# Package ‘corr2D’

October 12, 2022

Type Package

Title Implementation of 2D Correlation Analysis in R

Version 1.0.3

Date 2022-07-14

Description Implementation of two-dimensional (2D) correlation analysis based on the Fourier-transformation approach described by Isao Noda (I. Noda (1993) <DOI:10.1366/0003702934067694>). Additionally there are two plot functions for the resulting correlation matrix: The first one creates colored 2D plots, while the second one generates 3D plots.

Imports doParallel (>= 1.0.8), foreach (>= 1.4.3), parallel (>= 3.0.2), fields (>= 8.2-1), mmand (>= 1.3.0), stats (>= 3.0.2), grDevices (>= 3.0.2), graphics (>= 3.0.2), utils (>= 3.0.2), colorspace (>= 1.3-0)

License GPL-3

RoxygenNote 7.2.0

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Suggests knitr, rmarkdown, R.rsp, testthat, rgl (>= 0.93.996-1), profr, xtable

VignetteBuilder R.rsp

NeedsCompilation no

Author Robert Geitner [cre, aut], Robby Fritzsch [aut], Thomas Bocklitz [aut], Juergen Popp [ctb, cph]

Maintainer Robert Geitner <robert.geitner@tu-ilmenau.de>

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**R topics documented:**

  - codis2d
**codis2d**

Two-dimensional codistribution spectroscopy.

**Description**

codis2d calculates the synchronous and asynchronous codistribution spectra.

**Usage**

codis2d(
  Mat,
  Ref = NULL,
  Wave = NULL,
  Time = NULL,
  Int = stats::splinefun,
  N = 2^ceiling(log2(NROW(Mat))),
  Norm = 1/(NROW(Mat) - 1),
  scaling = 0,
  corenumber = parallel::detectCores(),
  preview = FALSE
)

**Arguments**

- **Mat** (Numeric matrix containing the data which will be correlated; 'spectral variable' by columns and 'perturbation variables' by rows.)
- **Ref** (Numeric vector containing a single spectrum, which will be subtracted from Mat to generate dynamic spectra for 2D correlation analysis. Default is NULL in which case the colMeans() of Mat is used as reference. The length of Ref needs to be equal to the number of columns in Mat. 2D codistribution spectroscopy is only strictly defined using the perturbation-mean spectrum as reference spectrum. Thus, any deviation from this definition can lead to unexpected results.)
- **Wave** (Numeric vector containing the spectral variable. Needs to be specified if column names of Mat are undefined.)
- **Time** (Numeric vector containing the perturbation variables. If specified, Mat will be interpolated to N equally spaced perturbation variables using Int.)
Function specifying how the dataset will be interpolated to give \( N \) equally spaced perturbation variables. \texttt{splinefun} (default) or \texttt{approxfun} can for example be used.

\( N \): Positive, non-zero integer specifying how many equally spaced perturbation variables should be interpolated using \texttt{Int}. \( N \) should be higher than 4. \texttt{corr2d} is fastest if \( N \) is a power of 2.

\( \text{Norm} \): A number specifying how the correlation matrix should be normalized.

\( \text{scaling} \): Positive real number used as exponent when scaling the dataset with its standard deviation. Defaults to 0 meaning no scaling. 0.5 (Pareto scaling) and 1 (Pearson scaling) are commonly used to enhance weak correlations relative to strong correlations. 2D codistribution spectroscopy is only strictly defined without the usage of any scaling techniques. Thus, any deviation from this definition can lead to unexpected results.

\( \text{corenumber} \): Positive, non-zero integer specifying how many CPU cores should be used for parallel fft computation.

\( \text{preview} \): Logical: Should a 3D preview of the asynchronous codistribution spectrum be drawn at the end? Uses \texttt{persp3d} from \texttt{rgl} package.

### Details

codis2d calculates the the synchronous 2D correlation spectrum and uses the 2D spectrum to calculate the synchronous and asynchronous codistribution spectra. For parallelization the \texttt{parCapply} function is used. Large input matrices (> 4000 columns) can lead to long calculation times depending on the number of cores used. Also note that the resulting matrix can become very large, adjust the RAM limit with \texttt{memory.limit} accordingly. For a detailed description of the underlying math see references.

### Value

codis2D returns a list of class "corr2d" containing the complex codistribution matrix ($\text{FT}$), the synchronous correlation spectrum ($\text{corr}$), the used reference spectrum $\text{Ref1}$ and $\text{Ref2}$, the spectral variables $\text{Wave1}$ and $\text{Wave2}$ as well as the (interpolated) perturbation variables ($\text{Time}$).

### References

I. Noda (2014) <DOI:10.1016/j.molstruc.2014.01.024>

### See Also

For plotting of the resulting list containing the 2D codistribution spectra see \texttt{plot_corr2d} and \texttt{plot_corr2din3d}.

### Examples

```r
testdata <- sim2ddata(C = NULL, Camp = NULL)
codis <- codis2d(testdata, corenumber = 1)
plot_corr2d(codis, Im(codis$FT),
```

```r
codis2d
```

```r
codis2d
```
corr2d

Two-dimensional correlation analysis.

description

corr2d calculates the synchronous and asynchronous correlation spectra between Mat1 and Mat1 (homo correlation) or between Mat1 and Mat2 (hetero correlation).

Usage

corr2d(
  Mat1,
  Mat2 = NULL,
  Ref1 = NULL,
  Ref2 = NULL,
  Wave1 = NULL,
  Wave2 = NULL,
  Time = NULL,
  Int = stats::splinefun,
  N = 2^ceiling(log2(NROW(Mat1)) ),
  Norm = 1/(pi * (NROW(Mat1) - 1)),
  scaling = 0,
  corenumber = parallel::detectCores(),
  preview = FALSE
)

Arguments

Mat1, Mat2
  Numeric matrix containing the data which will be correlated; 'spectral variable' by columns and 'perturbation variables' by rows. For hetero correlations Mat1 and Mat2 must have the same number of rows.

Ref1, Ref2
  Numeric vector containing a single spectrum, which will be subtracted from Mat1 (or Mat2, respectively) to generate dynamic spectra for 2D correlation analysis. Default is NULL in which case the colMeans() of Mat1 (or Mat2, respectively) is used as reference. The length of Ref1 (or Ref2) needs to be equal to the number of columns in Mat1 (or Mat2).

Wave1, Wave2
  Numeric vector containing the spectral variable. Needs to be specified if column names of Mat1 (or Mat2) are undefined.

Time
  Numeric vector containing the perturbation variables. If specified, Mat1 (and Mat2 if given) will be interpolated to N equally spaced perturbation variables using Int to speed up the fft algorithm.
 corr2d

<table>
<thead>
<tr>
<th>Int</th>
<th>Function specifying how the dataset will be interpolated to give ( N ) equally spaced perturbation variables. <code>splinefun</code> (default) or <code>approxfun</code> can for example be used.</th>
</tr>
</thead>
<tbody>
<tr>
<td>( N )</td>
<td>Positive, non-zero integer specifying how many equally spaced perturbation variables should be interpolated using <code>Int</code>. ( N ) should be higher than 4. <code>corr2d</code> is fastest if ( N ) is a power of 2.</td>
</tr>
<tr>
<td>Norm</td>
<td>A number specifying how the correlation matrix should be normalized.</td>
</tr>
<tr>
<td>scaling</td>
<td>Positive real number used as exponent when scaling the dataset with its standard deviation. Defaults to 0 meaning no scaling. 0.5 (Pareto scaling) and 1 (Pearson scaling) are commonly used to enhance weak correlations relative to strong correlations.</td>
</tr>
<tr>
<td>corenumber</td>
<td>Positive, non-zero integer specifying how many CPU cores should be used for parallel ( \text{fft} ) computation.</td>
</tr>
<tr>
<td>preview</td>
<td>Logical: Should a 3D preview of the synchronous correlation spectrum be drawn at the end? Uses <code>persp3d</code> from <code>rgl</code> package.</td>
</tr>
</tbody>
</table>

**Details**

corr2d uses a parallel fast Fourier transformation (`fft`) to calculate the complex correlation matrix. For parallelization the `foreach` function is used. Large input matrices (> 4000 columns) can lead to long calculation times depending on the number of cores used. Also note that the resulting matrix can become very large, adjust the RAM limit with `memory.limit` accordingly. For a detailed description of the underlying math see references.

**Value**

corr2D returns a list of class "corr2d" containing the complex correlation matrix (`$FT`), the used reference spectra (`$Ref1`, `$Ref2`), the spectral variables (`$Wave1`, `$Wave2`), the Fourier transformed data (`$ft1`, `$ft2`), the (interpolated) perturbation variables (`$Time`) and logical variable (`$Het`) indicating if homo (FALSE) or hetero (TRUE) correlation was done.

**References**

I. Noda (1993) <DOI:10.1366/0003702934067694>
R. Geitner et al. (2019) <DOI:10.18637/jss.v090.i03>

**See Also**

For plotting of the resulting list containing the 2D correlation spectra see `plot_corr2d` and `plot_corr2din3d`.

**Examples**

data(FuranMale, package = "corr2D")
twod <- corr2d(FuranMale, Ref1 = FuranMale[, 1], corenumber = 1)

plot_corr2d(twod, xlab = expression(paste("relative Wavenumber" / cm^-1)),
            ylab = expression(paste("relative Wavenumber" / cm^-1)))
corr2t2d  Two-trace two-dimensional (2T2D) correlation spectroscopy

## Description

corr2t2d compares a pair of spectra in the form of a cross correlation analysis.

## Usage

corr2t2d(Sam, Ref, Wave = NULL, preview = FALSE)

## Arguments

- **Sam**: Numeric vector containing the sample spectrum to be correlated. Can contain the spectral variable of the sample and reference spectrum as names.
- **Ref**: Numeric vector containing the sample spectrum to be correlated. Can contain the spectral variable of the sample and reference spectrum as names.
- **Wave**: Numeric vector containing the spectral variable. Needs to be specified if names of Sam and Ref are undefined.
- **preview**: Logical: Should a 3D preview of the asynchronous codistribution spectrum be drawn at the end? Uses persp3d from rgl package.

## Details

corr2t2d implements the Two-trace two-dimensional (2T2D) approach as described by I. Noda (2018) <DOI:10.1016/j.molstruc.2018.01.091>. The idea is to compare two spectra in a 2D correlation-like approach which was previously not possible as 2D correlation analysis usually needs at least three spectra.

## Value

corr2t2d returns a list of class "corr2d" containing the complex correlation matrix ($FT$), the correlation and disrelation coefficient as a complex matrix ($coef$), the sample $Ref1$ and reference spectrum $Ref2$ as well as the spectral variable $Wave1$ and $Wave2$.

## References


## See Also

For plotting of the resulting list containing the 2D correlation spectra or correlation coefficient see plot_corr2d and plot_corr2din3d.
Examples

testdata <- sim2ddata()
twodtest <- corr2t2d(testdata[4, ], testdata[5, ])
plot_corr2d(twodtest, Im(twodtest$FT))

FuranMale

FT-Raman spectra of furan maleimide based self-healing polymer

Description

Six preprocessed FT-Raman spectra of a self-healing polymer. The wavenumber region shows the C=C vibrations of furan, maleimide and their respective Diels-Alder adduct. The row names show the measurement temperature in degree Celsius, while the column names show the relative wavenumber.

Format

A matrix containing 6 spectra by rows with 145 wavenumbers by columns.

Source


is.corr2d

Check for object class "corr2d"

Description

The function checks if an object is of class "corr2d".

Usage

is.corr2d(x)

Arguments

x An object which should be check if it is of class "corr2d".

Details

The function uses the inherits function.
**plot_corr2d**

**Plot two-dimensional correlation spectra.**

**Description**

`plot_corr2d` plots two-dimensional correlation spectra either as an image or a contour plot. Red color indicates positive correlations, while blue color shows negative ones.

**Usage**

```r
plot_corr2d(
  Obj,
  what = Re(Obj$FT),
  specx = Obj$Ref1,
  specy = Obj$Ref2,
  xlim = NULL,
  ylim = NULL,
  xlab = expression(nu[1]),
  ylab = expression(nu[2]),
  Contour = TRUE,
  axes = 3,
  Legend = TRUE,
  N = 20,
  zlim = NULL,
  Cutout = NULL,
  col = par("col"),
  lwd = par("lwd"),
  lwd.axis = NULL,
  lwd.spec = NULL,
)
```

**Value**

A logical scalar

**References**

R. Geitner et al. (2019) <DOI:10.18637/jss.v090.i03>

**Examples**

```r
data(FuranMale, package = "corr2D")
twod <- corr2d(FuranMale, Ref1 = FuranMale[1, ], corenumber = 1)

# TRUE
is.corr2d(twod)
# FALSE
is.corr2d(2)
```
Arguments

Obj
List from `corr2d` containing the 2D correlation data.

what
Real numeric matrix containing the z-values that should be plotted.

specx, specy
Numeric vector containing the data that should be plotted on top (`specx`) and/or on the left (`specy`) of the 2D spectrum. Mat, `specx` and/or `specy` should have the same dimensions, respectively. If `NULL` nothing will be plotted.

xlim, ylim
Numeric vector with two values indicating the borders of the 2D plot. Also truncates `specx` and/or `specy` to match the new plot range.

xlab, ylab
Character or expression containing the text that will be plotted on the bottom (`xlab`) and/or to the right (`ylab`) of the 2D plot. Labels can be suppressed with `NA`.

Contour
Logical: Should a contour (`TRUE`) or image (`FALSE`) be drawn?

axes
Integer ranging from 0 to 3. Should the axis of the 2D plot be drawn? "0" means no axes, "1" only bottom axis, "2" only right axis and "3" both axes are drawn.

Legend
Logical: Should a color legend be plotted in the top right corner?

N
Positive, non-zero integer indicating how many contour or image levels should be plotted.

zlim
Numeric vector with two values defining the z-range of the 2D plot.

Cutout
Numeric vector with two values defining which z-values should not be plotted. Use with care, because this can generate misleading 2D plots.

col
A specification for the plotting color of the reference spectra (top and left), axes, axes ticks and the central plot surrounding box. See `par` and `contour` for additional information.

lwd
A numeric value which sets the line width in the contour plot. See `par` and `contour` for additional information.

lwd.axis
A numeric value which sets the line width for axes and the central plot surrounding box. See `par` and `axis` for additional information.

lwd.spec
A numeric value which sets the line width in the reference spectra on top and to the left. See `par` and `plot.default` for additional information.

cex.leg
A numerical value giving the amount by which numbers at the legend should be magnified. See `par` and `image.plot` for additional information.

at.xaxs, at.yaxs
The points at which tick-marks are to be drawn at the x- and y-axis, respectively. See `axis` for additional information.
label.xaxs, label.yaxs
This can either be a logical value specifying whether (numerical) annotations are to be made at the tickmarks of the x- and y-axis, or a character or expression vector of labels to be placed at the tickpoints of the x- and y-axis. See axis for additional information.

line.xlab, line.ylab
Numeric value on which MARgin line the x- and y-label is plotted, respectively, starting at 0 counting outwards. See mtext for additional information.

Details
For the synchronous correlation spectrum the real component (Re) of the complex correlation matrix must be plotted. The asynchronous spectrum is the respective imaginary component (Im). Cutout can be used to leave out smaller (noise) contributions, but should be used with care as it can be used to create misleading 2D correlation plots. See references for interpretation rules (so called Noda rules).

References
R. Geitner et al. (2019) <DOI:10.18637/jss.v090.i03>

See Also
See plot_corr2din3d for 3D plots.

Examples
```r
data(FuranMale, package = "corr2D")
twod <- corr2d(FuranMale, Ref1 = FuranMale[1, ], corenumber = 1)

plot_corr2d(twod, xlab = expression(paste("relative Wavenumber" / cm^-1)),
            ylab = expression(paste("relative Wavenumber" / cm^-1)))

plot_corr2d(twod, at.xaxs = c(1560, 1585, 1610),
            label.xaxs = c(1560, 1585, 1610),
            col = 2, lwd = 3, col.axis = 3, col.lab = 4, Legend = FALSE,
            cex.lab = 3, xlab = "Large x label", ylab = "Large y label",
            line.xlab = 5, line.ylab = 5)
```
plot_corr2din3d 3D plot of two-dimensional correlation spectra.

Description

plot_corr2din3d plots two-dimensional correlation spectra as an 3D surface.

Usage

plot_corr2din3d(
  Mat,
  specx = NULL,
  specy = NULL,
  scalex = NULL,
  scaley = NULL,
  Col = colorspace::diverge_hcl(64, h = c(240, 0), c = 100, l = c(20, 100), power = 0.4),
  reduce = NULL,
  zlim = NULL,
  projection = FALSE,
  ...
)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mat</td>
<td>Real numeric matrix containing the z-values to plot.</td>
</tr>
<tr>
<td>specx, specy</td>
<td>Numeric vector containing the data, that will be plotted at the x and y axis. Can be any data and does not need to have the same dimensions as Mat.</td>
</tr>
<tr>
<td>scalex, scaley</td>
<td>A real number which describes how specx (or specy) get scaled. Positive numbers lead to a spectrum plotted inside the box, while negative numbers lead to a spectrum plotted outside the box.</td>
</tr>
<tr>
<td>Col</td>
<td>Vector containing colors used to plot the 3D plot and the respective projection.</td>
</tr>
<tr>
<td>reduce</td>
<td>Non-zero rational number describing how to resample the data values. Can reduce the computational demand and can be used for fast previews.</td>
</tr>
<tr>
<td>zlim</td>
<td>Numeric vector with two values indicating the z-range of the 3D plot.</td>
</tr>
<tr>
<td>projection</td>
<td>Logical: Should a 2D projection of the 3D surface be plotted a the bottom of the box?</td>
</tr>
<tr>
<td>...</td>
<td>Additional arguments passed to drape.plot.</td>
</tr>
</tbody>
</table>

Details

For the synchronous correlation spectrum the real component (Re) of the complex correlation matrix must be plotted. The asynchronous spectrum is the respective imaginary component (Im).
sim2ddata

Simulate kinetic data from two-step sequential first-order reactions

Description

sim2ddata simulates kinetic data for the sequential reaction A -> B -> C with the time constants k1 and k2.

Usage

```r
sim2ddata(
  L = 400,
  t = 0:10,
  k1 = 0.2,
  k2 = 0.8,
  X = c(1000, 1400),
  A = c(1080, 1320),
  Aamp = c(3, 8),
  B = c(1120, 1280),
  Bamp = c(5, 15),
  C = c(1160, 1240),
  Camp = c(4, 9)
)
```
Arguments

L
Positive, non-zero integer specifying how many spectral variables should be used to describe the kinetic dataset.

t
Numeric vector containing non-negative real numbers describing at which reaction times the kinetic data should be sampled.

k1, k2
Positive, non-zero real numbers describing the time constants used to simulate the reactions A -> B (k1) and B -> C (k2).

X
Numeric vector with two values specifying the range of the simulated spectral variables.

A, B, C
Numeric vector with two real values specifying the two signal positions of species A, B and C, respectively. It’s the mean used in \texttt{dnorm} to simulate the signal. C and Camp may be NULL in which case only the reaction A -> B is simulated and sampled.

Aamp, Bamp, Camp
Numeric vector with two values specifying the signal width of species A, B and C, respectively. It’s the standard deviation (sd) used in \texttt{dnorm} to simulate the signal. C and Camp may be NULL in which case only the reaction A -> B is simulated and sampled.

Details

The simulation assumes 2 spectral signals for each of the 3 species A, B and C. The sequential reaction is defined by 2 time constants k1 and k2. The spectral information can be sampled at every point during the reaction to get an arbitrary profile of the kinetic data. The signals of the three species are modeled by a normal distribution. In addition the spectral variable is assumed to be equidistant and the number of spectral variables can also be chosen arbitrary.

Value

\texttt{sim2ddata} returns a matrix containing the kinetic data. The matrix contains the sampled reaction times by rows and the spectral variables by columns. The reaction times are the row names while the spectral variables are saved as the column names. The matrix has the ideal format to be analyzed by \texttt{corr2d}.

References

The default values are inspired by: I. Noda (2014) <DOI:10.1016/j.molstruc.2014.01.024>

Examples

 testData <- sim2ddata()

twodtest <- corr2d(testdata, corenumber = 1)

plot_corr2d(twodtest)
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