NTS: An R Package for Nonlinear Time Series Analysis

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Abstract  Linear time series models are commonly used in analyzing dependent data and in forecasting. On the other hand, real phenomena often exhibit nonlinear behavior and the observed data show nonlinear dynamics. This paper introduces the R package NTS that offers various computational tools and nonlinear models for analyzing nonlinear dependent data. The package fills the gaps of several outstanding R packages for nonlinear time series analysis. Specifically, the NTS package covers the implementation of threshold autoregressive (TAR) models, autoregressive conditional mean models with exogenous variables (ACMXs), functional autoregressive models, and state-space models. Users can also evaluate and compare the performance of different models and select the best one for prediction. Furthermore, the package implements flexible and comprehensive sequential Monte Carlo methods (also known as particle filters) for modeling non-Gaussian or nonlinear processes. Several examples are used to demonstrate the capabilities of the NTS package.

Introduction: nonlinear time series analysis in R

Time series analysis investigates the dynamic dependence of data observed over time or in space. While linear time series analysis has been extensively studied in the literature with many software packages widely available, nonlinear time series analysis only attracts limited attention. Although there exist some software packages for analyzing nonlinear time series focusing on different sets of tools, there are still significant gaps in capability. The NTS (Tsay et al., 2020), a recent R package, provides a number of functions for simulating, analyzing, and predicting nonlinear time series data. The available models include univariate and multivariate TAR models, conditional intensity models, nonlinear state-space models, and functional time series models. The package also features various nonlinearity tests and sequential Monte Carlo (SMC) methods. While NTS does not intend to be comprehensive, it fills the important missing parts of the existing packages, providing users valuable tools for analyzing dependent data with nonlinear dynamics. The package is now available from the Comprehensive R Archive Network at http://CRAN.R-project.org/package=NTS.

NTS incorporates the latest developments in statistical methods and algorithms for analyzing nonlinear time series data, and it makes the following contributions: (1) NTS offers various computational tools with a wide range of applications, and it fills the gaps left by the existing R functions. There are several R packages focusing on nonlinear time series. The nonlinearTseries (Garcia and Sawitzki, 2020) package implements the methods based on information theory, the NLinTS (Youssef, 2020) package introduces functions for causality detection and neural networks, and the nlts (Bjornstad, 2018) package emphasizes nonparametric autoregression and tests. NTS, providing computational tools for TAR models, ACMx models, convolutional functional autoregressive (CFAR) models, and non-Gaussian and nonlinear state-space models, consists of some of the missing pieces in the current coverage, hence making a more completed toolkit for nonlinear time series analysis in R. Other well-known modern methods for nonlinear data such as smoothing, deep learning and random forest that have been implemented in packages sm (Bowman and Azzalini, 2018), tree (Ripley, 2019) and randomForest (Breiman et al., 2018) can be adopted for nonlinear time series analysis, even though they are mainly designed for independent data. Hence, they are not included in this package. (2) NTS provides complete solutions with superior performance for the nonlinear models entertained. For example, NTS implements estimation, prediction, model building and model comparison procedures for TAR models. It allows the threshold variable in the model to be a lag variable or an exogenous variable, while the TAR (Zhang and Nieto, 2017) package, using Markov Chain Monte Carlo and Bayesian methods aiming to deal with missing values, assumes the threshold variable is exogenous. Threshold estimation methods in NTS, which perform recursive least squares or nested sub-sample searching, are more computationally efficient than the conditional least squares methods implemented in package tsDyn (Narzo et al., 2020). Furthermore, the threshold nonlinearity test proposed by Tsay (1989) in NTS is specifically designed for self-exciting TAR (SETAR) models while the existing R package nonlinearTseries just conducts general nonlinearity tests. In addition, NTS utilizes the out-of-sample forecasting to evaluate different TAR models to avoid overfitting, while other R packages such as tsDyn just compare TAR models based on AIC and residuals. (3) NTS offers additional options to existing packages with more flexibility. Specifically, NTS offers R functions to fit the ACMx model for time series analysis of count data, which allow the conditional distribution to be double Poisson, while the tscount (Liboschik et al., 2020) package uses the generalized linear models and only considers Poisson and negative binomial distributions. Another example is that
NTS implements the estimation and prediction procedures of CFAR models proposed by Liu et al. (2016), which give an intuitive and direct interpretation for functional time series analysis and provide more flexibility for estimation to deal with irregular observation locations compared to functional autoregressive models developed by Bosq (2000) introduced in the ftsa (Hyndman and Shang, 2020) package. (4) NTS provides easy access to SMC methods with various options for statistical inference. It contains different R functions which can be easily implemented for filtering and smoothing and are much more user-friendly, while the SMC (Goswami, 2011) package only writes a generic function for SMC and requires more effort from users.

The goal of this paper is to highlight the main functions of the NTS package. In the paper, we first consider different models for nonlinear time series analysis, and provide an overview of the available functions for parameter estimation, prediction and nonlinearity tests in the NTS package. Then we discuss the functions for SMC methods and demonstrate their applications via an example. Conclusions are given at the end.

Models and methods available in NTS

TAR models

TAR models are a piecewise extension of the autoregressive (AR) model proposed by Tong (1978). It has been widely used in many scientific fields, such as economics (Tong and Lim, 1980; Tiao and Tsay, 1989), finance (Domian and Louton, 1997; Narayan, 2006), among others (Chen, 1995). The models are characterized by partitioning the Euclidean space into non-overlapping regimes via a threshold variable and fitting a linear AR model in each regime (Li and Tong, 2016). The partition is by various thresholds in the domain of the threshold variable.

Let \( \{ r_j | i = 0, \ldots, m \} \) be a sequence of real numbers satisfying

\[
 r_0 = -\infty < r_1 < r_2 < \ldots < r_{m-1} < r_m = \infty.
\]

A time series \( \{ y_t | t = 1, \ldots, n \} \) follows an \( m \)-regime TAR model with threshold variable \( z_t \), threshold delay \( d > 0 \), and order \( (p_1, \ldots, p_m) \), if

\[
 y_t = \begin{cases} 
 \phi_{0,1} + \sum_{i=1}^{p_1} \phi_{1,i} y_{t-i} + \epsilon_t, & \text{if } z_{t-d} \leq r_1, \\
 \phi_{0,2} + \sum_{i=1}^{p_2} \phi_{2,i} y_{t-i} + \epsilon_t, & \text{if } r_1 < z_{t-d} \leq r_2, \\
 \vdots & \\
 \phi_{0,m} + \sum_{i=1}^{p_m} \phi_{m,i} y_{t-i} + \epsilon_t, & \text{if } r_{m-1} < z_{t-d},
\end{cases}
\]

where \( \phi_{j,i} \) are real numbers, \( \epsilon_1, \ldots, \epsilon_m \) are positive real numbers, and \( \epsilon_t \) are i.i.d random variables with mean 0 and variance 1. If the threshold variable \( z_t = y_t \) for \( t = 1, \ldots, n \), Model (1) is called a SETAR model with delay \( d \). The coefficients \( \phi_{j,i} \) must satisfy certain conditions for the stationarity of \( y_t \). These conditions are complicated in general, but some special cases are available in the literature. See, for instance, Chen and Tsay (1991) and the references therein. In particular, it is interesting to point out that the stationarity of each marginal model in (1) is not needed for the stationarity of \( y_t \). As a matter of fact, Model (1) would become more interesting when some of the marginal models are nonstationary.

Threshold estimation for two-regime TAR models

In this subsection, we introduce three algorithms for estimation of two-regime TAR models.

The two-regime TAR model can be rewritten as

\[
 y_t = (\beta_1^t x_{1,t} + \epsilon_t) I(z_{t-d} \leq r_1) + (\beta_2^t x_{2,t} + \epsilon_t) I(z_{t-d} > r_1),
\]

where \( I(\cdot) \) is the indicator function, \( x_{j,t} = (y_{t-1}, \ldots, y_{t-p})' \), and \( \beta_j = (\phi_{0,j}, \phi_{1,j}, \ldots, \phi_{p,j})' \) collects the AR coefficients in regime \( j \), \( j = 1, 2 \).

Define \( p = \max\{p_1, p_2, d\} \), and \( x_j = (x_{p+1,j}, \ldots, x_{n,j})' \) for \( j = 1, 2 \). Write \( x_1(r) = x_1 * I(z_{t-d} \leq r) \), and \( x_2(r) = x_2 * I(z_{t-d} > r) \), where * denotes the Hadamard product operator of matrices. Then Equation (2) can be re-expressed in a matrix form

\[
 y = x_1(r_1) \beta_1 + x_2(r_1) \beta_2 + \epsilon,
\]

where \( y = (y_{p+1}, \ldots, y_n)' \), \( \epsilon = (\epsilon_{p+1}, \ldots, \epsilon_n)' \), and \( \epsilon_t = [I(z_{t-d} \leq r_1) \epsilon_t + I(z_{t-d} > r_1) \epsilon_t] \) for \( t = p+1, \ldots, n \).
(Conditional) least squares: For each fixed threshold candidate \( r \), least squares method can be used to estimate the AR coefficients \( \beta_1 \) and \( \beta_2 \).

\[
\hat{\beta}_1(r) = [x_1(r)'x_1(r)]^{-1}x_1(r)'y, \quad \hat{\beta}_2(r) = [x_2(r)'x_2(r)]^{-1}x_2(r)'y.
\]

(4)

It yields the following error function

\[
S_n(r) = y'y - \hat{\beta}_1(r)'x_1(r)'x_1(r)\hat{\beta}_1(r) - \hat{\beta}_2(r)'x_2(r)'x_2(r)\hat{\beta}_2(r).
\]

(5)

To get sufficient number of observations in each regime for estimation, we assume that the threshold value \( r_i \) lies in a bounded interval \([\underline{r}, \overline{r}]\). Then it can be estimated as

\[
r_i = \text{arg} \min_{r \in \{z_{q_{i+1}}, ..., z_{n-d}\} \cap [\underline{r}, \overline{r}]} S_n(r).
\]

(6)

Recursive least squares: Recursive least squares method provides an efficient way to update the least squares solution with new observations, and is much less computationally expensive than the ordinary least squares method. When we traverse all possible thresholds and calculate \( S_n(r) \) in (5), recursive least squares can be used to estimate \( \beta_1 \) and \( \beta_2 \) in (4) helping us effectively reduce the computational cost.

Let \( S = \{z_{q_{i+1}}, ..., z_{n-d}\} \cap [\underline{r}, \overline{r}] \) be the set containing all candidates for the threshold value, \( n_0 \) be the number of elements in \( S \), \( z(j) \) be the \( j \)-th largest value in set \( S \), and \( t(j) \) be the time index for \( z(j) \). In other words, \( z(j) = z(t(j)) \).

Here is the algorithm of recursive least squares for TAR model estimation:

1. When \( z(1) \) is used as a tentative threshold value to estimate \( \beta_1 \),

\[
P_1(1) = [x_1(1)'x_1(1)]^{-1}, \quad \hat{\beta}_1(z(1)) = P_1(1)x_1(z(1))'y.
\]

When \( z(n_0) \) is used as a tentative threshold value to estimate \( \beta_2 \),

\[
P_2(n_0) = [x_2(z(n_0))'x_2(z(n_0))]^{-1}, \quad \hat{\beta}_2(z(n_0)) = P_2(n_0)x_2(z(n_0))'y.
\]

2. For \( k = 2, ..., n_0 \), we estimate the AR coefficients in regime 1 with the following

\[
K_1(k) = P_1(k-1)x_{1(1),d1}/[1 + P_1(k-1)x_{1(1),d1}P_1(k-1)],
\]

\[
P_1(k) = P_1(k-1) - K_1(k)x_{1(1),d1}P_1(k-1),
\]

\[
\hat{\beta}_1(z(k)) = \hat{\beta}_1(z(k-1)) + K_1(k)y_{1(1),d1} - \hat{\beta}_1(z(k-1))x_{1(1),d1}.
\]

For \( k = n_0 - 1, ..., 1 \), we estimate the AR coefficients in regime 2 with the following

\[
K_2(k) = P_2(k+1)x_{2(1),d2}/[1 + P_2(k+1)x_{2(1),d2}P_2(k+1)],
\]

\[
P_2(k) = P_2(k+1) - K_2(k)x_{2(1),d2}P_2(k+1),
\]

\[
\hat{\beta}_2(z(k)) = \hat{\beta}_2(z(k+1)) + K_2(k)y_{2(1),d2} - \hat{\beta}_2(z(k+1))x_{2(1),d2}.
\]

3. With \( \hat{\beta}_1(z(j)) \) and \( \hat{\beta}_2(z(j)) \) for \( j = 1, ..., n_0 \), we can obtain \( S_n(z(j)) \) and then estimate \( r_i \) with (6).

Nested sub-sample search (NeSS) algorithm: NeSS algorithm proposed by Li and Tong (2016) produces a much faster way to search threshold candidates, and reduce the computational complexity dramatically.

Li and Tong (2016) shows that there exists a positive constant \( C \) depending only on \( y, p_1 \) and \( p_2 \), such that

\[
\sup_{r \in [\underline{r}, \overline{r}]} \frac{|C - S_n(r)|}{n} \leq 0,
\]

where \( j(r) \) is a non-stochastic continuous function over \([\underline{r}, \overline{r}]\), and it is strictly monotonically increasing in \([\underline{r}, r_1] \) and strictly monotonically decreasing in \([r_2, \overline{r}] \). It implies that \( S_n(r) \) may have only one minimum value over the set \( \{k\Delta : k \in \mathbb{Z}\} \cap [\underline{r}, \overline{r}] \) for some \( \Delta > 0 \). This provides theoretical support for the following NeSS algorithm to seek the minimizer of \( S_n(r) \).

NeSS algorithm:

0. Get the initial feasible set \( S = \{z_{p-d+1}, ..., z_{n-d}\} \cap [\underline{r}, \overline{r}] \) for the threshold value estimation.

1. Obtain the 25th, 50th, and 75th percentiles of the feasible set, and define them as \( q_1, q_2 \) and \( q_3 \), respectively. Calculate \( S_n(q_1), S_n(q_2) \), and \( S_n(q_3) \).
Table 1: Comparison among various R functions for SETAR model estimation (200 replicates)

| Function                  | Sample size 200 | Sample size 2000 |
|---------------------------|-----------------|------------------|
|                           | Elapsed time    | MSE              | Elapsed time    | MSE              |
| uTAR with recursive least squares | 2.802s          | 0.0017           | 44.020s         | 2.08e-05         |
| uTAR with NeSS algorithm  | 20.360s         | 0.0017           | 25.878s         | 2.08e-05         |
| setar                     | 7.147s          | 0.0017           | 286.226s        | 2.08e-05         |

2. If \( S_n(q_1) \leq S_n(q_2) \) and \( S_n(q_1) \leq S_n(q_3) \), the feasible set is updated as \( S \cap (-\infty, q_2] \).

3. Minimize \( S_n(r) \) over the new feasible set and get \( r_1 \).

Comparing to the standard search algorithm which traverses all the threshold candidates, NeSS algorithm reduces the number of least squares operations from \( O(n) \) to \( O(\log n) \).

R functions for TAR models in NTS

In the R package NTS, the function uTAR implements recursive least squares estimation or the NeSS algorithm for TAR model estimation. The two methods both have lower computational complexity than the existing R function setar designed for SETAR model estimation in the tsDyn package, which performs least squares estimation and adopts a single grid search algorithm.

To illustrate, we use the following data generating process to compare the performance of the three methods\(^1\).

\[
y_t = \begin{cases}
1 - 0.3y_{t-1} + 0.5y_{t-2} + \epsilon_t, & \text{if } y_{t-2} \leq 0.2, \\
-1 + 0.6y_{t-1} + 0.3y_{t-2} + \epsilon_t, & \text{if } y_{t-2} > 0.2.
\end{cases}
\]  

Table 1 summarizes the average elapsed time and mean squared error (MSE) of the estimated threshold value for 200 replications. Recursive least squares method and NeSS algorithm implemented by uTAR both take shorter time than setar when sample size is large. It is also seen that when sample size is large, NeSS algorithm is the fastest, but when the sample size is relatively small, the recursive least squares method is the fastest.

Besides threshold value estimation for univariate time series, the NTS package implements data generating, forecasting, model checking, and model comparison procedures for both univariate and multivariate time series into user-friendly computational tools. Table 2 lists these functions of NTS related to TAR models. In the following we will demonstrate the usage of functions for univariate time series through the data generating process in Model (7).

Table 2: List of R functions about TAR models in the package NTS

| Function   | Description                                                                 |
|------------|-----------------------------------------------------------------------------|
| uTAR.sim   | Generate a univariate SETAR process for up to 3 regimes                     |
| uTAR       | Estimate univariate two-regime TAR models including threshold               |
| uTAR.est   | Estimate multiple regimes TAR models with known threshold(s)               |
| uTAR.pred  | Predict a fitted univariate TAR model                                       |
| thr.test   | Test for threshold nonlinearity of a scalar series                         |
| mTAR.sim   | Generate a multivariate two-regime SETAR process                           |
| mTAR       | Estimate multivariate two-regime TAR models including threshold             |
| mTAR.est   | Estimate multivariate multiple-regime TAR models                            |
| ref.mTAR   | Refine a fitted multivariate two-regime TAR model                           |
| mTAR.pred  | Predict a fitted multivariate TAR model                                     |

The function uTAR.sim generates data from a given univariate SETAR model for up to three regimes with following arguments: nob is the sample size of the generated data, ar.order specifies the AR orders for different regimes, phi is a real matrix containing the AR coefficients with one row for a

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\(^1\)The program is run on a personal computer with a 2.30GHz Intel Core(TM)i5-8259U CPU, 16GB RAM and 64-bit Operating system.
regime, \( d \) is the time delay, \( \text{cnst} \) is a vector of constant terms for the regimes, and \( \text{sigma} \) is a vector containing the standard deviations of the innovation process of the regimes. It also allows users to customize the burn-in period with option \( \text{ini} \). The function returns a list of components including the generated data from the specified TAR model (\( \text{series} \)) and the innovation series (\( \text{at} \)).

We simulate the data generating process in Model (7) with the following code. Figure 1 shows the time series plot of the first 200 observations of the simulated data.

```R
R> set.seed(1687)
R> y <- uTAR.sim(nob = 2000, arorder = c(2,2), phi = t(matrix(c(-0.3, 0.5, 0.6, + -0.3), 2, 2)), d = 2, thr = 0.2, cnst = c(1, -1), sigma = c(1, 1))
```

![Time series plot of a SETAR process](image)

**Figure 1:** Time series plot of the first 200 observations generated from the SETAR model in Equation (7).

Estimation of the threshold value of the two-regime SETAR process can be done via the function `uTAR` as illustrated below:

```R
R> thr.est<- uTAR(y = y$series, p1 = 2, p2 = 2, d = 2, thrQ = c(0, 1), Trim = c(0.1, + 0.9), include.mean = T, method = "NeSS", k0 = 50)
Estimated Threshold: 0.1951103
Regime 1:

| Estimate | Std. Error | t value | Pr(>|t|) |
|----------|------------|---------|----------|
| X1       | 1.0356009  | 0.04902797 | 21.12265 | 8.946275e-85 |
| X2       | -0.3017810 | 0.01581242  | -19.08506 | 2.383743e-71 |
| X3       | 0.4890477  | 0.02707987  | 18.05945  | 7.230880e-65 |

nob1 & sigma1: 1236 1.017973

Regime 2:

| Estimate | Std. Error | t value | Pr(>|t|) |
|----------|------------|---------|----------|
| X1       | -1.1352678 | 0.07222915 | -15.717585 | 2.107275e-48 |
| X2       | 0.5560001  | 0.03177212  | 17.499922 | 7.360494e-58 |
| X3       | -0.2122922 | 0.04641671  | -4.573616 | 5.596852e-06 |

nob2 & sigma2: 762 1.034592

Overall MLE of sigma: 1.024343
Overall AIC: 101.8515

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R> thr.est <- uTAR(y = y$series, p1 = 2, p2 = 2, d = 2, thrQ = c(0,1), Trim = c(0.1, + 0.9), include.mean = T, method = "RLS")
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```
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uTAR has the following arguments: y is a vector of observed time series, p1 and p2 are the AR order of regime 1 and regime 2, respectively, d is the delay, and thrV contains the external threshold variable z_t which should have the same length as that of y. For SETAR models, thrV is not needed and should be set to NULL. thrQ determines the lower and upper quantiles to search for threshold value. Trim defines the lower and upper trimmings to control the minimum sample size in each regime and determines [r, r] for estimation. include.mean is a logical value for including the constant term in each linear model. method decides the way to search the threshold value, and there are two choices, "RLS" for recursive least squares and "NeSS" for NeSS algorithm. k0 is only used when NeSS algorithm is selected to control the maximum sub-sample size.

From the output, the estimated threshold value is 0.195, which is close to the true value 0.2. The estimated constant terms for regime 1 and regime 2 are 1.036 and −1.135, respectively. The estimated AR coefficients for regime 1 and regime 2 are −0.302, 0.489, 0.556, and −0.212, respectively. The estimated standard deviations of the innovation processes in two regimes are 1.018 and 1.035. As expected, all estimates are significant and close their true parameters.

Here we provide an incomplete list of the returned values of the function uTAR:

- residuals: estimated innovations or residuals series.
- coefs: a 2-by-(p + 1) matrix. The first row and second row show the estimated coefficients in regime 1 and 2, respectively.
- sigma: estimated covariances of the innovation process in regime 1 and regime 2.
- thr: estimated threshold value.

Estimation of a multiple-regime TAR model with pre-specified threshold values can be done by the function uTAR.est.

R> est <- uTAR.est(y = y$series, arorder = c(2, 2), thr = thr.est$thr, d = 2,  
+ output = FALSE)

Here aroder is a row vector of positive integers containing the AR orders of all the regimes. thr collects the threshold values whose length should be the number of regimes minus 1. output is a logical value for printing out the estimation results with default being TRUE. The function uTAR.est returns the following components: coefs is a matrix with m rows in which each row contains the estimated parameters for one regime, sigma contains the estimated innovation variances for different regimes, residuals collects the estimated innovations, and sresi shows the standardized residuals.

The following R code provides one-step-ahead prediction with function uTAR.pred.

R> set.seed(12)  
R> pred <- uTAR.pred(model = est, orig = 2000, h = 1, iteration = 100, ci = 0.95,  
+ output = TRUE)  
Forecast origin: 2000  
Predictions: 1-step to 1-step  
   step forecast  
[1,] 1 -1.429635  
Pointwise 95 % confident intervals  
   step Lowb  Uppb  
int 1 -2.991667 0.6531542

The output above shows that the one-step ahead prediction for y_{2001} is −1.43. Various options in the function uTAR.pred provide users the flexibility to customize the forecasting origin with orig, forecast horizon with h, number of iterations with iterations, and confidence level with ci. The function uTAR.pred returns the prediction with pred.

The R function thr.test in the NTS package implements the F test designed for SETAR models and proposed by Tsay (1989). The test helps users detect the existence of nonlinear dynamics in the data. Below is the R code and output when we perform the nonlinearity tests with thr.test.

R> thr.test(y$series, p = 2, d = 2, ini = 40, include.mean = T)  
SETAR model is entertained  
Threshold nonlinearity test for (p,d): 2 2  
F-ratio and p-value: 213.0101 1.511847e-119

ini is the initial number of data to start the recursive least square estimation. The output shows that p-value is very small, and it indicates that there is nonlinearity in the series y$series.
Back-testing can be used to evaluate the forecasting performance of a model and to conduct model comparison between different models. Back-testing for a univariate SETAR model is implemented through the function `backTAR` with syntax:

```R
R> backTAR(model, orig, h = 1, iter = 3000)
```

where `model` is an object returned by `uTAR` or `uTAR.est`, `h` is the forecast horizon, and `iter` controls the number of simulation iterations in prediction.

The function returns the model, out-of-sample rolling prediction errors and predicted states. It also provides information for model comparison. The following example shows the out-of-sample forecasting performance of SETAR models with delay 2 and 1, respectively. It shows that the root MSE, mean absolute error, and biases of the model with delay 2 are all smaller than those of the model with delay 1. Hence, as expected, the model with delay 2 is preferred.

```R
R> set.seed(11)
R> backTAR(est, 50, 1, 3000)
Starting forecast origin: 50
1-step to 1-step out-sample forecasts
RMSE: 1.02828
MAE: 0.8172728
Bias: -0.001337478
Performance based on the regime of forecast origins:
Summary Statistics when forecast origins are in State: 1
Number of forecasts used: 1204
RMSEj: 1.029292
MAEj: 0.8172963
Biasj: 0.00259177
Summary Statistics when forecast origins are in State: 2
Number of forecasts used: 746
RMSEj: 1.026645
MAEj: 0.817235
Biasj: -0.007679051
```

The usage of functions for multivariate two-regime TAR models listed in Table 2, including `mTAR.sim`, `mTAR`, `mTAR.pred`, is similar to that of the univariate counterpart functions discussed before. The only exception is that these multivariate functions take different arguments to define the vector autoregressive(VAR) coefficients:

- `phi1, phi2`: VAR coefficient matrices of regime 1 and regime 2.
- `sigma1, sigma2`: innovation covariance matrices of regime 1 and regime 2.
- `c1, c2`: constant vectors of regime 1 and regime 2.
- `delay`: two elements `(i, d)` with `"i"` being the index of the component to be used as the threshold variable and `"d"` the delay for threshold variable.

```R
R> thr.est2 <- uTAR(y = y$series, p1 = 2, p2 = 2, d = 1, thrQ = c(0, 1),
+ Trim=c(0.1, 0.9), include.mean = T, method = "RLS")
R> est2 <- uTAR.est(y = y$series, arorder = c(2, 2), thr = thr.est2$thr, d = 1)
R> set.seed(11)
R> backTAR(est2, 50, 1, 3000)
Starting forecast origin: 50
1-step to 1-step out-sample forecasts
RMSE: 1.38731
MAE: 1.105443
Bias: -0.006635381
Performance based on the regime of forecast origins:
Summary Statistics when forecast origins are in State: 1
Number of forecasts used: 1112
RMSEj: 1.360347
MAEj: 0.909089
Biasj: 0.2462278
Summary Statistics when forecast origins are in State: 2
Number of forecasts used: 838
RMSEj: 1.4223
MAEj: 1.124622
Biasj: -0.3421769
```
The function \texttt{mTAR} conducts the nested sub-sample search algorithm and provides different choices of criterion for threshold selection with the option score, namely (AIC, det(RSS)). It has less computational cost, but only applies to two-regime models. \texttt{mTAR.est} can handle multiple regimes. They both return a list of components with the estimated VAR coefficients in \( \beta \), estimated innovation covariance matrices in \( \sigma \), and estimated innovations in residuals.

Analysis of non-Gaussian time series

Autoregressive conditional mean (ACM) models are designed for time series of count data, starting with the autoregressive conditional Poisson models, and various extensions of ACM models were investigated. The \texttt{NTS} includes a function \texttt{ACMx} for the estimation of ACMx models. Let \( y_t \) be the time series of interest, \( x_t \) be a vector containing the exogenous variables, and \( F_t = \{ y_{t-1}, y_{t-2}, \ldots; x_t, x_{t-1}, \ldots \} \). The ACMx models postulate

\[
y_t | F_t \sim F(\cdot | \mu_t),
\]

where \( \mu_t = E(y_t | F_t) = \exp(x_t' \beta) \lambda_t \), and \( \lambda_t \) follows the model

\[
\lambda_t = \omega + \sum_{i=1}^{p} \alpha_i \left[ \frac{y_{t-i}}{\exp(x_{t-i}' \beta)} \right] + \sum_{j=1}^{q} \gamma_j \lambda_{t-j},
\]

\( p \) and \( q \) are nonnegative integers, \( \omega > 0 \), and \( \alpha_i \) and \( \gamma_j \) are parameters satisfying certain conditions so that \( \lambda_t \) is always positive and finite. The conditional distribution \( F(y_t | F_t) \) can be Poisson, negative binomial, or double Poisson (Tsay and Chen, 2018).

The estimation of ACMx models is implemented via the function \texttt{ACMx} with syntax:

\[
\text{R> ACMx(y, order = c(1, 1), X = NULL, cond.dist = "po", ini = NULL)}
\]

where \( y \) is the series of count data, \( X \) is the matrix of exogenous variables, \( \text{order} \) specifies the values for \( p \) and \( q \), \( \text{cond.dist} \) determines the conditional distribution with options: ‘po’ for Poisson, ‘nb’ for negative binomial, and ‘dp’ for double Poisson, and \( \text{ini} \) collects initial parameter estimates designed for use with ‘nb’ or ‘dp’.

We illustrate the function \texttt{ACMx} with an example below:

\[
\text{R> set.seed(12)}
\]
\[
\text{R> x <- rnorm(1000)*0.1}
\]
\[
\text{R> y <- matrix(0, 1000, 1)}
\]
\[
\text{R> y[1] <- 2}
\]
\[
\text{R> lambda <- matrix(0, 1000, 1)}
\]
\[
\text{R> for (i in 2:1000){}
+ \quad lambda[i] <- 2 + 0.2*y[i-1]/exp(x[i-1]) + 0.5*lambda[i-1]
+ \quad set.seed(i)
+ \quad y[i] <- rpois(1, exp(x[i]) * lambda[i])
+ }
\]
\[
\text{R> ACMx(y, order = c(1, 1), x, "po")}
\]

Initial estimates: 1.056732 1.738874 0.05 0.5

loB: -1.056732 1e-06 1e-06 1e-06

upB: 3.170195 19.12762 0.5 0.999999

Maximized log-likelihood: -2373.08

Coefficient(s):

\[
\begin{array}{cccc}
\text{Estimate} & \text{Std. Error} & \text{t value} & \text{Pr(>|t|)} \\
\text{beta} & 1.0562836 & 0.1274853 & 8.28553 & 2.2204e-16 *** \\
\text{omega} & 2.6696378 & 0.5569954 & 4.79293 & 1.6437e-06 *** \\
\text{alpha} & 0.1579050 & 0.0265997 & 5.93634 & 2.9145e-09 *** \\
\text{gamma} & 0.4427157 & 0.0913361 & 4.84711 & 1.2528e-06 *** \\
\end{array}
\]

Signif. codes: 0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

Here a time series following the ACMx model with Poisson conditional distribution, order (1,1), \( \beta = 1, \omega = 2, \alpha = 0.2 \) and \( \gamma = 0.5 \) is generated. The R output reports the estimated coefficients which are all significant and close to their true values.
Functional time series

Functional time series analysis has received much attention since the pioneering work of Bosq (2000), and has been widely applied in many fields, including environmental science (Hormann and Kokoszka, 2010), social science (Hyndman and Shang, 2009), and finance (Diebold and Li, 2006; Horváth et al., 2013). Liu et al. (2016) proposed a new class of models called the CFAR models, which has an intuitive and direct interpretation of the dynamics of a stochastic process. The NTS encompasses functions to implement the method proposed by Liu et al. (2016).

Before presenting these R functions, we briefly introduce the CFAR model and its estimation procedure. A sequence of square integrable random functions \( \{X_t \mid t = 1, \ldots, T\} \) defined on \([0,1]\) follows a CFAR model of order \( p \) if

\[
X_t(s) = \sum_{i=1}^{p} \int_0^1 \phi_i(s-u)X_{t-i}(u)du + \epsilon_t(s), \quad s \in [0,1],
\]

where \( \phi_i(\cdot) \) are square integrable and defined on \([-1,1] \) \( i = 1, \ldots, p \) and are called the convolutional coefficient functions, and \( \epsilon_t(\cdot) \) are i.i.d. Ornstein-Uhlenbeck (O-U) processes defined on \([0,1]\) satisfying the stochastic differential equation, \( d\epsilon_t(s) = -\rho\epsilon_t(s)ds + \sigma dW_t \), \( \rho > 0 \), and \( W_t \) is a Wiener process.

In practice, \( X_t(\cdot) \) is usually observed only at discrete points, \( s_i = i/N, i = 0, \ldots, N \) for time \( t = 1, \ldots, T \). Liu et al. (2016) recovers the function \( X_t(\cdot) \) by linear interpolation,

\[
\tilde{X}_t(s) = (s_i - s)X_t(s_{i-1}) + (s - s_{i-1})X_t(s_i),
\]

for \( s_{i-1} \leq s < s_i \), and approximates \( \phi_i(\cdot) \) by cubic B-splines,

\[
\hat{\phi}_i(\cdot) \approx \phi_i(\cdot) = \sum_{j=1}^{k} \hat{\beta}_{k,i,j}B_{k,i,j}(\cdot), \quad \text{for } i = 1, \ldots, p,
\]

where \( \{B_{k,i,j}, j = 1, \ldots, k\} \) are uniform cubic B-spline basis functions with \( k \) degrees of freedom.

With the above approximation, the B-spline coefficients \( \beta = \{\hat{\beta}_{k,i,j}\}, \rho, \text{ and } \sigma^2 \) can be estimated by maximizing the approximated log-likelihood function. Specifically

\[
(\hat{\beta}, \hat{\rho}, \hat{\sigma}^2) = \arg \max Q(\beta, \rho, \sigma^2),
\]

where

\[
Q(\beta, \rho, \sigma^2) = C + \frac{(N + 1)(T - p)}{2} \ln \left( \frac{\pi \sigma^2}{\rho} \right) - \frac{N(T - p)}{2} \ln(1 - e^{-2\rho/N}) - \frac{1}{2} \sum_{t=1}^{T} e_t^\top \Sigma^{-1} e_t,
\]

where \( C \) is a constant,

\[
e_t = (e_{t,0}, \ldots, e_{t,N})^\top, \quad e_{t,\ell} = X_t(\ell/N) - \sum_{j=1}^{p} \sum_{j=1}^{k} \hat{\beta}_{k,i,j} \int_0^{1} B_{k,j}(\ell/N - u) \tilde{X}_t(s)du \quad \text{for } \ell = 0, \ldots, N,
\]

and \( \Sigma \) is an \((N + 1)\)-by-\((N + 1)\) matrix with \( \sigma^2 e^{-\rho|\ell - j|/N} \) as its \((i,j)\)-th entry.

The convolutional functions are estimated by

\[
\hat{\phi}_i(\cdot) = \sum_{j=1}^{k} \hat{\beta}_{k,i,j}B_{k,i,j}(\cdot).
\]

In the NTS package, model specification and estimation of a CFAR process can be carried out by the functions \texttt{F_test_cfar} and \texttt{est_cfar} with the syntax:

\[\texttt{R} > \texttt{F_test_cfar}(f, \text{p.max} = 6, \text{df.b} = 10, \text{grid} = 1000)\]

\[\texttt{R} > \texttt{est_cfar}(f, \text{p} = 3, \text{df.b} = 10, \text{grid} = 1000)\]

The observed functional time series is stored in \( f \), which is a \( T \)-by-\((N + 1) \) matrix, where \( T \) is the length of time, and \( (N + 1) \) is the number of discrete observations of the functional data at a given time period. \( p \) specifies the CFAR order. \( \text{df.b} \) determines the degrees of freedom \( k \) for the natural cubic splines, and \( \text{grid} \) is the number of grid points used to construct the functional time series and noise process.

The function \texttt{F_test_cfar} returns the test statistics and the \( p \)-values for a sequence of \( F \)-tests with \( H_0 : \phi_i(\cdot) = 0, i = 1, \ldots, k - 1 \) versus \( H_2 : \phi_i(\cdot) \neq 0, i = 1, \ldots, k \) for \( k = 1, \ldots, \text{p.max} \). The function \texttt{est_cfar} returns the following values: \texttt{phi_coef} collects the estimated spline coefficients \( \hat{\beta}_{k,i,j} \) for the convolutional function(s) which is a \((\text{df.b} + 1) \times p \) matrix, and \texttt{phi_func} contains the estimated convolutional function values which is a \((2 \times \text{grid} + 1) \times p \) matrix. \texttt{rho} is the estimated \( \rho \) in the O-U process, and \texttt{sigma} is estimated standard deviation of the noise process, respectively.
Figure 2: Plot of true convolution function $\phi(\cdot)$ and estimated convolution function $\hat{\phi}(\cdot)$ of the functional time series generated from Equation (8).

The function `g_cfar` in the NTS package generates a CFAR process with argument `tmax` being the length of time, `rho` is the parameter for the O-U process, `sigma` is the standard deviation of the O-U process, `phi_list` is the convolutional function(s), and `ini` is the burn-in period. It returns a list with two components. One is `cfar`, a `tmax`-by-`(grid+1)` matrix following the CFAR(p) model, and the other one `epsilon` is the innovation at time `tmax`. Function `p_cfar` provides the forecasts of a CFAR model with argument `model` as a result of an `est_cfar` fit and argument `m` as the forecasting horizon.

Let us consider a CFAR(1) process with the following convolutional coefficient function $\phi(s) = \frac{10}{\sqrt{2\pi}}e^{-50s^2}, \quad s \in [-1, 1]$,

where $\phi(\cdot)$ is the probability density function of a Gaussian random variable with mean 0 and standard deviation 0.1 truncated in the interval $[-1, 1]$. For the O-U process, $\rho = 5$ and its standard deviation is 1. The following R code simulates such a CFAR(1) process specified in Equation (8) with $N = 50$, $T = 1000$, and burn-in period 100, conducts an $F$ test, performs the estimation procedure, and provides a one-step ahead prediction.

```r
R> phi_func <- function(x){
+   return(dnorm(x, mean = 0, sd = 0.1))
+ }
R> t <- 1000; N <- 50
R> x <- g_cfar(t, rho = 5, phi_func, sigma = 1, ini = 100)
R> f <- x$cfar[, seq(1, 1001, 1000/N)]
R> F_test_cfar(f, p.max = 2, df_b = 10, grid = 1000)
Test and p-value of Order 0 vs Order 1:
[1] 1368.231 0.000
Test and p-value of Order 1 vs Order 2:
[1] 0.6848113 0.7544222
R> model <- est_cfar(f, p = 1, df_b = 10, grid = 1000)
R> print(c(model$rho, model$sigma))
[1] 4.940534 1.005315
R> pred <- p_cfar(model, f, m = 3)
```

From the output, the $p$-values for $F$ tests suggest that we choose CFAR(1) model for the data. $\hat{\rho} = 4.941$ and the standard deviation of the noise process is estimated as 1.005, and they are both close to their true values. Figure 2 plots the estimated convolutional coefficient function (dashed line) and true function $\phi(\cdot)$ (solid line). It can be seen that `est_cfar` performs well.

F_test_cfarh and est_cfarh in NTS can deal with heteroscedasticity and irregular observation locations, while the existing R package ftsa designed for functional time series assumes that the functional data are observed at regular locations. The two functions come with following arguments: `weight` is the heteroscedasticity weight function of the noise process with `grid+1` elements, `num_obs` is a `t`-by-1 vector and collects the numbers of observations at different times, and `x_pos` is a `t`-by-$(N+1)$ matrix and shows the observation locations, where $(N + 1)$ is the maximum number of observation at a time. The R code below will yield the same results as the previous one does. Hence, the output is omitted.

```r
R> num_obs <- rep(N+1, t); x_pos <- matrix(rep(seq(0, 1, 1/N), each = t), t, N+1);
R> weight0 <- function(x)(return(rep(1, length(x))))
```
State-space modeling via SMC methods

It is challenging to derive analytic solutions for filtering or smoothing of nonlinear or non-Gaussian state-space models. The SMC approach fully utilizes the dynamic nature of the model, and is an effective way to solve such complex problems (Isay and Chen, 2018). Consider the state-space model:

\[
\text{State equation: } x_t = s_t(x_{t-1}, \epsilon_t) \quad \text{or } x_t \sim q_t(\cdot \mid x_{t-1}),
\]

\[
\text{Observation equation: } y_t = h_t(x_t, \epsilon_t) \quad \text{or } y_t \sim f_t(\cdot \mid x_t),
\]

where \(x_t\) is the unobservable state variable and \(y_t\) is the observation (\(t = 1, \ldots, T\)). The underlying states evolve through the known function \(s_t(\cdot)\) and the state innovation \(\epsilon_t\), following a known conditional distribution \(q_t(\cdot)\). The information on the underlying states is observed indirectly through \(y_t\) via the known function \(h_t(\cdot)\) and with observational noise \(\epsilon_t\). The function \(h_t(\cdot)\) and the distribution of \(\epsilon_t\) are known with possibly unknown parameters to be estimated.

In general, there are four main statistical inference objectives associated with a state-space model:

1. Filtering: obtain the marginal posterior distribution of the current state \(x_t\) given the entire history of the observations up to the current time, that is, \(p(x_t \mid y_1, \ldots, y_T)\).
2. Prediction: obtain the marginal posterior distribution of the future state given the current available information, that is, \(p(x_{t+1} \mid y_1, \ldots, y_T)\).
3. Smoothing: obtain the posterior distribution of the state at the time \(t\) given the entire available information, that is, \(p(x_t \mid y_1, \ldots, y_T)\) for \(t < T\).
4. Likelihood and parameter estimation. SMC uses a set of weighted samples \(\{x_t^{(j)}, w_t^{(j)}\}\) to evaluate the likelihood function \(L(\theta) = p(y_1, \ldots, y_T \mid \theta) = \int p(x_1, \ldots, x_T, y_1, \ldots, y_T \mid \theta)dx_1 \ldots dx_T\).

SMC is a recursive procedure with three components:

- **Propagation step**: At time \(t\) for \(j = 1, \ldots, m\): draw \(x_t^{(j)}\) from a trial distribution \(g_t(x_t \mid x_{t-1}^{(j)}, y_t)\), where \(m\) is the Monte Carlo sample size. Attach it to \(x_{t-1}^{(j)}\) to form \(x_t^{(j)} = (x_{t-1}^{(j)}, x_t^{(j)})\). Compute the new weight for \(x_t^{(j)}\).
- **Resampling step**: Sample a set of indices \(\{I_1, \ldots, I_m\}\), where \(I_k \in \{1, \ldots, m\}\) according to a set of priority scores \(\pi^{(j)}\), \(j = 1, \ldots, m\). Replace the sample with \(\{x_t^{(I_1)}, w_t^{(I_1)} = w_t^{(I_2)} = \ldots = w_t^{(I_m)}\}\).
- **Inference Step**: Estimation of \(E_x[\{h(x_t)\}]\) for some integrable function \(h(\cdot)\) using the generated weighted samples \(\{x_t^{(j)}, w_t^{(j)}\}, j = 1, \ldots, m\), where \(\pi_t(\cdot)\) is the target distribution.

The selection of the propagation trial distribution \(g_t(x_t \mid x_{t-1}, y_t)\) plays a key role for an efficient implementation of SMC. Since the efficiency of SMC is determined by the variance of weight distribution, one would naturally want to choose \(g_t(\cdot)\) so that the incremental weight is as close to a constant as possible. Liu and Chen (1995, 1998) proposed the trial distribution

\[
g_t(x_t \mid x_{t-1}) = p(x_t \mid x_{t-1}, y_t) \propto q_t(x_t \mid x_{t-1})f_t(y_t \mid x_t).
\]

The proposed distribution utilizes information from both the state and observation equations, hence is termed a full information propagation step.

**Algorithm: Full information propagation step**

At time \(t\), for \(j = 1, \ldots, m\):

1. Draw \(x_t^{(j)}\) from the local posterior distribution,

\[
g_t(x_t \mid x_{t-1}^{(j)}, y_t) = p(x_t \mid x_{t-1}^{(j)}, y_t) \propto f_t(y_t \mid x_t)q_t(x_t \mid x_{t-1}^{(j)}).
\]

2. Compute the incremental weight

\[
u_t^{(j)} = \frac{1}{\sqrt{\pi_t(x_t)}}f_t(y_t \mid x_t)q_t(x_t \mid x_{t-1}^{(j)})dx_t,
\]

and the new weight \(w_t^{(j)} = w_{t-1}^{(j)}u_t^{(j)}\).
Because using full information propagation allows for more efficient estimation procedure with Rao-Blackwellization, in NTS we provide a specific function for it, different from the more general function using any user designed propagation function.

It is shown that the variance of the weight distributions stochastically increases over time. Continuing to propagate the samples with small weights forward is a waste of computational resources since inference is based on weighted average of the samples. One solution is to use resampling schemes to duplicate the importance samples and to eliminate the ones with low weights. Hence resampling steps are essential in SMC implementations. Another issue to consider is how to make inference as efficient as possible. Rao-Blackwellization is one of the effective tools that can be used.

Smoothing is another important inference to make when we analyze data with state-space models. Delayed estimation (e.g., making inference on $p(x_{t-d} | y_1, \ldots, y_t)$) is a special case of smoothing. It can be achieved simply by using the weighted sample $\{(x_{t-d}^{(i)}, w_t^{(i)})\}$. However, when $d$ is large, this sampling approach does not work well because resampling reduces the number of unique ancestors and thus increases the estimation errors. A more efficient algorithm is to calculate the backward smoothing weights, after obtaining the forward filtering weighted samples $\{(x_t^{(i)}, w_t^{(i)}) : j = 1, \ldots, m\}$. The resulting weighted sample $\{(x_t^{(j)}, w_t^{(j)}) , j = 1, \ldots, m\}$ is properly weighted.

**Algorithm: Weight marginalization SMC smoother**

Let $\tilde{x}_t^{(j)} = w_t^{(j)}, j = 1, \ldots, m$. For $t = T - 1, T - 2, \ldots, 1$ and $j = 1, \ldots, m$:

Calculate

$$\tilde{x}_t^{(j)} = \frac{\sum_{i=1}^m q_{t+1}(x_{t+1}^{(i)} | x_{t}^{(j)}) w_{t+1}^{(i)}}{\sum_{i=1}^m q_{t+1}(x_{t+1}^{(i)} | x_{t}^{(i)}) w_{t+1}^{(i)}} \tilde{x}_{t+1}^{(j)}$$

and the smoothing weight $\tilde{w}_t^{(j)} = w_t^{(j)} \tilde{x}_t^{(j)}$.

Table 3 lists functions in NTS that implement the aforementioned SMC procedures. Compared to the existing R package SMC coming with one generic function, NTS provides various functions for statistical inference and are much more user-friendly.

**Table 3: List of R functions about SMC in package NTS**

| Usage       | Function      | Description                                       |
|-------------|---------------|---------------------------------------------------|
| Generic function | SMC           | SMC method with delay but not using a full information propagation step |
|             | SMC.Smooth    | SMC smoothing method                              |
|             | SMC.Full      | SMC method using a full information propagation step |
|             | SMC.Full.RB   | SMC method using a full information propagation step with Rao-Blackwellization estimate and no delay |

The SMC function can be called by:

```r
R> SMC(Sstep, nobs, yy, mm, par, xx.init, xdim, ydim, resample.sch, + delay = 0, funH = identity)
```

The following arguments need to be specified for SMC.

- **Sstep**: A function that performs one step propagation using a proposal distribution. Its input variables include ($mm, xx, logww, yyyy, par, xdim, ydim$), where $xx$ and $logww$ are the prior iteration samples and their corresponding log weights, and $yyyy$ is the observation at current time step. It returns a list that contains $xx$ (the sample $x_t$) and $logww$ (their corresponding log weights).
- **nobs**: the number of observations, $T$.
- **yy**: the observations with $T$ columns and $ydim$ rows.
- **mm**: the Monte Carlo sample size $m$.
- **par**: a list of parameter values to pass to Sstep.
- **xx.init**: the initial samples of $x_0$.
- **xdim, ydim**: the dimension of the state variable $x_t$ and the observation $y_t$.
- **resample.sch**: a binary vector of length $nobs$, reflecting the resampling schedule. `resample.sch[i]=1` indicates resample should be carried out at step $i$.
- **delay**: the maximum delay lag for delayed weighting estimation. Default is zero.
- **funH**: a user supplied function $h(\cdot)$ for estimating $E(h(x_t) | y_{t+d})$. Default is identity function for estimating the mean with no delay. The function should be able to take vector or matrix as input and operates on each element of the input.

The function returns $xhat$, an array with dimensions ($xdim, nobs, delay+1$) and the scaled log-likelihood value `loglike`. The functions SMC.Smooth, SMC.Full and SMC.Full.RB have similar inputs and outputs, except that SMC.Smooth needs another input function for the backward smoothing step and funH is a function for estimating $E(h(x_t) | y_1, \ldots, y_T)$. 

Here is an example demonstrating how to implement SMC methods using the NTS package. Passive sonar is often used in military surveillance systems to track a target and to reduce the chance to be detected by the target. Without using an active sonar, it collects the signals generated by the target. Where multiple detectors, the location of the target can be identified (Peach, 1995; Kronhamn, 1998; Arulampalam et al., 2004; Tsay and Chen, 2018).

Suppose the target is moving in a two-dimensional plane and there are two stationary detectors located on the same plane at \((\eta_1, \eta_2)\) \((i=1,2)\) corresponding to a Cartesian coordinate. At each time \(t\) the observations consist of two angles \(\phi_{it}(i=1,2)\) of the target related to the detectors with noise. Assume that the target is moving with random acceleration in both directions. We use \(d_{it}\) and \(d_{2it}\) to denote the true locations at time \(t\) in x axis and y axis respectively, and \(s_{it}\) and \(s_{2it}\) to denote the speed at time \(t\) in x axis and y axis respectively. Let \(\Delta T\) be the time duration between two consecutive observations and we assume that the target maintains a random but constant acceleration between two consecutive observations. \(e_{it}\) and \(e_{2it}\) are the total acceleration within the period in \(x\) and \(y\) directions which are assumed to follow \(N(0, q_i^2)\) and \(N(0, q_{i2}^2)\), respectively. The motion model is

\[
d_{it} = d_{it-1} + s_{it-1} \Delta T + 0.5 \Delta T e_{it}, \quad e_{it} \sim N(0, q_i^2),
\]
\[
s_{it} = s_{it-1} + e_{it}, \quad \text{for } i=1,2.
\]

The unobserved state variable is \(x_t = (d_{1t}, d_{2t}, s_{1t}, s_{2t})'\). One observes only the directions of the target with observational errors. Assume \(\Delta T = 1\), the system can be rewritten as

**State equation** \(x_t = H x_{t-1} + W w_t, \) \(i=1,2\)

**Observation equation** \(\phi_{it} = \arctan \left( \frac{d_{2it} - \eta_{2it}}{d_{1it} - \eta_{1it}} \right) + e_{it}, \) for \(i=1,2\),

where

\[
H = \begin{pmatrix}
1 & 0 & 1 & 0 \\
0 & 1 & 0 & 1 \\
0 & 0 & 1 & 0 \\
0 & 0 & 0 & 1
\end{pmatrix}, \quad W = \begin{pmatrix}
0.5 q_1 & 0 & 0 \ 0.5 q_2 \\
q_1 & 0 & 0 \ q_2
\end{pmatrix},
\]
\[
w_t = [w_{1t} w_{2t}]', \quad w_t \sim N(0, 1) \quad \text{and} \quad e_{it} \sim N(0, r_i^2) \quad \text{for } i=1,2.
\]

This problem is highly nonlinear.

In this example, the sensors are placed as \((\eta_1, \eta_2) = (0,0)\) and \((\eta_{21}, \eta_{22}) = (20,0)\), and the measurement errors \(e_{1t}\) and \(e_{2t}\) follow \(N(0,0.02^2)\). A target moves with an initial value \(x_0 = (10,0,0.01,0.01)\) and a random acceleration with variance \(q_1^2 = q_2^2 = 0.03^2\) in both directions.

We use the following code to generate data.

```R
R> simPassiveSonar <- function(nn = 300, q, r, W, V, s2, start, seed){
+ set.seed(seed)
+ x <- matrix(nrow = 4, ncol = nn)
+ y <- matrix(nrow = 2, ncol = nn)
+ for(ii in 1:nn){
+ if(ii == 1) x[, ii] <- start
+ if(ii > 1) x[, ii] <- H %*% x[, ii - 1] + W %*% rnorm(2)
+ y[, ii] <- atan(x[, ii]/x[, ii - 1] + V %*% rnorm(2)
+ + y[, ii] <- (y[, ii] + V %*% rnorm(2) + 0.5*pi)%%pi - 0.5*pi
+ }
+ return(list(xx = x, yy = y, H = H, W = W, V = V))
+ }
R> simu_out <- simPassiveSonar(nobs = 300, q = c(0.03, 0.03), r = c(0.02, 0.02),
+ start = c(10, 10, 0.01, 0.01), H = c(1, 0, 1, 0, 0, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0),
+ W = c(0.5*q[1], 0, 0, 0.5*q[2], q[1], 0, 0, q[2]),
+ V = diag(r))
R> plot(simu_out$xx[1, ], simu_out$xx[2, ], xlab = 'x', ylab = 'y', type = 'l')
```
Figure 3 shows a simulated trajectory for \( t = 1, \ldots, 300 \), which contains two sharp turns in the middle of the observational period.

We consider using the state equation as the proposal distribution. Specifically, given \( x_t^{(j)} \), generate \( x_t^{(j)} \) using (9). The incremental weight \( u_t^{(j)} \) becomes

\[
u_t^{(j)} \propto p(y_t | x_t^{(j)}) = \sum_{i=0}^{m} p(y_t | x_t^{(i)}) \exp \left\{ - \frac{(\phi_1 - \hat{\phi}_1^{(j))})^2 + (\phi_2 - \hat{\phi}_2^{(j))})^2}{2\tau^2} \right\}.
\]

The following statements are user generated functions for SMC implementation.

```r
R> SISstep.Sonar <- function(mm, xx, logww, yy, par, xdim = 1, ydim = 1){
+ H <- par$H; W <- par$W; V <- par$V; s2 <- par$s2;
+ xx <- H%*%xx + W%*%matrix(rnorm(2*mm), nrow = 2, ncol = mm)
+ y1 <- atan(xx[2,]/xx[1,])
+ y2 <- atan(xx[2,]/(xx[1,] - s2))
+ res1 <- (yy[1] - y1 + 0.5*pi)%%pi - 0.5*pi
+ res2 <- (yy[2] - y2 + 0.5*pi)%%pi - 0.5*pi
+ uu <- -res1**2/2/V[1, 1]**2 - res2**2/2/V[2, 2]**2
+ logww <- logww + uu
+ return(list(xx = xx, logww = logww))
+ }

R> SISstep.Smooth.Sonar <- function(mm, xxt, xxt1, ww, vv, par){
+ H <- par$H; W <- par$W;
+ uu <- 1:mm
+ aa <- 1:mm
+ xxt1p <- H%*%xxt
+ for(i in 1:mm){
+ res1 <- (xxt1[, i] - xxt1p[, i])/W[1,1]
+ res2 <- (xxt1[, i] - xxt1p[, i])/W[2, 2]
+ aa[i] <- sum(exp(-0.5*res1**2 - 0.5*res2**2)*ww)
+ }
+ for(j in 1:mm){
+ res1 <- (xxt1[, j] - xxt1p[, j])/W[1,1]
+ res2 <- (xxt1[, j] - xxt1p[, j])/W[2, 2]
+ uu[j] <- sum(exp(-0.5*res1**2 - 0.5*res2**2)*vv/aa)
+ }
+ vv <- ww*uu
+ return(list(vv = vv))
+ }
```

Now we are ready to run the Monte Carlo sample with size \( m = 10000 \).
The top left panel in Figure 4 shows the delayed estimation using the delay weighting method with delay \( d = 0, 5, \) and \( 10 \), and Monte Carlo sample size \( m = 10000 \). The bottom panels in Figure 4 plot the estimation error in the \( x \) and \( y \) directions. The benefit of using delayed estimation can be clearly seen.

SMC smoothing can be implemented with the following R code. The top right panel in Figure 4 plots the smoothing results, and it shows that the SMC smoothing function performs very well.

\begin{verbatim}
R> set.seed(1)
R> par <- list(H = H, W = W, V = V, s2 = s2)
R> delay <- 10
R> out <- SMC(SISstep.Sonar, nobs, yy, mm, par, xx.init, xdim, ydim, resample.sch, delay)
R> tt <- 100:nobs
R> plot(simu_out$xx[1, tt], simu_out$xx[2, tt], xlab = 'x', ylab = 'y')
R> for(dd in c(1, 6, 11)){
+ tt <- 100:(nobs - dd)
+ lines(out$xhat[1, tt, dd], out$xhat[2, tt, dd], lty = 23 - 2*dd, lwd = 1)
+ }
R> legend(25, 22.5, legend = c("delay 0", "delay 5", "delay 10"), lty = c(21, 11, 1))
\end{verbatim}

Figure 4: Delayed filtering and smoothing results for the SMC example generated from Equation (9) and Equation (10).
Conclusion

The paper introduces the R package NTS which offers a broad collection of functions for the analysis of nonlinear time series data. We briefly review various nonlinear time series models, including TAR models, ACMx models, CFAR models, and state-space models. The associated estimation, identification, and forecasting procedures are discussed. The NTS package provides computational tools to fit these models, to evaluate their performance, and to provide predictions. Furthermore, the functions can be used, extended, and modified within the package to analyze larger univariate/multivariate, Gaussian/non-Gaussian time series. These features enable users to carry out a comprehensive and complex analysis of time series without the constraints from software availability.

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