ACCURATE UNCERTAINTY QUANTIFICATION USING INACCURATE COMPUTATIONAL MODELS

PHAEDON-STELIOS KOUTSOURELAKIS *

Abstract. This paper proposes a novel uncertainty quantification framework for computationally demanding systems characterized by a large vector of non-Gaussian uncertainties. It combines state-of-the-art techniques in advanced Monte Carlo sampling with Bayesian formulations. The key departure from existing works is the use of inexpensive, approximate computational models in a rigorous manner. Such models can readily be derived by coarsening the discretization size in the solution of the governing PDEs, increasing the time step when integration of ODEs is performed, using fewer iterations if a non-linear solver is employed or making use of lower order models. It is shown that even in cases where the inaccurate models provide very poor approximations of the high-fidelity response, statistics of the latter can be quantified accurately with significant reductions in the computational effort. Multiple approximate models can be used and rigorous confidence bounds of the estimates produced are provided at all stages.

Key words. uncertainty quantification, Monte Carlo, Bayesian, nonparametric regression

AMS subject classifications. 65C05,62F15,62G08

1. Introduction and examples. Scientists have come to recognize the stochastic aspects inherent in several physical systems and processes and seek ways to quantify the probabilistic characteristics of their behavior. Their analysis tools are usually restricted to elaborate legacy codes which have been developed over a long period of time and are generally well-tested. They do not however include any stochastic components and their alteration is commonly impossible or ill-advised. In many problems of engineering or physical interest the only feasible solution for uncertainty quantification is provided by non-intrusive methodologies.

Traditionally, two approaches have have attracted most attention. On one hand methods based on polynomial chaos expansions (PC, [41]) and on the other techniques anchored around Monte Carlo simulations. PC models, although originally developed as intrusive techniques ([18]), have grown into prominence in recent years with the development of non-intrusive, stochastic collocation approaches ([31, 42, 15]). They are based on a representation of the random input by a finite number of uncorrelated random variables (usually normally distributed) and orthogonal polynomials (usually Hermite). The solution or output process is expressed with respect to the same basis and the coefficients of the expansion are determined by calculating weighted residuals or using a collocation approach. Although mathematically elegant, PC-based approaches utilize a second-order matching (up to the autocovariance function) of the input processes which does not account for important higher order statistics that might affect the system’s response. The computational effort grows with the number of random variables used to approximate the input which also adversely affects the accuracy, especially in the stochastic collocation version, as an interpolation in a very high dimensional space is required.

Standard Monte Carlo simulations require a minimal implementation overhead as the coupling with existing deterministic solvers is trivial. Most often than not however, in systems of physical interest, each of the runs of the forward solver requires several CPU-hours and multiple processors. Even though the convergence rate $O(1/\sqrt{N})$...
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(Where \( N \) is the number of independent samples) is independent of the dimensionality of the random input, it is sufficiently slow to constitute the method impractical or infeasible as several calls to the forward model have to be made to achieve good accuracy. Another disadvantage of classical Monte Carlo is that it does not directly provide confidence intervals for the estimates produced as those are usually based on asymptotic results (i.e. when \( N \to \infty \)) and the central limit theorem ([36]). Recent years have seen significant progress in the development of advanced Monte Carlo techniques which employ evolutionary strategies in combination with Markov Chain Monte Carlo (MCMC) and Importance Sampling ([17, 33, 25]) but also techniques making use of sensitivity derivatives to increase the accuracy of sampling methods ([5, 6]). In many cases this has led to algorithms which require 10 or 100 times less samples in order to produce estimates of the same accuracy ([2]).

Despite these significant improvements, the associated computational effort can still be tremendous for systems of practical interest, which are characterized by a large number of input uncertainties and when even a modest number of 100 or 1000 runs can be infeasible. It is obvious that a new perspective is needed. In the author’s opinion this can be achieved if analysis goes beyond the black-box solver as the only means of probing the problem of interest. Indeed in many situations, several other pieces of knowledge and structural elements of the problem at hand, are readily available but left unexploited. For example, quite frequently the computational models of interest involve the solution of a system of PDEs using Finite Elements (FE) or Finite Differences (FD). These imply the spatio-temporal discretization of the governing differential equations and quite often the mesh sizes or time steps have to be particularly small in order to capture the salient features of the solution. Without recourse to rigorous mathematical proofs, it is well-known that an FE solver that operates on a coarser spatio-temporal grid can give an approximate solution at a lower computational cost as the system of equations that need to be solved are smaller. The deviation from the high-fidelity (or accurate) solution can be significant but in principle this approximate solver can be used to obtain some, inaccurate of course, information about our accurate model. As it will be demonstrated in the sections to come, it is not important if the solutions of the approximate solver deviate significantly from the accurate, but it suffices that they exhibit some sort of dependence. It is this dependence that we will exploit in a general and rigorous computational framework in conjunction with a few, carefully selected runs of the full, high-fidelity model. It should be noted that the same idea has also been advocated in [22, 24].

We will make no claims about the optimality of the approximate solvers selected. In fact as it will be demonstrated in the examples even crude approximations can yield impressive increases in computational efficiency. Furthermore, the framework proposed allows for the introduction of several such approximate models similar to the way one would elicit opinions from multiple experts before making a decision. In that respect even low-order, fast PC models can be utilized. Such an approach can be employed even in cases where no accurate computational model exists but rather we have to rely on experiments in order to collect the necessary information about the system. Since conducting experiments can be costly and time consuming it is desirable to minimize them by making use of approximate computational models that might be available ([38, 39, 14]).

To that end we investigate Bayesian alternatives to classical uncertainty quantification techniques with emphasis on problems with a large number of input uncertainties. In particular we formulate a regression problem that establishes the connection
between the response values from the approximate and high-fidelity models. This is achieved using a flexible, non-parametric Bayesian model that employs a very efficient Sequential Monte Carlo (SMC) inference algorithm. In contrast to [22], the proposed scheme makes sole use of approximate model outputs and therefore is highly suited for problems with high-dimensional input uncertainties which precludes the derivation of statistical emulators. We make no a priori assumptions about the relation between inaccurate and accurate model outputs which might limit our ability to mine any information the former provide. An added advantage of this approach is that prior knowledge or expertise of the analyst regarding the relationship between approximate and accurate solvers can be readily incorporated in the prior distributions.

Once the relation between approximate (inexpensive) and high-fidelity (expensive) models is established, the posterior distribution can be readily used to obtain estimates and credible intervals on the output statistics of interest. These can in turn be used to adaptively improve the accuracy of the regression by performing runs of the expensive solver in regions that contribute most significantly to these uncertainties. It should be noted that these posterior intervals incorporate uncertainty about the regression model parameters as a fully Bayesian inference scheme is employed. Outputs from multiple approximate solvers can be used simultaneously in order to improve the accuracy and computational efficiency of the model. This resembles mixtures of experts models that are commonly used in various statistical applications, as each approximate solver provides some, generally incomplete, information about the high-fidelity model which is then aggregated in order to obtain the best possible estimate.

2. Proposed Approach. Let \((\Omega, F, P)\) be a complete probability space, where \(\Omega\) is the event space, \(F\) the \(\sigma\)-algebra, and \(P\) the probability measure. Consider the following stochastic differential equation:

\[
L(u(z, t); \xi(\omega)) = f(z, t; \xi(\omega)), \quad z \in \mathcal{D}, \; t > 0
\]

(2.1)

defined on the domain \(\mathcal{D} \subset \mathbb{R}^q (q = 1, 2, 3)\) with appropriate initial/boundary conditions which might also depend on the vector of uncertainties represented by \(\xi(\omega) : \Omega \rightarrow \mathbb{R}^d\). We are particularly interested in the most general and difficult case where \(\xi\) is of very large dimension (i.e. \(d \gg 1\)) and it should not or cannot be condensed using any of the standard dimension reduction techniques (e.g. PCA). Let \(u(z, t; \xi(\omega))\) denote the solution process which satisfies Equation (2.1) for \(P\)-almost everywhere and naturally depends on \(\xi\). We are interested in the statistics of the solution itself or of a function thereof which we denote by \(y(\xi) : \mathbb{R}^d \rightarrow \mathbb{R}^r\) emphasizing the dependence on the vector of input uncertainties \(\xi\). We further postulate the existence of a forward solver of the linear/nonlinear equation in Equation (2.1) that corresponds to a deterministic version of the differential operator \(L\) (for fixed \(\xi\)). In general however one might consider reduced versions of the above problem (i.e. dependence only on time or space) or even systems (and associated computational models) which are not governed by SPDEs but nevertheless characterized by a high-dimensional vector of input uncertainties \(\xi\). For illustration purposes we will restrict the presentation to the case that \(y\) is scalar (i.e. \(r = 1\)). Naturally \(y\) may depend on other deterministic parameters which are omitted for economy of notation.

The input uncertainties \(\xi\) are characterized by a probability density \(\pi_\xi\). In order for the problem to be well-posed, \(\pi_\xi\) need not be known analytically but it suffices to be able to draw samples from \(\pi_\xi\). Our goal is to calculate statistics of the response,
e.g.:

\[ Pr[y \in A] = \int 1_A(y(\xi)) \pi_\xi(\xi) \, d\xi \]  \hspace{1cm} (2.2)

where \( 1_A \) is the indicator function of a \( \pi_\xi \)-measurable subset \( A \), or:

\[ E[h(y)] = \int h(y(\xi)) \pi_\xi(\xi) \, d\xi \]  \hspace{1cm} (2.3)

where \( h \) is any \( \pi_\xi \)-integrable function.

Quite often the statistics of interest involve very rare events (i.e. \( Pr[y \in A] << 1 \)), as is the case for example in molecular dynamics simulations where transition to a local minimum of the free-energy landscape happens infrequently or in estimating reliability of mechanical components. In other cases we are interested in expectations of multimodal functions \( h \) as is the case in nonlinear dynamical systems where small perturbations in the input \( \xi \) can lead to significant differences in the (long-term) response. Due to the large variance of the integrands in Equations (2.2) or (2.3), several calls to the forward solver have to be made (to calculate the response \( y \) for various \( \xi \)’s) which imply a significant or insurmountable computational burden, particularly in cases where each of these calls requires several CPU-hours on multiple processors.

To address these issues we postulate the existence of \( M \) approximate forward solvers \( x_m(\xi) : \mathbb{R}^d \to \mathbb{R}^r \), for \( m = 1, \ldots, M \), as those discussed in the introduction and in the numerical examples to follow. Each of those provides approximations to the output of interest at a fraction of the computational cost. The latter requirement is the key in increasing the overall computational efficiency in the proposed framework, whereas the former condition can be interpreted very loosely. In fact it is acceptable that the \( x_m \)’s provide very poor estimates (i.e. \( y(\xi) - x_m(\xi) \) is relatively large) as long as there is some statistical dependence between them. In that sense \( x_m \) might not even correspond to the same output quantities, although in such cases the selection of reasonable approximate solvers can be less straightforward. For the purposes of this work, \( x_m \)’s are viewed as (potentially) biased and partial predictors of the output \( y \). Our goal is to quantify the information these predictors provide for the purposes of estimating statistics of \( y \) at a fraction of the computational cost. In particular, if \( x = (x_1, x_2, \ldots, x_m) \), Equation (2.2) can be rewritten as:

\[ Pr[y \in A] = E_x [Pr[y \in A \mid x]] \]
\[ = \int Pr[y \in A \mid x] \pi_x(x) \, dx \]
\[ = \int \left( \int 1_A(y) \, p(y \mid x) \, dy \right) \pi_x(x) \, dx \]  \hspace{1cm} (2.4)

and Equation (2.3):

\[ E[h(y)] = E_x [E[h(y) \mid x]] \]
\[ = \int E[h(y) \mid x] \pi_x(x) \, dx \]
\[ = \int \left( \int h(y) \, p(y \mid x) \, dy \right) \pi_x(x) \, dx \]  \hspace{1cm} (2.5)

Hence it is apparent that estimates of \( Pr[y \in A] \) and \( E[h(y)] \) can be obtained as long as the density \( \pi_x(x) = \int \delta(x - x(\xi)) \, \pi_\xi(\xi) \, d\xi \) and conditional \( p(y \mid x) \) are
known. Given that calls to the approximate solvers are computationally inexpensive in relative terms as it will be seen in the examples of section 3, \( \pi_x(x) \) can be readily evaluated using direct or advanced Monte Carlo techniques as those discussed in the introduction. The pivotal component is the conditional density \( p(y \mid x) \) which probabilistically quantifies the information that the predictors \( x \) provide about \( y \).

In one extreme, if \( y \) and \( x \) are statistically independent, knowledge of \( x \) is completely useless in furnishing information about \( y \) and \( p(y \mid x) = p(y) \). Hence the proposed framework cannot offer any improvement. In the other extreme, there exists an injective, deterministic mapping between the two quantities, i.e. \( y = g(x) \) and therefore knowing \( x \) and its statistics translates straightforwardly to \( y \) since \( p(y \mid x) = \delta(y - g(x)) \). The proposed methodology is applicable to all cases except the one of independence between \( x \) and \( y \).

Critical to the feasibility of proposed framework is establishing a quantitative link between the reference \( y \) and approximate \( x \) outputs as described by \( p(y \mid x) \). This will be accomplished using computationally generated data that consist of pairs of \( \{(x_i = x(\xi_i), y_i = y(\xi_i))\}_{i=1}^n \) obtained by running low and high-fidelity solvers for the same \( \xi_i \). This is discussed in detail in the next 3 sub-sections. The task of utilizing the inferred models for the purposes of estimating Equation (2.4) or Equation (2.5) is discussed in sub-section 2.4 and illustrated in the examples of section 3.

### 2.1. Hierarchical Bayesian model.

We assume that the data have been rescaled so that \( x_i \in [0,1]^M \) and consider regression models of the form:

\[
y(\xi_i) = y_i = f(x(\xi_i); \theta) + \sigma Z_i \tag{2.6}
\]

where \( f \) is a function of the predictors \( x = (x_1, \ldots, x_M) \) and model parameters \( \theta \), and \( Z_i \) are i.i.d standard normal variates i.e. \( Z_i \sim \mathcal{N}(0,1) \) (if \( y \in \mathbb{R}^r \) then \( f, Z_i \in \mathbb{R}^r \)). Equation (2.6) postulates that, given the model parameters \( \theta \), for an input \( \xi_i \) for which the outputs of the approximate models are \( x_m(\xi_i) \), the target response \( y(\xi_i) \) is normally distributed with mean \( f(x(\xi_i); \theta) \) and standard deviation \( \sigma \), i.e.:

\[
y_i \mid x(\xi_i), \theta, \sigma \sim \mathcal{N}(f(x(\xi_i); \theta), \sigma^2) \tag{2.7}
\]

At first glance such a model seems highly restrictive as it is unlikely that \( y \) is normally distributed (given \( x \) and model parameters \( \theta \)). For that purpose we adopt a Bayesian formulation in which the model parameters \( \theta \) are assumed random and equipped with a distribution. This allows us to actually formulate a family of such models (each corresponding to a particular \( \theta \)) and even though conditionally on \( \theta \), \( y \) is normally distributed, marginally (when \( \theta \) are integrated out) non-Gaussian distributions can be considered.

Bayesian formulations differ from classical statistical approaches (frequentist) in that all unknown parameters are treated as random. As a consequence, the results of the inference process are not point estimates but distribution functions and multiple are possible. In fact any \( \theta \) in the support of the likelihood and the prior is admissible and the relative plausibility of various values is quantified by the posterior. Credible or confidence intervals can be readily estimated from the posterior which quantify inferential uncertainties about the unknowns.

The crucial ingredient is of course the prior specification, not only in terms of the functional form of \( p(\theta) \) but primarily in terms of the structural characteristics of the relation between \( y \) and \( x \) that is implied in Equation (2.6). It is easily understood, that any parameterization that depends on a finite number of \( \theta \) will be restrictive
no matter how large the family of models that it contains. Furthermore, in order
to be consistent with the principle of parsimony, prior models should make as few
assumptions as possible and allow their complexity to be inferred from the data.
To satisfy the aforementioned desiderata and overcome the shortcomings of existing
approaches, we propose the use of nonparametric priors ([40, 28]). As the term can be
misleading, we note that this does not imply lack of parameters but rather that the
number of parameters is not a priori fixed and can change as the data dictates. At
the core of such representations, lie simple basis functions, whose shape and location
are controlled by a few parameters. The key unknown is the cardinality of the model,
i.e. the number of such terms that are needed to provide a good interpretation of the
data. Consider the expansion:

$$f(x; \theta) = a_0 + \sum_{j=1}^{k} a_j K_j(x; \phi_j) \quad x \in D$$  \hspace{1cm} (2.8)

where $\theta = (k, \{\phi_j\})$, $K_j$ are kernels that serve as the basis functions of our represen-
tation and $\phi_j$ associated parameters. Expression (2.8) is motivated by the representer
theorem of Kimeldorf and Wahba ([23]), which states that the solution to the problem
of minimizing a goodness-of-fit loss function subject to a Reproducing Kernel Hilbert
Space norm penalty lies in a subspace represented as in Equation (2.8). Overcomplete
representations as in Equation (2.8) have been advocated because they have greater
robustness in the presence of noise, can be sparser, and can have greater flexibility in
matching structure in the data ([27, 1, 28]). One possible selection for the functional
form of $K_j$, that also has an intuitive parameterization, is isotropic, Gaussian kernels:

$$K(x; \phi_j = (\nu_j, \tau_j)) = \exp\{-\tau_j \|x - \nu_j\|^2\}$$  \hspace{1cm} (2.9)

The parameters $\tau_j$ directly correspond to the scale of variability of $f(x)$. Large $\tau_j$'s
imply narrowly concentrated fluctuations and large values slower varying fields. The
center of each kernel is specified by the location parameter $\nu_j$.

The parameters of the prior model adopted consist of:

- $k$: the number of kernel functions needed,
- $\{a_j\}_{j=0}^{k}$, the coefficients of the expansion in Equation (2.8). Each of those
can be a scalar or vector depending on the dimensionality of the high-fidelity
output $y$.
- $\{\tau_j\}_{j=1}^{k}$ the precision parameters of each kernel which pertain to the scale of
the unknown field(s), and
- $\{\nu_j\}_{j=1}^{k}$ the center locations of the kernels which are points in $[0, 1]^M$.

Let $\theta_k = \{(a_j)_{j=0}^{k}, \{\tau_j\}_{j=1}^{k}, \{\nu_j\}_{j=1}^{k}\} \in \Theta_k$ denote the vector containing all
the unknown parameters and $\theta = (k, \theta_k)$. If $k$ is also assumed unknown and al-
lowed to vary, then the dimension of $\theta_k$ is variable as well and $\Theta_k \triangleq (\mathbb{R}^{k+1})^r \times
(\mathbb{R}^{+})^k \times ([0, 1]^M)^k$. For example, in the case of two approximate solvers $(x_1, x_2)$
$(M = 2)$ and a scalar $y \ (r = 1)$, $\theta_k$ is of dimension $(k + 1) + k + (2k) = 1 + 4k$, i.e. $\Theta_k \triangleq \mathbb{R}^{1+k} \times (\mathbb{R}^{+})^k \times [0, 1]^{2k}$. In accordance with the Bayesian paradigm, all
unknowns are considered random and are assigned prior distributions which quantify
any information, knowledge, physical insight, mathematical constraints that is avail-
able to the analyst before the data is processed. Naturally, if specific information
about the relation between accurate $y$ and approximate $x$ outputs is available it can
be reflected on the prior distributions. We consider prior distributions of the following

...
form (excluding hyperparameters):
\[
p(k, \{a_j\}_{j=0}^k, \{\tau_j\}_{j=1}^k, \{x_j\}_{j=1}^k) \propto p(k) \\
\times p(\{a_j\}_{j=0}^k | k) \\
\times p(\{\tau_j\}_{j=1}^k | k) \\
\times p(\{\nu_j\}_{j=1}^k | k) \quad (2.10)
\]

In order to increase the robustness of the model and exploit structural dependence we adopt hierarchical prior models ([16]) as discussed in the sequence.

2.2. Prior Distribution. Pivotal to the robustness and expressivity of the model is the selection of the model size, i.e. of the number of kernel functions \(k\) in Equation (2.8). This number is unknown a priori and in the absence of specific information, sparse representations should be favored. This is not only advantageous for computational purposes, as the number of unknown parameters is proportional to \(k\), but also consistent with the parsimony of explanation principle or Occam’s razor ([21, 35, 32]). For that purpose, we propose a Poisson prior for \(k\):
\[
p(k \mid \lambda) = e^{-\lambda} \frac{\lambda^k}{k!} \quad k = 0, 1, \ldots, \infty \quad (2.11)
\]

For computational purposes, the aforementioned distribution is truncated beyond \(k_{\text{max}}\). The latter is selected based on computational limitations and defines the support of the prior. This prior allows for representations of various cardinalities to be assessed simultaneously with respect to the data. As a result the number of unknowns is not fixed and the corresponding posterior has support on spaces of different dimensions as discussed in more detail in the sequence. In this work, an exponential hyper-prior is used for the hyper-parameter \(\lambda\) to allow for greater flexibility and robustness i.e. \(p(\lambda \mid s) = s \exp\{-\lambda s\}\). After integrating out \(\lambda\) we obtain:
\[
p(k \mid s) \propto \frac{1}{(s + 1)^{k+1}}, \quad \text{for } k = 0, 1, \ldots, k_{\text{max}} \quad (2.12)
\]

The parameters \(\{\tau_j\}_{j=1}^k\) control the scale of variability in the relation between \(y\) and \(x\). If prior information about this is available then it can be readily accounted for by appropriate prior specification. In the absence of such information however multiple possibilities exist. We assumed \(\tau_j\) are a priori independent i.e. \(p(\{\tau_j\}_{j=1}^k) = \prod_{j=1}^k p(\tau_j)\) and a \(\text{Gamma}(a_\tau, b_\tau)\) prior was used for each \(\tau_j\):
\[
p(\{\tau_j\}_{j=1}^k \mid k, a_\tau, b_\tau) = \prod_{j=1}^k \frac{b_\tau^{a_\tau}}{\Gamma(a_\tau)} \tau_j^{a_\tau-1} \exp(-b_\tau \tau_j) \quad (2.13)
\]

This has a mean \(a_\tau / b_\tau\) and coefficient of variation \(1 / \sqrt{a_\tau}\). Diffuse versions can be adopted by selecting small \(a_\tau\). A non-informative prior \(p(\tau_j) \propto 1/\tau_j\) arises as a special case for \(a_\tau = 2\) and \(b_\tau = 0\) which is invariant under rescaling. Furthermore, it offers an interesting physical interpretation as it favors “slower” varying representations (i.e. smaller \(\tau\)’s). In order to automatically determine the mean of the Gamma prior, we express \(b_\tau = \mu_j a_\tau\) where \(\mu_j\) is a location parameter for which an Exponential hyper-prior is used with a hyper-parameter \(a_\mu\) i.e. \(p(\mu_j) = e^{-\mu_j / a_\mu}\). Integrating out the
controls the spread of the prior is modeled by the standard inverse gamma distribution for

distributions this can be incorporated in the model. Naturally if prior information is available about subregions with sig-

resulting to the following expression:

\[ \mu_j = \frac{1}{\tau_j} \]

where \( \Gamma(z) \) is the \((k + 1) \times (k + 1)\) identity matrix. The hyper-parameter \( \sigma_n^2 \) which

controls the spread of the prior is modeled by the standard inverse gamma distribution \( \text{Inv} - \text{Gamma}(a_0, b_0) \). It can readily be marginalized leading to the following prior

for \( a_j \)’s:

\[ p(a_j) \propto \frac{1}{(2\pi)^{k/2}} \sqrt{a_j} \exp\left(-\frac{a_j}{2}\right) \]

(2.14)

For the unknown kernel center locations \( \nu_j \), a uniform prior in \([0, 1]^M\) was used. Naturally if prior information is available about subregions with significant fluctuations this can be incorporated in the model.

Based on the aforementioned equations, the complete prior model is given by:

\[ p(\theta \mid s, a_\tau, a_\mu, a_\sigma) = \frac{1}{(s + 1)^{k+1}} \prod_{j=1}^{k} \frac{\Gamma(a_\tau + 1)}{\Gamma(a_\tau)} \frac{a_\tau^{a_\tau}}{\tau_j^{a_\tau+1}} \frac{1}{a_\mu} \frac{1}{(a_\tau \tau_j + a_\mu^{-1})^{(a_\tau+1)}} \]

(2.15)

\[ \times \frac{1}{(2\pi)^{(k+1)/2}} \left( b_0 + \frac{1}{2} \sum_{j=0}^{k} a_j^2 \right)^{(a_0 + (k+1)/2)} \]

(2.16)

Given \( n \) data pairs, \((x_i, y_i)_{i=1}^n\) the likelihood \( p(y_{1:n} \mid x_{1:n}, \theta) \) is:

\[ p(y_{1:n} \mid x_{1:n}, \theta) = \prod_{i=1}^{n} p(y_i \mid x_i, \theta) \]

\[ = \frac{1}{(2\pi)^{n/2}} \frac{1}{\sigma^n} \exp\left(-\frac{1}{2\sigma^2} \sum_{i=1}^{n} (y_i - f(x_i; \theta))^2\right) \]

(2.17)

A \text{Gamma}(a, b)\ prior was used for the variance \( \sigma^{-2} \) of the Gaussian error in Equation (2.6), which is conjugate to the likelihood above and can be readily marginalized resulting to the following expression:

\[ L_n(\theta) = p(\theta \mid (x_{1:n}, y_{1:n})) = \frac{\Gamma(a + n/2)}{(b + \frac{1}{2} \sum_{i=1}^{n} (y_i - f(x_i; \theta))^2)^{a+n/2}} \]

(2.18)

where \( \Gamma(z) = \int_{0}^{\infty} t^{z-1} e^{-t} dt \) is the gamma function.

The combination of the prior \( p(\theta) \) with the likelihood \( L_n(\theta) \) corresponding to \( n \) data points, give rise to the posterior density \( \pi_n(\theta) \) which is proportional to:

\[ \pi_n(\theta) = p_n(\theta \mid (x_{1:n}, y_{1:n})) \propto L_n(\theta) \ p(\theta) \]

(2.19)
Even though several parameters have been marginalized from the pertinent expressions, the corresponding posteriors can be readily obtained, or rather be sampled from, once the posteriors $\pi_n(\theta)$ has been determined. Of particular interest for prediction purposes is the variance $\sigma^2$ of the error term (Equation (2.6)). From Equation (2.18) and the conjugate prior model adopted for $\sigma^2$, it can readily be shown that the conditional posterior is given by a Gamma distribution:

$$
\pi_n(\sigma^{-2}, \theta) = p(\sigma^{-2}, \theta | (x_{1:n}, y_{1:n}))
= \pi_n(\sigma^{-2} | \theta) \pi_n(\theta | (x_{1:n}, y_{1:n})) \tag{2.21}
$$

and:

$$
\pi_n(\sigma^{-2} | \theta) = p(\sigma^{-2} | \theta, (x_{1:n}, y_{1:n}))
= \text{Gamma} \left( a + \frac{n}{2}, b + \frac{\sum_{i=1}^{n} (y_i - f(x_i; \theta))^2}{2} \right) \tag{2.22}
$$

In the context of Monte Carlo simulation, this trivially implies that once samples $\theta$ from $\pi_n$ have been obtained, samples of $\sigma^{-2}$ can also be drawn from the aforementioned Gamma.

It is worth pointing out, that Equation (2.20) defines a sequence of posterior densities with support on $\bigcup_{k=0}^{k_{max}} \{k\} \times \Theta_k$. Each $\pi_n$ corresponds to $n$ data points. It is easily understood that for small datasets, i.e. small $n$, the effect of the likelihood function $L_n$ will be subdued and the associated posterior $\pi_n$ will have fewer modes as it is dominated by the prior. As more data points are added and $n$ increases the contribution of the likelihood becomes more pronounced and the posterior will potentially exhibit more idiosyncratic characteristics. As a result the task of identifying these posteriors becomes increasingly more difficult for larger $n$. It is this feature that we propose of exploiting in the next section in order to increase the accuracy and improve on the efficiency of the inference process.

### 2.3. Bayesian Computation - Determining the Posterior

The posterior defined above is analytically intractable. For that reason, Monte Carlo methods provide essentially the only accurate way to infer $\pi_n$. Traditionally Markov Chain Monte Carlo techniques (MCMC) have been employed to carry out this task ([40, 1]). These are based on building a Markov chain that asymptotically converges to the target density (in this case $\pi_n$) by appropriately defining a transition kernel. While convergence can be assured under weak conditions ([29, 37]), the rate of convergence can be extremely slow and require a lot of likelihood evaluations. Particularly in cases where the target posterior has multiple modes, very large mixing times are required.

In this work we propose a recursive inference algorithm based on Sequential Monte Carlo techniques (SMC, [30, 12]) that ingests the data one at a time or in larger batches and independently of the order of presentation. The sequential incorporation of data points introduces a tempering effect in the sense described in the previous paragraph. As a result, the global problem of identifying a potentially multi-modal posterior is decomposed to a series of easier, tractable problems. More importantly, the inferences made can be readily updated if more data becomes available. As with MCMC, in SMC samplers the target distribution(s) need only be known up to a constant. The basis of the approximation is a set of random samples (commonly referred to as particles), which are propagated using a combination of importance sampling, resampling and MCMC-based rejuvenation mechanisms ([10, 9]). Each of these particles is associated with an importance weight which is proportional to the
posterior value of the respective particle. These weights are updated sequentially along with the particle locations. Hence if \( \{ \theta_n^{(i)}, w_n^{(i)} \}_{i=1}^N \) represent \( N \) such particles and associated weights for distribution \( \pi_n(\theta) \) then:

\[
\pi_n(\theta) \approx \sum_{i=1}^{N} W_n^{(i)} \delta_{\theta_n^{(i)}}(\theta)
\]

where \( W_n^{(i)} = w_n^{(i)}/\sum_{i=1}^{N} w_n^{(i)} \) are the normalized weights and \( \delta_{\theta_n^{(i)}}(.) \) is the Dirac function centered at \( \theta_n^{(i)} \). Furthermore, for any function \( h(\theta) \) which is \( \pi_n \)-integrable ([8, 7]):

\[
\sum_{i=1}^{N} W_n^{(i)} h(\theta_n^{(i)}) \to \int h(\theta) \pi_n(\theta) \, d\theta \quad \text{almost surely}
\]

In order to facilitate the transition between two successive posteriors \( \pi_n \) and \( \pi_{n+1} \) (particularly for small \( n \)), we can introduce a series of bridging distributions, based on a modified annealing scheme. In particular, if \( \pi_0 \) is the prior \( p(\theta) \) (Equation (2.17)) we define a family of artificial, auxiliary distributions \( \pi_{n,\gamma}(\theta) \) as follows:

\[
\pi_{n,\gamma}(\theta) \propto L_{n,\gamma}(\theta) \, p(\theta)
\]

based on the modified likelihood:

\[
L_{n,\gamma}(\theta) = \frac{\Gamma(a + (n + \gamma)/2)}{(b + \frac{1}{2} \sum_{i=1}^{n}(y_i - f(x_i; \theta))^2 + \gamma(y_{n+1} - f(x_{n+1}; \theta))^2)^{a+(n+\gamma)/2}} , \quad \gamma \in [0, 1]
\]

where \( \gamma \) plays the role of reciprocal temperature. Trivially for \( \gamma = 0 \) we recover \( \pi_n \) and for \( \gamma = 1, \pi_{n+1} \). The role of these auxiliary distributions is to bridge the gap between \( \pi_n \) and \( \pi_{n+1} \) and provide a smooth transition path where importance sampling can be efficiently applied. In this process, inferences based on \( n \) data points are transferred and updated to conform with the additional \( (n+1)^{th} \) datum. Starting with a particulate approximation for \( \pi_0(\theta) = p(\theta) \) (which trivially involves drawing samples from the prior with weights \( w_0^{(i)} = 1 \)), the goal is to gradually update the importance weights and particle locations in order to approximate the target posteriors \( \pi_n \).

We propose an adaptive SMC algorithm, that extends existing versions ([9, 10]) in that it automatically determines the number of intermediate bridging distributions needed. In this process we are guided by the Effective Sample Size \( ESS = 1/\sum_{i=1}^{N}(W_{s+1}^{(i)})^2 \) which provides a measure of degeneracy in the population of particles. Let \( s \) denote the number of intermediate bridging distributions between \( \pi_n \) and \( \pi_{n+1} \) and \( \gamma_s \) the associated reciprocal temperature. If \( ESS_s \) is the \( ESS \) of the population after the step \( s \), then in the most favorable scenario that the next bridging distribution \( \pi_{n,\gamma_{s+1}} \) is very similar to \( \pi_{n,\gamma_s} \), then \( ESS_{s+1} \) should not be that much different from \( ESS_s \). On the other hand if that difference is pronounced then \( ESS_{s+1} \) could drop dramatically. Hence in determining, the next auxiliary distribution, we define an acceptable reduction in the \( ESS \), i.e. \( ESS_{s+1} \geq \zeta \, ESS_s \) (where \( \zeta < 1 \)) and prescribe \( \gamma_{s+1} \) (Equation (2.25)) accordingly. The proposed Adaptive SMC algorithm is summarized in Table 2.1.

The role of the \( \text{Reweighing} \) step is to correct for the discrepancy between the two successive distributions in exactly the same manner that importance sampling
Adaptive SMC algorithm:

1. Initialize population \( \{ \theta_{0,0}^{(i)}, w_{0,0}^{(i)} \}_{i=1}^{N} \) where \( \theta_{0,0}^{(i)} \) are i.i.d draws from the prior \( \pi_{0} \) and \( w_{0,0}^{(i)} = 1 \) (\( \text{ESS}_0 = 0 \)). Set \( l = 0 \) and \( s = 0 \) and \( \gamma_0 = 0 \).
2. For \( l < n \):
   a) Set \( s = s + 1 \).
   b) Reweight: If \( w_{l,s}^{(i)}(\gamma_s) = w_{l,s-1}^{(i)} - \frac{\pi_{l,\gamma_s}(\theta_{l,s-1})}{\pi_{l,\gamma_{s-1}}(\theta_{l,s-1}^{(i)})} \) are the updated weights as a function of \( \gamma_s \) then determine \( \gamma_s \in (\gamma_{s-1}, 1] \) so that the associated \( \text{ESS}_s = \zeta \text{ESS}_{s-1} \) (the value \( \zeta = 0.95 \) was used in all the examples). Calculate \( w_{l,s}^{(i)} \) for this \( \gamma_s \).
   c) Resample: If \( \text{ESS}_s \leq \text{ESS}_{\text{min}} \) then resample (the value \( \text{ESS}_{\text{min}} = N/2 \) was used in all the examples).
   d) Rejuvenate: Use an MCMC kernel \( P_{l,s}(\cdot,\cdot) \) that leaves \( \pi_{l,\gamma_s} \) invariant to perturb each particle \( \theta_{l,s-1}^{(i)} \rightarrow \theta_{l,s}^{(i)} \).
   e) The current population \( \{ \theta_{l,s}^{(i)}, w_{l,s}^{(i)} \}_{i=1}^{N} \) provides a particulate approximation of \( \pi_{l,\gamma_s} \) in the sense of Equations (2.23), (2.24).
   f) If \( \gamma_s = 1 \) set \( l = l + 1 \), \( \theta_{l,0}^{(i)} = \theta_{l-1,s}^{(i)} \), \( w_{l,0}^{(i)} = w_{l-1,0}^{(i)} \), \( s = 0 \) and \( \gamma_0 = 0 \).

is employed. The Resampling step aims at reducing the variance of the particulate approximation by eliminating particles with small weights and multiplying the ones with larger weights. The metric that we use in carrying out this task is the Effective Sample Size (ESS) defined earlier. If this degeneracy exceeds a specified threshold, resampling is performed. As it has been pointed out in several studies (\cite{11}), frequent resampling can deplete the population of its informational content and result in particulate approximations that consist of even a single particle. Throughout this work \( \text{ESS}_{\text{min}} = N/2 \) was used. Although other options are available, multinomial resampling is most often applied and was found sufficient in the problems examined.

A critical component involves the perturbation of the population of samples by a standard MCMC kernel in the Rejuvenation step as this determines how fast the transition takes place. Although there is freedom in selecting the transition kernel \( P_{l,s}(\cdot,\cdot) \) (the only requirement is that it is \( \pi_{l,\gamma_s} \)-invariant), there is a distinguishing feature that will be elaborated in the next sub-section (see 2.3.1). The target posteriors \( \pi_n \) (as well as the intermediate bridging distributions in Equation (2.25)) live in spaces of varying dimensions as previously discussed. Hence an exploration of the state space must involve trans-dimensional proposals. Pairs of such moves can be defined in the context of Reversible-Jump MCMC (RJMCMC, \cite{19}) such as adding/deleting a kernel in the expansion of Equation (2.8), or splitting/merging kernels (see 2.3.1).

It should be noted that the framework proposed is directly parallelizable, as the evolution (reweighting, rejuvenation) of each particle is independent of the rest. The particulate approximations obtained at each step, provide a concise summary of the posterior distribution based on the respective forward solver. This can be readily updated in the manner explained above, if more data become available, i.e. more runs of the approximate and high-fidelity solvers are invoked.
An advantageous feature of the Bayesian framework is that the confidence in the estimates made can be readily quantified by establishing posterior (or credible) intervals from the particulate approximations (Equation (2.23)). It is these credible intervals (or in general measures of the variability in the estimates such as the posterior variance) that can guide adaptive acquisition of data. Since we want to minimize calls to the accurate solver $y$, we can utilize these inferences in order to perform runs in regions that will be most informative of the sought output and therefore make near-optimal use of the computational resources available. This will be discussed in more detail in the numerical examples.

2.3.1. Trans-dimensional MCMC. As mentioned earlier, a critical component in the SMC framework proposed is the MCMC-based rejuvenation step of the particle locations $\theta$. It should be noted that the kernel $P_{l,s}(\cdot,\cdot)$ in the rejuvenation step (Step 2.d) of the SMC algorithm in Table 2.1) does not need to be known explicitly as it does not enter in any of the pertinent equations. It is sufficient that it is $\pi_{l,\gamma}$-invariant which is the target density. For the efficient exploration of the state space, we employ a mixture of moves which involve fixed dimension proposals (i.e., proposals for which the cardinality of the representation $k$ is unchanged) as well as moves which alter the dimension $k$ of the vector of parameters $\theta$. We consider a total of $M = 7$ such moves, each selected with a certain probability. The fixed-dimension moves involve updates of either $a_j$, $\tau_j$ or $\nu_j$ of a randomly selected kernel $j$ in the expansion of Equation (2.8). In addition four trans-dimensional proposals are used which due to space limitations are just enumerated here. The interested reader can find more details in [26].

Birth-Death: A new kernel term is added or a kernel selected at random in the expansion of Equation (2.8) is removed.

Split-Merge A kernel term in the expansion of Equation (2.8) is split into two (with appropriate amplitudes, scales and locations) or two existing kernels (located sufficiently close) are merged into one (see [26]).

2.4. Prediction - Calculating statistics of high-fidelity solver output. We return to the original problem of estimating statistics of the output $y$ of the high-fidelity solver using the relationship with the outputs $x$ of the approximate solvers. The aforementioned Bayesian model is able to not only provide estimates but also quantify the level of confidence one can assign to the predicted outcome. Let $(\theta, \sigma)$ denote a pair of values for the model parameters in Equation (2.6). For these parameter values, the conditional density $p(y \mid x)$ can be obtained based on the regression model adopted and Equation (2.7):

$$p(y \mid x) \approx p(y \mid x, \theta, \sigma) = \frac{1}{\sqrt{2\pi} \sigma} \exp\left\{ -\frac{1}{2\sigma^2} (y - f(x; \theta))^2 \right\}$$

(2.27)

which upon substitution in Equation (2.4) yields:

$$Pr[y \in A; \theta, \sigma] = \int q_A(x; \theta, \sigma) \pi_x(x) \, dx$$

(2.28)

where $q_A(x; \theta, \sigma) = \int 1_A(y) \, p(y \mid x, \theta, \sigma) \, dy$. The latter expresses the probability that the accurate response $y \in A$ for fixed approximate response $x$ (and model parameters). Consider for example the case that we are interested in calculating a probability of exceeding a threshold $y_0 \in \mathbb{R}$, i.e. $A = (y_0, +\infty)$. Then:

$$q_A(x; \theta, \sigma) = \Phi \left( \frac{f(x; \theta) - y_0}{\sigma} \right)$$

(2.29)
where $\Phi(z) = \int_{-\infty}^{z} \frac{1}{\sqrt{2\pi}} e^{-\frac{w^2}{2}} dw$ is the standard normal CDF.

Given a number of training data $(x_{1:n}, y_{1:n})$, the plausibility of various parameter values is quantified by the posterior $\pi_n(\theta) = p(\theta \mid (x_{1:n}, y_{1:n}))$. Hence, one can obtain point estimates of $q_A$ based for example on the maximum a posteriori values (MAP) $\theta_{MAP} = \arg\max \pi_n(\theta)$ or the posterior mean $E_n[\theta] = \int \theta \pi_n(\theta) \, d\theta$. More importantly, due to its dependence of $\theta$ (and $\sigma$), $q_A$ is also random and its distribution can be determined from the posterior distribution of $\theta$ and $\sigma$. This distribution therefore indirectly depends on the training data upon which posterior inferences were based. In particular, one can estimate the posterior mean of $q_A$ for the case of Equation (2.29) as follows:

$$
\hat{q}_A(x) = E_n[q_A] = \int \Phi \left( \frac{f(x; \theta) - y_0}{\sigma} \right) \pi_n(\theta, \sigma^{-2}) \, d\theta \, d\sigma^{-2} \\
= \int \Phi \left( \frac{f(x; \theta) - y_0}{\sigma} \right) \pi_n(\theta) \pi_n(\sigma^{-2} \mid \theta) \, d\theta \, d\sigma^{-2} \\
\approx \sum_{i=1}^{N} W_n(i) \Phi \left( \frac{f(x; \theta_n(i) - y_0)}{\sigma_n(i)} \right) \tag{2.30}
$$

where the particulate approximation $\{\theta_n, W_n(i)\}_{i=1}^N$ of $\pi_n$ obtained through the proposed SMC scheme was used and $\{\sigma_n(i)\}_{i=1}^N$ are corresponding draws from the conditional posterior in Equation (2.22). Equation (2.30) expresses how, on average, based on the available data $(x_{1:n}, y_{1:n})$ the probability of interest depends on the approximate solver values $x$. This can provide an estimate of the probability of interest by substitution in Equation (2.28).

Furthermore, the weighted samples $q_A^{(i)}(x) = \Phi \left( \frac{f(x; \theta_n(i) - y_0)}{\sigma_n(i)} \right)$ provide a particulate approximation of the distribution of $q_A(x)$, $\forall x$. p-Quantiles $q_A,p(x)$ can be readily estimated as:

$$
Pr[q_A(x) \leq q_A,p(x)] \approx \sum_{i=1}^{N} W_n(i) H \left( q_A^{(i)}(x) - q_A,p(x) \right) = p \tag{2.31}
$$

where $H(\cdot)$ is the Heaviside function. These can readily yield confidence bounds for the probability of interest when substituted in Equation (2.28). More importantly perhaps, these bounds can serve as the basis for active learning i.e. determining where more training samples need to be generated in order to refine the estimates produced. Consider for example the $p = 1\%$ and $p = 99\%$ quantiles $q_A,0.01(x)$ and $q_A,0.99(x)$. Then based on Equation (2.28) an ordering of $x$ can be constructed based on:

$$
(q_A,0.99(x) - q_A,0.01(x)) \pi_x(x) \tag{2.32}
$$

Thus $x$ (or regions in the $x$-space) for which the aforementioned value is large contribute more in the uncertainty about the probability of interest $Pr[y \in A]$ and therefore could serve as the best candidates for generating an additional training pair $(x_{n+1}, y_{n+1})$. This is particularly important in the cases considered where each run for the evaluation of the high-fidelity response $y$ can be extremely expensive and therefore optimal use of the computational resources is crucial. These additional training
samples can be readily incorporated based on the SMC scheme adopted and updates of the particulate approximation that reflect the new data can be produced. These in turn can lead to updates in the estimates made as well as the confidence bounds. Naturally other measures of variability of \( q_A(x) \), such as the variance (or coefficient of variation) can be used in place of \((q_{A,0.99}(x) - q_{A,0.01}(x))\) in Equation (2.32). The variance for each \( x \) can be estimated as follows:

\[
Var_n[q_A] \approx \sum_{i=1}^{N} W_n^{(i)} \left( q_A^{(i)}(x) - \hat{q}_A(x) \right)^2 \tag{2.33}
\]

For estimates of expectations of functions \( h(.) \) of the accurate output \( y \) as in Equation (2.5), the same results apply if in place of \( q_A(x) \) we use:

\[
q_h(x) = \int h(y)p(y \mid x) \, dy \tag{2.34}
\]

3. Numerical results. In the examples presented, the following values for the hyperparameters of the prior model were used (Equation (2.17)):

- \( k_{max} = 100 \) and \( s = 1.0 \) (Equation (2.12))
- \( a_\tau = 2.0 \) (Equation (2.13)) and \( a_\mu = 0.01 \) (Equation (2.14))
- \( a_\theta = 1.0 \) and \( b_\theta = 1.0 \) (Equation (2.16))
- \( a = 2. \) and \( b = 1. \times 10^{-6} \) (Equation (2.18))

Furthermore, \( N = 1,000 \) particles were employed in the adaptive SMC scheme described in section 2.3. As in most systems of practical interest, the computational cost is dominated by the number of calls to the forward solver, we report results on computational effort in terms of the number of runs of the high-fidelity, expensive solver.

3.1. Example 1. The first example involves a problem from fracture mechanics where it is known that small scale stochastic fluctuations can have a significant impact in the macroscale response. We consider cohesive interface of unit length that is pulled apart in Mode I fracture and is modeled with cohesive zone elements (Figure 3.1). These are line (or surface in 3D) elements which are located at the interface and govern the separation process in accordance with a cohesive law. The concept of cohesive laws was pioneered by Dugdale ([13]) and Barenblatt ([3]) in order to model fracture processes and has been successfully used in a Finite Element setting by several researchers ([43, 4, 34]). According to these models, fracture is initiated when the interface traction exceeds a threshold \( T_c \) and progresses gradually as the separation takes place across an extended crack tip or cohesive zone and is resisted by cohesive tractions. We assume herein a simple constitutive law relating interface traction-separation as seen in Figure 3.1. Under monotonic loading the normal interface traction decays as \( T = T_c \left( 1 - \frac{\delta}{\delta_c} \right) \) for \( \delta \leq \delta_c \) and \( T = 0 \) for \( \delta > \delta_c \). The fracture energy \( G_c \) is given by \( G_c = T_c \delta_c / 2 \). The constitutive rate equations are:

\[
\dot{T} = \begin{cases} 
-\frac{T_c}{\delta_c} & \text{if } \delta > 0 \\
\frac{T}{\delta} & \text{if } \delta < 0 \\
0 & \text{if } \delta > \delta_c
\end{cases}
\tag{3.1}
\]

At the microstructural level, the cohesive properties exhibit random variability. We adopt the following simple random field descriptions for the model parameters:

\[
T_c(z) = T_0 + \Delta T_0 \, U_1(z)
\]

\[
G_c(z) = G_0 + \Delta G_0 \left( \rho U_1(z) + U_2(z) \right) \quad z \in [0,1]
\tag{3.2}
\]
where:

\[ U_i(z) = 2\Phi(h_i(z)) - 1, \quad i = 1, 2 \quad (3.3) \]

and \( h_1(z) \) is a zero-mean, unit variance Gaussian process with autocorrelation \( R_h(\Delta z) = E[h(z)h(z + \Delta z)] = \exp\{-\frac{|\Delta z|}{\rho} \} \) (\( \Phi \) is the standard normal CDF). The parameter \( z_0 \) controls the length scale of heterogeneity and it was taken equal to 0.1. The field \( h_2(z) \) was assumed to represent a Discretized white noise process. The parameter \( \rho \) controls the autocorrelation between the two properties and was taken equal to 0.9 which implies that areas with high \( T \) are more likely to have high \( G_c \) as well. Also, the values \( T_0 = 1.0, \Delta T_0 = 0.5, G_c = 10^{-3} \) and \( \Delta G_0 = 0.5 \times 10^{-3} \) were used.

The reference, high-fidelity solver was a detailed finite element model consisting of 1,000 cohesive elements of equal length with properties assigned based on the values of \( T_c(z) \) and \( G_c(z) \) at their midpoint. The mesh size is much smaller than the length scale of variability of the cohesive properties as determined by the correlation length \( z_0 \) defined above. The output of interest \( y \) was the fracture energy released when a uniform separation \( \delta = 0.5 \times 10^{-3} \) was applied at the interface. The quasi-static, nonlinear calculation of the output of interest was carried out by applying separation increments of \( 0.5 \times 10^{-6} \) in order to capture accurately the traction-separation history (i.e. a total of 1,000 iterations).

We considered a single approximate solver with a much coarser mesh consisting of only 10 cohesive elements of equal length, i.e. each macro-element takes the place of 100 micro-elements of the accurate model. The assigned cohesive strength \( T_c \) in each macro-element was set equal to the minimum of the cohesive strengths of the 100 corresponding micro-elements, and the fracture energy \( G_c \) equal to the average of the fracture energies of the the 100 corresponding micro-elements. In addition a displacement increment of \( 0.5 \times 10^{-5} \) (in contrast to the \( 0.5 \times 10^{-6} \) for the high-fidelity solver) was used in order to carry out the quasi-static, nonlinear integration of the equations of equilibrium and the constitutive model (Equation (3.1)). As a result of these crude simplifications the approximate solver was 1,069 faster than the accurate. Figure 3.2(a) compares the approximate and accurate model outputs where significant discrepancies can be observed (e.g. when \( x = 4 \times 10^{-4}, y \approx 5 \times 10^{-4} \), i.e. a 25% difference). It is also observed that the mapping from \( x \) to \( y \) is one-to-many, as the coarse model crudely smears some of the fine details that affect the response.

In order to assess the performance of the method proposed we considered the event \( y > y_0 = 5.601 \times 10^{-4} \) which corresponds to a probability \( 10^{-3} \). This was found using an advanced Monte Carlo simulation procedure as discussed in the introduction ([2, 25]). The computational effort amounted to 1,500 calls to the accurate, expensive model and the coefficient of variation (c.o.v) of the estimate was 0.23 (based on an asymptotic, and rather optimistic, bound in [2]). This roughly implies that with probability 0.95, the actual probability of the event of interest is in the interval \([0.55 \times 10^{-3}, 1.5 \times 10^{-3}]\). In order to achieve the same c.o.v with standard Monte Carlo, 18,500 calls to the high-fidelity model would have been needed.

The density \( \pi_x \) of \( x = x_1 \) was estimated using the same advanced Monte Carlo scheme that required 5,000 calls to the approximate solver. The corresponding CDF is depicted in Figure 3.2(b) for probabilities as low as \( 10^{-5} \). Note that due to the reduced computational effort associated with calls to the approximate solver, the computational time for this task amounted to (approximately) 5 runs of the high-fidelity model.

The crucial task, that of estimating \( p(y \mid x_1) \) involves the nonparametric Bayesian
Cohesive interface - Cohesive Law: $T_c$ denotes ultimate interfacial tension (when the stress reaches $T_c$ the cohesive element is activated), $\delta_c$ denotes the ultimate separation interface (when the separation reaches $\delta_c$ the interface tension becomes zero) and $G_c$ denotes the fracture energy which is equal to the area under the tension-separation curve.

Fig. 3.2. Accurate (expensive) $y$ vs. Approximate (inexpensive) $x$ model output (a) and Cumulative distribution function for the approximate solver output $x$ (b).

regression model discussed previously. Figure 3.3 depicts posterior statistics of the regression model for various training sample sizes and Figure 3.4 the posterior mean and posterior quantiles of $q_A(x)$ based on Equations (2.30) and (2.31) for various sample sizes. Table 3.1 summarizes the estimates based on the posterior mean $\hat{q}_A(x)$ and confidence bounds established with $q_{A,0.01}(x)$ and $q_{A,0.99}(x)$. It is noted that even with a small number of calls to the high-fidelity, expensive solver, the estimates obtained are reasonably good and most importantly the lower and upper confidence bounds always include the reference value. This is particularly important for engineering purposes since if the analyst decides that these confidence bounds are not satisfactory, additional runs of the high-fidelity model can be performed in order to refine these estimates. As the number of training samples increases the posterior mean approaches the true value and the credible intervals become more concentrated. A complete view of the the cdf of the accurate model output $y$ is depicted in Figure 3.5 based on 150 training samples.

3.2. Example 2. We consider a problem in nonlinear solid mechanics that illustrates the capabilities of the proposed methodology and the the significant im-
Fig. 3.3. Posterior mean of $f(\mathbf{x}; \theta)$ and posterior density of $\sigma$ (Equation (2.6)) for various training sample sizes

Table 3.1
Estimates of $Pr[y > y_0 = 5.601 \times 10^{-4}]$ and computational effort (the latter is measured in equivalent number of calls to the high-fidelity, expensive model)

<table>
<thead>
<tr>
<th>Number of samples</th>
<th>Posterior mean</th>
<th>Posterior quantile 1%</th>
<th>Posterior quantile 99%</th>
<th>Computational Effort</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>$6.36 \times 10^{-3}$</td>
<td>$5.91 \times 10^{-4}$</td>
<td>$7.17 \times 10^{-2}$</td>
<td>15</td>
</tr>
<tr>
<td>50</td>
<td>$1.75 \times 10^{-3}$</td>
<td>$7.39 \times 10^{-4}$</td>
<td>$3.55 \times 10^{-3}$</td>
<td>55</td>
</tr>
<tr>
<td>150</td>
<td>$1.01 \times 10^{-3}$</td>
<td>$7.07 \times 10^{-4}$</td>
<td>$1.42 \times 10^{-3}$</td>
<td>155</td>
</tr>
</tbody>
</table>
provements in computational efficiency even when very crude approximate solvers are selected. A random two-phase medium consisting of two elastic-perfectly-plastic materials occupies the unit square in 2D and is subjected to plane stress loading conditions. It was assumed that the matrix phase (white in Figure 3.6) had a yield stress $\sigma_{\text{matrix}}^{\text{yield}} = 0.1$ and the inclusion phase (black in Figure 3.6) a yield stress $\sigma_{\text{inclusion}}^{\text{yield}} = 1.0$. The same elastic properties were assumed for both phases (elastic modulus $E = 1.0$ and Poisson’s ratio $\nu = 0.3$) and the von-Mises yield criterion was used. The stochasticity in the problem is introduced by the distribution of the black disks of diameter $d = 0.0859375$ whose centers are assumed to follow a Poisson point process on the unit square. The intensity of the point process is selected so that the volume fraction of the inclusion phase is 65%. This was intentionally chosen to be close to the percolation threshold of 68% ([44]) in order to have realizations where the inclusion phase was connected and disconnected. In the former case, the load-bearing capacity (in the horizontal direction) of the specimen is high as the strong, inclusion phase forms a network that carries the load, whereas in the disconnected case, the load-bearing capacity is lower and determined by the yield stress of the matrix phase. Hence the stochastic geometry completely determines the mechanical behavior of the system. The vector of uncertain parameters $\xi$ (Equation (2.2)) consists of the number of inclusion disks and the coordinates of their centers. As the former is a random variable (following a Poisson distribution) the corresponding $\pi_\xi(\xi)$ has
support in spaces of varying dimension and non-zero probability for arbitrarily large $d$ where $\xi \in \mathbb{R}^d$. It should be noted however that on average there are 181 disks and $\text{dim}(\xi) = 1 + 2 \times 181 = 363$. Usual dimension reduction techniques based on second order properties would be extremely misleading in this case as higher order statistics of the random medium (relating to connectivity) dominate mechanical response.

![Figure 3.6. Example 2](image)

The high-fidelity, expensive model corresponds to a Finite Element solver with $128 \times 128$ elements i.e. 11 elements per inclusion diameter and $\approx 33,000$ dof, of the governing equations of equilibrium:

$$\nabla \cdot \hat{\sigma} = 0 \quad (3.4)$$

where $\hat{\sigma}$ is the Cauchy stress tensor and $\dot{\sigma} = c : (\dot{\varepsilon} - \dot{\varepsilon}^p)$ is the nonlinear constitutive rate equation with $\varepsilon$ and $\varepsilon^p$ being the total and plastic strain tensors respectively.

The yield stress was assumed constant within each element and equal to the yield stress of the the phase occupying the majority of its area. The response of interest $y$ was the ultimate strength of the specimen in the horizontal direction and the average computational time was $\approx 700$ sec on a single CPU.

A single approximate model was used (i.e. $M = 1$) which corresponds to a Finite Element solver on a uniform $8 \times 8$ mesh and $128$ dof. Constant yield stress was assigned to each of the 64 elements based on the log-average of the yield stress within each element (Figure 3.7(b)). Hence if $D_e$ is the subdomain occupied by element $e$, its yield stress $\sigma_{\text{yield},e} = \exp\left\{\frac{1}{|D_e|} \int_{D_e} \log (\sigma_{\text{yield}}(s)) \, ds\right\}$. The computational time for calculating the ultimate strength $x = x_1$ was 0.15 sec, i.e. $\approx 4,700$ times faster than the high-fidelity model. It is obvious that such a solver introduces a significant error as it does not sufficiently resolve the governing PDEs and smears out the connectivity details that determine the ultimate strength of the specimen. Furthermore, the log-average rule used to determine the yield stress of the elements does not represent a consistent upscaling scheme of the material model. This discrepancy can be seen in Figure 3.8(a) which depicts 100 pairs of approximate $x_1$ vs. accurate response $y$. It is also observed that the mapping from $x_1$ to $y$ is one-to-many as the approximate model blurs some of the important microstructural details.

In order to compare the performance of the method we considered the event $y > y_0 = 0.521$ which corresponds to a probability $10^{-3}$. This was found using
an advanced simulation procedure based on Sequential Monte Carlo as discussed in detail in ([33, 17, 25]). The computational effort amounted to 1,500 calls to the high-fidelity solver and the coefficient of variation (c.o.v) of the estimate was 0.23 (based on an asymptotic, and rather optimistic, bound in [2]). This roughly implies that with probability 0.95, the actual probability of the event of interest is in the interval $[0.55 \times 10^{-3}, 1.5 \times 10^{-3}]$. It should be noted that to achieve the same c.o.v with standard Monte Carlo, 18,500 calls to the high-fidelity solver would have been needed.

The density $\pi_x$ of $x = x_1$ was estimated using the same advanced Monte Carlo scheme that required 5,000 calls to the approximate solver. The corresponding cdf is depicted in Figure 3.8(b) for probabilities as low as $10^{-5}$. Note that due to the reduced computational effort associated with calls to the approximate solver, the effective computational time for this task amounted to (approximately) 1 call to the high-fidelity solver.
As with the previous example, Figure 3.9 depicts posterior statistics of this model for various training sample sizes. Figure 3.10 depicts the posterior mean and posterior quantiles of $q_A(x)$ based on Equations (2.30) and (2.31) for various sample sizes. Table 3.2 summarizes the estimates based on the posterior mean $\hat{q}_A(x)$ and confidence bounds established with $q_{A,0.01}(x)$ and $q_{A,0.99}(x)$. It is noted that good estimates of the actual output statistic can be obtained at a fraction of the computational cost. Furthermore, good confidence bounds are readily obtained for all sample sizes. The same good quality in the results is also observed in the estimates of the whole cdf of the accurate model output which is depicted in Figure 3.11 for 100 training samples.

<table>
<thead>
<tr>
<th>Number of samples</th>
<th>Posterior mean</th>
<th>Posterior quantile 1%</th>
<th>Posterior quantile 99%</th>
<th>Computational Effort</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>$1.47 \times 10^{-2}$</td>
<td>$2.33 \times 10^{-4}$</td>
<td>$3.80 \times 10^{-1}$</td>
<td>11</td>
</tr>
<tr>
<td>20</td>
<td>$6.24 \times 10^{-3}$</td>
<td>$3.56 \times 10^{-3}$</td>
<td>$1.90 \times 10^{-2}$</td>
<td>21</td>
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</tr>
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<td>$4.25 \times 10^{-4}$</td>
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<td>$4.75 \times 10^{-4}$</td>
<td>$2.14 \times 10^{-3}$</td>
<td>101</td>
</tr>
</tbody>
</table>

We examined the same problem using $M = 2$ approximate solvers. In addition to the approximate solver discussed earlier, we considered a second FE model with the same mesh of 64 elements (Figure 3.7(b)). The yield stress $\sigma_{\text{yield},e}$ for each element $e$ was now determined by averaging i.e. $\sigma_{\text{yield},e} = \frac{1}{|D_e|} \int_{D_e} \sigma_{\text{yield}}(s) \, ds$. This is again an inconsistent rule with the accurate (reference) constitutive model which yields approximate solutions. Furthermore, due to the concavity of the log-function and Jensen’s inequality, the assigned yield stresses $\sigma_{\text{yield},e}$ were smaller in the first model and as a result the predicted approximate outputs $x_1 < x_2$.

Figure 3.12 depicts 50 triplets $(x_1, x_2, y)$ comparing the accurate model output with the approximate solutions provided by the two reduced models. It is expected that the addition of the second model will yield more information about $y$ that can be readily taken into account by the Bayesian framework presented. The joint pdf $\pi_x(x_1, x_2)$ was estimated using 5,000 calls to each solver (i.e. total 10,000) which due to their reduced computational cost amounted to the equivalent of $\approx 2$ runs of the high-fidelity solver.

Figure 3.13 depicts posterior statistics of the regression model for 50 training samples. Finally Figure 3.14 illustrates the complete cdf of the accurate model output $y$ based on the same number of training samples. The computational effort for obtaining this result is equivalent to 52 calls to the high-fidelity solver (i.e. 2 for estimating $\pi_x$ and 50 for obtaining the training data). The addition of the second predictor $x_2$ offers a significant improvement w.r.t. Figure 3.11 not only in terms of accuracy but also in terms computational efficiency since the latter result required effectively 101 calls to the high-fidelity solver.

4. Conclusions & Discussion. The majority of systems of physical and engineering interest are characterized by a large number of uncertainties. These are non-Gaussian distributed and quite frequently their higher-order properties play a decisive role in the statistics of the response/output. While Monte Carlo techniques provide the only general method for uncertainty quantification in such systems and
Despite the significant progress of recent years, they might still imply significant computational effort. The present paper introduced a Bayesian framework where outputs from approximate, inexpensive solvers can be rigorously utilized in order to accelerate the solution process. We made use of a flexible, non-parametric Bayesian model and a general SMC-based inference engine that is able to establish a quantitative link between inaccurate and accurate outputs.

The advantages offered by the proposed strategy hinge on the computational cost associated with the inaccurate solvers and the information they furnish about the output of the expensive computational model. In general these requirements are in competition as increased accuracy generally implies an increased computational cost. As we argued earlier and demonstrated in the examples, what actually matters is not the difference between accurate and inaccurate outputs in absolute terms but the correlation (or more generally dependence) that is exploited by our model. It is generally much easier to identify inexpensive models that are somewhat correlated with the accurate output in a wide class of problems. The proposed framework is flexible enough to allow for the simultaneous and rigorous incorporation of a number of such approximate models to account for peculiarities of each problem and varying insight of the analyst. In deriving such approximate models, the following guidelines can be useful although not restrictive and can be used separately or in combinations:

- **Discretization in space.** This is one of the most common scenarios encountered as a lot of systems imply the solution of PDEs and therefore require some sort of spatial discretization. Considering coarser meshes (or a hierarchy of those) and/or reducing the order of the basis functions is an obvious way of obtaining less-accurate models while reducing the computational cost.

- **Discretization in time.** Considering larger time steps is an apparent way of obtaining less accurate models. In several situations higher order integration schemes are used to ensure greater accuracy (e.g. Runge-Kutta methods). Utilizing lower-order schemes will reduce the computational effort and in a lot of cases provide some information about the higher-order, expensive solver output. Reduced-order models based on a truncated modal-decomposition can also be employed at least in linear systems and linear projections (e.g. POD schemes) in more general cases.

- **Nonlinearities.** Nonlinear constitutive laws are frequently employed to achieve greater fidelity in the representation of various physical phenomena. These can be (partially) approximated by cheaper, linear models at a fraction of the computational cost. Furthermore, in most cases the solution of the governing equations involves Newton-Raphson iterations of the discretized system. The number of such iterations can be reduced and quasi-Newton-Raphson schemes can be employed (which calculate the Jacobian less frequently).

- **Iterative solvers.** For large systems, iterative (Gauss-Seidel, Krylov etc) solvers are usually preferred due to favorable scaling with the dimensionality of the problem. Obviously the computational cost increases with the number of iterations which in turn is determined by a tolerance parameter. Allowing for greater tolerances will of course decrease the accuracy of the solution but also decrease the computational effort.

In the presence of a multitude of options, it is important to have quantitative estimates of the information various approximate models provide. In particular it is important to assess whether the estimates produced using a particular set of approximate outputs is sufficient or further refinement is needed. To that end, the proposed
methodology is capable of providing rigorous credible intervals. When the approximate outputs are not informative (i.e., the correlation with the accurate output is small - see discussion at the end of section 2), these credible intervals will be large indicating high variability in the model prediction. If this variability is not acceptable, incorporating outputs from additional approximate solvers would be necessary. It should be noted that the proposed technique does not provide a priori any indication on how to construct approximate models as it is based on “learning from data” obtained from computer simulations (or experiments). It can however mine the data to extract quantitative information about the accurate (expensive) model output. Deciding on the relative importance of various solvers is something that can definitely be handled in the Bayesian framework proposed ([20]) and is the subject of current investigations.

While this capability was not utilized in the examples presented, the credible intervals produced, can also assist in minimizing the number of calls to the high-fidelity, expensive solver by performing runs in selected regions that will be most informative. Furthermore it offers the capability of utilizing multiple approximate solvers and it opens the door for designing or optimizing systems in the presence of uncertainties based on establishing functions of the output statistics with respect to the design variables.

Acknowledgments. The author would like to acknowledge the two anonymous reviewers whose valuable comments and suggestions improved the overall quality of the paper.

REFERENCES

Fig. 3.9. Posterior mean of $f(x; \theta)$ and posterior density $\sigma$ for various training sample sizes
Fig. 3.10. Posterior mean and quantiles for $Pr[y \geq 0.521 | x]$ based on various sample sizes

Fig. 3.11. Posterior for $Pr[y \geq y_0] \forall y_0$ based on 100 training samples
Fig. 3.12. Accurate (expensive) $y$ vs. Approximate (inexpensive) $x = (x_1, x_2)$ model outputs

Fig. 3.13. Posterior mean of $f(x; \theta)$ and posterior density $\sigma$ for 50 training samples

Fig. 3.14. Posterior mean and quantiles for $Pr[y \geq y_0]$, $\forall y_0$ based on 50 training samples