



Factor extraction using Kalman filter and smoothing: This is not just another survey[☆]

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ARTICLE INFO

Keywords:

Dynamic factor model
Expectation maximization algorithm
Identification
Macroeconomic forecasting
State-space model

ABSTRACT

Dynamic factor models have been the main “big data” tool used by empirical macroeconomists during the last 30 years. In this context, Kalman filter and smoothing (KFS) procedures can cope with missing data, mixed frequency data, time-varying parameters, non-linearities, non-stationarity, and many other characteristics often observed in real systems of economic variables. The main contribution of this paper is to provide a comprehensive updated summary of the literature on latent common factors extracted using KFS procedures in the context of dynamic factor models, pointing out their potential limitations. Signal extraction and parameter estimation issues are separately analyzed. Identification issues are also tackled in both stationary and non-stationary models. Finally, empirical applications are surveyed in both cases. This survey is relevant to researchers and practitioners interested not only in the theory of KFS procedures for factor extraction in dynamic factor models but also in their empirical application in macroeconomics and finance.

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1. Introduction

In recent decades, dynamic factor models (DFMs) have been widely used to represent comovements within large systems of macroeconomic and financial variables where the cross-sectional dimension is often relatively large compared with the time dimension; see [Stock and Watson \(2017\)](#) for the importance of DFMs in time series

econometrics research. DFMs generally assume the existence of a small number of unobserved factors capturing the comovements in the system, being the main “big data” tool used by empirical macroeconomists during the last 20 years. [Diebold \(2003\)](#) points out that, although DFMs do not “really” analyze big data, they represent a movement of macroeconomics in this direction. [Stock and Watson \(2016b\)](#) also describe the DFM as a “big data” tool.

Two main types of procedures for factor extraction are popular in the related literature. First, many applications consider **factors extracted using non-parametric procedures based on principal components (PCs)** which are computational simple and have well-known asymptotic properties. For example, PCs are consistent under mild conditions and robust to the underlying dependence of common factors and idiosyncratic components, provided that the factors are pervasive and the idiosyncratic dependence is weak; see, for example, [Bai \(2003\)](#) and [Lettau and Pelger \(2020\)](#). As a consequence, PC procedures are very

[☆] Financial support from the Spanish Government Project PID2019-108079GB-C22/AEI/10.13039/501100011033 (MINECO/FEDER) is gratefully acknowledged by Pilar Poncela. Esther Ruiz and Karen Miranda acknowledge financial support from project PID2019-108079GB-C21 (MINECO/FEDER). We are also very thankful for useful comments by Ganics Gergely Akos, Máximo Camacho, Frank Diebold, Laurent Ferrara, Peter Fuleky, Marc Hallin, Siem Jan Koopman, and two anonymous referees. All of them have been a great help to improve the presentation of this survey suggesting several relevant references. Obviously any remaining errors or omissions are our own responsibility.

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popular for factor estimation and several excellent surveys are available in the literature; see [Bai and Ng \(2008\)](#) for a technical survey on the econometric theory for PCs. However, PC procedures do not use all the information available in the data when the common factors are serially dependent and, consequently, they are not efficient. Furthermore, as there is not a particular specification of the dynamic dependence of the factors, one cannot obtain their out-of-sample forecasts.

Second, factors can be extracted using Kalman Filter and Smoothing (KFS) procedures that cast the DFM as a state-space model (SSM). One important feature of KFS procedures is that they open the door to maximum likelihood (ML) estimation of the model parameters. Thus, if the assumed model specification is correct, KFS procedures are efficient for factor extraction. Furthermore, factor extraction based on KFS procedures allows missing data and mixed frequencies to be handled in a simple way; see [Camacho, Perez-Quiros, and Poncela \(2013\)](#) and [Luciani \(2017\)](#) who survey the literature on missing observations and mixing frequencies. Moreover, KFS procedures are of interest in empirical applications because they allow for incorporating restrictions on the factor loadings as, for example, in [Reis and Watson \(2010\)](#) and [Coroneo, Giannone, and Modugno \(2016\)](#) who impose a block structure, or on the idiosyncratic components as in [Luciani \(2015\)](#). They are also attractive because it is possible to perform counterfactual exercises as in, for example, [Luciani \(2015\)](#) or to incorporate seasonal dependencies as in [Alonso, García-Martos, Rodríguez, and Sánchez \(2011\)](#), [Camacho, Lovcha, and Perez-Quiros \(2015\)](#) and [Nieto, Peña, and Saboya \(2016\)](#). In addition, KFS procedures have been extended to account for regime-switching non-linearities; see, for example, [Camacho, Perez-Quiros, and Poncela \(2015, 2018\)](#) and [Camacho, Leiva-Leon, and Perez-Quiros \(2016\)](#).¹ However, KFS procedures also have drawbacks. Historically, their implementation has been numerically challenging when the cross-sectional dimension of the system under analysis is large. Furthermore, KFS procedures require full specification of the dependence of the common and idiosyncratic components. These specifications introduce potential misspecification that is not reflected in the model-based inference.

The literature on factor extraction using KFS has only been partially reviewed; see [Barhoumi, Darné, and Ferrara \(2013\)](#), [Stock and Watson \(2011, 2016b\)](#), and, more recently, [Doz and Fuleky \(2020\)](#). In this paper, we update and complement previous surveys. Although this is, by necessity, a selective review of the literature, the main contribution of this paper is to provide a comprehensive updated summary of the literature on latent common factors extracted using KFS procedures in the context of DFMs, highlighting the potential limitations of these procedures and pointing out open issues that require further research. Signal extraction and parameter estimation issues are separately analyzed. Identification is also tackled

¹ Note that the references mentioned are just an illustration of the interesting problems related to factors extraction that can be easily handled using KFS procedures. Obviously, as we will see, the literature on KFS in the context of DFMs is much more extensive.

in both stationary and non-stationary models. Finally, empirical applications are surveyed in both cases. This survey is relevant to researchers and practitioners interested not only in the theory of KFS procedures for factor extraction in DFMs but also in their empirical applications, mainly in the areas of macroeconomics and finance.

The rest of the paper is organized as follows. Section 2 introduces notation by presenting the SSM and the KFS algorithms for factor extraction. Section 3 deals with the representation of DFMs as SSMs and how factor extraction can be performed in this context. We consider the effect on the extracted factors and their MSEs of assuming that the DFM is exact when it is not. We also consider the effect of the dependence and number of underlying factors and of the cross-sectional dimension on the properties of the extracted factors. In Section 4, we review estimation procedures of the parameters of the DFM and illustrate their performance when extracting the common factors, assuming that the model specification is known. Section 5 deals with model specification. In particular, we describe procedures for determining the number of factors and the lags of factors. In Section 6, we survey empirical applications implementing KFS in the context of stationary DFMs to describe and forecast the future evolution of variables of interest. Section 7 considers DFMs for non-stationary systems. Section 8 concludes the paper with some final remarks.

2. State-space models: Kalman filter and smoothing algorithms and estimation

In this section, we briefly describe SSMs, the algorithms to extract the unobserved states, and the parameter estimators; see [Durbin and Koopman \(2012\)](#) and [Harvey \(1989\)](#) for comprehensive treatments of the SSMs, KFS algorithms, and estimators described in this section.

2.1. State-space models

SSMs were originally developed by control engineers with the attention focused on a set of m unobserved state variables, α_t , that evolve over time and are related to an N -dimensional vector of variables, Y_t , observed at time t , for $t = 1, \dots, T$. An SSM specifies a full parametric model for both Y_t and α_t and can be formulated in a variety of ways. In what follows, we consider the following time-homogeneous linear Gaussian SSM without deterministic terms:

$$Y_t = Z\alpha_t + \varepsilon_t \quad (1)$$

$$\alpha_t = W\alpha_{t-1} + R\eta_t \quad (2)$$

where ε_t is an $N \times 1$ white noise vector with covariance matrix H , and η_t is a $g \times 1$ white noise vector with covariance matrix Q .² The disturbances ε_t and η_t are

² Apart from the already mentioned Markov switching models, there are interesting DFMs in which the system matrices are not time-invariant: for example, the DFMs with time-varying factor loadings of [Eickmeier, Lemke, and Marcellino \(2015\)](#) and [Stock and Watson \(2016b\)](#).

uncorrelated with each other for all t and s , and are uncorrelated with the initial state, α_0 . The system matrices, Z , H , W , R , and Q are $N \times m$, $N \times N$, $m \times m$, $m \times g$, and $g \times g$ non-stochastic.³ Eqs. (1) and (2) are known as the measurement equation and transition equation, respectively. The SSM in Eqs. (1) and (2) is fully specified when additional assumptions about the distribution of the initial state and the distribution of the disturbances are made. Usually, both disturbances and the initial state are assumed to be Gaussian vectors, the latter with mean a_0 and covariance matrix P_0 .

2.2. Kalman filter and smoothing algorithms

The SSM opens the way for the application of the Kalman filter, a recursive procedure for computing the optimal estimator, in the sense that it minimizes the mean square error (MSE), of the state vector at time t based on observations up to and including Y_t . The Kalman filter enables the estimate of the state vector to be continually updated as new observations become available. Furthermore, the Kalman filter provides the basis for one-step-ahead prediction and smoothing, as well as out-of-sample forecasts of the factors. Define $a_t = E(\alpha_t | Y_1, \dots, Y_t)$ and $P_t = E((\alpha_t - a_t)(\alpha_t - a_t)' | Y_1, \dots, Y_t)$. The Kalman filter updating equations are given as follows⁴:

$$a_t = a_{t|t-1} + P_{t|t-1} Z' \Sigma_t^{-1} (Y_t - Za_{t|t-1}), \quad (3)$$

$$P_t = P_{t|t-1} - P_{t|t-1} Z' \Sigma_t^{-1} Z P_{t|t-1}, \quad (4)$$

where $a_{t|t-1} = E(\alpha_t | Y_1, \dots, Y_{t-1})$ and $P_{t|t-1} = E((\alpha_t - a_{t|t-1})(\alpha_t - a_{t|t-1})' | Y_1, \dots, Y_{t-1})$ are obtained from the following prediction equations:

$$a_{t|t-1} = Wa_{t-1}, \quad (5)$$

$$P_{t|t-1} = WP_{t-1}W' + RQR', \quad (6)$$

and

$$\Sigma_t = ZP_{t|t-1}Z' + H. \quad (7)$$

Note that Σ_t is the covariance matrix of the innovations, $v_t = Y_t - E(Y_t | Y_1, \dots, Y_{t-1})$. Inverting Σ_t can be a difficult

(2002), or the time-varying parameters (volatilities and constants) model proposed by Delle Monache, Petrella, and Venditti (2016). Fiorentini, Sentana, and Shephard (2003) and Koopman, Mallee, and Van der Wel (2010) allow the volatilities to be time-varying. Koop and Korobilis (2014) also consider an SSM with time-varying parameters. Furthermore, there are models with deterministic components; see, for example, Grassi, Proietti, Frale, Marcellino, and Mazzi (2015) and Jungbacker and Koopman (2015), among many others. Most of the procedures described in this paper can be extended to these cases; see, for example, Koopman (1993) for a general specification of the SSM with time-varying system matrices. However, we focus on the homogeneous time-invariant SSM to simplify the exposition.

³ The SSM can also be written with time-varying system matrices as far as they depend on past observations and, consequently, they are known at time $t - 1$.

⁴ If the disturbances are not normally distributed, then $a_{t|t-1}$ and a_t are the minimum mean square linear estimators (projections) of α_t given Y_1, \dots, Y_{t-1} and Y_1, \dots, Y_t , respectively. See Sentana, Almuzara, and Amengual (2019) for a test for normality of the disturbances in the context of DFM's.

task when the cross-sectional dimension of Y_t , N , is large.⁵ Harvey (1989) suggests two alternative possible solutions. First, using the Woodbury identity, it is possible to see that

$$\Sigma_t^{-1} = H^{-1} - H^{-1}Z (P_{t|t-1}^{-1} + Z'H^{-1}Z)^{-1} Z'H^{-1}, \quad (8)$$

and

$$|\Sigma_t| = |H| \times |P_{t|t-1}| \times |P_{t|t-1}^{-1} + Z'H^{-1}Z|. \quad (9)$$

Expression (8) is easy to evaluate if H is diagonal. Furthermore, given that the covariance matrix of ε_t , H , is time-invariant, it only needs to be inverted once. The second solution proposed by Harvey (1989) to avoid inverting Σ_t is to use the information filter that gives a set of recursions for the information matrix, P_t^{-1} .

Note that using (8) requires the existence of $P_{t|t-1}^{-1}$. However, $P_{t|t-1}$ could be non-invertible if, for instance, there are moving-average components in the state vector; see, for example, Ansley and Kohn (1985). In the case of non-invertible matrices, one could use a pseudo-inverse or a generalized inverse as, for example, the Moore–Penrose generalized inverse.

Furthermore, Jungbacker and Koopman (2015) argue that (8) does not necessarily lead to computational gains, because when N is very large, Σ_t^{-1} and the Kalman filter recursions still remain high-dimensional. Alternatively, they propose a computationally efficient procedure for the Kalman filter recursions. The key insight is that the observed time series, Y_t , can be split into a low-dimensional vector series and a high-dimensional vector series as follows:

$$Y_t^* = AY_t, \quad (10)$$

where A is an $N \times N$ non-singular matrix such that $A = [A^L A^H]'$ with $A^L = Z^{\dagger} H^{-1}$ being an $n \times N$ matrix where $n \ll N$ is the rank of Z and Z^{\dagger} is a basis for the column space of Z . If Z is of full column rank and $n = m$, then $Z^{\dagger} = Z$ and, consequently, $A^L = ZH^{-1}$.⁶ Define $Y_t^L = A^L Y_t$. The measurement equation for Y_t^L is given by

$$Y_t^L = A^L Z \alpha_t + A^L \varepsilon_t. \quad (11)$$

For factor extraction, we need to apply the Kalman filter to the low-dimensional series, Y_t^L ; see Grassi et al. (2015) for an empirical application.

The Kalman filter is in a steady state if

$$P_{t+1|t} = \bar{P}. \quad (12)$$

The important fact about an SSM being in a steady state is that the recursion for the MSEs of the state is redundant. Consequently, the covariance matrix of the innovations, Σ_t , also converges to a steady-state $\bar{\Sigma}$, as follows:

$$\lim_{t \rightarrow \infty} \Sigma_t = \bar{\Sigma} = Z \bar{P} Z' + H, \quad (13)$$

and

$$\bar{\Sigma}^{-1} = H^{-1} - H^{-1}Z (\bar{P}^{-1} + Z'H^{-1}Z)^{-1} Z'H^{-1}, \quad (14)$$

⁵ We assume that the inverse of Σ_t exists.

⁶ The matrix A^H is not needed for filtering or estimation.

where \bar{P} is obtained by solving the algebraic Riccati equation and relates to results in optimal control theory; see Chow (1975). Harvey (1989) shows that, if the system is detectable⁷ and stabilizable,⁸ and if $P_{1|0}$ is positive semi-definite, then

$$\lim_{t \rightarrow \infty} P_{t|t-1} = \bar{P}, \quad (16)$$

with \bar{P} being independent of $P_{1|0}$. If the system is observable and if $P_{1|0} - \bar{P}$ is positive definite or $P_{1|0} = \bar{P}$, then the result in (16) holds, although this is not sufficient for the steady state to be reached exponentially fast.

As mentioned above, in addition to one-step-ahead predictions, $a_{t|t-1}$, and update filtered estimates of the state, a_t , one can also obtain smoothed estimates, given by $a_{t|T} = E[\alpha_t | Y_1, \dots, Y_T]$ together with their corresponding MSEs, $P_{t|T} = E[(\alpha_t - a_{t|T})(\alpha_t - a_{t|T})' | Y_1, \dots, Y_T]$. The smoothed estimates of the state can be computed backwards, for $t = T-1, \dots, 1$, by the following recursive formulae:

$$a_{t|T} = a_t + C_t (a_{t+1|T} - Wa_t) \quad (17)$$

$$P_{t|T} = P_t + C_t (P_{t+1|T} - P_{t+1|t}) C_t' \quad (18)$$

where $C_t = P_t W' P_{t+1|t}^{-1}$. Note that, from a computational point of view, Eqs. (17) and (18) are not preferred for smoothing; see the discussion in Durbin and Koopman (2012) for a description of more efficient state smoothers.

The prediction equations of the Kalman filter in (5) and (6) can also be used recursively to obtain h -step-ahead out-of-sample forecasts of the state by using them from $h = 2, \dots, H$ without the update step, as follows:

$$a_{T+h|T+h-1} = Wa_{T+h-1|T+h-2}, \quad (19)$$

$$P_{T+h|T+h-1} = WP_{T+h-1|T+h-2}W' + RQR', \quad (20)$$

with $a_{T+1|T}$ and $P_{T+1|T}$ given by (5) and (6).

2.3. Parameter estimation

The system matrices may depend on a set of unknown parameters. One of the main statistical tasks is often the estimation of these parameters. The Kalman filter is also important because it enables the likelihood function to be calculated through the prediction error decomposition,

⁷ The state vector is observable if it can be determined exactly given Y_t, \dots, Y_{t+m-1} . The system is observable if $\text{Rank}[Z', WZ', \dots, (W')^{m-1}Z'] = m$. Observability implies detectability.

⁸ Defining $Q = R^*R'$, the transition equation can be written as

$$\alpha_t = Wa_{t-1} + B\eta_t^*, \quad (15)$$

where $B = RR^*$, and $\eta_t^* = R^{*-1}\eta_t$ is such that $E(\eta_t^*) = 0$ and $E(\eta_t^*\eta_t^{*'}) = I_g$. The model is controllable if $\text{Rank}[B, WB, \dots, W^{m-1}B] = m$. If B is of rank m , the controllability condition is satisfied. This condition means that from any particular value of α_t , the noise η_t^* can be chosen in such a way that any desired value for α_{t+m} can be attained. When the model is not controllable, certain elements in the state vector can only be manipulated indirectly via other elements. Conditions that are sufficient for controllability are also sufficient for stabilizability.

opening the way for ML estimation of any unknown parameters of the model, Ψ . For a Gaussian model, the prediction error decomposition of the log-likelihood function is given by

$$\begin{aligned} \log L(Y; \Psi) = & -\frac{NT}{2} \log(2\pi) - \frac{1}{2} \sum_{t=1}^T \log |\Sigma_t| \\ & - \frac{1}{2} \sum_{t=1}^T v_t' \Sigma_t^{-1} v_t, \end{aligned} \quad (21)$$

where $Y = (Y_1, \dots, Y_T)$ and both v_t and Σ_t can be obtained from the Kalman filter. In multivariate models, it is not always clear how to compute appropriate starting values for the initial state; see de Jong (1991) for an easy-to-implement algorithm.

Very recently, Delle Monache and Petrella (2019) took advantage of the matrix representation of the SSM and derived closed-form expressions of the log-likelihood and the smoothed estimator of the state that are computationally feasible even for very large N , provided that the covariance matrices of the measurement and transition noise are invertible.

Finally, the log-likelihood can also be obtained using the transformation proposed by Jungbacker and Koopman (2015). In particular,

$$\begin{aligned} \log L(Y; \Psi) = & \log L(Y^L; \Psi) - \frac{T}{2} \log \frac{|\Sigma_\varepsilon|}{|A^L \Sigma_\varepsilon A^{L'}|} \\ & - \frac{1}{2} \sum_{t=1}^T e_t' \Sigma_\varepsilon^{-1} e_t, \end{aligned} \quad (22)$$

where $\log L(Y^L; \Psi)$ can be obtained as in (21) based on the low-dimensional vector $Y^L = (Y_1^L, \dots, Y_T^L)$, and e_t is the generalized least squares (GLS) residual of the regression of Y_t on Z with covariance matrix Σ_ε .

The numerical maximization of the likelihood can be a difficult task when N is large and the number of parameters in the model is also large. In this case, the Gaussian log-likelihood can be maximized using the expectation maximization (EM) algorithm of Dempster, Laird, and Rubin (1977) proposed by Shumway and Stoffer (1982) and Watson and Engle (1983) for ML estimation in SSMs; see Wu (1983) for the convergence properties of the EM algorithm. The computational cost of the EM algorithm can be rather high, since it requires a smoother for the state vector and the corresponding MSE matrices. However, if the only parameters to be estimated are in the covariance matrices, the EM algorithm can be modified to be based on the disturbance smoother of Koopman (1993), saving considerable computational effort.

A very important issue related to the estimation of the parameters of SSMs is the identification of the state vector. Given the model in (26) and (27), there is not a unique representation of the state vector. Define an arbitrary non-singular $m \times m$ matrix, B , and consider a new state vector $\alpha_t^* = B\alpha_t$. The following SSM is observationally equivalent to the SSM in Eqs. (1) and (2):

$$Y_t = Z^* \alpha_t^* + \varepsilon_t \quad (23)$$

$$\alpha_t^* = W^* \alpha_{t-1}^* + R^* \eta_t \quad (24)$$

where $Z^* = ZB^{-1}$, $W^* = BWB^{-1}$, and $R^* = BR$. Therefore, there are m^2 unknowns in matrix B . In order to identify the model, Harvey (1989) proposes to restrict $RQR' = I_m$, which means $\frac{m(m+1)}{2}$ restrictions, and the elements in Z to be such that $z_{ij} = 0$ for $j > i, i = 1, \dots, m - 1$, which implies the $\frac{m(m-1)}{2}$ additional restrictions needed to identify the state vector; see Hannan and Deistler (1988) and Trenkler and Weber (2016) for discussions on identification issues in SSMs.

Subject to certain regularity conditions, the ML estimator, $\hat{\psi}$, has a limiting multivariate normal distribution with mean ψ and covariance matrix $T^{-1}A^{-1}$, where $A = \lim_{T \rightarrow \infty} \frac{IM}{T}$ with IM being the information matrix; see Harvey (1989) for a discussion of the regularity conditions, and Hannan and Deistler (1988) for the frequentist properties of the ML estimator.

When the Kalman filter converges to a steady state exponentially fast, the properties of the ML estimator (conditional on the starting values) do not depend on the way in which the filter is started off. However, it is important to note that exact ML estimation requires exact initialization of the Kalman filter; see, for example, de Jong (1988) and Francke, Koopman, and de Vos (2010) for ML estimation of SSMs, and Koopman and Shephard (1992) for the expression of the exact log-likelihood.

Note that if the unknown parameters in $\bar{\Sigma}$, the steady state of Σ_t , are independent of the unknown parameters determining $E(Y_t | Y_{t-1}, \dots, Y_1)$, then maximizing the likelihood is equivalent to minimizing

$$S(\Psi) = \left| \sum_{t=1}^T v_t v_t' \right|. \quad (25)$$

3. Dynamic factor models and KFS factor extraction

DFMs are examples of the much larger class of SSMs, in which observable variables are expressed in terms of unobserved or latent variables, which in turn evolve according to some lagged dynamics. In this section, we describe how DFMs can be expressed as SSMs and how the KFS algorithms can be used for factor extraction.

Consider that $Y_t = (Y_{1t}, \dots, Y_{Nt})'$, $t = 1, \dots, T$, is a stationary zero mean $N \times 1$ vector time series generated by the following DFM⁹:

$$Y_t = \chi_t + \varepsilon_t, \quad (26)$$

where $\chi_t = (\chi_{1t}, \dots, \chi_{Nt})'$ and $\varepsilon_t = (\varepsilon_{1t}, \dots, \varepsilon_{Nt})'$ are $N \times 1$ vectors representing the common and idiosyncratic components, respectively.¹⁰ Depending on the definition of χ_t , there are two main versions of the DFM usually considered in the literature: the *static* and the *dynamic* versions.

⁹ We assume that all deterministic components have been removed from the series in Y_t previous to their analysis.

¹⁰ Note that the model in (26) together with the specifications of F_t and ε_t below, can be considered as an SSM in the signal-to-noise spirit; see Harvey (1989). We are very grateful to one referee for pointing out this analogy.

3.1. Static dynamic factor models

The *static* version of the DFM (S-DFM) establishes a contemporaneous relationship between each variable in the system and the unobserved underlying factors at time t as follows:

$$\chi_{it} = \lambda_i' F_t \quad (27)$$

where $\lambda_i = (\lambda_{i1}, \dots, \lambda_{ir})$ is the $r \times 1$ vector of factor loadings of the Y_i variable. It is popular to assume that F_t , the $r \times 1$ vector of common factors, evolves over time following a stationary VAR(p) model given by

$$F_t = \Phi_1 F_{t-1} + \Phi_2 F_{t-2} + \dots + \Phi_p F_{t-p} + u_t, \quad (28)$$

where $\Phi_i, i = 1, \dots, p$, are matrices containing the autoregressive parameters, and u_t is an $r \times 1$ white noise vector with covariance matrix Σ_u . This specification of the factors has been considered in many empirical studies with the values of r and p depending on the particular application; see, for example, Proietti (2011), who specified $r = 6$ and $p = 1$, and Camacho and Perez-Quiros (2010) and Scotti (2016), both with $r = 1$ and $p = 11$.¹¹ In what follows, we consider $p = 1$ to simplify notation. However, all results can be easily extended to models with $p > 1$.

Finally, the idiosyncratic components, ε_{it} , are often specified as AR(p_i^*) processes as follows:

$$\varepsilon_{it} = \theta_{1i} \varepsilon_{it-1} + \theta_{2i} \varepsilon_{it-2} + \dots + \theta_{p_i^* i} \varepsilon_{it-p_i^*} + e_{it}, \quad (29)$$

where $\theta_{1i}, \dots, \theta_{p_i^* i}$ are autoregressive parameters, and $e_t = (e_{1t}, \dots, e_{Nt})$ is the vector of idiosyncratic noise, assumed to be white noise with covariance matrix Σ_e . The autoregressive order in (29) depends on the particular application. In many studies, it is considered the same for all $i = 1, \dots, N$. In this case, denote $p^* = p_i^*$. For example, Scotti (2016) assumes that $p^* = 1$, while Camacho and Perez-Quiros (2010) assume that $p_i^* = 1$ for all monthly variables and $p_i^* = 5$ for the quarterly variables (GDP and unemployment) of their system. Also, García-Ferrer and Poncela (2002) allow p_i^* to vary between 1 and 6. Note that these two models are small in scale, with $N = 13$ and $N = 5$, respectively. As for the factors, we henceforth assume that $p^* = p_i^* = 1$ to simplify the analysis. Finally, if the idiosyncratic components, ε_t , are assumed to be cross-sectionally uncorrelated, i.e. Σ_e is diagonal, the DFM is known as *exact*, whereas if the idiosyncratic noise terms are weakly cross-correlated, the DFM is called *approximate*.¹² In an exact DFM, for the

¹¹ Some authors also consider factors generated by moving-average processes; see, for example, Bai and Ng (2007) and Otrok and Whiteman (1998). However, these are the exception.

¹² With respect to what are called exact or approximate DFMs in the related literature, the specification about whether the idiosyncratic noise terms are temporally correlated is not clear. This distinction was originally given in the context of PC factor extraction and, consequently, the dynamic dependence of the idiosyncratic noise was irrelevant. As a consequence, in the original definition of exact or approximate DFMs, nothing is specified about whether the idiosyncratic components were serially uncorrelated. Afterwards, in the literature several authors defined exact DFMs as those in which the idiosyncratic component is cross-sectionally uncorrelated regardless of whether it has serial correlation; see, for example, Bańbura and Modugno (2014) and Stock and Watson (2016a).

purposes of explaining contemporaneous movements and making forecasts, once you know the factors, the other series provide no additional useful information.

Therefore, the S-DFM is given by

$$Y_t = \Lambda F_t + \varepsilon_t \quad (30)$$

where $\Lambda = (\lambda_1, \dots, \lambda_N)'$ is the $N \times r$ matrix of factor loadings, and F_t and the elements of ε_t are defined in (28) and (29), respectively.

Consider first the S-DFM with serially uncorrelated idiosyncratic noise, i.e. $\theta_i = 0$ for $i = 1, \dots, N$. In this case, it is straightforward to write the S-DFM as an SSM in Eqs. (1) and (2) by considering $m = g = r$, $Z = \Lambda$, $\alpha_t = F_t$, $H = \Sigma_e$, $W = \Phi_1$, $R = I$, and $Q = \Sigma_u$. If the S-DFM is further exact, assuming that r and p as well as all parameters in the model are known, the KFS algorithms described in Section 2 can be implemented to extract the factors, regardless of the cross-sectional dimension, N , by applying the expression of the inverse of the innovation covariance matrix in (8). In the more realistic case in which Σ_e is not diagonal, one can still invert Σ_t using expression (8) by inverting Σ_e using the Cholesky decomposition $\Sigma_e = DD'$, where D is a lower triangular matrix. Inverting D is feasible even if N is very large. Alternatively, the factors can be extracted using the KFS algorithms as if Σ_e were diagonal although it is not. Obviously, in this latter case, the factors extracted using KFS and the corresponding MSEs obtained are not the true conditional means (projections) and MSEs of the factors.

As an illustration of the effects of misspecification on factor extraction using KFS, we simulate a system with $r = 1$ factor characterized by an AR(1) model with autoregressive parameter $\phi = 0.7$ and variance of the noise $1 - \phi^2$. The idiosyncratic noise terms are serially uncorrelated, homoscedastic, with variances of 0.5, and cross-sectionally correlated with the correlation matrix given by a Toeplitz matrix whose (i, j) element is given by $\tau^{|i-j|}$ with $\tau = 0.5$. To analyze the misspecification effects as a function of the cross-sectional dimension, we simulate systems with $T = 200$ observations and $N = 5$, 50, and 150 variables, which represent small, medium, and large systems, respectively. The accuracy of the point factor estimates is measured by computing the sample MSEs, given by $\frac{1}{T} \sum_{t=1}^T (\hat{F}_t - F_t)^2$, while the accuracy of the KFS MSEs is measured by the sample coverages computed as the percentage of times the true factor is included in the 95% confidence interval constructed using the KFS's MSEs. Note that a correct evaluation of the extracted factor requires a measure of its uncertainty as, for example, the MSE; see, for instance, the discussions in Bai (2003), Jackson, Kose, Otkrok, and Owyang (2016), and Thorsrud (2020). A formal analysis of the effects of misspecification of the idiosyncratic noise on the point and interval estimation of the factors is beyond the objectives of this survey. Table 1, which reports the sample MSEs and coverages for the particular simulations described above, shows that for these particular simulated systems, the increases in the MSEs of the misspecified models are 11.87%, 13.33%, and 5% when compared with the corresponding MSEs of the true specification, for $N = 5, 50$,

and 150, respectively. As expected, the loss of accuracy of the point-estimated factors, when assuming falsely that the idiosyncratic components are uncorrelated, is negligible when the cross-sectional dimension, N , is large. However, when looking at the coverages reported in Table 1, we can observe that the coverages of the intervals constructed with the misspecified smoother are below nominal. When $N = 5$, the coverage is 90%, while the coverage is 75.5% when $N = 150$. Therefore, the undercoverage seems to be larger as N increases. The MSEs of the smoothed factors are clearly affected by the misspecification. In this particular example, they are smaller than they should be. To have a visual plot of this simulation, Fig. 1 plots the true simulated factor together with the factor extracted using the smoothing algorithm—assuming that all parameters are known (left column)—and the smoothed factor extracted when all the parameters are known but Σ_e is assumed to be diagonal (right column), together with their corresponding 95% confidence bounds. Fig. 1 shows that the differences between both point smooth factors are only visually appreciable when $N = 5$. However, the intervals of the misspecified smoother are wider than those obtained with the correct model. When $N = 50$, the point estimates of the factors are nearly the same, with the main differences appearing in the MSEs, which are smaller than they should be. When N is large, the MSEs are already so close to zero that it is indifferent whether the filter is run with the true covariance matrix of the idiosyncratic components or with a diagonal matrix; see Poncela and Ruiz (2020) for further illustrations with other cross-sectional dimensions when the idiosyncratic components are treated as if they were cross-sectionally uncorrelated, and Luciani (2014) who shows that accounting for cross-correlation rarely boosts the forecasting accuracy.

Consider now the S-DFM in which the idiosyncratic noise is serially correlated according to (29), and denote by Θ the matrix with $\theta_i, i = 1, \dots, N$ in its main diagonal. In this case, the DFM can be reformulated by augmenting the state vector with lags of the factors as follows:

$$Y_t = \Theta Y_{t-1} + [\Lambda \quad -\Theta \Lambda] \begin{bmatrix} F_t \\ F_{t-1} \end{bmatrix} + \varepsilon_t \quad (31)$$

$$\begin{bmatrix} F_t \\ F_{t-1} \end{bmatrix} = \begin{bmatrix} \Phi_1 & 0 \\ I_r & 0 \end{bmatrix} \begin{bmatrix} F_{t-1} \\ F_{t-2} \end{bmatrix} + \begin{bmatrix} u_t \\ 0 \end{bmatrix}, \quad (32)$$

where I_r is the $r \times r$ identity matrix; see Bai and Li (2016), Jungbacker and Koopman (2015), Jungbacker, Koopman, and van der Wel (2011), Pinheiro, Rua, and Dias (2013), Poncela and Ruiz (2015), Reis and Watson (2010), Stock and Watson (2005) and Watson and Engle (1983) for implementations of the model in (31) and (32). Defining the observations as $Y_t - \Theta Y_{t-1}$, the model in (31) and (32) can be directly cast in state space form by setting $m = 2r$, $g = m$, $Z = [\Lambda \quad -\Theta \Lambda]$, $\alpha_t = [F_t \quad F_{t-1}]$, $W = \begin{bmatrix} \Phi_1 & 0 \\ I_r & 0 \end{bmatrix}$, $R = [I_r \quad 0]$ and $\eta_t = u_t$, in the SSM in (1) and (2).¹³

¹³ Alternatively, one can deal with the autocorrelation of the idiosyncratic noise by augmenting the state vector by ε_t ; see, for example,

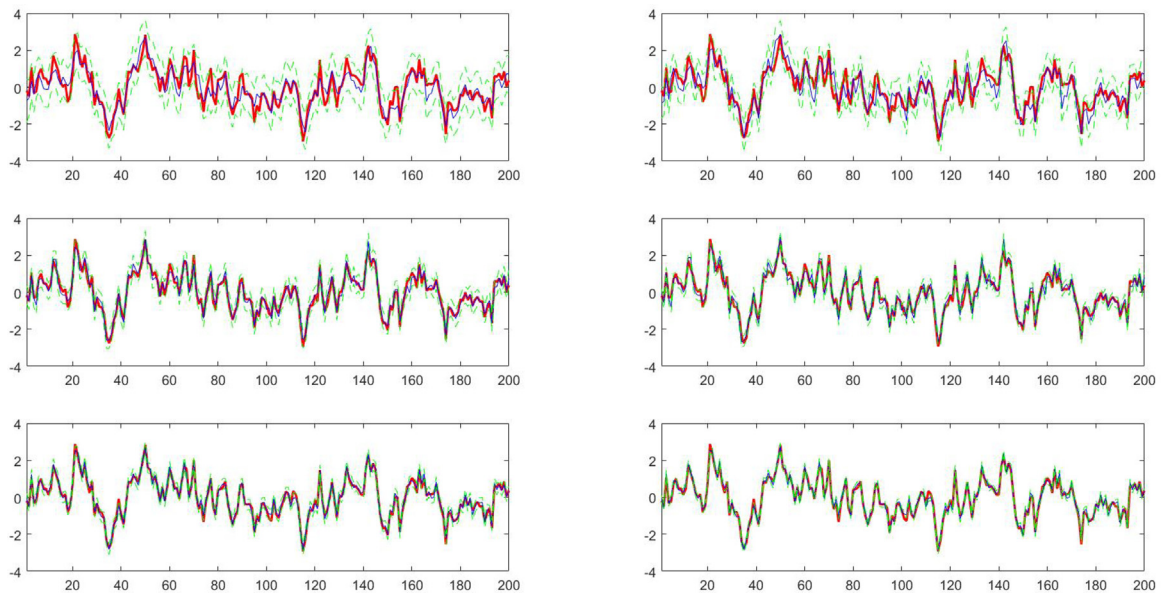


Fig. 1. Simulated factor (red line) together with the factor extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification (left column) and the true model, assuming that the idiosyncratic component is serially and cross-sectionally uncorrelated (right column). The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row) and weakly cross-correlated idiosyncratic components. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Table 1

Mean Square Errors (MSEs) of factors extracted using the true and misspecified DFM with known and estimated parameters. The coverages correspond to the intervals constructed using the KFS's MSEs with a nominal coverage of 95%.

	Known parameters				Estimated parameters			
	True model		Misspecified model		True model		Misspecified model	
	MSE	Coverage	MSE	Coverage	MSE	Coverage	MSE	Coverage
Cross-sectionally correlated idiosyncratic noise								
$N = 5$	0.219	0.965	0.245	0.900	0.218	0.945	0.266	0.670
$N = 50$	0.045	0.960	0.051	0.805	0.204	0.525	0.073	0.730
$N = 150$	0.020	0.965	0.021	0.755	4.879	0.045	0.226	0.375
Autocorrelated idiosyncratic noise								
$N = 5$	0.173	0.970	0.174	0.965	0.197	0.899	0.226	0.870
$N = 50$	0.019	0.985	0.019	0.970	0.071	0.894	0.052	0.865
$N = 150$	0.008	0.965	0.008	0.955	0.230	0.397	0.234	0.330
Cross-sectionally correlated and autocorrelation idiosyncratic noise								
$N = 5$	0.302	0.985	0.348	0.865	–	–	–	–
$N = 50$	0.051	0.990	0.055	0.755	–	–	–	–
$N = 150$	0.022	0.970	0.023	0.715	–	–	–	–

Even if the idiosyncratic components, ε_t , are serially correlated, the KFS can be run as if they were uncorrelated. Fig. 2 illustrates the results by plotting the true factor simulated by the same model described above but with the cross-sectionally uncorrelated idiosyncratic components generated by independent AR(1) models, all of them with autoregressive parameter 0.5 as in Doz, Giannone, and Reichlin (2012). The factors are extracted

by the KFS based on the true model and assuming that the idiosyncratic components are serially uncorrelated. Note that, in this case, the difference between the point-estimated factors is even smaller than in Fig. 1. Table 1, which reports the sample MSEs and coverages for these particular simulations, shows that both are nearly the same, regardless of whether the autocorrelation of the idiosyncratic component is taken into account. It seems that the effects of the misspecification of the serial correlation in the idiosyncratic components are milder than the effects of misspecification of cross-sectional correlations.¹⁴

Baíbarúa and Modugno (2014), Coroneo et al. (2016) and Jungbacker et al. (2011). The main problem associated with this alternative is that the state vector dimension, $m = g = r + N$, increases with N and can be infeasible from a computational point of view for large cross-sectional dimensions. Both formulations lead to the same results when the initialization issues are properly accounted for.

¹⁴ This result could be expected when extracting the factors. However, misspecification of the serial correlation of the idiosyncratic

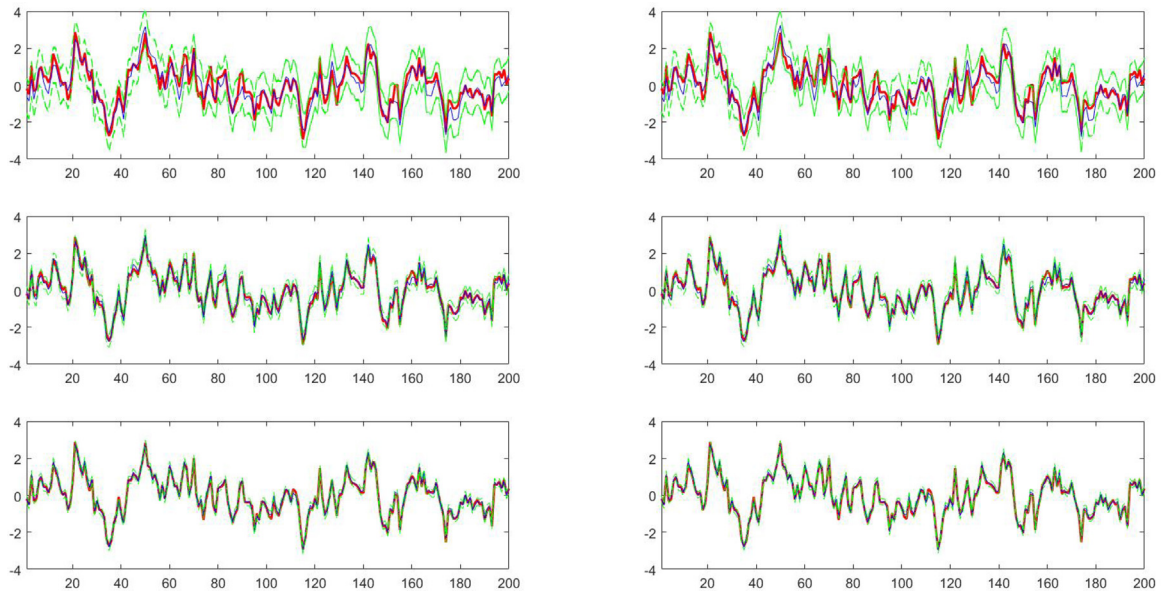


Fig. 2. Simulated factor (red line) together with the factor extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification (left column) and the true model, assuming that the idiosyncratic component is serially and cross-sectionally uncorrelated (right column). The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row) and serially correlated idiosyncratic components. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Finally, the same conclusions can be obtained from Fig. 3, which plots the same quantities described above when the idiosyncratic noise is both serially and cross-sectionally correlated; see also the quantities reported in Table 1, which shows that the undercoverage can be mainly attributed to the lack of consideration of the cross-correlation of the idiosyncratic components.

Note that these conclusions are obtained in a very simple stationary model with a single factor that is moderately serially dependent. It is possible that if the factor and idiosyncratic noise have other alternative structures of serial correlations, the conclusions could be different.

3.2. Dynamic version of the dynamic factor model

In the dynamic version of the DFM (D-DFM), each variable in the system at time t is related to the unobserved factors at time t , $t - 1, \dots, t - s$.¹⁵ Following Bai and Ng (2007), and Stock and Watson (2016b), among others, we express the common component of the D-DFM as follows:

$$\chi_t = \Lambda(L)G_t, \quad (33)$$

where $\Lambda(L) = \Lambda_0 + \Lambda_1L + \dots + \Lambda_sL^s$ and G_t is the $q \times 1$ vector of unobserved factors. Assuming for simplicity that the factors, G_t , follow a VAR(1) and $s = 1$, the D-DFM with

components could have implications for forecasting; see D'Agostino and Giannone (2012) and Poncela, Senra, and Sierra (2020) who conclude that the effect of the idiosyncratic dynamics on forecasting macroeconomic inflation and commodity inflation, respectively, are negligible.

¹⁵ Bai and Ng (2007) claim that, for forecasting purposes, little is to be gained from a distinction between static and dynamic factors.

the common component defined as in (33) can be written as an S-DFM with restrictions as follows¹⁶:

$$Y_t = \begin{bmatrix} \Lambda_0 & \Lambda_1 \end{bmatrix} \begin{bmatrix} G_t \\ G_{t-1} \end{bmatrix} + \varepsilon_t \quad (34)$$

$$\begin{bmatrix} G_t \\ G_{t-1} \end{bmatrix} = \begin{bmatrix} \Phi_1 & 0 \\ I_q & 0 \end{bmatrix} \begin{bmatrix} G_{t-1} \\ G_{t-2} \end{bmatrix} + \begin{bmatrix} v_t \\ 0 \end{bmatrix}, \quad (35)$$

where $F_t = \begin{bmatrix} G_t \\ G_{t-1} \end{bmatrix}$ can be treated as an $r = q(s + 1)$ vector of static factors with the errors given by $u_t = \begin{bmatrix} v_t \\ 0 \end{bmatrix}$; see Bai and Ng (2007) for the expression of the D-DFM as an S-DFM in a more general context. The covariance matrix of the disturbances of the static factors, Σ_u has rank q .

Alternatively, the D-DFM can be written as an S-DFM as follows

$$Y_t = (\Lambda_0\Phi + \Lambda_1)G_{t-1} + \Lambda_0v_t + \varepsilon_t \quad (36)$$

$$\begin{bmatrix} G_{t-1} \\ v_t \end{bmatrix} = \begin{bmatrix} \Phi & I_q \\ 0 & 0 \end{bmatrix} \begin{bmatrix} G_{t-2} \\ v_{t-1} \end{bmatrix} + \begin{bmatrix} 0 \\ I_q \end{bmatrix} v_t \quad (37)$$

Once more, the D-DFM is written as an S-DFM with two common factors. It is important to point out that, when the D-DFM is written as an S-DFM, as in Eqs. (34) and (35), it is not possible to assume that the static factors are orthogonal. Alternatively, if the D-DFM is written as an S-DFM, as in Eqs. (36) and (37), the factors can be assumed

¹⁶ The cases in which $p > 1$ and/or $s > 1$ follow straightforwardly but they are notationally more cumbersome.

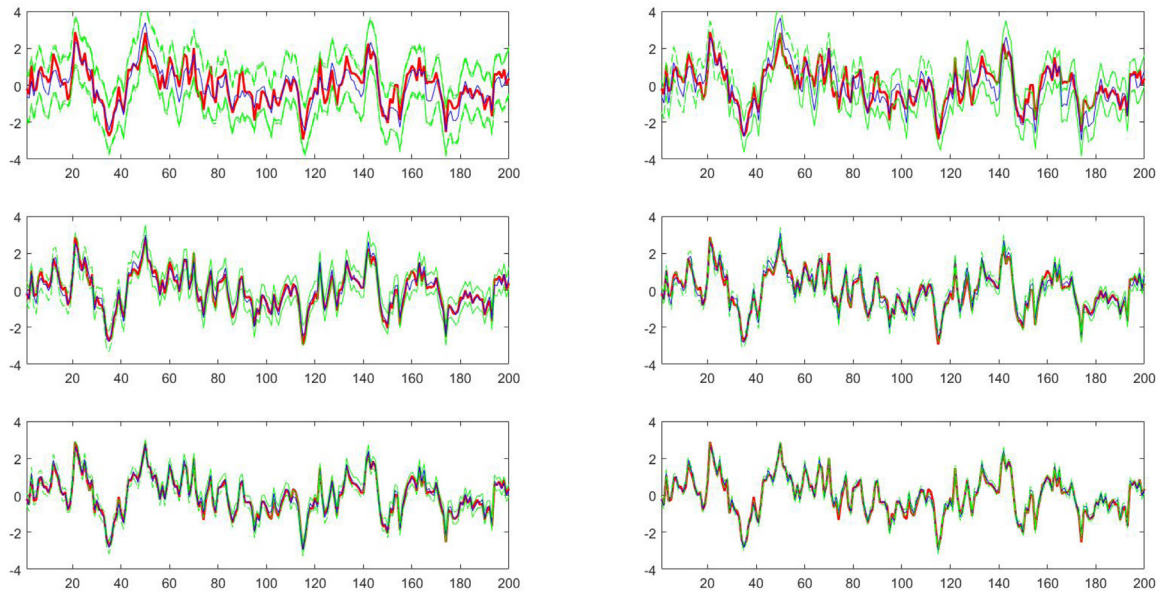


Fig. 3. Simulated factor (red line) together with the factor extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification (left column) and the true model, assuming that the idiosyncratic component is serially and cross-sectionally uncorrelated (right column). The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row) and serially and cross-sectionally correlated idiosyncratic components. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

to be orthogonal but the matrix of loading is of rank q and, consequently, it is not possible to find a rotation such that $\Lambda'\Lambda$ is diagonal.¹⁷

If the idiosyncratic noise terms are serially uncorrelated, then the model in (34) and (35) can be cast as an SSM and the KFS can be implemented to extract the factors; see Pinheiro et al. (2013) for an empirical implementation. If the idiosyncratic noise is serially correlated as in (29), then the D-DFM with $s = p = 1$ can be expressed as follows:

$$Y_t = \Theta Y_{t-1} + \Lambda_0 G_t + (\Lambda_1 - \Theta \Lambda_0) G_{t-1} - \Theta \Lambda_1 G_{t-2} + e_t \quad (38)$$

$$\begin{bmatrix} G_t \\ G_{t-1} \\ G_{t-2} \end{bmatrix} = \begin{bmatrix} \Phi_1 & 0 & 0 \\ I_r & 0 & 0 \\ 0 & I_r & 0 \end{bmatrix} \begin{bmatrix} G_{t-1} \\ G_{t-2} \\ G_{t-3} \end{bmatrix} + \begin{bmatrix} v_t \\ 0 \\ 0 \end{bmatrix}. \quad (39)$$

The D-DFM in (38) and (39) can be cast as an SSM and the factors extracted using KFS without further issues.

As explained above, when describing the KFS factor extraction in S-DFMs, the factors of a D-DFM can also be extracted using KFS as if the idiosyncratic noise terms are serially and cross-sectionally uncorrelated even if they are not.

4. Estimating parameters of DFM

The KFS factor extraction described in the previous section assumes known parameters. However, in practice, the parameters are unknown and need to be estimated

before running KFS algorithms. In this section, we survey the estimators of the parameters of the DFM based on ML when the model specification, i.e. s , r , and p , is known.¹⁸ The first issue faced when estimating the parameters of a DFM is related to parameter identification. In this section, we first describe the parameter identification and then their ML-based and Least Squares (LS)-based estimation.

4.1. Identification

As explained in Section 2, in an SSM such as that in Eqs. (1) and (2), one needs to impose m^2 restrictions to identify the m unobserved states; see Anderson and Rubin (1956) and Geweke and Singleton (1981) for excellent discussions on identification issues. Consequently, in the S-DFM with serially uncorrelated idiosyncratic noise, one needs to impose r^2 restrictions to identify the factors. For many applications, including macro-monitoring and forecasting, it is necessary only to identify the space spanned by the factors. Consequently, it is popular to solve the lack of identification by imposing mathematically convenient normalizations. It is common to assume that the factor noise covariance matrix, $\Sigma_u = I_r$ or the covariance of

¹⁷ Note that many procedures based on PCs require that the factors and loadings are simultaneously orthogonal.

¹⁸ In this paper, we focus on ML-based estimators. Other estimators have been proposed in the literature as Bayesian estimators (Jackson et al. (2016), Kaufmann and Schumaker (2019), Kose, Otrok, and Whiteman (2003), Lopes and West (2004) and Otrok and Whiteman (1998), MCMC procedures as in Moench, Ng, and Potter (2013), and the frequency-domain version of the EM algorithm (Fiorentini, Galesi, & Sentana, 2018). Finally, Kapetanios and Marcellino (2009) propose estimating the parameters using subspace algorithms; see Eickmeier and Ziegler (2008) for a comparison of the predictive performance of factors estimated using subspace methods with alternative estimators of the factors.

the factors, $\Sigma_F = I_r$ that amounts to $\frac{r(r+1)}{2}$ restrictions and that $\lambda_{ij} = 0, j > i$ that imposes the $\frac{r(r-1)}{2}$ additional restrictions needed. For example, Jungbacker and Koopman (2015) assume instead that Σ_u is diagonal and the diagonal elements of the r top rows of the loading matrix Λ are restricted to be one, and Solberger and Spanberg (2020) assume that $\Sigma_F = I_r$ with additional restrictions on the matrix of loadings. Alternatively, many authors assume that the matrix composed by the first r rows of the matrix Λ is the identity matrix (r^2 restrictions); see, for example, Bai and Ng (2013), Proietti (2011), Stock and Watson (2011), and Coulombe et al. (this issue). This latter restriction is denoted as the *named factor* restriction by Stock and Watson (2011).¹⁹ Alternatively, Reis and Watson (2010) assume that $\Lambda' \Lambda$ is diagonal and that the columns of Λ sum up to zero. Note that these are $\frac{r(r+1)}{2}$ restrictions. Although, they did not say it explicitly, additional $\frac{r(r-1)}{2}$ restrictions are needed for identification; see Bai and Li (2012, 2016) who consider five sets of restrictions and discuss how the distribution of an ML estimator depends on the identification restrictions.

Note that, when the idiosyncratic errors are correlated and/or the DFM has dynamic factors, even if the state vector is extended, the number of restrictions is still r^2 given that all new elements in the transition matrix, W , and in the covariance matrix, Q , are known. However, it is important to note that when dealing with the DDFM, there is a further identification issue that affects the specification of the model, as discussed below.

4.2. Estimation

Estimating the parameters of the DFM depends crucially on whether the model is static or dynamic, and on the specification of the idiosyncratic components. We first describe the ML estimator and then the two-step estimators based on LS.

4.2.1. Maximum likelihood

First, consider the S-DFM in (30) with serially and contemporaneously uncorrelated idiosyncratic errors. In this case, estimation of the parameters can be carried out, for example, assuming normality, by ML. Given that, in this case, Σ_ϵ is diagonal, the innovation covariance matrix, Σ_t can be easily inverted using (8) and the Kalman filter (KF) can be used to compute the innovation decomposition form of the Gaussian likelihood in (21), which can be maximized using numerical optimization algorithms; see, for example, Aruoba, Diebold, and Scotti (2009) and Engle and Watson (1981). As explained above, instead of numerical maximization of the likelihood obtained using the recursions of the Kalman filter, one can use the matrix form proposed by Delle Monache and Petrella (2019) who give details about how to implement it to the S-DFM with serially uncorrelated errors and show results for N up

to 200. Alternatively, given that numerically finding the maximum of the log-likelihood can be infeasible when N is large due to the very large number of parameters to be estimated, the Gaussian log-likelihood can be maximized using the EM algorithm proposed by Shumway and Stoffer (1982) and Watson and Engle (1983) for ML estimation in SSMs.²⁰ Although a complete treatment of the EM algorithm for SSMs can be found in, for example, Durbin and Koopman (2012) and Harvey (1989), we describe it next for completeness. The EM algorithm is based on the following decomposition of the log-likelihood conditional on the initial conditions for the factors:

$$\begin{aligned} \log L = & -\frac{T}{2} \log |\Sigma_u| - \frac{1}{2} \sum_{t=1}^T (F_t - \Phi F_{t-1})' \\ & \times \Sigma_u^{-1} (F_t - \Phi F_{t-1}) - \\ & \frac{T}{2} \log |R| - \frac{1}{2} \sum_{t=1}^T (Y_t - \Lambda F_t)' \Sigma_\epsilon^{-1} (Y_t - \Lambda F_t). \end{aligned}$$

The expected value of the log-likelihood conditional on the observations Y_1, \dots, Y_T is given by

$$\begin{aligned} E(\log L | Y_1, \dots, Y_T) = & -\frac{T}{2} \log |\Sigma_u| \\ & - \frac{1}{2} \sum_{t=1}^T \text{tr}\{E[(F_t - \Phi F_{t-1})(F_t - \Phi F_{t-1})' | Y_1, \dots, Y_T] \Sigma_u^{-1}\} - \\ & \frac{T}{2} \log |R| - \frac{1}{2} \sum_{t=1}^T \text{tr}\{E[(Y_t - \Lambda F_t)(Y_t - \Lambda F_t)' | Y_1, \dots, Y_T] \Sigma_\epsilon^{-1}\}, \end{aligned}$$

where

$$\begin{aligned} E[(Y_t - \Lambda F_t)(Y_t - \Lambda F_t)' | Y_1, \dots, Y_T] &= Y_t Y_t' + \Lambda P_{t|T} \Lambda' + \Lambda f_{t|T} f_{t|T}' \Lambda' - 2 Y_t f_{t|T}' \Lambda' \\ &= P_{t|T} + f_{t|T} f_{t|T}' + \Phi P_{t-1|T} \Phi' + \\ &\quad \Phi f_{t-1|T} f_{t-1|T}' \Phi' - 2 \Phi [f_{t-1|T} f_{t|T}' + C_t], \end{aligned}$$

and where $f_{t|T}$ is the smoothed estimate of F_t obtained using (17), $P_{t|T}$ is its MSE given in (18), and $C_t = E[(F_t - f_{t|T})(F_{t-1} - f_{t-1|T})' | Y_1, \dots, Y_T]$ can be obtained by the Kalman smoother if the state vector is augmented to include F_{t-1} .

The conditional expectation of the log-likelihood is maximized by

$$\hat{\Lambda} = \sum_{t=1}^T Y_t f_{t|T}' \left(\sum_{t=1}^T (f_{t|T} f_{t|T}' + P_{t|T}) \right)^{-1} \quad (40)$$

$$\hat{\Phi} = \sum_{t=1}^T (f_{t|T} f_{t-1|T}' + C_t) \left(\sum_{t=1}^T (f_{t-1|T} f_{t-1|T}' + P_{t-1|T}) \right)^{-1}. \quad (41)$$

¹⁹ Note that, in this type of identification restriction, the variable ordering could matter for parameter estimation given that the leading series determine the factors; see Chan, Leon-Gonzalez, and Strachan (2018) and Lopes and West (2004) in the context of a factor model in which the factors have no dynamic dependence.

²⁰ Comparing the properties of parameter estimates obtained by maximizing the likelihood based on the EM algorithm and on the procedure proposed by Delle Monache and Petrella (2019) could be of interest for further research.

Note that the estimators in (40) and (41) can be substituted when necessary by restricted versions after imposing the adequate restrictions. Furthermore, if the VAR order $p > 1$, the estimator can be modified accordingly. Finally, the corresponding estimators of the covariance matrices are given by

$$\hat{\Sigma}_e = \text{diag} \left\{ \frac{1}{T} \sum_{t=1}^T \hat{e}_t \hat{e}'_t \right\} \quad (42)$$

$$\hat{\Sigma}_u = \frac{1}{T} \sum_{t=1}^T \hat{u}_t \hat{u}'_t \quad (43)$$

where $\hat{e}_t = Y_t - \hat{A}F_t$ and $\hat{u}_t = F_t - \hat{\Phi}F_{t-1}$.

The EM algorithm works iteratively. Given starting values for the parameters, Ψ^0 , the expectation (E) step consists in computing the smoothed estimates of the factors and their MSEs. Once these smoothed expectations are obtained, the maximization (M) step consists in estimating the parameters using (40) to (43). These steps are iterated until convergence. Note that the EM algorithm can also be adopted to obtain ML estimates based on the likelihood decomposition proposed by Jungbacker and Koopman (2015) in (22). For each EM step, the Kalman smoothing is based on the low-dimensional model; see Grassi et al. (2015) for an empirical implementation. The parameters of the exact S-DFM can be estimated by ML using the EM algorithm regardless of N ; see, among many others, Stock and Watson (1989, 1991) and Coulombe et al. (this issue) with $N = 4$, Baíbura, Giannone, Modugno, and Reichlin (2013) with $N = 24$, Quah and Sargent (1993) with $N = 60$, and Proietti (2011) with $N = 148$. The EM algorithm is convenient because, in SSMs, it is straightforward to compute the expected value of α_t conditional on Y_T and to maximize the log-likelihood using standard regression formulae. However, the EM algorithm also has some disadvantages. First, the matrix of second partial derivatives is not available and, consequently, standard errors of the parameter estimates cannot be obtained directly. However, these partial derivatives can be approximated by perturbing the likelihood function in the neighborhood of the maximum. Computation of the information matrix via recursions is also possible, as in Harvey (1989) or Cavanaugh and Shumway (1996). Versions of the information matrix, obtained from outputs arising naturally in the EM algorithm, such as in Meng and Rubin (1991) or Oakes (1999), are either hard to compute, as in the former, or will involve relatively intractable derivatives, as in the latter. Jamshidian and Jennrich (2000) analyze the performance of several procedures to compute the standard errors of the parameters estimated using the EM algorithm, including procedures based on the numerical differentiation of the Fisher score vector.²¹

²¹ A compromise that is easy to apply and will be robust toward distributional assumptions is the bootstrap, as derived in Stoffer and Wall (1991). However, it is not clear whether bootstrapping can be

Finally, note that the EM estimators in (40) and (41) can be substituted when necessary by restricted versions after imposing the adequate restrictions. Furthermore, if the VAR order is $p > 1$, the EM estimator can also be modified accordingly.

If the idiosyncratic noise terms are weakly cross-correlated, ML estimation is still feasible if N is not very large; see, for example, Coulombe et al. (this issue). However, if the cross-sectional dimension is large, ML is not feasible due to the extremely large number of parameters. In any case, Doz et al. (2012) prove the consistency, when both N and T diverge to infinity, of smoothed factors extracted when the DFM parameters are substituted by estimates obtained using the EM algorithm assuming wrongly that the idiosyncratic components have neither cross-sectional nor temporal correlations.

This estimator is known in the related literature as quasi-ML (QML). The $\min(\sqrt{N}, \sqrt{T})$ -consistency and asymptotic normality of the estimates of the loadings, factors, and common components have been proved by Barigozzi and Luciani (2020) who derived the conditions under which the asymptotic distribution can still be used for inference in case of misspecification.²² As an illustration of the performance of the factors extracted using KFS with estimated parameters instead of the true parameters in a DFM with cross-sectionally correlated errors, we consider the same systems generated in the previous section with $N = 5, 50$, and 150 series and $T = 200$ temporal observations and implement the EM algorithm to estimate the parameters. It is important to point out that, in this paper, our objective is to illustrate the effects of misspecification on factor extraction. However, when the model parameters are estimated, misspecification also affects the estimated parameters. If the focus were on parameters instead of factors, then an analysis of the effects of misspecification on the estimated parameters would be warranted. Table 1, which reports the sample MSEs and coverages, shows that when the true model is estimated with $N = 5$, the results are very similar to those obtained with known parameters. Although the MSEs of the point estimates of the misspecified model are only slightly worse, the coverage in this case is very low (only 67% when the nominal is 95%).

As N increases, and given that T is fixed, the quality of the estimated parameters decreases, as does the quality of the point and interval estimates of the factors extracted with the estimated true model. For example, when $N = 50$, the sample MSE of the factor extracted with the estimated true model increases by 353.3% with respect to that of the factor extracted using the true parameters. Furthermore, the coverage is 52.5% instead of 96%. When $N = 150$, it is not possible to estimate all the parameters

implemented in the context of a large cross-sectional dimension, N . In the context of non-stationary DFMs, Peña and Poncela (2006) estimate the model parameters using the EM procedure with a final pass using the scoring algorithm to obtain uncertainty measures of the estimated parameters.

²² Barigozzi and Luciani (2020) compare the loadings, factors, and common components estimated using PC and QML estimators and conclude that, in static DFMs, both procedures are rather similar.

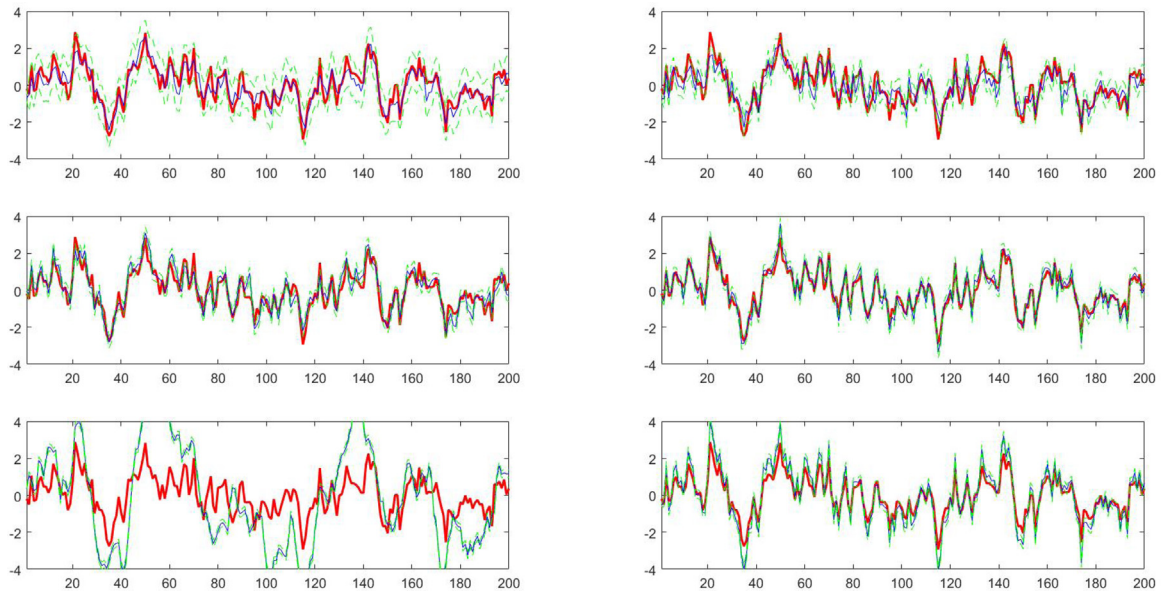


Fig. 4. Simulated factor (red line) together with the factor extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification (left column) and the true model assuming that the idiosyncratic component is serially and cross-sectionally uncorrelated (right column), both with the parameters estimated by ML using the EM algorithm. The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row) and serially and cross-sectionally correlated idiosyncratic components. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

in the model using only $T = 200$ observations. The MSE is large while the coverage is 4.5%. The true factor is almost always outside the interval. Fig. 4 plots the true factor together with the factor extracted when the parameters are estimated, assuming that the true model specification is known (left column), together with the 95% intervals for the factor obtained when the MSE of the smoothed estimates is obtained using the estimated parameters.

The first conclusion from Fig. 4 is that, if one wants to estimate the full covariance matrix of the idiosyncratic errors, the sample size, T , should be large enough when compared with the cross-sectional dimension. When $N = 150$ and $T = 200$, there is not enough information as to estimate the full covariance matrix of the idiosyncratic errors and, consequently, the extracted factor is far away from the true factor. However, when T is large enough, the factor extracted with estimated parameters behaves similarly to the factor extracted with known parameters; compare the first column in Fig. 1 when $N = 5$ or 50.

Regardless of N , the parameters of the exact S-DFM in Eqs. (31) and (32), with serially correlated idiosyncratic errors specified as a diagonal VAR model, can still be estimated by ML using the EM algorithm modified with Cochrane–Orcutt iterations to estimate Θ conditional on Λ , and Λ conditional on Θ ; see Reis and Watson (2010) for an empirical implementation in a system with $N = 187$. Grassi et al. (2015) consider a system with $N = 170$ series, and Bai and Li (2016) conjecture what the limiting distribution of the parameters and factors should be.²³

²³ The procedure implemented by Reis and Watson (2010) is only valid when no missing values occur in Y_t ; see Jungbacker et al. (2011)

Note that, in this case, the innovation covariance matrix can still be inverted easily using (8) due to the diagonality of Σ_e .²⁴

Table 1 reports the sample MSE of the factor extracted from the same simulated systems described above generated by the DFM with autocorrelated idiosyncratic components when the parameters are estimated. Regardless of whether the true or misspecified models are estimated, we can observe that, if $N = 5$, the sample MSEs (coverages) are only slightly larger (smaller) than those of the corresponding models with known parameters. However, the sample MSEs (coverages), which are similar regardless of whether the true or the misspecified models are estimated, increase (decrease) with respect to those of the corresponding models with known parameters. When $N = 150$, the sample MSEs are very large and the coverage of the confident intervals is very low (recall that we are estimating the parameters with just $T = 200$ observations); see Fig. 5, which illustrates the figures reported in Table 1.²⁵

for a detailed discussion and computationally feasible solution of the missing value problem.

²⁴ The parameters of the S-DFM with autocorrelated idiosyncratic noise—written, as in footnote 9, by extending the state vector with the idiosyncratic noise—have been estimated by Coroneo et al. (2016) by EM, adding a small noise term to the measurement equation.

²⁵ We do not consider the estimation of the parameters of the DFM with cross-sectionally and serially correlated idiosyncratic noise because it is infeasible, due to the very large number of parameters to be estimated with just $T = 200$ observations.

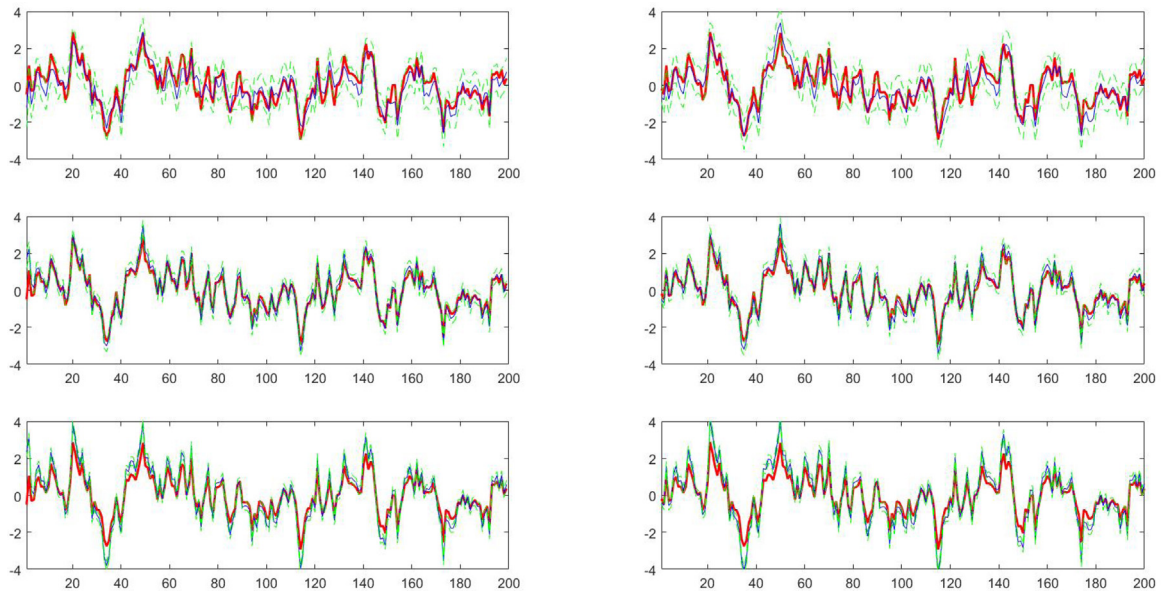


Fig. 5. Simulated factor (red line) together with a factor extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification (left column) and the true model, assuming that the idiosyncratic component is serially and cross-sectionally uncorrelated (right column), with both the parameters estimated by ML using the EM algorithm, and with the first based on the Cochrane–Orcutt specification. The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row) and serially and cross-sectionally correlated idiosyncratic components. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Following Doz et al. (2012), extracting the factors using the KFS based on QML parameter estimates is very popular in empirical applications; see, for example, Baíbura, Giannone, and Reichlin (2011) and den Reijer and Johansson (2019). This procedure has been extended by Baíbura and Modugno (2014) to deal with missing observations and idiosyncratic dynamics; see Scotti (2016) for an empirical application.²⁶ Marcellino and Sivec (2016) extend the procedure proposed by Doz et al. (2012) to a mixed-frequency factor-augmented VAR (MF-FAVAR) model.

Moving now to the D-DFM in Eqs. (34) and (35), note that QML estimation based on the EM algorithm is still possible, provided that the regressions in the maximization step are conveniently restricted; see, for example, Jungbacker and Koopman (2015) and Pinheiro et al. (2013). Barigozzi and Luciani (2020) extend the asymptotic results to the D-DFM written as an S-DFM in which the covariance matrix of the factor noise is singular.²⁷ Stock and Watson (2016a) also suggest estimating the D-DFM by imposing the restriction that Σ_u has rank q .

Finally, closely related to the QML estimator of Bai and Li (2016), Doz et al. (2012) propose a three-step estimator to derive the limiting distribution of the smoothed factors and estimate the dynamics of the idiosyncratic components. In the first step, Bai and Li (2016) propose

the joint estimation of the loadings and idiosyncratic variances, avoiding the specification of the factor dynamics by maximizing the following log-likelihood²⁸:

$$\log L(Y; \Psi) = \frac{1}{2N} \log |\Sigma_Y| - \frac{1}{2N} \text{tr}(M_Y \Sigma_Y^{-1}) \quad (44)$$

where $\Sigma_Y = \Lambda M_F \Lambda' + \Omega$ with $M_F = \frac{1}{T} \sum_{t=1}^T (F_t - \bar{F})(F_t - \bar{F})'$ and $\Omega = \text{diag}(E(\varepsilon\varepsilon'))$, where ε is the $N \times T$ matrix given by $\varepsilon = (\varepsilon_1, \dots, \varepsilon_T)$, and $M_Y = \frac{1}{T} \sum_{t=1}^T (Y_t - \bar{Y})(Y_t - \bar{Y})'$. Note that Σ_Y is an approximation of $E(M_Y)$ because Ω is restricted to be diagonal. The parameters to be estimated are $\Psi = (\Lambda, \Omega, M_Y)$. In the second step, the factors are estimated by GLS as follows:

$$\hat{F} = Y' \hat{\Omega}^{-1} \hat{\Lambda} \left(\hat{\Lambda}' \hat{\Omega}^{-1} \hat{\Lambda} \right)^{-1} \quad (45)$$

and the parameters of the VAR(p) model are estimated based on \hat{F} . Finally, in the last step, the Kalman smoother is evaluated using the previous parameter estimates.

4.2.2. Least squares estimation

Doz, Giannone, and Reichlin (2011) prove the consistency, when both N and T diverge to infinity, of the smoothed factors extracted when the parameters of the stationary S-DFM are substituted by two-step estimates. In the first step, the r factors are extracted using PCs. Denote the PC-extracted factors by \hat{f}_t^{PC} and the corresponding estimated loadings by $\hat{\Lambda}^{PC}$. In the second step,

²⁶ See Jungbacker et al. (2011) and Reis and Watson (2010) for early applications of DFM with missing observations.

²⁷ Barigozzi and Luciani (2020) show that, when the number of shocks is smaller than the number of factors, the QML estimates of the loadings are worse when estimated by PCs than when estimated by QML, while the factors are hardly affected.

²⁸ Bai and Li (2016) point out that, under fixed N , if cross-sectional heteroscedasticity exists but it is not allowed in the estimation, then the estimated factor loadings are inconsistent.

assuming that the dynamic dependence of the factors is described by a VAR(1) process, the autoregressive parameters can be estimated by OLS as follows:

$$\hat{\Lambda}^{PC} = \sum_{t=1}^T Y_t \hat{f}_t^{PC'} \left(\sum_{t=1}^T \hat{f}_t^{PC} \hat{f}_t^{PC'} \right)^{-1} \quad (46)$$

$$\hat{\Phi}^{PC} = \sum_{t=1}^T \hat{f}_t^{PC} \hat{f}_{t-1}^{PC'} \left(\sum_{t=1}^T \hat{f}_{t-1}^{PC} \hat{f}_{t-1}^{PC'} \right)^{-1} \quad (47)$$

The corresponding estimators of the covariance matrices are given by Eqs. (42) and (43), where $\hat{e}_t = Y_t - \hat{\Lambda}^{PC} \hat{f}_t^{PC}$ and $\hat{u}_t = \hat{f}_t^{PC} - \hat{\Phi}^{PC} \hat{f}_{t-1}^{PC}$. Doz et al. (2011) show that the smoothed factors extracted using the PC estimates of the parameters are consistent, even if the idiosyncratic component is wrongly assumed to be temporal and cross-sectionally uncorrelated due to the misspecification error vanishing as N and T diverge to infinity; see Angelini, Camba-Mendez, Giannone, Reichlin, and Rünstler (2011), Baïbura and Rünstler (2011), Giannone, Reichlin, and Sala (2005) and Giannone, Reichlin, and Small (2008) for implementations of this estimator. Koop and Korobilis (2014) extend the procedure proposed by Doz et al. (2011) to deal with time-varying parameters and stochastic volatility.

It is important to point out that when estimating using this two-step procedure, the restrictions imposed to identify the factors in PCs and those imposed in the SSM representation of the DFM should be the same. For example, very recently, Solberger and Spanberg (2020) implemented in E-Views the KFS factor extraction based on the two-step estimator of the parameters of the S-DFM. Although Solberger and Spanberg (2020) say that they can estimate the D-DFM, they are not taking into account the specification in the static representation of the D-DFM in (34) and (35); see, for example, Bai and Ng (2007) for the relation between the static factors extracted using PC and the dynamic factors.

Finally, note that the two-step estimator of the parameters has often been used to obtain starting values of the parameters in the EM algorithm; see, for example, Doz et al. (2012) and Proietti (2011). Joseph, Kalamara, Potjagailo, and Kapetanios (this issue) also use PC factors as starting values in the context of the estimator proposed by Kapetanios and Marcellino (2009).

5. Model specification

For KFS factor extraction to be efficient, it is crucial that the SSM is correctly specified. In this section, we describe some selected procedures used in the literature to specify the model, in particular, to determine the number of static and dynamic factors and their lags. We also describe procedures to decide the dependence structure of factors and idiosyncratic noise. In many works that implement KFS to extract latent factors in the context of DFMs, the specification of the factors is *ad hoc*. For example, in many works the factors and idiosyncratic components are assumed to follow $VAR(p_1)$ and $VAR(p_2)$ models, respectively, with p_1 and p_2 fixed *a priori*; see,

for example, Aruoba et al. (2009) and Proietti (2011) who consider DFMs with $p_1 = 1$ and $p_2 = 0$, or Frale, Marcellino, Mazzi, and Proietti (2011) who are interested in just one common factor, i.e. $r = 1$. Alternatively, p_1 can be determined by using information criteria as, for example, in Joseph et al. (this issue). In other works, the number of factors is determined *a priori* without using any specific criteria; see, for example, Aruoba et al. (2009) and Camacho and Perez-Quiros (2010) who, in the context of small-scale DFMs, fix $r = 1$, or Chauvet and Senyuz (2016) who fix $r = 2$. Delle Monache et al. (2016) and Koopman et al. (2010) also choose an *ad hoc* number of factors. Joseph et al. (this issue) determine the number of factors as the lowest one that explains 50% of the variance in the data. Also, in many works the specification of the model is chosen by minimizing a root mean square error criterion; see, for example, Hindrayanto, Koopman, and de Winter (2016) and Rünstler et al. (2009).

Alternatively, many authors implement methods for determining the number of factors designed in the context of PC methods. For example, in the context of S-DFMs, the number of static factors can be selected by using one of several procedures proposed in the literature; see Breitung and Eickmeier (2006), Stock and Watson (2011), and Barhoumi et al. (2013), among others, for more detailed and complete reviews. The most widely used methods to determine the number of factors in S-DFMs are due to Bai and Ng (2002); see, for example, Proietti (2011). The Bai and Ng (2002) criteria are based on modifications of the Akaike and Bayesian information criteria, taking into account the cross-sectional and temporal dimensions of the dataset as arguments of the function penalizing overparameterization. However, these criteria usually detect too many factors, as they are sensitive to the choice of r_{\max} , i.e. the maximum number of factors; see, for example, the Monte Carlo results in Ahn and Horenstein (2013) and the arguments by Hindrayanto et al. (2016). Alternatively, Alessi, Barigozzi, and Capasso (2010) refine the Akaike and Bayesian criteria proposed by Bai and Ng (2002) by multiplying the penalty function by a constant that tunes the penalizing power of the function itself. Furthermore, Alessi et al. (2010) suggest estimating the number of factors using different subsamples. These criteria are linked to the eigenvalues of the sample covariance matrix of the variables in the system. In particular, the number of factors is selected as the number of eigenvalues larger than a threshold specified by a penalty function.

Another useful procedure to select the number of static factors is due to Onatski (2010), who proposes selecting r using the difference between the eigenvalues of $\frac{1}{T} Y'Y$ and proves that this selection is consistent. Note that, in spite of the lack of consistency of the eigenvalues when both N and T go together to infinity, as pointed out by Lam and Yao (2012) through a simulation exercise, the empirical evidence and simulations results obtained by Corona, Poncela, and Ruiz (2017) indicate that the difference between adjacent eigenvalues is a good estimator of the number of common factors. The procedure works even when the proportion of the variance attributed to the factors is small relative to the variance,

due to idiosyncratic noise, or when the noise terms are substantially correlated. Under the assumption of normality, both cross-sectional and temporal dependence in the idiosyncratic noise are allowed. In a way, this procedure formalizes the eyeball decisions taken when examining the scree plot of the eigenvalues, as proposed by Cattell (1966). The intuition behind the method is as follows: the line connecting two adjacent zero eigenvalues has zero slope, while that connecting eigenvalues linked to common factors should have a slope different from zero. This procedure is based on determining a sharp threshold that consistently separates the bounded and diverging eigenvalues of the sample covariance matrix of the observed series.

Finally, Lam and Yao (2012) propose a procedure to determine the number of factors based on separating strong and weak factors that require the idiosyncratic components to be white noise. They propose determining the number of factors by the number of non-zero eigenvalues of the following matrix:

$$S_Y = \sum_{k=1}^K C_Y(k)C'_Y(k), \quad (48)$$

where $C_Y(k) = \text{Cov}(Y_t, Y_{t-k})$ and K should be small, as the autocorrelation is often at its strongest at the small lags.

Finally, Choi and Jeong (2019) propose additional criteria to determine r . Further, they carried out detailed Monte Carlo experiments to compare the performance of the more popular criteria available in the literature. They show that it is difficult to conclude which criterion performs best, and they advise that, in empirical applications, one should consider several criteria at the same time. Along the same line, Stock and Watson (2011) point out that, when dealing with empirical systems, different methods frequently determine a different number of static factors, with limited research comparing the performance of the different methods. Consequently, they suggest augmenting the statistical estimators of r by inspecting the scree plots and with judgment informed by the application at hand; see, for example, Hindrayanto et al. (2016) and Schiavoni, Palm, Smeekes, and van der Brakel (2019) for empirical implementations.

In the context of D-DFMs, the first issue faced when trying to identify the number of dynamic factors is related to the simultaneous identification of the lag order of the VAR model for the factors, p , and the number of lags, s . To illustrate this problem, consider the following D-DFM, in which the common component is given in Eq. (28):

$$Y_t = \Lambda(L)G_t + \varepsilon_t \quad (49)$$

$$G_t = \Phi_1 G_{t-1} + \cdots + \Phi_p G_{t-p} + v_t. \quad (50)$$

The D-DFM can be written as follows:

$$Y_t = \Pi(L)v_t + \varepsilon_t \quad (51)$$

where $\Pi(L) = \Lambda(L)\Phi(L)^{-1}$ with $\Phi(L) = I_r - \Phi_1 L - \Phi_2 L^2 - \cdots - \Phi_p L^p$; see Sargent and Sims (1977) for an early proposal of the model in (51). The model in Eq. (51) is known as the generalized DFM (G-DFM) and the parameters of the infinite lag polynomial $\Pi(L)$ matrix cannot be

estimated by ML or QML without further restrictions. Several authors propose estimating them using dynamic PCs based on frequency-domain procedures; see Forni, Hallin, Lippi, and Reichlin (2004, 2005) and Forni, Hallin, Lippi, and Zaffaroni (2015, 2017). Recently, Peña, Smuler, and Yohai (2019) proposed estimating the G-DFM by linear generalized PCs in the time domain.²⁹ The identification problem appears because, from $\Pi(L)$, it is not possible to recover the polynomials $\Lambda(L)$ and $\Phi(L)$ in a unique form without imposing restrictions; see, for example, the discussion in Lütkepohl (2005). In order to identify the model, Forni, Hallin, Lippi, and Reichlin (2000) consider restrictions on the eigenvalues of the spectral density matrix, and Hallin and Liska (2007) and Onatski (2009) propose procedures to determine q in the context of the generalized DFM.

Alternatively, Stock and Watson (2005) propose determining the number of common dynamic factors, q , by considering the static representation. Consider, for example, the D-DFM in Eqs. (34) and (35), which can be written as follows:

$$Y_t = \Lambda F_t + \varepsilon_t \quad (52)$$

$$F_t = \Phi F_{t-1} + B v_t. \quad (53)$$

where $\Lambda = [\Lambda_0 \Lambda_1]$ and $\Phi = \begin{pmatrix} \Phi_1 & 0 \\ I_q & 0 \end{pmatrix}$ and $B = \begin{pmatrix} I_q \\ 0 \end{pmatrix}$. After substituting (53) into (52), define $X_t = Y_t - \Lambda \Phi F_{t-1}$, which can be written as follows:

$$X_t = \Pi v_t + \varepsilon_t, \quad (54)$$

where $\Pi = \Lambda B$. In Eq. (54), X_t is represented as a factor model with q serially uncorrelated factors that correspond to the shocks, v_t . Were X_t observed data, Stock and Watson (2005) argue that q could be consistently determined using the Bai and Ng (2002) information criteria. However, given that this is infeasible, they propose applying Bai and Ng (2002) to $\hat{X} = Y_t - \hat{\Sigma} \hat{F}_{t-1}$, where $\hat{\Sigma}$ is an estimator of $\Lambda \Phi$, and \hat{F}_{t-1} is an estimator of F_{t-1} . Stock and Watson (2005) propose estimating $\hat{\Sigma}$ and \hat{F}_{t-1} by first estimating Λ and F_t by PCs, and then Φ by the OLS estimator of the regression of \hat{F}_t onto $(\hat{F}_{t-1}, \dots, \hat{F}_{t-p})$, where p can be chosen using the Bayesian information criterion.³⁰ Amengual and Watson (2007) prove the consistency of this procedure. Alternatively, Bai and Ng (2007) propose determining q by computing the eigenvalues of the covariance matrix of the residuals of the regression of \hat{F}_t

²⁹ Stock and Watson (2005) point out that dynamic PCs produce two-sided estimates of the factors and, consequently, these estimates are not suitable for forecasting. Rünstler et al. (2009) compare the forecast performance of G-DFMs with those of alternative DDFMs, and find evidence in favor of the G-DFM with the gains in forecast precision against the alternative factor models being rather small and statistically insignificant. D'Agostino and Giannone (2012) also compare static and dynamic PCs by forecasting inflation, and conclude that both are similar. Recently, Peña et al. (2019) proposed one-sided dynamic PCs that are appropriate for forecasting.

³⁰ Alternatively, Stock and Watson (2005) propose directly estimating \hat{X}_t using the OLS estimator from the regression of Y_t onto $(\hat{F}_{t-1}, \dots, \hat{F}_{t-p})$.

onto $(\hat{F}_{t-1}, \dots, \hat{F}_{t-p})$ and testing whether these eigenvalues satisfy an asymptotically shrinking bound that reflects sampling error. Note that the results of the test may depend on this shrinking bound.³¹ Breitung and Pigorsch (2013) propose determining the number of dynamic factors, q , using canonical correlation analysis of the current and past values of the common factors in the static representation instead of PCs. The procedure proposed by Breitung and Pigorsch (2013) also depends on tuning parameters. Zhao, Cui, and Wang (2017) carried out a Monte Carlo comparison of alternative procedures to determine the number of dynamic factors, and concluded that the procedures proposed by Bai and Ng (2007) and Hallin and Liska (2007) perform better.

These procedures are very popular. For example, Angolini et al. (2011) determine r using Bai and Ng (2002), p using Schwarz information criteria, and q using Bai and Ng (2007).³² However, it is important to recall that the procedures for determining q have been designed in the context of PC factor extraction and that the assumptions behind PCs could not be satisfied when the D-DFM is written as an S-DFM. Furthermore, when the data follow a D-DFM and we fit an S-DFM, the number of factors increase from q to $r = q(s + 1)$, but these r factors are not mutually independent at all lags. Consequently, it could be expected that the number of "static" factors determined by the procedures designed for truly static S-DFMs is going to be such that $r < q(s + 1)$; see the arguments in Peña and Tsay (2020).

The discussion about the determination of r and q shows that, in practice, this is a difficult problem.³³ Furthermore, in the case of the D-DFM, even if we were able to chose r and q , we do not have enough information to recover the specification in Eqs. (34) and (35). The procedures to determine the number of factors, r , when the D-DFM is written as a "static" DFM, described above, are based on a different specification, in which the autoregressive matrices of the factors are not restricted. Therefore, when estimating the parameters and running the KFS algorithms, the estimated model is not exactly the true specification.

The difficulty in choosing the correct specification of the DFM that is then used for factor extraction (after estimating the unknown parameters) calls for adequate procedures for testing for the specification of the model. Harvey and Koopman (1992) propose some specification tests based on auxiliary residuals. However, these tests were implemented to univariate models. Extensions to multivariate models, such as those of interest in this survey, are needed in the literature. Recently, Fiorentini and

Sentana (2019) derived score tests of misspecification in DFM based on frequency domain techniques.

6. Forecasting using KFS in stationary DFM: Empirical applications

The number of empirical applications of stationary DFM using KFS for factor extraction is very extensive. In this subsection, we survey some empirical applications to show the large range of possibilities of these models and procedures. We apologize for those important works that have not been mentioned in this subsection.

One of the first implementations of KFS algorithms to extract factors in the context of DFM was Engle and Watson (1981), who estimated the unobserved metropolitan wage rate for Los Angeles based on observations of sectoral wages within the Standard Metropolitan Statistical Area. Afterwards, in the last 40 years, the number of applications of KFS algorithms in DFM has been increasing over time. In this section, we describe just some of the main contributions.

Among the many applications of forecasting using KFS procedures within the context of factor extraction in DFM, one of the most active areas is real-time macro-monitoring; see Stock and Watson (2016a) for a description of empirical applications. The first relevant contribution in this area is due to Stock and Watson (1989, 1991) who, within the context of a small-dimensional DFM, construct the experimental coincident economic indicator (CEI), which was released monthly through the National Bureau of Economic Research from May 1989 to December 2003. The CEI was the Kalman filter estimate of the common factors among the big monthly indicators often used in dating the business cycle. The DFM was estimated by ML using an SSM; see Carriero and Marcellino (2007) who compare several parametric and non-parametric procedures to extract a single common factor in the context of estimating the composite coincident and leading indexes for the UK. Since then, several applications have appeared in the literature. Aruoba et al. (2009) track the high-frequency evolution of real activity by constructing an index based on indicators measured at different frequencies, even very high-frequency (daily) data. This index, updated daily by the Federal Reserve Bank of Philadelphia, has been added to Bloomberg's real-time data and can be followed on its platform. Based on the index proposed by Aruoba et al. (2009), Scotti (2016) constructs uncertainty indexes based on smoothed weights estimated using the method proposed by Koopman and Harvey (2003). Camacho and Perez-Quiros (2010) evaluate the short-term forecasts from a factor model in a truly real-time setup for the euro area. They deal with ragged edges using the proposal by Giannone et al. (2008), with mixed frequencies using the filter proposed by Mariano and Murasawa (2003) and with data revisions using Evans (2005). Aruoba and Diebold (2010) propose an extension to the DFM suggested by Stock and Watson (1991). Grassi et al. (2015) construct monthly indicators of economic activity for the euro area and its largest member countries. Very recently, Diebold (2020) implemented the index proposed

³¹ The procedure proposed by Bai and Ng (2007) is a useful cross-check of the more informal procedure proposed by Giannone et al. (2005).

³² In an application to euro area data, they chose $r = 5$, $q = 3$, and $p = 1$.

³³ It is important to point out that, in an empirical exercise, D'Agostino and Giannone (2012) conclude that there is no evident improvement in the forecast accuracy when allowing for the number of dynamic factors to be smaller than the number of static factors.

by Aruoba et al. (2009) to monitor the evolution of US growth during the Covid-19 pandemic.

Closely related to the construction of business-cycle indexes is the problem of nowcasting, which is important since forecasting improvements of GDP with respect to naive models are mainly limited to the current quarter; see Baíbura et al. (2011) for a survey on nowcasting. Factors extracted using KFS algorithms can be used to forecast in the context of the diffusion indexes of Stock and Watson (2002). Baíbura and Rünstler (2011) propose obtaining prediction weights of the individual time series in a system by an extension of the Kalman filter using the estimation procedure of Doz et al. (2011). Rünstler (2016) proposes using these prediction weights to refine the data set by eliminating uninformative series when forecasting using DFM. KFS has been implemented to exploit large information to bridge monthly and quarterly variables with different publication lags, by combining predictors in few common factors, which are then used as regressors in bridge equations via the Kalman filter. The procedure, originally proposed by Giannone et al. (2008), was first applied to US data at the Board of Governors of the Federal Reserve and is also regularly implemented at the ECB; see Rüstler et al. (2009) for a comparison of forecasts of GDP obtained in ten countries of the euro area, Angelini et al. (2011), Matheson (2010) for early estimates of current-quarter GDP in the euro area, D'Agostino, McQuinn, and O'Brien (2012) for Irish quarterly GDP, and Aastveit and Trovik (2012) who investigate the properties of financial assets to forecast growth in Norway. Hindrayanto et al. (2016) compare the performance of four different estimators of the factors in a pseudo-real-time competition to forecasts GDP growth in the euro area and five of its largest countries. In particular, they compare forecasts when the factors are estimated using PC and the two procedures of Doz et al. (2011, 2012), and Bräuning and Koopman (2014). These last authors propose adopting a low-dimensional SSM that jointly deals with the target variable and the factors. More recently, Jansen and de Winter (2018) explored the effects of combining model-based nowcasting GDP forecasts and quarterly judgmental consensus forecasts for the G7 countries. It is important to note that, due to the difficulty in choosing the correct specification of the model, several authors propose pooling the forecasts obtained from different models with a different number of factors; see, for example, Hindrayanto et al. (2016), Kuzin, Marcellino, and Schumacher (2013) and Jansen, Jiu, and de Winter (2016). Camacho and Martinez-Martin (2014) describe an application in which a stationary small-scale DFM is implemented to forecast US GDP with a system of variables that includes survey data and financial indicators.

Apart from monitoring and forecasting real economic activity, central banks, researchers, and analysts have widely used DFM to track and forecast the evolution of inflation, or to detect comovements in prices, especially in goods or commodities. To cite just a few applications, Aruoba and Diebold (2010) use small-scale DFM to monitor both real economic activity and inflation, taking into account their interactions. Reis and Watson (2010)

use quarterly inflation for $N = 187$ sectors in US personal consumption expenditures to extract what they call “pure” inflation and separate it from other components of inflation. Inflation in commodities has been analyzed in Delle Chiaie, Ferrara, and Giannone (2017) and Poncela et al. (2020).

Baíbura and Modugno (2014) and Giannone et al. (2008) use SSMs for nowcasting—the problem of predicting the present, the very near future, and the very recent past, which is related to both missing observations and mixing frequencies. Marcellino and Schumacher (2010) and Foroni and Marcellino (2014) compare different methods to deal with mixed frequencies when forecasting German GDP and nowcasting euro area macroeconomic aggregates, respectively. Bok, Giannone, and Tambalotti (2018) fit a DFM with 37 predictors at different frequencies for nowcasting US GDP.

Another important application of factor extraction in forecasting is the factor-augmented VAR (FAVAR) model. FAVAR models have enjoyed increasing popularity in forecasting macroeconomic variables. Koop and Korobilis (2014) propose a FAVAR model with time-varying parameters (TVP-FAVAR) to construct a financial-conditions index to track expectations about growth in key US macroeconomic variables. Marcellino and Sivec (2016) implement an MF-FAVAR model to evaluate the effects of monetary, oil, and fiscal shocks. Very recently, Joseph et al. (this issue) implemented FAVAR models to forecast UK inflation and compared several alternative procedures.

KFS procedures have also been implemented when forecasting in the context of financial variables. One of the most popular applications is to forecast the yield curve. Notable examples include Diebold and Li (2006) and Diebold, Rudebusch, and Aruoba (2006) who describe the yield curve by framing the Nelson–Siegel model into an SSM with three unobserved factors that represent the level, slope, and curvature and that are modeled as VAR processes with the loadings that depend on a single parameter. Koopman et al. (2010) forecast interest rates over different maturities based on this model extended by allowing the loading parameter to vary over time and the variances to follow GARCH models.³⁴ Jungbacker, Koopman, and van der Wel (2014) impose smoothness restrictions on factor loadings by using a cubic spline function that depends on time to maturity, and Koopman and Van der Wel (2013) include macroeconomic factors in the dynamic yield curve model and conclude that macroeconomic variables can lead to more accurate yield curve forecasts.³⁵ In a different context, Coroneo et al. (2016) implement the methodology of Doz et al. (2012) to forecast monthly excess bond returns based on out-of-sample forecasts of the underlying yield and macro-economic factors. Another strand of the literature focuses on extracting

³⁴ Note that models with time-varying loadings could be considered as non-stationary. However, given that the factors are stationary, their treatment, from the point of view of factor extraction, is the same as in stationary models.

³⁵ Poncela (2013) note that this increase in accuracy may be the effect of the model specification instead of the effect of the added macroeconomic variables.

information from the components of the yield curve to forecast business-cycle turning points; see Chauvet and Senyuz (2016) who fit a DFM with $r = 2$ with this purpose. An application of the QML procedure is proposed by Doz et al. (2012) to construct financial conditions indexes as, for example, the Chicago Fed's National Conditions Index (NFCI). The NFCI provides a weekly update on US financial conditions in money markets, debt, and equity markets and the traditional and “shadow” banking systems; see Brave and Butters (2012) for a description of the NFCI. Matheson (2012) also extracts financial indexes for the US and the euro area using the estimation procedure proposed by Doz et al. (2011), while Menden and Proaño (2017) use the EM algorithm to estimate a financial index based on a DFM for the US. Finally, a very interesting application of KFS to extract factors is carried out by Haugh and Ruiz Lacedelli (2020) who carry out scenario analysis to measure the risk associate with derivative portfolios.

Camacho, dal Bianco, and Martínez-Martín (2015a, 2015b), Camacho and Perez-Quiros (2010), Jungbacker et al. (2011), and Mariano and Murasawa (2010) also implement SSMS with missing data. Very recently, Schiavoni et al. (2019) incorporated Google Trends information into the estimation of unobserved components to improve official unemployment statistics using KFS with mixed frequencies. Martínez-Martín and Rusticelli (in press) also implement the methodology of Mariano and Murasawa (2010).

Proietti (2011) considers a large-scale ($N = 149$, $T = 150$) static stationary exact DFM to extract factors for the euro area in which the series in the system are observed monthly and quarterly to deal with the aggregation problem.

Finally, there are also a large number of applications for non-economic data. Just to name one included in this special issue, Coulombe, Diebold, Gobel, Rudebusch, and Zhang (this issue) extract the common factor from a system of four sea-ice measures to obtain an optimal composite measure.

7. Non-stationary DFMs

So far, DFMs have been assumed to be stationary. However, panel time series are often non-stationary due to the presence of unit roots. In this case, it is very popular to differentiate individually each series in a univariate fashion in the system until stationarity is reached. Then, the corresponding DFM is fitted to the differentiated system to extract the factors and the accumulated factors are obtained if needed; see, for example, Mariano and Murasawa (2003, 2010) and Grassi et al. (2015), among many others. This solution might lead to a loss of information, missclassifications of factors as stationary/non-stationary, and inefficient estimations of the parameters and factors; see, for instance, Barigozzi and Luciani (2019), and Corona, Poncela, and Ruiz (2020). Alternatively, the DFM can be implemented directly to the original non-stationary system. This section surveys some relevant issues specific to non-stationary DFMs when the KFS algorithms are implemented to extract the factors. First,

we consider the case of known models. Second, we consider that the DFM specification is known but the parameters need to be estimated. Finally, we deal with specification issues. The last subsection reviews selected empirical applications of non-stationary DFMs in which KFS procedures are implemented to extract the factors.

7.1. Factor extraction in non-stationary DFMs with known parameters

If the model specification and the parameters are known, the KFS can be implemented to extract the factors in non-stationary DFMs without new issues on top of those already described for their stationary counterparts. There are two considerations worth mentioning. First, the role of the starting values for the state vector (factors) should be different from those usually used in stationary models. Second, it is important to investigate the performance of KFS extraction when the idiosyncratic noise is non-stationary but temporal dependence is ignored when filtering. Consider first that $Y_t = (Y_{1t}, \dots, Y_{Nt})$, $t = 1, \dots, T$ is a non-stationary $I(1)$ vector of time series generated by the DFM in (30),

where F_t is an $r \times 1$ vector of $I(1)$ common factors and the idiosyncratic components are stationary.³⁶ Therefore, there are N non-stationary $I(1)$ time series generated by $r < N$ non-stationary $I(1)$ common trends. For simplicity, consider that the VAR model for the r common non-stationary factors is as in Eq. (28) with $r = r_1$, $p = 1$, and $\Phi_1 = I_r$. The error u_t is white noise with diagonal variance matrix Σ_u . We also assume that the idiosyncratic components are white noise with diagonal variance matrix R . The Kalman filter can be applied in order to extract the common factors using the initial conditions for the state vector suggested in Peña and Poncela (2006) with $f_0 = \Lambda^{-}Y_1$ being Λ^{-} , a generalized inverse of Λ , and with $P_0 = kI_r$, where the scalar k is such that $k \rightarrow \infty$. Additional possibilities for initializing the Kalman filter in the presence of non-stationary latent variables are suggested, for instance, in Ansley and Kohn (1985), de Jong (1991), and Koopman (1997). To illustrate the performance of the KFS in this case, we simulate a system with $r = 1$ factor characterized by a random walk with zero initial condition, $F_0 = 0$, and $\sigma_u^2 = 1$. The idiosyncratic noise is serially and cross-sectionally uncorrelated and homoscedastic with variances equal to 1. As in the stationary case, we simulate systems with $T = 200$ observations and $N = 5, 50$, and 150 variables to replicate small, medium, and large systems, respectively.

³⁶ We assume that there are not deterministic components. In systems where there are, Bai (2004) suggests eliminating all the deterministic components before applying non-stationary factor analysis to the series in levels. Furthermore, Banerjee and Macellino (2017) argue that the assumption of stationary idiosyncratic noise is in accordance with macroeconomic systems that exhibit a high rejection of the hypothesis of a unit root of the estimated idiosyncratic components; see, for example, Poncela and García-Ferrer (2014), and Schiavoni et al. (2019) for empirical implementations with stationary idiosyncratic components. Finally, Barigozzi and Luciani (2019) also contemplate the possibility of lags of the common factors in the measurement equation. This can be dealt with as in the stationary case.

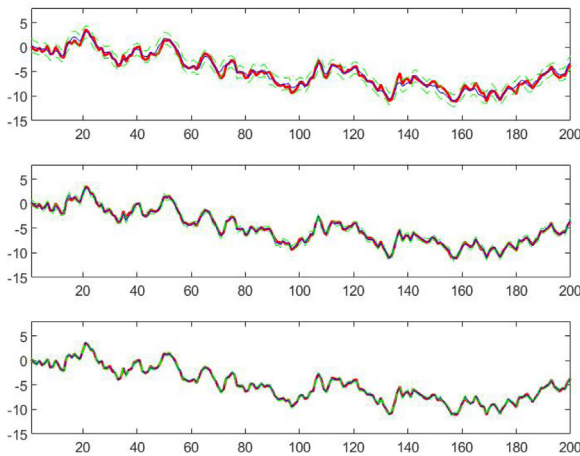


Fig. 6. Simulated factor (red line) together with the factor extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification and known parameters for a random walk common factor. The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

The factor loadings are drawn from a uniform distribution $U(0, 1)$. Fig. 6 plots the true simulated factor together with the factor extracted using KFS assuming all parameters are known and using the initial conditions obtained as suggested by Peña and Poncela (2006). As for the stationary case, the differences between the extracted and population factors are only visually appreciable when $N = 5$.

Instead of considering a DFM with all factors being $I(1)$, it is possible to face situations in which some factors are non-stationary while others are stationary; see Barigozzi and Luciani (2017), García-Ferrer and Poncela (2002) and Peña and Poncela (2006) among others. Assume that there are r_1 factors that are $I(1)$ and r_0 stationary factors, with $r = r_1 + r_0$ being the total number of common factors.³⁷ The model is given by (26) to (28) with r_1 roots of the determinantal equation, $|\Phi(L)| = |I_r - \Phi_1 L - \dots - \Phi_p L^p| = 0$, on the unit circle, and r_0 outside it. Usually, the variance associated with the common non-stationary factors is higher than that linked to the stationary ones. In order to check how both types of common factors are extracted, we perform a simulation considering $r = 2$ with one factor generated by a random walk with $\sigma_{u_1}^2 = 1$ and the other by a stationary AR(1) model with autoregressive parameter $\phi = 0.5$ and $\sigma_{u_2}^2 = 1$. Therefore, in (28), $p = 1$ and $\Phi_1 = \text{diag}(1; 0.5)$. Fig. 7 plots the true simulated factors together with their corresponding extracted factors and 95% confidence bounds. Notice that, as before, if the model is known, the differences between the extracted and population factors are only visually appreciable when $N = 5$ for both factors.

³⁷ Higher orders of integration have been considered, for instance, in Peña and Poncela (2006).

Finally, we consider other sources of non-stationarity in the system besides the $I(1)$ common factors, which may render spurious common factors if they are not properly taken into consideration. In particular, we consider the case of just $r = r_1$ non-stationary factors with some or all of the idiosyncratic components also being random walks. If all the idiosyncratic components are $I(1)$, the observed time series are not cointegrated, while if just $g_1 < N$ idiosyncratic components are $I(1)$, there is cointegration among some of the observed series. Notice that if all the idiosyncratic components are random walks, the observed time series can be univariately differenced and the analysis can be performed in first differences. For illustrative purposes consider now the model given in Eqs. (26)–(29), where the VAR for the common non-stationary factors is as in (28) with $r = r_1$, $p = 1$, and $\Phi_1 = I_r$. The error u_t associated with the common factors is white noise with diagonal variance matrix Σ_u . As for the stationary case, from now on, we assume that $p^* = p_i^* = 1$ in Eq. (29) in order to simplify the analysis, where either $\theta_{1i} = 1$ when the idiosyncratic components are also non-stationary, or $|\theta_{1i}| < 1$ if they are stationary.

We assume that the idiosyncratic components are cross-sectionally uncorrelated. Barigozzi, Lippi, and Luciani (2020) argue that when $\theta_{1i} < 1$, the dynamics of the $I(0)$ idiosyncratic components do not need to be specified in order to obtain consistent estimates of the common components. For illustrative purposes, we simulate a system generated by only one common random walk ($r = r_1 = 1$) where the first $N/2$ idiosyncratic components are also random walks if N is even, and the first $N/2 - 1$ if N is odd. The remaining idiosyncratic components are stationary with parameter $\theta_{1i} = 0.5$. The model can be written as in Eqs. (31) and (32) with $\Phi_1 = 1$ and $\Theta = \text{diag}(\theta_{11}, \dots, \theta_{1N})$. We extract the common factor assuming that we know the true model. The results are presented in Fig. 8, left column. Only for $N = 5$, we can see by visual inspection the difference between the estimated and the true common factor.³⁸ In Fig. 8, centered column, we replaced the true AR parameter of the idiosyncratic component whenever it was stationary (that is, when $\theta_{1i} = 0.5$) by 0. As it can be seen, differences between the true and estimated common factors are only visually appreciable for $N = 5$, as for the case of using the true model. Finally, we ignore all dynamics in the idiosyncratic component, stationary or not, and extract the common nonstationary factor using a further misspecified model, considering that the specific components are all stationary white noise. As pointed out by Bai (2004) and Onatski and Wang (2020) for principal components, and Barigozzi and Luciani (2019) for KFS methods, the extracted factor in this case is not consistent, as it is reflected in the illustration presented in Fig. 8, right column, where we can see important differences between the true and estimated common factors for all cross-sectional dimensions $N = 5, 50$, and 150 . Therefore,

³⁸ Barigozzi and Luciani (2019) argue that estimating a misspecified model while ignoring the dynamics of the idiosyncratic components when they are stationary does not affect the consistency results for the estimated common components.

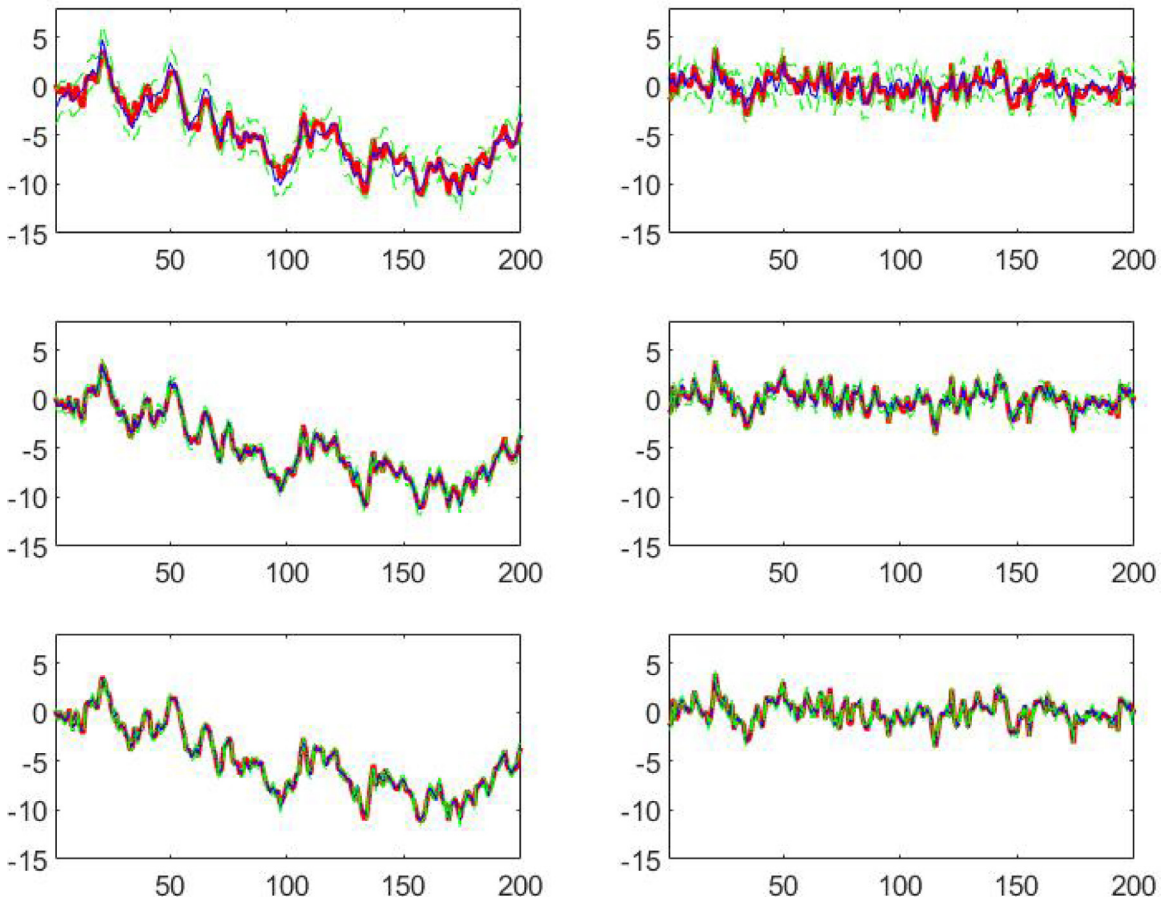


Fig. 7. Simulated factors (red line) together with factors extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification and known parameters for a random walk (left column) and a stationary AR(1) (right column) common factor. The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

misspecification of the idiosyncratic components still renders consistent estimates of the common factor as long as the misspecification is only associated to the stationary part, while misspecifications related to the nonstationary idiosyncratic components break the consistency of the extracted common factors. The adverse effects seem to be less important as the cross-sectional dimension increases in the simulated example.

7.2. Factor extraction in non-stationary DFMs with unknown parameters

As for the stationary case, usually the parameters of the model are unknown and need to be estimated. However, there is an important difference between the identification restrictions in DFMs with non-stationary factors and those described above for the stationary case. If we are interested in extracting a particular set of nonstationary common factors, the identification restrictions need to ensure the superconsistency of the estimators of the factor loading matrix. Johansen and Tabor (2017) analyze an SSM with an unobserved multivariate random walk and check whether the common components extracted with

known parameters and the corresponding components extracted with estimated parameters are cointegrated. They conclude that, although cointegration holds for the common components, $\chi_{i,t}$, $i = 1, \dots, N$, it is not always the case for the common trends. In the context of the ML estimator of the parameters of an autoregressive representation of the SSM model, they show that the extracted trends obtained with the true and estimated parameters are cointegrated if and only if the estimators of the factor loadings converge to the population parameters at a faster rate than \sqrt{T} .

Once the non-stationary factors are properly identified, the DFM parameters can be estimated as in the stationary case.³⁹ Most works dealing with KFS procedures for factor extraction in non-stationary DFMs obtain parameter estimates based on ML methods with the log-likelihood maximized either using Newton-type optimization algorithms as in Fraile et al. (2011) or using the EM algorithm as in

³⁹ Obviously, the non-stationary nature of the data should be taken into account with the unit roots imposed before estimating the parameters. Recall that, in this subsection, we consider that the DFM specification is known.

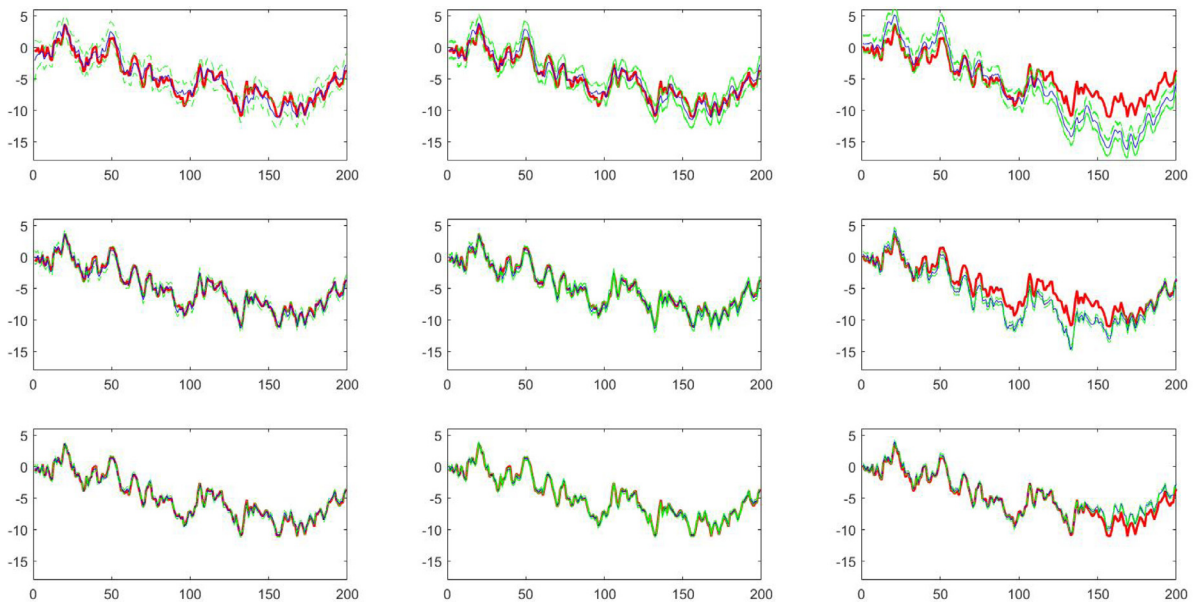


Fig. 8. Simulated factors (red line) together with factors extracted by the Kalman smoother (blue lines) and 95% confidence bounds (green lines) obtained using the true specification and known parameters for a random walk and possibly nonstationary idiosyncratic errors when using the true model (left column), a misspecified model only for the stationary idiosyncratic components (center column) assuming that they are white noise, and a misspecified model for the idiosyncratic components, stationary or nonstationary, assuming that they are white noise (right column). The systems are simulated by a DFM with $N = 5$ (first row), $N = 50$ (second row), and $N = 150$ (third row). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Peña and Poncela (2004), Zuur, Fryer, Jolliffe, Dekker, and Beukema (2003) and Seong, Ahn, and Zadrozny (2013) who consider systems with small cross-sectional dimension, in particular, $N = 12$ in the first work, and $N = 5$ in the last two works. The EM algorithm is also able to handle large N systems; see, for example, Barigozzi and Luciani (2017, 2019) and Quah and Sargent (1993). Barigozzi and Luciani (2019) provide consistency results for the common components for the model estimated using the EM algorithm and KFS when both N and T go to infinity.

There are two important issues related to implementing the EM algorithm in non-stationary DFMs. First, it is important to point out that the estimation procedures described above rely on the idiosyncratic noise being stationary. However, even if the factors were observed, if unit roots are allowed in the idiosyncratic components, it is no longer possible to “regress” the observed time series over the factors to estimate the factor loadings as in Eq. (40) of the EM algorithm. Alternatively, if the idiosyncratic component for the i th series is non-stationary, Barigozzi and Luciani (2019) propose replacing $Y_{i,t}$ by $Y_{i,t} - \varepsilon_{i,t}$ in order to remove the sources of non-stationarity not coming from the common factors from $Y_{i,t}$, and adding some additional noise.

The second issue relates to the parameters used to initialize the EM algorithm. First, consider the initial values for the loading matrix, Λ . Peña and Poncela (2006) assume that the non-stationarity is only present in the common factors and suggest as initial values of the columns of

the factor loading matrix, Λ , the eigenvectors of the first-lag autocovariance or uncentered lagged second-moments matrix of the observed series properly normalized by T^{2d} , instead of T where d is the order of integration of the observed series. The idea is that, if there is no serial correlation in the idiosyncratic noise or it is very weak, the eigenvectors of the lagged covariance matrices will be a better estimate of the factor loading matrix since they would not be contaminated by the zero lag correlation present in the noise. Alternatively, Barigozzi, Lippi, and Luciani (2021) and Barigozzi and Luciani (2019) allow for $I(1)$ idiosyncratic noise and suggest using the eigenvectors of the covariance matrix of the model in first differences in order to avoid spurious effects due to unit roots in the idiosyncratic noise.

The initial values for the parameters of the state equation needed for the ML estimator can be obtained by applying LS to the consistent PC common factors. For instance, Barigozzi et al. (2021) use data on first differences to estimate the factor loading matrix but project the levels of the series over those estimates of the factor loading matrix to get pre-estimators of the common factors.

Alternatively, Johansen and Tabor (2017) consider an LS estimator based on using the cointegration implications of the non-stationary DFM; the original LS estimator for multivariate non-stationary systems was proposed by Harvey and Koopman (1997).

7.3. Specification of non-stationary DFM

As in the stationary case, we need to specify the model and write it in state-space form prior to estimating the parameters and factor extraction. The main issue in non-stationary DFM is to determine the number of non-stationary and stationary common factors; see Barhoumi et al. (2013) who review procedures for the determination of the number of factors in non-stationary systems. It is important to note that, in empirical applications, it is often the case that the variance associated to the non-stationary common factors is much larger than that linked to the stationary factors and, consequently, the later ones may be difficult to detect; see Peña and Poncela (2006). In practice, if the number of stationary and non-stationary factors cannot be determined simultaneously, one should first determine the latter and, afterwards, the former. In order to determine the number of factors, it is important to consider three possible scenarios regarding the idiosyncratic components: (i) all of them are non-stationary, (ii) all of them are stationary, and (iii) a mixture of both, some of them are non-stationary while some others are stationary. If all idiosyncratic components were non-stationary, then the number of common factors could be determined after differentiation of the original systems of variables. In what follows we review some of the procedures to set up the number of common factors when some of them are non-stationary.⁴⁰

First, assuming that the idiosyncratic component is stationary, Peña and Poncela (2006) determine the number of common non-stationary factors as the number of nonzero eigenvalues of the following generalized covariance matrices:

$$C_Y(k) = \frac{1}{T^\alpha} \sum_{t=k+1}^T (Y_{t-k} - \bar{Y})(Y_t - \bar{Y})' \quad (55)$$

where $\alpha = 2d$ if $d \geq 1$, while $\alpha = 1$ when $d = 0$, and $\bar{Y} = \frac{1}{T} \sum_{t=1}^T Y_t$ is the time average of the vector of observed series. The procedures for determining the number of stationary common factors can then be implemented to $Y_t^* = Y_t - \Lambda_1(\Lambda_1' \Lambda_1)^{-1} \Lambda_1' Y_t$, where Λ_1 is the matrix of weights corresponding to the non-stationary factors. Although Peña and Poncela (2006) analyze the properties of this procedure in small cross-sectional dimension, Lam and Yao (2012) suggest that it could also be applied in models with large N . Peña and Poncela (2006) also argue that, instead of the generalized covariance matrices, the following generalized sample second moment matrices

⁴⁰ Notice that, in cases (ii) and (iii), there are cointegration relationships among the series in the system Y_t . For finite N , Escribano and Peña (1994) show that, if the idiosyncratic components are stationary and there are r_1 common trends or non-stationary factors, there must be $N - r_1$ cointegration relations among the observed series. So, the number of common trends, r_1 , can be determined by estimating the number of cointegration relations $N - r_1$. Note that Bai and Wang (2016) show that the link between cointegration and common trends is broken in the setup of large N, T . Cointegration among the factors themselves is considered in Barigozzi et al. (2021) and Barigozzi and Luciani (2019). For simplicity, we assume that the non-stationary common factors are not cointegrated.

can be used to determine r_1 :

$$A_y(k) = \frac{1}{T^\alpha} \sum_{t=k+1}^T Y_{t-k} Y_t' \quad (56)$$

This matrix is also used by Barigozzi and Trapani (2018), who propose a procedure based on randomization tests that allows for non-stationary idiosyncratic components and that can be implemented in settings with large cross-sectional dimension.

If the idiosyncratic component is further assumed to be not only stationary but also white noise, Peña and Poncela (2006) propose testing for the number of common factors, stationary or not, using the following test based on canonical correlations:

$$s = -(T - k) \sum_{j=0}^{N-r-1} \log(1 - \lambda_{N-j}) \quad (57)$$

where $\lambda_1 > \dots > \lambda_N$ are the ordered eigenvalues of the matrix $M(k) = C_Y'(k) C_Y^{-1}(0) C_Y(k) C_Y^{-1}(0)$. The statistic s is asymptotically a $\chi_{(N-r)^2}^2$ when T goes to ∞ . The advantage of canonical correlation analysis is that you may detect the total number of common factors independently of their order of integration. The disadvantage of the procedure is that any remaining serial correlation present in the idiosyncratic noise is detected as additional common factors. This problem has been very recently solved in Bolívar (2020), who modified the test by taking into account the presence of correlated idiosyncratic components; see also Pan and Yao (2008) for methods of detecting the number of common factors based on the same principle.

Apart from the previous methods that rely on analyzing the eigenstructure of certain matrices, Bai and Ng (2004) propose a procedure, PANIC, for the number of common non-stationary factors based on PC while allowing for non-stationary idiosyncratic components as well. The procedure is as follows: First, apply PC to the data in first differences and extract the common factors. The number of common factors for stationary data can be determined applying the information criteria of Bai and Ng (2002). Denote them by f_t . Then recumulate to obtain the common factors in levels $\hat{f}_t = \sum_{s=2}^t f_s$, $t = 2, \dots, T$. In this way, we have consistent estimates of the common factors even though there might be unit roots in the idiosyncratic components. If there is only one common factor, PANIC performs a simple unit root test. If there are multiple factors, Bai and Ng (2004) suggest the tests for common trends proposed by Stock and Watson (1988) to determine the number of stochastic trends in the estimated r common factors; see Harvey (1989) who also suggests using Stock and Watson (1988) to determine the number of common trends. As regards the non-stationarity of the idiosyncratic noise, univariate unit root tests may have low power, so they proposed using panel unit root tests. In particular, they propose the following test:

$$S = \frac{-2 \sum_{i=1}^N \log s_i - 2N}{\sqrt{4N}} \quad (58)$$

where s_i is the p -value corresponding to the Dickey-Fuller test of the i th idiosyncratic residual. Pooled tests cannot be used in the original data because of strong cross-correlation due to the common factors, but they can be used in the specific components since this strong cross-correlation has been removed after extracting the common factors.

Finally, Nyblom and Harvey (2000) propose a test for the number of common trends based on the Lagrange multiplier principle. The asymptotic distribution of their test statistics depends only on the rank r_1 , the number of common non-stationary factors, of the covariance matrix of the disturbances driving the N random walks.

7.4. Empirical applications of non-stationary DFM

The number of empirical applications of non-stationary DFM using KFS for factor extraction is much more scarce than in the stationary case.

Many important applications of DFM deal with real economic activity. Although the vast majority of papers using DFM to track and forecast the real economy use stationary DFM, there are a few papers that fit non-stationary DFM using KFS procedures to extract the factors. In general, analysts are interested in: (i) detecting the turning points of the business cycle through indexes of economic activity, and (ii) forecasting real GDP or other variables related to real economic activity. One of the pioneering papers is Quah and Sargent (1993) who use information from a large set of cross-sectional data to track the temporal evolution of aggregate fluctuations. They use a non-stationary DFM with annual data ($T = 48$) over a cross-section of $N = 60$ sectoral US unemployment data. This seminal paper has spurred a vast body of literature regarding DFM, stationary or not, for the analysis of business cycles. García-Ferrer and Poncela (2002) use a small-scale factor model with stationary and non-stationary common factors to forecast real GNP of a set of euro area countries that represent about 80% of the euro area GDP. They find the DFM outperform other forecasting alternatives. The same data set is used in Peña and Poncela (2004) to illustrate the smaller MSEs obtained with DFM over the pooled term in García-Ferrer, Highfield, Palm, and Zellner (1987) and univariate alternatives. They also show that the common factor estimated as the pooling term exhibits minimum MSE if the common factor follows a random walk. Barigozzi and Luciani (2017) use non-stationary DFM to separate long-run from short-run comovements in real economic activity, using a large data set of US macroeconomic indicators. Frale et al. (2011) are interested in an index of economic activity and use mixed frequency non-stationary DFM to build EURO-MIND, an index to track business cycles in the euro area. Seong et al. (2013) use the data of the four big US monthly indicators (industrial production, employment, income, and sales) and quarterly GDP to compute in-sample monthly smoothed estimates and out-of-sample monthly forecasts of GDP. Bujosa, García-Ferrer, and de Juan (2013), Bujosa, García-Ferrer, de Juan,

and Martín-Arroyo (2020) determine non-stationary common trends based on the identification procedure in Peña and Poncela (2006) to build monthly coincident and leading indicators for the Spanish economy. They keep only the lower frequency component in order to obtain smooth indexes, with the aim of reducing the uncertainty of the state of the economy. When building indexes of economic activity, one-factor models are usually estimated. However, Martínez, Nieto, and Poncela (2016) ask which common factor should be used as the index of interest in multifactor models. They complement the analysis comparing the peaks of the estimated common factors with those of a reference series using Fisher's randomization test for matching pairs. They apply it to real economic activity data in Colombia and to financial series from Colombian stock markets. Combes and Doz (2018) extract the non-stationary common factors using the two-step procedure of Doz et al. (2011) to forecast French GDP, and run a pseudo-real time forecasting exercise to compare these predictions with those obtained using stationary DFM. They conclude that the differences are not relevant. A different application related to GDP can be found in Corona et al. (2020) where non-stationary DFM are used to detect risk sharing or the amount of GDP shocks that are smoothed, and therefore not passed into consumption, through cross-border international flows.

In the non-stationary case there are also DFM to monitor inflation; see, for instance, Delle Monache et al. (2016). Barnichon and Mesters (2018) also fit a non-stationary DFM with the factors extracted using KFS to separate aggregate labor market forces and demographic-specific trends to construct a business-cycle indicator based on unemployment.

Another set of applications is more related to finance. Harvey, Ruiz, and Shephard (1994) find that two common trends explain the evolution of volatilities of four exchange rates in the context of multivariate stochastic volatility models. Peña and Poncela (2006) use stationary and non-stationary common factors to model interest rates at different maturities. Broto and Perez-Quiros (2015) work with the sovereign credit default swap spreads of ten OECD economies. These are decomposed into a global common random walk, a specific random walk linked to European peripheral countries, and an idiosyncratic component, in order to study contagion.

Other applications include hourly electricity prices, as in Alonso et al. (2011) and Carpio and Juan (2014). In demography, Ortega and Poncela (2005) use non-stationary DFM for $I(2)$ time series in order to forecast fertility rates of southern European countries. In environmetrics, Zuur et al. (2003) apply non-stationary DFM to model the biomass of marine species, and Nieto et al. (2016) extend the non-stationary DFM to seasonal data and apply it to monthly measures of rainfall measured at six airports in Colombia.

Finally, comparing the predictive performance of small-scale models based on levels or on differenced data, Fuleky and Bonham (2015) conclude that the former have better properties in the context of some macroeconomic variables.

8. Conclusions

Using KFS procedures for factor extraction is a powerful instrument that generates efficient estimates of the underlying latent factors, to deal in a direct way with different types of data anomalies and non-stationarities. In recent years, some issues dealing with the computational complexity of KFS factor extraction have been solved and the EM algorithm is now predominant in the estimation of the parameters of DFM even when the cross-sectional dimension is very large. When fitting DFM written as an SSM to large systems of time series, it is crucial to take special care of the identification of the parameters and the specification of the model (mainly in cases of dynamic and non-stationary factors). Due to the importance of a correct specification of the factors, there is a need for more evaluation and testing of the model specification.

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