Package 'GPoM'

January 20, 2025

Type Package

Title Generalized Polynomial Modelling

Version 1.4

Date 2023-06-16

Maintainer Mireille Huc <mireille.huc@u-paris2.fr>

Description Platform dedicated to the Global Modelling technique. Its aim is to obtain ordinary differential equations of polynomial form directly from time series. It can be applied to single or multiple time series under various conditions of noise, time series lengths, sampling, etc. This platform is developped at the Centre d'Etudes Spatiales de la Biosphere (CESBIO), UMR 5126 UPS/CNRS/CNES/IRD, 18 av. Edouard Belin, 31401 TOULOUSE, FRANCE. The developments were funded by the French program Les Enveloppes Fluides et l'Environnement (LEFE, MANU, projets GloMo, SpatioGloMo and MoMu). The French program Defi InFiNiTi (CNRS) and PNTS are also acknowledged (projects Crops'IChaos and Musc & SlowFast). The method is described in the article : Mangiarotti S. and Huc M. (2019) <doi:10.1063/1.5081448>.

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LazyData TRUE

RoxygenNote 7.1.1

Depends R (>= 3.6), deSolve, rgl

Imports float

Suggests signal, knitr, rmarkdown

VignetteBuilder knitr

Encoding UTF-8

NeedsCompilation no

Author Sylvain Mangiarotti [aut], Mireille Huc [cre, aut], Flavie Le Jean [ctb], Malika Chassan [ctb], Laurent Drapeau [ctb], Institut de Recherche pour le Développement [fnd], Centre National de la Recherche Scientifique [fnd] Repository CRAN Date/Publication 2023-06-16 08:10:10 UTC

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GPoM-package

GPoM package: Generalized Polynomial Modelling

Description

GPoM is a platform dedicated to the Global Modelling technique. Its aim is to obtain deterministic models of Ordinary Differential Equations from observational time series. It applies to single and to multiple time series. With single time series, it can be used: to detect low-dimnesional determinism and low-dimensional (deterministic) chaos. It can also be used to characterize the observed behavior, using the obtained models as a proxy of the original dynamics, as far as the model validation could be checked. With multiple time series, it can be used: to detect couplings between observed variables, to infer causal networks, and to reformulate the original equations of the observed system (retro-modelling). The present package focuses on models in Ordinary Differential Equations of polynomial form. The package was designed to model weakly predictable dynamical behavior, either linear or nonlinear. Several vignettes are associated to the package which can be used as a tutorial, and it also provides an overlook of the diversity of applications and at the performances of the tools. Users are kindly asked to quote the corresponding references when using the package (see hereafter).

Note

FOR USERS

This package was developped at Centre d'Etudes Spatiales de la Biosphere (Cesbio, UMR 5126, UPS-CNRS-CNES-IRD, http://www.cesbio.ups-tlse.fr). An important part of the developments were funded by the French program Les Enveloppes Fluides et l'Environnement (LEFE, MANU, projets GloMo, SpatioGloMo and MoMu). The French program Défi InFiNiTi (CNRS) and PNTS are also acknowledged (projects Crops'IChaos and Musc & SlowFast).

If you apply this package to single time series, please quote [6]. If you apply it to multivariate time series, please quote [10]. If you apply it to infer couplings among time series, please quote [8]. If you apply it to classification, please quote [11].

HISTORICAL BACKGROUND

The global modelling technique was initiated during the early 1990s [1-3]. It takes its background from the Theory of Nonlinear Dynamical Systems. Earlier investigations can also be found in the fields of Electrical Engineering and Statistics but these mainly focused on linear problems [4]. The approach became applicable to the analysis of real world environmental behaviours by the end of the 2000s [5-7]. Recent works have shown that the approach could be applied to numerous other dynamical behaviors [8-10]. Global modelling aims to obtain deterministic models directly from observed time series.

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Author(s)

Sylvain Mangiarotti, Flavie Le Jean, Malika Chassan, Laurent Drapeau, Mireille Huc.

Maintainer: M. Huc <mireille.huc@u-paris2.fr>

References

[1] J. P. Crutchfield and B. S. McNamara, 1987. Equations of motion from a data series, Complex Systems. 1, 417-452.

[2] Gouesbet G., Letellier C., 1994. Global vector-field reconstruction by using a multivariate polynomial L2 approximation on nets, Physical Review E, 49 (6), 4955-4972.

[3] C. Letellier, L. Le Sceller, E. Marechal, P. Dutertre, B. Maheu, G. Gouesbet, Z. Fei, and J. L. Hudson, 1995. Global vector field reconstruction from a chaotic experimental signal in copper electrodissolution, Physical Review E, 51, 4262-4266.

[4] L. A. Aguirre & C. Letellier, Modeling nonlinear dynamics and chaos: A review, Mathematical Problems in Engineering, 2009, 238960.

C. Letellier, L. Le Sceller, E. Marechal, P. Dutertre, B. Maheu, G. Gouesbet, Z. Fei, and J. L. Hudson, 1995. Global vector field reconstruction from a chaotic experimental signal in copper electrodissolution, Physical Review E 51, 4262-4266.

[5] J. Maquet, C. Letellier, and L. A. Aguirre, 2007. Global models from the Canadian Lynx cycles as a first evidence for chaos in real ecosystems, Juornal of Mathematical Biology. 55(1), 21-39.

[6] Mangiarotti S., Coudret R., Drapeau L., & Jarlan L., 2012. Polynomial search and global modeling : Two algorithms for modeling chaos, Physical Review E, 86, 046205.

[7] Mangiarotti S., Drapeau L. & Letellier C., 2014. Two chaotic models for cereal crops observed from satellite in northern Morocco. Chaos, 24(2), 023130.

[8] Mangiarotti S., 2015. Low dimensional chaotic models for the plague epidemic in Bombay (1896-1911). Chaos, Solitons and Fractals, 81A, 184-186.

[9] Mangiarotti S., Peyre M. & Huc M., A chaotic model for the epidemic of Ebola Virus Disease in West Africa (2013-2016). Chaos, 26, 113112, 2016.

[10] Mangiarotti S., 2014. Modelisation globale et Caracterisation Topologique de dynamiques environnementales - de l'analyse des enveloppes fluides et du couvert de surface de la Terre a la caracterisation topolodynamique du chaos. Habilitation to Direct Research, University of Toulouse 3, France.

[11] Mangiarotti S., Sharma A.K., Corgne S., Hubert-Moy L., Ruiz L., Sekhar M., Kerr Y., Can the global modelling technique be used for crop classification? Chaos, Solitons & Fractals, in press.

allMod_nVar3_dMax2 data set

Numerical description of a list of eighteen three-dimensional chaotic sytems (see vignette 7_Retro-Modelling)

Description

A list named allMod_nVar3_dMax2 of matrix providing the numerical description of eighteen threedimensional chaotic systems:

Lorenz-1963 (\$L63), Rössler-1976 (\$R76), Burke & shaw 1981 (\$BS81), Lorenz-1984 (\$L84), Nosé

allToTest

& Hooer 1986 (\$NH86), Genesio & Tosi 1992 (\$GT92), Spott systems 1994 (\$SprF, \$SprH, \$SprK, \$SprO, \$SprP, \$SprG, \$SprM, \$SprQ, \$SprS), Chlouverakis & Sprott 2004 (\$CS2004), Li 2007 (\$Li2007) and the Cord system by Aguirre & Letellier 2012 (\$Cord2012). Each dynamical system is provided as a matrix: each column corresponds to one equation, each lines to the polynomial coefficients which order is following the convetion defined by function poLabs(nVar = 3, dMax = 2).

Usage

allMod_nVar3_dMax2

Format

An object of class list of length 18.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

References

All the references are provided in vignette 7_retro-modelling.

allToTest	A list providing the description of six models tested by the function
	autoGPoMoTest.

Description

List of 6 models available for tests (by autoGPoMoTest). Each model (mToTest1, mToTest2, etc.) is provided as a matrix of dimension 10 * 3. Each column corresponds to one equation. The order of the coefficients follows the conventions defined by poLabs(nVar = 3, dMax = 2).

Usage

allToTest

Format

An object of class list of length 6.

Author(s)

Sylvain Mangiarotti, Mireille Huc

Examples

autoGPoMoSearch Automatic search of polynomial Equations

Description

This algorithm aims to get an ensemble of possible models which integrability will be tested later with function autoGPoMoTest. By default, all the terms are considered available (Some of the terms can be excluded intentionally using the option filterReg). The maximum size of the equation depends on the model dimension nVar, and on the maximum polynomial degree dMax. The algorithm removes polynomial terms one by one using a leave-one-out method.

Usage

```
autoGPoMoSearch(
   data,
   dt,
   nVar,
   dMax,
   dMin = 0,
   weight = NULL,
   show = 0,
   underSamp = NULL,
   filterReg = NULL
```

```
)
```

Arguments

data	Input Time series: Each column is one time series that corresponds to one variable.
dt	Time sampling of the input series.
nVar	Number of variables considered in the polynomial formulation.
dMax	Maximum degree of the polynomial formulation.

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autoGPoMoTest

dMin	The minimum negative degree of the polynomial formulation (0 by default).
weight	A vector providing the binary weighting function of the input data series (0 or 1). By default, all the values are set to 1.
show	Provide (2) or not (0-1) visual output during the running process.
underSamp	Number of points used for undersampling the data. For undersamp = 1 the com- plete time series is used. For undersamp = 2, only one data out of two is kept, etc.
filterReg	A vector that specifies the template for the equation structure (for one single equation). The convention defined by poLabs is used. Value is 1 if the regressor is available, 0 if it is not.

Value

A list of two matrices:

\$filtMemo describes the selected terms (1 if the term is used, 0 if not)

\$KMemo provides the corresponding coefficients

Author(s)

Sylvain Mangiarotti, Flavie Le Jean

See Also

autoGPoMoTest, gPoMo, findAllSets, poLabs

Examples

autoGPoMoTest

Tests the numerical integrability of models and classify their dynamical regime

Description

Tests the numerical integrability of provided models (these may have been obtained with function autoGPoMoSearch), and classify these models as Divergent, Fixed Points, Periodic or not Unclassified (potentially chaotic).

Usage

```
autoGPoMoTest(
 data,
 nVar,
 dMax,
 dMin = 0,
 tin = NULL,
 dt = NULL,
 show = 1,
 verbose = 1,
 allKL = allKL,
 numValidIC = 1,
 weight = NULL,
 IstepMin = 10,
 IstepMax = 10000,
 tooFarThr = 4,
 FxPtThr = 1e-08,
 LimCyclThr = 1e-06,
 method = "rk4"
)
```

Arguments

data	Input Time series: Each column is one time series that corresponds to one variable.
nVar	Number of variables considered in the polynomial formulation.
dMax	Maximum degree of the polynomial formulation.
dMin	The minimum negative degree of the polynomial formulation (0 by default).
tin	Input date vector which length should correspond to the input time series.
dt	Sampling time of the input time series.
show	Provide (2) or not (0-1) visual output during the running process.
verbose	Gives information (if set to 1) about the algorithm progress and keeps silent if set to 0.
allKL	A list of all the models \$mToTest1, \$mToTest2, etc. to be tested. Each model is provided as a matrix.
numValidIC	Line number of the first valid initial conditions, that is, such as weight is not equal to zero.
weight	A vector providing the binary weighting function of the input data series (0 or 1). By default, all the values are set to 1.
IstepMin	The minimum number of integration step to start of the analysis (by default IstepMin = 10).
IstepMax	The maximum number of integration steps for stopping the analysis (by default IstepMax = 10000).

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tooFarThr	Divergence threshold, maximum value of the model trajectory compared to the data standard deviation. By default a trjactory is too far if the distance to the center is larger than four times the variance of the input data.
FxPtThr	Threshold used to detect fixed points.
LimCyclThr	Threshold used to detect the limit cycle.
method	The integration technique used for the numerical integration. By default, the fourth-order Runge-Kutta method (method = 'rk4') is used. Other methods such as 'ode45' or 'lsoda' may also be chosen. See package deSolve for details.

Value

A list containing:

\$okMod A vector classifying the models: diverging models (0), periodic models of period-1 (-1), unclassified models (1).

\$okMod A matrix classifying the model variables: diverging variable (0), period-1 variable (-1), period-2 variable (-2), fixed point variable (2), unclassified models (1).

\$coeff A matrix with the coefficients of one selected model

\$models A list of all the models to be tested \$mToTest1, \$mToTest2, etc. and of all selected models \$model1, \$model2, etc.

\$tout The time vector of the output time series (vector length corresponding to the longest numerical integration duration)

\$stockoutreg A list of matrices with the integrated trajectories (variable X1 in column 1, X2 in 2, etc.) for all the models \$model1, \$model2, etc.

Author(s)

Sylvain Mangiarotti, Flavie Le Jean

See Also

autoGPoMoSearch, gPoMo, poLabs

Examples

bDrvFilt

Description

Build the Savitzky-Golay derivative filter (Savitzky-Golay, 1964).

Usage

bDrvFilt(nDrv, tstep, winL = 9)

Arguments

nDrv	The number of derivatives to be computed.
tstep	Sampling time.
winL	The local window length to be used for computing the derivatives [1].

Value

dFlt A matrix of size (nDrv+1) * winL

Author(s)

Sylvain Mangiarotti

References

[1] Savitzky, A.; Golay, M.J.E., Smoothing and Differentiation of Data by Simplified Least Squares Procedures. Analytical Chemistry 36 (8), 1627-1639, 1964.

cano2M

cano2M: Converts a model in canonical form into a matrix form

Description

Converts the vectorial formulation of canonical models into a matrix formulation (that is, including explicitely all the equations). For both input, the list of terms follows the convention defined by poLabs.

Usage

```
cano2M(nVar, dMax, poly, dMin = 0)
```

combiEq

Arguments

nVar	The number of variables
dMax	The maximum degree allowed in the formulation
poly	A vector of coefficients corresponding to the regressor of the canonical function
dMin	The minimum negative degree of the polynomial formulation (0 by default).

Value

Kmod A matrix with nVar columns of the complete description of the equations. The first columns relates to the canonical part dX1/dt = X2, dX2/dt = X3 etc. and the column is the polynomial term itself

Author(s)

Sylvain Mangiarotti, Mireille Huc

See Also

drvSucc, gPoMo, poLabs

Examples

```
# A vector of polynomial terms corresponding to a canonical form:
polyTerms <- c(0.2,0,-1,0.5,0,0,0,0,0)
# Convert this vector into a matrix formulation with all the equations:
K <- cano2M(3,2,polyTerms)
# Visualize the equations:
visuEq(K,3,2)
```

combiEq

combiEq : Combine Equations from different sources

Description

Combines equations of different sources into a single system. During this combination, the polynomial maximal degree can be either imposed or optimized to reduce the model size. All the input have to follow the convention defined by poLabs.

Usage

```
combiEq(allKL, eqOrder = NULL, dMaxOut = NULL)
```

Arguments

allKL	A list of models, each provided as a matrix. A single matrix can also be pro- vided, it will be transformed into a list containing a single matrix.
eqOrder	A list of vector, providing each the equations number (relating to the input mod- els) to be kept in the output equation system. If not provided, all the equations are kept. A single matrix can also be provided, it will be transformed into a list containing a single matrix.
dMaxOut	The maximal polynomial degree of the output equation system (if not provided, this degree is deduced from the input models)

Value

KLout A matrix of the combined model

Author(s)

Sylvain Mangiarotti

See Also

gPoMo, poLabs

Examples

```
# Load models
data("allMod_nVar3_dMax2")
# Display equations of system 1
visuEq(nVar = 3, dMax = 2, K = allMod_nVar3_dMax2$NH86, substit = 1)
# Display equations of system 2
visuEq(nVar = 3, dMax = 2, K = allMod_nVar3_dMax2$R76, substit = 1)
# put the two systems in a list
allK <- list()
allK[[1]] <- allMod_nVar3_dMax2$NH86
allK[[2]] <- allMod_nVar3_dMax2$R76</pre>
# Example 1: reformulate two autonomous system in a single matrix
visuEq(K = allK[[1]], substit = c('u', 'v', 'w'))
visuEq(K = allK[[2]], substit = c('X', 'Y', 'Z'))
Knew <- combiEq(allK)</pre>
visuEq(K = Knew, substit = c('u', 'v', 'w', 'X', 'Y', 'Z'))
# Example 2
inXnote = list()
inXnote[[1]] <- c('u', 'v', 'w')
inXnote[[2]] <- c('X', 'Y', 'Z')
visuEq(K = allK[[1]], substit = inXnote[[1]])
visuEq(K = allK[[2]], substit = inXnote[[2]])
XnoteOut = c('X', 'Y', 'Z', 'u', 'v', 'w')
Knew2 <- combiEq(allK, eqOrder=c(4,5,6,1,2,3))</pre>
visuEq(K = Knew2, substit = XnoteOut)
```

compDeriv

```
# Example 3
inXnote = list()
inXnote[[1]] <- c('u', 'v', 'w')
inXnote[[2]] <- c('X', 'Y', 'Z')
visuEq(K = allK[[1]], substit = inXnote[[1]])
visuEq(K = allK[[2]], substit = inXnote[[2]])
XnoteOut = c('u', 'X', 'v', 'Y', 'w', 'Z')
Knew3 <- combiEq(allK, eqOrder=c(1,4,2,5,3,6), dMaxOut = 3)
visuEq(K = Knew3, substit = XnoteOut)</pre>
```

compDeriv

Computes the successive derivatives of a time series

Description

Computes the successive derivatives from one single time series, with the Savitzky-Golay approach (1964).

Usage

compDeriv(TS, nDrv, tstep, winL = 9)

Arguments

TS	A single time series provided as a single vector.
nDrv	The number of derivatives to be computed from the input series. The resulting number of outpout time series will thus be $nVar = nDrv + 1$.
tstep	Sampling Time of the input time series TS.
winL	The local window length used for computing the derivatives [1-2].

Value

drv A matrix containing the original variable (smoothed by the filtering process) in the first comlumn and its nDrv+1 first derivatives in the next columns (note that winL values of the original time series will be lost both at the begining and the end of the time series due to boundary effect).

Author(s)

Sylvain Mangiarotti

References

 Savitzky, A.; Golay, M.J.E., Smoothing and Differentiation of Data by Simplified Least Squares Procedures. Analytical Chemistry 36 (8), 1627-1639, 1964.
 Steinier J., Termonia Y., Deltour, J. Comments on smoothing and differentiation of data by sim-

concat

See Also

gloMoId, gPoMo, poLabs

Examples

```
# load data:
data(NDVI)
```

```
# Compute the derivatives:
drv <- compDeriv(NDVI[,1], nDrv = 3, tstep = 1/125)</pre>
```

concat

Concat Concatenates separated time series

Description

The aim of this code is to provide, from a set of multiple time series, a single concatenated time series for applying the global modeling technique to all the time time series in association.

Usage

concat(svrlTS, winL = 9)

Arguments

svrlTS	All separated time series.
winL	Total number of points used for computing the derivatives of the input time series. This parameter will be used as an input in function drvSucc to compute the derivatives.

Value

concaTS The concatenated time series.

Author(s)

Sylvain Mangiarotti, Mireille Huc

References

S. Mangiarotti, F. Le Jean, M. Huc & C. Letellier, 2016. Global modeling of aggregated and associated chaotic dynamics, Chaos, Solitons & Fractals, 83, 82-96.

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concatMulTS

Examples

```
# load data
data("svrlTS")
# Concatenate the data set into a single time series
winL = 55
concaTS <- concat(svrlTS, winL = winL)</pre>
# Plot the concatenated time series
plot(concaTS$sglTS$TS[,1], concaTS$sglTS$TS[,2],
     main = 'Concatenated time series',
     xlab = 'Time (concatenated)', ylab = 'y(t)',
     type = 'l', col = 'gray')
lines(concaTS$sglTS$TS[concaTS$sglTS$W == 1,1],
      concaTS$sglTS$TS[concaTS$sglTS$W == 1,2], type = 'p', col = 'green', cex = 0.5)
lines(concaTS$sglTS$TS[concaTS$sglTS$W == 0,1],
      concaTS$sglTS$TS[concaTS$sglTS$W == 0,2], type = 'p', col = 'red', cex = 0.5)
lines(concaTS$sglTS$TS[,1], concaTS$sglTS$W, type = 'l')
# The concatenated data set can be used for global modelling:
GPout1 <- gPoMo(data = concaTS$sglTS$TS[,2], tin = concaTS$sglTS$TS[,1],</pre>
                dMax = 2, nS = 3, winL = winL, weight = concaTS$sglTS$W, show = 1,
                IstepMin = 10, IstepMax = 6000, nPmin = 11, nPmax = 11, method = 'rk4')
```

concatMulTS	ConcatMulTS Concatenates separated time series (of single or multi-
	ples variables)

Description

The aim of this code is to provide, from multiple sets of (single or multiple) time series, a single concatenated set of time series for applying the global modeling technique to all the time time series in association.

Usage

```
concatMulTS(svrlTS, winL = 9)
```

Arguments

svrlTS	All separated sets of time series.
winL	Total number of points used for computing the derivatives of the input time series. This parameter will be used as an input in function drvSucc to compute the derivatives.

Value

concaTS A single set of concatenated time series.

Author(s)

Sylvain Mangiarotti, Mireille Huc

References

S. Mangiarotti, F. Le Jean, M. Huc & C. Letellier, 2016. Global modeling of aggregated and associated chaotic dynamics, Chaos, Solitons & Fractals, 83, 82-96.

Examples

```
# load data
data("svrlTS")
# Concatenate the data set into a single time series
winL = 55
concaTS <- concat(svrlTS, winL = winL)</pre>
# Plot the concatenated time series
plot(concaTS$sglTS$TS[,1], concaTS$sglTS$TS[,2],
     main = 'Concatenated time series',
     xlab = 'Time (concatenated)', ylab = 'y(t)',
     type = 'l', col = 'gray')
lines(concaTS$sglTS$TS[concaTS$sglTS$W == 1,1],
      concaTS$sglTS$TS[concaTS$sglTS$W == 1,2], type = 'p', col = 'green', cex = 0.5)
lines(concaTS$sglTS$TS[concaTS$sglTS$W == 0,1],
      concaTS$sglTS$TS[concaTS$sglTS$W == 0,2], type = 'p', col = 'red', cex = 0.5)
lines(concaTS$sglTS$TS[,1], concaTS$sglTS$W, type = 'l')
# The concatenated data set can be used for global modelling:
GPout1 <- gPoMo(data = concaTS$sglTS$TS[,2], tin = concaTS$sglTS$TS[,1],</pre>
                dMax = 2, nS = 3, winL = winL, weight = concaTS$sglTS$W, show = 1,
                IstepMin = 10, IstepMax = 6000, nPmin = 11, nPmax = 11, method = 'rk4')
```

d2pMax

Provides the number of polynomial terms pMax given dMax and nVar

Description

Computes the number of polynomial terms pMax used to formulate an equation given the maximal polynomial degree dMax and the number of variables nVar following the conventions as defined by fuction poLabs.

Usage

d2pMax(nVar, dMaxKnown, dMin = 0)

Arguments

nVar	Number of variables considered in the polynomial formulation.
dMaxKnown	The maximum polynomial degree dMax
dMin	The minimum negative degree of the polynomial formulation (0 by default).

Value

The number pMax of polynomial terms used to code a polynomial equation

Author(s)

Sylvain Mangiarotti

See Also

gloMoId, gPoMo, poLabs

Examples

Description

To reduce the computation time, the outputs of the simulations presented in vignette VI have been run beforehand and saved in this file.

Usage

data_vignetteIII

Format

An object of class list of length 12.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

data_vignetteVI data set

Output of the vignette VI_Sensitivity

Description

To reduce the computation time, the outputs of the simulations presented in vignette VI have been run beforehand and saved in this file.

Usage

```
data_vignetteVI
```

Format

An object of class list of length 6.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

data_vignetteVII data set

Output of the vignette VII_Retro-Modelling

Description

To reduce the computation time, the outputs of the simulations presented in vignette VII have been run beforehand and saved in this file.

Usage

```
data_vignetteVII
```

Format

An object of class list of length 29.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

derivODE2	A subfonction for the numerical integration of polynomial equations provided in a generic form following the convetion defined by function poLabs.

Description

This function provides the one step integration of polynomial Ordinary Differential Equations (ODE). This function requires the function ode (deSolve package).

Usage

derivODE2(t, x, K, dMin = 0, regS = NULL)

Arguments

t	All the dates for which the result of the numerical integration of the model must be provided
х	Current state vector (input from which the next state will be estimated)
К	A matrix providing the model description: each column corresponds to one equation which polynomial organisation is following the convention defined by function poLabs.
dMin	The minimum negative degree of the polynomial formulation (0 by default).
regS	Current states of each polynomial terms used in poLabs. These states can be deduced from the current state vector x (using the function regSeries). When available, it can be provided as an input to avoid uncessary computation.

Author(s)

Sylvain Mangiarotti

See Also

numicano, numinoisy

derivODEwMultiX	deriveODEwMultiX : A Subfonction for the numerical integration of
	polynomial equations in the generic form defined by function $poLabs$ and with External Forcing $F(t)$

Description

This function provides the one step integration of polynomial Ordinary Differential Equations (ODE). This function requires the function ode ("deSolve" package). This function has to be run with the Runge-Kutta method (method = 'rk4')

Usage

derivODEwMultiX(t, x, K, extF, regS = NULL)

Arguments

t	All the dates for which the result of the numerical integration of the model will have to be provided
x	Current state vector (input from which the next state will be estimated)
К	is the model: each column corresponds to one equation which organisation is following the convention given by function $poLabs$ which requires the definition of the model dimension $nVar$ (i.e. the number of variables) and the maximum polynomial degree dMax allowed. The last Equation correspond to the forcing variable that is artificially set to 0.
extF	is the external forcing. It is defined by two columns. The first colomn corre- spond to time t. The second column to $F(t)$ the forcing at time t. Note that when launching the integration function ode, the forcing $F(t)$ should be provided with a sampling time twice the sampling time used in t (because rk4 method will always use an intermediate time step).
regS	Current states of each polynomial terms used in poLabs. These states can be deduced from the current state vector x (using function regSeries). When available, it can be provided as an input to avoid uncessary computation.

Value

XXX

Author(s)

Sylvain Mangiarotti

Examples

```
# build a non autonomous model
nVar = 4
dMax = 3
omega = 0.2
gamma = 0.05
KDf=matrix(0, nrow = d2pMax(nVar = nVar, dMax = dMax), ncol = nVar)
KDf[11,1] = 1
KDf[2,2] = 1
KDf[5,2] = 1
KDf[11,2] = -gamma
KDf[35,2] = -1
KDf[2,3] = NA
KDf[2,4] = NA
visuEq(K = KDf, substit = c('x', 'y', 'u', 'v'))
#
# Prepare the external forcing
# number of integration time step
```

detectP11imCyc1

```
Istep <- 500
# time step
smpl <- 1 / 20
# output time vector
dater <- (0:Istep) * smpl</pre>
# hald step time vector (for Runge-Kutta integration)
daterdbl <- (0:(Istep*2 + 1)) * smpl / 2</pre>
# generate the forcing (here variables u and v)
extF = cbind(daterdbl, -0.1 * cos(daterdbl * omega), 0.05 * cos(daterdbl * 16/3*omega))
# Initial conditions to be used (external variables can be set to 0)
etatInit <- c(-0.616109362 , -0.126882584 , 0, 0)
#
# Numerical integration
reconstr2 <- ode(etatInit, dater, derivODEwMultiX,</pre>
                 KDf, extF = extF, method = 'rk4')
# Reconstruction of the output
nVarExt <- dim(extF)[2] - 1
reconstr2[,(nVar - nVarExt + 2):(nVar + 1)] <- extF[(0:Istep+1)*2, 2:(nVarExt+1)]</pre>
```

detectP1limCycl Detection of limit cycles of period-1

Description

This algorithm aim to detect period-1 limit cycles from trajectories in the phase sapce considered in a bidimensional projection.

Usage

```
detectP1limCycl(data, LimCyclThreshold = 0.01, show = 2)
```

Arguments

data	A matrix of the trajectory in a 2D space (if more than two columns are provided,
	only the two first columns are considered)
LimCyclThreshold	
	The detection threshold
show	Indicates the deepness of the feedback (from 0 to 2)

Value

Indicates if a limit cycle is detected (1) or not (0)

Author(s)

Sylvain Mangiarotti

See Also

autoGPoMoTest

drvSucc

drvSucc : Computes the successive derivatives of a time series

Description

Computes the successive derivatives from one single time series, using the Savitzky-Golay algorithm (1964).

Usage

drvSucc(tin = NULL, serie, nDeriv, weight = NULL, tstep = NULL, winL = 9)

Arguments

tin	Input date vector which length should correspond to the input time series.
serie	A single time series provided as a single vector.
nDeriv	The number of derivatives to be computed from the input time series. The re- sulting number of time series obtained in output will be nDeriv + 1.
weight	A vector providing the binary weighting function of the input data series (0 or 1). By default, all the values are set to 1.
tstep	Sampling time of the input time series. Used only if time vector tin is not provided.
winL	Number (exclusively odd number) of points of the local window used for com- puting the derivatives along the input time series. The Savitzky-Golay filter is used for this purpose [1,2].

Value

A list containing:

\$serie The original time serie

\$tin The time vector containing the dates corresponding to the original time series

\$tstep The time step (assumed to be regular)

\$tout The time vector of the output series

seriesDeriv A matrix containing the original time series (smoothed by the filtering process) in the first column and its nDeriv + 1 successive derivatives in the next ones. Note that winL values of the original time series will be lost, that is (winL - 1)/2 at the begining and (winL - 1)/2 at the end of the time series due to a computation boundary effect).

Author(s)

Sylvain Mangiarotti, Mireille Huc

drvSucc

References

[1] Savitzky, A.; Golay, M.J.E., Smoothing and Differentiation of Data by Simplified Least Squares Procedures. Analytical Chemistry 36 (8), 1627-1639, 1964.

[2] Steinier J., Termonia Y., Deltour, J. Comments on smoothing and differentiation of data by simplified least square procedure. Analytical Chemistry 44 (11): 1906-1909, 1972.

See Also

gloMoId, gPoMo, poLabs, compDeriv

Examples

```
##############
# Example 1 #
##############
# Generate a time series:
tin <- seq(0, 5, by = 0.01)
data <- 2 * sin(5*tin)</pre>
dev.new()
oldpar <- par(no.readonly = TRUE)</pre>
on.exit(par(oldpar))
par(mfrow = c(3, 1))
# Compute its derivatives:
drv <- drvSucc(tin = tin, nDeriv = 2, serie = data, winL = 5)</pre>
#
# plot original and filtered series
plot(tin, data, type='l', col = 'black', xlab = 't', ylab = 'x(t)')
lines(drv$tout, drv$seriesDeriv[,1], lty = 3, lwd = 3, col = 'green')
# analytic 1st derivative
firstD <- 10 * cos(5 * tin)
# plot both
plot(tin, firstD, type = 'l', col = 'black', xlab = 't', ylab = 'dx/dt')
lines(drv$tout, drv$seriesDeriv[,2], lty = 3, lwd = 3, col = 'green')
#
# analytic 2nd derivative
scdD <- -50 * sin(5 * tin)</pre>
# plot both
plot(tin, scdD, type = 'l', col = 'black', xlab = 't', ylab = 'd2x/dt2')
lines(drv$tout, drv$seriesDeriv[,3], lty=3, lwd = 3, col = 'green')
##############
# Example 2 #
##############
# load data:
data("Ross76")
tin <- Ross76[,1]</pre>
data <- Ross76[,2]</pre>
# Compute the derivatives
drvOut <- drvSucc(tin, data, nDeriv=4)</pre>
```

```
dev.new()
oldpar <- par(no.readonly = TRUE)</pre>
on.exit(par(oldpar))
par(mfrow = c(3, 1))
# original and smoothed variable:
plot(drvOut$tin, drvOut$serie,
     type='p', cex = 1, xlab = 'time', ylab = 'x(t)')
lines(drvOut$tout, drvOut$seriesDeriv[,1], type='p', col='red')
lines(drvOut$tout, drvOut$seriesDeriv[,1], type='l', col='red')
# 1st derivative:
plot(drvOut$tout, drvOut$seriesDeriv[,2],
     type='p', col='red', xlab = 'time', ylab = 'dx(t)/dt')
lines(drvOut$tout, drvOut$seriesDeriv[,2], type='1', col='red')
# 2nd derivative:
plot(drvOut$tout, drvOut$seriesDeriv[,3],
     type='p', col='red', xlab = 'time', ylab = 'd2x(t)/dt2')
lines(drvOut$tout, drvOut$seriesDeriv[,3], type='l', col='red')
```

extractEq

extractEq : Extracts Equations from one system

Description

Combines equations of different sources into a single system. During this combination, the polynomial maximal degree can be either imposed or optimized to reduce the model size. All the input have to follow the convention defined by poLabs.

Usage

```
extractEq(KL, eqVect)
```

Arguments

KL	A model, provided as a matrix.
eqVect	A vector of integers, providing the equations numbers to be kept in the output equation system.

Author(s)

Mireille Huc

findAllSets

Find all possible sets of equation combinations considering an ensemble of possible equation.

Description

For each equation to be retrieved, an ensemble of potential formulation is given. For instance, if three possible formulations are provided for equation (1), one for equation (2) and two for equation (3). In this case, six (i.e. 3*1*2) possible sets of equations can be obtained from these potential formulations. The aim of this program is to formulate all the potential systems from the individual formulations provided of the individual equations.

Usage

findAllSets(allFilt, nS = c(3), nPmin = 1, nPmax = 14)

Arguments

allFilt	A list with: (1) A matrix allFilt\$Xi of possible formulations for each equation (corresponding to variable Xi); And (2) a vector allFilt\$Npi providing the number of polynomial terms contained in each formulation.
nS	A vector providing the number of dimensions used for each input variables (see Examples 1 and 2). The dimension of the resulting model will be nVar = sum(nS).
nPmin	Corresponds to the minimum number of parameters (and thus of polynomial term) allowed.
nPmax	Corresponds to the maximum number of parameters (and thus of polynomial) allowed.

Value

SetsNp A list of two matrices \$Sets A matrix defining all the sets the equation combination (each line provides a combination, for instance, a line with 1,2,2 means the first equation of allFilt\$X1, the second one of allFilt\$X2 and the second one of allFilt\$X3) \$Np A matrix providing the number of parameters of all equation combination (each line provides the number of parameter of the selected equations)

Author(s)

Sylvain Mangiarotti

See Also

autoGPoMoSearch

Examples

```
##############
# Example 1 #
##############
# We build an example
allFilt <- list()</pre>
# For equation 1 (variable X1)
allFilt$Np1 <- 1
                         # only one formulation with one single parameter
# For equation 2 (variable X2)
allFilt$Np2 <- c(3, 2) # two potential formulations, with respectively three and four parameters
# For equation 3 (variable X3)
allFiltP3 <- c(4, 2) # two potential formulations, with respectively two and four parameters
# Formulations for variables Xi:
# For X1:
allFilt$X1 <- t(as.matrix(c(0,0,0,1,0,0,0,0,0,0)))
# For X2:
allFilt$X2 <- t(matrix(c(0,-0.85,0,-0.27,0,0,0,0.46,0,0,
                         0, -0.64, 0, 0, 0, 0, 0, 0.43, 0, 0),
                       ncol=2, nrow=10))
# For X3:
allFilt$X3 <- t(matrix(c(0, 0.52, 0, -1.22e-05, 0, 0, 0.99, 5.38e-05, 0, 0,
                          0, 0.52, 0, 0, 0, 0, 0.99, 0, 0, 0),
                       ncol=2, nrow=10))
# From these individual we can retrieve all possible formulations
findAllSets(allFilt, nS=c(3), nPmin=1, nPmax=14)
# if only formulations with seven maximum number of terms are expected:
findAllSets(allFilt, nS=c(3), nPmin=1, nPmax=7)
```

gloMoId

Global Model Identification

Description

Algorithm for global modelling in polynomial and canonical formulation of Ordinary Differential Equations. Univariate Global modeling aims to obtain multidimensional models from single time series (Gouesbet & Letellier 1994, Mangiarotti et al. 2012). An example of such application can be found in Mangiarotti et al. (2014) For a multivariate application, see GPoMo (Mangiarotti 2015, Mangiarotti et al. 2016).

Example: For a model dimension nVar=3, the global model will read: dX1/dt = X2dX2/dt = X3dX3/dt = P(X1, X2, X3).

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gloMoId

Usage

```
gloMoId(
  series,
  tin = NULL,
  dt = NULL,
  nVar = NULL,
  dMax = 1,
  dMin = 0,
  weight = NULL,
  show = 1,
  filterReg = NULL,
  winL = 9
)
```

Arguments

series	The original data set: either a single vector corresponding to the original variable; Or a matrix containing the original variable in the first column and its successive derivatives in the next columns. In the latter case, for the construction of n-dimensional model, series should have $nVar + 1$ columns since one more derivative will be necessary to identify the model parameters. Variable nVar will be set equal to n. In the former case, that is when only a single vector is provided, the derivatives will be automatically recomputed. Therefore, the dimension nVar expected for the model has to be provided.
tin	Input date vector which length should correspond to the input time series.
dt	Sampling time of the input time series.
nVar	Number of variables considered in the polynomial formulation.
dMax	Maximum degree of the polynomial formulation.
dMin	The minimum negative degree of the polynomial formulation (0 by default).
weight	A vector providing the binary weighting function of the input data series (0 or 1). By default, all the values are set to 1.
show	Provide (2) or not (0-1) visual output during the running process.
filterReg	A vector that specifies the template for the equation structure (for one single equation). The convention defined by poLabs is used. Value is 1 if the regressor is available, 0 if it is not.
winL	Total number of points used for computing the derivatives of the input time series. This parameter will be used as an input in function drvSucc to compute the derivatives.

Value

A list of five elements :

\$init The original time series and the successive derivatives used for the modeling.

\$filterReg The structure of the output model. Value is 1 if the regressor is available, 0 if it is not. The terms order is given by function poLabs.

\$K Values of the identified coefficients corresponding to the regressors defined in filterReg.

\$resTot The variance of the residual signal of the model.

\$resSsMod The variance of the residual signal of the closer submodels.

\$finalWeight Weighting series after boundary values were removed.

Author(s)

Sylvain Mangiarotti, Laurent Drapeau, Mireille Huc

References

[1] Gouesbet G., Letellier C., Global vector-field reconstruction by using a multivariate polynomial L2 approximation on nets, Physical Review E, 49 (6), 4955-4972, 1994.

[2] Mangiarotti S., Coudret R., Drapeau L., & Jarlan L., Polynomial search and global modeling : Two algorithms for modeling chaos, Physical Review E, 86, 046205, 2012.

[3] Mangiarotti S., Drapeau L. & Letellier C., Two chaotic models for cereal crops observed from satellite in northern Morocco. Chaos, 24(2), 023130, 2014.

[4] Mangiarotti S., Low dimensional chaotic models for the plague epidemic in Bombay (1896-1911), Chaos, Solitons & Fractals, 81(A), 184-196, 2015.

[5] Mangiarotti S., Peyre M. & Huc M., A chaotic model for the epidemic of Ebola Virus Disease in West Africa (2013-2016). Chaos, 26, 113112, 2016.

See Also

gPoMo, autoGPoMoSearch, autoGPoMoTest, poLabs

Examples

###############

Example 2

gPoMo

```
data(NDVI)
# Definition of the Model structure
terms <- c(1, 0, 0, 0, 1, 1, 1, 1, 0, 1, 1, 1, 0, 1, 1, 1, 1, 1, 1, 1)
poLabs(3,3)[terms==1]
reg <- gloMoId(NDVI [,1:1], dt=1/125, nVar=3, dMax=3,</pre>
               show=0, filterReg=terms==1)
##############
# Example 3 #
##############
# load data
data("Ross76")
# time vector
tin <- Ross76[1:500,1]
# single time series
series <- Ross76[1:500,3]</pre>
# some noise is added
series[1:100] <- series[1:100] + 0.01 * runif(1:100, min = -1, max = 1)</pre>
series[301:320] <- series[301:320] + 0.05 * runif(1:20, min = -1, max = 1)</pre>
# weighting function
W <- tin * 0 + 1
W[1:100] <- 0 # the first hundred values will not be considered
W[301:320] <- 0 # twenty other values will not be considered either
reg <- gloMoId(series, dt=1/100, weight = W, nVar=3, dMax=2, show=1)</pre>
visuEq(reg$K, 3, 2, approx = 4)
# first weight which value not equal to zero:
i1 = which(reg$finalWeight == 1)[1]
v0 <- reg$init[i1,1:3]</pre>
reconstr <- numicano(nVar=3, dMax=2, Istep=5000, onestep=1/250, PolyTerms=reg$K,</pre>
                      v0=v0, method="ode45")
plot(reconstr$reconstr[,2], reconstr$reconstr[,3], type='1', lwd = 3,
                        main='phase portrait', xlab='time t', ylab = 'x(t)', col='orange')
# original data:
lines(reg$init[,1], reg$init[,2], type='1',
      main='phase portrait', xlab='x', ylab = 'dx/dt', col='black')
# initial condition
lines(v0[1], v0[2], type = 'p', col = 'red')
```

Description

Algorithm for a Generalized Polynomial formulation of multivariate Global Modeling. Global modeling aims to obtain multidimensional models from single time series [1-2]. In the generalized (polynomial) formulation provided in this function, it can also be applied to multivariate time series [3-4].

Example:

Note that nS provides the number of dimensions used from each variable

case I

For nS=c(2,3) means that 2 dimensions are reconstructed from variable 1: the original variable X1 and its first derivative X2), and 3 dimensions are reconstructed from variable 2: the original variable X3 and its first and second derivatives X4 and X5. The generalized model will thus be such as: dX1/dt = X2

dX2'/dt = P1(X1, X2, X3, X4, X5)dX3/dt = X4dX4/dt = X5dX5/dt = P2(X1, X2, X3, X4, X5).

case II

For nS=c(1,1,1,1) means that only the original variables X1, X2, X3 and X4 will be used. The generalized model will thus be such as:

dX1/dt = P1(X1, X2, X3, X4)dX2/dt = P2(X1, X2, X3, X4)dX3/dt = P3(X1, X2, X3, X4)dX4/dt = P4(X1, X2, X3, X4).

Usage

```
gPoMo(
  data,
  tin = NULL,
  dtFixe = NULL,
  dMax = 2,
  dMin = 0,
  nS = c(3),
  winL = 9,
  weight = NULL,
  show = 1,
  verbose = 1,
  underSamp = NULL,
  EqS = NULL,
  AndManda = NULL,
  OrMandaPerEq = NULL,
  IstepMin = 2,
  IstepMax = 2000,
  nPmin = 1,
  nPmax = 14,
```

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gPoMo

```
tooFarThr = 4,
FxPtThr = 1e-08,
LimCyclThr = 1e-06,
nPminPerEq = 1,
nPmaxPerEq = NULL,
method = "rk4"
)
```

Arguments

data	Input Time series: Each column is one time series that corresponds to one variable.
tin	Input date vector which length should correspond to the input time series.
dtFixe	Time step used for the analysis. It should correspond to the sampling time of the input data. Note that for very large and very small time steps, alternative units may be used in order to stabilize the numerical computation.
dMax	Maximum degree of the polynomial formulation.
dMin	The minimum negative degree of the polynomial formulation (0 by default).
nS	A vector providing the number of dimensions used for each input variables (see Examples 1 and 2). The dimension of the resulting model will be $nVar = sum(nS)$.
winL	Total number of points used for computing the derivatives of the input time series. This parameter will be used as an input in function drvSucc to compute the derivatives.
weight	A vector providing the binary weighting function of the input data series (0 or 1). By default, all the values are set to 1.
show	Provide (2) or not (0-1) visual output during the running process.
verbose	Gives information (if set to 1) about the algorithm progress and keeps silent if set to 0.
underSamp	Number of points used for undersampling the data. For undersamp = 1 the complete time series is used. For undersamp = 2, only one data out of two is kept, etc.
EqS	Model template including all allowed regressors. Each column corresponds to one equation. Each line corresponds to one polynomial term as defined by function poLabs.
AndManda	AND-mandatory terms in the equations (all the provided terms should be in the equations).
OrMandaPerEq	OR-mandatory terms per equations (at least one of the provided terms should be in each equation).
IstepMin	The minimum number of integration step to start of the analysis (by default IstepMin = 10).
IstepMax	The maximum number of integration steps for stopping the analysis (by default IstepMax = 10000).

nPmin	Corresponds to the minimum number of parameters (and thus of polynomial term) allowed.
nPmax	Corresponds to the maximum number of parameters (and thus of polynomial) allowed.
tooFarThr	Divergence threshold, maximum value of the model trajectory compared to the data standard deviation. By default a trjactory is too far if the distance to the center is larger than four times the variance of the input data.
FxPtThr	Threshold used to detect fixed points.
LimCyclThr	Threshold used to detect the limit cycle.
nPminPerEq	Corresponds to the minimum number of parameters (and thus of polynomial term) allowed per equation.
nPmaxPerEq	Corresponds to the maximum number of parameters (and thus of polynomial) allowed per equation.
method	The integration technique used for the numerical integration. By default, the fourth-order Runge-Kutta method (method = 'rk4') is used. Other methods such as 'ode45' or 'lsoda' may also be chosen. See package deSolve for details.

Value

A list containing:

\$tin The time vector of the input time series

\$inputdata The input time series

\$tfiltdata The time vector of the filtered time series (boudary removed)

\$filtdata A matrix of the filtered time series with its derivatives

\$okMod A vector classifying the models: diverging models (0), periodic models of period-1 (-1), unclassified models (1).

\$coeff A matrix with the coefficients of one selected model

\$models A list of all the models to be tested \$mToTest1, \$mToTest2, etc. and all selected models \$model1, \$model2, etc.

\$tout The time vector of the output time series (vector length corresponding to the longest numerical integration duration)

\$stockoutreg A list of matrices with the integrated trajectories (variable X1 in column 1, X2 in 2, etc.) of all the models \$model1, \$model2, etc.

Author(s)

Sylvain Mangiarotti, Flavie Le Jean, Mireille Huc

References

[1] Gouesbet G. & Letellier C., 1994. Global vector-field reconstruction by using a multivariate polynomial L2 approximation on nets, Physical Review E, 49 (6), 4955-4972.

[2] Mangiarotti S., Coudret R., Drapeau L. & Jarlan L., Polynomial search and Global modelling:

gPoMo

two algorithms for modeling chaos. Physical Review E, 86(4), 046205.
[3] Mangiarotti S., Le Jean F., Huc M. & Letellier C., Global Modeling of aggregated and associated chaotic dynamics. Chaos, Solitons and Fractals, 83, 82-96.
[4] S. Mangiarotti, M. Peyre & M. Huc, 2016. A chaotic model for the epidemic of Ebola virus disease in West Africa (2013-2016). Chaos, 26, 113112.

See Also

gloMoId, autoGPoMoSearch, autoGPoMoTest

autoGPoMoSearch, autoGPoMoTest, visuOutGP, poLabs, predictab, drvSucc

Examples

visuEq(out3\$models\$model5, approx = 3, substit = 1) # the original Rossler system is thus retrieved

```
#Example 4
data("Ross76")
tin <- Ross76[,1]
data <- Ross76[,2:3]</pre>
# model template:
EqS <- matrix(1, ncol = 3, nrow = 10)
EqS[,1] <- c(0,0,0,1,0,0,0,0,0,0)
EqS[,2] <- c(1,1,0,1,0,1,1,1,1,1)
EqS[,3] <- c(0,1,0,0,0,0,1,1,0,0)
visuEq(EqS, substit = c('X', 'Y', 'Z'))
dev.new()
out4 <- gPoMo(data, tin=tin, dMax = 2, nS=c(2,1), show = 1,</pre>
      EqS = EqS, IstepMin = 10, IstepMax = 2000,
      nPmin = 9, nPmax = 11)
visuEq(out4$models$model2, approx = 2, substit = c("Y", "Y2", "Z"))
#Example 5
# load data
data("TSallMod_nVar3_dMax2")
#multiple (six) time series
tin <- TSallMod_nVar3_dMax2$SprK$reconstr[1:400,1]</pre>
TSRo76 <- TSallMod_nVar3_dMax2$R76$reconstr[,2:4]</pre>
TSSprK <- TSallMod_nVar3_dMax2$SprK$reconstr[,2:4]</pre>
data <- cbind(TSRo76,TSSprK)[1:400,]</pre>
dev.new()
# generalized Polynomial modelling
out5 <- gPoMo(data, tin = tin, dMax = 2, nS = c(1,1,1,1,1,1),
              show = 0, method = 'rk4',
              IstepMin = 2, IstepMax = 3,
              nPmin = 13, nPmax = 13)
# the original Rossler (variables x, y and z) and Sprott (variables u, v and w)
# systems are retrieved:
visuEq(out5$models$model347, approx = 4,
       substit = c('x', 'y', 'z', 'u', 'v', 'w'))
# to check the robustness of the model, the integration duration
# should be chosen longer (at least IstepMax = 4000)
```

GSproc

Gram-Schmidt procedure

Description

Computes regressors coefficients using the Gram-Schmidt procedure.

NDVI

Usage

GSproc(polyK, ivec, weight = NULL)

Arguments

polyK	One list including \$Y and \$phy with: \$Y a matrix for which the ith column will be used to add one orthogonal vector to the (i-1)th vectors of the current orthogonal base; and \$phy such as the current orthogonal base is given by the (i-1)th first columns of matrix polyK\$phy.
ivec	Defines i, the current vector of polyK\$Y and the current orthogonal base of pParam\$phy.
weight	The weighing vector.

Value

uNew The model parameterization, that is: The residual orthogonal vector that can be included into the current orthogonal base. If the current base is empty, uNew is equal to the input vector of \$Y; if the base is complete, uNew equals 0.

Author(s)

Sylvain Mangiarotti

NDVI

A time series of vegetation index measured from satellite

Description

A time series of 28 years of Normalized Difference Vegetation Index measured from space by the Advanced Very High Resolution Radiometer (AVHRR) sensor from 1982 to 2008 (see reference [1] for details).

Usage

NDVI

Format

An object of class data. frame with 1000 rows and 4 columns.

Author(s)

Sylvain Mangiarotti, Flavie Le Jean

References

[1] Mangiarotti S., Drapeau L. & Letellier C., 2014. Two chaotic models for cereal crops observed from satellite in northern Morocco.

numicano

Description

Function for the numerical integration of Ordinary Differential Equations of polynomial form.

Usage

```
numicano(
    nVar,
    dMax,
    dMin = 0,
    Istep = 1000,
    onestep = 1/125,
    KL = NULL,
    PolyTerms = NULL,
    v0 = NULL,
    method = "rk4"
)
```

Arguments

nVar	Number of variables considered in the polynomial formulation.
dMax	Maximum degree of the polynomial formulation.
dMin	The minimum negative degree of the polynomial formulation (0 by default).
Istep	The number of integration time steps
onestep	Time step length
KL	Matrix formulation of the model to integrate numerically
PolyTerms	Vectorial formulation of the model (only for models of canonical form)
v0	The initial conditions (a vector which length should correspond to the model dimension nVar)
method	The integration method (See package deSolve), by default method = 'rk4'.

Value

A list of two variables:

\$KL The model in its matrix formulation

\$reconstr The integrated trajectory (first column is the time, next columns are the model variables)

Author(s)

Sylvain Mangiarotti
numicano

See Also

derivODE2, numinoisy

```
##############
# Example 1 #
# For a model of general form (here the rossler model)
# model dimension:
nVar = 3
# maximal polynomial degree
dMax = 2
# Number of parameter number (by default)
pMax <- d2pMax(nVar, dMax)
# convention used for the model formulation
poLabs(nVar, dMax)
# Definition of the Model Function
a = 0.520
b = 2
c = 4
Eq1 <- c(0, -1, 0, -1, 0, 0, 0, 0, 0, 0)
Eq2 <- c(0, 0, 0, a, 0, 0, 1, 0, 0, 0)
Eq3 <- c(b,-c, 0, 0, 0, 0, 0, 1, 0, 0)
K <- cbind(Eq1, Eq2, Eq3)</pre>
# Edition of the equations
visuEq(K, nVar, dMax)
# initial conditions
v0 <- c(-0.6, 0.6, 0.4)
# model integration
reconstr <- numicano(nVar, dMax, Istep=1000, onestep=1/50, KL=K,</pre>
                      v0=v0, method="ode45")
# Plot of the simulated time series obtained
dev.new()
plot(reconstr$reconstr[,2], reconstr$reconstr[,3], type='1',
      main='phase portrait', xlab='x(t)', ylab = 'y(t)')
```

numicano

```
##############
# Example 3 #
# For a model of general form (here the rossler model)
# model dimension:
nVar = 3
# maximal polynomial degree
dMax = 2
dMin = -1
# Number of parameter number (by default)
pMax <- regOrd(nVar, dMax, dMin)[2]</pre>
# convention used for the model formulation
poLabs(nVar, dMax, dMin)
# Definition of the Model Function
a = 0.520
b = 2
c = 4
Eq1 <- c(0,-1, 0,-1, 0, 0, 0, 0, 0, 0, 1, 0, 0, 0, 0)
Eq2 <- c(0, 0, 0, a, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0)
Eq3 <- c(b, -c, 0, 0, 0, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0)
K <- cbind(Eq1, Eq2, Eq3)</pre>
# Edition of the equations
#visuEq(K, nVar, dMax)
# initial conditions
v0 <- c(-0.6, 0.6, 0.4)
# model integration
reconstr <- numicano(nVar, dMax, dMin, Istep=1000, onestep=1/50, KL=K,</pre>
                      v0=v0, method="ode45")
# Plot of the simulated time series obtained
dev.new()
plot(reconstr$reconstr[,2], reconstr$reconstr[,3], type='1',
      main='phase portrait', xlab='x(t)', ylab = 'y(t)')
```

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numiMultiX

Description

Function for the numerical integration of Ordinary Differential Equations of polynomial form including single or Multiple external forcing

Usage

```
numiMultiX(
    nVar,
    dMax,
    Istep = 1000,
    onestep = 1/125,
    KDf,
    extF = extF,
    v0 = NULL,
    method = "rk4"
)
```

Arguments

nVar	Number of variables considered in the polynomial formulation.
dMax	Maximum degree of the polynomial formulation.
Istep	The number of integration time steps. By default, Istep = 1000
onestep	The time step to be used for numerical integration
KDf	The nonautonomous model in its matrix formulation, NA (i.e. not available) values should be provided for forcing variables provided as an external signal
extF	A matrix providing the time vector in the first column, and time series of each forcing in the next ones
vØ	The initial conditions. Its length should be in agreement with the dynamical system dimension. Therefore, 0 or NA can be provided for external forcing
method	integration method. By default 'rk4' is used

Value

A list of two variables:

\$KDf The nonautonomous model in its matrix formulation

\$reconstr The integrated trajectory (first column is the time, next columns are the model variables)

Author(s)

Sylvain Mangiarotti

See Also

derivODE2, numicano, numinoisy

```
##############
# Example 1 #
##############
# build a non autonomous model
nVar = 4
dMax = 3
gamma = 0.05
KDf=matrix(0, nrow = d2pMax(nVar = nVar, dMax = dMax), ncol = nVar)
KDf[11,1] = 1
KDf[2,2] = 1
KDf[5,2] = 1
KDf[11,2] = -gamma
KDf[35,2] = -1
KDf[2,3] = NA
KDf[2,4] = NA
visuEq(K = KDf, substit = c('x', 'y', 'u', 'v'))
# build an external forcing
# number of integration time step
Istep <- 500
# time step
smpl <- 1 / 20
# output time vector
tvec <- (0:(Istep-1)) * smpl</pre>
# angular frequency (for periodic forcing)
omega = 0.2
# half step time vector (for Runge-Kutta integration)
tvecX <- (0:(Istep*2-2)) * smpl / 2</pre>
# generate the forcing (here variables u and v)
extF = cbind(tvecX, -0.1 * cos(tvecX * omega), 0.05 * cos(tvecX * 16/3*omega))
# decimate the data
extFrs <- extF[seq(1,dim(extF)[1],by=50),]</pre>
extFrs <- rbind(extFrs,extF[dim(extF)[1],])</pre>
# Initial conditions to be used (external variables can be set to 0)
etatInit <- c(-0.616109362 , -0.126882584 , NA, NA)
# model integration
out <- numiMultiX(nVar, dMax, Istep=Istep, onestep=smpl, KDf=KDf,</pre>
```

numinoisy

```
extF,
                        v0=etatInit, method="rk4")
outrs <- numiMultiX(nVar, dMax, Istep=Istep, onestep=smpl, KDf=KDf,</pre>
                        extFrs,
                        v0=etatInit, method="rk4")
dev.new
oldpar <- par(no.readonly = TRUE)</pre>
on.exit(par(oldpar))
par(mfrow = c(2, 2), # 2 \times 2 pictures on one plot
    pty = "s")
plot(out$reconstr[,2],out$reconstr[,3],
    xlab = 'x(t)', ylab = 'y(t)', type = 'l', col = 'red')
lines(outrs$reconstr[,2],outrs$reconstr[,3],
    xlab = 'x(t)', ylab = 'y(t)', type = 'l', col = 'green')
plot(out$reconstr[,2],out$reconstr[,4],
    xlab = 'x(t)', ylab = 'u(t)', type = 'l', col = 'red')
plot(out$reconstr[,4],out$reconstr[,5],
    xlab = 'u(t)', ylab = 'v(t)', type = 'l', col = 'red')
```

numinoisy

Generates time series of deterministic-behavior with stochatic pertubations (measurement and/or dynamical noise)

Description

Generates time series from Ordinary Differential Equations perturbed by dynamical and/or measurement noises

Usage

```
numinoisy(
  x0,
  t,
  K,
  varData = NULL,
  txVarBruitA = NULL,
  txVarBruitM = NULL,
  varBruitA = NULL,
  varBruitM = NULL,
  taux = NULL,
  freq = NULL,
  variables = NULL,
  method = NULL
)
```

Arguments

×0	The initial conditions. Should be a vector which size must be equal to the model dimension $\dim(K)[2]$ (the number of variables of the model defined by matrix K).
t	A vector providing all the dates for which the output are expected.
К	The Ordinary Differential Equations used to model the dynamics. The number of column should correspond to the number of variables, the number of lines to the number of parameters following the convention defined by poLabs(nVar,dMax).
varData	A vector of size nVar providing the caracteristic variances of each variable of the dynamical systems in ODE defined by matrix K. If not provided, this variance is automatically estimated.
txVarBruitA	A vector defining the ratio of ADDITIVE noise for each variable of the dynam- ical system in ODE. The additive noise is added at the end of the numerical integration process. The ratio is defined relatively to the signal variance of each variable.
txVarBruitM	A vector defining the ratio of DYNAMICAL noise for each variable of the dy- namical system in ODE. This noise is a perturbation added at each numerical integration step. The ratio is defined relatively to the signal variance of each variable.
varBruitA	A vector defining the variance of ADDITIVE noise for each variable of the dy- namical system in ODE. The additive noise is added at the end of the numerical integration process.
varBruitM	A vector defining the variance of DYNAMICAL noise for each variable of the dynamical system in ODE. This noise is a perturbation added at each numerical integration step.
taux	Generates random gaps in time series. Parameter taux defines the ratio of data to be kept (e.g. for $taux = 0.75$, 75 percents of the data are kept).
freq	Subsamples the time series. Parameter freq defines the periodicity of data kept (e.g. for $freq = 3$, 1 data out of 3 is kept).
variables	Defines which variables must be generated.
method	Defines the numerical integration method to be used. The fourth-order Runge- Kutta method is used by default (method = 'rk4'). Other method may be used (such as 'ode45' or 'lsoda'), see function ode from package deSolve for details.

Value

A list of two variables:

\$donnees The integrated trajectory (first column is the time, next columns are the model variables)

\$bruitM The level of dynamical noise

numinoisy

\$bruitA The level of additive noise

\$vectBruitM The vector of the dynamical noise used to produce the time series

\$vectBruitA The vector of the additive noise used to produce the time series

\$ecart_type The level standard deviation

Author(s)

Sylvain Mangiarotti, Malika Chassan

```
##############
# Example 1 #
# Rossler Model formulation
# The model dimension
nVar = 3
# maximal polynomial degree
dMax = 2
a = 0.520
b = 2
c = 4
Eq1 <- c(0,-1, 0,-1, 0, 0, 0, 0, 0, 0)
Eq2 <- c(0, 0, 0, a, 0, 0, 1, 0, 0, 0)
Eq3 <- c(b,-c, 0, 0, 0, 0, 0, 1, 0, 0)
K <- cbind(Eq1, Eq2, Eq3)</pre>
# Edit the equations
visuEq(K, nVar, dMax)
# initial conditions
v0 <- c(-0.6, 0.6, 0.4)
# output time required
timeOut = (0:800)/50
# variance of additive noise
varBruitA = c(0,0,0)^{2}
# variance of multiplitive noise
varBruitM = c(2E-2, 0, 2E-2)^2
# numerical integration with noise
intgr <- numinoisy(v0, timeOut, K, varBruitA = varBruitA, varBruitM = varBruitM, freq = 1)</pre>
# Plot of the simulated time series obtained
dev.new()
plot(intgr$donnees[,2], intgr$donnees[,3], type='1',
      main='phase portrait', xlab='x(t)', ylab = 'y(t)')
dev.new()
oldpar <- par(no.readonly = TRUE)</pre>
on.exit(par(oldpar))
par(mfrow = c(3, 1))
plot(intgr$donnees[,1], intgr$donnees[,2], type='1',
```

```
main='phase portrait', xlab='x(t)', ylab = 'y(t)')
lines(intgr$donnees[,1], intgr$vectBruitM[,2]*10, type='l',
    main='phase portrait', xlab='x(t)', ylab = 'e(t)*10', col='red')
plot(intgr$donnees[,1], intgr$vectBruitM[,3]*10, type='l',
    main='phase portrait', xlab='x(t)', ylab = 'e(t)*10', col='red')
lines(intgr$donnees[,1], intgr$vectBruitM[,3]*10, type='l',
    main='phase portrait', xlab='x(t)', ylab = 'e(t)*10', col='red')
plot(intgr$donnees[,1], intgr$donnees[,4], type='l',
    main='phase portrait', xlab='x(t)', ylab = 'y(t)')
lines(intgr$donnees[,1], intgr$vectBruitM[,4]*10, type='l',
    main='phase portrait', xlab='x(t)', ylab = 'e(t)*10', col='red')
```

odeBruitMult2

For the numerical integration of ordinary differential equations with dynamical noise.

Description

A subfunction for the numerical integration of Ordinary Differential Equations provided in a generic polynomial form. Model formulation follows the convention defined by function poLabs.

Usage

```
odeBruitMult2(
   x0,
   t,
   K,
   varData = NULL,
   txVarBruitM = NULL,
   varBruitM = NULL,
   method = NULL
)
```

Arguments

x0	Initial conditions
t	All the dates for which the result of the numerical integration of the model must be provided
К	A matrix providing the model description: each column corresponds to one equation which polynomial organisation is following the convention defined by function poLabs.
varData	A vector of size nVar providing the caracteristic variances of each variable of the dynamical systems in ODE defined by matrix K. If not provided, this variance is automatically estimated.

P1FxCh

txVarBruitM	A vector defining the ratio of DYNAMICAL noise for each variable of the dy- namical system in ODE. This noise is a perturbation added at each numerical integration step. The ratio is defined relatively to the signal variance of each variable.
varBruitM	A vector defining the variance of DYNAMICAL noise for each variable of the dynamical system in ODE. This noise is a perturbation added at each numerical integration step.
method	Numerical method used in the integration process. (see ode function in deSolve package for details).

Author(s)

Sylvain Mangiarotti, Malika Chassan

See Also

numinoisy

P1FxCh

A data set for testing periodicity

Description

A matrix of 6 columns corresponding to six time series, two resulting from a Period-1 limit cycle, two from regime converging to fixed point, and two relating to a chaotic behavior

Usage

P1FxCh

Format

An object of class matrix (inherits from array) with 1000 rows and 6 columns.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

P1FxChP2

Description

Trajectories for testing periodicity. The following regimes are made available: Period-1 in columns 1:2, Fixed Point in 3:4, chaotic in 5:6, Period-2 in 7:8

Usage

P1FxChP2

Format

An object of class matrix (inherits from array) with 1000 rows and 8 columns.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

p2dMax	p2dMax : provides the maximum polynomial degree dMax given the
	number of variables nVar and the number of possible polynomial
	<i>terms</i> pMax.

Description

Find the maximum polynomial degree dMax given the number of polynomial terms pMax and the system dimension nVar.

Usage

```
p2dMax(nVar, pMaxKnown, dMin = 0)
```

Arguments

nVar	Number of variables considered in the polynomial formulation.
pMaxKnown	The number of polynomial terms
dMin	The minimum negative degree of the polynomial formulation (0 by default).

Value

dMax The maximum polynomial degree

paramId

Author(s)

Sylvain Mangiarotti, Laurent Drapeau

See Also

gloMoId, gPoMo, poLabs

Examples

```
p2dMax(6,462)
# indeed:
length(poLabs(nVar=6, dMax=5))
```

paramId

For parameter Identification

Description

Estimate the polynomial coefficients.

Usage

```
paramId(allForK, drv, weight)
```

Arguments

allForK	The list of input parameters
drv	The derivative (on the equation left hand)
weight	The weighting series

Value

allForK The initial list completed with the model parameters.

poLabs

Author(s)

Sylvain Mangiarotti

poLabs

Polynomial labels order

Description

Defines the order of the polynomial labels given the number of variables nVar and the maximum polynomial degree dMax.

Usage

poLabs(nVar, dMax, dMin = 0, findIt = NULL, Xnote = "X")

Arguments

nVar	The number of variables
dMax	The maximum degree allowed in the formulation
dMin	The minimum negative degree of the polynomial formulation (0 by default).
findIt	A vector of selected terms.
Xnote	Enables to defines the notation used for the variable, by default $Xnote = 'X'$.

Value

lbls A vector of characters. Each element is the expression of one polynomial term, such as $X_1^2 X_3 X_4$

Author(s)

Sylvain Mangiarotti

See Also

visuEq

Examples

```
#Regressor order for three variables \eqn{(X1,X2,X3)} (nVar = 3) for a maximum
#polynomial degree equal to 2 (dMax = 2): poLabs(3,2)
#and for two variables only : poLabs(2,2)
```

For a quadratic equation of two variables, # the polynomial \deqn{P(X1,X2) = 0.5 + 0.3 X1 -0.25 X1 X2} # could thus be written as a vector Pvec such as: Pvec = c(0.5, 0, 0, 0.3, -0.25, 0) # considering the convention corresponding to

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predictab

```
poLabs(2,2)
# Indeed:
poLabs(2, 2, findIt = Pvec!=0)
# An alternative notation can be used with parameter Xnote
poLabs(2, 2, findIt = Pvec!=0, Xnote = 'w')
# or also
poLabs(2, 2, findIt = Pvec!=0, Xnote = c('x','y'))
```

	tab

Estimate the models performance obtained with GPoMo *in term of predictability*

Description

The algorithm aims to estimate automatically the forecasting performances of the models obtained with gPoMo.

Usage

```
predictab(
    ogp,
    fullt = NULL,
    fulldata = NULL,
    hp = NULL,
    Nech = 50,
    intSimStep = NULL,
    show = 1,
    selecmod = NULL,
    id = 1,
    selV = 1,
    na.rm = FALSE
)
```

Arguments

ogp	The output list obtained from function gPoMo.
fullt	Time vector of the data set for which predictability will be tested
fulldata	Data set for which predictability will be tested
hp	Time vector of the horizon of prediction
Nech	Number of simulations
intSimStep	Internal number of simulation steps
show	Provide (2) or not (0-1) visual output during the running process.
selecmod	A vector of the model selected.
id	The type of model to identify. id = 1 corresponds to unidentified models, that is, potentialy chaotic.

predictab

selV	Selected variable for the analysis
na.rm	Indicates if the NA should be removed ($na.rm = TRUE$) or not ($na.rm = FALSE$).

Value

ErrmodAll A list of matrix \$Predmod1, \$Predmod2, etc. and \$Errmod1, \$Errmod2, etc. providing respectively the forecasting and the forecasting error of models 1, 2, etc. Each column corresponds to one simulation starting from a specific initial condition. Each line corresponds to one horizon of prediction. Vectors corresponding to the initial condition time tE and the horizon of prediction hpE are also provided in \$tE and \$hpE, respectively. The percentiles of the distributions of error growth are provided in qt (0.05, 0.1, 0.25, 0.5, 0.75, 0.9, 0.95) and of absolute error growth in qt2 (0.5, 0.75, 0.9, 0.95, 0.98, 0.99).

Author(s)

Sylvain Mangiarotti, Mireille Huc

```
# load data
data("Ross76")
# time vector
tin <- Ross76[seq(1, 3000, by = 8), 1]
# single time series
data <- Ross76[seq(1, 3000, by = 8), 3]
# dev.new()
# plot(tin, data, xlab = 'time', ylab = 'y(t)')
# global modelling
# results are put in list outputGPoM
outputGPoM <- gPoMo(data[1:300], tin = tin[1:300], dMax = 2, nS=c(3),</pre>
                  show = 0, method = 'rk4',
                   nPmin = 10, nPmax = 12,
                  IstepMin = 150, IstepMax = 151)
#
visuOutGP(outputGPoM)
# and test predictability #
outpred <- predictab(outputGPoM, hp = 15, Nech = 30)
# manual visualisation of the outputs (e.g. for model 1):
dev.new()
image(outpred$tE, outpred$hpE, t(outpred$Errmod1),
xlab = 't', ylab = 'hp', main = 'Errmod1')
```

pTimEv

Description

Estimate the parameters variations of a model of canonical form considering a sliding window on an external dataset.

Usage

```
pTimEv(
 TS,
 nVar,
 dMax,
 TSdate,
 whatTerms = NULL,
 weight = NULL,
 wlength = 1000,
 onestep = 100,
 removeExtr = 1
)
```

Arguments

TS	The time series to be tested
nVar	Number of variables considered in the polynomial formulation.
dMax	Maximum degree of the polynomial formulation.
TSdate	The time vector
whatTerms	The terms to be considered in the analysis. Note that these are organised follow- ing the convention defined by poLabs(nVar,dMax). Since only the structure is required, if coefficients are provided, these are transformed to 1.
weight	A vector providing the binary weighting function of the input data series (0 or 1). By default, all the values are set to 1.
wlength	The window length
onestep	Step length between two estimations
removeExtr	Ratio of estimated values to be removed (if chosen equal to 0.1, only 90 disersion will be kept)

Value

A list containing:

slidingoutGM An n*(pMax+1) matrix presenting the pMax estimated parameters p1(t), p2(t) etc. column by column. The residual signal epsilon(t) is provided in the last (i.e. pMax + 1) column. Each line correspond to one date provided in TSdate

\$TSdate A time vector relating to the estimates presented in \$slidingoutGM

\$W A vector providing the output values that can kept (=1) or must be removed (=0)

\$whatTerms A vector recalling the terms taken into account in the analysis (their order refers to poLabs(nVar,dMax) function)

\$param A vector with the parameter values used to apply the function: nVar, dMax, wlength, onestep, removeExtr

Author(s)

Sylvain Mangiarotti

See Also

autoGPoMoSearch, gPoMo, poLabs

Examples

```
#Example
data(TS)
plot(TS[,1], TS[,2], type='1')
nVar <- 3
dMax <- 2
pMax <- choose(nVar+dMax,dMax)</pre>
whatTerms <- c(1,1,0,1,1,1,1,1,1,1)
# apply pTimEv
statio <- pTimEv(TS[,2], nVar, dMax, TS[,1], whatTerms = whatTerms,</pre>
                 wlength = 1000, onestep = 20, removeExtr = 0.15)
# Plot the results
dev.new()
 layout(matrix(1:12, nrow=4, ncol=3, byrow = TRUE))
 what <- which(statio$whatTerms!=0)</pre>
 for (i in what) {
      plot(statio$TSdate[statio$W==1], statio$slidingoutGM[statio$W==1,i],
           xlab='TSdate', ylab='coeff', main=poLabs(nVar,dMax)[i])
     }
 plot(statio$TSdate[statio$W==1], statio$slidingoutGM[statio$W==1,pMax+1],
       xlab='date', ylab='Epsilon', main='Resid', log = 'y')
```

reg0rd

Generate the conventional order for polynomial terms in a the polynomial formulation

Description

Generate the conventional order of the polynomial terms for the polynomial description. It is formulated as a matrix of exponents: Each column of the matrix (a,b,c, ...) corresponds to a product of the nVar available variables X1, X2, X3, etc., that is, $X1^aX2^bX3^c$, etc.

regSeries

Usage

regOrd(nVar, dMax, dMin = 0)

Arguments

nVar	The number of variables
dMax	The maximum degree allowed in the formulation
dMin	The minimum negative degree of the polynomial formulation (0 by default).

Value

A matrix of exponents. Each column corresponds to one polynomial term. Each line correspond to the exponent of one variable. For example, a column of three exponents (0,2,1) corresponds to the monomial $X1^0 \times X2^2 \times X3^1$, that is $X2^2X3$.

Author(s)

Sylvain Mangiarotti

See Also

poLabs

regSeries

Estimates the monomial time series

Description

Creates time series by multiplying given time series among them.

Usage

regSeries(nVar, dMax, series, dMin = 0, pReg = NULL)

Arguments

nVar	Number of variables considered in the polynomial formulation.
dMax	Maximum degree of the polynomial formulation.
series	A matrix containing the original time series from which the monomials are built. Each column corresponds to one given variable.
dMin	The minimum negative degree of the polynomial formulation (0 by default).
pReg	A matrix filled, for each column, with powers of time series used to create.

Value

rpFull A matrix of time series. Each column corresponds to one regressor such as $X_1^2 X_3 X_4$

Author(s)

Sylvain Mangiarotti

Examples

```
data(TSallMod_nVar3_dMax2)
sprottK <- as.matrix(TSallMod_nVar3_dMax2$SprK$reconstr)[,2:4]
dMax <- 2
nVar <- dim(sprottK)[2]
#Example 1
polySeries1 <- regSeries(nVar, dMax, sprottK)
#Example 2
p <- c(1,3,1)
polySeries2 <- regSeries(nVar, dMax, sprottK, pReg=p)</pre>
```

Rossler-1976 data set Time series of the Rossler-1976 system

Description

The Rössler system is the 3-dimensional chaotic system dx/dt = -y - z dy/dt = x + ay dz/dt = b + z(x - c), discovered by Otto E. Rössler in 1976 [1]. The following parameters and initial conditions were used to produce the present data set: a = 0.520, b = 2, c = 4and (x0, y0, z0) = (-0.04298734, 1.025536, 0.09057987). The following four columns are provided: (1) time t, (2) x(t), (3) y(t) and (4) z(t). For this parameterization, the Rössler system produces a chaotic behavior characterized by a regime non-coherent in phase (oscillations duration can be very different from one oscillation to another).

Usage

Ross76

Format

An object of class deSolve (inherits from matrix) with 4000 rows and 4 columns.

Author(s)

Sylvain Mangiarotti, Flavie Le Jean, Malika Chassan, Laurent Drapeau, Mireille Huc.

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RosYco

References

[1] O. Rössler, 1976. An Equation for Continuous Chaos, Physics Letters, 71A, 2-3, 155-157.

RosYco

Twelve Rossler-1976 time series (exclusively variable y)

Description

Twelve independant Rossler-1976 time series (variable y). The parameters used to generate the time series correspond to a phase coherent behavior. Details can be found in [1]

Usage

RosYco

Format

An object of class matrix (inherits from array) with 3000 rows and 12 columns.

Details

Another set of time series of the Rossler-1976 chaotic system

Author(s)

Sylvain Mangiarotti, Flavie Le Jean.

References

[1] Mangiarotti S., Le Jean F., Huc M. & Letellier C., Global Modeling of aggregated and associated chaotic dynamics. Chaos, Solitons and Fractals, 83, 82-96.

subSysD

subSysD : Sub-systems Disentangling

Description

Detect, disentangle and reformulate Sub-systems from an ensemble of equations.

Usage

subSysD(inK, inXnote = NULL)

Arguments

inK	A list of models, each provided as a matrix. A single matrix can also be pro- vided, it will be transformed into a list containing a single matrix.
inXnote	A vector with the names of the input variables. If not provided, default notation is used: "X1", "X2", etc.

Value

subS A matrix with the extracted subsystem

Author(s)

Sylvain Mangiarotti

See Also

gPoMo, poLabs, combiEq

```
# Load models
data("allMod_nVar3_dMax2")
# Display equations of system 1
visuEq(nVar = 3, dMax = 2, K = allMod_nVar3_dMax2$NH86, substit = 1)
# Display equations of system 2
visuEq(nVar = 3, dMax = 2, K = allMod_nVar3_dMax2$R76, substit = 1)
# put the two systems in a list
allK <- list()
allK[[1]] <- allMod_nVar3_dMax2$NH86</pre>
allK[[2]] <- allMod_nVar3_dMax2$R76
# Example 1 (two independent subsystems)
# take two separate systems and mix them
inXnote = list()
inXnote[[1]] <- c('u', 'v', 'w')
inXnote[[2]] <- c('X', 'Y', 'Z')
visuEq(K = allK[[1]], substit = inXnote[[1]])
visuEq(K = allK[[2]], substit = inXnote[[2]])
XnoteOut = c('u', 'X', 'v', 'Y', 'w', 'Z')
Knew3 <- combiEq(allK,dMaxOut = 3, eqOrder = c(1,4,2,5,3,6))
visuEq(K = Knew3, substit = XnoteOut)
# Disentangle the subsystems from the mixed equations
dstgl <- subSysD(Knew3, inXnote = XnoteOut)</pre>
## Optional
# library(igraph)
# g1<-graph.adjacency(dstgl$FM);</pre>
# l <- layout_with_fr(g1)</pre>
# plot(g1, edge.arrow.siez = .4, edge.curved=.4, vertex.label=XnoteOut, layout = 1)
# Example 2 (one subsystem included in the other)
Kduff <- matrix(0, ncol = 4, nrow = 35)
```

svrlTS

```
Kduff[11,1] <- Kduff[5,2] <- Kduff[2,3] <- 1
Kduff[35,2] <- -1
Kduff[11,2] <- -0.05
Kduff[5,4] <- 2 * acos(-1) / 6.2
Xnote <- c("x", "y", "u", "v")
visuEq(Kduff, substit = Xnote)
dstgl2 <- subSysD(Kduff, inXnote = Xnote)</pre>
```

svrlTS

A data set for the global modeling of time series in association

Description

This data set aims to test the global modelling technique when several time series of different sizes are available. Four time series are provided, all derived from the Rössler-1976 system.

Usage

svrlTS

Format

An object of class list of length 4.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

References

S. Mangiarotti, F. Le Jean, M. Huc & C. Letellier, 2016. Global modeling of aggregated and associated chaotic dynamics, Chaos, Solitons & Fractals, 83, 82-96.

testP

Periodic solution test

Description

Tests if a trajectory is periodic.

Usage

```
testP(data, wthresh = 0.1, fxPtThresh = 1e-04, show = 0)
```

Arguments

Input Time series: Each column is one time series that corresponds to one variable.
Threshold used to detect the limit cycle.
Threshold used to detect fixed points.
Provide (2) or not (0-1) visual output during the running process.

Value

periodic An integer classifying the models: diverging or unclassified trajectory (0), period-1 trajectory (-1), period-2 trajectory (-2) and fixed Point (2).

Author(s)

Sylvain Mangiarotti, Flavie Le Jean

See Also

autoGPoMoTest, gPoMo

Examples

```
#Example
# Load data:
data('P1FxChP2')
# Test a period-1 trajectory
testP(P1FxChP2[,1:2], wthresh=0.1, fxPtThresh = 1e-6, show=0)
# Test a Fixed Point trajectory
testP(P1FxChP2[,3:4], wthresh=0.1, fxPtThresh = 1e-6, show=0)
# Test a chaotic trajectory
testP(P1FxChP2[,5:6], wthresh=0.1, fxPtThresh = 1e-6, show=0)
# Test a period-2 trajectory
testP(P1FxChP2[,7:8], wthresh=0.1, fxPtThresh = 1e-6, show=0)
```

ΤS

Time series resulting from the integration of a non stationary system

Description

A 2*6001 matrix with the time vector in column one and a time series resulting from the integration of a non stationary Rössler system – parameter a varying in time: a(t) – in column two.

Usage

ΤS

Format

An object of class matrix (inherits from array) with 6001 rows and 2 columns.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

TSallMod_nVar3_dMax2 data set

Time series of three-dimensional chaotic sytems (for vignette VII_Retro-Modelling)

Description

A list of matrix providing the time series in a list named TSallMod_nVar3_dMax2 of eighteen three-dimensional chaotic systems: Lorenz-1963 (\$L63), Rössler-1976 (\$R76), Burke & shaw 1981 (\$BS81), Lorenz-1984 (\$L84), Nosé & Hooer 1986 (\$NH86), Genesio & Tosi 1992 (\$GT92), Spott systems 1994 (\$SprF, \$SprH, \$SprK, \$SprO, \$SprP, \$SprG, \$SprM, \$SprQ, \$SprS), Chlouverakis & Sprott 2004 (\$CS2004), Li 2007 (\$Li2007) and the Cord system by Aguirre & Letellier 2012 (\$Cord2012). Time series are provided in a matrix in which each column corresponds to one variable of the dynamical systems.

Usage

TSallMod_nVar3_dMax2

Format

An object of class list of length 18.

Author(s)

Sylvain Mangiarotti, Mireille Huc.

References

References for the systems are provided in vignette 'VII_retro-modelling'.

visuEq

Description

Displays the model equations for a polynomial model which description is provided as a matrix K, each column corresponding to one equation. The coefficients of the polynomial terms are given following the order defined by function poLabs. The matrix can also be provided in a list K, in this case, the matrix should be located in K\$model[[selecmod]] where selecmod should be provided as input parameter.

Usage

```
visuEq(
   K,
   nVar = NULL,
   dMax = NULL,
   dMin = 0,
   substit = 0,
   approx = FALSE,
   selecmod = NULL
)
```

Arguments

К	A matrix providing the model description: each column corresponds to one equation which polynomial organisation is following the convention defined by function poLabs.
nVar	The number of variables
dMax	The maximum degree allowed in the formulation
dMin	The minimum negative degree of the polynomial formulation (0 by default).
substit	Applies subtitutions to the default values: for substit = 0 (default value), variables are chosen as X1, X2, for substit = 1, variable X1, X2, will be replaced by x, y, z, for substit = 2, the codes provides a LaTex-like formulation of the model. The variables name can also be defined explicitely as follows: for substit = $c('x', 'H', 'T1')$, variables X1, X2, X3 will be replaced by x, H and T1.
approx	The number of extra digits to be used: for approx = FALSE (default value) digits are edited with double precision; for approx = TRUE, only the minimum number of digits is edited (in order to have all the terms different from 0) for approx = 1, 2, etc. then respectively 1, 2, etc. digits are added to the minimum number of digits corresponding to approx = TRUE.
selecmod	An integer providing the number in the sublist when the model matrix is pro- vided in a list. Should not be provided (or NULL) if the model matrix is provided directly.

visuOutGP

Value

N A list of Nvar elements presenting a set of equtions, each equation being an element of this list and written as a character strings

Author(s)

Sylvain Mangiarotti

Examples

```
#EQUATIONS VISUALISATION
# number of variables:
nVar <- 3
# maximum polynomial degree:
dMax <- 2
# polynomial organization:
poLabs(nVar,dMax)
# model construction
KL = matrix(0, ncol = 3, nrow = 10)
KL[1,1] <- KL[2,2] <- 1
KL[4,1] <- -1
KL[5,3] <- -0.123456789
# Equations visualisation:
# (a) by default, variables names X1, X2, X3 are used
visuEq(KL, nVar, dMax)
# (b) for susbstit=1, variables names x, y, y are used instead
visuEq(KL, nVar, dMax, approx = TRUE, substit=1)
# (c) the name of the variables can also be chosen manualy
visuEq(KL, nVar, dMax, approx = 3, substit=c('U', 'V', 'W'))
# A canonical model can be provided as a single vector
polyTerms <- c(0.2,0,-1,0.5,0,0,0,0,0,0)
visuEq(polyTerms, 3,2)
```

visuOutGP

visuOutGP : get a quick information of gPoMo output

Description

The algorithm aims to get a quick information about the outputs obtained with gPoMo.

Usage

```
visuOutGP(
    ogp,
    selecmod = NULL,
    id = 1,
    prioMinMax = "data",
```

```
opt3D = "TRUE",
maxPages = NULL,
seeEq = 1
)
```

Arguments

ogp	The output list obtained from gPoMo.
selecmod	A vector of the selected model. Maximum 24 models can be presented at the same time.
id	The type of model to identify. id = 1 corresponds to the unidentified models, that is, potentialy chaotic models).
prioMinMax	Gives the priority for the plots among: "data", "model", "dataonly" and "modelonly".
opt3D	Provides a 3D plot (x,y,z) when opt = 'TRUE' (the rgl library is required).
maxPages	The maximum of pages to be displayed (4 by default, but this may be insufficient when too many models remain)
seeEq	Indicates if equations should be displayed (see Eq = 1, by default) or not (see Eq = 0).

Value

A Matrix describing the terms composing each model by row. The first row corresponds to the model detection (1 unclarified, 2 diverging, 0 is fixed point, -n with n an integer, is period-n cycle')

Author(s)

Sylvain Mangiarotti

wInProd

Description

Computes weighted inner products.

Usage

wInProd(A1, A2, weight = NULL)

Arguments

A1	The input matrix 1.
A2	The input matrix 2.
weight	The weighting vector.

Value

inP The weighted inner product.

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