Dynamic Dependence Networks: Financial Time Series Forecasting & Portfolio Decisions

Zoey Yi Zhao∗, Meng Xie† and Mike West‡

Original technical report, December 2015
The final revised paper appears (with invited discussion) in the journal:

Abstract

We discuss Bayesian forecasting of increasingly high-dimensional time series, a key area of application of stochastic dynamic models in the financial industry and allied areas of business. Novel state-space models characterizing sparse patterns of dependence among multiple time series extend existing multivariate volatility models to enable scaling to higher numbers of individual time series. The theory of these dynamic dependence network models shows how the individual series can be decoupled for sequential analysis, and then recoupled for applied forecasting and decision analysis. Decoupling allows fast, efficient analysis of each of the series in individual univariate models that are linked--for later recoupling--through a theoretical multivariate volatility structure defined by a sparse underlying graphical model. Computational advances are especially significant in connection with model uncertainty about the sparsity patterns among series that define this graphical model; Bayesian model averaging using discounting of historical information builds substantially on this computational advance. An extensive, detailed case study showcases the use of these models, and the improvements in forecasting and financial portfolio investment decisions that are achievable. Using a long series of daily international currency, stock indices and commodity prices, the case study includes evaluations of multi-day forecasts and Bayesian portfolio analysis with a variety of practical utility functions, as well as comparisons against commodity trading advisor benchmarks.

Keywords: Bayesian forecasting; discount model averaging; dynamic graphical model; graphical model uncertainty; multiregression dynamic model; portfolio optimization; sparse dynamics

∗Statistical Arbitrage Researcher, Citadel LLC, Chicago, IL 60603, USA. zoeyzhao1010@gmail.com
†PhD student, Department of Statistical Science, Duke University, Durham, NC 27708, USA. mengaxie@gmail.com
‡The Arts & Sciences Professor of Statistics & Decision Sciences, Department of Statistical Science, Duke University, Durham, NC 27708, USA. mw@stat.duke.edu
1 Introduction

Applied time series analysis, forecasting and accompanying methods of decision analysis using increasingly sophisticated stochastic models of time series is nowadays central to many companies, non-profit organizations, research groups and individuals in the business of investment management, as well as in the broader financial services industries. Among frontier applied research questions is a central challenge of scaling statistical analysis addressing dynamics in cross-series relationships of multiple time varying indices—i.e., of usefully characterizing complex patterns of multivariate volatility to apply to forecasting and decisions with higher-dimensional time series. This is the focus of this paper, addressed in terms of modelling and methodological advances coupled with a detailed case study in finance.

Dynamic dependence network models are extensions of multiregression dynamic models (MDMs—[2, 17–19]). MDMs incorporate directed graphical model structure into a multivariate time series, allowing contemporaneous values of some univariate series to appear as predictors of other series. Originally introduced to preserve certain conditional independence structures related to causality over time [18], MDMs have been developed and applied to multivariate time series in areas such as forecasting of brand sales and traffic flows [e.g. 2, 17], and as empirical models of dynamic network structures generating inter-related time series in areas such as neuroscience, engineering signal processing and financial econometrics [e.g. 5, 10, 13]. While these previous works have illustrated the effectiveness of some particular MDMs in inference and forecasting of multivariate time series, our interests here are defined by needs for several extensions of the modelling ideas and methodology. Beginning with the basic MDM framework, we are motivated to extend and explore more general methodology to capture and quantify time-variations in patterns of conditional independence structures. We do this via the concept of sparsity in conditional dependence networks and develop analysis to enable dynamic modelling of these sparse networks over time. This is overlaid with innovations in Bayesian model uncertainty analysis relative to conditional independence structure, in a dynamic/adaptive strategy that also deals with model parameter uncertainty. Further, we link this extended MDM framework to the increasingly adopted Cholesky-style approach to modelling multivariate stochastic volatility [e.g. 10–14, 16, 23]. Then, we are interested in extensions to include time-varying autoregressions in predictive model components. Such natural extensions of MDMs have not, to date, been exploited, in part due to the lack of extension of existing theoretical results for forecast distributions more than one-step ahead; we address this in the context of an overall simulation-based analysis that immediately allows forecasting multi-steps ahead as required in many applications including our portfolio studies. Finally, DDNMs inherit the MDM feature that sequential time series analysis and forecasting can be decoupled into that of a set of univariate dynamic linear models (DLMs)—so enabling fast, parallel processing—and then recoupled for forecasting and decisions.

Section 2 discusses MDMs and links to Cholesky-style multivariate volatility models, then develops the decouple/recouple feature of sequential analysis that enables parallel processing of multivariate time series. This section notes some new and practically relevant technical developments that are detailed in the Appendix. Section 3 defines a class of DDNMs that extend MDMs to include predictive dynamic model components with time-varying autoregressive (TVAR) structure. The forward-filtering and forecasting analysis is discussed, with required extensions to the existing MDM theory. In this context, we develop model structure uncertainty analysis via sequential Bayesian mixture modelling with implied model averaging for inference and forecasting. Developed in detail in Section 4, this addresses uncertainty about, and learning on, structural model components including the predictor variable uncertainty that defines the “network” structure; that is, the patterns of sparsity in contemporaneous relationships among series as well...
as potential links to lagged values of the series. Embedding these in an overall framework of Bayesian model uncertainty and model averaging leads to inference on the time-varying structure of the implicit network of interconnections. The analysis also includes uncertainty about key model hyper-parameters, including TVAR lags and discount factors defining rates of change of state vectors and volatility processes. A practically important element of the work is the use of annealed structure learning via power discounting that acts to limit the degeneracy of posterior model probabilities over time, and so enhance model adaptability to new data and changing circumstances. Discussed in Section 4, this is shown in the case study of Section 6 to be both statistically supported—in terms of enhancing model fit and forecast performance—and to underlie improved decisions in resulting financial portfolio evaluations.

With brief background on Bayesian decision analysis in dynamic portfolio allocation in Section 5, the case study in Section 6 concerns a 13−dimensional time series of daily prices (in $US) of several international currencies, commodities and stock indices over a time-span of 11 years. The section summarizes key aspects of DDNM specification, assessment and use in both 1− and 5−day ahead forecasting, and explores the outcomes of a range of portfolio studies. Key technical details are in the series of sections of the Appendix. Section 7 concludes the paper with some summary comments.

2 MDM Framework

The framework is that of structured state-space modelling and general notation follows that of standard Bayesian dynamic linear models [e.g. 15, 26]. The $m \times 1$-vector time series $y_t = (y_{1t}, \ldots, y_{mt})'$ is observed over time $t = 1, 2, \ldots$. Denote by $D_0$ information available at $t = 0$, and by $D_t = \{D_{t-1}, y_t\}$ the time $t$ information set; the latter are sequentially updated as observations are made over time.

2.1 Cholesky-Style Multiregression Dynamic Models

Consider each univariate series $y_{jt}$; for $j = 1 : m − 1$, let $pa(j) \subseteq \{j + 1 : m\}$ be a subset of indices of those series higher than $j$ in the selected order, and set $pa(m) = \emptyset$, the empty set. Then $y_{pa(j),t}$ is the $|pa(j)| \times 1$−vector of time $t$ values on the series in the parental set $pa(j)$. The $m$ independent, univariate DLMs of eqn. (1) define a triangular system that, by composition, yields a full multivariate model for $y_t$.

Assuming sparsity of parental sets—i.e., that some or all of the $pa(j)$ contain fewer than the full number of potential parental indicators—this is a dynamic graphical model [4, 8, 17, 18, 24]; the graphical modelling terminology reflects the construction of the model from a set of conditional distributions in a directed, acyclic graph format resulting from the triangular/Cholesky-style specification. This is a general example of a multiregression dynamic model.

The coupled set of univariate DLMs is, over times $t = 1, 2, \cdots$,

$$y_{jt} = x_{jt}' \phi_{jt} + y_{pa(j),t}' \gamma_{jt} + \nu_{jt} = F_{jt}' \theta_{jt} + \nu_{jt}, \quad j = 1 : m, \quad (1)$$

with components as follows:

- $x_{jt}$, a known column vector of predictors or constants, with corresponding dynamic regression coefficients in the column state vector $\phi_{jt}$, each of dimension $p_{j\phi}$.

- $\gamma_{jt}$, a vector of dynamic regression coefficients $\gamma_{jht}, h \in pa(j)$, linking contemporaneous values of some of the other series to series $j$; the number of parents and dimension of $\gamma_{jt}$ is $p_{j\gamma} = |pa(j)|$.
• The observation errors are conditionally independent over $j$ with $\nu_{jt} \sim N(0, 1/\lambda_{jt})$ independently of $F_{jt}, \theta_{jt}$, with possibly time-varying precisions $\lambda_{jt}$. Define $\Lambda_t = \text{diag}(\lambda_{1t}, \ldots, \lambda_{mt})$.

• The full dynamic state and regression vectors, each of dimension $p_j = p_{j\phi} + p_{j\gamma}$, are

$$
\theta_{jt} = \begin{pmatrix} \phi_{jt} \\ \gamma_{jt} \end{pmatrix} \quad \text{and} \quad F_{jt} = \begin{pmatrix} x_{jt} \\ y_{pa(j),t} \end{pmatrix}.
$$

Complete model specification involves time evolution models for the $\theta_{jt}$ and $\lambda_{jt}$. We develop these below, building on traditional Bayesian dynamic linear model specifications [e.g. 26].

With notation $\tilde{\gamma}_{j,pa(j),t} \equiv \gamma_{jt}$ and $\tilde{\gamma}_{jht} = 0$ for $h \notin pa(j)$, collect the effective coefficients $\gamma_{jt}$ and implicit zero values in the matrix

$$
\Gamma_t = \begin{pmatrix}
0 & \tilde{\gamma}_{12t} & \cdots & \tilde{\gamma}_{1mt} \\
0 & 0 & \tilde{\gamma}_{23t} & \cdots & \tilde{\gamma}_{2mt} \\
\vdots & \vdots & \ddots & \vdots & \vdots \\
0 & 0 & \cdots & 0 & \tilde{\gamma}_{m-1,mt} \\
0 & 0 & \cdots & 0 & 0
\end{pmatrix}.
$$

The set of $m$ coupled models of eqn. (1) can then be written as

$$(I - \Gamma_t)y_t = \mu_t + \nu_t$$

where $\mu_t = (\mu_{1t}, \ldots, \mu_{mt})'$ with $\mu_{jt} = x_{jt}'\phi_{jt}$, and $\nu_t \sim N(0, \Lambda_t^{-1})$ with precision matrix $\Lambda_t$. Then

$$y_t \sim N((I - \Gamma_t)^{-1}\mu_t, \Omega_t^{-1}) \quad \text{with} \quad \Omega_t = (I - \Gamma_t)'\Lambda_t(I - \Gamma_t).$$

Hence $I - \Gamma_t$ is the Cholesky of $\Lambda_t$ subject to row-scaling by the square roots of the diagonal entries $\lambda_{jt}$.

The parental sets reflect the contemporaneous conditional dependence structure across the series; conditional on state parameters and predictors, for any $i > j$ we see that $y_{it} \perp \perp y_{jt}|y_{pa(j),t}$ if $i \notin pa(j)$. This class of MDMs naturally yields a path to flexible modelling of multivariate volatility, as we can use any state-space evolution for the $\gamma_{jt}$ and univariate volatility models for the $\lambda_{jt}$, and they together induce the stochastic dynamics of the implied $\Omega_t$. Also, sparse structuring will be based on small, parsimonious choices of parental sets, so yielding structure and, typically, sparsity in the resulting dynamic graphical model. The desirability of this– and benefits in terms of potential to improve forecasts and resulting decisions in areas such as financial portfolio analysis– has been highlighted in earlier uses of MDMs and other approaches to dynamic graphical models [e.g. 4, 8, 17, 18, 24].

### 2.2 Tractable MDMs: Forward-Filtering and Forecasting

The models of eqn. (1) are completed by specifying dynamic model forms for the state vectors $\theta_{jt}$ and precisions $\lambda_{jt}$ over time. Standard DLM classes provide ranges of models for structured, linear and conditionally Gaussian evolutions of the $\theta_{jt}$ coupled with tractable discount specifications for evolution noise levels as well as for the residual volatilities $\lambda_{jt}$ [15, 26]. We use one of the simplest such specifications in our case study below– multivariate random walk evolutions for each of the $\theta_{jt}$ over time coupled with discount specifications– and so restrict discussion here to that specific model form.
Specifically, we adopt random walk state evolution models \( \theta_{jt} = \theta_{j,t-1} + \omega_{jt} \) where the evolution error \( \omega_{jt} \) is zero-mean normal, independent over time and across series, and has a time-dependent evolution error variance matrix defined via a single discount factor \( \delta_j \in (0, 1] \). Coupled with this is a standard random walk volatility model \( \lambda_{jt} = \lambda_{j,t-1}\eta_jt/\beta_j \) where the \( \eta_jt \) are independent beta random variates with time-dependent beta parameters defined via the single discount factor \( \beta_j \in (0, 1] \) for series \( j \). Again, these are standard models and full details appear in the above references. We include summary details in Appendix A of this paper, together with summaries of the resulting prior, posterior and forecast distributions as they are updated over time. Critically, these analyses apply in parallel, the series being decoupled for forward-filtering and forecasting within-series; forecast distributions are then coupled together for multivariate forecasting, as summarized below.

Key elements of the forward-filtering analyses are sequentially updated versions of the following distributions. These are specific to each series \( j \) and conditionally independent across \( j \). See Appendix A for the full technical and notational details.

**Posterior and priors at time \( t-1 \):** At each time \( t-1 \) information \( D_{t-1} \) is sufficiently summarized in terms current normal/gamma posteriors

\[
(\theta_{j,t-1}, \lambda_{j,t-1}|D_{t-1}) \sim NG(m_{j,t-1}, C_{j,t-1}, n_{j,t-1}, n_{j,t-1}s_{j,t-1}),
\]

where the notation represents the conditional normal and marginal gamma

\[
\theta_{j,t-1}|\lambda_{j,t-1}, D_{t-1} \sim N(m_{j,t-1}, C_{j,t-1}/(s_{j,t-1}\lambda_{j,t-1}))
\]
\[
\lambda_{j,t-1}|D_{t-1} \sim G(n_{j,t-1}/2, n_{j,t-1}s_{j,t-1}/2)
\]

and where \( s_{j,t-1} \) is a current point estimate of the residual variance \( 1/\lambda_{j,t-1} \). These imply 1-step ahead prior distributions for states of the same normal/gamma form

\[
(\theta_{jt}, \lambda_{jt}|D_{t-1}) \sim NG(a_{jt}, R_{jt}, r_{jt}s_{j,t-1})
\]

where \( a_{jt} \equiv m_{j,t-1}, R_{jt} = C_{j,t-1}/\delta_j \) and \( r_{jt} = \beta_jn_{j,t-1} \) based on the specified discount factors \( \delta_j, \beta_j \).

**1-step forecasts at time \( t-1 \):** The implied predictive distribution is \( T \) with \( r_{jt} \) degrees of freedom,

\[
(y_{jt}|y_{pa(j),t}, D_{t-1}) \sim T_{r_{jt}}(f_{jt}(y_{pa(j),t}), q_{jt}(y_{pa(j),t}))
\]

where \( y_{pa(j),t} \) appears linearly in \( f_{jt}(y_{pa(j),t}) \) and quadratically \( q_{jt}(y_{pa(j),t}) \).

**\( k \)-step forecasts:** More than 1-step ahead forecast distributions are similarly given by conditional \( T \) distributions; the conditioning requires known values of future independent predictor variables and explicitly involves future values of parental series for each \( j \). The practical approach to utilizing this theory for \( k \)-step forecasting is detailed below.
2.3 Multivariate Predictive Distributions 1–Step Ahead

The compositional nature of the triangular/Cholesky-style MDM yields access to nice analytics in evaluating 1–step ahead forecasts and various relevant aspects of the full multivariate predictive distribution. Applied work will require computation of predictive means and variance matrices, and other summaries, as well as evaluations of the joint density function. Some specific comments are given here, with full theoretical details in Appendix B of the paper.

First, note that the full 1–step ahead predictive density for \( y_t \) is, via composition, simply

\[
p(y_t|D_{t-1}) = \prod_{j=1}^{m} p(y_{jt}|y_{pa(j),t}, D_{t-1}). \tag{3}
\]

As the univariate conditionals \( p(y_{jt}|y_{pa(j),t}, D_{t-1}) \) are T densities noted above, the multivariate distribution is a product of Ts. The p.d.f. is easily evaluated based on observed data. This point is important in model assessment, comparison and combination, as the product over time of the joint predictive densities under any chosen model defines the model marginal likelihood. In extended models below, we use this in comparing model structures— in terms of ranges of parental sets— as well as model hyper-parameters, including discount factors and other aspects of model specification.

Second, we typically require 1–step ahead predictive moments as well as, in some cases, other summaries. For portfolio applications based on mean-variance optimisations and trade-offs— such as in our case study of this paper— we are interested in forecast mean vectors, variance matrices and precision matrices

\[
\begin{align*}
f_t &= E(y_t|D_{t-1}), & Q_t &= V(y_t|D_{t-1}), & K_t &= V(y_t|D_{t-1})^{-1},
\end{align*}
\]

under (3). Assume that, for all \( j, t \), the degrees-of-freedom parameters \( r_{jt} \) exceed 2, so that the variances exist. The recursive form of the compositional model means that we have access to analytically tractable recursions to enable the calculations of \( f_t \) and \( Q_t \); full details appear in Appendix B. This explicitly recognizes the appearance of contemporaneous values of the \( y_{pa(j),t} \) in the conditioning of forecasts for \( y_{jt} \).

Further, in some applied work we are also interested in the predictive precision matrix \( K_t \). This directly reflects the conditional dependence structure between variables and thus plays a crucial component in various types of analysis. In particular, the precision matrix has a determining effect on the allocation portfolio weights in financial portfolio studies. It turns out that the recursive evaluation of joint predictive means and variance matrices has an analytically nice parallel for recursive computation of the precision matrix. As well as a new theoretical result for MDMs, this is a key practical note since it allows us to avoid direct matrix inversion. Again, details are given below in Appendix B.

Finally, we note that the computation of moments and precision matrices of \( k \)—step forecast distributions for \( k > 1 \) follow very similar lines, so details are omitted here.

3 Dynamic Dependence Network Models

We use the term dynamic dependence network model (DDNM) for an MDM that has been extended to allow for time-varying autoregressive (TVAR) components in each univariate series. This broader model class extends the practical utility of MDMs, while requiring extensions of the methodology to enable Bayesian forecasting in the resulting time-varying, vector autoregressions (TV-VARs) coupled with
Cholesky-style multivariate volatility. The “network” terminology is relevant in that DDNMs have dynamic linkages both across series and at lagged values that can represent—and be interpreted as—both contemporaneous and lagged network interconnections. Indeed, variants of these models that utilize latent thresholding concepts for the dynamics of state vectors have recently been explored in contexts where network structure is a key interest [13]. Here that is not a key focus, but the inclusion of TVAR model components is of central interest in terms of improving forecast accuracy, and resulting characterizations of cross-series patterns in multivariate volatility.

The DDNM class modifies the basic MDM of eqn. (1) as follows. For each \( j = 1 : m \),

\[
y_{jt} = c_{jt} + \sum_{i=1:p_{j\lambda}} y_{jt-i} \phi_{j\lambda i} + y_{pa(j),t} \gamma_{jt} + \nu_{jt},
\]

where \( c_{jt} \) is a time-varying intercept, each \( \phi_{j\lambda i} \) is a \( m \)-vector of TV-VAR coefficients for lag \( i \) = 1, \ldots, \( p_{j\lambda} \) for some maximum lag \( p_{j\lambda} \), and \( \nu_{jt} \) the observation noise. It is clear that we could add dynamic effects of additional independent variables to enrich the class of models; our case study does not do that, but the methodological details are directly extensible.

Let \( p_{\lambda} = \max(p_{1\lambda}, \ldots, p_{m\lambda}) \) and write \( \tilde{\phi}_{j\lambda} \) for the \( p_{\lambda} \times 1 \)-vector that extends \( \phi_{j\lambda} \) with zeros for the elements of subscript larger than \( p_{j\lambda} \). Then (4) can be written in the vector form

\[
(I - \Gamma_t) y_t = c_t + \phi_{1\lambda} y_{t-1} + \cdots + \phi_{p_{\lambda} \lambda} y_{t-p_{\lambda}} + \nu_t,
\]

where \( \phi_{j\lambda} = (\tilde{\phi}_{1\lambda\lambda}, \ldots, \tilde{\phi}_{p_{\lambda} \lambda\lambda})' \) and \( \Gamma_t \) is as in eqn. (2). In our portfolio application, we consider the simple but practically central TV-VAR model where the autoregressive predictor variables of series \( j \) only contains its own lags and the intercept; that is,

\[
y_{jt} = c_{jt} + \sum_{i=1:p_{j\lambda}} \phi_{j\lambda i} y_{jt-i} + y_{pa(j),t} \gamma_{jt} + \nu_{jt},
\]

where \( \phi_{j\lambda i} \) is the time-varying autoregressive coefficient of series \( j \) at lag \( i \), \( (i = 1, \ldots, p_{j\lambda}) \). This can be written as an MDM in which \( x_{jt} = (1, y_{jt-1}, \cdots, y_{jt-p_{j\lambda}})' \) and \( \phi_{j\lambda} = (c_{jt}, \phi_{j\lambda 1}, \ldots, \phi_{j\lambda p_{j\lambda}})' \). As a result, much of the theory and methodology of MDMs applies. In particular, the general results on forward-filtering and 1-step ahead forecasting of Section 2.2 hold for these DDNMs. However, for forecasting more than one-step ahead, the MDM theory is inapplicable. This arises as the existing theoretical results for forecasting in the MDM framework require knowledge of the future predictor variables; in autoregressive contexts, the future predictors include lagged values of the \( y_{jt} \) which are unknown at the time of forecasting. For any \( k > 1 \), forecasting \( k \)-steps ahead requires an ability to deal with uncertainty about the then-required predictors that are values of \( y_{t-k+1} \) now in the dynamic linear regressions representing both lagged values of the current series \( j \) as well as the parental predictors.

The solution to this is simulation: Given a current, time \( t \) set of posterior for state vectors and volatilities across the \( m \) series, we can trivially simulate each model to time \( t + 1 \), and conditional on the value of the sampled \( y_{t+1} \) vector, continue to sample \( y_{t+2}, y_{t+3}, \) and so forth up to whatever lead time required. Repeating this independently will define a Monte Carlo sample from the full set of posterior predictive distributions over each \( j \), and hence from the full predictive distribution \( p(y_{t+1}, \ldots, y_{t+k} | D_t) \). Direct summarization then leads to Monte Carlo approximations to predictive mean vectors, variance matrices and other quantities of interest. Critically, we can simulate samples as large as desired very efficiently, since this uses the analytically tractable set of \( m \) DLMs analyzed and simulated in parallel. The forward-filtering updates and simulation computations are standard and technically/computationally trivial.
4 Model Structure and Hyperparameter Uncertainty

Application of DDNMs requires addressing questions of model uncertainty about key defining parameters: the structural parental sets $pa(j)$, and the hyperparameters comprising TVAR model orders $p_{j\lambda}$ and discount factors $(\delta_j, \beta_j)$ for each $j = 1 : m$. We address this using multiple DDNMs, each defined by selected parameters, evaluating and sequentially revising posterior model probabilities across this discrete set of models, and then averaging over models for inferences and predictions. This basic mixture modelling approach has been central to Bayesian forecasting and dynamic models for decades, predating its more recent popularization in static models under the name Bayesian model averaging (BMA) [e.g. 7, 26, chapt 12, and references therein].

4.1 Discrete Sets of Models and Model Probabilities

For each $j = 1 : m$, define $\mathcal{M}_j = \{pa(j), p_{j\lambda}, \delta_j, \beta_j\}$ for any specific choice of these parameters. Here: (i) $pa(j) \subseteq \{j + 1 : m\}$ can take any of the $2^{m-j}$ possible values (though we may decide to restrict the possibilities based on exploratory analysis of initial training data or on substantive grounds); (ii) $p_{j\lambda} \in 1 : d$ for some specified maximum lag $d$; (iii) the discount factor pair $(\delta_j, \beta_j)$ takes a value from a discrete set of $k$ points on a grid in $(0, 1]^2$. Allowing the maximum set of possible parents, this defines a class of $n_j = 2^{m-j}(d+1)k$ possible DDNMs for series $j$.

Given specific values for each $\mathcal{M}_j$, we have one DDNM for $y_t$ whose parameters are denoted by $\mathcal{M}_{1:m} = \{\mathcal{M}_1, \ldots, \mathcal{M}_m\}$. The number of such models is $n = \prod_{j=1:m} n_j = 2^{m(m-1)/2}(d+1)^mk^m$; in any realistic application, this will be too large a class of models to evaluate. For example, our case study has $m = 13$, $d = 2$ and $k = 25$, yielding more than $7 \times 10^{47}$ possibilities. Fortunately, the compositional structure of DDNMs leads to a massive reduction based on assumption of independent priors over model structures across the set of $m$ univariate models. See this as follows.

At any given model $\mathcal{M}_{1:m}$, extend our earlier notation for $1$–step forecast densities of eqn. (3) to explicitly note the dependence on the values in $\mathcal{M}_{1:m}$; at time $t$, the p.d.f. is $p(y_t | D_{t-1}, \mathcal{M}_{1:m})$. Then, under any prior distribution giving initial probabilities $p(\mathcal{M}_{1:m} | D_0)$ to each of the large set of possible models, the posterior model probability based on observed data $D_t = \{D_0, y_{1:t}\}$ is

$$
p(\mathcal{M}_{1:m} | D_t) \propto p(\mathcal{M}_{1:m} | D_0)p(y_{1:t} | D_{t-1}, \mathcal{M}_{1:m}) = p(\mathcal{M}_{1:m} | D_0) \prod_{r=1:t} p(y_r | D_{r-1}, \mathcal{M}_{1:m})$$

$$
= p(\mathcal{M}_{1:m} | D_0) \prod_{j=1:m} \prod_{r=1:t} p(y_r | D_{r-1}, \mathcal{M}_j) 
$$

(6)

where the last step inserts the product of univariate T densities of eqn. (3) at each time $r = 1 : t$. Now suppose that prior at $t = 0$ has model parameters independent across series $j$, with $p(\mathcal{M}_{1:m} | D_0) = \prod_{j=1:m} p(\mathcal{M}_j | D_0)$. This implies that the full model posterior in eqn. (6) is the product of $m$ independent model posteriors

$$
p(\mathcal{M}_j | D_t) \propto p(\mathcal{M}_j | D_0) \prod_{r=1:t} p(y_r | D_{r-1}, \mathcal{M}_j) = p(\mathcal{M}_j | D_{t-1})p(y_t | y_{pa(j),1:t-1}, D_{t-1}, \mathcal{M}_j)
$$

(7)

This last equation also shows how these model probabilities are sequentially updated over time as successive observations are made. Each of the $1$–step conditional densities is a univariate T, so the computations
are trivial for each \( j, t \). The factorization over series \( j \) implies that the analyses can be run in parallel. This exploitation of the decoupling inherent in DDNMs thus reduces the computation to that of evaluating \( \sum_{j=1}^{m} 2^{m-j}(d+1)k = (2^{m-1})(d+1)k \) univariate DLMs in parallel and then combining the results across series to deliver model probabilities over all \( \mathcal{M}_{1:m} \). In our case study with \( m = 13, d = 2 \) and \( k = 25 \), this yields a very manageable number of just over 300,000 models. Then, as a result, we have overall model probabilities updated sequentially via the resulting dynamic/time \( t \) version of eqn. (6), namely

\[
p(\mathcal{M}_{1:m}|D_t) \propto p(\mathcal{M}_{1:m}|D_{t-1})p(y_t|D_{t-1}, \mathcal{M}_{1:m}).
\]  

The analysis requires specification of model uncertainty priors. We have prior independence over \( j = 1 : m \) as noted above, and take each series \( j \) – specific model prior as \( p(\mathcal{M}_j) = p(pa(j))p(p_{j\lambda})p(\delta_j, \beta_j) \) with independent components as follows:

(i) Our interest in sparse models favors priors on the parental sets \( pa(j) \) that penalize large values of \( p_{j\gamma} = |pa(j)| \). We use a traditional Bayesian variable inclusion prior in which parents are included independently with probability \( \rho \). Thus, a parental set \( pa(j) \) with \( p_{j\gamma} = c \) elements has prior probability \( p(pa(j)) = \rho^c(1-\rho)^{m-j-c} \), with an expected parental set size of \( (m-j)\rho \).

The latter provides insight into prior specification of \( \rho \). (ii) Discount factors \( (\delta_j, \beta_j) \) are selected from a rectangular grid of \( k \) specified values with a uniform discrete prior on the grid. (iii) The TVAR model order \( p_{j\lambda} \) is assigned a uniform prior on the range \( 0 : d \), given the chosen maximum possible lag \( d \).

### 4.2 Extended Model Uncertainty Analysis using Power Discounting

We make one additional extension of the model uncertainty framework. It is well known that, with sufficient data accrued, posterior model probabilities concentrate around a smaller set of models, eventually favouring a single model [e.g. 26, chapt 12]. This theoretically guaranteed behaviour can often lead to significantly down-weighting many models that may be of possible future interest, and degrade predictive performance as a result. This has led to interest in discounting past data to allow model probabilities to depend more on recent and current behaviour of the time series, and to adapt more adequately to incoming observations. A particular method of power discounting, used historically in Bayesian forecasting [25, p.445] has shown promise in portfolio studies [27] and has recently received attention in other applied areas [9, 22] (linking to a parallel historical literature where discount factors are called “forgetting” factors). The basic idea and resulting implementation is simple.

Extend the discrete set of models to \( \mathcal{M}_j = \{pa(j), p_{j\lambda}, \delta_j, \beta_j, \alpha\} \) where \( \alpha \in (0, 1] \) is a model probability power discount factor. Then the computation of posterior model probabilities is modified from the standard Bayesian update of eqn. (8) to the extended form

\[
p(\mathcal{M}_{1:m}|D_t) \propto p(\mathcal{M}_{1:m}|D_{t-1})^{\alpha} p(y_t|D_{t-1}, \mathcal{M}_{1:m}),
\]  

then being normalized to sum to 1 over all possibilities \( \mathcal{M}_{1:m} \). The \( \alpha \) – power applied to the time \( t - 1 \) model probabilities acts to increase the dispersion of this time \( t - 1 \) posterior, somewhat down-weighting the information content of past data. Smaller values of \( \alpha \) discount history to a greater extent, “flattening” the time \( t - 1 \) posterior relative to the standard update when \( \alpha = 1 \).

The prior specification now extends to add a prior \( p(\alpha) \). We take \( p(\alpha) \) to be discrete uniform on a chosen grid of points in \( (0, 1] \). This extends the analysis to include the power discount, so extends the model size in one additional dimension. Resulting conditional posterior model probabilities are still computed as above and then combined for evaluation of eqn. (9) at each time \( t \).
Finally note that, given the full set of model probabilities at each time $t$, we simply marginalize via summation to deduce implied marginal posteriors on any subset of elements of $M_{1:m}$. This is the route to evaluating over time the posterior support for different values of each discount factor, now including the power discount factor, as well as parental set membership and TVAR model orders. We use this extensively in the case study below.

### 4.3 Forecasting in DDNMs under Model Uncertainty

Under the general discrete model space, 1-step ahead forecast distributions are discrete mixtures over models $M_{1:m}$ of the product-form DDNM forecast distributions whose structure is discussed in Section 2.3. The 1-step ahead forecast mean vectors and variance matrices required for portfolio studies can then be evaluated by Bayesian model averaging using extensions of the nice, analytic recursions of that section and Appendix B. Several changes are needed to account for the model averaging, and a new theoretical element for computing covariance terms between pairs of series $y_{ht}$, $y_{jt}$—detailed in Appendix C of this paper—is key. Readers can refer to that appendix for additional details.

For forecasting more than 1-step ahead, the involvement of TVAR terms in DDNMs means that we do not have easily implemented analytic forms for forecast moments. Hence, for $k > 1$ we explore the $k$-step-ahead predictive distribution $p(y_{t+k-1}|D_{t-1})$ via direct and straightforward Monte Carlo simulation. Drawing a large Monte Carlo sample from this distribution—as detailed in Appendix C—yields relevant Monte Carlo estimates of the predictive mean and variance matrix. The simulations can be run in parallel and computationally cheap per sample.

In our case study, as in other financial applications, we adopt models in which the $y_{jt}$ are logged values of FX prices, commodity prices, or stock prices. Hence, even if we had access to analytic forms of predictive moments, they would be of limited interest as our portfolio decision analyses require—as inputs at each decision making stage—the predictive mean vectors and variance matrices of the implied returns, i.e., non-linear transformations of differences of log prices. Here the use of Monte Carlo simulations comes into play positively, as we can of course simply transform all simulated price series to returns, and hence directly compute sample estimates of the forecast means and variance matrices on the returns scale.

### 5 Dynamic Portfolio Allocation

Our case study concerns Bayesian forecasting and decision analysis for financial time series, and follows standard approaches in utilizing extensions of traditional Markowitz portfolio optimisation [1, 20, 21]. The basic methodology for our portfolio decision analysis is summarized here.

We consider $k$-step ahead portfolio optimisation with on DDNM-based forecasts; in the study below we evaluate cases with $k = 1$ and $k = 5$ on daily data. Focus here on the 1-step case; the development for $k > 1$ is the same but for the fact that it uses $k$-step ahead predictive distributions and assessments of portfolio characteristics with rolling 5-day horizons; readers can impute the omitted details. As we are interested in financial returns from investment decisions, any model of financial series that does not directly uses observed returns as the time series $y$ will have to enable computation of the implied mean vectors and variance matrices of future returns themselves. As noted earlier, our case study adopts our generally preferred approach of modelling log prices of FX series, commodities, and stock market indices. Given observed or simulated log prices $y_{jt}$ for one series $j$ over any period of time, the returns are simply
evaluated via exponentiating the differences of log prices. Hence, in our simulation-based analysis of DDNMs, it is trivial to map Monte Carlo samples of predictive distributions for future log prices to those of future returns, and then compute Monte Carlo approximations of the required moments and other aspects of the distributions.

For our development here, we drop the time and forecast horizon in the notation for clarity. Whatever the model form, step ahead desired, and nature of computation, we will use the running notation but now explicitly for returns: looking ahead our desired horizon, $y$ is the $m$–vector of future returns, and we suppose that at the current time point we have evaluated the forecast mean vector $f$ and variance matrix $Q$, denoted by $y \sim (f, Q)$. We now reallocate existing investments at the current time according to a portfolio weight vector $w$ that redistributes investments among the $m$ indices; the eventual return will then be $w'y$ when we move ahead in time and learn $y$. The weight vector is chosen via Bayesian decision analysis to optimise a specific portfolio loss function: generally, minimizing expected portfolio risk while aiming for good realized returns, subject to additional constraints. Under the forecast moments $y \sim (f, Q)$, the implied return on the portfolio for any given portfolio weight vector $w$ then has mean and variance $w'y \sim (w'f, w'Qw)$. Portfolio risk is taken as the standard deviation $\sqrt{w'Qw}$; in financial terminology this is referred to as the projected risk (PR) of the portfolio. We take the traditional closed portfolio approach in which the portfolio weights sum to 1, i.e., $w'1 = 1$, so that we are simply reinvesting existing resources (not increasing from external sources or reducing the overall investment level) in order to make fair comparisons across models and utility functions.

We examine variants of three commonly used portfolio allocation rules, as follows. These all depend critically on the forecast precision matrix, denoted by $K = Q^{-1}$.

1. **Target portfolio**: Given a specified return target $r$, optimise the portfolio weights by minimizing the ex ante portfolio variance among the restricted set portfolios with expected return $w'f = r$. The investor decision problem reduces to choosing the vector of portfolio weights $w$ to minimize $w'Qw$, subject to constraints $w'f = r$ and $w'1 = 1$. Direct analysis using Lagrange multipliers yields analytic forms of the optimising vector $w_1$ [e.g. 1, 15, chapt 10].

2. **Target constraint portfolio**: This modifies the first rule by adding the constraints that each element of $w$ must be nonnegative. The solution $w_2$ has no closed analytic form, but can easily computed using one of many standard non-linear optimisation algorithms; we use the quadratic programming tools in Matlab in our case study.

3. **Benchmark uncorrelated target portfolio**: This strategy involves an additional “benchmark” time series $z$. The forecasting model is then fitted to the extended $(m + 1)$–dimensional series, with the benchmark now included. This will lead to forecast moments for the returns on the original $m$ series together with the benchmark. At our current time point, suppose this leads to forecast benchmark mean and variance $z \sim (s, v)$ and with forecast covariance vector between our original series and the benchmark of $C(y, z) = q$ for some covariance vector $q$. This strategy modifies the target portfolio with the additional constraints $w'f = r + s$ and $w'q = 0$. That is, we aim at a return that exceeds that of the benchmark by the specified target $r > 0$, while being uncorrelated with the benchmark. Direct analysis using Lagrange multipliers yields an analytic form for the optimising vector $w_3$.

The traditional target portfolio defines a trade-off between risk and expected return. Adding non-negativity constraints will naturally increase the risk for a given target return, and is a rule that comes closer to
representing realistic constraints on individual investors and some mutual funds, for example, which are inherently constrained to be long-only. The benchmark uncorrelated rule defines a decoupling and risk diversification strategy; a core idea is that it aims to exceed the benchmark whether it rises or falls. The benchmark can be any asset; we choose the S&P 500 index as benchmark in our case study.

6 Case Study: Financial Forecasting and Decisions

One main goal is to examine and illustrate the utility of DDNMs, using our extended model uncertainty framework. Within that, a key applied interest is the utility in practical portfolio decision making. It has been empirically demonstrated by many that more accurate forecasts do not necessarily lead to better investment performance; hence, in addition to describing aspects of the Bayesian analysis in terms of inference on model structure and forecasting, we evaluate a number of performance measures of more practical relevance to the financial investment management context.

For higher-dimensional portfolios, appropriately structuring and constraining forecast distributions is critical; even very small changes in variances and covariances among assets can have important consequences on resulting portfolios [e.g. 4]. If conditional dependence structures can be appropriately captured by sparse DDNMs, the reduced parameter dimension can improve accuracy and stability of estimation and hence forecasting, and so lead to improved portfolios. The extended DDNM uncertainty analysis offers the ability to focus on classes of sparse models, and additional stability may then arise via the Bayesian averaging over multiple sparse models for prediction.

6.1 Data

We analyze $m = 13$ financial time series with $y_t$ being logged values of daily closing prices of 9 currency exchange rates relative to the US dollar, 2 commodities prices, and 2 U.S. stock indices; see Table 1. The time period of 2,979 working-week days from August 1st 2000 to December 31st 2011 includes periods of major growth as well as recession in the US and worldwide economies. The series represent major liquid benchmark securities across 3 asset classes to approximately reflect the global macroeconomic conditions. We model log prices directly, building on our experience that this is a surer route to useful predictive models than the traditional approach using returns [e.g. 11]; the basic point here is that moving to returns can “difference away” small changes in time that can be important in influencing short-term predictions if allowed via models on prices (or log prices) that have some opportunity to capture them. We split the data into a training and test data period: from August 1st 2000 to April 14th 2006 (1,489 observations) as the training data set, and then from April 17th 2006 to December 31st 2011 (1,490 log-prices) as the test data set to evaluate step ahead forecasting and portfolio outcomes.

6.2 Model Setup and Training Data Analysis

We first apply the DDNM with model uncertainty to analyze the training data. With a little over 300,000 possible models $M_{1:m}$, this is computationally accessible. Priors are as discussed above, with specific settings shown in the first 5 rows of Table 2. This defines a symmetric bivariate grid of $k = 5 \times 5 = 25$ values of the model discount factors $(\delta_j, \beta_j)$ for each $j$, a grid of values for the power discount factor $\alpha$.
Table 1: Financial time series in case study

<table>
<thead>
<tr>
<th>(j)</th>
<th>Name</th>
<th>Asset</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CHF</td>
<td>Swiss Franc</td>
</tr>
<tr>
<td>2</td>
<td>EUR</td>
<td>Euro</td>
</tr>
<tr>
<td>3</td>
<td>NSD</td>
<td>NASDAQ Composite Index</td>
</tr>
<tr>
<td>4</td>
<td>S&amp;P</td>
<td>S&amp;P 500 Index</td>
</tr>
<tr>
<td>5</td>
<td>NOK</td>
<td>Norwegian Krone</td>
</tr>
<tr>
<td>6</td>
<td>GBP</td>
<td>British Pound</td>
</tr>
<tr>
<td>7</td>
<td>AUD</td>
<td>Australian Dollar</td>
</tr>
<tr>
<td>8</td>
<td>NZD</td>
<td>New Zealand Dollar</td>
</tr>
<tr>
<td>9</td>
<td>ZAR</td>
<td>South African Rand</td>
</tr>
<tr>
<td>10</td>
<td>GOL</td>
<td>Gold</td>
</tr>
<tr>
<td>11</td>
<td>CAD</td>
<td>Canadian Dollar</td>
</tr>
<tr>
<td>12</td>
<td>JPY</td>
<td>Japanese Yen</td>
</tr>
<tr>
<td>13</td>
<td>OIL</td>
<td>Crude Oil</td>
</tr>
</tbody>
</table>

over \((0.95, 1]\) (following experiences in [27]), maximum TVAR model order \(d = 2\), and prior parental set inclusion probability of \(\rho = 0.3\).

Following this analysis, we then modify the candidate model set prior to the sequential analysis over the test data period. The point here is simply to recognise that many models in the full set of models have such low posterior probability based on the training data that it is justifiable— and then computationally efficient— to cut-back to a smaller space of potential models for further use in sequential forecasting and portfolio studies over the test data period. We remove models whose posterior probabilities after the training period are lower than \(th\), a small threshold; this study takes \(th = 0.001\) as per Table 2.

Table 2: Hyper-parameter values and control settings

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Setting</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\delta_j)</td>
<td>0.975 : 0.005 : 0.995</td>
<td>(th)</td>
<td>0.001</td>
</tr>
<tr>
<td>(\beta_j)</td>
<td>0.975 : 0.005 : 0.995</td>
<td>(nmc)</td>
<td>10,000</td>
</tr>
<tr>
<td>(\alpha)</td>
<td>0.950 : 0.005 : 1.000</td>
<td>Target return</td>
<td>0.5% (daily)</td>
</tr>
<tr>
<td>(d)</td>
<td>2</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(\rho)</td>
<td>0.3</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

For the test data analysis period, we run the extended DDNM analyses with model uncertainty, sequentially filtering and updating posterior model probabilities and conditional posteriors for model state vectors and volatilities within each model, and then evaluating 1– to 5-step ahead forecast distributions at each time point. Each forecasting exercise generates samples of size \(nmc = 10,000\) for evaluation of forecast moments, giving point forecasts and variance matrices that feed into the portfolio decision analyses under each of the three classes of portfolio loss functions. This decision analysis uses a daily base target return on portfolios of 0.1%, i.e., a 5—day target of 0.5%; this corresponds to an appropriately aggressive annual target return of 30%. We evaluate several risk characteristics of the optimised portfolios, as well as realised portfolio returns, over the test data period.

6.3 Forecast-based Model Assessment

Figure 1 displays the time trajectories of the marginal posterior probabilities \(p(\alpha|D_t)\), covering both training and testing periods. We can see that models with \(\alpha = 1\), are clearly dominated by those with \(\alpha < 1\),
confirming the relevance of power discounting of model probabilities. Also, the posterior probabilities of $\alpha < 1$ stay relatively stable over time with insignificant differences over the range of values less than 1 specified here; that is, $\alpha = 1$ is ruled out, but values on the 0.95 – 0.995 ranges are otherwise hardly discriminated.

Figure 1: Trajectories over time $t$ of posterior probabilities $p(\alpha|D_t)$ for each of the gridded values of $\alpha \in \{0.950, 0.955, \ldots, 0.995, 1.000\}$.

Figure 2 shows overall forecast accuracy measured by out-of-sample root mean squared error (RMSE) and mean absolute deviation (MAD) of the 1− and 5−step ahead point forecasts under model-averaged predictive distributions conditional on $\alpha$ over the test data period. As shown in both figures, higher $\alpha$ over this range leads to lower prediction accuracy. The relevance of $\alpha < 1$ is reinforced here. In some later summaries, we look at outcomes based on models with $\alpha = 0.98$ as an example; note that both the forecast accuracy and model posterior probabilities are relatively high at this power discounting level. These substantial improvements in forecasting at both 1− and 5−days ahead strongly support the strategy of power discounting in sequential updating of probabilities over models on purely statistical grounds; later we show additional support for values $\alpha < 1$ in terms of portfolio decision outcomes.

Some insights into the adequacy of model structure can be gained by viewing plots such as in Figure 3. This shows the trajectories of log prices with the 5−day ahead forecast mean and 90% credible intervals for the Oil series. The lower frames show scatter and QQ plots of the corresponding standardized 5−step forecast errors. There is evidence of slightly heavy-tailed departure from normal—as expected—and overall excellent conformance to the model. This typifies exploratory residual plots across the series—overall indicating no strong evidence of model inadequacies.
Figure 2: Out-of-sample prediction accuracy measured by RMSE and MAD of 1− and 5− step ahead point forecasts plotted against the power discount factor $\alpha$. This shows the dominance of smaller values of $\alpha$ at both forecast horizons on these two raw forecast accuracy measures, consistent with the preference for values less than 1 based on the posterior model probabilities in Figure 1. Note that use of much lower values of $\alpha$ than explored here would lead to low posterior probabilities and increasing RMSE and MAD measures; the range 0.95 − 0.98 appears to be a “sweet-spot” for this key parameter.

Figure 3: The log Oil price series and 5−day ahead forecasts (upper frame), with corresponding scatter and QQ plots (lower frames) of standardized 5−day ahead forecast errors against standard normal quantiles.
6.4 Dynamic Posterior Inferences on Components of Model Structure

6.4.1 Lags in Time-varying Autoregressive Components

Figure 4 shows the time trajectories of posterior means $E(p_{j\lambda}|D_t)$ for the effective TVAR lags for each series $j = 1 : 13$, together with the probabilities that $(p_{j\lambda} = 2|D_t)$. Note variation over time across all series as the model adapts to time-varying patterns in the data. To key out some example features, note that the posterior on lag 1 is high and stable over time for CHF, but for a burst of volatility during the early months of the global recession in early 2008; S&P shows somewhat more volatile patterns over time and favours higher lagged structure.

![Figure 4: Inferences on TVAR lag order.](image)

Figure 4: Inferences on TVAR lag order. The heat-maps show, for each series $j = 1 : 13$, the trajectories over time of posterior means $E(p_{j\lambda}|D_t)$ (upper frame) and probabilities on $(p_{j\lambda} = 2|D_t)$ (lower frame).

6.4.2 Series-specific Discount Factors

Figure 5 shows trajectories of posterior means for the discount factors, $E(\delta_j|D_t)$ and $E(\beta_j|D_t)$ for each series $j$ over the time frame. There are notable changes over time on each, reflecting adaptation of the underlying posterior model probabilities. For instance, we see clear shifts to favouring lower volatility discount factors $\beta_j$ across basically all series beginning around September 2008 onwards, i.e. as the global recession escalates. The model recognizes the need for increased volatility across the entire system, and appropriately adapts to the major changes experienced at that time. Later, in early 2010 as global markets
are stabilizing, the posterior shifts mass towards higher $\beta_j$ values as global and series-specific patterns lead to reduced volatility levels. Two other highlights are that the posterior for the S&P state discount factor $\delta_4$ favours high values throughout, reflecting the innate stability of relationships of this major aggregate index with predictors, and the posterior for the volatility discount factor $\beta_5$ on NOK also naturally reflects lower volatility in price fluctuations of this strong and stable currency relative to the other series.

![Figure 5: Inferences on series-specific discount factors. The heat-maps show, for each series $j = 1 : 13$, the trajectories over time of posterior means of discount factors for the model state vectors, $E(\delta_j|D_t)$ (upper frame), and of those for the residual volatility in observations, $E(\beta_j|D_t)$ (lower frame).](image)

6.4.3 DDNM Parental Sets

Trajectories of posterior probabilities of parental set membership for each series are shown in Figure 6. That is, for each series $j$ and potential parental series $i \in (j + 1) : m$, the posterior probability that $i \in pa(j)$ conditional on $D_t$ over time $t$. These figures exemplify the abilities of the model to: (i) focus on data-supported sparse models, as many such posterior probabilities are low across the entire time period, or for major time periods; (ii) identify strong predictive relationships through evaluation of high posterior probabilities of some contemporaneous parents being included; and (iii) adapt to changing circumstances, with probabilities showing more dynamics during the recessionary years in some cases. Inherently also, the variations over time formally accommodate patterns of collinearity among potential parental predictors for each series, and time variation in such patterns.
Figure 6: Heat-maps showing time trajectories of posterior parental set inclusion posterior probabilities for each asset over the test data period. The colorbars to the right indicate the probability scale.
Figure 6 (continued)
Keying out some examples, the posterior probability that S&P is a parent of NSD is high throughout the whole sample period, a relationship in accord with the common market sense of strong and sustained relationships among the two stock indices. Similarly, NZD is naturally a sustained parental predictor for AUD, with posterior probability close to 1 throughout the whole sample period. One example of changing parental structure is the case of Oil as a potential parent for CAD; the posterior probability is generally low during the later years of the great moderation, up to 2008 when it increased to higher levels in the global recession, and then maintains generally higher levels to the end of the time period.

One summary of the complexity/sparsity of model structure is the size of each parental $p_{j\gamma} = |pa(j)|$. By averaging across models with respect to model probabilities at any time, we can evaluate summaries such as the posterior mean $E(p_{j\gamma}|D_t)$. Such calculations show that—while there are clear dynamics over time in the posterior probabilities of individual parents—there is strong stability in terms of the effective parent set sizes. This stability in part reflects collinearities among potential parental predictors, and hence the positive relevance of a sparsity-inducing prior for parent inclusion. On the latter, the posterior selects “out” many potential parental predictors across the series for much of the time, again reflecting data-based support for relative sparsity. Rough summaries of posterior mean parental set sizes are that CHF, EUR, NSD have around 5-6 parents; OIL (of course, as the last in the order) has none while S&P, much higher in the order, also has just around 0-1; NOK, GBP have around 2-3, AUD has about 4-5, and the remaining indices have around 1.

### 6.5 Bayesian Portfolio Decision Analysis

#### 6.5.1 Performance Measures

We evaluate portfolio characteristics and outcomes, comparing results based on the three portfolio utility structures described in Section 5. For each, we evaluate separately in the contexts of the DDNM-based forecasts for both 1— and 5—day ahead portfolio rebalancing. The quantitative measures we use are the
following standard performance indicators:

- **Cumulative return (CR)**: Write $RR_t$ for the realized return of the portfolio at each period $t$. The cumulative return over a time period $\tau = 1 : t$ is then $CR_t = \prod_{\tau=1:t} (1 + RR_\tau)$.

- **Mean realized return (MRR)**: Over any time period $1 : t$, this is simply $MRR_t = t^{-1} \sum_{\tau=1:t} RR_\tau$.

- **Risk (R)**: The realized risk over any period $1 : t$ is simply the sample standard deviation $R_t$ of realized returns $RR_{1:t}$.

- **Projected Risk (PR)**: Projected risk is the finance term for the theoretical standard deviation of the forecast distribution of the optimised portfolio, as noted in Section 5. At any time $t$, if $(f_t, Q_t)$ are the mean vector and variance matrix of the forecast returns and $w_t$ the optimal weight vector, then $PR_t = \sqrt{w_t^\prime Q_t w_t}$.

- **Sharpe ratio (SR)**: This compares realized returns to realized risk via $SR_t = MRR_t / R_t$ over any period $1 : t$, typically converted to and reported on an annualized basis.

- **Projected Sharpe ratio**: This is the theoretical analogue of the realized Sharp ratio, given by $PSR_t = (w_t^\prime f_t) / PR_t$ for forecasts made at time $t$ and with resulting optimal weight vector $w_t$.

Further comparisons are made with a professional investment community benchmark, the Newedge CTA Index [3]. Professional money managers and commodity trading advisors (CTA) typically monitor managed futures accounts, which generally have exposure to a number of markets such as stocks, derivatives, commodities, energy, agriculture and currency. The CTA Index is designed to track the largest 20 CTAs (largest in terms of assets under management) and to be representative of the managed futures markets broadly. Since the portfolio of the 13 assets here is comparable to that managed by CTAs, we compare the performance of our portfolios with the publicly available Newedge CTA index as a key benchmark.

### 6.5.2 1−Day ahead forecasting and decisions

We set daily target returns to 0.1% corresponding to a target of 30% on an annual basis. Figure 7 shows results across the test data period based on 1−day ahead forecasts for model averaged DDNMs using differing values of the model probability power discount $\alpha$. The plot shows cumulative returns, Sharpe ratios and risks from each of the three portfolios, together with the CTA index. It can be seen that the target and benchmark neutral portfolios perform similarly well and beat the CTA index consistently in terms of both raw (CR) and risk-adjusted (SR) returns. Relative to the CTA Index, the excess risk incurred by these strategies are modest and outweighed by the improved returns. The target constraint portfolio, in contrast, has substantially poorer performance on all three metrics on this short-term, 1−day ahead basis.

### 6.5.3 5−Day ahead forecasting and decisions

With a 5−day target return of 0.5%, consistent with the 30% annual return, the parallel 5−day results appear in Figure 8. It is evident that when $0.96 \leq \alpha \leq 0.98$, both return measures are higher than for $\alpha$ outside this range, while the risk is only slightly higher. Forecasting and decisions using standard model uncertainty analysis ($\alpha = 1$) is evidently dominated in terms of portfolio performance by models...
Figure 7: Cumulative returns (CR), Sharpe ratios (SR) and risk (R) based on 1–day ahead forecasts from models differing only through the value of the power discount \( \alpha \). This shows similar performance of the target and benchmark neutral portfolios on all measures, and their dominance over the CTA index in terms of raw and risk-adjusted returns.

Figure 8: Cumulative returns (CR), Sharpe ratios (SR) and risk (R) based on 5–day ahead forecasts from models differing only through the value of the power discount \( \alpha \), with conclusions generally paralleling those under the 1–day analysis. One specific point to note is that the target constraint portfolio outperforms others at 5–days ahead but under-performs relatively at the 1–day horizon, a point linked to reversal and momentum effects in assets: long-only portfolios like the target constraint here tend to show reversal effects at shorter horizons while benefiting from momentum effects at longer horizons.
with $\alpha$ in this range, while models with smaller $\alpha$ clearly suffer degraded performance (due to over-discounting historical data and hence over-fitting more recent data). We also now see that target constraint portfolio shows generally superior performance in this longer-term, 5–day ahead analysis than at the shorter 1–day horizon, achieving generally higher returns at the cost of higher risk incurred by its “no-shorting” constraints.

Compared with the CTA index, all portfolios with $0.96 \leq \alpha \leq 0.98$ have higher raw and risk-adjusted returns, while the risks are also larger but better-compensated by higher returns. If we take $\alpha = 0.98$ as an example, more detailed performance comparisons are shown in Figure 9 and Table 3. The target constraint portfolio empirically realizes largest cumulative return over the full test data period, as a trade-off from also incurring the largest risk among the three portfolios. A point of practical interest is that, in terms of cumulative return, the portfolio with target constraint outperforms others at 5–days ahead but underperforms relatively at the 1–day horizon. This is consistent with the increased role of reversal rather than momentum effects on assets in the short-term, and vice versa in the long-term. That is, long-only portfolios—such as our target constraint portfolios—tend to show reversal effects at shorter horizons while benefiting from momentum effects at longer horizons.

![Figure 9: Trajectories over time of the cumulative returns (CR), projected risk (PR) and projected Sharpe ratio (PSR) based on 5–day ahead forecasts using the model based on power discount $\alpha = 0.98$.](image)
Table 3: Summary portfolio performance of 5−day analysis using $\alpha = 0.98$ (with CTA Index and Sharpe ratios).

<table>
<thead>
<tr>
<th></th>
<th>CTA Index</th>
<th>Target constrained</th>
<th>Benchmark neutral</th>
</tr>
</thead>
<tbody>
<tr>
<td>MRR</td>
<td>0.0008</td>
<td>0.0023</td>
<td>0.0012</td>
</tr>
<tr>
<td>Risk</td>
<td>0.0113</td>
<td>0.0201</td>
<td>0.0119</td>
</tr>
<tr>
<td>Sharpe ratio</td>
<td>0.4977</td>
<td>0.8150</td>
<td>0.7451</td>
</tr>
</tbody>
</table>

7 Summary Comments

Bayesian statistical thinking has had an enormous impact on sequential analysis and forecasting across a wide spectrum of areas—from core finance, to econometrics, to neuroscience, to IT, among others. The methodology of DDNMs developed here, building on the core existing theory of MDMs with a set of major extensions, aims to advance this impact by enriching the time series and forecasting toolbox with models that are increasingly flexible and useful. The keys to this are (a) the ability to customize model specifications for each univariate series separately, (b) overlaying this with flexible models of multivariate stochastic volatility, and (c) dealing adaptively over time with issues of model specification and uncertainty, in terms of both structural and parameter uncertainties. The class of DDNMs builds on MDMs to addresses point (a) with time-varying autoregressive model components as well as dynamic regressions in individual series. This is coupled with time-varying, Cholesky-style MDM structures to address point (b). The development of sequential learning and discount Bayesian model uncertainty analysis– applied to predictive and contemporaneous model structure as well as sets of defining model parameters– addresses point (c).

Faced with increasingly high-dimensional data with needs for increasingly fast data processing—coupled with basic interests in coherent summary inferences and predictions being rapidly computed revised—the continued emphasis on building bigger, more customized and hence more elaborate statistical models stresses us in what we know works at all, let alone what works well, in higher dimensions. Hence the interest in— and imperative to consider—models that are defined by sets of decoupled univariate models as starting points, but that are then properly recoupled for formal inferences and forecasting. The analysis developed here is an example of this: it utilises the inherent conditional independence structure of DDNMs to decouple the series into the set of univariate models to be run in parallel for forward filtering and one-step forecasting— with or without model uncertainty analysis; these are then recoupled for full, coherent multivariate forecasting.

The application to financial time series forecasting and decision analysis exemplifies the DDNM approach and the utility of the discount Bayesian model uncertainty analysis overlaid on the core model framework. The extended model uncertainty analysis highlights time-varying patterns in effective autoregressive lag structures, discount factors for stochastic volatilities, and in conditional contemporaneous dependencies among the series. Moreover, the decoupling property allows for time-variation in any of these features to be series-specific, providing flexibility in modelling individual series. Finally, the applied study clearly and substantially demonstrates the practical value and utility of the dynamic dependence network framework, evidencing its ability to extend and develop the portfolio of Bayesian forecasting and decision analysis methodology relevant in day-to-day applications in finance and business.

One specific point of current interest and for future exploration relates to the fact that the ordering of
the individual time series in the data vector is part of the model structure and specification, and one that
can have a practical impact on forecasts and decisions. Here we have focused on illustrating the impacts
on forecasting of AR lag order, series-specific discount factors, and sparse parental sets on forecasting; it
can be argued that the specification of the order of the series is an additional part of the model structure
that might be regarded as uncertain, and treated this way as well. A somewhat different view is that the
series order be regarded as a decision variable, and that Bayesian decision analysis be applied to guide
its choice. We note that recent work with related models take a third path in which the series order is
irrelevant, but that leads to needs for creativity in approximation of posterior and forecast distributions,
and more challenging computational methods for model fitting and forecasting [6]. These comments all
touch on open questions for future research linked to the series order question in DDNMs.

Appendix A: MDM Filtering and Forecasting Summaries

We give summary details of the equations defined by evolution and updating steps in the set of \( m \) univariate
DLMs in the MDM framework of Section 2.1, and in which each of the \( j = 1 : m \) decoupled univariate
DLMs have state and volatility evolutions as defined in Section 2.2.

As noted, we adopt random walk state evolution models \( \theta_{jt} = \theta_{j,t-1} + \omega_{jt} \) where the evolution error
\( \omega_{jt} \) is zero-mean normal, independent over time and across series, and has a time-dependent evolution
error variance matrix defined via a single discount factor \( \delta_j \in (0, 1] \). Coupled with this is a standard
random walk volatility model \( \lambda_{jt} = \lambda_{j,t-1}\eta_{jt}/\beta_j \) where the \( \eta_{jt} \) are independent beta random variates with
time-dependent beta parameters defined via the single discount factor \( \beta_j \in (0, 1] \) for series \( j \). These are
standard models and full details appear in [26] and [15]. The summary details here provide the bases
for sequential analyses. These apply in parallel over series \( j \) for forward-filtering and forecasting within-
series, and are then coupled together for multivariate forecasting.

A1. Posterior at time \( t-1 \)

Standing at time \( t-1 \), we have series \( -j \) specific normal/gamma posteriors

\[
(\theta_{j,t-1}, \lambda_{j,t-1}|D_{t-1}) \sim NG(m_{j,t-1}, C_{j,t-1}, n_{j,t-1}, n_{j,t-1}s_{j,t-1}).
\]

A2. Evolution from time \( t-1 \) to \( t \)

Evolving to time \( t \), the state vector \( \theta_{j,t-1} \) undergoes a linear state evolution and the precision \( \lambda_{j,t-1} \) under-
goes a coupled gamma:beta discount evolution. The implied prior for the next time point is then

\[
(\theta_{jt}, \lambda_{jt}|D_{t-1}) \sim NG(a_{jt}, R_{jt}, r_{jt}, r_{jt}s_{j,t-1})
\]

with \( a_{jt} \equiv m_{j,t-1}, \ R_{jt} = C_{j,t-1}/\delta_j \) and \( r_{jt} = \beta_j n_{jt} \). Here the \( \delta_j \) and \( \beta_j \) are in \( (0, 1] \) and, typically, take
larger values in this range [e.g. 26, chaps 6, 10].

A3. 1-step ahead forecasts at time \( t-1 \).

The implied predictive distribution is T with \( r_{jt} \) degrees of freedom, namely

\[
(y_{jt}|y_{pa(j),t}, D_{t-1}) \sim T_{r_{jt}}(f_{jt}(y_{pa(j),t}), q_{jt}(y_{pa(j),t}))
\]

25
where \( f_{jt}(y_{pa(j),t}) = F'_{jt}a_{jt} \) and \( q_{jt}(y_{pa(j),t}) = s_{j,t-1} + F'_{jt}R_{jt}F_{jt} \). Defining
\[
\alpha_{jt} = \begin{pmatrix} a_{j\phi t} \\ a_{j\gamma t} \end{pmatrix} \quad \text{and} \quad R_{jt} = \begin{pmatrix} R_{j\phi t} \\ R_{j\gamma t} \end{pmatrix},
\]
we have
\[
f_{jt}(y_{pa(j),t}) = x'_{jt}a_{j\phi t} + y'_{pa(j),t}a_{j\gamma t}
\]
\[
q_{jt}(y_{pa(j),t}) = s_{j,t-1} + y'_{pa(j),t}R_{jt}y_{pa(j),t} + 2y'_{pa(j),t}R'_{j\phi t}x_{jt} + x'_{jt}R_{j\phi t}x_{jt}.
\]

A4. Updating equations at time \( t \)

With the normal/gamma prior above, the implied normal/gamma posterior is
\[
(\theta_{jt}, \lambda_{jt}|D_t) \sim NG(m_{jt}, C_{jt}, n_{jt}, n_{jt}s_{jt})
\]
with defining parameters computed using standard updating equations [15, sect. 14.3], as follows:

First, compute the following:
\[
\begin{align*}
e_{jt} &= y_{jt} - F'_{jt}a_{jt} \\
q_{jt} &= s_{j,t-1} + F'_{jt}R_{jt}F_{jt} \\
A_{jt} &= R_{jt}F_{jt}/q_{jt} \\
z_{jt} &= (r_{jt} + e^2_{jt}/q_{jt})/(r_{jt} + 1)
\end{align*}
\]

Then, compute the posterior parameters:
\[
\begin{align*}
m_{jt} &= \alpha_{jt} + A_{jt}e_{jt} \\
C_{jt} &= (R_{jt} - A_{jt}A'_{jt}q_{jt})z_{jt} \\
n_{jt} &= r_{jt} + 1 \\
s_{jt} &= s_{j,t-1}z_{jt}
\end{align*}
\]

A5. Multi-step forecasting from any time point \( t \)

For \( k \)-step ahead forecasting at any time \( t \), we clearly need the future values of parental variables for each model, so make that explicit in notation. At time \( t \), assuming knowledge of future \( x_{j,t+k} \) and being explicit about the need for values of the parental series, we have the following.

First, the \( k \)-step ahead prior at time \( t \) is
\[
(\theta_{j,t+k}, \lambda_{j,t+k}|D_t) \sim NG(a_{jt}(k), R_{jt}(k), r_{jt}(k), r_{jt}(k)s_{jt})
\]
where \( a_{jt}(k), R_{jt}(k) \) are updated inductively from \( a_{jt}(k-1), R_{jt}(k-1) \) and \( r_{jt}(k) = \beta_j n_{jt} \). Based on this, we have forecast distribution
\[
(y_{j,t+k}|y_{pa(j),t+k}, D_t) \sim T_{r_{jt}}(f_{j,t+k}(y_{pa(j),t+k}), q_{j,t+k}(y_{pa(j),t+k}))
\]
where \( f_{j,t+k}(y_{pa(j),t+k}) = F'_{j,t+k}a_{j,t}(k) \) and \( q_{j,t+k}(y_{pa(j),t+k}) = s_{j,t-1} + F'_{j,t+k}R_{j,t}(k)F_{j,t+k} \). Write
\[
\alpha_{j,t}(k) = \begin{pmatrix} a_{j\phi t}(k) \\ a_{j\gamma t}(k) \end{pmatrix} \quad \text{and} \quad R_{jt}(k) = \begin{pmatrix} R_{j\phi t}(k) & R_{j\phi t}(k) \\ R_{j\gamma t}(k) & R_{j\gamma t}(k) \end{pmatrix}.
\]
Then
\[ f_{j,t+k}(y_{pa(j),t+k}) = a'_{j,t+k}a_{j\phi t}(k) + y'_{pa(j),t+k}a_{j\gamma t}(k) \]
\[ q_{j,t+k}(y_{pa(j),t+k}) = s_{j,t-1} + y'_{pa(j),t+k}R_{j\gamma t}(k)y_{pa(j),t+k} + 2y'_{pa(j),t+k}R_{j\phi t}(k)x_{j,t+k} + x'_{j,t+k}R_{j\phi t}(k)x_{j,t+k}. \]  
(13)

**Appendix B: Joint Predictive Moments and Precision Matrices in MDMs**

**B1. Joint Predictive Mean and Variance Matrix**

As discussed in Section 2.3, we are interested in the moments
\[ f_t = E(y_t|\mathcal{D}_{t-1}) \quad \text{and} \quad Q_t = V(y_t|\mathcal{D}_{t-1}) \]
under (3). Assume that, for all \( j, t, r_{jt} > 1 \) so that these moments exist. The compositional model form allows for recursive moment computation that recognizes the appearance of contemporaneous values of the \( y_{pa(j),t} \) in the conditioning of forecasts for \( y_{jt} \). Details now follow (note also that \( k\)—step ahead computations are very similar, and so details are omitted here).

For each \( j = 1 : m - 1 \), denote the mean vector and variance matrix \((m - j)\)—vector \( y_{j+1:m,t} \) by
\[ f_{j+1:m,t} = E(y_{j+1:m,t}|\mathcal{D}_{t-1}) \quad \text{and} \quad Q_{j+1:m,t} = V(y_{j+1:m,t}|\mathcal{D}_{t-1}). \]  
(14)

**i. Start at \( j = m \):** compute the univariate mean and variance of \( y_{mt} \),
\[ f_{mt} = f_{mt}(\emptyset), \quad q_{mt} = q_{mt}(\emptyset)r_{mt}/(r_{mt} - 2) \]
using the implied simplified forms of (11) with all terms in \( y_{pa(m)} \) set to zero. Insert \( f_{mt} \) as the \( m \)—the element of \( f_t \) and \( q_{mt} \) as the \((m, m)\)—element of \( Q_t \).

**ii. For \( j = m - 1, m - 2, \ldots, 1 \) in turn:**

- At this point, we have already computed the values of the moments of (14) from the previous steps. These are used in the following calculations. First, write \( f_{pa(j),t} \) for the subvector of means in \( f_{j+1:m,t} \) on elements in \( pa(j) \) only, and \( Q_{pa(j),t} \) for the corresponding submatrix of \( Q_{j+1:m,t} \). Then, the marginal mean \( f_{jt} \) and variance \( q_{jt} \) of \( y_{jt} \) are computed as follows.
\[ f_{jt} = a'_{jt}a_{j\phi t} + f'_{pa(j),t}a_{j\gamma t}, \]
\[ q_{jt} = (s_{j,t-1} + u_{jt})r_{jt}/(r_{jt} - 2) + a'_{j\gamma t}Q_{pa(j),t}a_{j\gamma t} \]  
(15)

where
\[ u_{jt} = f'_{pa(j),t}R_{j\gamma t}f_{pa(j),t} + tr(R_{j\gamma t}Q_{pa(j),t}) + 2a'_{jt}R_{j\phi t}f_{pa(j),t} + x'_{jt}R_{j\phi t}x_{jt}. \]
Insert \( f_{jt} \) as the \( j \)—th element of \( f_t \) and \( q_{jt} \) as the \((j, j)\)—element of \( Q_t \), respectively.

- Compute the covariance vector \( C(y_{jt}, y_{j+1:m,t})|\mathcal{D}_{t-1} \) as follows. Write \( a_{j+1:m\gamma t} \) for the \((m - j)\)—vector that extends \( a_{j\gamma t} \) with zeros for elements \( h \notin pa(j) \). Then
\[ C(y_{jt}, y_{j+1:m,t}|\mathcal{D}_{t-1}) = Q_{j+1:m,t}a_{j+1:m\gamma t} = q_{j,j+1:m,t}. \]
Insert element \( h \) of this vector as the \((j, h + j) \) and \((h + j, j) \) entries of \( Q_t \), for \( h = 1 : m - j. \)
iii. End: at this point, \( j = 1 \) and we have filled in all elements of the \( m \times m \) matrix \( \mathbf{Q}_t \).

**B2. Predictive precision matrix**

Consider now the predictive precision matrix \( \mathbf{K}_t = \mathbf{V}(\mathbf{y}_t|\mathbf{D}_{t-1})^{-1} \). For each \( j = 1 : m - 1 \), denote the precision matrix of the \((m - j)\)-vector \( \mathbf{y}_{j+1:m,t} \) by \( \mathbf{K}_{j+1:m,t} = \mathbf{V}(\mathbf{y}_{j+1:m,t}|\mathbf{D}_{t-1})^{-1} \), in parallel to the subvector means and variance matrices in eqn. (14) above; again, at \( j = m - 1 \) these are scalars.

To compute the precision matrix at each time \( t \), we can avoid matrix inversion by utilizing the interim products, namely the covariance vectors \( \mathbf{q}_{j,j+1:m,t} \) that have been already calculated above. This block-wise inversion decreases computational instability and reduce complexity to \( O(m^2) \), and is especially efficient under sparse models for larger \( m \). The computation of \( \mathbf{K}_t \) is as follows.

i. **Start at** \( j = m \): Compute the precision of \( \mathbf{y}_{m,t} \),

\[
\mathbf{K}_{m:m,t} = 1/\mathbf{q}_{m,t}.
\]

ii. **For** \( j = m - 1, m - 2, \ldots, 1 \) **in turn:**

- Given the computed \( \mathbf{q}_{j,j+1:m,t} \), compute the precision matrix \( \mathbf{K}_{j:m,t} \) via its partition as

\[
\begin{pmatrix}
\mathbf{k}_{jt} & \mathbf{h}_{jt} \\
\mathbf{h}_{jt}' & \mathbf{H}_{j,t}
\end{pmatrix}
\]

with entries

\[
\mathbf{k}_{jt}^{-1} = \mathbf{q}_{jt} - \mathbf{q}_{j,j+1:m,t}\mathbf{K}_{j+1:m,t}\mathbf{q}_{j,j+1:m,t}',
\]

\[
\mathbf{h}_{jt} = -\mathbf{k}_{jt}\mathbf{q}_{j,j+1:m,t}\mathbf{K}_{j+1:m,t},
\]

\[
\mathbf{H}_{j,t} = \mathbf{K}_{j+1:m,t} + \mathbf{k}_{jt}^{-1}\mathbf{h}_{jt}'\mathbf{h}_{jt}.
\]

Here \( \mathbf{k}_{jt} \) is a scalar, so this recursive computation of \( \mathbf{K}_t \) avoids matrix inversions.

iii. **End:** at this point, \( j = 1 \) and we have filled in all elements of the precision matrix \( \mathbf{K}_t \), i.e. \( \mathbf{K}_{1:m,t} \).

**Appendix C: Forecast Moments in Mixtures of DDNMs**

**C1. 1–Step Ahead Forecast Moments in Discrete Mixtures of DDNMs**

With reference to 1–step ahead forecasting in mixtures of DDNMs arising via Bayesian model uncertainty analysis as in Section 4.3, basic technical details are noted here. This gives the analytic forms for 1–step ahead joint predictive means, covariance and precision matrices, via direct extension of the analytic results in Appendix B.

Standing at time \( t - 1 \), we introduce the following notation for marginal and model-conditional forecast means and variances within each univariate series \( j = 1 : m \). First, label the full set of possible series \( j \) models as \( \mathcal{M}_j = \mu \) (in an arbitrary order), and denote the number of such models by \( m_j \) so that \( \mu = 1 : m_j \) indexes the set.
Means and variances: Write
\[
E(y_{jt}|D_{t-1}) \equiv E(y_{jt}|D_{j,t-1}) = f_{jt}, \quad V(y_{jt}|D_{t-1}) \equiv V(y_{jt}|D_{j,t-1}) = q_{jt},
\]

\[
E(y_{jt}|D_{j,t-1}, M_j = \mu) = f_{j\mu}, \quad V(y_{jt}|D_{j,t-1}, M_j = \mu) = q_{j\mu},
\]

for \( j = 1 : m \) and \( \mu = 1 : m_j \). Then standard results for moments of mixtures [e.g. 26, sect 12.2] give
\[
f_{jt} = E(y_{jt}|D_{j,t-1}) = \sum_{\mu=1:m_j} f_{j\mu} p(M_j = \mu|D_{t-1}), \quad (16)
\]
\[
q_{jt} = V(y_{jt}|D_{j,t-1}) = \sum_{\mu=1:m_j} [ (f_{j\mu} - f_{jt})^2 + q_{j\mu} ] p(M_j = \mu|D_{t-1}). \quad (17)
\]

Covariances: Now consider two series \( y_{ht}, y_{jt} \) where \( 1 \leq h < j \leq m \). The 1-step forecast covariance at time \( t \) can be evaluated as
\[
C(y_{ht}, y_{jt}|D_{t-1}) = \sum_{\mu=1:m_h} C(y_{ht}, y_{jt}|D_{t-1}, M_h = \mu) p(M_h = \mu|D_{t-1}). \quad (18)
\]
This can be seen as follows. First, since \( h < j \) we have \( p(y_{jt}|M_h = \mu, D_{t-1}) = p(y_{jt}|D_{t-1}) \) and thus
\[
E(y_{jt}|M_h = \mu, D_{t-1}) = E(y_{jt}|D_{t-1}) \quad \text{for all} \mu \in \{1, \cdots, m_h\}.
\]
Then
\[
C(y_{ht}, y_{jt}|D_{t-1}) = E(y_{ht}y_{jt}|D_{t-1}) - E(y_{ht}|D_{t-1})E(y_{jt}|D_{t-1})
= \sum_{\mu=1:m_h} [ E(y_{ht}y_{jt}|D_{t-1}, M_h = \mu) - E(y_{ht}|D_{t-1}, M_h = \mu)E(y_{jt}|D_{t-1}) ] p(M_h = \mu|D_{t-1})
= \sum_{\mu=1:m_h} [ E(y_{ht}y_{jt}|D_{t-1}, M_h = \mu) - E(y_{ht}|D_{t-1}, M_h = \mu)E(y_{jt}|D_{t-1}, M_h = \mu) ] p(M_h = \mu|D_{t-1})
= \sum_{\mu=1:m_h} C(y_{ht}, y_{jt}|D_{t-1}, M_h = \mu) p(M_h = \mu|D_{t-1}),
\]
as stated.

Recursive evaluation of full mean vector and variance matrix: The joint predictive mean \( f_t = E(y_t|D_{t-1}) \) and variance matrix \( Q_t = V(y_t|D_{t-1}) \) can now be calculated recursively, as follows.

i. Start at \( j = m \): For each \( \mu = 1 : m_m \), compute the univariate mean and variance of \( y_{mt} \) under each \( M_m = \mu \), namely
\[
f_{j\mu t} = f_{\mu | m_{jt}}(\emptyset), \quad q_{j\mu t} = q_{\mu | m_{jt}}(\emptyset)r_{m_{jt}}/(r_{m_{jt}} - 2)
\]
using the implied simplified forms of (15) with all terms in \( y_{pa(m)} \) set to zero. Then calculate the marginal predictive mean and variance of \( y_{jt}|D_{t-1} \) using eqn. (16). Insert \( f_{\mu t} \) as the \( m \)-th element of \( f_t \) and \( q_{\mu t} \) as the \( (m, m) \)-element of \( Q_t \).
ii. For each \( j = m - 1, m - 2, \ldots, 1 \) in turn: Visit each model \( \mathcal{M}_j = \mu \) in \( 1 : m_j \). Make explicit in the notation that the parental set for series \( j \) generally depends on the model by writing \( pa(j|\mu) \) here. Then, compute the following quantities in parallel:

\[
\begin{align*}
    f_{jmt} &= E[ E(\mathbf{x}'_{jmt} \mathbf{a}_{jmu}\theta + \mathbf{y}'_{pa(j|\mu),t} \mathbf{a}_{jmu\gamma t}| \mathbf{y}_{pa(j|\mu),t}, \mathcal{D}_{t-1}, \mathcal{M}_j = \mu)| \mathcal{D}_{t-1}, \mathcal{M}_j = \mu] \\
    &= \mathbf{x}'_{jmt} \mathbf{a}_{jmu\theta} + f'_{pa(j|\mu),t} \mathbf{a}_{jmu\gamma t}, \\
    q_{jmt} &= (s_{jmu,t-1} + u_{jmt})r_{jmt}/(r_{jmt} - 2) + a'_{jmu\gamma t} \mathbf{Q}_{pa(j|\mu),t} \mathbf{a}_{jmu\gamma t},
\end{align*}
\]

where \( u_{jmt} = f'_{pa(j|\mu),t} \mathbf{R}_{jmu\gamma t} f_{pa(j|\mu),t} + \text{tr}(\mathbf{R}_{jmu\gamma t} \mathbf{Q}_{pa(j|\mu),t}) + 2 \mathbf{x}'_{jmt} \mathbf{R}_{jmu\gamma t} f_{pa(j|\mu),t} + \mathbf{x}'_{jmt} \mathbf{R}_{jmu\theta} \mathbf{x}_{jmt} \). These two moments are obtained from the conditional distribution given by (11).

Then calculate \( q_{jmt} \) and \( f_{jmt} \) according to (16), and insert \( f_{jmt} \) as the \( j \)-th element of \( \mathbf{f}_t \) and \( q_{jmt} \) as the \( (j,j) \)-element of \( \mathbf{Q}_t \), respectively. Write \( \mathbf{a}_{j+1:m,\mu\gamma t} \) for the \((m-j)\)-vector that extends \( \mathbf{a}_{j\mu\gamma t} \) with zeros for elements \( h \notin pa(j|\mu) \). Then

\[
C(y_{jt}, y_{j+1:m,t}| \mathcal{D}_{t-1}, \mathcal{M}_j) = \mathbf{Q}_{j+1:m,t} \mathbf{a}_{j+1:m,\mu\gamma t}.
\]

Then calculate \( C(y_{jt}, y_{j+1:m,t}| \mathcal{D}_{t-1}) = q_{j:j+1:m,t} \) according to (18), and insert element \( h \) of this vector as the \((j,h+j)\) and \((h+j,j)\) entries of \( \mathbf{Q}_t \), for \( h = 1 : m - j \).

iii. End: at this point, \( j = 1 \) and we have filled in all elements of the \( m \)-vector \( \mathbf{f}_t \), \( m \times m \) matrix \( \mathbf{Q}_t \).

The precision matrix can be calculated recursively in a similar fashion, paralleling Section 7.

C2: \( k \)-step Forecasting via Simulation

The \( k \)-step-ahead predictive mean and variance matrix at any time \( t \) cannot be directly evaluated analytically unless \( k = 1 \) (above). Hence we utilize direct and straightforward Monte Carlo simulations. Specify a Monte Carlo sample size \( nmc \) and proceed with the following steps. This generates random samples from the full predictive \( p(y_{t+1:t+k}| \mathcal{D}_t) \), i.e., giving synthetic future trajectories over all time points \( t + r \) for \( r = 1 : k \).

Monte Carlo averaging at each time point then provides approximations to predictive means and variance matrices, and any other quantities of interest. These can be tuned for accuracy by simply increasing \( nmc \) as the simulations are both parallelizable and computationally cheap per sample.

In the details below, we again extend the parental set notation so that, for each series \( j \) and any specific model \( \mathcal{M}_j = \mu \) in \( 1 : m_j \), the parental set is now denoted by \( pa(j|\mu) \). Notice that when \( j = m \), we have \( pa(j|\mu) = \emptyset \) for all \( \mathcal{M}_m = \mu \in \{1, \ldots, m_m\} \).

For each series \( j = m, m-1, \ldots, 1 \) in turn, we simulate a Monte Carlo sample of size \( nmc \) of values of \( y_{j,t+r} \) over \( r = 1, 2, \ldots, k \). At each series index \( j < m \), the values of any required parental predictors \( l \in pa(j|\mu) \) for the generated model \( \mathcal{M}_j = \mu \) will have been simulated at previous step \( l \), and so are available as conditional predictors for series \( j \). The process is as follows.

Start: For each \( j = m, m-1, \ldots, 1 \):

1. For the current series index \( j \), sample the discrete posterior \( p(\mathcal{M}_j| \mathcal{D}_{t-1}) \) over models \( \mathcal{M}_j \) to generate a sample of \( nmc \) models \( \mathcal{M}_j = \mu^i, i = 1 : nmc \).
For $r = 1$—step ahead, generate samples $y_{j,t+1}$ by simulating from the 1—step ahead T distribution $p(y_{j,t+1}|y_{pa(j),t+1}, D_t, M_j = \mu^i)$; this is just eqn. (12) at $k = 1$. Each sample $i$ is generated from this conditional based on the $i$—th sample value of the parental set.

For each of $r = 2, \cdots, k$—steps ahead in sequence, repeat the above computation to generate sample values $y_{j,t+r}$; at step $r$, this again simulates T distributions as in eqn. (12) at $k = r$. Each sample $i$ is based on such a T distribution whose parameters involve the recently simulated values of all needed predictors to evaluate both the parental vector $y_{pa(j),t+r}$ and the vector $x_{j,t+r}^i$ if it includes lagged values of any of the series.

2. Based on these Monte Carlo samples now running over series $m, m-1, \ldots, j$, compute series $j$ moments and update the saved Monte Carlo information, as follows.

For each step ahead $r = 1 : k$ in this order:

- Compute the sample mean and variance of $\{y_{j,t+r}, i = 1 : nmc\}$ as the Monte Carlo approximations to the predictive moments of $y_{j,t+r}$, i.e., the values $f_{jt}(r)$ and $q_{jt}(r)$. Insert $f_{jt}(r)$ as the $j$—th element of $f_t(r)$ and $q_{jt}(r)$ as the $(j, j)$ element of $Q_t(r)$.
- Compute the sample covariance vector of $\{y_{j,t+r}, y_{j+1:t+r}, i = 1 : nmc\}$ as the Monte Carlo approximation to predictive covariance vector $C(y_{j,t+r}, y_{j,j+1:t+r}|D_t)$, insert element $n$ of this vector as the $(j, n)$ and $(n, j)$ entries of $Q_t(r)$, for $n = j + 1 : m$.

3. If $j > 1$, move to the next series $j \leftarrow j - 1$ and repeat; otherwise, stop and save the complete set of Monte Carlo samples, if desired, as well as the complete predictive mean vectors $f_t(r)$ and variance matrices $Q_t(r)$ over $r = 1 : k$.

Acknowledgements

We thank the editor-in-chief for guidance in revising the paper for publication, and the positive and constructive comments of two anonymous referees. This work was completed while Zoey Zhao was a PhD student in the Department of Statistical Science at Duke University, while aspects of the work were developed while Meng Xie was an undergraduate student at Duke. The research was partly supported by a grant from the National Science Foundation [DMS-1106516]. Any opinions, findings and conclusions or recommendations expressed in this work are those of the authors and do not necessarily reflect the views of the NSF.

References


32


