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# 12 Abstract

The consideration of imprecise probability in engineering analysis to account for missing, vague or incom-13 plete data in the description of model uncertainties is a fast-growing field of research. Probability-boxes 14 (p-boxes) are of particular interest in an engineering context, since they offer a mathematically straight-15 forward description of imprecise probabilities, as well as allow for an intuitive visualisation. In essence, 16 p-boxes are defined via lower and upper bounds on the cumulative distribution function of a random 17 variable whose exact probability distribution is unknown. However, the propagation of p-boxes on model 18 inputs towards bounds on probabilistic measures describing the uncertainty on the model responses is nu-19 merically still very demanding, and hence is subject of intensive research. In order to provide an overview 20 on the available methods, this paper gives a state-of-the art review for the modelling and propagation of 21 p-boxes with a special focus on structural reliability analysis. 22

# 23 Keywords: imprecise probability, p-boxes, literature review, reliability analysis, surrogate modelling

# 24 1. Introduction

Numerical models give an unparalleled insight into the response of the structure under consideration 25 to a set of predefined loading conditions, and hence, allow for a largely virtualized design optimization 26 workflow. Examples of such models include finite element models of structures or thermal systems, but 27 also other numerical schemes aimed at approximating complex multi-physical systems from the nanoscopic 28 to the largest possible level can be considered. However, despite the highly detailed numerical predictions 20 that can be obtained, these results often do not achieve a satisfactory level of agreement with 'reality', 30 i.e., the actual physical behaviour of the considered continuum in the effective operational environment. 31 This discrepancy is caused by epistemic (reducible) and aleatory (caused by variation) uncertainty in 32 the model. Usually, a distinction between model form and parametric uncertainty is made, where the 33

former describes possibly unwarranted approximations of the mathematical description of reality, whereas the latter refers to discrepancies in the parameters of these models with respect to reality. This paper solely focuses on parametric uncertainties. In recent years, several highly performing methods based on stochastic analysis [1], fuzzy set theory and interval analysis [2] have been introduced in literature to account for these type of uncertainties in the model parameters  $\boldsymbol{x}$ . Also several authors compared the applicability of a selection of these techniques in applications such as Geotechnical engineering [3] or inverse uncertainty quantification for stochastic dynamics [4, 5].

#### 41 1.1. Probabilistic analysis

Probabilistic analysis is a powerful and mature tool to deal with aleatory uncertainties in numerical 42 analyses. In order to express aleatory uncertainty in the model parameters, they are usually modelled 43 as random variables, denoted by  $\mathbf{X} = (X_1, \ldots, X_{n_x})$  with support domain  $D_{\mathbf{X}} \subseteq D_x$ . Their values are 44 outcomes of a random experiment where a probability P can be assigned to X taking a value within 45 a specific measurable set that is a subset of  $D_{\mathbf{X}}$ . The probability that  $\mathbf{X}$  is less than or equal to  $\mathbf{x}$  is 46 modelled as a joint cumulative distribution function (CDF)  $F_{\mathbf{X}}(\mathbf{x}) = P(X_1 \leq x_1, \ldots, X_{n_x} \leq x_{n_x})$  for 47  $x \in D_X$ . Its derivative is denoted by  $f_X$  and is known as the joint probability density function (PDF). 48 Since the inputs of the model are represented by a random vector, it follows that the model responses 49 become random variables Y, which are distributed according to the (generally unknown) CDF  $F_Y$ . Note 50 that  $F_X$  and  $F_Y$  in general do not belong to the same family of distribution functions. 51

Let  $\mathcal{M}$  represent a function that maps a set of  $n_x$  input parameters  $\boldsymbol{x} \in D_x \subseteq \mathbb{R}^{n_x}$ , with  $D_x$  a set of feasible input parameters (e.g., non-negative Young's moduli or contact stiffness values), to a set of  $n_y$ output parameters  $\boldsymbol{y} \in \mathbb{R}^{n_y}$  via following relationship:

$$\boldsymbol{y} = \mathcal{M}(\boldsymbol{x}),\tag{1}$$

where  $\mathcal{M}$  may represent numerical model that provides a discretized approximation of the continuum 55 physics that describe the modelling problem at hand. Usually, given  $f_X$ , an analyst is then interested in 56 computing the expected value of some random variable  $\mathcal{H}(\mathbf{X})$ , i.e.,  $E[\mathcal{H}(\mathbf{X})]$ . Here, E is the expected 57 value operator and  $\mathcal{H}$  is a function defined on  $D_{\mathbf{X}}$ . Typically, in this context,  $\mathcal{H}$  is used to compute the 58 nth central moments of  $\mathbf{Y}$ , with  $n \in \mathbb{N}$ . Hereto,  $\mathcal{H}$  represents the component-wise exponentiation of the 59 model responses  $\boldsymbol{y} = \mathcal{M}(\boldsymbol{x})$ , i.e.,  $\mathcal{H}(\boldsymbol{x}) = \boldsymbol{y}^n$ , or  $\mathcal{H}(\boldsymbol{x}) = (\boldsymbol{y} - \boldsymbol{\mu}_{\boldsymbol{Y}})^n$  with  $\boldsymbol{\mu}_{\boldsymbol{Y}} = E[\boldsymbol{Y}]$ . In an engineering 60 context, an analyst is mostly interested into whether their design, be it a structure, system or a complex 61 network, will perform reliably given the uncertainties in their manufacturing and operating conditions. 62 Usually, the probability of failure is estimated in this context to assess the reliability of their design. 63

The probability of failure can be computed as  $p_{\rm f} = P(\mathcal{M}(\mathbf{X}) \leq 0)$ , where  $\mathcal{M}$  with  $n_y = 1$  represents a performance function that indicates whether the design failed  $(\mathcal{M}(\mathbf{x}) \leq 0)$  or not  $(\mathcal{M}(\mathbf{x}) > 0)$  for  $\mathbf{x} \in D_{\mathbf{X}}$ . In this context,  $\mathcal{H}(\mathbf{x})$  is defined as  $\mathcal{H}(\mathbf{x}) = I_{\mathcal{M}}(\mathbf{x})$  with  $I_{\mathcal{M}}$  the indicator function that has value 1 in case  $\mathcal{M}(\mathbf{x}) \leq 0$ ,  $\mathbf{x} \in D_{\mathbf{X}}$ , and 0 otherwise. Overall, the expected value of  $\mathcal{H}(\mathbf{X})$  is determined by evaluating the integral of the following form:

$$\mathcal{P} = \int_{D_{\boldsymbol{X}}} \mathcal{H}(\boldsymbol{x}) f_{\boldsymbol{X}}(\boldsymbol{x}) \, \mathrm{d}\boldsymbol{x}, \tag{2}$$

where the physical interpretation of  $\mathcal{P} = E[\mathcal{H}(\mathbf{X})]$  depends on the definition of  $\mathcal{H}$ . For the remainder of 69 the paper, the notation  $\mathcal{H}$  is used to abstract the specific application (i.e., calculation of the moments or 70 failure probability approximation) from the method that is being discussed. While at first sight it might 71 be tempting to evaluate this integral using numerical quadrature schemes, such solutions become quickly 72 unfeasible with respect to the non-linearity of the limit state function and/or the number of considered 73 random variables [6], even though lower/upper bounds [7] or approximate solutions [8] exist in certain 74 cases. In general, even integrating just the joint PDF (i.e.,  $\mathcal{H} = 1$ ) is not so trivial by quadrature, as they 75 tend to be extremely non-linear, especially when the random variables are highly correlated. Therefore, 76 Eq. (2) is usually solved by asymptotic approximations [9] or advanced simulation methods such as subset 77 simulation [10], directional importance sampling [11] or the probability density evolution method [12] in 78 case of stochastic dynamics. 79

#### 80 1.2. Imprecise probabilistic analysis

In most real-life applications, an analyst has only partial information about  $F_{\mathbf{X}}$  or  $f_{\mathbf{X}}$  due to the 81 presence of epistemic uncertainty. This is a result of the often imprecise, diffuse, fluctuating, incomplete 82 or vague nature of the available information. Moreover, the available information might be objective or 83 subjective and consist of collected data (e.g., via experiments or data mining) and theoretical knowledge 84 on the considered problem, but also expert opinions with different levels of trustworthiness [13]. Some 85 illustrations of such situations can be found in the benchmark study presented in [14]. In engineering 86 analysis, the main challenge is then to formulate suitable models that incorporate these various sources 87 of data in an objective way, without introducing unwarranted conclusions and/or ignoring significant 88 information to ensure that the calculated results do not deviate from reality. The class of imprecise prob-89 abilistic approaches attempts to solve this general problem and includes a plethora of different methods, 90 including Bayesian methods [15, 16, 17, 18], random sets [19, 20, 21], sets of probability measures [22], 91 evidence theory-based methods (such as Dempster-Shafer Theory) [23, 24, 25, 26] and interval probabil-92 ities [27] of which probability bounds methods [28] and fuzzy stochastic methods [29, 30] are extensions. 93



Figure 1: Illustration of parametric and distribution-free p-boxes. The black lines indicate the graphs of  $\overline{F}_X$  and  $\underline{F}_X$ , being the bounds on the p-boxes. The colored lines illustrate a set of admissible distribution functions for  $F_X$  that constitute the p-box.

Furthermore, a study of Monte Carlo methods for the general case of propagating imprecise probabilities 94 is given for instance in [31] or [32]. Answering the question on which of these methods is the most ap-95 propriate method from this broad class of techniques is in general not possible as the most appropriate 96 mathematical framework depends on the nature of the information that is available to the analyst. It 97 should be noted that the application of the general framework of imprecise probability theory requires 98 complex mathematical descriptions and methods. Furthermore, due to several restricting assumptions 99 that are required, the methods are sometimes also very hard to translate to engineering practice. For a 100 thorough treatment on the selection of the most appropriate method, the reader is referred to [13]. 101

In many engineering applications, simplified imprecise probability models are often preferable for sim-102 pler utilization and representation. A popular representative thereof are probability-boxes (p-boxes), 103 which provide a set of possible probability distributions for  $F_{\mathbf{X}}$  bounded by a lower CDF  $\underline{F}_{\mathbf{X}}$  and an 104 upper CDF  $\overline{F}_{\mathbf{X}}$ . This type of credal set encompassing the unknown CDF is computationally efficient [33], 105 easy to construct [34], and offers a simple graphical representation, see Fig. 1. This figure shows the 106 two main types of p-boxes, being parametric and distribution-free p-boxes. Distribution-free p-boxes 107 consider only the upper and lower CDF, and any CDF that complies with these bounds is admissible. 108 Parametric p-boxes on the other hand impose additional constraints on admissible distribution functions, 109 for instance by defining a family of distribution functions. A rigorous and more detailed definition of 110 both types of p-boxes is given in Section 2.1. 111

<sup>112</sup> Their simpler utilization and representation make the application of p-boxes particularly interesting

for engineering analysis. However note that even with all their benefits over other, more general, imprecise 113 probability models, computations involving p-boxes still require large computational budgets as they 114 incorporate effectively a set of probability distributions that all need to be accounted for. Hence, advanced 115 methods for p-box propagation have been subject to intense research over the past decades and various 116 efficient methods addressing numerous applications of different complexity were proposed. This paper 117 aims at giving an overview of a selection of promising approaches for the propagation of p-boxes in 118 engineering analysis. This is complemented by an introduction to p-boxes showing their relationship to 119 related imprecise probability models including their translation, and capabilities how to construct p-boxes 120 based on given information. 121

#### 122 1.3. A guideline to read this paper

Depending on the need of the reader, this paper can be used in several ways. For instance, a newcomer 123 in the field of imprecise probabilities and/or p-boxes might use the entire manuscript to get the overall 124 ideas on the methods, as well as obtain the references to recent key works in the field. In this case, it is 125 recommended to consider all sections of the paper. On the other hand, an analyst that is knowledgeable 126 with imprecise probabilities, but is unsure how to model them based on available data will gain most 127 from the information in Section 3. Conversely, if an analyst is unsure which state-of-the-art propagation 128 method is best applicable for their problem, they are kindly referred to Section 4 and the references 129 therein included. To give the full overview; Section 2 describes the theoretical foundations of p-boxes 130 and their analysis. Section 3 discusses the construction of p-boxes based on various sources of information. 131 Section 4 highlights a selection of developments for the propagation of p-boxes, published during the last 132 few years and ends with a summarizing table. Finally, Section 5 lists the conclusions of this paper. 133

#### 134 2. Probability boxes

In the following two sections, the case  $n_x = 1$  is considered for notational simplicity. This is furthermore warranted since most engineering literature on the subject, as will be clear from Section 3, either considers the univariate case of  $n_x = 1$ , or when  $n_x > 1$  full independence among all  $X_i$ ,  $i = 1, ..., n_x$ , with  $F_{\mathbf{X}}(\mathbf{x}) = \prod_{i=1}^{n_x} F_{X_i}(x_i)$ ,  $\mathbf{x} \in D_{\mathbf{X}}$ . For more information on the general modeling of multivariate p-boxes including dependence, the reader is referred to [35, 36].

### 140 2.1. Theoretical background

The main idea of a p-box is that there exist an unknown CDF  $F_X$  of the random variable X for which only bounds can be provided. Thus, a p-box is described by a lower CDF  $\underline{F}_X \in \mathbb{F}$  and an upper CDF  $\overline{F}_X \in \mathbb{F}$ , where  $\mathbb{F}$  expresses the set of all CDFs on  $D_X \subseteq \mathbb{R}$ . These CDFs are collected as a pair

 $[\underline{F}_X, \overline{F}_X]$  which yields a set of possible CDFs  $\{F_X \in \mathbb{F} \mid \underline{F}_X(x) \leq F_X(x) \leq \overline{F}_X(x), x \in D_X\}$  for the 144 unknown CDF of X. The definition of a p-box corresponds to defining a lower probability  $\underline{P}$  and upper 145 probability  $\overline{P}$  on events  $\{X \leq x\} = (-\infty, x] \cap D_X$ , i.e.,  $\underline{P}(X \leq x) = \underline{F}_X(x)$  and  $\overline{P}(X \leq x) = \overline{F}_X(x)$ 146 for  $x \in D_X$ , which yields a credal set of probability measures. Via the p-box framework, the epistemic 147 uncertainty that comes for example from incomplete data on  $F_X(x)$  is accounted for by assigning an 148 interval  $[\underline{F}_X(x), \overline{F}_X(x)]$  for each value of  $x \in \mathbb{R}$ , see [34]. In case sufficient high quality information over 149 the entire range of possible values for x is available to the analyst,  $[\underline{F}_X(x), \overline{F}_X(x)]$  will be a tight interval, 150 and the p-box will be close to a crisp (deterministic) distribution. Otherwise, when less information is 151 available, the bounds may become wider to acknowledge weaker confidence in the results. In case no 152 further assumptions are made concerning the set of possible CDFs, this type of p-box is also denoted a 153 distribution-free p-box. This is the most general type of p-box, which allows for the highest flexibility 154 when modelling parameters subject to aleatory and epistemic uncertainty, since any non-decreasing and 155 right-continuous function that is consistent with these bounds is admissible. Indeed, it can be shown 156 that crisp values, intervals and crisp probability distributions are all special cases of the distribution-free 157 p-box [28]. As a final note, since distribution-free p-boxes are so general in their definition, also CDFs 158 that are questionable from a physical perspective are explicitly included in the definition. 159

Besides distribution-free p-boxes, there are **parametric p-boxes**, which are described by a family 160 of CDFs whose parameters  $\theta_i \in \mathbb{R}$  are unknown up to the property that they must be contained within 161 intervals  $[\underline{\theta}_i, \overline{\theta}_i], i = 1, \ldots, n_{\theta}$ . These parameters describe specific distribution properties and are collected 162 in the vector  $\boldsymbol{\theta} \in \mathbb{R}^{n_{\theta}}$ . The Cartesian product of the intervals is also denoted as  $D_{\boldsymbol{\theta}}$ , i.e., it holds  $\boldsymbol{\theta} \in D_{\boldsymbol{\theta}}$ . 163 Hence, a parametric p-box yields the set of possible CDFs  $\{F_X(\cdot, \theta) \in \mathbb{F} \mid \theta \in D_{\theta}\}$  for the unknown CDF 164 of the random variable X. An example of a parametric p-box can be defined as the Gaussian distribution 165 family with parameters  $\boldsymbol{\theta} = (\mu, \sigma)$  contained in  $D_{\boldsymbol{\theta}} = [\underline{\mu}_X, \overline{\mu}_X] \times [\underline{\sigma}_X, \overline{\sigma}_X]$ . Parametric p-boxes have the 166 property to clearly distinguish between aleatory uncertainty, represented by the distribution family, and 167 epistemic uncertainty, represented by the intervals for the parameters  $\theta$ . The upper and lower bounding 168 CDFs of a parametric CDF can be computed as 169

$$\underline{F}_X(x) = \min\{F_X(x, \theta) \mid \theta \in D_{\theta}\},\tag{3}$$

$$\overline{F}_X(x) = \max\{F_X(x, \theta) \mid \theta \in D_\theta\}$$
(4)

for  $x \in D_X$ . Note that the distribution-free p-box defined by these bounds does not correspond to the parametric p-box as the latter one is more restrictive in general, i.e. there are CDF within these bounds not belonging to the family of the parametric p-box. Both types of p-boxes are illustrated in Fig. 1.

<sup>173</sup> In order to account for more information about the shape of CDFs, such as an admissible distribu-

tion family, symmetry, or about bounds on one or more statistical moments of  $F_X$ , a p-box can also be described by a quintuple  $(\overline{F}_X, \underline{F}_X, \mu_X^I, \sigma_X^I, \mathcal{F})$ , see [13]. Here, the confidence interval of the mean value  $\mu_X^I \subseteq [-\infty, \infty]$ , the confidence interval  $\sigma_X^I \subseteq [0, \infty]$  of the standard deviation, and the family of admissible CDFs  $\mathcal{F} \subseteq \mathbb{F}$  can be specified. Note that a distribution-free p-box can also be represented as a quintuple, noted  $(\overline{F}_X, \underline{F}_X, [-\infty, +\infty], [0, \infty], \mathbb{F})$ . Furthermore, the p-box framework was also recently extended to account for imprecision in stochastic processes by explicitly accounting for additional epistemic uncertainty in the process' autocorrelation structure [37, 38].

In the following subsections, the connection of p-boxes to some closely related uncertainty models for imprecise probabilities is demonstrated. This may help the reader in both understanding the similarities and differences between p-boxes and these models and converting them into p-boxes or vice versa.

#### 184 2.2. Hierarchical probabilistic models

An alternative approach to deal with parametric p-boxes is to apply hierarchical probabilistic models. 185 Following this approach, the epistemic uncertainty related to the parameters  $\boldsymbol{\theta}$  of the CDF  $F_X(\cdot, \boldsymbol{\theta})$  are 186 represented using a random variable  $\Theta$  with distribution  $F_{\Theta}$ . On the one hand, hierarchical probabilistic 187 models can be regarded as a special case of a p-box where intervals are used to bound possible values of 188  $\theta$ . According to possibility theory, these intervals encode the set of all distribution functions bounded by 189 the interval. As such, selecting a single distribution function out of this set introduces knowledge into the 190 analysis that might not be fully objective. On the one hand, parametric p-boxes might be constructed 191 using credible intervals from Bayesian methods along with hierarchical probabilistic models, see Section 192 3.4. In this case, p-boxes describe an excerpt of this modelling where the tails of  $F_{\Theta}$  are neglected. 193

Using hierarchical probabilistic models, the effect of the epistemic uncertainty on the probabilis-194 tic measure under consideration depends on the applied propagation schemes. For instance, when re-195 weighting schemes such as presented in [39, 40, 41] are applied to infer the bounds, this is not problematic 196 since they allow for a clear separation between aleatory and epistemic uncertainty. In these types of meth-197 ods, the distribution  $F_{\Theta}$  is a purely instrumental tool to determine a functional relationship between  $\mathcal{P}$ 198 and  $\theta$ , the influence of which is integrated out of the result in later stages of the analysis, see Section 4. 199 However, when this single distribution is used to make strong inference on the bounds of  $\mathcal{P}$ , e.g., via 200 sampling, this will lead to inherent bias on the results of the analysis. 201

#### 202 2.3. Random sets

A p-box can be regarded as a special case of a random set, which has important implications for some of the propagation methods explained in Section 4. To see this, consider a probability space  $(\Omega, \mathcal{F}_{\Omega}, P_{\Omega})$  and a subset  $\mathcal{K}_X$  of the power set of  $D_X \subseteq X$ . A random set  $\Gamma_X$  is then a mapping <sup>206</sup>  $\Gamma_X : \Omega \to \mathcal{K}_X, \alpha \mapsto \Gamma_X(\alpha)$ , where each  $\Gamma_X(\alpha) \in \mathcal{K}_X, \alpha \in \Omega$ , is called a focal element. When distribution-<sup>207</sup> free p-boxes are defined as  $\Gamma_X(\alpha) = \left[\overline{F}_X^{-1}(\alpha), \underline{F}_X^{-1}(\alpha)\right]$  for  $\alpha \in \Omega$  and  $\Omega = [0, 1]$  with uniform probability <sup>208</sup> distribution, they are a specific case of random sets, see [42]. Furthermore note that for finite  $\mathcal{K}_X$ , random <sup>209</sup> sets correspond to a Demspter-Shafer structures, see also [42].

Since a random set is not capable of representing a single parameterized distribution family, a direct relationship with parametric p-boxes cannot be established [43, 44]. Conversion is possible however by first converting the parametric p-box into a distribution-free p-box, see Eq. (3) and (4). Moreover,  $\Gamma_X(\alpha)$ can also be defined directly here via the inverse distributions of the family  $F_X(\cdot, \boldsymbol{\theta}), \boldsymbol{\theta} \in D_{\boldsymbol{\theta}}$ , i.e.,

$$\Gamma_X(\alpha) = \left[\min_{\boldsymbol{\theta}\in D_{\boldsymbol{\theta}}} F_X^{-1}(\alpha, \boldsymbol{\theta}), \max_{\boldsymbol{\theta}\in D_{\boldsymbol{\theta}}} F_X^{-1}(\alpha, \boldsymbol{\theta})\right],\tag{5}$$

 $_{214}$  as shown in [44].

# 215 2.4. Fuzzy probabilities

An extension to the p-box is provided by fuzzy probabilities, which allow for considering a fuzzy set of 216 probability models, each having their own level of plausibility according to the available information [3]. 217 According to this framework, the fuzzy membership function serves as an instrument to combine various 218 plausible intervals  $[\underline{F}_X^{\alpha}(x), \overline{F}_X^{\alpha}(x)], \ \alpha \in [0, 1], \text{ for } x \in D_X \text{ to define distribution-free p-boxes in a single$ 219 scheme, and hence, allows for assessing the sensitivity of the bounds  $\underline{\mathcal{P}}^{\alpha}$  and  $\overline{\mathcal{P}}^{\alpha}$  of  $\mathcal{P}$ . Indeed, sensitivities 220 of  $\mathcal{P}$  are found by considering the rate of change of the bounds on the interval with respect to the size of the 221 input intervals represented in the fuzzy numbers. It holds  $[\underline{F}_X^{\alpha_i}(x), \overline{F}_X^{\alpha_i}(x)] \subseteq [\underline{F}_X^{\alpha_j}(x), \overline{F}_X^{\alpha_j}(x)], x \in D_X$ 222 and therefore  $[\underline{\mathcal{P}}^{\alpha_i}, \overline{\mathcal{P}}^{\alpha_i}] \subseteq [\underline{\mathcal{P}}^{\alpha_j}, \overline{\mathcal{P}}^{\alpha_j}]$  for  $0 \leq \alpha_j \leq \alpha_i \leq 1$ . Furthermore, the concept can be also applied 223 to parametric p-boxes, see [45]. Here, the fuzzy membership function is used to assign an  $\alpha$ -level to the 224 parameters  $\boldsymbol{\theta}$  of  $F_X(\cdot, \boldsymbol{\theta})$ . Then, the same analysis can be conducted as for distribution-free p-boxes. As 225 the methods discussed further in the paper, which are developed for p-boxes, can always be applied to 226 fuzzy probabilities in an  $\alpha$ -cut sense, the latter are not discussed in more detail. 227

# 228 3. Construction of p-boxes for engineering analysis

This section provides an overview how distribution-free and parametric p-boxes can be constructed based on given information. Here, a distinction is made between the three types of information: incomplete or imprecise distribution properties, datasets, or multiple sources of p-boxes. In the following, the focus is put on distribution-free p-boxes first. They are recommended when there is no knowledge in favour of a particular distribution family. If this information is available but the parameters  $\boldsymbol{\theta}$  of  $F_X(\cdot, \boldsymbol{\theta})$  are unknown, parametric p-boxes are preferred. A guide to find an appropriate construction method isprovided in Table 1.

Furthermore note that distribution-free p-boxes can be always constructed as an approximation or an actual conversion of uncertainty models yielding lower and upper probabilities for events  $X \le x$ , see Section 2. For a general introduction on the construction of p-boxes, the reader is referred to [34], where most of the approaches presented in the following are included. A comparison of selected methods can be found, e.g. in [46, 47].

Table 1: Overview of which sections in the paper provide an appropriate p-box construction method depending on the available information and p-box type.

type	distribution-free p-box	parametric p-box	
incomplete distribution information	mean, variance, support: Sec. 3.1	parameters: Sec. 3.4.1	
dataset	Sec. 3.2	Sec. 3.4.2	
multiple sources	Sec. 3.3		

#### 241 3.1. Incomplete distribution properties

In the case that only a limited number of distribution properties are known, like its shape or support, moments, or quantiles, various methods to construct a p-box are available, see [34]. These methods use the information about the distribution properties to derive proper bounds on the distribution. Often, they are based on well-known statistical inequalities. In the following, three methods addressing the support  $D_X$  and the first two moments of a random variable X are presented exclusively. These assume limited but precisely known distribution properties.

# 248 3.1.1. Support

If only the support of a distribution is known, the interval  $D_X = [\underline{x}, \overline{x}]$  can be used as a representation in case the support is bounded. This corresponds to a p-box described by two unit step functions  $H_{\underline{x}}$  and  $H_{\overline{x}}$  at its minimum and maximum values  $\underline{x}$  and  $\overline{x}$ , i.e.,  $\underline{F}_X(x) = H_{\overline{x}}(x)$  and  $\overline{F}_X(x) = H_{\underline{x}}(x)$  for  $x \in D_X$ .

### 252 3.1.2. Mean and variance

If the values of the mean  $\mu_X$  and the variance  $\sigma_X^2$  are known, the two-sided Chebyshev's inequality can be used to construct a p-box as described in [48], i.e.,

$$\underline{F}_X(x) = \begin{cases} 0 & \text{for } x < \mu + \sigma, \\ 1 - \frac{\sigma^2}{(x-\mu)^2}, & \text{for } x \ge \mu + \sigma, \end{cases}$$
(6)

$$\overline{F}_X(x) = \begin{cases} \frac{\sigma^2}{(x-\mu)^2}, & \text{for } x < \mu - \sigma, \\ 1, & \text{for } x \ge \mu - \sigma. \end{cases}$$
(7)

for  $x \in D_X$ . Instead of Chebyshev's inequality, Cantelli's inequality is used to construct a p-box based on the mean and variance in [49].

#### 257 3.1.3. Mean, variance, and support

If both its bounded support  $D_X$  and its first two moments are known, the p-box bounds can be formulated as

$$\underline{F}_{X}(x) = \begin{cases}
0 & \text{for } x \leq \mu + \frac{\sigma^{2}}{\mu - \overline{x}}, \\
1 - \frac{b(1+a)-c-b^{2}}{a} & \text{for } \mu + \frac{\sigma^{2}}{\mu - \overline{x}} < x < \mu + \frac{\sigma^{2}}{\mu - \underline{x}}, \\
\frac{1}{1 + \frac{\sigma^{2}}{(x-\mu)^{2}}} & \text{for } \mu + \frac{\sigma^{2}}{\mu - \underline{x}} \leq x < \overline{x}, \\
1 & \text{for } x \geq \overline{x}, \\
1 & \text{for } x \leq \underline{x}, \\
\frac{1}{1 + \frac{(x-\mu)^{2}}{\sigma^{2}}} & \text{for } \underline{x} < x \leq \mu + \frac{\sigma^{2}}{\mu - \overline{x}}, \\
1 - \frac{b^{2}-ab+c}{1-a} & \text{for } \mu + \frac{\sigma^{2}}{\mu - \overline{x}} < x < \mu + \frac{\sigma^{2}}{\mu - \underline{x}}, \\
1 & \text{for } x \geq \mu + \frac{\sigma^{2}}{\mu - \underline{x}}, \\
\end{cases} \tag{9}$$

where  $a = \frac{x-x}{\overline{x}-\underline{x}}$ ,  $b = \frac{\mu-x}{\overline{x}-\underline{x}}$ ,  $c = \frac{\sigma^2}{(\overline{x}-\underline{x})^2}$ , see e.g. [50]. Eq. (8) and (9) are based on the one-sided Chebyshev's inequalities and are tighter compared to the bounds in Section 3.1.1 and 3.1.2.

#### 262 3.2. Dataset

In case limited information about the probability distribution is available in form of a dataset  $\mathcal{X} \subset$ 263  $\mathbb{R}^{n_x}$ , the properties that are used in the methods of Section 3.1 can be estimated. In order to account for 264 this estimation however, these methods need to be slightly adapted to inform the p-boxes, as described 265 in [51, 49] for the sample mean and sample variance. Moreover, there are also methods which do not 266 require an estimation of distribution properties for a given dataset: the methods of Kolmogorov-Smirnoff 267 confidence bounds and robust Bayes. They are widely used in literature and are described briefly below. 268 Note that data-based methods generally do not provide absolute bounds for p-boxes due to their nature, 269 e.g. by using a confidence level smaller than 1 to avoid conservatism. 270

### 271 3.2.1. Distribution support estimation

In case very few data-points are available, estimating the bounds of the support of the p-box might be the only option for an analyst. This estimation can for instance be based on worst-case likelihood estimation [52], potentially in combination with Bayesian approaches [53]. Scenario optimization [54] can also be used in this context to obtain bounds with a proven degree of robustness under mild assumptions.

# 276 3.2.2. Kolmogorov-Smirnoff confidence bounds

Given a dataset  $\mathcal{X}$  with N samples, an empirical distribution  $F_{\mathcal{X}}$  can be computed. Then, Kolmogorov-Smirnoff (K-S) confidence bounds for  $F_{\mathcal{X}}$  define the bounds of a p-box as proposed in [34]. For  $x \in D_X$ , it holds

$$\underline{F}_X(x) = \min(1, \max(0, F_{\mathcal{X}}(x) - D_N^{\alpha})), \tag{10}$$

$$\overline{F}_X(x) = \min(1, \max(0, F_{\mathcal{X}}(x) + D_N^{\alpha})), \tag{11}$$

where  $D_N^{\alpha}$  is a K-S critical value at significance level  $\alpha$  for a dataset with N samples which can be found in [55].

# 282 3.2.3. Robust Bayes

Furthermore, a p-box can be obtained by using robust Bayes methods, introduced by [56]. Here, the basic idea is to consider the parameters  $\boldsymbol{\theta}$  also as random variables expressed in  $\boldsymbol{\Theta}$  and to apply standard Bayesian inference to all plausible likelihood functions  $L(\cdot, \mathcal{X})$  and all plausible prior distributions. Here,  $f_{\boldsymbol{\Theta}}$  denotes the PDF of the prior and  $L(\cdot, \mathcal{X})$  is the likelihood of observing  $\mathcal{X}$  depending on the incorporated distribution family  $F_X(\cdot, \boldsymbol{\theta})$ . This implies a class of posterior PDFs of  $\boldsymbol{\Theta}$ , denoted by  $f_{\boldsymbol{\Theta}}(\cdot|\mathcal{X})$ , via Bayes theorem

$$f_{\Theta}(\boldsymbol{\theta}|\mathcal{X}) = \frac{L(\mathcal{X}, \boldsymbol{\theta})}{\int_{D_{\Theta}} L(\mathcal{X}, \boldsymbol{\theta}) f_{\Theta}(\boldsymbol{\theta}) \,\mathrm{d}\boldsymbol{\theta}} f_{\Theta}(\boldsymbol{\theta})$$
(12)

and pairwise combination. Then, a p-box can be constructed by the envelope of all resulting CDFs using
Bayesian point estimates, see [57], or credible intervals/regions like discussed in Section 3.4.2, see [46].
Moreover, a Bayesian pointwise approach that considers specific percentiles of all resulting CDFs can be
used for the construction of a p-box as well, see [58].

# 293 3.3. Aggregation of p-boxes

In the methods above, the intention was to obtain a p-box based-on given information. If there are already  $n_{\rm p}$  p-boxes  $[\underline{F}_X^{(j)}, \overline{F}_X^{(j)}]$  available to describe a single quantity, aggregation methods can be used. In the following, three popular methods, namely the envelope, intersection, and mixture strategy, are reviewed. For further methods, the reader is once again referred to [34].

#### <sup>298</sup> 3.3.1. Envelope and intersection

If there are multiple p-boxes of which it at least one encompasses the unknown CDF of X, but there is no information which p-boxes really encompass it, the envelope strategy can be used. Here, an envelope 301 p-box is defined as

$$\underline{F}_X(x) = \min\{\underline{F}_X^{(j)}(x) \mid j = 1, \dots, n_{\rm s}\},\tag{13}$$

$$\overline{F}_X(x) = \max\{\overline{F}_X^{(j)}(x) \mid j = 1, \dots, n_{\rm s}\}$$
(14)

for  $x \in D_X$ . This corresponds to a conservative modelling. Opposite to the envelope strategy, there is the intersection strategy for which all available p-boxes are considered as reliable. Here, the intersection of all p-boxes is used, see [34]. For this strategy, the min and max operators in Eq. (13) and (14) are exchanged.

#### 306 3.3.2. Mixture

If there are multiple p-boxes which were constructed for specific situations that suffer under variability, the mixture strategy can be used for the condensation in a single p-box. Here, the idea is to use weights  $w_j > 0$  with  $W = \sum_{j=1}^{n_s} w_j$  to express the relative frequencies. Then, the mixture p-box is defined as

$$\underline{F}_X(x) = \frac{1}{W} \sum_{j=1}^{n_{\rm s}} w_j \underline{F}_X^{(j)}(x), \tag{15}$$

$$\overline{F}_X(x) = \frac{1}{W} \sum_{j=1}^{n_{\rm s}} w_j \overline{F}_X^{(j)}(x)$$
(16)

for  $x \in D_X$ . A special case are even weights, e.g.,  $w_j = 1, j = 1, ..., n_s$  with  $W = n_s$ , which correspond to an arithmetic averaging of the p-boxes.

#### 312 3.4. Parametric p-box construction

In order to construct a parametric p-box, the distribution family must be known. Hence, the problem of constructing a p-box reduces to establishing bounding intervals for the corresponding parameters  $\boldsymbol{\theta}$  of  $F_X(\cdot, \boldsymbol{\theta})$ . Usually, these intervals are assumed or estimated for a given dataset, see the methods below. Note that all methods to obtain a parametric p-box can be also used to build a distribution-free p-box by using Eq. (3) and (4) which yield the envelope of the parametric p-box.

#### 318 3.4.1. Bounds on distribution parameters

In case bounds on the parameters  $\theta$  are available, e.g., from expert knowledge, the intervals for these parameters can be specified directly. For lower bounds  $\underline{\theta}_i$  and an upper bounds  $\overline{\theta}_i$ ,  $i = 1, \ldots, n_{\theta}$ their domain is denoted by  $D_{\theta}$  (see Section 2.1). If there are  $n_{\rm s}$  sources that provide different intervals, aggregation methods similar to Section 3.3 could be used, e.g., an envelope of all candidate domains  $D_{\theta}^{(j)}$ , where  $\underline{\theta}_i = \min{\{\underline{\theta}_i^{(j)} \mid j = 1, \ldots, n_{\rm s}\}}$  and  $\overline{\theta}_i = \max{\{\overline{\theta}_i^{(j)} \mid j = 1, \ldots, n_{\rm s}\}}$ ,  $i = 1, \ldots, n_{\theta}$ .

# 324 3.4.2. Dataset

Given a dataset  $\mathcal{X}$ , there are several methods to obtain interval estimates for the parameters  $\theta$ of  $F_X(\cdot, \theta)$ . Popular methods comprise confidence intervals from classical statistics, which cover the unknown, but deterministic parameters with a probability  $\alpha$ , or credible intervals from Bayesian statistics, in which the random vector  $\Theta$ , representing the parameters of the CDF, can be found with a probability  $\alpha$ , see [59] for further information on their computation. Note that in general independence between the parameters  $\theta$  needs to be assumed for  $n_{\theta} > 1$  in order to obtain interval regions.

# 331 4. Propagation methods for p-boxes

This section discusses commonly applied numerical schemes for propagating p-boxes towards bounds on the  $n^{th}$  central moment of the model response to a load and/or the probability of failure of the designed structure, system or complex network. In the case where X is represented as a p-box, a direct calculation of  $\mathcal{P}$ , as introduced in Eq. (2), is no longer possible since a set of PDFs that are consistent with the definition of the p-box has to be considered. Indeed, the consideration of a set of  $f_X$  causes the probabilistic measure  $\mathcal{P}$  to become set-valued, too. The solution of this problem requires dedicated numerical procedures, which are described in the proceeding sections.

#### 339 4.1. Double loop approaches

In case X represents a distribution-free p-box, the lower and upper bounds  $\underline{\mathcal{P}} \leq \mathcal{P} \leq \overline{\mathcal{P}}$  can be obtained by solving the following optimization problems:

$$\underline{\mathcal{P}} = \min_{f_{\mathbf{X}}} \int_{D_{X}} \mathcal{H}(\mathbf{x}) f_{\mathbf{X}}(\mathbf{x}) \,\mathrm{d}\mathbf{x}$$
(17)

342 and:

$$\overline{\mathcal{P}} = \max_{f_{\mathbf{X}}} \int_{D_{X}} \mathcal{H}(\mathbf{x}) f_{\mathbf{X}}(\mathbf{x}) \, \mathrm{d}\mathbf{x}.$$
(18)

Note that these optimization problems are potentially very complicated since the optimization has to be carried out over the set of all possible  $f_X$  consistent with the definition of the p-box. Hence, this constitutes a non-convex, discontinuous optimization problem, which are notoriously difficult so solve exactly. In certain cases, tighter bounds on  $\mathcal{P}$  can be obtained by means of linear programming, without having to construct the probability bounds of the input random variables [60, 61].

A first approach to simplify the optimization problems is to *slice* the p-box in order to transform the above problem into the propagation of a large number of intervals, each having a corresponding probability mass, which are then propagated through  $\mathcal{M}$  to infer bounds on  $\mathcal{P}$ . The propagation of intervals is a well-understood problem in the context of uncertainty propagation [2]. However, following this approach the required number of evaluations of Eq. (1) scales exponentially with  $n_x$  [62]. This led to the development of methods such as interval Monte Carlo simulation [62] or interval-Quasi Monte Carlo simulation [63]. These methods manage to break this exponential scaling of the computational cost by bounding  $\mathcal{P}$  using following formulations:

$$\underline{\mathcal{P}} = \frac{1}{n} \sum_{k=1}^{n} \overline{\mathcal{H}}(\boldsymbol{r}_k), \tag{19}$$

$$\overline{\mathcal{P}} = \frac{1}{n} \sum_{k=1}^{n} \underline{\mathcal{H}}(\mathbf{r}_k) \tag{20}$$

with  $\overline{\mathcal{H}}(\boldsymbol{r}_k)$  and  $\underline{\mathcal{H}}(\boldsymbol{r}_k)$  defined as:

$$\overline{\mathcal{H}}(\boldsymbol{r}_k) = \max\{\mathcal{H}(\boldsymbol{x}) \mid \overline{F}_{\boldsymbol{X}}^{-1}(\boldsymbol{r}_k) \le \boldsymbol{x} \le \underline{F}_{\boldsymbol{X}}^{-1}(\boldsymbol{r}_k)\},$$
(21)

$$\underline{\mathcal{H}}(\boldsymbol{r}_k) = \min\{\mathcal{H}(\boldsymbol{x}) \mid \overline{F}_{\boldsymbol{X}}^{-1}(\boldsymbol{r}_k) \le \boldsymbol{x} \le \underline{F}_{\boldsymbol{X}}^{-1}(\boldsymbol{r}_k)\}.$$
(22)

The parameters  $\mathbf{r}_k, j = 1, \dots, N$  are realisations of a sample of N independent and identically dis-357 tributed (i.i.d.) random variables according to a multivariate standard uniform distribution. As is clear 358 from these equations, a large number of model evaluations is still required to estimate of the bounds on 359  $\mathcal{P}$  with sufficiently small variance, especially since an interval propagation problem (Eq. (21)) has to be 360 solved for each  $r_k$ . Note that in the general case, this interval problem has to be solved using global 361 optimization approaches to accommodate possible non-convexity in  $\mathcal{M}$  with respect to x and/or  $\theta$  [64]. 362 Further improvement in computational efficiency can be obtained by resorting to efficient interval prop-363 agation schemes such as those based on Bernstein polynomials [65, 66], Cauchy deviates [28] (as recently 364 applied in [67] and [68]), the transformation method [69] or Taylor series expansion methods [70, 71]. 365 Further improvements in terms of efficiency can be obtained by using saddle-point approximations, as 366 introduced in [72]. A more general version of the interval Monte Carlo approach was introduced by 367 Alvarez in [42, 73] based on random sets (see also subsection 2.3). The main advantage of considering 368 the full random set is that this representation is more general, and hence, intervals and Dempster-Shafer 369 structures can be considered as well in the same framework [74]. Furthermore, the framework allows for 370 including efficient sampling schemes, such as e.g., subset simulation [74]. 371

In the case of parametric p-boxes, the extrema represented by Eq.(17) and Eq.(18) can be determined directly since the set of all possible  $f_{\mathbf{X}}$  is readily parameterized. In this case, for each realisation of these parameters of  $f_{\mathbf{X}}$ , a reliability problem is solved, for instance for linear limit-state functions using FORM as presented in [75], or in more general cases using simulation methods. Using simulation methods, even in the simplest case where the p-box describes a set of possible  $f_{\mathbf{X}}$  by means of interval-valued statistical

moments, such calculation can be prohibitively demanding from a numerical standpoint. On the one 377 hand, the calculation of the failure probability for a fixed value of the parameters associated with the 378 stochastic process is quite costly. On the other hand, solving the associated optimization problems 379 in this simple case is far from trivial, as it constitutes a double loop problem, where the inner loop 380 comprises probability estimations, leading to possibly non-smooth behaviour of the objective function 381 due to the inherent variance on the estimator of  $\mathcal{P}$ . Hence, apart from considering near-trivial simulation 382 models, the propagation of p-box-valued parameters towards the bounds on the probabilistic measure 383  $\mathcal{P}$  is computationally intractable. Note that in some very specific cases, analytical solutions are also 384 available [76]. 385

#### 386 4.2. Decoupling methods

The class of decoupling methods aims at decoupling the double loop, presented in Eqs. (17) and (18) by separating the propagation of aleatory and epistemic uncertainties. This class of methods includes techniques based on importance sampling and operator norm theory. Both methods are restricted to parametric p-boxes, more precisely, p-boxes that are constructed by defining some parameters  $\boldsymbol{\theta}$  of the distribution  $F_{\boldsymbol{X}}(\boldsymbol{x}|\boldsymbol{\theta})$  to be interval valued.

# 392 4.2.1. Importance sampling-based methods

The core idea of importance sampling based methods is to propagate a single, well-chosen realisation  $\hat{f}_{X}$  of a parameterized p-box (where  $\hat{f}_{X}$  is optimal with respect to a predefined measure), and reweigh the obtained samples of y to infer bounds on  $\mathcal{P}$ .

A first such method is Extended Monte Carlo simulation, as introduced by [39], which is applicable to the propagation of parameterized p-boxes subjected to epistemic uncertainty in their first two moments, as well as the probability of failure. As a first step, the parameters  $\boldsymbol{\theta}$  of the p-box, which account for  $\mu_x$  and  $\sigma_x$  in the quintuple description, are represented by a subjective probability model  $f_{\Theta}(\boldsymbol{\theta}) = \prod_{i=1}^{n_{\theta}} f_{\Theta_i}(\theta_i)$ . Then, a local estimation for  $\mathcal{P}$ , being  $\hat{p}_{\rm f}$ , is derived as:

$$\hat{\mathcal{P}}(\boldsymbol{\theta}) = \frac{1}{N} \sum_{k=1}^{N} \mathcal{H}(\boldsymbol{x}_k) \frac{f_{\boldsymbol{X}}(\boldsymbol{x}_k \mid \boldsymbol{\theta})}{f_{\boldsymbol{X}}(\boldsymbol{x}_k \mid \boldsymbol{\theta}^*)}$$
(23)

which is an unbiased estimator, but highly affected by the selection of  $\theta^*$ . 'Local' in this context denotes that the estimator is derived for a fixed value of  $\theta$  inside its support  $\theta^I$ . This fixed value,  $\theta^*$ , should be selected such that it minimizes the variance on the estimator  $\hat{\mathcal{P}}(\theta)$  [77], similarly to conventional Importance Sampling, as:

$$\boldsymbol{\theta}^* = \operatorname{argmin} \int_{D_{\boldsymbol{\theta}}} T(\boldsymbol{\theta}, \boldsymbol{\theta}^*) f_{\boldsymbol{\Theta}}(\boldsymbol{\theta}) \,\mathrm{d}\,\boldsymbol{\theta}$$
(24)

with  $T(\boldsymbol{\theta}, \boldsymbol{\theta}^*) = V [\mathcal{H}(\boldsymbol{X}) f_{\boldsymbol{X}}(\boldsymbol{X} \mid \boldsymbol{\theta}) / f_{\boldsymbol{X}}(\boldsymbol{X} \mid \boldsymbol{\theta}^*)]$  and V is the variance operator with respect to  $f_{\boldsymbol{X}}(\cdot \mid \boldsymbol{\theta}^*)$   $\boldsymbol{\theta}^*$ ). The global version of this approach is based on realizations  $(\boldsymbol{x}_k, \boldsymbol{\theta}_k), k = 1, \dots, N$  of a joint sample distributed according to a joint PDF  $f_{\boldsymbol{X},\boldsymbol{\Theta}}$ . The estimator  $\hat{\mathcal{P}}(\boldsymbol{\theta})$  is in this case expressed as:

$$\hat{\mathcal{P}}(\boldsymbol{\theta}) = \frac{1}{N} \sum_{k=1}^{N} \mathcal{H}(\boldsymbol{x}_k) \frac{f_{\boldsymbol{X}}(\boldsymbol{x}_k \mid \boldsymbol{\theta})}{f_{\boldsymbol{X}}(\boldsymbol{x}_k \mid \boldsymbol{\theta}_k)}$$
(25)

where  $x_k$  and  $\theta_k$  are generated by applying the correct inverse probabilistic transform to the corresponding variables of a multivariate standard uniform distribution. The global estimator gives a better estimation of  $\mathcal{P}$  over the entire support of  $\theta$ , at the cost of lower accuracy around  $\theta^*$  and a higher computational cost, since in this case, also convergence in terms of the effect of  $\theta$  has to be ensured.

An alternative optimal sampling density to propagate parameterized p-boxes following a reweighted sampling scheme was proposed by [78, 79]. Following the approach of [78, 79], the optimal density should obtained by minimizing the total expected squared Hellinger distance between  $f_{\mathbf{X}}(\cdot \mid \boldsymbol{\theta})$  and the optimal sampling density  $f_{\mathbf{X}}(\cdot \mid \boldsymbol{\theta}^*)$  under an isoperimetric constraint that ensures that the derived optimal sampling density is a valid density function. The main difference with optimal sampling density presented in Eq. (24) is that this approach is not aimed at minimizing the variance, but rather that the sampling density is a solution to the target density.

# 419 4.2.2. Advanced Line Sampling

As an alternative decoupling strategy to deal with p-box valued uncertainty, Advanced Line Sampling 420 was recently introduced [80]. Opposed to 'conventional' line sampling [81], this approach adaptively 421 looks for the so-called important direction in standard normal space. Furthermore, due to this adaptive 422 refinement, the same important direction can be used for the entire p-box analysis. Additionally, the 423 method allows for reusing samples that are generated within the inner loop to be re-used during other 424 iterations of the outer loop, significantly increasing the computational efficiency [80]. Based on these 425 properties, a gain in computational efficiency of a factor of 4 over regular line-sampling approaches can 426 be obtained, as reported in [80]. 427

#### 428 4.2.3. Operator norm theory

Operator norm theory provides an alternative pathway to decouple the double loop in Eq. (17) and (18), as presented first in [82] for the case of the class of linear models  $\mathcal{M}$ . In case an affine formulation of the imprecise random variables in terms of their parameters is possible, the propagation of the imprecise stochastic load can be performed in a two-step procedure. First, the values of the epistemic parameters that yield an extremum for  $\mathcal{P}$  are determined by maximizing/minimizing the operator norm. Specifically, the operator norm is computed over the product of the linear mapping provided by the numerical model  $\mathcal{M}$  with a basis  $\boldsymbol{B}$  that represents the auto-correlation of the load on the model:

$$\boldsymbol{\theta}^{\underline{*}} = \underset{\boldsymbol{\theta} \in \boldsymbol{\theta}^{I}}{\operatorname{argmin}} \|\boldsymbol{A}(\boldsymbol{\theta})\|_{p^{(1)}, p^{(2)}}$$
(26)

436

$$\boldsymbol{\theta}^{\overline{*}} = \underset{\boldsymbol{\theta} \in \boldsymbol{\theta}^{I}}{\operatorname{argmax}} \|\boldsymbol{A}(\boldsymbol{\theta})\|_{p^{(1)}, p^{(2)}}$$
(27)

with  $A = \mathcal{M}B$ , where B can for instance be determined following the well-known Karhunen-Loève expansion [82]. The operator norm  $||A||_{p^{(1)},p^{(2)}}$  is generally defined as [83]:

$$||\boldsymbol{A}||_{p^{(1)},p^{(2)}} = \inf\left\{c \ge 0 : ||\boldsymbol{A}\boldsymbol{v}||_{p^{(1)}} \le c \cdot ||\boldsymbol{v}||_{p^{(2)}} \quad \forall \boldsymbol{v} \in \mathbb{R}^{n_{KL}}\right\},\tag{28}$$

and gives a measure for how much A lengthens vector v in the maximum case. The practical calculation of 439 the operator norm is case dependent. For instance, when considering first excursion problems [84, 85, 86], 440 i.e.,  $\mathcal{P} \equiv p_{\rm f}$ , the selection of  $p^{(1)} \to \infty$  and  $p^{(2)} = 2$  has been found to be a good choice [87]. In this case, the 441 operator norm corresponds to the maximum  $\mathcal{L}_2$  norm of a row of A [83]. Then, two failure probabilities, 442 corresponding to  $p_{\rm f}(\theta^{\underline{*}})$  and  $p_{\rm f}(\theta^{\overline{*}})$  have to be computed to determine the bounds on  $\mathcal{P}$ . As such, the 443 double loop is effectively replaced by two deterministic optimizations and two crisp reliability estimations. 444 Gains in computational efficiency with several orders of magnitude have been reported [82, 87]. The main 445 drawback of the method is the limited scope, since the approach is only applicable to uncertain linear 446 models with epistemic uncertain structural parameters, subjected to imprecisely defined load conditions. 447

# 448 4.3. Surrogate modelling for p-boxes

Surrogate models approximate well-selected 'regions' of  $\mathcal{M}$  by a computationally more efficient sur-449 rogate model  $\hat{\mathcal{M}}(\cdot \mid \boldsymbol{a})$ . For instance, in the specific case of reliability analysis,  $\hat{\mathcal{M}}(\cdot \mid \boldsymbol{a})$  is designed 450 to be highly accurate in the region around the limit state function (i.e.,  $\mathcal{M}(\mathbf{x}) = 0$ . This surrogate 451  $\mathcal{M}$ , which is parameterized by a vector  $\boldsymbol{a} \in \mathbb{R}^{n_a}$ , is usually *trained* by means of a set of training data 452  $\{(\boldsymbol{x}_i, \boldsymbol{y}_i) \mid i = 1, ..., N\}$  via a supervised learning approach as to minimize the discrepancy between 453  $\hat{y}_i = \hat{\mathcal{M}}(x_i \mid a)$  and  $y_i$ , according to a predefined measure (e.g., in an  $L_2$  sense). These training data 454 are generated either a priori (e.g., in case of sensitivity analysis) or enriched following active learning 455 approaches [88, 89], which is most commonly applied in the field of reliability analysis. Examples of 456 such maps to represent  $\hat{\mathcal{M}}$  that have been used in the context of propagating p-boxes include Gaus-457 sian process models [90] (also known as Kriging), polynomial response surface models [91] or techniques 458 based on Taylor series expansions [92]. Also adaptive schemes based on Kriging have been introduced 459 in literature [57] that are applicable to both parametric and distribution-free p-boxes. In this section, 460

three classes of methods are explained in detail that are highly promising from an engineering point of view due to their 'black-box' nature (i.e., they require no interaction with the inner operations of  $\mathcal{M}$ ), theoretical implications and numerical efficiency. Note that in essence, each type of surrogate model can be used in combination with a double-loop approach since they are very cheap to evaluate. The selection of the appropriate type of surrogate model in fact only depends on  $\mathcal{M}$ .

#### 466 4.3.1. Polynomial Chaos Expansions & Kriging models

Polynomial chaos expansion (PCE) and Kriging are two widely used surrogate modelling techniques that approximate  $\mathcal{M}$  via intricate regression schemes. In general, PCE and Kriging have different fields of application in the propagation of uncertainties. On the one hand, if the analyst is interested in propagating uncertainty in general (e.g., when  $\mathcal{H}(\boldsymbol{x}) = (\boldsymbol{y} - \boldsymbol{\mu}_{\boldsymbol{Y}})^n$ ) PCE generally is better suited. Conversely, when considering reliability analysis, Kriging approaches are generally more performing since they allow for performing active learning [89, 93, 94], even though active learning approaches for PCE have also been introduced [95].

474 A sparse PCE surrogate model is given by:

$$\hat{\mathcal{M}}(\boldsymbol{x} \mid \boldsymbol{a}) = \sum_{\alpha \in \mathcal{A}} a_{\alpha} \boldsymbol{\phi}_{\alpha}(\boldsymbol{x}),$$
(29)

where  $\phi_{\alpha}$  are multivariate orthonormal polynomials and  $\mathcal{A} \subset \mathbb{N}^{n_x}$  is a finite set of multi-indices that is obtained by sparse decomposition. In [96], distribution-free p-boxes are propagated in a two-level approach in which first  $\mathcal{M}$ , and second  $\mathcal{M}$  and  $\overline{\mathcal{M}}$  (in the sense of Eq (21) and (22)) are substituted using sparse PCE. The training set is generated for an auxiliary input vector X and least angle regression (LARS) is used for training. In case of parametric p-boxes, it is proposed in [97] to model the sparse PCE coefficients  $a_{\alpha}$  as quadratic polynomial functions of the parameters  $\boldsymbol{\theta}$  of the p-box and using a double-loop sampling for the propagation.

Whereas PCE methods focus on the global behaviour of  $\mathcal{M}$  and are therefore suitable for a general propagation of p-boxes, Kriging methods focus on a local behaviour of  $\mathcal{M}$  and are therefore often preferred for reliability analysis. Indeed, in this context, a high accuracy in the vicinity where { $\mathcal{M} = 0$ } is especially crucial. Using Kriging, a surrogate  $\hat{\mathcal{M}}$  for the limit-state function is considered to be a realization of a Gaussian process. It is:

$$\hat{\mathcal{M}}(\boldsymbol{x} \mid \boldsymbol{a}) = \boldsymbol{\beta}_{\boldsymbol{a}}^{\mathrm{T}} \boldsymbol{\psi}(\boldsymbol{x}) + Z_{\boldsymbol{a}}(\boldsymbol{x}, \omega), \qquad (30)$$

where the first term, consisting of coefficients  $\beta_a$  and regression functions  $\psi$ , is the mean value of the process, and the second term is a zero-mean, stationary Gaussian process, characterized by a variance and an auto-correlation function depending on a. Similar to above, a two-level approach in which first  $\mathcal{M}$ , and second  $\underline{\mathcal{M}}$  and  $\overline{\mathcal{M}}$  are substituted is considered for distribution-free p-boxes in [57]. Here, adaptive Kriging Monte Carlo simulation (AK-MCS) is used for an accurate estimation of the failure probabilities and random slicing is used to obtain  $\underline{\mathcal{P}}$  and  $\overline{\mathcal{P}}$ , see Eq. (19) and (20). Also in [57], a failure probability  $\mathcal{P}(\theta)$  which depends on the parameters  $\theta$  is estimated via AK-MCS and efficient global optimization (EGO) for parametric p-boxes. A similar, but more detailed, Kriging-based procedure for parametric p-boxes is also described in [98].

# 496 4.3.2. High-dimensional model representation based methods

The Extended Monte Carlo framework, as introduced in Section 4.2.1 allows for propagating parametrized p-boxes by a single probabilistic simulation and a reweighting step. Nonetheless, still a considerable number of evaluations of  $\mathcal{M}$  are required, which might impede practical applications. Therefore, in [39], both the local and global Extended Monte Carlo methods were integrated with a high-dimensional model representation (HDMR) decomposition of  $\mathcal{M}$  as a surrogate modelling strategy. Following a HDMR decemposition,  $\mathcal{P}$  can be represented as:

$$\mathcal{P}(\boldsymbol{\theta}) = p_{\mathrm{f},0} + \sum_{i=1}^{n_{\theta}} p_{\mathrm{f},i}(\theta_i) + \sum_{1 \le i < j \le d} p_{\mathrm{f},ij}([\theta_i, \theta_j]) + \dots + p_{\mathrm{f},12\dots n_{\theta}}(\boldsymbol{\theta}).$$
(31)

Note that HDMR decompositions are more widely applicable than to represent  $\mathcal{P}$ . In the context 503 of propagating p-boxes, in [39], it is proposed to apply a cut-HDMR strategy in combination with 504 the local Extended Monte Carlo Method, allowing for a rigorous estimation of the variances of the 505 estimators, as well as an estimation of the sensitivity of the parameters in  $\theta$ . Similarly, it is proposed 506 to perform a Random Slicing HDMR decomposition in combination with the Global Method. For the 507 details concerning the implementation of these techniques, as well as the corresponding proofs, the reader 508 is referred to [39]. These methods were recently also extended to be applied in combination with Line 509 Sampling in [99]. 510

An alternative application of the Sobol-Hoeffding decomposition in the context of propagating imprecise probabilities through numerical models is given by [100]. In [100], the authors apply a fuzzy probabilistic approach in the study of designing cylindrical shells under geometric imperfections, which are modelled as a random field. Specifically, imprecision in the auto-correlation structure of the random field is accounted for by means of fuzzy arithmetic, and the S-H decomposition is applied to speed up the corresponding  $\alpha$ -level optimization.

# 517 4.3.3. Interval predictor models

An interval predictor model (IPM), as introduced in [101], is a type of surrogate model that approximates  $\mathcal{M}$  by means of an interval-valued map  $\hat{\mathcal{M}}_I(\cdot, \boldsymbol{\theta}) : \mathbb{R}^{n_x} \to \mathbb{IR}$ , where  $\mathbb{IR}$  is the set of all intervals in  $\mathbb{R}$ . This map can be constructed with a minimal number of assumptions on the mapping provided by  $\mathcal{M}_I(\boldsymbol{x}, \boldsymbol{\theta})$  given by:

$$\hat{\mathcal{M}}_{I}(\boldsymbol{x},\boldsymbol{\theta}) = \left\{ y = \boldsymbol{\theta}^{\mathrm{T}} \boldsymbol{\phi}(\boldsymbol{x}) \mid \boldsymbol{\theta} \in \boldsymbol{\theta}^{I} \right\}$$
(32)

with  $\phi$  a basis (e.g., polynomial or trigoniometric),  $\theta$  the fitting parameters of the IPM and  $\theta^{I} = [\underline{\theta}, \overline{\theta}]$  an  $n_{\theta}$ -dimensional hyper-rectangular set. An optimal IPM is constructed by minimizing  $E[(\overline{\theta} - \underline{\theta})|\phi(x)|]$ . Scenario Optimization [54] can be used to judge the generalization properties of the IPM. In case the corresponding optimization problem is convex, the reliability R of the IPM (i.e., the probability that a future unobserved data point will be contained in the IPM) is bounded by:

$$P(R \ge 1 - \epsilon) > 1 - \beta, \tag{33}$$

where  $\epsilon$  and  $\beta$  are the confidence and reliability parameters, which for our hyper-rectangular model can be obtained from

$$\beta \ge \binom{k+n_{\theta}-1}{k} \sum_{i=0}^{k+n_{\theta}-1} \binom{N}{i} \epsilon^{i} (1-\epsilon)^{N-i}, \tag{34}$$

where k is the number of data points discarded by some algorithm and  $\epsilon$  can be chosen as a very small 529 number (e.g.,  $\epsilon = 1 \cdot 10^{-06}$ ). An approach to apply IPMs in the context of propagating parametrized 530 p-boxes is introduced by [102]. They show that IPMs can be used as surrogate model to speed up the 531 calculation of Eq.(17) and Eq.(18), including a strategy to intelligently construct the set  $\{(x_i, y_i) \mid i =$ 532 1, ..., N. Furthermore, they show that the IPM can also be used as a surrogate model for g, which in 533 its turn can be used in combination with importance sampling to determine  $[\underline{\mathcal{P}}, \overline{\mathcal{P}}]$ . Other applications 534 include estimating the bounds on  $\mathcal{P}$  resulting from the surrogate model inaccuracy in a deterministic 535 case [103]. 536

The main advantages of these techniques are that (1) they are completely black-box as they don't require any assumption on  $\mathcal{M}$  and (2) that under the mild assumption of convexity of the training guaranteed reliability bounds on the accuracy are obtained based on the rigorous framework of Scenario Optimization, which was recently extended to non-convex optimization problems too [54]. Unfortunately, active learning of this type of surrogate models is not feasible, since this violates the required assumptions on independence between the training samples [103].

# 543 4.4. Concluding discussion

As an attempt to create some clarity in the applicability of the multitude of available methods for 544 the propagation of p-boxes, Table 2 summarizes the discussed methods, including their class, limitations 545 and to which type of p-box they are applicable. Note that no precise statements on accuracy and/or 546 numerical efficiency are given, as these depend fully on the problem under consideration. For instance, for 547 linear models, the operator norm will undoubtedly give the best results from all 'direct' solution methods, 548 as it reduces the solution of the problem to two deterministic optimization problems and two reliability 549 analyses. On the other hand, for highly nonlinear problems, this method will fail, and potentially methods 550 based on surrogate modelling will outperform the other methods. To make a fully fair comparison between 551 these methods in this respect, a dedicated benchmark is study is required, which falls outside the scope of 552 this paper. It should be noted, however, that in case there is no prerogative to use the numerical model, 553 the computational efficiency of propagating imprecise probabilities with surrogate modelling approaches 554 is orders of magnitude higher as compared to the approaches that directly use the numerical model. 555 This is particularly true when advanced active learning methods such as AK-MCS [89] are applied in the 556 context of reliability analysis. 557

Method	class	Type p-box	Limitation	ref
Double loop	Direct	Both	Computational cost	
Interval Monte Carlo	Direct	Free	Computational cost	[62]
Random set methods	Direct	$\mathrm{Both}^1$	Computational cost	[74]
Advanced Line Sampling	Decoupling	Param.	Moderate linearity	[80]
Extended Monte Carlo	Decoupling	Param.	Stochastic hyper-parameters	[39]
Operator norm	Decoupling	Param.	Linear models	[82]
PCE	Surrogate	$\operatorname{Both}$	Global approximation of $\mathcal{M}$	[57]
Kriging	Surrogate	$\operatorname{Both}$	Local approximation of $\mathcal{M}$	[104]
HDMR	Surrogate	Param.	Dimension of $\boldsymbol{x}$	[100]
IPM	Surrogate	Param.	No adaptive refinement	[102]

Table 2: Summary of black-box propagation schemes for p-boxes

Generally, optimization approaches such as double loop or sampling methods provide inner approximations of the bounds on  $P_f$  as they generate realisations within  $[\underline{F}_X(x), \overline{F}_X(x)]$  and try to approach  $\mathcal{P}$ , respectively  $\overline{\mathcal{P}}$  from the inside-out [43]. Note that, in case distribution-free p-box methods such as those based on random sets are applied to parametric p-boxes, this effectively constitutes an outer approximation.

<sup>&</sup>lt;sup>1</sup>More general imprecise probability models can be considered too, please refer to Section 2.3 for more information

### 563 5. Conclusions

The development of highly efficient approaches to perform engineering computations with imprecise probabilities, represented as p-boxes, is a quickly expanding field of research. The main challenge in this context is to overcome the required double loop propagation framework to estimate the bounds on probabilistic measures of the structure under consideration (such as, e.g., the probability of failure). Apart from near-trivial numerical simulation models, such double loop calculations are computationally intractable without resorting to high-performance computing facilities.

This problem is currently being tackled from two sides: (1) by improving the propagation efficiency of 570 p-boxes aimed at breaking the double loop and (2) developing efficient surrogate models for the numerical 571 models to be used in the double loop. Concerning the former set of solutions, highly efficient propagation 572 schemes have been introduced in recent years. However, these methods are either limited in terms of 573 the admissible descriptions of the uncertainty, or the non-linearity of the underlying numerical model. 574 Future developments in these areas should concentrate on expanding the scope of applicability of these 575 techniques. Concerning the latter, surrogate models usually only require some smoothness constraints 576 on the underlying numerical model, which allows for a greater flexibility. Nonetheless, the accuracy of 577 the calculation of the bounds on the probabilistic measures is limited to the accuracy of the underlying 578 surrogate model. Furthermore, also the training of these surrogate models can entail a non-negligible 579 numerical cost, which is commonly mitigated by resorting to active learning. 580

As such, to conclude, the last 15 years brought many highly performing approaches to compute with 581 imprecise probabilities in general, and p-boxes in specific. The main challenge at this point appears 582 to translate this set of highly performing methods to industrial applications involving multi-physical 583 and/or million degree-of-freedom numerical models. Further work in this domain will include continuing 584 developments on a theoretical side (e.g., constrained distribution-free p-boxes) and propagation aspects 585 (e.g., operator norm theory for nonlinear dynamics). We expect that the current rapid developments 586 in the domain of machine learning and big data can play a pivotal role in (1) the characterization of 587 uncertainties, where the uncertainty characteristics are added by the machine learning algorithm, (2) 588 the propagation and inverse identification of p-boxes, much alike active learning surrogate models, (3) 589 performing dimension reduction by finding optimal representations of the uncertainty and (4) detecting 590 dependencies in very high-dimensional datasets. 591

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