Nonlinear autoregressive time series models in R using tsDyn version 0.6

last revised 22/02/2007 by Antonio, Fabio Di Narzo

## 1 Introduction

tsDyn is an R package for the estimation of a number of nonlinear time series models. The package is at an early stage, and may presumably change significantly in the near future. However, it is quite usable in the current version.

Each function in the package has at least a minimal help page, with one or more working examples and detailed explanation of function arguments and returned values. In this document we try to give an overall guided tour of package contents, with some additional notes which are generally difficult to put in the context of a manual.

This guide is divided into 3 main sections:

- Explorative analysis tools
- Nonlinear autoregressive models
- A case study

## 2 Explorative analysis

## 2.1 Bivariate and trivariate relations

A first explorative analysis should include inspecting the distribution of  $(x_t, x_{t-l})$ and that of  $(x_t, x_{t-l_1}, x_{t-l_2})$  for some lags  $l, l_1, l_2$ . This can be done easily in R in a variety of ways. The tsDyn package provide functions autopairs and autotriples for this purpose.

The **autopairs** function displays, in essence, a scatterplot of time series  $x_t$  versus  $x_{t-lag}$ . The main arguments to the function are the time series and the desired lag. The scatterplot may be also processed to produce bivariate kernel density estimations, as well as nonparametric kernel autoregression estimations. The type of output is governed by the argument type. Possibile values, along with their meanings, are:

lines	directed lines
points	simple scatterplot
levels	iso-density levels
persp	density perspective plot
image	density image map

regression kernel autoregression line superposed to scatterplot For kernel density and regression estimation, you can specify also the kernel window h. A typical call to that function can be: All arguments (except the time series x) have default values.

By default, if running in an interactive environment, the function displays a simple experimental cross-platform GUI, where you can change function parameters and watch interactively how the plot changes.

Similar to autopairs, there is the autotriples function. This shows  $x_t$  versus  $(x_{t-lag1}, x_{t-lag2})$ , so that the user has to specify time series x and lags lag1 and lag2. The scatterplot can be processed to produce kernel regression estimates. Plotting possibilities are:

levels iso-values lines
persp perspective plot
image image map
lines directed lines
points simple scatterplot

#### 2.2 Linearity

An interesting tool for inspecting possible nonlinearities in the time series is the *locally linear autoregressive fit* plot, proposed by Casdagli [1]. Suppose you think that the dynamical system underlying your time series is best reconstructed with embedding dimension m and time delay d. Then the locally linear autoregressive fit plot displays the relative error made by forecasting time series values with linear models of the form:

$$x_{t+s} = \phi_0 + \phi_1 x_t + \ldots + \phi_m x_{t-(m-1)d}$$

estimated on points in the sphere of radius  $\epsilon$  around  $\mathbf{x}_t^m$  for a range of values of  $\epsilon$ . A minimum attained at relatively small values of  $\epsilon$  may indicate that a global linear model would be inappropriate for the approximation of the time series dynamics.

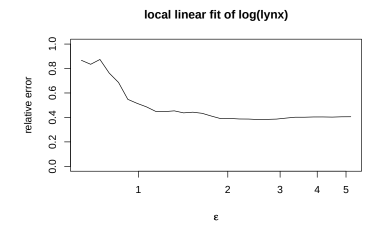
For this analysis tsDyn proposes the function llar which accepts, among others, the following arguments:

#### **x** time series

m, d, steps embedding parameters (see the above model formulation) The function returns a 'llar' object, which can be plotted with the generic plot method. So, a typical usage would be:

\_ R code \_

obj <- llar(log(lynx), m = 3) plot(obj)

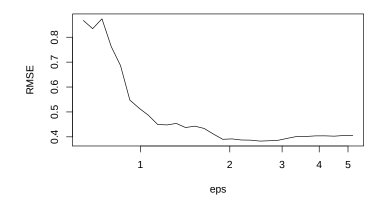


However, the obj object can be explicitely converted in an ordinary data.frame:

	_ R code
obj <- data.frame(obj)	
with variables:	
	_ R code
names(obj)	- 17 0000
	_ output
[1] "RMSE" "eps" "frac"	"avfound"

where 'RMSE' stands for Relative Mean Square Error, and eps is enough selfexplaining. You can explore this object with usual R commands dedicated to data.frames, such as:

	B. code	
plot(RMSE ~	eps, data = obj, type = "1", log = "x")	



### 2.3 Tests (experimental)

tsDyn implements conditional mutual independence and linearity tests as described in Manzan [5]. Function implementations are rather basic, and little tested. Use them carefully!

The delta.test function performs a bootstrap test of independence of  $x_t$  versus  $x_{t-md}$  conditional on intermediate observations  $\{x_{t-d}, \ldots, x_{t-(m-1)d}\}$ . The test statistic, available with the function delta, is based on the sample correlation integral, and calls internally the d2 function provided by the tseriesChaos package. Among others things, the test requires the specification of a neighborhood window  $\epsilon$ .

Function arguments are the time series x, a vector of embedding dimensions m, time delay d, a vector of neighborhood windows eps, the number of bootstrap replications B. However, default values are available for m, d, eps and B, so that a typical call can be:

	к соде	
delta.test(x)		

The return value is a matrix of p-values, labelled with their associated embedding dimensions and neighborhood windows (normally multiple values are tried simultaneously).

The delta.lin.test function performs a bootstrap test of linear dipendence of  $x_t$  versus  $x_{t-md}$  conditional on intermediate observations  $\{x_{t-d}, \ldots, x_{t-(m-1)d}\}$ . The test statistic is available with the function delta.lin. The function arguments and returned values are the same as those of delta.test.

### 3 Nonlinear autoregressive time series models

Consider the discrete-time univariate stochastic process  $\{X_t\}_{t \in T}$ . Suppose  $X_t$  is generated by the map:

$$X_{t+s} = f(X_t, X_{t-d}, \dots, X_{t-(m-1)d}; \theta) + \epsilon_{t+s}$$
(1)

with  $\{\epsilon_t\}_{t\in T}$  white noise,  $\epsilon_{t+s}$  indipendent w.r.t.  $X_{t+s}$ , and with f a generic function from  $\mathbf{R}^m$  to  $\mathbf{R}$ . This class of models is frequently referenced in the literature with the acronym NLAR(m), which stands for *NonLinear AutoRegressive* of order m.

In (1), we have implicitely defined the *embedding dimension* m, the *time delay* d and the *forecasting steps* s. The vector  $\theta$  indicates a generic vector of parameters governing the shape of f, which we would estimate on the basis of some empirical evidence (i.e., an observed time series  $\{x_1, x_2, \ldots, x_N\}$ ).

In tsDyn some specific NLAR models are implemented. For a list of currently available models, type:

	R code
<pre>availableModels()</pre>	
	output
[1] "linear" "nnetTs" "setar" "]	±

Each model can be estimated using a function which takes the name of the model as indicated by availableModels. I.e., use linear for fitting a linear model.

All those functions returns an object of base class **nlar**, from which informations can be extracted using some common methods. Among others:

print(obj) #prints basic infos on fitted model and estimated parameters summary(obj) #if possible, shows more detailed infos and diagnostics on estimated model plot(obj) #shows common diagnostic plots

Another method that can be useful for inspecting the estimated model properties is the **predict** method:

	R code	
<pre>x.new &lt;- predict(obj, n.ahead = )</pre>	)	

This function attempts to extend of n.ahead observations of the original time series used for estimating the model encapsulated in obj using the so called skeleton of the fitted model. Assuming that from (1) we estimated f as  $\hat{f} = f(\cdot; \hat{\theta})$ , using the time series  $\mathbf{x} = \{x_1, x_2, \dots, x_N\}$ , we have:

$$\hat{x}_{N+1} = \hat{f}(x_{N-s}, x_{N-s-d}, \dots, x_{N-s-(m-1)d})$$

$$\hat{x}_{N+2} = \hat{f}(x_{N-s+1}, x_{N-s+1-d}, \dots, x_{N-s+1-(m-1)d})$$

$$\dots$$

$$\hat{x}_{N+S} = \hat{f}(x_{N-s+(S-1)}, x_{N-s+(S-1)-d}, \dots, x_{N-s+(S-1)-(m-1)d})$$

A detailed description of some actually implemented models follows.

#### 3.1 Linear models

$$X_{t+s} = \phi + \phi_0 X_t + \phi_1 X_{t-d} + \dots + \phi_m X_{t-(m-1)d} + \epsilon_{t+s}$$
(2)

It's a classical AR(m) model, and its specification doesn't require additional hyper-parameters. Estimation is done via CLS (Conditional Least Squares). The summary command returns asymptotics standard errors for the estimated coefficients, based on the normality assumption of residuals.

Note that in R there are plenty of functions for AR (and, more generally, ARMA) models estimation, with different estimation methods, such as ML, CLS, Yule-Walker, .... If you really need to fit linear models, use these methods directly.

#### 3.2 SETAR models

$$X_{t+s} = \begin{cases} \phi_1 + \phi_{10}X_t + \phi_{11}X_{t-d} + \dots + \phi_{1L}X_{t-(L-1)d} + \epsilon_{t+s} & Z_t \le \text{th} \\ \phi_2 + \phi_{20}X_t + \phi_{21}X_{t-d} + \dots + \phi_{2H}X_{t-(H-1)d} + \epsilon_{t+s} & Z_t > \text{th} \end{cases}$$
(3)

with  $Z_t$  a threshold variable. How is one to define  $Z_t$ ? Strictly speaking, in SETAR models  $Z_t$  should be one of  $\{X_t, X_{t-d}, X_{t-(m-1)d}\}$ . We can define the threshold variable  $Z_t$  via the *threshold delay*  $\delta$ , such that

$$Z_t = X_{t-\delta d}$$

Using this formulation, you can specify SETAR models with:

<i>R code</i>		
obj <- setar(x, m = , d = , steps = , thDelay = )		

where thDelay stands for the above defined  $\delta$ , and must be an integer number between 0 and m-1.

For greater flexibility, you can also define the threshold variable as an arbitrary linear combination of lagged time series values:

$$Z_t = \beta_1 X_t + \beta_2 X_{t-1} + \ldots + \beta_m X_{t-(m-1)d}$$

In R this is implemented as follows:

<i>R code</i>		
obj <- setar(x, m = , d = ,	, steps = , $mTh$ = )	

where mTh stands for  $\beta$ , and takes the form of a vector of real coefficients of length m.

Finally,  $Z_t$  can be an external variable. This is obtained with the call:

<i>R</i> code		
obj <- setar(x, m = , d = , steps = , thVar = )		

where thVar is the vector containing the threshold variable values.

The threshold variable isn't the only additional parameter governing the SETAR model. One can specify the *low* and *high* regime autoregressive orders L and H. These can be specified with the arguments mL and mH, respectively:

<i>R_ code</i>	
obj <- setar(x, m = , d = , steps = , thDelay = , mL = , mH = )	

If not specified, mL and mH defaults to m.

Another hyper-parameter one can specify is the threshold value th, via the additional argument **th**. If not specified, this is estimated by fitting the model for a grid of different, reasonable values of th, and taking the best fit as the final th estimate.

Note that, conditional on  $\{Z_t \leq th\}$ , the model is linear. So, for a fixed threshold value, the CLS estimation is straightforward.

The summary command for this model returns asymptotic standard errors for the estimated  $\phi$  coefficients, based on the assumption that  $\epsilon_t$  are normally distributed.

#### 3.3 LSTAR models

The LSTAR model can be viewed as a generalization of the above defined SE-TAR model:

$$X_{t+s} = (\phi_1 + \phi_{10}X_t + \phi_{11}X_{t-d} + \dots + \phi_{1L}X_{t-(L-1)d})(1 - G(Z_t, \gamma, \operatorname{th})) + (\phi_2 + \phi_{20}X_t + \phi_{21}X_{t-d} + \dots + \phi_{2H}X_{t-(H-1)d})G(Z_t, \gamma, \operatorname{th}) + \epsilon_{t+s}$$

with G the logistic function, and  $Z_t$  the threshold variable. For  $Z_t$ , L and H specification, the same convention as that of SETAR models is followed. In addition, for LSTAR models one has to specify some starting values for all the parameters to be estimated:  $(\phi, \gamma, \text{th})$ .

Once again, estimation is done using CLS, but here an exact solution does not extist, so a numerical minimization algorithm has to be used. If not provided, starting values for  $(\gamma, \text{th})$  are obtained by rough grid search.

### 3.4 Neural Network models

A neural network model with linear output, D hidden units and activation function g, is represented as:

$$x_{t+s} = \beta_0 + \sum_{j=1}^{D} \beta_j g(\gamma_{0j} + \sum_{i=1}^{m} \gamma_{ij} x_{t-(i-1)d})$$
(4)

For the implementation the **nnet** package is used, so please refer to the **nnet** package documentation for more details.

The only additional argument for specifying this model is the number of hidden units size, which stands for the above defined D:

R code
obj <- nnetTs(x, m = , d = , steps = , size = )

The estimation is done via CLS. No additional summary informations are available for this model.

#### 3.5 Additive Autoregressive models

A non-parametric additive model (a GAM, Generalized Additive Model), of the form:

$$x_{t+s} = \mu + \sum_{i=1}^{m} s_i(x_{t-(i-1)d})$$
(5)

where  $s_i$  are smooth functions represented by penalized cubic regression splines. They are estimated, along with their degree of smoothing, using the mgcv package [8].

No additional parameters are required for this model:

R code	
obj <- aar(x, m = , d = , steps = )	

Some diagnostic plots and summaries are provided for this model, adapted from those produced by mgcv.

### 3.6 Model selection

A common task in time series modelling is *fine tuning* of the hyper-parameters. R is a complete programming language, so the user can easily define his error criterion, fit a set of models and chose the best between them. However, the **tsDyn** package provides some helpful functions for this task.

For SETAR models, there is the selectSETAR function. The time series, the embedding parameters and a vector of values for each provided hyper-parameter is passed to this function. The routine then tries to fit the model for the full grid of hyper-parameter values, and gives as output a list of the best combinations found. So, for example:

	R code					
х	x <- log10(lynx)					
s	selectSETAR(x, m = 3, mL = 1:3, mH = 1:3, thSteps = 5, thDelay = 0:2)					
						output
	thDelay		th	mL	mH	pooled.AIC
1	2	2.	886993	3	2	-40.83722
2	2	2.	886993	3	3	-39.72782
3	2	2.	886993	2	2	-34.36648
4	1	3.	276573	3	2	-34.04683
5	2	2.	886993	2	3	-33.25708
6	2	2.	886993	1	2	-32.42552
7	1	3.	276573	3	3	-32.16499
8	2	2.	886993	1	3	-31.31612
9	1	3.	276573	2	2	-30.36645
10	1	2.	886993	3	2	-28.84706

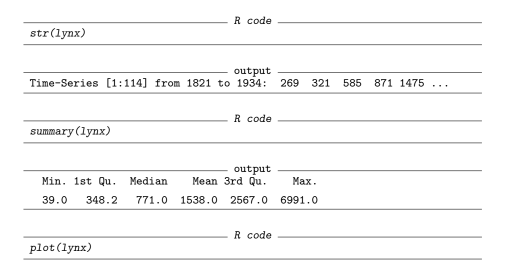
tries to fit  $3 \times 3 \times 3 \times 5$  models, one for each combination of mL,mH,thDelay and th, and returns the best combinations w.r.t. the AIC criterion.

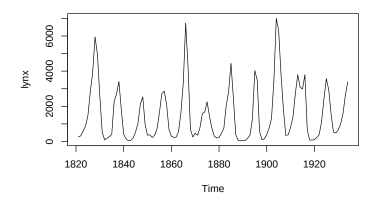
Totally analogous are the selectLSTAR and selectNNET functions, for which we refer to the online documentation.

## 4 Case study

We herein analyse the Canadian lynx data set. This consists of annual records of the numbers of the Canadian lynx trapped in the Mackenzie River district of North-west Canada for the period 1821-1934.

The time series, named lynx, is available in a default R installation, so one can type directly, in an R session:





Here we will roughly follow the analysis in Tong [7].

## 4.1 Explorative analysis

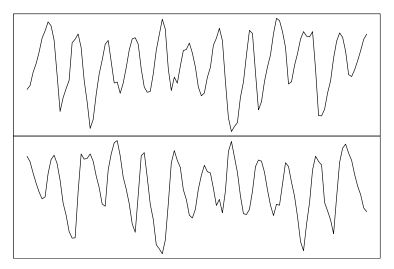
First, we log transform the data:

x <- log10(lynx)

\_\_\_\_\_ R code \_\_\_\_

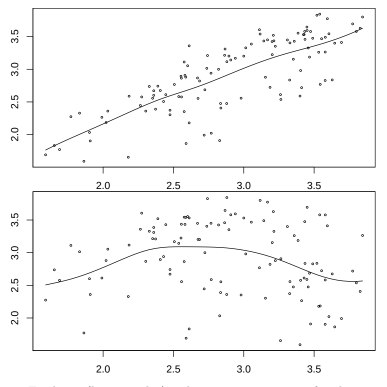
Plot of the time series and time-inverted time series:

```
R code ______
par(mfrow = c(2, 1), mar = c(0, 0, 0, 0))
plot(x, ax = F)
box()
plot(x[length(x):1], type = "l", ax = F)
box()
```



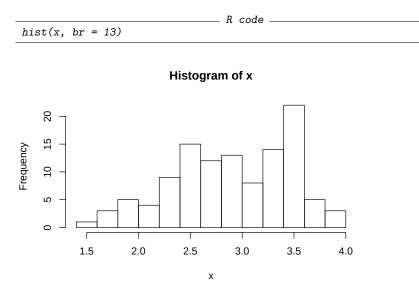
Nonparametric regression function of  $X_t$  versus  $X_{t-1}$  and of  $X_t$  versus  $X_{t-3}$  (kernel estimation):

```
\begin{array}{c} R \ code \\ \hline par(mfrow = c(2, 1), \ mar = c(2, 2, 0, 0)) \\ autopairs(x, lag = 1, type = "regression", GUI = FALSE) \\ autopairs(x, lag = 3, type = "regression", GUI = FALSE) \end{array}
```

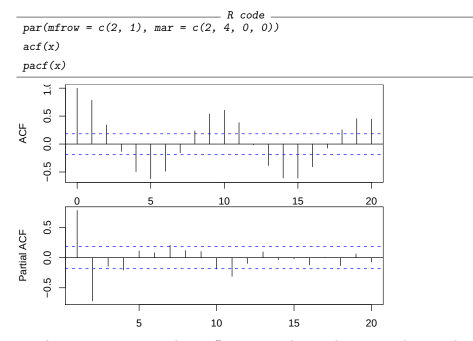


For lag 3 (bottom plot), a linear approximation for the regression function may be questionable.

The marginal histogram of data shows bimodality:

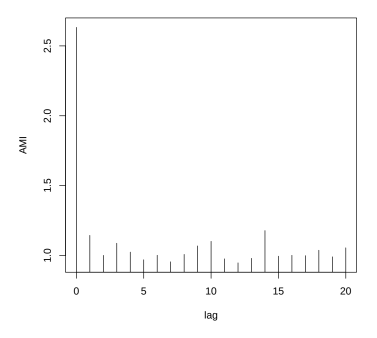


Global and partial autocorrelation:



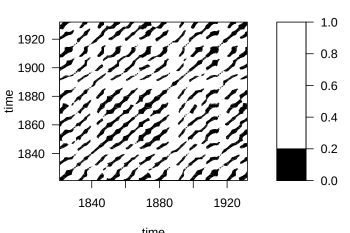
The tseriesChaos package offers some other explorative tools tipycal of nonlinear time series analysis. The Average Mutual Information (see online help for further explanation):

#### **Average Mutual Information**



Recurrence plot (see online help):

R code \_\_\_\_\_\_ R code \_\_\_\_\_\_\_ R code \_\_\_\_\_\_ R code \_\_\_\_\_\_ R code \_\_\_\_\_\_ R code \_\_\_\_\_\_ R



**Recurrence plot** 

time

From this plot, deterministic cycles appears from the embedding-reconstructed underlying dynamics.

Directed lines are a tipycal tool for time series explorations. The lag.plot function in the base stats package does this well:

R code lag.plot(x, lags = 3, layout = c(1, 3))2.0 2.5 3.0 3.5 2.0 2.5 3.0 3.5 4.0 3.5 3.0 2.5 2.0 1.5 0 lag 2 2.5 3.0 lag 3 2.0 3.5

Especially for lag 2, a cycle is again evident. Moreover, the void space in the middle is a typical argument for rejecting the bivariate normality of  $(X_t, X_{t-l})$ .

What follows is the application of still-experimental code for testing the conditional mutual independence and linearity for lags 2 and 3:

					_ R code
de	lta.tes	t(x)			
					_ output
	eps				1
m	0.2792	0.5584	0.8376	1.1168	
2	0.02	0.02	0.08	0.02	
3	0.28	0.04	0.02	0.02	
					_ R code
de	lta.lin	.test(x)			
					_ output
	eps				-
m	0.2792	0.5584	0.8376	1.1168	
2	0.42	0.12	0.60	0.72	
3	0.22	0.22	0.24	0.56	
m2	eps 0.2792 0.42	0.5584	0.8376 0.60	1.1168 0.72	

P-values are reported, labelled with their embedding dimensions m and window values  $\epsilon$ . We reject conditional independence quite easily. There is some trouble instead for deciding to reject or not linearity. See Manzan [5] for a detailed discussion on these tests.

#### 4.2 Model selection

The first model proposed in the literature for these data was an AR(2):

$$X_t = 1.05 + 1.41X_{t-1} - 0.77X_{t-2} + \epsilon_t$$

with  $v(\epsilon_t) = \sigma^2 = 0.04591$ .

This can be estimated with tsDyn using the command:

	_ R code
<pre>mod.ar &lt;- linear(x, m = 2)</pre>	
mod.ar	

Non linear au	itoregressiv		
AR model			
Coefficients	:		
(intercept)	phi.1	phi.2	
1.0576005	1.3842377	-0.7477757	

As an improvement to the AR model, we may consider applying a SETAR(2; 2,2) model with threshold delay  $\delta = 1$ . In R:

```
mod.setar <- setar(x, m = 2, mL = 2, mH = 2, thDelay = 1)
mod.setar</pre>
```

```
_ output _
Non linear autoregressive model
SETAR model (2 regimes)
Coefficients:
Low regime:
   phi1.0
              phi1.1
                          phi1.2
0.5884369 1.2642793 -0.4284292
High regime:
  phi2.0
            phi2.1
                       phi2.2
1.165692 1.599254 -1.011575
Threshold
Variable: Z(t) = + (0) X(t) + (1) X(t-1)
Value: 3.318
Proportion of points in low regime: 69.6%
```

So, the fitted model may be written as:

$$X_{t+1} = \begin{cases} 0.588 + 1.264X_t - 0.428X_{t-1} & X_{t-1} \le 3.318\\ 1.166 + 1.599X_t - 1.012X_{t-1} & X_{t-1} > 3.318 \end{cases}$$

For an automatic comparison, we may fit different linear and nonlinear models and directly compare some measures of their fit:

```
R code ______
mod <- list()
mod[["linear"]] <- linear(x, m = 2)
mod[["setar"]] <- setar(x, m = 2, thDelay = 1)
mod[["lstar"]] <- lstar(x, m = 2, thDelay = 1)</pre>
```

```
output _______
Using maximum autoregressive order for low regime: mL = 2
Using maximum autoregressive order for high regime: mH = 2
Performing grid search for starting values...
Starting values fixed: gamma = 10 , th = 3.345661 ; SSE = 4.338199
Optimization algorithm converged
```

```
______ R code _____
mod[["nnetTs"]] <- nnetTs(x, m = 2, size = 3)
```

```
output ______

# weights: 13

initial value 1123.766049

iter 10 value 7.776995

iter 20 value 5.136185

iter 30 value 4.711834

iter 40 value 4.563658

iter 50 value 4.438878

iter 60 value 4.422297

iter 70 value 4.421858

iter 80 value 4.421757

iter 90 value 4.421303

final value 4.421303

stopped after 100 iterations
```

\_\_\_\_\_ R code \_\_\_\_\_\_ R c

Now the mod object contains a labelled list of fitted nlar models. As an example, we can compare them in term of the AIC and MAPE index:

			_ R code		
sapply(mod	d, AIC)				
			output		
linear	setar	lstar	-		
-121.8737 -	-358.3740 -	356.6509 -3	44.4731 -32	8.0813	
			_ R code		
sapply(mod	d, MAPE)				
			_ output		
linear	setar	lstar	-		
0.06801955	0.05648596	0.05580283	0.05709281	0.05951108	

From this comparison, the SETAR model seems to be the best. More detailed diagnostics can be extracted:

summary(mod[["setar"]])

```
_ R code _
```

```
output -
Non linear autoregressive model
SETAR model (2 regimes)
Coefficients:
Low regime:
    phi1.0
               phi1.1
                          phi1.2
 0.5884369 1.2642793 -0.4284292
High regime:
   phi2.0
            phi2.1
                       phi2.2
 1.165692 1.599254 -1.011575
Threshold
Variable: Z(t) = + (0) X(t) + (1) X(t-1)
Value: 3.318
Proportion of points in low regime: 69.6%
Residuals:
     Min
                       Median
                 1Q
                                     ЗQ
                                              Max
-0.571121 -0.109431 0.017641 0.116468 0.516270
Fit:
residuals variance = 0.03814, AIC = -358, MAPE = 5.649\%
```

```
Estimate Std. Error t value Pr(>|t|)

phi1.0 0.588437 0.143307 4.1061 7.844e-05 ***

phi1.1 1.264279 0.065256 19.3741 < 2.2e-16 ***

phi1.2 -0.428429 0.077487 -5.5291 2.260e-07 ***

phi2.0 1.165692 0.876606 1.3298 0.1863928

phi2.1 1.599254 0.108966 14.6767 < 2.2e-16 ***

phi2.2 -1.011575 0.265011 -3.8171 0.0002255 ***

----

Signif. codes: 0

Threshold

Variable: Z(t) = + (0) X(t) + (1) X(t-1)

Value: 3.318

Proportion of points in low regime: 69.6%
```

More diagnostic plots can be displayed using the command:

```
plot(mod[["setar"]])
```

Coefficient(s):

# \_ R code \_

### 4.3 Out-of-sample forecasting

Fit models on first 104 observations:

```
R code
set.seed(10)
mod.test <- list()
x.train <- window(x, end = 1924)
x.test <- window(x, start = 1925)
mod.test[["linear"]] <- linear(x.train, m = 2)
mod.test[["setar"]] <- setar(x.train, m = 2, thDelay = 1)
mod.test[["lstar"]] <- lstar(x.train, m = 2, thDelay = 1, trace = FALSE,
+ control = list(maxit = 1e+05))
mod.test[["nnet"]] <- nnetTs(x.train, m = 2, size = 3, control = list(maxit = 1e+05))</pre>
```

\_\_\_\_ output \_\_

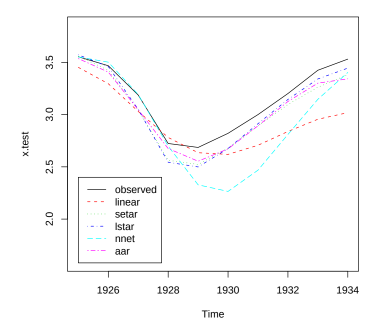
# weights: 13
initial value 1009.386247
iter 10 value 5.610020

```
iter 20 value 5.422071
iter 30 value 5.408916
iter 40 value 5.319344
iter 50 value 5.080814
iter 60 value 4.862847
iter 70 value 4.596344
iter 80 value 4.436638
iter 90 value 4.296703
iter 100 value 4.266607
final value 4.266607
stopped after 100 iterations
```

\_\_\_\_\_\_ R code \_\_\_\_\_\_ mod.test[["aar"]] <- aar(x.train, m = 2)

Compare forecasts with real last 10 observed values:

```
R code
frc.test <- lapply(mod.test, predict, n.ahead = 10)
plot(x.test, ylim = range(x))
for (i in 1:length(frc.test)) lines(frc.test[[i]], lty = i +
+ 1, col = i + 1)
legend(1925, 2.4, lty = 1:(length(frc.test) + 1), col = 1:(length(frc.test) +
+ 1), legend = c("observed", names(frc.test)))
```



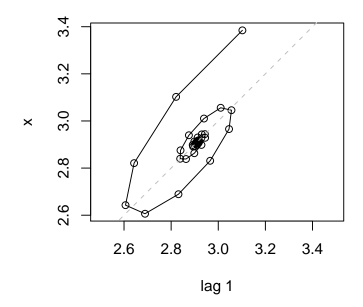
From this visual comparison, the SETAR(2; 2, 2) model seems to be one of the bests.

## 4.4 Inspecting model skeleton

An interesting task can be inspecting the fitted model skeleton.

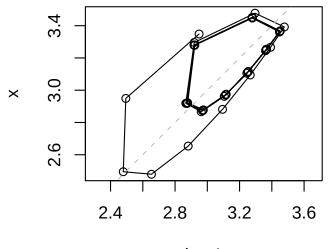
This can be achieved by comparing the forecasting results under each model.

```
R code ______ R code ______
x.new <- predict(mod[["linear"]], n.ahead = 100)
lag.plot(x.new, 1)
```



A fixed point, i.e. the only possible stationary solution with a linear model.

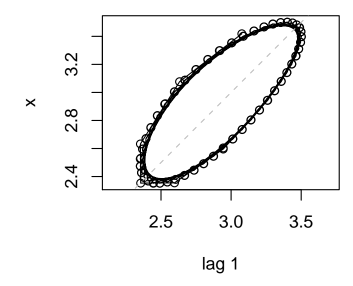
R code	
<pre>x.new &lt;- predict(mod[["setar"]], n.ahead = 100)</pre>	
lag.plot(x.new, 1)	



lag 1

A stable periodic cycle.

x.new <- predict(mod[["nnetTs"]], n.ahead = 100)
lag.plot(x.new, 1)</pre>



Appears to be a quasiperiodic cycle lying on an invariant curve.

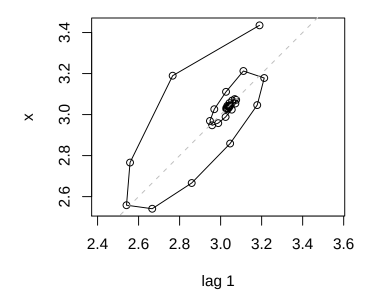
## 5 Sensitivity on initial conditions

In the previous section we observed skeletons with cyclical or limit fixed point behaviour.

Neural networks and SETAR models can explain also different types of attractors. For this data set, Tong [7] showed that particular types of SETAR models can yeld to fixed limit points as well as unstable orbits and *possibly chaotic* systems.

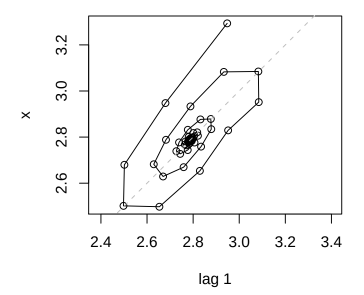
For example, a fixed limit point:

R code mod.point <- setar(x, m = 10, mL = 3, mH = 10, thDelay = 0, th = 3.12) lag.plot(predict(mod.point, n.ahead = 100))



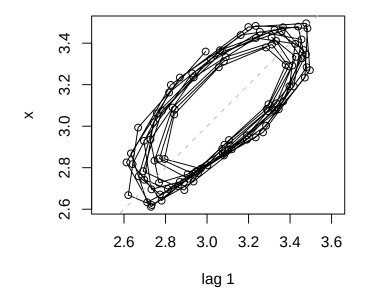
Unstable orbit:

mod.unstable <- setar(x, m = 9, mL = 9, mH = 6, thDelay = 4, + th = 2.61) lag.plot(predict(mod.unstable, n.ahead = 100))

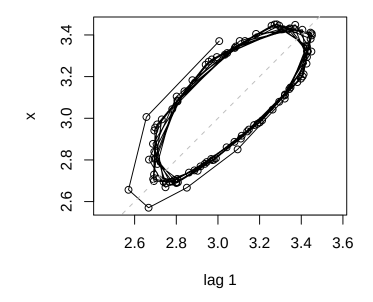


Possibly chaotic systems:

 $\begin{array}{l} R \ code \\ \hline mod.chaos1 <- \ setar(x, \ m = 5, \ mL = 5, \ mH = 3, \ thDelay = 1, \ th = 2.78) \\ lag.plot(predict(mod.chaos1, \ n.ahead = 100)) \end{array}$ 



 $\begin{array}{c} R \ code \\ \hline mod.chaos2 <- \ setar(x, \ m = 5, \ mL = 5, \ mH = 3, \ thDelay = 1, \ th = 2.95) \\ lag.plot(predict(mod.chaos2, \ n.ahead = 100)) \end{array}$ 



For a given fitted model, we can try estimating the maximal Lyapunov exponent with the Kantz algorithm using the lyap\_k function in the tseriesChaos package [6, 4]. This function takes as input an observed time series, so we can proceed as follows:

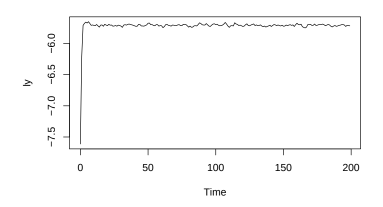
- 1. generate N observations from the model
- 2. add a little observational noise (otherwise the Kantz algorithm will fail)
- 3. apply the lyap\_k function to the generated time series

Follows the R code for analysing the selected SETAR(2; 2,2) model of the previous paragraph and the possibly chaotic SETAR(2; 5,3) just seen above.

R code N <- 1000 x.new <- predict(mod[["setar"]], n.ahead = N) x.new <- x.new + rnorm(N, sd = sd(x.new)/100) ly <- lyap\_k(x.new, m = 2, d = 1, t = 1, k = 2, ref = 750, s = 200, + eps = sd(x.new)/10)

\_ output \_

Finding nearests Keeping 741 reference points Following points plot(ly)



There is no scaling region, so the maximal Lyapunov exponent can assumed to be  $\leq 0$ .

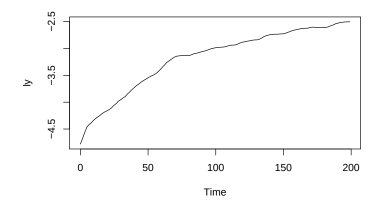
R code x.new <- predict(mod.chaos2, n.ahead = N) x.new <- x.new + rnorm(N, sd = sd(x.new)/100) ly <- lyap\_k(x.new, m = 5, d = 1, t = 1, k = 2, ref = 750, s = 200, + eps = sd(x.new)/10)

\_ output .

Finding nearests Keeping 732 reference points Following points

plot(ly)

\_ R code \_



Here there is a scaling region. The final  $\lambda$  estimate for this time series is the slope of the plotted curve in that region:

lyap(ly, st	art = 6, end = 70)	_ R code		
		output		
(Intercept) lambda		Output		
-4.53873645	0.01979556			

At this point a natural question can be: why not use directly the original time series as input to  $lyap_k$  instead of model-generated observations? The answer here is that we have a too short time series for succesfully applying the Kantz algorithm, so a preliminary modelling for generating more observations is necessary.

## References

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