

ExCALIBUR

Selection of techniques for Model Order Reduction

M2.5.2

The report describes work for ExCALIBUR project NEPTUNE at Milestone 2.5.2. This binds the reports 2047352 2-TN-01[\[1\]](#page-2-0) and 2047352 2-TN-02[\[2\]](#page-2-1) as of August 27, 2021, which are presented in composite form: sections 2 and 3 (pp.1-7 of total 20 pp) comprise the original 2047352 2-TN-01.

The aim of this report is to provide a preliminary description of potential reduction of size of inputs, size of design, and size of outputs, and associated computational benefits of reduced order modelling for fusion codes, with a focus on relevant proxyapps. Due to time constraints, a single proxyapp - describing anisotropic heat transport - is considered, though this is supported by toy examples and further references from the oeuvre of the authors (who are world experts in the field), including illustrative examples of coupled models.

The report stresses the need for reduced order models (hereafter ROMs) in situations mandating large ensembles of expensive simulations (as are expected to arise in NEPTUNE for e.g. uncertainty quantification or data assimilation aspects). A bird's eye perspective on the subject area is provided and justification is given for the decisions to focus primarily on non-intrusive ROMs meaning that there is no need to alter the subject simulation code; the ROM is then 'data-driven' in that it is constructed using solely the outputs of the subject - a decision entirely consistent with the NEPTUNE separation of concerns ethos, and more specifically to use the family of Gaussian process surrogates (hereafter GP-ROM). Some relevant aspects of the current state-of-the-art are indicated, with reference to the authors' recent works.

The report details a particular non-intrusive ROM as applied to the anisotropic diffusion problem (viz. a two-dimensional diffusion equation in which the diffusivity is represented by a spatiallyvarying anisotropic tensor). The numerical representation is via the finite-element package Firedrake, giving a solution comprising 78961 degrees of freedom. This model space dimensionality is reduced to 25 using the technique of proper orthogonal decomposition (POD); the reduced system is then used to fit a Gaussian process using a constant mean (trend function) and a Matérn kernel for the prior covariance. The efficacy of the resulting model, in terms of predicted mean and confidence interval, is shown to be reasonable by comparing some of the ROM outputs to the corresponding full numerical solutions. This section contains also a technical exposition of the offline regression used to construct the Gaussian process, in which the main computational bottleneck a cubic scaling in the number of data samples - is emphasized.

A subsequent section provides a description of active learning (AL) for constructing a GP-ROM. AL entails choosing the next sampling position dynamically based on the current ROM - various algorithms can be used: the MacKay strategy (giving ALM) is to minimize the local maximum posterior predictive variance, leading to the problem of over-sampling at boundaries, while the alternate Cohn scheme (ALC) involves minimizing the same variance averaged over the domain (the need for additional model evaluations during these steps is avoided by a property of Gaussians under Bayesian combination). The AL schemes are applied to the anisotropic diffusion problem, demonstrating that (1) the active learning saturates at large sample numbers giving there similar performance to a static Latin hypercube design (LHD); and (2) the ALM performs worse than ALC or a LHD at large sample number because of the boundary over-sampling problem. For realistic fusion problems, the samples are likely to be sufficiently expensive that the saturated regime is not encountered, hence AL is expected to be useful. The authors propose also a strategy for identifying and excluding regions of flat response surface in the offline phase, in order to mitigate against the cubic scaling problem in fitting a GP-ROM with a large number of samples - this involves a dense exploration of the response surface and then the selection from these outputs of an appropriate set of initial data, followed by active learning using the data already generated (thus giving a procedure that is explicitly parallelizable). This approach provides arguably some protection against the problem of missing extreme outlying events - one must bear in mind that the quality of the ROM is to a large extent dependent on the experimental design. As a caution, however, it must be noted that, for the NEPTUNE use case, a dense sampling of the response surface is likely to represent a very large computational challenge.

The report closes with a discussion of possible future directions in a section in which the recommendations interspersed at appropriate locations serve in lieu of a formal concluding section. A proposal to test some intrusive methods in order to mitigate the problem of excluding extreme events when applying dimension reduction techniques (e.g. POD) is included; the output here would be a physics-informed ROM (references in the context of machine learning are given).

A subsection presents deep GPs (meaning that the outputs of a GP are used sequentially as the inputs to another GP) where the motivation is to construct ROMs for systems with regime change; the deep GP is able to incorporate multiple kernels, as evidenced by the example of fitting a toy dataset containing distinct quiescent and active regions.

A further subsection focuses on active subspace methods for reducing the *input* dimension (cf. the POD in Section 3, which reduces the number of internal model states). A technique called *sufficient dimension reduction* is briefly outlined (with further reference provided), giving a general method for detecting the linear combinations of inputs to which a model is most sensitive.

A final subsection outlines, with the aid of a toy example, linked GPs, which differ from deep GPs in that they model a system that can be explicitly decomposed into component parts, the simpler components being more amenable to GP emulation than is the whole (the divide-andconquer principle applies). This gives also a semi-intrusive procedure for constructing a GP-ROM for a coupled system. It is explained that there remain challenges associated to the dimension reduction of the intermediate data in cases where the intermediate states have input dimension much higher than that of the global input. It is clear that active sampling in the case of a linked GP can mitigate the problem of under-exploration of the input parameter space of intermediate components in a coupled model i.e. the fact that the space-filling property of a LHD is lost once the inputs have been propagated through one or more component models (thus giving, amongst other problems, the potential to miss extreme events).

Acknowledgement

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References

- [1] D. Ming and S. Guillas. Report on suitability and potential of ROM to fusion models : A Nonintrusive ROM for Solvers with High-dimensional Outputs. Technical Report 2047352 2-TN-01, UKAEA Project Neptune, 2021.
- [2] D. Ming and S. Guillas. Report on suitability and potential of Reduced Order Modelling (ROM) to fusion models: Gaussian Process ROM for Solvers with High-dimensional Outputs. Technical Report 2047352 2-TN-02, UKAEA Project Neptune, 2021.

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¹ Report on suitability and potential of Reduced Order Modelling ² (ROM) to fusion models

Gaussian Process ROM for Solvers with High-dimensional Outputs

Deyu Ming and Serge Guillas

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1 Disclaimer

 We are very grateful to Dr Patrick Farrell for the provision of the proxyApp modelling the anisotropic heat transport problem. It is the only fusion model we could access over the short period of the funded project (4 January 2021 - 31 July 2021). We discussed with the NEPTUNE team (Benjamin Dudson and Patrick Farrell) the possibility of using another model to couple two models in a one-way coupling for UQ using ROM: the anisotropic heat transport model and the isotropic heat conduction to the solid wall. But the wall boundary proxyApp is not yet available. As a result, we could not examine in practice the possibility of implementing ROM for UQ in the context of nuclear fusion modelling where models are coupled. We nevertheless [p](#page-21-0)rovided some examples of UQ coupling at the end of this report from the paper [Ming and](#page-21-0) [Guillas](#page-21-0) [\(2021\)](#page-21-0) and discussed future directions below.

¹⁹ 2 Introduction

 Many modern physical computer models involve solving PDEs with numerical solvers, such as finite element methods (FEM), which can be computationally expensive due to

- **•** ever more complex and larger-scale models;
- high-dimensional input and output;
- large demands on computational resources.

 These create challenges to efficient uncertainty quantification of computer models, such as the fusion models, as we often need to run the models many times for tasks such as sensitivity analysis, uncertainty propagation and model calibration. To tackle these challenges, reduced order models (ROM) are needed to

- serve as low-dimensional replacements with comparable accuracy;
- reduce evaluation time of original solvers;
- save storage, e.g., for high-dimensional output.

 Traditional reduced order models, also known as intrusive reduced order models, often are constructed using reduced basis methods [\(Quarteroni et al., 2015\)](#page-21-1), among which the Proper ³⁴ Orthogonal Decomposition (POD) is perhaps the most popular technique. The intrusive reduced order models for original high-fidelity models with high-dimensional output are typically built using a two-phase procedure called offline-online decomposition:

³⁷ • *offline phase*: high-fidelity solutions/outputs are obtained and reduced basis is calculated;

 • online phase: the original problems are projected onto the reduced space for efficient computation of solutions at new inputs.

 However, the online phase of the intrusive reduced order modelling is challenging in practice because:

 • expertise and domain knowledge are required to project the equations and physics of the original high-fidelity problems to constructed reduced space;

• dimensionality reduction techniques are largely constrained by the problem formulation;

• uncertainty is not incorporated.

 For these reasons, in this report we focus on non-intrusive reduced order models for problems with high dimensional outputs, utilising the family of Gaussian process (GP) surrogates (also known as emulators). GP emulators have been successfully implemented for dimension reduction of either outputs or inputs. For instance:

- [Guillas et al.](#page-21-2) [\(2018\)](#page-21-2) used Functional Principal Components Analysis (FPCA) as an equiv- alent approach to POD for time series outputs of tsunami waves, and [Chang et al.](#page-20-0) [\(2019\)](#page-20-0) used Spherical Harmonics and Gaussian Markov Random Fields for optimal reduction of surfaces outputs.
- For inputs, [Liu and Guillas](#page-21-3) [\(2017\)](#page-21-3) employed a kernel-based approach to extract the few input field directions of most influence for the outputs in order to build GPs with few input dimensions (orders of magnitude gain in dimension).

 The report is organised as follows. In Section [3,](#page-4-0) a non-intrusive ROM with GP surrogates and POD is described and applied in a anisotropic heat transport problem. We then propose and discuss an active learning procedure to construct the introduced non-intrusive ROM with an illustrative example in Section [4.](#page-9-0) Future directions are discussed in Section [5.](#page-15-0)

3 Non-intrusive ROM with Gaussian Process Surrogates

 The non-intrusive reduced order modelling is a data-driven approach that uses a statistical surrogate model to mimic the functional relations between the model input and constructed reduced output space in the online phase of the offline-online decomposition. The utilisation of statistical surrogates alleviates the difficulties involved in reformulating the original high-fidelity problems under the intrusive reduced order modelling. In particular, with GP surrogates we

⁶⁷ are able to quantify uncertainty of the high-dimensional outputs predicted at unobserved input ⁶⁸ positions.

69 Let $\mathbf{X} \in \mathbb{R}^{N \times D}$ contain N sets of D dimensional input of a computer model, which produces ⁷⁰ N corresponding sets of K dimensional output $\mathbf{Y} \in \mathbb{R}^{N \times K}$ accordingly. Then, one can mimic ⁷¹ the functional relationships between the input **X** and each output dimension $\mathbf{Y}_k \in \mathbb{R}^{N \times 1}$ by a 72 GP surrogate \mathcal{GP}_k independently for $k = 1, \ldots, K$ without considering the dependence between ⁷³ output dimensions [\(Gu and Berger, 2016\)](#page-20-1). Ignoring the potential cross-dependence does not ⁷⁴ pose a serious issue unless we are interested in the joint distribution of the output, and it can be ⁷⁵ shown [\(Kyzyurova, 2017\)](#page-21-4) that the independently constructed GP surrogates correspond to the ⁷⁶ marginal GPs of a joint GP surrogate under certain dependence structures. The GP surrogate τ GP_k is formally defined as a multivariate normal distribution with respect to \mathbf{Y}_k :

$$
\mathbf{Y}_k \sim \mathcal{N}(\boldsymbol{\mu}_k(\mathbf{X}), \sigma_k^2 \mathbf{R}_k(\mathbf{X})),
$$

⁷⁹ in which the *i*-th element of $\mu_k(\mathbf{X}) \in \mathbb{R}^{N \times 1}$ is often specified by a trend function $f_k(\mathbf{X}_i)$ with 80 $\mathbf{X}_i \in \mathbb{R}^{1 \times D}$ being the *i*-th row of **X**, and the *ij*-th element of $\mathbf{R}_k(\mathbf{X}) \in \mathbb{R}^{N \times N}$ is given by ⁸¹ $c_k(\mathbf{X}_i, \mathbf{X}_j)$, where c_k is a given kernel function. The trend function f_k can be formulated as a 82 linear combination of a set of basis functions of X_i and we assume a constant trend function 83 $f_k(\mathbf{X}_i) = b_k$ in this report.

⁸⁴ There are various choices for c_k (see [Rasmussen and Williams](#page-21-5) [\(2006\)](#page-21-5)). In this report, we use ⁸⁵ the separable kernel function:

$$
c_k(\mathbf{X}_i, \mathbf{X}_j) = \prod_{d=1}^D c_{k,d}(X_{id}, X_{jd}),
$$

 87 where $c_{k,d}$ is a one-dimensional kernel function. A typical choice for $c_{k,d}$ in computer model ⁸⁸ emulation is the squared exponential (SExp) kernel:

$$
c_{k,d}(X_{id}, X_{jd}) = \exp\left\{-\frac{(X_{id} - X_{jd})^2}{\gamma_{k,d}^2}\right\},\,
$$

90 where $\gamma_{k,d} > 0$ is the range parameter. However, the SExp kernel has been criticised for its ⁹¹ over-smoothness [\(Stein, 1999\)](#page-22-0) for physical problems as well as its associated ill-conditioned 92 problems [\(Dalbey, 2013;](#page-20-2) [Gu et al., 2018b\)](#page-21-6). Another popular kernel choice is the Matérn ker-⁹³ nel [\(Rasmussen and Williams, 2006\)](#page-21-5):

$$
c_{k,d}(X_{id}, X_{jd}) = \exp\left(-\frac{\sqrt{2p+1}r_{ij,d}}{\gamma_{k,d}}\right) \frac{p!}{(2p)!} \sum_{i=0}^p \frac{(p+i)!}{i!(p-i)!} \left(\frac{2r_{ij,d}\sqrt{2p+1}}{\gamma_{k,d}}\right)^{p-i},
$$

95 where $r_{ij,d} = X_{id} - X_{jd}$. The Matérn kernel is known to be less prone to ill-conditioning ⁹⁶ issues and provides a reasonably adequate smoothness to the GP surrogates. In particular, the 97 Matérn-2.5 kernel, which is defined as the Matérn kernel with $p = 2$:

98
$$
c_{k,d}(X_{id}, X_{jd}) = \left(1 + \frac{\sqrt{5}|X_{id} - X_{jd}|}{\gamma_{k,d}} + \frac{5(X_{id} - X_{jd})^2}{3\gamma_{k,d}^2}\right) \exp\left\{-\frac{\sqrt{5}|X_{id} - X_{jd}|}{\gamma_{k,d}}\right\},\,
$$

⁹⁹ [i](#page-22-1)s the default kernel choice for many computer model emulation packages, such as DiceKriging [\(R](#page-22-1)ous-

100 [tant et al., 2012\)](#page-22-1) and RobustGaSP [\(Gu et al., 2018a\)](#page-21-7). Therefore, we employ the Matérn-2.5 kernel ¹⁰¹ in this report.

The posterior predictive distribution $\mathcal{N}(\widehat{\mu}_k(\mathbf{x}^*), \widehat{\sigma}_k^2(\mathbf{x}^*))$ of \mathcal{GP}_k with respect to the output ¹⁰³ $Y_k^*(\mathbf{x}^*)$ at an unobserved input position \mathbf{x}^* is given in different analytical forms depending 104 on how the model parameters b_k , σ_k^2 and $\{\gamma_{k,d}\}_{d=1,\dots,D}$ are estimated. Different maximumlikelihood-based estimation approaches and the corresponding expressions for $\hat{\mu}_k(\mathbf{x}^*)$ and $\hat{\sigma}_k^2(\mathbf{x}^*)$ ¹⁰⁶ are discussed in [Roustant et al.](#page-22-1) [\(2012\)](#page-22-1); [Gu et al.](#page-21-6) [\(2018b\)](#page-21-6).

 The main computational bottlenecks of the GP surrogate construction are the number of data points N and the dimension K of the output of a computer model. Since the inference 109 of GP surrogates involve inversions of the correlation matrix $\mathbf{R}_k \in \mathbb{R}^{N \times N}$ with computational 110 complexity of $\mathcal{O}(N^3)$, it soon becomes computationally prohibitive to build GP surrogates in practice when N is more than several thousands. In such a case, one may need sparse approx- imations [\(Liu et al., 2020\)](#page-21-8) to the GP to reduce the computational complexity induced by the big data.

¹¹⁴ In computer model experiments, one often does not have big data (i.e., realisations from ¹¹⁵ the underlying computer model) due to the limited computational budget. However, if the $_{116}$ input dimension D is large, then small data are insufficient to explore adequately the whole ¹¹⁷ input domain and thus the resulting GP surrogates can be inaccurate. High input dimension ¹¹⁸ also causes challenges to the model estimation because a large number of range parameters $_{119}$ { $\gamma_{k,d}$ } $_{d=1,...,D}$ need to be estimated for each output dimension. To alleviate this issue, one can 120 reduce the input dimension D to P such that $P \ll D$ by dimension reduction techniques such as ¹²¹ POD, kernel dimension reduction [\(Liu and Guillas, 2017\)](#page-21-3), and active subspace [\(Tripathy et al.,](#page-22-2) $122 \quad 2016$).

 A high output dimension K creates the issue that it can be computational burdensome to build K independent GP surrogates: without parallel implementation the training and validation of a huge amount of GP surrogates are practically infeasible. This report tackles the latter issue on high-dimensional outputs (e.g., a snapshot where each point on the snapshot represents a FE solution and contributes to the output dimensionality) produced by computer models. Perhaps the most straightforward approach to address the issue is to reduce the output dimension K to 129 L such that $L \ll K$ by POD.

- 130 The POD of $\mathbf{Y} \in \mathbb{R}^{N \times K}$ can be done with following steps:
- 131 1. Compute the sample mean $\mu_Y \in \mathbb{R}^{1 \times K}$ of Y and obtain the centred output matrix $Y_c =$ 132 $Y - \mu_Y;$
- 133 2. Implement the eigendecomposition of $\mathbf{G} = \frac{1}{N} \mathbf{Y}_c \mathbf{Y}_c^\top$ such that $\mathbf{G} = \mathbf{V} \mathbf{\Lambda} \mathbf{V}^\top$, where the columns of $V \in \mathbb{R}^{N \times N}$ contains the eigenvectors of G and the diagonal of $\Lambda \in \mathbb{R}^{N \times N}$ 135 contains the corresponding eigenvalues $(\lambda_1, \ldots, \lambda_N)$ in descending order;
- 136 3. Compute $\tilde{\mathbf{V}} = \mathbf{Y}_c^{\top} \mathbf{V} \in \mathbb{R}^{K \times N}$, which contains the eigenvectors of sample covariance matrix $\mathbf{C} = \frac{1}{N} \mathbf{Y}_c^{\top} \mathbf{Y}_c;$
- 138 4. Choose $L \leq N$ and obtain the low dimensional output $\hat{\mathbf{Y}} = \mathbf{Y}_c \tilde{\mathbf{V}}_L \in \mathbb{R}^{N \times L}$, where

¹³⁹ $\tilde{\mathbf{V}}_L \in \mathbb{R}^{K \times L}$ contains the first L eigenvectors included in $\tilde{\mathbf{V}}$.

140 One can also obtain \tilde{V} by performing the singular value decomposition (SVD) of Y_c that is im-¹⁴¹ plemented, e.g., in the PCA function of Python package scikit-learn [\(Pedregosa et al., 2011\)](#page-21-9). 142 After obtaining the low dimensional data $\mathbf{\hat{Y}}$, we then construct L independent GP surrogates 143 of each of L dimensions of $\hat{\mathbf{Y}}$. Let $\mathcal{N}(\hat{\mu}_{l}(\mathbf{x}^{*}), \hat{\sigma}_{l}^{2}(\mathbf{x}^{*}))$ be the posterior predictive distribution ¹⁴⁴ of $\widehat{Y}_l^*(\mathbf{x}^*)$, the *l*-th dimension of the low dimensional output, predicted at an unobserved input 145 position x^* . Then the posterior predictive distribution of the corresponding high dimensional ¹⁴⁶ output $\mathbf{Y}^*(\mathbf{x}^*) \in \mathbb{R}^{1 \times K}$ is given by

$$
\mathcal{N}\left(\widehat{\boldsymbol{\mu}}(\mathbf{x}^*)\tilde{\mathbf{V}}_L^\top + \boldsymbol{\mu}_{\mathbf{Y}}, \tilde{\mathbf{V}}_L\widehat{\boldsymbol{\Sigma}}(\mathbf{x}^*)\tilde{\mathbf{V}}_L^\top\right),
$$

¹⁴⁸ where $\hat{\boldsymbol{\mu}}(\mathbf{x}^*) = (\hat{\mu}_1(\mathbf{x}^*), \dots, \hat{\mu}_L(\mathbf{x}^*))$ and $\hat{\boldsymbol{\Sigma}}(\mathbf{x}^*) = \text{diag}(\hat{\sigma}_1^2(\mathbf{x}^*), \dots, \hat{\sigma}_L^2(\mathbf{x}^*))$.

 Figure [1](#page-7-0) demonstrates the procedure to build non-intrusive reduced order model with GP surrogates. In the offline phase, dimension-reduction techniques, e.g., POD, are applied to reduce the high-dimensional output to a low-dimensional space. Then in the online phase, GP surrogates are constructed independently on each reduced dimension. Using the constructed GP surrogate and reduced basis, one can obtain the predicted low-dimensional and in turn the high-dimensional output at new input positions with little computational efforts.

Figure 1: The workflow to construct non-intrusive ROM with GP. The black arrows represent the offline phase; the blue arrows represent the online phase; the red arrows represent the prediction procedure using the constructed non-intrusive ROM with GP.

¹⁵⁵ 3.1 Example: 2-D model of anisotropic heat transport

 In this section, we explore the non-intrusive ROM with GP to mimic the FE solver to the 2-D problem "Open field lines with oscillating anisotropy directions" in [Deluzet and Narski](#page-20-3) [\(2019\)](#page-20-3). 158 The problem has two key inputs m and α that control the anisotropy of the solution field, i.e., the anisotropy direction is defined by

$$
\mathbf{b} = \frac{\mathbf{B}}{|\mathbf{B}|}, \quad \mathbf{B} = \begin{pmatrix} \alpha(2y-1)\cos(m\pi x) + \pi \\ \pi\alpha m(y^2 - y)\sin(m\pi x) \end{pmatrix},
$$

161 where $m/2$ is the number of oscillation periods in the computational domain and α is the ¹⁶² amplitude. The output is a high-dimensional 2-D field defined on the square computational 163 domain $[0, 1] \times [0, 1]$ and allows a closed form solution.

¹⁶⁴ 3.1.1 Experimental Setup

¹⁶⁵ To construct the reduced basis via the POD and the GP surrogate, N=40 samples are arranged 166 in a Latin hypercube over $m \in [0, 12]$ and $\alpha \in [0, 3]$ (see the left plot in Figure [2\)](#page-8-0). We then run ¹⁶⁷ the FE solver (implemented in Firedrake [\(Rathgeber et al., 2016\)](#page-21-10)) of the toy problem to obtain 168 the corresponding 2-D outputs, each of which contains FE solutions on $K = 78961$ nodes. These 169×78961 high-dimensional outputs are then reduced to 40 low-dimensional outputs (40×25) ¹⁷⁰ using POD by retaining the first 25 principal components out of the total 40 components, see the right plot in Figure [2,](#page-8-0) where the cumulative explained variance is defined as $\frac{\sum_{i=1}^{L} x_i}{\sum_{i=1}^{N} x_i}$ ¹⁷¹ the right plot in Figure 2, where the cumulative explained variance is defined as $\frac{\sum_{i=1}^{L} \lambda_i}{\sum_{i=1}^{N} \lambda_i}$ with L ¹⁷² be the number of components.

Figure 2: (Left): Training and designing points generated for the inputs m and α . The blue points are design input locations generated from the Latin hypercube design and the red points are testing input locations; (Right): cumulative explained variance given by the POD.

¹⁷³ GP surrogates are then constructed independently for each of the 25 dimensions of the 174 reduced order data. GP surrogates are trained with the Matérn-2.5 kernel using the RobustGaSP ¹⁷⁵ package in R.

¹⁷⁶ 3.1.2 Experimental Results

177 We test the constructed non-intrusive ROM at four testing input positions $(m, \alpha) = (6, 2)$, $(10, 2), (1, 2)$ $(10, 2), (1, 2)$ $(10, 2), (1, 2)$ and $(10, 0)$ (see the left plot of Figure 2). The FE solutions (from the Firedrake) and the predicted solutions from the built ROM are compared in Figure [3.](#page-9-1) The normalised (to the range of FE solutions) errors between the FE solutions and the predicted solutions from the built ROM are shown in Figure [4.](#page-9-2) The coverage of the ROM (i.e., the instances that the FE solutions fall within the predictive bounds of GP-based ROM) are also given in Figure [5.](#page-10-0)

¹⁸³ It can been seen from these results that the constructed ROM with GP could predict well ¹⁸⁴ the FE solutions of the anisotropic problem at input locations that are not realised. Among 185 the four testing positions, the final case with $m = 10$ and $\alpha = 0$ presents the largest normalised 186 errors up to 13%. This is not a surprising result because m has no effect on the FE solution of 187 the problem when $\alpha = 0$. However, this information is not fully captured in the training data ¹⁸⁸ and thus not gained by the non-intrusive ROM with GP, which is pure data-driven method that 189 only understands the functional relation between m , α and the solution field from the training ¹⁹⁰ set. As a result, we could observe 5 blurred oscillation periods in the predicted solutions from

Figure 3: Comparisons of FE solutions to the predicted solutions given by the constructed GPbased ROM. The first row gives the FE solutions. The second row gives the predicted solutions from the GP-based ROM. The columns from left to right correspond to testing input positions $(m, \alpha) = (6, 2), (10, 2), (1, 2)$ and $(10, 0)$ respectively.

Figure 4: The normalised errors between FE solutions and the predicted solutions from the ROM with GP surrogate. The plots from left to right correspond to testing input positions $(m, \alpha) = (6, 2), (10, 2), (1, 2)$ and $(10, 0)$ respectively.

 ROM in Figure [3.](#page-9-1) However, the predictive interval (whose upper and lower bounds are given 192 at two standard deviations $2\hat{\sigma}$ above and below the predictive mean $\hat{\mu}$) of the GP-based ROM covers the FE solutions sufficiently in this case, demonstrating that one can benefit from the predictive uncertainty embedded in the non-intrusive ROM coupled with GP emulation.

¹⁹⁵ 4 Active learning for Non-intrusive ROM with Gaussian Pro-¹⁹⁶ cess Surrogates

¹⁹⁷ 4.1 Why Active Learning?

¹⁹⁸ Active learning, also known as sequential design, is a collection of approaches that adaptively ¹⁹⁹ enrich the training points for surrogate modelling of computer solvers. In comparison to one-

Figure 5: The coverage of constructed ROM with GP, giving the instances that FE solutions fall within the predictive bounds provided by the ROM with GP. 1 indicates that the FE solution is covered by the predictive interval (whose upper and lower bounds are given at two standard deviations $2\hat{\sigma}$ above and below the predictive mean $\hat{\mu}$) and 0 indicates otherwise. The plots from left to right correspond to testing input positions $(m, \alpha) = (6, 2), (10, 2), (1, 2)$ and $(10, 0)$ respectively.

 shot designs, such as Latin-hypercube designs (LHD), the active learning is preferred in many cases:

- One wants a proper utilisation of computational resources. Active learning allows one to choose computer model input locations adaptively, and therefore can monitor the quality of the resulting surrogate model while the active learning is in progress and determine whether to pause or continue the model evaluations;
- More computer model evaluations are needed in the input region of interest. Unlike static space-filling designs, such as LHD, active learning, depending on the quality of the under- lying surrogate model (as we will discuss in Section [4.4\)](#page-13-0), could direct the computer models to evaluate at input locations where the model response exhibits more variations and thus are more of interest;
- There are existing computer model evaluations, but are potentially large in size and/or not produced with a careful design. It can be computationally inefficient to generate a new design, e.g., a static space-filling design, if one has an existing set of model evaluations because one could utilise the data available. However, it can be both numerically inefficient (e.g., the design formed by the existing data is poor) and computationally burdensome (e.g., the existing data is of large size) to use the whole existing model realisations for surrogate modelling. Thus, one can use active learning to choose training data adaptively from the existing model evaluations from a small design size while at the same time prevent from the numerical instabilities induced by poor designs;
- There is a system of coupled computer models. It has been shown in [Ming and Guillas](#page-21-0) [\(2021\)](#page-21-0) that active learning is essential to construct Gaussian process (GP) based surrogate models in a computationally efficient and effective manner. Static designs of global inputs can produce poor designs, and thus numerical issues, to sub-models of a computer system,

²²⁴ and can also waste computational resources over input regions of sub-models that are not

²²⁵ contributing to the global outputs (that correspond to the global input region of interest).

²²⁶ 4.2 Implementation

227 Assume that we have data $\mathcal{D}_n = \{\mathbf{X}_n, \mathbf{Y}_n\}$ that consists of input $\mathbf{X}_n \in \mathbb{R}^{n \times D}$ and the responding 228 high-dimensional computer model output $\mathbf{Y}_n \in \mathbb{R}^{n \times K}$. Then, a generic active learning procedure 229 that selects the next input position x_{n+1} to be evaluated by the computer model for refinement ²³⁰ of GP based non-instrusive ROM (abbreviated as GP-ROM in the remainder of the report) 231 introduced in Section [3](#page-4-0) is given in Algorithm [1.](#page-11-0) Once x_{n+1} is determined, one can then obtain 232 the augmented data $\mathcal{D}_{n+1} = {\mathbf{X}_{n+1}, \mathbf{Y}_{n+1}}$ by concatenating \mathbf{x}_{n+1} and its corresponding high-233 dimensional output y_{n+1} to \mathcal{D}_n and update GP-ROM $\{\mathcal{GP}_l\}$ by re-invoking Algorithm [1.](#page-11-0)

Input: (i) $\mathcal{D}_n = {\mathbf{X}_n, \mathbf{Y}_n}$; (ii) a candidate set C of input locations ${\mathbf{x}_i}_{i=1,\dots,M}$.

Output: The next input position x_{n+1} to be evaluated by the computer model.

- 1: Compute the low-dimensional output $\hat{\mathbf{Y}}_n \in \mathbb{R}^{n \times L}$ of \mathbf{Y}_n and the corresponding eigenvalues $\lambda_{l=1,...,L}$ using POD;
- 2: Construct GP-ROM $\{\mathcal{GP}_l\}$ using $\{\mathbf{X}_n, \hat{\mathbf{Y}}_n\};$
- 3: Calculate the criterion $I_l(\mathbf{x})$ at each input locations in C using \mathcal{GP}_l for all l;
- 4: Choose for the next input position x_{n+1} by solving

$$
\mathbf{x}_{n+1} = \operatorname*{argmax}_{\mathbf{x} \in C} \sum_{l=1}^{L} w_l I_l(\mathbf{x}) \quad \text{with} \quad w_l = \frac{\lambda_l}{\sum_{i=1}^{n} \lambda_i}
$$

We present two candidates for the criterion $I_l(\mathbf{x})$ based on the Active Learning MacKay (ALM) [\(MacKay, 1992\)](#page-21-11) and the Active Learning Cohn (ALC) [\(Cohn, 1996\)](#page-20-4) respectively for the selection of x_{n+1} . ALM aims to find the next input location that corresponds to the maximum predictive variance exhibited by the GP-ROM. Thus, $I_l(\mathbf{x})$ is defined by

$$
I_l(\mathbf{x}) = \widehat{\sigma}_l^2(\mathbf{x}),
$$

where $\hat{\sigma}_l^2(\mathbf{x})$ is the posterior predictive variance of \mathcal{GP}_l evaluated at **x**. However, ALM has a well-know issue that it selects excessive input locations around boundaries of the input region because of the lack of data beyond boundaries. To alleviate this issue, ALC aims to select the input position such that the integrated predictive variance of GP-ROM over the input region is minimised after augmenting x_{n+1} to X_n . Formally, $I_l(x)$ under ALC is defined by

$$
I_l(\mathbf{x}) = -\int_{\mathbf{x}^*\in\mathcal{X}} \widehat{\sigma}_l^2\left(\mathbf{x}^*|[\mathbf{X}_n^\top,\mathbf{x}^\top]^\top\right) \mathrm{d}\mathbf{x}^*.
$$

²³⁴ where $\hat{\sigma}_l^2(\mathbf{x}^*|[\mathbf{X}_n^\top,\mathbf{x}^\top]^\top)$ is interpreted as the posterior predictive variance of \mathcal{GP}_l evaluated at 235 \mathbf{x}^* given the input data \mathbf{X}_n being augmented by \mathbf{x}_n . It is worth noting that the computation of ²³⁶ $\hat{\sigma}_l^2(\mathbf{x}^*|[\mathbf{X}_n^\top,\mathbf{x}^\top]^\top)$ does not require evaluations of the associated computer model at \mathbf{x}_n because ²³⁷ the predictive variance of GP does not depend on the output data. In practice, the integral 238 involved in ALC can be approximated by the Monte Carlo integration over a reference set \mathcal{X} ²³⁹ (that can be the same as the candidate set C) generated by the LHD. To implement a full active ²⁴⁰ learning procedure, one often starts with a small data set that is generated by a static design, ²⁴¹ such as LHD, and then execute T iterations of Algorithm [1](#page-11-0) to enrich the initial data set with T ²⁴² additional realisations from the computer model.

²⁴³ 4.3 Active learning for the GP-ROM emulation of the 2-D anisotropic heat ²⁴⁴ transport model

²⁴⁵ In this section, we demonstrate how efficiency gains can be made using active learning for the ²⁴⁶ GP-ROM of the FE solver to the 2-D problem described in Section [3.1](#page-7-1)

²⁴⁷ 4.3.1 Experimental Setup

To initiate the active learning to build GP-ROM, N=20 initial training data points, whose input locations are generated via the LHD over $m \in [0, 12]$ and $\alpha \in [0, 3]$ with the corresponding 2-D output (that contains $K = 78961$ solution nodes) determined by running the FE solver (implemented in Firedrake [\(Rathgeber et al., 2016\)](#page-21-10)). We then iterate Algorithm [1](#page-11-0) for both ALM and ALC 80 times to augment additional 80 training data points to the initial data set. At each iteration of the active learning, we choose the number of components L (in Line [1](#page-11-1) of Algorithm [1\)](#page-11-0) to be retained from POD based on the following criteria:

$$
L = \underset{L^* \in \{1, ..., n\}}{\text{argmin}} \left| \frac{\sum_{i=1}^{L^*} \lambda_i}{\sum_{i=1}^{n} \lambda_i} - 0.9998 \right|,
$$

where $\lambda_1 > \lambda_2 > \cdots > \lambda_n$. To take into account the effects of initial data set on the active learning, we repeat both ALM- and ALC-based active learning 10 times. For the comparison between ALM and ALC, we generate 2500 testing data points over $m \in [0, 12]$ and $\alpha \in [0, 3]$ and compute the Normalised Root Mean Squared Error (NRMSE) at each active learning iteration by

$$
\text{NRMSE} = \frac{1}{2500} \sum_{i=1}^{2500} \frac{\sqrt{\frac{1}{K} (\widetilde{\mathbf{z}}_i - \mathbf{z}_i)(\widetilde{\mathbf{z}}_i - \mathbf{z}_i)^\top}}{\max(\mathbf{z}_i) - \min(\mathbf{z}_i)} \times 100\%,
$$

²⁴⁸ where $\tilde{\mathbf{z}}_i \in \mathbb{R}^{1 \times K}$ and $\mathbf{z}_i \in \mathbb{R}^{1 \times K}$ are 2-D FE solution fields generated by the GP-ROM and 249 Firedrake at the *i*-th testing input location, respectively.

²⁵⁰ In terms of implementation, we construct GP-ROM and compute corresponding ALM and ²⁵¹ ALC criterions at each iteration of the active learning using the laGP package in R.

²⁵² 4.3.2 Experimental Results

 Figure [6](#page-13-1) presents the NRMSEs of GP-ROMs built with ALM- and ALC-based active learning over 80 iterations, in comparison to those constructed with the static LHD at various design sizes. It can be observed that for design size less than 50, GP-ROMs trained using the active learning, regardless of ALM or ALC, provide higher accuracy than those trained using the static LHD. However, as the design sizes increases, the accuracy of GP-ROMs built by the active learning and LHD are comparable. This is because with a large design size, the input domain is

densely space-filled by the LHD and thus the NRMSE of the corresponding GP-ROM converges

to that of the GP-ROM trained with the active learning.

Figure 6: Comparison of NRMSEs of GP-ROM constructed using the ALM-based active learning, the ALC-based active learning, and the static LHD.

 We also observe from Figure [6](#page-13-1) that for design size larger than 60 GP-ROMs constructed by LHD perform better (in terms of overall lower NRMSE) than those built by ALM-based active learning. This observation can be explained by the fact that ALM-based active learning has the tendency to choose excessive input locations around boundaries of the input domain (see $_{265}$ Figure [7\(a\)\)](#page-13-2) and thus could fail to achieve a satisfactory design, in which input locations are preferred to be scattered within the input domain of interest (see Figure [7\(b\)\)](#page-13-3).

Figure 7: Designs produced by a random trial (out of 10 repeated trials) of ALM- and ALC-based active learning.

4.4 Discussion

 In this section, we introduce a simple and effective procedure to implement the active learning for GP-ROM construction. Although the active learning may eventually produce a space-filling design, it gives the computer model experimenters more controls over their computational re sources. One may criticise that active learning is not computationally efficient in the sense that it directs model runs sequentially and thus can be time-consuming in comparison to static one- shot designs in which model runs can be done in parallel. This statement is sensible when one posses sufficient computational power (for parallel computing) and active learning also produces a space-filling design. However, in real-world data these conditions may not be fulfilled. Our computational resources may not permit us to obtain model realisations that cover adequately the input region of interest (for an accurate surrogate model) and a space-filling design may not capture sufficiently (without tremendous computational efforts) the input regions where the model response exhibits abrupt changes, even if we have an advanced surrogate model (that is suitable for both stationary and non-stationary data). On the contrary, active learning has the ability to focus on input regions where the corresponding output surfaces show more variations, given that the underlying surrogate model provides a satisfactory uncertainty quantification (e.g., highlighting the regions with higher predictive standard deviations). A fact often forgotten in computer model experiments is that design and surrogate modelling are not two separate tasks. Good designs produce good surrogates with less numerical issues and more reliable uncertainty quantification, which in turn induces designs that better represent the functional behaviours of computer models under the consideration. These are the reasons why active learning could be preferred to static space-filling designs, which could cause the surrogate modelling challenging (e.g., a large number of realisations that are needed to capture well the computer model can cause the GP-ROM computationally prohibitive) and do not utilise the uncertainties quantified by surrogate models for design improvement.

 It is worth noting that active learning does not guarantee the locations of (possibly very small but important) input regions of a computer model that correspond to abrupt changes to the model responses. The design produced by the active learning depends on the quality of the underlying surrogate model, which in turn depends on the information contained in the training data (assuming that the surrogate represents the training data adequately and produces sensible uncertainty quantification). Therefore, whether active learning could find input regions that has very localised and important features depends on if the information of the regions exists in the training data. For this reason, it is vital to have a good initial design that incorporates such information for the active learning. However, in practice this can be difficult to achieve, particularly for high-dimensional cases, even we have some prior knowledge that such non-stationary features exist in the computer model, and as a consequence we may obtain a surrogate that completely ignores these regions with significant computational costs being wasted. To alleviate this issue, one could simply evaluate the computer model with a high- resolution design using the parallel computing. In this way, the local behaviours of a computer model can be captured within a reasonable amount of time. Nevertheless, it is not advisable to use all model evaluations for surrogate modelling, especially for GP-based surrogates because the large amount of data can cause GP surrogates computationally prohibitive and some evaluations (e.g., that form a flat response surface) are redundant for surrogate improvement. As a result, we propose the following hybrid static-active learning procedure to address the scenario in which we aim to construct efficiently (in terms of computation and time) a surrogate model that could mimic the underlying computer model with localised behaviours:

- 1. Generate a data set by evaluating the computer model over a dense space-filling design in parallel;
- 2. Choose a subset of the produced data set as the initial design for the active learning;
- 3. Implement the active learning that adaptively refines the design and the surrogate model, e.g., GP-ROM, by selecting data points from the data set produced in Step 1.

 There are several benefits provided by the above procedure. Firstly, the high-resolution design provides some guarantees that our data contain information of localised behaviours embedded in the underlying computer model. In addition, unlike typical active learning that evaluates models sequentially, active learning in Step 3 uses the data set already generated with a parallelisable strategy and thus could save a considerate amount of time (especially when computer models are very expensive to run). Furthermore, with active learning one is able to pick (potentially a small amount of) data points (from the generated data set) that contribute most to the surrogate quality, instead of naively pouring the whole data set into the surrogate construction (causing computational difficulties). Perhaps the most decisive and challenging step of the above procedure is Step 2 because, as discussed, one expects to incorporate some information of localised behaviours of a computer model into the initial design such that the resulting surrogate is less likely to overlook these features. How to integrate experts' knowledge about the localised features into the initial design is worth exploring in the future, but the procedure above indicates a potentially brutal but simply implementation for Step 2: choose multiple random subsets of the data set, then proceed to Step 3 for multiple surrogate constructions, and choose the surrogate that gives the best predictive accuracy (e.g., lowest overall predictive error against the generated data set). This implementation is computationally efficient because active learnings in Step 3 initiated by different random designs can be executed in parallel and do not involve computer model evaluations.

337 5 Future Directions

 We demonstrate in this report that a GP-ROM could be used to replace computationally expen- sive computer solvers for problems with high-dimensional output, in one of the building blocks of nuclear fusion modelling. However, dimension reduction techniques such as POD lose informa- tion when the original data are projected onto a lower dimensional space, and thus some extreme but important events could be masked in the low dimensional data, a scenario called masking effect. As a result, if the surrogate is built on the low dimensional data one may not be able to recover these outlying events using the constructed non-intrusive ROM. Therefore, other dimen- sion reduction methods that may be more resistant to the masking effect could be examined. In addition, although GP-ROM requires no domain knowledge and access to the source code of original problems, it ignores the physics implied by the underlying problem and thus may be inaccurate comparing to the its intrusive counter-party. Therefore, it would be worth exploring the trade-off between the speed and accuracy of intrusive and non-intrusive MOR, especially in context of UQ. It would also be interesting to find a middle ground where one could exploit the benefits (e.g., accuracy, speed and uncertainty) of both intrusive and non-intrusive ROM,

 producing a physics-informed non-intrusive ROM. Some relevant literature on physics-informed machine learning (say using a boundary condition or other approaches) include [Vernon et al.](#page-22-3)

[\(2019\)](#page-22-3); [Kashinath et al.](#page-21-12) [\(2021\)](#page-21-12); [Watson-Parris](#page-22-4) [\(2021\)](#page-22-4).

 Recommendation: Investigate how to apply physics-informed GP-ROM in key nuclear fusion models. Examine how to build new types of GP-ROM for the case of particle-based models (PIC) whose outputs need to be understood as a continuum.

5.1 Deep GP for Non-intrusive ROM

 In this report we explored how to construct GP-ROM using active learning. Active learning is particularly useful when the underlying computer model exhibits non-stationary features as it has the ability to produce a non-uniform design that appreciates the non-stationarity. However, the success of the active learning relies on the quality of uncertainty quantified by the surrogate model. Since conventional GP surrogates assume stationarity, more advanced non-stationary GP models, such as deep Gaussian processes [\(Damianou and Lawrence, 2013\)](#page-20-5), would be good candidates for non-intrusive ROM of fusion models that exhibits non-stationarity. Deep Gaussian processes (DGPs) are feed-forward compositions of conventional stationary GPs with flexible model expressiveness, particularly for non-stationary data. However, training and prediction of DGP based emulators are challenging due to the non-linearity induced by the kernel functions involved in GPs. Various inference methods thus are introduced to tackle this issue. [V](#page-22-5)ariational inferences, such as Doubly Stochastic Variational Inference (DSVI) [\(Salimbeni and](#page-22-5) [Deisenroth, 2017\)](#page-22-5), is computationally thrifty but is not accurate because simplified assumptions over the latent variables in DGP hierarchy are assumed. On the contrary, the fully-Bayesian approach introduced by [Sauer et al.](#page-22-6) [\(2020\)](#page-22-6) gives a comprehensive uncertainty quantification of DGPs, but at the expense of computation. The stochastic imputation approach recently proposed by [Ming et al.](#page-21-13) [\(2021\)](#page-21-13) is a DGP inference method that enjoys both computational speed and the predictive accuracy, and could be a competitive and potential candidate for DGP emulations of non-stationary fusion models. It is implemented in the dgpsi package^{[1](#page-3-0)}.

 Figure [8](#page-17-0) showcases the ALM-based active learning using a two-layered DGP surrogate (i.e., composition of two stationary GPs) trained with the stochastic imputation in comparison to that using a stationary GP. It can be observed that DGP surrogate outperforms the GP surrogate in both mean predictions and uncertainty quantification. In addition, with DGP the active learning could produce a non-uniform design that appreciates the non-stationarity of the underlying data. While the active learning essentially produces a quite space-filling design under GP, it assigns three time more number of design points to the rough (and more interesting) regime over [0, 0.5] than the flat regime over $(0.5, 1]$ under DGP. Although this is a simple 1-D example, it gives motivations why DGP surrogate should be seriously considered if the reduced-order output of a fusion model is non-stationary and the active learning is employed.

 Recommendation: Investigate how to reduce dimensionality of outputs for key nuclear fusion models whose behaviour may present sharp transitions or various regimes, such as turbulence models. The key question is then how to understand and represent the continuum of outputs

https://github.com/mingdeyu/DGP

Figure 8: ALM-based active learning using GP and DGP emulations. Solid line represents the underlying true function; Dashed line is the mean prediction; Shaded area represents 95% predictive interval; Dots (6 in total) are initial training points and triangles (14 in total) are training points enriched by the active learning procedure using GP and DGP surrogates. The vertical dashed line indicates a visual split of the underlying true function into a rough regime over $[0, 0.5]$ and a flat regime over $(0.5, 1]$.

³⁹¹ features across regimes. Indeed these features shown in [3](#page-4-0) can vary across regimes and must

³⁹² be made consistent by some form of joint augmentation possibly at a small cost but with large

³⁹³ benefits for emulation.

394 5.2 Active subspace for efficient dimension reduction of inputs

 The efficiency of reducing dimensions in the inputs was demonstrated in [Liu and Guillas](#page-21-3) [\(2017\)](#page-21-3). Gains of orders of magnitude can be achieved. For instance, the application to a surface of inputs (a mesh of 3200 elements) enabled a reduction from dimension 3200 to 5 with fast and accurate emulation. Only about 100 simulations were needed to come up with 5 key dimensions as a recombination of the original 3200 dimensions. A summary of the method is presented ⁴⁰⁰ below. It is implemented in the Alan Turing Institute Package *Multi-Output Gaussian Process* 401 Emulator (MOGP)^{[2](#page-3-0)}. The context is:

- Simulator input X (high dimension \mathbb{R}^p) and output $Y = f(X)$ (one dimension \mathbb{R}^1)
- 403 GP emulation: fit an GP and predict $f(x_{new})$ using a sample of simulations $f(X_1),..., f(X_n)$

• Find a reduced space (known as sufficient dimension reduction SDR) $R(X) \in \mathbb{R}^d$, $d < p$, ⁴⁰⁵ such that there is (nearly) no loss of information in predicting Y by providing $R(X)$ instead 406 of X

-
- To achieve SDR, employ the gradient-based Kernel Dimension Reduction (gKDR) approach [\(Fukumizu and Leng, 2014\)](#page-20-6):

$$
R(X) = B^T X, \quad B^T B = I_d, \quad d < p.
$$

²https://github.com/alan-turing-institute/mogp-emulator

- 407 Estimate B from simulation samples $(X_1, Y_1), ..., (X_n, Y_n)$. Note that no strong assumption ⁴⁰⁸ are made on the variables (type, distribution, dimension).
- ⁴⁰⁹ The specific technical steps in gKDR involve two Reproducing kernel Hilbert spaces (RKHS):
- 410 Prepare kernels $k_{\mathcal{X}}$ and $k_{\mathcal{Y}}$, with the associated (RKHS) $\mathcal{H}_{\mathcal{X}}$ and $\mathcal{H}_{\mathcal{Y}}$

⁴¹¹ • The quantities of interest are the gradients $\frac{\partial E[g(Y)|X]}{\partial X}$ for any $g \in \mathcal{H}_{\mathcal{Y}}$ as their evaluation ⁴¹² is the ingredient for the identification of the reduced subspace, by looking at the most ⁴¹³ influential directions.

• Estimate (see [Fukumizu and Leng](#page-20-6) [\(2014\)](#page-20-6) for details)

$$
\hat{M}_n = \frac{1}{n} \sum_{i=1}^n \nabla \mathbf{k}_X (X_i)^T (G_X + n\epsilon_n I)^{-1} G_Y (G_X + n\epsilon_n I)^{-1} \nabla \mathbf{k}_X (X_i)
$$

where G_X and G_Y are the Gram matrices $(k_X(X_i, X_j))$ and $(k_Y(Y_i, Y_j))$, and $\nabla \mathbf{k}_X(x) =$ 415 $(\partial k_{\mathcal{X}}(X_1,x)/\partial x,...,\partial k_{\mathcal{X}}(X_n,x)/\partial x)^T \in \mathbb{R}^{n \times m}$ for any $x \in \mathbb{R}^m$.

• Eigen-decompose \hat{M}_n into $\hat{M}_n = \hat{Q} \hat{\Lambda} \hat{Q}^T$ and partition

$$
\hat{\Lambda} = \begin{bmatrix} \hat{\Lambda}_1 & \\ & \hat{\Lambda}_2 \end{bmatrix}, \quad \hat{Q} = [\hat{B} \ \hat{C}],
$$

⁴¹⁶ where $\hat{\Lambda}_1 = diag(\hat{\lambda}_1, ..., \hat{\lambda}_d)$ consisting of the first d largest eigenvalues, to ultimately ⁴¹⁷ provide the dimension reduction.

⁴¹⁸ The emulation with dimension reduction can be carried out and its loss quantified [\(Liu and](#page-21-3) ⁴¹⁹ [Guillas, 2017\)](#page-21-3):

- $f(X) \approx \hat{f}(\hat{B}^T X)$
-

$$
\left\|f - \hat{f}\right\|_{L_2} =
$$
\n
$$
O_p\left(\frac{4}{\lambda_d - \lambda_{d+1}} n^{-\min\{\frac{1}{3}, \frac{2\beta+1}{4\beta+4}\}} \left(\sum_{i=1}^d c_i \hat{\lambda}_i^2\right)^{\frac{1}{2}} + \left(\sum_{i=d+1}^m c_i \hat{\lambda}_i^2\right)^{\frac{1}{2}}\right)
$$

- ⁴²³ ► Build emulator $\tilde{f} \approx \hat{f}$ on low dimensional space $\hat{B}^T X$
	- Approximation procedure:

$$
f(X) \approx \hat{f}(\hat{B}^T X) \approx \tilde{f}(\hat{B}^T X)
$$

 424 The choice of remained dimension d and hyperparameters is performance based (e.g. in the ⁴²⁵ [q](#page-21-3)uality of the predictions in a leave-one-out strategy) and can result in very large gains [\(Liu](#page-21-3) ⁴²⁶ [and Guillas, 2017\)](#page-21-3).

⁴²⁷ Recommendation: Investigate how to reduce dimensionality of inputs of key nuclear fusion ⁴²⁸ models such as the magnetic field modelled input of the anisotropic heat transfer model.

⁴²⁹ 5.3 Linked GP for Non-intrusive ROM

 Since fusion models are often multi-disciplinary and multi-physics, the recent advances on linked Gaussian process surrogates [\(Ming and Guillas, 2021\)](#page-21-0) must be considered. The linked GP is $_{432}$ $_{432}$ $_{432}$ implemented in the dgpsi package³. As an illustration, consider a toy system that consists of two feed-forward connected computer models shown in Figure [9.](#page-19-0) By directly applying con-434 ventional GP emulation, one fails to capture the local feature (over $[-1, 1]$) of the underlying system with ten system runs, see Figure [10\(d\).](#page-19-1) However, if the linked GP is employed to the system by constructing GP surrogates on sub-models individually with active learning (see Fig- ure [10\(a\)](#page-19-2) and [10\(b\)\)](#page-19-3), one could capture the local feature of the overall system sufficiently (see 438 Figure [10\(c\)\)](#page-19-4). This is because the local feature of the entire system over $[-1, 1]$ is created by the composition of simpler individual sub-models, and thus constructing system surrogate on the basis of elementary emulators could achieve better emulation performance. Besides, using the active learning one could optimise the designs for individual sub-models, and thus obtain better corresponding GP surrogates, which in turn produce system surrogate with higher accuracy.

Figure 9: An illustrative example of a system of two computer models f_1 and f_2 . Note this is only for illustration. Linked GP in [Ming and Guillas](#page-21-0) [\(2021\)](#page-21-0) can work on any feed-forward computer systems.

Figure 10: Linked GP (in (c)) and linked GP (in (d)) emulators of a feed-forward system $(f_2 \circ f_1)$ of two computer models f_1 and f_2 connected as shown in Figure [9.](#page-19-0) The filled circles in (d) are training points for conventional GP, while in (a) and (b) the filled circles represent the initial design of the active learning to build GP surrogates of individual computer models f_1 and f_2 for linked GP in (c); the filled triangles in (a) and (b) are training points created by the active learning; the solid line is the underlying true function; the dashed line is the mean prediction; the shaded area represents 95% prediction interval.

 The toy example motivates further explorations of linked GP in constructing non-intrusive ROM for fusion systems by linking non-intrusive ROM of individual sub-models. For example, to construct the ROM of the two-layered system in Figure [9,](#page-19-0) one could first build GP-based 446 non-intrusive ROM (as demonstrated above) for all individual sub-models $(f_1 \text{ and } f_2)$ and then

³https://github.com/mingdeyu/DGP

447 construct the non-intrusive ROM of the whole system by linking the non-intrusive ROM of f_1 to that of f_2 through the reduced space w analytically. One key benefit of this approach for system-wise reduced order modelling is that one only needs to do dimensionality reduction to the outputs of sub-models. Whereas, to build intrusive ROM, one has to make extra challenging 451 efforts to reformulate the original high-fidelity model f_2 under reduced input w and output y.

 Implementing the active learning for linked GP surrogates for systems of computer models with high-dimensional outputs is also challenging. In comparison to the static design (in which the training input data of one sub-model matches the training output data produced by the feeding sub-models), the active learning (e.g., the adaptive design introduced in [Ming and Guillas](#page-21-0) [\(2021\)](#page-21-0)) could lose the input/output data matching, and thus further explorations are required to examine how to conduct dimension reductions for the internal sub-model input/output so that all information contained in the training data of linked sub-models are utilised.

 Recommendation: Investigate how to jointly reduce dimensionality of outputs that are inputs of key nuclear fusion models, such as the heat from the anisotropic heat transfer model propa- gated to the wall heat transfer model. Emulation with high-dimensional outputs (GP-ROM) of the first simluator and active subspace for dimension reduction of the subsequent inputs of the following simulator should be used in synergy. To establish such a combined strategy will require examining carefully how to weigh variations in outputs of the first model and the influence of inputs for the second. The sampling approach of [4](#page-9-0) needs to be tailored to this new context as well. It is necessary to carry out such combination of methods and strategies due to the very high dimensions, heavy data transfers, and extremely costly simulations.

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