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Design optimization of a scramjet under uncertainty using probabilistic learning on manifolds

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Abstract

We demonstrate, on a scramjet combustion problem, a constrained probabilistic learning approach that augments physics-based datasets with realizations that adhere to underlying constraints and scatter. The constraints are captured and delineated through diffusion maps, while the scatter is captured and sampled through a projected stochastic differential equation. The objective function and constraints of the optimization problem are then efficiently framed as non-parametric conditional expectations. Different spatial resolutions of a large-eddy simulation filter are used to explore the robustness of the model to the training dataset and to gain insight into the significance of spatial resolution on optimal design.

Keywords: Scramjet simulations, Machine learning, Uncertainty quantification, Optimization under uncertainty, Sampling on manifolds, Diffusion maps

1. Introduction

Model-based design optimization is a significant capability for systems whose behavior has not yet been catalogued but where functional dependence of system behavior on component behavior can be described through conservation or other such fundamental principles. These systems typically include new designs with no corresponding legacy knowledge and complex systems for which perturbations in any component has consequences on system-level performance. Predictive models for these systems are typically discretizations of mathematical formulations of the underlying fundamental principles resulting in computational models with very significant computational requirements. It is typical for such systems, given their novelty or complexity, to invoke reductions that could be either physical, mathematical, or algorithmic in nature. The impact of these reductions on the predicted performance of any given design can put into question the optimal character of solutions obtained through traditional design optimization procedures. One rational formalism for accounting for these assumptions throughout the design process is to explore the robustness of an optimal solution to perturbations in these assumptions. Probabilistic modeling provides an effective procedure for characterizing these assumptions and has typically been implemented through either parametric procedures where model parameters are described as random variables [1, 2, 3, 4, 5] or within a non-parametric framework where the model itself is described as a random operator [6]. Irrespective of how a non-deterministic problem is implemented, it requires the numerical exploration of a statistical ensemble, thus quickly exacerbating the computational burden of an already massive exercise. Clearly, new perspectives on characterizing and evaluating performance and optimal designs of complex systems are required if physics-based modeling is indeed a necessary ingredient of the prediction and optimization process.

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In this paper we tackle the above challenge in the context of a hypersonic combustion process. Large-eddy simulation (LES) modeling is used to discretize the Navier-Stokes equations, with several of the filtering parameters and operational parameters described as random variables. In the process, we rely on a new methodology developed by the authors that conjoins diffusion maps and projected Itô sampling to augment a small statistical ensemble (the training set) with a large number of realizations that are consistent, in a useful sense, with both underlying mechanisms and evidence. Diffusion maps [7] are used to extract nonlinear manifolds from the training set, while the Itô samplers [8] are used to sample on these manifolds from a specified target probability measure. We construe these manifolds as encoding non-parametric characterization of constraints on the data, including physics-based constraints.

The optimization problem is typically formulated in terms of several quantities of interest (QoIs) that are themselves functionals of the field variables constituting the solution of the LES problem. The map from input parameters to QoIs is a composition of two maps. First is the map represented by the numerical solver, which encodes conservation laws as described by the LES model. This is followed by a postprocessing map that evaluates the QoIs, which enter in the evaluation of objectives and constraints. The LES map is challenged with a computational burden that limits its ability to efficiently explore parameter space, a key requirement for design optimization. The postprocessing map faces its own challenges, namely the lack of apparent physics-based constraints on the QoIs, which forces the characterization of the QoIs in terms of the input parameters and the LES map. These challenges notwithstanding, it can be expected that these two maps do indeed constrain the QoIs, albeit in an unknown and intractable manner. By discovering these constraints, the task of input space exploration would be reduced to exploring a low-dimensional manifold. For that, we propose to use the probabilistic learning on manifolds [9, 4] that relies on diffusion maps (DMAPS), with the corresponding manifold characterized through a positive definite operator and its associated projections. Then, statistical sampling on this manifold is readily accomplished as described above, via a projected Itô equation whose invariant measure is a nonparametric statistical estimate of the measure of the training data, represented as a Gaussian mixture model. We compare optimization results obtained from three LES models representing different spatial resolutions, and explore the impact of the size of the training set on the resulting optimal designs.

The paper is organized as follows. In the next section, an overview is presented on the physics problem representing scramjet combustion. Following that, probabilistic learning on manifolds using diffusion maps and Itô sampling is described. The optimization problem under uncertainty is then illustrated and its solution analyzed. Some remarks are presented in a concluding section.

2. Overview of the hypersonic reactive flow model

Our physical application of interest stems from the HIFiRE (Hypersonic International Flight Research and Experimentation) program [10, 11], which has cultivated a mature experimental campaign with accessible data through its HIFiRE Flight 2 (HF2) project [12, 13]. The HF2 payload involves a cavity-based hydrocarbon-fueled dual-mode scramjet and was tested under flight conditions of Mach 6–8+. A ground test rig, designated the HIFiRE Direct Connect Rig (HDCR) (Figure 1(a)), was developed to duplicate the isolator/combustor layout of the flight test hardware, and to provide ground-based data for comparisons with flight measurements, verifying engine performance and operability, and designing fuel delivery schedule [14, 15]. While data from flight tests are not publicly released, HDCR ground test data are available [14, 16]. Therefore, we aim to simulate and assess behavior of reactive flows inside the HDCR, in order to facilitate future computational developments that can make use of these experimental datasets.

The computational domain for the HDCR is highlighted by red lines in Figure 1(b). The rig consists of a constant-area isolator (planar duct) attached to a combustion chamber. It includes primary injectors that are mounted upstream of flame stabilization cavities on both the top and bottom walls. Secondary injectors along both walls are positioned downstream of the cavities. Flow travels from left to right in the $x_s$-direction (streamwise), and the geometry is symmetric about the centerline in the $y_s$-direction. Numerical simulations take advantage of this symmetry by considering a domain that covers only the bottom half of this configuration. The fuel supplied through the injectors is a gaseous mixture containing 36% methane and 64% ethylene by volume, which acts as a surrogate with similar combustion properties as JP-7 [17]. A reduced, three-step mechanism [18, 19] is initially employed to characterize the combustion process. Arrhenius formulations of
The kinetic reaction rates are adopted, and the parameters are selected to retain robust/stable combustion in the current simulations.

LES calculations are then performed using the RAPTOR code framework developed by Oefelein [20, 21, 22]. RAPTOR solves the fully coupled conservation equations of mass, momentum, total energy, and species for a chemically reacting flow. It is designed to handle high Reynolds number, high pressure, real gas and liquid conditions over a wide range of Mach numbers while accounting for detailed thermodynamics and transport processes at the molecular level. The numerical schemes in RAPTOR involve non-dissipative, discretely conservative, staggered, finite volume differencing. This scheme is well-adapted to LES simulations as it eliminates numerical contamination of the subfilter due to artificial dissipation and ensures proper discrete conservations. The subfilter closure is obtained using a mixed Smagorinsky model [23, 24].

In our numerical studies, we designate five input variables as design variables that we can directly control during the design optimization process. These design variables are the global equivalence ratio \( \phi_G \), ratio of primary to secondary injector equivalence ratios \( \phi_R \), location of the primary injector \( x_1 \), location of the secondary injector \( x_2 \), and inclination of the primary injector \( \theta_1 \). Specifically, if \( \phi_1 \) and \( \phi_2 \) denote the equivalence ratios from the primary and secondary injectors, respectively, then \( \phi_G = \phi_1 + \phi_2 \) and \( \phi_R = \phi_1 / \phi_2 \).

The feasible design ranges are shown in Table 5. Furthermore, we allow a total of 11 model parameters to be uncertain, shown in Table 1 along with their uncertainty distributions. These distributions are assumed uniform across the ranges indicated.

The data utilized in the current analysis are from two-dimensional simulations of the scramjet computation, using grid resolutions where cell sizes are 1/8, 1/16, and 1/32 of the injector diameter \( d = 3.175 \text{ mm} \) (respectively denoted by “d/8”, “d/16”, and “d/32” cases in this paper), corresponding to respectively around 63 thousand, 250 thousand, and 1 million grid points. The number of time steps for each run are selected to maintain an approximately equal wall-clock time, while timestep sizes are determined adaptively based on the Courant-Friedrichs-Lewy (CFL) criterion. The simulations are warm-started on solutions engineered from a quasi-steady state nominal condition simulation, and take around \( 1.7 \times 10^3 \), \( 1.1 \times 10^4 \), and \( 3.9 \times 10^4 \) CPU hours per run for d/8, d/16, and d/32, respectively. The intense computational demand thus limits us with a total of 1053, 222, and 23 simulation runs for d/8, d/16, and d/32, respectively, where the simulation inputs are uniformly randomized jointly in the parameter and design spaces. Several QoIs are considered in this paper, either computed directly in the LES code or evaluated subsequently through postprocessing. All QoIs are time-averaged variables, where the instantaneous solutions corresponding to the second half for each run are time-averaged to generate one solution per run. The entire set of QoIs are described below, and while only a subset (in Table 4) directly enter the optimization problem definition, the rest (in Table 2) are retained to assist the discovery of the probabilistic manifold.

- **Combustion efficiency** \( (\eta_{\text{comb}}) \), a critical performance indicator for engines, is defined based on static enthalpy quantities [15, 25]:
  \[
  \eta_{\text{comb}} = \frac{H(T_{\text{ref}}, Y_e) - H(T_{\text{ref}}, Y_{\text{ref}})}{H(T_{\text{ref}}, Y_{e,\text{ideal}}) - H(T_{\text{ref}}, Y_{\text{ref}})}.
  \]
Here $H$ is the total static enthalpy, the “ref” subscript indicates a reference condition derived from the inputs, the “e” subscript is for the exit, and the “ideal” subscript is for the ideal condition where all fuel is burnt to completion. The reference condition corresponds to that of a hypothetical non-reacting mixture of all inlet air and fuel at thermal equilibrium. The numerator, $H(T_{ref}, Y_e) - H(T_{ref}, Y_{ref})$, thus reflects the global heat released during the combustion, while the denominator represents the total heat release available in the fuel-air mixture.

- **Burned equivalence ratio** ($\phi_{burn}$) is defined to be equal to $\phi_{burn} \equiv \phi_C \eta_{comb}$. It represents the air excess, and high values of $\phi_{burn}$ results from a combination of high thermal efficiencies and stoichiometric to rich equivalence ratios, and are associated with conditions away from blowout regimes.

- **Stagnation pressure loss ratio** ($P_{stagloss}$) is defined as
  \[
  P_{stagloss} = 1 - \frac{P_{s,e}}{P_{s,i}},
  \]
  where $P_{s,e}$ and $P_{s,i}$ are the wall-normal-averaged stagnation pressure quantities at the exit and inlet planes, respectively. Higher values of $P_{stagloss}$ illustrate pressure loss across the combustor and are associated with a decrease in efficiency.

- **Maximum root-mean-square (RMS) pressure** ($\max P_{rms}$) is the maximum RMS pressure across the spatial domain:
  \[
  \max P_{rms} = \max_{x,y} \sqrt{P(x,y)^2 - [P(x,y)]^2},
  \]
  with $P$ indicating time-averaged quantity. This QoI reflects the maximum pressure oscillation amplitude, which is useful for engine structural considerations.

- **Turbulence kinetic energy** (TKE) is characterized by the RMS velocity at a given location:
  \[
  \text{TKE} = \frac{1}{2} (u_{rms}^2 + v_{rms}^2).
  \]
  In the numerical investigations of this paper, we will look at TKE from multiple streamwise locations at $x_s/d = 5, 50, 85, 110, 140, 190, 220$. TKE captures statistical signatures of the spatial heterogeneity of turbulence.

- **Initial shock location** ($x_{shock}$) is the most upstream shock location, which we currently compute by detecting a rapid pressure change. More upstream locations of the initial shock train are linked with higher chances of unstart and vibration, and degraded safety of the scramjet engine.

### 3. Probabilistic Learning on Manifolds

The initial data set, denoted by $[x_d]$, consists of $N$ samples of $\mathbb{R}^n$-valued vectors, and is represented as an $n \times N$ matrix. Here, $n$ denotes the number of observables for each sample. The data can be thought of as $N$ points residing on a manifold in a $n$-dimensional ambient space. Alternatively, the data can be viewed as $n$ points in $N$-dimensional space. Through a diffusion maps analysis [7, 26, 27], the manifold localization in $\mathbb{R}^n$ is restated as subspace localization in $\mathbb{R}^N$, providing a more structured context for analysis. We first apply an affine transformation on the data using the eigenvectors of the empirical covariance matrix, transforming matrix $[x_d]$ into a new matrix $[\eta_d]$ whose rows are orthogonal. The purpose of this transformation is not to reduce dimensionality, but rather to improve the numerical conditioning of the data, and to permit a probabilistic treatment with uncorrelated variables. In some instances, dimension reduction may ensue, reflecting very strong linear correlations in the data. This step of the analysis incurs minimal computational burden while permitting us to leverage prior knowledge in the form of the empirical covariance of the data. Thus in general, matrix $[\eta_d]$ will be an $\nu \times N$ matrix, with $\nu \leq n$. In the next subsection, we provide a brief overview of the diffusion maps construction, followed by a brief overview of the projected Itô equation to sample on the associated manifolds.
with the data \[7\]. The right eigenvector 
\( g \) 
localized within the span of the first few 
\( m \) 
λ 
sorting the eigenvalues such that 
\( 1 = \lambda_1 \) 
\( \leq \lambda_2 \leq \cdots \leq \lambda_m \), it can be shown that the original data set \( \{\eta_d\} \) is localized within the span of the first few \( m \) of these eigenvectors \( \{g^\alpha, \alpha = 1, \ldots, m\} \) \[28, 7\]. We also denote by \( [g] \) the matrix whose \( \alpha^{th} \) column is \( g^\alpha \in \mathbb{R}^N \). The value of \( m \) is chosen to correspond to a noticeable reduction in the magnitude of the eigenvalues, specifically

\[
m = \arg \min_{\alpha} \left\{ \frac{\lambda_\alpha}{\lambda_2} < L \right\}
\]

with \( L \) chosen to be equal to 0.1. Typically, the ratio \( \lambda_m/\lambda_2 \) is of the order of 0.1. Clearly, these eigenvalues and hence the optimal value of \( m \) depend on \( \epsilon \), the width of the diffusion kernel. A maximum entropy argument \[29\] is used to simultaneously select the values of \( \epsilon \) and \( m \).
3.2. Itô Sampler on Manifolds

We construe localization to the manifold as satisfying underlying constraints, including the prevailing physics. This is justified by the fact that the information used to delineate the manifold is obtained from LES simulations. Any structure that relates the observables is thus inherited from the underlying physics. Our next task is to sample on this manifold in a manner that is consistent with the scatter originally observed in the data. In order to generate additional samples on the manifold, we first construct a probability model from the initial training data set, \([\eta_d]\). Two assumptions are required for this construction [9]. We first assume that the \(N\) columns of \([\eta_d]\) are independent realizations of an \(\mathbb{R}^d\)-valued random vector \(H\) whose empirical covariance matrix is the identity matrix, and estimate its probability density function (PDF) as a Gaussian mixture in the form,

\[
p_H(\eta) = \frac{1}{N} \sum_{j=1}^{N} \pi \left( \frac{\hat{s}_\nu}{s_\nu} \eta^{j}\eta - \eta \right), \tag{9}
\]

where \(\pi\) is the positive function from \(\mathbb{R}^\nu\) into \([0, +\infty]\) defined, for all \(\eta\) in \(\mathbb{R}^\nu\), by

\[
\pi(\eta) = \frac{1}{(\sqrt{2\pi} \hat{s}_\nu)^\nu} \exp \left\{ -\frac{1}{2\hat{s}_\nu^2} \|\eta\|^2 \right\},
\]

(10)

with \(\|\eta\|\) denoting the Euclidean norm in \(\mathbb{R}^\nu\) and where the positive parameters \(s_\nu\) and \(\hat{s}_\nu\) are defined by

\[
s_\nu = \left\{ \frac{4}{N(2 + \nu)} \right\}^{1/(\nu+4)}, \quad \hat{s}_\nu = \frac{s_\nu}{\sqrt{s_\nu^2 + \frac{N-1}{N}}}. \tag{11}
\]

With this choice of \(s_\nu\) and \(\hat{s}_\nu\), the mean-squared error is minimized [30] and realizations of random vector \(H\) are normalized, a requirement consistent with their construction through an eigen-decomposition [8]. This is the PDF of random vector \(H\) characterized by equation (9), we now consider the joint occurrence of the \(\nu\) data points. This joint behavior is significant as it carries a signature of intrinsic structure not available from the initial training data set, \([\eta_d]\) as realized in the data. In order to generate additional samples on the manifold, we first construct a probability model, next task is to sample on this manifold in a manner that is consistent with the scatter originally observed in the data. This approach consists of constructing an Itô equation that is constrained to the manifold, through projections, and whose invariant measure has the density specified by Equation (12). The next describe the procedure for generating samples of \([H]\) from the PDF specified in Equation (12).

The approach consists of constructing an Itô equation that is constrained to the manifold, through projections, and whose invariant measure has the density specified by Equation (12). First, we introduce the projection \(g\) on the subspace spanned by the \([g]\),

\[
[a] = [g] (\langle [g]^T [g] \rangle)^{-1}, \tag{13}
\]

the \(\nu \times N\) matrix \([N]\) whose entries are independent standard gaussian variables, the \(\nu \times N\) random matrix \([H_d]\) with known realization is \([\eta_d]\), and the \(\nu \times N\) matrix \([dW(r)]\) \((r \geq 0)\) whose \(i^{th}\) column is \(dW^i\) with \(\{W^i, i = 1, \cdots, N\}\) being independent copies of the \(\nu\)-dimensional normalized Wiener process. It can then be shown that solutions \(\{Z(r), r \geq 0\}\) of the following Itô stochastic differential equations [9]

\[
d[Z(r)] = [Y(r)] dr,
\]

\[
d[Y(r)] = [L([Z(r)] [g]^T)] [a] dr - \frac{1}{2} f_0 [Y(r)] dr + \sqrt{f_0} [dW(r)] [a], \tag{15}
\]

be shown that solutions \(\{Z(r), r \geq 0\}\) of the following Itô stochastic differential equations [9]
with the initial condition
\[ Z(0) = [H_x][a] , \quad Y(0) = [N][a] \quad \text{a.s.}, \]
are samples from the \( \nu \times N \) random matrix \( [H] = [Z][g]^T \) with PDF \( p_H([u]) = c q([u]) \) in which \( c \) is a constant of normalization, and where
\[
[L([u])]_{k\ell} = \frac{\partial}{\partial u_k^\ell} \log \{q(u^\ell)\} , \quad [u] = [u^1, \ldots, u^N].
\] (17)

Given our choice of Gaussian mixture model for \( q \), the expression for \( [L] \) can be expanded as follows,
\[
[L([u])]_{k\ell} = \frac{1}{q(u^\ell)} \frac{1}{2} \frac{1}{N} \sum_{j=1}^{N} \left( \frac{s_{\nu}}{s_{\nu}^2} \eta_{d,j} - u^\ell \right) \exp \left\{ -\frac{1}{2s_{\nu}^2} \left\| \frac{s_{\nu}}{s_{\nu}} \eta_{d,j} - u^\ell \right\|^2 \right\} .
\] (18)

The Itô equations specified by Equations (14) and (15) are solved using a Störmer-Verlet algorithm, a symplectic scheme well-adapted to Hamiltonian non-dissipative systems [31]. In that scheme, we use a value of \( f_0 \) equal to 0.23, and an integration step of 0.02.

### 3.3. Conditional Expectations

All the observables are used to characterize the basis vectors for the diffusion maps, and they all contribute to its construction. The greater the number of observables, the more distinct the interdependence between them, and the more robust is the associated manifold. Only a subset of these observables (QoIs), however, is relevant to the design optimization problem. Thus, once the manifold has been characterized and the initial training set augmented as described in the previous section, new observables relevant to the objective is obtained in the form [4, 30],

\[
E\{R|W = w_o\} \simeq \sum_{\ell=1}^{\nu_s} \frac{1}{(\nu_s(2 + m_w + 1))^{1/(4m_w + 1)}} \exp \left\{ -\frac{1}{2s^2} \left\| \hat{w}^\ell - \hat{w}_o \right\|^2 \right\} \sigma_R + \bar{R}.
\] (23)
In the above, $\hat{\tau}^f$ and $\hat{\mathbf{w}}^f$ are realizations of $\hat{R}$ and $\hat{\mathbf{W}}$, respectively. In evaluating chance constraints, it is often required to compute the probability of a random variable $R$ exceeding some threshold value, $q_f$, conditional on a specified values $w_o$ of the control variables. This probability can be expressed as the following integral,

$$h(q_f, w_o) = \mathbb{P}[R > q_f \mid \mathbf{W} = w_o] \simeq \int_{q_f}^{+\infty} p_R(r \mid w_o) dr,$$

which can be estimated as [5]

$$h(q_f, w_o) \simeq \sum_{\ell=1}^{\nu_s} h^\ell(\hat{q}_f) \exp \left\{ -\frac{1}{2s^2} \| \hat{\mathbf{w}}^\ell - \hat{\mathbf{w}}_o \|^2 \right\}$$

$$= \sum_{\ell=1}^{\nu_s} \exp \left\{ -\frac{1}{2s^2} \| \hat{\mathbf{w}}^\ell - \hat{\mathbf{w}}_o \|^2 \right\}$$

where

$$\hat{q}_f = (q_f - \mathcal{R})/\sigma, \quad h^\ell(\hat{q}_f) = \frac{1}{2} \left( 1 - \text{erf} \left( \frac{\hat{q}_f - \hat{\tau}^\ell}{s\sqrt{2}} \right) \right)$$

and $\text{erf}(y) = \frac{2}{\sqrt{\pi}} \int_0^y e^{-t^2} dt$ is the error function.

The augmented datasets were generated for each of the three spatial resolutions, $d/8$, $d/16$ and $d/32$. For each of these three cases, different values of $N$, the size of the training set, were used to delineate different manifolds and integrate forward the associated projected Itô sampler. The various cases are shown in Table (3). For $d/8$ and $d/16$, the 28 observables shown in Tables (1), (2), (4), and (5) are included in the manifold detection process. For the case $d/32$, given the small number of training samples (maximum of 23), only 9 observables were included in the analysis, consisting of the 4 QoIs and 5 controls, shown in Tables (4) and (5), respectively. The value of $\nu$, representing the dimension of the decorrelated observations is also shown in table (3). A threshold of 99% was used for truncation in this representation, and the ensuing slight reduction in the dimension of observables is indicative of strong linear dependence of one or two variables on the remaining variables. The value of $m$ shown in Table (3) represents the number of eigenvectors retained in the characterization of the diffusion manifold following the criterion specified by Equation (8). We note that the dimension of the DMAP embedding, $m$, is often larger than the dimension $\nu$ of the decorrelated observations. This reinforces the notion that the reduction achieved via this embedding is relative to the size of the data, $N$, and not relative to the size of the ambient space. This is consistent with the procedure, whereby the choice of observables in the training dataset is not necessarily exhaustive, and the ability of the DMAP construction to detect an intrinsic dimensionality irrespective of such limitations.

We also note that the number of sufficient points, $N$, depends on the level of fluctuations across several sets of $N$ points. If the manifold coordinates fluctuate noticeably across $N$-point samples, then the size of the sample is not sufficient. Our analysis, therefore, does not only consider the embedded dimensionality, but also the uncertainty which could be due to unobserved dynamics.

The reactive flows used in this paper to describe scramjet combustion are sufficiently complex that higher spatial resolutions result in distinctly new behaviors (that are arguably closer to reality) and not mere numerical refinements. With this in mind, it is expected that the coarser discretizations should exhibit greater fluctuations and necessitating more samples to be sufficiently-well characterized statistically. Indeed, an analysis of $d/8$ and $d/16$ models using training sets of size $N$ smaller than 50 did not converge during the optimization task.

<table>
<thead>
<tr>
<th>Resolution</th>
<th>$N$=Size of training set</th>
<th>$\epsilon$</th>
<th>$n$</th>
<th>$\nu$</th>
<th>$m$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$d/8$</td>
<td>50 100 200 500 1053</td>
<td>441</td>
<td>28</td>
<td>24</td>
<td>(25-29)</td>
</tr>
<tr>
<td>$d/16$</td>
<td>50 100 150 222</td>
<td>441</td>
<td>28</td>
<td>25</td>
<td>27</td>
</tr>
<tr>
<td>$d/32$</td>
<td>10 12 14 16 18 20 23</td>
<td>159</td>
<td>9</td>
<td>8</td>
<td>9</td>
</tr>
</tbody>
</table>

Table 3: Spatial resolution and size of training set for all cases; $\epsilon$=width of diffusion kernel; $n$= number of observables per sample; $\nu$=dimension of de-correlated variables; $m$=dimension of diffusion map.
For each of the cases shown in this table, a total of $n_{\text{NMC}}$ samples of $[Z(r)]$ were generated according to the manifold sampling scheme [9] summarized earlier in the paper, such that $\nu \times N \times n_{\text{NMC}} = 5 \times 10^6$. Conditional expectations and conditional probabilities expressed in Equations (23) and (25) respectively, are then estimated using these samples.

Figure (2) shows the marginal PDFs for resolution $d/8$ as the size $N$ of the training set is increased monotonically over the set (50, 100, 200, 500, 1053). The vertical marker in Figure (2)-(a) indicates the lower bound on the value of $\phi_{\text{burn}}$ as specified subsequently for design optimization. The vertical markers in Figures (2b) and (2-c) indicate the upper bounds on $P_{\text{stagloss}}$ and $\max P_{\text{rms}}$, respectively. The vertical bar in Figure (2-d) shows the range of the computed objective function specified in the optimization problem. This is further explained in the next section. The apparent convergence of these PDFs as $N$ increases suggests that the probability measure is eventually supported by the manifold, with the projection inducing only a small discrepancy. Figures (3) and (4) show similar results for spatial resolutions of $d/16$ and $d/32$, respectively. It is noted that there are clear differences between the PDFs at different spatial resolutions. This indicates that further mesh refinement, beyond $d/32$, is necessary to arrive at a fully converged mesh solution. Some of these PDFs, for instance, exhibit bimodal behavior, symmetry, or skewness at one resolution but not at the others.

Figure (5-a) shows a scatter plot of the 2nd and 4th diffusion coordinates, synthesized for 3 samples each of size $N=20$ for the $d/32$ case. Figure (5-b) shows a similar plot for the 3rd and 4th coordinates. We note the stability of the geometric structure under statistical perturbations of the underlying dataset. Similar stability is observed for other coordinates at $N=20$, but they do not exhibit a recognizable geometric structure. The diffusion coordinate were not stable for values of $N$ smaller than 20, although the optimal solutions (discussed in the following section) were not sensitive to these statistical perturbations.

4. Optimization Problem of Scramjet with Uncertainty

Our objective is to maximize the combustion efficiency $\eta_{\text{comb}}$, subject to constraints on the burned equivalence ratio $\phi_{\text{burn}}$, the stagnation pressure loss across the combustor $P_{\text{stagloss}}$, and the maximum pressure RMS $\max P_{\text{rms}}$. These QoIs, which we denote by $Q_i$, are shown in Table (4) together with their respective ranges.

The five control variables, denoted by $w_i$, used to attain the target design are shown in Table (5) together with the respective feasible ranges. We also denote by $P(E \mid Q_2)$ the probability of event $E$ conditional on $Q_2$ associated with the current iteration in the optimization process. We reinterpret the bounds on the QoIs, shown in Table (4) to be chance constraints to be satisfied with a specified probability greater than $1 - \alpha$. 

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<thead>
<tr>
<th>Variable name</th>
<th>Symbol</th>
<th>Lower/Upper bound</th>
</tr>
</thead>
<tbody>
<tr>
<td>$Q_1$ = Burned equivalence ratio</td>
<td>$\phi_{\text{burn}}$</td>
<td>$\geq L_1 = 0.1900$</td>
</tr>
<tr>
<td>$Q_2$ = Combustion efficiency</td>
<td>$\eta_{\text{comb}}$</td>
<td>to be maximized</td>
</tr>
<tr>
<td>$Q_3$ = Stagnation pressure loss</td>
<td>$P_{\text{stagloss}}$</td>
<td>$\leq U_3 = 0.3700$</td>
</tr>
<tr>
<td>$Q_4$ = Maximum RMS pressure</td>
<td>$\max P_{\text{rms}}$</td>
<td>$\leq U_4 = 0.2705$</td>
</tr>
</tbody>
</table>

Table 4: Quantities of interest

<table>
<thead>
<tr>
<th>Variable name</th>
<th>Symbol</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>$w_1$ = Global equivalence ratio</td>
<td>$\phi_G$</td>
<td>[0.5-1.0]</td>
</tr>
<tr>
<td>$w_2$ = Primary-secondary ratio</td>
<td>$\phi_R$</td>
<td>[0.25-0.35]</td>
</tr>
<tr>
<td>$w_3$ = Primary injector location</td>
<td>$x_1$</td>
<td><a href="m">0.231-0.2564</a></td>
</tr>
<tr>
<td>$w_4$ = Secondary injector location</td>
<td>$x_2$</td>
<td><a href="m">0.40755-0.43295</a></td>
</tr>
<tr>
<td>$w_5$ = Primary injector angle</td>
<td>$\theta_1$</td>
<td>[5-25°]</td>
</tr>
</tbody>
</table>

Table 5: Control variables
Figure 2: Probability density functions of four quantities of interest for resolution d/8. Each figure shows results for a range of values of $N$, the training-set size. Vertical markers indicate lower bound for optimization in subfigure a), upper bounds in subfigures b) and c) (Table (4)); vertical marker in subfigure d) shows computed range of optimal objective function as $N$ varies.
Figure 3: Probability density functions of four quantities of interest for resolution d/16. Each figure shows results for a range of values of $N$, the training-set size. Vertical markers indicate lower bound for optimization in subfigure a) , upper bounds in subfigures b) and c) (Table (4)); vertical marker in subfigure d) shows computed range of optimal objective function as $N$ varies.
Figure 4: Probability density functions of four quantities of interest for resolution d/32. Each figure shows results for a range of values of $N$, the training-set size. Vertical markers indicate lower bound for optimization in subfigure a) , upper bounds in subfigures b) and c) (Table (4)); vertical marker in subfigure d) shows computed range of optimal objective function as $N$ varies.
and replace these bounds with the following equations,
\[
\begin{align*}
  & P\{Q_1(w) > L_1 \mid Q_2\} > 1 - \alpha \quad (27a) \\
  & P\{Q_3(w) < U_3 \mid Q_2\} > 1 - \alpha \quad (27b) \\
  & P\{Q_4(w) < U_4 \mid Q_2\} > 1 - \alpha \quad (27c)
\end{align*}
\]

We are thus dealing with the following optimization problem:
\[
\begin{align*}
  w_{\text{opt}} &= \arg \max_{w \in C_w \subset \mathbb{R}^5} J(w) \quad (28a) \\
  \text{subject to} & \quad c_i < 0 \quad i = 1, 2, 3 \quad (28b)
\end{align*}
\]
where the objective function is given by the following expectation,
\[
J(w) = \mathbb{E}[Q_2], \quad (29)
\]
the feasible domain, \(C_w\), is the 5-dimensional cube specified by the following ranges,
\[
\begin{align*}
  & w_1 \in [0.5, 1] \\
  & w_2 \in [0.25, 0.35] \\
  & w_3 \in [0.231, 0.2564] \\
  & w_4 \in [0.40755, 0.43295] \\
  & w_5 \in [5, 25] \quad (30)
\end{align*}
\]
and the constraints are specified as
\[
\begin{align*}
  & c_1(w_{\text{opt}}) = 1 - \alpha - P\{Q_1(w_{\text{opt}}) > L_1 \mid Q_2\} \quad (31a) \\
  & c_2(w_{\text{opt}}) = 1 - \alpha - P\{Q_3(w_{\text{opt}}) < U_3 \mid Q_2\} \quad (31b) \\
  & c_3(w_{\text{opt}}) = 1 - \alpha - P\{Q_4(w_{\text{opt}}) < U_4 \mid Q_2\} \quad (31c)
\end{align*}
\]
The optimization problem is formulated and solved for each of the cases shown in Table (3). It is noted from Figures (2)-(4) that for \(d/16\), the lower bound on \(\phi_{\text{burn}}\) and the upper bound on \(P_{\text{stagloss}}\) are both in the
tail area, with a small likelihood of being satisfied. For d/8 and d/32, on the other hand, the bounds are situated centrally within the support of the distribution.

To solve this non-convex constrained optimization problem we rely on GA [32], a genetic algorithm package within the R language [33]. We set the population size within each generation to 50, and the maximum number of iterations to 1000. We also set the number of unchanged iterations at convergence to 100. The probability of mutation in a parent chromosome and the number of best fitness individuals to survive at each generation (elitism) are set to 0.8 and 2.5, respectively. Finally the number of consecutive generations without improvement that characterizes the optimal solution is set to 100. In all the results shown below, an optimal solution was attained without activating a stopping criterion based on the maximum number of iterations (set to 1000). The optimization problem was solved on multicore CPUs taking advantage of the multicore feature of GA.

Following a first set of GA optimization runs using a randomly selected initial population, a second set of GA runs was performed on all cases in Table (3) using all the results from the first set as an initial population for all cases. This was done in order to test whether a unique optimal solution exists across all resolutions, which was not the case. Figure (6) shows the values of the optimal solutions as evaluated by the genetic optimization algorithm. Each of the 5 subfigures there shows the values of one of the 5 control variables. Within each subfigure, the red circles indicate solutions associated with d/32 resolution, the green squares show solutions for the d/16 resolution, and the blue losanges correspond to the d/8 resolution. Within each resolution, results are shown for all the values of N (size of initial training set) indicated in Table (3).

We note immediately that in most cases, the optimal values of the control variables are evaluated in the neighborhood of their bounds. This is indicative that the attained optimal solutions are efficient relative to the constraints.

The vertical markers in Figures (2(d)-4(d)) indicate the range of the objective function $E(\eta_{comb})$ at the optimal solution. This range corresponds to the various values of N. It is noted that, while the optimal value of the objective function is robust to changes in N for both the d/8 and d/32 resolutions, it is very sensitive to N for the d/16 resolution; this is despite the seemingly converged behavior of the marginal density functions, as function of N, in Figure (3(d)). It is clear from the foregoing observations that performance and convergence criteria based on marginal density functions may be misleading when exploring high-dimensional problems. In general, the conditional expectations used in the optimization problem probe higher order joint density functions of the QoI and the control variables. The accurate estimation of these requires significantly larger number of samples, which is facilitated by the manifold sampling approach used in the present paper.

5. Conclusion

The paper presents the application of a recent machine learning algorithm to a very large scale LES-based optimization problem associated with scramjet combustion. The approach used relies on learning a joint statistical model of the observables which include QoIs, input parameters (including design parameters), as well as other observables. The scatter is construed to occur around an intrinsic structure to this dataset that relates its components. The variance of this scatter around this structure is much smaller than in ambient space, and can thus be explained with a smaller dataset. The combination of diffusion maps, Itô sampling, and non-parametric statistics come together in developing efficient algorithms that encapsulate these ideas.

Three different LES spatial resolutions are explored, each with a different number of samples in the associated training set. Good convergence of the optimal solution is observed as function of the size of the training set, although the optimal solutions are different for each of the three resolutions. It is noted that while the optimal solution in each of the three cases tends to occur at a corner of the feasible domain (i.e. several design variables lie along their respective lower or upper bound), the specific corner depends on the specific spatial resolution. This highlights some of the challenges facing multi-fidelity/multi-resolution optimization strategies.

6. Acknowledgment

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Figure 6: Optimal values of the five control parameters for \(\frac{d}{8}\), \(\frac{d}{16}\), and \(\frac{d}{32}\) resolutions and different sizes of training set.
7. References

References


