (log)-cutoff are not displayed
digits the number of digits to be displayed
... further parameters to internally-called functions
Details

The relative loadings of components allow a direct interpretation of the effects of principal components. For acomp/aplus classes the relation is induced by a ratio, which can optionally be log-transformed. For the rcomp/rplus-classes the relation is induced by a difference, which is meaningless when the units are different.

Value

The value is a matrix of type "relativeLoadings.princomp.*", containing the ratios in the compositions represented by the loadings (optionally scaled by the standard deviation of the components and scale.sdev).

Author(s)


See Also

princomp.acomp, princomp.aplus, princomp.rcomp, princomp.rplus, barplot

Examples

data(SimulatedAmounts)
pc <- princomp(acomp(sa.lognormals5))
pc
summary(pc)
relativeLoadings(pc,log=TRUE)
relativeLoadings(pc)
relativeLoadings(pc,scale.sdev=1)
relativeLoadings(pc,scale.sdev=2)

plot(relativeLoadings(pc,log=TRUE))
plot(relativeLoadings(pc))
plot(relativeLoadings(pc,scale.sdev=1))
plot(relativeLoadings(pc,scale.sdev=2))

rcomp Compositions as elements of the simplex embedded in the D-dimensional real space

Description

A class providing a way to analyse compositions in the philosophical framework of the Simplex as subset of the $R^D$. 
Usage

```r
rcomp(X, parts=1:NCOL(oneOrDataset(X)), total=1, warn.na=FALSE,
    detectionlimit=NULL, BDL=NULL, MAR=NULL, MNAR=NULL, SZ=NULL)
```

Arguments

- **X**: composition or dataset of compositions
- **parts**: vector containing the indices xor names of the columns to be used
- **total**: the total amount to be used, typically 1 or 100
- **warn.na**: should the user be warned in case of NA, NaN or 0 coding different types of missing values?
- **detectionlimit**: a number, vector or matrix of positive numbers giving the detection limit of all values, all columns or each value, respectively
- **BDL**: the code for 'Below Detection Limit' in X
- **SZ**: the code for 'Structural Zero' in X
- **MAR**: the code for 'Missing At Random' in X
- **MNAR**: the code for 'Missing Not At Random' in X

Details

Many multivariate datasets essentially describe amounts of D different parts in a whole. This has some important implications justifying to regard them as a scale on its own, called a "composition". The functions around the class "rcomp" follow the traditional (often statistically inconsistent) approach regarding compositions simply as a multivariate vector of positive numbers summing up to 1. This space of D positive numbers summing to 1 is traditionally called the D-1-dimensional simplex.

The compositional scale was in-depth analysed by Aitchison (1986) and he found serious reasons why compositional data should be analysed with a different geometry. The functions around the class "acomp" follow his approach. However the Aitchison approach based on log-ratios is sometimes criticized (e.g. Rehder and Zier, 2002). It cannot deal with absent parts (i.e. zeros). It is sensitive to large measurement errors in small amounts. The Aitchison operations cannot represent simple mixture of different compositions. The used transformations are not uniformly continuous. Straight lines and ellipses in Aitchison space look strangely in ternary diagrams. As all uncritical statistical analysis, blind application of logratio-based analysis is sometimes misleading. Therefore it is sometimes useful to analyse compositional data directly as a multivariate dataset of portions summing to 1. However a clear warning must be given that the utilisation of almost any kind of classical multivariate analysis introduce some kinds of artifacts (e.g. Chayes 1960) when applied to compositional data. So, extra care and considerable expert knowlegde is needed for the proper interpretation of results achieved in this non-Aitchison approach. The package tries to lead the user around these artifacts as much as possible and gives hints to major pitfalls in the help. However meaningless results cannot be fully avoided in this (rather inconsistent) approach.

A side effect of the procedure is to force the compositions to sum to one, which is done by the closure operation `clo`.

The classes rcomp, acomp, aplus, and rplus are designed in a fashion as similar as possible, in order to allow direct comparison between results achieved by the different approaches. Especially the
acomp logistic transforms \texttt{clr, alr, ilr} are mirrored by analogous linear transforms \texttt{cpt, apt, ipt} in the rcomp class framework.

**Value**

a vector of class "rcomp" representing a closed composition or a matrix of class "rcomp" representing multiple closed compositions, by rows.

**Missing Policy**

Missing and Below Detection Limit Policy is explained in deeper detail in \texttt{compositions.missing}.

**Author(s)**

Raimon Tolosana-Delgado, K.Gerald v.d. Boogaart \url{http://www.stat.boogaart.de}

**References**


**See Also**

\texttt{cpt, apt, ipt, acomp, rplus, princomp.rcomp, plot.rcomp, boxplot.rcomp, barplot.rcomp, mean.rcomp, var.rcomp, variation.rcomp, cov.rcomp, msd, convex.rcomp, +.rcomp}

**Examples**

data(SimulatedAmounts)
plot(rcomp(sa.tnormals))
Description
The real compositions form a manifold of the real vector space. The induced operations +, -, *, / give results valued in the real vector space, but possibly outside the simplex.

Usage
convex.rcomp(x, y, alpha=0.5)
## Methods for class "rcomp"
## x+y
## x-y
## -x
## x*r
## r*x
## x/r

Arguments
x an rcomp composition or dataset of compositions
y an rcomp composition or dataset of compositions
r a numeric vector of size 1 or nrow(x)
alpha a numeric vector of size 1 or nrow(x) with values between 0 and 1

Details
The functions behave quite like +.rmult.
The convex combination is defined as: x*alpha + (1-alpha)*y

Value
rmult-objects containing the given operations on the simplex as subset of the $R^D$. Only the convex combination convex.rcomp results in an rcomp-object again, since only this operation is closed.

Note
For * the arguments x and y can be exchanged.

Author(s)

See Also
+.rmult, +.acomp, cpt, rcomp, rmult
Examples

```r
rcomp(1:5)* -1 + rcomp(1:5)
data(SimulatedAmounts)
cdata <- rcomp(sa.lognormals)
plot( tmp <- (cdata-mean(cdata))/msd(cdata) )
class(tmp)
mean(tmp)
msd(tmp)
var(tmp)
plot(convex.rcomp(rcomp(c(1,1,1)),sa.lognormals,0.1))
```

---

**rcompmargin**  
*Marginal compositions in real geometry*

**Description**

Compute marginal compositions by amalgamating the rest (additively).

**Usage**

```r
rcompmargin(X,d=c(1,2),name="+",pos=length(d)+1,what="data")
```

**Arguments**

- `X`: composition or dataset of compositions
- `d`: vector containing the indices xor names of the columns to be kept
- `name`: The new name of the amalgamation column
- `pos`: The position where the new amalgamation column should be stored. This defaults to the last column.
- `what`: The role of X either "data" for data (or means) to be transformed or "var" for variances to be transformed.

**Details**

The amalgamation column is simply computed by adding the non-selected components after closing the composition. This is consistent with the `rcomp` approach and is widely used because of its easy interpretation. However, it often leads to difficult-to-read ternary diagrams and is inconsistent with the `acomp` approach.

With the argument what="var" the function transformes an rcomp variance to the resulting variance of the resulting composition.

**Value**

A closed compositions with class "rcomp" containing the selected variables given by d and the the amalgamation column.
**Missing Policy**

MNAR has the highest priority, MAR next and WZERO (BDL, SZ), zero values are considered as 0 and reported as BDL in the End.

**Author(s)**


**References**

References missing

**See Also**

acompmargin, rcomp

**Examples**

```r
data(SimulatedAmounts)
plot.rcomp(sa.tnormals5,margin="rcomp")
plot.rcomp(rcompmargin(sa.tnormals5,c("Cd","Zn")))
plot.rcomp(rcompmargin(sa.tnormals5,c(1,2)))
```

---

**rDirichlet**

*Dirichlet distribution*

**Description**

The Dirichlet distribution on the simplex.

**Usage**

```r
rDirichlet.acomp(n,alpha)
```

**Arguments**

- `n` number of datasets to be simulated
- `alpha` parameters of the Dirichlet distribution

**Details**

The Dirichlet distribution is the result of closing a vector of equally-scaled Gamma-distributed variables. It is the conjugate prior distribution for a vector of probabilities of a multinomial distribution. Thus, it generalizes the beta distribution for more than two parts.
Value

a generated random dataset of class "acomp" or "rcomp", drawn from a Dirichlet distribution with the given parameter alpha. The names of alpha are used to name the parts.

Author(s)


References


See Also

rnorm.acomp

Examples

tmp <- rDirichlet.acomp(10, alpha=c(A=2, B=0.2, C=0.2))
plot(tmp)

Read standard data files

Reads a data file in a geoeas format

Description

Reads a data file, which must be formatted either as a geoEAS file (described below).

Usage

read.geoeas(file)
read.geoEAS(file)

Arguments

file a file name, with a specific format
Details

The data files must be in the adequate format: "read.geoEAS" and "read.geoeas" read geoEAS format.

The geoEAS format has the following structure:

- a first row with a description of the data set
- the number of variables (=nvars)
- "nvars" rows, each containing the name of a variable
- the data set, in a matrix of "nvars" columns and as many rows as individuals

Value

A data set, with a "title" attribute.

Note

Labels and title should not contain tabs. This might produce an error when reading.

Author(s)

Raimon Tolosana-Delgado

References

Missing references

See Also

read.table

Examples

# Files can be found in the test-subdirectory of the package
#
## Not run:
read.geoeas("TRUE.DAT")
read.geoEAS("TRUE.DAT")

## End(Not run)
replot

Modify parameters of compositional plots.

Description
Display only a subset of the plots.

Usage
replot(..., dev=dev.cur(), plot=TRUE, envir=NULL, add=FALSE)
replotable(expr, add=FALSE)
noreplot(expr, dev=dev.cur())

Arguments
expr
A (unquoted) expression that does the plotting. replotable will make the generated plot replotable and noreplot will do the inverse and avoid that the plots overwrites the current database entry.

... Plot parameters to be modified. E.g. onlyPanel

dev
The device that currently contains the plot. It will be plotted in the current device.

plot
logical or call or list of calls. If plot is TRUE, the new version of the plot is plotted in the current environment (and typically stores itself here). If plot is FALSE the modified plot is simply stored, rather than actually plotted (in its own old plotting environment). If the parameter is something else, it is stored to the internal plot database for the given device (but not plotted or evaluated).

envir
a new environment to be assigned to the plot. Rarely needed.

add
either a logical to indicating that the plot adds something to the plot. Or a number / name of the added thing to be modified.

Details
Some of the plot routines of compositions internally store their call as a mean for replaying the plot when information is added or parameters are modified. The stored call can be modified by this function, which pretty much works like a simplified version of update.

replot allows to redo the plot typically in a different device or with different parameters. The function provides this functionality at a totally different level than dev.copy and allows for the modification of high level parameters on the fly.

Plots can be stored in the internal database by calling replot with a parameter plot set to the call of that plot. Plotting functions without this functionality can be filtered through replotable(). However in this case all parameter names should be given explicitly.

There are actually three levels of possible replay: The dev.copy level on which graphic actions are replayed. The gsi.pairs function level that organizes panels plots and uses an internal replotting facility to allow modification of the parameter, e.g. addings lines .... And than there is the high level of the actual function call generating the plot.
Value

replot returns an invisible copy of the modified call. replotable and noreplot return the result of expression.

Note

The function works by reevaluating the call in its environment. Thus the plot will change!!! if the data has changed.

The function always handles the latest plot from the package. If another plot ignorant of the replot system has meanwhile be used it will be ignored.

Author(s)


See Also

plot.acomp, plot.aplus, boxplot.acomp

Examples

data(SimulatedAmounts)
plot(acomp(sa.lognormals5))
straight(acomp(c(1,1,1,1,1)),acomp(c(1,2,3,4,5)))
replot(onlyPanel=c(2,3))
oldPlot <- replot(plot=FALSE) # get the plotting call
replotable(plot(x=1:10)) # To make a graphic replottable
replot(col=1:10)
replot(plot=oldPlot) # Restore the old plot (without plotting)
replot(onlyPanel=NULL) # View the whole plot again
replot(pch=20) # Actually plot it
replot(col=20) # since the actual plot is gsi.pairs not a plot.acomp

## Not run:
# The following line in a plotting function stores the plot for reploting.
replot(plot=match.call()) # Store current call as plot
replot() # simply plot once again
replot(dev=otherdev) # redo a plot from an other device here.
replot(onlyPanel=c(3,4)) # modify the plot (and replot it)
replot(onlyPanel=c(3,4),dev=7,plot=FALSE) # modify a stored plot

## End(Not run)
**rlnorm**

*The multivariate lognormal distribution*

**Description**

Generates random amounts with a multivariate lognormal distribution, or gives the density of that distribution at a given point.

**Usage**

```
rlnorm.rplus(n, meanlog, varlog)
dlnorm.rplus(x, meanlog, varlog)
```

**Arguments**

- `n` number of datasets to be simulated
- `meanlog` the mean-vector of the logs
- `varlog` the variance/covariance matrix of the logs
- `x` vectors in the sample space

**Value**

- `rlnorm.rplus` gives a generated random dataset of class "rplus" following a lognormal distribution with logs having mean `meanlog` and variance `varlog`.
- `dlnorm.rplus` gives the density of the distribution with respect to the Lesbesgue measure on R+ as a subset of R.

**Note**

The main difference between `rlnorm.rplus` and `rnorm.aplus` is that `rlnorm.rplus` needs a logged mean. The additional difference for the calculation of the density by `dlnorm.rplus` and `dnorm.aplus` is the reference measure (a log-Lebesgue one in the second case).

**Author(s)**


**References**


**See Also**

`rnorm.acomp`
Examples

```r
MyVar <- matrix(c(0.2,0.1,0.0,
                   0.1,0.2,0.0,
                   0.0,0.0,0.2),byrow=TRUE,nrow=3)
MyMean <- c(1,1,2)
plot(rlnorm.rplus(100,log(MyMean),MyVar))
plot(rnorm.aplus(100,MyMean,MyVar))
x <- rnorm.aplus(5,MyMean,MyVar)
dnorm.aplus(x,MyMean,MyVar)
dlnorm.rplus(x,log(MyMean),MyVar)
```

\[r\text{Mahalanobis}\]

**Description**

Decisions about outliers are often made based on Mahalanobis distances with respect to robustly estimated variances. These functions deliver the necessary distributions.

**Usage**

```r
rEmpiricalMahalanobis(n,N,d,...,sorted=FALSE,pow=1,robust=TRUE)
pEmpiricalMahalanobis(q,N,d,...,pow=1,replicates=100,resample=FALSE,robust=TRUE)
qEmpiricalMahalanobis(p,N,d,...,replicates=1000,resample=FALSE,pow=1,robust=TRUE)
rMaxMahalanobis(n,N,d,...,pow=1,robust=TRUE)
pMaxMahalanobis(q,N,d,...,replicates=998,resample=FALSE,robust=TRUE)
qMaxMahalanobis(p,N,d,...,replicates=998,resample=FALSE,pow=1,robust=TRUE)
rPortionMahalanobis(n,N,d,cut,...,pow=1,robust=TRUE)
pPortionMahalanobis(q,N,d,cut,...,replicates=1000,resample=FALSE,pow=1,robust=TRUE)
qPortionMahalanobis(p,N,d,cut,...,replicates=1000,resample=FALSE,pow=1,robust=TRUE)
pQuantileMahalanobis(q,N,d,p,...,replicates=1000,ulimit=TRUE,pow=1,robust=TRUE)
```

**Arguments**

- **n** Number of simulations to do.
- **q** A vector giving quantiles of the distribution.
p  A vector giving probabilities. (only a single probility for pQuantileMahalanobis)
N  Number of cases in the dataset.
d  degrees of freedom (i.e. dimension) of the dataset.
cut A cutting limit. The random variable is the portion of Mahalanobis distances
lower equal to the cutting limit.
... further arguments passed to MahalanobisDist
pow  the power of the Mahalanobis distance to be used. Higher powers can be used
to stretch the outlierregion visually.
robust logical or a robust method description (see robustnessInCompositions) speci-
fying how the center and covariance matrix are estimated, if not given.
sorted Specifies a transformation to be applied to the whole sequence of Mahalanobis
distances: FALSE is no transformation; TRUE sorts the entries in ascending or-
der, a numeric vector picks the given entries from the entries sorted in ascending
order; alternatively a function such as max can be given to directly transform the
data.
replicates the number of datasets in the Monte-Carlo-Computations used in these routines.
resample a logical forcing a resampling of the Monte-Carlo-Sampling. See details.
ulimit logical: is this an upper limit of a joint confidence bound or a lower limit.

Details

All the distribution correspond to the distribution under the Null-Hypothesis of multivariate joint
Gaussian distribution of the dataset.

The set of empirically estimated Mahalanobis distances of a dataset is in the first step a random
vector with exchangeable but dependent entries. The distribution of this vector is given by the
rEmpiricalMahalanobis if no sorted argument is given. Please be advised that this is not a fixed
distribution in a mathematical sense, but an implementation dependent distribution incorporating
the performance of underlying robust spread estimator. As long as no sorted argument is given
pEmpiricalMahalanobis and qEmpiricalMahalanobis represent the distribution function and the
quantile function of a randomly picked element of this vector.
If a sorted attribute is given, it specifies a transformation is applied to each of the vector prior to pro-
cessing. Three important special cases are provided by separate functions. The MaxMahalanobis
functions correspond to picking only the largest value. The PortionMahalanobis functions corre-
spond to reporting the portion of Mahalanobis distances over a cutoff. The QuantileMahalanobis
distribution corresponds to the distribution of the p-quantile of the dataset.
The Monte-Carlo-Simulations of these distributions are rather slow, since for each datum we need
to simulate a whole dataset and to apply a robust covariance estimator to it, which typically itself
involves Monte-Carlo-Algorithms. Therefore each type of simulations is only done the first time
needed and stored for later use in the environment gsi.pStore. With the resampling argument a
resampling of the cached dataset can be forced.

Value

The r* functions deliver a vector (or a matrix of row-vectors) of simulated value of the given distri-
butions. A total of n values (or row vectors) is returned.
The p* functions deliver a vector (of the same length as x) of probabilities for random variable of
the given distribution to be under the given quantil values q.  
The q* functions deliver a vector of quantiles corresponding to the length of the vector p providing the probabilities.

Note

Unlike the mahalanobis function this function does not be default compute the square of the mahalanobis distance. The pow option is provided if the square is needed.
The package robustbase is required for using the robust estimations.

Author(s)


See Also

dist, OutlierClassifier1

Examples

rEmpiricalMahalanobis(10, 25, 2, sorted=TRUE, pow=1, robust=TRUE)
pEmpiricalMahalanobis(qchisq(0.95, df=10), 11, 1, pow=2, replicates=1000)
(xx <- pMaxMahalanobis(qchisq(0.95, df=10), 11, 1, pow=2))
qEmpiricalMahalanobis(0.95, 11, 2)
rMaxMahalanobis(10, 25, 4)
qMaxMahalanobis(xx, 11, 1)

rmult    Simple treatment of real vectors

Description

A class to analyse real multivariate vectors.

Usage

rmult(X, parts=1:NCOL(oneOrDataset(X)), orig=attr(X,"orig"),
      missingProjector=attr(X,"missingProjector"))
## S3 method for class 'rmult'
print(x,...)

Arguments

   X    vector or dataset of numbers considered as elements of a R-vector
   parts vector containing the indices xor names of the columns to be used
   x    an rmult object
   orig the original untransformed dataset
missingProjector
the Projector on the observed subspace
...further generic arguments passed to print.default

Details

The rmult class is a simple convenience class to treat data in the scale of real vectors just like data in the scale of real numbers. A major aspect to take into account is that the internal arithmetic of R is different for these vectors.

Value

a vector of class "rmult" representing one vector or a matrix of class "rmult", representing multiple vectors by rows.

Author(s)


See Also

+.rmult, scalar, norm.rmult, %*%.rmult, rplus, acomp.

Examples

plot(rnorm.rmult(30, mean=0:4, var=diag(1:5)+10))
Arguments

- **x**: an rmult vector or dataset of vectors
- **y**: an rmult vector or dataset of vectors
- **r**: a numeric vector of size 1 or nrow(x)

Details

The operators try to mimic the parallel operation of R on vectors of real numbers on vectors of vectors represented as matrices containing the vectors as rows.

Value

an object of class "rmult" containing the result of the corresponding operation on the vectors.

Author(s)


See Also

- `rmult`
- `%*%` for inner products

Examples

```r
x <- rmult(matrix(sqrt(1:12), ncol=3))
x
x + rmult(1:3)
x * 1:4
1:4 * x
x / 1:4
x / 10
```

Description

An rmult object is considered as a sequence of vectors. The `%*%` is considered as the inner multiplication. An inner multiplication with another vector is the scalar product. An inner multiplication with a matrix is a matrix multiplication, where the rmult-vectors are either considered as row or as column vector.

Usage

```r
## S3 method for class 'rmult'
x %*% y
```
Arguments

x  an rmult vector or dataset of vectors, a numeric vector of length (gsi.getD(y)), or a matrix

y  an rmult vector or dataset of vectors, a numeric vector of length (gsi.getD(x)), or a matrix

Details

The operators try to mimic the behavior of %*% on c() - vectors as inner product applied in parallel to all vectors of the dataset. Thus the product of a vector with another rmult object or unclassed vector v results in the scalar product. For the multiplication with a matrix each vector is considered as a row or column, whatever is more appropriate.

Value

an object of class "rmult" or a numeric vector containing the result of the corresponding inner products.

Note

The product x %*% A %*% y is associative.

Author(s)


See Also

rmult, %*%.rmult

Examples

x <- rmult(matrix( sqrt(1:12), ncol = 3 ))
x%*%x
A <- matrix( 1:9, nrow = 3 )
x %*% A %*% x
x %*% A
A %*% x
x %*% 1:3
x %*% 1:3
1:3 %*% x
**Description**

`rnorm.X` generates multivariate normal random variates in the space `X`.

**Usage**

```r
rnorm.acomp(n, mean, var)
rnorm.rcomp(n, mean, var)
rnorm.aplus(n, mean, var)
rnorm.rplus(n, mean, var)
rnorm.rmult(n, mean, var)
rnorm.ccomp(n, mean, var, lambda)
dnorm.acomp(x, mean, var, withJacobian=FALSE)
dnorm.aplus(x, mean, var, withJacobian=FALSE)
dnorm.rmult(x, mean, var)
```

**Arguments**

- `n`: number of datasets to be simulated
- `mean`: The mean of the dataset to be simulated
- `var`: The variance covariance matrix
- `lambda`: The expected total count
- `x`: vectors in the sampling space
- `withJacobian`: should the Jacobian of the log or logratio transformation be included in the density calculations? defaults to FALSE (see details)

**Details**

The normal distributions in the various spaces dramatically differ. The normal distribution in the `rmult` space is the commonly known multivariate joint normal distribution. For `rplus` this distribution has to be somehow truncated at 0. This is here done by setting negative values to 0, i.e. this simulation function produces a sort of multivariate tobit model.

The normal distribution of `rcomp` is seen as a normal distribution within the simplex as a geometrical portion of the real vector space. The variance is thus forced to be singular and restricted to the affine subspace generated by the simplex. The necessary truncation of negative values is currently done by setting them explicitly to zero and reclosing afterwards, again in the fashion of a tobit model.

The "acomp" and "aplus" are themselves metric vector spaces and thus a normal distribution is defined in them just as in the real space. The resulting distribution almost correspond to multivariate lognormal in the case of "aplus" and Aitchison normal distribution in the simplex in the case of "acomp". These models are equivalent in probability to the multivariate lognormal distribution and the additive logistic normal distribution respectively, albeit without including the Jacobian of the
log or the logratio transformation. If you are interested in the density of the additive logistic normal model, give the extra argument withJacobian=TRUE. If you are interested in the multivariate log-normal density you can either do the same, or better call `dlnorm.rplus`. Densities are only provided for the models constructed for `rmult`, `aplus` and `acomp` because they do exist with respect to the Lebesgue measure of each of these spaces. In the other cases it is not possible to compute a measure, since the truncation at zero values produce distributions that are not absolutely continuous with respect to the real, conventional Lebesgue measure. For count compositions `ccomp` a `norm.acomp` is realized and used as a parameter to a Poisson distribution (see `rpois.ccomp`). So, this is in reality no normal model, but a double stochastic counting process.

**Value**

A random dataset of the given class generated by a normal distribution with the given mean and variance in the given space. For the density functions `d*`, the value of the probability density at the values of `x` provided

**Author(s)**


**References**


**See Also**

`runif.acomp`, `rlnorm.rplus`, `rDirichlet.acomp`

**Examples**

```r
MyVar <- matrix(c(0.2, 0.1, 0.0, 0.1, 0.2, 0.0, 0.0, 0.0, 0.2), byrow=TRUE, nrow=3)
MyMean <- c(1, 1, 2)

plot(rnorm.acomp(100, MyMean, MyVar))
plot(rnorm.rcomp(100, MyMean, MyVar))
plot(rnorm.aplus(100, MyMean, MyVar))
plot(rnorm.rplus(100, MyMean, MyVar))
```
robustnessInCompositions

Handling robustness issues and outliers in compositions.

Description

The seamless transition to robust estimations in library(compositions).

Details

A statistical method is called nonrobust if an arbitrary contamination of a small portion of the dataset can produce results radically different from the results without the contamination. In this sense many classical procedures relying on distributional models or on moments like mean and variance are highly nonrobust.

We consider robustness as an essential prerequirement of all statistical analysis. However in the context of compositional data analysis robustness is still in its first years.

As of Mai 2008 we provide a new approach to robustness in the package. The central idea is that robustness should be more or less automatic and that there should be no necessity to change the code to compare results obtained from robust procedures and results from there more efficient nonrobust counterparts.

To achieve this all routines that rely on distributional models (such as e.g. mean, variance, principle component analysis, scaling) and routines relying on those routines get a new standard argument of the form:

`fkt(...,robust=getOption("robust"))`

which defaults to a new option "robust". This option can take several values:

- FALSE The classical estimators such as arithmetic mean and persons product moment variance are used and the results are to be considered nonrobust.
- TRUE The default for robust estimation in the package is used. At this time this is `covMcd` in the `robustbase`-package. This default might change in future.
- "pearson" This is a synonym for FALSE and explicitly states that no robustness should be used.
- "mcd" Minimum Covariance Determinant. This option explicitly selects the use of `covMcd` in the `robustbase`-package as the main robustness engine.

More options might follow later. To control specific parameters of the model the string can get an attribute named "control" which contains additional options for the robustness engine used. In this moment the control attribute of mcd is a control object of `covMcd`. The control argument of "pearson" is a list containing addition options to the mean, like trim.

The standard value for getOption("robust") is FALSE to avoid situation in which the user thinks he uses a classical technique. Robustness must be switched on explicitly. Either by setting the option...
with options(robust=TRUE) or by giving the argument. This default might change later if the authors come to the impression that robust estimation is now considered to be the default.

For those not only interested in avoiding the influence of the outliers, but in an analysis of the outliers we added a subsystem for outlier classification. This subsystem is described in outliersInCompositions and also relies on the robust option. However evidently for these routines the factory default for the robust option is always TRUE, because it is only applicable in an outlier-aware context.

We hope that in this way we can provide a seamless transition from nonrobust analysis to a robust analysis.

**Note**

IMPORTANT: The robust argument only works with the classes of the package. Only your compositional analysis is suddenly robust.

The package robustbase is required for using the robust estimations and the outlier subsystem of compositions. To simplify installation it is not listed as required, but it will be loaded, whenever any sort of outlierdetection or robust estimation is used.

**Author(s)**


**See Also**

var.acomp, mean.acomp, robustbase, compositions-package, missings, outlierplot, OutlierClassifier1, ClusterFinder1

**Examples**

```r
A <- matrix(c(0.1,0.2,0.3,0.1),nrow=2)
Mvar <- 0.1*ilrvar2clr(A%*%t(A))
Mcenter <- acomp(c(1,2,1))
typicalData <- rnorm.acomp(100,Mcenter,Mvar) # main population
colnames(typicalData)<-c("A","B","C")
data5 <- acomp(rbind(unclass(typicalData)+outer(rbinom(100,1,p=0.1)*runif(100),c(0.1,1,2))))

mean(data5)
mean(data5,robust=TRUE)
var(data5)
var(data5,robust=TRUE)
Mvar
biplot(princomp(data5))
biplot(princomp(data5,robust=TRUE))
```
rplus

Amounts i.e. positive numbers analysed as objects of the real vector space

Description

A class to analyse positive amounts in a classical (non-logarithmic) framework.

Usage

```
rplus(X, parts=1:NCOL(oneOrDataset(X)), total=NA, warn.na=FALSE, 
   detectionlimit=NULL, BDL=NULL, MAR=NULL, MNAR=NULL, SZ=NULL)
```

Arguments

- `X` vector or dataset of positive numbers considered as amounts
- `parts` vector containing the indices xor names of the columns to be used
- `total` a numeric vectors giving the total amount of each dataset
- `warn.na` should the user be warned in case of NA,NaN or 0 coding different types of missing values?
- `detectionlimit` a number, vector or matrix of positive numbers giving the detection limit of all values, all columns or each value, respectively
- `BDL` the code for 'Below Detection Limit' in X
- `SZ` the code for 'Structural Zero' in X
- `MAR` the code for 'Missing At Random' in X
- `MNAR` the code for 'Missing Not At Random' in X

Details

Many multivariate datasets essentially describe amounts of D different parts in a whole. When the whole is large in relation to the considered parts, such that they do not exclude each other, and when the total amount of each componenten is actually determined by the phenomenon under investigation and not by sampling artifacts (such as dilution or sample preparation) then the parts can be treated as amounts rather than as a composition (cf. `rcomp, aplus`).

In principle, amounts are just real-scaled numbers with the single restriction that they are nonnegative. Thus they can be analysed by any multivariate analysis method. This class provides a simple access interface to do so. It tries to keep in mind the positivity property of amounts and the special point zero. However there are strong arguments why an analysis based on log-scale might be much more adapted to the problem. This log-approach is provided by the class `aplus`.

The classes `rcomp, acomp, aplus, and rplus` are designed in a fashion as similar as possible in order to allow direct comparison between results obtained by the different approaches. In particular, the `aplus` logistic transform `ilt` is mirrored by the simple identity transform `iit`. In terms of computer science, this identity mapping is actually mapping an object of type "rplus" to a class-less datamatrix.
Value

A vector of class "rplus" representing a vector of amounts or a matrix of class "rplus" representing multiple vectors of amounts, by rows.

Missing Policy

Missing and Below Detection Limit Policy is more detailed explained in compositions.missing.

Author(s)


References


See Also

iit.rcomp, aplus.princomp.rplus.plot.rplus.boxplot.rplus, barplot.rplus, mean.rplus, var.rplus, variation.rplus, cov.rplus, msd

Examples

data(SimulatedAmounts)
plot(rplus(sa.lognormals))

---
rplusarithm vectorial arithmetic for data sets with rplus class

Description

The positive quadrant forms a manifold of the real vector space. The induced operations +,-,*,/ give results valued in this real vector space (not necessarily inside the manifold).

Usage

mul.rplus(x,r)
## Methods for class rplus
## x+y
## x-y
## -x
## x*r
## r*x
## x/r
rpois

Arguments

x  
an rplus composition or dataset of compositions

y  
an rplus composition or dataset of compositions

r  
a numeric vector of size 1 or nrow(x)

Details

The functions behave quite like +.rmult.

Value

rmult-objects containing the given operations on the rcomp manifold as subset of the $R^{D}$. Only the addition and multiplication with positive numbers are internal operation and results in an rplus-object again.

Note

For * the arguments x and y can be exchanged.

Author(s)


See Also

+.rmult, +.acomp, cpt, rcomp, rmult

Examples

rplus(1:5)* -1 + rplus(1:5)
data(SimulatedAmounts)
cdata <- rplus(sa.lognormals)
plot( tmp <- (cdata-mean(cdata))/msd(cdata) )
class(tmp)
mean(tmp)
msd(tmp)
var(tmp)

rpois  
Simulate count compositions without overdispersion

Description

Generates multinominal or multi-Poission random variates based on an Aitchison composition.
**Usage**

- `rpois.ccomp(n, p, lambda)`
- `rmultinom.ccomp(n, p, N)`

**Arguments**

- **n**: number of datasets to be simulated
- **p**: The composition representing the probabilities/portions of the individual parts
- **lambda**: scalar or vector giving the expected total count
- **N**: scalar or vector giving the total count

**Details**

A count composition is a realisation of a multinomial or multivariate Poisson distribution.

**Value**

A random dataset ccount dataset

**Author(s)**


**See Also**

- `rnorm.ccomp`

**Examples**

```r
p <- acomp(c(3,3,3))
rpois.ccomp(10, p, 40)
rmultinom.ccomp(10, p, 40)
```

---

**Description**

Generates random compositions with a uniform distribution on the (rcomp) simplex.

**Usage**

- `runif.acomp(n, D)`
- `runif.rcomp(n, D)`
**scalar**

**Arguments**

- `n`: number of datasets to be simulated
- `D`: number of parts

**Value**

A generated random dataset of class "acomp" or "rcomp" drawn from a uniform distribution on the simplex of `D` parts.

**Note**

The only difference between both routines is the class of the dataset returned.

**Author(s)**


**References**


**See Also**

- `rDirichlet.acomp`

**Examples**

```r
plot(runif.acomp(10,3))
plot(runif.rcomp(10,3))
```

---

**Description**

Computes scalar products of datasets of vectors or vectorial quantities.

**Usage**

```r
scalar(x,y)
## Default S3 method:
scalar(x,y)
```
Arguments

- **x**: a vector or a matrix with rows considered as vectors
- **y**: a vector or a matrix with rows considered as vectors

Details

The scalar product of two vectors is defined as:

\[ \text{scalar}(x, y) := \sum (x_i y_i) \]

Value

A numerical vector containing the scalar products of the vectors given by `x` and `y`. If both `x` and `y` contain more than one vector the function uses parallel operation like it would happen with an ordinary product of vectors.

Note

The computation of the scalar product implicitly applies the `cdt` transform, which implies that the scalar products corresponding to the given geometries are returned for `acomp`, `rcomp`, `aplus`, `rplus`-objects. Even a useful scalar product for factors is induced in this way.

Author(s)


Examples

```r
classical(vector(x[1:10]))
classical(vector(x[1:10]), method = "spline")
classical(vector(x[1:10]), method = "loess")
classical(vector(x[1:10]), method = "three")
```

Description

The dataset is standardized by optional scaling and centering.

Usage

```r
scale(x, center = TRUE, scale = TRUE,...)
## Default S3 method:
scale(x,center=TRUE, scale=TRUE,...)
## S3 method for class 'acomp'
scale(x,center=TRUE, scale=TRUE,...,robust=getOption("robust"))
## S3 method for class 'rcomp'
scale(x,center=TRUE, scale=TRUE,...,robust=getOption("robust"))
```
## S3 method for class 'aplu' scale(x, center=TRUE, scale=TRUE,..., robust=getOption("robust"))

## S3 method for class 'rplu' scale(x, center=TRUE, scale=TRUE,..., robust=getOption("robust"))

## S3 method for class 'rmult' scale(x, center=TRUE, scale=TRUE,..., robust=getOption("robust"))

### Arguments

- **x**: a dataset or a single vector of some type
- **center**: logical value or the center to be subtracted.
- **scale**: logical value or a scaling factor to for multiplication.
- **robust**: A robustness description. See robustnessInCompositions for details.
- **...**: added for generic generality

### Details

scaling is defined in different ways for the different data types. It is always performed as an operation in the enclosing vector space. In all cases an independent scaling of the different coordinates is not always appropriate. This is only done for rplus and rmult geometries. The other three geometries are treated with a global scaling, keeping the relative variations of every part/amount.

The scaling factors can be a matrix (for cdt or idt space), a scalar, or for the r* geometries vector for scaling the entries individually. However scaling the entries individually does not make sense in the a* geometries. The operation achieve in the r*-geometries is indeed the centering of the a*-geometries.

### Value

a vector or data matrix, as x and with the same class, but acordingly transformed.

### Note

Note that the "rcomp" and "rplus" objects does not preserve their geometry during scaling and are therefore reported as "rmult" objects.

See the documentation in package base for details on scale and scale.default. These functions are only modified to allow the additional robustness parameter.

### Author(s)


### See Also

split(base)
Sediments

Sediments

Proportions of sand, silt and clay in sediments specimens

Description

Data provide sand, silt and clay compositions of 21 sediments specimens, 10 of which are identified as offshore, 7 as near shore and 4 new samples.

Usage

data(Sediments)

Format

• sandnumeric: the portion of sand
• siltnumeric: the portion of silt
• claynumeric: the portion of clay
• typenumeric: 1 for offshore, 2 for near shore and 3 for new samples

Details

The data comprise 21 cases: 10 offshore, 7 near shore and 4 new samples, and 4 variables: sand, silt and clay proportions and in addition the type of sediments specimens – 1 for offshore, 2 for near shore and 3 for new samples.

All 3-part compositions sum to one.

Note

Courtesy of J. Aitchison
Source

Aitchison: CODA microcomputer statistical package, 1986, the file name YATQUAD.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


segments

Draws straight lines from point to point.

Description

The function draws lines from a points x to a point y in the given geometry.

Usage

```r
## S3 method for class 'acomp'
segments(x0, y, ..., steps = 30, aspanel = FALSE)
## S3 method for class 'rcomp'
segments(x0, y, ..., steps = 30, aspanel = FALSE)
## S3 method for class 'aplus'
segments(x0, y, ..., steps = 30, aspanel = FALSE)
## S3 method for class 'rplus'
segments(x0, y, ..., steps = 30, aspanel = FALSE)
## S3 method for class 'rmult'
segments(x0, y, ..., steps = 30, aspanel = FALSE)
```

Arguments

- `x0` dataset of points (of the given type) to draw the line from
- `y` dataset of points (of the given type) to draw the line to
- `...` further graphical parameters
- `steps` the number of discretisation points to draw the segments, since the representation might not visually be a straight line
- `aspanel` Logical, indicates use as slave to do acutal drawing only.

Details

The functions add lines to the graphics generated with the corresponding plot functions.

Adding to multipaneled plots redraws the plot completely, and is only possible when the plot has been created with the plotting routines from this library.
Author(s)


See Also

plot.acomp, lines.acomp

Examples

data(SimulatedAmounts)

plot(acomp(sa.lognormals))
segments.acomp(acomp(c(1,2,3)), acomp(c(2,3,1)), col="red")  
segments.rcomp(acomp(c(1,2,3)), acomp(c(2,3,1)), col="blue")

plot(aplus(sa.lognormals[,1:2]))
segments.aplus(aplus(c(10,20)), aplus(c(20,10)), col="red")
segments.rplus(rplus(c(10,20)), rplus(c(20,10)), col="blue")

plot(rplus(sa.lognormals[,1:2]))
segments.aplus(aplus(c(10,20)), aplus(c(20,10)), col="red")
segments.rplus(rplus(c(10,20)), rplus(c(20,10)), col="blue")
Details

The data consist of 36 cases: 14 with known disease A, 16 with known disease B, and 6 new cases and 5 v variables: a, b, c, and d for 4 serum proteins and Type for the diseases: 1 for disease A, 2 for disease B, and 3 for new cases. All row serum proteins proportions sums to 1 except some rounding errors.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name SERPROT.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


---

**ShiftOperators**  
*Shifts of machine operators*

**Description**

Compositions of eight-hours shifts of 27 machine operators.

**Usage**

```r
data(ShiftOperators)
```

**Details**

A study of the activities of 27 machine operators during their eight-hours shifts has been conducted, and proportions of time spend in the following categories:

A: high-quality production,  
B: low-quality production,  
C: machine setting,  
D: machine repair,

are recorded. Of particular interest are any insights which such data might give of relationships between productive and nonproductive parts of such shifts. All compositions sum up one except for rounding error.

Note

Courtesy of J. Aitchison
Source

Aitchison: CODA microcomputer statistical package, 1986, the file name SHIFT.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


Ternary diagrams

Description

Displaying compositions in ternary diagrams

Usage

```r
simpleMissingSubplot(loc, portions, labels=NULL, 
  col=c("white","yellow","red","green","blue"),
  ..., border="gray60", vertical=NULL, xpd=NA)
```

Arguments

- `loc`: a vector of the form c(x1,x2,y1,y2) giving the drawing rectangle for the subplot in coordinates as in `par("usr")`. I.e. if the plot is logrithmic the base 10 logarithm is to be used:
  - x1 left boundary of drawing rectangle
  - y1 lower boundary of drawing rectangle
  - x2 right boundary of drawing rectangle
  - y2 upper boundary of drawing rectangle
- `portions`: The portions of different missing categories
- `labels`: The labels for the categories.
- `col`: The colors to plot the different categories.
- `...`: further graphical parameters passed to `text`
- `border`: The color to draw the borders of the rectangles.
- `vertical`: Should a horizontal or a vertical plot be produced. If NULL the choice is done automatically according to the size of the recangle provided.
- `xpd`: extended plot region. See `par("xpd")`.

Details

This function is typically not called directly, however it could in principle be used to add to plots. The user will modify the function call only to modify the appearance of the missing plot.

The labels are only plotted for nonzero portions. In this way it is always possible to realize the presence of a given missing type, even if it is a too small portion to be actually displayed. In case of overplotting of different labels a further investigation using `missingSummary` should be used.
SimulatedAmounts

Author(s)

See Also
plot.aplus

Examples

data(SimulatedAmounts)
plot(apcomp(sa.missings))
plot(apcomp(sa.missings),mp=simpleMissingSubplot(c(0,0.1,0.2,1),
    missingInfo[c(1,3:5,2)],
    c("Not Missing",paste("Missing Only:",cn),"Totally Missing"),
    col=c("gray","red","green","blue","darkgray"))
)
ms <- missingSummary(sa.missings)
for( i in 1:3 )
    simpleMissingSubplot(c(0.9+0.03*(i-1),0.9+0.03*i,0.2,1), ms[i,])

SimulatedAmounts  Simulated amount datasets

Description
Several simulated datasets intended as reference examples for various conceptual and statistical
models of compositions and amounts.

Usage
data(SimulatedAmounts)

Format
Data matrices with 60 cases and 3 or 5 variables.

Details
The statistical analysis of amounts and compositions is set to discussion. Four essentially different
approaches are provided in this package around the classes "rplus", "aplus", "rcomp", "acomp". There is no absolutely "right" approach, since there is a conection between these approaches and the
processes originating the data. We provide here simulated standard datasets and the corresponding
simulation procedures following these several models to provide "good" analysis examples and to
show how these models actually look like in data.

The data sets are simulated according to correlated lognormal distributions (sa.lognormals, sa.lognormal5),
winsorised correlated normal distributions (sa.tnormals, sa.tnormal5), Dirichlet distribution on the
simplex (sa.dirichlet, sa.dirichlet5), uniform distribution on the simplex (sa.uniform, sa.uniform5),
and a grouped dataset (sa.groups, sa.groups5) with three groups (given in sa.groups.area and sa.groups5.area) all distributed accordingly with a lognormal distribution with group-dependent means.

We can imagine that amounts evolve in nature e.g. in part of the soil they are diluted and transported in a transport medium, usually water, which comes from independent source (the rain, for instance) and this new composition is normalized by taking a sample of standard size. For each of the datasets sa.X there is a corresponding sa.X.dil dataset which is build by simulating exactly that process on the corresponding sa.X dataset. The amounts in the sa.X.dil are given in ppm. This idea of a transport medium is a major argument for a compositional approach, because the total amount given by the sum of the parts is induced by the dilution given by the medium and thus non-informative for the original process investigated.

If we imagine now these amounts flowing into a river and sedimenting, the different contributions are accumulated along the river and renormalized to a unit portion on taking samples again. For each of the dataset sa.X.dil there is a corresponding sa.X.mix dataset which is built from the corresponding sa.X dataset by simulating exactly that accumulation process. Mixing of different compositions is a major argument against the log based approaches (aplus, acomp) since mixing is a highly nonlinear operation in terms of log-ratios.

Author(s)


Source

The datasets are simulated for this package and are under the GNU Public Library Licence Version 2 or newer.

References

http://www.stat.boogaart.de/compositions/data


Examples

data(SimulatedAmounts)
plot.acomp(sa.lognormals)
plot.acomp(sa.lognormals.dil)
SimulatedAmounts

plot.acomp(sa.lognormals.mix)
plot.acomp(sa.lognormals5)
plot.acomp(sa.lognormals5.dil)
plot.acomp(sa.lognormals5.mix)
plot(acomp(sa.missings))
plot(acomp(sa.missings5))

#library(MASS)
plot.rcomp(sa.tnormals)
plot.rcomp(sa.tnormals.dil)
plot.rcomp(sa.tnormals.mix)
plot.rcomp(sa.tnormals5)
plot.rcomp(sa.tnormals5.dil)
plot.rcomp(sa.tnormals5.mix)

plot.acomp(sa.groups,col=as.numeric(sa.groups.area),pch=20)
plot.acomp(sa.groups.dil,col=as.numeric(sa.groups.area),pch=20)
plot.acomp(sa.groups.mix,col=as.numeric(sa.groups.area),pch=20)
plot.acomp(sa.groups5,col=as.numeric(sa.groups.area),pch=20)
plot.acomp(sa.groups5.dil,col=as.numeric(sa.groups.area),pch=20)
plot.acomp(sa.groups5.mix,col=as.numeric(sa.groups.area),pch=20)

plot.acomp(sa.uniform)
plot.acomp(sa.uniform.dil)
plot.acomp(sa.uniform.mix)
plot.acomp(sa.uniform5)
plot.acomp(sa.uniform5.dil)
plot.acomp(sa.uniform5.mix)

plot.acomp(sa.dirichlet)
plot.acomp(sa.dirichlet.dil)
plot.acomp(sa.dirichlet.mix)
plot.acomp(sa.dirichlet5)
plot.acomp(sa.dirichlet5.dil)
plot.acomp(sa.dirichlet5.mix)

# The data was simulated with the following commands:

#library(MASS)
dilution <- function(x) {clo(cbind(x,exp(rnorm(nrow(x),5,1))))[,1:ncol(x)]*1E6}
seqmix   <- function(x) {clo(apply(x,2,cumsum))*1E6}

vars <- c("Cu","Zn","Pb")
vars5 <- c("Cu","Zn","Pb","Cd","Co")

sa.lognormals <- structure(exp(matrix(rnorm(3*60),ncol=3) %*% chol(matrix(c(1,0.8,-0.2,0.8,1,
-0.2,-0.2,-0.2,1),ncol=3))+matrix(rep(c(1:3),each=60),ncol=3)),
dimnames=list(NULL,vars))
```r
# Simulated Amounts

plot.acomp(sa.lognormals)
pairs(sa.lognormals)

sa.lognormals.dil <- dilution(sa.lognormals)
plot.acomp(sa.lognormals.dil)
pairs(sa.lognormals.dil)

sa.lognormals.mix <- seqmix(sa.lognormals.dil)
plot.acomp(sa.lognormals.mix)
pairs(sa.lognormals.mix)

sa.lognormals5 <- structure(exp(matrix(rnorm(5*60),ncol=5) %*
chol(matrix(c(1,0.8,-0.2,0.8,0,
0.8,1,-0.2,0,0,
-0.2,-0.2,1,0,0,
0,0,0,5,4.9,
0,0,0,4.9,5),ncol=5)) +
matrix(rep(c(1,3,-2,-2),each=60),ncol=5)),
dimnames=list(NULL,vars5))

plot.acomp(sa.lognormals5)
pairs(sa.lognormals5)

sa.lognormals5.dil <- dilution(sa.lognormals5)
plot.acomp(sa.lognormals5.dil)
pairs(sa.lognormals5.dil)

sa.lognormals5.mix <- seqmix(sa.lognormals5.dil)
plot.acomp(sa.lognormals5.mix)
pairs(sa.lognormals5.mix)

sa.groups.area <- factor(rep(c("Upper","Middle","Lower"),each=20))
sa.groups <- structure(exp(matrix(rnorm(3*20*3),ncol=3) %*
chol(0.5*matrix(c(1,0.8,-0.2,0.8,1,
-0.2,-0.2,-0.2,1),ncol=3)) +
matrix(rep(c(1,2,2.5,2,2.9,5,4,2,5),
each=20),ncol=3)),
dimnames=list(NULL,c("clay","sand","gravel")))

plot.acomp(sa.groups,col=as.numeric(sa.groups.area),pch=20)
pairs(sa.lognormals,col=as.numeric(sa.groups.area),pch=20)

sa.groups.dil <- dilution(sa.groups)
plot.acomp(sa.groups.dil,col=as.numeric(sa.groups.area),pch=20)
pairs(sa.groups.dil,col=as.numeric(sa.groups.area),pch=20)

sa.groups.mix <- seqmix(sa.groups.dil)
plot.acomp(sa.groups.mix,col=as.numeric(sa.groups.area),pch=20)
pairs(sa.groups.mix,col=as.numeric(sa.groups.area),pch=20)
```
sa.groups5.area <- factor(rep(c("Upper","Middle","Lower"),each=20))
sa.groups5 <- structure(exp(matrix(rnorm(5*20*3),ncol=5) **%
    chol(matrix(c(1,0.8,-0.2,0,0,
    0.8,1,-0.2,0,0,
    -0.2,-0.2,1,0,0,
    0,0,0,5,4.9,
    0,0,0,4.9,5),ncol=5))+
    matrix(rep(c(1,2,2.5,
    2,2.9,5,
    4,2.5,0,
    -2,-1,-1,
    -1,-2,-3),
    each=20),ncol=5)),
    dimnames=list(NULL,
    vars5))

plot.acomp(sa.groups5,col=as.numeric(sa.groups5.area),pch=20)
pairs(sa.groups5,col=as.numeric(sa.groups5.area),pch=20)

sa.groups5.dil <- dilution(sa.groups5)
plot.acomp(sa.groups5.dil,col=as.numeric(sa.groups5.area),pch=20)
pairs(sa.groups5.dil,col=as.numeric(sa.groups5.area),pch=20)

sa.groups5.mix <- seqmix(sa.groups5.dil)
plot.acomp(sa.groups5.mix,col=as.numeric(sa.groups5.area),pch=20)
pairs(sa.groups5.mix,col=as.numeric(sa.groups5.area),pch=20)

sa.tnormals <- structure(pmax(matrix(rnorm(3*60),ncol=3) **%
    chol(matrix(c(1,0.8,-0.2,0,0.8,1,
    -0.2,-0.2,-0.2,1),ncol=3))+
    matrix(rep(c(0:2),each=60),ncol=3),0),
    dimnames=list(NULL,c("clay","sand","gravel")))

plot.rcomp(sa.tnormals)
pairs(sa.tnormals)

sa.tnormals.dil <- dilution(sa.tnormals)
plot.acomp(sa.tnormals.dil)
pairs(sa.tnormals.dil)

sa.tnormals.mix <- seqmix(sa.tnormals.dil)
plot.acomp(sa.tnormals.mix)
pairs(sa.tnormals.mix)

sa.tnormals5 <- structure(pmax(matrix(rnorm(5*60),ncol=5) **%
    chol(matrix(c(1,0.8,-0.2,0,0,
    0.8,1,-0.2,0,0,
    -0.2,-0.2,1,0,0,
    0,0,0,5,4.9,
    0,0,0,4.9,5),ncol=5))+
    matrix(rep(c(1,2,2.5,
    2,2.9,5,
    4,2.5,0,
    -2,-1,-1,
    -1,-2,-3),
    each=20),ncol=5)),
    dimnames=list(NULL,
    vars5))
-0.2,-0.2,1,0,0,
0,0,0,0.05,0.049,
0,0,0,0.049,0.05),ncol=5)) +
matrix(rep(c(0:2,0.1,0.1),each=60),ncol=5),0),
dimnames=list(NULL,
vars5))

plot.rcomp(sa.tnormals5)
pairs(sa.tnormals5)

sa.tnormals5.dil <- dilution(sa.tnormals5)
plot.acomp(sa.tnormals5.dil)
pairs(sa.tnormals5.dil)

sa.tnormals5.mix <- seqmix(sa.tnormals5.dil)
plot.acomp(sa.tnormals5.mix)
pairs(sa.tnormals5.mix)

sa.dirichlet <- sapply(c(clay=0.2,sand=2,gravel=3),rgamma,n=60)
colnames(sa.dirichlet) <- vars
plot.acomp(sa.dirichlet)
pairs(sa.dirichlet)

sa.dirichlet.dil <- dilution(sa.dirichlet)
plot.acomp(sa.dirichlet.dil)
pairs(sa.dirichlet.dil)

sa.dirichlet.mix <- seqmix(sa.dirichlet.dil)
plot.acomp(sa.dirichlet.mix)
pairs(sa.dirichlet.mix)

sa.dirichlet5 <- sapply(c(clay=0.2,sand=2,gravel=3,humus=0.1,plant=0.1),rgamma,n=60)
colnames(sa.dirichlet5) <- vars5
plot.acomp(sa.dirichlet5)
pairs(sa.dirichlet5)

sa.dirichlet5.dil <- dilution(sa.dirichlet5)
plot.acomp(sa.dirichlet5.dil)
pairs(sa.dirichlet5.dil)

sa.dirichlet5.mix <- seqmix(sa.dirichlet5.dil)
plot.acomp(sa.dirichlet5.mix)
pairs(sa.dirichlet5.mix)

sa.uniform <- sapply(c(clay=1,sand=1,gravel=1),rgamma,n=60)
colnames(sa.uniform) <- vars
plot.acomp(sa.uniform)
pairs(sa.uniform)

sa.uniform.dil <- dilution(sa.uniform)
plot.acomp(sa.uniform.dil)
pairs(sa.uniform.dil)

sa.uniform.mix <- seqmix(sa.uniform.dil)
plot.acomp(sa.uniform.mix)
pairs(sa.uniform.mix)

sa.uniform5 <- sapply(c(clay=1,sand=1,gravel=1,humus=1,plant=1),rgamma,n=60)
colnames(sa.uniform5) <- vars5
plot.acomp(sa.uniform5)
pairs(sa.uniform5)

sa.uniform5.dil <- dilution(sa.uniform5)
plot.acomp(sa.uniform5.dil)
pairs(sa.uniform5.dil)

sa.uniform5.mix <- seqmix(sa.uniform5.dil)
plot.acomp(sa.uniform5.mix)
pairs(sa.uniform5.mix)

tmp<-set.seed(1400)
A <- matrix(c(0.1,0.2,0.3,0.1),nrow=2)
Mvar <- 0.1*ilrvar2clr(A %*% t(A))
Mcenter <- acomp(c(1,2,1))
typicalData <- rnorm.acomp(100,Mcenter,Mvar) # main population
colnames(typicalData)<-c("A","B","C") # A dataset without outliers
sa.outliers1 <- acomp(rnorm.acomp(100,Mcenter,Mvar))
# A dataset with 10% data with a large error in the first component
sa.outliers2 <- acomp(rbind(typicalData+rbinom(100,1,p=0.1)*rnorm(100)*acomp(c(4,1,1))))
# A dataset with a single outlier
sa.outliers3 <- acomp(rbind(typicalData,acomp(c(0.5,1.5,2))))
colnames(sa.outliers3)<-colnames(typicalData)
tmp<-set.seed(30)
rcauchy.acomp <- function (n, mean, var){
  D <- gsi.getD(mean)-1
  perturbe(ilrInv(matrix(rnorm(n*D)/rep(rnorm(n),D), ncol = D) %*% chol(clrvar2ilr(var))), mean)
}
# A dataset with a Cauchy type distribution
sa.outliers4 <- acomp(rcauchy.acomp(100,acomp(c(1,2,1)),Mvar/4))
colnames(sa.outliers4)<-colnames(typicalData)
# A dataset with like sa.outlier2 but a differently strong distortions
sa.outliers5 <- acomp(rbind(unclass(typicalData)+outer(rbinom(100,1,p=0.1)*runif(100),c(0.1,1,2))))
# A dataset with a second population
sa.outliers6 <- acomp(rbind(typicalData,rnorm.acomp(20,acomp(c(4,4,1)),Mvar)))
# Missings
sa.missings <- simulateMissings(sa.lognormals, dl=0.05, MAR=0.05, MNAR=0.05, SZ=0.05)
sa.missings[5,2] <- BDLvalue

sa.missings5 <- simulateMissings(sa.lognormals5, dl=0.05, MAR=0.05, MNAR=0.05, SZ=0.05)
sa.missings5[5,2] <- BDLvalue

objects(pattern="sa.*")

---

**simulatemissings**

*Artificial simulation of various kinds of missings/polluted data*

**Description**

These are simulation mechanisms to check that missing techniques perform in sensible ways. They just generate additional missings of the various types in a given dataset, according to a specific process.

**Usage**

```r
simulateMissings(x, dl=NULL, knownlimit=FALSE,
                 MARprob=0.0, MNARprob=0.0, mnarity=0.5, SZprob=0.0)
```

```r
observeWithAdditiveError(x, sigma=dl/dlf, dl=sigma*dlf, dlf=3,
                          keepObs=FALSE, digits=NA, obsScale=1,
                          class="acomp")
```

**Arguments**

- **x**: a dataset that should get the missings
- **dl**: the detection limit described in `clo`, to impose an artificial detection limit
- **knownlimit**: a boolean indicating whether the actual detection limit is still known in the dataset.
- **MARprob**: the probability of occurrence of 'Missings At Random' values
- **MNARprob**: the probability of occurrence of 'Missings Not At Random'. The tendency is that small values have a higher probability to be missed.
- **mnarity**: a number between 0 and 1 giving the strength of the influence of the actual value in becoming a MNAR. 0 means a MAR like behavior and 1 means that it is just the smallest values that is lost
- **SZprob**: the probability to obtain a structural zero. This is done at random like a MAR.
- **sigma**: the standard deviation of the normal distributed extra additive error
- **dlf**: the distance from 0 at which a datum will be considered BDL
- **keepObs**: should the (closed) data without additive error be returned as an attribute?
- **digits**: rounding to be applied to the data with additive error (see Details)
**Details**

Without any additional parameters no missings are generated. The procedure to generate MNAR affects all variables.

Function "simulateMissings" is a multipurpose simulator, where each class of missing value is treated separately, and where detection limits are specified as thresholds.

Function "observeWithAdditiveError" simulates data within a very specific framework, where an additive error of \( \text{sd}=\sigma \) is added to the input data \( x \), and BDLs are generated if a datum is less than \( \text{dfl} \times \sigma \). Afterwards, the resulting data are rounded as \( \text{round(data/obsScale,digits)} \times \text{obsScale} \), i.e. a certain observation scale \( \text{obsScale} \) is chosen, and at that scale, only some digits are kept. This framework is typical of chemical analyses, and it generates both BDLs and pollution/rounding of (apparently) "right" data.

**Value**

A dataset like \( x \) but with some additional missings.

**Author(s)**

K.Gerald van den Boogaart

**References**


See compositions.missings for more details.

**See Also**

compositions.missings

**Examples**

```r
data(SimulatedAmounts)
x <- acomp(sa.lognormals)
xnew <- simulateMissings(x,d1=0.05,MAR=0.05,MNAR=0.05,SZ=0.05)
acomp(xnew)
plot(missingSummary(xnew))
```
Skulls  

Measurement of skulls

Description

Usage
data(Skulls)

Details
As a part of a study of seventeenth-century English skulls three angles of a triangle in the cranium

\[
\begin{align*}
N: & \text{ nasial angle}, \\
A: & \text{ alveolar angle}, \\
B: & \text{ basilar angle},
\end{align*}
\]

were measured for 22 female and 29 male skulls. These, together with similar measurement of 22 female and 29 male skulls of the Naqada race are presented. The general objective is to investigate possible sex and race differences in skull shape. The angles sums in the row are all equal to 180 degrees.

Note
Courtesy of J. Aitchison

Source
Aitchison: CODA microcomputer statistical package, 1986, the file name SKULLS.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References

---

SkyeAFM  

AFM compositions of 23 aphyric Skye lavas

Description
AFM compositions of 23 aphyric Skye lavas. AFM diagrams formed from the relative proportions of A: alkali or Na2O + K2O, F: Fe2O3, and M: MgO, are common in geochemistry.
split

Usage

```r
data(SkyeAFM)
```

Details

AFM compositions of 23 aphyric Skye lavas. AFM diagrams formed from the relative proportions
of tive proportions of A: alkali or Na2O + K2O, F: Fe2O3, and M: MgO. Adapted from Thompson,
Esson and Duncan: Major element chemical variations in the Eocene lavas of the Isle of Skye,
Scotland. All row percentage sums to 100.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name SKYEAFM.DAT, here
included under the GNU Public Library Licence Version 2 or newer.

References

Thompson, Esson and Duncan: Major element chemical variations in the Eocene lavas of the Isle

See Also

Skye

---

### split

#### Splitting datasets in groups given by factors

**Description**

Splits data sets of compositions in groups given by factors, and gives the same class as the data to
the result.

**Usage**

```r
## S3 method for class 'acomp'
split(x,f,drop=FALSE,...)
## S3 method for class 'rcomp'
split(x,f,drop=FALSE,...)
## S3 method for class 'aplus'
split(x,f,drop=FALSE,...)
## S3 method for class 'rplus'
split(x,f,drop=FALSE,...)
```
## S3 method for class 'rmult'

```r
split(x,f,drop=FALSE,...)
```

## S3 method for class 'ccomp'

```r
split(x,f,drop=FALSE,...)
```

### Arguments

- **x**: a dataset or a single vector of some type
- **f**: a factor that defines the grouping or a list of factors
- **drop**: drop=FALSE also gives (empty) datasets for empty categories
- **...**: Further arguments passed to `split.default`. Currently (and probably) without any use.

### Value

A list of objects of the same type as `x`.

### Author(s)


### See Also

`split`

### Examples

```r
data(SimulatedAmounts)
split(acomp(sa.groups),sa.groups.area)
lapply( split(acomp(sa.groups),sa.groups.area), mean)
```

---

### Description

The function draws lines in a given direction `d` through points `x`.

### Usage

```r
straight(x,...)
```

```r
# S3 method for class 'acomp'
straight(x,d,...,steps=30,aspanel=FALSE)
```

```r
# S3 method for class 'rcomp'
straight(x,d,...,steps=30,aspanel=FALSE)
```

```r
# S3 method for class 'aplus'
```
straight(x,d,...,steps=30,aspanel=FALSE)
## S3 method for class 'rplus'
straight(x,d,...,steps=30,aspanel=FALSE)
## S3 method for class 'rmult'
straight(x,d,...,steps=30,aspanel=FALSE)

### Arguments

- **x**: dataset of points of the given type to draw the line through
- **d**: dataset of directions of the line
- **...**: further graphical parameters
- **steps**: the number of discretisation points to draw the segments, since the representation might not visually be a straight line
- **aspanel**: Logical, indicates use as slave to do actual drawing only.

### Details

The functions add lines to the graphics generated with the corresponding plot functions. Adding to multipaneled plots redraws the plot completely, and is only possible when the plot has been created with the plotting routines from this library. Lines end when they leave the space (e.g. the simplex), which sometimes leads to the impression of premature end (specially in rcomp geometry).

### Author(s)


### See Also

plot.acomp, lines.acomp

### Examples

```r
data(SimulatedAmounts)
plot(acomp(sa.lognormals))
straight(mean(acomp(sa.lognormals)),
         princomp(acomp(sa.lognormals))$Loadings[1,],
         col="red")
straight(mean(rcomp(sa.lognormals)),
         princomp(rcomp(sa.lognormals))$loadings[,1],
         col="blue")

plot(aplus(sa.lognormals[,1:2]))
straight(mean(aplus(sa.lognormals[,1:2])),
         princomp(aplus(sa.lognormals[,1:2]))$Loadings[1,],
         col="red")
straight(mean(rplus(sa.lognormals[,1:2])),
         princomp(rplus(sa.lognormals[,1:2]))$loadings[,1],
```
summary.acomp

Summarizing a compositional dataset in terms of ratios

Description

Summaries in terms of compositions are quite different from classical ones. Instead of analysing each variable individually, we must analyse each pair-wise ratio in a log geometry.

Usage

## S3 method for class 'acomp'
summary(object, ..., robust=getOption("robust"))

Arguments

- object: a data matrix of compositions, not necessarily closed
- ...: not used, only here for generics
- robust: A robustness description. See robustnessInCompositions for details. The parameter can be null for avoiding any estimation.

Details

It is quite difficult to summarize a composition in a consistent and interpretable way. We tried to provide such a summary here, based on the idea of the variation matrix.

Value

The result is an object of type "summary.acomp"

- mean: the mean.acomp composition
- mean.ratio: a matrix containing the geometric mean of the pairwise ratios
- variation: the variation matrix of the dataset (variation.acomp)
- expsd: a matrix containing the one-sigma factor for each ratio, computed as exp(sqrt(variation.acomp(W))). To obtain a two-sigma-factor, one has to take its squared value (power 1.96, actually).
invexpsd
the inverse of the preceding one, giving the reverse bound. Additionally, it can be "almost" interpreted as a correlation coefficient, with values near one indicating high proportionality between the components.

min
a matrix containing the minimum of each of the pairwise ratios

q1
a matrix containing the 1-Quartile of each of the pairwise ratios

median
a matrix containing the median of each of the pairwise ratios

q3
a matrix containing the 3-Quartile of each of the pairwise ratios

max
a matrix containing the maximum of each of the pairwise ratios

Author(s)

References

See Also
acomp

Examples

data(SimulatedAmounts)
summary(acomp(sa.lognormals))

Description
Summary of a vector of amounts, according to its underlying geometry.

Usage

## S3 method for class 'aplus'
summary( object, ..., digits=max(3, getOption("digits")-3), robust=NULL)
## S3 method for class 'rplus'
summary( object, ..., robust=NULL)
## S3 method for class 'rmult'
summary( object, ..., robust=NULL)
summary.rcomp

Arguments

object
an aplus/rplus set of amounts

digits
the number of significant digits to be used. The argument can also be used with rplus/rmult.

... not used, only here for generics

robust
A robustness description. See robustnessInCompositions for details. The option is currently not supported. If support is added the default will change to getOption(robust).

Details

The obtained value is the same as for the classical summary summary, although in the case of aplus objects, the statistics have been computed in a logarithmic geometry, and exponentiated afterwards (which just changes the mean, equivalent to the geometric mean of the data set).

Value

A matrix containing summary statistics (minimum, the three quantiles, the mean and the maximum) of each component.

Author(s)


See Also

aplus,rplus,summary.acomp,summary.rcomp

Examples

```r
data(SimulatedAmounts)
summary(aplus(sa.lognormals))
summary(aplus(sa.tnormals))
summary(rplus(sa.lognormals))
summary(rplus(sa.tnormals))
summary(rmult(sa.lognormals))
```

summary.rcomp  Summary of compositions in real geometry

Description

Compute a summary of a composition based on real geometry.
Usage

```r
## S3 method for class 'rcomp'
summary( object, ... , robust=NULL)
```

Arguments

- **object**: an `rcomp` dataset of compositions
- **...**: further arguments to `summary`
- **robust**: A robustness description. See `robustnessInCompositions` for details. The option is currently not supported. If support is added the default will change to `getOption(robust)`.

Details

The data is applied a `clo` operation before the computation. Note that the statistics obtained will not keep any consistency if computed with all the parts available or only with a subcomposition.

Value

A matrix containing summary statistics. The value is the same as for the classical summary `summary` applied to a closed dataset.

Author(s)


See Also

- `rcomp`, `summary.aplus`, `summary.acomp`

Examples

```r
data(SimulatedAmounts)
summary(rcomp(sa.lognormals))
summary(rcomp(sa.tnormals))
```

Description

Routines to compute the global projector to the observed subspace, down-weighting the subspaces with more missing values.
Usage

sumMissingProjector(x,...)
## S3 method for class 'acomp'
sumMissingProjector(x,has=is.NMV(x),...)
## S3 method for class 'aplus'
sumMissingProjector(x,has=is.NMV(x),...)
## S3 method for class 'rcomp'
sumMissingProjector(x,has=!(is.MAR(x)|is.MNAR(x)),...)
## S3 method for class 'rplus'
sumMissingProjector(x,has=!(is.MAR(x)|is.MNAR(x)),...)
## S3 method for class 'rmult'
sumMissingProjector(x,has=is.finite(x),...)

Arguments

x a dataset of some type containing missings
has the values to be regarded as non missing
... further generic arguments that might be useful for other functions.

Details

The function missingProjector generates a list of N square matrices of dimension DxD (with N and D respectively equal to the number of rows and columns in x). Each of these matrices gives the projection of a data row onto its observed sub-space. Then, the function sumMissingProjector takes all these matrices and sums them in an efficient way, generating a "summary" of observed sub-spaces.

Value

The matrix of rotation/re-weighting of the original data set, down-weighting the subspaces with more missing values. This matrix is useful to obtain estimates of the mean (and variance, in the future) still unbiased in the presence of lost values (only of type MAR, strictly-speaking, but anyway useful for any type of missing value, when used with care). This matrix is the Fisher Information in the presence of missing values.

Missing Policy

No missing policy is given by the routine itself. Its treatment of missing values depends on the "has" argument.

Author(s)

Supervisor

References

See Also
missingProjector, clr, rcomp, aplus, princomp, acomp, plot.acomp, barplot.acomp, mean.acomp, var.acomp, variation.acomp, cov.acomp, msd

Examples
data(SimulatedAmounts)
sumMissingProjector(acomp(sa.lognormals))
sumMissingProjector(acomp(sa.tnormals))

Supervisor

Proportions of supervisor's statements assigned to different categories

Description
The results of a study of a single supervisor in his relationship to three supervisee are recorded. Instructions in a technical subject took place in sessions of one hour and with only one supervisee at the time. Each supervisee attended six sessions (once every two weeks in a twelve-week period). All of 18 sessions were recorded and for each session the 'statements' of the supervisor were classified into four categories. Thus for each session the proportion of statements in the four categories are set out in a two-way table according to the fortnight (6) and the supervisee (3).

Usage
data(Supervisor)

Format
A 18x13 matrix

Details
For each session the 'statements' of the supervisor were classified into four categories
• C: commanding, posing a specific instruction to the supervisee,
• D: demanding, posing a specific question to the supervisee,
• E: exposing, providing the supervisee with an explanation,
• F: faulting, pointing out faulty technique to the supervisee.

Thus for each session the proportion of statements in the four categories are set out in a two-way table according to the fortnight (6) and the supervisee (3). The C, D, E, F values in the rows sum mostly to 1, except for some rounding errors.
ternaryAxis

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name SUPERVIS.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


ternaryAxis

Axis for ternary diagrams

Description

Displaying compositions in ternary diagrams

Usage

ternaryAxis(side=1:3, at=seq(0.2, 0.8, by=0.2),
labels=if(is.list(at)) lapply(at, format) else format(at),
..., tick=TRUE, pos=0,
font.axis=par("font.axis"),
font.lab=par("font.lab"),
ltb="solid", lwd=1,
len.tck=0.025, dist.lab=0.03,
dist.axis=0.03,
ltb.tck="solid",
col.axis=par("col.axis"),
col.lab=par("col.lab"),
cex.axis=par("cex.axis"),
cex.lab=par("cex.lab"),
Xlab=NULL, Ylab=NULL, Zlab=NULL, small=TRUE,
xpd=NA, aspanel=FALSE)

Arguments

side a vector giving the sides to draw the axis on.  1=under the plot, 2=the upper right axis, 3=the upper left axis.  -1 is the portion axis of the first component, -2 is the portion axis of the second component, -3 is the portion axis of the third component.  An empty vector or 0 suppresses axis plotting, but still plots the Xlab, Ylab and Zlab parameters.

at a vector or a list of vectors giving the positions of the tickmarks.
ternaryAxis

labels

tick

pos

font.axis

font.lab

lty

lty.tck

len.tck

dist.axis

dist.lab

lwd

col.axis

col.lab

cex.axis

cex.lab

Xlab

Ylab

Zlab

small

xpd

aspanel

...
totals

Author(s)


See Also

plot.aplus, plot3D (for 3D plot), kingTetrahedron (for 3D-plot model export), qnorm.acomp.boxplot.acomp

Examples

data(SimulatedAmounts)
plot(acomp(sa.lognormals), axes=TRUE)
ternaryAxis(side=1:3, pos=0, col.axis="red", col.lab="green")
ternaryAxis(side=1:3, at=1:9/10, 
        labels=expression(9:1, 4:1, 7:3, 3:2, 1:1, 2:3, 3:7, 1:4, 1:9),
        pos=0, col.axis="red", col.lab="green")
ternaryAxis(side=rep(-1:-3,3), labels=paste(seq(20,80,by=20),"%"),
        pos=rep(c(0,0.5,1),each=3), col.axis=1:3, col.lab="green")
ternaryAxis(side=rep(1:3,3), at=1:9/10, 
        labels=expression(9:1, 4:1, 7:3, 3:2, 1:1, 2:3, 3:7, 1:4, 1:9),
        pos=rep(c(0,0.5,1),each=3))

plot(acomp(sa.lognormals5), axes=TRUE)
ternaryAxis(side=1:3, pos=0, col.axis="red", col.lab="green")
ternaryAxis(side=1:3, at=1:9/10, 
        labels=expression(9:1, 4:1, 7:3, 3:2, 1:1, 2:3, 3:7, 1:4, 1:9),
        pos=0, col.axis="red", col.lab="green")

totals

Total sum of amounts

Description

Calculates the total amount by summing the individual parts.

Usage

totals(x,...)
## S3 method for class 'acomp'
totals(x,..., missing.ok=TRUE)
## S3 method for class 'rcomp'
totals(x,..., missing.ok=TRUE)
## S3 method for class 'aplus'
totals(x,..., missing.ok=TRUE)
## S3 method for class 'rplus'
totals(x,..., missing.ok=TRUE)
## S3 method for class 'ccomp'
totals(x,..., missing.ok=TRUE)
Transformations from 'mixtures' to 'compositions' classes

Arguments

x an amount/amount dataset
... not used, only here for generic purposes
missing.ok if TRUE ignores missings; if FALSE issues an error if the total cannot be calculated due to missings.

Value

a numeric vector of length equal to ncol(x) containing the total amounts

Missing Policy

if missing.ok=TRUE missings are just regarded as 0, if missing.ok=FALSE WZERO values is still regarded as 0 and other sorts lead to NA in the respective totals.

Author(s)


See Also

aplus

Examples

data(SimulatedAmounts)
totals(acomp(sa.lognormals))
totals(rcomp(sa.lognormals,total=100))
totals(aplus(sa.lognormals))
totals(rplus(sa.lognormals))
aplus(acomp(sa.lognormals),total=totals(aplus(sa.lognormals))))
tryDebugger

Usage

mix.2aplus(X)
mix.2acomp(X)
mix.2rcomp(X)
mix.2rplus(X)
mix.2rmult(X)

Arguments

X mixture object to be converted

Details

A 'compositions' object is obtained from the mixture object m, having the same data matrix as mixture object m i.e. m$mat.

Value

A 'compositions' object of the class 'aplus', 'acomp', 'rcomp', 'rplus' or 'rmult'.

See Also

aplus acomp rcomp rplus rmult

Examples

## Not run:
m <- mix.Read("Glac.dat")  # reads the Glacial data set from Aitchison (1986)
m <- mix.Extract(m,c(1,2,3,4))  # mix object with closed four parts subcomposition
ap <- mix.2aplus(m)  # ap is a 'compositions' object of the aplus class
ac <- mix.2acomp(m)  # ac is a 'compositions' object of the acomp class

## End(Not run)

tryDebugger

Empirical variograms for compositions

Description

An R-debugger that also works with errors in parameters.

Usage

tryDebugger(dump = last.dump)
**Arguments**

- `dump`: An R dump object created by 'dump.frames'.

**Details**

Works like debugger, with the small exception that it also works in situations of nasty errors, like recursive parameter evaluation, missing parameters, and additional errors in arguments.

**Value**

Nothing.

**Author(s)**


**See Also**

`debugger`

**Examples**

```r
## Not run:
f <- function(x,y=y) {y}
f(1)
tryDebugger() # works
debugger() # Does not allow to browse anything
## End(Not run)
```

---

**ult**

*Uncentered log transform*

**Description**

Compute the uncentered log ratio transform of a (dataset of) composition(s) and its inverse.

**Usage**

- `ult(x,...)`
- `ultInv(z,...)`
- `Kappa(x,...)`
Arguments

x a composition or a data matrix of compositions, not necessarily closed
z the ult-transform of a composition or clr-transforms of compositions (or a data matrix), not necessarily centered
... for generic use only

Details

The ult-transform is simply the elementwise log of the closed composition. The ult has some important properties in the scope of Information Theory of probability vectors (but might be mostly misleading for exploratory analysis of compositions).

Value

ult gives the uncentered log transform,
ultInv gives closed compositions with the given ult/clr-transforms
Kappa gives the difference between the clr and the ult transforms. It is quite linked to information measures.

Author(s)


See Also

ilr,alr,apt

Examples

(tmp <- ult(c(1,2,3)))
ultInv(tmp)
ultInv(tmp) - clo(c(1,2,3)) # 0
data(Hydrochem)
cdata <- Hydrochem[,6:19]
pairs(ult(cdata),pch="."
Kappa(c(1,2,3)))

var.acomp Variances and covariances of amounts and compositions

Description

Compute the (co)variance matrix in the several approaches of compositional and amount data analysis.
Usage

\[
\text{var}(x, \ldots)
\]

## Default S3 method:
\[
\text{var}(x, y = \text{NULL}, \text{na.rm} = \text{FALSE}, \text{use}, \ldots)
\]

## S3 method for class 'acomp'
\[
\text{var}(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'rcomp'
\[
\text{var}(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'aplus'
\[
\text{var}(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'rplus'
\[
\text{var}(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'rmult'
\[
\text{var}(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

cov(x, y = x, \ldots)

## Default S3 method:
\[
cov(x, y = \text{NULL}, \text{use} = \text{"everything"}, \text{method} = \text{c("pearson", "kendall", "spearman")}, \ldots)
\]

## S3 method for class 'acomp'
\[
cov(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'rcomp'
\[
cov(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'aplus'
\[
cov(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'rplus'
\[
cov(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

## S3 method for class 'rmult'
\[
cov(x, y = \text{NULL}, \ldots, \text{robust} = \text{getOption("robust")}, \text{use} = \text{"all.obs"}, \text{giveCenter} = \text{FALSE})
\]

Arguments

- **x**: a dataset, eventually of amounts or compositions
- **y**: a second dataset, eventually of amounts or compositions
- **na.rm**: see var
- **use**: see var
- **method**: see cov
... further arguments to \texttt{var} e.g. use 

\textbf{robust}  
A description of a robust estimator. FALSE for the classical estimators. See \texttt{robustnessInCompositions} for further details.

\textbf{giveCenter}  
If TRUE the center used in the variance calculation is reported as a "center" attribute. This is especially necessary for robust estimations, where a reasonable center can not be computed independently for the me variance calculation.

\section*{Details}
The basic functions of \texttt{var, cov} are turned to S3-generics. The original versions are copied to the default method. This allows us to introduce generic methods to handle variances and covariances of other data types, such as amounts or compositions.

If classed amounts or compositions are involved, they are transformed with their corresponding transforms, using the centered default transform (\texttt{cdt}). That implies that the variances have to be interpreted in a log scale level for \texttt{acomp} and \texttt{aplus}.

We should be aware that variance matrices of compositions (\texttt{acomp} and \texttt{rcomp}) are singular. They can be transformed to the corresponding nonsingular variances of ilr or ipt-space by \texttt{clrvar2ilr}.

In R versions older than v2.0.0, \texttt{var} and \texttt{cov} were defined in package “base” instead of in “stats”. This might produce some misfunction.

\section*{Value}
The variance matrix of \texttt{x} or the covariance matrix of \texttt{x} and \texttt{y}.

\section*{Author(s)}
K.Gerald v.d. Boogaart \url{http://www.stat.boogaart.de}

\section*{See Also}
\texttt{cdt, clrvar2ilr, clo, mean.acomp, acomp, rcomp, aplus, rplus, variation}

\section*{Examples}
\begin{verbatim}
data(SimulatedAmounts)
meanCol(sa.lognormals)
var(acomp(sa.lognormals))
var(rcomp(sa.lognormals))
var(aplus(sa.lognormals))
var(rplus(sa.lognormals))

cov(acomp(sa.lognormals5[,1:3]),acomp(sa.lognormals5[,4:5]))
cov(rcomp(sa.lognormals5[,1:3]),rcomp(sa.lognormals5[,4:5]))
cov(aplus(sa.lognormals5[,1:3]),aplus(sa.lognormals5[,4:5]))
cov(rplus(sa.lognormals5[,1:3]),rplus(sa.lognormals5[,4:5]))
cov(acomp(sa.lognormals5[,1:3]),aplus(sa.lognormals5[,4:5]))

svd(var(acomp(sa.lognormals)))
\end{verbatim}
Description

Compute the variation matrix in the various approaches of compositional and amount data analysis. Pay attention that this is not computing the variance or covariance matrix!

Usage

```r
variation(x, ...)    # S3 method for class 'acomp'
variation(x, ..., robust=getOption("robust"))    # S3 method for class 'rcomp'
variation(x, ..., robust=getOption("robust"))    # S3 method for class 'aplus'
variation(x, ..., robust=getOption("robust"))    # S3 method for class 'rplus'
variation(x, ..., robust=getOption("robust"))    # S3 method for class 'rmult'
variation(x, ..., robust=getOption("robust"))    # S3 method for class 'rmult'
```

Arguments

- `x`: a dataset, eventually of amounts or compositions
- `...`: currently unused
- `robust`: A description of a robust estimator. FALSE for the classical estimators. See `robustnessInCompositions` for further details.

Details

The variation matrix was defined in the `acomp` context of analysis of compositions as the matrix of variances of all possible log-ratios among components (Aitchison, 1986). The generalization to rcomp objects is simply to reproduce the variance of all possible differences between components. The amount (`aplus`, `rplus`) and rmult objects should not be treated with variation matrices, because this was intended to skip the existence of a closure (which does not exist in the case of amounts).

Value

The variation matrix of `x`.

Author(s)

variograms

See Also
cdt, clrvar2ilr, clo, mean.acomp, acomp, rcomp, aplus, rplus

Examples
data(SimulatedAmounts)
meanCol(sa.lognormals)
variation(acomp(sa.lognormals))
variation(rcomp(sa.lognormals))
variation(aplus(sa.lognormals))
variation(rplus(sa.lognormals))
variation(rmult(sa.lognormals))

Description
Valid scalar variogram model functions.

Usage
vgram.sph( h , nugget = 0, sill = 1, range= 1,... )
vgram.exp( h , nugget = 0, sill = 1, range= 1,... )
vgram.gauss( h , nugget = 0, sill = 1, range= 1,... )
vgram.cardsin( h , nugget = 0, sill = 1, range= 1,... )
vgram.lin( h , nugget = 0, sill = 1, range= 1,... )
vgram.pow( h , nugget = 0, sill = 1, range= 1,... )
vgram.nugget( h , nugget = 1,...,tol=1E-8 )

Arguments
h a vector providing distances, a matrix of distance vectors in its rows or a data.frame of distance vectors.
nugget The size of the nugget effect (i.e. the limit to 0). At zero itself the value is always 0.
sill The sill (i.e. the limit to infinity)
range The range parameter. I.e. the distance in which sill is reached or if this does not exist, where the value is in some sense nearly the sill.
... not used
tol The distance that is considered as nonzero.
Details

The univariate variograms are used in the CompLinCoReg as building blocks of multivariate variogram models.

- sphSpherical variogram
- expExponential variogram
- gaussThe Gaussian variogram.
- gaussThe cardinal sine variogram.
- linLinear Variogram. Increases over the sill, which is reached at range.
- powThe power variogram. Increases over the sill, which is reached at range.
- nuggetThe pure nugget effect variogram.

Value

A vector of size NROW(h), giving the variogram values.

Author(s)


References

Tolosana, van den Boogaart, Pawlowsky-Glahn (2009) Estimating and modeling variograms of compositional data with occasional missing variables in R, StatGis09

See Also

vgram2lrvgram, CompLinModCoReg, vgmFit

Examples

```r
## Not run:
data(juraset)
X <- with(juraset,cbind(X,Y))
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
lrv <- logratioVariogram(comp,X,maxdist=1,nbins=10)
plot(lrv)
## End(Not run)
```
var.lm  
Residual variance of a model  

Description

Computes the unbiased estimate for the variance of the residuals of a model.

Usage

## S3 method for class 'lm'
var(x, ...)

Usage

## S3 method for class 'lm'
var(x, ...)

Arguments

x  
a linear model object

...  
Unused, for generic purposes only.

Details

The difference of this command to var(resid(X)) is that this command correctly adjusts for the degrees of freedom of the model.

Value

var.lm  returns a scalar giving the estimated variance of the residuals

var.mlm  returns a the estimated variance covariance matrix of the residuals

Author(s)


See Also

vcov

Examples

data(Orange)
var(lm(circumference~age, data=Orange))
var(lm(cbind(circumference, age)~age, data=Orange))
vcovAcomp

Variance covariance matrix of parameters in compositional regression

Description

The variance covariance tensor structured according of linear models with ilr(acomp(...)) responses.

Usage

vcovAcomp(object,...)

Arguments

object a statistical model
... further optional parameters for vcov

Details

The prediction error in compositional linear regression models is a complicated object. The function should help to organize it.

Value

An array with 4 dimensions. The first 2 are the index dimensions of the ilr transform. The later 2 are the index of the parameter.

Author(s)


See Also

vcov

Examples

data(SimulatedAmounts)
model <- lm(ilr(sa.groups)~sa.groups.area)
vcovAcomp(model)[,,1,1]
Compositional variogram model fitting

Description

Fits a parametric variogram model to an empirical logratio-Variogram

Usage

```r
vgmFit2lrv(emp, vg, ..., mode = "log", psgn = rep(-1, length(param)), print.level = 1)
vgmFit(emp, vg, ..., mode = "log", psgn = rep(-1, length(param)), print.level = 1)
vgmGof(p = vgmGetParameters(vg), emp, vg, mode = "log")
vgmGetParameters(vg, envir = environment(vg))
vgmSetParameters(vg, p)
```

Arguments

- `emp`: An empirical logratio-Variogram as e.g. returned by `logratioVariogram`
- `vg`: A compositional clr-variogram (or ilt-vagriogram) model function.
- `...`: further parameters to `nlm`
- `mode`: either "ls" or "log" for selection of either using either least squares or least squares on logarithmic values.
- `psgn`: Contains a parameter code for each of the parameters. -1 means the parameter should be used as is. 0 means the parameter is nonnegative and 1 means the parameter is strictly positive. This allows to provide parameter limits if the fitting procedure fails.
- `print.level`: The print.level of `nlm`. 0 for no printing. 1 for a rough information about the success and 2 for step by step printing.
- `p`: Is the parameter of the variogram model in linearized form as e.g. returned by `vgmGetParameters`.
- `envir`: The environment the default parameters of the model should be evaluated in.

Details

The function is mainly a wrapper to `nlm` specifying the an objective function for modelling fitting, taking the starting values of fitting procedure from the default arguments and writing the results back. Variogram model fitting is more an art than a straight forward procedure. Fitting procedures typically only find a right optimum if reasonable starting parameters are provided. The fit should be visually checked afterwards.

The meaning of `psgn` is subject to change. We will probably provide a more automatic procedure later.

`vgmFit` is a copy of `vgmFit2lrv`, but deprecated. The name will later be used for other functionality.
WhiteCells

Value

vgmFit2lr returns a list of two elements.

nlm
The result of nlm containing convergence codes.

vg
A version of vg but with default parameters modified according to the fitting.

gmGof returns a scalar quantifying the goodness of fit of a model and an empirical variogram.
gmGetParameters extracts the default values of a variogram model function to a parameter vector.
It returns a numeric vector.
gmSetParameters does the inverse operation and modifies the default according to the new values in p. It returns vg with modified default parameter values.

Author(s)

See Also

vgm2lrgram, CompLinModCoReg, logratioVariogram

Examples

## Not run:
data(juraset)
X <- with(juraset,cbind(X,Y))
comp <- acomp(juraset,c("Cd","Cu","Pb","Co","Cr"))
lrv <- logratioVariogram(comp,X,maxdist=1,nbins=10)
fff <- CompLinModCoReg(~nugget()+sph(0.5)+R1*exp(0.7),comp)
fit <- vgmFit(lrv,fff)
fit
fff(1:3)
plot(lrv,lrvg=vgm2lrgram(fit$vg))

## End(Not run)

WhiteCells

White-cell composition of 30 blood samples by two different methods

Description

In 30 blood samples portions of three kinds of white cells

- G: granulocytes,
- L: lymphocytes,
- M: monocytes,

were determined with two methods, time-consuming microscopic and automatic image analysis.
The resulting 30 pairs of 3-part compositions are recorded.
Usage

data(WhiteCells)

Format

A 30x6 matrix

Details

In an experiment each of 30 blood samples was halved, one half being assigned randomly to one method, the other half to the other method. We have 60 cases of 3-part compositions but these are essentially 30 pairs of related compositions. All 3-part portions sums to one, except for some rounding errors.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name WCELLS.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


Yatquat

Yatquat fruit evaluation

Description

The quality of yatquat tree fruit is assessed in terms of relative proportions by volume of flesh, skin and stone. In an experiment an arboriculturist uses 40 trees, randomly allocated 20 to the hormone treatment and leaves untreated the remaining 20 trees. Data provides fruit compositions of the present season and the preceding season, as well as the treatment: 1 for the treated trees, -1 for untreated trees.

Usage

data(Yatquat)

Format

A 40x7 data matrix
Details

The yatquat tree produces each season a single large fruit. Data provides fruit compositions of the present season, the compositions of the fruit of the same 40 trees for the preceding season when none of the trees were treated, and in addition the Type: 1 for the treated trees, -1 for untreated trees. For each of the 40 cases we have two 3-part composition on flesh, skin and stone. The column names are:

- prFL portion of fruit flesh in the present season,
- prSK portion of fruit skin in the present season,
- prST portion of fruit stone in the present season,
- Type 1 for treated, $-1$ for untreated trees,
- paFL portion of fruit flesh in the preceding season,
- paSK portion of fruit skin in the preceding season,
- paST portion of fruit stone in the preceding season,

All 3-part compositions sum to one.

Note

Courtesy of J. Aitchison

Source

Aitchison: CODA microcomputer statistical package, 1986, the file name YATQUAT.DAT, here included under the GNU Public Library Licence Version 2 or newer.

References


Examples

```r
#data(Yatquat)
#plot(acomp(Yatquat[,1:3]),col=as.numeric(Yatquat[,4])+2)
#plot(acomp(Yatquat[,5:7]),col=as.numeric(Yatquat[,4])+2)
```

zeroreplace  Zero-replacement routine

Description

A function to automatically replace rounded zeroes/BDLs in a composition.

Usage

```r
zeroreplace(x,d=NULL,a=2/3)
```
Arguments

- **x**: composition or dataset of compositions
- **d**: vector containing the detection limits of each part
- **a**: fraction of the detection limit to be used in replacement

Details

If `d` is given, zeroes from each column of `x` are replaced by the corresponding detection limit contained there, scaled down by the value of `a` (usually a scalar, although if it is a vector it will be recycled with a warning). The variable `d` should be a vector of length equal to `ncol(x)` or a matrix of the same shape as `x`. If `d=NULL`, then the detection limit is extracted from the data set, if it is available there (i.e., if there are negative numbers). If no negative number is present in the data set, and no value is given for `d`, the result will be equal to `x`. See `compositions.missings` for more details on the missing policy.

Value

an object of the same class as `x`, where all `WZERO` values have been replaced. Output contains a further attribute (named `Losts`), with a logical array of the same dimensions as `x`, showing which elements were replaced (TRUE) and which were kept unchanged (FALSE).

References


http://ima.udg.es/Activitats/CoDaWork03

http://ima.udg.es/Activitats/CoDaWork05

See Also

`compositions.missings`, `getDetectionlimit`

Examples

```r
data(SimulatedAmounts)
x <- acomp(sa.lognormals)
xnew <- simulateMissings(x,dl=0.05,knownlimit=FALSE)
xnew
xrep <- zeroreplace(xnew,0.05)
xrep
```
Index

*Topic IO
  mix.Read, 127
  Read standard data files, 196
*Topic NA
  getdetectionlimit, 84
  missingProjector, 124
  missingsummary, 126
*Topic aplot
  replot, 198
  ternaryAxis, 242
*Topic classes
  acomp, 13
  ccomp, 45
  is.acomp, 103
  print.acomp, 179
  transformations from 'mixtures' to 'compositions' classes, 245
*Topic cluster
  ClusterFinder1, 57
*Topic datagen
  simulatemissings, 230
*Topic datasets
  Aar, 12
  Activity10, 19
  Activity31, 20
  AnimalVegetation, 24
  ArcticLake, 29
  Bayesite, 37
  Blood23, 41
  Boxite, 42
  ClamEast, 51
  ClamWest, 52
  Coxite, 71
  DiagnosticProb, 73
  Firework, 79
  Glacial, 85
  Kongite, 90
  HouseholdExp, 92
  Hydrochem, 92
  jura, 107
  Kongite, 110
  Metabolites, 120
  PogoJump, 165
  Sediments, 218
  SerumProtein, 220
  ShiftOperators, 221
  SimulatedAmounts, 223
  Skulls, 232
  SkyeAFM, 232
  Supervisor, 241
  WhiteCells, 257
  Yatquat, 258
*Topic debugging
  tryDebugger, 246
*Topic distribution
  rAitchison, 185
  rMahalanobis, 201
  rnorm, 207
  rpois, 213
  runif, 214
*Topic dynamic
  kingTetrahedron, 108
*Topic hplot
  CoDaDendrogram, 59
  pairwiseplot, 146
  plot.acomp, 151
  pwlrPlot, 181
  simplemissingplot, 222
*Topic htest
  ccompgof, 46
  fitdirichlet, 80
  fitSameMeanDifferentVarianceModel, 81
  gausstest, 82
  gof, 86
  NormalTests, 135
*Topic logic
  binary, 38
**Topic multivariate**

acomparith, 15
acompmargin, 17
acompscalarproduct, 18
alr, 22
aplus, 24
aplusarithm, 26
apt, 28
arrows3D, 30
as.data.frame, 31
axis3D, 32
balance, 33
barplot.acomp, 35
biplot3D, 40
boxplot, 43
cdt, 49
clo, 53
clr, 54
clr2ilr, 56
coloredBiplot, 62
colorsForOutliers, 64
CompLinModCoReg, 65
compOKriging, 67
ConfRadius, 68
cor.acomp, 69
cpt, 72
dist, 74
dratplot, 75
endmemberCoordinates, 77
groupparts, 89
HotellingsTsq, 91
idt, 94
iit, 96
ilr, 97
ilrBase, 99
ilt, 100
ipt, 101
IsMahalanobisOutlier, 104
isPortionLines, 105
lines, 111
logratioVariogram, 112
lrvgram, 114
MahalanobisDist, 115
matmult, 117
mean.acomp, 118
missing.compositions, 121
mvar, 129
names, 131

norm, 132
normalize, 134
oneOrDataset, 136
outlierclassifier, 137
outlierplot, 139
parametricMat, 148
perturbe, 149
plot.aplus, 154
plot3D, 156
plot3Dacomp, 157
plot3Daplus, 159
plot3Drmult, 160
plot3Drplus, 161
plotlogratioVariogram, 163
plotmissingsummary, 164
powerofpsdmatrix, 166
princomp.acomp, 167
princomp.aplus, 170
princomp.rcomp, 173
princomp.rmult, 175
princomp.rplus, 177
qqnorm, 183
R2, 184
ratioLoadings, 188
rcomp, 190
rcomparithm, 193
rcompmargin, 194
rDirichlet, 195
rlnorm, 200
rmult, 203
rmultarithm, 204
rmultmatmult, 205
rplus, 211
rplusarithm, 212
scalar, 215
scale, 216
segments, 219
split, 233
straight, 234
summary.acomp, 236
summary.aplus, 237
summary.rcomp, 238
sumprojector, 239
totals, 244
ult, 247
var.acomp, 248
variation, 251
variograms, 252
INDEX

varmlm, 254
vcovAcomp, 255
vgmFit, 256
zeroreplace, 259
*Topic package
 compositions-package, 5
*Topic robust
 outliersInCompositions, 143
 robustnessInCompositions, 209
*Topic univar
  geometricmean, 83
  meanrow, 119
  *acomp (acomparith), 15
  *aplus, 150
  *aplus (aplusarithm), 26
  *rcomp (rcomparith), 193
  *rmult (rmultarithm), 204
  *rplus (rplusarithm), 212
  +.acomp, 193, 213
  +.acomp (perturbe), 149
  +.aplus (aplusarithm), 26
  +.rcomp, 192
  +.rcomp (rcomparith), 193
  +.rmult, 26, 193, 204, 213
  +.rmult (rmultarithm), 204
  +.rplus, 150
  +.rplus (rplusarithm), 212
  -.acomp (perturbe), 149
  -.aplus (aplusarithm), 26
  -.rcomp (rcomparith), 193
  -.rmult (rmultarithm), 204
  -.rplus (rplusarithm), 212
  /.*acomp (acomparith), 15
  /.*aplus (aplusarithm), 26
  /.*rcomp (rcomparith), 193
  /.*rmult (rmultarithm), 204
  /.*rplus (rplusarithm), 212
  **% (matmult), 117
  **%.acomp (acompscalarproduct), 18
  **%.aplus (acompscalarproduct), 18
  **%.rmult (rmultmatmult), 205
  **%, 117
  **%.rmult, 19, 27, 117, 204–206

Aar, 12
acomparith, 10, 13, 16–18, 25, 26, 36, 50, 53, 54,
61, 95, 103, 118, 119, 123, 124, 130,
150, 168, 169, 180–182, 191, 192,
BDL (missing.compositions), 121
BDLvalue (missing.compositions), 121
binary, 38
binary2logical (binary), 38
biplot, 40, 63
biplot3D, 40
bit (binary), 38
bit<- (binary), 38
bitCount (binary), 38
Blood23, 41
Boxite, 42
boxplot, 43, 182
boxplot.acomp, 15, 36, 153, 156, 180, 184, 199, 241, 244
boxplot.aplus, 26
boxplot.rcomp, 192
boxplot.rplus, 212
bxp, 44
ccomp, 45
compgof, 46
ccompMultinomialGOF.test, 46
ccompMultinomialGOF.test (ccompgof), 46
ccompPoissonGOF.test, 46
ccompPoissonGOF.test (ccompgof), 46
cdt, 49, 74, 95, 118, 130, 216, 250, 252
cdt.ccomp, 46
cdtInv, 95
cdtInv (cdt), 49
cdtInv .ccomp, 46
cgram2vgram (lrvgram), 114
ClamEast, 51
ClamWest, 52
clo, 14, 53, 119, 191, 230, 239, 250, 252
clr, 14–16, 23, 25, 34, 50, 54, 54, 73, 98, 100, 158, 169, 180, 192, 241
clr2irlr, 56
clrInv (clr), 54
clrvar2irlr, 250, 252
clrvar2irlr (clr2irlr), 56
ClusterFinder1, 57, 105, 138, 142, 144, 210
CoDaDendrogram, 59
coloredBiplot, 62, 141
colorsForOutliers, 64
colorsForOutliers (colorsForOutliers), 64
colorsForOutliers (colorsForOutliers), 64
CompLinModCoReg, 65, 66, 68, 113, 115, 149, 163, 253, 257
compOKriging, 67
composition.missing
(missing.compositions), 121
composition.missings
(missing.compositions), 121
compositions (compositions-package), 5
compositions-package, 5, 12, 124, 144, 210
compositions.missing, 14, 26, 46, 54, 119, 127, 192, 212
compositions.missings
(missing.compositions), 121
compositions.missings, 85, 165, 180, 231, 260
compositions.missings
(missing.compositions), 121
ConfRadius, 68, 91
cov.rcomp, 192
cov.rcomp (rcomparithm), 193
cor, 70
cor (cor.acomp), 69
cor.acomp, 69
cov, 130, 249, 250
cov (var.acomp), 248
cov.acomp, 15, 241
cov.aplus, 26
cov.rcomp, 192
cov.rplus, 212
covMcd, 118, 209
Coxite, 71
cpt, 28, 29, 50, 72, 102, 103, 175, 192, 193, 213
cptInv (cpt), 72
dAitchison (rAitchison), 185
Data01 (Hongite), 90
Data02 (Kongite), 110
Data03 (Boxite), 42
Data04 (Coxite), 71
Data05 (ArcticLake), 29
Data06 (SkyeAFM), 232
Data07 (Supervisor), 241
DATA08 (HouseholdExp), 92
Data09 (Metabolites), 120
DATA10 (Activity10), 19
Data11 (WhiteCells), 257
Data12 (Yatquat), 258
Data13 (Firework), 79
<table>
<thead>
<tr>
<th>Function</th>
<th>Page(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>is.MNAR (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>is.MNV (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>is.NMV (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>is.rcomp (is.acomp)</td>
<td>103</td>
</tr>
<tr>
<td>is.rmult (is.acomp)</td>
<td>103</td>
</tr>
<tr>
<td>is.rplus (is.acomp)</td>
<td>103</td>
</tr>
<tr>
<td>is.SZ (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>is.WMNAR (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>IsMahalanobisOutlier</td>
<td>104</td>
</tr>
<tr>
<td>isoPortionLines</td>
<td>105</td>
</tr>
<tr>
<td>isoProportionLines</td>
<td>105</td>
</tr>
<tr>
<td>jura</td>
<td>107</td>
</tr>
<tr>
<td>jura259 (jura)</td>
<td>107</td>
</tr>
<tr>
<td>juraset (jura)</td>
<td>107</td>
</tr>
<tr>
<td>Kappa (ult)</td>
<td>247</td>
</tr>
<tr>
<td>kingTetrahedron</td>
<td>108, 153, 158, 159, 244</td>
</tr>
<tr>
<td>kmeans</td>
<td>59</td>
</tr>
<tr>
<td>Kongite</td>
<td>110</td>
</tr>
<tr>
<td>lines</td>
<td>111</td>
</tr>
<tr>
<td>lines.acomp</td>
<td>220, 235</td>
</tr>
<tr>
<td>lm</td>
<td>69, 185</td>
</tr>
<tr>
<td>lmrob</td>
<td>147, 181</td>
</tr>
<tr>
<td>logratioVariogram</td>
<td>112, 115, 163, 256, 257</td>
</tr>
<tr>
<td>lrvgram</td>
<td>114</td>
</tr>
<tr>
<td>mahalanobis</td>
<td>116, 203</td>
</tr>
<tr>
<td>MahalanobisDist</td>
<td>115, 202</td>
</tr>
<tr>
<td>MAR (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>MARvalue (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>matmult</td>
<td>117</td>
</tr>
<tr>
<td>maxBit (binary)</td>
<td>38</td>
</tr>
<tr>
<td>mcov (mvar)</td>
<td>129</td>
</tr>
<tr>
<td>mean</td>
<td>118, 120</td>
</tr>
<tr>
<td>mean.acomp</td>
<td>15, 70, 118, 124, 130, 169, 180, 210, 236, 241, 250, 252</td>
</tr>
<tr>
<td>mean.aplus</td>
<td>26, 172</td>
</tr>
<tr>
<td>mean.aplus (mean.acomp)</td>
<td>118</td>
</tr>
<tr>
<td>mean.ccomp</td>
<td>46</td>
</tr>
<tr>
<td>mean.ccomp (mean.acomp)</td>
<td>118</td>
</tr>
<tr>
<td>mean.col (meanrow)</td>
<td>119</td>
</tr>
<tr>
<td>mean.rcomp</td>
<td>192</td>
</tr>
<tr>
<td>mean.rcomp (mean.acomp)</td>
<td>118</td>
</tr>
<tr>
<td>mean.rmult (mean.acomp)</td>
<td>118</td>
</tr>
<tr>
<td>mean.row (meanrow)</td>
<td>119</td>
</tr>
<tr>
<td>mean.rplus</td>
<td>84, 120, 212</td>
</tr>
<tr>
<td>mean.rplus (mean.acomp)</td>
<td>118</td>
</tr>
<tr>
<td>meanCol</td>
<td>118, 119</td>
</tr>
<tr>
<td>meanCol (meanrow)</td>
<td>119</td>
</tr>
<tr>
<td>meanRow (meanrow)</td>
<td>119</td>
</tr>
<tr>
<td>meanrow</td>
<td>119</td>
</tr>
<tr>
<td>Metabolites</td>
<td>120</td>
</tr>
<tr>
<td>missing.compositions</td>
<td>121</td>
</tr>
<tr>
<td>missingProjector</td>
<td>124, 240, 241</td>
</tr>
<tr>
<td>missings</td>
<td>210</td>
</tr>
<tr>
<td>missings (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>missingsInCompositions</td>
<td>12, 124, 125, 144</td>
</tr>
<tr>
<td>mix.2acomp (transformations from 'mixtures' to 'compositions' classes)</td>
<td>245</td>
</tr>
<tr>
<td>mix.2aplus (transformations from 'mixtures' to 'compositions' classes)</td>
<td>245</td>
</tr>
<tr>
<td>mix.2rcomp (transformations from 'mixtures' to 'compositions' classes)</td>
<td>245</td>
</tr>
<tr>
<td>mix.2rmult (transformations from 'mixtures' to 'compositions' classes)</td>
<td>245</td>
</tr>
<tr>
<td>mix.2rplus (transformations from 'mixtures' to 'compositions' classes)</td>
<td>245</td>
</tr>
<tr>
<td>mix.Read</td>
<td>127</td>
</tr>
<tr>
<td>MNAR (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>MNARvalue (missing.compositions)</td>
<td>121</td>
</tr>
<tr>
<td>msd</td>
<td>15, 26, 192, 212, 241</td>
</tr>
<tr>
<td>msd (mvar)</td>
<td>129</td>
</tr>
<tr>
<td>mul.rplus (rplusarithm)</td>
<td>212</td>
</tr>
<tr>
<td>mvar</td>
<td>69, 129, 185</td>
</tr>
<tr>
<td>na.fail</td>
<td>120</td>
</tr>
<tr>
<td>na.omit</td>
<td>120</td>
</tr>
<tr>
<td>na.pass</td>
<td>120</td>
</tr>
<tr>
<td>names</td>
<td>131</td>
</tr>
<tr>
<td>names.ccomp</td>
<td>46</td>
</tr>
<tr>
<td>names&lt;-.acomp (names)</td>
<td>131</td>
</tr>
</tbody>
</table>
INDEX

names<-.aplus(names), 131
names<-.ccomp(names), 131
names<-.rcomp(names), 131
names<-.rmult(names), 131
names<-.rplus(names), 131
nlm, 256, 257
NMV (missing.compositions), 121
noreplot (replot), 198
norm, 132
norm.rmult, 134, 204
normalize, 133, 134
NormalTests, 135

observeWithAdditiveError
(simulatemissings), 230
observeWithDetectionLimit
(simulatemissings), 230
observeWithDetectionLimit
(simulatemissings), 230
oneOrDataset, 136
outlierclassifier, 137
OutlierClassifier1, 64, 65, 104, 105, 116,
142–144, 203, 210
OutlierClassifier1 (outlierclassifier), 137
outlierplot, 39, 105, 138, 139, 143, 144, 210
outliersInCompositions, 12, 58, 104, 124,
138, 140, 143, 144, 210

pairs, 153
pairwisePlot, 182
pairwisePlot (pairwiseplot), 146
pairwiseplot, 146
par, 59, 60, 147, 151, 155, 243
parameterPosdefClrMat (parametricMat),
148
parameterPosdefMat (parametricMat), 148
parameterRank1ClrMat (parametricMat),
148
parameterRank1Mat (parametricMat), 148
parametricMat, 148
parametricPosdefClrMat (parametricMat),
148
parametricPosdefMat (parametricMat), 148
parametricRank1ClrMat (parametricMat),
148
parametricRank1Mat (parametricMat), 148
pchForOutliers1 (colorsForOutliers), 64
pEmpiricalMahalanobis (rMahalanobis),
201
perturbe, 14, 149
perturbe.aplus, 25
perturbe.aplus (aplusarithm), 26
pf, 91
pHotellingsTsq (HotellingsTsq), 91
plot, 31, 33, 157, 159–162
plot.acomp, 15, 36, 45, 63, 76, 107, 110, 112,
124, 151, 180, 184, 199, 220, 235, 241
plot.aplus, 26, 147, 153, 154, 156, 182, 199,
223, 244
plot.ccomp, 46
plot.ccomp (plot.acomp), 151
plot.default, 182
plot.logratioVariogram
(plotlogratioVariogram), 163
plot.missingSummary
(plotmissingsummary), 163
plot.princomp.acomp (princomp.acomp),
167
plot.princomp.aplus (princomp.aplus),
170
plot.princomp.rcomp (princomp.rcomp),
173
plot.princomp.rplus (princomp.rplus),
177
plot.rcomp, 192
plot.rcomp (plot.acomp), 151
plot.relativeLoadings.princomp.acomp
(ratioLoadings), 188
plot.relativeLoadings.princomp.aplus
(ratioLoadings), 188
plot.relativeLoadings.princomp.rcomp
(ratioLoadings), 188
plot.relativeLoadings.princomp.rplus
(ratioLoadings), 188
plot.rmult (plot.aplus), 154
plot.rplus, 212
plot.rplus (plot.aplus), 154
plot3D, 31, 33, 153, 156, 159, 160, 162, 244
plot3D.acomp, 157, 159–162
plot3D.acomp (plot3Dacomp), 157
plot3D.aplus, 157, 159–162
plot3D.aplus (plot3Daplus), 159
plot3D.rcomp, 157, 159–162
plot3D.rcomp (plot3Dacomp), 157
plot3D.rmult, 157, 159–162
plot3D.rmult(plot3Drmult), 160
plot3D.rplus, 157, 159–161
plot3D.rplus(plot3Drplus), 161
plot3Dacomp, 157
plot3Daplus, 159
plot3Drmult, 160
plot3Drplus, 161
plotlogratioVariogram, 163
plotmissingsummary, 164
pMaxMahalanobis(rMahalanobis), 201
PogoJump, 165
points3d, 31, 33, 157, 159–162
PoissonGOF.test, 46
PoissonGOF.test(ccompgof), 46
power.acomp, 14
power.acomp(acomparith), 15
power.aplus, 25
power.aplus(aplusarithm), 26
powerofpsdmatrix, 166
pPortionMahalanobis(rMahalanobis), 201
pQuantileMahalanobis, 144
pQuantileMahalanobis(rMahalanobis), 201
predict.princomp.acomp
(princomp.acomp), 167
predict.princomp.aplus
(princomp.aplus), 170
predict.princomp.rcomp
(princomp.rcomp), 173
predict.princomp.rplus
(princomp.rplus), 177
princomp.acomp, 15, 167, 171, 172, 175, 189, 190, 241
princomp.aplus, 26, 169, 170, 179, 189, 190
princomp.default, 175
princomp.rcomp, 169, 173, 179, 189, 190, 192
princomp.rmult, 175
princomp.rplus, 172, 175, 176, 177, 189, 190, 212
print.acomp, 179
print.aplus(print.acomp), 179
print.princomp.acomp(princomp.acomp), 167
print.princomp.aplus(princomp.aplus), 170
print.princomp.rcomp(princomp.rcomp), 173
print.princomp.rplus(princomp.rplus), 177
print.rcomp(print.acomp), 179
print.relativeLoadings.princomp.acomp
(ratioLoadings), 188
print.relativeLoadings.princomp.aplus
(ratioLoadings), 188
print.relativeLoadings.princomp.rcomp
(ratioLoadings), 188
print.relativeLoadings.princomp.rplus
(ratioLoadings), 188
print.rmult(rmult), 203
print.rplus(print.acomp), 179
pwlrPlot, 147, 181
pwlrplot(pwlrPlot), 181
qEmpiricalMahalanobis, 143
qEmpiricalMahalanobis(rMahalanobis), 201
qHotellingsTsq(HotellingsTsq), 91
qMaxMahalanobis, 143
qMaxMahalanobis(rMahalanobis), 201
qPortionMahalanobis(rMahalanobis), 201
qnorm, 183
qnorm.acomp, 45, 153, 156, 244
R2, 184
rAitchison, 185
ratioLoadings, 188
rcomparith, 193
rcompmargin, 18, 151, 153, 194
rDirichlet, 48, 81, 88, 136, 195
rDirichlet.acomp, 188, 208, 215
Read standard data files, 196
read.geoEAS, 129
read.geoEAS(Read standard data files), 196
read.geoeas, 129
read.geoeas(Read standard data files), 196
read.table, 129, 197
relativeLoadings, 168, 169, 171, 172, 174, 175, 178, 179
relativeLoadings(ratioLoadings), 188
rEmpiricalMahalanobis(rMahalanobis), 201
replot, 198
replotable (replot), 198
rlnormal, 30, 32, 40, 156, 158–160, 162
rlnorm, 200
rlnorm.rplus, 208
rlnorm.rplus (rlnorm), 200
rMahalanobis, 201
rMaxMahalanobis (rMahalanobis), 201
rmt, 27, 78, 124, 178, 193, 203, 205, 206, 213, 246
rmultarithm, 204
rmultinom.ccomp, 46
rmultinom.ccomp (rpois), 213
rmultmatmult, 205
rmult, 207
rnorm, 207
rnorm.acomp, 48, 81, 88, 136, 184, 188, 196, 200
rnorm.acomp (rnorm), 207
rnorm.aplus, 184
rnorm.aplus (rnorm), 207
rnorm.ccomp, 46, 214
rnorm.ccomp (rnorm), 207
rnorm.rmult, 184
rnorm.rmult (rnorm), 207
rnorm.rplus, 184
rnorm.rplus (rnorm), 207
robust (robustnessInCompositions), 209
robustnessInCompositions, 12, 104, 115, 124, 140, 143, 144, 152, 156, 167, 202, 209, 217, 236, 238, 239, 250, 251
rplus, 10, 25, 26, 36, 50, 95–97, 103, 118, 119, 123, 130, 178, 179, 192, 204, 211, 216, 238, 246, 250–252
rplusarithm, 212
rpois, 213
rpois.ccomp, 46, 208
rpois.ccomp (rpois), 213
rPortionMahalanobis (rMahalanobis), 201
Rsquare (R2), 184
runif, 214
runif.acomp, 48, 81, 88, 136, 188, 208
sa.dirichlet (SimulatedAmounts), 223
sa.dirichlet5 (SimulatedAmounts), 223
sa.groups (SimulatedAmounts), 223
sa.groups5 (SimulatedAmounts), 223
sa.lognormals (SimulatedAmounts), 223
sa.lognormals5 (SimulatedAmounts), 223
sa.missings (SimulatedAmounts), 223
sa.missings5 (SimulatedAmounts), 223
sa.outliers1 (SimulatedAmounts), 223
sa.outliers2 (SimulatedAmounts), 223
sa.outliers3 (SimulatedAmounts), 223
sa.outliers4 (SimulatedAmounts), 223
sa.outliers5 (SimulatedAmounts), 223
sa.outliers6 (SimulatedAmounts), 223
sa.tnormals (SimulatedAmounts), 223
sa.tnormals5 (SimulatedAmounts), 223
sa.uniform (SimulatedAmounts), 223
sa.uniform5 (SimulatedAmounts), 223
scalar, 204, 215
scale, 152, 155, 216, 217
scale.default, 217
Sediments, 218
segments, 219
SerumProtein, 220
shapiro.test, 183
ShiftOperators, 221
simplemissingplot, 222
simpleMissingSubplot (simplemissingplot), 222
SimulatedAmounts, 223
simulatemissings (simulatemissings), 230
simulatemissings, 230
Skulls, 232
Skye, 233
SkyeAFM, 232
solve, 115
spineplot, 182
split, 217, 233, 234
split.ccomp, 46
straight, 112, 234
stripchart, 141
summary, 238, 239
summary.acomp, 236, 238, 239
summary.aplus, 237, 239
summary.ccomp, 238, 238
summary.rmult (summary.aplus), 237
summary.rplus (summary.aplus), 237
sumMissingProjector, 125
sumMissingProjector (sumprojector), 239
sumprojector, 239
Supervisor, 241
SZ (missing.compositions), 121
SZvalue (missing.compositions), 121
t.test, 83
ternaryAxis, 242
text, 222
totals, 244
totals.comp, 46
transformations from 'mixtures' to 'compositions' classes, 245
tryDebugger, 246
uciptInv (ipt), 101
ult, 247
ultInv (ult), 247
unbinary (binary), 38
update, 198

var, 130, 249, 250
var (var.acomp), 248
var.acomp, 15, 70, 137, 169, 180, 210, 241, 248
var.aplus, 26
var.lm (var.lm), 254
var.lm (var.lm), 254
var.rcomp, 192
var.rmult, 171, 173, 176, 177
var.rplus, 212
variation, 250, 251
variation.acomp, 15, 180, 236, 241
variation.aplus, 26
variation.rcomp, 192
variation.rplus, 212
variograms, 252
var.lm, 254
vcov, 254, 255
vcovAcomp, 255

vgmFit, 66, 68, 113, 115, 149, 253, 256
vgmFit2lrv (vgmFit), 256
vgmGetParameters (vgmFit), 256
vgmGof (vgmFit), 256
vgmSetParameters (vgmFit), 256
vgram.cardsin (variograms), 252
vgram.exp (variograms), 252
vgram.gauss (variograms), 252
vgram.lin (variograms), 252
vgram.nugget (variograms), 252
vgram.pow (variograms), 252
vgram.sph (variograms), 252
vgram2lrvgram, 66, 68, 113, 149, 163, 253, 257
vgram2lrvgram (lrvgram), 114

vp.boxplot (boxplot), 43
vp.logboxplot (boxplot), 43
vp.qnorm (qnorm), 183

whichBits (binary), 38
WhiteCells, 257

xsimplex, 187
Yatquat, 258
zeroreplace, 85, 123, 124, 180, 259