SCALABLE BAYESIAN REDUCED-ORDER MODELS FOR SIMULATING HIGH-DIMENSIONAL MULTISCALE DYNAMICAL SYSTEMS

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Abstract. While existing mathematical descriptions can accurately account for phenomena at microscopic scales (e.g. molecular dynamics), these are often high-dimensional, stochastic and their applicability over macroscopic time scales of physical interest is computationally infeasible or impractical. In complex systems, with limited physical insight on the coherent behavior of their constituents, the only available information is data obtained from simulations of the trajectories of huge numbers of degrees of freedom over microscopic time scales. This paper discusses a Bayesian approach to deriving probabilistic coarse-grained models that simultaneously address the problems of identifying appropriate reduced coordinates and the effective dynamics in this lower-dimensional representation. At the core of the models proposed lie simple, low-dimensional dynamical systems which serve as the building blocks of the global model. These approximate the latent, generating sources and parameterize the reduced-order dynamics. We discuss parallelizable, online inference and learning algorithms that employ Sequential Monte Carlo samplers and scale linearly with the dimensionality of the observed dynamics. We propose a Bayesian adaptive time-integration scheme that utilizes probabilistic predictive estimates and enables rigorous concurrent simulation over macroscopic time scales. The data-driven perspective advocated assimilates computational and experimental data and thus can materialize data-model fusion. It can deal with applications that lack a mathematical description and where only observational data is available. Furthermore, it makes non-intrusive use of existing computational models.

Key words.

AMS subject classifications.

1. Introduction. The present paper is concerned with the development of probabilistic coarse-grained models for high-dimensional dynamical systems with a view of enabling multiscale simulation. We describe a unified treatment of complex problems described by large systems of deterministic or stochastic ODEs and/or large number of data streams. Such systems arise frequently in modern multi-physics applications either due to the discrete nature of the system (e.g. molecular dynamics) or due to discretization of spatiotemporal models (e.g. PDEs):

\[ \frac{dy_t}{dt} = f(y_t), \ y \in \mathcal{Y} \]  

(1.1)

where \( \dim(\mathcal{Y}) >> 1 \) (e.g. \( \mathbb{R}^d, d >> 1 \)). Stochastic versions are also frequently encountered:

\[ \frac{dy_t}{dt} = f(y_t; u_t) \]  

(1.2)

where \( u_t \) is a driving stochastic process (i.e. Wiener process). Uncertainties could also appear in the initial conditions that accompany the aforementioned systems of equations.

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Even though the numerical solution of (stochastic) ODEs is a well-studied subject and pertinent computational libraries are quite mature, traditional schemes are impractical or infeasible for systems which are high-dimensional and exhibit a large disparity in scales. This is because most numerical integrators must use time-steps of the order of the fastest scales which precludes solutions over long time ranges that are of interest for physical and engineering purposes. In the context of atomistic simulations, practically relevant time scales exceed typical integration steps of $\sim 1\text{fs}$ by several orders of magnitude. Furthermore, when numerical solutions of transient PDEs are sought, resolution and accuracy requirements give rise to systems with more than $10^9$ degrees of freedom where the integration time steps are slaved by fast reaction rates or high oscillation frequencies. This impedes their solution and frequently constitutes computationally infeasible other important tasks such as stability analysis, sensitivity, design and control.

Multiscale dynamical systems exist independently of the availability of mathematical models. Large numbers of time series appear in financial applications, meteorology, remote sensing where the phenomena of interest unfold also over a large range of time scales. A wealth of time series data is also available in experimental physics and engineering which by themselves or in combination with mathematical models can be useful in analyzing underlying phenomena by deriving reduced, predictive descriptions.

Quite frequently the time evolution of all the observables is irrelevant for physical and practical purposes and the analysis is focused on a reduced set of variables or reaction coordinates $\hat{y}_t = P(y_t)$ obtained by an appropriate mapping $P : \mathcal{Y} \rightarrow \hat{\mathcal{Y}}$. The goal is then to identify a closed, deterministic or stochastic system of equations with respect to $\hat{y}_t$, e.g.:

$$\frac{d\hat{y}_t}{dt} = \hat{f}(\hat{y}_t), \quad \hat{y}_t \in \hat{\mathcal{Y}} \quad (1.3)$$

In the context of equilibrium thermodynamics where ensemle averages with respect to the invariant distribution of $\hat{y}_t$ are of interest, coarse-graining amounts to free-energy computations. In the nonequilibrium case and when an invariant distribution exists, a general approach for deriving effective dynamics is based on Mori-Zwanzig projections. Other powerful numerical approaches to identify the dynamical behavior with respect to the reduced coordinates include transition path sampling, the transfer operator approach, the nudged elastic band, the string method, Perron cluster analysis and spectral decompositions. Marked efforts in chemical kinetics have led to an array of computational tools such as computational singular perturbation, the intrinsic low-dimensional manifold approach and others. Notable successes in overcoming the timescale dilemma have also been achieved in the context of MD simulations (or Hamiltonian systems in general).

In several problems, physical or mathematical arguments have led analysts to identify a few, salient features and their inter-dependencies that macroscopically describe the behavior of very complex systems consisting of a huge number of individuals/agents/components/degrees of freedom. These variables parameterize a low-dimensional, attracting, invariant, “slow” manifold characterizing the long-term process dynamics. Hence the apparent complexity exhibited in the high-dimensionality and the multiscale character of the original model is a pretext of a much simpler, latent structure that, if revealed, could make the aforementioned analysis tasks much more tractable. The emergence of macroscopic, coherent behavior has been the foundation
of coarse-grained dynamic models that have been successful in a wide range of applications. The coarse-grained parameterization and associated model depend on the analysis objectives and particularly on the time scale one wishes to make predictions. Modern approaches with general applicability such as the Equation-free method [92] or Heterogeneous Multiscale Method (HeMM, [47]) are also based on the availability of reduced coordinates and in the case of HeMM of a macroscopic model which is informed and used in conjunction with the microscale model.

Largely independently of the developments in the fields of computational physics and engineering, the problem of deriving, predictive reduced-order models for a large number of time series that potentially exhibit multiple scales has also been addressed in statistics and machine learning communities [140, 53] with applications in network analysis [33], environmetrics [141], sensor network monitoring [142, 117], moving object tracking [4], financial data analysis [5], computer model emulation [103]. Significant advances have been achieved in modeling [131], forecasting [143] and developing online, scalable algorithms [51, 126, 91, 94, 89, 145] that are frequently based on the discovery of hidden variables that provide insight to the intrinsic structure of streaming data [51, 35, 110, 109, 85].

The present paper proposes a data-driven alternative that is able to automatically coarse-grain high-dimensional systems without the need of preprocessing and availability of physical insight. The data is most commonly obtained by simulations of the most reliable, finest-scale (microscopic) model available. This is used to infer a lower-dimensional description that captures the dynamic evolution of the system at a coarser scale (i.e. a macroscopic model). The majority of available techniques address separately the problems of identifying appropriate reduced coordinates and the effective dynamics in this lower-dimensional representation. It is easily understood that the solution of one affects the other. We propose a general framework where these two problems are simultaneously solved and coarse-grained models are built from the ground up. We propose procedures that concurrently infer the macroscopic dynamics and their mapping the high-dimensional, fine-scale description. As a result no errors or ambiguity are introduced when the fine-scale model needs to be reinitialized in order to obtain additional simulation data. To that end, we advocate a largely unexplored in computational physics perspective based on the Bayesian paradigm which provides a rigorous foundation for learning from data. It is capable of quantifying inferential uncertainties and, more importantly, uncertainty due to information loss in the coarse-graining process.

We present a Bayesian state-space model where the reduced, coarse-grained dynamics are parametrized by tractable, low-dimensional dynamical models. These can be viewed as experts offering opinions on the evolution of the high-dimensional observables. Each of these modules could represent a single latent regime and would therefore be insufficient by itself to explain the whole system. As is often the case with real experts, their opinions are valid under very specific regimes. We propose therefore a framework for dynamically synthesizing such models in order to obtain an accurate global representation that retains its interpretability and computational tractability.

An important contribution of the paper, particularly in view of enabling simulations of multiscale systems, is online inference algorithms based on Sequential Monte Carlo that scale linearly with the dimensionality of the observables \( d \) (Equation (1.1)). These allow the recursive assimilation of data and re-calibration of the coarse-grained dynamics. The Bayesian framework adopted provides probabilistic predictive esti-
mates that can be employed in the context of adaptive time-integration. Rather than determining integration time steps based on traditional accuracy and stability metrics, we propose using measures of the predictive uncertainty in order to decide how long into the future the coarse-grained model can be used. When the uncertainty associated with the predictive estimates exceeds the analyst’s tolerance, the fine-scale dynamics can be consistently re-initialized in order to obtain additional data that sequentially update the coarse-grained model.

In agreement with some recently proposed methodologies \[92, 93\], the data-driven strategy can seamlessly interact with existing numerical integrators that are well-understood and reliably implemented in several legacy codes. In addition, it is suitable for problems where observational/experimental data must be fused with mathematical descriptions in a rigorous fashion and lead to improved analysis and prediction tools.

The structure of the rest of the paper is as follows. In Section 2 we describe the proposed framework in relation with the state-of-the-art in dimensionality reduction. We provide details of the probabilistic model proposed in the context of Bayesian State-Space models in Section 2.2. Section 2.3 is devoted to inference and learning tasks which involve a locally-optimal Sequential Monte Carlo sampler and an online Expectation-Maximization scheme. The utilization of the coarse-grained dynamics in the context of a Bayesian (adaptive) time-integration scheme is discussed in 2.4 and numerical examples are provided in Section 3.

2. Proposed Approach.

2.1. From static-linear to dynamic-nonlinear dimensionality reduction.

The inherent assumption of all multiscale analysis methods is the existence of a lower-dimensional parameterization of the original system with respect to which the dynamical evolution is more tractable at the scales of interest. In some cases these slow variables can be identified a priori and the problem reduces to finding the necessary closures that will give rise to a consistent dynamical model. In general however one must identify the reduced space \( \hat{Y} \) as well as the dynamics within it.

A prominent role in these efforts has been held by Principal Component Analysis (PCA) -based methods. With small differences and depending on the community other terms such as Proper Orthogonal Decomposition (POD) or Karhunen-Loève expansion (KL), Empirical Orthogonal Functions (EOF) have also been used. PCA finds its roots in the early papers by Pearson \[111\] and Hotelling \[84\] and was originally developed as a static dimensionality reduction technique. It is based on projections on a reduced basis identified by the leading eigenvectors of the covariance matrix \( C \). In the dynamic case and in the absence of closed form expressions for the actual covariance matrix, samples of the process \( y_t \in \mathbb{R}^d \) at \( N \) distinct time instants \( t_i \) are used in order to obtain an estimate of the covariance matrix:

\[
C \approx C_N = \frac{1}{N-1} \sum_{i=1}^{N} (y_{t_i} - \mu)(y_{t_i} - \mu)^T \tag{2.1}
\]

where \( \mu = \frac{1}{N} \sum_{i=1}^{N} y_{t_i} \) is the empirical mean. If there is a spectral gap after the first \( k \) eigenvalues and \( V_K \) is the \( d \times K \) matrix whose columns are the \( K \) leading normalized eigenvectors of \( C_N \) then the reduced-order model is defined with respect to \( \hat{y}_t = V_K y_t \). The reduced dynamics can be identified by a Galerkin projection (or a Petrov-Galerkin projection) of the original ODEs in Equation (1.1):

\[
\frac{d\hat{y}_t}{dt} = V_K^T f(V_K^T \hat{y}_t) \tag{2.2}
\]
Bayesian reduced-order models

Fig. 2.1. The phase space is assumed two-dimensional for illustration purposes i.e. \( y_t = (y_{t(1)}, y_{t(2)}) \). Each black circle corresponds to a realization \( y_{ti} \). \( P : Y \to \hat{Y} \) is the projection operator from the original high-dimensional space \( Y \) to the reduced-space \( \hat{Y} \).

Hence the reduced space \( \hat{Y} \) is approximated by a hyperplane in \( Y \) and the projection mapping \( P \) linear (Figure 2.1(a)). While it can be readily shown that the projection adopted is optimal in the mean square sense for stationary Gaussian processes, it is generally not so in cases where non-Gaussian processes or other distortion metrics are examined. The application of PCA-based techniques, to high-dimensional, multiscale dynamical systems poses several modeling limitations. Firstly, the reduced space \( \hat{Y} \) might not be sufficiently approximated by a hyperplane of dimension \( K << d \). Even though this assumption might hold locally, it is unlikely that this will be a good global approximation. Alternatively, the dynamics of the original process might be adequately approximated on \( K \)-dimensional hyperplane but this hyperplane might change in time. Secondly, despite the fact that the projection on the subspace spanned by the leading eigenvectors captures most of the variance of the original process, in cases where this variability is due to the fast modes, there is no guarantee that \( \hat{y}_t \) will account for the long-range, slow dynamics which is of primary interest in multiscale systems. Thirdly, the basic assumption in the estimation of the covariance matrix, is that the samples \( y_{ti} \) are drawn from the same distribution, i.e. that the process \( y_t \) is stationary. A lot of multiscale problems however involve non-stationary dynamics (e.g. non-equilibrium MD [78, 30]). Hence even if a stationary reduced-order process provides a good, local, approximation to \( y_t \), it might need to be updated in time. Apart from the aforementioned modeling issues, significant computational difficulties are encountered for high-dimensional systems (\( d = \text{dim}(Y) >> 1 \)) and large datasets (\( N >> 1 \)) as the \( K \) leading eigenvectors of large matrices (of dimension proportional to \( d \) or \( N \)) need to be evaluated. This effort must be repeated, if more samples become available (i.e. \( N \) increases) and an update of the reduced-order model is desirable. Recent efforts have concentrated on developing online versions [137] that circumvent this problem.

The obvious extension to the linear projections of PCA is nonlinear dimensionality reduction techniques. These have been the subject of intense research in statistics and machine learning in recent years (121 119 130 144 123 10 9) and fairly recently have found their way to computational physics and multiscale dynamical systems (e.g. 82 96 107 59). They are generally based on calculating eigenvectors of an affinity matrix of a weighted graph. While they circumvent the limiting, linearity assumption of standard PCA, they still assume that the underlying process is stationary (Figure 2.1(b)). Even though the system's dynamics might be appropriately tracked on a
lower-dimensional subspace for a certain time period, this might not be invariant across the whole time range of interest. The identification of the dynamics on the reduced-space $\hat{Y}$ is not as straightforward as in standard PCA and in most cases, a deterministic or stochastic model is fit directly to the projected data points \cite{31, 50, 58}. More importantly since the inverse mapping $P^{-1}$ from the manifold $\hat{Y}$ to $Y$ is not available analytically, approximations have to be made in order to find pre-images in the data-space \cite{11, 50}. From a computational point of view, the cost of identifying the projection mapping is comparable to standard PCA as an eigenvalue problem on a $N \times N$ matrix has to be solved. Updating those eigenvalues and the nonlinear projection operator in cases where additional data become available implies a significant computational overhead although recent efforts \cite{122} attempt to overcome this limitation.

A common characteristic of the aforementioned techniques is that even though the reduced coordinates are learned from a finite amount of simulation data, there is no quantification of the uncertainty associated with these inferences. This is a critical component not only in cases where multiple sets of reduced parameters and coarse-grained models are consistent with the data, but also for assessing errors associated with the analysis and prediction estimates. It is one of the main motivations for adopting a probabilistic approach in this project. Statistical models can naturally deal with stochastic systems that frequently arise in a lot of applications. Most importantly perhaps, even in cases where the fine-scale model is deterministic (e.g. Equation (1.1)), a stochastic reduced model provides a better approximation that can simultaneously quantify the uncertainty arising from the information loss that takes place during the coarse-graining process \cite{52, 95}.

A more general perspective is offered by latent variable models where the observed data (experimental or computationally generated) is augmented by a set of hidden variables \cite{13}. In the case of high-dimensional, multiscale dynamical systems, the latent model corresponds to a reduced-order process that evolves at scales of practical relevance. Complex distributions over the observables can be expressed in terms of simpler and tractable joint distributions over the expanded variable space. Furthermore, structural characteristics of the original, high-dimensional process $y_t$ can be revealed by interpreting the latent variables as generators of the observables.

In that respect, a general setting is offered by Hidden Markov Models (HMM, \cite{64}) or more generally State-Space Models (SSM) \cite{18, 65, 80}. These assume the existence of an unobserved (latent) process $\hat{y}_t$ described by a (stochastic) ODE:

$$\frac{d\hat{y}_t}{dt} = \hat{f}(\hat{y}_t; w_t) \quad \text{(transition equation)} \quad (2.3)$$

which gives rise to the observables $y_t \in \mathbb{R}^d$ as:

$$y_t = h(\hat{y}_t, v_t) \quad \text{(emission equation)} \quad (2.4)$$

where $w_t$ and $v_t$ are unknown stochastic processes (to be inferred from data) and $\hat{f}: \mathbb{R}^K \rightarrow \mathbb{R}^K, \ h: \mathbb{R}^K \rightarrow \mathbb{R}^d$ are unknown measurable functions. The transition equation defines a prior distribution on the coarse-grained dynamics whereas the emission equation, the mapping that connects the reduced-order representation with the observable dynamics. The object of Bayesian inference is to learn the unobserved (unknown) model parameters from the observed data. Hence the coarse-grained model and its relation to the observable dynamics are inferred from the data.
The form of Equations (2.3) and (2.4) affords general representations. Linear and nonlinear PCA models arise as special cases by appropriate selection of the functions and random processes appearing in the transition and emission equations. Note for example that the transition equation (Equation (2.3)) for $\hat{y}_t$ in the case of the PCA-based models reviewed earlier is given by Equation (2.2) and the emission equation (Equation (2.4)) that relates latent and observed processes is linear, deterministic and specified by the matrix of $K$ leading eigenvectors $V_K$.

An extension to HMM is offered by switching-state models \[74, 24, 72, 124\] which can be thought of as dynamical mixture models \[23, 66\]. The latent dynamics consist of a discrete process that takes $M$ values, each corresponding to a distinct dynamical behavior. This can be represented by an $M$-dimensional vector $z_t$ whose entries are zero except for a single one $m$ which is equal to one and represents the active mode/cluster. Most commonly, the time-evolution of $z_t$ is modeled by a first-order stationary Markov process:

$$z_{t+1} = T z_t$$

where $T = [T_{m,n}]$ is the transition matrix and $T_{m,n} = Pr[z_{m,t+1} = 1 | z_{n,t} = 1]$. In addition to $z_t$, $M$ processes $x^{(m)}_t \in \mathbb{R}^K$, $m = 1, \ldots, M$ parameterize the reduced-order dynamics (see also discussion in section 2.2). Each is activated when $z_{m,t} = 1$. In the linear version (Switching Linear Dynamic System, SLDS \[1\]) and conditioned on $z_{m,t} = 1$, the observables $y_t$ arise by a projection from the active $x^{(m)}_t$ as follows:

$$y_t = P^{(m)} x^{(m)}_t + v_t, \quad v_t \sim N(0, \Sigma) \quad (i.i.d)$$

where $P^{(m)}$ are $d \times K$ matrices $(K << d)$ and $\Sigma$ is a positive definite $d \times d$ matrix. Such models provide a natural, physical interpretation according to which the behavior of the original process $y_t$ is segmented into $M$ regimes or clusters, the dynamics of which can be low-dimensional and tractable. From a modeling point of view, the idea of utilizing a mixture of simple models provides great flexibility \[16, 14, 125, 70\] as it can be theorized that given a large enough number of such components, any type of dynamics can be sufficiently approximated. In practice however, a large number might be needed, resulting in complex models containing a large number of parameters.

Such mixture models have gained prominence in recent years in the machine learning community. In \[15\] for example, a dynamic mixture model was used to analyze a huge number of time series, each corresponding to a word in the English vocabulary as they appear in papers in the journal Science. The latent discrete variables represented topics and each topic implied a distribution on the space of words. As a result, not only a predictive summary (dimensionality reduction) of the high-dimensional observables was achieved but also an insightful deconstruction of the original time series was made possible. In fact current research in statistics has focused on infinite mixture models where the number of components can be automatically inferred from the data \[129, 21, 12, 56, 57\]. In the context of computer simulations of high-dimensional systems, such models have been employed by \[55, 82, 80, 79, 81\] where maximum likelihood techniques were used to learn the model parameters.

In the next sections we present a novel model that generalizes SLDS. Unlike mixture models which assume that $y_t$ is the result of a single reduced-order process at a time, we propose a partial-membership model (referred to henceforth as

\[1\] sometimes referred to as jump-linear or conditional Gaussian models
Partial-Membership Linear Dynamic System (PMLDS) which allows observables to have fractional memberships in multiple clusters. The latent, building blocks are experts [86, 87, 77, 75] which, on their own, provide an incomplete, biased prediction but when their “opinions” are appropriately synthesized, they can give rise to a highly accurate aggregate model.

From a modeling perspective such an approach has several appealing properties. The integrated coarse-grained model can be interpretable and low-dimensional even for large, multiscale systems as its expressive ability does not hinge upon the complexity of the individual components but rather is a result of its factorial character (67). Intricate dynamical behavior can be captured and decomposed in terms of simple building blocks. It is highly-suited for problems that lack scale separation and where the evolution of the system is the result of phenomena at a cascade of scales. Each of these scales can be described by a latent process and the resulting coarse-grained model will account not only for the slow dynamics but also quantify the predictive uncertainty due to the condensed, fast-varying features.

From an algorithmic point of view we present parallelizable, online inference/learning schemes, which can recursively update the estimates produced as more data become available i.e. if the time horizon \( t \) of the observables \( y_{1:t} \) increases. Unlike some statistical applications where long time series are readily available, in the majority of problems involving computational simulations of high-dimensional, multiscale systems, data is expensive (at least over large time horizons) as they imply calls to the microscopic solvers. The algorithms presented are capable of producing predictive estimates “on the fly” and if additional data is incorporated, they can readily update the model parameters. In addition, such schemes can take advantage of the natural tempering effect of introducing the data sequentially which can further facilitate the solution of the global estimation problem. More importantly perhaps, the updating schemes discussed have linear complexity with respect to the dimensionality \( d \) of the original process \( y_t \).

2.2. Partial-Membership Linear Dynamic System. We present a hierarchical Bayesian framework which promotes sparsity, interpretability and efficiency. The framework described can integrate heterogeneous building blocks and allows for physical insight to be introduced on a case-by-case basis. When dealing with high-dimensional molecular ensembles for example, each of these building blocks might be an (overdamped) Langevin equation with a harmonic potential [17, 80, 83]. It is obvious that such a simplified model would perhaps provide a good approximation under specific, limiting conditions (e.g. at a persistent metastable state) but definitely not across the whole time range of interest. Due to its simple parameterization and computational tractability, it can easily be trained to represent one of the “experts” in the integrated reduced-model. It is known that these models work well under specific regimes but none of them gives an accurate global description. In the framework proposed, they can however be utilized in a way that combines their strengths, but also probabilistically quantifies their limitations.

A transient, nonlinear PDE can be resolved into several linear PDEs whose simpler form and parameterization makes them computational tractable over macroscopic time scales and permits a coarser spatial discretization. Their combination with time-varying characteristics can give rise to an accurate global approximation. Their simplicity and limited range of applicability would preclude their individual use. In the framework proposed however, these simple models would only serve as good local approximants and their inaccurate predictions would be synthesized into an accurate,
and\n\textbf{ics} isotropic Ornstein-Uhlenbeck (OU) processes are used to model the hidden dynam-
particular, the following parameterization is employed:

This equation essentially implies a prior distribution on the space of hidden processes parameterized by a set of (unknown a priori) parameters $\theta^{(m)}$. It should be noted that while the proposed framework allows for any type of process in Equation (2.7), it is desirable that these are simple, in the sense that the parameters are low-dimensional and can be learned swiftly and efficiently. We list some desirable properties of the prior models [125]:

1. Stationarity: Unless specific prior information is available, it would be unreasonable to impose a time bias on the evolution of any of the reduced dynamics processes. Hence it is important that the models adopted are a priori stationary. Note that the posterior distributions might still exhibit non-stationarity.

2. Correlation Decay: It is easily understood that for any $m$ and $t_1, t_2$, the correlation $x^{(m)}_{t_1}$ and $x^{(m)}_{t_2}$ should decay monotonically as $|t_2 - t_1|$ goes to $+\infty$. This precludes models that do not explicitly account for the time evolution of the latent processes and assume that hidden states are not time-dependent (e.g. static PCA models).

3. Other: Although this is not necessary, we adopt a continuous time model in the sense of [135] with an analytically available transition density which allows inference to be carried out seamlessly even in cases where the observables are obtained at non-equidistant times. As a result the proposed framework can adapt to the granularity of the observables and also provide exact probabilistic predictions at any time resolution.

Although more complex models can be adopted we assume here that independent, isotropic Ornstein-Uhlenbeck (OU) processes are used to model the hidden dynamics $x^{(m)}_t$. The OU processes used comply with the aforementioned desiderata. In particular, the following parameterization is employed:

$$
\frac{dx^{(m)}_t}{dt} = g_m(x^{(m)}_t; \theta^{(m)}_x), \quad m = 1, \ldots, M
$$

This equation essentially implies a prior distribution on the space of hidden processes parameterized by a set of (unknown a priori) parameters $\theta^{(m)}_x$. It should be noted that while the proposed framework allows for any type of process in Equation (2.7), it is desirable that these are simple, in the sense that the parameters $\theta^{(m)}_x$ are low-dimensional and can be learned swiftly and efficiently. We list some desirable properties of the prior models [125]:

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$$
\frac{dx^{(m)}_t}{dt} = -b^{(m)}_x(x^{(m)}_t - q^{(m)}_x)dt + (S^{(m)}_x)_{1/2} dW^{(m)}_t
$$

where $W^{(m)}_t$ are Wiener processes (independent for each $m$), $b^{(m)}_x > 0, q^{(m)}_x \in \mathbb{R}^K$ and $S^{(m)}_x$ are positive definite matrices of dimension $K \times K$. The aforementioned

![Fig. 2.2. Realizations of two hidden (M = 2) one-dimensional (K = 1) Ornstein-Uhlenbeck processes were used: $dx^{(m)}_t = -b^{(m)}_x(x^{(m)}_t - q^{(m)}_x)dt + (S^{(m)}_x)_{1/2} dW^{(m)}_t$, with $b^{(1)}, q^{(1)}_x, S^{(1)} = (1, 3, 2)$ (fast) and $b^{(2)}, q^{(2)}_x, S^{(2)} = (0.01, -3, 0.02)$ (slow).]
FIG. 2.3. The logistic normal distribution was used to model the weights associated with each of the hidden processes depicted in Figure 2.2. In particular, an isotropic Ornstein-Uhlenbeck process \( \frac{d\hat{z}_t}{dt} = -b_{\hat{z}}(\hat{z}_t - q_{\hat{z}})dt + \Sigma_{\hat{z}}^{1/2}dW_t \) with \( b_{\hat{z}} = 0.001, q_{\hat{z}} = [0, 0]^T \) and \( \Sigma_{\hat{z}} \). Graphs depict the resulting observable time history (left column) and its histogram (right column) arising from the unobserved processes in Figure 2.2 and for three values of \( \rho \). It is noted that time histories exhibit fast and slow scales of the processes in Figure 2.2. Furthermore, by changing a single parameter (i.e. \( \rho \)) one can obtain two, three or a single metastable state (right column - peaks of the histogram).

model has a Gaussian invariant (stationary) distribution \( \mathcal{N}(q_x^{(m)}, \frac{1}{2b_{z_x}^2}, S_x^{(m)}) \). The transition density denoted by \( p(x_t^{(m)} | x_{t-1}^{(m)}) \) for time separation \( \delta t \) is also a Gaussian \( \mathcal{N}(\mu_{t,\delta t}, S_{\delta t}) \) where:

\[
\begin{align*}
\mu_{t,\delta t} &= x_{t-1}^{(m)} - (1 - e^{-b_{z_x}^{(m)} \delta t})(x_{t-1}^{(m)} - q_x^{(m)}) \\
S_{\delta t} &= \frac{1 - e^{-2b_{z_x}^{(m)} \delta t}}{2b_{z_x}^{(m)}} S_x^{(m)}
\end{align*}
\]  

(2.9)

It is not expected that simple processes on their own will provide good approximations to the essential dynamics exhibited in the data \( y_t \). In order to combine the dynamics implied by the \( M \) processes in Equation (2.7), we consider an \( M \)-dimensional process \( z_t \) such that \( \sum_{m=1}^{M} z_{m, t} = 1 \) and \( z_{m, t} > 0, \forall t \) and define an appropriate prior. The coefficients \( z_{m, t} \) express the weight or fractional membership to each process/expert \( x_t^{(m)} \) at time \( t \). We use the logistic-normal model as a prior for \( z_t \). It is based on a Gaussian process, \( \hat{z}_t \) whose dynamics are also prescribed by an isotropic OU process:

\[
d\hat{z}_t = -b_{z}(\hat{z}_t - q_{z})dt + S_z^{1/2}dW_t
\]  

(2.10)

and the transformation:

\[
z_{m, t} = \frac{e^{z_{m, t}} + 1/\sum_{m=1}^{M} e^{z_{m, t}}}{1/\sum_{m=1}^{M} e^{z_{m, t}} + 1}, \forall m, t
\]  

(2.11)
The invariant and transition densities of $\hat{z}_t$ are obviously identical to the ones for $x_t^{(m)}$ with appropriate substitution of the parameters. The hidden processes $\{x_t^{(m)}\}_{m=1}^M$ and associated weights $z_t$ give rise to the observables $y_t$ as follows (compare with Equation (2.10)):

$$y_t = \sum_{m=1}^M z_{m,t} \ P^{(m)} x_t^{(m)} + v_t, \quad v_t \sim \mathcal{N}(0, \Sigma) \quad (i.i.d)$$

(2.12)

where $P^{(m)}$ are $d \times K$ matrices ($K << d$) and $\Sigma$ is a positive definite $d \times d$ matrix. The aforementioned equation implies a series of linear projections on hyperplanes of dimension $K$. The dynamics along those hyperplanes are dictated by a priori independent process $x_t^{(m)}$. It is noted however the reduced dynamics are simultaneously described by all the hidden processes (Figure 2.3). This is in contrast to PCA methods where a single such projection is considered and mixture PCA models where even though several hidden processes are used, at each time instant it is assumed that a single one is active. Due to the factorial nature of the proposed model, multiple dynamic regimes can be captured by appropriately combining a few latent states. While mixture models (Figure 2.1(c)) provide a flexible framework, the number of hidden states might be impractically large. As it is pointed out in [67], in order to encode for example a time sequence with 30 bits of information one would need $k = 2^{30}$ distinct states. It is noted that even though linear projections are implied by Equation (2.12) and Gaussian noise $v_t$ is used, the resulting model for $y_t$ is nonlinear and non-Gaussian as it involves the factorial combination of $M$ processes $\{x_t^{(m)}\}_{m=1}^M$ with $z_t$ which are a posteriori non-Gaussian.

The parameters $z_{m,t}$ express the relative importance of the various reduced models or equivalently the degree to which each data point $y_t$ is associated with each of the $M$ reduced dynamics $x_t^{(m)}$. It is important to note that the proposed model allows for time varying weights $z_{m,t}$ and can therefore account for the possibility of switching between different regimes of dynamics. Figure 2.3 depicts a simple example ($d = 1$) which illustrates the flexibility of the proposed approach.

The unknown parameters of the coarse-grained model consist of:

- **dynamic variables** denoted for notational economy by $\Theta_t$ (i.e. $\{x_t^{(m)}\}_{m=1}^M$, $z_t$ for $t = 1, 2, \ldots$).

- **static variables** denoted by $\Theta$ (i.e. $\theta_1^{(m)} = (b_z^{(m)}, q_z^{(m)}, S_z^{(m)})$ in Equation (2.8), $\theta_z = (b_z, q_z, S_z)$ in Equation (2.10) and $\{P^{(m)}\}_{m=1}^M, \Sigma$ in Equation (2.12)).

2.3. Inference and learning. Inference in the probabilistic graphical model described involves determining the probability distributions associated with the unobserved (hidden) static $\Theta$ and dynamic parameters $\Theta_t$ of the model. In the Bayesian setting adopted this is the posterior distribution of the unknown parameters of the coarse-grained model. Given the observations (computational or experimental) of the original, high-dimensional process $y_{1:}\pi = \{y_t\}_{t=1}^T$, we denote the posterior by $p(\Theta, \Theta_{1:T})$:

$$p(\Theta, \Theta_{1:T} | y_{1:T}) = \frac{p(y_{1:T} | \Theta, \Theta_{1:T}) \ p(\Theta_{1:T})}{p(y_{1:T})}$$

(2.13)
The normalization constant \( p(y_{1:\tau}) \) is not of interest when sampling for \( \Theta \) or \( \Theta_{1:\tau} \) but can be quite useful for model validation purposes.

A telescopic decomposition holds for the likelihood which according to Equation (2.12) is given by:

\[
p(y_{1:\tau} | \Theta, \Theta_{1:\tau}) = \prod_{t=1}^{\tau} p(y_t | \Theta, \Theta_t) \tag{2.14}
\]

where the densities in the product are described in Equation (2.17). Equation (2.12) defines the likelihood \( p(y_t | \Theta, \Theta_t) \) which is basically the weighted product of the likelihoods under each of the hidden processes/experts \( x_t^{(m)} \):

\[
p(y_t | \Theta, \Theta_t) \propto \frac{1}{c(\Theta, \Theta_t)} \prod_{m=1}^{M} p_{m}^{z_{m,t}}(y_t | \Theta, \Theta_t) \tag{2.15}
\]

where the normalizing constant \( c(\Theta, \Theta_t) \) ensures that the density integrates to one with respect to \( y_t \). According to Equation (2.12):

\[
p_m(y_t | \Theta, \Theta_t) \propto \frac{1}{|\Sigma|^{1/2}} \exp \left\{ -\frac{1}{2} (y_t - P^{(m)} x_t^{(m)})^T \Sigma^{-1} (y_t - P^{(m)} x_t^{(m)}) \right\} \tag{2.16}
\]

The likelihood can be written in a more compact form as:

\[
p(y_t | \Theta, \Theta_t) \propto \frac{1}{|\Sigma|^{1/2}} \exp \left\{ -\frac{1}{2} (y_t - W_t X_t)^T \Sigma^{-1} (y_t - W_t X_t) \right\} \tag{2.17}
\]

where:

\[
X_t^T = \left[ (x_t^{(1)})^T, (x_t^{(2)})^T, \ldots, (x_t^{(M)})^T \right]^T \tag{2.18}
\]

and:

\[
W_t = \begin{bmatrix} z_{1,t} P^{(1)} & z_{2,t} P^{(2)} & \cdots & z_{M,t} P^{(M)} \end{bmatrix} \tag{2.19}
\]

The first-order Markovian processes adopted for the prior modeling of the dynamic parameters \( \Theta_t \) (Equations (2.8), (2.10), 2.9) imply that:

\[
p(\Theta_{1:\tau}, \Theta) = p(\Theta) \prod_{t=1}^{\tau} p(\Theta_t | \Theta_{t-1}, \Theta) \tag{2.20}
\]
where \( p(\Theta_1 \mid \Theta_0, \Theta) = p(\Theta_1 \mid \Theta) = \nu_0(\Theta_1 \mid \Theta) \) is the prior on the initial condition which in this work is taken to be the stationary distribution of the underlying OU processes (see discussion in Section 2.2.1) and denoted for notational economy by \( \nu_0(\cdot \mid \Theta) \).

The posterior encapsulates uncertainties arising from the potentially stochastic nature of the original process \( y_t \) as well as due to the fact that a finite number of observations were used. The difficulty of the problem is that both the dynamic (\( \Theta_{1:t} \)) and the static parameters (\( \Theta \)) are unknown. We adopt a hybrid strategy whereby we sample from the full posterior for the dynamic parameters \( \Theta_t \) and provide point estimates for the static parameters \( \Theta \). If uniform priors are used for \( \Theta \) then the procedure proposed reduces to a maximum likelihood estimation. Non-uniform priors have a regularization effect which can promote the identification of particular features.

While the hybrid strategy proposed is common practice in pertinent models (64), in the current framework it is also necessitated by the difficulty in sampling in the high-dimensional state space of \( \Theta \) (note that the projection matrices \( P^{(m)} \) in particular are of the dimension of the observables \( d \) and \( d \gg 1 \)) as well as the need for scalability in the context of high-dimensional systems. The static parameters \( \Theta \) are estimated by maximizing the log-posterior.

\[
L(\Theta) = \log \pi(\Theta \mid y_{1:t}) = \log \int \pi(\Theta, \Theta_{1:t} \mid y_{1:t}) \, d\Theta_{1:t} \tag{2.21}
\]

Maximization of \( L(\Theta) \) is more complex than a standard optimization task as it involves integration over the unobserved dynamic variables \( \Theta_{1:t} \). While maximization can be accelerated by using gradient-based techniques (e.g. gradient ascent), the dimensionality of \( \Theta \) makes such an approach impractical as it can be extremely difficult to scale the parameter increments. We propose therefore adopting an Expectation-Maximization framework (EM) which is an iterative, robust scheme that is guaranteed to increase the log-posterior at each iteration (41, 64). It is based on constructing a series of increasing lower bounds of the log-posterior using auxiliary distributions \( q(\Theta_{1:t}) \):

\[
L(\Theta) = \log \pi(\Theta \mid y_{1:t}) = \log \int q(\Theta_{1:t}) \pi(\Theta, \Theta_{1:t} \mid y_{1:t}) \, d\Theta_{1:t} = \log \int q(\Theta_{1:t}) \frac{\pi(\Theta, \Theta_{1:t} \mid y_{1:t})}{\pi(\Theta_{1:t} \mid y_{1:t})} \, d\Theta_{1:t} \geq \int q(\Theta_{1:t}) \log \frac{\pi(\Theta, \Theta_{1:t} \mid y_{1:t})}{q(\Theta_{1:t} \mid y_{1:t})} \, d\Theta_{1:t} \tag{2.22}
\]

It is obvious that this inequality becomes an equality when in place of the auxiliary distribution \( q(\Theta_{1:t}) \) the posterior \( \pi(\Theta_{1:t} \mid \Theta, y_{1:t}) \) is selected. Given an estimate \( \Theta^{(s)} \) at step \( s \), this suggests iterating between an Expectation step (E-step) whereby we average with respect to \( q^{(s)}(\Theta_{1:t}) = \pi(\Theta_{1:t} \mid \Theta^{(s)}, y_{1:t}) \) to evaluate the lower bound:

\[
\text{E-step:} \quad F^{(s)}(q^{(s)}, \Theta) = \int q^{(s)}(\Theta_{1:t}) \log \pi(\Theta, \Theta_{1:t} \mid y_{1:t}) \, d\Theta_{1:t} - \int q^{(s)}(\Theta_{1:t}) \log q^{(s)}(\Theta_{1:t}) \, d\Theta_{1:t} \tag{2.23}
\]

and a Maximization step (M-step) with respect to \( F^{(s)}(q^{(s)}, \Theta) \) (and in particular the
first part in Equation (2.23) since the second does not depend on $\Theta$:

\begin{equation}
M\text{-step: } \Theta^{(s+1)} = \arg \max_{\Theta} F^{(s)}(q^{(s)}, \Theta) \\
= \arg \max_{\Theta} E_q^{(s)}(\Theta_{1:\tau}) \left[ \log \pi(\Theta, \Theta_{1:\tau} | y_{1:\tau}) \right] \\
= \arg \max_{\Theta} Q(\Theta^{(s)}, \Theta) \tag{2.24}
\end{equation}

As the optimal auxiliary distributions $q^{(s)}(\Theta_{1:\tau}) = \pi(\Theta_{1:\tau} | \Theta^{(s)}, y_{1:\tau})$ are intractable, we propose employing a Sequential Monte Carlo (SMC or particle filter, [45, 37]) scheme for estimating the expectations in the M-Step, i.e. Equation (2.24). SMC samplers provide a parallelizable framework for non-linear, non-Gaussian filtering problems whereby the target distribution $q^{(s)}(\Theta_{1:\tau}) = \pi(\Theta_{1:\tau} | \Theta^{(s)}, y_{1:\tau})$ is represented with a population of $N$ particles $\Theta_{1:\tau}^{(s,i)}$ and weights $W^{(s,i)}$ such that the expectation in Equation (2.24) can be approximated by:

\begin{equation}
E_q^{(s)}(\Theta_{1:\tau}) \left[ \log \pi(\Theta, \Theta_{1:\tau} | y_{1:\tau}) \right] \approx \sum_{i=1}^{N} W^{(s,i)} \log \pi(\Theta, \Theta_{1:\tau}^{(s,i)}) | y_{1:\tau} \tag{2.25}
\end{equation}

In section 2.3.1 we discuss a particle filter that takes advantage of the particular structure of the posterior and employs the locally optimal importance sampling distribution. Nevertheless, SMC samplers involve sequential importance sampling, and their performance decays with increasing $\tau$ as the dimension of the state space $\Theta_{1:\tau}$ increases even when resampling and rejuvenation mechanisms are employed ([7]). Recent efforts based on exponential forgetting have shown that the accuracy of the approximation can be improved (while keeping the number of particles $N$ fixed) by employing smoothing ([24]) over a fixed-lag in the past ([24]).

In this paper we make use of an approximate but highly efficient alternative proposed in [6, 7, 8]. This is based on the so-called split-data likelihood (SDL) first discussed in [110], which consists of splitting the observations into blocks (overlapping or non-overlapping) of length $L < \tau$ and using the pseudo-likelihood which arises by assuming that these blocks are independent. It is shown in [7] that this leads to an alternative Kullback-Leibler divergence contrast function and under some regularity conditions that the set of parameters optimizing this contrast function includes the true parameter. Because the size of the blocks is fixed, the degeneracy of particle filters can be averted and the quality of the approximations can be further improved by applying a backward smoothing step over each block ([79]). Let $k$ denote the index of the block of length $L$ considered and $\bar{y}_k = y_{|(k-1)L+1:kL}$ and $\bar{\Theta}_k = \Theta_{|(k-1)L+1:kL}$. If $\tau = r\ L$, the likelihood is approximated by:

\begin{equation}
p(y_{1:\tau} | \Theta, \Theta_{1:\tau}) \approx \prod_{k=1}^{r} p(\bar{y}_k | \Theta, \bar{\Theta}_k) \tag{2.26}
\end{equation}

When $\Theta_t$ has reached a stationary regime with invariant density $\nu_0(\cdot | \Theta)$, then for any $k$, $(\Theta, \bar{\Theta}_k, \bar{y}_k)$ are identically distributed according to:

\begin{equation}
\bar{p}(\Theta, \bar{\Theta}_k, \bar{y}_k) = \frac{\pi(\Theta)\nu_0(\Theta_{(k-1)L} | \Theta)p(y_{(k-1)L} | \Theta_{(k-1)L}, \Theta)}{\prod_{k=1}^{r} L} p(\Theta_t | \Theta_{t-1}, \Theta)p(y_t | \Theta_t, \Theta) \tag{2.27}
\end{equation}

In a batch EM algorithm using the split-data likelihood and the $k^{th}$ block of data, the M-step would involve maximization with respect to $\Theta$ (see also Equation
Note that in order to simplify the notation we consider an arbitrary block of length \(2.31\) with respect to the density \(\bar{p}\) we discuss Monte Carlo approximations of the expectations appearing in Equation (119) while the M-step is left unchanged. In particular, at iteration \(k \equiv s\):

\[
\text{online E-step } \bar{Q}((\theta^{(1:k-1)}), \Theta) = (1 - \gamma_k)\bar{Q}((\theta^{(1:k-2)}), \Theta) + \gamma_k \int \bar{p}(\Theta_k | \Theta^{(k-1)}, \bar{y}_k) \log \bar{p}(\Theta, \bar{y}_k) d\Theta_k
\]  

and update the value of the parameters \(\Theta\) as:

\[
\text{online M-step } \Theta^{(k)} = \arg \max_\Theta \bar{Q}((\theta^{(1:k-1)}), \Theta)
\]  

The algorithm relies on a non-increasing sequence of positive stepsizes \(\{\gamma_k\}_{k \geq 0}\) such that \(\sum_k \gamma_k = +\infty\) and \(\sum_k \gamma_k^2 < +\infty\). In this work we adopted \(\gamma_k = \frac{1}{k^a}\) with \(a = 0.51\). Naturally the integrals above over the hidden dynamic variables \(\Theta_k\) are estimated using SMC-based, particulate approximations of \(\bar{p}(\Theta_k | \Theta^{(k-1)}, \bar{y}_k)\). For small \(L\) the convergence will in general be slow as the split-block likelihood assumption will be further from the truth. For larger \(L\), convergence is faster but the performance of the filter decays. For that purpose we also employed a backward smoothing filter over each block using the algorithm described in [69]. The computational cost of the smoothing algorithm is \(O(N^2L)\).

In practice, and in particular for the exponential distributions utilized in the proposed model (e.g. Equation (2.17)), the EM iterations reduce to calculating a set of (multivariate) sufficient statistics \(\Phi\). In particular, instead of the log-posterior lower bound \(\bar{Q}((\theta^{(1:k-1)}), \Theta)\) in Equation (2.29) we update the sufficient statistics as follows:

\[
\Phi^{(k)} = (1 - \gamma_k)\Phi^{(k-1)} + \gamma_k \int \bar{p}(\Theta_k | \Theta^{(k-1)}, \bar{y}_k) \phi(\Theta_k) d\Theta_k
\]  

where \(\int \bar{p}(\Theta_k | \Theta^{(k-1)}, \bar{y}_k) \phi(\Theta_k) d\Theta_k\) denotes the set of sufficient statistics associated with the block of data \(\bar{y} = y^{(k-1)L+1:kL}\). Specific details are provided in the Appendix. It is finally noted, that learning tasks in the context of the probabilistic model proposed, should also involve identifying the correct structure (e.g. the number of different experts \(M\)). While this problem poses some very challenging issues which are currently the topic of active research in various contexts (e.g. nonparametric methods), this paper is exclusively concerned with parameter learning. In section 3 we discuss Bayesian validation techniques for assessing quantitatively the correct model structure which are computationally feasible due to the efficiency of the proposed algorithms.

2.3.1. Locally optimal Sequential Monte Carlo samplers. In this section we discuss Monte Carlo approximations of the expectations appearing in Equation (2.31) with respect to the density \(\bar{p}(\Theta_k | \Theta, y_{1:L}) = p(\Theta_{(k-1)L+1:kL} | \Theta, y_{(k-1)L+1:kL})\). Note that in order to simplify the notation we consider an arbitrary block of length \(L\)
(e.g. $k = 1$) and do not explicitly indicate the iteration number of the EM algorithm. Hence the target density is:

$$p(\Theta_{1:L} \mid \Theta, y_{1:L}) = \frac{1}{p(y_{1:L} \mid \Theta)} \nu_0(\Theta_1 \mid \Theta)p(y_1 \mid \Theta_1, \Theta) \prod_{t=2}^L p(\Theta_t \mid \Theta_{t-1}, \Theta)p(y_t \mid \Theta_t, \Theta)$$  

(2.32)

where the dynamic variables are $\Theta_t = (X_t, z_t)$ (Equation 2.18). Based on earlier discussions, the evolution dynamics of these variables are independent i.e.:

$$\nu_0(\Theta_1 \mid \Theta) = \nu_0(X_1 \mid \Theta)\nu_0(z_1 \mid \Theta)$$  

(2.33)

and:

$$p(\Theta_t \mid \Theta_{t-1}, \Theta) = p(X_t \mid X_{t-1}, \Theta)p(z_t \mid z_{t-1}, \Theta)$$  

(2.34)

Since there is a deterministic relation between $\bar{z}_t$ and $z_t$ (Equation 2.11) we use them interchangeably. In particular we use $\bar{z}_t$ in the evolution equations since the initial and transition densities are Gaussian (Equation 2.11) and $z_t$ in the likelihood densities as the expressions simplify in Equation 2.12. The initial and transition densities for $X_t$ are also Gaussian. Given that $x_t^{(m)}$ are a priori independent, we have that:

$$p(X_t \mid X_{t-1}, \Theta) = \prod_{m=1}^M p(x_t^{(m)} \mid x_{t-1}^{(m)}, \Theta) = \mathcal{N}(X_t \mid \mu_t, S_X)$$  

(2.35)

where the mean $\mu_t = \mu_t(X_{t-1})$ is given by Equation 2.18 and $S_X = \text{diag}(S_{x,1}, \ldots, S_{x,M})$ (from Equation 2.9 as well).

SMC samplers operate on a sequence of target densities $p(\Theta_{1:t} \mid y_{1:t}, \Theta)$ which, for any $t$, are approximated by a set of $n$ random samples (or particles) $\{\Theta_{1:t}^{(i)}\}_{i=1}^n$. These are propagated forward in time using a combination of importance sampling, resampling and MCMC-based rejuvenation mechanisms [38, 37, 138, 133]. Each of these particles is associated with an importance weight $W_t^{(i)} (\sum_{i=1}^n W_t^{(i)} = 1)$ which is updated sequentially along with the particle locations in order to provide a particulate approximation:

$$p(\Theta_{1:t} \mid y_{1:t}, \Theta) \approx \sum_{i=1}^n W_t^{(i)} \delta_{\Theta_{1:t}^{(i)}}(\Theta_{1:t})$$  

(2.36)

where $\delta_{\Theta_{1:t}^{(i)}}(.)$ is the Dirac function centered at $\Theta_{1:t}^{(i)}$. Furthermore, for any measurable $\phi(\Theta_{1:t})$ (as in Equation 2.31) and $\forall t$ [38, 26, 20]:

$$\sum_{i=1}^n W_t^{(i)} g(\Theta_{1:t} \mid y_{1:t}, \Theta) \rightarrow \int \phi(\Theta_{1:t}) p(\Theta_{1:t} \mid y_{1:t}, \Theta)d\Theta_{1:t} \quad \text{(almost surely)}$$  

(2.37)

The particles are constructed recursively in time using a sequence of importance sampling densities $q_t(\Theta_t \mid \Theta_{t-1}, y_t, \Theta_t)$. The importance weights are determined from the fact that:

$$p(\Theta_{1:t} \mid y_{1:t}, \Theta) = p(\Theta_{1:t-1} \mid y_{1:t-1}, \Theta) \frac{p(\Theta_t \mid \Theta_{t-1}, \Theta)p(y_t \mid \Theta_t, \Theta)}{p(y_t \mid y_{1:t-1}, \Theta)}$$  

(2.38)

Let $\{W_t^{(i)}, \Theta_{1:t-1}^{(i)}\}_{i=1}^N$ the particulate approximation of $p(\Theta_{1:t-1} \mid y_{1:t-1}, \Theta)$. Note that for $t = 1$ and for the Gaussian initial densities $\nu_0$ of the proposed model, this consists of exact draws and weights $W_t^{(i)} = 1/N$. At time $t$ we proceed as follows [38]:

$$p(\Theta_{1:t} \mid y_{1:t}, \Theta) \approx \sum_{i=1}^N W_t^{(i)} \delta_{\Theta_{1:t}^{(i)}}(\Theta_{1:t})$$  

(2.39)
This implies using the prior to draw for the future state of the system
\[ \hat{\text{mean}} \bar{\text{mean}} \]
In practice, it is usually impossible to sample from \( q_{\Theta} \) density, at least with respect to the time-varying parameters. Results do not include just point estimates but rather samples from the posterior and propose an importance sampling density of the form:
\[ q_{t}^{\text{opt}}(\Theta_t | \Theta_{t-1}, y_t, \Theta) = \frac{p(\Theta_t | \Theta_{t-1}, \Theta)p(y_t | \Theta_t, \Theta)}{\int p(\Theta_t | \Theta_{t-1}, \Theta)p(y_t | \Theta_t, \Theta) \, d\Theta_t} \quad (2.41) \]
In practice, it is usually impossible to sample from \( q_{t}^{\text{opt}} \) and/or calculate the integral in the denominator. As a result, approximations are used which nevertheless result in non-zero variance in the estimators. In this paper we take advantage of the fact that the transition density of \( X_t \) as well as the likelihood, conditioned on \( z_t \) are Gaussians and propose an importance sampling density of the form:
\[ q_{t}(X_t, \hat{z}_t | X_{t-1}, \hat{z}_{t-1}, y_t, \Theta) = p(\hat{z}_t | \hat{z}_{t-1}, \Theta) \frac{p(X_t | X_{t-1}, \Theta)p(y_t | X_t, z_t, \Theta)}{\int p(X_t | X_{t-1}, \Theta)p(y_t | X_t, z_t, \Theta) \, dX_t} \quad (2.42) \]
This implies using the prior to draw \( \hat{z}_t \) and the locally optimal distribution (conditioned on \( \hat{z}_t \) or equivalently \( z_t \)) for \( X_t \). The latter will also be a Gaussian whose mean \( \bar{\mu}_t \) and covariance \( \bar{S}_X \) can be readily be established (e.g. using Kalman filter formulas):
\[ \bar{S}_X = \left( S_X^{-1} + W_t^T \Sigma^{-1} W_t \right)^{-1} \]
\[ \bar{\mu}_t = S_X \left( S_X^{-1} \mu_t + W_t^T \Sigma^{-1} y_t \right) \quad (2.43) \]
As a result the incremental weights \( u_t \) are given by:
\[ u_t = |\bar{S}_X|^{1/2} \exp \left\{ \frac{1}{2} \mu_t^T \bar{S}_X^{-1} \mu_t - \frac{1}{2} \mu_t^T S_X^{-1} \mu_t \right\} \quad (2.44) \]

2.4. Prediction and Bayesian adaptive time-integration. Bayesian inference results do not include just point estimates but rather samples from the posterior density, at least with respect to the time-varying parameters \( \Theta_t \). The inferred posterior can be readily used to make probabilistic predictions about the future evolution of the high-dimensional, multiscale process \( y_t \). Given observations \( y_{1:t} = \{ y_i \}_{i=1}^t \), the predictive posterior for the future state of the system \( y_{t+1:t+T} \) over a time horizon \( T \)

can be expressed as:

\[
p(y_{\tau+1:T} \mid y_{1:T}) = \int p(y_{\tau+1:T} \mid \Theta_{\tau+1:T}, \Theta_{1:T}) \, d\Theta_{T+1:T} \, d\Theta_{\tau+1:T} \\
= \int p(y_{\tau+1:T} \mid \Theta_{\tau+1:T}) \, p(\Theta_{\tau+1:T} \mid \Theta_{1:T}) \, d\Theta_{T+1:T} \, d\Theta_{\tau+1:T} \tag{2.45}
\]

The integral above can be approximated using Monte Carlo. In particular given the particulate approximation of the posterior \( p(\Theta_{\tau+1:T} \mid y_{1:T}) \) (which consists of samples of the dynamic variables \( \Theta_{\tau} \) and the MAP estimate of \( \Theta_{\tau+1:T} \) \), samples from the prior \( p(\Theta_{\tau+1:T} \mid \Theta_{1:T}, \Theta_{\tau}) \) and subsequently the likelihood \( p(y_{\tau+1:T} \mid \Theta_{\tau+1:T}) \) can readily be drawn. In fact, given that the latter is a multivariate Gaussian, the predictive posterior will consist of a mixture of Gaussians, one for each sample of \( \Theta_{\tau+1:T} \) drawn.

The important consequence of the Bayesian framework advocated is that predictive estimates are not restricted to point estimates but whole distributions which can readily quantify the predictive uncertainty. This naturally gives rise to Bayesian, adaptive, time-integration scheme that allows bridging across timescales while providing quantitative, probabilistic estimates of the accuracy of the coarse-grained dynamics (Figure 2.5). The distribution of Equation (2.45) is used to probabilistically predict the evolution of the system. The time range over which the reduced model is employed does not have to be specified a priori but can be automatically determined by the variance of the predictive posterior (Figure 2.6). Once this exceeds the allowable tolerance specified by the analyst, the fine-scale process is reinitialized and more data are obtained, that can in turn be used to update the coarse-grained model. It is emphasized that due to the generative character of the model proposed, the reinitialization can be performed consistently based on general on the emission equations Equation (2.20).

In contrast to existing techniques such as projective and coarse-projective integration as well as Heterogeneous Multiscale Methods, there is no need to prescribe lifting and restriction operators and no ambiguity exists with regards to the appropriateness of the reinitialization scheme. Furthermore, the probabilistic coarse-grained model provides quantitative estimates for its predictive ability and automatically identifies the need for more information from the fine-scale model.

3. Numerical experiments. The first examples is intended to validate the accuracy of the proposed online EM scheme and utilizes a synthetic dataset. The second example uses actual data and illustrates the superiority of the proposed PMLD model over existing SLDS models. Finally the third example provides an application in the context of multiscale simulations for the time-dependent diffusion equation.

3.1. Synthetic data. We generated data from the proposed model in order to investigate the ability of the inference and learning algorithms discussed. In particular, we considered a mixture of two \( M = 2 \), one-dimensional OU processes \((K = 1)\) as in Equation (2.8) with \((b^{(1)}, q_x^{(1)}, \Sigma^{(1)}) = (0.1, -5.0, 0.2)\) (slow) and \((b^{(2)}, q_x^{(2)}, \Sigma^{(2)}) = (1.0, +5.0, 2.0)\) (fast). The logistic normal distribution was used to model the weights.
Bayesian reduced-order models

prior $p(\Theta, \Theta_1: \tau)$

likelihood $p(y_1: \tau | \Theta, \Theta_1: \tau)$

posterior $p(\Theta, \Theta_1: \tau | y_1: \tau)$

predictive post. $p(y_{\tau+1: \tau+T} | y_1: \tau)$

Bayesian model

data $y_{1: \tau}$

coarse-grained model

fine-scale model

99% post. quantile

1% post. quantile

acceptable predictive uncertainty

consistent reinitialization

Fig. 2.5. Bayesian adaptive time-integration and data-model fusion illustrated for a two-dimensional flow. The data generated from computational simulations $y_{1: \tau}$ and/or experiments $y_{1: \tau'}$ are sequentially incorporated in the Bayesian model and the posterior $p(\Theta, \Theta_1: \tau | y_1: \tau)$ over dynamic and static parameters is updated. The predictive posterior $p(y_{\tau+1: \tau+T} | y_1: \tau)$ is over the time horizon $T$ used to efficiently produce probabilistic predictions of the evolution of the high-dimensional process $y_t$ in the future. When the uncertainty associated with those predictions exceeds the analysts’ tolerance, the original system is consistently reinitialized and more data are generated. These are used to update the (predictive) posterior and to produce additional predictive estimates. It is noted that the tolerance in the predictive uncertainty can also be measured with respect to (low-dimensional) observables which are usually of interest in practical applications.

associated with each of the hidden processes using an isotropic Ornstein-Uhlenbeck process $dz_t = -b_z(z_t - q_z)dt + \Sigma_z^{1/2}dW_t$ with $b_z = 1.0$, $q_z = [0, 0]^T$ and $\Sigma_z = \begin{bmatrix} 10. & 0 \\ 0 & 10. \end{bmatrix}$. Two $10 \times 1$ projection vectors $P^m, m = 1, 2$ were generated from the prior $\mathcal{N}(0, 100I)$ (see Appendix) and $(d = 10)$ time series $y_t$ were produced based on Equation (2.12) with idiosyncratic variances $\Sigma = 0.1^2I$ and time step $\delta t = 1$. The resulting time series exhibit multimodal, non-Gaussian densities as can be seen in Figure 3.1(a) as well as two distinct time scales as it can be seen in the autocovariances plotted in Figure 3.1(b).

Figure 3.2 depicts the convergence of the proposed online EM scheme to the reference values of $b_x^{(m)}$, $m = 1, 2$, for various block sizes $L$ and particle populations $N$. Figure 3.3 depicts the evolution of the log-likelihood per iteration of the EM algorithm. Figure 3.4(a) depicts the normalized error in the identified $P^{(m)}$, $m = 1, 2$ and idiosyncratic variances $\Sigma$ pre coordinate after 20,000 iterations. In all cases the algorithm exhibits good convergence to the reference values.

3.2. Temperature Dataset. The goal of this numerical experiment is to illustrate the interpretability of the proposed model and compare with the switching-state linear model discussed in section 2.1 (Equation (2.6)). For that purpose we utilized the temperature data (in degrees Fahrenheit) of the capitals of the 50 states in the U.S.A
P.S. KOUTSOURELAKIS, E. BILIONIS

Fig. 3.1. Densities (a)) and Autocovariances (b)) of times series \( y_{t,1} \) (black), \( y_{t,2} \) (red) and 
\( y_{t,4} \) (blue) (solid lines). With (− ◦ −) the densities and autocovariances of the same times series
generated using the learned model parameters using the proposed online EM scheme with \( L = 200 \)
and \( N = 200 \) (see Figures 3.2 and 3.3).

Fig. 3.2. Convergence of \( b_{k}^{(1)} \) (black) and \( b_{k}^{(2)} \) (red) using the online EM algorithm for three
different combinations of \( L \) and \( N \). (− ◦ −) corresponds to \( L = 100 \), \( N = 100 \), (− ⊠ −) to \( L = 200 \),
\( N = 200 \) and (−△−) to \( L = 20 \), \( N = 1000 \) (\( d = 50 \)). The data was obtained from http://www.engr.udayton.edu/weather/citylistUS.htm
and it represents the average daily temperatures from 01/01/1996 until 01/13/2009
(i.e. 5,127 daily observations).

Figure 3.5 depicts the posterior memberships corresponding to the SLDS and
PMLDS models based on a reduced model with two hidden states (\( M = 2 \)) described
by one-dimensional OU processes (\( K = 1 \)). The former assumes that at each time
instant the observables \( y_{t} \) arises from a single hidden process. Hence a single entry
of \( z_{t} = [z_{1,t}, z_{2,t}, \ldots, z_{M,t}] \) is equal to 1 and the rest are all equal to 0. The top
part of Figure 3.5 shows the posterior mean of \( z_{m,t}, m=1,2 \). As one would expect
the two-states correspond to cold-winter (blue) and hot-summer (red) and alternate
periodically (roughly the cold-winter state is active between early November until
mid-April and the hot-summer state in the remainder of the calendar year). The top
part of Figure 3.7 depicts the corresponding \( P^{(m)}, m=1,2 \) where southern states have
higher values and northern states lower. Naturally, winter and summer represent the
two extremes but several intermediate states are also present. The proposed partial-membership model can account for those states without increasing the cardinality of the reduced-order dynamics. As it can be seen in the bottom part of Figure 3.5 which depicts the particulate approximation of the posterior of \( z_{m,t}, m = 1, 2 \), the two hidden states can also be attributed to the two extremes but the actual temperatures arise by a weighted combination of these two. Naturally during spring-summer months the weight of the “red” state is higher and during autumn-winter months the weight of the “blue” state takes over. The posterior of the hidden processes \( \mathbf{z}^{(m)}_t, m = 1, 2 \) is depicted in Figure 3.6.

In order to quantitatively compare the two models we calculated the average,
Fig. 3.5. (Top) Posterior mean of $z_{m,t}, m = 1,2$ based on the SLDS and (Bottom) particulate approximation of the posterior of $z_{m,t}, m = 1,2$ PMLDS. Both results were obtained using the previously discussed online EM scheme with $L = 200$ and $N = 100$

| $M$ | $K$ | SLDS         | PMLDS         |
|-----|-----|--------------|--------------|
| 2   | 1   | $-179.97 \pm 37.31$ | $-171.11 \pm 37.20$ |
| 2   | 2   | $-170.68 \pm 36.95$ | $-141.11 \pm 27.82$ |
| 4   | 1   | $-176.40 \pm 34.36$ | $-143.81 \pm 25.56$ |
| 4   | 2   | $-166.05 \pm 30.57$ | $-117.67 \pm 21.15$ |

Table 3.1

One-step-ahead predictive log-likelihood (Equation (3.1)) of SLDS and PMLDS models for various $M$, $K$. The table reports the average value plus/minus one standard deviation in bits. All results were obtained using the previously discussed online EM scheme with $L = 200$ and $N = 100$

The latter integral is approximated by Monte Carlo using the MAP estimate of $\Theta$ and the particulate approximation of the posterior for the dynamic variables $\Theta_t$. This provides a measure of how well the model generalizes to a novel observation from the same distribution as the training data and higher values imply a better model. Table 3.1 reports the average values (in bits) plus/minus the standard deviation (over $t \in [0,T]$).

$$
p(y_{t+1} \mid y_{1:t}) = \int \log p(y_{t+1} \mid \Theta, \Theta_{t+1} y_{1:t}) p(\Theta, \Theta_{t+1} \mid y_{1:t}) d\Theta d\Theta_{t+1}
$$

$$
= \int \log p(y_{t+1} \mid \Theta, \Theta_{t+1} y_{1:t}) \left[ \underbrace{p(\Theta_{t+1} \mid \Theta_t, \Theta)}_{\text{prior}} \right] \left[ \underbrace{p(\Theta, \Theta_t \mid y_{1:t})}_{\text{posterior}} \right] d\Theta d\Theta_{t+1} \tag{3.1}
$$

The latter integral is approximated by Monte Carlo using the MAP estimate of $\Theta$ and the particulate approximation of the posterior for the dynamic variables $\Theta_t$. This provides a measure of how well the model generalizes to a novel observation from the same distribution as the training data and higher values imply a better model. Table 3.1 reports the average values (in bits) plus/minus the standard deviation (over $t \in (100, T = 5127)$). Similar calculations were carried out for other model cardinalities (i.e. $M, K$) and in all cases the proposed model exhibited superior performance. This superiority becomes more pronounced as $M$ and $K$ increased which can be attributed to the factorial character of PMLDS.
3.3. Transient Heat equation. We finally apply the proposed analysis scheme to the one-dimensional transient heat equation:

\[
\begin{align*}
\frac{\partial u}{\partial t} &= \frac{\partial}{\partial x} \left( a(x) \frac{\partial u}{\partial x} \right), \quad x \in [0, 1] \\
u(0, t) &= u(1, t) = 0, \quad \forall t
\end{align*}
\]

(3.2)

The spatial domain was discretized with 1,000 finite elements of equal length and we considered a “rough” conductivity profile shown in Figure 3.8(a). The conductivity \(a(x)\) in each finite element was assumed constant and its value was drawn independen-
ently from a uniform distribution\footnote{We considered a single realization of the conductivity profile and solved for it as a deterministic problem. The stochastic PDE where \(a(x)\) is random is not considered here.}. For \(x \in [0, 0.5]\) we used the uniform \(U[0.01, 0.1]\) and for \(x \in (0.5, 1)\), \(U[0.51, 0.6]\). This naturally resulted in the jump observed in Figure 3.8(a), which as a consequence gave rise to two distinct slow time scales in the solution profile \(u(x, t)\) depicted in Figure 3.8(b). A rough profile of initial conditions was also used (as can be seen Figure 3.8(b) \(t = 0\)). In particular at each node \(x_i = 0.001i\), \(i = 1, \ldots, 1001\) we set \(u(x_i, 0) = 10x_i(1 - x_i)(1 + 0.1Z_i)\) where \(Z_i \sim N(0, 1)\) (i.i.d).

Upon spatial discretization, we obtain a coupled system of ODEs:

\[
y_i + Ky_i = 0
\]

where \(y_i \in \mathbb{R}^{1001}\) represents the solution at the nodes \(x_i\), i.e. \(y_i = [u(x_1, t), u(x_2, t), \ldots, \ldots, u(x_{1001}, t)]^T\). In contrast to existing approaches for the same diffusion equation (e.g. 2 [118]) we do not exploit mathematical properties of the PDE in specifying the coarse-grained model but rely on data. This data is obtained upon temporal discretization of Equation (3.3) where a time step \(\delta t = 0.0001\) was used. As a result at each time step we obtained a vector of observables \(y_i\) of dimension \(d = 1001\). This data was incorporated in the Bayesian model proposed using two hidden OU process (\(M = 2\)) of dimension \(K = 2\) each. In particular we employed the online EM scheme previously discussed over blocks of length \(L = 10\) time steps and \(N = 100\) particles. In particular (see also Figure 2.3):

- data over 20 times steps \(\delta t\), \(y_{1:20}\) (i.e. corresponding to total time \(20\delta t = 0.002\)) were ingested by the Bayesian reduced model, and
- the latter was used to predict the evolution of the system over 500 time steps (i.e. total time \(T = 500\delta t = 0.05\)).
- The original solver of the governing PDE was then re-initialized using the posterior mean estimate of the state of the system \(y_{20}\) and was run for further 20 time steps. Using the additional data obtained \(y_{20:520}\), the Bayesian model was updated and the process described was repeated.

It is noted that the proposed Bayesian prediction scheme results in a reduction of the number of fine scale integration time steps by a factor of 25 (\(T/20\delta t = 0.05/0.002\)) leading to a significant acceleration of the simulation process. Figure 3.9 depicts the posterior estimates of the solution at various time instants. In all cases these approximate very accurately the exact solution and these estimates improve as more are accumulated. One of the main advantages of the proposed approach is that apart from single-point estimates one can readily obtain credible intervals that quantitify predictive uncertainties due to information loss by the use of the reduced-order dynamic model and the finite amount of data used to learn that model. As it is seen in Figure 3.9 these envelop the exact solution and become tighter at larger times. As one would expect, when a larger predictive horizon \(T = 0.1\) (i.e. 1,000 time steps \(\delta t\) is used, as it can be seen in Figures 3.10 and 3.11 the predictive uncertainty grows. Such a scheme however is twice as efficient leading to a reduction of computational effort by a factor of 50 (i.e. \(T/20\delta t = 0.1/0.002\)). Hence if the analyst is willing to tolerate the additional uncertainty, efficiency gains can be achieved. This supports the arguments made previously with regards to an adaptive Bayesian scheme where, the level of predictive uncertainty would be specified and the algorithm would automatically revert to the fine scale model in order to obtain more data and improve the predictive estimates.
Bayesian reduced-order models

Fig. 3.8. Dynamic heat equation. Spatial discretization with 1,000 finite elements. Time discretization using $\delta t = 1 \times 10^{-4}$.

Fig. 3.9. Comparison of predictive posterior estimates (posterior mean and 5% and 95% quantiles) with exact solution $u(x, t)$ at various $t$.

4. Conclusions. The proposed modeling framework can extract interpretable reduced representations of high-dimensional systems by employing simple, low-dimensional processes. It simultaneously achieves dimensionality reduction and learning of reduced dynamics. The Bayesian framework adopted provides a generalization over single-point estimates obtained through maximum-likelihood procedures. It can quantify uncertainties associated with learning from finite amounts of data and produce probabilistic predictive estimates. The latter can be used to rigorously perform concurrent
Fig. 3.10. Comparison of predictive posterior estimates (posterior mean and 5% and 95% quantiles) with exact solution $u(x, t)$ at various $t$. These results were obtained with a prediction horizon $T = 0.1$ ($\delta t = 0.0001$) in contrast to Figure 3.9 which were obtained for $T = 0.05$.

Fig. 3.11. Comparison of predictive posterior estimates (posterior mean and 5% and 95% quantiles) with exact solution $u(x, t)$ at $t = 1.0$. These results were obtained with a prediction horizon (a) $T = 0.05$ and (b) $T = 0.1$.

Critical to the efficacy of the proposed techniques is scalability particularly with regards to the large dimension $d$ of the original process. The algorithms proposed imply $O(d)$ order of operations. Furthermore they can dynamically update the coarse-grained models as more data become available. In a typical scenario, the fine-scale model is reinitialized several times in order to obtain additional information about the system’s evolution that is incorporated in the coarse-grained dynamics “on the fly”.

The Bayesian, statistical perspective can readily be extended to the modeling stochastic dynamical systems. This would require generating more than one realizations of the original dynamics which can nevertheless be incorporated in the coarse-grained models using the same online EM scheme. In fact the loss of information that unavoidably takes place during the coarse-graining, results in probabilistic reduced-order models even if the original model was deterministic. A critical question that offers opportunity for future research on the topic relates to structural learning and in particularly with the dimensionality of the representation, i.e. the number of hid-
den processes $x_{t}^{(m)}$ needed (denoted by $M$ in Equation (2.7)). Treating this as a model selection problem as it was done in the examples, assumes that there is a single, optimal finite-dimensional representation. Current research activities are centered around nonparametric Bayesian priors over infinite combinatorial structures based on the Dirichlet process paradigm and infinite latent features models (e.g. [129, 70, 75]). These offer an alternative perspective by assuming that the number of building blocks is potentially unbounded, and that the observables only manifest a sparse subset of those. As a result, the cardinality of the coarse-grained model can be automatically determined from the data. Another aspect that warrants further investigation is prior modeling of the static parameters. Apart from the regularization effect this offers, it can promote the discovery of desirable features, such as slow-varying essential dynamics, sparse factors (e.g. $P^{(m)}$ in Equation (2.12)) which can advance the interpretability of the results and facilitate the inference process.
Appendix. This appendix discusses the sufficient statistics and update equations for the static parameters $\Theta$ used in the probabilistic model proposed. In the first section we discuss parameters appearing in the reduced-order dynamic models and in the second those appearing in the likelihood.

Sufficient statistics for parameters appearing in the prior. As discussed in section \ref{sec:2.2.1}, independent, isotropic OU processes are used as prior models for the latent, coarse-grained dynamics $x_{t}^{(m)} \in \mathbb{R}^{K}$ as well the process $z_{t} \in \mathbb{R}^{M}$ that models the frictional memberships to each process $m$. We therefore discuss the essential elements for the online EM computations described in section \ref{sec:2.3} \[\text{[6, 7, 8]}\] for a general isotropic OU process in $\mathbb{R}^{n}$ of the form:

$$dx_{t} = -b(x_{t} - q)dt + S^{1/2}dW_{t} \tag{4.1}$$

It is of interest to determine the parameters $\theta = (b, q, S)$. Let also $\pi(\theta)$ denote the prior on $\theta$. The readers can adjust the expressions below to any $x_{t}^{(m)}$ or $z_{t}$ since independent priors were used. Note that the stationary distribution of $x_{t}$ is a Gaussian:

$$\nu_{0}(x) = N(x | q, C = \frac{1}{2b}S) \tag{4.2}$$

and the transition density $p(x_{t} | x_{t-1})$ assuming that equidistant time instants with time step $\delta t$ are considered, is given by:

$$p(x_{t} | x_{t-1}) = N(x_{t} | \mu_{\delta t}(x_{t-1}), S_{\delta t}) \tag{4.3}$$

where:

$$\mu_{\delta t}(x_{t-1}) = x_{t-1} - (1 - e^{-b\delta t})(x_{t-1} - q) \tag{4.4}$$

and:

$$S_{\delta t} = \frac{1 - e^{-2b\delta t}}{2b}S \tag{4.5}$$

Given a block of length $L$ with observables $y_{1:L}$ and according to Equations \[\text{[2.27, 2.28]}\] we have that:

$$\hat{Q}(\theta^{(k-1)}, \Theta) = \left\langle -\frac{1}{2} \log |C| - \frac{1}{2}(x_{1} - q)^{T}C^{-1}(x_{1} - q) + \sum_{t=2}^{L} -\frac{1}{2} \log |S_{\delta t}| - \frac{1}{2}(x_{t} - q)^{T}C^{-1}(x_{t} - q) \right\rangle + \log \pi(\theta)$$

where the brackets $\left\langle . \right\rangle$ imply expectation with respect to $\hat{p}(x_{1:L} | \theta^{(k-1)}, y_{1:L})$ as in Equation \[\text{[2.28]}\]. In order to maximize $\hat{Q}(\Theta^{(1:k-1)}, \Theta)$ as in Equation \[\text{[2.30]}\] one needs to solve the system of equations arising from $\frac{\partial \hat{Q}(\Theta^{(1:k-1)}, \Theta)}{\partial \theta} = 0$. These equations equations with respect to $\theta$ are solved with fixed point iterations. They depend on
the following 7 sufficient statistics  \( \Phi = \{ \Phi_j \}_{j=1}^7 \):

\[
\begin{align*}
\Phi_1 &= <x_1> \\
\Phi_2 &= <x_1x_1^T> \\
\Phi_3 &= \left\langle \sum_{t=2}^{L} x_{t-1} \right\rangle \\
\Phi_4 &= \left\langle \sum_{t=2}^{L} x_t - x_{t-1} \right\rangle \\
\Phi_5 &= \left\langle \sum_{t=2}^{L} x_{t-1}x_{t-1}^T \right\rangle \\
\Phi_6 &= \left\langle \sum_{t=2}^{L} (x_t - x_{t-1})x_{t-1}^T \right\rangle \\
\Phi_7 &= \left\langle \sum_{t=2}^{L} (x_t - x_{t-1})(x_t - x_{t-1})^T \right\rangle
\end{align*}
\]

(4.6)

**Sufficient statistics for parameters appearing in the likelihood.** The process a bit more involved in the case of the parameters appearing in the likelihood Equation (2.14) i.e. the projection matrices \( \{ P^{(m)} \}_{m=1}^M \) of dimension \( d \times K \) and the covariance \( \Sigma \) which is a (positive definite) matrix of \( d \times d \). In order to retain scalability in high-dimensional problems (i.e. \( d \gg 1 \)) we assume a diagonal form of \( \Sigma = \text{diag}(\sigma_1^2, \sigma_2^2, \ldots, \sigma_d^2) \) which implies learning \( d \) parameters rather than \( d(d+1)/2 \).

Denoting now by \( \theta = ((P^{(m)})_{m=1}^M, \{ \sigma_j^2 \}_{j=1}^d) \), \( \pi(\theta) \) the prior and according to Equations (2.27) and (2.28) we have that:

\[
\hat{Q}(\theta^{(k-1)}, \theta) = \left\langle \sum_{t=1}^{L} -\frac{1}{2} \log |\Sigma| - \frac{1}{2}(y_t - W_tX_t)^T \Sigma^{-1}(y_t - W_tX_t) \right\rangle
\]

(4.7)

Differentiation with respect to \( P^{(m)} \) reveals that the stationary point must satisfy:

\[
A^{(m)} = \sum_{n=1}^{M} P^{(n)} B^{(n,m)}
\]

(4.8)

where the sufficient statistics are:

\[
\begin{align*}
\underline{A}^{(m)}_{d \times K} &= \left\langle \sum_{t=1}^{L} z_{t,m}y_t (x_t^{(m)})^T \right\rangle, \quad m = 1, 2, \ldots, M
\end{align*}
\]

(4.9)

and:

\[
\begin{align*}
\underline{B}^{(n,m)}_{K \times K} &= \left\langle \sum_{t=1}^{L} z_{t,n}z_{t,m}x_t^{(n)} (x_t^{(m)})^T \right\rangle
\end{align*}
\]

(4.10)

In the absence of a prior \( \pi(\theta) \) and if \( P^{(m)}_j \) and \( A_j^{(m)} \) represent the \( j^{th} \) rows \( (j = 1, \ldots, d) \) of the matrices \( P^{(m)} \) and \( A^{(m)} \) respectively, then Equation (4.9) implies:

\[
\begin{pmatrix}
A_j^{(1)} & A_j^{(2)} & \cdots & A_j^{(M)}
\end{pmatrix}
= \begin{pmatrix}
P_j^{(1)} & P_j^{(2)} & \cdots & P_j^{(M)}
\end{pmatrix}
\begin{pmatrix}
P_j^{(1,1)} & P_j^{(1,2)} & \cdots & P_j^{(1,M)} \\
P_j^{(2,1)} & P_j^{(2,2)} & \cdots & P_j^{(2,M)} \\
\vdots & \vdots & \ddots & \vdots \\
P_j^{(M,1)} & P_j^{(M,2)} & \cdots & P_j^{(M,M)}
\end{pmatrix}
\]

(4.11)
This leads to the following update equations for $P_j^{(m)}$, $\forall j, m$:

$$P_j = A_j B^{-1}$$

(4.12)

Note that the matrix $B$ to be inverted is independent of the dimension of the observables $d$ ($d >> 1$) and the inversion needs to be carried out once for all $j = 1, \ldots, d$. Hence the scaling of the update equations for $P^{(m)}$ is $O(d)$ i.e. linear with respect to the dimensionality of the original system.

Furthermore, in the absence of a prior $\pi(\theta)$, differentiation with respect to $\sigma_j^{-2}$ ($j = 1, \ldots, d$) leads to the following update equation:

$$L \sigma_j^2 = \sum_{t=1}^L y_{t,j}^2 - 2A_j P_j^T + P_j BP_j^T$$

(4.13)

In summary the sufficient statistics needed are the ones in Equations (4.9) and (4.12).

In the numerical examples in this paper a diffuse Gaussian prior was used for $P^{(m)}$ with variance 100 for each of the entries of the matrix. This leads to the addition of the term $1/100$ in the diagonal elements of the $B$ in Equation (4.12). No priors were used for $\sigma_j^2$. 

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