Reliability diagrams for non-parametric density forecasts of continuous variables: Accounting for serial correlation

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Reliability is seen as a primary requirement when verifying probabilistic forecasts, since a lack of reliability would introduce a systematic bias in subsequent decision-making. Reliability diagrams comprise popular and practical diagnostic tools for the reliability evaluation of density forecasts of continuous variables. Such diagrams relate to the assessment of the unconditional calibration of probabilistic forecasts. A reason for their appeal is that deviations from perfect reliability can be visually assessed based on deviations from the diagonal. Deviations from the diagonal may, however, be caused by both sampling effects and serial correlation in the forecast-verification pairs. We build on a recent proposal, consisting of associating reliability diagrams with consistency bars that would reflect the deviations from the diagonal that are potentially observable even if density forecasts are perfectly reliable. Our consistency bars, however, reflect potential deviations originating from the combined effects of limited counting statistics and serial correlation in the forecast-verification pairs. They are generated based on an original surrogate consistency resampling method. Its ability to provide consistency bars with a significantly better coverage against the independent and identically distributed (i.i.d.) resampling alternative is shown from simulations. Finally, a practical example of the reliability assessment of non-parametric density forecasts of short-term wind-power generation is given. Copyright © 2010 Royal Meteorological Society

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1. Introduction

Over the past few decades, one of the major breakthroughs in forecasting meteorological variables for applications such as weather derivatives and renewable energy generation comes from the transition from point to probabilistic forecasting (Gneiting, 2008a). One has to acknowledge the significant contribution of some of the leading meteorological centres, e.g. the National Centers for Environmental Prediction (NCEP) and European Centre for Medium-Range Weather Forecasts (ECMWF) in developing ensemble forecasting systems, as well as a probabilistic view of meteorological forecasting. For a detailed overview of ensemble forecasting and the underlying probabilistic forecasting philosophy developed in the meteorological community, the reader is referred to Leutbecher and Palmer (2008) and references therein. From a decision-making perspective, it has been
shown that pricing weather derivatives based on density forecasts would bring significant benefits (Taylor and Buizza, 2006). In parallel for the example case of renewable energy, the production of which is a direct function of meteorological conditions, it is argued that optimal management and trading of generated energy should be based on probabilistic forecasts (Pinson et al., 2007a). The transition from point to probabilistic forecasts is not only observed in the meteorological literature; probabilistic forecasts are also becoming customary products in economics and finance (Abramson and Clemen, 1995; Tay and Wallis, 2000; Timmermann, 2000) or more generally in management sciences. When considering continuous variables such as wind speed, the most complete information about the expected realization for a given lead time takes the form of a density forecast (equivalently referred to as predictive distribution), giving the probability density function of the corresponding random variable. Having a broader view of decision-making for real-world problems, Gneiting (2008b) argues that for a large class of cost functions of forecast users, optimal decisions relate directly to given quantiles of predictive densities of the variable of interest.

In parallel to these developments towards probabilistic forecasting and subsequent optimal decision-making, the issue of probabilistic forecast verification has attracted significant attention. For a recent overview of verification methods for probabilistic forecasts of categorical and continuous variables in atmospheric sciences, see Jolliffe and Stephenson (2003). A primary requirement for probabilistic forecasts relates to their calibration (equivalently referred to as their reliability), which corresponds to their probabilistic correctness. We hereby follow the paradigm introduced by Gneiting et al. (2007), i.e. based on maximizing the sharpness of predictive distributions subject to calibration. Note that this framework for probabilistic forecast evaluation is different from that based on testing for correct conditional coverage of density forecasts, as described in Christoffersen (1998) and Diebold et al. (1998), where focus is given to one-step-ahead forecast only, in a time-series framework. The calibration requirement mentioned above calls for a thorough reliability assessment prior to proceeding with sharpness. Even if considering the use of proper scoring rules (see Bröcker and Smith, 2007b, among others), calculating scores alone does not permit one to see whether better score values come from higher reliability or increased sharpness. A decomposition of these proper scores into their reliability and sharpness components should then be performed. A reliability assessment is necessary in order to make sure that systematic bias is not introduced in further decision-making.

A popular and straightforward way of assessing the calibration of probabilistic forecasts is via the use of reliability diagrams (Atger, 1998, 2004). The reliability diagrams we consider here for the reliability assessment of non-parametric density forecasts of continuous variables consist of an equivalent cumulative version of the popular probability integral transform (PIT) histograms, otherwise called Talagrand diagrams, used for the verification of ensemble forecasts (Hamill, 2001). An extensive presentation of reliability diagrams for density forecasts of continuous variables (and equivalently for ensembles) is given in section 2.

Recently, for the case of probability forecasts of binary events, Bröcker and Smith (2007a) have explained how reliability diagrams may be misinterpreted, since even for perfectly reliable probabilistic forecasts there will always be deviations from the diagonal originating from sampling effects. They have proposed an elegant framework permitting one to easily integrate information about the impact of sampling effects directly in reliability diagrams. While their point is highly relevant, it is argued here that it is not only counting statistics but also serial correlation in forecast-verification pairs that affects the interpretation of reliability diagrams produced from finite-size datasets. Somehow, the methodology proposed by Bröcker and Smith (2007a) is based on the fallacy that for reliable forecasts the random variables with realizations given by the probability integral transforms should be independent and identically distributed (i.i.d.), U[0, 1]. It is true that, by definition, a necessary condition for density forecasts to be reliable is that such random variables are distributed as U[0, 1]. It is not a necessary condition, however, for successive realizations to be independent. In the case of reliability diagrams for probabilistic forecasts of binary events, it might be acceptable to assume independence, as forecast-verification pairs corresponding to a given probability class may be sparsely and randomly distributed over the verification period. It cannot be the case when assessing the reliability of probabilistic forecasts of multicategorical variables or density forecasts of continuous variables. This will be illustrated by a simple practical example based on climatology density forecasts in section 3. The role of serial correlation and sampling effects has already been hinted at by Pinson et al. (2007b) for the case of the verification of wind-power density forecasts, or by Hamill (2001) when considering the verification of ensemble forecasts of meteorological variables. Section 4 introduces a methodology to provide consistency bars in the spirit of Bröcker and Smith (2007a), but accounting for serial correlation effects, using an original surrogate consistency resampling method. Simulations in section 5 show how serial correlation adds to the sampling effect, and demonstrate the pitfall stemming from inference of the confidence one may have in reliability diagrams based on an i.i.d. assumption for probability integral transforms of reliable density forecasts. It also evaluates the validity and accuracy of the methodology introduced. The calibration assessment of short-term forecasts of wind-power generation serves as an illustrative application in section 6. Concluding remarks end the article in section 7.

2. Reliability diagrams for non-parametric density forecasts of continuous variables

Using t as the time index, we denote the stochastic process of interest by \( \{Y_t \} \) and the time series of observed realizations by \( \{y_t \} \). Each random variable \( Y_t \) can take values in \( \mathcal{C} \subset \mathbb{R} \), with for instance \( \mathcal{C} = \mathbb{R}^+ \) for the case of wind speed. For simplicity, it is assumed that random variables and corresponding realizations are equally spaced in time. Also, it is considered that focus is on this stochastic process at a single location only, as reliability assessment of probabilistic forecasts for spatial fields would also be impacted by spatial sampling and correlation effects that are not treated here. Discussions on these spatial effects can be found in Wilks (1995) and Jolliffe and Stephenson (2003).

We denote the density forecast for the stochastic process of interest issued at time t for lead time t + k by \( f_{k,t}(y) \), i.e. based on maximizing the sharpness of predictive distributions subject to calibration.
and the related cumulative distribution function by $F_{t+k|t}(y)$. $\hat{F}_{t+k|t}(y)$ is a forecast of the probability distribution function of $Y_{t+k}$ given the information set available at time $t$, which may be derived from ensemble forecasts or from some non-parametric statistical forecasting techniques, e.g. quantile regression. In a general manner, if no assumption is made about the shape of predictive distributions, a non-parametric density forecast $\hat{f}_{t+k|t}(y)$ can be summarized by a set of $m$ quantile forecasts:

$$\hat{f}_{t+k|t}(y) = \left\{ \hat{q}_{i|t+k}^{(a)} \right\} \left| 0 \leq a_1 < \ldots < a_i < \ldots < a_m \leq 1 \right\},$$

(1)

that is, with chosen nominal proportions $a_i$ spread on the unit interval. In turn, a quantile forecast $\hat{q}_{i|t+k}^{(a)}$ with nominal proportion $a_i$ is defined by

$$P_{\hat{F}_{t+k|t}} \left[ Y_{t+k} < \hat{q}_{i|t+k}^{(a)} \right] = a_i, \quad a_i \in [0,1].$$

(2)

Both forecasts and observations are available for a limited time period $T \subset \mathbb{N}^+$ used for forecast verification. Note that for ensemble forecasts with exchangeable members, non-parametric density forecasts can also be defined as in (1). In such cases, by ordering the $j$ ensemble members in ascending order, the $j$th member gives the quantile with nominal proportion $a_j = j/(J + 1)$. The (continuous) density forecast $\hat{f}_{t+k|t}(y)$ can then be built by interpolation within the set of quantiles.

For a given forecast horizon $k$, the core concept behind the use of reliability diagrams for evaluating density forecasts of continuous variables is that the series of predictive densities $\{\hat{F}_{t+k|t}(y)\}$ are reliable if and only if the random variable $Z_{t+k} = \hat{F}_{t+k|t}(Y_{t+k})$ is distributed U[0,1]. In practice this is performed by studying the realizations of $Z_{t+k}$ that are given by the sequence $\{z_{t+k}\}$ of probability integral transforms, with $Z_{t+k} = F_{\hat{F}_{t+k|t}}(Y_{t+k})$.

Such a definition of reliability actually corresponds to an unconditional calibration of the density forecasts, since no distinction is made between different time points in the dataset, or other conditions that may affect density forecast reliability. This contrasts with the idea of conditional calibration introduced by Christoffersen (1998) and Diebold, et al. (1998) for the case of density forecasts in a time-series context. It also contrasts with the idea of conditional reliability assessment presented by Pinson, et al. (2007b) and Bröcker (2009). Both Pinson, et al. (2007b) and Bröcker (2009) base their argument on the fact that for the case of nonlinear processes the reliability of probabilistic forecasts may be influenced by a set of external factors, or may even simply vary as a function of the forecasts themselves. We refer to as unconditional calibration thus corresponds to the overall probabilistic bias of the density forecasts as discussed by Murphy (1993) and Taylor (1999), for instance, or to what Gneiting, et al. (2007) refer to as probabilistic calibration.

A consequence of the fact that reliability diagrams relate to an unconditional calibration assessment is that one cannot assume that the sequence of probability integral transforms $\{z_{t+k}\}$ for a sequence of reliable forecast densities and corresponding verifications is i.i.d. This would be the case only if, for each given time of the evaluation set, density forecasts were equal to the true conditional densities of the stochastic data generating process (Diebold, et al., 1998; Gneiting, et al., 2007).

In practice, since non-parametric density forecasts as defined by (1) consist of a collection of quantile forecasts for which the nominal proportions are known, evaluating the reliability of density forecasts is achieved by verifying the reliability of each individual quantile forecast. Let us introduce in a first stage the indicator variable $\xi_{t+k}^{(a_i)}$. Given a quantile forecast $\hat{q}_{i|t+k}^{(a_i)}$ issued at time $t$ for lead time $t+k$, and the verification $Y_{t+k}$, $\xi_{t+k}^{(a_i)}$ is given by

$$\xi_{t+k}^{(a_i)} = \begin{cases} 1 & \text{if } Y_{t+k} < \hat{q}_{i|t+k}^{(a_i)} \leq \hat{q}_{i+1|t+k}^{(a_i)} \ , \\ 1 & \text{if } Y_{t+k} < \hat{q}_{i|t+k}^{(a_i)}, \\ 0 & \text{otherwise}. \end{cases}$$

(3)

It can be appealing to project into the standard Gaussian domain by using an inverse normal transformation, i.e. one based on the inverse probit function $\Phi^{-1}$,

$$\Phi^{-1} : p \rightarrow \sqrt{2}\text{erf}^{-1}(2p - 1),$$

(4)

with erf$^{-1}$ the inverse error function. The resulting random variables and corresponding realizations are denoted $\tilde{Z}_{t+k}$ and $\tilde{z}_{t+k}$, respectively, with simply $\tilde{Z}_{t+k} = \Phi^{-1}(Z_{t+k})$ and $\tilde{z}_{t+k} = \Phi^{-1}(z_{t+k})$. Considering $\{\tilde{z}_{t+k}\}$ instead of $\{z_{t+k}\}$ for assessing the calibration of density forecasts has some advantages discussed in the literature (Berkowitz, 2001) for hypothesis testing or for studying potential serial correlation. In the present case, it will permit us to apply classical tools from linear time-series analysis. Consequently, the indicator variable introduced in (3) can be equivalently defined as

$$\xi_{t+k}^{(a_i)} = 1(\tilde{z}_{t+k} < \tilde{q}_{i|t+k}^{(a_i)}),$$

(5)

with $\tilde{q}_{i|t+k}^{(a_i)}$ the quantile with proportion $a_i$ of a standard Gaussian distribution. In the following, we will equivalently consider the possibility of working with the indicator variable definitions of (3) and (5) in order to assess the reliability of density forecasts.

The time series $\{\xi_{t+k}^{(a_i)}\}$ is a binary sequence that corresponds to the series of hits (if the verification $Y_{t+k}$ lies below the quantile forecast) and misses (if otherwise) over the evaluation set. It is by studying $\{\xi_{t+k}^{(a_i)}\}$ that one can assess the reliability of a time series of quantile forecasts. Indeed, an estimate $\hat{a}_{t+k}$ of the actual proportion $a_{t+k} = \mathbb{E}[\xi_{t+k}^{(a_i)}]$, for a given horizon $k$, is obtained by calculating the mean of the $\{\xi_{t+k}^{(a_i)}\}$ time series over the test set

$$\hat{a}_{t+k} = \frac{1}{N} \sum_{t=1}^{N} \xi_{t+k}^{(a_i)} = \frac{n_{t+k|0}}{n_{t+k|0} + n_{t+k|1}},$$

(6)

where $N$ is the number of time indices in $T$, and $n_{t+k|0}$ and $n_{t+k|1}$ correspond to the sum of hits and misses, respectively. They are calculated with

$$n_{t+k|1} = \#(\xi_{t+k}^{(a_i)} = 1) = \sum_{t=1}^{N} \xi_{t+k}^{(a_i)},$$

(7)

$$n_{t+k|0} = \#(\xi_{t+k}^{(a_i)} = 0) = N - n_{t+k|1},$$

(8)

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This measure of observed proportion serves as a basis for drawing reliability diagrams for density forecasts of continuous variables, which give the observed proportions \( \hat{a}_{i,j} \) of the quantile forecasts against the nominal ones. They therefore consist of quantile–quantile (Q–Q) plots, which are a classical diagnostic tool in the statistical literature. A particular feature of this definition of reliability diagrams is that each quantile is evaluated individually, which will allow us to define a consistency bar for each nominal proportion independently of the other quantiles. Our consistency bars are then pointwise consistency bars. Also, note that the reliability diagrams we consider here are for density forecasts of continuous variables and thus somewhat different from those considered for probabilistic forecasting of binary/categorical variables (Bröcker and Smith, 2007a). The argument developed in the present article regarding the fact that serial correlation effects should be accounted for is still valid, however. In parallel, the methodology described below can be straightforwardly applied for the case of the verification of ensemble forecasts, owing to the definition employed for non-parametric density forecasts.

When visually inspecting the calibration from reliability diagrams, a common intuitive thought is that the closer the observed proportions to the diagonal the better. This is because asymptotically, as the number of forecast-verification pairs tends towards infinity, one wishes that

\[
\lim_{N \to \infty} \hat{a}_{i,j} = \alpha_i, \quad \forall i,
\]

implying that observed proportions asymptotically equal nominal ones. In practice, however, evaluation sets consisting of forecast-verification pairs are of finite (and often quite limited) size, and it is not expected that observed proportions lie exactly along the diagonal, even if the density forecasts are perfectly reliable. This issue is discussed in detail in Jolliffe and Stephenson (2003) and Bröcker and Smith (2007a), while a more general discussion on the uncertainty of verification measures can be found in Jolliffe (2007). Our contribution concerns the fact that not only sampling effects but also serial correlation in sequences of forecast-verification pairs may affect the observed reliability of even perfectly reliable density forecasts of continuous variables. A simple mathematical proof of that effect is given in Appendix A.

3. Example of serial correlation in probability integral transforms for reliable probabilistic forecasts

Consider here the issue of density forecasting of some continuous meteorological variable, say wind speed, at a forecast horizon \( k \). It is common knowledge that climatology comprises a benchmark density forecast, which has the nice property of being well calibrated, and the characteristic of having no resolution because it consists of unconditional density forecasts (see for instance the discussion in Pinson and Madsen, 2009a). Figure 1(a) depicts a time series of mean hourly wind speed at the Horns Rev wind farm in Denmark over a period of almost 1000 h. This offshore wind farm has been the first large-scale offshore wind farm worldwide, and has hence motivated a number of studies for e.g. the characterization of local wind characteristics (Vincent, et al., 2009) or the (probabilistic) forecasting of its power output (Pinson and Madsen, 2009a). The time series of wind speed is normalized by the maximum wind speed observed over the period, consequently taking values in \([0, 1]\).

For any time \( t \) in this dataset, a perfectly reliable climatology density forecast \( \tilde{f}_{t+k|t}(y) \) for a given look-ahead time \( k \) can be obtained by defining it from the distribution of wind-speed hourly averages over the period of 10 000 h. The corresponding cumulative distribution function \( \tilde{F}_{t+k|t} \) is represented in Figure 1(b). This is obviously not the way an actual climatology density forecast would be built, since it would be based on a long-record time series of past measurements. For the purpose of our example, however, the climatology density forecast we build is similar in essence to actual climatology forecasts based on long records of data because it is a unique unconditional density based on recorded data; we ensure it is perfectly calibrated by relying on the measurements themselves. As a consequence, if one were producing a reliability diagram for evaluating the calibration of such density forecasts, the observed proportions would exactly lie on the diagonal.

Following the discussion of section 2, the calibration assessment of density forecasts with reliability diagrams is based on studying the distribution of the random

\[\text{normalized wind speed} = \frac{y}{\text{maximum wind speed observed over the period}}\]

\[\text{Cumulative probability} = \frac{\text{cumulative distribution function}}{\text{total number of observations}}\]

\[\text{Normalized wind speed} = \frac{\text{cumulative distribution function}}{\text{total number of observations}}\]

\[\text{Time [hours]} = \frac{\text{time [hours]}}{\text{total number of hours in the dataset}}\]

\[\text{Normalized wind speed} = \frac{\text{cumulative distribution function}}{\text{total number of observations}}\]

\[\text{Cumulative probability} = \frac{\text{cumulative distribution function}}{\text{total number of observations}}\]
variables $Z_{tk} = \tilde{F}_{tk}(Y_{tk})$, the realizations of which are given by the probability integral transforms $\tilde{z}_t = \tilde{F}_{t}(y_{tk})$. Intuitively, since it is expected that time series of hourly wind-speed averages will exhibit a significant autocorrelation pattern (see discussion in Vincent, et al., 2009), and since such time series are transformed through a monotonic (strictly) increasing function, an autocorrelation pattern is also expected to be present in the time series $\{z_t\}$ of probability integral transforms. This argument also applies to the corresponding time series $\{\tilde{z}_t\}$.

The density forecasts defined by $\hat{F}_{tk}(\cdot)$ are by construction perfectly reliable: we thus have $Z_{tk} \sim U[0,1]$. For random variables distributed $U[0,1]$, it appears more relevant to define autocorrelation in terms of rank correlation for various lags. It is depicted here in Figure 2(a). In parallel, Figure 2(b) shows the linear correlogram for the time series $\{\tilde{z}_t\}$. One sees from Figure 2 that the rank and linear correlograms, in the uniform and Gaussian domains, respectively, look very similar. This is in agreement with the comment of de Oliveira (2003) that, surprisingly, transformations may not significantly affect an observed (serial or spatial) correlation structure. Most importantly, one notices that autocorrelation values appear to be significantly positive (at a 5% significance level) for lags up to 24 h, and significantly different from zero for other lags. This confirms the statement made in section 1 that, even if a necessary condition for density forecasts to be reliable is that $Z_{tk} \sim U[0,1]$, successive random variables (and corresponding probability integral transforms) do not have to be independent. A consequence of this result is that such correlations should be taken into account when performing hypothesis testing about the reliability of density forecasts, or alternatively when issuing consistency or confidence bars whilst inferring the observed reliability. Note also that, as mentioned by Diebold, et al. (1998), one could look at the correlograms of centred probability integral transforms at the power $j$, $j \geq 2$ in order to observe dependences in higher order moments, i.e. mainly for variance, skewness and kurtosis.

4. Consistency bars using surrogate consistency resampling

Let us recall Brocker and Smith (2007a), following Smith (1997), who introduced the proposal of generating consistency bars for reliability diagrams. The idea of consistency resampling is extended here to also account for a correlation pattern in the $\{z_t\}$ time series. The aim of our consistency bars is then, for each of the quantile forecasts that make up density forecasts, to reflect the possible range (for a given confidence level $1 - \beta$) of observed proportions for this quantile if it were indeed reliable, given the temporal dependences observed in the forecast-verification pairs induced by the process and forecasting system of interest. From a hypothesis-testing point of view, one could say that having the observed proportion of quantile forecasts (for a given nominal proportion) within the range of consistency bars implies that one cannot reject the hypothesis of these quantile forecasts being reliable. This paradigm translates to accepting that there may be a number of perfectly reliable forecast systems for the time series of measurements of interest, and that the combination of forecasts and verifications may induce different types of dependence structure in the time series $\{z_t\}$ of probability integral transforms.

The surrogate consistency resampling method belongs to the more general class of resampling methods for dependent data. For an overview of those methods, see e.g. Lahiri (2003). An important advantage of this surrogate approach is that complete time series are simulated based on their spectrum, instead of considering subsamples or blocks. An alternative to the use of surrogate consistency resampling would be to employ a model-based consistency resampling approach, where the general class of autoregressive moving average (ARMA($p,q$)) models could be envisaged for modelling the underlying process in the $\{\tilde{z}_t\}$ time series. A subtle issue in this case would relate to the optimal selection of the autoregressive and moving-average orders $p$ and $q$. Model mis-specification could have a dramatic impact on the quality of the generated consistency bars. An additional advantage of the surrogate approach is that by construction it can be ensured that surrogate time series $\{\tilde{z}_t\}$ will be...
distributed $\mathcal{N}(0, 1)$; this would not be the case if we employed a model-based approach.

4.1. Basics of the surrogate method

The basis of our surrogate consistency resampling method is related to the amplitude-adjusted Fourier transform (AAFT) algorithm described by Theiler, et al. (1992). This algorithm is based on the assumption that the observed time series are a monotonic transformation of realizations from a linear Gaussian process. It is indeed the case that for reliable density forecasts the time series $\{z_k\}$ are a monotonic transformation of $\{\tilde{z}_k\}$, for which each of the realizations are generated from $\mathcal{N}(0, 1)$. The consecutive steps of the AAFT algorithm can be described as follows.

1. The time series $\{z_k\}$ is rescaled and transformed (while ensuring that ranks in the data are conserved) to obtain a linear Gaussian process with the time series of realizations $\{\tilde{z}_k\}$.
2. A Fourier-transform-based algorithm is employed to obtain a Gaussian surrogate time series $\{\tilde{z}'_{k,j}\}$, by randomizing the phases. The Gaussian surrogate time series $\{\tilde{z}'_{k,j}\}$ has the same marginal distribution and correlogram as the time series $\{z_k\}$.
3. $\{\tilde{z}'_{k,j}\}$ is transformed back to the domain of the original time series, leading to the surrogate time series $\{\tilde{z}_{k,j}\}$, but with a slightly different autocorrelation function and power spectrum.

A sound property of the AAFT algorithm is that the surrogate time series have the same periodogram and marginal distribution as the original time series $\{z_k\}$ of probability integral transforms, hence also ensuring they have a similar correlogram. For a more detailed description of the surrogate data method and the AAFT algorithm, the reader is referred to Theiler, et al. (1992). Potential deviations in the marginal distribution or in the spectrum can be corrected by employing the iteratively refined surrogate method of Schreiber and Schmitz (2000). The core of the method, i.e. the phase scrambling, is further discussed in Davison and Hinkley (1997).

One can consider directly employing this surrogate method for generating a set of surrogates that are distributed $\mathcal{U}(0, 1)$ and have the same correlogram as the sequence of observed probability integral transforms. This would be done by first transforming the sequence $\{z_k\}$ to $\mathcal{U}(0,1)$ (while preserving the ranks) and then employing the AAFT algorithm for generating a set of surrogate Gaussian time series. The last step, consisting of transforming back the Gaussian surrogates to $\mathcal{U}(0,1)$, can actually be avoided, owing to the equivalence in the definition of the indicator variable from (3) and (5). The counting necessary for building consistency bars can then be equivalently performed using the surrogate time series $\{\tilde{z}_{k,j}\}$ or $\{\tilde{z}'_{k,j}\}$.

If one directly employs the above method for generating surrogate time series, however, the resulting surrogates will not be sampled from the true process behind $\{\tilde{z}_k\}$, but merely from the periodogram obtained from a limited sample. Consistency bars generated from this method would thus not be valid. Consequently, a proposal method for surrogate consistency resampling from the true process behind $\{\tilde{z}_k\}$ is described below. It consists of first identifying the true process behind $\{\tilde{z}_k\}$ with spectral analysis, and then generating surrogates having periodograms sampled from the smooth spectrum of the process.

4.2. Smooth spectrum and periodogram sampling

Since the core of the surrogate data method relates to the phase scrambling of a linear Gaussian process, a spectral analysis framework appears relevant for characterizing the process $\{\tilde{z}_k\}$. We use $g(\omega)$ to denote the spectrum of the linear Gaussian process $\{\tilde{z}_k\}$, with $\omega$ the angular frequency and $I_\omega(\omega)$ the periodogram of the time series $\{\tilde{z}_k\}$. The periodogram $I_\omega(\omega)$ corresponds to the sample spectrum observed from $\{\tilde{z}_k\}$ and is not a consistent (though unbiased) estimate of $g(\omega)$. A consistent estimate can be obtained instead as a smooth spectrum based on truncated periodograms and lag windows. While only the main equations and results are given here, the reader is referred to Madsen (2007) for more extended developments.

The smooth-spectrum estimate $\hat{g}(\omega)$ of $g(\omega)$ based on the time series $\{\tilde{z}_k\}$ is given as

$$g(\omega) = \frac{1}{2\pi} \sum_{k=-N+1}^{N-1} \lambda_k C_k \exp(-i\omega k),$$

where $C_k$ is the sample autocorrelation for lag $k$ and $\lambda_k$ is a lag window, permitting a reduction in the influence of further lags on the estimate of the spectrum. While there exist a large number of potential lag windows, the class of general Tukey windows is considered here, defined by

$$\lambda_k = \begin{cases} 1 - 2\alpha + 2\alpha \cos(\pi k/M), & |k| \leq M, \\ 0, & |k| > M, \end{cases}$$

where $M$ is the truncation point and $\alpha$ the parameter controlling the shape of the lag window. In particular here we employ the Tukey–Hanning window, defined by $\alpha = 1/4$. Deciding on an appropriate truncation point $M$ may be quite difficult, as this relies on the expertise of the practitioner. It will be shown from the simulations below that the surrogate consistency resampling method proposed is not that sensitive to the choice of $M$. A limited expertise in relation to the expected serial correlation structure present in the $\{\tilde{z}_k\}$ time series can provide a relevant guess. When the smooth-spectrum estimate $\hat{g}(\omega)$ is obtained, it can be used for generating a number of realistic periodograms for the surrogate time series $\{\tilde{z}_{k,j}\}$. For that, let us recall here some properties of the periodogram for linear Gaussian processes:

1. the periodogram values $I_\omega(\omega_p)$, with $\omega_p = (2\pi/N)p$, $p = 1, \ldots, N/2$, the so-called fundamental frequencies, are independent;
2. $2I_\omega(\omega_p)/g(\omega_p) \sim \chi^2(2), p \neq 1, N/2$;
3. $I_\omega(\omega_p)/g(\omega_p) \sim \chi^2(1), p = 1, N/2$.

Property (i), relating to the independence of periodogram values for the fundamental frequencies $\omega_p$ (i.e. those defining the orthogonal basis of the Fourier series), is actually a crucial property beyond the phase scrambling in the surrogate data method introduced above (Davison and Hinkley, 1997; Theiler, et al., 1992). In parallel, the smooth-spectrum estimate $\hat{g}(\omega)$ can be plugged into properties (ii) and (iii) in order to simulate periodograms $I_\omega(\omega_p)$ for the surrogate time series $\{\tilde{z}_{k,j}\}$, from independent random draws of $\chi^2(1)$ and $\chi^2(2)$ random variables.
4.3. Surrogate consistency resampling

A full description of the surrogate consistency resampling method is given here based on the elements introduced in the above paragraphs. Again, consider the question of reliability assessment of a sequence \( \{ \hat{Y}_{t+k|i}(y) \} \) of density forecasts (for a given forecast horizon \( k \)), for a corresponding time series \( \{ y_{t+k} \} \) of observations. Following the paradigm introduced above, one accepts that even if this set of density forecasts were reliable, the forecast-verification pairs would induce a temporal correlation structure in the time series \( \{ z_{t,k} \} \) of probability integral transforms.

The first step of the surrogate consistency resampling approach is to rescale the time series \( \{ z_{t,k} \} \) so that they have a marginal distribution \( U[0,1] \) with a rank-preserving transformation, thus not altering its rank corregrogram. The rescaled time series are then projected into the standard Gaussian domain by using the inverse probit function. It is assumed that the obtained time series \( \{ \tilde{z}_{t,k} \} \) is a sample of what would be the sequence of realizations coming from the forecast-verification pairs if the forecasting system were indeed reliable. The smooth spectrum \( g(\omega) \) of the related linear Gaussian process \( \{ \tilde{z}_{t,k} \} \) is estimated given the choice of a truncation point \( M \). The \( j \)th resampling cycle then consists of the following steps.

1. Generate a surrogate periodogram \( \tilde{I}_N^0(\omega) \) from the smooth spectrum \( g(\omega) \).
2. Employ a Fourier-transform-based algorithm to generate a surrogate time series \( \{ \tilde{Z}_{t,k} \} \) by randomizing the phases of the surrogate periodogram \( \tilde{I}_N^0(\omega) \).
3. Calculate the observed proportions for each of the quantiles with nominal proportions \( \alpha_i, i = 1, \ldots, m \) of the density forecasts using (6), based on the surrogate time series \( \{ \tilde{Z}_{t,k} \} \), yielding the surrogate observed proportions \( \hat{\alpha}_i^{(0)}, i = 1, \ldots, m \). Note that one can obtain surrogate time series of probability integral transforms \( \{ \tilde{z}_{t,k}^{(0)} \} \) and of verifications \( \{ y_{t+k}^{(0)} \} \) by inverse transformation.

This resampling cycle is repeated a number of times \( B \) and yields an empirical distribution of surrogate observed proportions for each quantile of the density forecasts. Let us define \( \tilde{G}_{h,k} \) as the cumulative version of this empirical distribution. \( \tilde{G}_{h,k} \) is a non-parametric estimate of what would be the distribution of proportions that could be observed for the dataset considered, for the quantile with nominal proportion \( \alpha_i \), if it were reliable and given the serial correlation structure induced by the forecast-verification pairs. Given the chosen confidence level \( (1 - \beta) \), the lower and upper bounds of the consistency bars are given by

\[
\begin{align*}
\tilde{a}_{k,i} &= \tilde{G}_{h,k}^{-1}(\beta/2), \\
\tilde{a}_{k,i} &= \tilde{G}_{h,k}^{-1}(1 - \beta/2).
\end{align*}
\]

The same argument as that developed by Bröcker and Smith (2007a) applies here, implying that, by construction, the surrogate time series of probability integral transforms directly relate to a hypothetical sequence of forecast densities \( \{ \hat{f}_{t+k|i}(y) \} \) that would be reliable in view of the corresponding time series \( \{ y_{t+k}^{(0)} \} \) of verifications. This is since it is imposed that surrogate time series \( \{ \tilde{z}_{t,k}^{(0)} \} \) are drawn from a \( N(0,1) \) distribution. In addition, consistency bars can be generated for all quantile forecasts in parallel – they are pointwise consistency bars, since as explained in section 2 the calibration assessment is individually performed for each each quantile with a given nominal proportion. There is, finally, no binning effect to be considered, as the calibration of all quantiles is verified against the same number of observations, corresponding to the number of time indices in \( T \).

5. Simulations

In this section simulations are performed, allowing us to demonstrate the pitfalls stemming from inference of the confidence one may have in reliability diagrams based on an i.i.d. assumption when issuing consistency bars. Simulations are also employed to demonstrate the validity of our approach before applying it to real-world test cases and data. The simulation set-up is described first, followed by simulation results and related comments.

5.1. Simulation set-up

The simulations are performed based on \( \{ \tilde{z}_{t,k} \} \) time series only (i.e. based on linear Gaussian processes), since the conversions from \( N(0,1) \) to \( U[0,1] \) (using the probit function \( \Phi \)) and from \( U[0,1] \) to the original domain of the observations are strictly monotonic, thus preserving ranks and counts. As a consequence, observed proportions of quantile forecasts are equivalent if calculated in the original, uniform or Gaussian domains.

Imagine generating a time series of \( \{ \tilde{z}_{t,k} \} \) of length \( N \) for which each realization is drawn from a standard Gaussian distribution \( N(0,1) \) and with a linear correlogram \( \rho(h), h \) being the difference between time indices. By definition, such a time series of probability integral transforms projected into a standard Gaussian domain corresponds to that for a reliable density forecasting system. Two types of correlograms are considered: on the one hand a damped exponential correlogram \( \rho_d(h) \), corresponding to a stationary first-order Markovian process, and on the other hand a correlogram \( \rho(h) \) taking the form of a damped exponential with oscillations, reflecting a seasonality in the sequence of probability integral transforms in the standard Gaussian domain. The damped exponential is simply given by

\[
\rho_d : h \rightarrow \exp(-\tau h), \quad \tau > 0, \tag{14}
\]

with \( \tau \) the parameter controlling the steepness of the exponential decay. In parallel, the damped exponential with oscillations is defined by

\[
\rho_o : h \rightarrow \frac{1}{2} \left[ \cos \left( \frac{2\pi h}{p} \right) + 1 \right] \exp \left( -\frac{2\pi h}{p} \right), \quad \tau, \rho > 0, \tag{15}
\]

with \( \tau \) being the same type of parameter, while \( \rho \) controls the period of oscillations. For the case of \( \rho_o(h), \tau \) is set to \( \tau = 0.3 \), while \( \rho_d(h) \) is parametrized with \( \tau = 0.6 \) and \( \rho = 12 \). The corresponding correlograms are depicted in Figure 3.

As explained in the above section, when one is calculating observed proportions of quantiles composing density forecasts from a time series \( \{ \tilde{z}_{t,k} \} \) of limited size, with...
For both correlograms, one notices that for certain draws the observed proportions may lie fairly close to the diagonal, while for some other draws they may be quite far from this same diagonal. This is while they all relate in the same way to reliable forecast systems. It can also be seen from Figure 4 that for a stronger correlation pattern like $\rho_s$, deviations from the diagonal may be larger. This suggests that serial correlation in forecast-verification pairs magnifies the sampling effects.

Given a chosen correlogram, a number $m$ of linear Gaussian time series \{\tilde{z}_{t,k}\} (of length $N$) are generated. Consistency bars are produced based on i.i.d. consistency resampling, and are based on the surrogate consistency resampling method described above. It is arbitrarily chosen to focus on 90% consistency bars (that is, for $(1 - \beta) = 0.9$), partly because it is the nominal coverage rate considered by Bröcker and Smith (2007a) and partly because this is a quite common choice for a number of real-world applications.

Note that similar simulations could be performed for other nominal coverage rates $(1 - \beta)$ in order to verify the quality of generated consistency bars. Over the $m$ time series, one counts the number of times the observed proportions for quantile forecasts that make up reliable density forecasts (i.e. the quantile of a standard Gaussian
of length \( N = 400 \) time steps with correlograms \( \rho_d \) or \( \rho_s \). Counts performed on such short time series will clearly be affected by the correlation patterns, as illustrated in Figure 4. In order to verify the coverage rate of the consistency bars, \( m = 1000 \) different time series are generated for each type of correlogram. Comparison is made between consistency bars generated from an i.i.d. consistency resampling method and those generated from the surrogate consistency resampling method introduced above. For this latter case, a benchmark consists of consistency bars generated from the true spectrum of the generated time series. In parallel, consistency bars generated from the estimated smooth spectrum (with various values of the truncation point \( M \)) are also evaluated. A number of \( B = 1000 \) surrogate time series are used for deriving consistency bars for each of the \( m \) time series. As is the case for any computer-intensive resampling method, the number of replications \( B \) should be chosen sufficiently large in order to obtain realistic confidence bounds, though not too large, in order to keep computational time reasonable. The values of the truncation points are chosen from an expert guess based on the analysis of the periodograms of the time series \( \{z_t\} \), as would be done for real-world applications. For the case of the correlogram \( \rho_d \), potential expert guesses could be \( M = \{12, 24, 36\} \) while for the case of the \( \rho_s \) correlogram, they could consist of \( M = \{24, 36, 48, 60\} \). It is often said that a reasonable choice for \( M \) is such that \( M = 2\sqrt{N} \) (Upton and Cook, 2002, pp. 324-325), which would translate to \( M = 40 \) here. The corresponding results, consisting of the observed coverage of consistency bars, are gathered in Tables I and II for the correlograms \( \rho_d \) and \( \rho_s \), respectively.

In both cases, one clearly sees that there is a significant difference between i.i.d. and surrogate consistency resampling methods in terms of observed coverage of the generated consistency bars. As expected, the lack of coverage is more significant as the correlation pattern is stronger, i.e. for the correlogram \( \rho_s \). This is an illustration of the pitfall stemming from assuming independence when serial correlation is indeed present in the sequence of forecast-verification pairs. Such lack of coverage could translate to concluding a lack of calibration of density forecasts over the period considered, while in fact the observed deviation from perfect reliability cannot be deemed significant.

When employing the surrogate consistency method with the true spectrum of the time series \( \{z_t\} \) in order to generate surrogate time series, the observed coverage of the consistency bars is very close to the target 90% nominal coverage. Furthermore, when employing the estimated smooth spectrum instead, the coverage of generated consistency bars is also close to the target 90% nominal coverage. Choosing a truncation point that is too small results in underrepresenting the correlation structure present in the sequence of forecast-verification pairs, leading to the generation of consistency bars that are too narrow. By making a reasonable guess for the truncation point, i.e. \( M = \{24, 36\} \) and \( M = \{48, 60\} \) for the case of the correlograms \( \rho_d \) and \( \rho_s \), respectively, the generated consistency bars appear to have an acceptable coverage. It is true that for practical real-world verification studies it might be difficult to pick an optimal truncation point (as would be the case for any spectral analysis study in any case), but a reasonable guess from the practitioner should provide a sufficiently accurate estimate of the spectrum, leading to appropriate consistency bars.

6. Application to the reliability assessment of density forecasts of wind-power generation

Wind power is the renewable energy with the fastest growth over the last few years. It has a significant share in the electricity generation mix in a number of European countries, most notably in Denmark and Spain. The optimal integration of this renewable energy into the existing electricity system requires forecasts for various ranges of horizons depending on the decisions to be made, i.e. from a few minutes ahead for the control of wind-farm output to several days ahead for offshore maintenance planning. The forecasts that are used most today have an hourly resolution up to 48–72 h ahead, are employed for the trading and management of the wind-power generation and are issued based on one or several forecasts of relevant meteorological variables for the site(s) of interest. If considering lead times from a few minutes up to 2 h ahead, forecasts are then generated from purely statistical methods relying on local measurements only, as for instance in Gneiting, et al. (2006) and Pinson and Madsen (2009b). For an overview of motivations, techniques and practical experience with wind-power forecasting, the reader is referred to Giebel, et al. (2003) and Costa, et al. (2008).

Among the various types of forecasting products employed for wind-energy management, maintenance planning and trading, non-parametric density forecasts are becoming more and more popular, since benefits from their use have been demonstrated (Pinson, et al., 2007a; Matos and Bessa, 2009). This is because the loss functions of forecast users commonly differ from the classical quadratic loss function, for which point forecasts relating to the conditional expectation are optimal. Such loss functions may also vary in time due to the changes in the structure and dynamics of electricity markets. As wind-power generation is a nonlinear and bounded process, predictive densities may be highly skewed and with heavy tails (Lange, 2005), and hence be difficult to model accurately with known parametric families of density functions (see the discussion by Pinson, 2006). This has motivated the development of a large number of non-parametric methods for wind-power density forecasting, based on statistical methods and/or ensemble forecasts (see Bremnes, 2006; Møller, et al., 2008; Nielsen, et al., 2006; Pinson and Madsen, 2009a, among others).

We consider here non-parametric density forecasts of wind-power generation for the whole installed capacity in Western Denmark, which approximately represents \( P_n = 2.5 \) GW over the period considered. All forecasts and measurements are normalized by this nominal capacity, and therefore expressed in percentages of \( P_n \). Forecasts are issued hourly, and have an hourly temporal resolution up to a forecast length of 43 h. The point forecasts of wind-power generation were provided by the wind-power prediction tool (WPPT) as described in Nielsen, et al. (2002), while
Table I. Summary of the observed coverage rate of the 90% consistency bars generated with either i.i.d. or surrogate consistency resampling methods, for time series with correlogram $\rho_d$. Consistency bars are for a set of quantiles defining non-parametric density forecasts. Surrogate consistency resampling based on the true spectrum of the time series is used as a benchmark. For the surrogate consistency resampling method using the estimated smooth spectrum, several expert guesses on the truncation point $M$ are considered. Note that asymptotically as $m$ and $B$ tend towards infinity the observed coverage rate for proportions $\alpha_i$ and $(1 - \alpha_i)$ should be the same. The differences here are due to sampling effects.

<table>
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<tr>
<th>$\alpha_i$</th>
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<th>Surrogate ($M = 24$)</th>
<th>Surrogate ($M = 36$)</th>
<th>Surrogate (true)</th>
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</thead>
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Table II. As Table I, but for $\rho_s$.

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<th>$\alpha_i$</th>
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<th>Surrogate ($M = 36$)</th>
<th>Surrogate ($M = 48$)</th>
<th>Surrogate ($M = 60$)</th>
<th>Surrogate (true)</th>
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the non-parametric density forecasts were generated based on the adapted resampling method initially described in Pinson (2006). The period for which both measurements and forecasts are available runs from the beginning of January 2006 until mid-November 2007. Figure 5 depicts an example with wind-power point forecasts issued on 8 January 2007 at noon and related non-parametric density forecasts, as well as the corresponding measurements. Density forecasts take the form of a set of central prediction intervals (centred in probability around the median) with increasing nominal proportions from 10% to 90%. They thus are defined by 18 quantile forecasts with nominal proportions from 5% to 95% with a 5% increment, except for the median.
Reliability Diagrams for Nonparametric Density Forecasts

Figure 5. Example of non-parametric density forecasts of wind-power generation for the whole of Western Denmark (issued on 8 January 2007 at noon) in the form of a river-of-blood fan chart. Density forecasts are represented as a set of central prediction intervals with increasing nominal proportions. Power values are normalized by the total wind capacity $P_n$ for the region. Measurements and point forecasts are also depicted.

Three different sets of forecast series are arbitrarily selected from the two years that were available. These three periods span autumn 2006 and spring and summer 2007. The first two sets consist of 600 forecast series, while the latter one consists of 800 forecast series. The reliability of non-parametric density forecasts is studied based on reliability diagrams in the form of those presented in section 2. Each lead time is considered individually. Inspection of the periodograms and correlograms of the $\{\tilde{z}_t, k\}$ time series suggests that truncation points between 36 and 60 h could be relevant for the first two sets of forecast series, while $M$ may take values between 72 and 96 h for the case of the last set, for all forecast horizons. A common value of 48 h is selected for the first two sets, while a value of 84 h is picked for the third one. As shown and discussed in section 5, the width of the generated consistency bars is not highly sensitive to the choice for $M$. A sufficiently large number $N$ of surrogate time series is chosen as $B = 1000$.

Let us now interpret these reliability diagrams. If consistency bars were not available, one would subjectively appreciate the observed deviations from the diagonal and decide on acceptable reliability (or not) of the various quantile forecasts that make up non-parametric density forecasts. One would then certainly accept all quantile forecasts to be reliable over set 3. In contrast, quantile forecasts with nominal proportions between 0.45 and 0.8 for set 1 and with nominal proportions between 0.1 and 0.65 for set 2 would be deemed as non-reliable owing to an increased deviation between the ideal diagonal line and observed proportions. Now consider the sets of consistency bars. For sets 1 and 3, the observed proportions of all quantiles that make up density forecasts lie within the consistency bars, even though deviations from the diagonal are of different magnitudes. This does not tell us that the quantile forecasts are reliable, but inversely that it cannot be concluded that they are not reliable (for a 10% level of significance). This goes against the subjective evaluation given previously. In contrast, for set 2 the fact that observed proportions for quantile forecasts with nominal proportions between 0.1 and 0.55 lie outside the consistency bars confirms that quantile forecasts for such nominal proportions should not be considered as reliable.
Figure 6. Example results from the reliability assessment of 42 hour ahead non-parametric density forecasts of wind-power generation. The smooth-
spectrum estimates related to the three sets of forecast series are gathered in (a). Reliability diagrams with consistency bars for a 90% confidence level are
depicted in (b), (c) and (d) for sets 1, 2 and 3.

7. Concluding remarks

Focus has been given to the question of the calibration
assessment of density forecasts of continuous variables,
originating from ensemble forecasts or statistical methods,
with reliability diagrams. It has been explained that
employing such reliability diagrams relates to the evaluation
of unconditional calibration. In many applications one
should acknowledge the presence of serial correlation in
the sequence of probability integral transforms even for
reliable density forecasts, in turn induced by the sequence
of forecast-verification pairs.

We have built on an interesting proposal by Bröcker and
Smith (2007a) consisting of associating reliability diagrams
with consistency bars. Consistency bars here reflect the
potential impact of both limited counting statistics and
serial correlation on what would be the observed reliability
of a perfectly reliable forecast system over the set of available
observations. An original surrogate consistency resampling
method has been introduced and evaluated for that purpose.
The presence of serial correlation clearly increases the width
of consistency bars. Even though the serial correlation
pattern has to be estimated (here in the frequency domain),
it has been demonstrated from simulations that the actual
coverage of our consistency bars is close to their intended
nominal coverage.

The consistency bars that have been considered in
the present article are pointwise consistency bars. This means
that they relate to the individual reliability assessment of
the quantile forecasts (for a given nominal proportion) that
make up non-parametric density forecasts. As a possible
extension of the work presented here, one may consider the
definition of consistency envelopes, which in contrast would
relate to the simultaneous reliability assessment of quantile
forecasts with various nominal proportions (thus for the full
densities) in a multiple hypothesis-testing framework.

Our most important message here is not that one should
mandatorily use the approach introduced for generating
consistency bars, but instead that one must consider the
potential effect of serial correlation in reliability assessments.
Indeed, it has been shown that assuming independence
of the sequence of probability integral transforms clearly
leads to an underestimate of the range of potentially
observed proportions for a perfectly reliable probabilistic
forecasting system over the period of interest. One may
decide on one’s own method of modelling or accounting for
For the surrogate consistency resampling method developed, only one parameter, i.e. the truncation point $M$, has to be selected for the estimation of the smooth spectrum. This is the price to pay for capturing the interdependence structure in the sequence of probability integral transforms. Even though the selection of $M$ may call for some statistical (and/or signal processing) expertise from the practitioner, the actual coverage of consistency bars is not highly sensitive to the choice for $M$, especially if consistency bars are to be used for visual assessment of density-forecast calibrations and not for thorough hypothesis testing. In parallel, the number $B$ of replications of the consistency resampling method, which corresponds to the number of surrogate time series to be generated, should be chosen sufficiently large, e.g. $B \geq 1000$. Since computational power is rapidly increasing, picking a large value for $B$ should not be a problem. For reference, only 4 min were necessary for generating the reliability diagrams with consistency bars of section 6 with Matlab, i.e. for time series of length 600 and 800 with the number of surrogates $B = 1000$.

Unconditional calibration is only one aspect of probabilistic forecast verification. It is a crucial aspect, however, as a probabilistic bias in density forecasts would directly translate to a bias in decisions to be made from such probabilistic forecasts. If one is strict about forecast verification, density forecasts that cannot be deemed as reliable should not be considered further for decision-making. Fortunately, one can easily correct for a lack of unconditional calibration, with e.g. conditional parametric models (Nielsen, et al., 2006) or smoothed bootstrap (Hall and Rieck, 2001). For the case of the application considered, the use of consistency bars has permitted us to enhance the article. Acknowledgements are finally due to Henrik Aa. Nielsen and James W. Taylor, as well as Tilmann Gneiting, for fruitful discussion about the reliability of ensemble and density forecasts.

Appendix

A mathematical proof of the effect of serial correlation on the size of consistency bars

In this appendix a simple mathematical proof is given of the effect of serial correlation on the size of consistency bars. More precisely, we show here that for any type of correlation in the sequence of forecast-verification pairs, the consistency bars are wider than in the i.i.d. case.

Let us focus, without loss of generality, on a given nominal proportion $\alpha$. The forecast horizon $k$ is omitted in the developments below in order to lighten the notations. In view of the definition of the indicator variable $\xi_i^{(\alpha)}$ in (3), it appears that $\xi_i^{(\alpha)}$ is the realization at time $t$ of a Bernoulli random variable with parameter $\alpha$. As a consequence, the observed proportion $\hat{\alpha}$, is a realization of a random variable defined as the sum of Bernoulli trials, scaled by the number of trials $N$, which here is the length of the evaluation period. Below, we will denote this sum by $X_i$, or $\tilde{X}_i$, for the i.i.d. and correlated cases, respectively.

In the case where there is no serial correlation present in the sequence of forecast-verification pairs, the corresponding Bernoulli trials are i.i.d. By definition, the sum $\tilde{X}_i$ of $N$ i.i.d. Bernoulli trials with chance of success $\alpha$ follows a binomial distribution, $X_i \sim \text{Bin}(N, \alpha)$. The first two moments of the distribution of the proportion $X_i/N$ are then given by

$$E[X_i/N] = \alpha_i, \quad \text{(A.1)}$$

and

$$\text{var}[X_i/N] = \frac{\alpha(1 - \alpha)}{N}. \quad \text{(A.2)}$$

In contrast, when serial correlation is present in the sequence of forecast-verification pairs, the corresponding Bernoulli trials cannot be independent. In such a case, it is known that the sum $\tilde{X}_i$ of $N$ dependent Bernoulli trials can be modelled with a beta-binomial distribution, see e.g. Ahn and Chen (1995) or Tsai, et al. (2003). This distribution is defined as

$$\tilde{X}_i \sim \text{Bin}(N, \theta) \quad \text{(A.3)}$$

with

$$\theta \sim \text{Beta}(\alpha_i, \sigma_\theta). \quad \text{(A.4)}$$

Note that, for the sake of simplicity, the beta distribution $\text{Beta}(\alpha_i, \sigma_\theta)$ in the above is characterized by its mean $\alpha_i$ and variance $\sigma_\theta$, instead of its two shape parameters. Consequently, the first two moments of the distribution of the proportion $\tilde{X}_i/N$ are given by

$$E[\tilde{X}_i/N] = \frac{\alpha_i}{N}, \quad \text{(A.5)}$$

and

$$\text{var}[\tilde{X}_i/N] = \frac{\alpha(1 - \alpha)}{N^2} \left(1 + \frac{N - 1}{\sigma_\theta + 1}\right). \quad \text{(A.6)}$$

Since necessarily $N > 1$ and $\sigma_\theta > 0$, one has

$$\text{var}[\tilde{X}_i/N] > \text{var}[X_i/N], \quad \text{(A.7)}$$

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meaning that, whatever $\alpha_i$, the distribution of proportions in the case of serial correlation will have a higher second-order moment than if there were no correlation. Such distributions will in any case be centred on $\alpha_i$ and symmetric around it. Therefore, for any confidence level $(1 - \beta)$, consistency bars will be wider if serial correlation is present in the sequence of forecast-verification pairs.

References


