TRANSPORT MAP ACCELERATED ADAPTIVE IMPORTANCE SAMPLING, AND APPLICATION TO INVERSE PROBLEMS ARISING FROM MULTISCALE STOCHASTIC REACTION NETWORKS

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Abstract. In many applications, inverse problems arise where where there are complex correlations between the different parameters which we wish to infer from data. The correlations often manifest themselves as lower dimensional manifolds on which the likelihood function is invariant, or varies very little. This can be due to trying to infer unobservable parameters, or due to sloppiness in the model which is being used to describe the data. In such a situation, standard sampling methods for characterising the posterior distribution which do not incorporate information about this structure will be highly inefficient. Moreover, most methods are inherently serial in nature, and as such are not expoiting the parallelised nature of modern computer infrastructure. In this paper, we seek to develop a method to tackle this problem, using optimal transport maps to simplify posterior distributions which are concentrated on lower dimensional manifolds.

We demonstrate the approach by considering inverse problems arising from partially observed stochastic reaction networks. In particular, we consider systems which exhibit multiscale behaviour, but for which only the slow variables in the system are observable. We demonstrate that certain multiscale approximations lead to more consistent approximations of the posterior than others.

1. Introduction. The importance of Markov chain Monte Carlo (MCMC) methods is becoming increasingly apparent, in a world replete with datasets which need to be combined with complex dynamical systems in order for us to be able make progress in a range of scientific fields. Different MCMC methods have been designed with different challenges in mind; high dimensional targets****CITE FASTER PA-PER****, large data sets ****zigzag****, other methods which exploit piecewise deterministic Markov processes ***bouncey particle***, methods which are able to make very large moves informed by the data****stochastic mala, HMC****, methods which are able to exploit the geometry of the target***riemannian****. Population Monte Carlo (pMC) methods uses efficient importance sampling proposals informed by an ensemble of particles which have already learned about the structure of the target***cite standard papers including AIS etc****. These methods are particularly useful in the context of low dimensional target distributions which have complex structure, for which traditional Metropolis-Hastings methods tend to struggle, for example when the target is multimodal or if the density is concentrated on a lower-dimensional manifold. Moreover, they are an excellent candidate for parallelisation, since the task of computing the likelihood for each particle in the ensemble can be distributed across a set of processors. This is of utmost importance in an age where increases in computational power are most likely to come from large-scale parallelisation as opposed to the large speed-ups in individual processors that we saw throughout the 20th Century. In ****cite PAIS***, we explored ideas from adaptive MCMC within a pMC-type

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structure, in order to construct an adaptive importance sampler which could automatically adapt to optimise efficiency. Alongside this, we explored the use of state-of-theart resamplers****cite sebastian***, and a greedy approximation (the approximate multinomial resampler) in order to improve the quality of the importance sampler proposals.

Other approaches to challenging posterior structure have been proposed, including the work of Marzouk and Parnod***cite***, who demonstrated how invertible transport maps can be constructed which map the target distribution close to an easily-explored reference density, such as a d-dimensional Gaussian. Since the map is invertible, Gaussian proposal densities on the reference space can be mapped to highly informed moves on the parameter space.

In this paper, we particularly concern ourselves with the problem of inverse problems in stochastic reaction networks. This type of model is used in situations where ODE or diffusion****Chem langevin**** approximations of chemical reaction kinetics are not viable, usually because of the low concentration of one or more chemical species****Gillespie****. Biochemical networks typically fall into this category, because of the small volume of the reactors involved (a cell).

One challenge with networks arising from these applications is that they often have some reactions which are occurring many times on a timescale in which other reactions in the system are unlikely to occur. This multiscale behaviour throws up two main challenges, both in observability in an experimental setting, but also in the context of modelling, where multiscale systems can make the simulation of the continuous time Markov chain computationally intractable.

A range of numerical methods have been suggested to tackle this latter problem, including tau-leaping***cite****, diffusion approximations ***radek and Yannis, cotter and Yannis****, quasi-steady state approximations ****slow scale and nested****, and other averaging arguments ****cotter comp. phys****. Some of these approaches are more computationally intensive than others, but this can be mitigated with the use of analytical results about the invariant densities of certain types of fast subsystems***product-form papers****.

We would not expect the fast processes in such a system to be experimentally observable. In such a situation where only the slow variables in a system are identifiable, in an ideal world, we would be able to integrate over the space of all possible trajectories of the fast processes. Since this is an insurmountable computational task, we might instead wish to approximate the likelihoods arising from such a set of observations by the use of an appropriate multiscale methodology.

The partial observability of these systems leads to only partial-observability of the unknown parameters in the system, for example the reaction rates of the fast reactions. The resulting multiscale approximations, and the resulting approximations of the likelihoods arising from the inverse problem for such a network, then often exhibit insensitivity to changes in certain directions of parameter space. This causes problems when aiming to sample from such a target, since the density is concentrated on a manifold whose tangential vectors point in directions of insensitivity. This sloppiness***cite*** of the likelihood is a challenging problem that appears in many applications in science and engineering***active subspaces literature here****.

In this work, we aim to improve the PAIS machinery, which is limited, as the same proposal kernel is used for each particle in the ensemble. This homogeneity of proposal kernels means that complex structures in the density can only be well represented with a large increase in the ensemble size, increasing computational cost. If computa-

tional resources do not allow for such an increase, this can lead to stability issues in the algorithm, due to the proposal distribution in the importance sampler not representing well the target density, leading to high variance in the weights, causing slow convergence.

As shown in ***cite marzouk****, transport maps can be used to create proposal densities which are highly informed by the structure of the target density. In this paper, we demonstrate how such maps can also be constructed using importance weights arising from a pMC-type algorithm in order to accelerate and stabilise convergence. We will then apply the methodology to some complex posterior distributions arising from Bayesian inversion of multiscale (bio-)chemical networks.

In Section 2, we will briefly reintroduce PAIS. In Section 3 we show how an appropriate transport map can be constructed from importance samples which maps the posterior close to a reference Gaussian measure. In Section 4 we show how such a map can be incorporated into a sophisticated parallel MCMC infrastructure in order to accelerate mixing, and improve stability. In Section 5 we seek to show the advantages of this approach through the analysis of a challenging test problem. In Section 6 we consider how likelihoods can be approximated using multiscale methodologies in order to carry out inference for multiscale and/or partially observed stochastic reaction networks. In Section 7 we present some numerical examples, which serve to demonstrate the increased efficiency of the described sampling methodologies, as well as investigating the posterior approximations discussed in the previous section. We conclude with a discussion in Section 8.

2. Parallel Adaptive Importance Sampling. Parallel adaptive importance sampling (PAIS) [9] is a variant of PMC [7], which is a family of methods which are based on importance sampling. In importance sampling, we attempt to characterise a target density through sampling from that density. However, the target density π itself is often too complex to sample from directly, so we instead sample from a proposal density χ . Each sample $\theta^{(k)} \sim \chi$ is then weighted by $w_k = \frac{\pi(\theta^{(k)})}{\chi(\theta^{(k)})}$, to take account of the bias of sampling from a different distribution to π . Monte Carlo estimates using a sample of size N of a function f with respect to π can then be made through the formula

$$\mathbb{E}_{\pi}(f) \approx \frac{1}{\bar{w}} \sum_{k=1}^{N} w_k f(\theta^{(k)}).$$

This method works well when π and χ are close, but can be excruciatingly slow when they are not. The idea behind PMC methods is to construct a good proposal distribution, either from the entire history of the algorithm up to the current point, or to use the current state of a whole ensemble of M particles in the system.

In PAIS, the proposal distribution χ is chosen to be the equally weighted mixture of any choice of MCMC proposal kernel, evaluated at each of the current particles in the system. If $\theta^{(k)} = [\theta_1^{(k)}, \theta_2^{(k)}, \dots, \theta_M^{(k)}]^{\top}$ is the current state of the ensemble, and we wish to use an MCMC proposal density $q(\cdot; \cdot, \beta)$ then

$$\chi^{(k)} = \frac{1}{M} \sum_{i=1}^{M} q(\cdot; \theta_i^{(k)}.$$

Often the variance of the MCMC proposal kernels can be tuned using their respective algorithmic parameters β . Good values for these algorithmic parameters can be found

by optimising for the effective sample size of the importance sample that is produced (see [9, 29] for more details).

If the ensemble is large enough, and the chain has entered probabilistic stationarity, then the current state of the ensemble is a good rough discrete approximation of the target density, and in turn $\chi^{(k)}$ is close enough to π to produce an efficient importance sample $\{\hat{\theta}^{(k)}, w^{(k)}\}$. It can be advantageous to use stratified sampling of the mixture in order to ensure that the sample made is as representative as possible of the target density, i.e.

$$\hat{\theta}_i^{(k)} \sim q(\cdot; \theta_i^{(k)})$$

for each $i=1,2,\ldots,M$. We now have a weighted sample, and it would be inadvisable to use an equally weighted mixture of proposal distributions from each of these points. Therefore, before starting the next iteration, the importance sample $\{\hat{\theta}^{(k)}, w^{(k)}\}$ is resampled to produce an equally weighted sample, ready for the next iteration of the algorithm. In PAIS, a state-of-the-art resampler is used, which uses optimal transport methods to find the equally weighted discrete sample which best represents the statistics of the importance sample [27]. For larger ensemble sizes M this can become expensive, in which case a greedy approximation of this algorithm, the Approximate Multinomial Resampler (AMR) can be implemented [9]. The output of the resampler is then denoted $\theta^{(k+1)}$ and the algorithm is ready for the next iteration. The importance samples $\{\hat{\theta}^{(k)}, w^{(k)}\}$ The PAIS is summarised in Algorithm 1.

Algorithm 1: The PAIS Algorithm.

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 \begin{array}{ll} \text{1 Initialise } \theta^{(1)} \sim \mu_0. \\ \textbf{2 for } k = 1, \dots, N \text{ do} \\ \textbf{3} & \text{Sample } \hat{\theta}_i^{(k)} \sim q(\cdot; \theta_j^{(k)}, \beta), \text{ for } i = 1, \dots, M. \\ \textbf{4} & \text{Calculate } w^{(k)} = (w_1^{(k)}, \dots, w_M^{(k)})^\top, \text{ where} \\ & w_i^{(k)} = \frac{\mu(\hat{\theta}_i^{(k)})}{\chi^{(k)}(\hat{\theta}_i^{(k)}; \theta^{(k)}, \beta)}. \\ \textbf{5} & \text{Resample } \theta^{(k+1)} \leftarrow \|w^{(k)}\|_1^{-1} \sum_{j=1}^M w_j^{(k)} \delta_{\hat{\theta}_j^{(k)}}(\cdot). \\ \textbf{6} & \text{Output } \{(w^{(n)}, \hat{\theta}^{(n)})\}_{n=1}^N. \end{array}
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One problem with the PAIS and other PMC methods can become apparent if the target density has very strong correlations in its structure, in particular if that correlation is not global but only local. In this case, unless the proposal densities q are informed by this local structure, the mixture distribution proposal may not well approximate π without a very large large ensemble size M, which can become inhibitively expensive. Some methods have been proposed [13] which use samples local to each particle to inform local covariance structure.

In this paper, we investigate the use of transport maps to learn local covariances across the whole of the domain, in order to stabilise PMC-type methods, and make these methods more applicable to a wider range of more challenging inference problems.

3. Construction of transport maps in importance sampling. In this Section, we describe the construction of transport maps which allow for the simplification

of complex posterior distributions in order to allow for improved sampling, in particular for methods based on importance sampling. In [14] the transport map was introduced to provide a transformation from the prior distribution to the posterior distribution, the idea being that one could draw a moderately sized sample from the prior distribution and use this sample to approximate a map onto the target space. Once this map was known to the desired accuracy a larger sample from the prior could be used to investigate the posterior distribution. This methodology was adapted in [25] to form a new proposal method for MH algorithms. In this case, rather than transforming a sample from the prior into a sample from the target distribution, the map transforms a sample from the posterior onto a reference space. The reference density is chosen to allow efficient proposals using a simple proposal distribution such as a Gaussian centred at the previous state. Proposed states can then be mapped back into a sample from the posterior by applying the inverse of the transport map.

Proposing new states in this way allows us to make large steps around complex probability distributions. It is also feasible in this framework to assume that the reference density is close enough to a standard Gaussian that we can efficiently propose moves using a proposal distribution which is independent of the current state, e.g. choose $q(\theta) = \mathcal{N}(0, I_n)$.

In this Section we outline the methodology in [25] for approximately coupling the target, μ_{θ} , with the reference distribution, μ_{r} , and show how the map can be constructed using a weighted sample and hence how we can incorporate the map into importance sampling schemes.

DEFINITION 3.1 (Transport Map T). A transport map T is a function $T: \mathcal{X} \to \mathbb{R}^d$ such that the pullback of the reference measure with density $\phi(\cdot)$,

$$\tilde{\pi}(\theta) = \phi(T(\theta))|J_T(\theta)|,\tag{3.1}$$

is equal to the target density $\pi(\theta)$ for all $\theta \in \mathcal{X}$. The pullback is defined in terms of the determinant of the Jacobian of T,

$$|J_T(\theta)| = \det \begin{bmatrix} \partial_{\theta_1} T_1(\theta) & \dots & \partial_{\theta_d} T_1(\theta) \\ \vdots & \ddots & \vdots \\ \partial_{\theta_1} T_d(\theta) & \dots & \partial_{\theta_d} T_d(\theta) \end{bmatrix}.$$

DEFINITION 3.2 (Target and Reference Space). The transport map pushes a particle from a target space \mathcal{X} , that is a subset of \mathbb{R}^d equipped with a target measure μ_{θ} , onto a reference space, R, again a subset of \mathbb{R}^d equipped with the reference measure μ_r .

Armed with such a map, independent samples can be made of the target measure, using the pullback of the reference density ϕ through T^{-1} . Clearly the pullback only exists when T is monotonic, i.e. has a positive definite Jacobian, and has continuous first derivatives. Not all maps satisfy these conditions, so we define a smaller space of maps, $\mathcal{T}^{\uparrow} \subset \mathcal{T}$ which contains all feasible maps. This space does not necessarily contain an exact coupling between target and reference space, and so we are motivated to formulate an optimisation problem to find the map $\tilde{T} \in \mathcal{T}^{\uparrow}$ which most closely maps the target density to the reference density.

As in previous work in [25], we can ensure invertibility if we restrict the map to be

lower triangular, i.e. $\tilde{T} \in \mathcal{T}^{\wedge} \subset \mathcal{T}^{\uparrow}$. This lower triangular map has the form,

$$T(heta_1,\ldots, heta_n) = egin{bmatrix} T_1(heta_1) \ T_2(heta_1, heta_2) \ dots \ T_n(heta_1,\ldots, heta_n) \end{bmatrix},$$

where $T_i : \mathbb{R}^i \to \mathbb{R}$.

3.1. The optimisation problem. Our aim is now to find the lower triangular map $\tilde{T} \in \mathcal{T}^{\triangleright}$ such that the difference between target density and the pullback of the reference density is minimised. As in [25], we choose the cost function to be the Kullback-Leibler (KL) divergence between the posterior density and the pullback density,

$$D_{\mathrm{KL}}(\pi \| \tilde{\pi}) = \mathbb{E}_{\pi} \left[\log \left(\frac{\pi(\theta)}{\tilde{\pi}(\theta)} \right) \right].$$

This divergence results in some nice properties which we will explore in the following derivation. The KL divergence is not a true metric since it is not symmetric, however it is commonly used to measure the distance between probability distributions due to it's relatively simple form, and because it provides a bound for the square of the Hellinger distance by Pinsker's inequality [26],

$$D_{KL}(p||q) \ge D_H^2(p,q),$$

which is a true metric between probability distributions p and q. Given the form of the pullback in Equation (3.1), now taken through an approximate map \tilde{T} , the divergence becomes

$$D_{\mathrm{KL}}(\pi \| \tilde{\pi}) = \mathbb{E}_{\pi} \left[\log \pi(\theta) - \log \pi_r(\tilde{T}(\theta)) - \log |J_{\tilde{T}}(\theta)| \right].$$

We note the posterior density is independent of \tilde{T} , and so it is not necessary for us to compute it when optimising this cost function. This expression is a complicated integral with respect to the target distribution, for which the normalisation constant is unknown. However this is exactly the scenario for which we would turn to MCMC methods for a solution.

To find the best coupling, $\tilde{T} \in \mathcal{T}^{\mathbb{A}}$, we solve the optimisation problem,

$$\tilde{T} = \arg\min_{T \in \mathcal{T}_{k}} \mathbb{E}_{\pi} \left[-\log \pi_{r}(T(\theta)) - \log |J_{T}(\theta)| \right]$$

which has a unique solution since the cost function is convex. We also include a regularisation term, which is required for reasons which will become clear later. The optimisation problem now takes the form

$$\tilde{T} = \arg\min_{T \in \mathcal{T}^{\perp}} \left[\mathbb{E}_{\pi} \left[-\log \pi_r(T(\theta)) - \log |J_T(\theta)| \right] + \beta \mathbb{E}(T(\theta) - \theta)^2 \right]. \tag{3.2}$$

The parameter $\beta > 0$ does not need to be tuned, as experimentation has shown that the choice $\beta = 1$ is sufficient for most problems. This expectation can be approximated by using an MCMC approximation. The form of the penalisation term promotes maps which are closer to the identity, and so prevents overfitting when the quality or size of the current sample from the posterior is not sufficient.

3.2. The structure of the map. Before we continue with the derivation of the optimisation problem, we consider the structure of the map in more detail. The lower triangular structure of the map not only guarantees monotonicity, it also allows for efficient calculation of the pullback density, as well as the inverse of the map, \tilde{T}^{-1} . The Jacobian of \tilde{T} is a lower triangular matrix,

$$J_{T}(\theta) = \begin{bmatrix} \partial_{\theta_{1}} \tilde{T}_{1}(\theta) & \dots & \partial_{\theta_{d}} \tilde{T}_{1}(\theta) \\ \vdots & \ddots & \vdots \\ \partial_{\theta_{1}} \tilde{T}_{d}(\theta) & \dots & \partial_{\theta_{d}} \tilde{T}_{d}(\theta) \end{bmatrix} = \begin{bmatrix} \partial_{\theta_{1}} \tilde{T}_{1}(\theta) & \dots & 0 \\ \vdots & \ddots & \vdots \\ \partial_{\theta_{1}} \tilde{T}_{d}(\theta) & \dots & \partial_{\theta_{d}} \tilde{T}_{d}(\theta) \end{bmatrix}$$

since $\partial_{\theta_n} \tilde{T}_k(\theta) = 0$ for all n > k. This lower triangular structure means that the determinant of the Jacobian is a product of the diagonal elements which, when we take logs, becomes

$$\log |J_{\tilde{T}}(\theta)| = \sum_{i=1}^{d} \log \partial_{\theta_i} \tilde{T}_i(\theta), \tag{3.3}$$

where we note that this term is separable in terms of the dimension i.

Inverting \tilde{T} at a point r is simplified by the lower triangular structure of the map. The map component $\tilde{T}_1(\theta)$ is a univariate polynomial in θ_1 , so we can find the inverse of this function by solving the equation $T_1(\theta_1) = r_1$. This inversion tells us the value of θ_1 , which means the next component is again a univariate polynomial, $T_2(\theta_2; \theta_1) = r_2$. We can then perform d root finding problems instead of a full d dimensional non-linear solve.

We require that the first derivatives of the map are continuous, which is easy to enforce by the choice of basis functions. Here we assume that the map will be built from a family of orthogonal polynomials, $\mathcal{P}(\theta)$, not necessarily orthogonal with respect to the target distribution. Each component of the map is defined as a multivariate polynomial expansion,

$$\tilde{T}_i(\theta; \gamma_i) = \sum_{\mathbf{j} \in \mathcal{J}_i} \gamma_{i, \mathbf{j}} \psi_{\mathbf{j}}(\theta). \tag{3.4}$$

The parameter $\gamma_i \in \mathbb{R}^{M_i}$ is a vector of coefficients. Each component of γ_i corresponds to a basis function $\psi_{\mathbf{j}}$, indexed by the multi-index $\mathbf{j} \in \mathbb{N}_0^d$. These multi-indices are elements of the multi-index set \mathcal{J}_i . A multi-index defines a product of univariate polynomials in θ_k ,

$$\psi_{\mathbf{j}}(\theta) = \prod_{k=1}^{i} \varphi_{j_k}(\theta_k), \quad \text{for} \quad \mathbf{j} \in \mathcal{J}_i,$$

and where $\varphi_{j_k}(\theta_k) \in \mathcal{P}(\theta_k)$. Since T is lower triangular, a multi-index $\mathbf{j} \in \mathcal{J}_i$ only contains entries for univariate polynomials in θ_k for $k \leq i$.

The cardinalities of the multi-index sets, $M_i = \text{card}(\mathcal{J}_i)$, give the number of unknowns in our optimisation problem, and so we would like to keep this number as small as possible. One option is to use polynomials of total order p,

$$\mathcal{J}_i^{\text{TO}} = \left\{ \mathbf{j} : \|\mathbf{j}\|_1 \le p, j_k = 0 \ \forall k > i \right\},\,$$

which is optimal in terms of the amount of information captured by the map about the target. The cardinality of $\mathcal{J}_i^{\text{TO}}$ is $M_i = \binom{i+p}{p}$ which increases rapidly in d and p,

where i = 1, ..., d. Smaller optimisation problems can be produced by constructing subsets of $\mathcal{J}_i^{\text{TO}}$. These index sets are discussed in [25]. Increased information with a slower increase in the number of map parameters can be achieved with the composition of maps discussed in [24]. Here we stick with polynomials of total order p since we work with low dimensional problems with the PAIS algorithm.

3.3. Implementation of the optimisation problem. We now discuss how we can evaluate the cost function in Equation (3.2). In [25], this expectation is approximated using an MCMC estimator, such that

$$C(T) = \mathbb{E}_{\pi} \left[-\log \pi_r(T(\theta)) - \log |J_T(\theta)| \right] + \beta \mathbb{E}(T(\theta) - \theta)^2$$

$$\approx \frac{1}{K} \sum_{i=1}^{d} \sum_{k=1}^{K} \left[-\log \pi_r(T_i(\theta^{(k)})) - \log \left| \frac{\partial T_i}{\partial \theta_i}(\theta^{(k)}) \right| + \beta (T_i(\theta^{(k)}) - \theta^{(k)})^2 \right]. \quad (3.5)$$

Here we diverge from previous work, as we aim to build a map from samples from an importance sampling scheme. Such samples no longer carry equal weight, and as such the Monte Carlo estimator becomes

$$C(T) = \frac{1}{\bar{w}} \sum_{i=1}^{d} \sum_{k=1}^{K} w_k \left[-\log \pi_r(T_i(\theta^{(k)})) - \log \left| \frac{\partial T_i}{\partial \theta_i}(\theta^{(k)}) \right| + \beta (T_i(\theta^{(k)}) - \theta^{(k)})^2 \right],$$
(3.6)

where w_k are the weights associated with each sample $\theta^{(k)}$, and \bar{w} is the sum of all these weights. Optimisation of this cost function results in a map from π to some reference density π_r . By choosing the reference density to be a Gaussian density, we can simplify this expression greatly. Substitution of the Gaussian density into Equation (3.6) leads to

$$C(T) = \frac{1}{\bar{w}} \sum_{i=1}^{d} \sum_{k=1}^{K} w_k \left[\frac{1}{2} T_i^2(\theta^{(k)}) - \log \frac{\partial T_i}{\partial \theta_i} (\theta^{(k)}) + \beta (T_i(\theta^{(k)}) - \theta^{(k)})^2 \right], \quad (3.7)$$

Note that since we assume that the map is monotonic, the derivatives of each component are positive and so this functional is always finite. In practice it is infeasible to enforce this condition across the whole parameter space. We instead enforce this condition by ensuring that the derivatives are positive at each sample point. This means that when we sample away from these support points while in reference space, it is possible to enter a region of space where the map is not monotonic.

We now return to the structure of the map components given in Equation (3.4). Since the basis functions are fixed, the optimisation problem in (3.2) is really over the map components $\bar{\gamma} = (\gamma_1, \dots, \gamma_d)$ where $\gamma_i \in \mathbb{R}^{M_i}$. Note that C(T) is the sum of d expectations, and these expectations each only concern one dimension. Therefore we can rewrite (3.2) as d separable optimisation problems.

$$\arg\min_{\gamma_i \in \mathbb{R}^{M_i}} \frac{1}{\bar{w}} \sum_{k=1}^K w_k \left[\frac{1}{2} T_i^2(\theta^{(k)}; \gamma_i) - \log \frac{\partial T_i}{\partial \theta_i} (\theta^{(k)}; \gamma_i) + \beta (T_i(\theta^{(k)}; \gamma_i) - \theta^{(k)})^2 \right], \tag{3.8}$$

subject to
$$\frac{\partial T_i}{\partial \theta_i}(\theta^{(k)}; \gamma_i) > 0$$
 for all $k = 1, \dots, K, i = 1, \dots, d$.

The sum in Equation (3.4) is an inner product between the vector of map coefficients, and the evaluations of the basis function at a particular $\theta^{(k)}$. If we organise our basis

evaluations into two matrices,

$$(F_i)_{k,\mathbf{j}} = \psi_{\mathbf{j}}(\theta^{(k)}), \quad \text{and} \quad (G_i)_{k,\mathbf{j}} = \frac{\partial \psi_{\mathbf{j}}}{\partial \theta_i}(\theta^{(k)}),$$

for all $\mathbf{j} \in \mathcal{J}_i^{\text{TO}}$, and $k = 1, \dots, K$, then we have that

$$T_i(\theta^{(k)}) = (F_i)_k \cdot \gamma_i$$
 and $\frac{\partial T_i}{\partial \theta_i}(\theta^{(k)}; \gamma_i) = (G_i)_k \cdot \gamma_i$,

so (3.8) becomes

$$\arg\min_{\gamma_i \in \mathbb{R}^{M_i}} \frac{1}{2} (F_i \gamma_i)^\top W(F_i \gamma_i) - \mathbf{w}^\top \log(G_i \gamma_i) + \frac{\beta}{\bar{w}} \sum_{k=1}^K w_k (F_i \gamma_i - \theta^{(k)})^\top (F_i \gamma_i - \theta^{(k)}),$$
(3.9)

subject to $G_i \gamma_i > 0$.

In this expression, the vector $\mathbf{w} = [w_1, w_2, \dots, w_K]^{\top}$ is the vector of the weights, W is the diagonal matrix $W = \operatorname{diag}(w)$ and $\operatorname{log}(G_i\gamma_i)$ is to be evaluated element-wise. As more importance samples are made, new rows can be appended to the F_i and G_i matrices, and $F_i^{\top}WF_i$ can be efficiently updated via the addition of rank-1 matrices. The regularisation term in Equation (3.9) can be approximated using Parseval's identity,

$$\frac{1}{\bar{w}} \sum_{k=1}^{K} w_k (F_i \gamma_i - \theta^{(k)})^{\top} (F_i \gamma_i - \theta^{(k)}) \xrightarrow[K \to \infty]{} \int_{\mathbb{R}^n} |T(\theta) - \theta|^2 d\mu_{\theta} = \sum_{\mathbf{j} \in \mathcal{J}_i^{\text{TO}}} (\gamma_{i,\mathbf{j}} - \iota_{\mathbf{j}})^2,$$

where ι is the vector of coefficients for the identity map. This is of course only true when the polynomial family $\mathcal{P}(\theta)$ is chosen to be orthonormal with respect to μ_{θ} ; however this approximation prevents the map from collapsing onto a Dirac when the expectation is badly approximated by a small number of samples.

These simplifications result in the efficiently implementable, regularised optimisation problem for computing the map coefficients,

$$\arg \min_{\gamma_i \in \mathbb{R}^{M_i}} \frac{1}{2\bar{w}} \gamma_i^\top F_i^\top W F_i \gamma_i - \frac{w^\top}{\bar{w}} \log(G_i \gamma_i) + \beta \|\gamma_i - \iota\|^2,$$
subject to $G_i \gamma_i > 0$, (3.10)

This optimisation problem can be efficiently solved using Newton iterations. It is suggested in [25] that this method usually converges in around 10-15 iterations, and we have seen no evidence that this is not a reasonable estimate. When calculating the map several times during a Monte Carlo run, using previous guesses of the optimal map to seed the Newton algorithm results in much faster convergence, usually taking only a couple of iterations to satisfy the stopping criteria.

The Hessian takes the form

$$HC_i(\gamma_i) = \frac{1}{\bar{w}} \left[F_i^\top W F_i + G_i^\top W \operatorname{diag}([G_i \gamma_i]^{-2}) G_i \right] + \beta I, \tag{3.11}$$

where $[G_i\gamma_i]^{-2}$ is to be taken element-wise, and I is the $M_i \times M_i$ identity matrix. The first derivative of $C_i(T)$ is

$$\nabla C_i(\gamma_i) = \frac{1}{\bar{w}} \left[F_i^\top W F_i \gamma_i - G_i^\top W [G_i \gamma_i]^{-1} \right] + \beta (\gamma_i - \iota),$$

again $[G_i\gamma_i]^{-1}$ is taken element-wise.

Algorithm 2: PAIS algorithm with adaptive transport map. Option 1.

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1 Initialise state \theta_i^{(1)} = \theta_0, \quad i = 1, \dots, M.
  2 Initialise map \bar{\gamma}^{(1)} = \iota.
  3 for k \leftarrow 1, \ldots, L-1 do
              Compute r_i = \tilde{T}(\theta_i^{(k)}; \bar{\gamma}^{(k)}), \quad i = 1, \dots, M.
Sample r_i' \sim q_r(\cdot; r_i).
Invert \hat{\theta}_i^{(k)} = \tilde{T}^{-1}(r_i'; \bar{\gamma}^{(k)}).
  4
  5
  6
                                                     w_i^{(k)} = \frac{\pi(\hat{\theta}_i^{(k)})}{\left(\sum_{j=1}^{M} q_r(r_i'; r_j)\right) |J_{\tilde{T}}(\hat{\theta}_i^{(k)}; \bar{\gamma}^{(k)})|}.
              Resample \theta^{(k+1)} \leftarrow \|w^{(k)}\|^{-1} \sum\limits_{i=1}^{M} w_j^{(k)} \delta_{\hat{\theta}_i^{(k)}}(\cdot).
  8
               if k \mod K_U = 0 and k < K_{stop} then
  9
                       for i \leftarrow 1, \ldots, n do
10
                              Solve (3.10) with \{(w^{(1)}, \hat{\theta}^{(1)}), \dots, (w^{(k+1)}, \hat{\theta}^{(k+1)})\} and update
11
12
                   \bar{\gamma}^{(k+1)} = \bar{\gamma}^{(k)}.
13
```

4. Transport map usage in PAIS and other PMC algorithms. Given importance samples from the target distribution, we have demonstrated how to construct an approximate transport map from the target measure to a reference measure. We now consider how to implement an importance sampling-based MCMC algorithm which uses these maps to propose new states. In [25] it was shown how approximate transport maps can be used to accelerate Metropolis-Hastings methods, with the map being periodically updated with the samples produced from the target measure. Convergence of this adaptation is shown in [25]. In this Section, we will show how similarly, these maps can be used to construct highly efficient importance sampling schemes.

In particular, we will show how we can use the transport map derived in Equation (3.10) to design a proposal scheme for the PAIS algorithm. In this case we have a choice in how to proceed; we propose new samples on reference space and resample on target space, or we both propose and resample on reference space, mapping onto target space to output the samples. The first option allows us to reuse much of the framework from the standard PAIS algorithm and in the numerics later we see that this performs better than both the Transport MH algorithm, and the standard PAIS algorithm. The second option requires some restructuring but results in improved performance from the resampler.

The first option is given in Algorithm 2. We denote the ensembles of states in target space $\theta^{(k)} = \{\theta_1^{(k)}, \dots, \theta_M^{(k)}\}$, and the states in the reference space, $r = \{r_1, \dots, r_M\}$, where M is the ensemble size. Similarly, the proposal states are denoted $r' = \{r'_1, \dots, r'_M\}$ and $(w^{(k)}, \hat{\theta}^{(k)}) = \{(w_1^{(k)}, \hat{\theta}^{(k)}_1), \dots, (w_M^{(k)}, \hat{\theta}^{(k)}_M)\}$, where these pairs are

Algorithm 3: PAIS algorithm with adaptive transport map. Option 2.

```
1 Initialise state \theta_i^{(1)} = \theta_0, \quad i = 1, \dots, M.
  2 Initialise map \bar{\gamma}^{(1)} = \iota.
  3 for k \leftarrow 1, \ldots, N-1 do
             Compute r_i = \tilde{T}(\theta_i^{(k)}; \bar{\gamma}^{(k)}), \quad i = 1, \dots, M.
  4
             Sample r'_i \sim q_r(\cdot; r_i).

Invert \hat{\theta}_i^{(k)} = \tilde{T}^{-1}(r'_i; \bar{\gamma}^{(k)}).
  5
  6
                                               w_i^{(k)} = \frac{\pi(\hat{\theta}_i^{(k)})}{\left(\sum_{j=1}^{M} q_r(r_i'; r_j)\right) |J_{\tilde{T}}(\hat{\theta}_i^{(k)}; \bar{\gamma}^{(k)})|}.
             Resample r^* \leftarrow \|w^{(k)}\|^{-1} \sum_{j=1}^M w_j^{(k)} \delta_{r'_j}(\cdot).
  8
             Invert \theta_i^{(k+1)} = \tilde{T}^{-1}(r_i^*).
  9
             if k \mod K_U = 0 and k < K_{stop} then
10
                    for i \leftarrow 1, \ldots, n do
11
                           Solve (3.10) with \{(w^{(1)}, \hat{\theta}^{(1)}), \dots, (w^{(k+1)}, \hat{\theta}^{(k+1)})\} and update
12
13
                \bar{\gamma}^{(k+1)} = \bar{\gamma}^{(k)}.
```

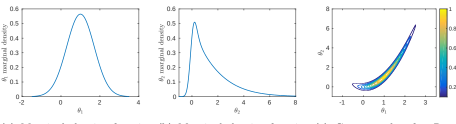
the states which together form our sample from the target distribution. As in the standard version of the PAIS algorithm we use the deterministic mixture weights. The second option, Algorithm 3, is similar to the first except on Line 8 where rather than resampling in target space we resample in reference space. In reference space the dimensions are roughly uncorrelated, and the Gaussian marginals are easy to approximate with fewer ensemble members. This means that the resampling step will be more efficient in moderately higher dimensions, which we discuss in Section 8.1.

5. Convergence of the transport proposal based MCMC algorithms. In this Section we study the convergence of the transport based proposal distributions which we have described in Section 4. We take as a test problem the ubiquitous Rosenbrock banana-shaped density. This target density is given by

$$\pi(\theta) = \frac{\sqrt{10}}{\pi} \exp\left\{-(1-\theta_1)^2 - 10(\theta_2 - \theta_1^2)^2\right\}. \tag{5.1}$$

A contour plot of the target density is given in Figure 5.1. This problem is challenging to sample from since it has a highly peaked and curved ridge, and is often used as a test problem in optimisation and MCMC communities.

5.1. Implementation details. Before looking at the performance of the MCMC algorithms, we demonstrate some properties of the transport maps we will be using in our MCMC algorithms. We draw 1 million samples from the density in (5.1), and use this sample in the framework of Section 3 to build a transport map. We use this map



(a) Marginal density function (b) Marginal density function (c) Contour plot for Rosenfor θ_2 .

Fig. 5.1. Visualisation of the Rosenbrock density as given in Equation (5.1).

to push forward the original sample onto the reference space, where we will be able to see how well the map has performed at converting the original sample to a standard Gaussian. We then pull the sample back on to target space using the inverse map to check that our map is invertible and well behaved.

For this example, we use an index set of total order 3 with monomial basis functions. It is important that total order is an odd number, since otherwise the map will not be surjective. This results in a map of the form

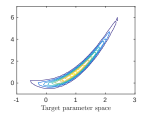
$$T(\theta_1, \theta_2) = \begin{bmatrix} T_1(\theta_1) \\ T_2(\theta_1, \theta_2) \end{bmatrix},$$

where

$$T_1(\theta_1) = \gamma_{1,1} + \gamma_{1,2}\theta_1 + \gamma_{1,3}\theta_1^2 + \gamma_{1,4}\theta_1^3,$$

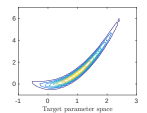
$$T_2(\theta_1, \theta_2) = \gamma_{2,1} + \gamma_{2,2}\theta_1 + \gamma_{2,3}\theta_1^2 + \gamma_{2,4}\theta_1^3 + \gamma_{2,5}\theta_2 + \gamma_{2,6}\theta_1\theta_2 + \gamma_{2,7}\theta_1^2\theta_2 + \gamma_{2,8}\theta_2^2 + \gamma_{2,9}\theta_1\theta_2^2 + \gamma_{2,10}\theta_3^3.$$

Clearly even with only basis functions of total order 3, we have a large number of unknowns in our optimisation problem, $\bar{\gamma} \in \mathbb{R}^{14}$. If we were to increase the dimension of θ further we would need to reduce the number of terms we include in the expansion by, for example, removing all the "cross" terms. This reduces the quality of our map but since we only require an approximate map we can afford to reduce the accuracy.



(b) Push forward of θ onto ref-

Reference



(c) Pull back of reference sample onto target space.

(a) Original sample θ from MH-RW algorithm. erence space.

Fig. 5.2. The effect of the approximate transport map \tilde{T} on a sample from the Rosenbrock target density as described in Equation (5.1).

Figure 5.2 shows the output of the approximate transport map. Even though we have truncated the infinite expansion in the monomial basis down to 4 and 10 terms in

respective dimensions, the push forward of the sample is still a unimodal distribution centred at the origin with standard deviation 1. As you move out into the tails of the reference density more non-Gaussian features are clearly visible. However, overall, the push forward of the target density does not look a challenging one to sample from, with even relatively simple MCMC methods such as MH-RW. The pullback from reference space, in Figure 5.2, is an exact match of the original sample since we have not perturbed the sample in reference space. This inversion is well defined in the sampling region, although not necessarily outside [25].

5.2. Numerical results for convergence of transport map based algorithms on the Rosenbrock density. We first find the optimal scaling parameters for the individual algorithms. This is done, as in [9], by optimising for the effective sample size in the PAIS algorithm, and by tuning the relative L^2 error in the MH algorithm. There is currently no guidance on the best way of tuning the MH algorithm with transport map proposals although one might expect results similar to the standard MH results, especially if adaptation of the map is stopped after a given point. As in the PAIS algorithm, optimising for the effective sample size might be the best option.

Statistic /	Algorithm	Transport M-H	Alg. 2	Alg. 3
$\overline{\delta_{L^2}}$		1.0e-0	1.1e-1	3.5e-1
$\delta_{\mathrm{ESS}}^{-}$		-	1.0e-1	5.2e-1
Acc. rate		0.23	-	_
ESS ratio		-	0.62	0.71

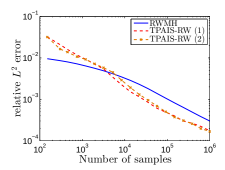
Table 5.1

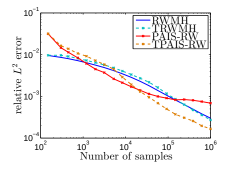
Optimal scaling parameters for the transport map based algorithms applied to R_1 , optimising for the L^2 error and the effective sample size (ESS) for PAIS algorithms, and for average acceptance rate for MH algorithms.

The optimal scaling parameters are given in Table 5.1. Here we see that the effective sample size is much lower than we see in the one-dimensional examples with the PAIS algorithms. However, in the Rosenbrock density (5.1) we are dealing with a much more complicated correlation structure, as well as a very slowly decaying tail in θ_2 . From our experiments, we have observed that the standard PAIS-RW required an ensemble size of M=500 to overcome the problems in this density, however the transport map transforms the tails to be more like those of a Gaussian which can be approximated well by a smaller ensemble size of M=150.

The convergence of the three algorithms is displayed in Figure 5.3. Figure (a) shows that the two variations of the transport based PAIS algorithms converge with similar rates. The second version, which performs the resampling stage in reference space rather than target space, has a slightly higher ESS, and is more stable than option (1). This version also has a property that we can exploit in Section 8.1.

6. Multiscale Methods for Stochastic Chemical Reaction Networks. In this Section, we discuss some recent advances in multiscale methods for stochastic reaction networks. Inverse problems arising in this area often lead to highly correlated and complex posterior distributions, which traditional MCMC methods can struggle to sample from. We will then go on to solve some inverse problems related to this in Section 7, using the transport map versions of the PAIS algorithm, as described in Section 4.





(a) Comparison of the two Transport PAIS (b) Comparison of Transport PAIS option (2) options. with the standard algorithms.

Fig. 5.3. Convergence of algorithms; Transport M-H, algorithm 2, algorithm 3 for density (5.1). Ensemble size M=150, resampling performed using the AMR algorithm.

We consider chemical reaction networks of N_s chemical species $\{S_j\}_{j=1}^{N_s}$, with population numbers given by $X(t) = [X_1(t), X_2(t), \dots, X_{N_s}(t)]^{\top} \in \mathbb{N}_0^{N_s}$ reacting in a small reactor, through N_r different reaction channels. When population numbers of one or more of the chemical species in the reactor is small, as is the case with chemical reactions occurring within living cells, the sporadic and discrete way in which reactions occur can not be well modelled by deterministic continuous models, such as ODEs. In such a situation, we may wish to model the dynamics through a discrete stochastic model, such as a continuous time Markov chain.

For each reaction R_j , for $j=1,2,\ldots N_r$, there is a propensity or hazard function $\alpha_j(X(t))$ which indicates how likely that reaction is to fire, defined by

$$\alpha_j(X(t)) = \lim_{dt \to 0} \mathbb{P}(\text{Reaction } R_j \text{ in the time interval } \quad s \in [t, t+dt]).$$

If a system satsifies what is called *mass action kinetics*, then the form of the function α_i is determined, up to a rate constant, by the reactants involved in that reaction:

$$\alpha_j(X) = k_j \prod_{m=1}^{N_s} \prod_{n=0}^{\nu_{j,m}-1} (X_m - n), \tag{6.1}$$

where $\nu_{j,m}$ is the *m*th component of the stoichiometric vector ν_j , X_m is the *m*th component of the state vector X, and the k_j are rate constants.

Following each reaction R_j there is an instantaneous change in the current state, as the reactants of the reaction are consumed, and the products produced. This is modelled by a change in the state vector $X(t) = X(t) + \nu_j$ where ν_j is that stochiometric vector for reaction R_j .

The model can be represented as the following expression involving N_r different unit rate Poisson processes [4] Y_j , given by:

$$X(t) = X(0) + \sum_{i=1}^{N_r} \nu_j \int_0^t \alpha_j(X(s)) ds.$$
 (6.2)

The master equation for this model is only solvable in certain situations, for example monomolecular networks [20], or for steady-state solutions certain deficiency zero

networks [1,2]. Trajectories for this system can be sampled exactly, for instance using the Gillespie SSA [17], or its variants [3,6,16,18]. However, if the system is stiff, i.e. there are some reactions which are firing many times on a timescale for which others are unlikely to fire at all, then trajectories can become prohibitively expensive to simulate, since these methods simulate every single reaction event with the same cost. In such a system, one might employ multiscale methods in order to approximate trajectories at a lower cost than the exact algorithms.

One common approach is to induce the quasi-steady-state approximation QSSA. This approximation makes the assumption that fast reactions enter quasi-equillibrium on a timescale which is negligible with respect to the timescale on which the slow dynamics in the system are evolving. This amounts to taking the assymptotic limit that the propensities of the "slow" reaction channels are equal to zero. This allows us to sample approximate trajectories of the slow reactions without the need to compute many fast reaction events.

This approach can work well when there is a large timescale gap between the fast and slow reactions in a system, but where the timescale gap is less pronounced, it can lead to significant errors [12]. Another approach is the constrained multiscale method (CMA) [10,11], based in part on the equation-free approach to multiscale computations [15,23]. This approach also assumes quasi-equilibrium in the system, but takes into account the effect that the slow reactions can have on the invariant distribution of the fast variables in the system. For a more detailed description of this method, please refer to the literature [10].

Multiscale methods are not only of use when forward evaluations of the dynamics are costly, but can also be of use where we attempt to solve an inverse problem where the fast variables are unobservable. One approach in this situation would be to integrate over all possible trajectories of the fast variables, but this would almost always be prohibitively expensive. Another approach would be to use an appropriate multiscale approximation, so that the effective dynamics of the slow variables can be approximate without the need to consider the rapid fluctuations of the fast variables in the system.

6.1. Likelihoods arising from stochastic reaction networks. Suppose that we are able to exactly observe the number of molecules of each chemical species in a system which satisfies mass action kinetics, and which can be well approximated by (6.2). Suppose that we wish to be able to infer the value of the rate constants of each reaction from these observations. Even with no observational noise, since the dynamics of the system are stochastic, this still leads to a Bayesian inverse problem where we can only infer a joint probability distributions on the reaction parameters. Suppose that we are in state $X(t) = X_0$. There are two independent univariate random variables which decide when and what the next reaction in the system is. There is the jth waiting time τ_j to the next reaction, which is given by an exponential random variable

$$\tau_j \sim \exp\left(\frac{1}{\alpha_0(X(t))}\right),$$

where $\alpha_0(X(t)) = \sum_i^{N_r} \alpha_i(X(t))$ is the total propensity in the system. The second is a multinomial random variable r_j which dictates which reaction has occurred during the jth reaction, which takes the value $r \in \{1, 2, ..., N_r\}$ with probability

$$\mathbb{P}(r=i) = \frac{\alpha_i(X(t))}{\alpha_0(X(t))}.$$

As such, in order to compute the likelihood of a particular trajectory given a possible realisation of the reaction parameters, it is sufficient to have access to the total time spent in each state, and the frequency of each reaction which led to leaving each state. From this formulation, we see that the random variables (τ_j, r_j) only depend on the states $\mathbf{X}(t_{j-1})$ and so are Markovian. This conditional independence means that we can group events together by what state the system was in when the event happened. We define two new random variables which depend on a state $\mathbf{Y} \in \mathcal{S}$, first the total time spent in state \mathbf{Y} ,

$$T(\mathbf{Y}) = \sum_{j=1}^{M} \tau_j \ \mathrm{I}(\mathbf{X}(t_{j-1}) = \mathbf{Y}).$$

This random variable, $T(\mathbf{Y})$, is a sum of exponentially distributed random variable, each with the rate $\alpha_0(\mathbf{Y}; \mathbf{k})$, and hence follows the Gamma distribution,

$$T(\mathbf{Y}) \sim \text{Gamma} (\alpha = K(\mathbf{Y}), \ \beta = \alpha_0(\mathbf{Y}; \mathbf{k})),$$
 (6.3)

where
$$K(\mathbf{Y}) = \sum_{j=1}^{M} I(\mathbf{X}(t_{j-1}) = \mathbf{Y}).$$

Similarly, we can define the reactions which occurred when the system was in state \mathbf{Y} as $\mathbf{r}(\mathbf{Y}) = [r_1(\mathbf{Y}), \dots, r_{N_r}(\mathbf{Y})]^{\top}$ where

$$r_i(\mathbf{Y}) = \sum_{j=1}^M \mathrm{I}(r_j = i ext{ and } \mathbf{X}(t_{j-1}) = \mathbf{Y}).$$

Here all the random variables r_j follow the same multinomial distribution, and so

$$\mathbf{r}(\mathbf{Y}) \sim \text{Multinomial}(K(\mathbf{Y}), \ \mathbf{p}(\mathbf{Y})).$$
 (6.4)

The random variables defined in Equations (6.3) and (6.4) are sufficient statistics for the posterior distribution $\pi(\mathbf{k}|D)$. With these definitions we define two new structures

$$\mathbf{T} = [T(\mathbf{Y}_1), \dots, T(\mathbf{Y}_K)]^{\top}, \text{ and } \mathbf{R} = [\mathbf{r}(\mathbf{Y}_1), \dots, \mathbf{r}(\mathbf{Y}_K)]^{\top},$$

where $K = |\mathcal{S}|$, the number of states in \mathcal{S} , and each state $\mathbf{Y}_i \in \mathcal{S}$ has been enumerated. We use these structures to define shorter notation,

$$\mathbf{T}_i = T(\mathbf{Y}_i), \quad \mathbf{R}_{ij} = r_j(\mathbf{Y}_i), \quad \text{and} \quad \mathbf{K}_i = K(\mathbf{Y}_i).$$

To construct the posterior distribution for the reaction rates, \mathbf{k} , in the chemical system, we formulate the likelihood using the sufficient statistics derived in the previous section. Due to the non-negativity of these reaction rates, we assign a Gamma prior distribution to each rate. Given the distributions in Equations (6.3) and (6.4) for our data, the likelihood of observing the data \mathbf{R} and \mathbf{T} is

$$\ell(\mathbf{R}, \mathbf{T}|\mathbf{k}) \propto \prod_{i=1}^{K} \text{Multi}(\mathbf{r}(\mathbf{Y}_i); \mathbf{K}_i, \mathbf{p}(\mathbf{Y}_i)) \text{Gamma}(\mathbf{T}_i; \mathbf{K}_i, \alpha_0(\mathbf{Y}_i)),$$

where again $\mathbf{Y}_i \in \mathcal{S}$ and the propensities α_i and probabilities p_i depend on the reaction rates $\mathbf{k} = [k_1, k_2, \dots k_{N_r}]^{\top}$ through the concept of mass action kinetics given in (6.1).

Our choice of Gamma prior distributions with hyper-parameters (a_i, b_i) results in the posterior distribution of the form

$$\pi(\mathbf{k}|\mathbf{R}, \mathbf{T}) \propto \ell(\mathbf{R}, \mathbf{T}|\mathbf{k}) \prod_{i=1}^{N_r} \operatorname{Gamma}(k_i; a_i, b_i)$$

$$\propto \exp\left\{ \sum_{i=1}^K \left[\mathbf{K}_i \log \alpha_0(\mathbf{Y}_i; \mathbf{k}) - \mathbf{T}_i \alpha_0(\mathbf{Y}_i; \mathbf{k}) + \sum_{j=1}^{N_r} \mathbf{R}_{ij} \log \mathbf{p}_j(\mathbf{Y}_i; \mathbf{k}) \right] + \sum_{i=1}^{N_r} ((a_i - 1) \log k_i - b_i k_i) \right\}.$$
(6.5)

6.2. Approximation of likelihoods in multiscale chemical networks. Suppose now that we are only observing the slower variables in a multiscale chemical system of this type. Suppose that $n_r < N_r$ is the number of slow reactions in the system, and the slow reactions are given by $\{R_{s_1}, R_{s_2}, \dots R_{s_{n_r}}\} \subset \{R_1, R_2, \dots, R_{N_r}\}$. The propensities of the slow reactions α_{s_i} may depend on the value of the fast variables which is unknown. However, the invariant distribution of the fast variables conditioned on the slow variables in the system can be approximated using a multiscale method, such as the QSSA or the CMA, as described briefly in Section 6. Once the approximations have been made, we arrive at approximate effective propensities $\bar{\alpha}_{s_i}$, which have been averaged over the computed invariant distributions of the fast variables. Then the approximate effective dynamics on the slow variable S(t) are given by

$$S(t) = S(0) + \sum_{i=1}^{n_r} \nu_{s_j} \int_0^t \bar{\alpha}_{s_j}(S(s)) ds.$$
 (6.6)

In turn, the approximate posterior distribution on the reaction parameters \mathbf{k} is then given by

$$\pi(\mathbf{k}|\mathbf{R},\mathbf{T}) \propto \ell(\mathbf{R},\mathbf{T}|\mathbf{k}) \prod_{i=1}^{n_r} \operatorname{Gamma}(k_i; a_i, b_i)$$

$$\propto \exp\left\{ \sum_{i=1}^K \left[\mathbf{K}_i \log \bar{\alpha}_0(\mathbf{Y}_i; \mathbf{k}) - \mathbf{T}_i \bar{\alpha}_0(\mathbf{Y}_i; \mathbf{k}) + \sum_{j=1}^{n_r} \mathbf{R}_{ij} \log \bar{\mathbf{p}}_j(\mathbf{Y}_i; \mathbf{k}) \right] + \sum_{i=1}^{n_r} ((a_i - 1) \log k_i - b_i k_i) \right\},$$
(6.7)

where $\bar{\alpha}_0(\mathbf{Y}_i; \mathbf{k}) = \sum_{j=1}^{n_r} \bar{\alpha}_j(\mathbf{Y}_i; \mathbf{k})$ is the total of the effective propensities, n_r is the number of slow reactions, $\bar{\mathbf{p}}(\mathbf{Y}_i; \mathbf{k})$ is the multiscale approximation of the expected proportion of each slow reaction at state \mathbf{Y}_i and with reaction rates \mathbf{k} and the data is limited only to changes in the slow variable due to the occurrence of slow reactions.

7. Numerical Examples. In this section we look at two examples of chemical systems to demonstrate the effectiveness of the Bayesian approach we have proposed, along with the transport PAIS algorithms for sampling the challenging posterior distribution on the reaction parameters that arise.

7.1. A Multiscale Chemical System. First we consider a simple multiscale example involving two chemical species S_1 and S_2 involved in only zeroth and first order reactions:

$$\emptyset \xrightarrow{k_1} S_1 \xrightarrow{k_2} S_2 \xrightarrow{k_4} \emptyset. \tag{7.1}$$

Each arrow represents a reaction from a reactant to a product, with some rate constant k_i , and where mass action kinetics is assumed. The parameters k_i are non-negative, and $\mathbf{k} = [k_1, \dots, k_4]^{\top} \in \mathbb{R}^4_+ = \mathcal{X}$. We denote the population count of species S_i by $X_i \in \mathbb{N}_0$. We consider this system with the following parameters:

$$k_1 = 100, k_2 = k_3 = 10, k_4 = 1, (7.2)$$

In this parameter regime the reactions $R_2: S_1 \to S_2$ and $R_3: S_2 \to S_1$ occur more frequently than the other reactions. Notice that both chemical species are involved in fast reactions. However, the quantity $S = X_1 + X_2$, is conserved by both of the fast reactions, and as such, this is the slowly changing quantity in this system.

We observe this system over the time period $t \in [0, 500]$ with initial condition ******, but we assume that we are only able to observe the slow variable S(t). The effective dynamics of S can be approximated by a system of only two reactions:

$$\emptyset \xrightarrow{\alpha_1(S)} S \xrightarrow{\bar{\alpha}_4(S)} \emptyset, \tag{7.3}$$

where here the propensities are shown as opposed to the rates.

The effective propensity $\bar{\alpha}_4(S)$ can be approximated through application of a multiscale approximation, as detailed in Section 6, and in more detail in [10]. These effective propensities can often be computed without the need for expensive simulations, in particular when the fast subsystem is deficiency zero, which is often the case [1,2]. For this very simple example, the dependance of these effective propensities on the original rate parameters can be explicitly understood. Under the QEA, the effective rate of degradation of the slow variable S is given by

$$\bar{\alpha}_4^{\text{QEA}}(s) = \frac{k_2 k_4 s}{k_2 + k_3}.$$

Similarly, the analysis of the constrained system as discussed in [10] yields the effective propensity

$$\bar{\alpha}_4^{\text{CMA}}(s) = \mathbb{E}_{\text{CMA}}\left[k_4 X_2 | S = s\right] = \frac{k_2 k_4 s}{k_2 + k_3 + k_4}.$$
 (7.4)

Our observations are uninformative about the reaction rates k_2, k_3 , and k_4 , as for each multiscale approximation there is a manifold $\mathcal{M} \subset \mathcal{X}$ along which the effective propensity $\hat{\alpha}_4$ is invariant, leading to a highly ill-posed inverse problem. This type of problem is notoriously difficult to sample from using standard MH algorithms, as the algorithms quickly gravitate towards the manifold \mathcal{M} but then exploration around \mathcal{M} is slow.

We aim to recover three different posterior distributions. In the first, which is of the form (6.5), we assume that we can fully and exactly observe the whole state vector for the whole of the observation time window $t \in [0, T]$. In the second and third, the same data are used, but restricted to observations of the slow variable $S = X_1 + X_2$.

Parameter\Dimension i	1	2	3	4
α_i	150	5	5	3
eta_i	$\frac{15}{9}$	$\frac{5}{12}$	$\frac{5}{12}$	1

Hyper-parameters in the prior distributions for the multiscale problem described in Section 7.1.

In these last two examples, we approximate the posterior using (6.7) with QEA and CMA approximations of the effective dynamics respectively.

In all three cases, we find the posterior distribution for $\mathbf{k} \in \mathcal{X}$. These four parameters are assigned Gamma prior distributions,

$$k_i \sim \text{Gamma}(\cdot; \alpha_i, \beta_i), \text{ for } i = 1, \dots, 4.$$

The hyper-parameters corresponding to each of these prior distributions are given in Table 7.1. These priors are the same for each of the three posterior distributions.

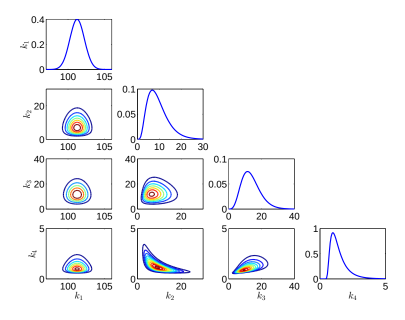


Fig. 7.1. CMA approximation (6.7) of the posterior arising from observations of the slow variable $S = X_1 + X_2$ for system (7.1) with true rates given by (7.2).

Figure 7.1 shows how this posterior looks when we use the CMA to model the effective degradation propensity $\bar{\alpha}_4$.

We consider several different algorithms for sampling from these distributions. First we implement both the PAIS and MH algorithms with a Gaussian proposal distribution. In the case of the PAIS algorithm, this is a Gaussian mixture proposal distribution, and in the MH algorithm this is equivalent to a vanilla random walk. The proposal distribution uses a covariance matrix which has been constructed using the sample covariance of a sample produced using a standard MH-RW algorithm. We also compare the PAIS and MH algorithms when using a transport map proposal distribution. This proposal method was discussed in detail in Chapter 4.

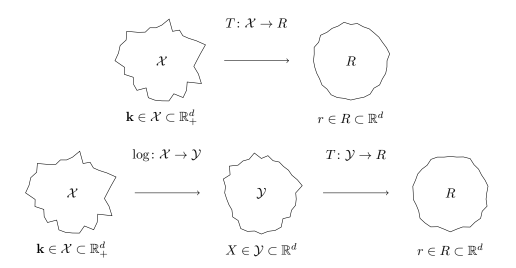


Fig. 7.2. Couplings between parameter space and reference when using the transport map with (bottom) and without (top) a log preconditioner.

Figure 7.2 (top) shows how we define a bijective map, \hat{T} , between parameter space \mathcal{X} and a reference space R. When using the transport map proposal distribution, we prefix the proposal method with a T, e.g. MH-RW and MH-TRW, as well as PAIS-RW and PAIS-TRW. In practice we cannot ensure that the approximate map T is uniquely invertible over the whole of R and so \tilde{T} is not truly bijective. This leads to problems for our strictly positive state space, $\mathcal{X} \subset \mathbb{R}^d_+$, since proposals in R do not necessarily map back onto \mathcal{X} .

Therefore we also consider how these algorithms perform when the transport map is preconditioned by the logarithmic function applied to the parameters. We choose to use this log preconditioner since it converts our strictly positive parameter space, \mathcal{X} , into an intermediate space $\mathcal{Y} \subset \mathbb{R}^d$. This allows us to define T between two subsets of \mathbb{R}^d , which means that all proposals made on the reference space are mapped to valid proposals on the true parameter space. It also reduces the work required of the map, which since we use only a finite dimensional approximation, can improve performance and stability. As before, the proposal distributions are labelled with a T for transport map and when using the intermediate space we prepend 'log' to the proposal method, e.g. MH-logRW and MH-logTRW, with, PAIS-logRW and PAIS-logTRW.

Figure 7.2 (bottom) displays the composition of maps when we include this log preconditioner leading to an intermediate space \mathcal{Y} . The inclusion of this additional map means that we must again alter our importance weight definition to reflect the pullback from R through \mathcal{Y} . The weight is now

$$w_i(\theta) = \frac{\pi(\theta|\mathbf{R}, \mathbf{T})}{\chi(\tilde{T} \circ \log(\theta)|\tilde{T} \circ \log(\theta^{(i-1)}))|J_{\tilde{T} \circ \log}(\theta)|},$$

where θ is a proposed new state, $\theta^{(i-1)}$ is the ensemble of states from the previous iteration, and $J_{\tilde{T} \circ \log}(\theta)$ is the Jacobian of the composition of the two maps. This Jacobian is straightforward to calculate,

$$|J_{\tilde{T}\text{olog}}(\theta')| = |J_{\tilde{T}}(\log(\theta'))||J_{\log}(\theta')|,$$

where the first determinant is computed as in (3.3), and the second is given by

$$|J_{\log}(\theta')| = \prod_{i=1}^d \frac{1}{\theta_i'}.$$

For this problem, we continue to use monomials in each dimension in our transport map construction. We use polynomials of total order p=3 as the basis functions, i.e.

$$T_i(\theta) = \sum_{\mathbf{j} \in \mathcal{J}_i^{\text{TO}}(p)} \gamma_{i,\mathbf{j}} \psi_{\mathbf{j}}(\theta) \text{ where } \psi_{\mathbf{j}}(\theta) = \prod_{k=1}^i \theta_k^{j_k},$$

and

$$\mathcal{J}_i^{\text{TO}}(p) = \{ \mathbf{j} \in \mathbb{N}_0^d \mid ||\mathbf{j}||_1 \le p, \text{ and } j_k = 0 \ \forall k > i \}.$$

We will use the AMR [29] resampler with an ensemble size of M = 500. This increase in ensemble size in comparison with previous example in Section 5 is to allow for the increase in parameter dimension.

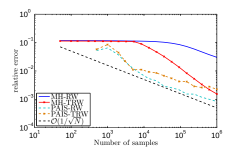
To measure the convergence of the sampling methods in this section, we will compare the convergence of the mean of each parameter. We approximate $\mathbb{E}(\mathbf{k})$ using 2.4 billion samples from the MH-RW algorithm. We do this for each of the eight algorithms we have so far discussed. Convergence is shown only for the CMA approximation of the posterior (6.7) with effective rate for the degradation of S given by (7.4), but we expect very similar results for the other posterior distributions discussed.

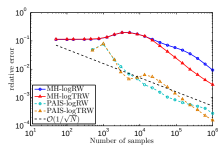
Algorithm	MH	PA	PAIS		
	δ	δ	ESS		
RW	5.5e-3	1.0e-0	9.0e-3		
TRW	1.2e-0	4.0e-1	1.2e-3		
logRW	1.7e-1	1.2e-0	6.0e-2		
$\log TRW$	2.7e-2	1.5e-1	3.5e-1		

Table 7.2

Optimal scaling parameters for the MH and PAIS algorithms applied to the constrained multiscale problem in Section 7.1. MH parameters optimised by acceptance rate, and PAIS parameters optimised using effective sample size.

The optimal scaling parameters for the proposal distributions are given in Table 7.2. These are precomputed by running the algorithms for different parameter values. MH-RW based algorithms are tuned to an acceptance rate close to the optimal 23.4%, and PAIS algorithms are tuned to maximise the effective sample size (ESS). The optimal parameter values for the MH algorithms are not particularly informative about their performance, since they relate to algorithms operating on different transformations. The ESS is highest for the logTRW variant of the PAIS algorithm, as we would hope. Convergence of the eight algorithms for this example is shown in Figure 7.3. We first note the poor performance of the MH based algorithms, each of them taking roughly 10,000 samples to begin converging. Only the MH-TRW is at all competitive with the PAIS algorithms. During the simulation interval, the MH-TRW algorithm has not settled down to the expected $\mathcal{O}(1/\sqrt{N})$ rate which means that the estimate is still



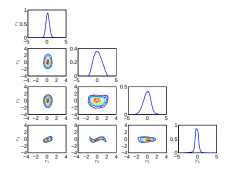


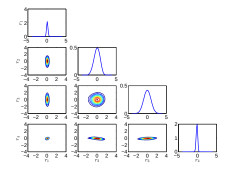
(a) Sampling algorithms with no preconditioner (b) Sampling algorithms with a log preconditioner for \tilde{T} .

Fig. 7.3. Convergence of eight different MCMC algorithms for the CMA approximation of the posterior arising from the example in Section 7.1.

biased by the long burn-in time. As we have seen in previous examples, the burn-in time for the PAIS algorithm is negligible.

The PAIS variants with RW and TRW proposals perform similarly on both sample spaces. When sampling without a preconditioner, the transport map is not quite as efficient, largely due to the difficulties discussed in the previous section i.e. many proposals are made which do result in negative reaction rates. Sampling with transport maps with a log preconditioner leads to more comparable Monte Carlo errors, with the logTRW being apparently slightly less stable. This proposal method becomes more stable as we increase either the ensemble size, or the number of iterations between updates of the transport map, T. Overall we see the smallest Monte Carlo errors for a given amount of computational effort coming from the PAIS-logTRW algorithm.





(a) Push forward of the posterior sample (b) Push forward of the posterior sample through \hat{T} with no log preconditioner. through \hat{T} with log preconditioner.

Fig. 7.4. Push forward of the target measure on reference space for the the CMA approximation of the posterior arising from the example in Section 7.1, found using the TRW and logTRW proposal distributions. Components are linked by the relation $r_i = T_i(k_i)$ in (a) and $r_i = T_i \circ \log(k_i)$ in (b).

We now look at the proposal distributions of the transport map accelerated algorithms. In Figure 7.4, we see the reference spaces found by; in (a) mapping the posterior through the map \hat{T} , and in (b) by mapping the posterior through $T \circ \log$. For the most part, each of these marginal distributions can be recognised as a Gaussian. However, with the exception of $\mathbb{P}(r_2, r_3)$, we would not consider them to be

close to a standardised $\mathcal{N}(0, I)$ distribution. Before thinking that the transport map has not helped us to find a 'nicer' space on which to propose new values we should consider that the dimensions are now (1) largely uncorrelated, and (2) the variances in each dimension are much more similar than they are in Figure 7.1.

In Figure 7.4 (b) we see that $var(r_1)$ and $var(r_4)$ are much smaller than $var(r_2)$ and $var(r_3)$. To combat this we have a number of choices. We might wish to learn the covariance structure of the push forward of the target onto the reference space, and incorporate this information into the covariances of the mixture components in an adaptive scheme. Another option is to increase the total order of our index set. For these numerics we have chosen p = 3, but we can obtain reference spaces which are closer to $\mathcal{N}_d(0, I)$ by choosing a larger p.

7.1.1. Comparison of the Constrained and QEA approaches. The convergence analysis has been performed for the constrained approximation of the posterior distribution. We now look at the differences in the approximations of the posterior distribution arising in this problem between the CMA and QEA. Recall that the approaches differed only in the form of the effective degradation propensity $\hat{\alpha}_4$,

$$\hat{\alpha}_4^{\text{QEA}}(s) = \frac{k_2 k_4 s}{k_2 + k_3}$$
 and $\hat{\alpha}_4^{\text{CMA}}(s) = \frac{k_2 k_4 s}{k_2 + k_3 + k_4}$.

This difference in the denominator causes a shift in the manifold on which the posterior is concentrated, as can be seen in Figure 7.5. The figure shows the difference in posteriors densities

$$\operatorname{diff}(\mu^{\text{CMA}}, \mu^{\text{QEA}}) = \pi^{\text{CMA}}(\mathbf{k}|\mathbf{R}, \mathbf{T}) - \pi^{\text{QEA}}(\mathbf{k}|\mathbf{R}, \mathbf{T}). \tag{7.5}$$

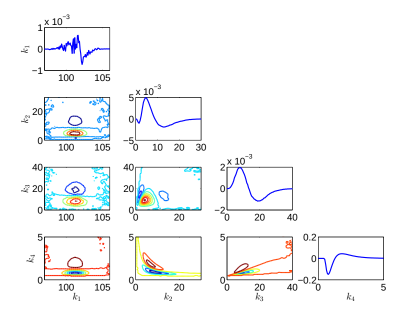


Fig. 7.5. Difference between the CMA and QEA posteriors as defined in Equation (7.5).

Since the two posteriors have been approximated using an MCMC sample, there is a significant amount of noise, particularly in the tails of the distributions. We can see

that the differences in the marginals for k_1, k_2 , and k_3 are relatively small, but the marginal for k_4 varies by a significant amount.

In the QEA approximation, the effective rate of degradation of S is given by $\hat{k}_4^{\text{QEA}} = k_2 k_4/(k_2 + k_3)$, and as such, our observations should be informative about this quantity. Using the CMA, this effective rate is approximated by $\hat{k}_4^{\text{CMA}} = k_2 k_4/(k_2 + k_3 + k_4)$. To validate our inferences on k_2 , k_3 and k_4 we would like to discover which model is most accurate and informative about these quantities, and which model gives us results which are most similar to that which can be obtained from the full model, with all reactions (including fast reactions) observed.

A conventional way to compare two models under a Bayesian framework is to calculate the Bayes factors [8]. The Bayes factor, $B_{1,2}$, between two models, \mathcal{M}_1 and \mathcal{M}_2 , can be interpreted as a ratio of the normalisation constants of the posterior distributions given each model,

$$B_{1,2} = \frac{\mathbb{P}(D|\mathcal{M}_1)}{\mathbb{P}(D|\mathcal{M}_2)}, \quad \text{where} \quad \mathbb{P}(D|\mathcal{M}_k) = \int_{\mathcal{X}} \mathbb{P}(D|\theta_k, \mathcal{M}_k) \mathbb{P}(\theta_k|\mathcal{M}_k) \, \mathrm{d}\theta_k.$$

Under the PAIS framework, it is straightforward to calculate these factors using the Monte Carlo estimator for the normalisation constants. From [28] these normalisation constants take the form

$$Z_k \approx \hat{Z}_k \frac{1}{NM} \sum_{i=1}^N \sum_{j=1}^M w_j^{(i)}(\mathcal{M}_k),$$

where $w_j^{(i)}(\mathcal{M}_k)$ is the weight under model \mathcal{M}_k corresponding to the j-th ensemble member on the i-th iteration. Hence, $B_{1,2} \approx \hat{Z}_1/\hat{Z}_2$. We can compare more than two models in this way by selecting the model with the largest marginal distribution as the best model.

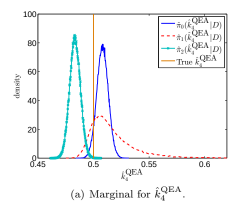
We now label the constrained model as \mathcal{M}_1 , the model arising from the QEA as \mathcal{M}_2 , and the full data model as \mathcal{M}_0 . Model \mathcal{M}_1 is dependent on the parameter $\theta_1 = (k_1, \hat{k}_4^{\text{CMA}})^{\top}$, and model \mathcal{M}_2 is dependent on the parameters $\theta_2 = (k_1, \hat{k}_4^{\text{QEA}})^{\top}$. The full model \mathcal{M}_0 is dependent on all four parameters $\theta_0 = (k_1, k_2, k_3, k_4)^{\top}$. The marginal densities for the data, evaluated at the observed data, given the models are displayed in Table 7.3.

k	$\mathbb{P}(D = (\mathbf{R}, \mathbf{T})^{\top} \mathcal{M}_k)$
0	6.8e-3
1	3.2e-3
2	1.7e-3

Table 7.3

Marginal distributions for the data $(\mathbf{R}, \mathbf{T})^{\top}$ for each model considered in Section 7.1.

Computing the Bayes factors from Table 7.3 we see that we should of course prefer the model \mathcal{M}_0 in which we observe all reactions perfectly. However this model is not significantly better than the CMA model ($B_{0,1}=2.09<3.2$, [22]). The Bayes factor $B_{0,2}=4.1>3.2$ tells us that we should significantly prefer the full model to the QEA approximation of the posterior. These Bayes factors present a relatively weak argument that the constrained approximation of the posterior is superior to the QEA approximation.



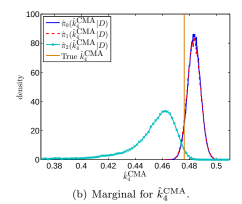


FIG. 7.6. Comparison of the approximate marginal densities for the quantities \hat{k}_4^{QEA} and \hat{k}_4^{CMA} across the three posterior densities $\pi_i = \mathbb{P}(\theta_i|D,\mathcal{M}_i)$ for i=0,1,2.

Figure 7.6 displays more compelling evidence in this direction. Here we show the marginal distributions for the quantities \hat{k}_4^{QEA} and \hat{k}_4^{CMA} with respect to each of the three posterior distributions, $\pi_i(\cdot|D,\mathcal{M}_i)$, i=0,1,2.

Firstly, in Figure 7.6 (a), displaying the marginals for \hat{k}_4^{QEA} we see that π_0 , the density arising from the full model with all reactions observed, is peaked a distance away from π_2 , QEA approximation. The CMA approximation π_1 , in contrast, is peaked close to peak of π_0 , but there is more uncertainty here since we do not observe the fast reactions.

In contrast, in Figure 7.6 (b), the marginal densities for \hat{k}_4^{CMA} for π_0 and π_1 are barely distinguishable, even with the vastly reduced data available in the posterior π_1 with CMA approximation. The QEA approximation π_2 is again peaked far away from both of the other distributions. This appears to demonstrate that the data regarding the slow variable S is only informative about k_1 and \hat{k}_4^{CMA} , as predicted by the CMA multiscale approximation.

7.2. Gene Regulatory Network. In this example, we look at a model of a gene regulatory network (GRN) mechanism [5, 19, 21] used by a particular cell to regulate the amount of a protein, P, present. Proteins can bind in pairs to form a dimer D. The dimer can bind to the gene G to a "switched off" state G', in which the production of MRNA molecules M is inhibited. The MRNA encodes for production of the protein P, and both P and M can degrade. The eight reactions are given in Equation (7.6).

$$P + P \xrightarrow{k_1} D \qquad G + D \xrightarrow{k_3} G'$$

$$G \xrightarrow{k_5} G + M \qquad M \xrightarrow{k_6} M + P$$

$$P \xrightarrow{k_7} \emptyset \qquad M \xrightarrow{k_8} \emptyset$$

$$(7.6)$$

We create trajectories of this system with the following parameters:****

$$k_1 = ?etc (7.7)$$

over the time period **** $t \in [0, ?]$.

T-cells can be manufactured which bind to proteins of a certain type, and cause them to phosphoresce, allowing for approximate observations of the concentration levels of that protein. However, the T-cells will not necessarily differentiate between monomers and dimers of the protein. We replicate such a scenario, in a zero-observational noise setting, by assuming that we can only observe T = P + 2D + 2G', the total number of protein molecules in the system, alongside M, the concentration of MRNA. The number of dimers, and the status of the genetic switch, are assumed unobservable. In order to sample from the posterior distribution on the reaction parameters in the system given these observations, we can either integrate over all possible trajectories of the fast variables (which is computationally intractable), or as in Section 7.1 we can use a multiscale method to approximate the posterior.

The effective dynamics of T and M are then given by:

$$\emptyset \xrightarrow{\hat{\alpha}_5} M \qquad M \xrightarrow{\alpha_6} M + T$$

$$T \xrightarrow{\hat{\alpha}_7} \emptyset \qquad M \xrightarrow{\alpha_8} \emptyset$$

$$(7.8)$$

Here, the propensities are displayed, as opposed to the rates, with approximations of the propensities $\hat{\alpha}_5$ and $\hat{\alpha}_7$ to be computed. We omit the derivation of these effective propensities, using the CMA methodology, for brevity, but interested readers can see more details in [29].

7.2.1. Target Distribution. As alluded to in the previous section, we aim to sample from the CMA approximation of the posterior distribution as defined in Equation (6.5). This example is more involved than the previous one, since we have an eight dimensional parameter space corresponding to the eight reaction rates in the system (7.6). However half of those reactions are unobservable, and their effect on the posterior is only felt through their effect on the effective propensities $\hat{\alpha}_5$ and $\hat{\alpha}_7$. The priors for this problem were chosen to be fairly uninformative with respect to the likelihood function for each reaction rate. A list of the hyper-parameters corresponding to each Gamma prior can be found in Table 7.4.

Dimension i	1	2	3	4	5	6	7	8
α_i	2	100	100	3	3	3	2	2
β_i	50	0.02	1	1	0.6	1	50	50

Table 7.4

Hyper parameters for the Gamma priors on each of the reaction rates in the GRN example in Section 7.2.

The one- and two-dimensional marginal distributions for the full posterior distribution are displayed in Figure 7.7. The posterior does not contain many interesting correlation structures, however several dimensions have leptokurtic tails which are difficult for the standard PAIS algorithm to sample from. The marginal densities also vary over very different scales, which could lead to problems with the algorithms which do not benefit from the transport map.

7.2.2. Implementation. We apply the eight RW and logRW proposal distributions, with and without transport map acceleration, which were discussed in Section 7.1 to this posterior. As with this example, the log conditioner is required so that our transport map is a function $T: \mathbb{R}^d \to \mathbb{R}^d$, rather than a map from \mathbb{R}^d_+ to \mathbb{R}^d . The transport map setup has not changed from the previous example. However, due to the scaling of the weight we have had to be more careful about which samples we use

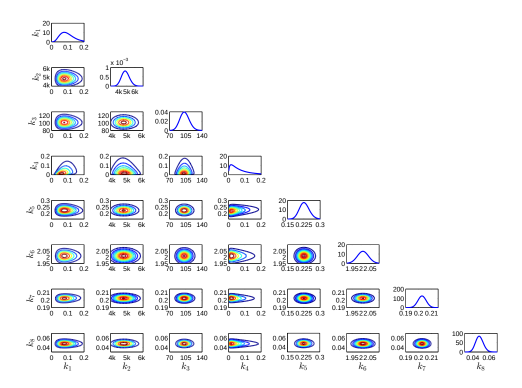


Fig. 7.7. Marginal densities of the CMA approximation of the posterior distribution for the GRN example in Section 7.2.

to build our map. We require that the objective function C(T), from Equation (3.7) is convex, which requires that the second derivative, given in Equation (3.11), is positive definite. This positive definite property depends on all weights in our sample being strictly positive, which is not always possible on a computer. For this reason we filter out any samples from the optimisation problem where the weight is a numeric zero. This does not affect the validity of the map since a weight zero would not contribute to the expectation.

In all eight algorithms we adapt the proposal variances during the burn-in phase of the algorithms to avoid the need for a huge ensemble size to approximate the posterior with a mixture of isotropic Gaussian kernels. We use the AMR resampler, this time with an ensemble size of M=2500. The increase in ensemble size compensates for the increase in dimension from four to eight.

As in the previous chemical reaction example, we measure convergence of the algorithms only through the convergence of the mean. We use the sample Mahalanobis distance, with 'true' covariance and 'true' mean built from a sample of size 2.4 billion using the MH-RW algorithm.

7.2.3. Convergence results. The scaling parameters used are selected by optimising the acceptance rate for the MH algorithms, and optimising the effective sample

size for the PAIS algorithms. The optimal parameters are given in Table 7.5.

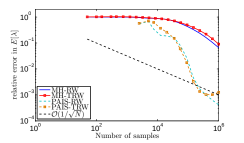
	МН				PAIS			
	RW	logRW	TRW	log-TRW	RW	logRW	TRW	log-TRW
$\beta_{\%}^*$	1.0e-1	3.9e-1	1.0e-0	6.0e-1	-	-	-	-
$eta_{\%}^* \ eta_{\mathrm{ESS}}^*$	-	-	-	-	1.0e-0	1.3e-0	8.0e-1	5.0e-1
ESS/M	-	-	-	-	$3.5\mathrm{e}\text{-}4$	4.7e-2	4.8e-2	1.6e-1

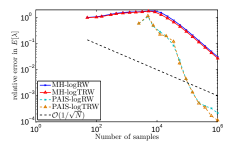
Table 7.5

Scaling parameters for the sampling algorithms applied to the GRN example in Section 7.2.

From the effective sample sizes shown in Table 7.5 we can see the improvement to the efficiency of the algorithm both by transforming the parameter space into something closer to Gaussian, and additionally by utilising a log preconditioner.

Due to the numerical cost of calculating the full relative L^2 errors for this posterior, we quantify the convergence speeds of these algorithms using the sample Mahalanobis distance between the sample mean and the 'true' mean. This convergence analysis is shown in Figure 7.8.





(a) Sampling algorithms with no preconditioner (b) Sampling algorithms with a log preconditioner for \tilde{T} .

FIG. 7.8. Convergence of eight different MCMC algorithms for the CMA approximation of the posterior arising from the example in Section 7.2.

The first thing to note is that the first PAIS measurement happens after 2500 samples, whereas the first MH measurement occurs after 50 samples. This is due to the much larger ensemble size required to sample in higher dimensions. We believe that an ensemble size of 2500 is over-cautious in this example and we could have used a smaller ensemble size. We also believe that the required ensemble size to sample from these posteriors is reduced by the use of the transport map.

The second obvious feature of these convergence plots is that the PAIS algorithms outperform the MH algorithms by a large margin - roughly a reduction of two orders of magnitude in the relative error over the same number of samples. A less positive result is that for this example the PAIS-TRW algorithm has found a map which has prevented further convergence of the mean. This is likely because one dimension of the transport map is sending proposals onto the negative real line and so all samples are being assigned a zero weight. This is behaviour which was anticipated, and it is always suggested that appropriate preconditioners are used where the posterior is not absolutely continuous with respect to Lebesgue measure, as is the case when the posterior is only non-zero on the positive quadrant of \mathbb{R}^d as in these examples.

The large increase in the effective sample size observed between the PAIS-logRW and PAIS-logTRW algorithms is converted into an estimate which is twice as accurate after 1 million samples.

A similar pattern is seen in the MH algorithms, where the MH-TRW performs the worst, and the MH-logTRW algorithm performs the best, although only marginally.

- 8. Discussion. One major issue with PMC-type algorithms is the curse of dimensionality. This curse is two-fold, since as the target distribution's dimension is increased, a larger ensemble size is often required in order to capture the addition complexity of the density that the higher dimension enables. Furthermore, the cost of the resampling step within the algorithm grows with the ensemble size. For example, the ETPF algorithm is $\mathcal{O}(n^2)$. However, the simiplification that the transport map makes has some potential to go some way to tackling this problem.
- **8.1. Sampling in (moderately) higher dimensions.** Here, we will briefly discuss how transport maps could aid with making moderately higher dimensional problems accessible to this family of methods.

Algorithm 3 allows us to decorrelate the dimensions of our random parameter on reference space, where we then can resample and map the resulting ensemble back onto target space. Since, on reference space, the dimensions are approximately uncorrelated, we are able to resample in each dimension separately. Resampling in a single dimension allows for optimisations in resampling code, and also means that the resampler is not affected by the curse of dimensionality.

In one dimension the ETPF algorithm can be implemented very efficiently. As described in [27], the coupling matrix has all non-zero entries in a staircase pattern when the state space is ordered. We can exploit this knowledge to produce Algorithm 4, which is much faster than using the simplex algorithm to minimise the associated cost function, and faster than the AMR algorithm [9]. Moreover, the task of running this resampler can then be parallelised over d processors. This could be an area for further study.

Algorithm 4: ETPF algorithm in one dimension.

```
1 Sort the states, \{(w_i, x_i)\}_{i=1}^M, into ascending order.
 2 Normalise the weights p_i = w_i/\|w\|_1.
 3 Set y_i \leftarrow 0 for all i = 1, \ldots, M.
 4 Set c \leftarrow 0
 5 for i \leftarrow 1, \ldots, M do
 6
        Set t \leftarrow p_i
         while j \leq M and t > 0 do
 7
             Set s \leftarrow (M^{-1} - c) \wedge t
 8
             Increase y_i by M \times s \times x_i.
 9
             Decrease t by s.
10
             Increase c by s.
11
             if t > 0 then
12
                 Increase j by 1.
13
                  Set c \leftarrow 0.
14
15 Return y.
```

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