Some Tools for Identification of Nonlinear Time Series

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Abstract

In classical time series analysis the sample autocorrelation function (SACF) and the sample partial autocorrelation function (SPACF) has gained wide application for structural identification of linear time series models. For non-linear time series these tools are not applicable since they only address variation which can be explained by linear models, and for this reason they may completely fail to detect non-linear dependencies. We suggest generalizations, founded on smoothing techniques, applicable for structural identification of non-linear time series models. A similar generalization of the sample cross correlation function is discussed. Furthermore, a measure of the departure from linearity is suggested. It is shown how bootstrapping can be applied to test for independence and for linearity. The generalizations do not prescribe a particular smoothing technique. In fact, when the smoother are replaced by a linear regression the generalizations of SACF and SPACF reduce to a close approximation of their linear counterparts. For this reason a smooth transition form the linear to the non-linear case can be obtained by varying the bandwidth of a local linear smoother. By adjusting the

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flexibility of the smoother the power of the tests for independence and linearity against specific alternatives can be adjusted. The generalizations allow for graphical presentations, very similar to those used for SACF and SPACF. In this report the generalizations are tested on some simulated data sets and on the Canadian lynx data. The generalizations seem to perform well and the measure of the departure from linearity proves to be an important additional tool.

KEY WORDS: Lagged scatter plot; Non-linear time series; Smoothing; Non-parametric; Independence; Bootstrap.

1 Introduction

The sample autocorrelation function and the sample partial autocorrelation function have gained wide application for structural identification of linear time series models. For non-linear time series these tools are not sufficient because they only address linear dependencies.

During the last couple of decades a number of results on properties of, and estimation and testing in, nonlinear models have been obtained. For an overview (Priestley, 1988; Tong, 1990; Tjøstheim, 1994) can be consulted. However, considerable fewer results have been seen on the problem of structural identification. Tiøstheim and Auestad (1994) have suggested a method based on kernel estimates to select the significant lags in a nonlinear model, and Granger and Lin (1994) used the mutual information coefficient and Kendall's τ as generalizations of the correlation coefficient and Kendall's partial τ as a generalization of the partial correlation coefficient. Chen and Tsay (1993) have considered a best subset modelling procedure and the ACE and BRUTO algorithms, see e.g. (Hastie and Tibshirani, 1990), for identification of non-linear additive ARX models. Recently, Lin and Pourahmadi (1998) have used the BRUTO algorithm to identify the lags needed in a semi-parametric non-linear model. Multivariate adaptive regression splines (Friedman, 1991) was introduced for modelling of non-linear autoregressive time series by Lewis and Stevens (1991). Teräsvirta (1994) suggested a modelling procedure for non-linear autoregressive time series in which a (parametric) smooth threshold auto regressive model is used in case a linear model proves to be inadequate. For the case of non-linear transfer functions Hinich (1979) considered the case where the impulse response function of the transfer function depends linearly on the input process.

In this report we define the new tools LDF (Lag Dependence Function), PLDF (Partial Lag Dependence Function), and NLDF (Non-linear Lag Dependence Function) for structural identification of non-linear time series. The tools can be applied in a way very similar to the sample autocorrelation function and the sample partial autocorrelation function. The tools are based on smoothing techniques, but they are not dependent on any particular smoother, see e.g. (Hastie and Tibshirani, 1990, Chapter 3) for an overview of smoothing techniques. For some smoothers an (almost) continuous transition from the linear to the non-linear case can be obtained by varying the smoothing parameter. Also, smoothers applying optimal selection of the bandwidth may be used; however, see e.g. (Chen and Tsay, 1993) for a discussion of the potential problems in applying criteria such as generalized cross validation to time series data. Under a hypothesis of independence bootstrap confidence intervals (Efron and Tibshirani, 1993) of the lag dependence function are readily calculated, and we propose that these can also be applied for the partial lag dependence function. Furthermore the non-linear lag dependence function can be used to test specific linear hypothesis. A modification of the partial lag dependence function is suggested that seemingly is more appropriate for identifying models for prediction. This also applies when linear models are used. The lag dependence function and the non-linear lag dependence function are readily calculated in that only univariate smoothing are needed, whereas multivariate smoothing or backfitting are required for the application of the remaining tools.

The suggested tools are illustrated both by using simulated linear and non-linear time series models, and by considering the Canadian lynx data (Moran, 1953), which have attained a bench-mark status in time series literature. Using the Canadian lynx data results very similar to those found by Lin and Pourahmadi (1998) are obtained.

In Section 2 the study is motivated by considering a simple deterministic non-linear process for which the sample autocorrelation function is nonsignificant. Section 3 describes the relations between multiple linear regression, correlation, and partial correlation with focus on aspects leading to the generalization. The proposed tools are described in Sections 4, 5, 6, and 7 and bootstrapping is considered in Section 8. Examples of application by considering simulated linear and non-linear processes and the Canadian lynx data (Moran, 1953) are found in Section 9. In Section 10 a generalization of the sample cross correlation function is briefly discussed. Finally, in Section 11 some further remarks are given.

2 Motivation

The sample autocorrelation function (Brockwell and Davis, 1987), commonly used for structural identification in classical time series analysis, measures only the degree of linear dependency. In fact deterministic series exists for which the sample autocorrelation function is almost zero, see also (Granger, 1983). One such example is $x_t = 4x_{t-1}(1 - x_{t-1})$ for which Figure 1 shows 1000 values using $x_1 = 0.8$ and the corresponding sample autocorrelation function SACF together with an approximative 95% confidence interval of the estimates under the hypothesis that the underlying process is i.i.d. Furthermore lagged scatter plots for lag one and two are shown. From the plot of the series and the SACF the deterministic structure is not revealed. However, the lagged scatter plots clearly reveals that the series contains a non-linear dynamic dependency.



Figure 1: The time series (top), SACF (bottom, left), x_t versus x_{t-1} (bottom, middle), and x_t versus x_{t-2} (bottom, right) for 1000 values from the recursion $x_t = 4x_{t-1}(1 - x_{t-1})$.

In practice the series will often be contaminated with noise and it is then

difficult to judge from the lagged scatter plots whether any dependence is present. Smoothing the lagged scatter plots will aid the interpretation but different smoothing parameters may result in quite different estimates. Therefore it is important to separate the variability of the smooth from the underlying dependence.

From Figure 1 it is revealed that, in principle, x_t can be regarded as a function of x_{t-k} for any k > 0, but k = 1 is sufficient, since x_t can be predicted exactly from x_{t-1} alone. This indicates that there may exist a non-linear equivalent to the partial autocorrelation function (Brockwell and Davis, 1987) and reveals that substantial information can be obtained by adjusting for the dependence of lag $1, \ldots, k-1$ when x_t and x_{t-k} are addressed. The sample partial autocorrelation function amounts to a linear adjustment.

3 Preliminaries

Estimates of correlation and partial correlation are closely related to values of the squared degree of determination (R-squared) obtained using linear regression models. The generalizations of the sample autocorrelation function SACF and the sample partial autocorrelation function SPACF are based on similar R-squared values obtained using non-linear models. In this section the relations between multiple linear regression, correlation, and partial correlation are presented.

Consider the multivariate stochastic variable (Y, X_1, \ldots, X_k) . The squared multiple correlation coefficient $\rho_{0(1...k)}^2$ between Y and (X_1, \ldots, X_k) can be written (Kendall and Stuart, 1961, p. 334, Eq. (27.56))

$$\rho_{0(1\dots k)}^2 = \frac{V[Y] - V[Y | X_1, \dots, X_k]}{V[Y]}.$$
(1)

If the variances are estimated using a maximum likelihood estimator, assuming normality, it then follows that an estimate of $\rho_{0(1...k)}^2$ is

$$R_{0(1\dots k)}^2 = \frac{SS_0 - SS_{0(1\dots k)}}{SS_0},\tag{2}$$

where $SS_0 = \sum (y_i - \bar{y})^2$ (where $\bar{y} = \sum y_i/N$) and $SS_{0(1...k)}$ is the sum of squares of the least squares residuals when regressing y_i linearly on

 x_{1i}, \ldots, x_{ki} $(i = 1, \ldots, N)$. $R^2_{0(1...k)}$ is also called the squared degree of determination of the regression and can be interpreted as the relative reduction in variance due to the regressors.

Hence it follows that when regressing y_i linearly on x_{ki} the squared degree of determination $R_{0(k)}^2$ equals the squared estimate of correlation between Y and X_k , and furthermore it follows that $R_{0(k)}^2 = R_{k(0)}^2$.

The partial correlation coefficient $\rho_{(0k)|(1...k-1)}$ between Y and X_k given X_1, \ldots, X_{k-1} measures the extend to which, by using linear models, the variation in Y, which cannot be explained by X_1, \ldots, X_{k-1} , can be explained by X_k . Consequently, the partial correlation coefficient is the correlation between $(Y | X_1, \ldots, X_{k-1})$ and $(X_k | X_1, \ldots, X_{k-1})$, see also (Rao, 1965, p. 270). Using (Whittaker, 1990, p. 140) we obtain

$$\rho_{(0k)|(1...k-1)}^{\rho_{(0k)|(1...k-1)}} = \frac{V[Y \mid X_1, \dots, X_{k-1}] - V[Y \mid X_1, \dots, X_k]}{V[Y \mid X_1, \dots, X_{k-1}]}.$$
(3)

For k = 1 it is readily seen that $\rho_{(0k)|(1...k-1)}^2 = \rho_{0(1)}^2$. If the variances are estimated using the maximum likelihood estimator, assuming normality, it follows that an estimate of $\rho_{(0k)|(1...k-1)}^2$ is

$$R_{(0k)|(1...k-1)}^{2} = \frac{SS_{0(1...k-1)} - SS_{0(1...k)}}{SS_{0(1...k-1)}}.$$
(4)

Besides an estimate of $\rho_{(0k)|(1...k-1)}^2$ this value can also be interpreted as the relative decrease in the variance when including x_{ki} as an additional predictor in the linear regression of y_i on $x_{1i}, \ldots, x_{k-1,i}$. Note that (4) may also be derived from (Ezekiel and Fox, 1959, p. 193).

Interpreting $R_{0(1...k)}^2$, $R_{0(k)}^2$, and $R_{(0k)|(1...k-1)}^2$ as measures of variance reduction, these can be calculated and interpreted for a wider class of models. In the remaining part of this report "~" will be used above values of SS and R^2 obtained from models other than linear models.

4 Lag Dependence

Assume that observations $\{x_1, \ldots, x_N\}$ from a stationary stochastic process $\{X_t\}$ exists. It is readily shown that the estimate of the autocorrelation

function in lag k is approximately equal to the estimate of the correlation coefficient between X_t and X_{t-k} using the observations $\{x_1, \ldots, x_N\}$. Hence, the squared SACF(k) can be approximated by the squared degree of determination when regressing x_t linearly on x_{t-k} , i.e. $R^2_{0(k)}$.

This observation leads to a generalization of SACF(k), based on $\hat{R}^2_{0(k)}$ obtained from a smooth of the k-lagged scatter plot, i.e. a plot of x_t against x_{t-k} . The smooth is an estimate of the conditional mean $f_k(x) = E[X_t | X_{t-k} = x]$. We then define the Lag Dependence Function in lag k, LDF(k), as

$$LDF(k) = \text{sign}\left(\hat{f}_k(b) - \hat{f}_k(a)\right)\sqrt{(\tilde{R}_{0(k)}^2)_+}$$
 (5)

where a and b is the minimum and maximum over the observations and the subscript "+" indicates truncation of negative values. The truncation is necessary to ensure that (5) is defined. However, the truncation will only become active in extreme cases. For some smoothers and lags it becomes active for the series considered in Figure 1.

Due to the sign, when $f_k(\cdot)$ is restricted to be linear, LDF(k) is a good approximation of SACF(k) and, hence, it can be interpreted as a correlation. In the general case LDF(k) can be interpreted as (the signed square-root of) the part of the overall variation in x_t which can be explained by x_{t-k} . Generally, *R*-squared for the non-parametric regression of x_t on x_{t-k} , $\tilde{R}_{0(k)}$ do not equal *R*-squared for the corresponding non-parametric regression of x_{t-k} on x_t , and consequently, unlike SACF(k), the lag dependence function is not an even function. In this report only causal models will be considered and (5) will only be used for k > 0 and by definition LDF(0) will be set equal to one.

5 Strictly Non-Linear Lag Dependence

The lag dependence function described in Section 4 measures both linear and non-linear dependence. If, in the definition of $\tilde{R}^2_{0(k)}$, the sum of squares from a overall mean SS_0 is replaced by the sum of squares from fitting a strait line to the k-lagged scatter plot, a measure of non-linearity is obtained. In this report this will be called the strictly Non-linear Lag Dependence Function in lag k, or NLDF(k).

6 Partial Lag Dependence

For the time series $\{x_1, \ldots, x_N\}$ the sample partial autocorrelation function in lag k, denoted SPACF(k) or $\hat{\phi}_{kk}$, is obtainable as the Yule–Walker estimate of ϕ_{kk} in the AR(k) model

$$X_t = \phi_{k0} + \phi_{k1} X_{t-1} + \ldots + \phi_{kk} X_{t-k} + e_t, \tag{6}$$

where $\{e_t\}$ is i.i.d. with zero mean and constant variance, see also (Brockwell and Davis, 1987, p. 235). An additive, but non-linear, alternative to (6) is

$$X_t = \varphi_{k0} + f_{k1}(X_{t-1}) + \ldots + f_{kk}(X_{t-k}) + e_t.$$
(7)

This model may be fitted using the backfitting algorithm (Hastie and Tibshirani, 1990), see also Section 6.1. The function $f_{kk}(\cdot)$ can be interpreted as a partial dependence function in lag k when the effect of lags $1, \ldots, k-1$ is accounted for. If the functions $f_{kj}(\cdot)$, $(j = 1, \ldots, k)$ are restricted to be linear then $\hat{f}_{kk}(x) = \hat{\phi}_{kk}x$ and the function can be uniquely identified by its slope $\hat{\phi}_{kk}$.

However, since the partial autocorrelation function in lag k is the correlation between $(X_t | X_{t-1}, \ldots, X_{t-(k-1)})$ and $(X_{t-k} | X_{t-1}, \ldots, X_{t-(k-1)})$, the squared SPACF(k) may also be approximated by $R^2_{(0k)|(1...k-1)}$, based on linear autoregressive models of order k-1 and k. Using models of the type (7) SPACF(k) may then be generalized using an R-squared value obtained from a comparison of models (7) of order k-1 and k. This value is denoted $\tilde{R}^2_{(0k)|(1...k-1)}$ and we define the Partial Lag Dependence Function in lag k, PLDF(k), as

$$PLDF(k) = sign\left(\hat{f}_{kk}(b) - \hat{f}_{kk}(a)\right) \sqrt{(\tilde{R}^2_{(0k)|(1...k-1)})_+}.$$
(8)

When (7) is replaced by (6) PLDF(k) is a good approximation of SPACF(k). As for LDF(k), generally, PLDF(k) cannot be interpreted as a correlation. However, PLDF(k) can be interpreted as (the signed square-root of) the relative decrease in one-step prediction variance when lag k is included as a predictor. For k = 1 the model (7) corresponding to k - 1 reduce to an overall mean and the R-squared value in (8) is thus $\tilde{R}^2_{0(1)}$, whereby PLDF(1) = LDF(1) if the same smoother is used for both

functions. It can be noticed that the same relation exists between the partial autocorrelation function and the autocorrelation function. For k = 0the partial lag dependence function is set equal to one.

Except for the sign PLDF(k) may also be based on the completely general autoregressive model

$$x_t = g_k(x_{t-1}, \dots, x_{t-k}) + e_t \tag{9}$$

where $g : \mathcal{R}^k \to \mathcal{R}$. However, the estimation of $g_k(\cdot, \ldots, \cdot)$ without other than an assumption of smoothness is not feasible in practice for k larger than, say, three, see also (Hastie and Tibshirani, 1990). Recently, alternatives to (9) has been considered by Lin and Pourahmadi (1998).

6.1 Fitting the Additive Models

To fit the non-linear additive autoregressive model (7) the backfitting algorithm (Hastie and Tibshirani, 1990) is suggested. However, concurvity (Hastie and Tibshirani, 1990) between the lagged values of the time series may exist and, hence, the estimates may not be uniquely defined. For this reason it is suggested to fit models of increasing order, starting with k = 1 and ending with the highest lag K for which PLDF(k) is to be calculated. In the calculation of the residual sum of squares only residuals corresponding to t = K + 1, ..., N should be used.

For the numerical examples considered in this report local polynomial regression (Cleveland and Devlin, 1988) is used for smoothing. The convergence criterion used is the maximum absolute change in any of the estimates relative to the range of the fitted values. Also, an iteration limit is applied.

For k = 1 the estimation problem reduces to local polynomial regression and hence convergence is guaranteed. If for any k = 2, ..., K convergence is not obtained, or if the residual sum of squares increases compared to the previous lag, we put $\hat{f}_{jk}(\cdot) = 0$, (j = k, ..., K) and $\hat{f}_{kj}(\cdot) = \hat{f}_{k-1,j}(\cdot)$, (j = 1, ..., k - 1). This ensures that convergence is possible for k + 1.

7 Partial *R*-squared for Non-Linear Relations

Basicly, PLDF(k) compares the fit of a model containing lags $1, \ldots, k$ relatively to fit of a model containing lags $1, \ldots, k - 1$. For prediction purposes it may seem more appropriate to compare the reduction in the one-step prediction variance. From the definition of $\tilde{R}^2_{0(1...k)}$ it is seen that

$$\Delta \tilde{R}^{2}_{0(k)} = \tilde{R}^{2}_{0(1...k)} - \tilde{R}^{2}_{0(1...k-1)}$$
$$= \frac{\widetilde{SS}_{0(1...k-1)} - \widetilde{SS}_{0(1...k)}}{SS_{0}}, \qquad (10)$$

is the normalized reduction in the (in-sample) one-step prediction error variance when including lag k as a predictor. The Partial *R*-Squared Function in lag k, PRSF(k), is then defined as

$$PRSF(k) = \text{sign}\left(\hat{f}_{kk}(b) - \hat{f}_{kk}(a)\right) \sqrt{(\Delta \tilde{R}_{0(k)}^2)_+}.$$
 (11)

As for PLDF(k) it is possible, except for the sign, to define PRSF(k)using models of the type (9). For k = 1 the model (7) corresponding to k-1 reduces to an overall mean and, hence, $\Delta \tilde{R}^2_{0(1)} = \tilde{R}^2_{0(1)}$. Consequently, PRSF(1) is equal to PLDF(1) and LDF(1), assuming the same smoother is used in all cases.

8 Testing Hypothesis of Independence or Linearity

Smoothers usually require one or more smoothing parameters to be selected, see e.g. (Hastie and Tibshirani, 1990, Chapter 3). Therefore, in principle, smoothing parameters can be selected to obtain *R*-squared values arbitrarily close to one, also when the underlying process is i.i.d. (assuming no ties are present in the data). For this reason it is important to obtain confidence limits of, e.g., the lag dependence function under the hypothesis that the underlying process is i.i.d. and for a given set of smoothing parameters. Furthermore, it seems applicable to use the strictly non-linear lag dependence function to test for linearity. These aspects are considered in this section.

8.1 Testing for Independence

Under the hypothesis that the time series $\{x_1, \ldots, x_N\}$ is observations from an i.i.d. process the distribution of any of the quantities discussed in the previous sections, except NLDF(k), can be approximated by generating a large number of i.i.d. time series of length N from an estimate of the density function of the process and recalculating the quantities for each of the generated time series. In this report the empirical density function will be used. However, for short time series it may be more appropriate to condition on a parametric form of the density function.

Methods as outlined above are often denoted bootstrap methods and in this context various approaches to the calculation of approximate confidence intervals have been addressed extensively in the literature, see e.g. (Efron and Tibshirani, 1993).

If local polynomial smoothers with degree one or larger is used, then at every point the smoother will be more flexible than a globally linear model. Since LDF(k) and PLDF(k) reduce to close approximations of SACF(k) and SPACF(k), respectively, when linear models are used instead of smoothers, $\pm 2/\sqrt{N}$ will be a lower bound on the 95% confidence interval of LDF(k) and PLDF(k).

Calculation of LDF(k) involves only scatter plot smoothing and, thus, it is faster to calculate than, e.g., PLDF(k). For this reason it is suggested to base a test for independence on LDF(k) for some range k = 1, ..., K. For an i.i.d. process it is obvious that the distribution of LDF(k) will depend on k only due to the fact that k affects the number of points on the k-lagged scatter plot. Hence, when $k \ll N$ the distribution of LDF(k) under the hypothesis of independence is approximately independent of k.

The sign in the definition of LDF(k) is included only to establish an approximate equality with SACF(k) when linear models are used and to include information about the sign of the average value of the slope. When the observations originates from an i.i.d. process LDF(k) will be positive with probability 1/2. Consequently, when the smoother is flexible enough the null-distribution of LDF(k) will be bimodal, since in this case $\tilde{R}^2_{0(k)}$ will be strictly positive. The most efficient way of handling this problem is to base the bootstrap calculations on the absolute value of LDF(k). Hence, an upper confidence limit on |LDF(k)| is to be approximated.

Below the standard, percentile, and BC_a methods, all defined in (Efron and Tibshirani, 1993, Chapters 13 and 14), will be briefly discussed. For the series considered in Figure 1 the LDF(k) were calculated for $k \leq 12$ using a local linear smoother and a nearest neighbour bandwidth of 1/3. The result is shown in Figure 2 together with 95% bootstrap confidence limits

calculated separately for each lag and based on 1000 bootstrap replicates, generated under the hypothesis of independence. The BC_a limit could not be calculated for lags 1 to 4, since all the bootstrap replicates were either smaller or larger than the actual value of |LDF(k)|. Results corresponding to Figure 2 when the true process is standard Gaussian i.i.d. are shown in Figure 3. For practical purposes an equality of the standard and percentile methods are observed (no difference is visible on the plots), whereas the results obtained using the BC_a method is highly dependent on the lag through the value of |LDF(k)|. Hence, the BC_a method cannot be used when the confidence limit is only calculated for one lag and used for the remaining lags as outlined above. The high degree of correspondence between the standard and percentile method indicates that sufficient precision can be obtained using the standard method on fever bootstrap replicates. This is highly related to the approximate normality of |LDF(k)| and it is suggested that this is investigated for each application before a choice between the standard and percentile method is made.



Figure 2: Absolute value of the Lag Dependence Function of the deterministic series presented in Figure 1. The dots indicate the maximum over the 1000 bootstrap replicates. Standard, percentile, and BC_a 95% confidence limits are indicated by lines (BC_a dotted).

The underlying model of the BC_a method assumes that the estimate in question may be biased and that the variance of the estimate depends linearly on an increasing transformation of the true parameter (Efron and Tibshirani, 1993, p. 326-8), and furthermore the estimate is assumed to be normally distributed. The bias and the slope of the line are then estimated from the data. With λ being the fraction of the bootstrap replicates strictly below the original estimate, the bias is $\Phi^{-1}(\lambda)$ (Φ is the cumula-



Figure 3: Absolute value of the Lag Dependence Function of 1000 observations from a standard Gaussian i.i.d. process. The dots indicate the maximum over the 1000 bootstrap replicates. Standard, percentile, and $BC_a 95\%$ confidence limits are indicated by lines (BC_a dotted).

tive standard Gaussian distribution function). This explains why the BC_a limit is non-existing for lags 1-4 of the deterministic series. The slope is estimated by use of the jackknife procedure (Efron and Tibshirani, 1993, p. 186). It seems that, although the underlying model of the BC_a method is a superset of the underlying model of the standard method, the estimation of bias and slope induces some additional variation in the confidence limit obtained. As a consequence it may be advantageous to average the BC_a limits over the lags and use this value instead of the individual values. However, the standard and percentile methods seem to be appropriate for this application and since significant savings of computational effort can be implemented by use of these methods it is suggested that only these are applied on a routine basis.

8.2 Testing for Linearity

Assuming a specific linear model this can be used for simulation and an approximate bootstrap confidence limit for |NLDF(k)| can be obtained given this model. Consequently, the alternative contains both linear and non-linear models. To make the approach sensible the linear model needs to be appropriately selected, i.e. using the standard time series tools of identification, estimation, and validation. Alternatively, the simulations can be performed using autocovariances only and assuming these to be zero after a specific lag. In this case an estimator of autocovariance must

be used that ensures that the autocovariance function used for simulation is non-negative definite, see e.g. (Brockwell and Davis, 1987, p. 27). Note that using this approach on the series considered in Figure 1 will, essentially, result in a test for independence.

Hjellvik and Tjøstheim (1996) consider a similar test for linearity and uses Akaike's information criterion (Brockwell and Davis, 1987) to select an appropriate AR(p)-model under which the bootstrap replicates are generated. In (Theiler, Eubank, Longtin, Galdrikian and Farmer, 1992) a range of alternative linear null hypotheses is considered. Especially, the random sampling in the phase spectrum described in Section 2.4.1 of this reference seems to be a relevant linear null hypothesis.

8.3 Confidence Limit for |PLDF(k)|

In Section 8.1 it is shown how bootstrapping can be used to construct an approximative confidence limit for |LDF(k)|. There is some indication that this limit can be used also for |PLDF(k)| if the same smoother is used for calculation of LDF(k) and $\hat{f}_{k1}(\cdot), \ldots, \hat{f}_{kk}(\cdot)$ (Sections 4 and 6).

For (linear) autoregressive models of order p, with i.i.d. $N(0, \sigma^2)$ errors, and fitted using N observations it holds approximately that the residual sum of squares is distributed as $\sigma^2 \chi^2 (N - p)$ (Brockwell and Davis, 1987, p. 251 and 254). Therefore if the true process is i.i.d. with variance σ^2 the following approximations apply when linear autoregressive models are used

$$SS_0 \sim \sigma^2 \chi^2 (N-1) \tag{12}$$

$$SS_{0(k)} \sim \sigma^2 \chi^2 (N-2) \tag{13}$$

$$SS_{0(1...k-1)} \sim \sigma^2 \chi^2 (N-k)$$
 (14)

$$SS_{0(1...k)} \sim \sigma^2 \chi^2 (N-k-1)$$
 (15)

For $N \gg k$ the distribution of all four sums of squares are approximately equal.

For locally weighted regression Cleveland and Devlin (1988) stated that the distribution of the residual sum of squares can be approximated by a constant multiplied by a χ^2 variable, see also (Hastie and Tibshirani, 1990, Section 3.9). Furthermore, for generalized additive models Hastie and Tibshirani (1990, Section 8.1) uses a χ^2 distribution with degrees of freedom equal to the number of observations minus a quantity depending on the flexibility of the smoothers used.

For these reasons we conjecture that when $N \gg k$ and when the same smoother is used for LDF(k) and PLDF(k), as outlined in the beginning of this section, then the sum of squares SS_0 , $\widetilde{SS}_{0(k)}$, $\widetilde{SS}_{0(1...k-1)}$, and $\widetilde{SS}_{0(1...k)}$ will follow approximately the same distribution.

This conjecture leads to approximate equality of means and variances of the sums of squares. Since for both LDF(k) and PLDF(k) the compared models differ by an additive term, estimated by the same smoother in both cases, we also conjecture that for an i.i.d. process.

$$Cor[\widetilde{SS}_{0(k)}, SS_0] \approx Cor[\widetilde{SS}_{0(1\dots k)}, \widetilde{SS}_{0(1\dots k-1)}].$$
(16)

Using linearizations about the mean of the sums of squares it then follows from the approximate equality of means that

$$E[|LDF(k)|] \approx E[|PLDF(k)|], \tag{17}$$

and from both conjectures that

$$V[|LDF(k)|] \approx V[|PLDF(k)|].$$
(18)

Eqs. (17) and (18) tell us that the approximate i.i.d. confidence limit obtained for |LDF(k)| can be used also as an approximate limit for |PLDF(k)|. In Section 9 (Canadian lynx data) an example of the quality of the approximation is given, and the mentioned arguments seems to be confirmed by the bootstrap limits obtained in that example.

9 Examples

9.1 Linear processes

Below it is briefly illustrated how LDF, PLDF, and PRSF behaves compared to SACF and SPACF in case of simple linear processes. The AR(2) process

$$X_t = 1.13X_{t-1} - 0.64X_{t-2} + e_t \tag{19}$$

and the MA(2) process

$$X_t = e_t + 0.6983e_{t-1} + 0.5247e_{t-2} \tag{20}$$

are considered, where in both cases $\{e_t\}$ is i.i.d. N(0, 1).

Figures 4 and 5 contain plots based on 100 simulated values from (19) and (20), respectively (the default random number generator of S-PLUS version 3.4 for HP-UX were used). Each figure show SACF and SPACF. The remaining plots are LDF, PLDF, and PRSF for local linear smoothers using a nearest neighbour bandwidth of 1.00 (2nd row), 0.50 (3rd row), and 0.1 (bottom row). 95% confidence intervals are indicated by dotted lines. The confidence intervals obtained for LDF are included on the plots of PLDF.

For the calculation of PLDF and PRSF a convergence criterion (see Section 6.1) of 0.01 and an iteration limit of 20 is used. Standard bootstrap intervals are calculated for LDF under the i.i.d. hypothesis using 200 replicates. For LDF the agreement with SACF is large for nearest neighbour bandwidths 1.0 and 0.5. As expected, the range of the confidence interval increases with decreasing bandwidth, and, using the smallest bandwidth, it is almost not possible to reject the i.i.d. hypothesis.

When a nearest neighbour bandwidth of 1.0 is used PLDF agrees well with SPACF for the lower half of the lags, whereas PLDF is exactly zero for most of the larger half of the lags. Similar comments apply for nearest neighbour bandwidths 0.5 and 0.1. This is due to the function estimates being set equal to zero when the iteration limit is exceeded. The alternative to PLDF, namely PRSF, behaves similar, but for lags two, or larger, the absolute value of PRSF is smaller than for the absolute value of PLDF. This is because PRSF addresses decreases in (normalized) prediction error variances, whereas PLDF compares the fit relative to the fit when the last lag is excluded.



Figure 4: Plots of autocorrelation functions and their generalizations for 100 observations from the AR(2) process (19).



Figure 5: Plots of autocorrelation functions and their generalizations for 100 observations from the MA(2) process (20).

9.2 Non-linear processes

Three non-linear processes are addressed, namely (i) the non-linear autoregressive process (NLAR(1))

$$X_t = \frac{1}{1 + \exp(-5X_{t-1} + 2.5)} + e_t,$$
(21)

where $\{e_t\}$ i.i.d. $N(0, 0.1^2)$, and (ii) the non-linear moving average process (NLMA(1))

$$X_t = e_t + 2\cos(e_{t-1}),$$
(22)

where $\{e_t\}$ i.i.d. N(0, 1) and (iii) the non-linear and deterministic process described in Section 2, called DNLAR(1) in the following. For all three cases 1000 observations are generated. The starting value for NLAR(1) is set to 0.5 and for DNLAR(1) it is set to 0.8. Plots of the series NLAR(1)and NLMA(1) are shown in Figure 6. The plot of DNLAR(1) is shown in Figure 1.



Figure 6: Plots of the series NLAR(1) (top) and NLMA(1) bottom.

For the calculation of LDF, PLDF, and NLDF a local linear smoother with a nearest neighbour bandwidth of 0.5 is used. Actually, lagged scatter plots indicate that a local quadratic smoother should be applied, at least for NLMA(1) and DNLAR(1), but to avoid a perfect fit for the deterministic series a local linear smoother is used. Confidence intervals are constructed using standard normal intervals, since normal QQ-plots of the absolute values of the 200 bootstrap replicates showed this to be appropriate. The confidence interval obtained for LDF is included on the plots of PLDF. Figure 8 shows SACF, SPACF, LDF, and PLDF for the three series. For NLMA(1) and DNLAR(1) the linear tools, SACF and SPACF, indicate independence and LDF shows that lag dependence is present. From these observations it can be concluded that NLMA(1) and DNLAR(1)are nonlinear processes. From the plots of LDF and PLDF it cannot be inferred whether NLMA(1) is of the autoregressive or of the moving average type. For DNLAR(1) the autoregressive property is more clear since PLDF drops to exactly zero after lag two. In case of DNLAR(1)a more flexible smoother will result in values of LDF being significantly different from zero for lags larger than one independent of the flexibility of the smoother used. This is an indication of DNLAR(1) being of the autoregressive type and NLMA(1) being of the moving average type.

For NLAR(1) the linear tools indicate that the observations come from an AR(1) process. This is not seriously contradicted by LDF or PLDF, although LDF decline somewhat slower to zero than SACF. To investigate if the underlying process is linear a Gaussian AR(1) model is fitted to the data and this model is used as the hypothesis under which 200 (parametric) bootstrap replicates of NLDF are generated. Figure 7 shows NLDF and a 95% standard normal interval, constructed under the hypothesis mentioned above. A normal QQ-plot show that the absolute values of the bootstrap replicates are approximately Gaussian. From the plot it is concluded that the underlying process is not the estimated AR(1)-model, and based on PLDF it is thus concluded that the observations originate from a nonlinear process of AR(1) type.



Figure 7: NLDF for NLAR(1), including a 95% confidence interval under the assumption of an AR(1) process (dotted).



Figure 8: SACF, SPACF, LDF, and PLDF for series NLMA(1), DNLAR(1), and NLAR(1) (columns, left to right).

9.3 Canadian lynx data

Recently, Lin and Pourahmadi (1998) analyzed the Canadian lynx data (Moran, 1953) using non-parametric methods quite similar to the methods presented in this report. The data is included in the software S-PLUS (version 3.4 for HP-UX) and described in (Tong, 1990, Section 7.2). In this report a thorough analysis of the data will not be presented, but the data will be used to illustrate how the methods suggested can be applied. As in (Lin and Pourahmadi, 1998) the data is \log_{10} -transformed prior to the analysis.

For the transformed data LDF, PLDF, and NLDF are computed using a local quadratic smoother and nearest neighbour bandwidths of 0.5 and 1. For LDF 200 bootstrap replicates are generated under the i.i.d. hypothesis and QQ-plots indicate that standard normal intervals are appropriate. The same apply for NLDF with the exception that the bootstrap replicates are generated under the hypothesis that the AR(2) model of Moran (1953), also described by Lin and Pourahmadi (1998), is true. Confidence intervals are computed also for PLDF for the nearest neighbour bandwidth of 1.0. The intervals are based one hundred bootstrap replicates of PLDF generated under the i.i.d. hypothesis. QQ-plots indicate that the percentile method should be applied to the absolute values of PLDF.

In Figure 9 plots of LDF, NLDF, and PLDF are shown. Dotted lines indicates 95% confidence intervals under the i.i.d. hypothesis (LDF) and under the AR(2) model of Moran (1953) (NLDF). The intervals obtained for LDF are also shown on the plots of PLDF. Furthermore, for the nearest neighbour bandwidth of 1.0, a 95% confidence interval for white noise is included on the plot of PLDF (solid lines).

From the plots of LDF it is clearly revealed that the process is not i.i.d. The plots of NLDF for a nearest neighbour bandwidth of 0.5 show hardly any significant values, but when a nearest neighbour bandwidth of 1.0 is used lags three and four show weak significance. This indicates that a small departure from linearity is present in the data. Finally, the plots of PLDFclearly illustrate that lag one and two are the most important lags and that other lags are, practically, non-significant. In conclusion, an appropriate model seems to be a non-linear autoregressive model containing lag one and two, i.e. a model of the type (7) with k = 2.

Estimation in this model using local quadratic smoothers and a nearest

neighbour bandwidth of 1.0 yields the results shown in Figure 10. The response for lag one seems to be nearly linear. This aspect should be further investigated. The results agree well with the results of Lin and Pourahmadi (1998).



Figure 9: Canadian lynx data (\log_{10} -transformed). Plots of *LDF*, *NLDF*, and *PLDF* using local quadratic smoothers and nearest neighbour bandwidths 0.5 (top row) and 1.0 (bottom row).

10 Lagged Cross Dependence

Given two time series $\{x_1, \ldots, x_N\}$ and $\{y_1, \ldots, y_N\}$ the Sample Cross Correlation Function between processes $\{X_t\}$ and $\{Y_t\}$ in lag k (SCCF_{xy}(k)) is an estimate of the correlation between X_{t-k} and Y_t . It is possible to generalized this in a way similar to the way LDF is constructed. Like SCCF this generalization will be sensible to autocorrelation, or lag dependence, in $\{X_t\}$ in general. For SCCF this problem is (approximately) solved by prewhitening (Brockwell and Davis, 1987, p. 402). However, prewhitening is very dependent on the assumption of linearity, in that it relies on the impulse response function from the noise being independent on the level. For this reason, in the non-linear case, it is not possible to use prewhitening and the appropriateness of the generalization of SCCF depend on $\{X_t\}$ being i.i.d.



Figure 10: Non-linear additive autoregressive model for the \log_{10} -transformed Canadian lynx data $(\hat{f}_{21}(\cdot) \text{ solid}, \hat{f}_{22}(\cdot) \text{ dotted})$. The estimate of the constant term is 2.76 and the MSE of the residuals is 0.0414.

11 Final Remarks

The generalizations of the sample correlation functions reduce to their linear counterpart when the smoothers are replaced by linear models. Hence, if a local linear smoother is applied an almost continuous transition from linear to non-linear measures of dependence is obtainable via the bandwidth of the smoother.

In the test for an i.i.d. process the alternative contain both linear and non-linear models. The degree of departure from linearity that can be detected by the test can be adjusted by selection of the flexibility of the smoother. If, e.g., a local quadratic smoother with a nearest neighbour bandwidth of 100% is applied, the test will have large power against an "almost quadratic" alternative. If, on the other hand, a more flexible smoother is used, the power against an "almost quadratic" alternative will be lower.

Optimal bandwidth selection is not addressed in this report. However, the methods can still be applied in this case, but the power against specific alternatives cannot be adjusted.

The estimation in the non-linear additive autoregressive model, which is used in the generalization of the sample partial autocorrelation function, breaks down in case of concurvity. Hence, the lags enter the model sequentially and in case of non-convergence of the backfitting iterations the last lag is, essentially, excluded from all subsequent models, c.f. Section 6.1. Furthermore, slow convergence have been observed in some cases. Hence, the particular procedure is probably not optimal. The procedure may be replaced by a modified BRUTO algorithm (Hastie and Tibshirani, 1990, p. 262-3) in which the smoothing parameters are fixed but where the null fit is still a possibility. However, due to the autoregression then e.g. local regression smoothers will be non-linear and the approximation to the degrees of freedom used in the BRUTO algorithm may not be applicable. Inspired by the work of Ye (1998) the degrees of freedom may be defined as the sum of the sensitivities of the fitted values with respect to the observations. Using this definition the degrees of freedom will equal the trace of the smoother matrix plus a quantity involving partial derivatives of elements of the smoother matrix with respect to the observations.

If the conditional mean of the series can be modelled the methods described in this report can be applied to the series of squared residuals and the conditional variance can, possibly, be addressed in this way. This approach is quite similar to the approach by Tjøstheim and Auestad (1994, Section 5).

The first author has created an experimental S-PLUS library implementing most of the methods described in this report, using local polynomial smoothers. The software has been used for most of the numerical calculations in this report. A copy of the software can be obtained by contacting the first author.

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