The White Reality Check and Some of Its Recent Extensions

Valentina Corradi\textsuperscript{1} and Norman R. Swanson\textsuperscript{2}

\textsuperscript{1}University of Warwick and \textsuperscript{2}Rutgers University

September 2011

Abstract

Halbert White’s \textit{A Reality Check for Data snooping} (\textit{Econometrica}, 2000) is a seminal contribution to the literature on comparing a benchmark model against multiple competitors. In the paper, he suggests a novel approach for controlling the overall error rate, thus circumventing the data snooping problem arising when comparing multiple different models. In this chapter, we discuss several recent extensions of the Reality Check approach. In particular, we begin by reviewing some recent techniques for constructing valid bootstrap critical values in the case of non-vanishing parameter estimation error, in the context of recursive estimation schemes, drawing on Corradi and Swanson (2007a). We then review recent extensions of the Reality Check to the evaluation of multiple confidence intervals and predictive densities, for both the case of a known conditional distribution (Corradi and Swanson 2006a,b) and of an unknown conditional distribution (Corradi and Swanson 2007b). Finally, as an original contribution of this chapter, we introduce a novel approach in which forecast combinations are evaluated via the examination of the quantiles of the expected loss distribution. More precisely, we compare models looking at cumulative distribution functions (CDFs) of prediction errors, for a given loss function, via the principle of stochastic dominance; and we choose the model whose CDF is stochastically dominated, over some given range of interest.

\textit{JEL classification}: C22, C51.

\textit{Keywords}: block bootstrap, recursive estimation scheme, Reality Check, parameter estimation error.

* Valentina Corradi, Department of Economics, University of Warwick, Department of Economics, Coventry CV4 7AL, UK, email: v.corradi@warwick.ac.uk. Norman R. Swanson, Department of Economics, Rutgers University, 75 Hamilton Street, New Brunswick, NJ 08901, USA, nswanson@con.rutgers.edu. This chapter has been prepared for the Festschrift in honor of Halbert L. White in the event of the conference celebrating his 60th birthday, entitled "Causality, Prediction, and Specification Analysis: Recent Advances and Future Directions", and held at the University of California, San Diego on May 6-7, 2011. Swanson thanks the Rutgers University Research Council for financial support.


1 Introduction

The development of econometrics over the last 50 years has been immutably tied to three key topics, model specification, estimation, and inference. In any of these areas, Hal White has been one of the most important players, if not the most important. To see this, it is enough to cite three of his most path-breaking books: *Asymptotic Theory for Econometrician* (1984), *A Unified Theory of Estimation and Inference for Nonlinear Dynamic Models* (with Ron Gallant, 1988), and *Estimation, Inference and Specification Analysis* (1994), each of which contains numerous chapters that are key to the development of modern econometric theory. One of the fundamental intuitions of Hal White is that models, as an approximation of Reality, are generally wrong. Nevertheless, we can learn important things, even from a wrong model.

To allow for misspecification is absolutely crucial in the context of out-of-sample prediction. In this chapter, we begin by assuming that we are given multiple predictions, arising from multiple different models. Our objective is either to select the model(s) producing the more accurate predictions, for a given loss function, or alternatively, to eliminate the models giving the least accurate predictions. Furthermore, in many such situations, we can choose a benchmark or reference model. This can be a model suggested by economic theory, or the winner of past competitions, or simply a model commonly used by practitioners. The key challenge, thus, is to asses whether there exists a competing model outperforming the benchmark. However, if we sequentially compare the reference model with each of its competitors, we may run into problems. In fact, as the number of competitors increases, the probability of picking an alternative model just by ”luck”, and not because of its intrinsic merit, increases and eventually will reach one. This is the well known problem of data-mining or data snooping.

Hal White’s *A Reality Check for Data Snooping* (*Econometrica*, 2000) is a seminal contribution to the issue of comparing a benchmark model against multiple competitors. The null hypothesis is that no competing model can produce a more accurate prediction than the benchmark model, at least for a given loss function. Hal recognizes that this is a composite null hypothesis, formed by the intersection of many individual hypotheses. The data snooping issue arises if we treat a composite null as a sequence of individual independent null hypotheses. In this case, we run into a sequential test bias problem, and we will eventually pick a competing model simply by chance. White’s Reality Check suggests a novel approach for controlling the overall probability of rejecting
the null hypothesis when it is true. Of note is that if we fail to reject the null hypothesis that no competitor outperforms the benchmark model, the obvious consequence is to base prediction on only the benchmark model. This is somewhat in contrast with the alternative approach based on forecast combination (see e.g. Elliott and Timmermann, 2004). The main original contribution of this chapter is that we suggest a novel approach which applies the Reality Check to forecast combinations. This is particularly useful, when we want to select among ways of aggregating forecasts from different agents or from a panel of professional forecasters.

The prediction errors used to construct Reality Check type tests arise in at least two ways. First, there are situations in which we have series of prediction errors, although we do not know the models used to generate the underlying predictions. For example, this situation arises when we have forecasts from different agents, or professional forecasters. Alternatively, we may have a sequence of Sharpe ratios or returns from different trading rules, as in the financial applications of Sullivan, Timmermann and White (1999, 2001). Second, there are situations in which we are interested in comparing estimated models. For example, we may want to decide whether to predict tomorrow’s inflation rate using an autoregressive model, a threshold model, or a Markov switching model. The parameters of these models are generally estimated. If the number of data used to estimate the model is larger than the number of observations used for forecast evaluation, or if the same loss function is used for in sample estimation and out of sample prediction, e.g. estimation by ordinary least squares (OLS) and a quadratic loss function, then the contribution of estimated parameters can be ignored. Otherwise, it has to be taken into account. The 2000 Reality Check mainly deals with the case in which parameters estimation error can be ignored. In particular, the suggested bootstrap procedure does not consider the issue of bootstrap estimation of the underlying parameters. Corradi and Swanson (2006a and 2007a) develop bootstrap procedures which properly capture the contribution of parameters estimation errors in the case of rolling or recursive estimation schemes, respectively.

The White Reality Check concerns comparison of point forecasts (and forecast errors) from multiple models. For example, we want to pick the model producing the most accurate point predictions of the inflation rate. However, there are situations in which we are instead interested in finding the model producing the most accurate interval predictions (e.g. that inflation will be within a given interval). Predictive interval accuracy is particularly important in the management of financial risk, in the insurance and banking industries, where confidence intervals or entire
conditional distributions are often examined. Evaluation of Value at Risk and Expected Shortfall are two main examples (see Duffie and Pan (1997) for further discussion). Corradi and Swanson (2005, 2006a, 2006b) extend the Reality Check to the case of intervals and conditional distributions.

The rest of the chapter is organized as follows. In Section 2 we discuss the Reality Check, and outline how to construct valid bootstrap $p$-values in the case of non-vanishing parameter estimation error, with both recursive and rolling estimation schemes. In Section 3 we extend the Reality Check to the evaluation of multiple confidence intervals and predictive densities. While Sections 2 and 3 survey some recent developments in the forecasting literature, Section 4 takes the approach one step further by combining forecast combination with multiple model evaluation, and introducing a new approach in which forecast combinations are evaluated via the examination of the quantiles of the expected loss distribution, using a Reality Check type testing approach. More precisely, we compare models by prediction error CDFs, for given loss functions, via the principle of stochastic dominance; and we choose the model whose CDF is stochastically dominated, over some given range of interest.

2 The White (2000) Reality Check

2.1 The Case of Vanishing Estimation Error

We begin by outlining the White (2000) Reality Check when parameter estimation error is asymptotically negligible. Consider a collection of $K + 1$ models, where model 0 is treated as the benchmark or reference model and models $k = 1, \ldots, K$ compose the set of competing models. The $h$–step ahead forecast error associated with model $k$, is $u_{i,t+h} = y_{t+h} - \phi_k(Z_t, \theta_k^{\dagger})$. As $\theta_k^{\dagger}$ is unknown, we don’t observe the prediction error $u_{k,t+h}$, but we only observe $\hat{u}_{k,t+h} = y_{t+h} - \phi_k(Z_t, \hat{\theta}_{k,t})$, where $\hat{\theta}_{k,t}$ is an estimator of $\theta_k^{\dagger}$ based on observations available at time $t$.

The common practice in out-of-sample prediction is to split the total sample of $T$ observations into two sub samples of length $R$ and $P$, with $R + P = T$. One uses the first $R$ observations to estimate a candidate model, and construct the first $h$–step ahead prediction error. Then, one uses $R+1$ observations to re-estimate the model and compute the second $h$–step ahead prediction error, and so on, until one has a sequence of $(P - h + 1) h$–step ahead prediction errors.$^1$ If we use this

$^1$Here, we use a recursive estimation scheme, where data up to time $t \geq R$ are used in estimation. West and McCracken (1998) also consider a rolling estimation scheme, in which a rolling windows of $R$ observations is used for
recursive estimation scheme, at each step the estimated parameters are given by

\[ \hat{\theta}_{k,t} = \arg \max_{\theta_k} \left\{ \frac{1}{t} \sum_{j=1}^{t} q_{k,j} (X_{k,t}, \theta_k) \right\} \text{ for } t \geq R, \]

(1)

where \( q_{k,j} \) can be thought of as the quasi-likelihood function associated with model \( k \). Under stationarity, \( \hat{\theta}_k^* = \arg \max_{\theta_k} E(q_{k,j} (X_{k,t}, \theta_k)) \).

Hereafter, for notational simplicity, we consider only the case of \( h = 1 \). Now, for a given loss function, \( g \), the Reality Check tests the following hypotheses:

\[ H_0 : \max_{k=1,\ldots,K} E \left( g(u_{0,t+1}) - g(u_{k,t+1}) \right) \leq 0 \]

versus

\[ H_A : \max_{k=1,\ldots,K} E \left( g(u_{0,t+1}) - g(u_{k,t+1}) \right) > 0. \]

The null hypothesis is that no competing model outperforms the benchmark, for a given loss function, while the alternative is that at least one competitor outperforms the benchmark. By jointly considering all competing models, the Reality Check controls the family-wise error rate (FWER), and circumvents the data snooping problem. In fact, the Reality Check procedure ensures that the probability of rejecting the null when it is false is smaller than or equal to a fixed nominal level, \( \alpha \). Needless to say, in the case of \( K = 1 \), the null hypothesis is a single hypothesis (i.e. equal forecast accuracy of models 0 and 1), and coincides with the Diebold Mariano (1995) test. The Reality Check statistic is given by:

\[ \hat{S}_P = \max_{k=1,\ldots,K} \hat{S}_P(0,k), \]

(2)

where

\[ \hat{S}_P(0,k) = \frac{1}{\sqrt{P}} \sum_{t=1}^{T-1} \left( g(\tilde{u}_{0,t+1}) - g(\tilde{u}_{k,t+1}) \right), \quad k = 1,\ldots,K. \]

\[ ^2 \text{If we instead use a rolling estimation scheme, then} \]

\[ \hat{\theta}_{k,t} = \arg \max_{\theta_k} \left\{ \frac{1}{R} \sum_{j=t-R+1}^{t} q_{k,j} (X_{k,t}, \theta_k) \right\} \quad R \leq t \leq T. \]

\[ ^3 \text{See Christoffersen and Diebold (1996,1997) and Elliott and Timmermann (2004,2005) for a detailed discussion of loss functions used in predictive evaluation.} \]
Letting $S_P(0,k) = \frac{1}{\sqrt{P}} \sum_{t=R}^{T-1} (g(u_{0,t+1}) - g(u_{k,t+1}))$, it is immediate to see that,

$$
\hat{S}_P(0,k) - S_P(0,k) = E(\nabla_{\theta_0} g(u_{0,t+1})) \frac{1}{\sqrt{P}} \sum_{t=R}^{T} \left( \hat{\theta}_{0,t} - \theta_0^T \right)
- E(\nabla_{\theta_0} g(u_{k,t+1})) \frac{1}{\sqrt{P}} \sum_{t=R+1}^{T} \left( \hat{\theta}_{k,t} - \theta_k^T \right) + o_P(1).
$$

Now, if $g = q_k$, then by the first order conditions, $E(\nabla_{\theta_0} g(u_{k,t+1})) = 0$. Thus, if we use the same loss function for estimation and prediction (e.g. we estimate the model by OLS and use a quadratic loss function), then parameter estimation error is asymptotically negligible. Furthermore, if $P/R \to 0$, as $P, R, T > \infty$ (i.e. the sample used for estimation grows at a faster rate than the sample used for forecast evaluation), then parameter estimation is again asymptotically negligible. Otherwise, it has to be taken into account.

Proposition 2.2 in White (2000) establishes that

$$
\max_{k=1,\ldots,K} \frac{1}{\sqrt{P}} \sum_{t=R}^{T-1} ((g(\tilde{u}_{0,t+1}) - g(\tilde{u}_{k,t+1})) - \mu_k) \xrightarrow{d} \max_{k=1,\ldots,K} Z_k,
$$

where $\mu_k = E(g(\tilde{u}_{0,t+1}) - g(\tilde{u}_{k,t+1}))$, $Z = (Z_1, \ldots, Z_K)^T$ is distributed as $N(0, V)$ and $V$ has typical element

$$
v_{j,k} = \lim_{P \to \infty} \frac{1}{\sqrt{P}} \sum_{t=R+1}^{T} (g(\tilde{u}_{0,t+1}) - g(\tilde{u}_{k,t+1})) \cdot \frac{1}{\sqrt{P}} \sum_{t=R+1}^{T} (g(\tilde{u}_{0,t+1}) - g(\tilde{u}_{k,t+1})).
$$

Because the maximum of a Gaussian process is not a Gaussian process, the construction of $p$-values for the limiting distribution above is not straightforward. White proposes two alternatives: (i) a simulation-based approach and (ii) a bootstrap-based approach. The first approach starts from a consistent estimator of $V$, say $\tilde{V}$. Then, for each simulation $s = 1, \ldots, S$, we construct

$$
\tilde{d}_P^{(s)} = \begin{pmatrix} \tilde{d}_{1,P}^{(s)} \\ \vdots \\ \tilde{d}_{K,P}^{(s)} \end{pmatrix} = \begin{pmatrix} \tilde{v}_{1,1} & \cdots & \tilde{v}_{1,K} \\ \vdots & \ddots & \vdots \\ \tilde{v}_{K,1} & \cdots & \tilde{v}_{K,K} \end{pmatrix}^{1/2} \begin{pmatrix} \tilde{\eta}_1^{(s)} \\ \vdots \\ \tilde{\eta}_K^{(s)} \end{pmatrix},
$$

where $(\tilde{\eta}_1^{(s)}, \ldots, \tilde{\eta}_K^{(s)})^T$ is drawn from a $N(0, I_K)$. Then, we compute $\max_{k=1,\ldots,K} |\tilde{d}_P^{(s)}|$, and the $(1-\alpha)$-percentile of its empirical distribution. This simulation based approach requires the estimation of $V$. Note that we can use an estimator of $V$ which captures the contribution of parameter estimation error, along the lines of West (1996) and West and McCracken (1998). However, if $K$
is large, and forecasting errors exhibit a high degree of time dependence, estimators of the long-run variance become imprecise and ill-conditioned, making inference unreliable, especially in small samples. This problem can be overcome using bootstrap critical values.

White (2000) outlines the construction of bootstrap critical values when the contribution of parameter estimation error to the asymptotic covariance matrix is asymptotically negligible. In this case, we resample blocks of \( g(\hat{u}_{0,t+1}) - g(\hat{u}_{k,t+1}) \) and, for each bootstrap replication \( b = 1, \ldots, B \), calculate

\[
\tilde{S}_P^*(b) = \frac{1}{\sqrt{P}} \sum_{t=R+1}^T (g^*(\hat{u}_{0,t+1}) - g^*(\hat{u}_{k,t+1})).
\]

Then, we compute the bootstrap statistic as \( \max_{k=1,\ldots,K} \left| \tilde{S}_P^*(0,k) - \tilde{S}_P(0,k) \right| \) and the \( (1-\alpha) \)-percentile of the empirical distribution of \( B \) such statistics is used for inference.

Before turning to the issue of constructing Reality Check \( p \)-values in the case of non-vanishing parameter estimation error, it is worthwhile to review some other recent developments in the Reality Check literature.

### 2.1.1 Controlling for Irrelevant Models

From the statistic in (2), it is immediate to see that any model which is strictly dominated by the benchmark does not contribute to the limiting distribution, simply because it does not contribute to the maximum. On the other hand, all models contribute to the limiting distribution of either the simulated or the bootstrap statistic. Thus, by introducing irrelevant models, the overall \( p \)-value increases. In fact, for a given level \( \alpha \), the probability of rejecting the null when it is false is \( \alpha \) when all models are as good as the benchmark (i.e. when \( E(g(u_{0,t+1}) - g(u_{k,t+1})) = 0 \) for \( k = 1,\ldots,K \)), otherwise the probability of rejecting the null is smaller than \( \alpha \), and decreases as the number of irrelevant models increases. While the Reality Check is able to control the family-wise error rate, and so avoids the issue of data snooping, it may thus be rather conservative.

For this reason, attempts have been made to modify the Reality Check in such a way as to control for both the family-wise error rate and the inclusion of irrelevant models. Hansen (2005) suggests a variant of the Reality Check, called the SPA (Superior Predictive Ability) test, which is less sensitive to the inclusion of poor models and thus less conservative. The SPA statistic is given...
by
\[ T_P = \max \left\{ 0, \max_{k=1, \ldots, K} \frac{1}{\sqrt{v_{k,k}}} \sum_{t=R+\tau}^{T} \hat{d}_{k,t} \right\}, \]

where \( \hat{d}_{k,t} = (g(\hat{u}_{0,t+1}) - g(\hat{u}_{k,t+1})) \) and \( \hat{v}_{k,k} \) is defined as in (4). The bootstrap counterpart to \( T_P \) at replication \( b \), \( T_P^{* (b)} \) is given by
\[ T_P^{* (b)} = \max \left\{ 0, \max_{k=1, \ldots, K} \left\{ \frac{1}{\sqrt{v_{k,k}}} \sum_{t=R+\tau}^{T} \left( \hat{d}_{k,t}^{(b)} - \hat{d}_{k,t} \mathbb{1}\{ \hat{m}_{k,b} > \hat{v}_{k,k} \sqrt{2 \ln \ln T/P} \} \right) \right\} \right\}. \]

Here, \( p \)-values for the SPA statistic are given by \( 1/B \sum_{b=1}^{B} 1\{ T_P^{* (b)} > T_P \} \). The logic underlying the construction of the SPA \( p \)-values is the following. When a model is too slack, and so it does not contribute to \( T_P \), the corresponding bootstrap moment condition is not recentered, and so the bootstrap statistic is also not affected by the irrelevant model. The fact that very poor models do not contribute to the bootstrap \( p \)-values makes the SPA \( p \)-values less conservative than the Reality Check \( p \)-values. Nevertheless, it cannot be established that the SPA test is uniformly more powerful than the Reality Check test. Corradi and Distaso (2011), using the generalized moment selection approach of Andrews and Soares (2010), derive a general class of superior predictive accuracy tests, which control for FWER and the contribution of irrelevant models. They show that Hansen’s SPA belongs to this class. Additionally, Romano and Wolf (2005) suggest a multiple step extension of the Reality Check which ensures tighter control of irrelevant models. A review of alternative ways of controlling for the overall error rate is provided in Corradi and Distaso (2011), and references contained therein.

2.1.2 Conditional Predictive Ability

In the Diebold-Mariano framework, as well as in the Reality Check framework, model \( k \) and model 0 are considered equally good, in terms of a given loss function, \( g \), if \( \mathbb{E}(g(u_{t,0}) - g(u_{t,k})) = 0 \). This is a statement about forecasting models. In fact, the null hypothesis is evaluated at the "pseudo-true" value for the parameters. Giacomini and White (GW: 2006) propose a novel approach in which model \( k \) and model 0 are considered equally good if \( \mathbb{E}(g(\hat{u}_{t,0}) - g(\hat{u}_{t,k}) | \mathcal{G}_t) = 0 \), where \( \mathcal{G}_t \) is an information set, containing (part of) the history available up to time \( t \). The two key differences between unconditional and conditional predictive accuracy tests are: (i) model comparison is based on estimated parameters in the GW approach, rather than on their probability limits, and (ii)
models in the GW approach are evaluated according to the expected loss conditional on a given information set \( G_t \), rather than unconditionally. The above is a statement about forecasting methods rather than forecasting models. The notion is that not only the model, but also the way it is estimated matters. Needless to say, if a large number of observations is used for estimation, the estimated parameters get close to their probability limits. For this reason, GW suggest using relatively short observation windows, whose length is fixed and does increase with the sample size. In this way, estimated parameters can be treated as strong mixing random variables.

### 2.2 Bootstrap Critical Values for Recursive Estimation Schemes

Whenever \( g \neq q_k \), for at least some \( k \), and \( P/R \to \pi \neq 0 \), then parameter estimation error contributes to the variance of the limiting distribution of the reality check test. One reason for using a different loss for estimation and prediction occurs when, for example, we use OLS for estimation, but then we want to use an asymmetric loss function which penalizes positive and negative errors in a different manner, when comparing predictive accuracy (see Zellner, 1986 and Christoffersen and Diebold, 1997). More specifically, when parameter estimation error does not vanish, we need to take into account the contribution of \( \frac{1}{\sqrt{P}} \sum_{t=R+1}^{T} \left( \hat{\theta}_{k,t} - \theta^1_k \right) \) to the asymptotic variance in (4). Hence, we need a bootstrap procedure which is valid for recursive \( m \)-estimators, in the sense that its use suffices to mimic the limiting distribution of \( \frac{1}{\sqrt{P}} \sum_{t=R}^{T-1} \left( \hat{\theta}_{k,t} - \theta^1_k \right) \).

One approach to the above issue of parameter estimation error is to use the block bootstrap for recursive \( m \)-estimators for constructing critical values. In this context, it is important to note that earlier observations are used more frequently than temporally subsequent observations, when forming test statistics. On the other hand, in the standard block bootstrap, all blocks from the original sample have the same probability of being selected, regardless of the dates of the observations in the blocks. Thus, the bootstrap estimator, say \( \hat{\theta}^*_t \), which is constructed as a direct analog of \( \hat{\theta}_t \), is characterized by a location bias that can be either positive or negative, depending on the sample that we observe. In order to circumvent this problem, Corradi and Swanson (2007a) suggest a re-centering of the bootstrap score which ensures that the new bootstrap estimator, which is no longer the direct analog of \( \hat{\theta}_{k,t} \), is asymptotically unbiased. It should be noted that the idea of re-centering is not new in the bootstrap literature for the case of full sample estimation. In fact, re-centering is necessary, even for first order validity, in the case of overidentified generalized method of moments (GMM) estimators (see e.g. Hall and Horowitz, 1996, Andrews, 2002, and...
Inoue and Shintani, 2006). This is due to the fact that, in the overidentified case, the bootstrap moment conditions are not equal to zero, even if the population moment conditions are. However, in the context of $m-$estimators using the full sample, re-centering is needed only for higher order asymptotics, but not for first order validity, in the sense that the bias term is of smaller order than $T^{-1/2}$ (see e.g. Andrews, 2002 and Goncalves and White, 2004). In the case of recursive $m-$estimators, on the other hand, the bias term is instead of order $T^{-1/2}$, so that it does contribute to the limiting distribution. This points to a need for re-centering when using recursive estimation schemes.

To keep notation simple, suppose that we want to predict, $y_t$ using one of its past lags, and one lag of vector of additional variables, $X_t$, and let $Z_t = (y_t, X_t)$. Using the overlapping block resampling scheme of Künsch (1989), at each replication, we draw $b$ blocks (with replacement) of length $l$ from the sample $W_t = (y_t, Z_{t-1})$, where $bl = T - 1$. Let $W_t^* = (y_t^*, Z_{t-1}^*)$ denote the resampled observations. As a bootstrap counterpart to $b_{θ,k,t}$, Corradi and Swanson (2007a) suggest constructing $b_{θ,k,t}^*$, defined as follows:

$$
\hat{θ}_{k,t}^* = \arg \min_{θ_k} \frac{1}{l} \sum_{j=1}^{l} \left( q_k(y_j^*, Z_{j-1}^*, θ_k) - θ_k' \left( \frac{1}{T} \sum_{h=1}^{T-1} \nabla_{θ_k} q_k(y_h, Z_{h-1}, \hat{θ}_{k,t}) \right) \right),
$$

where $R ≤ t ≤ T - 1$, $k = 0, 1, ..., K$.

Note that $\hat{θ}_{k,t}^*$ is not the direct analog of $\hat{θ}_{k,t}$ in (1). Heuristically, the additional recentering term in (5) has the role of offsetting the bias that arises due to the fact that earlier observations have a higher chance of being drawn than temporally subsequent observations. Theorem 1 in Corradi and Swanson (2007a) establishes that the limiting distribution of $\frac{1}{\sqrt{T}} \sum_{t=R}^{T-1} \left( \hat{θ}_{k,t}^* - \hat{θ}_{k,t} \right)$ is the same as that of $\frac{1}{\sqrt{T}} \sum_{t=R}^{T-1} \left( \hat{θ}_{k,t} - θ_k^\dagger \right)$, conditional on the sample, and for all samples except a set with probability measure approaching zero. We can easily see how this result allows for the construction of valid bootstrap critical values for the Reality Check. Let $\hat{u}_{k,t+1} = y_{t+1} - φ_k \left( Z_t, \hat{θ}_{k,t} \right)$ and $\hat{u}_{k,t+1}^* = y_{t+1}^* - φ_k \left( Z_t^*, \hat{θ}_{k,t}^* \right)$, so that the Reality Check statistic $S_P$ is defined as in (2). The bootstrap counterpart of $S_P$ is given by

$$
\hat{S}_P^* = \max_{k=1, ..., K} S_P^*(0, k),
$$
where
\[
\hat{S}_P^*(0, k) = \frac{1}{\sqrt{P}} \sum_{t=0}^{T-1} \left[ (g(y_{t+1}^* - \phi_0(Z_{t+1}^*, \hat{\theta}_{0,t}^*))) - g(y_{t+1}^* - \phi_k(Z_{t+1}^*, \hat{\theta}_{k,t}^*))) \right]
- \left\{ \frac{1}{T} \sum_{j=1}^{T-1} \left( g(y_{j+1}^* - \phi_0(Z_{j+1}^*, \hat{\theta}_{0,t}^*))) - g(y_{j+1}^* - \phi_k(Z_{j+1}^*, \hat{\theta}_{k,t}^*))) \right\} \right].
\] (6)

It is important to notice that the bootstrap statistic in (6) is different from the “usual” bootstrap statistic, which is defined as the difference between the statistic computed over the sample observations and over the bootstrap observations. In fact, in \(\hat{S}_P^*(0, k)\), the bootstrap (resampled) component is constructed only over the last \(P\) observations, while the sample component is constructed over all \(T\) observations. The percentiles of the empirical distribution of \(\hat{S}_P^*\) can be used to construct valid bootstrap critical values for \(\hat{S}_P\), in the case of non vanishing parameter estimation error. Their first order validity is established in Proposition 2 in Corradi and Swanson (2007a). Valid bootstrap critical values for the rolling estimation case are outlined in Corradi and Swanson (2006a).

3  Extending the Reality Check to Forecast Interval Evaluation

3.1  The Case of Known Distribution Function

Thus far, we have discussed pointwise predictive accuracy testing (i.e. wherein models are evaluated on the basis of selecting the most accurate pointwise forecasts of a given variable). However, there are several instances in which merely having a “good” model for the conditional mean and/or variance may not be adequate for the task at hand. For example, financial risk management involves tracking the entire distribution of a portfolio, or measuring certain distributional aspects, such as value at risk (see e.g. Duffie and Pan (1997)). In such cases, models of conditional mean and/or variance may not be satisfactory. A very small subset of important contributions that go beyond the examination of models of conditional mean and/or variance include papers which: assess the correctness of conditional interval predictions (see e.g. Christoffersen (1998)); assess volatility predictability by comparing unconditional and conditional interval forecasts (see e.g. Christoffersen and Diebold (2000)); and assess conditional quantiles (see e.g. Giacomini and Komunjer (2005)). A thorough review of the literature on predictive interval and predictive density evaluation is given in Corradi and Swanson (2006b).
Corradi and Swanson (2006a) extend the Reality Check to predictive density evaluation, and outline a procedure for assessing the relative out-of-sample predictive accuracy of multiple misspecified conditional distribution models, that can be used with rolling and recursive estimation schemes. The objective is to compare these models in terms of their closeness to the true conditional distribution, \( F_0(u|Z^t, \theta_0) = \Pr(y_{t+1} \leq u|Z^t) \).\(^4\) In the spirit of White (2000), we choose a particular conditional distribution model as the “benchmark” and test the null hypothesis that no competing model can provide a more accurate approximation of the “true” conditional distribution, against the alternative that at least one competitor outperforms the benchmark model. Following Corradi and Swanson (2005), accuracy is measured using a distributional analog of mean square error. More precisely, the squared (approximation) error associated with model \( k, k = 1, \ldots, K \), is measured in terms of the average over \( U \) of \[ E \left( \left( F_k(u|Z^t, \theta_k^1) - F_0(u|Z^t, \theta_0) \right)^2 \right), \] where \( u \in U \), and \( U \) is a possibly unbounded set on the real line. Additionally, integration over \( u \) in the formation of the actual test statistic is governed by \( \phi(u) \geq 0 \), where \( \int_U \phi(u) = 1 \). Thus, one can control not only the range of \( u \), but also the weights attached to different values of \( u \), so that more weight can be attached to important tail events, for example. We also consider tests based on an analogous conditional confidence interval version of the above measure. Namely, \[ E \left( \left( F_k(\underline{u}|Z^t, \theta_k^1) - F_k(\bar{u}|Z^t, \theta_k^1) \right) - (F_0(\bar{u}|Z^t, \theta_0) - F_0(\underline{u}|Z^t, \theta_0)) \right)^2, \] where \( \underline{u} \) and \( \bar{u} \) are “lower” and “upper” bounds on the confidence interval to be evaluated. For notational simplicity, in the sequel we focus on conditional forecast interval comparison, and set \( \underline{u} = -\infty \) and \( \bar{u} = u \). For example, we say that model 1 is more accurate than model 2, if \[ E \left( \left( F_1(u|Z^t, \theta_1^1) - F_0(u|Z^t, \theta_0) \right)^2 - \left( F_2(u|Z^t, \theta_2^1) - F_0(u|Z^t, \theta_0) \right)^2 \right) < 0. \] This measure defines a norm and it implies a standard goodness of fit measure.

Another measure of distributional accuracy available in the literature is the Kullback-Leibler Information Criterion, KLIC (see e.g. White, 1982, Vuong, 1989, Fernandez-Villaverde and Rubio-Ramirez, 2004, Amisano and Giacomini, 2007, and Kitamura, 2004). According to the KLIC approach, we should choose Model 1 over Model 2 if \[ E(\log f_1(y_{t+1}|Z^t, \theta_1^1) - \log f_2(y_{t+1}|Z^t, \theta_2^1)) > 0. \]\(^4\)With a slight abuse of notation, in this section the subscript 0 denotes the ”true” conditional distribution model, rather than the benchmark model; and the subscript 1 thus now denotes the benchmark model.
The KLIC is a sensible measure of accuracy, as it chooses the model which on average gives higher probability to events which have actually occurred. The drawback is the KLIC approach cannot be easily generalized to compare conditional intervals.

The hypotheses of interest are formulated as:

\[ H_0 : \max_{k=2,\ldots,K} \left( \mu_1^2(u) - \mu_k^2(u) \right) \leq 0 \]

versus

\[ H_A : \max_{k=2,\ldots,K} \left( \mu_1^2(u) - \mu_k^2(u) \right) > 0, \]

where \( \mu_k^2(u) = E\left( \left( 1\{y_{t+1} \leq u\} - F_k(u|Z^t, \theta^i) \right)^2 \right), k = 1, \ldots, K. \) Note that for any given \( u, \)
\( E(1\{y_{t+1} \leq u\}|Z^t) = Pr(y_{t+1} \leq u|Z^t) = F_0(u|Z^t, \theta_0). \) Thus, \( 1\{y_{t+1} \leq u\} - F_k(u|Z^t, \theta^i) \) can be interpreted as an “error” term associated with computation of the conditional expectation under \( F_k. \)

The statistic is:

\[ Z_P = \max_{k=2,\ldots,K} Z_{P,u,\tau}(1, k), \quad (7) \]

with

\[ Z_{P,u}(1, k) = \frac{1}{\sqrt{P}} \sum_{t=R}^{T-1} \left( \left( 1\{y_{t+1} \leq u\} - F_1(u|Z^t, \hat{\theta}_{1,t}) \right)^2 - \left( 1\{y_{t+1} \leq u\} - F_k(u|Z^t, \hat{\theta}_{k,t}) \right)^2 \right), \]

where, as usual, \( R + P = T, \) and \( \hat{\theta}_{k,t} \) can be either a recursive or a rolling estimator. The limiting distribution of (7) is established in Proposition 1(a) in Corradi and Swanson (2006a), who also suggest how to construct valid bootstrap critical values, for both the recursive and the rolling estimation cases.

### 3.2 The Case of Unknown Distribution Function

There are cases in which the distribution function is not known in closed form. This problem typically arises when the variable we want to predict is generated by highly nonlinear dynamic models. Very important examples are Dynamic Stochastic General Equilibrium (DSGE) Models, which generally cannot be solved in closed form (see Bierens, 2007, for a discussion of different ways of approximating DSGEs). Since the seminal papers by Kydland and Prescott (1982), Long and
Plosser (1983) and King, Plosser and Rebelo (KPR: 1988a,b), there has been substantial attention given to the problem of reconciling the dynamic properties of data simulated from DSGE models, and in particular from real business cycle (RBC) models, with the historical record. A partial list of advances in this area includes: (i) the examination of how RBC simulated data reproduce the covariance and autocorrelation functions of actual time series (see e.g. Watson, 1993); (ii) the comparison of DSGE and historical spectral densities (see e.g. Diebold, Ohanian and Berkowitz, 1998); (iii) the evaluation of the difference between the second order time series properties of vector autoregression (VAR) predictions and out-of-sample predictions from DSGE models (see e.g. Schmitt-Grohe, 2000); (iv) the construction of Bayesian odds ratios for comparing DSGE models with unrestricted VAR models (see e.g. Chang, Gomes and Schorfheide, 2002 and Fernandez-Villaverde and Rubio-Ramirez, 2004); (v) the comparison of historical and simulated data impulse response functions (see e.g. Cogley and Nason, 1995); (vi) the formulation of “Reality” bounds for measuring how close the density of an DSGE model is to the density associated with an unrestricted VAR model (see e.g. Bierens and Swanson, 2000); and (vii) loss function based evaluation of DSGE models (see e.g. Schorfheide, 2000).

The papers cited above evaluate the ability of a given DSGE model to reproduce a particular characteristic of the data. Corradi and Swanson (2007b) use a Reality Check approach to evaluate DSGEs in terms of their ability to match (with historical data) the joint distribution of the variables of interest, and provide an empirical application in terms of the comparison of several variants of the stochastic growth model of Christiano (1988). As the distribution function is not known in closed form, we replace it with its simulated counterpart.

To keep notation simple, as above, we consider the case of confidence intervals, setting $u = -\infty$, and $u = \infty$. Hereafter, $F$ represents the joint distribution of a variable of interest, say $Y_t$ (e.g. output growth and hours worked). The hypotheses are:

$$H_0 : \max_{k=2, \ldots, K} \left( \left( F_0(u; \theta_0) - F_1(u; \theta_k^1) \right)^2 - \left( F_0(u; \theta_0) - F_k(u; \theta_k^1) \right)^2 \right) \leq 0$$

$$H_A : \max_{k=2, \ldots, K} \left( \left( F_0(u; \theta_0) - F_1(u; \theta_k^1) \right)^2 - \left( F_0(u; \theta_0) - F_k(u; \theta_k^1) \right)^2 \right) > 0.$$ 

Thus, under $H_0$, no model can provide a better approximation of the joint CDF than model 1. In order to test $H_0$ versus $H_A$, the relevant test statistic is $\sqrt{T}Z_{T,S}$, where

$$Z_{T,S} = \max_{k=2, \ldots, K} \sqrt{T}Z_{k,T,S}(u),$$

13
\[
Z_{k,T,S}(u) = \frac{1}{T} \sum_{t=1}^{T} \left( 1\{Y_t \leq u\} - \frac{1}{S} \sum_{n=1}^{S} 1\{Y_{1,n}(\hat{\theta}_{1,T}) \leq u\} \right)^2 \\
- \frac{1}{T} \sum_{t=1}^{T} \left( 1\{Y_t \leq u\} - \frac{1}{S} \sum_{n=1}^{S} 1\{Y_{k,n}(\hat{\theta}_{k,T}) \leq u\} \right)^2,
\]

and \(Y_{k,n}(\hat{\theta}_{k,T})\) represents simulated counterparts of \(Y_t\) (i.e. the variables simulated under model \(k\) at simulation \(n\), using the estimated parameters \(\hat{\theta}_{k,T}\)). Heuristically, if \(S\) grows sufficiently fast with respect to \(T\), then \(\frac{1}{S} \sum_{n=1}^{S} 1\{Y_{k,n}(\hat{\theta}_{k,T}) \leq u\}\) can be treated as the "true" distribution of the data simulated under model \(k\). Broadly speaking, we are comparing different DSGE models, on the basis of their ability to match a given simulated joint CDF with that of the historical data. As we are comparing joint CDFs, there is no real predictive accuracy testing story here, and thus this is an in-sample test.

When constructing the bootstrap counterpart of \(Z_{k,T,S}\), we need to distinguish between the case in which \(T/S \to 0\) and that in which \(T/S \to \delta \neq 0\). Whenever \(T/S \to 0\), simulation error is asymptotically negligible, and thus there is no need to resample the simulated observations. In this case, the bootstrap statistic is given by \(\max_{k=2,\ldots,K} \sqrt{T}Z^*_{k,T,S}(u)\), where

\[
Z^*_{k,T,S}(u) = \frac{1}{T} \sum_{t=1}^{T} \left( 1\{Y^*_t \leq u\} - \frac{1}{S} \sum_{n=1}^{S} 1\{Y_{1,n}(\hat{\theta}^*_{1,T}) \leq u\} \right)^2 \\
- \frac{1}{T} \sum_{t=1}^{T} \left( 1\{Y^*_t \leq u\} - \frac{1}{S} \sum_{n=1}^{S} 1\{Y_{k,n}(\hat{\theta}^*_{k,T}) \leq u\} \right)^2.
\]

On the other hand, whenever \(T/S \to \delta \neq 0\), then simulation error contributes to the limiting distribution. In this case, one has to additionally resample the simulated observations, and thus \(Y_{1,n}(\hat{\theta}^*_{1,T})\) and \(Y_{k,n}(\hat{\theta}^*_{k,T})\) in (9) should be replaced by \(Y^*_{1,n}(\hat{\theta}^*_{1,T})\) and \(Y^*_{k,n}(\hat{\theta}^*_{k,T})\). In both cases, the validity of bootstrap critical values is been established in Proposition 2 of Corradi and Swanson (2007b).
4 Predictive Evaluation Based on Distribution of Loss

4.1 Outline

Central Banks and financial institutions have regular access to panels of forecasts for key macroeconomic variables that are made by professional forecasters. A leading example is the Survey of Professional Forecasts (SPF), which is compiled at the Philadelphia Federal Reserve Bank. Using this dataset, much focus has centered on how to combine predictions (see e.g. Capistran and Timmerman (2009)) and how to assess forecast rationality (see e.g. Elliott, Komunjjer and Timmermann (2008)). With regard to forecast combination, Capistran and Timmermann (2009), as well as Elliott and Timmermann (2004, 2005), estimate combination weights by minimizing a given loss function, ensuring that the weights converge to those minimizing expected loss. Wallis (2005) proposes combining forecasts using a finite mixture distribution, and Smith and Wallis (2009) suggest the use of simple averages. With regard to rationality assessment, Elliott, Komunjjer and Timmermann (2008) test whether forecasters taking part in the SPF are rational for some parameterization of a flexible loss function. This is clearly an important approach when testing for rationality. However, in many instances, users have a given loss function, and only assess the accuracy of available forecasts under their own loss. Here, we take the loss function as given, and discuss predictive combination and accuracy assessment of datasets such as the SPF. However, this is done via analysis of cumulative loss distributions rather than synthetic measures of loss accuracy such as mean square error and mean absolute error.

More specifically, the objective is to introduce an alternative criterion for predictive evaluation which measures accuracy via examination of the quantiles of the expected loss distribution. The criterion is based on comparing empirical CDFs of predictive error loss, using the principle of stochastic dominance. The heuristic argument underpinning our approach is that the preferred model is one for which the error loss CDF is stochastically dominated by the error loss CDF of every competing model, at all evaluation points. In this sense, a model that has smaller quantiles at all regions of the loss distribution is selected, rather than a model that minimizes a single criterion, such as the mean square error. If a model is not strictly dominated, then our approach allows us to pinpoint the region of the loss distribution for which one model is preferred to another.

Applications for which the criterion is designed include: generic predictive accuracy testing; forecast model selection; and forecast combination. For example, in the context of the SPF, a
panel of \( N_t \) forecasts for a given variable are made by professionals at each point in time, \( t \). Both the number of individuals taking part in the survey, as well as the specific individuals generally change, from period to period. In this context, the criterion can be applied as follows. Assume that objective is to select and combine forecasts from the SPF. A set of rules, including for example, the simple mean or median across all forecasters, and quantile based weighted combinations across forecasts are defined. Then, the loss function of the forecast errors implied by the rules are evaluated using tests based on the stochastic dominance criterion.

4.2 Set-Up

In each period \( t \), we have a panel of \( N_t \) forecasts. The objective is to choose among \( k \) possible combinations of the available forecasts, under a given loss function, \( g(\cdot) \). In order to allow for frequent possible entry and exit into the panel, the combinations are simple rules, which are applied each period, regardless of the composition of the panels. Examples are: (i) simple average, (ii) simple average over a given range, such as the 25th-75th percentiles, or (iii) assigning different weights to different interquantile groups from the panel, such as a weight of 0.75 for the average over the 25th-75th percentile and 0.125 for the average over the first and last quartiles.

Define \( e_{i,t} = y_t - y_{t,h,i}^f \), \( i = 1, \ldots, k \), to be the forecast error associated with the \( h \)-step ahead prediction constructed using combination \( i \). Let \( g_{i,t} = g(e_{i,t}) \), where \( g(\cdot) \) is a generic loss function. Also, let \( F_{g,i}(x) \) be the empirical distribution of \( g(e_{i,t}) \) evaluated at \( x \), and let \( \hat{F}_{g,i,T}(x) \) be its sample analog, i.e.

\[
\hat{F}_{g,i,T}(x) = \frac{1}{T} \sum_{t=1}^{T} 1 \{ g(e_{i,t}) \leq x \}.
\]

The hypotheses of interest are:

\[
H_0 : \max_{i > 1} \inf_{x \in \mathcal{X}} (F_{g,1}(x) - F_{g,i}(x)) \geq 0
\]

versus

\[
H_A : \max_{i > 1} \inf_{x \in \mathcal{X}} (F_{g,1}(x) - F_{g,i}(x)) < 0.
\]

For the sake of simplicity suppose that \( k = 2 \). If \( F_{g,1}(x) - F_{g,2}(x) \geq 0 \) for all \( x \), then the CDF associated with rule 1 always lies above the CDF associated with rule 2. Then, heuristically, \( g(e_{1,t}) \) is (first order) stochastically dominated by \( g(e_{2,t}) \) and rule 1 is the preferred combination. This is
because all of the quantiles of \( g(e_{1,t}) \) are smaller than the corresponding quantiles of \( g(e_{2,t}) \). More formally, for a given \( x \), suppose that

\[
F_{g,1}(x) = \theta_1 \text{ and } F_{g,2}(x) = \theta_2,
\]

then we choose rule 1 if \( \theta_1 > \theta_2 \). This is because \( x \) is the \( \theta_1 \)-quantile under \( F_{g,1} \) and the \( \theta_2 \)-quantile under \( F_{g,2} \) and, as \( \theta_1 > \theta_2 \), the \( \theta_2 \)-quantile under \( F_{g,1} \) is smaller than under \( F_{g,2} \). Thus, for all evaluation points smaller than \( x \), \( g(e_{1,t}) \) has more probability mass associated with smaller values than \( g(e_{2,t}) \) does.

It follows that if we fail to reject the null, rule 1 is selected. On the other hand, rejection of the null does not imply that rule 1 should be discarded. Instead, further analysis is required in order to select a rule. First, one needs to discriminate between the cases for which the various CDFs do not cross, and those for which they do cross. This is accomplished by proceeding sequentially as follows. For all \( i \neq j, i, j = 1, \ldots, k \), sequentially test

\[
H_{i,j}^0 : \sup_{x \in X} (F_{g,i}(x) - F_{g,j}(x)) \leq 0
\]

versus its negation. Eliminate rule \( i \), if \( H_{i,j}^0 \) is not rejected. Otherwise, retain rule \( i \). There are two possible outcomes.

I: If there is a rule which is stochastically dominated by all other rules, we eventually discard all the “dominating” rules and remain with only the dominated one. This is always the case when no CDFs cross; but also clearly occurs in cases when various CDFs cross, as long as the dominated CDF cross no other CDF.

II: Otherwise, we remain with a subset of rules, all of which have crossing CDFs, and all of which are stochastically dominated by the eliminated rules.

Note that the logic underlying the outlined sequential procedure is reminiscent of the idea underlying the Model Confidence Set approach of Hansen, Lunde and Nason (2010), in which the worst models are eliminated in a sequential manner, and one remains with a set of models that are roughly equally good, according to the given evaluation criterion.

In the case where there are crossings, further investigation is needed. In particular, in this case, some rules are clearly dominant over certain ranges of loss, and are dominated over others. At this point, one might choose to plot the relevant CDFs, and examine their crossing points. Then, one has to make a choice. For example, one can choose a rule which is dominant over small values of \( x \).
and is dominated over large values of $x$. This is the case in which one is concerned about making larger losses than would be incurred, were the other rule used, in a region where losses are large; while not being concerned with the fact that they are making larger losses, relative to those that would be incurred, were the other rule used, when losses are relatively small. Needless to say, one can also use a model averaging approach over the various survivor rules.

4.3 Statistic

In order to test $H_0$ versus $H_A$ construct the following statistic:

$$L_{g,T} = -\max_{i > 1} \inf_{x \in X} \sqrt{T} \left( \hat{F}_{g,1,T}(x) - \hat{F}_{g,i,T}(x) \right),$$

where $\hat{F}_{g,j,T}(x)$, $j \geq 1$, is defined above; and where the negative sign in front of the statistic ensures that the statistic does not diverge under the null hypothesis. On the other hand, in order to test $H_0^I$, we instead suggest the following statistic,

$$L_{i,j,g,T} = -\sup_{x \in X} \sqrt{T} \left( \hat{F}_{g,i,T}(x) - \hat{F}_{g,j,T}(x) \right).$$

In the context of testing for stochastic dominance, Linton, Maasoumi and Whang (2005) construct critical values via subsampling. Here we instead use the “m out of n” bootstrap. Proceed to construct critical values as follows:

(i) We have $T$ observations. Set $\Upsilon < T$.

(ii) Draw $b$ blocks of length $l$, where $bl = \Upsilon$. One block consists, simultaneously, of draws on the actual data as well as the rule based combination forecasts. Thus, if there are two rules, say, and the block length is 5, then a “block” consists of a 3x1 vector of length 5. This yields one bootstrap sample, which is used to construct a bootstrap statistic,

$$L_{g,\Upsilon}^* = -\max_{i > 1} \inf_{x \in X} \sqrt{\Upsilon} \left( \hat{F}_{g,1,\Upsilon}(x) - \hat{F}_{g,i,\Upsilon}(x) \right),$$

where

$$\hat{F}_{g,i,\Upsilon}(x) = \frac{1}{\Upsilon} \sum_{t=1}^{\Upsilon} 1 \left\{ g^* (e_{i,t}) \leq x \right\}$$

and

$$g^* (e_{i,t}) = g \left( y_{t}^* - y_{t,h,i}^* \right)$$

Note: The basic difference between subsampling and “m out of n” bootstrap is that in the latter case we resample overlapping blocks.
(iii) Construct $B$ bootstrap statistics and then compute their empirical distribution. The sample statistic is then compared against the percentile of this empirical distribution.
5 References


21


