ABSTRACT

Driven by the critical needs of biomanufacturing 4.0, we present a probabilistic knowledge graph hybrid model characterizing complex spatial-temporal causal interdependencies of underlying bioprocessing mechanisms. It can faithfully capture the important properties, including nonlinear reactions, partially observed state, and nonstationary dynamics. Given limited process observations, we derive a posterior distribution quantifying model uncertainty, which can facilitate mechanism learning and support robust process control. To avoid evaluation of intractable likelihood, Approximate Bayesian Computation sampling with Sequential Monte Carlo (ABC-SMC) is developed to approximate the posterior distribution. Given high stochastic and model uncertainties, it is computationally expensive to match process output trajectories. Therefore, we propose a linear Gaussian dynamic Bayesian network (LG-DBN) auxiliary likelihood-based ABC-SMC algorithm. Through matching observed and simulated summary statistics, the proposed approach can dramatically reduce the computation cost and improve the posterior distribution approximation.

Keywords Approximate Bayesian Computation, Auxiliary Likelihood-based Summary Statistics, Cell therapy manufacturing, Bioprocess hybrid model, Latent State

1 Introduction

The biomanufacturing industry is growing rapidly and it plays a critical role to ensure public health and support economy. However, biomanufacturing often faces critical challenges, including high complexity, high variability, and very limited process observations. As new biotherapeutics (e.g., cell/gene therapies) become more and more personalized, biomanufacturing requires more advanced manufacturing protocols. With cells (or other living organisms, such as bacteria and yeast) as factories, it involves a complex stochastic decision process (SDP) with output trajectory dynamics and variations influenced by biological/physical/chemical (a.k.a. biophysicochemical) reactions occurring at molecular, cellular, and system levels.

In general, there are two main categories of biomanufacturing process modeling methodologies in the existing literature: mechanistic and data-driven approaches. The ordinary/partial differential equations (ODE/PDE) mechanistic models are developed based on biophysicochemical mechanisms. They have good interpretability and show generally higher extrapolation power than data-driven models. However, existing mechanistic models often fail to rigorously account for uncertainties, i.e., inherent stochasticity and model uncertainty. For example, batch-to-batch variation, known as a major source of bioprocess uncertainty [1], is ignored in deterministic mechanistic models. Therefore, mechanistic models may not fit well with the observations from real systems in many situations, which also limits their power in terms of mechanism learning and optimal/robust/personalized process control to support on-demand manufacturing. On the other hand, data-driven approaches often use general statistical or machine learning approaches to capture process patterns observed in data. The prediction accuracy of these models largely depends on the the size of process data and their interpretability is limited.

Driven by the critical challenges of biomanufacturing and limitations of existing process modeling methods, we developed probabilistic knowledge graph (KG) hybrid model characterizing the risk- and science-based understanding...
of bioprocess spatiotemporal causal interdependences \[2,3,4\]. It can leverage the information from existing mechanistic models between and within each operation unit, as well as facilitating mechanism learning from heterogeneous data. \[3\] introduced KG-based reinforcement learning (RL) to guide customized decision making. Since the proposed model-based RL scheme on the Bayesian KG can provide an insightful prediction on how the effect of inputs propagates through mechanism pathways, impacting on the output trajectory dynamics and variations, it can find process control policies that are interpretable and robust against model risk, and overcome the key challenges of biopharmaceutical manufacturing.

A computational Bayesian inference approach, called Approximate Bayesian Computation (ABC), is used to approximate the posterior distribution for models with intractable likelihoods. It bypasses the evaluation of likelihood function by simulating model parameters and synthetic data sets, and retaining the parameter samples such that the associated data set are “similar enough” to the observed data set. For complex biomanufacturing processes with high stochastic and model uncertainties, it is computationally challenging to generate simulation trajectories close to real-world observations. Thus, the similarity can be measured based on summary statistics. Obviously, the choice of statistics is paramount. Recently, there has been much interest in formalizing an auxiliary likelihood based ABC, which uses a simpler and related model to derive summary statistics \[5,6,7\].

Following the spirit of the auxiliary likelihood-based ABC \[6\], we utilize a linear Gaussian dynamic Bayesian network (LG-DBN) auxiliary model to derive summary statistics for ABC-SMC that can accelerate online inference on hybrid models with high fidelity characterizing complex bioprocessing mechanisms. The proposed DBN auxiliary approach in conjunction with sequential importance sampling can efficiently approximate hybrid model posterior distribution. Therefore, the key contributions of this paper: (1) given very limited real world data, we propose DBN auxiliary likelihood-based ABC-SMC sampling to generate posterior samples of bioprocess hybrid model parameters quantifying model uncertainty; (2) this simple LG-DBN auxiliary model can capture the critical dynamics and variations of bioprocess trajectory, ensure the computational efficiency, and enable high quality of inference, which can facilitate mechanism online learning and support robust process control; and (3) the empirical study shows that our approach can outperform the original ABC-SMC approach given tight computational budget.

The remainder of the paper is organized as follows. We provide the problem description and summarize the proposed framework in Section 2. Then, we present a probabilistic KG hybrid model, capturing the important properties of biomanufacturing processes, and describe ABC for approximating the posterior distribution of model parameters in Section 3. We derive LG-DBN auxiliary likelihood based summary statistics, which can facilitate Bayesian inference on hybrid models with high fidelity in Section 4. We conduct the empirical study on cell therapy manufacturing in Section 5 and conclude the paper in Section 6.

2 Problem Description and Proposed Framework

Driven by the needs of biomanufacturing process online learning, monitoring, and control, we create a probabilistic KG hybrid model characterizing underlying mechanisms and causal interdependencies between critical process parameters (CPPs) and critical quality attributes (CQAs). It models how the effect of state and action at any time \( t \), denoted by \( \{s, a\} \), propagates through mechanism pathways impacting on the output trajectory dynamics and variations. Here we use cell culture process for illustration. The process state transition model is denoted by \( p(s_{t+1}|s_t, a_t; \theta) \) where \( s_t \in \mathcal{S} \subset \mathbb{R}^d \) denotes the partially observable bioprocess state (i.e., extra- and intra-cellular enzymes, proteins, metabolites, media), \( a_t \in \mathcal{A} \) denotes action (i.e., agitation rate, oxygen/nutrient feeding rates), \( \theta \) is a finite set of actions, and \( t \in \mathcal{H} \equiv \{1,2,\ldots,H+1\} \) denotes the discrete time index. At any time \( t \), the agent observes the state \( s_t \) and takes an action \( a_t \). Thus, given model parameters \( \theta \), the joint distribution of process trajectory \( \tau = (s_1, a_1, \ldots, s_{H}, a_{H}, s_H) \) becomes,

\[
p(\tau|\theta) = p(s_1) \prod_{t=1}^{H} p(s_{t+1}|s_t, a_t; \theta) p(a_t).
\]

The state transition \( p(s_{t+1}|s_t, a_t; \theta) \) is modeled by a hybrid (mechanistic/statistical) model. Its structure takes existing mechanistic models as prior. For example, since the key factors influencing process dynamics and variability in cell
culture are induced by cellular metabolisms [3], the probabilistic state transition of this KG hybrid model can incorporate cell metabolic networks and account for the variations of cell behaviors under heterogeneous micro-environments. Given limited historical trajectory observations, we focus on hybrid model Bayesian inference to support online mechanism learning, monitoring, and reliable interpretable prediction, accounting for both inherent stochasticity and model uncertainty.

There are key properties in biomanufacturing process, specially for personalized cell/gene therapies, including (1) partially observed state \( s_t \) that means only limited proportion of state observable; (2) stochastic state transition model \( p(s_{t+1} | s_t, a_t ; \theta) \) involves high inherent stochasticity; and (3) very limited and heterogeneous online and offline measurement data. In this paper, the posterior distribution will be derived to quantify model uncertainty. Due to the nature of biopharmaceutical manufacturing, the stochastic state transition models \( p(s_{t+1} | s_t, a_t ; \theta) \) are highly complex, non-linear, and nonstationary.

### 2.1 Hybrid Modeling for Biomanufacturing Process with Partially Observed State

At any time step \( t \), the process state \( s_t \) is composed of observable and latent state variables, i.e., \( s_t = (x_t, z_t) \) with \( x_t \in S_x \) and latent state variables \( z_t \in S_z \), where \( S_x \subseteq \mathbb{R}^{d_x} \) and \( S_z \subseteq \mathbb{R}^{d_z} \) with \( S_x = S_x \times S_z \) and \( d = d_x + d_z \). Denote the partially observed state trajectory as \( \tau_t \equiv (x_1, a_1, \ldots, x_H, a_H, x_{H+1}) \). Given model parameters \( \theta \), by integrating out the latent states \( (z_1, \ldots, z_{H+1}) \), the likelihood of observation \( \tau_t \) becomes,

\[
p(\tau_t | \theta) = \int \cdots \int p(\tau | \theta) dz_1 \cdots dz_{H+1}.
\]

To support bioprocess mechanism learning and decision making, this hybrid model characterizes the risk- and science-based understanding of underlying mechanisms and spatial-temporal causal interdependencies of CPPs/CQAs. It can connect heterogeneous online and offline measurements to infer unobservable state (such as assessing underlying metabolic state that determines cell product functional properties), support process monitoring, and facilitate real-time release.

Therefore, we model the bioprocess state transition with a hybrid model. Given the existing ODE-based mechanistic model, \( ds/dt = f(s, a, \phi) \), by using the finite difference approximation for derivatives, i.e., \( ds \approx \Delta t s = s_{t+1} - s_t \), and \( d\tau \approx \Delta t \), we construct the hybrid model for state transition,

\[
x_{t+1} = x_t + \Delta t \cdot \boldsymbol{f}_x(x_t, z_t, a_t; \phi) + \varepsilon_{x,t+1}^x \quad \text{and} \quad z_{t+1} = z_t + \Delta t \cdot \boldsymbol{f}_z(x_t, z_t, a_t; \phi) + \varepsilon_{z,t+1}^z
\]

with unknown kinetic coefficients \( \phi \in \mathbb{R}^{d\phi} \) (e.g., cell growth and inhibition rates). The function structures of \( \boldsymbol{f}_x(\cdot) \) and \( \boldsymbol{f}_z(\cdot) \) are the parts of \( \boldsymbol{f}(\cdot) \) associated to the observable state output \( x_{t+1} \) and the latent state output \( z_{t+1} \). By applying the central limit theorem, the residual terms, accounting for inherent stochasticity and other factors, are modeled by multivariate Gaussian distributions \( \varepsilon_{x,t+1}^x \sim \mathcal{N}(0, V^x) \) and \( \varepsilon_{z,t+1}^z \sim \mathcal{N}(0, V^z) \) with zero means and covariance matrices \( V^x \) and \( V^z \). Then, the state transition distribution becomes,

\[
x_{t+1} | x_t, z_t, a_t \sim \mathcal{N} \left( x_t + \Delta t \cdot \boldsymbol{f}_x(x_t, z_t, a_t), V^x_{t+1} \right) \quad \text{and} \quad z_{t+1} | x_t, z_t, a_t \sim \mathcal{N} \left( z_t + \Delta t \cdot \boldsymbol{f}_z(x_t, z_t, a_t), V^z_{t+1} \right)
\]

Therefore, the state transition model specified by parameters \( \theta = (\phi, V^x, V^z) \) characterizes the bioprocess dynamics, mechanisms, and inherent stochasticity.

### 2.2 Challenges of Hybrid Model Inference Under High Stochasticity and Limited Data

Given limited real-world data with size \( m \), denoted by \( D = \{ \tau_x^{(i)} : i = 1, 2, \ldots, m \} \), the model uncertainty is quantified by a posterior distribution derived through applying the Bayes’ rule,

\[
p(\theta | D) \propto p(\theta) p(D | \theta) = p(\theta) \prod_{i=1}^{m} p \left( \tau_x^{(i)} | \theta \right)
\]

where \( p(\theta) \) represents the prior distribution. It is challenging to directly derive or computationally assess the posterior distribution \( p(\theta | D) \) in eq. (1). First, there could exist large-dimensional latent state variables \( z_t \), especially for multi-scale bioprocess model developed to characterizing the cell response to environmental perturbation. It is computationally expensive to assess the likelihood of each observation, \( p(\tau_x^{(i)} | \theta) = \int \cdots \int p(\tau_x^{(i)} | \theta) dz_1 \cdots dz_{H+1} \), especially for biomanufacturing process with optical sensor assisted online monitoring, i.e., the value of \( H \) is large. Second, the mechanistic model \( f(s, a, \phi) \) can be a nonlinear function of state \( z \) as well as a nonlinear function of parameters \( \phi \). The kinetic coefficients can be random even though we start with fixed values in this paper. For example, the kinetic
coefficients (such as cell growth rate, oxygen/nutrient uptake rates, and metabolic waste excretion rates) can depend on the gene of seed cells, as well as cell culture environments. They can have batch-to-batch variations. Third, the amount of real-world process observations can be very limited (especially for personalized bio-drug manufacturing) even though inherent stochasticity and model complexity can be high. This leads to high model uncertainty.

Thus, in Section 3, ABC approach is considered to approximate the posterior distribution of hybrid model with high fidelity that can capture the key features of biomanufacturing processes. Since it is computationally expensive especially under the situations with high stochastic and model uncertainties, LG-DBN auxiliary ABC-SMC is used to facilitate the Bayesian inference. This auxiliary model can be accurate for biomanufacturing process online monitored with optical sensors (i.e., TPE and Raman sensors).

3 Sequential Importance Sampling based Hybrid Model Inference and Algorithm Development

When the assessment of likelihood for any observation is computationally intractable, such as \( p(\tau^{(i)}|\theta) = \int \cdots \int p(\tau^{(i)}|\theta)dz_1 \cdots dz_{h+1} \) for \( i = 1, 2, \ldots, m \), the ABC approach is recommended to approximate the posterior distribution \( \mathcal{P}^* \). In the naive ABC implementation, we draw a candidate sample from the prior \( \theta \sim p(\theta) \) and then generate a simulation dataset \( \mathcal{P}^* \) from the hybrid model. If the simulated dataset \( \mathcal{P}^* \) is “close” to the observed real-world observations \( \mathcal{P} \), we accept the sample \( \theta \); otherwise reject it. Thus, we approximate the posterior distribution \( p(\theta|\mathcal{P}) \) with \( p(\theta|d(\mathcal{P}, \mathcal{P}^*) \leq h) \), where \( d(\cdot) \) is a distance metric (e.g., Euclidean distance, likelihood distance) and \( h \) is an approximation tolerance level.

However, for any given small tolerance level \( h \), we often face very low accept rate for complex biomanufacturing processes with high stochastic and model uncertainties. The random discrepancy between multivariate process trajectories \( \mathcal{P} \) and \( \mathcal{P}^* \) could be large even when we have the parameter sample \( \theta \) equal to \( \theta^* \). In addition, given very limited real-world data for the complex hybrid model, the design space of \( \theta \) and the model uncertainty can be large.

To increase the accept rate and ensure the computational efficient generation of samples \( \theta \) with good approximation on the critical features occurring in the real-world data, we can define the distance measure \( d(\cdot) \) based on selected lower dimensional summary statistics, denoted by \( \eta(\mathcal{P}) \). That mean we accept samples \( \theta \) which lead to the summary statistics of simulated data, denoted by \( \eta^* = \eta(\mathcal{P}^*) \), close to the one of observations \( \eta_{obs} = \eta(\mathcal{P}) \). Thus, the standard ABC framework \( \beta \) becomes

\[
p_{ABC}(\theta|\eta_{obs}) \propto \int \mathbb{1}(d(\eta^*, \eta_{obs}) \leq h) p(\eta^*|\theta)p(\theta)d\eta^*.
\]

Therefore, as the distance tolerance \( h \) gradually decreases, we have

\[
\lim_{h \to 0} p_{ABC}(\theta|\eta_{obs}) \propto \int \delta_{\eta_{obs}}(\eta^*) p(\eta^*|\theta)p(\theta)d\eta^* = p(\eta_{obs}|\theta)p(\theta) \propto p(\theta|\eta_{obs}),
\]

where \( \delta_x(x) \) denotes the Dirac measure, defined as \( \delta_x(x) = 1 \) if \( x \in X \) and \( \delta_x(x) = 0 \) otherwise.

A good design of ABC summary statistics \( \eta \) should balance complexity v.s. informativeness. If the summary statistics \( \eta \) are sufficient for \( \theta \), then \( p(\theta|\eta_{obs}) \) will be equivalent to \( p(\theta|\mathcal{P}) \). Then, with small threshold \( h \), the ABC approximate \( p_{ABC}(\theta|\eta_{obs}) \) in (2) can provide a good approximation of the true posterior. However, in the most situation, it is challenging to specify the sufficient statistics since the KG hybrid model is built based on nonlinear mechanistic model and it accounts for the key features including (1) partially observed state; (2) heterogeneous offline and online measures; (3) nonlinear mechanisms and dynamics; and (4) batch-to-batch variations on mechanistic coefficients. Thus, in Section 4 we project the bioprocess hybrid model into linear Gaussian dynamic Bayesian Network (LG-DBN) based auxiliary model space that has tractable likelihood. We will use the LG-DBN likelihood to derive summary statistics accelerating the selection of samples \( \theta \). This LG-DBN can capture the key properties of process dynamics and variations to support robust and optimal control. Our study also shows that complex KG hybrid models will asymptotically converge to a LG-DBN model as time interval \( \Delta t \) becomes “smaller and smaller” by applying Taylor approximation [3]. This LG-DBN approximation holds well for many cases with online sensor measurements and the biological state of cells does not change quickly.

The basic ABC generates candidate samples from the prior \( p(\theta) \) and uses the accept/reject approach to keep those samples satisfying the approximation threshold requirement. This can be extremely ineffective especially for the situations using noninformative prior that has a wide sampling space. The ABC-sequential Monte Carlo (ABC-SMC) methods derived from the sequential importance sampling [9][10] can improve the sampling efficiency through generating candidate samples from updated posterior approximates. In specific, let \( g \) denote the index of ABC iterations.
used to improve the approximation of the posterior distribution \( p(\theta|D) \). We select a sequence of intermediate target distribution, denoted by \( \{\pi_g\} \) for \( g = 1, 2, \ldots, G \), converging to the posterior \( p(\theta|D) \), i.e.,

\[
\pi_g(\theta) = p(\theta) \mathbb{I}(d(\eta^*, \eta_{obs}) \leq h_g).
\]

Through gradually reducing the tolerance level \( h_g \), we can better approximate the posterior distribution \( p(\theta|D) \). Direct sampling from \( p(\theta) \) and having the accept/reject based on the condition \( \mathbb{I}(d(\eta^*, \eta_{obs}) \leq h_g) \) in (4) is not simulation efficient. The accept rate can be as low as \( h_g \) becomes smaller and smaller.

Thus, we use the *sequential importance sampling (SIS)* and select a sequence of proposal distribution, denoted by \( \{\zeta_g\} \) for \( g = 1, 2, \ldots, G \), to improve the sampling efficiency, i.e.,

\[
\zeta_g(\theta) = \mathbb{I}(\pi_g(\theta) > 0) \int \pi_{g-1}(\theta') K(\theta', \theta) d\theta',
\]

where \( K(\theta', \theta) \) is a Markov kernel. The proposal distribution \( \zeta_g(\theta) \) is defined as the perturbed previous intermediate distribution \( \pi_{g-1} \) through the perturbation kernel \( K(\cdot) \). After generating \( N \) samples from the proposal distribution \( \theta_n \sim \zeta_g(\theta) \) for \( n = 1, 2, \ldots, N \), we weight it by \( w_n^{(g)} = \pi_g(\theta_n)/\zeta_g(\theta_n) \). The condition, \( \mathbb{I}(\pi_g(\theta) > 0) \), in (5) is used to satisfy the importance sampling condition, i.e., \( \{\theta: \pi_g(\theta) > 0\} \subset \{\theta: \zeta_g(\theta) > 0\} \). This can avoid the weight becoming infinite, which will lead to high variance on the SIS estimator. We set the first proposal distribution to be the prior distribution, i.e., \( \zeta_1(\theta) = p(\theta) \).

**Algorithm 1: DBN auxiliary based ABC-SMC for hybrid model Bayesian inference.**

**Input:** the prior distribution \( p(\theta) \); the number of particles \( N \); process observations \( D = \{\tau_n^{(i)}\}_{i=1}^m \); the perturbation kernel function \( K(\cdot) \); the number of particles to keep at each iteration \( N_\alpha = \lfloor \alpha N \rfloor \) with \( \alpha \in [0, 1] \); and the minimal acceptance rate \( p_{acc_{min}} \).

**Output:** posterior distribution approximate \( \hat{p}(\theta|D) \).

for \( n = 1, \ldots, N \) do

1. Sample \( \theta_n^{(0)} \sim p(\theta) \);
2. Generate \( m \times L \) predicted trajectories \( \{\tau_n^{(i)}\}_{i=1}^L \) using \( \theta_n^{(0)} \);
3. Set \( q_n^{(0)} = d(\eta(\tau_n), \eta(\tau_n)) \) and \( w_n^{(0)} = 1 \);

4. Let \( h_1 \) be the first \( \alpha \)-quantile of \( q_n^{(0)} \) \( \{q_n^{(0)}\}_{n=1}^N \);
5. Let \( \{\theta_n^{(1)}, w_n^{(1)}, q_n^{(1)}\} = \{(\theta_n^{(0)}, w_n^{(0)}, q_n^{(0)}) | q_n^{(0)} \leq h_1, 1 \leq n \leq N\} \), \( p_{acc} = 1 \) and \( g = 2 \);

while \( p_{acc} > p_{acc_{min}} \) do

for \( n = N_\alpha + 1, \ldots, N \) do

6. Sample \( \theta_k^{(g-1)} \) from \( \theta_k^{(g-1)} \) with probability \( \frac{w_n^{(g-1)}}{\sum_{j=k}^N w_n^{(g-1)}}, 1 \leq k \leq N_\alpha \);
7. Perturb the particle to obtain \( \hat{\theta}_n^{(g-1)} \sim K(\theta_n^{(g-1)}); \)
8. Generate \( m \times L \) predicted trajectories \( \{\tau_n^{(i)}\}_{i=1}^L \) using \( \theta_n^{(g-1)} \);
9. Set \( q_n^{(g-1)} = d(\eta(\tau_n), \eta(\tau_n)) \);
10. Set \( w_n^{(g-1)} = \frac{p(\theta_n^{(g-1)}) d(\eta(\tau_n), \eta(\tau_n)) \mathbb{I}(d(\eta^*, \eta_{obs}) \leq h_{g-1})}{\sum_{j=1}^N w_n^{(j-1)} K(\theta_n^{(g-1)}, \theta_j^{(g-1)})} \);
11. Set \( p_{acc} = \frac{1}{N-N_\alpha} \sum_{k=N_\alpha+1}^N \mathbb{I}(q_k^{(g-1)} \leq h_{g-1}); \)
12. Let \( h_g \) be the first \( \alpha \)-quantile of \( q_n^{(g-1)} \) \( \{q_n^{(g-1)}\}_{n=1}^N \);
13. Let \( \{(\theta_n^{(g)}, w_n^{(g)}, q_n^{(g)})\} = \{(\theta_n^{(g-1)}, w_n^{(g-1)}, q_n^{(g-1)}) | q_n^{(g-1)} \leq h_g, 1 \leq n \leq N\} \) and \( g = g + 1 \);
14. Return the approximated posterior distribution, \( \hat{p}(\theta|D) = \frac{1}{\sum_{n=1}^N w_n^{(g-1)}} \sum_{n=1}^N w_n^{(g-1)} \delta_{\theta_n^{(g-1)}}(\theta) \);

The proposed LG-DBN auxiliary likelihood-based ABC-SMC sampling procedure is summarized in Algorithm 1. It incorporates an adaptive selection approach on the threshold \( h_g \) from \( [9][11][12] \). The initial set of parameter samples \( \{\theta_n^{(0)}\}_{n=1}^N \) is generated from the prior distribution \( p(\theta) \) in Step 1. The associated weights \( \{w_n^{(0)}\}_{n=1}^N \) and distances \( \{q_n^{(0)}\}_{n=1}^N \) are calculated in Steps 2-3. Considering the impact from stochastic uncertainty, we generate \( m \times L \) predicted trajectories, compute the DBN auxiliary based summary statistics \( \eta^* \), and then calculate the distance \( q_n^{(0)} \). The
tolerance level $h_g$ in any $g$-th iteration is determined online as the $\alpha$-quantile of the $\{q_{\alpha}^{(g)}\}_{g=1}^N$. The particles, satisfying this tolerance denoted by $\{\Theta^n\}_{n=1}^{N_{\alpha}}$, constitute the weighted empirical distribution to approximate the posterior distribution in Steps 5 and 13, where $N_{\alpha} = \lceil \alpha N \rceil$. The approximation accuracy is measured by the corresponding distances $\{q^n\}_{n=1}^{N_{\alpha}}$. Then, $N - N_{\alpha}$ new particles are drawn from the proposal distribution $\tilde{\zeta}_{\theta}(\Theta)$ in Steps 6-7. The associated weights and distances are calculated in Steps 8-10. The tolerance level $h_g$ and the posterior distribution approximate $p_{\theta}(\Theta)$ are updated in Steps 12-13. We repeat Steps 6-13 until the proportion of particles satisfying the tolerance level $h_{g-1}$ among the $N - N_{\alpha}$ new particles is below the pre-specified threshold $p_{\text{tune}}$. Finally, the ABC-SMC algorithm returns the weighted empirical distribution, denoted by $\tilde{p}(\Theta|\mathcal{D})$, as posterior distribution approximate in Step 14.

4 DBN Auxiliary Likelihood-based Summary Statistics for Distance Measure Development

Motivated by the studies [6, 5], in this section, we derive LG-DBN auxiliary likelihood-based summary statistics for ABC-SMC to capture the crucial features of the bioprocess trajectory, including dynamics and variations. Given a set of observations $\mathcal{D} = \{\mathbf{x}^{(i)}_t : i = 1, 2, \ldots, m\}$, we derive the MLE of LG-DBN auxiliary model, i.e., maximizing the log-likelihood $\hat{\beta}(\mathcal{D}) = \arg\max_{\beta} p(\mathcal{D}|\beta)$. Then we use it as the summary statistics $\eta \equiv \hat{\beta}^*$ to calculate the distance measure $q \equiv d(\hat{\beta}, \beta^*)$, where $\hat{\beta}^*$ is the summary statistics of simulated data. In the following, we first develop the LG-DBN model with only observable state transition in Section 4.1 and then discuss the parameter estimation in Section 4.2.

4.1 The development of LG-DBN Auxiliary Model

Let $\mathbf{x}^k_t \sim \mathcal{N}(\mu^k_t, (\psi^k_t)^2)$ with $k = 1, 2, \ldots, d$ model the variation in the $k$-th initial observed state. In practice, to ensure product quality, CPPs are strictly regulated by the specifications of ranges of values. Thus, we model $\mathbf{a}_t$ as a random variable. Let $d^k_t \sim \mathcal{N}(\lambda^k_t, (\sigma^k_t)^2)$ with $k = 1, 2, \ldots, d_{\text{new}}$ model the variation in the $k$-th action for $t = 1, 2, \ldots, H$. At any time $t$, LG-DBN auxiliary model has the state transition model,

$$x_{t+1} = \mu_{t+1}^* + \psi_{t}^*(x_{t} - \mu_{t}^*) + \psi_{t}^*(a_{t} - \mu_{t}^*) + (V_{t+1}^*)^\frac{1}{2} \omega_{t},$$

where $\mu_{t}^* = (\mu_{1}^*, \ldots, \mu_{d_{t}}^*)$, $\mu_{t}^* = (\lambda_{1}^*, \ldots, \lambda_{d_{t}}^*)$, $\omega_{t}$ is an $d_{t}$-dimensional standard normal random vector, and $V_{t+1}^*$ is a diagonal covariance matrix. The coefficients $\psi_{t}^*$ and $\psi_{t}^*$ measure the main effects of current observed state $x_{t}$ and action $a_{t}$ on the next observed state $x_{t+1}$. Let $\mathbf{a}_t = (\sigma_1^t, \ldots, \sigma_{d_{\text{new}}}^t)$ and $\mathbf{v}_t = (v_{1}^t, \ldots, v_{d_{\text{new}}}^t)$. Thus, the LG-DBN model, specified by parameters $\hat{\beta} = (\mu^t, \psi^t, \psi^t, \mu^t, \sigma^t, v^t) = \{(\mu_{1}^t, \mu_{2}^t, \psi_{1}^t, \psi_{2}^t, \sigma_{1}^t, v_{1}^t)| 1 \leq t \leq H\}$, has the joint distribution of bioprocess trajectory: $p(\mathbf{x}_{t}) = p(x_1, a_1, \ldots, x_H, a_H, x_{H+1}) = p(x_1) \prod_{t=1}^{H} p(x_{t+1}|x_{t}, a_{t})p(a_{t})$.

Let $\mu_{t} = [\mu_{1}^t, \mu_{2}^t, \ldots, \mu_{H}^t, \mu_{H+1}^t]^{T}$. Following [13], we rewrite (6) in the following form

$$\mathbf{a}_t = \mu_{t} + B(\mathbf{a}_t - \mu_{t}) + \Sigma^{\frac{1}{2}} \omega_t$$

where $\omega_t$ is an $(H + 1)d_{a} + Hd_{v})$-dimensional standard normal random vector, $\Sigma^{\frac{1}{2}} = \text{diag}(v_{1}^t, \sigma_{1}^t, \ldots, v_{H}^t, \sigma_{H}, v_{H+1}^t)$ is the diagonal matrix of the conditional standard deviations of observed state and actions, and the coefficient matrix of observed trajectory is written as

$$B = \begin{bmatrix}
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\psi_{1}^t & \psi_{1}^t & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & \psi_{2}^t & \psi_{2}^t & 0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
0 & 0 & 0 & 0 & 0 & 0 & \psi_{H}^t & \psi_{H}^t & 0 \\
0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0
\end{bmatrix}.$$ 

Thus, by rearranging (7) and letting $\mathbf{a}_t = \mathbf{B}(\mathbf{a}_t - \mu_{t}) + \Sigma^{\frac{1}{2}} \omega_t$, we have $\mathbf{a}_t \sim \mathcal{N}(\mu_{t}, (I - B)^{-1}\Sigma^{\frac{1}{2}}(I - B)^{-T})$ with mean $\mathbb{E}[\mathbf{a}_t] = \mathbf{B}(\mathbf{a}_t - \mu_{t})$ and covariance matrix $\text{Cov}(\mathbf{a}_t - \mu_{t}) = (I - B)^{-1}\Sigma^{\frac{1}{2}}(I - B)^{-T}$.

4.2 Linear Gaussian Dynamic Bayesian Network based Summary Statistics

Let $\tilde{\mathbf{a}}, \tilde{x}_{1}, \ldots, \tilde{x}_{H}, \tilde{a}_{H}, \tilde{x}_{H+1} = \mathbf{a}_t - \mu_{t}$, where $\tilde{x}_{t}$ and $\tilde{a}_{t}$ denote centered observable state and decision. Given $m$ observations $\mathcal{D} = \{\mathbf{a}_t^{(i)}\}_{i=1}^{m}$, the unbiased estimator $\tilde{\mu}_{\tau} = \frac{1}{m} \sum_{i=1}^{m} \tilde{\mathbf{a}}^{(i)}$ can.Given $m$ observations $\mathcal{D} = \{\mathbf{a}_t^{(i)}\}_{i=1}^{m}$, the unbiased estimator $\tilde{\mu}_{\tau} = \frac{1}{m} \sum_{i=1}^{m} \tilde{\mathbf{a}}^{(i)}$ can be easily obtained by using the fact $\mathbb{E}[$
The log-likelihood of the centered trajectory observations \( \{ \tilde{x}_i^{(0)} \}_{i=1}^m \) becomes,

\[
\max_{\psi^r, \psi^a, V} \ell \left( \tilde{x}_1^{(0)}, \ldots, \tilde{x}_m^{(0)}; \psi^r, \psi^a, V \right) = \max_{\psi^r, \psi^a, V} \log \prod_{i=1}^m p \left( \tilde{x}_i^{(0)} \right) \\
= \max_{V_i} \sum_{i=1}^m \log p \left( \tilde{x}_i^{(0)} \right) - \sum_{i=1}^m \log \sum_{j=1}^H \max_{\psi^r, \psi^a} \log p \left( \hat{a}_i^{(0)} | \tilde{x}_i^{(0)} \right)
\]

(8)

Since both initial state \( \tilde{x}_i^{(0)} \) and actions \( \hat{a}_i^{(0)} \) are normally distributed with mean zero, the MLEs of their variance are sample covariances:

\[
\hat{V}_x^{(1)} = \frac{1}{m} \sum_{i=1}^m \left( \tilde{x}_i^{(0)} \right)^2
\]

with \( k = 1, 2, \ldots, d_a \). In addition, at any time \( t \), we have the log likelihood of a sample \( \tilde{x}_i^{(0)} \)

\[
\log p \left( \tilde{x}_i^{(1)} | \tilde{x}_i^{(0)}, \hat{a}_i^{(0)} \right) \propto -\frac{m}{2} \log |\tilde{V}_x^{(1)}| - \frac{1}{2} \left( \tilde{x}_i^{(1)} - \tilde{V}_x^{(1)} \tilde{x}_i^{(0)} - \psi^r \tilde{a}_i^{(0)} \right)^{\top} \tilde{V}_x^{(1)} \left( \tilde{x}_i^{(1)} - \tilde{V}_x^{(1)} \tilde{x}_i^{(0)} - \psi^r \tilde{a}_i^{(0)} \right)
\]

(9)

Let \( \hat{x}_i^{(0)} \) and \( \hat{a}_i^{(0)} \) denote the \( i \)-th rows of output matrix \( X \) and the input matrix \( Y \). Let \( B_t = (\psi_r, \psi_a)^{\top} \) denote the coefficient vector. As a result, the MLEs of \( \psi_r \) and \( \psi_a \) are

\[
(\psi_r, \psi_a)^{\top} = \hat{B}_t = \arg \max_{B_t} -\frac{1}{2} (Y - XB_t)^{\top} (V_r^{(t)} - 1) (Y - XB_t) = (X^{(t)} (V_r^{(t)} - 1) X)^{\top} Y (V_r^{(t)} - 1) Y.
\]

(10)

The MLE of each standard deviation can be computed by \( \hat{\psi}_{r}^{\top} = \sqrt{\frac{1}{m} \sum_{i=1}^m } \left( \tilde{x}_i^{(0)} \right)^2 \) \([14]\). In summary, from observations \( S \), the auxiliary MLE can then be obtained as \( \hat{B} = (\hat{\mu}^r, \hat{\mu}^a, \hat{\psi}_r, \hat{\psi}_a, \hat{\sigma}_{\psi}, \hat{\sigma}_{\psi} \).

## 5 Empirical Study

In this section, we use the erythroblast cell therapy manufacturing example presented in [15] to assess the performance of the proposed LG-DBN auxiliary likelihood-based ABC-SMC approach.

### 5.1 Hybrid Modeling for Cell Therapy Manufacturing Process

The cell culture of erythroblast exhibits two phases: a relatively uninhibited growth phase followed by an inhibited phase. [15] identified that this reversible inhibition is caused by an unknown cell-driven factor rather than commonly known mass transfer or metabolic limitations. They developed an ODE-based mechanistic model describing the dynamics of an unidentified autocrine growth inhibitor accumulation and its impact on the erythroblast cell production process,

\[
\frac{d\rho_t}{dr} = \rho_t \left( 1 - \left( 1 + e^\left( h(k_e - k_i) \right) \right)^{-1} \right) \quad \text{and} \quad \frac{dI_t}{dr} = \frac{d\rho_t}{dr} - r_d I_t
\]

where \( \rho_t \) and \( I_t \) represent the cell density and the inhibitor concentration (i.e., latent state) at time \( t \). The kinetic coefficients \( \phi = \{ r_c, k_c, k_e, r_d \} \) denote the cell growth rate, inhibitor sensitivity, inhibitor threshold, and inhibitor decay. Then, we construct the hybrid model

\[
\rho_{t+1} = \rho_t + \Delta \cdot r_c \rho_t \left( 1 - \left( 1 + e^\left( h(k_e - k_i) \right) \right)^{-1} \right) + e_t \rho \quad \text{and} \quad I_{t+1} = I_t + \Delta \cdot \left( \frac{\rho_{t+1} - \rho_t - r_d I_t}{\Delta} \right) + e_t
\]

(11)

where the residuals follow the normal distributions \( e_t \sim \mathcal{N}(0, \sigma_{e_t}) \) and \( e_t \sim \mathcal{N}(0, \sigma_{e_t}) \) by applying CLT. Therefore, the hybrid model is specified by \( \theta = \{ r_c, k_c, k_e, r_d, v_p, v_t \} \). The prediction is made on a basis of three hours \( \Delta t = 3 \) from 0 hour to 30 hours (corresponding to time step \( t = 1, 2, \ldots, 11 \)).

We denote the “true” hybrid model with underlying parameters \( \theta^a \). Following [15], we specify the true mechanistic parameter values as \( \phi^a = \{ r_c, k_c, k_e, r_d \} = \{ 0.057, 3.4, 2.6, 0.005 \} \). We set the bioprocess noise level \( v = v_p = v_t \), the initial cell density \( 3 \times 10^6 \) cells/ml \( (\rho_1 = 3) \), and no initial inhibition \( (I_1 = 0) \). Based on the simulation data generated by the true hybrid model, we assess the performance of the proposed LG-DBN auxiliary ABC-SMC algorithm under different levels of bioprocess noise \( v = \{ 0.1, 0.2 \} \) and model uncertainty (i.e., different sizes of process observations \( m = 3, 6, 20 \) batches).
5.2 LG-DBN Auxiliary ABC-SMC Performance Assessment

We compare the performance of LG-DBN auxiliary ABC-SMC with naive ABC-SMC (i.e. without using the LG-DBN as auxiliary model) in terms of: (1) prediction accuracy, (2) computation cost, and (3) posterior concentration. The distance metric of naive ABC-SMC is $d(\mathcal{D}, \mathcal{D}^*)$. The results are estimated based 30 macro-replications. We set the number of particles $N = 400$, the ratio $\alpha = 0.5$, the number of replications $L = 60$, and the minimal accept rate $P_{acc_{min}} = 0.15$. The prior distributions of model parameters are set as: $r_g \sim U(0, 0.5), k_i \sim U(0, 5), k_c \sim U(0, 5), r_d \sim U(0, 0.05), v_p \sim U(0, 0.2), \text{ and } v_l \sim U(0, 0.2)$.

One of the major benefits induced by the LG-DBN auxiliary likelihood is that it provides an efficient way to measure the distance between simulated and observed samples, which quickly leads to posterior samples fitting well on dynamics and variations. To show the advantage of LG-DBN auxiliary ABC-SMC, we first study its computational efficiency improvement. For each $r$-th macro replication, let $T_w^{(r)}$ and $T_{wo}^{(r)}$ represent the computation cost of the ABC-SMC algorithm with and without LG-DBN auxiliary. The computational efficiency improvement after incorporating the LG-DBN auxiliary model is evaluated as the time consuming ratio, i.e., $C^{(r)} = T_{wo}^{(r)} / T_w^{(r)}$. We record the 95% confidence interval (CI) for improvement, denoted by $\bar{C}$.

| Process Noise | $m = 3$ | $m = 6$ | $m = 20$ |
|---------------|---------|---------|----------|
| $v = 0.1$    | 1.27 ± 0.11 | 1.43 ± 0.11 | 2.44 ± 0.15 |
| $v = 0.2$    | 1.39 ± 0.08 | 1.52 ± 0.17 | 2.63 ± 0.20 |

Table 1: Computational efficiency improvement ratio.

Then, we compare the prediction accuracy of the posterior predictive distributions obtained from ABC-SMC with and without LG-DBN auxiliary. We estimate the parameters $\theta = (r_g, k_i, k_c, r_d, v_p, v_l)$. Specifically, in each macro replication, we generate posterior samples $\{\theta^{(i)}\}_{i=1}^{N_{\alpha}}$ by LG-DBN auxiliary and naive ABC-SMC approaches to approximate the posterior predictive distribution,

$$p(\rho_i, l_i | \rho_1, l_1, \mathcal{D}) = \int p(\rho_i, l_i | \theta, \rho_1, l_1) p(\theta | \mathcal{D}) d\theta = \frac{1}{N_{\alpha}} \sum_{i=1}^{N_{\alpha}} p(\rho_i, l_i | \rho_1, l_1, \theta^{(i)})$$

where the probability density $p(\rho_i, l_i | \rho_1, l_1, \theta^{(i)})$ is computed by the hybrid model (11) at $\theta^{(i)}$. Given the “true” model parameters $\theta^*$, we can also construct the predictive distribution $p(\rho_i, l_i | \rho_1, l_1, \theta^*)$ from the model (11). Figure 1 shows posterior predictive distributions of cell density and inhibitor concentration at 30 hours or timestep $t = 11$ given a fixed initial state $(\rho_1, l_1) = (3, 0)$. The black dashed line represents the predictive distribution of “true” model $p(\rho_{11, l1} | \rho_1, l_1, \theta^*)$.

Table 2: The K-S statistics of cell density and inhibitor accumulation at 30-th hour (i.e., $t = 11$).

| State | Process Noise | $m = 3$ | $m = 6$ | $m = 20$ | $m = 3$ | $m = 6$ | $m = 20$ |
|-------|---------------|---------|---------|----------|---------|---------|----------|
| $\rho_i$ | $v = 0.1$ | 0.34 ± 0.04 | 0.31 ± 0.03 | 0.25 ± 0.02 | 0.26 ± 0.05 | 0.24 ± 0.04 | 0.23 ± 0.03 |
|       | $v = 0.2$ | 0.25 ± 0.05 | 0.22 ± 0.04 | 0.19 ± 0.02 | 0.36 ± 0.04 | 0.32 ± 0.03 | 0.28 ± 0.02 |
| $I_i$   | $v = 0.1$ | 0.45 ± 0.04 | 0.46 ± 0.03 | 0.44 ± 0.02 | 0.68 ± 0.04 | 0.69 ± 0.03 | 0.67 ± 0.02 |
|       | $v = 0.2$ | 0.38 ± 0.05 | 0.37 ± 0.05 | 0.36 ± 0.04 | 0.53 ± 0.07 | 0.55 ± 0.06 | 0.56 ± 0.04 |

By comparing Figure 1(a) and 1(b) to Figure 1(c) and 1(d), we observe that DBN auxiliary ABC-SMC shows more robust performance across macro-replications and the posterior predictive distributions are generally closer to the “true” predictive distribution than naive ABC-SMC. We further investigate Panel 1(a) and 1(c). In low noise level $v = 0.1$, the auxiliary based ABC-SMC tends to overestimate the variance $\nu_p$ causing the estimated posterior predictive distributions more flat than the “true” predictive distribution. However, in high noise level, the posterior predictive distribution of LG-DBN auxiliary ABC-SMC is more accurate than that from naive ABC-SMC which consistently underestimates the variance $\nu_p$. The auxiliary ABC-SMC shows consistently better predictions of inhibitor concentration from Figure 1(d) and 1(d).
Figure 1: Posterior predictive distributions of cell density and inhibitor concentration at 30 hours ($t = 11$) $p(p, I_1 | p_1, I_1)$ obtained from 6 macro-replications (simulated with common random numbers). The color filled areas under the probability density curve represent estimated posterior predictive distributions in different macro-replications. The black dashed line represents the predictive distribution of the “true” model, i.e. $p(p, I_1 | p_1, I_1, \theta^*)$. The rows of each panel represent noise levels (i.e. $v = 0.1, 0.2$) while the columns of each panel are sample sizes of observations (i.e. $m = 3, 6, 20$). Panel (a) and Panel (b) represent auxiliary based ABC-SMC algorithm. Panel (c) and Panel (d) represent naive ABC-SMC algorithm.

We further use the Kolmogorov–Smirnov (K-S) statistics to assess the performance of LG-DBN auxiliary ABC-SMC and naive ABC-SMC. The K-S statistics quantifies the distance between the empirical distribution functions of the samples from posterior predictive distribution and predictive distribution of “true” model. The K-S statistics is $D = \sup_s |F^s - F^p|$, where $F^s$ and $F^p$ are the empirical distribution functions of the samples from predictive distribution of “true” model and posterior predictive distribution respectively. The smaller value of K-S statistic means better approximation performance of posterior predictive distribution. The number of sample used to construct empirical distribution function is set as $K = 2000$ in each macro-replication for both distributions. We summarize 95% CIs of distances for both cell density and inhibitor accumulation at 30-th hour, denoted by $\bar{D}$ and $S_D$ with $\bar{D} = \frac{1}{30}\sum_{r=1}^{30} D(r)$ and $S_D = [\sum_{r=1}^{30} (D(r) - \bar{D})^2 / 29]^{1/2}$.

As shown in Table 2, the LG-DBN auxiliary ABC-SMC algorithm has better performance in inhibitor concentration prediction at all levels of model estimation uncertainty and stochastic uncertainty. It also provides better prediction on cell density under high stochastic uncertainty, which is common in bioprocess. The results are consistent with the observations obtained from Figure 1. The performance improvement can be further observed from the estimated posterior distributions of hybrid model parameters; see the representative plots of cell growth rate $r_d$ and inhibitor decay $r_d$ in Figure 2. The posterior distributions estimated by LG-DBN auxiliary ABC-SMC has better concentration (i.e., how much the posterior mass is close to the true value $I_0^*$) than naive ABC-SMC in all noise levels and sample sizes. Especially, the better inference on $r_d$ is inspiring due to its extremely small but positive true value (i.e., $r_d^* = 0.005$) and also it is only involved in the ODE mechanistic model of latent state $I_1$; see Figure 1.

In sum, compared with naive ABC-SMC, the proposed LG-DBN auxiliary ABC-SMC algorithm tends to have better prediction accuracy and computational efficiency especially under the situations with high stochastic and model uncertainties. This can benefit bioprocess mechanism learning and robust control.
6 Conclusion

To leverage the information from existing mechanistic models and facilitate learning from real-world data, we develop a probabilistic knowledge graph (KG) hybrid model that can faithfully capture the important properties of integrated biomanufacturing processes, including nonlinear reactions, partially observed state, and nonstationary dynamics. Since the likelihood is intractable, approximate Bayesian computation (ABC) sampling strategy is used to generate samples to approximate the posterior distribution. For complex biomanufacturing processes with high stochastic and model uncertainties, it is computationally challenging to generate simulated trajectories close to real-world observations. Therefore, in this paper, we utilize a simple linear Gaussian dynamic Bayesian network (LG-DBN) auxiliary model to design summary statistics for ABC-SMC that can accelerate Bayesian inference on the probabilistic KG hybrid model with high fidelity characterizing complex bioprocessing mechanisms. The empirical study demonstrates that the proposed LG-DBN auxiliary ABC-SMC can improve computational efficiency and prediction accuracy. In the future research, we will extend this approach to complex multi-scale bioprocess hybrid model in order to facilitate metabolic mechanism learning and support robust control on both cellular and system levels.

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