Data-driven uncertainty quantification using the arbitrary polynomial chaos expansion

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1. Introduction

The lack of information about the properties of physical systems, such as material parameters or boundary values, can lead to model uncertainties up to a level where the quantification of prediction uncertainties may become the dominant question in application tasks. Most physical processes appearing in nature are non-linear and, as a consequence, the required mathematical models are non-linear. Traditional and very well-known approaches for stochastic simulation are brute-force Monte Carlo simulation (e.g. [29]) and related approaches (e.g. latin hypercube sampling [15]). Unfortunately, for large and complex models, Monte Carlo techniques are inadequate. Even single deterministic simulations may require parallel high-performance computing. As a reasonably fast and attractive alternative, stochastic model reduction techniques based on the polynomial chaos expansion can be applied.

Polynomial chaos expansion. A large number of studies for diverse applications is based on the polynomial chaos expansion (PCE) introduced by Wiener [52] in 1938. The chaos expansion offers an efficient high-order accurate way of including non-linear effects in stochastic analysis. PCE can be seen, intuitively, as a mathematically optimal way to construct and obtain a model response surface in the form of a high-dimensional polynomial in uncertain model parameters. The chances and limitations of polynomial chaos and related expansion techniques were discussed in [3]. The paper [43] showed how to use PCE for robust design under uncertainty with controlled failure probability. Recently, the sensitivity analysis based on PCE decomposition [5,9,34] has received increased attention. The papers [42,45] demonstrate correspondingly how classical PCE and its new aPC version can deliver the information required for global sensitivity analysis at low computational costs. Also, FORM and SORM methods (e.g. [17]) could be extended to higher-order accuracy via PCE, however, this has not yet been achieved.

The PCE technique can mainly be sub-divided into intrusive and non-intrusive approaches for the involved projection integral. The intrusive approach requires manipulation of the governing equations and can sometimes provide semi-analytical solutions for stochastic analysis. The best-known method from this group is the stochastic Galerkin technique, which originated from structural...
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any attempt to construct probability density functions of any
tainty in quantifying statistical model output distribution. Also,
information on input statistics introduces its own type of uncer-
metric transformations [11]. Using transformed variables for
by an adequate transformation called Gaussian anamorphosis
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Polynomial chaos expansion for non-Gaussian distributions. The
original PCE is based on Hermite polynomials, which are optimal
for normally distributed random variables. Unfortunately, natural
phenomena and uncertainty in engineering are often not that
simple, and the distribution of physical or model parameters
often cannot be considered Gaussian. However, it is possible to
put into conformity a physical variable with a normal variable
by an adequate transformation called Gaussian anamorphosis

Orthogonal polynomials by [57,58]. The gPC extends PCE towards
a counted number of parametric statistical distributions (Gamma,
Beta, Uniform, etc.). However, application tasks demand further
adaptation of the chaos expansion technique to a larger spectrum of
distributions. In [50,52], the authors presented a multi-element
generalized polynomial chaos (ME-gPC) method. It is based on a
decomposition of the random space into local elements, and
subsequently implements gPC locally within the individual ele-
mants. An error control theory for the ME-gPC method was
developed in [51] for elliptic problems. The ME-gPC is the first
adaptive piecewise approach helping to deal with discontinuity of
distributions or of model responses, and provides the desired
adaptation to a wide spectrum of distributions. The ME-gPC
conception offered in [50] provides a flexible tool for stochastic
modeling, but interprets these data as an exactly known prob-
bility distribution, and considerably increases the computational
effort for multidimensional stochastic problems.

Limited availability of data. The methods discussed above
assume an exact knowledge of the involved probability density
functions. Unfortunately, information about the distribution of
data is very limited in realistic engineering applications, especially
when environmental influences or natural phenomena are
involved, or when predicting or engineering the environment
(see also [46]). Applied research on (partially) natural or complex
realistic systems often faces the problem of limited information
about the model parameters and even about their probability
distributions. For example, material properties of underground
reservoirs are insufficiently available to provide a full picture
of their distribution. Moreover, the statistical distribution of model
parameters can be nontrivial, e.g., bounded, skewed, multi-modal,
discontinuous, etc. Also, the dependence between several uncer-
tain input parameters might be unknown, compare [25]. Depend-
ing on the modeling task and circumstances, statistical information
on model parameters may be available either discrete, continuous,
or discretized continuous, they could exist analytically as PDF/CDF
or numerically as histogram. The key shortcoming of current PCE
approaches in this context is twofold. First, they are heavily
restricted in handling most of these conditions, and second they
assume that this information is complete and perfect.

Small samples or data sets do not contain perfect or complete
information on the probability distribution of model input para-
eters. For example, the study [46] demonstrated that limited
information on input statistics introduces its own type of uncer-
tainty in quantifying statistical model output distribution. Also,
any attempt to construct probability density functions of any
particular shape from samples of limited size or from sparse
information introduces additional subjectivity into the analysis,
which bears the severe risk of leading to biased results. In a
related application study [44], we illustrate that errors or addi-
tional (and mostly subjective) assumptions in data interpretation
can severely bias uncertainty quantification and risk assessment,
and hence could lead to failing designs. Methods of maximum
entropy [18], closely related to known as the exponential poly-
nomial method [13] in reliability engineering, and minimum
relative entropy [56] are often used in the engineering sciences
to construct a probability distribution from sparse information
(mostly in the form of a few statistical moments and bounds) that
may be available from different instances of the same object or
from different objects with supposedly similar properties or
conditions. Although these two methods are designed to mini-
mize subjectivity and even though they can preserve the sample
moments up to arbitrary order, they are heavily debated within
the statistical community (e.g. [33]). In fact, they still introduce
new assumptions and impose a specific assumption on distribu-
tion shape. The same is true for other typical methods to
construct PDFs from moments in the field of reliability engine-
ering, such as the Hermite polynomial transformation [53]. Such
methods, however, are more subjective than entropy-based
methods, since they cannot keep the original sample moments
up to higher orders unchanged. If one still desires to fit a PDF as
a pragmatic tool to filter raw data against noise, one should have
full freedom in the chosen distribution shapes, not restricted by
the technical constraints of PCE or gPC.

Approach and novelties. To overcome the first part of the
problem, we claim that it is not even necessary to cast the
available statistical information into probability density func-
tions. Instead, the available information can directly and most
purely be used in stochastic analysis, when using our data-driven
formulation of PCE, see Section 2. We argue that applied tasks
demand direct handling of arbitrary data distributions without
additional assumptions for stochastic analysis. To overcome the
second part of the problem, we suggest to perform a robustness
analysis around the PCE (here: aPC) to assess the impact of
incomplete statistical input information. Overall, we propose to
align the complexity level and order of analysis with the relia-
bility and detail level of statistical information on the input
parameters.

The concept we propose in the current paper is to approach
the problem in a highly parsimonious and yet fully data-driven
description of randomness. We draw attention to the arbitrary
polynomial chaos (aPC) that has recently been touched upon in a
few theoretical papers. Two studies focusing on proofs of exis-
tence were published in the mathematical stochastics community
[14,37]. Constructing the aPC polynomials by Gram–Schmidt
orthogonalization was presented in the field of aerospace engi-
neering [55,54]. These studies did not discuss the aPC in the light
of data availability, limited reliability of data and assumptions in
data interpretation.

The aPC extends chaos expansion techniques by employing a
global polynomial basis for arbitrary distributions of data. In a
certain sense, it allows to return back to a global basis with the
new freedom of arbitrary polynomial chaos that the ME-gPC
[50,51] uses only within piecewise local elements.

The most important property of the aPC that we will install
and exploit in the current paper is that the aPC can work with
probability measures that may be (if necessary) implicitly and
incompletely defined via their moments only, and that it requires
no additional information. In fact, our equations will show
explicitly (in closed form) that statistical moments are the
only source of information that is propagated in all polynomial
expansion-based stochastic approaches. Thus, exact probability
density functions do not have to be known and do not even have
to exist. For finite-order expansion, only a finite number of moments has to be known. This opens the path to data-driven applications, where data samples with limited size merely allow inference of a few statistical moments, but are not sufficient to support, without a substantial degree of subjectivity, the assumption of an exactly known probability measure (see discussion in Section 4.1). Fully in line with the demands of application tasks, the statistical data of modeling parameters can be specified either analytically (as probability density/cumulative distribution functions), numerically as histogram or as raw data sets.

In Section 2, we deliver the necessary mathematical material, and provide the necessary properties and proofs in Section 3. The convergence rate of the aPC will be illustrated in the context of an example problem in Section 4. How to address issues arising from the incomplete and inaccurate character of raw data sets used as statistical input information is discussed and illustrated in Section 5.

2. The arbitrary polynomial chaos expansion

2.1. One-dimensional aPC

We will consider a stochastic process in the probability space \((\Omega, A, \mathcal{F})\) with space of events \(\Omega\), \(\sigma\)-algebra \(A\) and probability measure \(\mathcal{F}\), see e.g. [22]. Let us consider a stochastic model \(Y = f(\xi)\) with model input \(\xi \in \Omega\) and model output \(Y\). For a stochastic analysis of \(Y\), the model \(f(\xi)\) may be expanded as follows:

\[
Y(\xi) \approx \sum_{i=1}^{d} c_i P_i(\xi),
\]

where \(d\) is the order of expansion, \(c_i\) are the expansion coefficients that are determined by Galerkin projection, numerical integration or collocation, and \(P_i(\xi)\) are the polynomials forming the basis \(\{P_0(\xi), \ldots, P_d(\xi)\}\) that is orthogonal (or even orthonormal) with respect to the measure \(\mathcal{F}\) (see Eq. (6)). The only difference between aPC and previous PCE methods is that the measure \(\mathcal{F}\) can have an arbitrary form, and thus the basis \(\{P_0(\xi), \ldots, P_d(\xi)\}\) has to be found specifically for the probability measure \(\mathcal{F}\) appearing in the respective application.

This opens the path to data-driven applications of aPC. If a function \(Y(\xi)\) is expanded in the orthonormal polynomial basis \(\{P_0(\xi), \ldots, P_d(\xi)\}\), then characteristic statistical quantities of \(Y(\xi)\) can be evaluated directly from the expansion coefficients \(c_i\). For example, the mean and variance of \(Y(\xi)\) are given by the following simple analytical relations:

\[
\mu_Y = c_1, \quad \sigma_Y^2 = \sum_{i=2}^{N} c_i^2.
\]

Notice that, in the current paper, we will focus on mono-dimensional stochastic input (i.e., only one uncertain parameter) for simplicity, but without loss of generality (see Section 2.2).

2.2. Multi-dimensional aPC

Most realistic applications feature multi-dimensional model input \(\xi\), i.e. \(\xi = \{\xi_1, \xi_2, \ldots, \xi_N\}\). Here, the total number of input parameters is equal to \(N\). The model parameters can be design or control parameters that can be chosen by the operator of a system, and uncertain parameters that describe our (incomplete) knowledge of the system properties. Hence, to investigate the influence of all input parameters \(\xi_1, \xi_2, \ldots\) on the model output \(Y\), the model output \(Y\) can be represented by a multivariate polynomial expansion as follows:

\[
Y(\xi_1, \xi_2, \ldots, \xi_N) \approx \sum_{i=1}^{M} c_i P_i(\xi_1, \xi_2, \ldots, \xi_N).
\]

Here, the coefficients \(c_i\) quantify the dependence of the model output \(Y\) on the input parameters \(\xi_1, \xi_2, \ldots, \xi_N\). The number \(M\) of terms in the expansion (3) depends on the total number of input parameters \(N\) and on the order \(d\) of the expansion, according to the combinatorial formula \(M = (N + d)!/(N!d!)\). The function \(\Phi_1\) is a simplified notation of the multi-variate orthogonal polynomial basis for \(\xi_1, \xi_2, \ldots, \xi_N\). Assuming that the input parameters within \(\xi_1, \xi_2, \ldots, \xi_N\) are independent (e.g. [20]), the multi-dimensional basis can be constructed as a simple product of the corresponding univariate polynomials

\[
\Phi_1(\xi_1, \xi_2, \ldots, \xi_N) = \prod_{j=1}^{N} (\xi_j - \mu_{\xi_j}) \Phi_j(\xi_j),
\]

where \(\mu_{\xi_j}\) is a multivariate index that contains the combinatoric information how to enumerate all possible products of individual univariate basis functions. In other words, the index \(x\) can be seen as \(M \times N\) matrix, which contains the corresponding degree (e.g. 0, 1, 2, etc.) for parameter number \(j\) in expansion term \(k\).

Let us mention that, in the current state of science for polynomial chaos expansions, the random variables have to be statistically independent or may be correlated in a linear fashion only. Linear correlation can be removed by adequate linear transformation, such as the KL-expansion [26], also called proper orthogonal decomposition [28] or principal component analysis [31] in other disciplines. Construction of a joint polynomial basis for statistically dependent random variables beyond linear dependence is a very important issue for future research.

2.3. Stochastic analysis based on PCE

Eqs. (1) and (3) can be interpreted as a model response surface for \(Y = f(\xi_1, \xi_2, \ldots, \xi_N)\), and represent the basic key element for: (I) uncertainty quantification; (II) robust design and (III) global sensitivity analysis.

(I) The simplest way to quantify uncertainty is via the analytical relations, see Eq. (2). However, in order to evaluate more complex statistical quantities, Monte Carlo simulation can be performed directly and immensely fast on the obtained polynomial given by Eq. (3) (see e.g. [44,49]). For Monte Carlo simulation in the absence of precise statistical information, Oladyshkin et al. [44] discuss and suggest the maximum entropy method for PDF estimation.

(II) Including design and control parameters together with uncertain parameters in expansion (3) provide an effective basis for robust design. The paper [43] showed how to use PCE for robust design under uncertainty with controlled failure probability.

(III) Expansion (3) also delivers the information required for global sensitivity analysis including simultaneous influences of different modeling parameters at low computational costs. For example, Sobol indices [35,36] or Weighted indices [45] can be computed directly from the coefficients \(a_j\).

3. Moment-based analysis

Let us define the polynomial \(p^{(k)}(\xi)\) of degree \(k\) in the random variable \(\xi \in \Omega\):

\[
p^{(k)}(\xi) = \sum_{i=0}^{k} p_{i}^{(k)} \xi^i, \quad k = 0, 1, 2, \ldots
\]

where \(p_{i}^{(k)}\) are coefficients in \(p^{(k)}(\xi)\).
Our goal is to construct the polynomials in Eq. (5) such that they form an orthonormal basis for arbitrary distributions. The arbitrary distributions for the framework presented in our paper can be either discrete, continuous, discretized continuous, specified analytically, as histograms, raw data sets or by their moments. In this paper, we exploit this freedom and show how to treat any given probability distribution solely defined by the statistical moments of \( \xi \). For limited-order expansion, this allows to work with arbitrary probability measures that are implicitly defined as moments. In this paper, we exploit this freedom and show how the system in Eq. (10) by substituting the first equation into the second, the first and the second into the third, and so on. In addition, we will apply condition (8). Hence, without loss of generality, the system in Eq. (10) can be reduced to

\[
\int_{\xi \in \Omega} \sum_{i=0}^{k} p^{(k)}_{i} \xi^{i} \: d\Gamma(\xi) = 0,
\]

\[
\int_{\xi \in \Omega} \sum_{i=0}^{k} p^{(k)}_{i} \xi^{i+1} \: d\Gamma(\xi) = 0,
\]

\[
\vdots
\]

\[
\int_{\xi \in \Omega} \sum_{i=0}^{k} p^{(k)}_{i} \xi^{i+k-1} \: d\Gamma(\xi) = 0,
\]

\[p^{(k)}_{k} = 1. \tag{11}\]

Note that this rearrangement defines the \( k \)th orthogonal polynomial independent of all other polynomials from the orthogonal basis. The \( k \)th raw moment of the random variable \( \xi \) is defined as

\[
\mu_{k} = \int_{\xi \in \Omega} \xi^{k} \: d\Gamma(\xi). \tag{12}\]

This allows to re-write Eq. (11) based on only the raw moments of \( \xi \):

\[
\sum_{i=0}^{k} p^{(k)}_{i} \mu_{i} = 0,
\]

\[
\vdots
\]

\[
\sum_{i=0}^{k} p^{(k)}_{i} \mu_{i+k-1} = 0,
\]

\[p^{(k)}_{k} = 1. \tag{13}\]

Alternatively, the system of linear equations (13) can be written in the more convenient matrix form:

\[
\begin{bmatrix}
\mu_{0} & \mu_{1} & \ldots & \mu_{k} \\
\mu_{1} & \mu_{2} & \ldots & \mu_{k+1} \\
\vdots & \vdots & \ddots & \vdots \\
\mu_{k-1} & \mu_{k} & \ldots & \mu_{2k-1} \\
0 & 0 & \ldots & 1
\end{bmatrix}
\begin{bmatrix}
p^{(k)}_{0} \\
p^{(k)}_{1} \\
p^{(k)}_{2} \\
p^{(k)}_{k-1} \\
p^{(k)}_{k}
\end{bmatrix}
= \begin{bmatrix}
0 \\
0 \\
\vdots \\
0 \\
1
\end{bmatrix}. \tag{14}\]

As a direct consequence, an orthogonal polynomial basis up to order \( d \) can be constructively defined for any arbitrary probability measure \( \Gamma \) under the following conditions: the coefficients \( p^{(k)}_{i} \) can be constructed if and only if the square matrix of moments in the left-hand side of Eq. (14) is not singular. In the Appendix, we provide a proof for this under the condition that the number of support points in the distribution of \( \xi \) is greater than \( k \) and that all moments up to order \( 2k-1 \) are finite. This holds for all continuous random variables, under the condition that \( 2k-1 \) moments exist. If the moments of \( \xi \) are evaluated directly from a data set of limited size or from a discrete probability distribution featuring a finite number of possible outcomes, there need to be \( k \) or more
distinct values in the data set or distribution. All moments are always finite if no element of the data set is infinite.

From the Eqs. (7) to (14), it becomes evident that moments are the only required form of information on input distributions for constructing the basis and thus to operate the aPC. For finite-order expansion, a finite number of moments is sufficient. Hence, if a raw data set is the only form of available input information, computing its moments is sufficient, and estimating a full PDF from the data is not necessary. The same is true if only a limited number of moments are provided as input characterization. Also, arbitrary parametric distribution can be addressed, simply by working with their moments. This implies that any difference between distributions that becomes visible only in moments of order higher than 2d−1 will be invisible to any order d polynomial expansion technique.

3.2. Explicit form of data-driven polynomial chaos

In this section, we present an analytical explicit form of the coefficients for moderate degrees of polynomials, which can be easily used for diverse data-driven application tasks, such as uncertainty quantification, global sensitivity analysis and probabilistic risk assessment. The coefficients for the higher degrees of polynomials can be obtained using the implicit scheme presented above (14), via recursive relations (see [1, Chapter 22]), via Gram–Schmidt orthogonalization (see [55,54]) or via the Stieljes procedure [39].

To simplify the explicit form of coefficients, we will assume a normalized distribution of data with zero mean and unit variance after linear transformation

$$z' = \frac{z - \mu}{\sigma},$$

which leads to a centralization and standardization of all moments. Thus, the orthogonal polynomial basis \(P^k(\zeta)\) \((k = 0, d)\) can be presented as

$$P^d(\zeta) = \sum_{l=0}^{d} p_{l}^{d} \left( \frac{z - \mu}{\sigma} \right)^{l},$$

where \(p_{l}^{d}\) are the coefficients of polynomial \(P^d\) defined explicitly through the raw moments of \(z\) from the relations below. Due to Eq. (15), the raw moments of \(\zeta\) are related to the central moments \(\mu_i(\zeta)\) of \(\zeta\) via:

$$\mu_i(\zeta) = \tilde{\mu}_i(\zeta) \psi^i(\zeta)^{-k/2}.$$

Coefficients for polynomial of 0 degree

$$p_0^{(0)} = 1.$$

Coefficients for polynomial of 1st degree

$$p_0^{(1)} = 0, \quad p_1^{(1)} = 1.$$

Coefficients for polynomial of 2nd degree

$$p_0^{(2)} = -1, \quad p_1^{(2)} = \mu_3, \quad p_2^{(2)} = 1.$$

Coefficients for polynomial of 3rd degree

$$p_0^{(3)} = \mu_2^2 - \mu_3^2 + \mu_3 \mu_4 - \mu_5, \quad p_1^{(3)} = -\mu_3 \mu_5 + \mu_2^2 - \mu_4 + \mu_3 \mu_4.$$

Coefficients for polynomial of 4th degree

$$p_0^{(4)} = \mu_3^2 \mu_2 \mu_4 + \mu_3^2 \mu_2 \mu_4 - \mu_3^2 \mu_4 + 2 \mu_3 \mu_2 \mu_4 \mu_6 + 2 \mu_3 \mu_2 \mu_4 \mu_6 - \mu_2^2 \mu_6 - \mu_3 \mu_2 \mu_4 \mu_6,$$

$$p_1^{(4)} = -\mu_3^2 \mu_2 \mu_4 + \mu_3^2 \mu_2 \mu_4 - \mu_3^2 \mu_4 + \mu_3 \mu_2 \mu_4 \mu_6 + \mu_3 \mu_2 \mu_4 \mu_6.$$

3.3. Normalization

The above orthogonal polynomial basis can be used directly for analysis. However, an orthonormal basis has more useful properties (see Eq. (2) and Section 3.4). Thus, the next step is to normalize the orthogonal basis. We will use the norm for the polynomial \(P^l\) introduced in Eq. (6):

$$\|P^l\|^2 = \int_{-1}^{1} (\psi^l(\zeta))^2 \lambda(\zeta).$$

Hence, a valid orthonormal polynomial basis \(\{\psi^{l(k)}(\zeta)\} (k = \overline{0,d})\) is:

$$\psi^{l(k)}(\zeta) = \frac{1}{\|P^l\|} \sum_{i=0}^{l} p_{i}^{l(k)} \zeta^i.$$

For normalization, the evaluation of \(\|P^l\|\) for \(k = d\) additionally requires finiteness and availability of the 2d th moment.

3.4. Summarized properties of the orthonormal basis

As consequence of the derivations in Sections 3.1 and 3.3, the polynomial basis in Eq. (23) has the following properties:

**Property I.** The orthonormal basis can be constructed without any hierarchical conditions or recurrence relations that are used in [1, Chapter 22] and in [54,55].

**Property II.** Existence of the moments \(\mu_0,\ldots,\mu_{2d}\) is the necessary and sufficient condition for constructing an orthonormal basis \(\{\psi^{l(0)},\ldots,\psi^{l(d)}\}\) up to degree d, together with the condition that the number of support points of \(\zeta\) is greater than d if \(\zeta\) is a discrete variable or is represented by a data set.

**Property III.** The orthonormal polynomial basis for arbitrary probability measures is based on the corresponding moments only, and does not require the knowledge (or even existence) of a probability density function.

**Property IV.** All the zeros of the orthogonal polynomials are real, simple and located in the interior of the interval of orthogonality [1]. This property is useful for numerical integration, especially for bounded distributions.

**Property V.** As particular cases, the Hermite, Laguerre, Jacobi polynomials, etc. from the Askey scheme and the polynomials for log-normal variables by Ernst et al. [14] can be reconstructed within a multiplicative constant.

**Property VI.** All distributions that share the same moments up to order 2d will also share the same basis, and thus will lead to identical results in an expansion up to order d.

4. Data-driven modeling

The arbitrary polynomial chaos expansion presented in Sections 2 and 3 provides a simple and efficient tool for analysing stochastic systems. We will consider a very simple model in order to focus all attention on our data-driven concept, which is based directly on the moments of sampled data without intermediate
steps of data reinterpretation. This avoids the subjectivity usually introduced when choosing among a small limited number of theoretical distributions to represent a natural phenomenon, and so avoids the problems of subjectivity under limited data availability is discussed in Section 1. These problems will be illustrated in Section 4.1. An application to a problem with a realistic level of complexity and a detailed discussion of expert’s subjectivity in uncertainty analysis is presented in [44]. That paper demonstrates how subjectivity of interpreting limited data sets can easily lead to substantial prediction bias, and that the subjective choice of distribution shapes has a similar relevance as uncertainties due to physical conceptualization, numerical codes and parameter uncertainty.

Here, for simplicity, we consider the exponential decay differential equation which was already used in [58] to illustrate the Askey scheme:

\[
\frac{dY(t)}{dt} = -\xi Y, \quad Y(0) = 1.
\]  

Let \(Y_{PC}\) be the solution obtained using the polynomial chaos expansion (1) for the problem defined in Eq. (24). We use a Monte Carlo simulation as reference solution and define the time dependent relative error \(\epsilon(t)\) between the polynomial chaos expansion solution \(Y_{PC}(t)\) and the Monte Carlo solution \(Y_{MC}(t)\) as

\[
\epsilon(t) = \frac{|Y_{PC}(t) - Y_{MC}(t)|}{Y_{MC}(t)}. \quad (25)
\]

### 4.1. Fidelity of data-driven interpretation

To illustrate the fidelity of the data-driven chaos expansion, we will consider a synthetic example for an empirical data distribution, see Fig. 1, and apply both aPC and the classical PCE for comparison. The illustrative data set (with sample size \(N=500\)) presented in the left plot of Fig. 1 was generated as superposition of normal and log-normal distributions and contains statistical noise introduced due to the small size of the data sample.

The classical approach would be to introduce a parametric probability distribution, e.g., with fitted mean and variance or with maximum likelihood parameters. Here, for illustration, we select the Normal, Lognormal and Gamma distribution, see the right plot of Fig. 1. Evidently, the list of possible candidate distributions for fitting to the considered data can be very long. Introducing a full probability density function (PDF) resembles a strong assumption on all higher moments up to infinite order, and claims to know the exact shape, e.g., also of the extreme value tails. Such assumptions on the alleged shape of the underlying probability density function, unfortunately, can lead to substantial errors in data interpretation. The data-driven approach strongly alleviates this situation, because it can directly handle a set of moments (e.g., the mean, variance, skewness, kurtosis, and so forth), without any further assumptions on higher-order moments and without having to introduce a PDF at all. This also provides the freedom to work with only a small number of moments obtained via expert elicitation, without asking for a full PDF.

We will apply different orders of the polynomial chaos expansion (1) to the test problem (24) using two sources of input information about the data distribution: (1) the three introduced assumptions on PDFs (right plot of Fig. 1) and (2) the pure raw data sample (left plot of Fig. 1). To observe the pure impact of data interpretation regardless of numerical techniques, we treat all four cases with the aPC, i.e., we construct an optimal orthonormal polynomial base (see Section 2) for each case. This avoids the non-linear transformations that are usually necessary to map the assumed PDFs onto the normal PDF. Such transformations would introduce additional errors, as discussed in Sections 4.2 and 4.3.

Technically, the coefficients \(c_i\) in the chaos expansion can be obtained, e.g., by Galerkin projection (e.g. [19,30,58]) or by the Collocation method (e.g. [16,26,60]). Both methods lead to the same result when using the optimal polynomial basis in the case of univariate analysis. Fig. 2 illustrates the convergence of the mean and variance (at time \(t=1\)) for the assumed PDFs (left plot of Fig. 1) and for the pure raw data (right plot of Fig. 1).

All considered cases reproduce an acceptable approximation with the linear expansion. Increasing the expansion order shows strong convergence for the data-driven polynomial chaos expansion. However, increasing the order does not assure convergence for the expansions based on interpreted data. The problem does not lie in poor numerical properties when treating these distributions, but in accurate convergence to a wrong value. This error is introduced only by fitting parametric PDFs to the data instead of letting the data fully manifest themselves. In fact, the PDF-based analysis are good only up to first order, because only moments up to the second are represented accurately by the matched PDFs. This means that all effort spent for higher-order expansion (especially for complex problems) is invested to negligible improvement, if not matched with an adequate effort for accurate data interpretation. The key advantage of aPC is this respect is that (1) it allows full freedom in the used type of input information, and (2) our analysis in Section 3 makes explicitly clear what amount of information (i.e., the moments up to a certain number)
intrusive approaches, the resulting values evaluate the integral in Eq. (26). In both intrusive and non-projection (intrusive) and Gauss quadrature (non-intrusive) to these two expansions techniques, we will apply both Galerkin mapping onto the normal distribution) to the example (24). For all tree techniques, we analyse the change with the distribution of the random variable \( x \) (here: Hermite) after transformation strongly enters the analysis at what expansion order. This allows to align the complexity and order of analysis with the reliability and detail level of statistical information on the input parameters.

4.2. Evidence of improved convergence

In this section we will illustrate the efficiency of analysis within an optimal (data-driven) polynomial basis. We wish to show the improved convergence rate of the arbitrary polynomial chaos compared to the classical PCE technique. The classical PC requires non-linear transformations to map non-normal input data distributions onto the normal PDF. In this technical aspect, the classical PCE does not differ from the gPC, which requires transformation onto one of possible PDF from Askey scheme. Eq. (24) can be expanded (see Eq. (1)) in the orthogonal polynomial basis \( \Psi_d(\xi) \). The projection coefficients are defined as

\[
q_i(t) = \int_{\xi \in \Omega} Y(t)\Psi_d(\xi) d\Gamma(\xi), \quad i = 0, \ldots, d. 
\] (26)

We will apply both the aPC and the classical PCE (including the mapping onto the normal distribution) to the example (24). For these two expansions techniques, we will apply both Galerkin projection (intrusive) and Gauss quadrature (non-intrusive) to evaluate the integral in Eq. (26). In both intrusive and non-intrusive approaches, the resulting values \( q_i(t) \) from Eq. (26) will change with the distribution of the random variable \( \xi \). For the optimal basis (used in aPC), however, the results from Galerkin projection and numerical integration coincide. This yields 3 distinguishable techniques. For all tree techniques, we analyse the performance for diverse exemplary distributions of the univariate input \( \xi \) (Rayleigh, Weibull, Log-normal). Detailed descriptions of these distributions can be found in [38]. For the classical PCE, the random variable \( \xi \) is not distributed in the same space as the polynomial basis \( \Psi_d(\xi) \), and an additional conversion is required. Thus we map the model variable \( \xi \) onto a corresponding normal variable \( \xi_N \) by Gaussian anamorphosis or normal score transform [48]. Fig. 3 illustrates the convergence of the mean and variance of \( Y \) (at time \( t = 1 \)) for our three exemplary distributions of the model input \( \xi \). As previously demonstrated for the gPC [57], expansion in the optimal polynomial basis without transformation shows at least an exponential convergence. Convergence with a non-optimal basis (here: Hermite) after transformation strongly depends on the nonlinearity of the required transformation from \( \xi \) to \( \xi_N \).

4.3. Clarification of error types

In Section 4.1, we demonstrated the possible errors introduced by subjective data interpretation when fitting parametric PDFs to raw data. In that analysis, we deliberately excluded the error by the different numerical accuracies of aPC and classical PCE. Now, we will demonstrate the faster numerical convergence of aPC compared to classical PCE with transformation, but this time excluding the error of data interpretation. We classify the cause of error into two types: I—transformation expansion error and II—numerical integration error. Modeling within the non-optimal basis using Galerkin projection leads to errors by expanding and truncating the transformation, which we denote here as transformation expansion error (type I). Modeling within the non-optimal basis using Gauss quadrature entails numerical integration error (type II). Using the optimal basis provides identical results for both intrusive and non-intrusive methods, because numerical integration is exact when using the roots of the \( d+1 \) order polynomial from the optimal basis, and because no transformation from \( \xi \) to \( \xi_N \) is necessary.

Transformation expansion error (type I). For intrusive manipulation, the anamorphosis transformation from \( \xi \) to \( \xi_N \) has to be expanded in \( \xi_N \). The finite number \( d \) of terms in this expansion causes the first type of error. The difference between expansion in an optimal basis and expansion of \( Y(\xi) \) after transformation to \( \xi_N \) is the so-called “aliasing error” (see [59]). Fig. 4 illustrates the nature of the this type of error. Expansion of the transformation (here \( \xi = \exp(\zeta_N) \)) at different orders is shown in the left plot of Fig. 4. The right plot of Fig. 4 demonstrates the corresponding mapping of a normal probability density function (PDF) back to physical space using the expanded and truncated expansion. In these examples, the normal PDF should transform to a log-normal PDF. The strong nonlinearity of the logarithmic transformation leads to a poor approximation with a finite number of terms. Thus, the choice of a non-optimal polynomial basis for the model input \( \xi \) leads to a wrong representation of the probability measure \( \Gamma_N \). This leads to the erroneous analysis of model output \( Y(\zeta) \) visible for Galerkin-based computations in Fig. 3.

Numerical integration error (type II). The accuracy of numerical integration (especially sparse) strongly depends on the choice of integration points. For example, in Gauss–Hermite integration, the polynomial basis defines the positions \( \zeta_i \) of integration points in the space of the input variable by the roots of the polynomial of degree \( d+1 \). Thus, using a non-optimal polynomial basis provides a non-optimal choice of the integration points, which causes the
second type of error. To illustrate this type of error, let us consider a stochastic model with a random variable $x$ that follows a non-Gaussian distribution. The selected model is an extremely simple non-linear one

$$Y(x) = x^6.$$  (27)

In our example, the input parameter $\xi$ is distributed according to the Chi-square distribution. We will construct two expansions: one based on Hermite polynomials with adequate Gaussian anamorphosis, and one based on optimal polynomials for the Chi-square distribution of the input data. In both cases, we employ Gauss quadrature and compare the results to a reference
solution. The supposedly optimal location of the integration points for Hermite polynomials correspond to the roots of the Hermite polynomial of order \(d+1\), back-transformed from \(\xi_N\) to \(\xi\) by anamorphosis. However, the truly optimal distribution of the integration points are the roots of the optimal polynomials that are orthogonal for \(\xi\) without further transformation. The obvious difference is shown in the left plot of Fig. 5. The transformed points are shifted against the optimal ones and thus cannot be considered as an optimal choice for numerical integration. Therefore, strong nonlinearity in the transformation leads to significant errors in PCE techniques that derive their numerical integration rules from the involved basis. Evidently, the example in Eq. (27) has an analytical solution, which can be reproduced by the expansion of 6th order within the optimal basis (see the right plot in Fig. 5). However, the transformed Hermite chaos combined with non-optimal Gauss quadrature does not converge to the known analytical solution even for the expansion degree \(d=6\) that should be, in theory, fully accurate construct \(Y = \xi^6\).

5. Robustness analysis for inaccurate input data

The presented approach can handle different forms of input information. In particular, it can directly handle raw data, which can be useful for practical applications. However, when the input data set is small, the sample moments are only uncertain estimates of real moments. Hence, a direct application of the method presented becomes less robust. In that case, it would be useful to apply some standard methods to assess the robustness in the estimation of moments, such as Jackknife or Bootstrapping (e.g. [12]). In the field of reliability engineering, Bootstrap methods have been applied to construct upper confidence limits for unreliability in [10]. Bootstrap-based confidence intervals caused by the uncertainty representing computationally demanding models by meta-models has been investigated in the paper [41] via regression based sensitivity analysis. The recent paper [4] explores sparse and partially random integration techniques for PCE, applied to sensitivity analysis, and provides Monte Carlo based estimates for the error introduced by the random character of the used integration rules. In this section, we focus on the robustness of data-driven expansions with respect to the limited size of a raw data set, that represents the underlying probability distributions of model input only inaccurately.

For that, we repeatedly (\(N=1000\)) generated raw data according to the assumed underlying theoretical distribution. Each time, we constructed a new data-driven basis and performed a projection of the model output \(Y\) (Eq. (24)) to the corresponding data-driven polynomial basis and computed the mean \(\mu_Y\) and variance.
of the model output $Y(\tau = 1)$ in each repetition. From this, we computed the variance of the mean $\sigma_Y^2$ and the variance of the variance $\sigma_{\sigma_Y^2}^2$. This entire nested Monte Carlo analysis was repeated for sizes of the raw data set ranging from $N=20$ to 1000. Fig. 6 shows the results for the distributions considered in Section 4.2. For this illustration, we used a 3rd degree of expansion. Other degrees of expansion (1–6) show similar results, all of them having error variances inversely proportional to the size of the sampled data set ($1/N$), i.e. having error standard deviations proportional to $1/\sqrt{N}$. This rate is visible as the slope of the scatter plots in Fig. 6.

The scatter is caused by the finite number of Monte Carlo repetitions used in the error estimation. It corresponds to the Monte Carlo error of the error estimate. Only visually, the scatter increases with increasing size $N$ of the data sets due to the logarithmic scale of the ordinate. The important aspect of the plots in the Fig. 6, however, is not the degree of scatter (i.e. the uncertainty of the error estimation), but the average slope (i.e. the error estimate itself).

Apparently, the data-driven chaos expansion has a convergence rate proportional to $1/\sqrt{N}$ for the standard deviation and confidence intervals of computed model output statistics. This convergence rate is well known for the variance of sample statics and from Monte Carlo techniques in general [6]. This means that the aPC does not guarantee the robustness and convergence properties with respect to insufficient sample size in comparison to moments from classical Monte Carlo techniques in general [7]. Karl Weierstrass established his approximation theorem in 1885, which states that every continuous function defined on an interval can be uniformly approximated as closely as desired by a polynomial function. A generalization of the Weierstrass theorem was proposed in the Stone-Weierstrass theorem where, instead of the interval, an arbitrary compact Hausdorff space is considered and, instead of the algebra of polynomial functions, approximation with elements from more general subalgebras were investigated [40]. Thus, if the random space is an arbitrary compact Hausdorff space, uniform convergence is guaranteed and the polynomial space is dense in the arbitrary compact Hausdorff space. However, it is not assured that any polynomial expansion (including the optimal orthonormal basis) will uniformly converge in any random space and the definition of the random space will define convergence in that space.

Formal knowledge about the convergence of the polynomial basis for diverse random spaces can be very useful for practical needs, such as the convergence for the Hermite basis in normal space [7]. Karl Weierstrass established his approximation theorem in 1885, which states that every continuous function defined on an interval can be uniformly approximated as closely as desired by a polynomial function. A generalization of the Weierstrass theorem was proposed in the Stone-Weierstrass theorem where, instead of the interval, an arbitrary compact Hausdorff space is considered and, instead of the algebra of polynomial functions, approximation with elements from more general subalgebras were investigated [40]. Thus, if the random space is an arbitrary compact Hausdorff space, uniform convergence is guaranteed and the polynomial space is dense in the arbitrary compact Hausdorff space. However, it is not assured that any polynomial expansion (including the optimal orthonormal basis) will uniformly converge in any random space and the definition of the random space will define convergence in that space.

Especially in such cases with very small data sets, expert opinion can be very useful to filter the data set, remove alleged outliers, fit a simple or complex PDF, and so forth. In our proposed approach, an expert will have total freedom of data interpretation (not restricted to the selection among standard PDFs) and can provide much more sophisticated information (e.g. lower and higher moments, complex and even non-parametric distributions, etc.). According to our approach, expert opinion (in a most general sense) will be incorporated directly without any additional transformation or additional subjectivity when translating it to the stochastic numerical framework. The presented methods allow experts to choose freely of technical constraints the shapes of their statistical assumptions.

6. Remaining issues for future research

The polynomial basis for continuous or discrete random variables can be constructed if and only if the number of support points (distinct values) in the distribution (within the available data set) is greater than the desired degree of the basis (see Property II in Section 3.4). However, for discrete cases we cannot guarantee that the integration points (see Section 4) will be distributed only within the space of a random variable. Still, it would be possible for each integration point to find a neighbouring point that belongs to the discrete space, but the convergence is not guaranteed, and this remains an open question for future research.

Formal knowledge about the convergence of the polynomial basis for diverse random spaces can be very useful for practical needs, such as the convergence for the Hermite basis in normal space [7]. Karl Weierstrass established his approximation theorem in 1885, which states that every continuous function defined on an interval can be uniformly approximated as closely as desired by a polynomial function. A generalization of the Weierstrass theorem was proposed in the Stone-Weierstrass theorem where, instead of the interval, an arbitrary compact Hausdorff space is considered and, instead of the algebra of polynomial functions, approximation with elements from more general subalgebras were investigated [40]. Thus, if the random space is an arbitrary compact Hausdorff space, uniform convergence is guaranteed and the polynomial space is dense in the arbitrary compact Hausdorff space. However, it is not assured that any polynomial expansion (including the optimal orthonormal basis) will uniformly converge in any random space and the definition of the random space will define convergence in that space.
uniquely solvable for the lognormal distribution. However, the mentioned works demand existence and precise knowledge about the probability density function, which is neither required nor desired for the arbitrary polynomial chaos (aPC). We observed convergence of the aPC in Section 4 for a counted number of useful cases, however research on a formal proof of convergence is a remaining question for future research.

As an outlook for future development, we point out the construction of a joint basis for parameters that have a complex statistical dependence beyond correlation. Following that direction, the aPC could be the first PCE family member that will allow to handle non-linear statistical dependence between input variables.

7. Summary and conclusions

In the current paper, we presented the arbitrary polynomial chaos expansion (aPC). The aPC conception provides a constructive and simple tool for uncertainty quantification, global sensitivity analysis and robust design. It offers a new data-driven approach for stochastic analysis that avoids the subjectivity of assigning parametric probability distributions that are not sufficiently supported by available data. We show that a global orthonormal polynomial basis for finite-order expansion demands the existence of a finite number of moments only, and does not require exact knowledge or even existence of a probability density function. Thus, the aPC can be constructed for arbitrary parametric and non-parametric distributions of data, even if the statistical model output characterization of input data is incomplete.

Also, the orthonormal basis can be constructed without using any hierarchical conditions or recurrence relations with polynomials of lower-order. For discrete random variables, the aPC can be constructed if and only if the number of discrete values of the random variable is greater than the largest considered degree of the basis. In case of continuous random variables, the aPC can be constructed from a number of moments which equals to two times the degree of the basis. If desired, the method can work directly with raw sampled data sets to represent the uncertainty and possible variational ranges of input data. The presented methods allow experts to choose freely of technical constraints the shapes of their statistical assumptions and makes explicitly clear what amount of information (i.e., the moments up to a certain order) enters the analysis at what expansion order. Overall, this allows to align the complexity and order of analysis with the reliability and detail level of statistical information on the input parameters.

We also provided numerical studies for diverse exemplary distributions, where we illustrated convergence rates for optimal and non-optimal polynomial bases using intrusive and non-intrusive methods. This analysis strictly illustrated that using a non-optimal polynomial basis provides slow convergence of the chaos expansion and therefore causes additional errors in subsequent analysis. Modeling results within our new data-driven aPC show, at least, an exponential convergence for all examined cases. We defined, discussed and illustrated the difference between numerical integration error and transformation expansion error. Both errors lead to wrong estimates of statistical characteristics when using a non-optimal basis. Thus, the aPC not only provides freedom for modeling physical systems with unknown probability distribution function, when only data sets of very limited size are available, but it also provides better convergence rates than conventional polynomial chaos techniques.

When the statistical information used as input is inaccurate or uncertain, i.e. when using small sets of raw data, a new form of uncertainty enters into the analysis and has to be considered in convergence assessments. We propose to apply Jackknife or Bootstrapping methods to assess the uncertainty of moments from small data sets, and then propagate that uncertainty through the aPC onto the output statistics to assess robustness. In an illustrative test case, we observe the classical Monte Carlo convergence rate for the results of the PCE analysis with respect to the size of the raw data set.

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Appendix: Non-singularity of the moments matrix

Let us write the square matrix of Eq. (14) in the following decomposed form:

\[
M = \begin{bmatrix}
H & B \\
C & D \\
\end{bmatrix},
\]

where

\[
H = \begin{bmatrix}
\mu_0 & \cdots & \mu_{k-1} \\
\vdots & \ddots & \vdots \\
\mu_{k-1} & \cdots & \mu_{2k-2} \\
\end{bmatrix}, \quad B = \begin{bmatrix}
\mu_k \\
\vdots \\
\mu_{2k-1} \\
\end{bmatrix},
\]

\[
C = [0 \ldots 0], \quad D = [1].
\]

Evidently, \( D \) is always invertible, and hence the determinant of \( M \) is given by:

\[
\det(M) = \det(D)\det(H - BD^{-1}C)
\]

Because \( \det(D) = 1 \) and \( C = [0, \ldots, 0] \), we obtain:

\[
\det(M) = \det(H).
\]

The matrix \( H \) is also known as the Hankel matrix of moments. The properties of its determinant were studied in the paper [27]. Moreover, Karlin [24] showed that \( \det(H) \) for \( \text{rank}(H) = k \) is zero if and only if the distribution of \( \xi \) has only \( k \) or fewer points of support. Thus, \( M \) is non-singular if and only if the number of support points in the distribution of \( \xi \) is greater than \( k \) and if all moments up to order \( 2k-2 \) are finite.

References
