Abstract. To date, the analysis of high-dimensional, computationally expensive engineering models remains a difficult challenge in risk and reliability engineering. We use a combination of dimensionality reduction and surrogate modelling termed partial least squares-driven polynomial chaos expansion (PLS-PCE) to render such problems feasible. Standalone surrogate models typically perform poorly for reliability analysis. Therefore, in a previous work, we have used PLS-PCEs to reconstruct the intermediate densities of a sequential importance sampling approach to reliability analysis. Here, we extend this approach with an active learning procedure that allows for improved error control at each importance sampling level. To this end, we formulate an estimate of the combined estimation error for both the subspace identified in the dimension reduction step and surrogate model constructed therein. With this, it is possible to adapt the design of experiments so as to optimally learn the subspace representation and the surrogate model constructed therein. The approach is gradient-free and thus can be directly applied to black box-type models. We demonstrate the performance of this approach with a series of low- (2 dimensions) to high- (869 dimensions) dimensional example problems featuring a number of well-known caveats for reliability methods besides high dimensions and expensive computational models: strongly nonlinear limit-state functions, multiple relevant failure regions, and small probabilities of failure.

Key words. Reliability Analysis, Rare event simulation, PLS-PCE, Dimensionality reduction, Active learning, Sequential importance sampling

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1. Introduction and previous work. An important challenge in the design, analysis and maintenance of engineering systems is the management of the associated uncertainties. It is common practice to analyse engineering systems by employing computational models that aim at representing the physical processes relevant to the system in consideration. These computational models take the form of an input-output mapping. Therein, uncertainty is represented by equipping the model input with an appropriate probabilistic model. Undesirable system responses are defined through a limit-state function (LSF). Reliability analysis is concerned with quantifying the probability of failure, which can be expressed as a d-fold integral of the input probability mass over the failure domain defined by non-positive values of the LSF, where d is the number of uncertain model inputs (see Section 2). In engineering, target failure probabilities are typically small; hence, reliability analysis requires the estimation of rare event probabilities. Reliability analysis approaches can be categorized into approximation (e.g. the first- and second-order reliability methods FORM and SORM [64, 26, 17]) and simulation methods. If the LSF is only weakly nonlinear and the input dimension of the model is moderate, FORM and SORM perform well even for small failure probabilities. The simplest simulation method is the Monte Carlo method [53]. The Monte Carlo method performs well independent of the problem input dimension, however its performance deteriorates as the failure probability decreases if the computational budget is fixed. Various techniques such as importance sampling (IS) [13, 23, 2] and line-sampling [29, 38] have been proposed to mitigate this dependence on the magnitude of the failure probability. More recently, sequential MC methods such as subset simulation [3] and IS-based sequential methods [40, 41, 81, 60, 66, 59] have been used successfully to efficiently solve high-dimensional reliability problems with small failure probabilities. If the computational model is expensive and a hierarchy of increasingly coarse and cheap models is accessible, multilevel and multi-fidelity [62] MC methods can help alleviate computational cost by performing most model evaluations on the cheaper models (e.g., a discretized differential equation with coarser resolution). In [77], multilevel MC is combined with subset simulation and recently [80] have introduced multilevel sequential IS based on the sequential IS approach in [60]. All of the above-mentioned approaches are designed to work with the probabilistic computational model directly. However, often this model encompasses a numerical solver for (sets of) partial differential equations such that a model evaluation is computationally expensive.

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This has increasingly lead researchers to turn towards surrogate model-based reliability methods. Such methods attempt to approximate the expensive computational model with a cheap surrogate model, whose coefficients are identified based on a set of original model evaluations: the design of experiments (DoE). [24] uses a polynomial response surface method for performing reliability analysis as early as 1989. [27] proposes an improved version of the response surface method. Since then, a variety of surrogate modelling techniques has been applied in the context of reliability analysis such as artificial neural networks [56, 33, 69], support vector machines [32, 12, 11], Gaussian process regression-based models [21, 20] and projection to polynomial bases including polynomial chaos expansions (PCE) [46, 44, 43, 71] and low-rank tensor approximations [37].

Static, global surrogate models suffer from a decrease in accuracy in the tails of the model response distribution such that they are of limited use for reliability analysis. In this context, static refers to surrogate models that are constructed based on a fixed DoE and global refers to surrogate models that are trained and evaluated on the entire input space (as opposed to locally con- and re-fined models). Thus, one can distinguish two strategies to overcome this limitation:

- **Locality**: Surrogate models are coupled with sequential sampling techniques which serve to focus the DoE and accuracy in the relevant regions around the failure hypersurface [55, 12, 11, 6, 57].

- **Adaptivity** (in the DoE): The DoE is augmented with points that are most informative with respect to the failure probability estimate according to an ‘in-fill criterion’. The refined surrogate model is then used to estimate the probability of failure with a sampling method and a large number of cheap samples. Such procedures are summarized under the terms active learning (AL) or optimal experimental design. AL in combination with crude Monte Carlo have been applied in reliability-based optimization and reliability analysis in [21, 52, 8, 63]. [69] investigates the performance of splines and neural networks in combination with directional sampling and IS and [20, 14] combine Gaussian process models with IS. [68] proposes a crude Monte Carlo procedure relying on a Gaussian process surrogate model with PCE-based mean trend (PCE-Kriging) along with a novel termination criterion for the AL.

Often, both AL and sequential sampling techniques are combined using various combinations of in-fill criteria and sequential sampling techniques such as adaptive IS [5] and subset simulation [12, 31, 6, 11]. [51] turns away from surrogate models that naturally provide a measure of prediction uncertainty such as Gaussian processes or support vector machines and demonstrate how an AL algorithm can be realized with PCE using a bootstrap estimator of the PCE prediction uncertainty.

In spite of a plethora of existing approaches to surrogate-assisted reliability analysis, the literature on high-dimensional problems ($d \geq 100$) in this context is scarce. [35, 45] propose to perform reliability analysis with a static, global Kriging model constructed in a low-dimensional linear subspace of the original model input space, which is identified by the active subspaces method [16] and autoencoders, respectively. Both [35, 45] apply their methods to moderate-dimensional problems with up to $d = 20$ and $d = 40$ input variables, respectively. [54] uses sliced inverse regression to identify a linear low-dimensional subspace and construct a static, global PCE in this space based on which they perform reliability analysis directly. [87] develops these ideas further by combining the active subspace-Kriging model with an AL approach and applies this combination to a high-dimensional analytical problem of $d = 300$ that possesses a perfectly linear low-dimensional structure.

In this work, we propose an importance sampler based on a dimensionality-reducing surrogate model termed partial least squares-driven PCE (PLS-PCE) [58] to efficiently solve high-dimensional reliability problems with underlying computationally expensive, nonlinear models and small target probabilities ($O(10^{-9})$). Similar to sliced inverse regression and active subspaces, PLS-PCE achieves dimensionality reduction by identifying a low-dimensional linear subspace of the original input space. Our method is based on [57] but introduces AL to refine the PLS-PCE approximation in each sequence of the IS procedure. In [57], PLS-PCE models are reconstructed in each level of a sequential importance sampling (SIS) scheme that is used to gradually shift the importance density towards the optimal importance density. In this work, we augment this approach with two novel contributions to rare event simulation of computationally expensive, potentially (but not necessarily) high-dimensional and nonlinear models:

1. We demonstrate how to perform active learning with PCE models by deriving an in-fill criterion from large-sample properties of the PCE coefficient estimates.

2. We show how to incorporate the sequential IS framework into the AL process for enhancing reliability and improving the PCE efficacy.
2. Reliability analysis. Consider a system represented by the computational model \( Y : \mathbb{D}_X \rightarrow \mathbb{R} \) with \( d \)-dimensional continuous random input vector \( X : \Omega \rightarrow \mathbb{D}_X \subseteq \mathbb{R}^d \), where \( \Omega \) is the sample space of \( X \) and by \( F_X(x) \), we denote its joint cumulative distribution function (CDF). \( Y \) maps to the system response \( Y = Y(x) \) with the model input \( x \in \mathbb{D}_X \). Based on the response \( Y \), unacceptable system states are defined by means of the limit-state function (LSF) \( g(Y) \). Defining \( g(x) = \tilde{g} \circ Y(x) \) and introducing the convention 

\[
g(x) = \begin{cases} 
\leq 0, \text{Failure} \\
> 0, \text{Safety}, 
\end{cases}
\]

the failure event of the system is defined as \( F = \{ x \in \mathbb{D}_X : g(x) \leq 0 \} \). The probability of failure is given by [18]

\[
p = \mathbb{P}(F) = \int_{\mathbb{D}_X} \mathbb{I}[g(x) \leq 0] f_X(x) \, dx = \mathbb{E}_{f_X}[\mathbb{I}(g(X) \leq 0)],
\]

where \( f_X(x) = \frac{\partial^d F}{\partial x_1 \ldots \partial x_d} \big|_x \) is the joint probability density function (PDF) of \( X \) and the indicator function \( \mathbb{I}[] \) equals 1 if the condition in the argument is true and 0 otherwise. Without loss of generality, one may formulate an equivalent reliability problem with respect to the standard-normal probability space using the random vector \( U : \Omega \rightarrow \mathbb{R}^d \). Given an isoprobabilistic transformation \( T : \mathbb{D}_X \rightarrow \mathbb{R}^d \), such that \( U = T(X) \), see, e.g., [28, 47], and defining \( G(U) = g(T^{-1}(U)) \), one can write (2.1) as

\[
p = \int_{\mathbb{R}^d} \mathbb{I}[G(u) \leq 0] \varphi_d(u) \, du = \mathbb{E}_{\varphi_d}[\mathbb{I}(G(U) \leq 0)],
\]

where \( \varphi_d \) denotes the \( d \)-dimensional independent standard-normal PDF. The crude Monte Carlo estimate of (2.2) is

\[
\hat{p}_{MC} = \frac{1}{n} \sum_{k=1}^{n} \mathbb{I}[G(u^k) \leq 0], \quad u^k \overset{i.i.d.}{\sim} \varphi_d,
\]

where \( u^k \overset{i.i.d.}{\sim} \varphi_d \) means that \( \{u^k\}_{k=1}^{n} \) are \( n \) samples that are independent and identically distributed according to \( \varphi_d \). This estimate is unbiased and has coefficient of variation (CoV)

\[
\delta_{MC} = \sqrt{\frac{1-p}{np}}.
\]

The number of samples required to compute \( \hat{p}_{MC} \) at a prescribed CoV \( \delta_0 \) reads

\[
n_0 = \frac{1-p}{\delta_0^2} \approx \frac{1}{\delta_0^2 p}.
\]

Therefore, crude Monte Carlo is inefficient for estimating rare event probabilities as, by definition, \( p \ll 1 \) and thus \( n_0 \) becomes large.
3. Sequential importance sampling for rare event estimation. Variance reduction techniques can be used to reduce the CoV of the probability estimate at a fixed budget of samples compared to crude Monte Carlo. One of the most commonly used variance reduction methods is the IS method. Let $h$ be a density, such that $h(u) > 0$ whenever $G(u) \leq 0$. Then, one can rewrite (2.2)

\begin{equation}
    (3.1) \quad p = \int_{\mathbb{R}^d} I(G(u) \leq 0) \frac{\varphi_d(u)}{h(u)} h(u) \, du = E_h[I(G(U) \leq 0)\omega(U)],
\end{equation}

which leads to the (unbiased) importance sampling estimator

\begin{equation}
    (3.2) \quad \hat{p}_{\text{IS}} = \frac{1}{n} \sum_{k=1}^{n} I(G(u^k) \leq 0)\omega(u^k), \quad u^k \sim h.
\end{equation}

The efficiency of IS depends intimately on the choice of the IS density $h$ and numerous techniques to construct it have been put forward. There exists an optimal importance density $h^*$ in the sense that it leads to

\begin{equation}
    (3.3) \quad h^*(u) = \frac{1}{p} I(G(u) \leq 0)\varphi_d(u).
\end{equation}

While this result is not immediately useful in estimating $p$ as it requires knowledge of $p$, it can be used to guide the selection of a suitable IS function $h$.

The SIS method proposed in [60] selects the IS density sequentially starting from a known distribution $h_0$ that is easy to sample from. It relies on a sequence of distributions $\{h_i(u)\}_{i=0}^{M}$,

\begin{equation}
    (3.4) \quad h_i(u) = \frac{\eta_i(u)}{p_i}, \quad i = 1, \ldots, M,
\end{equation}

where $\{\eta_i(u)\}_{i=0}^{M}$ are non-normalized versions of $\{h_i(u)\}_{i=0}^{M}$ and $\{p_i\}_{i=0}^{M}$ are the respective normalizing constants. The goal is to arrive at $h_M$, which is sufficiently close to $h^*$ based on some criterion, and perform importance sampling with $h_M$. To this end, it is necessary to estimate $p_M$ and obtain samples from $h_M$. Based on the likelihood ratio of two succeeding non-normalized distributions $\omega_i(u) = \eta_i(u) / \eta_{i-1}(u)$, we have

\begin{equation}
    (3.5) \quad s_i = \frac{p_i}{p_{i-1}} = \int_{\mathbb{R}^d} \frac{\eta_i(u)}{\eta_{i-1}(u)} h_{i-1}(u) \, du = E_{h_{i-1}}[\omega_i(u)].
\end{equation}

Therefore, an estimate of $p_M$ is given by

\begin{equation}
    (3.6) \quad \hat{p}_M = \prod_{i=1}^{M} s_i \text{ with } \hat{s}_i = \frac{1}{n} \sum_{k=1}^{n} \omega_i(u^k), \quad u^k \sim h_{i-1}.
\end{equation}

Samples from $h_i$ can be obtained using Markov Chain Monte Carlo (MCMC) methods given samples from $h_{i-1}$. More precisely, [60] proposes a resample-move scheme in which Markov chain seeds are obtained as samples from $h_{i-1}$ that are then reweighted (resampled with weights) according to $\omega_i(u)$. In this way, the seed samples are already approximately distributed according to the stationary distribution of the Markov chain $h_i$ and long burn-in periods can be avoided. We adopt an adaptive conditional MCMC sampler (aCS) to perform the move step due to its robust performance in high-dimensional settings. Details can be found in [60].

The $h_i$ are chosen as smooth approximations of $h^*$ using the standard-normal CDF $\Phi(\cdot)$ (compare Fig. 1):

\begin{equation}
    (3.7) \quad h_i(u) = \frac{1}{p_i} \Phi \left( -\frac{G(u)}{\sigma_i} \right) \varphi_d(u) = \frac{1}{p_i} \eta_i(u),
\end{equation}
where \( p_i = \mathbb{E}_{\phi_i} [\Phi(-G(U)/\sigma_i)] \) is a normalizing constant and \( \sigma_i \) is the smoothing parameter. Prescribing \( \sigma_0 > \sigma_1 > \cdots > \sigma_M \) ensures that the sequence \( \{h_i(u)\}_{i=0}^M \) approaches \( h^* \). In each level, to avoid degeneration of the weights \( \omega_i \) (meaning \( \omega_i \) assuming values close to 0 at all current samples), \( h_{i-1}(u) \) and \( h_i(u) \) cannot be too different in the sense that they share no support regions on which both have considerable probability mass. This is avoided by prescribing an upper bound for the estimated coefficient of variation of the weights \( \delta_{w,i} = \hat{\text{COV}}[\omega_i(U)] \), which provides a criterion for determining \( \sigma_i \):

\[
\sigma_i = \arg \min_{\sigma \in [0, \sigma_{i-1}]} \left( \delta_{\omega,i}(\sigma) - \delta_{\text{target}} \right)^2.
\]

[60] recommends \( \delta_{\text{target}} = 1.5 \). The algorithm terminates when \( h_i \) is close enough to \( h^* \) in the sense that

\[
\hat{\text{COV}} \left[ \frac{h^*(U)}{h_i(U)} \right] = \mathbb{E}_{\phi_i} \left[ \frac{\varphi_d(U) I(G(U) \leq 0)}{\Phi(-G(U)/\sigma_i)} \right] = \mathbb{E}_{\phi_i} \left[ \frac{I(G(U) \leq 0)}{\Phi(-G(u)/\sigma_i)} \right] \leq \delta_{\text{target}}.
\]

The final estimate of \( \mathbb{P}(F) \) reads

\[
\hat{\mathbb{P}}_{\text{SIS}} = \hat{\mathbb{P}}_M \hat{\mathbb{E}}_{\phi_d} \left[ \frac{I(G(U) \leq 0)}{\eta_M(U)} \right] = \left( \prod_{i=0}^M \hat{s}_i \right) \frac{1}{n} \sum_{k=1}^n \frac{I(G(u^k) \leq 0)}{\Phi(-G(u^k)/\sigma_M)}, \quad u^k \overset{i.i.d.}{\sim} h_M.
\]

Algorithm 3.1 summarizes the complete SIS-\( \alpha \text{CS} \) procedure.

4. Partial least squares-based polynomial chaos expansions.

4.1. Polynomial Chaos Expansions. Polynomial chaos expansions (PCEs) are a tool for forward modelling the relationship between an input \( X \) and an output \( Y = \mathcal{Y}(X) \). With \( \mathcal{H} \), we denote the Hilbert space of functions that are square-integrable with respect to \( f_X \), i.e., \( \{v : \mathbb{E}_{f_X} [v(X)^2] < \infty \} \). \( \mathcal{H} \) admits an inner product of two functions \( v, w \in \mathcal{H} \):

\[
\langle v, w \rangle_{\mathcal{H}} = \mathbb{E}_{f_X(x)} [v(X)w(X)] = \int_{\mathbb{R}^d} v(x)w(x)f_X(x)dx.
\]

Let \( \{\varphi_j(X), j \in \mathbb{N}\} \) be a complete and orthonormal basis of \( \mathcal{H} \) so that \( \langle v, \varphi_j \rangle_{\mathcal{H}} = \delta_{j\ell} \) and let \( \mathcal{Y} \in \mathcal{H} \). Then,

\[
\mathcal{Y}(X) = \sum_{j=0}^{\infty} b_j \varphi_j(X),
\]

where the coefficients \( b_j \) are defined by projecting \( \mathcal{Y} \) on the basis:

\[
b_j = \langle \mathcal{Y}, \varphi_j \rangle_{\mathcal{H}}, \quad j \in \mathbb{N}.
\]
\textbf{Algorithm 3.1 SIS-aCS [60]}

1: \textbf{Input} LSF \( G(u) \), target CoV \( \delta_{\text{target}} \), samples per level \( n \), input dimension \( d \), burn-in period \( b \), max. iterations \( i_{\text{max}} \)

2:

3: Set \( i = 0 \), \( \sigma_0 = \infty \), \( h_0(u) = \varphi_d(u) \)

4: Sample \( U_0 = \{ u^k, k = 1, \ldots, n \} \in \mathbb{R}^{n \times d} \)

5: Compute \( G_0 = G(U_0) \in \mathbb{R}^{n \times 1} \)

6: for \( i \leftarrow 1, i_{\text{max}} \) do

7: \( i \leftarrow i + 1 \)

8: Compute \( \sigma_i \) according to (3.8)

9: Compute weights \( \omega_i = \{ \Phi[-G_{i-1}/\sigma_i]/\Phi[-G_{i-1}/\sigma_{i-1}], k = 1, \ldots, n \} \in \mathbb{R}^{n \times 1} \)

10: Compute \( \hat{s}_i \) according to (3.6),

11: \( U_{i-1} \leftarrow \) draw weighted resample from \( U_{i-1} \) with weights \( \omega_i \) \quad \triangleright \text{sample with replacement}

12: \( (U_i, G_i) = \text{MCMC-aCS}(U_{i-1}, G_{i-1}, b) \) \quad \triangleright \text{Details on MCMC-aCS in [60]}

13: if (3.9) then

14: \quad break

15: Set \( M \leftarrow i \)

16: Estimate failure probability \( \hat{p}_{\text{SIS}} = \left( \prod_{i=1}^{M} \hat{s}_i \right) \frac{1}{n} \sum_{k=1}^{n} \frac{1(G_k \leq 0)}{\Phi[-G_k/\sigma_M]} \) \quad \triangleright (3.10)

17: return \( U_M, G_M, \hat{p}_{\text{SIS}} \).

Since \( \mathcal{Y} \in \mathcal{H} \), the truncation

\begin{equation}
\hat{\mathcal{Y}}_n(X) = \sum_{j=0}^{n} b_j \psi_j(X)
\end{equation}

asymptotically converges to \( \mathcal{Y} \) as \( n \to \infty \) in the mean-square sense. [85] demonstrates how to construct complete orthonormal bases of \( \mathcal{H} \) as polynomial families for various standard input distribution types. In particular, if \( F_X(x) = \Phi_d(x) \), where \( \Phi_d \) denotes the \( d \)-variate independent standard-normal CDF, the tensorized, normalized probabilist’s Hermite polynomials

\begin{equation}
\Psi_k(U) = \prod_{i=1}^{d} \psi_{k_i}(U_j)
\end{equation}

form a complete orthonormal basis of \( \mathcal{H} \). \( \{ \psi_j(U), j \in \mathbb{N} \} \) are the univariate, normalized (probabilist’s) Hermite polynomials and \( k = (k_1, \ldots, k_d) \in \mathbb{N}^d \). By means of the isoprobabilistic transformation \( T : X \to U \) introduced in the previous section, we define PCEs in standard-normal space for the remainder of the paper. The PCE of maximum total order \( p \) reads

\begin{equation}
\hat{\mathcal{Y}}_p(U) = \sum_{|k| \leq p} b_k \Psi_k(U).
\end{equation}

The total number of basis functions in the PCE, \( P \), depends on the input dimension \( d \) and the maximum total polynomial order \( p \):

\begin{equation}
P = \binom{d + p}{p}.
\end{equation}

The projection in (4.3) can be transformed into an equivalent ordinary least squares (OLS) problem [7]. PCEs become computationally intractable if \( d \) is large, i.e., they cannot be used for problems with high-dimensional input due to the sheer number of basis functions and corresponding coefficients. In particular, the computation is rendered infeasible by the necessary number of operations to compute the set of \( P \) multi-indices and the necessary number of model evaluations to obtain meaningful estimates of the coefficients. Solution strategies to overcome these limitations (at least partially) include a hyperbolic truncation of the

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index set (this means to replace the condition on the $\ell_1$-norm in (4.6), $|k| \leq p$, with one on a general $\ell_q$-norm of $|k|_q = (\sum_{i=1}^d p_i^q)^{1/q} \leq p$ with $q < 1$) or enforcing a maximum interaction order (i.e., a maximum number of non-zero entries in $k$) [9]. These approaches result in more parsimonious models and allow for PCEs to be applied in higher-dimensional problems, however do so at the cost of decreased model expressivity. Sparsity-inducing solvers have been proposed to relax the dimensionality constraint imposed by the size of the regression problem. Approaches may be based on a variety of solvers for the $\ell_1$-regularized least squares problem such as least-angle regression (LARS) that is used for PCEs in [10], compressive sensing [86] and orthogonal matching pursuit [61, 74, 19] as well as sparse Bayesian learning methods [73, 34, 67, 76]. For a comprehensive overview, the reader is referred to the recent literature review and benchmark study [50, 49].

4.2. Basis adaptation via partial least squares. In order to obtain a parsimonious yet expressive model, we turn to low-dimensional model representations rather than sparse solutions to the full-dimensional model. To achieve this, the PCE representation is rotated onto a new basis defined by the variables $Z = Q^T U$, where $Q \in \mathbb{R}^{d \times d}$ and $Q^T Q = I$, with $I$ denoting the identity matrix. This has first been proposed in [72]. The PCE with respect to the novel basis reads

$$\hat{\mathcal{J}}^Q_p(U) = \sum_{|k| \leq p} a_k \Psi_k(Z) = \sum_{|k| \leq p} a_k \Psi_k(Q^T U) .$$

Since $Z$ is a standard-normal random vector due to the orthogonality property of $Q$, the PCE basis elements remain unchanged. Merely, a new set of coefficients $a_k$ enters the formulation in the adapted basis. The columns of $Q$ define linear combinations of the original input. We seek to choose $Q$ such that most of the relevant information to construct an accurate surrogate $Y$ is captured in the first $m$ directions, where $m < d$ leads to dimensionality reduction. We retain only these first $m$ columns of $Q$ in the matrix $Q_m$ and define a corresponding PCE of reduced dimension as

$$\hat{\mathcal{J}}^Q_{m}(U) = \sum_{|k| \leq p} a_k \Psi_k(Q_m^T U) ,$$

where $k \in \mathbb{N}^m$. [72] computes the basis adaptation $Q_m$ by evaluating first- or second-order PCE coefficients only with a sparse-grid numerical quadrature. [75] couples this approach with compressive sensing to simultaneously identify $Q_m$ and the PCE coefficients in the subspace. In [58], we show that important directions can be identified efficiently based on a set of original function evaluations via partial least squares (PLS).

PLS establishes a linear relationship between variables $U$ and $Y$ based on $n_x$ observations of both quantities [83]. By $U_X \in \mathbb{R}^{n_x \times d}$, we denote the matrix of $n_x$ observations of $U$ and by $Y_X \in \mathbb{R}^{n_x \times 1}$ we denote the corresponding vector of scalar responses. PLS sequentially identifies $m$ latent components $\{t_j\}_{j=1}^m$, where $t_j \in \mathbb{R}^{n_x \times 1}$ such that they have maximum covariance with $Y_X$. After determining each $t_j$, PLS assumes a linear relationship between $t_j$ and $Y_X$ and evaluates the corresponding coefficient $a_j$ of $t_j$ by OLS. After each iteration, the matrices $U_X$ and $Y_X$ are deflated by the contribution of the $j$-th PLS-component. Components are extracted until a certain error criterion is met, which can be formulated, e.g., through the norm of the residual response vector or via cross-validation.

The nonlinear version of PLS in turn relaxes the assumption of a linear relationship between latent component and the response. A number of nonlinear PLS algorithms have been proposed [65]. Here we employ the approach of Refs. [82, 4] that introduces an additional loop into the algorithm for running a Newton-Raphson procedure iterating between the current latent component and the response. Ultimately, we are interested in computing the orthogonal transformation matrix $Q_m$ in (4.9). PLS produces two different matrices $R$ and $W$ that are suitable to this end, which motivates two different flavors of PLS-PCE. In PLS-PCE-R as proposed in [58] (see Subsection 4.3), each nonlinear relationship between the $\{t_j\}_{j=1}^m$ and the response is modelled as a univariate PCE. The coefficients of these univariate PCEs are computed simultaneously with the latent structure and the resulting model is a sum of univariate PCEs. Alternatively, the univariate PCEs are discarded after the PLS-PCE algorithm terminates and a multivariate (sparse) PCE is constructed in the subspace formed by the so-called weights $\{w_j\}_{j=1}^m$ leading to PLS-PCE-W (see Subsection 4.4).

4.3. PLS-PCE-R. PLS-PCE-R identifies $m$ latent components and for each component, it returns the direction $r_j$ and the univariate PCE along this direction. The univariate PCEs are defined by their
260 polynomial orders \( q_j \) \( j = 1 \) and the associated coefficient vectors \( \{ a_j \}^m_{j=1} \). The polynomial order is identified
261 with leave-one-out cross validation [15]. For each \((j\text{-th})\) latent component, the nonlinear PLS iteration is
262 repeated for different polynomial orders and \( q_j \) is chosen as the order minimizing the leave-one-out error.
263 The PLS-PCE-R model reads
264 \[
265 \hat{Y}(u) = \hat{a}_0 + \sum_{j=1}^{m} (a_j^q j) \psi_q \left[ r_j^T (u - \mu_U) \right],
266 \]
267 where \( \hat{a}_0 = \hat{E}[Y] \), \( \psi_q (U) \) is a vector function assembling the evaluations of the one-dimensional Hermite
268 polynomials up to order \( q_j \) and \( \mu_U \) is the columnwise sample mean of \( U_e \). The model structure is illustrated
269 in Fig. 2. The PLS directions \( r_j \) can be evaluated in terms of the PLS weights \( w_j \) and loads \( p_j \) through the
270 following recursive relation [30]
271 \[
272 r_1 = w_1
273 \]
274 \[
275 r_j = w_j - r_{j-1} (p_{j-1}^T w_j).
276 \]
277 \( R = [r_1, \ldots, r_m] \in \mathbb{R}^{d \times m} \) is a matrix collecting all PLS directions. \( R \) is not necessarily orthogonal, i.e., in
278 general \( R^T R \neq I \). However, in [58] it is shown that \( R^T R \approx I \) when \( n_e \) is large and \( U_e^T U_e \) is diagonal, which
279 is the case if \( U_e \) is drawn from \( \varphi_d \). In this case, (4.10) is equivalent to a PCE of the form (4.9), where only
280 main effects in the latent components are considered.
281
282 \[ 4.4. \text{PLS-PCE-W.} \] PLS-PCE-W defines \( W \) as basis of the subspace rather than \( R \), where \( W = \[
283 \]
284 \[ [w_1, \ldots, w_m] \in \mathbb{R}^{d \times m}. \] Within linear PLS, the columns of \( W \) form an orthogonal basis. Within nonlinear
285 PLS, the Newton-Raphson step may introduce deviations from orthogonality, which are however negligible in
286 all tested examples. The univariate PCEs obtained through the Newton-Raphson step will be optimal with
287 respect to \( R \), not \( W \). Thus, in PLS-PCE-W these univariate polynomials are discarded once \( W \) is identified
288 and a multivariate (sparse) PCE is constructed in the subspace defined by \( W \) using least-angle regression and
289 a hyperbolic truncation scheme for the multivariate PCE basis as proposed by [10]. In this way PLS-PCE-
290 W achieves more flexibility compared to PLS-PCE-R by including interactions of the latent components in
291 exchange for a departure from optimality in the match between latent component and surrogate model. In
292 analogy to (4.9), the PLS-PCE-W model reads
293 \[
294 \hat{Y}(u) = \hat{a}_0 + \sum_{k \in \alpha} \hat{a}_k \psi_k \left[ W^T (u - \mu_U) \right],
295 \]
296 where \( \alpha \in \mathbb{N}^{P \times d} \) is the multi-index set, which indicates the polynomial orders of the \( d \) univariate polynomials
297 in each of the \( P \) multivariate polynomials as obtained with LARS. Both PLS-PCE-R and PLS-PCE-W are
298 summarized in Algorithm 4.1. In the following, we will use the PLS-PCE-W model as we observed a superior
299 performance for this model compared to PLS-PCE-R models in the context of the proposed approach.
300
301 5. Learning PLS-PCE models in each SIS level.
302
303 \[ 5.1. \text{The sequential subspace importance sampler.} \] We recently proposed to reconstruct low-
304 dimensional PLS-PCE-W models in each level of SIS to improve the tractability of high-dimensional reliability
305 analysis with computationally expensive models [57]. We term this approach sequential subspace importance
306 sampling or SSIS. The efficiency of SIS benefits from surrogate modelling through a considerable reduction
307 of required model evaluations. The PLS-PCE model alone, being a global surrogate model, is a relatively
308 limited tool for reliability analysis. Combining it with SIS provides the means to sequentially move the DoE
309 towards relevant regions in the input space and thereby renders difficult reliability problems accessible to
310 surrogate modelling. At the \( i \)-th SSIS level, a new local DoE is sampled from the current importance density
311 \( h_i \) through a resampling step on the \( N \) available samples from \( h_i \). The new local DoE is appended to the
312 global DoE comprising earlier designs from levels 1 through \( i - 1 \). Based on the updated global DoE, a new
313 PLS-PCE model is constructed and SIS is rerun for \( i + 1 \) levels from \( h_0 \) to obtain samples for the next local
314 DoE. Due to this restart, it is sensible to let previously used local DoEs remain in the global DoE such that
315 the \( i \)-th surrogate model accurately predicts the LSF output along the entire path of samples moving from the
316 nominal distribution \( h_0 \) to \( h_i \). The restart itself incurs no additional LSF evaluations and serves to stabilize
Active learning has emerged in the late 1980s as a subfield of machine learning [70] and was known in the statistical theory of regression as optimal experimental design since the early 1970s [25]. At its heart is the idea that supervised learning algorithms can perform better if allowed to choose their training data. We consider a ‘pool-based sampling’ variant of active learning, in which a large pool of unlabeled data points are made available to the algorithm. Within SIS, one has \( n \) samples from \( h_i \) available in the \( i \)-th level. The algorithm then selects \( n_{\text{add}} \) points that are labeled (i.e. for which the LSF is evaluated) and added to the DoE based on a measure of information gain. This measure typically takes the form of a learning function \( L \) that is maximized over the sample pool to perform selection. The learning function employed in the context of SSIS is discussed in Subsection 5.2.

The probability of failure estimator for SSIS/ASSIS is analogous to (3.10) with the difference that SIS
Algorithm 4.1 PCE-driven PLS algorithm [58]

1: **Input** Input matrix $\mathbf{U}_\varepsilon$ and output vector $\mathbf{Y}_\varepsilon$, maximum polynomial order $p$
2: 3: Set $\mathbf{E} = \mathbf{U}_\varepsilon - \mu_\varepsilon$, $\mathbf{F} = \mathbf{Y}_\varepsilon - \mu_\varepsilon$, $\epsilon_w = 10^{-3}$, $\epsilon_y = 10^{-3}$, $j = 1$
4: repeat
5: | Compute weight $w_j^0 = \mathbf{E}^T \mathbf{F} / \|\mathbf{E}^T \mathbf{F}\|$
6: | for $q \leftarrow 1, p$ do
7: | | Set $w_j^q = w_j^0$
8: | repeat
9: | | Compute score $t_j^q = \mathbf{E} w_j^q$
10: | | Fit a 1D PCE of order $q$ $\hat{a}_j^q \leftarrow \text{fit} \left[ \mathbf{F} = (a_j^q)^T \psi_q(t_j^q) + \epsilon \right]$
11: | | Set $\hat{\mathcal{M}}_j^q(t) = (\hat{a}_j^q)^T \psi_q(t_j^q)(t)$
12: | | Compute the error $e = \mathbf{F} - (\hat{a}_j^q)^T \psi_q(t_j^q)$
13: | | Compute $\Delta w_j^q = (\mathbf{A}^T \mathbf{A})^{-1} \mathbf{A}^T e$ with $\mathbf{A} = \nabla_w (\hat{a}_j^q)^T \psi_q(\mathbf{E} w)$
14: | | Set $w_j^q \leftarrow w_j^q + \Delta w_j^q$
15: | | Normalize $w_j^q \leftarrow w_j^q / \|w_j^q\|$
16: | until $\|\Delta w_j^q\|$ is smaller than $\epsilon_w$
17: | Evaluate the relative leave-one-out error $\epsilon_j^{LOO}$ as in [10]
18: | Set $\{q_j, \hat{a}_j^q, w_j^q\}$ as the triple $\{q_j, \hat{a}_j^q, w_j^q\}$ with the smallest $\epsilon_j^{LOO}$
19: | Compute score: $t_j^{q_j} = \mathbf{E} w_j^{q_j}$
20: | Compute load: $p_j^{q_j} = \mathbf{E}^T t_j^{q_j} / ((t_j^{q_j})^T t_j^{q_j})$
21: | Deflate: $\mathbf{E} \leftarrow \mathbf{E} - t_j^{q_j} (p_j^{q_j})^T$, $\mathbf{F} \leftarrow \mathbf{F} - (\hat{a}_j^q)^T \psi_q(t_j^{q_j})$
22: | $j \leftarrow j + 1$
23: until change in $\|\mathbf{F}\|$ is smaller than $\epsilon_y$
24: Compute $\mathbf{R} = [\mathbf{r}_1, \mathbf{r}_2, \ldots, \mathbf{r}_n]$ according to (4.11) \hspace{1cm} \triangleright \text{For the R-based version of PLS-PCE}
25: Build $\hat{\mathbf{Y}}(\mathbf{u})$ according to (4.10)
26: Gather $\mathbf{W} = [w_1, w_2, \ldots, w_m]$ \hspace{1cm} \triangleright \text{For the W-based version of PLS-PCE}
27: Build $\hat{\mathbf{Y}}(\mathbf{u})$ according to (4.12) and [10]
28: return $\mathbf{R}/\mathbf{W}, \hat{\mathbf{Y}}(\mathbf{u})$

is performed with an LSF approximation $\hat{G}$ that is based on the final surrogate model:

\[
\hat{p} = \left( \prod_{i=1}^{M} \hat{s}_i \right) \frac{1}{n} \sum_{k=1}^{n} \mathbb{I}(\hat{G}(\mathbf{u}^k) \leq 0) \varphi_d(\mathbf{u}^k) \eta_M(\mathbf{u}^k), \quad \mathbf{u}^k \overset{i.i.d.}{\sim} \mu, \quad h_M.
\]  

The ratio of normalizing constants $\{\hat{s}_i\}_{i=1}^{M}$ are estimated as

\[
\hat{\xi}_t = \frac{1}{n} \sum_{k=1}^{n} \hat{\omega}_t(\mathbf{u}^k) = \frac{1}{n} \sum_{k=1}^{n} \frac{\Phi(-\hat{G}(\mathbf{u}^k) / \sigma_i)}{\Phi(-\hat{G}(\mathbf{u}^k) / \sigma_{i-1})}, \quad \mathbf{u}^k \overset{i.i.d.}{\sim} \mu, \quad h_i.
\]

The SSIS/ASSIS algorithms are stopped based on a similar criterion as for SIS given in (3.9):

\[
\mathbb{E} \left[ \mathbb{I}(\hat{G}(\mathbf{U}) \leq 0) \phi(-\hat{G}(\mathbf{U}) / \sigma) \right] \leq \delta_{\text{target}}.
\]

Fig. 3 depicts flow diagrams of the SSIS and ASSIS algorithms.

5.2. Active learning of low-dimensional model representations. In the context of SSIS, the learning function $\mathcal{L}$ should express the prediction uncertainty at each sample of the current IS density for a given PLS-PCE-W surrogate. This prediction uncertainty is due to the estimation of both the subspace and
the surrogate model with a finite-sized DoE. We describe this uncertainty with the variance of the LSF based
on the surrogate model conditional on \( u, V[G|U = u] \). Note that, whenever the distribution with respect to
which \( E[\cdot] \) or \( V[\cdot] \) are evaluated is not made explicit as a subscript, it is implicitly assumed as the distribution
of the argument. For example, \( V[G|U = u] = Vf_G|u \).

Let \( \xi_0 = a \in \mathbb{R}^{P \times 1} \) and \( \xi_j = w_j \in \mathbb{R}^{d \times 1}, j = 1, \ldots, m, \) such that \( \xi = [\xi_0^T, \xi_1^T, \ldots, \xi_m^T]^T \in \mathbb{R}^{md+P \times 1} \)
is the collection of all \( md + P \) model parameters. Further, let \( \xi^* \) denote their corresponding point estimates
returned by Algorithm 4.1. The first-order expansion of \( \tilde{V}[G|u] \) around \( \xi^* \) reads

\[
\tilde{\sigma}_G^2(u) = \tilde{V}[G|u] \approx \left[ \frac{\partial \tilde{G}}{\partial \xi} \right]_{\xi=\xi^*}^T \hat{\Sigma}_{\xi \xi} \left[ \frac{\partial \tilde{G}}{\partial \xi} \right]_{\xi=\xi^*},
\]

where \( \hat{\Sigma}_{\xi \xi} \) is an estimate of the parameter covariance matrix. Next, we neglect the pairwise cross-covariance
of PCE coefficients \( a \) and the subspace components \( w_j \) and consider

\[
\tilde{\sigma}_G^2(u) = \tilde{V}[G|u] \approx \sum_{j=0}^{m} \left[ \frac{\partial \tilde{G}(u, \xi)}{\partial \xi_j} \right]_{\xi_j=\xi_j^*}^T \hat{\Sigma}_{\xi_j \xi_j} \left[ \frac{\partial \tilde{G}(u, \xi)}{\partial \xi_j} \right]_{\xi_j=\xi_j^*}.
\]

This significantly reduces the number of \( \Sigma_{\xi \xi} \)-entries that have to be estimated, namely from \( P^2 + 2Pmd + m^2d^2 \)
to \( P^2 + md^2 \). More importantly, the coefficients of the PCE, \( \xi_0 \), are obtained with linear regression while
the subspace, \( \{\xi_j\}_{j=1}^m \), is obtained in the inner loop of Algorithm 4.1 with nonlinear regression. Under
some regularity conditions, the estimator \( \xi \) is consistent [84] and converges in distribution to a multivariate
Gaussian distribution with mean \( \xi \) and covariance \( \Sigma_{\xi \xi} \). In analogy with linear regression, an estimate of \( \Sigma_{\xi \xi} \)
is given through

\[
\hat{\Sigma}_{\xi_j \xi_j} = \frac{\tilde{\sigma}_j^2}{n_\xi} (A_j^T A_j)^{-1}
\]

with

\[
A_j = \left[ \frac{\partial \tilde{Y}(u, \xi)}{\partial \xi_j} \right]_{u=U_\xi} \in \mathbb{R}^{n_\xi \times d} \quad \text{and} \quad \tilde{\sigma}_j^2 = \frac{1}{n_\xi - md - P} \sum_{k=1}^{n_\xi} (Y^k - \hat{Y}(U_\xi^k))^2.
\]

\( \tilde{\sigma}_j^2 \) is the standard estimator for the error variance of the surrogate model. \( A_j \) is the gradient of the surrogate
model \( Y \) with respect to the model parameters evaluated at each of the \( n_\xi \) points in the DoE \( U_{\xi} \). \( A_0 \) is
merely the design matrix and does not require the computation of any derivatives. Note that computing the
gradients \( \{A_j\}_{j=0}^m \) does not require any model evaluations. For \( j = 0 \), it is

\[
\frac{\partial \tilde{Y}(u, \xi)}{\partial \xi_0} = [\Psi_1 (W^T (u - \mu_U))]_{i=1}^{P-1} \quad \text{with} \quad W = [\xi_1, \xi_2, \ldots, \xi_m].
\]

For \( j > 0 \) and recalling \( z = W^T (u - \mu_U) \), we have

\[
\frac{\partial \Psi_k(z)}{\partial \xi_j} = \frac{\partial}{\partial w_j} \Psi_k(W^T (u - \mu_U)) = \frac{\partial \Psi_k(z)}{\partial z_j} (u - \mu_U)
\]

\[
= (u - \mu_U) \prod_{\substack{i=1 \atop i \neq j}}^{m} \psi_{k_i}(w_i^T u) \frac{\partial \psi_{k_j}(w_j^T u)}{\partial z_j}
\]

\[
= (u - \mu_U) \prod_{\substack{i=1 \atop i \neq j}}^{m} \psi_{k_i}(w_i^T u) \sqrt{k_j \psi_{k_j - 1}(w_j^T u)}. \]
In the last equality, we have used the following expression for derivatives of univariate normalized Hermite polynomials:

\[
\frac{d\psi_n(x)}{dx} = \sqrt{n}\psi_{n-1}(x).
\]

(5.10)  

\[
\frac{\partial \hat{Y}(u, \xi)}{\partial \xi_j} \text{ for } j > 0 \text{ follows as }
\]

\[
\frac{\partial \hat{Y}(u, \xi)}{\partial \xi_j} = \frac{\partial \hat{Y}(z)}{\partial w_j} = \sum_{k \in \alpha} \hat{a}_k \frac{\partial \psi_k(z)}{\partial \xi_j}, \quad j > 0.
\]

(5.11)  

The partial derivative \( \frac{\partial \hat{G}}{\partial \xi_j} \) in (5.5) can be evaluated using the chain rule of differentiation, which yields

\[
\frac{\partial \hat{G}}{\partial \xi_j} = \frac{\partial \hat{G}}{\partial \hat{Y}} \frac{\partial \hat{Y}}{\partial \xi_j}.
\]

(5.12)  

The first term on the right-hand side is typically easy to compute and often equals \( \pm 1 \) (the sign is irrelevant as the gradient enters the quadratic form in (5.5)) if the LSF returns the difference between the model output and a prescribed threshold. In this case, the first factor on the right-hand side of (5.12) drops out. If, however, the LSF is not continuously differentiable with respect to the model, we may construct a surrogate model of \( \hat{G} \) directly by using a DoE containing LSF evaluations rather than model evaluations in Algorithm 4.1. The second term on the right-hand side can be obtained reusing the gradients from the \( A_j \) in (5.7) that — in this case — are not evaluated at the DoE and thus are functions of \( u \).

When setting up the learning function, there is a distinction to be made between an intermediate SIS level and the final SIS level: In the intermediate level, the goal is to accurately estimate the ratios of normalizing constants and to propagate the samples to the next level. In the final level, the goal is to build the probability of failure estimator and thus to accurately approximate the true limit-state hypersurface. With this in mind, the learning functions for adapting the surrogate models in levels \( i = 1, \ldots, M \), and after the final level are readily stated as

\[
L_G(u) = \begin{cases} 
\sigma_{\hat{G}}(u), & \text{intermediate SIS level} \\
\sigma_{\hat{G}}(u)/|\hat{G}(u)|, & \text{after final SIS level}.
\end{cases}
\]

(5.13)  

After the final level, SIS has converged and we are using samples from the final biasing density \( h_M \) to refit a surrogate model that captures the failure hypersurface well. The learning function in this case is defined in the spirit of the learning function put forward in [21]. The denominator penalizes samples whose image under \( \hat{G} \) is far away from 0 assuming that therefore they are themselves far away from the failure hypersurface. Such samples are unlikely to be misclassified as safe if located in the failure domain or vice versa. In all previous levels of SIS, there is no failure hypersurface to be approximated but only importance weights and the resulting ratio of normalizing constants. Here, the denominator in the learning function is dropped as there is no benefit to penalizing samples with large absolute image values under \( \hat{G} \).

In each AL iteration, the pool is searched for one or several points maximizing \( L(u) \). If \( n_{\text{add}} > 1 \) new points are added per AL iteration, the current sample pool is transformed to the low-dimensional subspace defined by \( W \) in order to identify \( n_{\text{add}} \) clusters (e.g., with k-means). Clustering in the subspace circumvents the performance deterioration most clustering methods experience in high dimensions [39]. The point maximizing (5.13) in each cluster is added to the DoE. In this way, the algorithm avoids a local concentration of the DoE in a single region and is also able to handle problems with multiple disconnected failure domains as long as these are contained in the subspace.

The active learning is terminated based on the maximum local standard deviation relative to the target average in the intermediate levels or based on the relative change of the probability of failure estimate after the final level:

\[
\left\{ \frac{\max_{k=1, \ldots, n} \left( \frac{\sigma_{\hat{G}}(u_k)}{|\hat{G}(u_k)|} \right) - \hat{\sigma}_{\text{final}}}{\hat{\sigma}_{\text{final}}} \leq \epsilon_{\text{AL}}, \quad \text{intermediate SIS level} \right\}, \\
\left\{ \frac{\hat{\sigma}_{\text{final}}}{p} \leq \epsilon_{\text{AL}}, \quad \text{after final SIS level} \right\},
\]

(5.14)  

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where appropriate choices for $\epsilon_{AL}$ lie in $[10^{-2}, 10^{-1}]$. $\hat{p}$ and $\hat{p}_{\text{last}}$ denote the probability of failure estimate based on the current and the last DoE within the AL loop. The probability of failure is estimated with a surrogate model-based run of SIS-aCS in each AL iteration. This causes no additional cost in terms of original model evaluations and ensures a reliable evaluation of the criterion even for extremely small failure probabilities. The active learning procedure is detailed in Algorithm 5.1 and the complete method is detailed in Algorithm 5.2.

**Algorithm 5.1 Active Learning**

1: **Input** LSF $G(\mathbf{u})$, AL error level $\epsilon_{AL}$, # of AL clusters $n_{\text{add}}$, Polynomial order $p$, DoE $\{U_\varepsilon, G_\varepsilon\}$,  
2: Sample pool $U_\text{pool}$
3: 
4: **while** true **do**
5: 
6: Run $[W, \hat{G}] = \text{PLS-PCE}(U_\varepsilon, G_\varepsilon, p, 'W')$  
7: **if** (5.14) **then**
8: break  
9: **end if**
10: Identify $n_{\text{add}}$ clusters among $U_\text{pool}W$  
11: for each cluster do
12: $U_\text{cluster} = \{\mathbf{u} \in U_\text{pool} : \mathbf{u} \in \text{cluster}\}$
13: Evaluate $\mathbf{u}^* = \text{argmax}\{\mathcal{L}(U_\text{cluster})\}$ according to (5.5)–(5.7), (5.12), and (5.13).
14: Append $U_\varepsilon \leftarrow [U_\varepsilon, \mathbf{u}^*]$  
15: Append $G_\varepsilon \leftarrow [G_\varepsilon, G(\mathbf{u}^*)]$
16: Remove $\mathbf{u}^*$ from $U_\text{pool}$
17: **end for**
18: return $U_\varepsilon, G_\varepsilon, \hat{G}$.

![Flowchart of Algorithm 5.1](image)

**Fig. 3:** Comparison of SIS-PLS-PCE with (right) and without (left) active learning.

6. **Numerical experiments.**

6.1. **Error measures.** In the following, we examine a series of examples of low to high input dimensionality characterized by varying degrees of nonlinearity of the LSF and varying number of disconnected failure regions. The computational cost of each approach is measured with the total number of required calls to the underlying computational model. The accuracy of the estimator is measured in terms of relative bias...
Algorithm 5.2 ASSIS (with PLS-PCE-W)

1: **Input** LSF $G(\mathbf{u})$, Target CoV $\delta_{\text{target}}$, Samples per level $n$, Input dimension $d$, DoE size $n_\epsilon$, AL error level $\epsilon_{\text{AL}}$, # of AL clusters $n_{\text{add}}$, Polynomial order $p$, $\epsilon_{\text{add}}$
2: 
3: Set $i = 0$, $\sigma_i = \infty$, $h_i(\mathbf{u}) = \varphi_d(\mathbf{u})$
4: Initialize $\mathbf{U}_\epsilon = [\cdot]$, $\mathbf{G}_\epsilon = [\cdot]$
5: Sample $\mathbf{U}_0 = \{\mathbf{u}_k\}_{k=1}^n \in \mathbb{R}^{n \times d}$
6: while true do
7: \hspace{1em} $i \leftarrow i + 1$
8: \hspace{1em} Sample $\mathbf{U}_{\text{tmp}} = \{\mathbf{u}_k\}_{k=1}^{n_\epsilon} \in \mathbb{R}^{n_\epsilon \times d}$
9: \hspace{1em} Compute $\mathbf{G}_{\text{tmp}} = G(\mathbf{U}_{\text{tmp}}) \in \mathbb{R}^{n_\epsilon \times 1}$
10: \hspace{1em} Append $\mathbf{U}_\epsilon \leftarrow \left[\mathbf{U}_\epsilon, \mathbf{U}_{\text{tmp}}\right]$
11: \hspace{1em} Append $\mathbf{G}_\epsilon \leftarrow \left[\mathbf{G}_\epsilon, \mathbf{G}_{\text{tmp}}\right]$
12: if $i > 1$ then
13: \hspace{1em} Run $\hat{G} = \text{PLS-PCE}(\mathbf{U}_\epsilon, \mathbf{G}_\epsilon, p, \ell W')$
14: \hspace{1em} Run $\mathbf{U}_{i-1}, \mathbf{G}_{i-1} = \text{SIS-aCS}(\hat{G}, \delta_{\text{target}}, n, d, i-1)$
15: \hspace{1em} Run $\mathbf{U}_\epsilon, \mathbf{G}_\epsilon, \hat{G} = \text{Active Learning}(G(\mathbf{u}), \epsilon_{\text{AL}}, n_{\text{add}}, p, \mathbf{U}_\epsilon, \mathbf{G}_\epsilon, \mathbf{U}_{i-1})$
16: Compute $\mathbf{G}_{i-1} = \hat{G}(\mathbf{U}_{i-1}) \in \mathbb{R}^{n \times 1}$
17: Compute $\sigma_i$ according to (3.8)
18: Compute $\hat{\sigma}_i$ and $s_i$ according to (5.2)
19: $\mathbf{U}_{i-1}, \mathbf{G}_{i-1} \leftarrow$ resample from $\mathbf{U}_{i-1}, \mathbf{G}_{i-1}$ with weights $\hat{\sigma}_i(\mathbf{U}_{i-1})$
20: Run $\mathbf{U}_i, \mathbf{G}_i = \text{SIS-aCS}(\mathbf{U}_{i-1}, \mathbf{G}_{i-1})$
21: if (5.3) then
22: \hspace{1em} Set $M \leftarrow i$
23: \hspace{1em} Run $\mathbf{U}_\epsilon, \mathbf{G}_\epsilon, \hat{G} = \text{Active Learning}(G(\mathbf{u}), \epsilon_{\text{AL}}, n_{\text{add}}, p, \mathbf{U}_\epsilon, \mathbf{G}_\epsilon, \mathbf{U}_{i-1})$
24: break
25: Run $\langle \mathbf{U}_M, \mathbf{G}_M, \hat{\sigma}_{\text{ASSIS}} \rangle = \text{SIS-aCS}(\hat{G}_M, \delta_{\text{target}}, n, d, M)$
26: return $M, \mathbf{U}_M, \mathbf{G}_M, \hat{\sigma}_{\text{ASSIS}}.$

and CoV
\begin{align*}
(6.1) \quad \text{relative Bias} &= \frac{p - \mathbb{E}[\hat{p}]}{p} \\
(6.2) \quad \text{CoV} &= \sqrt{\frac{\mathbb{V}[\hat{p}]}{\mathbb{E}[\hat{p}]}},
\end{align*}

where $p$ is the known exact probability of failure or a reference solution computed with a large number of samples as reported in the corresponding references in Table 1. Further, we compute the relative root-mean-squared error (RMSE) of the probability of any failure estimate $\hat{p}$, which combines bias and variability of the estimator as
\begin{align*}
(6.3) \quad \text{relative RMSE} &= \sqrt{\frac{\mathbb{E}[(p - \hat{p})^2]}{p^2}} = \sqrt{\text{relative Bias}^2 + \left(\frac{\mathbb{E}[\hat{p}]}{p}\right)^2 \text{CoV}^2}.
\end{align*}

The expectation and variance operators in the above equations are approximated by repeating each analysis 100 times. Additionally, the relative estimation error is defined as
\begin{align*}
(6.4) \quad \text{relative error} = \frac{\hat{p}}{p}.
\end{align*}

6.2. Low- and medium-dimensional examples. The subspace importance sampler is designed to tackle high-dimensional problems, yet its performance should not deteriorate as the problem dimension decreases. We first investigate its performance in eight examplatory problems with dimension $2 \leq d \leq 100$. 
Table 1: Low- to medium-dimensional investigated benchmark problems.

| Problem     | Failure probability | Inputs | Input Variables | Properties                        | References |
|-------------|---------------------|--------|-----------------|------------------------------------|------------|
| Hat         | $1.037 \cdot 10^{-4}$ | 2      | standard-normal | Strongly nonlinear                 | [68]       |
| Cantilever  | $3.94 \cdot 10^{-6}$ | 2      | Gaussian        | Strongly nonlinear                 | [6]        |
| 4-Branch    | $5.60 \cdot 10^{-9}$ | 2      | standard-normal | Multiple failure regions; extremely rare event | [6, 79]   |
| (acc. to [6]) |                   |        |                 |                                    |            |
| Borehole    | $1 \cdot 10^{-5}$   | 8      | Log-normal, Uniform | Strongly nonlinear, No underlying low-dimensional structure | [1]        |
| (276.7 m$^3$ year$^{-1}$) |            |        |                 |                                    |            |
| Truss       | $1.6 \cdot 10^{-3}$ | 10     | Log-normal, Gumbel | mildly nonlinear                   | [42]       |
| (0.12m)     |                     |        |                 |                                    |            |
| Rare Truss  | $1.02 \cdot 10^{-8}$ | 10     | Log-normal, nonlinear | Extremely rare event; (modified) | [42]       |
| (0.18m)     |                     |        |                 |                                    |            |
| Quadratic   | $6.62 \cdot 10^{-6}$ | 10     | standard-normal | Strongly nonlinear; Underlying low-dimensional structure | [23, 78] |
| ($\kappa = 5$) |                 |        |                 |                                    |            |
| Quadratic   | $6.62 \cdot 10^{-6}$ | 100    | standard-normal | Strongly nonlinear; Underlying low-dimensional structure | [23, 78] |
| ($\kappa = 5$) |                 |        |                 |                                    |            |

We demonstrate how both SSIS and ASSIS cope with multiple failure domains, strong nonlinearities and extremely small target failure probabilities. In the interest of brevity, the examples are listed in Table 1 along with the problem dimension, target probability of failure and key characteristics of the problem. The references provided in Table 1 may be consulted for detailed descriptions of the problem setups.

We solve the example problems with SIS-aCS with $n = 2 \cdot 10^3$ samples per level and a burn-in period of $b = 5$ samples within each MCMC chain. As suggested in [60], we choose $\delta_{\text{target}} = 1.5$ for the exit criterion (3.9) for SIS-aCS as well as our surrogate-based samplers. We compare this reference to SSIS and ASSIS for which we use an initial sample size of $n_E = 5d$. All underlying PLS-PCE-W models are computed with a maximum number of subspace directions of $m = 10$ and a maximum total polynomial degree of $|q| \ell_q \leq 7$, where $q = 0.75$. To achieve a fair comparison between ASSIS and SSIS, we first run ASSIS and then SSIS with $n_E$ for the latter chosen such that both methods use an approximately equal number of LSF evaluations. For both SSIS and ASSIS, we choose $n = 10^4$ with a burn-in period of $b = 30$. For ASSIS, we set $\epsilon_{\text{CAL}} = 0.02$. Within SSIS/ASSIS many samples per level and long burn-in periods are affordable as sampling is performed with the surrogate model. For ASSIS we select $n_{\text{add}} = 1$ unless prior knowledge of the problem structure suggests otherwise (the only exception in the set of examples considered here is the 4-branch function for which we select $n_{\text{add}} = 4$ as it features four relevant failure regions in the input space). Fig. 4 displays the performance of SIS, SSIS and ASSIS for the examples in Table 1 in terms of the error measures defined in Subsection 6.1 and the total number of LSF evaluations (with the original model).

For all showcased examples, ASSIS yields equally or more accurate estimates compared to SSIS at equal cost. It also either matches or outperforms SIS at significantly reduced costs. Except for the easiest problems, i.e., those featuring well-behaved (truss) or low-dimensional (2D hat) LSFs associated with comparatively large failure probabilities, the in-level adaptivity of ASSIS leads to significant bias correction (Fig. 4, bottom right) and variance reduction (Fig. 4, top right).

[60] discusses the choice of the MCMC sampler for SIS and find that aCS as employed here is outperformed by a Gaussian mixture proposal in low-dimensional problems, while the latter is the preferred choice as the problem dimension grows. Our method is designed for the solution of high-dimensional reliability...
problems and we thus consistently use aCS. However, we remark that in the 2-dimensional examples the relative RMSE produced by all considered SIS-based methods can be decreased by an order of magnitude by using the Gaussian mixture proposal at constant cost.

Comparing the truss and the rare truss models, the additional number of SIS levels required in the solution of the latter evidently leads to a deterioration of the SSIS estimate (Fig. 4, top left). This is due to single runs (less than 10\%) among the 100 repetitions in which the sampled DoEs lead to extreme outliers in the failure probability estimates (Fig. 5). While this effect vanishes when increasing the number of samples in the DoE, ASSIS offers a more cost-effective alternative to avoid such outliers by actively learning an informative augmentation of adverse DoEs. In this way, subspace identification and surrogate modelling errors cannot propagate and accumulate across the levels of SIS as they are controlled by the AL procedure. In fact, the phenomenon of rather rare but all the more severe outliers deteriorating the error mean and variability is a problem SSIS is facing not only in the rare truss example but also in the cantilever and both quadratic examples. Conversely, it is seen that in the 4-branch example, SSIS consistently and considerably overestimates the probability of failure while ASSIS captures the probability of failure rather well. The ASSIS-based probability of failure estimates each strive towards one of two accumulation points, one of which corresponds to the true probability of failure while the other represents a slightly underestimated failure probability. This second cumulation point corresponds to ASSIS runs where only two of the four important failure regions of the 4-branch functions have been identified. This behaviour can be alleviated by employing a Gaussian
mixture proposal in this example.

The two quadratic LSF models with 10 and 100 input dimensions demonstrate how the required num-
ber of LSF evaluations depends on the problem dimension in both surrogate-based approaches. This is due
to the fact that the PLS-PCE model requires at least \( d \) (often more) samples to identify a suitable subspace.
Thus, as described above, we choose \( n_E \) as a multiple of \( d \).

6.3. High-dimensional example: Steel plate. We consider a modified version of the example given
in [78, 48], which consists of a low-carbon steel plate of length 0.32 m, width 0.32 m, thickness \( t = 0.01 \) m,
and a hole of radius 0.02 m located at the center. The Poisson ratio is set to \( \nu = 0.29 \) and the density of the
plate is \( \rho = 7850 \text{ kg/m}^3 \). The horizontal and vertical displacements are constrained at the left edge. The
plate is subjected to a random surface load that acts on the right narrow plate side. The load is modelled as
a log-normal random variable with mean \( \mu_q = 60 \text{ MPa} \) and \( \sigma_q = 12 \text{ MPa} \). The Young’s modulus \( E(x,y) \) is
considered uncertain and spatially variable. It is described by a homogeneous random field with lognormal
marginal distribution, mean value \( \mu_E = 2 \times 10^5 \text{ MPa} \) and standard deviation \( \sigma_E = 3 \times 10^4 \text{ MPa} \). The
autocorrelation function of the underlying Gaussian field \( \ln E \) is modeled by the isotropic exponential model

\[
\rho_{\ln E}(\Delta x, \Delta y) = \exp \left\{ -\frac{\sqrt{\Delta x^2 + \Delta y^2}}{l_{\ln E}} \right\}
\]

with correlation length \( l_{\ln E} = 0.04 \text{ m} \). The Gaussian random field \( \ln E \) is discretized by a Karhunen-Loève-
expansion (KLE) with \( d_E = 868 \), which yields a mean error variance of 7.5% and reads

\[
E(x,y) = \exp \left\{ \mu_{\ln E} + \sigma_{\ln E} \sum_{i=1}^{d_E} \sqrt{\lambda_{i}^E} \varphi_i^E(x,y) \xi_i \right\}.
\]

\( \mu_{\ln E} \) and \( \sigma_{\ln E} \) are the parameters of the log-normal marginal distribution of \( E \), \( \{\lambda_{i}^E, \varphi_i^E\} \) are the eigenpairs
of the correlation kernel in (6.5) and \( \xi \in \mathbb{R}^{d \times 1} \) is a standard-normal random vector. The most influential
eigenfunctions (based on a global output-oriented sensitivity analysis of the plate model performed in [22])
are shown in Fig. 6 on the right.
The stress \( \sigma(x, y) = [\sigma_x(x, y), \sigma_y(x, y), \tau_{xy}(x, y)]^T \), strain \( \epsilon(x, y) = [\epsilon_x(x, y), \epsilon_y(x, y), \gamma_{xy}(x, y)]^T \) and displacement \( u(x, y) = [u_x(x, y), u_y(x, y)]^T \) fields of the plate are given through elasticity theory, namely the Cauchy-Navier equations [36]. Given the configuration of the plate, the model can be simplified under the plane stress hypothesis, which yields

\[
G(x, y) \nabla^2 u(x, y) + \frac{E(x, y)}{2(1-\nu)} \nabla (\nabla \cdot u(x, y)) + b = 0.
\]

Therein, \( G(x, y) := \frac{E(x, y)}{2(1+\nu)} \) is the shear modulus, and \( b = [b_x, b_y]^T \) is the vector of body forces acting on the plate. (6.7) is discretized with a finite-element method. That is, the spatial domain of the plate is discretized into 282 eight-noded quadrilateral elements, as shown in Fig. 6. The LSF is defined by means of a threshold for the the first principal plane stress

\[
\sigma_1 = 0.5(\sigma_x + \sigma_y) + \sqrt{(0.5(\sigma_x + \sigma_y))^2 + \tau_{xy}^2}
\]

evaluated at node 11 (see green marker Fig. 6, left). Node 11 indicates a location where maximum plane stresses occur frequently in this example. The LSF reads

\[
g(U) = \sigma_{\text{threshold}} - \sigma_1(U),
\]

where \( \sigma_{\text{threshold}} = 450 \text{ MPa} \). The target probability of failure is determined to \( p = 4.23 \cdot 10^{-6} \) with \( \text{CoV} = 0.0119 \) as the average of 100 repeated runs of subset simulation [3] with \( 10^4 \) samples per level.

SIS-aCS is run with \( n = 2 \cdot 10^4 \) samples per level and a burn-in period of \( b = 5 \) samples within each MCMC chain. SSIS and ASSIS are run with \( n = 10^5 \) samples per SIS level, a burn-in period \( b = 30 \) and an AL threshold of \( \epsilon_{\text{AL}} = 0.02 \). In the first level \( n_\xi = 900 \) and in each additional level only \( n_\xi = 100 \) samples are added in the initial sampling phase. Table 2 lists the average estimated probabilities of failure along with error measures and average number of required LSF evaluations. It is seen that both SSIS and ASSIS alleviate computational cost by more than an order of magnitude while at the same time reducing the relative RMSE by at least an order of magnitude. The decomposition of the RMSE in CoV and relative bias reveals that this is mostly due to variance reduction as SIS-aCS already yields a small bias.

A parameter study of important ‘tweakable’ parameters of ASSIS is depicted in Fig. 7. Parameters that are
Table 2: Accuracy and cost of SIS, SSIS & ASSIS for the plate example based on 100 repetitions of the analysis. The reference $p_{ref} = 4.23 \cdot 10^{-6}$ is computed with 100 repeated runs of subset simulation with $10^4$ samples per level with CoV = 0.0119 for the mean estimate.

| Method     | $E[p]$       | relative RMSE | CoV     | relative bias | avg. # LSF evaluations |
|------------|--------------|---------------|---------|---------------|------------------------|
| SIS-aCS    | $3.88 \cdot 10^{-6}$ | 0.576         | 0.625   | 0.083         | 17000                  |
| SSIS       | $3.99 \cdot 10^{-6}$ | 0.061         | 0.021   | 0.058         | 1300                   |
| ASSIS      | $4.10 \cdot 10^{-6}$ | 0.036         | 0.021   | 0.030         | 1318                   |

Fig. 7: Steel plate reliability using ASSIS: parameter influence studies.

not subject to a parametric study are chosen as above, with the exception of $n = 10^4$ instead of $n = 10^5$. The estimation error and computational cost of ASSIS is analyzed for varying active learning threshold $\epsilon_{AL}$, number of samples in the DoE $n_E$, the number of samples per SIS level $n$ and the target CoV $\delta_{target}$ used for the SIS procedure. The scaling of 10% between the initial DoE and all subsequent DoE samples samples is kept constant.

The parameters $\epsilon_{AL}$ and $n_E$ describe the behaviour of the surrogate modelling and active learning procedures while $n$ and $\delta_{target}$ describe SIS itself. Fig. 7 shows that increasing the target coefficient of variation leads to a reduced number of levels in the SIS procedure, which is directly associated with a reduction in computational cost. The reduction is relatively small here as most of the samples are added in the first level. By design, the number of required samples remains unaffected by varying the number of samples per SIS level, while the estimation error depends reciprocally on it. Conversely, and also by design, the computational cost depends monotonically on the choice of $n_E$. If a majority of the used original LSF evaluations are added during an AL procedure, this relationship may be nonlinear. For the plate example, however, the initially drawn DoE samples at each level makes up for the majority of used original LSF evaluations, hence the linear dependency. The estimation errors decrease slightly with increasing DoE size, although the effect is limited as high accuracy is already achieved with the first DoE of the lowest investigated DoE size. The fact...
that the subspace does not change significantly with increasing SIS level leaves little to be learned by adding more LSF evaluations to the DoE. This is also the reason for the competitive performance of SSIS in this example. The estimation errors (as well as the computational cost in this case) remain unaffected by varying AL thresholds $\epsilon_{AL}$, which is in line with the observation that a large fraction of the computational budget is spent on the initial DoE sampling rather than the AL-based DoE augmentation.

7. Concluding remarks. This paper proposes a method for the cost-efficient solution of high-dimensional reliability problems. We build on a recently introduced dimensionality reducing surrogate modelling technique termed partial least squares-driven polynomial chaos expansion (PLS-PCE) [58] and previous work, in which we use PLS-PCE surrogates to reconstruct biasing densities within a sequential importance sampling scheme [57] (sequential subspace importance sampling: SSIS). We refine this approach by devising an active learning procedure in each SIS level, which serves to effectively control the estimation error introduced by the surrogate-based importance density reconstructions. The learning procedure, i.e., the selection of new points for the DoE, is driven by an estimate of both the subspace and surrogate model estimation error. This criterion can be generally used in polynomial chaos expansion-based active learning procedures.

We showcase the performance of SSIS and ASSIS in nine example applications with input dimensionality ranging from $d = 2$ to 869. The examples feature different typical caveats for reliability methods such as multiple failure domains, strongly nonlinear limit-state functions and extremely small target probabilities of failure. Depending on the example, we achieve a cost reduction of one to over two orders of magnitude with ASSIS compared to the reference method (sequential importance sampling with the original model) at equal or lower estimation errors. It is shown that SSIS is susceptible to the randomness of the initial DoE occasionally producing outliers if the DoE is adverse. The active learning procedure (ASSIS) remedies this drawback and stabilizes the estimator by augmenting potentially adverse DoEs with informative additional samples.

The million dollar question, as with any surrogate model, is on the method’s ability to generalize. Certainly, there exist examples that do not possess a suitable linear subspace as required by PLS-PCE modelling. However, by means of coupling PLS-PCE with sequential importance sampling, this requirement is relaxed somewhat as only a locally accurate surrogate model is required to propagate samples from one intermediate biasing density to the next. Hence, ASSIS can still be expected to perform well for examples lacking a global low-dimensional structure as long as there exists a local low-dimensional structure that may vary across different regions in the input space.

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