

ISSN 1440-771X  
ISBN 0 7326 1082 6

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**Working Paper 2/2001**

**April 2001**

**DEPARTMENT OF ECONOMETRICS  
AND BUSINESS STATISTICS**

**MONASH**  
UNIVERSITY  
AUSTRALIA

# The Importance of Common Cyclical Features in VAR Analysis: A Monte-Carlo Study\*

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March 2001

## Abstract

Despite the commonly held belief that aggregate data display short-run comovement, there has been little discussion about the econometric consequences of this feature of the data. We use exhaustive Monte-Carlo simulations to investigate the importance of restrictions implied by common-cyclical features for estimates and forecasts based on vector autoregressive models. First, we show that the “best” empirical model developed without common cycle restrictions need not nest the “best” model developed with those restrictions. This is due to possible differences in the lag-lengths chosen by model selection criteria for the two alternative models. Second, we show that the costs of ignoring common cyclical features in vector autoregressive modelling can be high, both in terms of forecast accuracy and efficient estimation of variance decomposition coefficients. Third, we find that the Hannan-Quinn criterion performs best among model selection criteria in simultaneously selecting the lag-length and rank of vector autoregressions.

**Keywords:** Reduced rank models, model selection criteria, forecasting, variance decomposition.

**JEL Classification:** C32, C53.

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\***Acknowledgments:** We thank George Athanasopoulos for excellent research assistance. The authors are responsible for any errors in this paper. João Victor Issler acknowledges the support of CNPq-Brazil and PRONEX. Farshid Vahid is grateful for financial support provided by the Monash University Research Fund Grant No. 22.097.007, and CNPq-Brazil.

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## 1. Introduction

In this paper we argue that short-run dynamic restrictions should be taken seriously in vector autoregressive (VAR) modelling. We focus on common-cycle restrictions because of their importance in macroeconomics. Common cyclical movements in detrended economic variables have been so prevalent that they have acquired the status of “stylized facts.” Lucas (1977) states that the main regularities observed in cyclical fluctuations of economic time series are in their *comovement*. In empirical studies, common cycles have been shown to be a feature of a variety of macroeconomic data sets. For example, Campbell and Mankiw (1989) find a common cycle between consumption and income for most G-7 countries. Engle and Kozicki (1993) find common international cycles in GNP data for OECD countries. Using US data, Issler and Vahid (2001) find common cycles for macroeconomic aggregates, and Engle and Issler (1995) and Carlino and Sill (1998) find common cycles for sectoral and regional outputs respectively. Like most applied macroeconomic research in the last fifteen years, these studies have investigated common-cyclical features using vector-autoregressive (VAR) or vector error-correction (VEC) models.

We investigate the importance of restrictions implied by common-cyclical features for forecasts, impulse-response functions, and variance-decomposition analysis of economic time-series based on VAR models. VAR models are most useful for short term forecasting, and short run dynamic restrictions can improve short-run forecasts. However, relative to the considerable effort that has been spent on examining the importance of cointegration restrictions in VAR models (see, among others, Engle and Yoo 1987, Clements and Hendry 1995, and Lin and Tsay 1996), no work has examined the effects of short-run restrictions. As shown by Engle and Yoo, the forecasting gains of imposing long-run constraints are realized only when the forecast horizon becomes large. In fact, in their simulations, the unconstrained VAR models produce better short-horizon forecasts than the VEC models. Because forecasting uncertainty at long horizons can be large, time-series models are generally most useful for forecasting over short horizons. Hence, imposing short-run constraints might be a way of improving the effectiveness of time-series models at horizons where they are most useful.

Incorporating common-cycle restrictions can reduce the number of free parameters of a VAR model and help achieve parsimony, more than cointegrating restrictions can. For example, when dealing with post-war quarterly data, and a VAR with three variables and eight lags, there are seventy five mean parameters to be estimated from about two hundred data points on each variable. If the three-variable system has one known cointegrating vector, the number of free parameters falls from seventy five to sixty nine when estimating a VEC model. Common-cyclical features show more potential in reducing the number of conditional-mean parameters. If the three variables in the VEC model share one common cycle, then the number of mean parameters falls from sixty nine to twenty seven.

We assess the effects of common-cyclical features on VAR models using Monte-Carlo simulations. The focus here is on the accuracy of multi-step ahead out-of-sample forecasts, as well as the accuracy of estimates of impulse-response functions and variance-decomposition of forecast errors. We design the simulations so that the results would be relevant for an applied macroeconomist estimating a relatively large number of parameters using a limited number of data points. To that end, we consider a variety of Data Generating Processes (DGPs) and sample sizes, that are similar to the “typical” data sets that applied researchers encounter in practice.

VAR models with common cycles fall into the general category of reduced-rank multivariate models<sup>1</sup>. We can represent these models in a reduced-rank regression framework by  $z_t = \Phi x_t + \varepsilon_t$ , where  $z_t$  contain the  $n$ -series of interest,  $x_t$  contains  $p$  lags of  $z_t$  (and possibly error-correction terms), and  $\varepsilon_t$  is a multivariate white-noise process. The matrix  $\Phi$  is not full rank, reflecting the fact that there are linear combinations of  $z_t$  that are white noise. If common cycles are a true feature of the data, and if the lag order of the VAR (VEC) is known to be  $p$ , then theory tells us that the estimate of  $\Phi$  with the correct rank-restrictions imposed must be more efficient than the unrestricted estimate of  $\Phi$  (see Ahn and Reinsel 1988). Even so, researchers may be reluctant to incorporate these parameter restrictions because of the asymmetric consequences of over versus under-parametrization. Because the true rank of  $\Phi$  is not known, it may seem wiser to live with a possibly inefficient unconstrained model rather than with a misspecified inconsistent model. We argue here that the cost of ignoring common-cycle restrictions is more than the efficiency loss in estimating  $\Phi$ . We show that, if only full-rank models are considered, the lag length chosen by the usual model-selection criteria is severely misspecified. Standard criteria may find too small a lag length in reduced-rank VARs simply because this is the only possible way available to achieve parsimony. For such misspecified models one cannot tell from theory what the consequences of incorporating rank restrictions will be.

An alternative to the usual model-selection criteria is to choose the lag length and the rank of the VAR simultaneously. Lütkepohl (1993, page 202) presents a set of model-selection criteria that can be used for that purpose, and we refer to this set as  $IC(p, r)$ . Our simulations reveal that, when the true DGP is a reduced-rank VAR model, the lag length chosen by the standard model-selection criteria (which we refer to as  $IC(p)$ ) can be quite different from that chosen when rank and order are selected simultaneously. Standard model-selection criteria that place a strong penalty on over-parameterization, such as the Schwarz or Hannan-Quinn criteria, may choose too small a lag-length when the true model has common cycles. However, they improve if the rank order is simultaneously selected with the lag length. We find strong evidence in favor of Hannan-Quinn criterion for choosing the correct lag and rank order overall. Regarding the Akaike information criterion, we observe that its tendency to choose

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<sup>1</sup>Classic references on reduced-rank VAR’s include Velu, Reinsel and Wickern (1986), Ahn and Reinsel (1988), and Tiao and Tsay (1989).

an over-parameterized model when the lag order and rank are selected simultaneously is accentuated relative to the case when only the lag length is selected.

Users of VAR models are often interested in forecasts, rather than the true lag order. Hence, we compare models based on their forecasting accuracy measures. For horizons up to sixteen periods ahead, using several measures of forecasting accuracy, we find that the forecasts produced by the reduced-rank models selected by  $IC(p, r)$  are generally superior to those produced by the models selected by  $IC(p)$ . Indeed, on average, if the Hannan-Quinn criterion is used to select lag order and rank, the cumulative accuracy of one to four-step-ahead forecasts can be improved by up to 20%. This sizable effect illustrates the potential gain associated with considering common-cycle restrictions at the model selection stage. For variance decompositions, reduced-rank models selected by  $IC(p, r)$  only do better when samples are large (more than 200 observations)<sup>2</sup>.

The outline of the paper is as follows. Section 2 states the reduced-rank restrictions that common cyclical fluctuations impose on the parameters of VAR models, and presents the model-selection criteria for reduced-rank models. Section 3 describes our Monte-Carlo design. Section 4 presents the simulation results. Section 5 presents a small empirical example using coincident and leading business-cycle series. Finally, Section 6 presents the main conclusions of the paper, as well as a suggestion for further research.

## 2. Common cycles in VAR models

As in most applied macroeconomic research, we assume that the objective is to build a time series model for the growth rate of a vector of  $n$  economic variables. We denote the levels of these variables at time  $t$  by  $Y_t$ , their logarithms by  $y_t$ , and their growth rates (i.e. the first difference of the logarithm of  $Y_t$ ) by  $\Delta y_t$ . We make the reasonable assumption that  $\Delta y_t$  is stationary, add the simplifying assumption that  $\Delta y_t$  has mean zero (without any loss of generality), and start with the Wold representation of  $\Delta y_t$ , i.e.

$$\Delta y_t = C(L) \varepsilon_t, \tag{2.1}$$

where  $C(L) = \sum_{j=0}^{\infty} C_j L^j$  is a matrix polynomial in the lag operator and  $C_0 = I_n$ . From the work of Beveridge and Nelson (1981) and Stock and Watson (1988), it is possible to decompose the log-level series  $y_t$  into common trends and cycles (we refer to this as the Beveridge-Nelson-Stock-Watson – or BNSW – decomposition). Using the identity  $C(L) = C(1) + \Delta C^*(L)$ , ignoring the initial value of  $y_0$ , and integrating both sides of (2.1) we get<sup>3</sup>:

$$y_t = C(1) \sum_{j=1}^t \varepsilon_j + C^*(L) \varepsilon_t$$

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<sup>2</sup>Notice that in the textbook example Lütkepohl (1993, pp. 202-3) the selected lag was identical whether or not the rank was also chosen, and in that case, he observed that the forecasts and variance decompositions were quite similar for the reduced rank and full rank models.

<sup>3</sup>See Stock and Watson (1988) or Vahid and Engle (1993) for more details.

$$= \tau_t + c_t, \quad (2.2)$$

where  $\tau_t = C(1) \sum_{j=1}^t \varepsilon_j$  and  $c_t = C^*(L) \varepsilon_t$  represent the trend and cyclical components of  $y_t$  respectively. In the BNSW decomposition, the  $n$  variables in  $y_t$  are decomposed into  $n$  random-walk components (stochastic trends) and  $n$  stationary components (stochastic cycles). If  $C(1)$  has rank  $n - q$  ( $q > 0$ ), the stochastic trends in  $y_t$  can be characterized as linear combinations of only  $n - q$  common random walks, in which case  $y_t$  is said to be cointegrated, or to have common stochastic trends, with  $q$  linearly independent cointegrating vectors (see Engle and Granger, 1987). If  $C^*(L)$  has rank  $r$  ( $r < n$ ), then the stochastic cycles in  $y_t$  can be characterized as linear combinations of  $r$  common stochastic cycles, with  $n - r$  linearly independent cofeature vectors (see Vahid and Engle, 1993). In this paper, we investigate the costs of ignoring this singularity in the stochastic cycles  $c_t$ .

For ease of exposition we assume that there is no cointegration in the system ( $q = 0$ )<sup>4</sup>, in which case the appropriate model for  $\Delta y_t$  will be a VAR, i.e.,

$$\begin{aligned} \Delta y_t &= A_1 \Delta y_{t-1} + \dots + A_p \Delta y_{t-p} + \varepsilon_t \\ &= \begin{bmatrix} A_1 & \dots & A_p \end{bmatrix} \begin{bmatrix} \Delta y_{t-1} \\ \vdots \\ \Delta y_{t-p} \end{bmatrix} + \varepsilon_t \\ &= \Phi x_t + \varepsilon_t, \end{aligned} \quad (2.3)$$

where  $\Phi = \begin{bmatrix} A_1 & \dots & A_p \end{bmatrix}$  and  $x_t = [\Delta y'_{t-1}, \dots, \Delta y'_{t-p}]'$ . If there are  $r$  common stochastic cycles in  $y_t$ , then  $C^*(L)$  in (2.2) has rank  $r$ , and the  $n \times np$  matrix  $\Phi$  must have rank  $r$  ( $< n$ ). This shows that VAR models with common-cyclical features among their variables fall into the general category of reduced-rank regression models.

Common-cycle constraints imply important restrictions for the impulse-response functions, variance-decompositions, and multi-step ahead forecasts. The existence of  $r$  common cycles implies that there are  $n - r$  independent linear combinations of  $\Delta y_t$  that are white noise. Thus, from (2.1), all matrices  $C_i$ , for  $i = 1, 2, \dots$ , must have rank  $r$ . These matrices  $C_i$ , which are usually normalized to be consistent with orthogonal errors, form the basis of the impulse-response functions and the forecast-error variance decompositions. For example, when they are post-multiplied by the Choleski factor of the variance-covariance matrix of  $\varepsilon_t$ , they yield the so-called orthogonalized impulse responses. Hence, it becomes clear that the presence of common cycles implies that the impulse responses of different variables to the same shocks will be linearly dependent. Therefore, if the objects of interest are the impulse responses (or variance decompositions of the forecast errors) of  $\Delta y_t$ , then common-cycle restrictions can have important repercussions for efficient estimation.

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<sup>4</sup>If there is cointegration, then the appropriate error-correction term has to be added to the right-hand side of (2.3), which, of course, will add another source of uncertainty in model building. Here, we abstain from dealing with it, focusing only on the consequences of ignoring common-cyclical components of VAR models.

A similar argument applies to forecasts of  $\Delta y_t$  at horizon  $h$ . These can be recursively calculated from:

$$\Delta y_{t+h}^f = A_1 \Delta y_{t+h-1}^f + \dots + A_p \Delta y_{t+h-p}^f = \Phi x_t^f, \quad (2.4)$$

where the superscript  $f$  stands for forecasts which use information up to period  $t$ , and actual variables are used instead of forecasts on the right-hand side where available. Since common cycles imply that the matrix  $\begin{bmatrix} A_1 & \dots & A_p \end{bmatrix} = \Phi$  has reduced rank, equation (2.4) clearly shows that they will also imply that the forecasts of  $\Delta y_t$  at any horizon will be linearly dependent. Again, if forecasting is the objective of the multivariate model building exercise, common-cycle restrictions will have important consequences.

### 2.1. Model selection criteria for reduced-rank models

Our motivation is to build VAR-based models for  $\Delta y_t$  that can be used for forecasting, impulse-response or variance-decomposition analysis. A critical step in constructing these models is the selection of the lag length of the VAR. Model-selection criteria are often used in practice, and in principle they are useful because they do not favor any specific model against others (see the discussion in Granger, King and White, 1995). However, model selection criteria may choose different lag orders, depending on whether or not we allow for reduced-rank parameter matrices in the VAR model. We investigate the performance of widely used selection criteria when (i) only the lag length is selected, and (ii) when the lag length and rank order are jointly selected. We also consider the alternative strategy of choosing the lag length with model-selection criteria and then choosing the rank by the common-cycle test recommended by Vahid and Engle (1993). We then compare our results, so as to recommend a strategy for empirical work.

Following Lütkepohl (1993), we focus on the Akaike (AIC), Hannan-Quinn (HQ) and Schwarz (SC) criteria for the simultaneous selection of lag and rank orders in VAR models. The lag order  $p$  and the number of common cycles  $r$  (i.e. the rank of  $\Phi$ ), can be simultaneously chosen to minimize one of the following model selection criteria,

$$AIC(p, r) = \ln \left| \hat{\Sigma}_\varepsilon(p, r) \right| + \frac{2}{T} \times r \times (np + n - r) \quad (2.5)$$

$$HQ(p, r) = \ln \left| \hat{\Sigma}_\varepsilon(p, r) \right| + \frac{2 \ln \ln T}{T} \times r \times (np + n - r) \quad (2.6)$$

$$SC(p, r) = \ln \left| \hat{\Sigma}_\varepsilon(p, r) \right| + \frac{\ln T}{T} \times r \times (np + n - r) \quad (2.7)$$

where  $n$  is the dimension of the system,  $r$  is the rank of VAR model,  $p$  is the number of lagged differences in the model,  $\hat{\Sigma}_\varepsilon(p, r)$  is the estimated variance-covariance matrix of the errors of the VAR model with  $p$  lags and rank  $r$ , and  $T$  is the number of observations.

For full-rank models ( $r = n$ ), the model selection criteria in (2.5)-(2.7) collapse to the usual criteria, which we call  $AIC(p)$ ,  $HQ(p)$ , and  $SC(p)$ . Calculating them is straightforward, since full-rank models can be estimated, equation by equation, using OLS. However, the

estimation of reduced rank models is not straightforward, and an easier way to calculate these model-selection criteria is to use the following well-known lemma:

**Lemma 2.1.** *Under the assumption that  $\Phi = \begin{bmatrix} A_1 & \dots & A_p \end{bmatrix}$  has rank  $r$ , the minimum of  $\ln \left| \frac{1}{T} \sum_{t=1}^T \varepsilon_t \varepsilon_t' \right|$  is*

$$\ln \left| \frac{1}{T} \sum_{t=1}^T \Delta y_t \Delta y_t' \right| + \sum_{i=n-r+1}^n \ln(1 - \lambda_i),$$

where  $\lambda_1 < \lambda_2 < \dots < \lambda_n$  are the sample squared canonical correlations between  $\Delta y_t$  and the set of regressors  $x_t$ . The sample squared canonical correlations are the eigenvalues of

$$\left( \sum_{t=p+1}^T \Delta y_t \Delta y_t' \right)^{-1} \left( \sum_{t=p+1}^T \Delta y_t x_t' \right) \left( \sum_{t=p+1}^T x_t x_t' \right)^{-1} \left( \sum_{t=p+1}^T x_t \Delta y_t' \right).$$

**Proof.** See Tso (1981).

This lemma implies that, after dropping the common term  $\ln \left| \frac{1}{T} \sum_{t=1}^T \Delta y_t \Delta y_t' \right|$  in (2.5)-(2.7), the model-selection criteria can be expressed in terms of the eigenvalues ( $\lambda_i$ ) as:

$$AIC(p, r) = \sum_{i=n-r+1}^n \ln(1 - \lambda_i(p)) + \frac{2}{T} \times r \times (np + n - r) \quad (2.8)$$

$$HQ(p, r) = \sum_{i=n-r+1}^n \ln(1 - \lambda_i(p)) + \frac{2 \ln \ln T}{T} \times r \times (np + n - r) \quad (2.9)$$

$$SC(p, r) = \sum_{i=n-r+1}^n \ln(1 - \lambda_i(p)) + \frac{\ln T}{T} \times r \times (np + n - r). \quad (2.10)$$

Hence, for fixed  $p$ , the model-selection criteria for any rank can be easily calculated after the relevant eigenvalues are computed. These eigenvalues can be easily calculated using any statistical program which has a canonical correlation procedure<sup>5</sup>. Notice that, for fixed  $T$  and  $n$ , the model-selection criteria in (2.8)-(2.10) depend only on the lag length  $p$  and on the rank  $r$  of the VAR model.

### 3. Monte-Carlo design

If samples are “large”, our intuition tells us that ignoring the common-cycle restrictions will not be very harmful. This is based on the expectation that with “large” samples, lag-order selection is likely to be unambiguous and parameter estimates will be precise, so that the reduced rank constraints will be (approximately) true for the estimated parameters, even when they are not imposed at the estimation stage. Hence, the estimated models with or

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<sup>5</sup>Two examples include SAS and STATA. Alternatively one can use any matrix program such as GAUSS, or modify any of the plethora of computer programs that use this lemma to calculate the Johansen cointegration test statistics (see chapter 20 of Hamilton(1994)).



without common-cycle restrictions will be so close that their results for forecasting, impulse-response, and variance-decomposition analysis will be very similar.

This intuition should not, however, be carried over to the case of “small” samples. Indeed, efficiency gains are potentially much more relevant when samples are small and degrees of freedom are scarce. In this context, selecting the lag order after assuming full rank can yield a completely different result from selecting lag order and rank simultaneously. We investigate this issue using 1000 simulations of 100 reduced-rank VARs based on either 100 or 200 observations. We tabulate results for cases when only the lag length is chosen, and when the rank and lag length are chosen simultaneously.

To make the presentation manageable, we only present results for three-dimensional VARs<sup>6</sup>. Models that consider the real side of the economy are often three-dimensional. For example, King et al (1991) estimate a VAR including output, consumption, and investment in order to test the real-business-cycle model of King, Plosser and Rebelo (1988). Issler and Ferreira (1998) use a VAR in output, labor, and capital inputs to estimate long-run elasticities of the aggregate production function.

The first parameter we set in the Monte-Carlo design is the lag length  $p$ . It is chosen in order to allow for the possibility of either under or over-parameterization of the VAR model. Lütkepohl (1985) uses a DGP with a true lag order of 1 in his simulations, making under-parameterization virtually impossible. This favors model-selection criteria which heavily penalize over-parameterization, e.g., the Schwarz criterion. Nickelsburg (1982) sets the true lag order to four in some of his simulations, but the maximum lag allowed for in the estimation is also set to four. This makes over-parameterization impossible, favoring liberal criteria such as the AIC. To avoid both problems, we choose the true lag order of four and allow for models of up to lag eight.

The properties of estimated VARs are only invariant to scaling the variance-covariance matrix of the errors by a constant. However, the following lemma shows that in order to cover the entire space of reduced-rank VAR processes of order  $p$ , one can fix the variance-covariance matrix of the error to be the identity matrix without any loss of generality.

**Lemma 3.1.** *Any arbitrary full rank linear transformation of a reduced-rank VAR, generates another VAR with the same rank.*

**Proof.** See Vahid and Issler (1999).

This lemma allows one to transform a reduced-rank VAR with a non-diagonal covariance matrix into another VAR with the same rank and an identity covariance matrix. This means that in the Monte Carlo analysis, if we consider the entire space of reduced-rank models and

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<sup>6</sup>An earlier version of the present paper (Vahid and Issler 1999) includes results for six-dimensional VARs.

compare different methods with a measure that is invariant to linear transformations, then we can fix the variance covariance matrix of the errors of the DGP to be the identity matrix without loss of generality.

However, an exhaustive Monte-Carlo study over the entire model space is infeasible. It is customary, as in Lütkepohl (1985), to choose several sets of eigenvalues for the companion matrix<sup>7</sup> of the VAR, and to choose arbitrary parameter matrices which give rise to those eigenvalues, and then to average the results over all these DGPs. Although the results generated from such a design strategy might be useful for general time-series analysts, they are unsuitable for economists who work with aggregate macroeconomic data. This is because the cyclical structure of macroeconomic aggregates can be quite weak, especially for systems which do not contain a monetary sector. For example, the system<sup>8</sup>  $R^2$  for King et al.'s (1991) VEC model of US per-capita income, consumption, and investment is 0.44, whereas the system  $R^2$  for 160 out of the 200 DGPs in Lütkepohl (1985) are above 0.5, and 96 of these are greater than 0.8. Since this paper is intended primarily for applied macroeconomists, a design which gives too much weight to models with a high system  $R^2$  would be inappropriate.

Here, we start with a “typical” macroeconometric study in order to select the DGP and the system  $R^2$  associated with it. The data set used for choosing our parameter values is the same as in King et al.(1991)<sup>9</sup>. For the three-variable system, we first fitted rank one and rank two VARs of order four to the first-differences of the logarithms of US per-capita private income, consumption, and investment over the period 1947.1 to 1988.4, which resulted in estimates for the  $A_t$ 's and for  $E(\varepsilon_t \varepsilon_t')$ . Then we determined the parameter values for our DGPs by randomly making 100 draws from the estimated 95% confidence regions for the parameters. For all cases, we have been careful to verify that all of these randomly drawn DGPs satisfy the stationarity conditions for vector autoregressions<sup>10</sup>. By choosing our DGPs from this “plausible” subset of the parameter space, we believe that our results are directly relevant for applied macroeconomists. The median of the system  $R^2$  measure for our generated three-variable DGPs is between 0.5 and 0.6, with less than 5% larger than 0.7 and none greater than 0.8.

The Monte-Carlo procedure can be summarized as follows. Using each of our 100 DGPs, we generate 1000 samples (once with 100, and again with 200 observations), record the lag length chosen by traditional (full-rank)  $AIC(p)$ ,  $HQ(p)$  and  $SC(p)$  measures, and the lag length and rank order chosen by model selection criteria stated in (2.8)-(2.10). In all cases, to

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<sup>7</sup>The *companion matrix* of a VAR( $p$ ) is the coefficient matrix of its VAR(1) representation. The condition for a VAR( $p$ ) to be stationary is that all of the eigenvalues of its companion matrix are inside the unit circle.

<sup>8</sup>The system  $R^2$  is a generalization of the single-equation  $R^2$  for multivariate models. See the Appendix for its definition.

<sup>9</sup>King et al.(1991) chose a lag length of eight for their three variable model and a lag length of six for their six variable model. They chose these lag lengths on a-priori grounds, without any reference to data.

<sup>10</sup>The range of the absolute value of the maximum eigenvalues of the tri-variate rank-one DGPs is (0.49, 0.87). For the tri-variate rank-two DGPs this range is (0.63, 0.92).

reduce the impact of initial values on simulated series, we generated 1000 observations, but only used the last 124 or 224 observations in the analysis. We take the model chosen using each  $IC(p)$  criterion and compare it with the model chosen using the corresponding  $IC(p, r)$  criterion. Each pair of chosen models is compared with respect to (i) their out-of-sample forecasting accuracy up to 16 periods ahead; and (ii) their mean-squared-error in estimating variance decompositions of forecast errors for selected horizons up to 16 periods ahead.

We explain the measures we chose to compute the accuracy of forecasts, impulse responses and variance decompositions, before stating our results.

### 3.1. Measuring forecast accuracy

Appropriate evaluation of forecasts depends on the specific use that the forecasts are needed for, i.e., the “loss function” of the user. The fact that we have applied economists as our target audience does not suggest that we should evaluate the forecasts of alternative models in any specific way. A macroeconomist who models the growth rate of income, consumption and investment, might in fact be interested in the growth rates of income, *savings* and investment, or she might be interested in forecasting the levels, based on the growth rates. Therefore, it is important to evaluate the forecasting performance of different models on the basis of measures that are invariant to linear transformation of forecasts, at one horizon, or across different horizons. One measure that satisfies this invariance property is the *generalized forecast error second moment (GFESM)* introduced by Clements and Hendry (1993). *GFESM* is the determinant of the expected value of the outer product of the vector of stacked forecast errors of all future times up to the horizon of interest. For example, if forecasts up to  $h$  quarters ahead are of interest, this measure will be:

$$GFESM = \left| E \begin{pmatrix} \tilde{\varepsilon}_{t+1} \\ \tilde{\varepsilon}_{t+2} \\ \vdots \\ \tilde{\varepsilon}_{t+h} \end{pmatrix} \begin{pmatrix} \tilde{\varepsilon}_{t+1} \\ \tilde{\varepsilon}_{t+2} \\ \vdots \\ \tilde{\varepsilon}_{t+h} \end{pmatrix}' \right|$$

where  $\tilde{\varepsilon}_{t+h}$  is the  $n$ -dimensional forecast error at horizon  $h$  of our  $n$ -variable model. It is obvious that this measure is invariant to elementary operations that involve different variables, and also to elementary operations that involve the same variable at different horizons. In our Monte-Carlo, the above expectation is evaluated for every model, by averaging over the simulations.

We also consider two popular measures of forecasting accuracy. The first is the determinant of the mean squared forecast error matrix at different horizons ( $|MSFE|$ ), and the second is the trace of the mean squared forecast error matrix ( $TMSFE$ ). The determinant of the *MSFE* is invariant to elementary operations on the forecasts of different variables at a single horizon, but it is not invariant to elementary operations on the forecasts across different

horizons. The trace of the mean squared forecast error matrix is not invariant to either of these transformations.

There is one complication associated with simulating 100 different DGPs. Simple averaging across different DGPs is not appropriate, because the forecast errors of different DGPs do not have identical variance-covariance matrices. Lütkepohl (1985) normalizes the forecast errors by their true variance-covariance matrix in each case to get i.i.d. observations. Unfortunately, this would be a very time consuming procedure for a measure like *GFESM*, which involves stacked errors over many horizons. Instead, for each information criterion, we calculate the percentage change in forecasting measures, comparing the full-rank models selected by  $IC(p)$ , with the reduced-rank models chosen by  $IC(p, r)$ . This procedure is done at every iteration for every DGP, and the final results are then averaged.

### **3.2. Precision of impulse-response and variance-decomposition estimates**

Although many applied studies that use VAR models focus on impulse-response functions and variance-decompositions of forecast errors, most simulation studies in the literature simply focus on forecast comparisons. However, the impulse-response functions and variance-decomposition of forecast errors differ from multi-step forecasts of VAR models because they depend on the variance-covariance matrix of system errors as well as being non-linear functions of the mean parameters. Given this added dimension to the problem, one cannot expect *a priori* to get similar results to the forecasting exercise.

Moreover, there are a few issues that are specific to the analysis of impulse-response functions and variance decompositions. First, errors have to be orthogonal for results to be meaningful. As is well known, there are several techniques that yield orthogonal errors. Here, we orthogonalize our shocks by the Choleski decomposition of the variance-covariance matrix, since this method is the most popular. It is well known that the Choleski method is not invariant to the ordering of the variables in the VAR. Hence, we consider all possible orderings of the variables in the system, and the presented results are the average over all these orderings. Second, for a three-variable system, there are nine impulse-response and variance-component coefficients in each horizon. In order to report results in a compact way, the mean-squared errors of each is computed for the rank-restricted, and the unrestricted VAR models. Then, the percentage improvement in MSE of the restricted model relative to that of the unrestricted model is computed for each of these coefficients. Finally, for each horizon, the mean percentage improvement across all coefficients for that horizon is computed. In order to keep the size of our tables down to a minimum, we only report the variance-decomposition results, since results for impulse responses are similar.

## **4. Monte-Carlo simulation results**

The main objectives of our study are to address the following:

1. Whether a model chosen with an  $IC(p, r)$  criterion is just a reduced rank version of a model chosen with the corresponding  $IC(p)$  criterion, or they can be non-nested;
2. Whether differences in the models chosen by these two classes of model selection criteria lead to major differences in forecasting accuracy; and,
3. Whether differences in the models chosen by these two classes of model selection criteria lead to major differences in the accuracy of their estimated impulse-response and variance-decomposition coefficients.

In addition, we also compare models where rank is chosen by statistical testing (sequential LR tests) with those where rank is chosen by model-selection criteria. Finally, we investigate the relative performance of different model-selection criteria in choosing the best forecasting model.

First, however, we assume that the lag-length and rank order are known, and we compare the accuracy of forecasts and variance decomposition coefficients for the estimated unrestricted and reduced rank VAR models. Although these results do not have any direct implication for applied work because they do not include lag-rank uncertainty, they serve as a useful benchmark for a better understanding of other results.

#### 4.1. The benchmark case when the lag-rank order is known

As a natural benchmark, we compare the accuracy of the forecasts and variance decompositions of estimated unrestricted VARs of correct lag-length, with those of estimated reduced rank VARs of correct lag and rank order. Any differences between reduced-rank and full-rank VAR models reflect the efficiency gains resulting from imposing the rank restrictions.

Table 1 shows the percentage improvement in different measures of forecast accuracy and in the mean-squared error (MSE) of variance-decomposition coefficients, when we allow for rank deficiencies. Three interesting conclusions can be made from this table. First, all measures of forecast comparison tell us that the correct rank restrictions lead to sizable improvements in forecasts over short horizons. The determinant and the trace of the MSE matrix become very close to zero, and the GFESM, which is a cumulative measure, flattens out after quarter 8. Second, the improvements in forecasts due to rank restrictions are more pronounced in smaller samples. All measures of improvements in forecast accuracy are almost twice as large when the sample size is 100, than when the sample size is 200. Third, the pattern of improvements in the variance decompositions is not similar to that of the forecasts. In particular, the one-step-ahead forecast decomposition estimates are significantly worse, when the true rank restrictions are imposed. Noting that the one-step-ahead variance decomposition estimates are only functions of the estimated variance covariance matrix of the errors, and in particular that they are *ratios* of the elements of the Choleski factor of this matrix, we conclude that the efficiency gains in estimating the VAR parameters do not lead

to better estimates of these ratios. However, the gains in estimating the mean parameters are so large that there are sizable improvements in variance decompositions for all horizons longer than one.

These results quantify the size of the efficiency gains predicted from econometric theory when the lag-length and the rank of VAR models are both known. Although they serve as a benchmark, these gains are irrelevant for empirical studies, because lag lengths and rank orders must be estimated beforehand.

## 4.2. Selection of lag and rank order

Table 2.a shows the frequency of lag-order selection in 1000 simulations of 100 trivariate VAR(4) models with rank 1. Each of *AIC*, *HQ* and *SC* are considered, firstly assuming full rank, and secondly when rank and lag orders are determined simultaneously. The top half of this table corresponds to a sample size of 100, and the bottom half corresponds to samples of 200 observations. Table 2.b shows the analogous frequencies when the true DGP is a trivariate VAR(4) of rank 2.

These tables confirm that selecting the lag and rank order jointly, can lead to a model which is of higher lag-order than the model chosen with conventional (full rank) criteria. For example, the top half of Table 2.a shows that for samples of 100 observations, the modal choice of all three criteria is a VAR(1), with *AIC* choosing the true lag of 4 only 14 percent of the time. The other two criteria choose a VAR(4) with a frequency of less than 1 percent. However, when the lag and rank are chosen simultaneously, there is a large reduction in the number of times that the VAR(1) is chosen, regardless of the criterion used. Furthermore, the frequency of choosing the correct lag increases significantly. In both Tables 2.a and 2.b, *AIC* chooses the correct lag and rank more often than the other two criteria, with *HQ* being a close second. The modal choice of the Schwarz criterion stays at a VAR(1), even with 200 observations.

Two points are worth noting. First, even when the criteria choose the wrong lag-length, they are likely to choose the correct rank. The only exception is *SC* when the true rank is 2 and there are only 100 observations. This suggests that common cycles can be detected even if the wrong lag-length is chosen. This is plausible, because the property that a linear combination of variables has no correlation with the past (the necessary and sufficient condition for common cycles), is unrelated to what those cycles are and whether they are correctly specified. The second point is that once one chooses lag length and rank simultaneously, the probability of choosing the correct lag length increases for all three criteria, and the probability of their overestimating the lag length also increases. Although the chance of overpredicting the lag length remains quite small for *HQ* and *SC*, it shoots up to more than 10 percent (and even to approximately 20 percent in the rank 1 model) for *AIC*.

### 4.3. Forecasts

Tables 3.a and 3.b show the percentage improvement in the measures of forecast accuracy when the lag and rank are chosen simultaneously. A general conclusion is that there are no differences between forecasts beyond 8 periods, and most of the advantage of looking for common cycles is in forecasting one to four periods ahead. These tables show that there are non-trivial gains from considering reduced rank models for short-run forecasting. The *GFESM* and *MSFE* measures, although not as pronounced as our benchmark case, show sizeable improvements for all criteria at horizons 1 to 4. The trace of the *MSFE* improves remarkably for *HQ* and *SC* when lag and rank are chosen simultaneously.

The results in Tables 3.a and 3.b also show which model selection criterion produced models with best forecasting performance on average at each horizon. For each horizon, the criterion that provided the best forecast performance according to *TMSFE* is indicated by superscript *b* in the *TMSFE* column, and the criterion that provided the worst forecast performance is indicated by superscript *w*<sup>11</sup>. Not surprisingly, we observe that when the DGP is relatively parsimonious (i.e. when it has rank 1) and sample size is small, *AIC* chooses models with the worst forecasting performance. However, in all other cases, the Schwarz criterion chooses models that on average produce the worst forecasts. The remarkable result is that *HQ* produces the best forecasting models in almost all cases. In the few cases where models chosen by *HQ* criterion are not the best, they are a very close second best.

Our results do not support the conclusion made by Lütkepohl (1985) that *SC* leads to best forecasting models, and this leads us to believe that Lütkepohl's conclusion must be an artifact of the Monte Carlo design used in his paper. Our results show that even though the forecast performance of models chosen by *SC* improves significantly when we use this criterion to choose lag and rank simultaneously, they are far from being the best. For the best forecasting performance, our simulations make a strong case for using the *HQ* criterion to choose lag and rank simultaneously.

### 4.4. Selecting rank by testing vs. by model-selection criteria

An alternative strategy for selecting VARs with common-cyclical features was proposed in Vahid and Engle (1993). It consists of choosing the lag length by *IC*(*p*) and then performing sequential LR tests to determine the rank. Table 4 compares the forecasting performance of VAR models selected by this procedure with those selected by *IC*(*p*). As in Table 3, we report the percentage improvement of forecasts of reduce-rank models over their unrestricted VAR counterparts, making the results in these two tables directly comparable. Table 4 shows that testing for rank, conditional on lag length, produces forecasting improvement over full-rank VARs. However, only in the case of *AIC* with 100 observations are these improvements larger

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<sup>11</sup>Notice that this information is obtained by comparing the forecasting performance across different criteria, and cannot be inferred from the numerical entries of Table 3.

than those one would obtain when lag and rank are selected simultaneously. This suggests that in small samples, the strategy of choosing lag length by  $AIC$  and then choosing rank by a sequence of LR tests leads to good forecasting models. However, given our results in Table 3a, model selection by  $HQ(p, r)$  seems to be a superior strategy for building forecasting models.

#### 4.5. Variance-decomposition results

The percentage improvements of the estimated forecast-error variance decomposition coefficients are presented in Table 5. It is noticeable that there are virtually no significant gains at any horizon when the sample size is 100 observations. This is in sharp contrast with our benchmark case reported in Table 1, where there were gains of 20 to 74 percent for all reported horizons other than 1. This may be due to the fact that the variance contributions are ratios of estimated parameters. Although the allowance for rank restrictions improves the parameter estimates in a direction that leads to better forecasts, these improvements lead to worse estimates of the variance *ratios* when samples are small. When the sample size is 200, the quality of variance-decompositions based on models chosen by  $IC(p, r)$  is far superior to that of models chosen by  $IC(p)$ .

### 5. Empirical Example

The empirical analysis of the three-variable system that generates our simulated DGPs is discussed in Issler and Vahid (2001). There, we obtained a percentage reduction of 30.3% for the one-step ahead  $|MSFE|$  using the reduced-rank model. Here, we investigate a larger VAR which can be potentially useful for business-cycle analysis.

The “pulse” of the US economy is monitored every month by observing fluctuations in four “coincident” variables, which are: 1) Personal income less transfer payments; 2) Index of industrial production; 3) Number of employees on nonagricultural payrolls; and 4) Manufacturing and trade sales<sup>12</sup>. In this section, we build a time-series model to forecast these coincident variables. It is well-known that other series lead the coincident series and therefore help in forecasting them. See, for example, Stock and Watson (1989) or Zellner and Hong (1989). We follow Zellner and Hong and use measures of growth in real money balances and in the real rate of return of stocks as two leading indicator variables<sup>13</sup>. Since the coincident variables are not cointegrated (see Stock and Watson 1989), this constitutes a six-variable VAR for all of these log-differenced series, although our primary focus will be in forecasting the log-differences of the four coincident series alone<sup>14</sup>.

<sup>12</sup>The mnemonics for these variables in the DRI database are GMYXPQ, IP, LPNAG and MTQ respectively.

<sup>13</sup>We use M2 deflated by producer price index as a measure of real balances (FM2/PWFSA in DRI) and S&P500 index deflated by the same price index (FSPCOM/PWFSA in DRI) in computing stock returns.

<sup>14</sup>We obtain similar results qualitatively when we consider forecasts of all six variables together. But forecasting stock returns is not an objective of our empirical study.



To make the empirical example conformable to our simulation study, we use monthly data from 1980:01 to 2000:07, a total of 247 observations. We develop our models on the basis of the first 199 observations, leaving the last 48 observations for out-of-sample forecast evaluation. To be consistent with our simulation results, we select models using the Hannan-Quinn criterion. The full-rank version of the  $HQ$  criterion selects one lag for the six variable VAR. However, if we use the lag-rank version of  $HQ$ , the selected lag order is two and the selected rank is three. Therefore, we compare the performance of a full-rank VAR(1) with that of a reduced-rank VAR(2) in forecasting the four coincident variables. The estimated models are used to generate 48, 24, 16, and 12 non-overlapping one, two, three, and four-step ahead forecasts respectively, with results reported in Table 6. The out-of-sample forecasting results conform to those in our simulation study. For all four short-run horizons, the reduced-rank model outperforms the full-rank model, with the largest percentage improvement of 23.8% for the  $|MSFE|$  at horizon four. This is a sizable improvement in forecasting accuracy.

It is informative to compare the univariate processes for individual variables implied by the estimated full rank VAR(1) model with those implied by the estimated rank 3 VAR(2) model. A full-rank 6 variable VAR(1) implies univariate  $ARMA(6, q)$  processes for each of the variables, where  $q$  is less than or equal to 5 and the autoregressive polynomials for all variables are identical. A rank 3 VAR(2) model implies univariate  $ARMA(6, q)$  processes for individual variables, where  $q$  is less than or equal to 6 and autoregressive polynomials for all variables are identical<sup>15</sup>. All 6 roots of the implied autoregressive polynomial of the estimated full rank VAR(1) model turned out to be real, whereas there was a pair of complex conjugate roots among the 6 roots implied by the estimated rank 3 VAR(2) model. Because complex roots give rise to oscillatory components, and coincident and leading indicators are supposed to measure the cyclical oscillations in the economy, this gives further evidence in favor of the estimated reduced rank VAR(2) model.

Our conclusion from the empirical example is that if econometricians are interested in coincident and leading indicators, they should consider reduced-rank models at the model selection stage.

## 6. Conclusion

This paper argues that in multivariate macroeconomic modelling, the stylized fact that “macroeconomic aggregates move together over the business cycle” should be taken seriously. Time series macro-econometric models provide useful forecasts for short horizons (1 to 8 periods). It is for these horizons that our Monte-Carlo study shows substantial gains in forecast accuracy if reduced-rank structures are allowed for. These gains are higher if the uncertainty about the lag length is assumed away, but they are still non-trivial in the more realistic case in which lag length and rank are chosen simultaneously.

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<sup>15</sup>This is a direct implication of Vahid (1999).

The results of our Monte-Carlo analysis of model selection criteria that simultaneously select lag length and rank order can be summarized as follows. The tendency of *AIC* to choose overparameterized models is worsened (particularly in small samples) when simultaneously choosing the rank and the lag length. Hence, we conclude that *AIC* should not be used for this purpose in small samples. On the other hand, the tendency of *HQ* and *SC* criteria to choose an underparameterized model is somewhat remedied when they are allowed to pick the rank and lag-length simultaneously. The *SC* criterion, however, still selects severely underparameterized models.

Contrary to previous literature that compares forecasts of VAR models selected by alternative model selection criteria, we find no support for the claim that *SC* leads to models that produce the best forecasts. We attribute this previous finding to the simple Monte Carlo design with short lag structures, that previous researchers have used. Indeed, in our simulations, the models selected by the Schwarz criterion produced worse forecasts than models chosen by the other two criteria. This was particularly evident in our simulations for the six-dimensional system<sup>16</sup>. Therefore, we conclude that *SC* should not be used for model selection in high dimensional time series models, regardless of whether a reduced-rank structure is allowed for or not. Instead, we recommend the Hannan-Quinn criterion, which generally leads to models with the best forecast performance, especially when it is used for simultaneously choosing lag length and rank order.

Our analysis shows that there is a tension between efficiently estimating the mean parameters while allowing for reduced-rank structures, and efficiently estimating variance parameters. For samples of 100 observations we find no gains from reduced rank structures in estimating variance decomposition coefficients. This result is reversed for larger samples of 200 observations. For the latter, there are non-trivial benefits in considering reduced-rank models in the estimation of variance contributions. Even though we have used all possible orderings of variables in performing our variance decompositions, we qualify our findings in that the accuracy of the latter may not be invariant to the method of orthogonalizing the errors.

Finally, it should be stressed that the message of this paper is that short-run restrictions are likely to be more important than cointegrating restrictions, for forecasting at the business-cycle horizons. Here, we have only considered common-cycle restrictions because of their important macroeconomic implications. We leave the investigation of possible gains resulting from other restrictions, such as block exogeneity restrictions, codependence and other types of rank restrictions, for future research.

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<sup>16</sup>The results for the six-dimensional system are not reported here to save space. They are reported in an earlier version of the current paper (Vahid and Issler 1999).

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## Appendix

### A. System $R^2$ and signal-to-noise ratio

In a multiple regression with stochastic regressors and i.i.d. errors,  $y = X\beta + \varepsilon$ , the limiting signal-to-noise ratio ( $snr$ ) can be defined as:

$$snr = \frac{\beta' \lim_{T \rightarrow \infty} E \left( \frac{X'X}{T} \right) \beta}{\sigma_\varepsilon^2}, \quad (\text{A.1})$$

where  $E(\varepsilon\varepsilon') = \sigma_\varepsilon^2 \cdot I$ , and the proportion of the variation of dependent variable explained by the model, i.e. the population  $R^2$ , is:

$$R^2 = \frac{\beta' \lim_{T \rightarrow \infty} E \left( \frac{X'X}{T} \right) \beta}{\sigma_\varepsilon^2 + \beta' \lim_{T \rightarrow \infty} E \left( \frac{X'X}{T} \right) \beta} = \frac{snr}{(1 + snr)}.$$

Since the asymptotic variance of  $\sqrt{T}(\hat{\beta} - \beta)$  is  $AVAR(\hat{\beta}) = \sigma_\varepsilon^2 \left( \lim_{T \rightarrow \infty} E \left( \frac{X'X}{T} \right) \right)^{-1}$ , we can write (A.1) as:

$$snr = \beta' \left( AVAR(\hat{\beta}) \right)^{-1} \beta. \quad (\text{A.2})$$

Consider now a  $VAR(p)$ :

$$y_t = A_1 y_{t-1} + \dots + A_p y_{t-p} + \varepsilon_t. \quad (\text{A.3})$$

The analogous measure of  $snr$  for it is:

$$snr = \beta' \left( \Sigma \otimes \Omega^{-1} \right) \beta \quad (\text{A.4})$$

where  $\beta = \text{vec}(A)$ ,  $A = \begin{bmatrix} A_1 & \dots & A_p \end{bmatrix}$ ,  $E(\varepsilon_t \varepsilon_{t-j}') = \Omega$ , and:

$$\Sigma = \begin{pmatrix} \Gamma_0 & \Gamma_1 & \dots & \Gamma_{p-1} \\ \Gamma_1' & \Gamma_0 & \dots & \Gamma_{p-2} \\ \dots & \dots & \dots & \dots \\ \Gamma_{p-1}' & \Gamma_{p-2}' & \dots & \Gamma_0 \end{pmatrix},$$

where  $\Gamma_j = E(y_t y_{t-j}')$ . Notice that  $\Sigma$  is completely determined by  $(A, \Omega)$  via the Yule-Walker equations<sup>17</sup>. After some algebra, it can be shown that (A.4) is equal to:

$$snr = \beta' \left( \Sigma \otimes \Omega^{-1} \right) \beta = \text{trace} \left( \Gamma_0 \Omega^{-1} \right) - n.$$

Using this last result, one can then define the system  $R^2$  to be:

$$R^2 = \frac{\text{trace}(\Gamma_0 \Omega^{-1}) - n}{1 + \text{trace}(\Gamma_0 \Omega^{-1}) - n}.$$

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<sup>17</sup>See Hamilton (1994) Chapter 10, Lütkepohl (1993) Chapter 1, or Reinsel (1993) Chapter 2.

**Table 1: Percentage improvement in different forecast accuracy measures and in the MSE of forecast-error variance decompositions when the true rank restrictions are imposed**

horizon	True rank is one				True rank is two			
( <i>h</i> )	GFESM	MSFE	TMSFE	Var. Dec.	GFESM	MSFE	TMSFE	Var. Dec.
Sample size 100								
1	22.22	22.22	1.97	-10.68	12.08	12.08	0.98	-7.67
4	60.41	8.66	1.72	73.93	34.56	5.02	0.96	20.69
8	70.54	1.70	1.39	54.03	42.57	1.32	0.77	21.38
12	72.34	0.46	1.03	52.41	44.66	0.45	0.59	21.76
16	72.86	0.21	0.81	52.06	45.26	0.25	0.47	21.88
Sample size 200								
1	11.22	11.22	1.14	-6.00	6.55	6.55	0.60	-3.55
4	29.53	3.80	0.88	94.05	17.80	2.32	0.52	27.19
8	32.97	0.52	0.64	66.65	20.72	0.46	0.37	27.01
12	33.37	0.11	0.45	63.96	21.19	0.07	0.27	27.19
16	33.47	0.05	0.34	63.48	21.25	0.06	0.20	27.24

**Table 2.a: Frequency of lag ( $p$ ) and lag-rank ( $p, r$ ) choice by different criteria when the true models are (4,1)**

Selected lag	1			2			3			4			5			6			7			8		
Selected rank	1	2	3	1	2	3	1	2	3	1 <sup>T</sup>	2	3	1	2	3	1	2	3	1	2	3	1	2	3
Number of observations=100																								
$AIC(p)$	–	–	57.0	–	–	13.1	–	–	12.6	–	–	14.0	–	–	2.0	–	–	0.7	–	–	0.3	–	–	0.3
$AIC(p, r)$	10.8	2.5	0.4	7.4	2.0	0.1	14.4	2.4	0.1	32.7	3.4	*	8.3	1.1	*	5.0	0.6	*	3.8	0.4	*	4.0	0.5	*
$HQ(p)$	–	–	92.9	–	–	4.7	–	–	1.7	–	–	0.7	–	–	*	–	–	*	–	–	0	–	–	0
$HQ(p, r)$	39.2	1.9	0.2	13.3	0.3	*	17.0	0.1	*	24.3	0.1	*	2.4	*	0	0.7	*	0	0.3	0	0	0.1	0	0
$SC(p)$	–	–	99.6	–	–	0.4	–	–	*	–	–	*	–	–	0	–	–	0	–	–	0	–	–	0
$SC(p, r)$	73.8	0.4	*	10.7	*	0	8.4	0	0	6.6	0	0	0.1	0	0	*	0	0	0	0	0	0	0	0
Number of observations=200																								
$AIC(p)$	–	–	25.9	–	–	10.7	–	–	20.0	–	–	40.0	–	–	2.7	–	–	0.5	–	–	0.2	–	–	*
$AIC(p, r)$	2.2	0.7	0.1	3.3	0.8	*	12.1	1.8	*	56.4	4.1	0.1	9.1	0.8	*	4.1	0.3	0	2.3	0.1	0	1.6	0.1	0
$HQ(p)$	–	–	80.1	–	–	7.8	–	–	7.2	–	–	4.9	–	–	*	–	–	0	–	–	0	–	–	0
$HQ(p, r)$	16.1	0.6	0.1	8.9	0.1	*	20.7	0.1	0	51.3	*	0	1.9	0	0	0.2	0	0	*	0	0	*	0	0
$SC(p)$	–	–	98.6	–	–	1.0	–	–	0.3	–	–	0.1	–	–	0	–	–	0	–	–	0	–	–	0
$SC(p, r)$	49.4	0.1	*	11.1	*	0	17.1	0	0	22.3	0	0	0.1	0	0	*	0	0	0	0	0	0	0	0

**Table 2.b: Frequency of lag ( $p$ ) and lag-rank ( $p, r$ ) choice by different criteria when the true models are (4,2)**

Selected lag	1			2			3			4			5			6			7			8		
Selected rank	1	2	3	1	2	3	1	2	3	1	2 <sup>T</sup>	3	1	2	3	1	2	3	1	2	3	1	2	3
Number of observations=100																								
$AIC(p)$	–	–	19.9	–	–	10.2	–	–	21.3	–	–	41.3	–	–	4.6	–	–	1.5	–	–	0.7	–	–	0.5
$AIC(p, r)$	1.1	4.9	1.0	1.0	4.7	0.6	2.5	15.5	1.2	4.3	43.7	1.8	1.2	7.0	0.3	0.8	3.1	0.1	0.7	1.8	*	0.9	1.6	*
$HQ(p)$	–	–	64.1	–	–	13.1	–	–	12.7	–	–	9.9	–	–	0.1	–	–	*	–	–	0	–	–	0
$HQ(p, r)$	8.6	19.6	1.9	5.0	8.1	0.2	8.1	14.8	0.1	10.5	20.8	*	1.1	0.6	0	0.4	0.1	0	0.1	*	0	0.1	*	0
$SC(p)$	–	–	93.2	–	–	5.1	–	–	1.5	–	–	0.2	–	–	0	–	–	0	–	–	0	–	–	0
$SC(p, r)$	30.3	30.6	1.2	9.5	4.8	*	9.4	4.3	*	7.9	1.9	0	0.2	*	0	*	0	0	*	0	0	0	0	0
Number of observations=200																								
$AIC(p)$	–	–	3.3	–	–	2.7	–	–	16.3	–	–	72.2	–	–	4.3	–	–	0.8	–	–	0.2	–	–	0.1
$AIC(p, r)$	0.1	0.6	0.2	0.1	0.9	0.1	0.3	10.2	0.7	0.9	72.3	2.5	0.2	7.1	0.2	0.1	2.0	*	0.1	0.8	*	0.1	0.4	*
$HQ(p)$	–	–	27.9	–	–	9.6	–	–	23.3	–	–	39.2	–	–	*	–	–	0	–	–	0	–	–	0
$HQ(p, r)$	1.3	7.5	0.6	0.9	4.7	*	3.4	20.0	*	4.7	56.2	*	0.2	0.4	0	*	*	0	*	0	0	*	0	0
$SC(p)$	–	–	74.4	–	–	10.4	–	–	10.3	–	–	5.0	–	–	0	–	–	0	–	–	0	–	–	0
$SC(p, r)$	9.2	26.9	0.7	4.3	6.8	*	8.2	15.1	0	9.1	19.8	0	*	*	0	*	0	0	0	0	0	0	0	0

Numbers in each cell represent percentage times that the model selection criterion corresponding to that row chose the lag-rank order corresponding to that column in 100,000 simulations (1000 simulations of 100 different DGPs). The true lag-order is identified with superscript  $T$ . A \* corresponds to a non-zero value less than 0.05 percent. Numbers in a row may not add up to 100.0 exactly because of rounding.

**Table 3a: Percentage improvement in different measures of accuracy in forecasts generated by the possibly reduced rank VAR over the full rank VAR chosen by the same model selection criterion when the true models are trivariate (4,1)**

horizon ( $h$ )	AIC			HQ			SC		
	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE
Sample size 100									
1	6.6	6.6	0.0 <sup>w</sup>	6.8	6.8	2.8 <sup>b</sup>	5.3	5.3	1.6
4	10.8	2.3	1.1	16.1	6.1	4.8 <sup>b</sup>	10.9	4.1	3.0 <sup>w</sup>
8	4.0	-1.0	0.0	15.1	-0.3	2.7 <sup>b</sup>	11.0	-0.1	1.7 <sup>w</sup>
12	2.0	-0.6	-0.2 <sup>w</sup>	14.2	-0.2	1.7 <sup>b</sup>	10.7	-0.1	1.1
16	1.0	-0.3	-0.2 <sup>w</sup>	13.7	-0.2	1.2 <sup>b</sup>	10.5	-0.1	0.8
Sample size 200									
1	9.1	9.1	2.0 <sup>b</sup>	11.0	11.0	6.7	8.3	8.3	5.3 <sup>w</sup>
4	22.2	3.2	2.0 <sup>b</sup>	30.8	8.2	7.7	22.5	7.1	6.8 <sup>w</sup>
8	22.1	0.1	1.0 <sup>b</sup>	31.8	0.5	4.4	23.4	0.4	3.9 <sup>w</sup>
12	22.1	0.0	0.7	31.7	0.0	2.8 <sup>b</sup>	23.4	0.0	2.6 <sup>w</sup>
16	22.0	0.0	0.5	31.7	0.0	2.1 <sup>b</sup>	23.3	0.0	1.9 <sup>w</sup>

**Table 3b: Percentage improvement in different measures of accuracy in forecasts generated by the possibly reduced rank VAR over the full rank VAR chosen by the same model selection criterion when the true models are trivariate (4,2)**

horizon ( $h$ )	AIC			HQ			SC		
	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE
Sample size 100									
1	7.6	7.6	0.1	5.9	5.9	2.2 <sup>b</sup>	1.6	1.6	1.4 <sup>w</sup>
4	19.2	2.9	0.5	19.2	6.1	3.9 <sup>b</sup>	10.1	6.1	4.3 <sup>w</sup>
8	20.4	0.1	0.1	19.7	-0.0	2.2 <sup>b</sup>	10.0	-0.0	2.5 <sup>w</sup>
12	20.5	0.0	0.0	19.6	-0.1	1.4 <sup>b</sup>	9.4	-0.0	1.6 <sup>w</sup>
16	20.5	0.1	0.0	19.4	-0.1	1.0 <sup>b</sup>	9.1	-0.1	1.2 <sup>w</sup>
Sample size 200									
1	5.9	5.9	0.7 <sup>b</sup>	6.8	6.8	2.3	8.8	8.8	5.4 <sup>w</sup>
4	15.3	2.0	0.5 <sup>b</sup>	20.5	4.3	2.6	28.7	8.9	6.5 <sup>w</sup>
8	17.1	0.2	0.3 <sup>b</sup>	21.7	0.3	1.5	31.1	0.6	3.7 <sup>w</sup>
12	17.3	0.0	0.2 <sup>b</sup>	21.8	0.0	1.0	31.3	0.1	2.5 <sup>w</sup>
16	17.3	0.0	0.1 <sup>b</sup>	21.7	0.0	1.0	31.2	-0.0	1.8 <sup>w</sup>

GFESM is Clements and Hendry's generalized forecast error second moment measure, |MSFE| is the determinant of the mean squared forecast error matrix, and TMSFE is the trace of the MSFE matrix. Superscripts  $b$  and  $w$  denote respectively the best and the worst forecasting performance across all three information criteria based on TMSFE.



**Table 4a: Percentage improvement in different measures of accuracy in forecasts generated by the possibly reduced-rank VAR model chosen by sequential rank testing over that of the full rank VAR when the true models are trivariate (4,1)**

horizon ( $h$ )	AIC			HQ			SC		
	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE
Sample size 100									
1	8.4	8.4	0.7	3.7	3.7	0.3	3.1	3.1	0.2
4	18.3	2.1	0.5	4.8	0.1	0.1	3.4	-0.0	0.0
8	21.4	0.6	0.4	5.1	0.0	0.0	3.5	-0.0	0.0
12	22.2	0.2	0.3	5.2	0.0	0.0	3.5	-0.0	0.0
16	22.5	0.1	0.3	5.2	0.0	0.0	3.5	-0.0	0.0
Sample size 200									
1	7.1	7.1	0.7	2.7	2.7	0.2	1.8	1.8	0.1
4	17.4	1.9	0.5	4.7	0.2	0.1	1.9	-0.0	0.0
8	19.5	0.3	0.4	5.1	0.1	0.1	2.0	-0.0	0.0
12	19.8	0.1	0.3	5.1	0.0	0.1	2.0	-0.0	0.0
16	20.0	0.1	0.2	5.2	0.0	0.0	2.0	0.0	0.0

**Table 4b: Percentage improvement in different measures of accuracy in forecasts generated by the possibly reduced-rank VAR model chosen by sequential rank testing over that of the full rank VAR when the true models are trivariate (4,2)**

horizon ( $h$ )	AIC			HQ			SC		
	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE	GFESM	MSFE	TMSFE
Sample size 100									
1	7.7	7.7	0.4	2.9	2.9	0.0	0.2	0.2	-0.3
4	21.5	3.0	0.5	7.6	1.0	0.2	0.7	0.2	0.0
8	27.3	1.0	0.5	9.2	0.2	0.2	0.9	-0.0	0.0
12	28.9	0.4	0.4	9.7	0.1	0.1	0.9	0.0	0.0
16	29.6	0.2	0.3	10.0	0.1	0.1	0.9	0.0	0.0
Sample size 200									
1	5.5	5.5	0.5	3.9	3.9	0.3	1.3	1.3	0.1
4	14.6	1.9	0.4	9.8	1.2	0.3	2.9	0.3	0.1
8	17.1	0.4	0.3	11.4	0.2	0.2	3.3	0.1	0.1
12	17.6	0.1	0.2	11.7	0.1	0.1	3.4	0.0	0.0
16	17.7	0.1	0.2	11.7	0.1	0.1	3.4	0.0	0.0

GFESM is Clements and Hendry's generalized forecast error second moment measure, |MSFE| is the determinant of the mean squared forecast error matrix, and TMSFE is the trace of the MSFE matrix.

**Table 5: Percentage improvement in MSE of forecast-error variance decomposition generated by the possibly reduced rank VAR over the full rank VAR chosen by the same model selection criterion**

horizon	True rank is one			True rank is two		
( <i>h</i> )	AIC	HQ	SC	AIC	HQ	SC
Sample size 100						
1	-19.11	-3.63	2.50	-13.59	-6.52	-5.27
4	0.11	2.56	9.11	5.51	2.95	-5.51
8	-7.11	-3.84	3.57	3.66	2.04	-6.00
12	-8.69	-5.32	2.48	3.27	1.67	-6.26
16	-9.25	-5.84	2.10	3.04	1.47	-6.41
Sample size 200						
1	-7.41	13.10	20.99	-4.35	1.13	12.65
4	37.50	51.70	38.33	19.23	25.46	23.22
8	26.12	47.04	33.04	17.97	26.08	26.51
12	23.96	45.62	31.90	17.67	26.01	26.39
16	23.41	45.12	31.51	17.56	25.92	26.23

**Table 6: Forecasting performance of alternative models of coincident variables**

Model	Full-rank VAR(1)		Rank 3 VAR(2)	
Horizon	MSFE	TMSFE	MSFE	TMSFE
1 month ahead	$0.3437 \times 10^{-4}$	0.6241	$0.3107 \times 10^{-4}$	0.5647
2 months ahead	$0.1374 \times 10^{-4}$	0.5932	$0.1156 \times 10^{-4}$	0.5442
3 months ahead	$0.0683 \times 10^{-4}$	0.5534	$0.0651 \times 10^{-4}$	0.5166
4 months ahead	$0.0868 \times 10^{-4}$	0.5643	$0.0701 \times 10^{-4}$	0.4916

Models were chosen using the *HQ* criterion. |MSFE| is the determinant of the mean squared forecast error matrix, and TMSFE is the trace of the MSFE matrix.