High Performance Uncertainty Quantification with Parallelized Multilevel Markov Chain Monte Carlo

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Figure 1: Three levels of discretisation of a tsunami model used in a multilevel Markov chain Monte Carlo method.

ABSTRACT
Numerical models of complex real-world phenomena often necessitate High Performance Computing (HPC). Uncertainties increase problem dimensionality further and pose even greater challenges. We present a parallelization strategy for multilevel Markov chain Monte Carlo, a state-of-the-art, algorithmically scalable Uncertainty Quantification (UQ) algorithm for Bayesian inverse problems, and a new software framework allowing for large-scale parallelism across forward model evaluations and the UQ algorithms themselves. The main scalability challenge presents itself in the form of strong data dependencies introduced by the MLMCMC method, prohibiting trivial parallelization.

Our software is released as part of the modular and open-source MIT Uncertainty Quantification Library (MUQ), and can easily be coupled with arbitrary user codes. We demonstrate it using the Distributed and Unified Numerics Environment (DUNE) and the ExaHyPE Engine. The latter provides a realistic, large-scale tsunami model in which we identify the source of a tsunami from buoy-elevation data.

KEYWORDS
Multilevel Methods, ADER-DG, Bayesian inverse problems, Tsunami simulation

1 INTRODUCTION
Numerical simulation is an established tool driving innovation in many fields of science and engineering. Numerical solutions of mathematical models, for example in the form of partial differential equations (PDE), provide a prediction about a real-world process. More generally, deterministic models map a given set of parameters to a specific set of predicted values. There are, however, many applications where this approach is insufficient and a stochastic model is needed: The exact values of model parameters might be
Tsunami Warning Center operated by NOAA in the United States. 2-3 wavelengths away from the coast. The aforementioned uncertainty in parameters is typically expressed in terms of probability distributions. Finding the distribution of model predictions resulting from stochastic parameter distributions is referred to as a forward UQ problem. When in turn real-world measurements, also affected by uncertainty, are available and the underlying stochastic model parameters explaining those measurements are to be quantified, we speak of an inverse UQ problem. The resulting parameter distribution is called the posterior.

Here, we focus on inverse UQ problems. These are notoriously ill-posed, that is, the solution may lack existence, uniqueness, or continuous dependence on the data. As a result, specialized methods are required for inverse UQ problems. There is a wide range of methods available, and they vary significantly in how they balance expressivity (in the sense of how much information can be gained) against computational efficiency and against required knowledge about the model. For example, there are highly efficient optimization-based methods such as [19] for determining the maximum a posteriori probability (MAP) point when derivatives of the data misfit functional are available. On the other hand, stochastic collocation type methods can work with simple model evaluations and recover the posterior [24], but become inefficient for higher dimensional parameter spaces [33]. Markov Chain Monte Carlo (MCMC) type methods in turn also recover the full posterior distribution and in many cases only need simple forward model evaluations, but come at a very high computational cost in terms of numerous model evaluations.

In this paper, we choose MCMC as the “gold standard” type of method [20] in the sense of recovering the exact posterior with minimal assumptions on the model. In order to overcome the enormous computational burden incurred by MCMC on costly models, we employ a multilevel MCMC method and present a new parallelization strategy to employ multilevel MCMC on modern HPC systems. We then couple our UQ method to sophisticated PDE solvers in order to obtain efficient forward models. Since our multilevel MCMC method only assumes simple forward model evaluations, no modifications to the solver frameworks are required. By building efficient model hierarchies, we exploit properties of the forward solvers and show synergies between the PDE models and the multilevel UQ approach. Efficient parallelization is a particular focus of this work since, on the one hand, it is inevitable when solving large-scale PDE models, and on the other hand, multilevel MCMC introduces data dependencies that make parallelization non-trivial compared to plain-vanilla Monte Carlo (MC) methods.

To demonstrate the efficiency of our method in practical applications we use buoy data from coastal Japan to infer the location of initial displacements that led to the Tohoku tsunami in 2011. This method of using DART buoy data to predict tsunamis is commonly used in many early warning systems, such as the one at the Pacific Tsunami Warning Center operated by NOAA in the United States and at the National Tsunami Warning Center. These methods are known to work well for tsunamis which are initiated more than 2-3 wavelengths away from the coast.

We model the propagation of the tsunami by solving the shallow water equations. Our model is capable of modeling wetting and drying so that coastal regions can be included in the simulation [27]. For the numerical solution of the PDE, we apply an Arbitrary-order DERivative Discontinuous Galerkin Method (ADER-DG) implemented in the ExaHyPE framework [29]. The parallelized Multilevel Markov Chain Monte Carlo (MLMCMC) method was implemented in the MUQ library [26], is publicly available and fully model agnostic. To our knowledge, this is the first parallel MLMCMC implementation available; a sequential one [22] is available as part of PyMC3 [32].

We further show parallel scalability of our UQ algorithm and implementation by coupling to a model implemented in the DUNE framework [1], at the same time demonstrating how our UQ software framework can be coupled with arbitrary model codes.

2 EFFICIENT SOLUTION OF INVERSE PROBLEMS VIA MULTILEVEL MCMC

For a general model \( F(\theta) \) mapping parameters \( \theta \) to model predictions \( F(\theta) \), we formulate a Bayesian inverse problem. Its goal is to use the forward model \( F \) in order to infer the distribution of an uncertain model parameter \( \theta \) from measurement data \( y \):

\[
\nu \equiv \mathcal{P}(\theta|y) = \frac{\mathcal{L}(y|\theta)\pi(\theta)}{\mathcal{P}(y)} \propto \mathcal{L}(y|\theta)\pi(\theta). \tag{1}
\]

Here, \( \mathcal{P}(\theta|y) \) is the inferred parameter distribution given measurements \( y \). The likelihood \( \mathcal{L} \) is defined as the probability of observing the given measurements if \( \theta \) were the true parameter. Further, \( \pi \) is the prior distribution encoding a priori knowledge about the inferred parameters. We treat the distribution of measurements \( \mathcal{P}(y) \) as a practically unobtainable scaling factor, as it is irrelevant for MCMC methods. The final target is the mean of a Quantity of Interest (QOI) \( \mathbb{E}_y[Q] \) with respect to the above posterior distribution.

Evaluating the likelihood typically means comparing the model prediction \( F(\theta) \) to the measurements \( y \). For example, assuming Gaussian measurement errors with covariance \( \Sigma_f \), the likelihood will have the distribution \( N(F(\theta), \Sigma_f) \). In our applications, evaluating \( F \) means solving a PDE, so this is where the main cost of a Bayesian problem lies.

The main idea of MCMC is to generate a Markov chain, carefully designed to have a stationary distribution matching the posterior distribution we are looking for. Thus, for a sufficiently large number of steps, the chain will approximately draw samples from the otherwise inaccessible posterior. Those samples can in turn be used in a MC-like fashion to estimate \( \mathbb{E}_y[Q] \) or other statistics of the posterior.

Algorithm 1 shows one of the most common MCMC algorithms, namely Metropolis-Hastings Markov Chain Monte Carlo (MCMC) [25]. One of its main advantages is that it only requires a finite number of direct evaluations of the posterior and thereby of the forward model. No further information such as model derivatives or adjoints are needed.

Once we have these samples approximating the posterior, we can apply some post processing to compute \( \mathbb{E}_y[Q] \). In practice, since in the MCMC algorithm we already compute the forward model for
where theory typically guarantees that coarser meshes still deliver with fewer samples. A wide variety of improved proposals aims very closely related to Multilevel Monte Carlo (MLMC) methods expected value of the QOI by a telescoping sum and benefits to be gained in the MLMCMC setting. However, as we will see later, there are additional intricacies close. A more obvious and theoretically supported choice for a complex PDE model, as long as model evaluations are sufficiently differential equations (ODE) as rough approximations of a more hierarchy of models is defined, ranging from cheap-to-compute and many others.

Dimension-Independent Likelihood-Informed (DILI) MCMC [6, 7], and less correlated propos-

distribution, a significant number of samples may be required. Just from the model evaluations.

In order to accurately represent a multi-dimensional probability information to the posterior approximation. Less correlated propos-
ing good proposal distributions 𝑞 𝜇
take a different route: An entire

Algorithm 1: Metropolis Hastings MCMC

Result: Markov chain {θ𝑖}𝑖=0𝑁 with starting parameter θ0 ∈ ℝm;

Choose starting parameter θ0 ∈ ℝm;

for 𝑖 = 0, ..., 𝑁 − 1 do

Choose starting point 𝑇𝑖 ∈ ℝm;

Draw proposal θ′ from proposal distribution 𝑞(θ′|θ𝑖).

Compute acceptance probability

\[ α(θ′|θ𝑖) = \min \left\{ \frac{v(θ′)q(θ′|θ𝑖)}{v(θ𝑖)q(θ𝑖|θ′)} \right\}. \]

Draw a random number 𝑟 ∈ [0, 1].

if 𝑟 < α(θ′|θ𝑖) then

Accept proposal: θ𝑖+1 = θ′.

else

Reject proposal: θ𝑖+1 = θ𝑖.

end

end

each sample, we can immediately compute QOI samples derived from the model evaluations. In order to accurately represent a multi-dimensional probability distribution, a significant number of samples may be required. Just like basic MC methods, algorithm 1 requires a full model evaluation per sample. If the model itself is costly, the overall computational effort can easily become intractable.

Most effort to improve the efficiency of MHMCMC goes into finding good proposal distributions 𝑞(θ′|θ𝑖): MHMCMC produces correlated samples and strongly correlated samples barely contribute information to the posterior approximation. Less correlated proposals and high acceptance rates allow to achieve a good approximation with fewer samples. A wide variety of improved proposals aims to achieve this. Examples include preconditioned Crank-Nicolson [3, 5, 30], Adaptive Metropolis [16, 17], Hamiltonian MCMC [11], Dimension-Independent Likelihood-Informed (DILI) MCMC [6, 7], and many others.

Rather than minimizing the number of model evaluations for a single model, MLMCMC [9, 10, 18] takes a different route: An entire hierarchy of models is defined, ranging from cheap-to-compute rough approximations to the most accurate, yet expensive, full model. MLMCMC makes no assumptions on what exactly those coarse models could be. For example, we could use suitable ordinary differential equations (ODE) as rough approximations of a more complex PDE model, as long as model evaluations are sufficiently close. A more obvious and theoretically supported choice for a level hierarchy is varying mesh width in numerical PDE solvers, where theory typically guarantees that coarser meshes still deliver reasonable approximations. Note that the fundamental approach is very closely related to Multilevel Monte Carlo (MLMC) methods [15]. However, as we will see later, there are additional intricacies and benefits to be gained in the MLMCMC setting.

The basic idea of MLMCMC is to replace the estimation of the expected value of the QOI by a telescoping sum

\[ \mathbb{E}_{\nu_l}[Q_L] = \mathbb{E}_{\nu_0}[Q_0] + \sum_{i=1}^{L} \left( \mathbb{E}_{\nu_l}[Q_i] - \mathbb{E}_{\nu_{l-1}}[Q_{l-1}] \right). \]  (2)

where \( Q_l \) denotes the approximation of the QOI \( Q \) using model \( l \) of the model hierarchy. Working with a hierarchy of models not only affects the QOI: Each forward model \( F_l \) now implies its very own likelihood function \( L_l \) and corresponding posterior density \( \nu_l \) according to the Bayesian inverse problem formulated above.

This reformulation is clearly equivalent, but offers opportunities for significantly reducing computational cost:

- Coarser chains can be used as cheap-to-compute, well informed and nearly uncorrelated proposals for finer chains.
- Variance reduction between levels can be exploited: The coarser corrections in eq. (2) need many samples, but those are cheap to compute; fine corrections are more expensive per sample, but the variance in those corrections is reduced and thus only few samples are needed, especially when the models are converging as \( l \to \infty \).

An MLMCMC method implementing both ideas is shown in algorithm 2, as originally introduced by the authors of [9, 10].

Algorithm 2: Multilevel MCMC

Result: Markov chains \( \{\theta_i^l\}_{i=0}^{N_l} \) for all levels \( l \in \{0, ..., L\} \). On level 0, run a conventional MCMC chain, delivering samples \( \theta_0^0 \).

for level \( l = 1, \ldots, L − 1 \) do

Choose starting point \( \theta_0^l \) with coarse component from next coarser starting point \( \theta_{l-1}^{l-1} \).

for sample \( j = 1, \ldots, N_l \) do

Given \( \theta_j^l \), generate proposal \( \theta_j^l = \left\{ \begin{array}{ll}
\theta_j^l & \text{if } \mu_j = 0 \\
\theta_j^l & \text{if } \mu_j = 1 \\
\theta_j^l & \text{if } \mu_j = 2
\end{array} \right. \)

where

- \( \theta_j^l \) is drawn from a level \( l-1 \) chain with subsampling rate \( \mu_j \) and
- \( \theta_j^l \) is drawn from a proposal density \( q_{l}(\theta_j^l|\theta_j^{l-1}) \).

Compute acceptance probability

\[ α(\theta_j^l, \theta_i^l) = \min \left\{ \frac{v_l(\theta_j^l)q_l(\theta_j^l|\theta_j^{l-1})}{v_l(\theta_i^l)q_l(\theta_i^l|\theta_i^{l-1})}, \frac{v_{l-1}(\theta_j^l)q_{l-1}(\theta_j^l|\theta_j^{l-1})}{v_{l-1}(\theta_i^l)q_{l-1}(\theta_i^l|\theta_i^{l-1})} \right\}. \]

Draw a random number \( r \in [0, 1] \).

if \( r < α(\theta_j^l, \theta_i^l) \) then

Accept proposal: \( \theta_j^{l+1} = \theta_j^l \)

else

Reject proposal: \( \theta_j^{l+1} = \theta_i^l \)

end

end

end

Here we begin with a regular MCMC chain for the coarsest level. We then proceed to generate MCMC chains for finer levels, while using samples from coarser chains as proposals. Since we permit increasing parameter dimensions across levels, the coarse samples \( \theta_j^l \), from level \( l-1 \) used as proposals for level \( l \) may need to be complemented with a fine proposal density \( q_l \) in order to form a proposal \( \theta_j^l \) of suitable dimension. Further, compared to MHMCMC,
the acceptance probability also needs to be adapted to avoid bias from the coarse proposals.

The efficiency gain of this multilevel algorithm over MHMCMC can be analyzed theoretically for models where PDE theory provides approximation error bounds for given cost. In [9, Thm. 3.4], the following cost estimate was proven for a Poisson equation model in $d$ dimensions similar to the Poisson model we use later. It is based on the mean square error of the estimate of $\mathbb{E}_\mu[Q]$, either directly computed via MHMCMC or from the telescoping sum eq. (2) in the MLMCMC case.

The computational cost $C_\varepsilon$ required to achieve a mean square error below $\varepsilon$ is then bounded by

$$C_\varepsilon^{\text{MCMC}} \leq \varepsilon^{-(d+2) - \delta} \quad \text{and} \quad C_\varepsilon^{\text{MLMCMC}} \leq \varepsilon^{-(d+1) - \delta}$$

respectively, where $\delta$ is a model specific constant. In this setting, which should be representative of many PDE models with model hierarchies based on mesh width, the cost for the multilevel method is therefore one order below the single-level one.

The hidden constants in the bounds are proportional to the integrated autocorrelation times of the Markov chains. On the finest level in the multilevel approach this factor is essentially reduced to one [10], leading to significant additional computational gains.

It is beyond the scope of this work to also prove the assumptions underlying the theoretical error estimates for our more complex tsunami model. This would be a particularly challenging task since we do not only rely on mesh refinement, but change bathymetry data across levels for improved solver performance. However, since our numerical PDE solver should fundamentally exhibit comparable scaling behaviour, we can reasonably expect similar gains in efficiency by employing the multilevel method in that setting as well.

## 3 Models

We will apply parallelized MLMCMC to two different models: The widely used and cheap to compute Poisson equation modelling single-phase subsurface flow as well as a more expensive and realistic Tsunami model. The former serves as a well-known reference, and due to its low cost is also suitable to gauge the MLMCMC parallelization in relatively short scaling experiments. The latter, due to its more interesting properties, allows us to demonstrate the practical value of the method, as well as the opportunities in creating model-specific hierarchies.

### 3.1 Poisson Model

In this example, our forward model maps a parameter $\theta$ that models the uncertainty in the diffusion coefficient to the solution of the Poisson PDE evaluated at certain points. The inverse problem consists in estimating the underlying parameter $\theta$ from given synthetic data while taking uncertainty into account.

Specifically, we solve the PDE

$$\nabla \cdot (\kappa(x, \theta) \nabla u(x, \theta)) = 0 \quad \text{for} \quad x \in \Omega \text{ and } \theta \in \Theta,$$

where we choose the domain $\Omega := [0, 1]^2$ and $\Theta := \mathbb{R}^m$. As boundary conditions, we apply $u(x) = 0$ on the left, $u(x) = 1$ on the right and natural Neumann boundary conditions at the remainder of the boundary. We model $\log(\kappa)$, the logarithm of the diffusion coefficient, as a zero-mean Gaussian random field with correlation length 0.15 and variance 1. In order to arrive at a finite dimensional representation of the field, we truncate use a Karhunen Loève (KL) expansion which we truncate after $m$ terms, i.e.

$$\log(\kappa(x, \theta)) = \sum_{k=1}^{m} \phi_k(x) \theta_k,$$

where $\phi_1, ..., \phi_m$ are the KL modes of largest wave length. Consequently, $\theta$ is a vector of KL coefficients. We implement the model in the DUNE framework [1] with a $Q_1$, Finite Element discretization on simple structured grids. To form a three-level model hierarchy for our MLMCMC method, we choose mesh widths of $\frac{1}{10}, \frac{1}{20},$ and $\frac{1}{50}.$ Across all three levels, we choose an identical parameter dimension $m = 113$; this specific number carries not much significance beyond being suitable for implementation in the circulant embedding [8] based random field generator $\text{dune-randfield}$.

In order to form a Bayesian inverse problem, we generate synthetic data based on a random field $\kappa(x, \theta)$, where $\theta$ is a fixed sample drawn from $\mathcal{N}(0, I)$ (shown in fig. 2).

The actual vector of measurements $y$ is then defined by solving the above Poisson problem for $\theta = \theta$, evaluating the solution $u$ at a grid of points: $(\frac{1}{10}, \frac{1}{20}, \frac{1}{50})^2$. Based on that, we define our likelihood $\mathcal{L}(y|\theta)$ to be a Gaussian $\mathcal{N}(F(\theta), \sigma_F^2 I)$ with $\sigma_F = 0.01$. Complementing it with a Gaussian prior $\pi(\theta) = \mathcal{N}(0, 4I)$, we complete our Bayesian inverse problem.

Note that, by generating synthetic measurements directly from our forward model, we commit an ‘inverse crime’ [4, p. 179]. In realistic applications a model error is inevitable, making it significantly harder to recover the underlying parameters from data accurately. However, for this problem we intentionally accept this simplification: Our focus in this case is algorithmic scalability and not a fully realistic setting. Further, verifying the correctness of the UQ method is somewhat easier without a model error.

As our QOI, we define $Q(\theta) = \kappa(x_k, \theta)$ where the $x_k$ form a grid of width $\frac{1}{50}$. This captures the parameter field we seek and, as necessitated by the telescoping sum in eq. (2), allows for a consistent dimension in QOI even when varying the parameter dimension across levels.

![Figure 2: Random field realization $\log(\kappa(\cdot, \hat{\theta}))$ and parameter field $\kappa(\cdot, \hat{\theta})$ from random field realization used for synthetic data.](image-url)
3.2 Tsunami Model

Tsunami propagation is typically modelled by some variant of the shallow water equations [2, 21, e.g.], allowing for a simulation in only two dimensions. The shallow water equations are obtained via depth-averaging of quantities (esp. momentum) from the more complicated three-dimensional Navier-Stokes equations, based on the modeling assumption that horizontal length scales are considerably greater than the vertical length scales.

In this setting, we concentrate on the basic shallow water equations with bathymetry source terms (neglecting friction terms or more advanced models for with non-hydrostatic corrections). The resulting equations can be written in first-order hyperbolic form as

\[
\begin{pmatrix}
\frac{\partial}{\partial t} h \\
\frac{\partial}{\partial t} u \\
\frac{\partial}{\partial t} v
\end{pmatrix} + \nabla \begin{pmatrix}
u h \\
u u h \\
u v h
\end{pmatrix} + 
\begin{pmatrix}
h g \partial_x (b + h) \\
0 \\
0
\end{pmatrix} = 0,
\]

where \( h \) denotes the height of the water column, \((u, v)\) the horizontal flow velocity, \( g \) gravity and \( b \) denotes the bathymetry. This hyperbolic system of equations is supplemented by a set of suitable initial and boundary values.

We discretise with an ADER-DG method as proposed in [12]. It is essentially a predictor-corrector scheme. A high-order solution is found element-locally and then corrected to take into account neighbors by solving Riemann problems along element interfaces. To resolve known high-order issues such as the Gibbs phenomenon a corresponding a-posteriori finite volume sub-cell limiter is applied [13]. This limiter detects and revokes problematic ADER-DG solution candidates and recomputes them with a robust Finite-Volume scheme (cf. [27, 28] for details), following the approach by LeVeque et al. [21]. At coastlines the schemes relies entirely on the Finite-Volume limiter, to correctly treat inundation.

As a large-scale example, we invert data from the Tohoku tsunami, which occurred subsequent to an earthquake in the Japan trench in 2011. We assume that the only significant sources of the tsunami are the displacements of the sea floor. In order to initialise the tsunami, we can impose the displacements as an instantaneous deformation of the bathymetry in the resting-lake case – compare respective modeling approaches by Saito et al. [31] or Madden et al. [23]. By keeping the water column constant, the change of the bathymetry is directly translated to the sea surface and generates the tsunami. Gravity, as the main acting force, initiates the propagation of the wave. The tsunami then evolves as a gravity wave. As reference solution for the initial displacements of the ocean floor we use respective simulation results provided by Galvez et al. [14]. The bathymetry data has been obtained from GEBCO 1.

Our goal is to obtain the parameters describing the initial displacements from the data of two available buoys located near the Japanese coast. Some of these parameters are reasonably well-known already, these include location of the hypocenter, length, width, and to some extent depth. Other parameters such as uplift are more difficult to estimate. In these tests we estimate the location of the initial displacement. The prior cuts off all parameters which would lead to an initial placement which is too close to the domain boundary. Figure 3 shows the cut-off values used. Some parameters may lead to unstable models, e.g. a parameter which initialises the tsunami on dry land, in this case we have treated the parameter as unphysical and assigned an almost zero likelihood.

In order to compute a likelihood of a given set of parameters given the simulation results we use a weighted average of the maximal wave height and the time at which it is reached. The likelihood is given by a normal distribution \( N(\mu, \Sigma) \) with mean \( \mu \) given by maximum waveheight \( \max(h) \) and the time \( t \) at which it is reached for the the two DART buoys 21418 and 21419.\(^2\) The covariance matrix \( \Sigma \) depends on the level, but not the probe point.

\(^1\)https://www.gebco.net/
\(^2\)This data can be obtained from NDBC https://www.ndbc.noaa.gov/
While parallelization in classical MC is trivial due to the independence of the samples, MCMC introduces data dependency through proposals depending on the previous step. In the case of MLMCMC, however, we clearly need to provide a hierarchy of models. Further, a QOI may be provided. This is implemented as a separate function call, since discarded samples in MCMC do not contribute to the QOI computation, and potentially expensive evaluations can thus be skipped.

### 4 PARALLEL MLMCMC IMPLEMENTATION

In this section we describe the new, highly scalable parallel implementation of MLMCMC used for the experiments in this paper. While parallelization in classical MC is trivial due to the independence of the samples, MCMC introduces data dependency through proposals depending on the previous step. In the case of MLMCMC as in [9, 10], we use coarser chain samples as proposals, which also introduces data dependency between levels. There are, however, multiple opportunities for parallelizing MLMCMC:

- **Models**: The forward models themselves may be parallelized. In fact, for large models like the tsunami model we introduce in section 3.2, that is inevitable anyway due to memory constraints.
- **Chains**: Instead of running a single Markov chain, multiple chains can be run in parallel and their samples combined. It is beneficial to not purely rely on this approach though since each chain requires a burn-in phase.
- **Levels**: Contributions to the multilevel telescoping sum in eq. (2) can be evaluated in parallel. Exploiting all those clearly introduces significant technical complexity. Therefore we provide our implementation as part of the MUQ C++ library [26]. The main goals of this implementation are:
  - **Parallelism**: All of the above levels of parallelism are supported.
  - **Simple user interface**: We hide the intricate details of communication from the user. The algorithm can be tweaked, but to get started the defaults suffice.
  - **Model-agnosticity**: The MLMCMC algorithm as detailed in section 2 only requires simple forward evaluations of the model. This theoretically allows coupling to arbitrary forward models without any need for, e.g., derivatives of the model map. We retain this model agnosticism in the sense that any forward model that can be called (possibly through wrappers) from C++ can be used.
  - **Modularity**: MUQ is, from the ground up, designed as a modular framework. Its modularity is closely modelled after the respective mathematical objects. We extend this concept to our parallel MLMCMC implementation by building on top of MUQ’s existing MCMC stack and, as detailed in the following, constructing modular parallel units.

#### 4.1 Model interface

MUQ provides an abstract interface for sampling algorithms, including MCMC type methods. In its most basic form, a model can be provided by implementing an AbstractSamplingProblem (see fig. 6). Our parallel MLMCMC implementation supports the same model interface, allowing to quickly use various UQ methods on a single model implementation. The interface requires the implementation of the density to sample from; in case of our Bayesian inverse problem, that is the posterior density in eq. (1). Further, a QOI may be provided. This is implemented as a separate function call, since discarded samples in MCMC do not contribute to the QOI computation, and potentially expensive evaluations can thus be skipped.
When implementing the MIComponentFactory interface, the SamplingProblem method is to return a model for a given model index, while FinestIndex specifies the index of the finest model the user provides, corresponding to $L$ in algorithm 2. CoarseProposal specifies how proposals are being drawn from coarser chains, and Interpolation determines how these are combined with a finer chain proposal as defined in Proposal.

In order to support parallel models, the user may modify ParallelizableMIComponentFactory, which receives an MPI communicator. The user can pass a parcer communicator to the model, this is essentially just an MPI C++ wrapper used in MUQ.

Note that, while in this paper we entirely focus on the multi-level case, the implementation actually supports a generalization to a multiindex method. Therefore, components are named MI for multiindex rather than ML, and the model hierarchy is indexed by a more general multiindex structure rather than an integer index.

4.2 Internal architecture

Our parallel process layout is shown in fig. 8. We define the following roles:

- **Fixed roles:** These are assigned to specific processes at the start of the parallel method. All other processes wait to be assigned a dynamic role.
  - **Root:** This process is responsible for launching the parallel method, assigning tasks to other processes and requesting collectors to begin collecting a certain number of MCMC samples. It is also the best place for users to implement custom (possibly adaptive) sampling strategies.
  - **Phonebook:** The phonebook tracks what dynamic roles processes are currently assigned to. Most importantly, it tracks which chains are currently sampling and which ones hold new samples ready to be picked up by other processes. Further, the phonebook can infer from that the computational load on a given level, since the relation between requested samples and assigned resources is available here. Therefore, it is also the key component in dynamic load balancing across levels and chains.
  - **Dynamic roles:** These roles may be assigned and reassigned at any time. In particular, this permits dynamic load balancing and possibly more advanced sampling strategies. Workers and controllers solve forward models, where workers share the load of running a single model evaluation and controllers additionally run the inherently sequential MCMC chains; consequently, they are assigned (and, in case of dynamic load balancing, reassigned) synchronously. This is facilitated by Message Passing Interface (MPI) subcommunicators, which are passed through to and should be used by the user’s model.
    - **Worker:** Workers are responsible for running the user’s forward model, specifically the user’s implementations of AbstractSamplingProblem. Mathematically, they provide parallelized evaluations of the posterior and quantity of interest on a given level for a given parameter. They listen to their respective controller’s ParallelAbstractSamplingProblem to signal the beginning of a evaluation for a specific parameter $\theta$. Once the signal arrives, they execute the user-implemented LogDensity method. Since all workers of a work group are called synchronously, the user can easily supply models assuming lock step parallelism.
    - **Collector:** Each controller is responsible for running a multilevel MCMC chain according to algorithm 2. Specifically, a controller on level $l$ contains a chain on level $l$ and one on $l−1$, to compute the level $l$ correction of the telescoping sum eq. (2) (with the obvious exception of the coarsest level 0). An instance of ParallelAbstractSamplingProblem is set up for each model needed. It provides an intermediate layer between instances of the user-implemented AbstractSamplingProblem instances running on multiple workers and the controller, allowing to transparently distribute model executions. As a result, neither the user nor the inherently sequential MLMCMC chains need to concern themselves with synchronizing worker processes to run forward models in lock step. The chains themselves are implemented using existing MUQ components: They are SingleChainMCMC instances with MCMCKernel implementations matching the acceptance probability of algorithm 2. Drawing samples from coarser chains as proposals is implemented with an MCMCProposal requesting coarser samples from other controllers via the phonebook process.

Note that this architecture is defined on process level (more specifically, in terms of MPI ranks). Thread-level parallelism can easily be exploited by worker processes. In fact we make use of this through Intel Threading Building Blocks (TBB) in our Exascale Hyperbolic PDE Engine (ExaHyPE) Tsunami model, since ExaHyPE
exhibits better performance characteristics with few MPI ranks per node each making use of several threads.

In order to make the parallel architecture as modular as sequential MUQ code, each of the above roles provide an MPI interface based on requests mimicking function calls. This allows recombining the parallel components in analogy to object orientation in order to implement other algorithms as well. For example, work groups as introduced above are based on a ParallelAbstractSamplingProblem, which in turn can be used to easily employ any sequential sampling algorithm on a parallelized model. Likewise, the phonebook could be swapped out for an alternative implementation with identical MPI interface, allowing for alternative load balancing strategies.

4.3 Load balancing

Data dependencies in MLMC (see algorithm 2) introduce a load balancing problem, since coarser chains need to provide proposals to finer ones only until the desired number of fine samples is computed. Estimating the ideal distribution of computational resources across levels is far from trivial or outright impossible in realistic applications, especially when adaptively determining the number of samples per level.

Therefore, our parallel MLMC implementation provides a load balancing mechanism to reassign worker processes to different tasks once samples on another level are more critical to runtime. Figure 9 illustrates this load balancing mechanism for a small test run.

Load balancing is implemented as part of the phonebook rank, since it keeps track of how samples are passed around. Levels with low load are detected when samples on that level are provided but not quickly picked up, while a high load is in turn detected when sample requests remain queued. Unanswered sample requests originating from other chains are given a higher impact than requests originating from collector processes, since the first case implies chains waiting and therefore bad machine utilization.

Models may have strongly varying run times. A new group of processes assigned to a certain level only reduces that level’s load once it actually provides its first sample. This implies the danger of reassigning tasks too frequently or too infrequently. In order to stabilize the load balancer, the respective model run times are inferred by the phonebook process by the frequency of samples provided. Based on that, scheduling will only take place at the time scale of the respective model evaluations.

Note that this load balancer is unaware of the specific types of proposals or MCMC kernels being executed. As a result, it can, for example, also be applied in the MLMC setting. Also, all model specific tuning parameters are determined dynamically, so no user intervention is required.
Table 3: Multilevel properties of Poisson application. For each level $l$ of mesh width $h_l$ with associated degrees of freedom (DOFs), we show the computational cost $t_l$, the number of subsamples $\rho_l$, and variance for a single representative component of $Q$.

| Level $l$ | $h_l$ | DOFs | $t_l$ [ms] | $\rho_l$ | $\tau_l$ | $\mathbb{V}[Q_0]$ or $\mathbb{V}[Q_l - Q_{l-1}]$ |
|---|---|---|---|---|---|---|
| 0 | 10 | 289 | 3.35 | 206 | 137.3 | $1.501 \times 10^{-2}$ |
| 1 | 2 | 6425 | 45.64 | 17 | 11.2 | $1.121 \times 10^{-3}$ |
| 2 | 256 | 66049 | 931.81 | 0 | 1.05 | $4.165 \times 10^{-5}$ |

Figure 10: Synthetic “true” field (left) and expected value of multilevel estimator (right).

Figure 11: Scalability of the Poisson model problem for $10^4$, $10^5$ and $10^6$ samples on levels 0, 1 and 2 respectively. Subsampling rates etc. are chosen according to table 3. The problem setup remains constant as the number of processors is increased.

5 NUMERICAL RESULTS

5.1 Poisson application and scalability

In order to fully specify the MLMCMC algorithm for the given problem, it is enough to set a Gaussian proposal on the coarsest level. We choose $N(0, 3I)$ in order to roughly match the prior. Since we have identical parameter dimensions across levels, no fine level proposals are needed.

The MLMCMC method run with $10^4$, $10^5$ and $10^6$ samples on levels 0, 1 and 2 exhibits properties detailed in table 3 and captures the main features of the parameter field underlying our synthetic data (see fig. 10). Clearly some higher frequency detail is not recovered. This, however, is expected due to the limited number of KL modes we include in our parameter space. Note that we can only recover this up to a scaling factor, since the solution is only determined by the parameter field up to a factor. Here, the choice of prior essentially determines the scaling of the solution we observe.

For a more detailed analysis of Bayesian inverse problems based on Poisson equation in MLMCMC, we refer to the original MLMCMC publication [9, 10].

In order to investigate parallel scalability of our MLMCMC implementation, we conduct weak and strong scaling experiments on the BwForCluster MLS&WISO Production HPC system. The partition we used consists of nodes with two 16-core Intel Xeon E5-2630v3 CPUs and 64 gigabytes of memory.

As forward model, we use the Poisson model, since its low computational demand allows us to stress the parallelized MLMCMC algorithm itself by running a large number of chains and samples.

We use the same inverse problem detailed above, even though the particular inverse problem does not affect the algorithm’s communication patterns and therefore parallel scalability.

For the strong scaling setup, we draw $10^4$, $10^5$ and $10^6$ samples on levels 0, 1 and 2 respectively. We further set subsampling rates according to table 3, and enable dynamic load balancing. As the timing results in fig. 11 show, we achieve linear speedup until relatively large burnin phases and suboptimal load balancing due to few samples per chain occur.

Technically the observed speedup even slightly exceeds linear. That is simply due to the fact that a fixed number of the processes is reserved for book keeping tasks (i.e. the root, phonebook and collector processes). As a result, for increased number of processes, a larger fraction contributes to parallel speedup by generating samples.

In our weak scaling test, we begin with the same setup as in the strong scaling setting. In particular, we again choose to compute $10^4$, $10^5$ and $10^6$ samples on levels 0, 1 and 2, and solve this problem using 64 processes. We then expand to a range from 32 to 1024 processes while scaling the number of samples on each level linearly in accordance with the number of processes.

The parallel efficiency (given in blue in Figure 12) is measured here as $\frac{t_{ref}}{t_N} \cdot 100\%$, where $t_{ref}$ is the quickest time taken over all runs and $t_N$ is the time taken on $N$ ranks. The initial increase to over 100% efficiency is due to the overhead of phonebook and collector ranks. We achieve fairly consistent results of up to 80 seconds of total run time except for the largest run. The latter is a very extreme scenario though, since the extremely short run time of the coarsest model leads to a significant load on the communication
Figure 12: Weak scalability and parallel efficiency of the Poisson model problem. At 64 cores $10^4$, $10^3$ and $10^2$ samples are computed on levels 0, 1 and 2 respectively. The number of samples is modified linearly with the number of processors.

infrastructure. We therefore consider it reasonable to assume that exceeding the ideal range should only occur for significantly larger numbers of processes in more realistic applications.

5.2 Tsunami application

As a second example of the MLMCMC method we invert data from the Tohoku tsunami. We find the parameter distribution describing the initial displacements from the data of two available buoys located near the Japanese coast. The scenario is described in more detail in Section 3.2. The MLMCMC method with three levels computes 800 samples on level 0, 450 on level 1 and 240 on level 2 with a subsampling rate of 25 on level 0 and 5 on level 1.

These tests were run on up to 72 Intel Skylake Xeon Platinum 8174 nodes of SuperMUC-ng consisting of 48 cores each. The tests were run using Intel’s TBB for parallelisation over the 48 cores of each node and MPI for parallelisation across nodes. Each ExaHyPE run used exactly one full node. The runtime for each forward model evaluation was on average 7.38 seconds on level 0, 97.3 seconds on level 1 and 438.1 seconds on level 2. These runtimes have a large variability as the model’s timestep depends on the uncertain parameters, making it a challenging test for the scheduling infrastructure. In total 61, 250 level 0, 34,500 level 1 and 240 level 2 forward model evaluations were required for this test.

Again, due to constant parameter dimension across levels, we only need to choose a proposal density for the coarsest level. Like before, we choose Adaptive Metropolis [16, 17] provided by MUQ. As initial prior we set $N(0, 10I)$ and update every 100 steps.

Figure 13 shows the resulting samples on level 0 and 1. The expected values $E(Q_0)$, and $E(Q_l - Q_{l-1})$ are shown as dashed line. The red marker shows the point $(0, 0)$. The point $(0, 0)$ is the position of the initial displacements as estimated in [14].

In order to illustrate how MLMCMC links coarser and finer level posteriors in order to obtain fine level corrections, fig. 14 shows how samples on level $l$ relate to the level $l - 1$ samples that served as their respective coarse proposals. The result can be thought of as a transformation between corresponding coarser and finer distributions, even though only in a non-deterministic sense. Since

Figure 13: The three-level tsunami test case, each point represents an accepted sample at level $l = 0, 1, 2$. The dashed lines show the expected value $E(Q_0)$ or $E(Q_0) + \sum_l E[Q_l - Q_{l-1}]$ with the point $(0, 0)$ in red for reference.
in this application we choose the QOI to be the uncertain parameter itself, the estimate of the terms in the telescoping sum (eq. (2)) actually correspond to the mean of the corrections displayed here. Variances and expected values are given in Table 4. The relatively cheap samples on level 0 provide a good initial estimate of the posterior, which are improved by the more expensive models utilising the full bathymetric data. In contrast to the Poisson case, we do not observe variance reduction across levels. This is somewhat expected, as the modified bathymetry does not permit the construction of a level hierarchy fulfilling the theoretical assumptions made by MLMCMC based on a priori error estimates. We do, however, still have the benefit of well-informed proposals on finer levels driven by coarser level chains.

6 CONCLUSION

In this paper we have presented a new parallelization strategy for MLMCMC as well as a model-agnostic implementation as part of the open-source and modular MUQ library. We have demonstrated the effectiveness of algorithm and implementation at solving Bayesian inverse UQ problems on complex and large-scale PDE models, highlighting opportunities in model-specific coarsening strategies in the process.

In order to verify our parallelization approach, we presented both strong and weak scaling results. An important feature of the work is the applicability of the provided MUQ interface, which allows easy coupling to other models and software packages. This we have shown by applying it to a Poisson model problem implemented in the DUNE framework and a tsunami model implemented in the ExaHyPE-Engine.

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