ncertainty Modeling and Propagation for
oundwater Flow: A Comparative Study of
Surrogates
Dedicated to the memory of K. Andrew Cliffe (1953–2014)
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Abstract
e compare sparse grid stochastic collocation and Gaussian pro- ss emulation as surrogates for the parameter-to-observation map a groundwater flow problem related to the Waste Isolation Pilot ant in Carlsbad, NM. The goal is the computation of the prob- ility distribution of a contaminant particle travel time resulting om uncertain knowledge about the transmissivity field. The lat- r is modelled as a lognormal random field which is fitted by stricted maximum likelihood estimation and universal kriging to obser- tional data as well as geological information including site-specific end regression functions obtained from technical documentation. The sulting random transmissivity field leads to a random groundwater w and particle transport problem which is solved realization-wise ing a mixed finite element discretization. Computational surrogates, and constructed, allow sampling the quantities of interest in the certainty analysis at substantially reduced computational cost. Spe- al emphasis is placed on explaining the differences between the

047two surrogates in terms of computational realization and interpreta-048tion of the results. Numerical experiments are given for illustration.

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 Keywords: sparse grid stochastic collocation, Gaussian process emulation, uncertainty propagation, kriging, Darcy flow, mixed finite elements

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057 1 Introduction

058By their very nature, the Earth Sciences have had to cope with uncertainty 059 from early on, and scientists from this field such as Harold Jeffreys and Albert 060 Tarantola have had foundational and lasting impact on how uncertainty is 061 modeled and merged with physical models in the interdisciplinary field now 062known as uncertainty quantification (UQ). A current account of uncertainty 063 quantification in subsurface hydrology can be found in Linde et al (2017). Many 064 UQ studies involve a system governed by a partial differential equation (PDE) 065in which one or more input quantities are uncertain. When this uncertainty 066 is described in probabilistic terms, we arrive at a PDE with random data, or 067 random PDE for short, the solutions of which are stochastic processes, also 068 referred to in this context as random functions or random fields. The task of 069 determining the probability distribution of the solution of a random PDE, or 070 of quantities of interest derived from such solutions, is known as uncertainty 071propagation or forward UQ (cf. Ernst et al (2022)). Approximation methods 072 for random fields and their incorporation into PDE solution methods have 073 been actively developed in the engineering and numerical analysis communities 074 in the past two decades, and excellent surveys can be found in Ghanem and 075 Spanos (1991): Babuška et al (2010): Schwab and Gittelson (2011): Gunzburger 076 et al (2014). At the same time, sampling-based simulation techniques known as 077 Gaussian process emulators have gained popularity in the statistics community 078for solving similar problems, cf. Sacks et al (1989); Currin et al (1991); Kennedy 079 and O'Hagan (2001); O'Hagan (2006). Our objective in this work is the direct 080 comparison of these two approaches using Monte Carlo sampling as a reference 081 in a case study on the hydrogeological transport of radionuclides within the 082 site assessment for a nuclear waste repository. In doing so, we place particular 083 emphasis on the careful construction of a stochastic model using geostatistical 084 techniques. 085

The uncertainty propagation techniques we shall consider are based on generating realizations (samples) of the uncertain input parameters, solving the PDE for each realization and then determining the statistical properties of the quantities of interest in a post-processing step. As each PDE solution typically requires considerable computational resources, the mapping of random input parameters to quantities of interest is often substituted by *surrogate models*, which are considerably less costly to evaluate, thus speeding up the uncertainty

propagation analysis. The two surrogates we shall compare, sparse polynomial 093 collocation and Gaussian process emulation are interesting in that they were 094developed in different fields (numerical analysis and statistics), display differ-095 ent performance characteristics, and also differ in the interpretations of the 096 surrogates they produce. Our work is closest in spirit to Owen et al (2017). 097 where Gaussian process emulation is compared with polynomial chaos expan-098 sion surrogates for two black-box computer simulators. Although different 099 in construction, polynomial chaos surrogates yield a multivariate polynomial 100 approximation of the input-output map realized by the computer simulator as 101 does stochastic collocation, whereas the latter is considerably easier to inte-102grate into PDE solvers. In place of a small number of discrete parameters in 103the models considered in (Owen et al, 2017), the random input in our ground-104water model is a random field, i.e., its realizations are functions, which can 105be considered as parameterized by a countably infinite number of parameters. 106A comparison between uncertainty propagation techniques in aerodynamic 107 modeling can be found in Liu et al (2017). 108

The remainder of the paper is organized as follows: Section 2 presents the 109problem of predicting the travel or *exit time* of radionuclides transported by 110 groundwater flow through a horizontal layer above the Waste Isolation Pilot 111 Plant, an operational underground disposal site for nuclear waste, in a scenario 112where a hypothetical future accidental breach leads to the release of radioactive 113material. The physical as well as the probabilistic model are presented as well 114 as how observational data of hydraulic transmissivity is incorporated, leading 115to the generation of samples of the exit time quantity of interest. Section 3 116describes the computational realization for solving the Darcy flow equations, 117 the construction of the truncated Karhunen-Loève representation of the ran-118 dom transmissivity field as well as the estimation of the cumulative distribution 119function of the exit time quantity of interest. Section 4 gives detailed descrip-120tion of the two surrogate types to be compared, Gaussian process emulation 121and sparse polynomial collocation, emphasising their differences with respect 122to construction, computation and interpretation. In Section 5, we present the 123results of numerical computations with both surrogates using original data 124from the WIPP site, and present our conclusions in Section 6. 125

2 Uncertainty Propagation for a Groundwater Flow Problem

In this section we introduce the application setting, physical model, UQ task as well as the probabilistic model with which this is addressed. 130 131 132

2.1 The Waste Isolation Pilot Plant (WIPP)

The Waste Isolation Pilot Plant (WIPP) in Carlsbad, NM is a long-term deep135geologic storage facility for transuranic waste operated by the U.S. Depart-136ment of Energy since 1999. One of the issues investigated in the course of an137

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extensive performance assessment for WIPP was the risk of hazardous mate-139140 rials escaping to the biosphere in the event of a future accidental breach of the enclosure system. As the most likely pathway for such contaminants is trans-141 142port through the subsurface via groundwater, we are led to the objective of predicting the groundwater flow and transport of contaminants released from 143the storage site. In case of the WIPP, the disposal area lies within in the Salado 144145bedded salt formation. The Salado itself as well as the overlying formations are essentially impermeable to groundwater with the exception of a laterally exten-146147sive but narrow layer of rock known as the Culebra Dolomite. Details of the geological site characterization can be found in the extensive documentation¹ 148149in the WIPP certification and recertification applications (U.S. Department 150of Energy (DOE), 2004, 2014) which are produced every five years. Figure 1, 151taken from (U.S. Department of Energy (DOE), 2014), shows the location of 152the WIPP site within the UTM coordinate system, the location of boreholes 153where measurements of transmissivity and hydraulic head were obtained as 154well as the boundaries of areas with distinct geological features.

155One of the most relevant quantities of interest is the travel or *exit time* of radionuclides after release from a point within the Culebra layer above the 156157site to reach the boundary of the repository area, the computation of which requires simulating the groundwater flow and transport in the Culebra. As the 158precise transmissivity properties of the rock are uncertain, the same applies to 159160 the exit time. In the remainder of this section we describe a model for ground-161 water flow and contaminant transport in which the uncertain transmissivity 162is modeled stochastically, incorporating geological background information, 163standard geostatistical assumptions as well as available measurement data. 164

165 2.2 Darcy Flow and Particle Transport

We model the flow of groundwater through the Culebra dolomite geological unit by stationary single-phase Darcy flow. Denoting by p the hydraulic head (pressure) and by K the (scalar) hydraulic conductivity, the volumetric flux (Darcy flux) q is given by

$$\boldsymbol{q} = -K\nabla p. \tag{1}$$

171 172 If \boldsymbol{u} denotes the pore velocity of the groundwater, which is related to the Darcy 173 flux in terms of the *porosity* ϕ as $\boldsymbol{q} = \phi \boldsymbol{u}$, conservation of mass in the absence 174 of sources and sinks leads to the divergence-free condition

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$$\nabla \cdot \boldsymbol{u} = 0. \tag{2}$$

177 Since the aquifer under consideration is essentially horizontal with a much 178 larger lateral than vertical extent, we model the flow as two-dimensional and 179 consider the hydraulic *transmissivity* T = bK in place of conductivity, where 180 b denotes the aquifer thickness.

181 On the boundary ∂D of the bounded computational domain D, we dis-182 tinguish impermeable segments Γ_N along which the normal flux vanishes and 184 \Box

¹These can be found at https://wipp.energy.gov/epa-certification-documents.asp.

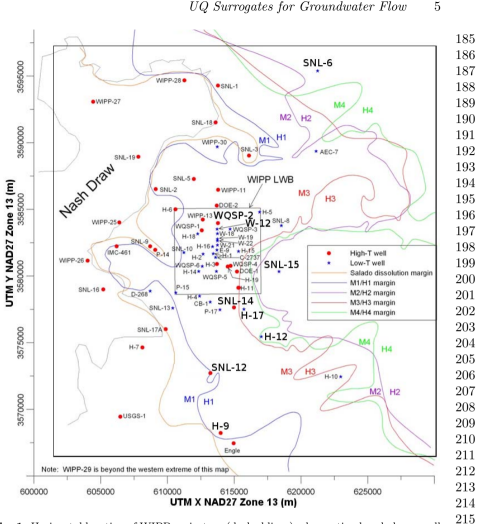


Fig. 1 Horizontal location of WIPP repisotory (dashed lines), observation boreholes as well as boundaries of distinct geological features. Source: (U.S. Department of Energy (DOE), 2014).

their complement $\Gamma_D = \partial D \setminus \Gamma_N$, where we prescribe the value of the hydraulic head p. Denoting by n the exterior unit normal vector along Γ_N and by g the prescribed head data along Γ_D , this leads to the boundary conditions

$$\boldsymbol{n} \cdot \boldsymbol{u} = 0 \text{ on } \Gamma_N, \qquad p = g \text{ on } \Gamma_D.$$
 (3)

The computational domain D as well as the boundary segments Γ_N and Γ_D are displayed in the left panel in Figure 2. The Dirichlet data q is obtained by evaluating a kriging interpolant (cf. Section 2.4.3) of observational hydraulic head data taken from (U.S. Department of Energy (DOE), 2014). As the flux 229

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variable \boldsymbol{u} is of primary interest in view of the subsequent transport calcula-231232tion we employ the usual mixed formulation of the boundary value problem presented by (1), (2) and (3). The associated variational formulation consists 233234in finding the pair $(\boldsymbol{u}, p) \in \mathcal{V} \times \mathcal{W}$ such that

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$$\left(\frac{\phi b}{T}\boldsymbol{u},\boldsymbol{v}\right) - (p,\nabla\cdot\boldsymbol{v}) = -\langle g,\boldsymbol{n}\cdot\boldsymbol{v}\rangle_{\Gamma_D} \qquad \forall \boldsymbol{v}\in\mathcal{V}, \qquad (4a)$$
$$(\nabla\cdot\boldsymbol{u},g) = 0 \qquad \forall g\in\mathcal{W} \qquad (4b)$$

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240with suitable boundary data $g \in H^{1/2}(\Gamma_D)$. Here (\cdot, \cdot) denotes the $L^2(D)$ 241inner product, the variational spaces are given by 242

 $\mathcal{V} = \{ \boldsymbol{v} \in \boldsymbol{H}(\operatorname{div}; D), \boldsymbol{n} \cdot \boldsymbol{v}|_{\Gamma_N} = 0 \}, \qquad \mathcal{W} = L^2(D)$

245and $\langle \cdot, \cdot \rangle_{\Gamma_D}$ denotes the duality pairing of $H^{1/2}(\Gamma_D) \times H^{-1/2}(\Gamma_D)$. Given the 246flux solution \boldsymbol{u} of (4), the trajectory of a particle from a release point $\boldsymbol{x}_0 \in D$ 247neglecting hydraulic dispersion is found as the solution of the initial value 248problem 249

$$\dot{\boldsymbol{x}}(t) = \boldsymbol{u}(\boldsymbol{x}(t)), \qquad t \ge 0, \quad \boldsymbol{x}(0) = \boldsymbol{x}_0.$$
(5)

(4b)

250A discussion of the regularity requirements for the Darcy flow problem (4) 251needed to ensure existence and uniqueness of the particle trajectory (5) can 252be found in (Graham et al, 2016, Section 5.3). As we shall see below, for the 253probabilistic model of transmissivity with finite-dimensional noise, which we 254shall employ in our calculations, these requirements are satisfied. As a *quantity* 255of interest derived from the solution of the random Darcy flow equations, 256we choose the logarithm of the travel or exit time of a particle released at 257a location x_0 inside the Culebra layer above the WIPP repository until it 258reaches the boundary of the subdomain $D_0 \subset D$ marking the edge of the 259WIPP site projected vertically up to the Culebra layer within the surrounding 260computational domain D, 261

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$$f_{\text{exit}} := \log \min\{t > 0 : \boldsymbol{x}(t) \notin D_0, \ \boldsymbol{x}_0 \in D_0\}.$$

264The location of the release point \boldsymbol{x}_0 , the perimeter of the WIPP site D_0 as well 265as a number of particle trajectory realizations from x_0 to ∂D_0 are displayed 266in Figure 2. 267

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2.3 Probabilistic Modeling of Uncertain Transmissivity 269

270The primary source of uncertainty in the modeling of flow and transport in 271the Culebra dolomite is the spatial variation of hydraulic conductivity, or, 272in our horizontal two-dimensional setting, transmissivity T. The prevailing 273mathematical description of uncertainty is probabilistic, i.e., the quantities in 274question are modeled as random variables following a given probability distri-275bution. The randomness thus introduced is an expression of uncertainty due 276

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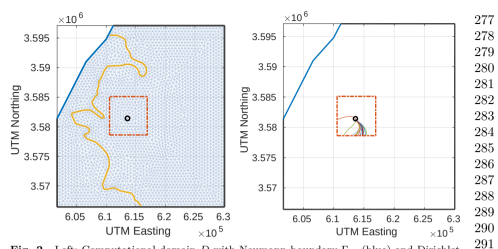


Fig. 2 Left: Computational domain D with Neumann boundary Γ_N (blue) and Dirichlet 292boundary Γ_D (black) as well as the perimeter of the WIPP site D_0 (red dashed), location of particle release point x_0 (black circle), and boundary of the Salado dissolution zone D_1 (yellow), cf. Section 2.4.1 below, respected by the triangular finite element mesh. Right: 294Simulation of several realizations of random particle trajectories from x_0 to ∂D_0 . 295

296to lack of knowledge of the precise spatial variation of transmissivity through-297out the domain D in the sense that some realizations of transmissivity across 298the domain are more likely than others. Rather than a deterministic value 299 $T = T(\boldsymbol{x})$, transmissivity at a point $\boldsymbol{x} \in D$ (scaled by porosity and thickness) 300 is thus expressed as a random variable $T(\boldsymbol{x}, \omega)$ governed by a probability mea-301 sure **P** defined on a probability space $(\Omega, \mathfrak{A}, \mathbf{P})$ with elementary outcome set 302 Ω carrying a σ -algebra \mathfrak{A} on which a probability measure **P** is defined. The 303 collection of all such random variables $\{T(\boldsymbol{x}, \omega) : \boldsymbol{x} \in D\}$ is known as a ran-304 dom field, i.e., a stochastic process for which the index variable x is a spatial 305coordinate. The most well-established probabilistic model for transmissivity in 306 the hydrology literature assumes that $T(\boldsymbol{x}, \cdot)$ follows a lognormal distribution, 307 i.e., that $Z(\boldsymbol{x}, \cdot) := \log T(\boldsymbol{x}, \cdot)$ is a Gaussian random field (cf. Freeze (1975); 308 Hoeksema and Kitanidis (1985) and (de Marsily, 1986, Chapter 11)). By con-309 sequence, realizations of $T = \exp(Z)$ are always positive. Such a Gaussian 310random field Z is completely specified by its mean and covariance function 311

$$\overline{Z}(\boldsymbol{x}) = \mathbf{E}[Z(\boldsymbol{x},\cdot)], \qquad \qquad \boldsymbol{x} \in D, \qquad \qquad \begin{array}{c} 312\\ 313 \end{array}$$

and
$$c(\boldsymbol{x}, \boldsymbol{y}) = \mathsf{E}\left[(Z(\boldsymbol{x}) - \overline{Z}(\boldsymbol{x}))(Z(\boldsymbol{y}) - \overline{Z}(\boldsymbol{y}))\right], \quad \boldsymbol{x}, \boldsymbol{y} \in D,$$
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respectively, where $\mathbf{E}\left[\cdot\right]$ denotes mathematical expectation with respect to \mathbf{P} . 316

We assume throughout that the covariance function of $Z = \log T$ isotropic 317and that the fluctuation $Z - \overline{Z}$ is wide-sense stationary such that we have 318 $c(\mathbf{x}, \mathbf{y}) = c(|\mathbf{x} - \mathbf{y}|)$, i.e., the covariance depends only on the (Euclidean) 319separation distance $r = |\mathbf{x} - \mathbf{y}|$. Moreover, we assume c(r) to belong to the 320

323 Matérn family of covariance models

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- $\begin{array}{c} 325\\ 326 \end{array}$

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$$c(r) = \frac{\sigma^2}{2^{\nu-1} \Gamma(\nu)} \left(\frac{2\sqrt{\nu} r}{\rho}\right)^{\nu} K_{\nu} \left(\frac{2\sqrt{\nu} r}{\rho}\right), \qquad r = |\boldsymbol{x} - \boldsymbol{y}|, \qquad (6)$$

where K_{ν} denotes the modified Bessel function of order $\nu > 0$. The quantity 328 ν is called the *smoothness parameter*, $\sigma^2 = c(0) = \text{Var } Z(\boldsymbol{x})$ is the (marginal) 329 330 variance (constant in \boldsymbol{x}) and the parameter $\rho > 0$ is called the *correlation* 331 *length*, a measure of how quickly the covariance decays with separation distance. An extensive justification for using the Matérn model as well as its 332 333 properties and scaling variants can be found in (Stein, 1999, pp. 48). For the 334 particular scaling (6), the Matérn covariance coincides with the exponential covariance for $\nu = \frac{1}{2}$, the Bessel covariance for $\nu = 1$ and the squared expo-335 nential covariance in the limit $\nu \to \infty$. The smoothness of the realizations of 336Z increases with ν , and the spatial scale of variation is described by ρ . We 337 338 determine the values of the hyperparameters (σ, ρ, ν) by statistical estimation 339 based on data published in the WIPP Compliance Recertification Assessment 340 U.S. Department of Energy (DOE) (2014) documents, which contain measure-341 ments of transmissivity in the Culebra dolomite at 62 boreholes throughout 342 the assessment site (cf. Figure 1). Figure 3 displays realizations of a Gaussian random field describing $Z = \log T$ throughout the computational domain 343 344 D representing the Culebra flow domain. It can be seen that larger values 345of ν result in realizations that are smoother, and smaller values of ρ lead to 346structures which decorrelate faster with separation distance.

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348 2.4 Statistical Estimation of Transmissivity Field

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$$Z := \log T = \overline{Z}(\boldsymbol{x}) + \tilde{Z}(\boldsymbol{x},\omega)$$
(7)

is Gaussian with (deterministic) mean \overline{Z} and (centered) residual field \tilde{Z} . Due to the complexity and irregular features of geological structures, it is crucial to merge stochastic models with available measurement data in a transparent fashion. Below we summarize the statistical techniques by which available data is incorporated into the stochastic model of uncertain transmissivity.

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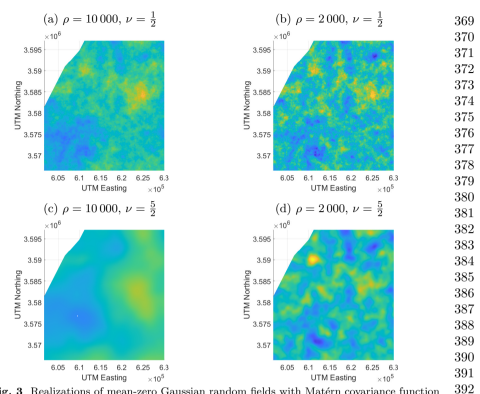


Fig. 3 Realizations of mean-zero Gaussian random fields with Matérn covariance function for different values of ρ and ν . All plots use the same color map and σ^2 was set to 1 in each case.

2.4.1 Regression Modeling of Mean Transmissivity

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The deterministic mean \overline{Z} of the log-transmissivity field is constructed as a linear regression model

$$\lceil h_1(\boldsymbol{x}) \rceil$$
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$$\overline{Z}(\boldsymbol{x}) = \sum_{j=1}^{n} \beta_j h_j(\boldsymbol{x}) = \boldsymbol{h}(\boldsymbol{x})^\top \boldsymbol{\beta}, \qquad \boldsymbol{h}(\boldsymbol{x}) = \begin{vmatrix} \vdots \\ h_k(\boldsymbol{x}) \end{vmatrix}, \qquad (8) \quad \begin{array}{c} 401 \\ 402 \\ 403 \\ 403 \end{vmatrix}$$

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in which the k components of h contain regression functions from which the 405trend behavior of Z can be obtained by linear combination. Known geological 406 features of the area under study can be incorporated by choosing the regression 407 functions as, e.g., indicator functions of subdomains possessing distinguishing 408 characteristics, linear or polynomial trends to be fitted as well as the variation 409of available quantities known or believed to affect the transmissivity field. 410Based on the available WIPP technical documents, model comparison was 411

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415 made using the five regression functions

416 $h_1(\boldsymbol{x}) \equiv 1$ (constant), $h_4(\boldsymbol{x}) = d(\boldsymbol{x})$ (overburden), $h_2(\boldsymbol{x}) = x_1$ (linear in x_1), $h_5(\boldsymbol{x}) = \mathbb{1}_{D_1}(\boldsymbol{x})$ (zone indicator). (9) $h_3(\boldsymbol{x}) = x_2$ (linear in x_2 .)

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421The first three regression functions allow to fit a basic affine trend. The over-422 burden $d(\mathbf{x})$ denotes the vertical distance between the ground surface and the 423 top of the Culebra layer above location \boldsymbol{x} . This is an indication of the extent 424to which erosion has led to stress relief on the underlying Culebra layer, possi-425bly causing new fracturing or the opening of pre-existing fractures and thereby 426enhancing transmissivity. Regression function h_5 is the indicator function of a 427 subdomain $D_1 \subset D$ to the north, south and west of the WIPP site, where dis-428solution of the upper Salado formation has led to strain in the rock overlying 429the Salado, including the Culebra, leading to larger apertures in existing frac-430tures, collapse and brecciation and thus to a generally higher transmissivity 431(cf. U.S. Department of Energy (DOE) (2004)). 432

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${}^{433}_{434}$ 2.4.2 Hyperparameter Estimation and Model Selection

For all combinations of the regression functions (9), a restricted maximum 435436likelihood (RML) estimation procedure detailed in Appendix A was used to 437 determine the hyperparameters σ, ρ and ν of the Matérn covariance model 438(6) based on the 62 transmissivity observations published in U.S. Department of Energy (DOE) (2014). Based on this calibrated covariance structure, a 439440 model comparison was carried out following a procedure proposed in Kitanidis 441(1997b), in which a significance test is used to determine whether adding fur-442 ther regression functions to a model better explains the data. In this way, we 443 arrived at a trend model (8) consisting of the regression functions $\{h_1, h_2, h_5\}$ from (9). In the following we refer to this parametrization of the mean as the 444 best model and to that containing only the constant trend function h_1 as the 445446constant model. The resulting estimates of the hyperparameters σ, ρ and ν for both models are given in Table 1. Note that we have fixed $\nu = 0.5$ in both cases 447 since the estimates for ν were sufficiently close to this value², which also allows 448a more efficient evaluation of the associated covariance function. The regres-449sion model estimated by the (weighted) least-squares method for the mean is 450451then

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$$Z(\boldsymbol{x}) = 143.98 - 2.55 \cdot 10^{-4} x_1 + 3.31 \, \mathbf{1}_{D_1}(\boldsymbol{x}).$$

453 Note that the values for x_1 (UTM Easting coordinates) are of order $6 \cdot 10^5$ for 454 the WIPP computational domain D.

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⁴⁵⁶ $\overline{}^{2}$ If we do not fix $\nu = 0.5$ but estimate it as well the RML results are $\hat{\sigma}^{2} = 6.14$, $\hat{\rho} = 2005.2$, 457 and $\hat{\nu} = 0.48$.

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Trend model	Sill σ^2	Range ρ	Smoothness ν	
h_1	17.12	6509.8	0.5	
h_1, h_2, h_5	6.15	1948.0	0.5	

Table 1 Restricted maximum likelihood estimation of hyperparameters σ^2 (variance or *sill*) and ρ (correlation length or *range*) for two trend models based on the 64 observations of transmissivity. The smoothness parameter was fixed at $\nu = 1/2$, which corresponds to the exponential covariance kernel. 466

2.4.3 Conditioning on Transmissivity Data

Once the mean and covariance functions of the Gaussian random field Z = 470log T have been determined, the log transmissivity measurements $\{z(\boldsymbol{x}_j)\}_{j=1}^N$ 471 may be used to further calibrate the stochastic model to fit the observations in a statistical sense using the technique known as kriging (cf. Cressie (1991); Kitanidis (1997a); Stein (1999)). Kriging refers to best linear unbiased prediction (BLUP) in which the value of the random field Z at an arbitrary location $\boldsymbol{x} \in D$ is estimated as an affine combination 475476477

$$\hat{Z} = \hat{Z}(\boldsymbol{x}, \omega) = \lambda_0(\boldsymbol{x}) + \boldsymbol{\lambda}(\boldsymbol{x})^\top \boldsymbol{Z}(\omega)$$
(10) 478
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of the (random) realizations $\mathbf{Z} = (Z(\mathbf{x}_1), \ldots, Z(\mathbf{x}_N))^{\top}$, with spatially varying coefficients $\lambda_0 : D \to \mathbb{R}$ and $\boldsymbol{\lambda} = (\lambda_1(\mathbf{x}), \ldots, \lambda_N(\mathbf{x})) : D \to \mathbb{R}^N$ chosen to make the estimator *unbiased* and *mean square optimal*, which requires that, for all $\mathbf{x} \in D$, we have 483

$$\mathsf{E}\left[\hat{Z}(\boldsymbol{x})\right] = \mathsf{E}\left[Z(\boldsymbol{x})\right] \quad \text{and} \quad \mathsf{E}\left[|Z(\boldsymbol{x}) - \hat{Z}(\boldsymbol{x})|^2\right] \to \min_{\lambda_0, \boldsymbol{\lambda}}. \tag{485}$$
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For a known mean function \overline{Z} the solution is given by the *(simple) kriging* 488 prediction or interpolation 489

where $\overline{Z} := [\overline{Z}(\boldsymbol{x}_1), \dots, \overline{Z}(\boldsymbol{x}_N)]^\top$, $\boldsymbol{c}(\boldsymbol{x}) := (c(\boldsymbol{x}, \boldsymbol{x}_1), \dots, c(\boldsymbol{x}, \boldsymbol{x}_N))^\top$ and $\boldsymbol{C} := 493$ $(c(\boldsymbol{x}_i, \boldsymbol{x}_j))_{i,j=1,\dots,N} \in \mathbb{R}^{N \times N}$, with mean square error given via the kriging 494 (error) covariance 495

$$\mathbf{E}\left[\left|Z(\boldsymbol{x}) - \hat{Z}(\boldsymbol{x})\right|^{2}\right] = \hat{c}(\boldsymbol{x}, \boldsymbol{x}), \qquad \hat{c}(\boldsymbol{x}, \boldsymbol{y}) \coloneqq c(\boldsymbol{x}, \boldsymbol{y}) - \boldsymbol{c}(\boldsymbol{x})^{\top} \boldsymbol{C}^{-1} \boldsymbol{c}(\boldsymbol{y}). \qquad \begin{array}{c} 497\\ 498\\ 499\end{array}$$

Note that for a Gaussian random field Z the kriging prediction \hat{Z} is again Gaussian and coincides with the conditioned random field $Z(\boldsymbol{x})|\boldsymbol{Z} = \boldsymbol{z}$, where $\boldsymbol{z} = (z_1, \dots, z_N)^{\top}$, so that $\hat{Z}(\boldsymbol{x}) \sim \mathsf{N}(\overline{\boldsymbol{Z}}(\boldsymbol{x}) + \boldsymbol{c}(\boldsymbol{x})^{\top} \boldsymbol{C}^{-1}(\boldsymbol{z} - \overline{\boldsymbol{Z}}), \hat{c}(\boldsymbol{x}, \cdot))$. It is easily verified that at the observation sites $\{\boldsymbol{x}_j\}_{j=1}^N$ we have $\hat{Z}(\boldsymbol{x}_j) = z(\boldsymbol{x}_j)$ and $\hat{c}(\boldsymbol{x}_j, \boldsymbol{x}_j) = 0$, hence the kriging estimate \hat{Z} of the random field Z interpolates the measurements.

In the variant called *universal kriging*, the mean \overline{Z} is not assumed known 507and instead modelled as in (8). Forming the least squares estimate $\hat{\beta}$ of β and 508proceeding as above with $\overline{Z}(\mathbf{x}) = \mathbf{h}(\mathbf{x})^{\top} \hat{\boldsymbol{\beta}}$ would fail to account for uncer-509510tainty in this estimate. Instead, we require that unbiasedness of the kriging estimate (10) hold for all $\boldsymbol{\beta} \in \mathbb{R}^k$, resp. for all possible mean functions. Apply-511ing unbiasedness as a contraint in the pointwise minimization over λ_0, λ via 512513Lagrange multipliers yields the *universal kriging prediction* or interpolation 514515

 $\hat{Z}(\boldsymbol{x}) = \begin{bmatrix} \boldsymbol{c}(\boldsymbol{x}) \\ \boldsymbol{h}(\boldsymbol{x}) \end{bmatrix}^{\top} \begin{bmatrix} \boldsymbol{C} & \boldsymbol{H} \\ \boldsymbol{H}^{\top} & \boldsymbol{0} \end{bmatrix}^{-1} \begin{bmatrix} \boldsymbol{Z} \\ \boldsymbol{0} \end{bmatrix}, \qquad (11)$

518 where 519

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 $oldsymbol{H} oldsymbol{H} = egin{bmatrix} h_1(oldsymbol{x}_1) & \ldots & h_k(oldsymbol{x}_1) \ dots & & dots \ h_1(oldsymbol{x}_N) & \ldots & h_k(oldsymbol{x}_N) \end{bmatrix} \in \mathbb{R}^{N imes k},$

522523 or, equivalently,

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 $\hat{Z}(\boldsymbol{x}) = \boldsymbol{h}(\boldsymbol{x})^{\top} \hat{\boldsymbol{\beta}} + \boldsymbol{c}(\boldsymbol{x})^{\top} \boldsymbol{C}^{-1} \left(\boldsymbol{Z} - \boldsymbol{H} \hat{\boldsymbol{\beta}} \right),$ (12)

527 where $\hat{\boldsymbol{\beta}} = (\boldsymbol{H}^{\top} \boldsymbol{C}^{-1} \boldsymbol{H})^{-1} \boldsymbol{H}^{\top} \boldsymbol{C}^{-1} \boldsymbol{Z}$, with mean square error 528 $\mathbf{E} \left[|Z(\boldsymbol{x}) - \hat{Z}(\boldsymbol{x})|^2 \right] = \hat{c}(\boldsymbol{x}, \boldsymbol{x})$ given in this case by the *universal kriging* 530 *(error) covariance*

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 $\hat{c}(\boldsymbol{x},\boldsymbol{y}) := c(\boldsymbol{x},\boldsymbol{y}) - \boldsymbol{c}(\boldsymbol{x})^{\top} \boldsymbol{C}^{-1} \boldsymbol{c}(\boldsymbol{y}) + \boldsymbol{\gamma}(\boldsymbol{x})^{\top} \boldsymbol{V} \boldsymbol{\gamma}(\boldsymbol{y}),$ (13)

533where $\boldsymbol{\gamma} = \boldsymbol{h}(\boldsymbol{x}) - \boldsymbol{H}^{\top} \boldsymbol{C}^{-1} \boldsymbol{c}(\boldsymbol{x})$ and $\boldsymbol{V} = (\boldsymbol{H}^T \boldsymbol{C}^{-1} \boldsymbol{H})^{-1}$. Thus, the univer-534sal kriging prediction (12) consists in obtaining the mean as the least squares 535estimate $h(x)^{\top}\hat{\beta}$ and proceeding as in simple kriging. However, the univer-536sal kriging mean square error contains the additional term $\boldsymbol{\gamma}(\boldsymbol{x})^{\top} \boldsymbol{V} \boldsymbol{\gamma}(\boldsymbol{x}) > 0$ 537compared to that of simple kriging, which accounts for the additional uncer-538tainty present in the estimated mean and β , respectively. Note further that, 539even for Gaussian Z, the universal kriging mean and (co)variance do not, in 540general, possess an interpretation as those of a conditioned Gaussian random 541field as is the case with simple kriging. 542

543 We now use the universal kriged Gaussian random field \hat{Z} obtained from 544 the available log transmissivity measurements $\boldsymbol{z} = \{z(\boldsymbol{x}_j)\}_{j=1}^N$ as our final 545 stochastic model for the uncertain transmissivity field, i.e.,

- 546 547
- $\hat{Z}(\boldsymbol{x}) \sim \mathsf{N}\left(\hat{z}(\boldsymbol{x}), \hat{c}(\boldsymbol{x}, \cdot)
 ight)$
- 548

549 with \hat{c} given in (13) and \hat{z} resulting by inserting the realization $\mathbf{Z} = \mathbf{z}$ in (11). 550 The resulting kriged mean \hat{z} and pointwise variance \hat{c} are displayed in Figure 4. 551 In summary, our approach for the stochastic modeling of the transmissivity 552 field $T = T(\mathbf{x}, \omega)$ is characterized by

- (2) obtaining the parameters σ , ν and ρ in the Matérn covariance function using restricted maximum likelihood estimation (RML);
- (3) conditioning the thus obtained lognormal field on the available observations of transmissivity at the WIPP site.

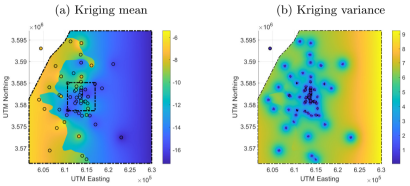


Fig. 4 Universal kriging prediction of $Z = \log T$ based on 62 available transmissivity observations. Left: kriged mean field $\hat{z}(\boldsymbol{x})$. Right: pointwise kriging variance $\hat{c}(\boldsymbol{x}, \boldsymbol{x})$. The circular markers indicate the locations (and values) of the observational log transmissivity data. The interpolation property of $\hat{z}(\boldsymbol{x})$ is apparent.

2.5 Uncertainty Propagation for the Quantity of Interest

For a random transmissivity field $T(\omega) = T(\cdot, \omega), \omega \in \Omega$, we consider individual 579 realizations of the associated random boundary value problem in its mixed 580 formulation (4), i.e., 581

$$\left(\frac{\phi b}{T(\omega)}\boldsymbol{u}(\omega),\boldsymbol{v}\right) - (p(\omega),\nabla\cdot\boldsymbol{v}) = -\langle g,\boldsymbol{n}\cdot\boldsymbol{v}\rangle_{\Gamma_D} \qquad \forall \boldsymbol{v}\in\mathcal{V},$$
(14a)

$$(\nabla \cdot \boldsymbol{u}(\omega), q) = 0 \qquad \qquad \forall q \in \mathcal{W}, \qquad (14b) \quad \frac{585}{586}$$

with random solution pair $(\boldsymbol{u}(\omega), p(\omega)) \in \mathcal{V} \times \mathcal{W}$. The equations (14) are now understood as holding **P**-almost surely. Under suitable assumptions (cf. Babuška et al (2007)) we have for the random solution pair, that $(\boldsymbol{u}, p) \in$ $L^2_{\mathbf{p}}(\mathcal{V} \times \mathcal{W})$, i.e., the norm of the solution is square integrable w.r.t. the probability measure **P**.

For the quantity of interest under consideration, the exit time for particle trajectories, each realization of the random flux yields a realization of the associated random initial value problem

$$\dot{\boldsymbol{x}}(t,\omega) = \boldsymbol{u}(\boldsymbol{x}(t,\omega),\omega), \quad t \ge 0, \qquad \boldsymbol{x}(0,\omega) = \boldsymbol{x}_0. \tag{15} \quad \begin{array}{c} 596\\ 597 \end{array}$$

599 **P**-almost surely, and hence, the quantity of interest becomes a random variable 600

601 602

$$f_{\text{exit}}(\omega) := \log \min\{t > 0 : \boldsymbol{x}(t,\omega) \notin D_0, \ \boldsymbol{x}_0 \in D_0\}.$$
 (16)

603 A complete characterization of the uncertainty in f is given by its cumulative 604 distribution function (CDF)

605

 $\begin{array}{c} 606 \\ 607 \end{array}$

$$F(s) := \mathbf{P}(f_{\text{exit}} \le s), \qquad F \colon \mathbb{R} \to [0, 1].$$

608 Due to the complexity of the problem, F cannot be given in analytic form and 609 has to be approximated. We comment on the computational aspects in the 610 next section.

611

⁶¹²₆₁₃ 3 Computational Realization

614 In this section we describe (i) the spatial discretization used for solving the 615 Darcy flow equations (4) or (14), respectively, given a realization of the trans-616 missivity field T, (ii) a discrete representation of the random model for the 617 transmissivity field T as well as (iii) a Monte Carlo approach for approximating 618 the CDF of the quantity of interest.

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620

²⁰ 3.1 Finite Element Solution of Darcy Flow Problem

⁶²¹ ⁶²² We solve the Darcy flow equations (4) – or individual realizations of their ⁶²³ random form (14) – using a mixed finite element discretization consisting of the ⁶²⁴ lowest order Raviart-Thomas space $\mathcal{V}_h \subset \mathcal{V}$ for the flux variable and piecewise ⁶²⁵ constant space $\mathcal{W}_h \subset \mathcal{W}$ for the hydraulic head with respect to a triangulation ⁶²⁶ \mathcal{T}_h of the domain D, where h > 0 is a measure of mesh resolution. This ⁶²⁷ discretization is known to be inf-sup-stable (cf. (Boffi et al, 2013, Chapter 7), ⁶²⁸ (Ern and Guermond, 2021, Chapter 51)).

We choose a fixed triangulation of the two-dimensional computational 629 domain with mesh width h chosen such that at least 10 elements correspond to 630 the correlation length of the random transmissivity field, resulting in a mesh 631 consisting of 28 993 triangles with the associated finite element spaces contain-632ing 72705 degrees of freedom (43712 for flux and 28993 for hydraulic head). 633 Note that a coarser mesh is depicted in Figure 2 for illustration purposes. The 634 particle tracking is performed by solving the ordinary differential equation 635 (15) for the given realization. For the lowest-order Raviart-Thomas discretiza-636 tion, the constraint of zero divergence results in an elementwise constant flux, 637 making this computation trivial and incurring no additional discretization 638 error. 639

640

641 3.2 Conditioned Karhunen-Loève Expansion

Various methods exist to generate realizations of random fields, like turning
 bands, circulant embedding and Karhunen-Loève expansion, see Lord et al

(2014). In this work, we generate approximate realizations of the Gaussian log 645 transmissivity field by truncating its Karhunen-Loève expansion, an orthogonal expansion of a random field based on the spectral decomposition of its 647 covariance operator 648

649

$$C: L^{2}(D) \to L^{2}(D), \qquad u \mapsto Cu, \qquad (Cu)(\boldsymbol{x}) = \int_{D} c(\boldsymbol{x}, \boldsymbol{y}) u(\boldsymbol{y}) \, \mathrm{d}\boldsymbol{y}, \quad (17) \quad \begin{array}{c} 650\\ 651\\ 652 \end{array}$$

which for continuous covariance functions is compact and selfadjoint, positive 653 definite and hence possesses a system of orthonormal eigenfunctions $(z_m)_{m=1}^{\infty}$ 654 which are complete in $L^2(D)$. Denoting by $\lambda_m \ge 0$ the eigenvalue associated 655 with eigenfunction z_m (ordered descending), a second-order random field Z on 656 D with mean \overline{Z} possesses the expansion 657

$$Z(\boldsymbol{x}) = \overline{Z}(\boldsymbol{x}) + \sum_{m=1}^{\infty} \sqrt{\lambda_m} z_m(\boldsymbol{x}) \xi_m, \qquad \boldsymbol{x} \in D,$$
(18)
$$\begin{array}{c} 658\\ 659\\ 660\\ 661\end{array}$$

converging in L^2 , where $(\xi_m)_{m\in\mathbb{N}}$ is a sequence of pairwise uncorrelated random variables and $(\lambda_m)_{m\in\mathbb{N}}$ is square summable. In the present setting, the log transmissivity field Z is Gaussian, as stated in Section 2.3, therefore we have $\xi_m \sim N(0, 1)$ for all m.

An approximation suited for computation is obtained by truncating the infinite expansion in (17) after a finite number M of terms, hence the resulting approximation 668

$$Z(\boldsymbol{x}) \approx \overline{Z}(\boldsymbol{x}) + \sum_{m=1}^{M} \sqrt{\lambda_m} z_m(\boldsymbol{x}) \xi_m$$
(19) 670

for fixed M will depend on the decay rate of the eigenvalues.

Once a truncation index M has been fixed, the random field can be regarded 673 as parameterized by the uncorrelated M-variate normal random vector $\boldsymbol{\xi} = 674$ $(\xi_1, \ldots, \xi_M)^\top \sim \mathsf{N}(\mathbf{0}, \boldsymbol{I})$, which takes values in \mathbb{R}^M . We may thus consider 675 all random quantities in (14), i.e., the transmissivity field T and the solution 676 (\boldsymbol{u}, p) of the Darcy flow equations as well as the particle trajectories (15) and 677 exit time f_{exit} in (16) as parameterized by realizations of this single random 678 vector. 679

Explicit closed-form solutions to the eigenvalue problem (17) are known 680 only for a small number of special cases, hence we approximate the eigenpairs 681 numerically. We approximate the covariance operator C, where the covariance 682 kernel is obtained from the universal kriging covariance \hat{c} in (13), by Galerkin 683 projection into a finite-dimensional subspace W_h of $L^2(D)$ consisting of piecewise constant functions with respect to a triangulation of the domain D, which 685 we assume to be polygonal for simplicity³. Denoting by $\{\phi_1, \ldots, \phi_N\}$ a basis 686

690

 $^{^{3}}$ We use the same finite element space as for the piecewise constant discretization of the hydraulic head p for convenience. 688

 $u(oldsymbol{x}) = \sum^N u_i \phi_i(oldsymbol{x})$

16 UQ Surrogates for Groundwater Flow

691 of \mathcal{W}_h , we represent functions in \mathcal{W}_h as

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⁶⁹⁶ ⁶⁹⁷ with coefficient vector $\boldsymbol{u} = (u_1, \ldots, u_N)^{\top}$. Substituting (20) into (17), multi-⁶⁹⁸ plying it by test functions ϕ_j and integrating over D we arrive at the discrete ⁶⁹⁹ generalized eigenvalue problem

700 701

$$Cu = \lambda Mu, \qquad (21)$$

(20)

 $\frac{702}{703}$ where \boldsymbol{C} is a symmetric positive semi-definite matrix with entries

$$\begin{array}{c} 704 \\ 705 \end{array}$$

 $[\boldsymbol{C}]_{i,j} = (C\phi_i, \phi_j)_{L^2(D)} = \int_D \phi_j(\boldsymbol{x}) \int_D c(\boldsymbol{x}, \boldsymbol{y}) \phi_i(\boldsymbol{y}) \,\mathrm{d}\boldsymbol{y} \,\mathrm{d}\boldsymbol{x}$ (22)

 $\begin{array}{c} 707\\ 708\\ 709 \end{array}$ and \pmb{M} is the symmetric positive definite Gram matrix of the piecewise constant basis with entries

- 710
- 711

 $[\boldsymbol{M}]_{i,j} = (\phi_j, \phi_i)_{L^2(D)} = \int_D \phi_j(\boldsymbol{x}) \phi_i(\boldsymbol{x}) \,\mathrm{d}\boldsymbol{x}.$ (23)

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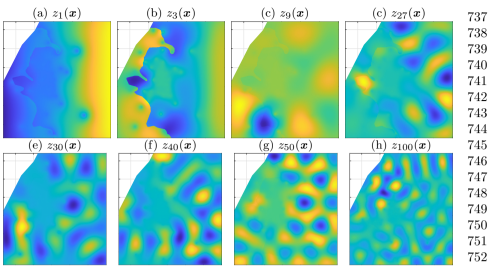
713An immediate difficulty with solving (21) is that C is a dense matrix due 714to the nonlocal nature of the integral operator C, hence generating and stor-715ing C is computationally expensive already for problems on two-dimensional 716domains, and even more so in three dimensions. Note that M is diagonal due 717to the disjoint supports of the ϕ_i . Moreover, even if generating and storing C 718were feasible, solving a dense eigenvalue problem by the standard symmetric 719QR algorithm results in excessive computation costs. We address this prob-720lem by first using an iterative method for approximating only the dominant 721M eigenvalues of C using a variant of the *thick-restart-Lanczos method* of Wu 722and Simon (2000), which requires only matrix vector products with C in the 723course of the iteration. Second, we represent C in *hierarchical matrix format* 724(cf. Hackbusch (2015)), which brings the cost of generating, storing and multu-725plying C by a vector from $\mathcal{O}(N^2)$ to a complexity $\mathcal{O}(N \log N)$. Further details 726on using hierarchical matrices in the context of random field generation with 727 the Galerkin method can be found in Eiermann et al (2007) and Khoromskij 728et al (2009).

Figure 5 shows a few computed eigenfunctions z_m for the kriging covariance function \hat{c} in (13) displayed in Figure 4.

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⁷³² 3.3 Empirical Estimation of the CDF ⁷³³

A common and convenient way to approximate the CDF F of the random quantity of interest $f_{\text{exit}}(\boldsymbol{\xi}) := \log \min\{t > 0 : \boldsymbol{x}(t, \boldsymbol{\xi}) \notin D_0, \ \boldsymbol{x}_0 \in D_0\}$ is by generating n samples f_1, \ldots, f_n of the random f_{exit} by sampling n different



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Fig. 5 Computed eigenfunctions of the kriging covariance function \hat{c} in (13), cf. Figure 4 .

realizations $\boldsymbol{\xi}_1, \dots, \boldsymbol{\xi}_n$ of the random coefficient vector $\boldsymbol{\xi}$ in the KL expansion 756 of the random field log T and solving the corresponding n boundary and initial 757 value problems to obtain $f_i = f_{\text{exit}}(\boldsymbol{\xi}_i)$. We then determine the empirical CDF 758 (ECDF) 759

$$F_n(s) = \frac{1}{n} \sum_{j=1}^n \mathbb{1}_{(-\infty, f_j]}(s).$$
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761
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762

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754 755

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The ECDF F_n is a random approximation to the CDF F of the quantity of interest f_{exit} due to the randomly drawn samples f_1, \ldots, f_n . We denote the error between the (random) ECDF and the true CDF by 763 764 765

$$D_n := \sup_{s \in \mathbb{R}} |F(s) - F_n(s)|.$$
(24) 766
767
768

For i.i.d. samples a classical result known as Donsker's theorem (Athreya and 769 Lahiri, 2006, Corollary 11.4.13) states 770

$$\sqrt{n}D_n \xrightarrow[n \to \infty]{} \sup_{t \in [0,1]} |B(t)|,$$
772
773

where B denotes a standard Brownian bridge on the unit interval [0, 1]. This theoretical result can be employed to compute the necessary minimal sample size n for a desired error criterion, which we fix here by requiring 777

$$\mathbf{P}(D_n > 0.01) \le 0.05. \tag{25} \begin{array}{c} 778\\ 779 \end{array}$$

Using the asymptotic result provided by Donsker's theorem as well as 780 $\mathsf{P}(||B||_{C[0,1]} > 1.36) \approx 0.05$, see (Williams, 2004, p. 343), we obtain for 782

783 $n \approx 20\,000$ that $\mathbf{P}(D_n > 0.01) \approx 0.05$. Hence, in the present setting this means 784 that, for this level of accuracy in approximating the CDF of the quantity of 785 interest, we need to solve $n = 20\,000$ Darcy flow equations and compute the 786 associated particle trajectories. Thus, the question arises whether we could 787 save computational work by employing surrogates for the mapping from the 788 random parameter vector $\boldsymbol{\xi}$ to the solution of the random PDE or the quantity 789 of interest f_{exit} itself.

790

791 Estimation of CDF based on surrogates

792 Assuming now that we have an approximation $\hat{f}_{exit}: \mathbb{R}^M \to \mathbb{R}$ to the quantity 793 of interest f seen as mapping from $\boldsymbol{\xi} \in \mathbb{R}^M \to \mathbb{R}$, the resulting approximate 794 ECDF $\hat{F}_n(s)$ based on n samples $\hat{f}_1, \ldots, \hat{f}_n$ of \hat{f}_{exit} resulting from n samples 795 $\boldsymbol{\xi}_i$ of the random KL parameter $\boldsymbol{\xi}$, where $\hat{f}_i = \hat{f}_{exit}(\boldsymbol{\xi}_i)$ is given by 796

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- 798

$$\hat{F}_N(s) = \frac{1}{N} \sum_{j=1}^N \mathbf{1}_{(-\infty,\hat{f}_j]}(s).$$

- 799
- 800

The question we investigate in this work is whether, for common surrogate 801constructions such as stochastic collocation and Gaussian process emulators, 802 the approximation error $||f_{\text{exit}} - \hat{f}_{\text{exit}}||$ (measured in a suitable norm) can 803 be made smaller than the sampling error D_n in the empirical estimation of 804 the CDF. To this end, we evaluate the quality of the surrogate f_{exit} by a 805 two-sample Kolmogorow-Smirnov (KS) test which is a well-known hypothe-806 sis test for checking whether sets of two samples—in our case $\hat{f}_1, \ldots, \hat{f}_n$ and 807 f_1, \ldots, f_n —are likely to have be drawn from the same distribution. Specifi-808 809 cally, in our case the KS test is passed at significance level $\alpha = 0.05$ if the 810 KS-statistic K satisfies

 $K := \sup_{s \in \mathbb{R}} \left| \hat{F}_n(s) - F_n(s) \right| \le 1.36 \frac{\sqrt{2}}{n},$

- 811
- 812
- 813

814 815 cf. Williams (2004).

816

817 4 Propagation Surrogates 818

819 In the following, we recall sparse grid polynomial collocation and Gaussian pro-820 cess emulators as surrogate techniques for approximating a function $f: \Xi \to \mathcal{Y}$ 821 of M (random or parametric) variables $\boldsymbol{\xi} \in \mathbb{R}^M$ taking values either in $\mathcal{Y} = \mathbb{R}$, 822 as for scalar quantities of interest such as the breakthrough time, or a function 823 space, e.g., $\mathcal{Y} = \mathcal{V} \times \mathcal{W}$, as for the solution of the mixed formulation (14) of 824 the Darcy flow equations with random conductivity.

We begin by illustrating the basic principles of polynomial collocation and Gaussian process emulation for the case of a single variable, i.e., $\Xi \subseteq \mathbb{R}$, before proceeding to the technical details for the multivariate case $\Xi \subseteq \mathbb{R}^M$, where we assume Ξ to be of product form $\Xi = \Xi^M$ with $\Xi \subseteq \mathbb{R}$.

4.1 Univariate Collocation and Emulation

As a simple example in the style of the GPE tutorial O'Hagan (2006), consider the function

$$y = f(\xi) := \xi + 3\sin\frac{3\xi}{4}, \qquad \xi \in \Xi := [0, 6].$$
 832

834 The presence of *input uncertainty*, i.e., uncertainty with regard to the precise 835 value of the independent variable ξ , is accounted for by modeling it as a random 836 variable $\xi \sim U[0,6]$. Suppose further that f is only accessible in the form of a 837 finite number of point evaluations $f(\xi)$, as is the case for the breakthrough time 838 in our WIPP case study, where each evaluation of the former requires solving 839 the Darcy flow problem followed by particle tracking up to the exit boundary. 840 The task is to construct a computationally inexpensive approximation $\hat{f}: \Xi \to I$ 841 \mathbb{R} of f given n evaluations

$$y_j = f(\xi_j), \qquad j = 1, \dots, n.$$
 843
844

The points of evaluation ξ_j are often called *design points* in the emulator literature and *nodes* or *knots* for collocation. Their choice depends on the type of surrogate being constructed. We begin with an elementary numerical analysis procedure and then contrast this with an approach rooted in the statistics community. 845 846 847 848 848 849 850

Polynomial Collocation

In the univariate case polynomial collocation simplifies to Lagrange interpolation by global polynomials, and the surrogate \hat{f} for f takes the familiar form

$$\hat{f}_n(\xi) := \sum_{j=1}^n f(\xi_j) \ell_j(\xi), \qquad \ell_j(\xi) = \prod_{k \neq j} \frac{\xi - \xi_k}{\xi_j - \xi_k}$$
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857

with $\{\ell_j\}_{j=1}^n$ the Lagrange fundamental polynomials associated with the nodes 858 $\{\xi_1, \ldots, \xi_n\}$. Although this expression is well-defined for any set of distinct 859 nodes, good approximation quality is only achieved if the points are chosen 860 with care. A classical choice for bounded intervals is the family of *Clenshaw*- 861 *Curtis nodes* (also called *Chebyshev nodes*). Scaled to the interval [0,6], the 862 set of *n* Clenshaw–Curtis nodes is given by 863

$$\xi_j = 3 + 3\cos\left(\frac{j-1}{n-1}\pi\right), \qquad j = 1, \dots, n.$$
 865
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Other common choices, particularly for UQ applications, are the roots of the *n*-868 th orthogonal polynomial associated with the probability density of ξ on Ξ , e.g., 869 Gauss–Legendre nodes for the uniform distribution or Gauss–Hermite nodes 870 for the normal distribution, cf. Babuška et al (2010). For optimal convergence 871 of the interpolants for smooth functions f it is well known that the spatial 872 distribution of the nodes $\xi_j \in \Xi$ should follow the equilibrium distribution 873 in the sense of logarithmic potential theory, which for the standard interval 874

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 $\Xi = [-1, 1]$ is given by $d\mu(\xi) = 1/\pi\sqrt{1-\xi^2}$, cf. (Trefethen, 2013, Chapter 12). 875 876 In particular, the nodes should cluster near the interval endpoints. Figure 6 shows two polynomial interpolation surrogates for f as well as the CDF of the 877 878 output $f(\xi)$.

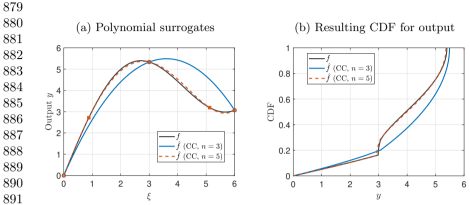


Fig. 6 The function $f(\xi) = \xi + 3\sin(3\xi/4)$ on $\Xi = [0,6]$ and its Lagrange interpolation 892 \hat{f}_n based on n=3 and n=5 Clenshaw-Curtis nodes (left) and the resulting CDF for the 893 output $y = f(\xi)$ and $\hat{y} = \hat{f}_n(\xi)$, resp., if $\xi \sim U(\Xi)$. 894

- 895 The approximation quality of polynomial interpolation depends not only 896 897 on the choice of interpolation nodes, but also on the *smoothness* of f. For example, we have for $f \in C^r(\Xi)$, $r \in \mathbb{N}$, that 898
- 899 900

$$||f - \hat{f}_n||_{\infty} \le c_r(f) n^{-r} (1 + \Lambda_{\xi_1, \dots, \xi_n})$$

901

where $||f - \hat{f}_n||_{\infty} = \sup_{\xi \in \Xi} |f(\xi) - \hat{f}_n(\xi)|$, $c_r(f)$ is a constant depending only on r and f, and $\Lambda_{\xi_1,\ldots,\xi_n}$ denotes the *Lebesgue constant* of the nodes ξ_1,\ldots,ξ_n . 902903 904 Thus, we should choose nodes which have a small Lebesgue constant, and one 905 which grows only slowly with n. This is the case for Chebyshev and Clenshaw– 906 Curtis nodes, for which 907

$$\Lambda_{\xi_1,\ldots,\xi_n}$$

908 Beside uniform convergence there are also classical results on convergence in 909 the L^p sense Nevai (1976, 1980, 1984), e.g., for Gauss-Legendre and Gauss-910Hermite nodes

 $\in \mathcal{O}(\log n).$

911 9

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913
$$\lim_{n \to \infty} \|f - \hat{f}_n\|_{L^p_{\mu}} = 0, \qquad \|f - \hat{f}_n\|_{L^p_{\mu}} = \left(\int_{\Xi} |f(\xi) - \hat{f}_n(\xi)|^p \,\mu(\mathrm{d}x)\right)^{1/p},$$
914

915where $\mu = \mathsf{U}(\Xi)$ or $\mu = \mathsf{N}(0,1)$, respectively. However, if f has low regularity 916or is discontinuous, then convergence may fail or it may take a very large 917number of nodes to approximate f with sufficient accuracy.

918In summary, polynomial collocation constructs a (deterministic) interpolat-919ing polynomial as a surrogate for f based on evaluations of f at n judiciously 920

chosen nodes, for which the error decays with n at a rate depending on the 921 smoothness of f. 922

Gaussian Process Emulation

925The GPE approach consists in applying a method originating in geostatistics, 926 namely the conditioning of Gaussian processes on observations (kriging), to 927 the input-output map of a computer code, which for simplicity we assume to 928 be represented by the scalar-valued function $f:\Xi\to\mathbb{R}$. Again, we are given 929 a finite number of input-output pairs and the goal is to approximate function 930 values at other locations. The uncertainty due to incomplete information at 931 these locations is accounted for in the GPE approach by modeling f as an 932 unknown realization of a Gaussian process indexed by Ξ , which is conditioned 933 on the given observations $y_i = f(\xi_i)$ —analogous to the conditioning of the 934 Gaussian log transmissivity on observational values in Section 2.4.3. In this 935 sense, GPE constructs not only a surrogate \hat{f} for f, but also a probabilistic 936 representation of its pointwise deviation $\hat{f}(\xi) - f(\xi)$. In this sense the term 937 *emulator* designates a statistical approximation of a function, which in this 938 context is referred to as the *simulator* (cf. O'Hagan (2006)). Before we provide 939 a more detailed discussion of this form of *output uncertainty quantification*, we 940 briefly describe how a GPE surrogate is derived. 941

Analogously, to Section 2.3 we first choose a Gaussian process model $G \sim \mathsf{N}(m,c)$ on Ξ with a (parametrized) mean function $m: \Xi \to \mathbb{R}$, e.g.,

$$m(\xi) = m(\xi; \boldsymbol{\beta}) = \sum_{k=1}^{p} \beta_k h_k(\xi), \qquad \boldsymbol{\beta} \in \mathbb{R}^p,$$

and a (parametrized) covariance function $c: \Xi \times \Xi \to \mathbb{R}$, e.g., a Matérn 948 covariance (6) or squared exponential covariance 949

$$c(\xi,\xi') = c(\xi,\xi';\ \sigma^2,\rho) = \sigma^2 \exp(-(\xi-\xi')^2/\rho), \qquad \xi,\xi' \in \Xi.$$
(26) 950
951

952In a true Bayesian approach, prior probability distributions are placed on the 953 hyperparameters β, σ^2, ρ of m and c. For now, however, we assume the covari-954ance c to be fixed and m to be given as linear regression model—in analogy to 955 Section 2.3. Conceptually, the Gaussian process describes our "prior beliefs" 956 about the unknown f in the form of, e.g., characteristic dependencies reflected 957 in the regression functions h_k in the mean model or smoothness properties 958 encoded in the choice of c. Given evaluations $f(\xi_i)$ of f at n design points ξ_i , 959we condition the Gaussian process G on this data and obtain $\hat{G}_n \sim \mathsf{N}(\hat{m}_n, \hat{c}_n)$ 960 with \hat{m}_n and \hat{c}_n determined by the relations for (simple or universal) krig-961 ing, see Section 2.4.3. The resulting surrogate f_n is the conditional mean (or 962

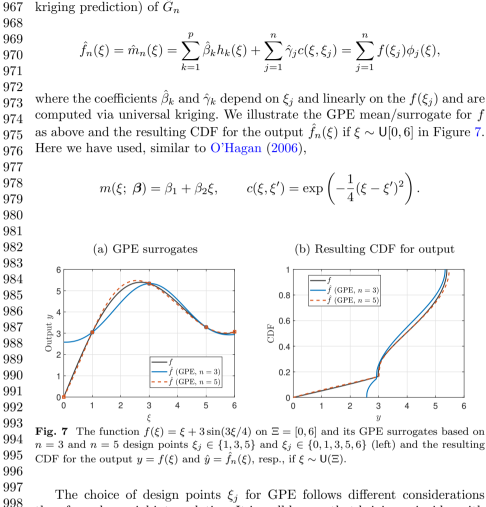
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⁹⁹⁸ The choice of design points ξ_j for GPE follows different considerations ⁹⁹⁹ than for polynomial interpolation. It is well known that kriging coincides with ¹⁰⁰⁰ kernel interpolation, see Scheuerer et al (2013). If we assume for simplicity ¹⁰⁰¹ that $m \equiv 0$ and c is given, then we can straightforwardly apply established ¹⁰⁰² approximation results from kernel interpolation theory by (Narcowich et al, ¹⁰⁰³ with $r \ge 1$ and suitable⁴ covariance functions c such as Matérn kernels (6)

 $\|f - \hat{f}_n\|_{\infty} < C_r(f) D_{\xi_1,\dots,\xi_r}(\Xi)^{r-\frac{1}{2}}$

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1008 where

1009
$$D_{\xi_1,...,\xi_n}(\Xi) := \max_{\xi \in \Xi} \min_{j=1,...,n} |\xi - \xi_j|$$

^{1011 &}lt;sup>4</sup> "Suitable" means here, that the *native* or *reproducing kernel Hilbert space* of *c* coincides with 1012 $H^r(\Xi)$. For more details we refer to Scheuerer et al (2013); Wendland (2004).

denotes the fill distance of the node set $\{\xi_1, \ldots, \xi_n\}$. For the Gaussian covariance function (26) we even obtain exponential convergence if the function f is 1014 analytic, see Wendland (2004), 1015

$$\|f - \hat{f}_n\|_{\infty} \le C(f) \ r^{\mathcal{D}_{\xi_1,\dots,\xi_n}(\Xi)}, \qquad r < 1.$$

 $1016 \\ 1017 \\ 1018$

Thus, for good approximation properties, GPE requires a space filling strat-1019egy for choosing design points, i.e., one which minimizes fill distance. In the1020univariate case that would be avhieved by equispaced points, in stark contrast1021to the optimal equilibrium distribution for interpolation nodes.1022

As mentioned, a GPE not only provides a surrogate f_n but also a 1023probabilistic quantification of the remaining pointwise error $f - \hat{f}_n$, which 10241025represents another important difference to (polynomial) collocation. In order 1026to understand this probabilistic error, recall that the conditioned Gaussian 1027 process G_n can be seen as our "posterior belief" about the unknown f given n evaluations $f(\xi_i)$. Thus, as for the transmissivity field in subsurface flow 1028 1029(which is deterministic but unknown) we model our uncertainty about the 1030true output $f(\xi)$ at a fixed input $\xi \in \Xi$ by $\hat{G}_n(\xi) \sim \mathsf{N}(\hat{f}_n(\xi), \hat{c}_n(\xi))$. This is 1031called *code* or *output uncertainty* in the GPE literature, and is distinct from 1032the *input uncertainty* modelled by random ξ : we have 1033

input uncertainty:
$$\xi$$
 random and $\xi \mapsto f(\xi)$ fixed1034output uncertainty: ξ fixedand $\xi \mapsto f(\xi)$ random10351036

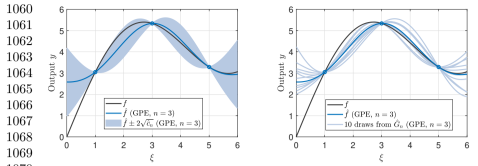
1037Of course, both uncertainty types can be superposed, as we shall see later. We 1038 illustrate the output uncertainty provided by the GPE in Figure 8: the left 1039panel shows f, \hat{f}_n as well as pointwise error estimates for $f - \hat{f}_n$ given by two 1040 times the standard deviation of $\hat{G}_n(\xi)$, which can be also understood as the 1041pointwise 95% credibility region for the unknown $f(\xi)$; the right panel shows 104210 realizations of the Gaussian process \hat{G}_n . Each of these could equally well 1043be used as a surrogate \hat{f}_n in place of \hat{m}_n , since they are also valid (random) 1044guesses for f. In this way, \hat{G}_n provides a random surrogate for f. 1045

Random draws from \hat{G}_n can then be used to quantify the effect of the output uncertainty about the value $f(\xi) \neq \hat{f}_n(\xi)$ within an uncertainty analysis for varying ξ , e.g., for estimating the CDF of $f(\xi)$ when $\xi \sim U(\Xi)$, see, e.g. Oakley and O'Hagan (2002). To explain this in more detail: Regarding the input uncertainty modelled by $\xi \sim U(\Xi)$ we would like to quantify its effect on the outcome by the CDF

$$F(y) = \mathbf{P}(f(\xi) < y).$$
1052
1053

This is a deterministic function for uncertainty analysis for random ξ . However, if we are not able to use f itself to compute F but rather use a GPE \hat{G}_n for f, we can, besides a deterministic approximation of F based on a deterministic 1054 1055 1056 1057

 $1059\,$ (a) GPE surrogate and credibility region $\,$ (b) 10 random draws/surrogates from GP $\,$



1070 Fig. 8 The function $f(\xi) = \xi + 3\sin(3\xi/4)$ on $\Xi = [0, 6]$, its GPE surrogate and the 1071 related 95% credibility region for f (left) as well as 10 paths (or surrogates) drawn from the 1072 conditioned GP \hat{G}_n .

1073

1074 surrogate \hat{f}_n for f

1075

$$F(y) \approx \mathbf{P}_{\xi}(f_n(\xi) < y)$$

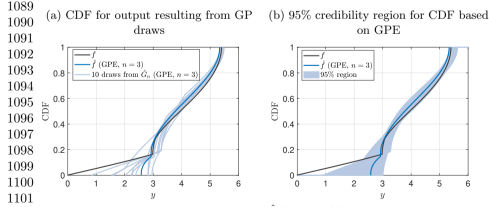
1076 also incorporate our remaining output uncertainty about f via the conditioned 1077 Gaussian process \hat{G}_n for f. This then yields a random CDF

1078

1079 $\hat{F}_n(y) = \mathbf{P}_{\xi}(\hat{G}_n(\xi) < y),$

1080

1081 due to the random \hat{G}_n where we emphasize that the CDF is only w.r.t. ran-1082 domness of the ξ . To illustrate this we show in Figure 9 the resulting CDFs 1083 for $\hat{f}_n(\xi), \xi \sim U(\Xi)$ using $\hat{f}_n = \hat{m}_n$ as well as \hat{f}_n set to be each of the 10 draws 1084 from \hat{G}_n (left) as well as the 95% credibility region for the true (but unknown) 1085 CDF values $F(y) = \mathbf{P}(f(\xi) < y)$ based on 10,000 draws from \hat{G}_n . The cred-1086 ibility region, thus, quantifies our uncertainty about the true CDF resulting 1087 from using a (random) surrogate instead of the true quantity of interest f. 1088



1102 Fig. 9 Resulting CDFs for the output $\hat{y} = \hat{f}(\xi), \xi \sim U(\Xi)$, based on the mean and 10 1103 random draws from the GPE \hat{G}_n (left), and the resulting 95% credibility region for the CDF 1104 of $y = f(\xi)$ derived from the GPE (right).

Discussion

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1106Polynomial collocation and Gaussian process emulators are well-established 1107 surrogate techniques using solely point evaluations of the underlying quantity 1108 of interest f, and both approaches rely on a certain smoothness of f. However, 1109they also differ in several aspects. This includes the type of basis functions 1110 from which each surrogate is constructed (polynomials vs. kernel functions or 1111 radial basis functions) as well as the selection strategies for nodes ξ_i (equilib-1112rium distribution in the sense of potential theory vs. space filling). Moreover, 1113 the GPE surrogate $\hat{f}_n = \hat{m}_n$ is based on minimizing the *average error* w.r.t. an 1114 assumed probability distribution over a function space, whereas interpolation 1115error bounds are obtained by a *worst-case error* analysis over a function class. 1116We refer to Ritter (2000) for more details on these two contrasting approaches. 1117In particular, for GPE we explicitly assume a probability distribution for the 1118 unknown function f, given by the prior Gaussian process model G, whereas 1119 for collocation we simply assume that f is sufficiently smooth. This prior prob-1120ability distribution for f is then updated given the data $f(\xi_i)$ in a Bayesian 1121fashion. Thus, GPE can be related to *Bayesian numerical analysis*, see Dia-1122conis (1988), or *probabilistic numerics*, see Hennig et al (2022), respectively, 1123and be seen as a Bayesian approach to kernel interpolation. In particular, the 1124conditioned (posterior) distribution for the unknown f provided by \hat{G}_n yields 1125an indicator for the remaining (output) uncertainty about f after its evalua-1126tion at n nodes ξ_i . Of course, the assumption of Gaussianity for this computer 1127output uncertainty is debatable. We refer to Bastos and O'Hagan (2009) for 1128diagnostics to validate the GP ansatz as well as to Kracker et al (2010) for a 1129performance study of GPE for "Gaussian" as well as "non-Gaussian" f. 1130

4.2 Polynomial Sparse Grid Collocation

Polynomial collocation in the context of UQ or parametric problems can roughly be described as computing an *M*-variate polynomial approximation to $f: \Xi \to \mathcal{Y}, \Xi \subseteq \mathbb{R}^d$, based on multivariate Lagrange interpolation. Sparse grid collocation uses sparse grids as multivariate interpolation node sets in order to mitigate the curse of dimensionality associated with straightforward tensor-product interpolation for high-dimensional parameter spaces. 1133

While more sophisticated sparse grid techniques have been developed in 1139 recent years, in this work we consider a basic and simple construction known 1140 as (Smolyak) sparse grid collocation introduced for UQ settings, e.g., in Xiu 1141 and Hesthaven (2005); Nobile et al (2008). To this end, assume $f \in C(\Xi; \mathcal{Y})$, 1142 i.e., the mapping f is continuous, and denote by 1143

$$\mathscr{P}_n(\Xi; \mathcal{Y}) = \left\{ \sum_{k=0}^n a_k \xi^k \colon a_k \in \mathcal{Y} \right\}$$
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the space of all \mathcal{Y} -valued univariate polynomials of degree at most n. Then 1147 for a given sequence of univariate node sets $\Xi_k := \{\xi_1^{(k)}, \ldots, \xi_{n_k}^{(k)}\} \subseteq \Xi, k \ge 1$, 1148 where we assume $n_1 = 1$ and $n_k < n_{k+1}$ throughout, we denote the associated 1149 1150

1151 univariate (Lagrange) interpolation operators by

 $\mathcal{I}_k \colon C(\Xi; \ \mathcal{Y}) \to \mathscr{P}_{n_k}(\Xi; \ \mathcal{Y}), \qquad (\mathcal{I}_k f)(\xi) \coloneqq \sum_{j=1}^{n_k} f\left(\xi_j^{(k)}\right) \ell_j^{(k)}(\xi), \quad \xi \in \Xi,$

1157 with $\ell_j^{(k)} \in \mathscr{P}_{n_k}(\Xi; \mathbb{R})$ the Lagrange fundamental polynomials associated 1158 with Ξ_k . The most immediate extension of the interpolation operator to the *M*-dimensional parameter domain Ξ would be the multivariate interpolation 1160 operator $\mathcal{I}_k : C(\Xi; \mathcal{Y}) \to \mathscr{P}_{n_k}(\Xi; \mathcal{Y})$ obtained by tensorization

$$(\mathcal{I}_{\boldsymbol{k}}f)(\boldsymbol{\xi}) := (\mathcal{I}_{k_1} \otimes \cdots \otimes \mathcal{I}_{k_M}) f(\boldsymbol{\xi}) = \sum_{\boldsymbol{j} \leq \boldsymbol{n}_{\boldsymbol{k}}} f\left(\boldsymbol{\xi}_{\boldsymbol{j}}^{(\boldsymbol{k})}\right) \ell_{\boldsymbol{j}}^{(\boldsymbol{k})}(\boldsymbol{\xi}),$$

1165 with multi-indices $\boldsymbol{j} = (j_1, \ldots, j_M), \boldsymbol{n}_{\boldsymbol{k}} = (n_{k_1}, \ldots, n_{k_M}) \in \mathbb{N}^M$, multivari-1166 ate nodes $\boldsymbol{\xi}_{\boldsymbol{j}}^{(k)} = (\boldsymbol{\xi}_{j_1}^{(k_1)}, \ldots, \boldsymbol{\xi}_{j_M}^{(k_M)}) \in \boldsymbol{\Xi}_{\boldsymbol{k}} := \boldsymbol{\Xi}_{k_1} \times \cdots \times \boldsymbol{\Xi}_{k_M}$, and tensorized ¹¹⁶⁷ Lagrange fundamental polynomials $\ell_{j}^{(k)}(\boldsymbol{\xi}) = \ell_{j_1}^{(k_1)}(\xi_1) \cdots \ell_{j_M}^{(k_M)}(\xi_M)$ for $\boldsymbol{\xi} =$ ¹¹⁶⁸ $(\xi_1, \dots, \xi_M) \in \boldsymbol{\Xi}$. However, this construction suffers heavily from the curse of dimensional dimension. 1109 dimensionality since the computational work for evaluating f at all points in 1170 d_{11} c_{12} the Cartesian product grid Ξ_k grows exponentially with dimension M.

Sparse grid constructions, which improve this to polynomial complexity in M, are based on the univariate detail operators

 $\Delta_i = \mathcal{I}_i - \mathcal{I}_{i-1}, \quad i \ge 1, \qquad \mathcal{I}_0 \equiv 0,$

so that $\mathcal{I}_k = \sum_{i=1}^k \Delta_i$, yielding the tensor product interpolation operator as

$$\mathcal{I}_k f = \sum_{i \leq k} \Delta_i f, \qquad \Delta_i = \Delta_{i_1} \otimes \cdots \otimes \Delta_{i_M}.$$

By contrast, the (Smolyak) sparse grid collocation operator is defined by Μ

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$$\mathcal{S}_{\ell,M}f := \sum_{|\boldsymbol{i}-\boldsymbol{1}|_1 \leq \ell} \Delta_{\boldsymbol{i}}f, \qquad |\boldsymbol{i}-\boldsymbol{1}|_1 := \sum_{j=1} |i_j-1|, \qquad \ell \geq 0.$$

By combinatorical arguments, one can obtain the equivalent combination technique representation

$$\mathcal{S}_{\ell,M}f = \sum_{\ell-M+1 \le |\mathbf{i}-\mathbf{1}| \le \ell} (-1)^{\ell+M-|\mathbf{i}|} \binom{M-1}{\ell+M-|\mathbf{i}|} \mathcal{I}_{\mathbf{i}}f,$$

which expresses the Smolyak operator as a linear combination of selected M-1197 variate tensor product interpolation operators. For the associated sparse grid 1198

$$\Xi_{\ell,M} := \bigcup \Xi_i$$
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$$\ell - M + 1 \le |i - 1| \le \ell$$
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consisting of all multivariate nodes occurring in these representations, the car-1203dinality $|\Xi_{\ell,M}|$ grows only polynomially w.r.t. M (cf. Novak and Ritter (1999)), 1204 1205 while the overall order of accuracy remains close to that of the full tensor product $\mathcal{I}_{(\ell+1,\ldots,\ell+1)}$. In particular, it can be shown (Bäck et al, 2011, Proposition 12061) that $\mathcal{S}_{\ell,M}$ is a projection on 1207

$$\mathscr{P}_{\ell,M}(\Xi; \ \mathcal{Y}) := \sum_{|i-1| \leq \ell} \mathscr{P}_{n_{i_1}}(\Xi; \ \mathcal{Y}) \otimes \cdots \otimes \mathscr{P}_{n_{i_M}}(\Xi; \ \mathcal{Y}).$$

1212 Note, however, that in general $S_{\ell,M}$ is not interpolatory unless the univariate 1213nodes sets are nested $\Xi_k \subset \Xi_{k+1}$ (Barthelmann et al, 2000, Proposition 6). 1214The latter is the case for Clenshaw–Curtis nodes with the "doubling sequence" $n_k = 2^k - 1$ ($k \ge 1$), or (weighted) Leja nodes with linear growth $n_k = k$ Ernst 1215et al (2021). In the following, we shall use the non-nested nodal sequence of 1217 *Gauss-Hermite* nodes, i.e., the roots of Hermite polynomials. This choice is 1218 common for collocation applied to functions of Gaussian random variables, see 1219 Babuška et al (2007); Nobile et al (2008); Ernst and Sprungk (2014). 1220

Convergence and Application

1222If f is sufficiently smooth then $S_{\ell,M}f$ can be shown to converge to f, 1223specifically

$$\|f - \mathcal{S}_{\ell,M}f\|_{L^2_{\mu}} \in \mathcal{O}\left(|\Xi_{\ell,M}|^{-r}\right),$$

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for an r < 1 using Gauss-Hermite nodes $\xi_i^{(k)}$ with linear growth $n_k = k$ or doubling growth $n_k = 2^{k-1} + 1$ $(k \ge 1)$, see, e.g., Ernst and Sprungk (2014); 12261227Ernst et al (2018). The rate of convergence r w.r.t. the number of collocation 1228points depends, of course, on the smoothness class of f. 1229

It was shown in (Ernst and Sprungk, 2014, Section 3) that the solution 1230 (\boldsymbol{u}, p) of the random/parametric mixed variational problem (4) allows for a 1231holomorphic extension into \mathbb{C}^M under suitable assumptions, which are satis-1232fied by truncated KL expansions (19) of a lognormal transmissivity field. Thus, 1233applying $\mathcal{S}_{\ell,M}$ to approximate the solution map $(\boldsymbol{u}, p) \colon \boldsymbol{\Xi} \to \mathcal{V} \times \mathcal{W}$ is justi-1234fied. By contrast, the quantity of interest given by the breakthrough time f_{exit} 1235may, in general, not even be a continuous function of the parameters $\boldsymbol{\xi}$, as 1236is immediate from considering the case of a particle grazing the exit bound-1237ary and returning into the domain for a particular parameter setting. Thus, 12381239applying $S_{\ell,M}$ to approximate f_{exit} directly may lead to inaccurate surrogate approximation or even divergence with increasing $|\Xi_{\ell,M}|$. 1240

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1243 However, a simple remedy is to use the surrogate

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- 1245 $\hat{f}_{\text{exit},\ell} = G_{\text{exit}} \left(\mathcal{S}_{\ell,M} \boldsymbol{u} \right)$
- 1246

1247 where $G_{\text{exit}}: \mathcal{V} \to \mathbb{R}$ denotes the mapping from a velocity field on D to the log 1248 breakthrough time of a particle following this field released at \mathbf{x}_0 at time t = 0, 1249 which is inexpensive to evaluate compared to solving the Darcy flow equations. 1250 Then, since L^2 -convergence implies convergence in distribution, assuming that 1251 the set of points of discontinuity of the mapping G_{exit} has probability measure 1252 zero, we have by the continuous mapping theorem

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1254 $\lim_{\ell \to \infty} \|F - \hat{F}_{\ell}\|_{\infty} = 0, \qquad \hat{F}_{\ell}(s) := \mathbf{P}_{\boldsymbol{\xi} \sim \mu} \left(G_{\text{exit}} \left(\mathcal{S}_{\ell,M} \boldsymbol{u}(\boldsymbol{\xi}) \right) \leq s \right),$ 1255

1256 where F denotes the true CDF of f_{exit} Thus, we are assured convergence of 1257 the CDF based on the surrogate $S_{\ell,M} u$ for the true velocity u to the true 1258 CDF for the breakthrough time.

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$\frac{1260}{1261}$ **4.3 Gaussian Process Emulators**

1262 Having described basic GPE methodology in Section 4.1, we now turn to 1263 the construction of GPEs for multivariate scalar-valued functions $f: \Xi \to \mathbb{R}$. 1264 Again, the approach is similar to multivariate geostatistics. We shall con-1265 sider the *full Bayesian* approach to GPE (cf. Kennedy and O'Hagan (2001); 1266 O'Hagan (2006)), which also entails specifying prior distributions for the 1267 hyperparameters contained in the mean and covariance functions which are 1268 also conditioned on the evaluations of f at the design points $\boldsymbol{\xi}_j$. As before, we 1269 start with a linear regression model for the mean

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$$m: \Xi \to \mathbb{R}, \qquad m(\boldsymbol{\xi}) = m(\boldsymbol{\xi}; \boldsymbol{\beta}) = \sum_{k=1}^{p} \beta_k h_k(\boldsymbol{\xi}), \quad \boldsymbol{\beta} \in \mathbb{R}^{p},$$

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¹²⁷⁴ with known regression functions $\boldsymbol{h} = (h_1, \ldots, h_p), h_k : \boldsymbol{\Xi} \to \mathbb{R} \ (h_1 \equiv 1 \text{ and} 1275 \ h_2(\boldsymbol{\xi}) = \boldsymbol{\xi}$ are common choices) and unknown coefficients $\boldsymbol{\beta} = (\beta_1, \ldots, \beta_p)^{\top}$. ¹²⁷⁶ For the emulator's covariance function $c : \boldsymbol{\Xi} \times \boldsymbol{\Xi} \to \mathbb{R}$ we fix the squared exponential kernel ¹²⁷⁸

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$$\begin{array}{l} 1279\\ 1280\\ 1280 \end{array} \quad c(\boldsymbol{\xi}, \boldsymbol{\xi}') = c(\boldsymbol{\xi}, \boldsymbol{\xi}'; \ \sigma^2, B) = \sigma^2 \exp(-(\boldsymbol{\xi} - \boldsymbol{\xi}')^\top B(\boldsymbol{\xi} - \boldsymbol{\xi}')), \qquad \boldsymbol{\xi}, \boldsymbol{\xi}' \in \Xi, \ (27) \end{array}$$

1281 where $\sigma^2 > 0$ is the marginal variance and $B = \text{diag}(b_1, \ldots, b_M) \in \mathbb{R}^{M \times M}$, 1282 $b_i > 0$ is a matrix of so-called *smoothness parameters*. For the squared exponen-1283 tial covariance (27) and choices for h_1 and h_2 mentioned above, it is known that 1284 the realizations of the Gaussian process are almost surely analytic w.r.t. $\boldsymbol{\xi}$. For 1286 1287

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other covariance functions, such as the family of Matérn kernels, one obtains 1289Gaussian processes with realizations of different smoothness orders.⁵ 1290

Thus, for fixed given β , σ , and B, the (prior) Gaussian process model for 1291the output of f for an arbitrary input $\boldsymbol{\xi} \in \boldsymbol{\Xi}$ is 1292

$$f(\boldsymbol{\xi}) \sim \mathsf{N}(m(\boldsymbol{\xi}; \boldsymbol{\beta}), c(\boldsymbol{\xi}, \boldsymbol{\xi}; \sigma^2, B)).$$
 1294

Similarly, for fixed $\boldsymbol{\beta}, \sigma$, and B, the vector $\boldsymbol{f} = (f(\boldsymbol{\xi}_1), \dots, f(\boldsymbol{\xi}_n))^{\top}$ of values 1296 of the Gaussian process at a set of design points $\{\boldsymbol{\xi}_1, \dots, \boldsymbol{\xi}_n\}$ has the *n*-variate 1297 1298 Gaussian distribution

$$\boldsymbol{f} = (f(\boldsymbol{\xi}_1), \dots, f(\boldsymbol{\xi}_n))^\top \sim \mathsf{N}(\boldsymbol{H}\boldsymbol{\beta}, \boldsymbol{C}_{\sigma^2, B})$$
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where $\boldsymbol{H} = (h_k(\xi_j)) \in \mathbb{R}^{n \times p}$ and $\boldsymbol{C}_{\sigma^2,B} = (c(\xi_i,\xi_j; \sigma^2,B)) \in \mathbb{R}^{n \times n}$. We 1302denote the probability density of this random vector $\mathbf{f} \in \mathbb{R}^n$ by 1303

$$p(\boldsymbol{f} \mid \boldsymbol{\beta}, \sigma, B) \propto \exp\left(-\frac{1}{2}(\boldsymbol{f} - \boldsymbol{H}\boldsymbol{\beta})^{\top} \boldsymbol{C}_{\sigma^{2}, B}^{-1}(\boldsymbol{f} - \boldsymbol{H}\boldsymbol{\beta})\right).$$

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Suitable values for the parameters β , σ , and B are usually not known a priori 1308and should be inferred based on the evaluations f. This is typically done in 1309a Bayesian fashion, i.e., we choose hyperpriors for these parameters which 1310 are then conditioned on the data $\boldsymbol{f} = (f(\boldsymbol{\xi}_1), \dots, f(\boldsymbol{\xi}_n))^{\top}$. Common choices 1311 for (β, σ^2) are a normal-inverse-gamma prior or a Jeffreys prior with density 1312 $p(\beta, \sigma^2) \propto \sigma^{-2}$ (cf. Oakley and O'Hagan (2002); Stone (2011)) since these 1313 allow for closed-form expressions for the resulting (marginal) posteriors. Given 1314 evaluations f, the resulting posterior for the parameters (β, σ^2) is then 1315

$$p(\boldsymbol{\beta}, \sigma^2 \mid \boldsymbol{f}, B) \propto p(\boldsymbol{f} \mid \boldsymbol{\beta}, \sigma, B) p(\boldsymbol{\beta}, \sigma^2).$$
 1317

For the estimation of the smoothness parameters B a "full" Bayesian infer-1319 ence based on data f would require Markov chain Monte Carlo simulations. 1320Instead, one often simply computes a point estimate based on maximizing 1321the marginal likelihood $p(\boldsymbol{f} \mid B) \propto \int p(\boldsymbol{\beta}, \sigma^2 \mid \boldsymbol{f}, B) p(\boldsymbol{\beta}, \sigma^2) \, \mathrm{d}\boldsymbol{\beta} \, \mathrm{d}\sigma^2$ for which 1322analytic formulas are available (Stone, 2011, Section 2.3.4). This often yields 1323competitive results to a full Bayesian inference Kracker et al (2010). 1324

Given f, the posterior density for the output $f(\boldsymbol{\xi})$ at new location $\boldsymbol{\xi}$ is then 1325

$$p(f(\boldsymbol{\xi}) \mid \boldsymbol{f}, \boldsymbol{\beta}, \sigma, B) \propto p(\boldsymbol{f} \mid \boldsymbol{\beta}, \sigma, B) p(\boldsymbol{\beta}, \sigma^2 \mid \boldsymbol{f}, B).$$

Marginalization by integrating out β and σ^2 can be done analytically for a 1329normal-inverse-gamma or Jeffreys prior $p(\beta, \sigma^2)$ and results in a Student-t 1330

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¹³³² ⁵We have also explored other covariance models such as the Matérn kernels for GPE surrogates; however, the overall conclusions in the numerical experiments were about the same as for the 1333 squared exponential (27).

1335 process (cf. Shah et al (2014)) for the prediction of the output of f, i.e., 1336

$$f(\boldsymbol{\xi}) \mid \boldsymbol{f} \sim t_{n-p} \left(\hat{m}_n(\boldsymbol{\xi}), \hat{\sigma}^2 \hat{c}_n(\boldsymbol{\xi}, \boldsymbol{\xi}) \right), \qquad (28)$$

 $\begin{array}{c} 1337\\ 1338 \end{array}$

1339 where \hat{m}_n and \hat{c}_n are the mean and covariance obtained by universal krig-1340 ing applied to f given the observations f (see (??) and (??)) with $\sigma^2 = 1$, 1341 respectively, and where $\hat{\sigma}^2$ is given by

$$\hat{\sigma}^{2} = \frac{1}{n-p} \boldsymbol{f}^{\top} \boldsymbol{C}^{-1/2} \left(\boldsymbol{I} - \boldsymbol{C}^{-1/2} \boldsymbol{H} \left(\boldsymbol{H}^{\top} \boldsymbol{C}^{-1} \boldsymbol{H} \right)^{-1} \boldsymbol{H}^{\top} \boldsymbol{C}^{-1/2} \right) \boldsymbol{C}^{-1/2} \boldsymbol{f}.$$

1345 1346 For the prediction of f at multiple new points we obtain a multivariate Student-1347 *t*-distribution with mean vector given by the evaluation of \hat{m}_n at those points 1348 and covariance matrix given by evaluating $\hat{\sigma}^2 \hat{c}_n$.

Regarding the choice of the design points for multivariate GPE we require again space filling designs. For compact $\Xi \subset \mathbb{R}^M$ these are, e.g., Sobol' points (Owen et al, 2017) or Latin hypercube designs (Viana, 2015). The latter extend also to $\Xi = \mathbb{R}^M$ w.r.t. $\mu = \mathsf{N}(0, \mathbf{I})$ as we require for the WIPP problem. As for the appropriate number $n \in \mathbb{N}$ of training points $\Xi_n = \{\xi_1, \ldots, \xi_n\} \subset \Xi$, a common rule of thumb calls for n = cM (Loeppky et al, 2009) with a factor $1355 c \geq 10$.

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$\frac{1350}{1357}$ Convergence and Application

1358 Since the GPE surrogate $\hat{f}_n = \hat{m}_n$ and its covariance \hat{c}_n are derived by uni-1359 versal kriging, we can again exploit the relation between kriging and kernel 1360 interpolation Scheuerer et al (2013). Again, assume $m \equiv 0$ for simplicity and 1361 c fixed as in (27). Then for compact $\Xi \subset \mathbb{R}^M$ and analytic $f : \Xi \to \mathbb{R}$ we have 1362

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$$||f - \hat{f}_n||_{\infty} \le C(f) \ r^{\mathsf{D}_{\boldsymbol{\xi}_1,\dots,\boldsymbol{\xi}_n}(\boldsymbol{\Xi})},$$

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1365 for a 0 < r < 1 as well as

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$$\hat{c}_n(x,x) \le C \ r^{2\mathrm{D}_{\boldsymbol{\xi}_1,\dots,\boldsymbol{\xi}_n}(\boldsymbol{\Xi})}$$

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1369 Thus, besides uniform convergence of the surrogate $\hat{f}_n \to f$, we also have 1370 vanishing output uncertainty regarding $f(\boldsymbol{\xi})$ as $n \to \infty$ —which is a consistency 1371 statement for the posterior for f here given by the Gaussian or Student-t1372 process \hat{G}_n . However, to our knowledge, no L^2 -convergence statements are 1373 available for the case of unbounded $\boldsymbol{\Xi} = \mathbb{R}^M$, as the setting of the WIPP 1374 problem would require.

1375 In the next section we will apply GPE to approximate the quantity of inter-1376 est f_{exit} directly. Thus, for convergence with $n \to \infty$, we require f_{exit} to be 1377 sufficiently smooth (see above) which may not be the case in general. How-1378 ever, it may well be that the surrogate \hat{f}_n and the related output uncertainty 1379 provided by the GPE for finite n = cM design points is sufficiently accurate 1380 for CDF estimation. We note that also vector-valued GPE are available, see

Álvarez et al (2012); Bilionis et al (2013); Cleary et al (2021); Higdon et al 1381(2008). Hence, we could apply a GPE to approximate the FE solution of the 1382random parametric variational problem (which depends analytically on $\boldsymbol{\xi}$, see 1383 comment above) and proceed as for polynomial collocation to provide approx-1384imate samples of $f_{\text{exit}}(\boldsymbol{\xi})$. We do not consider this option in this work, since 1385the FE space is very high dimensional (of order 10^4) and thus the GPE would 1386 involve too many parameters to estimate based on less than 20,000 design 13871388 points.

5 Numerical Results

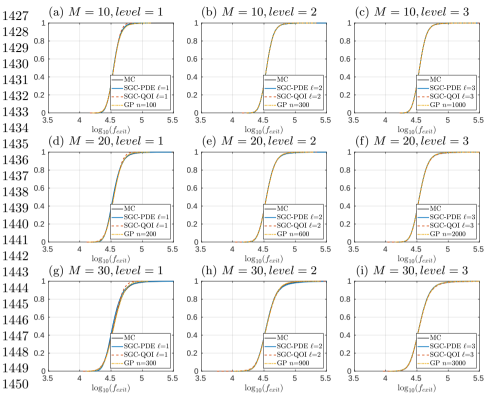
We now perform a numerical study comparing sparse grid polynomial collocation and Gaussian process emulators as surrogates for the task of approximating the CDF of the exit time $f_{\text{exit}}(\boldsymbol{\xi})$ using M terms or coefficients $\boldsymbol{\xi} \sim N(0, \boldsymbol{I})$ 1394 in the truncated KL expansion of the log transmissivity field $Z = \log T$. We vary M = 10, 20, 30 and apply the following three surrogate approaches: 1396

- SGC-PDE: We apply Smolyak sparse grid polynomial collocation $S_{\ell,M}$ 1397 to approximate the solution pair (\boldsymbol{u},p) of the weak mixed formulation 1398 and then obtain approximate samples $\hat{f}_{\text{exit}}(\boldsymbol{\xi}_i)$ of the exit time by solving 1399 the particle transport given the approximate velocity field $S_{\ell,M}\boldsymbol{u}(\boldsymbol{\xi}_i)$, i.e., 1400 $\hat{f}_{\text{exit}}(\boldsymbol{\xi}_i) = G_{\text{exit}}(S_{\ell,M}\boldsymbol{u}(\boldsymbol{\xi}_i))$ where $\boldsymbol{\xi}_i \sim N(0, \boldsymbol{I}), i = 1, \dots, N$ iid. 1401
- SGC-QoI: We apply Smolyak sparse grid polynomial collocation $S_{\ell,M}$ directly to approximate the exit time $f_{\text{exit}}(\boldsymbol{\xi}_i)$ and in this way obtain approximate samples via $\hat{f}_{\text{exit}}(\boldsymbol{\xi}_i) = S_{\ell,M} f_{\text{exit}}(\boldsymbol{\xi}_i)$ where $\boldsymbol{\xi}_i \sim \mathsf{N}(0, \boldsymbol{I})$, $i = 1, \dots, N$ iid.
- **GPE:** We apply Gaussian process emulation to approximate the exit time $f_{\text{exit}}(\boldsymbol{\xi}_i)$ and obtain approximate samples via $\hat{f}_{\text{exit}}(\boldsymbol{\xi}_i) = \hat{m}_n(\boldsymbol{\xi}_i)$ where $\boldsymbol{\xi}_i \sim \mathsf{N}(0, \boldsymbol{I}), i = 1, \dots, N$ iid and \hat{m}_n denotes the GPE mean. 1406 1407 1408 1409

For each surrogate we generate N = 20000 approximate samples of the 1410 quantity of interest and compare these to N = 20000 samples of the "true" 1411 f_{exit} evaluated by solving the Darcy flow equations and particle transport 1412problem each time (denoted MC for Monte Carlo in the following). The num-1413ber $N = 20\,000$ of samples is derived from the error criterion outlined in 1414Section 3.3. For SGC we use different levels $\ell = 1, 2, 3$, and for the GPE dif-1415ferent numbers of design points n = cM with c = 10, 20, 30, 50, 100. We show 1416the resulting empirical CDFs for the log exit time in Figure 10. It is apparent 1417 that, for each M = 10, 20, 30 all surrogate methods yield a very good fit to 1418 the reference ECDF obtained by the plain Monte Carlo approach. Slight devi-1419ations can be seen for the lowest level $\ell = 1$ for SGC-QoI, but at least for 1420 $\ell \geq 2$ it is difficult to distinguish the four ECDFs. Therefore, we take a closer 1421look at the performance of the surrogates in Table 2, where we report the 1422resulting values of the KS statistic $K = \sup_{s \in \mathbb{R}} \left| \hat{F}_n(s) - F_n(s) \right|$ of the empir-14231424ical CDF F_n obtained by Monte Carlo sampling of f_{exit} and the empirical 1425CDF F_n obtained by Monte Carlo sampling of the surrogate f_{exit} . Moreover, 1426

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1451 Fig. 10 Empirical CDFs obtained by Monte Carlo, SGC and GPE surrogates for different 1452 lengths M of the KLE.

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1454 we indicate by an asterisk that the error K in the ECDFs is negligible, i.e., 1455 that the Kolmogorov–Smirnov test is passed (at significance level $\alpha = 0.05$), 1456 and hence there is no indication that the samples were drawn from different 1457 distributions. We make the following observations:

1458• For M = 10,20 all three surrogates pass the KS-test at least for level 1459 $\ell > 2$ (SGC) or n > 30M design points (GPE). For M = 30 this is 1460also the case for SGC-PDE with $\ell \geq 2$ and GPE with n = 100M. Thus, by employing the considered surrogates we can obtain an ECDF for the 1461exit time which is essentially indistinguishable (for $\alpha = 0.05$) from the 14621463"true" ECDF but which required just a fraction of the computational cost of the latter. Indeed, compared to $N = 20\,000$ solutions of the Darcy 14641465flow equations, we require merely between $\approx 200 \ (M = 10)$ and ≈ 2000 1466(M = 30) PDE solves when a surrogate is used.

• For SGC-PDE we observe a steep increase in n with M but overall a robust and good performance.

• For the SGC-QoI approach we observe a significantly worse performance for M = 30 which may be due to insufficient (mixed) smoothness of f_{exit} . 1471 1472

Surrogate		M = 10		M = 20		M = 30	
		n	K	n	K	n	K
GC-PDE	$\ell = 1$	21	0.0128^{*}	41	0.0281	61	0.0495
GC-PDE	$\ell = 2$	241	0.0028^{*}	881	0.0045^{*}	1921	0.0118^{*}
GC-PDE	$\ell = 3$	2001	0.0019^{*}	13201	0.0023^{*}	41601	0.0052^{*}
GC-QOI	$\ell = 1$	21	0.0271	41	0.0293	61	0.0435
GC-QOI	$\ell = 2$	241	0.0065^{*}	881	0.0088^{*}	1921	0.0196
GC-QOI	$\ell = 3$	2001	0.0048*	13201	0.0089^{*}	41601	0.0138
PE	c = 10	100	0.0136	200	0.0245	300	0.0309
PE	c = 20	200	0.0092^{*}	400	0.0191	600	0.0228
PE	c = 30	300	0.0062^{*}	600	0.0116^{*}	900	0.0171
PE	c = 50	500	0.0041^{*}	1000	0.0070^{*}	1500	0.0141
PE	c = 100	1000	0.0031^{*}	2000	0.0064^{*}	3000	0.0087^{*}

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Table 2 Performance of the SGC and GPE surrogates for different lengths M of the KL expansion measured by the value of the resulting KS statistic K. Here, n refers to the number of PDEs to be solved for building the surrogate and an asterisk denotes that the KS-test was passed at significance level $\alpha = 0.05$.

• For the GPE approach we observe deteriorating performance for increasing M, i.e., we require a larger factor c for the numer of design points 1490 n = cM in order to pass the KS test and have small values of K (c = 20 1491 for M = 10, c = 30 for M = 20 and c = 100 for M = 30). This may be 1492 due to the curse of dimensionality for kernel interpolation methods. 1493

Changing the trend model for $\log T$

1496Despite the overall positive observations for the employed surrogates made so 1497 far we report how the outcome may change if we simply use a different trend 1498model for the mean of the log transmissivity field $\log T$. Instead of using the 1499constant, linear in x_1 , and zone indicator regression functions h_1 , h_2 , and h_5 , 1500respectively, see (9), we only use the constant h_1 . This leads to a different 1501Matérn covariance function used for $\log T$, see Table 1 and thus also to different 1502eigenvalues and eigenfunctions in the KL expansion. Moreover, the smooth-1503ness properties of the mapping $\boldsymbol{\xi} \mapsto f_{\text{exit}}(\boldsymbol{\xi})$ may change as well. In fact, in 1504Table 3 we observe a much diminished performance of all three surrogate tech-1505niques: Now only SGC-PDE passes the KS test and only for the shorter KL 1506truncation kength M = 10, 20. However, SGC-PDE and GPE provide a visu-1507ally acceptable fit to the reference ECDF in Figure 11, whereas we clearly see 1508a deterioration for the SGC-QoI surrogate. This distinctly worse performance 1509of SGC-QoI may be due to lacking smoothness of $\boldsymbol{\xi} \mapsto f_{\text{exit}}(\boldsymbol{\xi})$ in this case.

1510For the GPE surrogate we also evaluate to what extent the accompanying 1511Gaussian model for this surrogate's output uncertainty covers the deviation 1512from the reference CDF. To this end, we focus on the setting where the GPE 1513surrogate performs worst, i.e., M = 30 using n = 300 design points, and 1514compute a 95% credibility region for the CDF based on 10000 random draws 1515of surrogates from the trained GPE. The results are reported in Figure 12 for 1516both trend models. We observe that the Gaussian output uncertainty model 1517appears overconfident in the case of the constant trend model. Thus, this 1518

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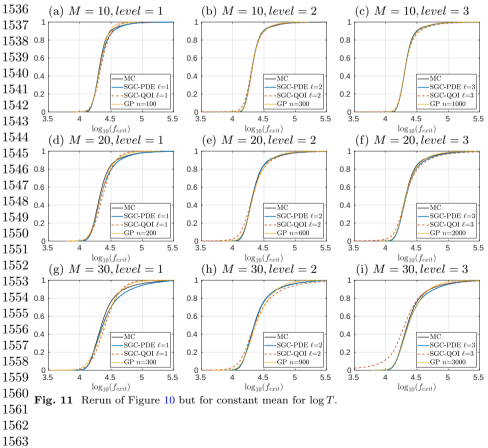
1519	Surrog	Surrogate		M = 10		M = 20		= 30
1520			n	K	n	K	n	K
	SGC-PDE	$\ell = 1$	21	0.0537	41	0.0653	61	0.0621
1521	SGC-PDE	$\ell = 2$	241	0.0123^{*}	881	0.0130^{*}	1921	0.0146
1522	SGC-PDE	$\ell = 3$	2001	0.0121^{*}	13201	0.0345	41601	0.0387
1523	SGC-QOI	$\ell = 1$	21	0.1099	41	0.1340	61	0.1301
1524	SGC-QOI	$\ell = 2$	241	0.0485	881	0.0798	1921	0.0697
-	SGC-QOI	$\ell = 3$	2001	0.0369	13201	0.0577	41601	0.1711
1525	GPE	c = 10	100	0.0366	200	0.0546	300	0.0815
1526	GPE	c = 20	200	0.0373	400	0.0415	600	0.0591
1527	GPE	c = 30	300	0.0153	600	0.0368	900	0.0615
1528	GPE	c = 50	500	0.0188	1000	0.0405	1500	0.0415
	GPE	c = 100	1000	0.0192	2000	0.0258	3000	0.0422
1529 <u>-</u>	Table 3 Rerun of	Table 2 b	ut for co	onstant me	an for log	g T.		
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34 UQ Surrogates for Groundwater Flow

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1532 experiment indicates that a sufficiently good performance of the surrogates for 1533 CDF estimation of exit times may depend on various aspects of the problem— 1534 such as the choice of the trend model for the log transmissivity field. 1535



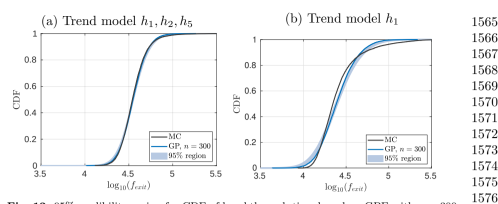


Fig. 12 95% credibility region for CDF of breakthrough time based on GPE with n = 300 for M = 30 KL terms for different trend models

Convergence Study

1580The negative results for the constant trend model raise the question whether 1581we simply did not use enough design points n or sparse collocation level ℓ 1582for the GPE and SGC surrogates, respectively, or whether the quantity of 1583interest is simply to rough to be approximated well by these methods. To 1584this end, we perform a convergence study for both scenarios: constant trend 1585model and "best" trend model using h_1 , h_2 , and h_5 in (9). We report the 1586associated L^2_{μ} -errors of the SGC surrogates for the flux \boldsymbol{u} and the quantity of 1587interest in Tables 4 and 5, respectively. We notice significantly larger errors for 1588 the constant trend model. In order to allow for a sufficiently high polynomial 1589degree for SGC to observe a significant error decay, we restrict ourselves to the 1590low-dimensional case of M = 2 and M = 5 KL terms. We report the resulting 1591errors of the velocity and the quantity of interest in Figure 13. There we clearly 1592observe a decaying error for increasing level ℓ and number of sparse grid nodes 1593 $|\Xi_{\ell,M}|$, respectively. Moreover, we observe that the rate of convergence for both 1594quantities is affected by the larger number of KL terms and the choice of the 1595trend model. The former was already observed in Ernst and Sprungk (2014). 1596The latter is also related to an observation made in Ernst and Sprungk (2014): 1597since the constant trend model yields a larger estimated value for the variance 1598 σ^2 , this in turn leads to a slower convergence rate of SGC. 1599

Regarding the application of GPE to approximate the quantity of interest, 1600we perform a similar study as for SGC using M = 2 and M = 5 KL terms. 1601The results are displayed in Figure 14. We observe that the L^2_{μ} -error (left 1602panel) does not decay with increasing number of design points, at least not 1603in the applied regime of up to n = 1000M design points. However, this is 1604mainly due to high approximation errors in the tail regions of the distribution 1605of $\boldsymbol{\xi} \sim \mu = \mathsf{N}(0, \boldsymbol{I})$. espite this, we observe a decay of the KS test statistic 1606value K, i.e., the L^{∞} -error of the ECDF for the quantity of interest, except 1607for M = 5 and the constant trend model.

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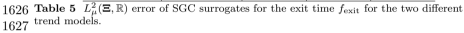
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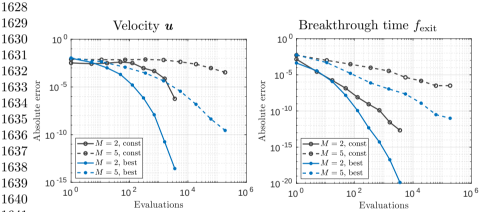
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1611	Trend model	Surrogate		M = 10	M = 20	M = 30
1612		SGC	$\ell = 1$	5.9897E-3	1.2933E-2	1.6810E-2
1613	h_1, h_2, h_5	SGC	$\ell = 2$	2.1354E-3	6.3868E-3	9.3400E-3
1614		SGC	$\ell = 3$	6.1168E-4	2.5686E-3	4.3738E-3
		SGC	$\ell = 1$	4.0723E-2	1.1149E-1	1.7963E-1
1615	h_1	SGC	$\ell = 2$	4.0331E-2	1.1113E-1	1.7329E-1
1616		SGC	$\ell = 3$	3.9595E-2	1.0598E-1	1.6928E-1

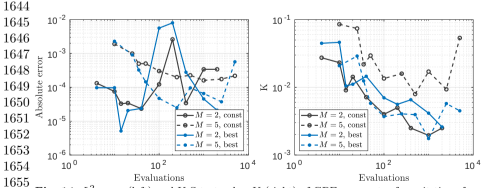
1617 Table 4 $\overline{L^2_{\mu}(\Xi, H(\text{div}; D))}$ error of SGC surrogates for the flux u for the two different 1618 trend models.

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1620	Trend model	Surrogate		M = 10	M = 20	M = 30
1621		SGC	$\ell = 1$	1.2296E-3	2.7434E-3	6.0602E-3
1622	h_1, h_2, h_5	SGC	$\ell = 2$	1.2699E-4	4.8917E-4	2.1426E-3
1623		SGC	$\ell = 3$	2.0075E-5	9.6514E-5	4.4401E-4
		SGC	$\ell = 1$	7.0990E-3	1.5259E-2	2.8396E-2
1624	h_1	SGC	$\ell = 2$	2.9464E-3	7.7314E-3	1.5502E-2
1625		SGC	$\ell = 3$	1.9730E-3	9.2632E-3	1.8001E-2





1641 Fig. 13 L^2_{μ} -error of SGC surrogates for the velocity \boldsymbol{u} (left) and exit time f_{exit} (right). 1642 For the flux we used the norm in $\boldsymbol{H}(\text{div}; D)$ to quantify the difference between $\boldsymbol{u}(\boldsymbol{\xi})$ and 1643 $\mathcal{S}_{\ell,M}\boldsymbol{u}(\boldsymbol{\xi})$.





6 Conclusion

In this work we have presented a complete uncertainty propagation workflow 1659for groundwater flow and particle transport simulations based on a real-world 1660 application related to the site performance assessment for a nuclear waste 1661 repository. We described in detail the construction of a stochastic model for 1662 an uncertain transmissivity field by geostatistical methods using the available 1663 observational data. Our main focus was the direct comparison of two estab-1664lished surrogate approaches for uncertainty propagation analysis: sparse grid 1665stochastic collocation and Gaussian process emulation. Both methods origi-1666 nate from different communities, i.e., numerical analysis and computational 1667 statistics, respectively. Our purpose was to describe the fundamental ideas and 1668 principles underlying both methods and compare their performance for the UQ 1669 problem under consideration, specifically for CDF estimation of scalar quanti-1670 ties of interest, in this case the travel time of groundwater-borne radionuclides. 1671 The overall conclusion is that both methods can achieve significant reduction 1672in computational cost, reducing the computational burden by a factor of 10 1673 to even 100 in some cases considerered. Moreover, we have observed that the 1674GPE surrogate seems to be more adversely affected by the high dimensionality 1675of the input space compared with sparse grid collocation, which is not surpris-1676 ing given the bad scaling of the filling distance with dimension. On the other 1677 hand, stochastic collocation must also be applied with care, since the quan-1678tity to be approximated has to depend sufficiently smoothly on the random 1679 inputs—such as the solution of the random PDE. However, the remarkable 1680 performance of both surrogates seems to be affected by modelling choices for 1681 the random log transmissivity field such as choice of the trend or regression 1682 model for the mean. Although this effect could be explained mathematically 1683in our case, it does place limitations on the practical benefits of UQ surrogate 1684methods for CDF estimation in groundwater flow applications. 1685

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A Restricted Maximum Likelihood Estimation

1945Under the models for the mean (8) and covariance structure (6), the Gaussian 1946 log-transmissivity field (7) has the form 1947

$$Z(\boldsymbol{x},\omega) = \boldsymbol{h}(\boldsymbol{x})^{\top} \boldsymbol{\beta} + \widetilde{Z}(\boldsymbol{x},\omega)$$
(29) 1948
1949

1950with $\boldsymbol{h}(\boldsymbol{x})^{\top} = [h_1(\boldsymbol{x}), \dots, h_k(\boldsymbol{x})]$ denoting the vector of regression functions 1951evaluated at $x \in D$ and $\beta \in \mathbb{R}^k$ the vector of regression coefficients. The resid-1952ual field \widetilde{Z} is Gaussian with zero mean. The covariance function $c_{\theta}(x, y)$ of 1953Z (and \widetilde{Z}) is given by (6), where $\boldsymbol{\theta} = (\sigma, \rho, \nu)$ denotes the triplet of parame-1954ters consisting of standard deviation σ , correlation length ρ and smoothness 1955parameter ν . The specification of the probabilistic model for the random field 1956 Z consists in determining the vector $\boldsymbol{\beta}$ of regression coefficients and the covari-1957 ance parameter vector $\boldsymbol{\theta}$. It is desired that estimation techniques for these 1958 based on observations be unbiased, i.e., that the average estimation error be 1959zero, and that this error be optimal in a least-squares sense. Another desirable 1960 property is *consistency* in the sense that the estimates converge to the true 1961values as more and more observations are added. 1962

The restriction of Z to a finite set of observation points $\{x_i\}_{i=1}^n \subset D$ is a multivariate Gaussian random vector, which we denote by

$$\begin{bmatrix} Z(\boldsymbol{x}_1, \omega) \end{bmatrix}$$
 1965
1966

$$\boldsymbol{Z} = \boldsymbol{Z}(\omega) = \begin{vmatrix} \vdots \\ Z(\boldsymbol{x}, \omega) \end{vmatrix} : \Omega \to \mathbb{R}^n. \tag{30}$$

$$\begin{array}{c} 1960\\ 1967\\ 1968 \end{aligned}$$

$$\begin{bmatrix} Z(\boldsymbol{x}_n, \omega) \end{bmatrix}$$
 1968
1969

In view of (29), its expectation is

$$\mathbf{E}[\mathbf{Z}] = \mathbf{H}\boldsymbol{\beta}, \qquad [\mathbf{H}]_{i,j} = h_j(\mathbf{x}_i), \quad i = 1, \dots, n, \quad j = 1, \dots, k,$$

and its joint probability density function is given by

$$p(\boldsymbol{\xi}; \boldsymbol{\beta}, \boldsymbol{\theta}) = \frac{1}{\sqrt{(2\pi)^n \det(\boldsymbol{C}_{\boldsymbol{\theta}})}} \exp\left(-\frac{1}{2}(\boldsymbol{\xi} - \boldsymbol{H}\boldsymbol{\beta})^\top \boldsymbol{C}_{\boldsymbol{\theta}}^{-1}(\boldsymbol{\xi} - \boldsymbol{H}\boldsymbol{\beta})\right), \qquad \boldsymbol{\xi} \in \mathbb{R}^{n, 1976}_{, 1977}$$

(31)1978

1935

1938

1939

1943

1944

1963

1964

1979 in which C_{θ} denotes the covariance matrix

1980

1981 1982

$$oldsymbol{C}_{oldsymbol{ heta}} = \mathsf{E}\left[oldsymbol{Z}oldsymbol{Z}^{ op}
ight] = \left[c_{oldsymbol{ heta}}(oldsymbol{x}_i,oldsymbol{x}_j)
ight]_{i,j=1}^n \in \mathbb{R}^{n imes n}$$

1983 of the random vector \boldsymbol{Z} .

1984 When the covariance parameters $\boldsymbol{\theta}$ are known, an unbiased, consistent and 1985 optimal estimate of $\boldsymbol{\beta}$, given a vector of observations $\boldsymbol{\zeta} \in \mathbb{R}^n$, is obtained by 1986 minimizing the weighted least-squares functional

1987

$$\|\boldsymbol{\zeta} - \boldsymbol{H}\boldsymbol{\beta}\|_{C_{\boldsymbol{\theta}}^{-1}}^2 := (\boldsymbol{\zeta} - \boldsymbol{H}\boldsymbol{\beta})^{\top} C_{\boldsymbol{\theta}}^{-1} (\boldsymbol{\zeta} - \boldsymbol{H}\boldsymbol{\beta}),$$

1989

1990 resulting in the estimate

1991

$$\hat{\boldsymbol{\beta}} = (\boldsymbol{H}^{\top} \boldsymbol{C}_{\boldsymbol{\theta}}^{-1} \boldsymbol{H})^{-1} \boldsymbol{H}^{\top} \boldsymbol{C}_{\boldsymbol{\theta}}^{-1} \boldsymbol{\zeta}$$

1993

1994 In maximum likelihood (ML) estimation, the joint probability density function 1995 (31) is maximized for the given observation vector $\boldsymbol{\xi} = \boldsymbol{\zeta}$ as a function of 1996 the parameters $\boldsymbol{\beta}$ and $\boldsymbol{\theta}$. To solve this nonlinear optimization problem one 1997 usually minimizes the negative logarithm $\ell(\boldsymbol{\zeta}; \boldsymbol{\beta}, \boldsymbol{\theta}) := -\log p(\boldsymbol{\zeta}; \boldsymbol{\beta}, \boldsymbol{\theta})$ of the 1998 likelihood given by

1999

$$\ell(\boldsymbol{\zeta}; \boldsymbol{\beta}, \boldsymbol{\theta}) = \frac{1}{2} \left[n \log(2\pi) + \log \det \boldsymbol{C}_{\boldsymbol{\theta}} + (\boldsymbol{\zeta} - \boldsymbol{H}\boldsymbol{\beta})^{\top} \boldsymbol{C}_{\boldsymbol{\theta}}^{-1} (\boldsymbol{\zeta} - \boldsymbol{H}\boldsymbol{\beta}) \right].$$
(32)

2002

2003 As is argued, e.g., in Kitanidis (1987), when random field hydrogeological 2004 parameters are estimated based on data from a finite region where the sep-2005 aration distance of the measurements is of the same order as the correlation 2006 length, the use of fitted means may introduce a bias in the estimation of the 2007 covariance parameters, resulting typically in an underestimation of both the 2008 variance and correlation length parameters. This bias is the result of strong 2009 correlations in the observations, preventing the estimation error from entering 2010 the asymptotic regime as more observations are added, since the number of 2011 independent measurements does not increase due to these strong correlations. A remedy known as restricted maximum likelihood estimation (RML) is to 2012 2013 apply a transformation to the data which filters out the mean. In the case of 2014 the linear model (8) for the mean, we consider the random vector \mathbf{Z}' obtained 2015 by projecting Z orthogonally onto the orthogonal complement of the range 2016 of H, hence removing any effect of the estimated regression coefficients β 2017 on the estimation of the covariance parameters. Indeed, if the columns of 2018 $\boldsymbol{Q} \in \mathbb{R}^{n \times (n-k)}$ form an orthonormal basis of range(\boldsymbol{H}), then $\boldsymbol{Q}^{\top} \boldsymbol{H} = \boldsymbol{O}$ and 2019 therefore the random vector

2020

2021

2022

2023

$$oldsymbol{Z}' := oldsymbol{Q}oldsymbol{Q}^ op oldsymbol{Z}$$

has expectation

$$\mathsf{E}\left[oldsymbol{Z}^{\prime}
ight] = \mathsf{E}\left[oldsymbol{Q}oldsymbol{Q}^{ op}(oldsymbol{H}eta+\widetilde{oldsymbol{Z}})
ight] = \mathsf{E}\left[\widetilde{oldsymbol{Z}}
ight] = \mathbf{0}$$

regardless of the value of β . Here \widetilde{Z} denotes the random vector obtained by restricting the residual random field \widetilde{Z} to the observation points. RML now maximizes the likelihood of the transformed random vector Z', which has an (n-k)-dimensional multivariate normal distribution with zero mean and covariance matrix $Q^{\top} C_{\theta} Q \in \mathbb{R}^{(n-k) \times (n-k)}$.