# Package 'ArArRedux'

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**Title** Rigorous Data Reduction and Error Propagation of Ar40 / Ar39 Data

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Description Processes noble gas mass spectrometer data to determine the isotopic composition of argon (comprised of Ar36, Ar37, Ar38, Ar39 and Ar40) released from neutron-irradiated potassium-bearing minerals. Then uses these compositions to calculate precise and accurate geochronological ages for multiple samples as well as the covariances between them. Error propagation is done in matrix form, which jointly treats all samples and all isotopes simultaneously at every step of the data reduction process. Includes methods for regression of the timeresolved mass spectrometer signals to t=0 ('time zero') for both single- and multi-collector instruments, blank correction, mass fractionation correction, detector intercalibration, decay corrections, interference corrections, interpolation of the irradiation parameter between neutron fluence monitors, and (weighted mean) age calculation. All operations are performed on the logs of the ratios between the different argon isotopes so as to properly treat them as 'compositional data', sensu Aitchison [1986, The Statistics of Compositional Data, Chapman and Hall].

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Imports methods

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2 average

X		27
	weightedmean	. 25
	timeresolved	
	summary.results	
	subset.timeresolved	
	results	
	redux	
	read	. 21
	process	. 21
	plotcorr	. 20
	plot.timeresolved	. 19
	PHdata	. 19
	param	. 18
	newredux	. 17
	Melbourne	. 17
	massfractionation	. 16
	logratios	. 16
	loadirradiations	. 15
	loaddata	. 14
	isoratios	. 14
	interference	. 13
	getmasses	. 11
	getJfactors	
	getages	
	get4039	
	fractionation	9
	fitlogratios	
	decaycorrection	
	concat	
	cleorrection	
	calibration	
	blankcorrected	
	averagebyday	
	averagebyday	. 1

average

Calculate the arithmetic mean

# Description

Calculate the arithmetic mean of some logratio data

```
average(x, i = NULL, newlabel = NULL)
```

averagebyday 3

### **Arguments**

an object of class redux or logratios
 (optional) vector of sample indices
 newlabel
 (optional) string with the new label to be assigned to the averaged values

### Value

an object of the same class as x

### **Examples**

```
data(Melbourne)
K <- average(Melbourne$X,grep("K:",Melbourne$X$labels),newlabel="K-glass")
plotcorr(K)</pre>
```

averagebyday

Average all the data collected on the same day.

### **Description**

This function is useful for grouping a number of replicate air shots or calibration experiments

### Usage

```
averagebyday(x, newlabel)
```

### **Arguments**

x an object of class timeresolved, logratios, PHdata or redux newlabel a string with the new label that should be given to the average

### Value

an object of the same class as x

```
dfile <- system.file("Calibration.csv",package="ArArRedux")
md <- loaddata(dfile)
ld <- fitlogratios(blankcorr(md))
d <- averagebyday(ld,"DCAL")
plotcorr(d)</pre>
```

4 blankcorr

blankcorr

Apply a blank correction

### **Description**

Applies a blank correction to some time-resolved mass spectrometer data

### Usage

```
blankcorr(x, ...)
## Default S3 method:
blankcorr(x, ...)
## S3 method for class 'timeresolved'
blankcorr(x, blanklabel = NULL, prefix = "", ...)
## S3 method for class 'PHdata'
blankcorr(x, blanklabel = NULL, prefix = "", ...)
## S3 method for class 'WiscAr'
blankcorr(x, blanklabel = NULL, prefix = "", ...)
```

### **Arguments**

```
    an object of class timeresolved or PHdata
    other arguments
    as string denoting the prefix of the blanks
    prefix
    a string to be prepended to the non-blanks
```

## Value

an object of class blankcorrected

```
samplefile <- system.file("Samples.csv",package="ArArRedux")
m <- loaddata(samplefile) # samples and J-standards
blanklabel <- "EXB#"
1 <- fitlogratios(blankcorr(m,blanklabel),"Ar40")
plotcorr(1)</pre>
```

blankcorrected 5

orrected <i>class</i>
-----------------------

### **Description**

An object class containing blank-corrected mass spectrometry data

### **Details**

Extends the class classes timeresolved and PHdata by adding an additional list item blankindices containg the index of the nearest blank. fitlogratios uses this information to group the samples during regression to 'time zero'.

calibration

Detector calibration

### **Description**

Apply the detector calibration for multicollector data

## Usage

```
calibration(x, ...)
## Default S3 method:
calibration(x, clabel, ...)
## S3 method for class 'WiscAr'
calibration(x, clabel, ...)
```

### **Arguments**

x an object of class redux optional arguments

clabel the label of the detector calibration data

### Value

an object of class redux

```
data(Melbourne)
C <- calibration(Melbourne$X,"DCAL")
plotcorr(C)</pre>
```

6 concat

clcorrection

Cl-interference correction

# Description

Apply the interference correction for the Cl-decay products

### Usage

```
clcorrection(X, irr)
```

# Arguments

X an object of class redux irr the irradiation schedule

#### Value

an object of class redux

# **Examples**

```
data(Melbourne)
Cl <- clcorrection(Melbourne$X, Melbourne$irr)
plotcorr(Cl)</pre>
```

concat

Merge a list of logratio data

# Description

Recursively concatenates a list of logratio data into one big dataset

### Usage

```
concat(lrlist)
## Default S3 method:
concat(lrlist)
## S3 method for class 'WiscAr'
concat(lrlist)
```

# Arguments

1rlist a list containing items of class logratios or redux

decaycorrection 7

#### Value

an object of the same class as x containing the merged dataset

### **Examples**

```
samplefile <- system.file("Samples.csv",package="ArArRedux")</pre>
kfile <- system.file("K-glass.csv",package="ArArRedux")</pre>
cafile <- system.file("Ca-salt.csv",package="ArArRedux")</pre>
dfile <- system.file("Calibration.csv",package="ArArRedux")</pre>
blanklabel <- "EXB#"
Jpos <- c(3,15)
m <- loaddata(samplefile) # samples and J-standards
mk <- loaddata(kfile) # K-interference data
mca <- loaddata(cafile) # Ca interference data</pre>
md <- loaddata(dfile) # detector intercalibrations</pre>
# form and fit logratios
1 <- fitlogratios(blankcorr(m,blanklabel),"Ar40")</pre>
lk <- fitlogratios(blankcorr(mk,blanklabel, "K:"), "Ar40")</pre>
k \leftarrow getmasses(lk, "Ar39", "Ar40") # subset on the relevant isotopes
lca <- fitlogratios(blankcorr(mca,blanklabel,"Ca:"),"Ar37")</pre>
ca <- getmasses(lca,c("Ar36","Ar39"),c("Ar37","Ar37")) # subset</pre>
ld <- fitlogratios(blankcorr(md))</pre>
d <- averagebyday(ld, "DCAL")</pre>
# merge all data (except air shots) into one big logratio structure
X <- newredux(concat(list(l,k,ca,d)),Jpos)</pre>
data(Melbourne)
if (isTRUE(all.equal(Melbourne$X,X))) {
   print("We just reconstructed the built-in dataset Melbourne$X")}
```

decaycorrection

Correct for radioactive decay occurred since irradiation

# Description

Correct for radioactive decay of neutron-induced 37Ar and 39Ar occurred since irradiation

# Usage

```
decaycorrection(x, irr, isotope)
```

### Arguments

x an objects of class redux or WiscAr

irr the irradiation schedule

isotope a string denoting the isotope that needs correcting

8 fitlogratios

### Value

```
an object of class redux
```

# Examples

```
data(Melbourne)
C <- calibration(Melbourne$X,"DCAL")
A <- massfractionation(C,Melbourne$fract)
D9 <- decaycorrection(A,Melbourne$irr,"Ar39")
plotcorr(D9)</pre>
```

fitlogratios

Extrapolation to 'time zero'

### Description

This function extrapolates time resolved mass spectrometer data to t=0. When fed with multicollector data, it forms the ratios of the raw signals, forms their logs and performs linear regression to t=0 When fed with single collector data, the function first takes their logs and extrapolates them to t=0 before taking ratios, unless denmass=NULL, in which case the logs of the raw signals are extrapolated.

### Usage

```
fitlogratios(x, ...)
## Default S3 method:
fitlogratios(x, ...)
## S3 method for class 'timeresolved'
fitlogratios(x, denmass = "Ar40", tmin = 0, ...)
## S3 method for class 'PHdata'
fitlogratios(x, denmass = NULL, tmin = 0, ...)
## S3 method for class 'WiscAr'
fitlogratios(x, denmass = "Ar39", ...)
```

### **Arguments**

```
    an object of class timeresolved or PHdata
    further arguments (see below)
    a string denoting the denominator isotope
    time stamp (in seconds) of 'time zero'
```

fractionation 9

### Value

```
an object of class logratios
```

# **Examples**

```
samplefile <- system.file("Samples.csv",package="ArArRedux")
m <- loaddata(samplefile) # samples and J-standards
blanklabel <- "EXB#"
l <- fitlogratios(blankcorr(m,blanklabel),"Ar40")
plotcorr(l)</pre>
```

fractionation

Compute the mass fractionation correction

### **Description**

Compares the measured 40Ar/36Ar ratio of an air shot on a given detector with the atmospheric ratio.

# Usage

```
fractionation(fname, detector, MS = "ARGUS-VI")
```

### **Arguments**

fname a .csv file with the air shot data detector the name of the ion detector MS the type of mass spectrometer

### Value

```
an object of class logratios
```

10 getages

get4039

Calculate the 40Ar\*/39ArK-ratios

### **Description**

Calculate the 40Ar\*/39ArK-ratios of interference corrected logratio intercept data

# Usage

```
get4039(X, irr)
```

### **Arguments**

X an object of class redux containing some interference corrected logratio inter-

cept data

irr the irradiation schedule

### Value

an object of class link{redux} containing the 40Ar\*/39ArK-ratios as intercepts and its covariance matrix as covmat

### **Examples**

```
data(Melbourne)
R <- get4039(Melbourne$X,Melbourne$irr)
plotcorr(R)</pre>
```

getages

Calculate 40Ar/39Ar ages

# Description

Calculate 40Ar/39Ar ages from a vector of 40Ar/39Ar-ratios and J-factors

### Usage

```
getages(RJ)
```

# Arguments

RJ

an object of class Redux containing the amalgamated \$^40\$Ar\$^\*\$/\$^39\$Ar\$\_K\$-ratios and J-factors with their covariance matrix

### Value

an object of class results containing the ages and their covariance matrix

getJfactors 11

### **Examples**

```
data(Melbourne)
R <- get4039(Melbourne$X,Melbourne$irr)
J <- getJfactors(R)
ages <- getages(J)
plotcorr(ages)</pre>
```

getJfactors

Calculate the irradiation parameter ('J factor')

### **Description**

Interpolate the irradiation parameters for the samples given the 40Ar\*/39ArK ratios of the samples and fluence monitors

# Usage

```
getJfactors(R)
```

### **Arguments**

R

a vector of 40Ar\*/39ArK ratios

### Value

an object of class redux containing, as intercepts, the  $40 \text{Ar}^*/39 \text{Ar} \text{K}$  ratios of the samples, the interpolated J-factors, and the 40 K decay constant; and as covmat: the covariance matrix. All other class properties are inherited from R.

# **Examples**

```
data(Melbourne)
R <- get4039(Melbourne$X,Melbourne$irr)
J <- getJfactors(R)
plotcorr(J)</pre>
```

getmasses

Select a subset of isotopes from a dataset

## **Description**

Extracts the intercepts, covariance matrix, etc. of a selection of isotopes from a larger dataset

12 getmasses

### Usage

```
getmasses(x, ...)
## Default S3 method:
getmasses(x, ...)
## S3 method for class 'timeresolved'
getmasses(x, mass, invert = FALSE, ...)
## S3 method for class 'WiscAr'
getmasses(x, mass, invert = FALSE, ...)
## S3 method for class 'logratios'
getmasses(x, num, den, invert = FALSE, ...)
## S3 method for class 'redux'
getmasses(x, num, den, invert = FALSE, ...)
```

### **Arguments**

X	an object of class logratios, timeresolved, PHdata or redux.
	other arguments
mass	a vector of strings denoting the masses of interest
invert	boolean parameter indicating whether the selection should be inverted (default = FALSE)
num	vector of strings indicating the numerator isotopes
den	vector of string indicating the denominator isotopes

### Value

an object of the same class as x

```
kfile <- system.file("K-glass.csv",package="ArArRedux")
mk <- loaddata(kfile)
lk <- fitlogratios(blankcorr(mk,"EXB#","K:"),"Ar40")
k <- getmasses(lk,"Ar39","Ar40") # subset of the relevant isotopes
plotcorr(k)</pre>
```

interference 13

interference	define the interference corrections	
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### **Description**

create a new object of class logratios containing the interferences from neutron reactions on Ca and K

### Usage

```
interference(intercepts, covmat, num, den, irr, label)
```

### **Arguments**

intercepts	a vector with logratios
covmat	the covariance matrix of the logratios
num	a vector of strings marking the numerator isotopes of intercepts
den	a vector of strings marking the denominator isotopes of intercepts
irr	an object of class irradiations
label	a string with a name which can be used to identify the interference data in subsequent calculations

### Value

an object of class logratios

```
samplefile <- system.file("Samples.csv",package="ArArRedux")</pre>
irrfile <- system.file("irradiations.csv",package="ArArRedux")</pre>
X <- read(samplefile,blabel="EXB#",Jpos=c(3,15))</pre>
irr <- loadirradiations(irrfile)</pre>
# assume log(36Ar/37Ar) = log(39Ar/37Ar) = 1 in co-irradiate Ca-salt
# with variances of 0.0001 and zero covariances
ca <- interference(intercepts=c(1,1),</pre>
                    covmat=matrix(c(0.001,0,0,0.001),nrow=2),
                    num=c("Ar39","Ar36"),den=c("Ar37","Ar37"),
                    irr=X$irr[1],label="Ca-salt")
# assume log(39Ar/40Ar) = 4.637788 in co-irradiate K-glass
# with variance 7.9817e-4
k <- interference(intercepts=4.637788,covmat=7.9817e-4,</pre>
                   num="Ar39",den="Ar40",irr=X$irr[1],
                   label="K-glass")
ages <- process(X,irr,ca=ca,k=k)</pre>
summary(ages)
```

14 loaddata

# Description

Calculate the Calculate the inverse isochron ratios of interference corrected logratio intercept data

# Usage

```
isoratios(X, irr, fract = NULL, ca = NULL, k = NULL, inverse = TRUE)
```

# Arguments

X	an object of class redux containing some interference corrected logratio intercept data
irr	the irradiation schedule
fract	list with air shot data (one item per denominator isotope)
ca	an object of class $logratios$ with Ca-interference data (not necessary if interferences are included in $\boldsymbol{X}$ )
k	an object of class $\log$ ratios with K-interference data (not necessary if interferences are included in $X$ )
inverse	logical. If TRUE, returns inverse isochron ratios ( $^{36}$ Ar/ $^{40}$ Ar vs. $^{39}$ Ar/ $^{40}$ Ar). Otherwise returns normal isochron ratios ( $^{40}$ Ar/ $^{36}$ Ar vs. $^{39}$ Ar/ $^{36}$ Ar).

### Value

an object of class link{isoratios} containing the isochron ratios and their covariance matrix.

# Examples

```
data(Melbourne)
IR <- isoratios(Melbourne$X,irr=Melbourne$irr,fract=Melbourne$fract)</pre>
```

loaddata	Load mass spectrometer data

# Description

Loads a file or directory with raw mass spectrometer data

```
loaddata(fid, MS = "ARGUS-VI")
```

loadirradiations 15

# **Arguments**

fid the file or directory name containing the data

MS one of either ARGUS-VI or WiscAr

### Value

an object of class timeresolved or PHdata and ArgusVI or WiscAr

# **Examples**

```
samplefile <- system.file("Samples.csv",package="ArArRedux")
masses <- c("Ar37","Ar38","Ar39","Ar40","Ar36")
m <- loaddata(samplefile,MS='ARGUS-VI') # samples and J-standards
plot(m,"MD2-1a","Ar40")</pre>
```

loadirradiations

Load the irradiation schedule

# **Description**

Loads a .csv file with the schedule of a multi-stage neutron irradiation

# Usage

```
loadirradiations(fname)
```

### **Arguments**

fname file name (in .csv format)

### Value

a list of irradiations, where each irradiation is a named list containing:

tin: vector with the start times of irradiations tout: vector with the end times of irradiations P: vector with the power of the irradiations

```
irrfile <- system.file("irradiations.csv",package="ArArRedux")
irr <- loadirradiations(irrfile)
str(irr)</pre>
```

16 massfractionation

logratios

The logratios class

### **Description**

An object class containing logratio intercepts

#### **Details**

A list with the following items:

labels: a vector of strings denoting the names of the runs num: a vector of strings denoting the numerator isotopes den: a vector of strings denoting the denominator isotopes intercepts: a vector of logratio intercepts or values covmat: the covariance matrix of intercepts

irr: a vector of strings denoting the irradiation runs

pos: a vector of integers with the positions in the irradiation stack thedate: a vector containing the acquisition dates and times

nlr: a vector with the number of logratios per run

massfractionation

Apply the mass fractionation correction

# Description

Applies the fractionation obtained from air shot data by fractionation to the denominator detector in order to correct it for the mass difference between the numerator and denominator isotopes.

# Usage

```
massfractionation(X, fract)
```

### Arguments

X an object of class redux

fract a list with fractionation data for Ar37, Ar39 and Ar40

### Value

an object of class redux

Melbourne 17

### **Examples**

```
graphics.off()
data(Melbourne)
C <- calibration(Melbourne$X,"DCAL")
A <- massfractionation(C,Melbourne$fract)
plotcorr(A)</pre>
```

Melbourne

An example dataset

# Description

Contains all the relevant information needed for the data reduction some ARGUS-IV data from the University of Melbourne

### Author(s)

David Philips <dphillip@unimelb.edu.au>

### **Examples**

```
data(Melbourne)
plotcorr(Melbourne$X)
```

newredux

Create a new redux object

### **Description**

Initialises a new redux object by packing a logratios dataset together with all the parameters needed for age calculation

```
newredux(x, ...)
## Default S3 method:
newredux(
    x,
    Jpos,
    detectors = list(Ar36 = "H1", Ar37 = "L2", Ar38 = "L1", Ar39 = "AX", Ar40 = "H1"),
    ...
)
## S3 method for class 'WiscAr'
newredux(x, Jpos, ...)
```

18 param

#### **Arguments**

x an object of class logratios

... optional arguments

Jpos a vector of integers denoting the positions of the fluence monitors in the irradi-

ation stack

detectors a list of strings denoting the detectors for each argon isotope

#### Value

an object of class redux

param

Set or get Ar-Ar\_Redux parameters

## **Description**

This function is used to query and modify the half lives, standard ages etc. associated with an object of class redux

### Usage

```
param(X, ...)
```

### **Arguments**

X an object of class redux

any combination of the parameters given below

#### **Details**

param grants access to the following parameters:

10: 40K decay constant (default value = 5.5492e-4 Ma-1, Renne et al. [2010])

s10: standard error of the 40K decay constant (default value = 0.0047e-4 Ma-1)

17: 37Ar decay constant (default value = 7.2438 yr-1, Renne and Norman [2001])

s17: standard error of the 37Ar decay constant (default value = 0.0083 yr-1)

19: 39Ar decay constant (0.002577 yr-1 Stoenner et al. [1965])

s19: standard error of the 39Ar decay constant (0.000014 yr-1)

16: 36Cl decay constant (default value = 2301.3e-9 yr-1)

s16: standard error of the 36Cl decay constant (default value = 7.6e-9 yr-1

pc1: (36Cl/38Cl)-production rate (default value = 252.7 for OSTR reactor, Renne et al. [2008])

spc1: standard error of the (36C1/38C1)-production rate (default value = 1.8)

ts: age of the fluence monitor (default = 28.201 Myr for the Fish Canyon Tuff, Kuiper et al. [2008])

sts: standard error of the fluence monitor age (default value = 0.023 Myr)

air: atmospheric 40Ar/36Ar ratio (default value = 298.56, Lee et al. [2006])

sair: standard error of the atmospheric 40Ar/36Ar ratio (default value = 0.155)

PHdata 19

### Value

returns the modified redux object OR the current parameter values if no optional arguments are supplied.

### **Examples**

```
data(Melbourne)
param(Melbourne$X)$air
Y <- param(Melbourne$X,air=295.5)
param(Y)$air</pre>
```

PHdata

The PHdata class

### **Description**

An object class containing time resolved 'peak-hopping' mass spectrometry data

#### **Details**

A list with the following items:

masses: a vector of strings denoting the isotopes monitored in each run signals: a list with objects of class timeresolved, each corresponding to a detector (i.e. length(signals)==1 for single collector instruments).

### See Also

loaddata

plot.timeresolved

Plot a time resolved mass spectrometry signal

### **Description**

Plots the raw signal of a given isotope against time.

```
## S3 method for class 'timeresolved'
plot(x, label = NULL, mass = NA, run = 1, hop = "101", tmin = 0, ...)
## S3 method for class 'WiscAr'
plot(x, hop = "101", ...)
## S3 method for class 'PHdata'
plot(x, ...)
```

20 plotcorr

### **Arguments**

x an object of class timeresolved or PHdata

label a string with the name of the run

mass a string indicating the isotope of interest

run the run number

hop identifier for mass channel in Noblesse instruments (if x has class WiscAr).

tmin time stamp (in seconds) of 'time zero'

... optional parameters

# **Examples**

```
samplefile <- system.file("Samples.csv",package="ArArRedux")
mMC <- loaddata(samplefile)
plot(mMC,"MD2-1a")
mPH <- loaddata(samplefile)
plot(mPH,"MD2-1a","Ar40")</pre>
```

plotcorr

Plot a matrix with correlation coefficients

# Description

Converts the covariance matrix to a correlation matrix and plots this is a coloured image for visual inspection.

### Usage

```
plotcorr(X)
```

# Arguments

X a data structure (list) containing an item called 'covmat' (covariance matrix)

```
graphics.off()
data(Melbourne)
plotcorr(Melbourne$X)
```

process 21

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Process logratio data and calculate 40Ar/39Ar ages

### **Description**

Performs detector calibration, mass fractionation correction, decay corrections, interference corrections, interpolates J-factors and calculates ages.

### Usage

```
process(X, irr, fract = NULL, ca = NULL, k = NULL)
```

### **Arguments**

Χ	an object of class redux
irr	the irradiation schedule
fract	list with air shot data (one item per denominator isotope)
ca	an object of class $logratios$ with Ca-interference data (not necessary if interferences are included in $\boldsymbol{X}$ )
k	an object of class $logratios$ with K-interference data (not necessary if interferences are included in $X$ )

# **Examples**

```
data(Melbourne)
ages <- process(Melbourne$X,Melbourne$irr,Melbourne$fract)
summary(ages)</pre>
```

read

Read mass spectrometer data

### **Description**

Reads raw mass spectrometer data and parses it into a redux format for further processing.

```
read(x, blabel, Jpos, kdat = NULL, cadat = NULL, ddat = NULL, MS = "ARGUS-VI")
```

22 redux

# **Arguments**

X	a .csv file with samples and fluence monitor data (if MS = 'ARGUS-VI') OR a directory with .RUN files (if MS = 'WiscAr')
blabel	a prefix string denoting the blanks
Jpos	a vector of integers denoting the positions of the fluence monitors in the irradiation stack
kdat	a .csv file with the K-interference monitor data (if MS = 'ARGUS-VI') OR a directory with .RUN files of K-interference monitor data (if MS = 'WiscAr') (optional)
cadat	a .csv file with the Ca-interference monitor data (if MS = 'ARGUS-VI') OR a directory with .RUN files of Ca-interference monitor data (if MS = 'WiscAr') (optional)
ddat	a .csv file with the detector calibration data (if MS = 'ARGUS-VI') OR the prefix of the standard gas measurements (if MS = 'WiscAr') (optional)
MS	a string denoting the type of mass spectrometer. At the moment there are two options: 'ARGUS-VI', for data that are acquired on the eponymous instrument following the measurement protocols at Melbourne University, and 'WiscAr' for the 5-detector Noblesse instrument at the University of Wisconsin. Please contact the author to add other file formats to Ar-Ar_Redux.

### Value

an object of class redux.

# **Examples**

redux The redux class

# Description

An object class that is used throughout Ar-Ar\_Redux

results 23

### **Details**

A list with the following items:

labels: a vector of strings denoting the names of the runs num: a vector of strings denoting the numerator isotopes den: a vector of strings denoting the denominator isotopes intercepts: a vector of logratio intercepts or values covmat: the covariance matrix of intercepts

irr: a vector of strings denoting the irradiation runs

pos: a vector of integers with the positions in the irradiation stack thedate: a vector containing the acquisition dates and times

nlr: a vector with the number of logratios per run

param: a list of global parameters

### See Also

param

results

The results class

### **Description**

A list with the following items:

### **Details**

labels: a vector of strings denoting the names of the runs

intercepts: a vector of ages

covmat: the covariance matrix of intercepts

thedate: a vector containing the acquisition dates and times

subset.timeresolved

Select a subset of some data

### **Description**

Extracts those intercepts, covariances etc. that match a given list of indices or labels.

24 summary.results

### Usage

```
## S3 method for class 'timeresolved'
subset(x, i = NULL, labels = NULL, invert = FALSE, include.J = FALSE, ...)
## S3 method for class 'logratios'
subset(x, i = NULL, labels = NULL, invert = FALSE, include.J = FALSE, ...)
## S3 method for class 'redux'
subset(x, i = NULL, labels = NULL, invert = FALSE, include.J = FALSE, ...)
## S3 method for class 'results'
subset(x, i = NULL, labels = NULL, invert = FALSE, ...)
```

### **Arguments**

an object of class timeresolved, logratios, redux or results

a vector with indices of the selected runs

labels a string or a vector of strings with sample names

boolean flag indicating whether the selection should be inverted, i.e. whether the selected indices or labels should be removed rather than retained include. J if TRUE, automatically adds the irradiation monitors to the selection other arguments

### Value

an object of the same class as x

## **Examples**

```
data(Melbourne)
ages <- process(Melbourne$X,Melbourne$irr,Melbourne$fract)
MD <- subset(ages,labels=c("MD2-1","MD2-2","MD2-3","MD2-4","MD2-5"))
plotcorr(MD)</pre>
```

summary.results

Summary table

### **Description**

Plots the ages and their standard errors

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

timeresolved 25

### **Arguments**

```
object an object of class results
... no other arguments
```

### **Examples**

```
data(Melbourne)
ages <- process(Melbourne$X,Melbourne$irr,Melbourne$fract)
summary(ages)[1:5,]</pre>
```

timeresolved

*The* timeresolved *class* 

### Description

An object class containing time resolved multi-collector mass spectrometry data

### **Details**

A list with the following items:

masses: a vector of strings denoting the isotopes monitored in each run

irr: a vector of strings denoting the irradiation runs

pos: a vector of integers with the positions in the irradiation stack thedate: a vector containing the acquisition dates and times

d: a data table

thetime: a matrix with the measurement times

### See Also

loaddata

weightedmean

Calculate the weighted mean age

## **Description**

Computes the error weighted mean and MSWD of some samples taking into covariances.

```
weightedmean(ages, prefix = NULL)
```

26 weightedmean

# Arguments

ages an object of class results

prefix is either a string with the prefix of the samples that need to be averaged, or a

vector of sample names.

### Value

a list with items:

avgt: the weighted mean age err: the standard error of avgt

MSWD: the Mean Square of the Weighted Deviates

```
data(Melbourne)
ages <- process(Melbourne$X,Melbourne$irr,Melbourne$fract)
weightedmean(ages,"MD2-")</pre>
```

# **Index**

```
average, 2
averagebyday, 3
blankcorr, 4
blankcorrected, 4, 5
calibration, 5
clcorrection, 6
concat, 6
decaycorrection, 7
fitlogratios, 5, 8
fractionation, 9, 16
get4039, 10
getages, 10
getJfactors, 11
getmasses, 11
interference, 13
isoratios, 14
loaddata, 14, 19, 25
loadirradiations, 15
logratios, 6, 9, 12, 14, 16, 17, 18, 21, 24
massfractionation, 16
Melbourne, 17
newredux, 17
param, 18, 18
PHdata, 4, 5, 12, 19, 20
plot.PHdata(plot.timeresolved), 19
plot.timeresolved, 19
plot.WiscAr (plot.timeresolved), 19
plotcorr, 20
process, 21
read, 21
redux, 6, 12, 17-19, 21, 22, 22, 24
```

```
results, 23, 24, 25

subset.logratios (subset.timeresolved), 23

subset.redux (subset.timeresolved), 23

subset.timeresolved, 23

summary.results, 24

timeresolved, 4, 5, 12, 19, 20, 24, 25

weightedmean, 25
```