Evaluating density forecasts: a comment

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Evaluating Density Forecasts: A Comment

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This is a comment on Mitchell and Wallis (2011) which in turn is a critical reaction to Gneiting et al. (2007). The comment discusses the notion of forecast calibration, the advantage of using scoring rules, the “sharpness” principle and a general approach to testing calibration. The aim is to show how a more general and explicitly stated framework can provide further insights into the theory and practice of probabilistic forecasting.

1 What is “forecast calibration”?

Both Gneiting et al. (2007) (hereafter GBR) and Mitchell and Wallis (2011) (hereafter MW) examine various important aspects of calibration, but they do not propose a consistent framework and do not give sufficiently universal definitions of “calibration” and the “ideal forecast”. This results in vague argumentation which can lead to confusion. So I start by clearing up the notion of calibration.

The general idea is that a well-calibrated probabilistic forecast must coincide with a suitable conditional distribution. The classical definition of calibration for dichotomous 0/1 outcomes requires that among the cases in which a forecast places probability $\pi$ on the event $X = 1$ this event must be observed with probability $\pi$: $P(X = 1|\pi) = \pi$. In other words, predicted frequencies must coincide with actual frequencies conditionally on the information contained in the forecast itself. More generally, let $F$ be a probabilistic forecast of $X$ stated in the form of a cumulative distribution function. It is calibrated if

$$F(x) = G(x|F),$$

where $G(x|F)$ is the conditional cumulative distribution function of $X$ given $F$. I call this type of forecast calibration \textit{auto-calibration}.

Now suppose that forecasting is based on some available information $\Omega$. Then $F$ is a function of $\Omega$: $F = F(\cdot;\Omega)$. Conditioning on $\Omega$ allows to strengthen the definition of calibration. The forecast $F = F(\cdot;\Omega)$ is \textit{calibrated with respect to $\Omega$} if

$$F(x) = G(x|\Omega).$$

Such a forecast can be called \textit{the ideal} one (of all the forecasts based on $\Omega$).

\footnote{E.g. Lichtenstein et al. (1982), p. 307: “Formally, a judge is calibrated if, over the long run, for all propositions assigned a given probability, the proportion true equals the probability assigned.”}

\footnote{A formal variant of this definition was proposed independently in Gneiting and Ranjan (2011). It is also similar to the definition of interval forecast efficiency with respect to the information set in Christoffersen (1998).}
Obviously, if a forecast is calibrated with respect to \( \Omega \) then it is auto-calibrated while the converse is not true in general. The conditional cumulative distribution function \( G(x|\Omega) \) need not coincide with \( G(x|F) \), because a non-ideal forecast utilize available information only partially.

Note that these definitions of calibration are made in terms of theoretical moments. GBR give their definitions of calibration in terms of empirical moments (for a potentially infinite sequence of forecasts).

2 Can independence and uniformity of PIT values be called complete calibration?

One point of MW’s criticism is that GBR are concerned with forecasting of white noise instead of more relevant forecasting of dependent time series. However, what is really important is that GBR do not pay due attention to the conditional nature of calibration.\(^3\) MW define “complete calibration” as independence and uniformity of the probability integral transform (PIT) values of a time series. Indeed, it is very true that the available history of a time series (including the PIT values) is an important information for judging calibration of forecasts of this series, but in general this view is unnecessarily narrow. The notion of forecast calibration is not about time series properties, it is about conditioning on forecasting information.

We first need to connect the definition of calibration given above with PIT. If the conditional distribution of \( X \) given \( \Omega \) is absolutely continuous then a probabilistic forecast \( F = F(\cdot;\Omega) \) is calibrated with respect to \( \Omega \) if and only if

\[
P|\Omega \sim U[0,1],
\]

where \( P = F(X|\Omega) \) is PIT of \( X \) based on the forecast \( F \).

Now consider a sequence \( F_t(x) \) of probabilistic forecasts of a univariate time series \( X_t \) based on its history \( \Omega_{t-1} = X_1, \ldots, X_{t-1}, \ t = 1, 2, \ldots \) (For \( t = 1 \) the forecast is unconditional). Define the corresponding PIT values as follows:

\[
P_t = F_t(X_t).
\]

This provides for each \( t \) a one-to-one transformation between \( P_1, \ldots, P_t \) and \( \Omega_t = X_1, \ldots, X_t \) whenever all the distributions are absolutely continuous. The series of forecasts is well-calibrated if and only if

\[
P_t|\Omega_{t-1} \sim U[0,1]
\]

or, equivalently,

\[
P_t|P_1, \ldots, P_{t-1} \sim U[0,1].
\]

MW concentrate on an equivalent property: the series of forecasts is well-calibrated if and only if all \( P_t \) are distributed as \( U[0,1] \) and are jointly independent.\(^4\) The problem with the PIT-based definition is that it is insufficiently generic; it does not generalize to other forecasting situations. Using MW’s words, it is important “to pay attention to the information set on which a forecast is based, its content and its timing”.

For example, independence and uniformity of a sequence of PIT values is not sufficient for calibration in the case when forecasts are based on information other than the history of \( X_t \) itself. In

\(^3\)Clements and Taylor (2003), p. 446: “Evaluating probability forecasts by calibration ignores the conditional aspect”.

\(^4\)Under assumption of calibration with respect to the past history what we have here is the Rosenblatt’s transformation \( P_t = G(X_t|\Omega_{t-1}) \) (Rosenblatt (1952)).
particular, consider predicting $X_t$ from $X_1, \ldots, X_{t-1}$ and the history of some other series $Z_1, \ldots, Z_{t-1}$. Then calibration with respect to $\Omega_{t-1} = X_1, \ldots, X_{t-1}, Z_1, \ldots, Z_{t-1}$ is equivalent to

$$P_t|\Omega_{t-1} \sim U[0, 1],$$

where $P_t = F_t(X_t)$. For a sequence of forecasts $F_t$ these conditions imply uniformity and independence of $P_t$, but in general uniformity and independence of $P_t$ is a weaker condition. The fact is fully recognized by MW, but they do not adjust their definition of “complete calibration” accordingly.

Another illustration of insufficiency of independence and uniformity of PIT values is the case of the unfocused forecast in the AR(2) example below. In that example forecasting information set includes not only the previous history of the series, but also random noise. Therefore there is no one-to-one correspondence between the previous history and the previous PIT values.

Further, independence of PIT values is not necessary for calibration in a situation of multi-step forecasting. An $h$-step-ahead forecast of $X_t$ would be based only on $X_1, \ldots X_{t-h}$ and the PIT series corresponding to the ideal forecast would in general be dependent.

3 Testing vs. scoring rules

Suppose that we need to choose between several probabilistic forecasts. When can we state that one forecast is better than another? This is a situation of decision making under risk. Let $u(x, a)$ be a utility function depending on an outcome $x$ and an action $a$. Suppose $a(F)$ is the best action given a probabilistic forecast $F$, that is,

$$a(F) \in \arg \max_a E[u(X, a)] \text{ for } X \sim F.$$

One can say that $F_1$ is (non-strictly) better than $F_2$ if it provides at least as large expected utility:

$$E[u(X, a(F_1))] \geq E[u(X, a(F_2))].$$

This is closely connected with the notion of a proper scoring rule (see Gneiting and Raftery (2007) for a comprehensive review). Define a scoring rule $S$ as the utility of outcome $x$ under action $a(F)$:

$$S(F, x) = u(x, a(F)).$$

This utility-based scoring rule would be a proper one since

$$S(F_1, F_1) = E[u(X, a(F_1))] \geq E[u(X, a(F_2))] = S(F_2, F_1) \text{ for } X \sim F_1.$$

Here

$$S(F_2, F_1) = E[S(F_2, X)] \text{ for } X \sim F_1$$

is the expected score of forecast $F_2$ under the assumption that $X$ is distributed as $F_1$.

Because of the link with utility maximization it is logical to base the theory of evaluating probabilistic forecasts on proper scoring rules. Then the choice between competing probabilistic forecasts is made on the basis of their expected scores. Expected scores can be estimated by the corresponding empirical average scores. Thus, one can simply select the forecast with the highest average score according to a suitable proper scoring rule. For a fixed set of forecasting methods there is no convincing reason to apply statistical testing.
MW argue that non-ideal forecasts can be distinguished from the ideal one by means of testing forecasting models against each other by equal predictive accuracy tests. However, proper scoring rules are no less suitable for the task of detecting the ideal forecast. It can be shown that the well-calibrated forecast is characterized by the maximal expected score for any proper scoring rule.\footnote{Diebold et al. (1998), p. 866: “…If a forecast coincides with the true data generating process, then it will be preferred by all forecast users, regardless of loss function.” See also Granger and Pesaran (2000).} Indeed, suppose that \( G = G(\cdot|\Omega) \) is such a forecast and \( F \) is some other forecast based on \( \Omega \). Then

\[
E[S(G,X)|\Omega] = S(G,G) \geq S(F,G) = E[S(F,X)|\Omega]
\]

and

\[
E[S(G,X)] \geq E[S(F,X)].
\]

Moreover, under appropriate additional conditions the inequality is strict here if the scoring rule \( S \) is strictly proper. This gives a reason for calling a forecast coinciding with the conditional distribution \( G(\cdot|\Omega) \) ideal.

The usefulness of using scoring rules for detecting the ideal forecast is illustrated by Table IV in MW’s paper. They use the logarithmic scoring rule \( S(f,x) = \log f(x) \), where \( f(x) = F'(x) \) is the probability density function corresponding to the c.d.f. \( F \). The logarithmic rule is known to be (strictly) proper. See also Table 5 in GBR’s paper which compare the average values of the logarithmic score and the continuous ranked probability score (another useful strictly proper scoring rule) of competing forecasts.

Scoring rules are straightforward and easy to deal with. Testing of one forecasting method against another is a less trivial operation while rarely providing much beyond the comparison of average scores. (An example of an understandable use is testing whether the average score of the best method is significantly greater than average scores of rival methods).

Of course, statistical testing is not something redundant in the context of probabilistic forecasting. Most important, various kinds of diagnostic tests can help to develop a forecasting method which is well enough calibrated by showing directions in which methods can be improved. However, discovering mis-calibration by diagnostic testing is not always a straightforward task. The following example is somewhat artificial and is not related to real forecasting problems, but it is suggestive.

Consider a series \( X_t \sim N(0,1), t = 1,2,\ldots \) To obtain \( X_t \) from \( X_{t-1} \) (1) transform \( X_t \) to \( U[0,1] \), (2) transform \( U[0,1] \) to \( U[0,1] \) by a highly nonlinear “tangling” transformation, (3) transform \( U[0,1] \) back to \( N(0,1) \), and finally (4) add an independent normal error and scale to unit variance. For example,

\[
\mu_t = \Phi^{-1}(\{ K \left| 2\Phi(X_{t-1}) - 1 \right| \}) \sqrt{1 - \lambda},
\]

\[
X_t|X_1,\ldots,X_{t-1} \sim N(\mu_t, \lambda).
\]

Here \( \Phi(\cdot) \) is the standard normal c.d.f., \( \{ \cdot \} \) is the fractional part function, \( K \) is a large enough integer and \( \lambda \in (0,1) \). One can see that \( X_t \) is a white noise series and is exactly distributed as \( N(0,1) \) for each \( t \) whenever \( X_1 \sim N(0,1) \) (though it is dependent and thus could not be strictly called a Gaussian white noise series). For large values of \( K \) ordinary PIT-based tests would not detect any dependence. Consequently, on the basis of PIT one would choose \( N(0,1) \) as a good forecasting distribution. However, \( N(\mu_t, \lambda) \) is a dramatically better forecasting distribution for a small \( \lambda \).

It can be concluded from the example that sometimes the direction of mis-calibration is not obvious. Popular PIT-based diagnostic tests can fail to detect departures from perfect calibration. At the same time comparison of forecasts by their average scores with proper scoring rules is feasible.
and works as expected irrespective of the kind of mis-calibration. (A reservation, however, should be made here. If there exists an alternative forecasting method then it can be utilized for the task of testing calibration. Such a test is discussed below.)

Actually MW’s paper is not an exception in its attention to statistical testing of forecasting models. A “test-test-test” bias is typical for econometric forecasting literature in general. Cf. Corradi and Swanson (2006) which is a survey of density forecast evaluation almost entirely devoted to various test procedures.

4 Maximizing sharpness subject to calibration

MW are critical of the conjecture that the problem of finding a good forecast can be viewed as the problem of maximizing sharpness subject to calibration which was stated by GBR. However, it can be shown that the conjecture is actually true provided that a vague “calibration” notion is replaced by auto-calibration introduced above.

If \( F \) is an auto-calibrated forecast (calibrated with respect to itself), that is, \( F = G(\cdot|F) \) and \( S \) is some scoring rule then

\[
E[S(F,X)|F] = S(F,F).
\]

Taking unconditional expectations of both sides gives

\[
E[S(F,X)] = E[S(F,F)].
\]

For a proper scoring rule \( S(F,F) \) can be viewed as a measure of sharpness\(^6\) of a probabilistic forecast \( F \). Hence the expected score for an auto-calibrated forecast equals its expected sharpness. This means that auto-calibrated forecasts can be compared on the basis of the levels of their expected sharpness. The ideal forecast is the sharpest of all auto-calibrated forecasts, because it is characterized by the maximal expected score.

This fact is seen from the classical partitioning of the Brier score for dichotomous outcomes into the sum of sharpness and calibration (“validity”) terms developed in Sanders (1963). For an auto-calibrated forecast the calibration term is zero. Thus, maximizing the expected sharpness among auto-calibrated forecasts is equivalent to maximizing the expected score of a proper scoring rule. Bröcker (2009) extended the decomposition to the case of an arbitrary discrete distribution and an arbitrary proper scoring rule. It can further be shown that the same decomposition holds even more generally and applies also to continuous distributions.

Given some proper scoring rule \( S \) define the divergence\(^7\) between distributions \( F_1 \) and \( F_2 \) as

\[
d(F_2,F_1) = S(F_1,F_1) - S(F_2,F_1)
\]

and denote \( G_F = G(\cdot|F) \). Then for the expected score we have

\[
E[S(F,X)] = E[E[S(F,X)|F]] = E[S(F,G_F)]
\]

which can be decomposed as follows:

\[
E[S(F,X)] = E[S(G_F,G_F)] - E[d(F,G_F)].
\]

\(^6\)For a proper scoring rule \( S(F,F) \) is a convex function of \( F \). Thus, according to DeGroot (1962) a concave function \( -S(F,F) \) can be viewed as a measure of uncertainty of a probability distribution \( F \). For the logarithmic scoring rule \( -S(F,F) \) is the familiar Shannon’s entropy measure.

\(^7\)The divergence \( d(F_2,F_1) \) is non-negative as long as \( S \) is proper and it is zero when the two distributions coincide. For the logarithmic scoring rule \( d \) is the Kullback–Leibler divergence.
The first term can be interpreted as the expected sharpness of $G_F$, which is the “recalibrated” version of forecast $F$, while the second term relates to the divergence between $F$ and $G_F$ (that is, it is a measure of mis-calibration of $F$ with respect to information contained in itself).

Although the sharpness conjecture proves to be correct, MW are right in pointing out that the criterion of sharpness per se seems somewhat excessive. First, obviously a proper scoring rule already combines sharpness and calibration in a balanced manner. Second, the principle of maximizing sharpness subject to calibration is difficult to apply in practice, because achieving perfect auto-calibration of a forecast can prove to be very difficult. However, the sharpness principle provides a useful insight into the essence of probabilistic forecasting.

### 5 A broader approach to calibration testing

MW rightly note that calibration can be evaluated by testing orthogonality conditions analogous to those known from the theory of point forecasting. It is important that their idea can be further generalized, which allows to design various kinds of diagnostic tests for forecast calibration. Many of the tests and criteria of calibration/efficiency developed in the literature can be shown to fall within this approach.

Consider a function $r = r(x, F)$ of an outcome $x$ and a c.d.f. $F$. Define

$$
\rho = \rho(F) = \mathbb{E}[r(X, F)] \quad \text{for } X \sim F.
$$

Auto-calibration of a probabilistic forecast $F$ is equivalent to

$$
\mathbb{E}[(r(X, F) - \rho(F))] = 0
$$

for any $r$. That is, $\rho$ must be an unbiased point forecast for $r$. The perfect calibration of $F$ with respect to $\Omega$ is a stronger property and requires conditional unbiasedness:

$$
\mathbb{E}[(r(X, F) - \rho(F))|\Omega] = 0.
$$

This conditional unbiasedness is equivalent to the following unconditional orthogonality:

$$
\mathbb{E}[(r(X, F) - \rho(F))q(\Omega)] = 0
$$

for any $q$ which is a function of available information $\Omega$. That is, $\rho$ must be an unbiased point forecast for $r$ and the forecast error must be uncorrelated with any function $q$ of the available information $\Omega$.

**Example 1.** For absolutely continuous case when $r = 1\{F(x) \leq p\}, \quad \rho = p$

for $p \in [0, 1]$, we obtain an extended version of the “probabilistic calibration” of GBR. The orthogonality condition $\mathbb{E}[(r - p)q] = 0$ for each $p \in [0, 1]$ and any function $q = q(\Omega)$ is another equivalent characterization of the perfect calibration with respect to $\Omega$. Note that unlike the formulation by GBR this one does not make direct use of the ideal forecast $G(\cdot; \Omega)$ and thus is more suitable for the construction of tests.

**Example 2.** Similarly, the “marginal calibration” of GBR corresponds to

$$
r = 1\{x \leq x_0\}, \quad \rho = F(x_0).
$$
Example 3. “Tests of efficiency” proposed by MW are also a special case of this approach with 
\( r = \Phi^{-1}(F(x)) \) and \( \rho = 0 \).

More generally, consider a function \( m = m(x, \Omega) \) depending on an outcome \( x \) and forecasting information \( \Omega \). Define

\[
\mu = \mu(\Omega) = E[m(X, \Omega)] \quad \text{for} \quad X \sim F = F(\cdot; \Omega).
\]

Calibration of \( F \) with respect to \( \Omega \) is equivalent to the following moment condition:

\[
E[(m(X, \Omega) - \mu(\Omega))] = 0
\]

for any \( m \).

Example 4. One more example parallels equal predictive accuracy tests based on the difference of the logarithmic scores (or Kullback–Leibler information criterion, KLIC) discussed by MW (see Amisano and Giacomini (2007)). The idea is to test calibration of one forecasting method against another one. Suppose that we want to test whether \( F_1 = F_1(\cdot; \Omega) \) is well-calibrated and \( F_2 = F_2(\cdot; \Omega) \) is an alternative forecast. Let \( m = S(F_2, X) - S(F_1, X) \) for some proper scoring rule \( S \). For a forecast \( F_1 \) which is calibrated with respect to \( \Omega \)

\[
\mu = E[m|\Omega] = S(F_2, F_1) - S(F_1, F_1).
\]

So we can test calibration of \( F_1 \) by testing that

\[
E[(S(F_2, X) - S(F_1, X)) - (S(F_2, F_1) - S(F_1, F_1))] = 0.
\]

(2)

Below we call this moment condition relative forecast calibration (RFC). A test based on it would have power against an alternative that \( F_2 \) is calibrated with respect to \( \Omega \), because then

\[
E[m - \mu] = E[(S(F_2, F_2) - S(F_1, F_2)) + E[(S(F_1, F_1) - S(F_2, F_1))] > 0,
\]

if the scoring rule is strictly proper.

For the case of the logarithmic score and two normal forecasting distributions, \( N(\alpha_1, \sigma_1^2) \) and \( N(\alpha_2, \sigma_2^2) \), the moment condition reduces to

\[
E\left[\frac{1}{2\sigma_2^2} (\sigma_1^2 + (\alpha_1 - \alpha_2)^2 - (X - \alpha)^2) - \frac{1}{2\sigma_1^2} (\sigma_2^2 - (X - \alpha_1)^2)\right] = 0.
\]

A pair of reciprocal RFC-based tests (\( F_1 \) against \( F_2 \) and \( F_2 \) against \( F_1 \)) can help to judge possible gains from combining two forecasts.

6 Example: Forecasting an autoregressive process

To illustrate the ideas I first adopt the simulation example from MW’s paper which relates to forecasting of AR(2) series \( Y_t = \phi_1 Y_{t-1} + \phi_2 Y_{t-2} + \epsilon_t \) with independent Gaussian disturbances \( \epsilon_t \sim N(0, 1) \). Table 1 lists six forecasting models to be compared. The actual data-generating process is represented by the ideal forecast. In this example there is no need to estimate parameters as they are known. I consider only the Case (2) with \( \phi_1 = 0.15, \phi_2 = 0.2 \). The length of the series is \( T = 150 \) if not specified otherwise.

I underline only two major aspects. First, I want to demonstrate using this setting that concentrating only on the unconditional uniformity of the PIT values and time-series properties is an unnecessary
narrowing of the opportunities for calibration testing. More general moment conditions can be fruitfully utilized. Second, I want to illustrate separation of model selection from diagnostic testing as opposed to mixing them in pairwise equal predictive ability tests.

Consider the unfocused forecast. Unconditionally it is probabilistically calibrated which leads to uniformity of PIT (cf. Table II of MW). Also the PIT series are serially independent (cf. Table III of MW). However, it is not auto-calibrated, which can be easily detected using a test of orthogonality between the inverse normal transformation of PIT $z_t = \Phi^{-1}(F_t(y_t))$ (which should be unconditionally distributed as $N(0,1)$ under auto-calibration) and the mean of the forecasting distribution. It is labeled “$z \times$ mean test” in Table 2. For an auto-calibrated forecast we have $E[\text{mean}(F_t)z_t] = 0$. The table shows that this condition is rejected in almost 100% of experiments in the case of the unfocused forecast. Here and below I use the usual t-ratios based on heteroskedasticity and autocorrelation consistent (HAC) standard errors for testing moment conditions. The results confirms also that the combined forecast (“Combo”) which is a weighted combination of AR1 and AR2 is not auto-calibrated (although the rejection rate is not very large), while for other forecasts there are no signs of mis-calibration (indeed, ideal, AR1 and AR2 forecasts are all auto-calibrated).

The row labeled “% best” shows the percent of experiments in which the corresponding model had the highest average logarithmic score. The ideal forecast was the best with probability of about 2/3. This moderately large value is explained by small values of the two autoregression coefficients ($\phi_1 = 0.15$, $\phi_2 = 0.2$), which makes the true data generating process rather close to AR1, AR2 and Combo alternatives in terms of the expected logarithmic score for relatively short series ($T = 150$). With $T = 1500$ the ideal forecast dominates the other ones (see row labeled “% best, 1500”). One can see that the average score is a sensible criterion for model selection which behaves in a predictable way. Testing is not really needed for the task of model selection. Of course, a test of whether the chosen model has significantly higher average score than the competitors in the spirit of White (2000) or Hansen (2005) can be useful.

To test calibration of the forecasting models against each other I employ a relative forecast calibration test based on the RFC property discussed above. The test is designed as one-sided to increase its power, because the test statistic is expected to be positive in situations when the alternative forecast can potentially be used to improve the forecast tested for calibration. Notable are the results of AR1 vs. AR2 and AR2 vs. AR1. The tests would frequently suggest the usefulness of combining the two models. The test against the ideal forecast can be seen to have high power. It can be compared to the equal predictive ability test based on the difference of the attained average scores $\Delta L_t = S(G_t,y_t) - S(F_t,y_t)$ (labeled “$\Delta L$ test vs. Ideal”). The later test is also implemented as one-sided to increase its power and make it comparable with the RFC test. The comparison clearly favors the RFC test as an instrument

<table>
<thead>
<tr>
<th>Forecast</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ideal</td>
<td>$N(\phi_1 Y_{t-1} + \phi_2 Y_{t-2}, 1)$</td>
</tr>
<tr>
<td>Clmt</td>
<td>$N(0, \sigma^2_{\text{clmt}})$</td>
</tr>
<tr>
<td>AR1</td>
<td>$N(\rho_1 Y_{t-1}, \sigma^2_{\text{ar1}})$</td>
</tr>
<tr>
<td>AR2</td>
<td>$N(\rho_2 Y_{t-2}, \sigma^2_{\text{ar2}})$</td>
</tr>
<tr>
<td>Combo</td>
<td>$0.5N(\rho_1 Y_{t-1}, \sigma^2_{\text{ar1}}) + 0.5N(\rho_2 Y_{t-2}, \sigma^2_{\text{ar2}})$</td>
</tr>
<tr>
<td>Unfocus</td>
<td>$0.5N(\phi_1 Y_{t-1} + \phi_2 Y_{t-2}, 1) + 0.5N(\phi_1 Y_{t-1} + \phi_2 Y_{t-2} + \tau, 1)$, where $\tau = -1$ or $1$ with equal probabilities</td>
</tr>
</tbody>
</table>

Note: $\rho_1 = \phi_1/(1 - \phi_2)$, $\rho_2 = \phi_1 \rho_1 + \phi_2$, $\sigma^2_{\text{clmt}} = 1/(1 - \phi_1 \rho_1 - \phi_2 \rho_2)$, $\sigma^2_{\text{ar1}} = (1 - \rho_1^2)\sigma^2_{\text{clmt}}$, $\sigma^2_{\text{ar2}} = (1 - \rho_2^2)\sigma^2_{\text{clmt}}$. |
<table>
<thead>
<tr>
<th></th>
<th>Ideal</th>
<th>Climit</th>
<th>AR1</th>
<th>AR2</th>
<th>Combo</th>
<th>Unfoc</th>
</tr>
</thead>
<tbody>
<tr>
<td>$z \times$ mean test</td>
<td>4.5</td>
<td>—</td>
<td>4.5</td>
<td>4.3</td>
<td>17.4</td>
<td>99.8</td>
</tr>
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<td>Exp. log. score</td>
<td>-1.418</td>
<td>-1.456</td>
<td>-1.438</td>
<td>-1.430</td>
<td>-1.425</td>
<td>-1.529</td>
</tr>
<tr>
<td>% best</td>
<td>66.5</td>
<td>1.2</td>
<td>6.8</td>
<td>12.2</td>
<td>13.2</td>
<td>0.1</td>
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<tr>
<td>% best, 1500</td>
<td>98.6</td>
<td>0.0</td>
<td>0.0</td>
<td>0.1</td>
<td>1.3</td>
<td>0.0</td>
</tr>
<tr>
<td>$\Delta L$ test vs. Ideal</td>
<td>—</td>
<td>45.7</td>
<td>32.5</td>
<td>25.4</td>
<td>16.9</td>
<td>90.9</td>
</tr>
<tr>
<td>RFC test vs. Ideal</td>
<td>—</td>
<td>92.6</td>
<td>78.3</td>
<td>62.4</td>
<td>25.8</td>
<td>100.0</td>
</tr>
<tr>
<td>RFC test vs. Climit</td>
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<td>—</td>
<td>4.9</td>
<td>5.0</td>
<td>0.7</td>
<td>100.0</td>
</tr>
<tr>
<td>RFC test vs. AR1</td>
<td>4.7</td>
<td>62.7</td>
<td>—</td>
<td>40.2</td>
<td>3.3</td>
<td>100.0</td>
</tr>
<tr>
<td>RFC test vs. AR2</td>
<td>3.4</td>
<td>86.6</td>
<td>58.6</td>
<td>—</td>
<td>9.4</td>
<td>100.0</td>
</tr>
<tr>
<td>RFC test vs. Combo</td>
<td>4.2</td>
<td>86.8</td>
<td>48.6</td>
<td>31.3</td>
<td>—</td>
<td>100.0</td>
</tr>
<tr>
<td>RFC test vs. Unfoc</td>
<td>3.9</td>
<td>37.0</td>
<td>17.8</td>
<td>9.9</td>
<td>6.0</td>
<td>—</td>
</tr>
</tbody>
</table>

Note: The table is based on 5000 simulations. The figures for the tests are percentages of rejection at 5% asymptotic significance level using the standard normal quantiles. The test statistics are t-ratios with Newey–West HAC standard errors and lag truncation 4. “% best, 1500” corresponds to 10 times longer series ($T = 1500$). The row labeled “Exp. log. score” shows approximate expected logarithmic scores. $S(F_2, F_1)$ functions for mixtures of normals needed for RFC statistics are computed by Monte Carlo with 100 simulations.

of identifying non-ideal forecasts.

7 Example: Forecasting a stock index

The second example is intended to show how the proposed framework can be utilized for the task of evaluating forecasts of real-world time series. The data are daily close levels of Russian stock market index (RTSI) and span the period from 1995-09-01 to 2011-06-08. The RTSI series is brought to stationarity by computing its growth rates in percent $y_t = (\log RTSI_t - \log RTSI_{t-1}) \times 100$. This provides a series of 3,928 observations. The forecasted variable is the growth rate 10 periods ahead. Thus, at time $t$ a forecast of $y_{t+1} + \cdots + y_{t+10} = (\log RTSI_{t+10} - \log RTSI_t) \times 100$ is obtained. (The ten-period horizon corresponds roughly to two weeks of physical time.) The following forecasts are considered.\(^8\)

1. The historical forecast based on the full history (“Hi-full”) uses previously observed 10-period growth rates. The historical observations are resampled to obtain a sample of size 1000.

2. “Hi-200” is also a historical forecast, but uses only a rolling span of the 200 most recent observations and does not use resampling.

3. “ExpSm” forecast is based on exponential smoothing for volatility $\sigma^2_t = \delta y^2_{t-1} + (1 - \delta) \sigma^2_{t-1}$ with the decay factor $\delta = 0.95$. The forecasting distribution is given by $N(0, 10h_t)$ and is derived from an assumption of a Gaussian random walk with the variance $h_t$. The recursion for volatility starts from the sample variance of the first 200 observations.

\(^8\)The RTSI series exhibits significant first-order serial correlation, but its impact on 10-period-ahead forecasts is small so I just ignore it.
Table 3: Forecasts of RTSI

<table>
<thead>
<tr>
<th></th>
<th>Hi-full</th>
<th>Hi-200</th>
<th>ExpSm</th>
<th>GARCH</th>
</tr>
</thead>
<tbody>
<tr>
<td>CRPS</td>
<td>−5.236</td>
<td>−5.232</td>
<td>−5.120</td>
<td>−5.092</td>
</tr>
<tr>
<td>Prob. calibr., 50%</td>
<td>−0.013</td>
<td>−0.004</td>
<td>−0.088°°°</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td>(0.020)</td>
<td>(0.020)</td>
<td>(0.021)</td>
<td>(0.020)</td>
</tr>
<tr>
<td>C. prob. calibr., 50%</td>
<td>0.132°°°</td>
<td>0.042°</td>
<td>−0.031</td>
<td>−0.003</td>
</tr>
<tr>
<td></td>
<td>(0.019)</td>
<td>(0.017)</td>
<td>(0.016)</td>
<td>(0.017)</td>
</tr>
<tr>
<td>RFC test vs. Hi-full</td>
<td>—</td>
<td>0.369°°°</td>
<td>0.273°°°</td>
<td>0.117°°</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(0.089)</td>
<td>(0.057)</td>
<td>(0.038)</td>
</tr>
<tr>
<td>RFC test vs. Hi-200</td>
<td>0.376°°°</td>
<td>—</td>
<td>0.353°°°</td>
<td>0.262°°°</td>
</tr>
<tr>
<td></td>
<td>(0.082)</td>
<td></td>
<td>(0.089)</td>
<td>(0.082)</td>
</tr>
<tr>
<td>RFC test vs. ExpSm</td>
<td>0.506°°°</td>
<td>0.579°°°</td>
<td>—</td>
<td>0.166°°°</td>
</tr>
<tr>
<td></td>
<td>(0.063)</td>
<td>(0.103)</td>
<td></td>
<td>(0.049)</td>
</tr>
<tr>
<td>RFC test vs. GARCH</td>
<td>0.406°°°</td>
<td>0.544°°°</td>
<td>0.222°°°</td>
<td>—</td>
</tr>
<tr>
<td></td>
<td>(0.049)</td>
<td>(0.098)</td>
<td>(0.049)</td>
<td></td>
</tr>
</tbody>
</table>

Note: Newey–West HAC standard errors with lag truncation 10 are shown in brackets. Statistical significance at 5% (1%, 0.1%) level is shown by ° (°°, °°°). RFC tests are CRPS-based.

4. GARCH forecast is based on the standard GARCH(1,1)-t model (GARCH with t distribution) with non-zero mean. The model is estimated recursively by the maximum likelihood method. The forecasting distribution is represented by a sample of 1000 simulated future trajectories.

All the forecasts are produced recursively for the forecasting period staring from the 200-th observation. They are compared by their observed averages of the continuous ranked probability score (CRPS). CRPS can be defined as

\[ S(F_t, y_t) = \frac{1}{2} E|Y - Y'| - E|Y - y_t|. \]

for a pair of independent \( Y, Y' \) from the forecasting distribution \( F_t \) and can be calculated in \( O(N \log N) \) operations needed for sorting if \( F_t \) is represented by a sample of size \( N \) (Gneiting and Raftery (2007)). One advantage of the CRPS over the logarithmic scoring rule is that it allows a forecast of a continuous random variable to be a discrete distribution. This scoring rule is also appealing, because it can be viewed as a generalization of the absolute distance loss which is a popular criterion for evaluation of point forecast.

The following statistics are summarized in Table 3.

1. “CRPS” is the average CRPS.

2. “Prob. calibr., 50%” is an unconditional probabilistic calibration statistic based on (1) for \( p = 0.5 \). This statistic relates to the location as measured by the median of the forecasting distribution.

3. “C. prob. calibr., 50%” is a central unconditional probabilistic calibration statistic with

\[ r = 1 \{ F(y) \in [(1 - p)/2, (1 + p)/2] \}, \quad \rho = p \]

for \( p = 0.5 \). This statistic relates to the coverage of the central 50% interval derived from the forecasting distribution.
4. “RFC vs. ⟨method⟩” is a one-sided RFC test, the same as in previous example, but based on the CRPS.

The selection of the calibration tests is somewhat arbitrary. The proposed framework allows to design many different tests, including other kinds of probabilistic calibration tests. Such tests are formal counterparts of visual inspection of PIT histograms, a method which is frequently employed for evaluation of density forecasts.\footnote{The problem with the histograms is that they can be potentially deceptive without error bands which are robust to heteroskedasticity and autocorrelation.}

The example pertains to a typical practical situation when none of the compared forecasts can be called “ideal”. All forecasts are not well-calibrated to different degrees (Table 3). For example “Hi-full” while having (unconditionally) correct location (see “Prob. calibr., 50%”) notably lack sharpness which is signaled by central probabilistic calibration: the actual values are too seldom found in the tails. “ExpSm” rigidly assumes zero mean which is not corroborated by the observed data and the probabilistic calibration test with \( p = 50\% \) indicates a negative bias accordingly; note also the skew of the histogram in Figure 1(a).

In general “GARCH” looks almost like a calibrated forecast if calibration is judged on the basis of ordinary PIT-based criteria. The histogram of PIT values at Figure 1(b) is not perfectly flat, but its unevenness is not very serious as confirmed by the two probabilistic calibration tests from Table 3. Also there are no serious signs of autocorrelation after lag 10 in both \( z_t = \Phi^{-1}(F_t(y_t)) \) and \( z'_t = \Phi^{-1}(|2F_t(y_t) - 1|) \) (calculated from the “folded” PIT values \( |2F_t(y_t) - 1| \) and designed to capture unaccounted volatility clustering). For example, the 11-th autocorrelation coefficient is 0.071 for \( z_t \) and −0.033 for \( z'_t \) while asymptotic standard errors are 0.045 and 0.031 respectively.\footnote{The Bartlett’s approximation for the variance of \( r_{11} \) is used which assumes no autocorrelation after lag 10 and is given by \((1 + 2r_1^2 + \cdots + 2r_{10}^2)/T\), where \( r_k \) is the \( k \)-th autocorrelation coefficient.}

GARCH model is the leader in terms of the average CRPS level, followed closely by exponential smoothing. However, all of the methods in pairwise comparisons by means of RFC tests show significant mis-calibration. For example, remarkably, GARCH is not able to encompass exponential smoothing, which can be considered as its “cheaper” substitute. The results show a potential for improving the forecasts by combining them.

8 Conclusions

Probabilistic forecasting steadily gains popularity and the two papers, Mitchell and Wallis (2011), Gneiting et al. (2007), make important contributions into this process. However, when discussing statistical procedures related to forecast calibration it seems natural to rely on a generic definition. My

Figure 1: Histograms of PIT values: (a) exponential smoothing; (b) GARCH
comment provides such a definition of calibration and delineates a general framework for calibration testing. The framework can facilitate construction of various diagnostic tests. An example of an interesting new test derived from these principles is a relative forecast calibration (RFC) test.

The current discussion accentuates several noteworthy points related to probabilistic forecasting.

- Uniformity and independence of PIT values as a condition of forecast calibration relates to an unreasonably narrow class of forecasting situations when forecasts of a time series are based on its own full history.

- When evaluating forecasts it is important to remember about possible goals of forecasting. Maximizing expected score corresponding to some proper scoring rule can be considered as a reduced representation of such goals.

- Statistical testing is not a substitute for using scoring rules when it is needed to select one method from a fixed set of forecasting methods. Consequently, equal predictive ability tests can be redundant in such a situation.

- Some types of mis-calibration cannot be detected by conventional PIT-based tests. In general calibration tests should be based the relevant information set.

- “Maximizing sharpness subject to (auto-)calibration” is a legitimate principle of probabilistic forecasting. However, its usefulness is limited, because in practice it is hard to achieve perfect auto-calibration.

- RFC-type tests can be rather sensitive to mis-calibration, which suggests that this class of tests is a natural replacement for equal predictive ability tests in the task of diagnosing imperfections of forecasting methods.

References


