RANDOMIZED LOCAL MODEL ORDER REDUCTION

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Abstract. In this paper we propose local approximation spaces for localized model order reduction procedures such as domain decomposition and multiscale methods. Those spaces are constructed from local solutions of the partial differential equation (PDE) with random boundary conditions, yield an approximation that converges provably at a nearly optimal rate, and can be generated at close to optimal computational complexity. In many localized model order reduction approaches like the generalized finite element method, static condensation procedures, and the multiscale finite element method local approximation spaces can be constructed by approximating the range of a suitably defined transfer operator that acts on the space of local solutions of the PDE. Optimal local approximation spaces that yield in general an exponentially convergent approximation are given by the left singular vectors of this transfer operator [I. Babuška and R. Lipton 2011, K. Smetana and A. T. Patera 2016]. However, the direct calculation of these singular vectors is computationally very expensive. In this paper, we propose an adaptive randomized algorithm based on methods from randomized linear algebra [N. Halko et al. 2011], which constructs a local reduced space approximating the range of the transfer operator and thus the optimal local approximation spaces. Moreover, the adaptive algorithm relies on a probabilistic a posteriori error estimator for which we prove that it is both efficient and reliable with high probability. Several numerical experiments confirm the theoretical findings.

Key words. localized model order reduction, randomized linear algebra, domain decomposition methods, multiscale methods, a priori error bound, a posteriori error estimation

AMS subject classifications. 65N15, 65N12, 65N55, 65N30, 65C20, 65N25

1. Introduction. Over the last decades (numerical) simulations based on partial differential equations (PDEs) have considerably gained importance in many (complex) applications. Model reduction is an indispensable tool for the simulation of complex problems where the use of standard methods such as finite elements (FE) and finite volumes is prohibitive. Examples for the latter are tasks where multiple simulation requests or real-time simulation response are desired, the (numerical) treatment of partial differential equations with rapidly varying and strongly heterogeneous coefficients, or simulations on very large or geometrically varying domains. Approaches developed to tackle such (complex) problems are localized model order reduction (localized MOR) approaches that are based on (combinations of) domain decomposition (DD) methods, multiscale methods, and the reduced basis method. This paper proposes local approximation spaces for interfaces or subdomains for local model order reduction procedures for linear, elliptic PDEs that yield a nearly optimally convergent approximation, are computationally inexpensive, and easy to implement.

Recently, local approximation spaces that are optimal in the sense of Kolmogorov [52] and thus minimize the approximation error among all spaces of the same dimension, have been introduced for subdomains $\Omega_{in}$ in [9] and for interfaces $\Gamma_{in}$ in [80]. To that end, an oversampling subdomain $\Omega$ which contains the target subdomain $\Omega_{in}$ or interface $\Gamma_{in}$ and whose boundary $\partial\Omega$ has a certain distance to the former is considered. Motivated by the fact that the global solution of the PDE satisfies

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the PDE locally, the space of harmonic functions — that means all local solutions of the PDE with arbitrary Dirichlet boundary conditions — is considered on the oversampling subdomain \( \Omega \). Note that in general we expect an exponential decay of the higher frequencies of the Dirichlet boundary conditions to \( \Omega_{in} \) or \( \Gamma_{in} \). Therefore, we anticipate that already a local ansatz space of very small size should result in a very accurate approximation of all harmonic functions on \( \Omega \). To detect the modes that still persist on \( \Omega_{in} \) or \( \Gamma_{in} \) a (compact) transfer operator is introduced that maps harmonic functions restricted to \( \partial \Omega \) to harmonic functions restricted to \( \Omega_{in} \) or \( \Gamma_{in} \), respectively.

The eigenfunctions of the “transfer eigenproblem” — the eigenvalue problem for the composition of the transfer operator and its adjoint — span the optimal space which yields in general a superalgebraically and thus nearly exponentially convergent approximation. Recently, in \([82, 81]\) the results in \([9, 80]\) have been generalized from linear differential operators whose associated bilinear form is coercive to elliptic, inf-sup stable ones.

However, computing say an FE approximation of these (continuous) optimal spaces by approximating the “transfer eigenproblem” requires first to solve the PDE on \( \Omega \) for each FE basis function as Dirichlet boundary conditions on \( \partial \Omega \) and subsequently to solve a dense eigenproblem of the size of the number of degrees of freedom (DOFs) on \( \partial \Omega \). This is prohibitively expensive for many applications, especially for problems in three space dimensions. Applying the implicitly restarted Lanczos method as implemented in ARPACK \([54]\) requires \( O(n) \) local solutions of the PDE in each iteration, where \( n \) denotes the desired size of the local approximation space.

In this paper we propose to build local approximation spaces adaptively from local solutions of the PDE with (Gaussian) random boundary conditions. To give an intuition why randomly generated local approximation spaces may perform very well, we note that if we draw say \( n \) independent random vectors which form the coefficients of FE basis functions on \( \partial \Omega \) and apply the transfer operator, due to the extremely rapid decay of higher frequencies from \( \partial \Omega \), the modes that still persist on \( \Omega_{in} \) or \( \Gamma_{in} \) will be very close to the optimal modes. In detail, based on methods from randomized linear algebra (randomized LA) \([38, 67]\) we propose an adaptive algorithm which iteratively enhances the reduced space by (local) solutions of the PDE for random boundary conditions and terminates when a probabilistic a posteriori error estimator lies below a given tolerance. We prove that after termination of the adaptive algorithm also the local approximation error is smaller or equal than the given tolerance with very high (given) probability. The respective probabilistic a posteriori estimator in this paper is an extension of a result in \([38]\) and we show in addition, as one contribution of this paper, that the effectivity of the a posteriori error estimator can be bounded by a constant with high probability. By using the matrix representation of the transfer operator we exploit results from randomized LA \([38]\) to prove that the reduced space produced by the adaptive algorithm yields an approximation that converges at a nearly optimal rate.\(^1\) Thanks to this excellent approximation capacity the adaptive algorithm proposed in this paper thus only requires very few local solutions of the PDE in addition to the minimal amount required and is therefore computationally very efficient. As one other (minor) contribution of this paper we extend the results for matrices in \([38]\) to finite dimensional linear operators. We consider in this article parameter-independent PDEs. However, the extension to parameterized PDEs can be realized straightforward (see \([80, 82]\)). Moreover, we assume here that the right-hand side of the PDE is given. If one wishes to construct local spaces for arbitrary

\(^1\)For a different analysis of the algorithm in \([38, 67]\) we refer to \([86]\).
right-hand sides prescribing random right-hand sides in the construction of local basis functions, as it is suggested in the context of numerical homogenization in [71, 72], seems to be an attractive option.

Algorithms from randomized LA have got a steadily growing deal of attention in recent years, especially for very large matrices for instance from problems in large-scale data analysis. Two of the most important benefits of randomization are that they can first result in faster algorithms, either in worst-case asymptotic theory and/or numerical implementation, and that they allow very often for (novel) tight error bounds [60]. Finally, algorithms in randomized LA can often be designed to exploit modern computational architectures better than classical numerical methods [60]. For open source software in randomized LA we refer for instance to [84, 55, 30]. A very popular algorithm in randomized LA is the randomized singular value decomposition (SVD) (see for instance [78, 67, 76]), which yields a very accurate approximation of the (deterministic) SVD, getting however along with only $O(n)$ applications of the matrix to random vectors. The randomized SVD can for instance rely on the matrix version of the adaptive algorithm we present in this paper (see [38, 67]). Moreover, the latter shares a close relationship with methods in randomized LA that are based on the concept of dimension reduction, relying on a random linear map that performs an embedding into a low-dimensional space (see e.g. [34, 73, 78, 67, 76]). Other randomized algorithms employ element-wise sampling of the matrix — for details we refer to the review in [25] and the references therein — or sampling of the columns or rows of the matrix [33, 34, 22, 77, 13, 26, 24, 25]. In both cases sampling is based on a certain probability distribution. In case of column sampling a connection to the low-rank approximations we are interested in in this paper can be set up via leverage scores [60, 24], where this probability distribution is based on (an approximation of) the space spanned by the best rank-$n$ approximation. In general this subcollection of columns or rows can then for instance be used to construct an interpolative decomposition or a CUR decomposition [67, 61, 23, 19, 26]. The matrix version of the adaptive algorithm we present in this paper can also be interpreted in the context of linear sketching: Applying the (input) matrix to a random matrix with certain properties results in a so-called “sketch” of the input matrix, which is either a smaller or sparser matrix but still represents the essential information of the original matrix (see for instance [25, 87] and references therein): for instance, one can show that under certain conditions on the random matrix, the latter is an approximate isometry [83, 87]. The computations can then be performed on the sketch (see e.g. [87, 78]). Using structured random matrices such as a subsampled random Fourier or Hadamard transform or the fast Johnson-Lindenstrauss transform [78, 88, 57, 24, 2, 20] is particularly attractive from a computational viewpoint and yields an improved asymptotic complexity compared to standard methods. Finally, randomization can also be beneficial to obtain high-performant rank-revealing algorithms [65, 27].

Using techniques from randomized LA has already been advocated in (localized) model order reduction approaches in other publications. In [85] Vouvakis et. al. demonstrated the potential of algorithms from randomized LA for domain decomposition methods by using adaptive, randomized techniques to approximate the range of discrete localized Dirichlet-to-Neumann maps in the context of a FETI-2A preconditioner. Regarding multi-scale methods the use of local ansatz spaces spanned by local solutions of the PDE with random boundary conditions is suggested in [17] for the generalized multiscale finite element method (GMSFEM). Here, the reduced space is selected via an eigenvalue problem restricted to a space consisting of local solutions of the PDE with random boundary conditions. Based on results in [66] an a priori
error bound is shown, however, in contrast to our approach, it depends in general on the square root of the number of DOFs on the outer boundary \( \partial \Omega \). Moreover, in contrast to [17] we can formulate our procedure as an approximation of the optimal local approximation spaces suggested in [9, 80] and are thus able to provide a relation to the optimal rate. Eventually, the method proposed in [17] either requires the dimension of the reduced space to be known in advance or the use of \( O(n) \) local solutions of the PDE in addition to the minimal amount required. Finally, we note that in [29] the local reduced space is constructed from local solutions of the PDE with a linear combination of discrete generalized Legendre polynomials with random coefficients as Dirichlet boundary conditions and in [16] \( \Omega \) functions on \( \partial \Omega \) with random coefficients are considered as boundary conditions. However, neither of the two articles takes advantage of the numerical analysis available for randomized LA techniques.

The potential of applying algorithms from randomized LA in model order reduction has also already been demonstrated: In [89] a method for the construction of preconditioners of parameter-dependent matrices is proposed, which is an interpolation of the matrix inverse and is based on a projection of the identity matrix with respect to the Frobenius norm. Methods from randomized LA are used to compute a statistical estimator for the Frobenius norm. In [42] a randomized SVD is employed to construct a reduced model for electromagnetic scattering problems. Finally, in [4] the authors suggest to employ a randomized SVD to construct a reduced basis for the approximation of systems of ordinary partial differential equations.

There are many other choices of local approximation spaces in localized MOR approaches. In DD methods reduced spaces on the interface or in the subdomains are for example chosen as the solutions of (local constrained) eigenvalue problems in component mode synthesis (CMS) [46, 11, 12, 41] or (generalized) harmonic polynomials, plane waves, or local solutions of the PDE accounting for instance for highly heterogeneous coefficients in the Generalized Finite Element Method (GFEM) [7, 6, 10, 9]. In the Discontinuous Enrichment Method (DEM) [31, 32] local FE spaces are enriched by adding analytical or numerical free-space solutions of the homogeneous constant-coefficient counterpart of the considered PDE, while interelement continuity is weakly enforced via Lagrange multipliers. In multiscale methods such as the multiscale FEM (MsFEM), the variational multiscale method (VMM), or the Local Orthogonal Decomposition Method (LOD) the effect of the fine scale on the coarse scale is either modeled analytically [45] or computed numerically by solving the fine-scale equations on local patches with homogeneous Dirichlet boundary conditions [44, 53, 63].

The reduced basis (RB) method has been introduced to tackle parameterized PDEs and prepares in a possibly expensive offline stage a low-dimensional reduced space which is specifically tailored to the considered problem in order to realize subsequently fast simulation responses for possibly many different parameters (for an overview see [75, 39, 37]). Combinations of the RB method with DD methods have been considered in [58, 59, 48, 5, 47, 29, 79, 49, 62, 64, 16]. Here, intra-element RB approximations are for instance coupled by either polynomial Lagrange multipliers [58, 59], generalized Legendre polynomials [47], FE basis functions [49], or empirical modes generated from local solutions of the PDE [29, 64, 16] on the interface. In order to address parameterized multiscale problems the local approximation spaces are for instance spanned by eigenfunctions of an eigenvalue problem on the space of harmonic functions in [28], generated by solving the global parameterized PDE and restricting the solution to the respective subdomain in [70, 3], or enriched in the online stage by local solutions of the PDE, prescribing the insufficient RB solution as Dirichlet
The remainder of this paper is organized as follows. In section 2 we present the problem setting and recall the main results for the optimal local approximation spaces introduced in [9, 80]. The main contributions of this paper are developed in section 3 where we propose an adaptive algorithm that generates local approximation spaces. Moreover, we prove a priori and a posteriori error bounds and show that the latter is efficient. Finally, we present numerical results in section 4 for the Helmholtz equation, stationary heat conduction with high contrast, and linear elasticity to validate the theoretical findings and draw some conclusions in section 5.

2. Optimal local approximation spaces for localized model order reduction procedures. Let $\Omega_{gl} \subset \mathbb{R}^d$, $d = 2, 3$, be a large, bounded domain with Lipschitz boundary and assume that $\partial \Omega_{gl} = \Sigma_D \cup \Sigma_N$, where $\Sigma_D$ denotes the Dirichlet and $\Sigma_N$ the Neumann boundary, respectively. We consider a linear, elliptic PDE on $\Omega_{gl}$ with solution $u_{gl}$, where $u_{gl} = g_D$ on $\Sigma_D$ and satisfies homogeneous Neumann boundary conditions on $\Sigma_N$. Note that we consider here homogeneous Neumann boundary conditions to simplify the notation; non-homogeneous Neumann boundary conditions can be taken into account completely analogous to non-homogeneous Dirichlet boundary conditions. To compute an approximation of $u_{gl}$ we employ a domain decomposition or multiscale method combined with model order reduction techniques, which is why we suppose that $\Omega_{gl}$ is decomposed into either overlapping or non-overlapping subdomains. Then, depending on the employed method, one may either require good reduced spaces for the subdomains, the interfaces, or both. To fix the setting we thus consider the task to find a good reduced space either on a subdomain $\Omega_{in} \subseteq \Omega \subseteq \Omega_{gl}$ with $\text{dist}(\Gamma_{\text{out}}, \partial \Omega_{in}) \geq \rho > 0$, $\Gamma_{\text{out}} := \partial \Omega \setminus \partial \Omega_{gl}$ or an interface $\Gamma_{\text{in}} \subset \partial \Omega$, where $\Omega_{\ast} \subseteq \Omega \subset \Omega_{gl}$ and $\text{dist}(\Gamma_{\text{out}}, \Gamma_{\text{in}}) \geq \rho > 0$. Possible geometric configurations are illustrated in Figure 2.1.

The challenge in constructing a good reduced space is the fact that although we know that $u_{gl}$ solves the PDE locally on $\Omega$ we do in general not know the trace of $u_{gl}$ on $\partial \Omega$ a priori. Therefore, we consider the following problem on $\Omega$: For given $f \in X_0'$ find $u \in X := \{w \in [H^1(\Omega)]^z : w = g_D \text{ on } \partial \Omega \cap \Sigma_D\}$ such that

$$\mathcal{A}u = f \quad \text{in } X_0',$$

for arbitrary Dirichlet boundary conditions on $\Gamma_{\text{out}}$, where $\mathcal{A} : [H^1(\Omega)]^z \to X_0'$, $z = 1, 2, 3$ is a linear, elliptic, and continuous differential operator and $X_0'$ denotes the dual space of $X_0 := \{v \in [H^1(\Omega)]^z : v|_{\Gamma_{\text{out}}} = 0, v|_{\Sigma_D \cap \partial \Omega} = 0\}$, $z = 1, 2, 3$. The latter is in turn equipped with the full $H^1$-norm.

By exploiting that the global solution $u_{gl}$ solves the PDE (2.1) locally, recently, optimal local approximation spaces have been introduced for subdomains in [9] and for interfaces in [80].\(^2\) As we aim at providing a good approximation for a whole set

\[^2\text{The key concepts of the construction of optimal local approximation spaces can be nicely illustrated by means of separation of variables in a simple example as in [80, Remark 3.3], see the supplementary materials section SM2.}\]
of functions, namely all functions that solve the PDE (2.1) locally, the concept of optimality of Kolmogorov [52] is used:

**Definition 2.1 (Optimal subspace in the sense of Kolmogorov).** Let $\mathcal{S}, \mathcal{R}$ be Hilbert spaces, $\mathcal{T} : \mathcal{S} \to \mathcal{R}$ a linear, continuous operator, and $\mathcal{R}^n$ an $n$-dimensional subspace of $\mathcal{R}$. Then the Kolmogorov $n$-width of the image of the mapping $\mathcal{T}$ applied to the unit ball of $\mathcal{S}$ in $\mathcal{R}$ is given by

$$d_n(\mathcal{T}(\mathcal{S}); \mathcal{R}) := \inf_{\mathcal{R}^n \subset \mathcal{R}} \sup_{\dim(\mathcal{R}^n)=n} \inf_{\psi \in \mathcal{S}, \zeta \in \mathcal{R}^n} \frac{\| \mathcal{T}\psi - \zeta \|_{\mathcal{R}}}{\| \psi \|_{\mathcal{S}}} = \inf_{\mathcal{R}^n \subset \mathcal{R}} \sup_{\psi \in \mathcal{S}, \zeta \in \mathcal{R}^n} \inf_{\| \psi \|_{\mathcal{S}} \leq 1} \| \mathcal{T}\psi - \zeta \|_{\mathcal{R}}.$$  

A subspace $\mathcal{R}^n \subset \mathcal{R}$ of dimension at most $n$ for which holds

$$d_n(\mathcal{T}(\mathcal{S}); \mathcal{R}) = \sup_{\psi \in \mathcal{S}} \inf_{\zeta \in \mathcal{R}^n} \frac{\| \mathcal{T}\psi - \zeta \|_{\mathcal{R}}}{\| \psi \|_{\mathcal{S}}}$$

is called an optimal subspace for $d_n(\mathcal{T}(\mathcal{S}); \mathcal{R})$.

Being interested in all local solutions of the PDE motivates considering the space of $\mathcal{A}$-harmonic functions on $\Omega$

$$\mathcal{H} := \{ v \in [H^1(\Omega)]^2 : \mathcal{A}v = 0 \text{ in } \lambda^0_0, \ w = 0 \text{ on } \Sigma_D \cap \partial \Omega \}, \ z = 1, 2, 3.$$  

Note that first we restrict ourselves here to the case $f = 0, g_D = 0, \partial \Omega_{in} \cap \partial \Omega = \emptyset$; the general case will be dealt with at the end of this subsection.

As in [9, 80] we may then introduce a transfer operator $\mathcal{T} : \mathcal{S} \to \mathcal{R}$ for Hilbert spaces $\mathcal{S}$ and $\mathcal{R}$, where $\mathcal{S} = \{ w|_{\Gamma_{\text{out}}} : w \in \mathcal{H} \}$. In order to define appropriate range spaces $\mathcal{R}$ that ensure compactness of $\mathcal{T}$ and allow equipping $\mathcal{R}$ with an energy inner product, we first introduce for a domain $D \subset \Omega$ an orthogonal projection $P_{\ker(\mathcal{A}), D} : [H^1(D)]^2 \to \ker(\mathcal{A})$ defined as $P_{\ker(\mathcal{A}), D} v := \sum_{k=1}^{\dim(\ker(\mathcal{A}))} (v, \eta_k)_{\text{quot}} \eta_k$. Here, $\eta_k$ is an orthonormal basis of $\ker(\mathcal{A})$ with respect to the $(\cdot, \cdot)_{\text{quot}}$ inner product, where the definition of the latter has to be inferred from the quotient space $\mathcal{H}|_{\Omega}/\ker(\mathcal{A})$. To illustrate those definitions note that for instance for the Laplacian $\ker(\mathcal{A})$ would be the constant functions and $(\cdot, \cdot)_{\text{quot}}$ would be the $L^2$-inner product on $D$. In the case of linear elasticity $\ker(\mathcal{A})$ would equal the six-dimensional space of the rigid body motions and $(\cdot, \cdot)_{\text{quot}}$ has to be chosen as the full $H^1$-inner product on $D$. We may then define the quotient space $\mathcal{H} := \{ v - P_{\ker(\mathcal{A}), \Omega}(v), \ v \in \mathcal{H} \}$ and specify the transfer operator. For $w \in \mathcal{H}$ we define $\mathcal{T}$ for interfaces or subdomains, respectively, as

$$\mathcal{T}(w|_{\Gamma_{\text{out}}}) = (w - P_{\ker(\mathcal{A}), \Omega}(w))|_{\Gamma_{\text{in}}} \quad \text{or} \quad \mathcal{T}(w|_{\Gamma_{\text{in}}}) = (w - P_{\ker(\mathcal{A}), \Omega_{\text{in}}}(w))|_{\Omega_{\text{in}}}$$

and set $\mathcal{R} = \{ w|_{\Gamma_{\text{in}}} : w \in \mathcal{H} \}$ or $\mathcal{R} = \{ (w - P_{\ker(\mathcal{A}), \Omega_{\text{in}}})|_{\Omega_{\text{in}}} : w \in \mathcal{H} \}$.

Some remarks are in order. In contrast to the definitions in [9, 80] we do not use a quotient space in the definition of the source space $\mathcal{S}$ as this would either significantly complicate the analysis of the randomized local spaces in section 3 or require the construction of a suitable basis in $\mathcal{S}$ or its discrete counterpart, which can become computationally expensive. Thanks to the Caccioppoli inequality (see supplementary materials section SM1), which allows us to bound the energy norm of $\mathcal{A}$-harmonic functions on $\Omega_{\text{in}}$ or $\Omega$, respectively, by their $L^2$-norm on $\Omega$, it can then be proved that the operator $\mathcal{T}$ is compact (see [9, 8, 80] for details), where $\Omega_{\text{in}}$
has been defined in the second paragraph of this section.\footnote{Note in this context that compactness of $\mathcal{T}$ as defined in (2.4) can be easily inferred from the compactness of the transfer operator acting on the quotient space $\mathcal{H}/\ker(A)$ as considered in [9, 8, 80] by employing that the mapping $\mathcal{K} : \mathcal{H}|_{\Gamma_{\text{out}}} \to (\mathcal{H}/\ker(A))|_{\Gamma_{\text{out}}}$ defined as $\mathcal{K}(v)|_{\Gamma_{\text{out}}} := (v - P_{\ker(A),\Omega})(v)|_{\Gamma_{\text{out}}}$ is continuous.} Let finally $\mathcal{T}^* : \mathcal{R} \to \mathcal{S}$ denote the adjoint operator of $\mathcal{T}$. Then, the operator $\mathcal{T}^*\mathcal{T}$ is a compact, self-adjoint, non-negative operator that maps $\mathcal{S}$ into itself, and the Hilbert-Schmidt theorem and Theorem 2.2 in Chapter 4 of [74] yield the following result:

**Theorem 2.2 (Optimal local approximation spaces [9, 80]).** The optimal approximation space for $d_n(\mathcal{T}(\mathcal{S}); \mathcal{R})$ is given by

\[
\mathcal{R}^n := \text{span}\{\phi^n_1, ..., \phi^n_n\}, \quad \text{where} \quad \phi^n_j = \mathcal{T}\varphi_j, \quad j = 1, ..., n,
\]

and $\lambda_j$ are the largest $n$ eigenvalues and $\varphi_j$ the corresponding eigenfunctions that satisfy the transfer eigenvalue problem: Find $(\varphi_j, \lambda_j) \in (\mathcal{S}, \mathbb{R}^+)$ such that

\[
(\mathcal{T}\varphi_j, w) = \lambda_j (\varphi_j, w)_{\mathcal{S}} \quad \forall w \in \mathcal{S}.
\]

Moreover, the following holds:

\[
d_n(\mathcal{T}(\mathcal{S}); \mathcal{R}) = \sup_{\xi \in \mathcal{S}} \inf_{\zeta \in \mathcal{R}^n} \frac{ ||\mathcal{T}\xi - \zeta||_\mathcal{R} }{ ||\xi||_\mathcal{S} } = \sqrt{\lambda_{n+1}}
\]

If we have $\partial\Omega_{\text{in}} \cap \Sigma_D \neq \emptyset$ we do not subtract the orthogonal projection on $\ker(A)$ either in the definition of the transfer operator in (2.4) or the definition of the range space for subdomains. Next, for $f \neq 0$ but still $g_D = 0$ we solve the problem: Find $u^f \in \mathcal{X}_0$ such that $Au^f = f$ in $\mathcal{X}_0^\text{in}$, and augment the space $\mathcal{R}^n$ either with $u^f|_{\Omega_{\text{in}}}$ or $u^f|_{\Gamma_{\text{in}}}$. To take non-homogeneous Dirichlet boundary conditions into account we consider the problem: Find $w^{g_D} \in \{w \in [H^1(\Omega)]^2 : w = g_D$ on $\partial\Omega \cap \Sigma_D$, $w = 0$ on $\Gamma_{\text{out}}\}$, $z = 1, 2, 3$, such that $Au^{g_D} = 0$ in $\mathcal{X}_0^\text{in}$. Finally, we may then define the optimal local approximation space for subdomains as

\[
\mathcal{R}^n_{\text{data,ker}} := \text{span}\{\phi^n_1, ..., \phi^n_n, u^f|_{\Omega_{\text{in}}}, u^{g_D}|_{\Omega_{\text{in}}}, \eta_1|_{\Omega_{\text{in}}}, ..., \eta_{\dim(\ker(A))}|_{\Omega_{\text{in}}}\}
\]

and for interfaces as

\[
\mathcal{R}^n_{\text{data,ker}} := \text{span}\{\phi^n_1, ..., \phi^n_n, u^f|_{\Gamma_{\text{in}}}, u^{g_D}|_{\Gamma_{\text{in}}}, \eta_1|_{\Gamma_{\text{in}}}, ..., \eta_{\dim(\ker(A))}|_{\Gamma_{\text{in}}}\},
\]

respectively, where $\{\eta_1, ..., \eta_{\dim(\ker(A))}\}$ denotes a basis for $\ker(A)$. In case there holds $\partial\Omega_{\text{in}} \cap \Sigma_D \neq \emptyset$ we do not augment the space $\mathcal{R}^n$ with a basis of $\ker(A)$.

**2.1. Approximation of the transfer eigenvalue problem with Finite Elements: matrix form of the transfer operator.** In this subsection we show how an approximation of the continuous optimal local spaces $\mathcal{R}^n_{\text{data,ker}}$ can be computed with the FE method and introduce the notation in this discrete setting required for the remainder of this paper.

To that end, we introduce a partition of $\Omega$ such that $\Gamma_{\text{in}}$ or $\partial\Omega_{\text{in}}$ do not intersect any element of that partition. In addition, we introduce an associated conforming FE space $X \subset [H^1(\Omega)]^2$, $z = 1, 2, 3$ with $\dim(X) = N$, a nodal basis $\{\psi_1, ..., \psi_N\}$ of $X$, the FE source space $\mathcal{S} := \{v|_{\Gamma_{\text{out}}} : v \in X\}$ of dimension $N_N$, and the FE range space $R := \{(v - P_{\ker(A),\Omega}(v))|_{\Gamma_{\text{in}}} : v \in X\}$ or $R := \{(v - P_{\ker(A),\Omega_{\text{in}}})|_{\Omega_{\text{in}}} : v \in X\}$ with $\dim(R) = N_N$. Next, we define the space of discrete $A$-harmonic functions

\[
\mathcal{H} := \{w \in X : Aw = 0 \text{ in } \mathcal{X}_0^\text{in}, w = 0 \text{ on } \Sigma_D \cap \partial\Omega\},
\]
where $A : X \to X'_0$ is the discrete counterpart of $A$ and $X'_0$ denotes the dual space of $X_0 := \{ v \in X : v|_{\Gamma_{\text{out}}} = 0, \, v|_{\partial \Omega_D} = 0 \}$. We may then define the discrete transfer operator $T : S \to R$ for $w \in \mathcal{H}$ as

$$
T(w|_{\Gamma_{\text{out}}}) = (w - P_{\text{ker}(A), \Omega}(w))|_{\Gamma_{\text{in}}} \quad \text{or} \quad T(w|_{\Gamma_{\text{out}}}) = (w - P_{\text{ker}(A), \Omega_{\text{in}}}(w))|_{\Omega_{\text{in}}}.
$$

In order to define a matrix form of the transfer operator we introduce DOF mappings $B_{S \to X} \in \mathbb{R}^{N_S \times N_X}$ and $B_{X \to R} \in \mathbb{R}^{N_X \times N_R}$ that map the DOFs of $S$ to the DOFs of $X$ and the DOFs of $X$ to the DOFs of $R$, respectively. Moreover, we introduce the stiffness matrix $A$ associated with the discrete operator $A$, where we assume that in the rows associated with the Dirichlet DOFs the non-diagonal entries are zero and the diagonal entries equal one. Note that in order to make the distinction between elements of the Hilbert spaces $S$ and $R$ and their coordinate representation in $\mathbb{R}^{N_S}$ and $\mathbb{R}^{N_R}$ explicit, we mark all coordinate vectors and matrices with an underline. By writing functions $\zeta \in S$ as $\zeta = \sum_{i=1}^{N_S} \zeta_i \psi_i|_{\Gamma_{\text{out}}}$ and defining $K_{\Omega_{\text{in}}}$ as the matrix of the orthogonal projection on $\text{ker}(A)$ on $\Omega_{\text{in}}$, we obtain the following matrix representation $T \in \mathbb{R}^{N_R \times N_S}$ of the transfer operator for domains

$$
T_{\zeta} = (1 - K_{\Omega_{\text{in}}}) B_{X \to R} A^{-1} B_{S \to X} \zeta.
$$

For interfaces, the projection on the quotient space is done before the index mapping. There, with $K_{\Omega}$ as the matrix of the orthogonal projection on $\text{ker}(A)$ on $\Omega$, the matrix representation of the transfer operator is given by

$$
T_{\zeta} = B_{X \to R} (1 - K_{\Omega}) A^{-1} B_{S \to X} \zeta.
$$

Finally, we denote by $M_S$ the inner product matrix of $S$ and by $M_R$ the inner product matrix of $R$. Then, the FE approximation of the transfer eigenvalue problem reads as follows: Find the eigenvectors $\zeta_j \in \mathbb{R}^{N_S}$ and the eigenvalues $\lambda_j \in \mathbb{R}^{N_R}$ such that

$$
T^t M_R T_{\zeta_j} = \lambda_j M_S \zeta_j.
$$

The coefficients of the FE approximation of the basis functions $\{\phi_1^{sp}, ..., \phi_n^{sp}\}$ of the optimal local approximation space

$$
R^n := \text{span}\{\phi_1^{sp}, ..., \phi_n^{sp}\}
$$

are then given by $\phi_j^{sp} = T_{\zeta_j}, \, j = 1, \ldots, n$. Adding the representation of the right-hand side, the boundary conditions, and a basis of $\text{ker}(A)$ yields the optimal space $R^n_{\text{data,ker}}$.

Note that we may also perform a singular value decomposition of the operator $T$, which reads

$$
T_{\zeta} = \sum_{j=1}^{\min\{N_S, N_R\}} \sigma_j \tilde{\phi}_j^{sp}(\chi_j, \zeta) s \quad \text{for} \quad \zeta \in S,
$$

Note that in the continuous setting the range space $\mathcal{R}$ is a subspace of the space $\mathcal{R} := \{(v - P_{\text{ker}(A), \Omega}(v))|_{\Gamma_{\text{in}}} : v \in [H^1(\Omega)]^2, \, z = 1, 2, 3 \}$ for interfaces and $\mathcal{R} := \{(v - P_{\text{ker}(A), \Omega_{\text{in}}}(v))|_{\Omega_{\text{in}}} : v \in [H^1(\Omega)]^2, \, z = 1, 2, 3 \}$ for subdomains. It can then be easily shown for the corresponding transfer operator $\tilde{T} : \mathcal{R} \to S$ which is defined identically as in (2.4) that there holds $d_{\text{opt}}(\tilde{T}(S); \mathcal{R}) = d_{\text{opt}}(\tilde{T}(S); \mathcal{R})$ and that the associated optimal approximation spaces are the same. This justifies the usage of the discrete range space as defined above.
with orthonormal bases \( \hat{\phi}^{sp}_{ij} \in R \), \( \chi_{j} \in S \), and singular values \( \sigma_j \in \mathbb{R}_+^j \), and define \( R^n := \text{span}\{\hat{\phi}^{sp}_1, ..., \hat{\phi}^{sp}_n\} \). Up to numerical errors this definition is equivalent to the definition in (2.15) and there holds \( \sigma_j = \sqrt{\lambda_j} \), \( j = 1, \ldots, \min\{N_S, N_R\} \), where \( \lambda_j \) are the eigenvalues of the discrete transfer eigenproblem (2.14). Note however that there holds \( (\hat{\phi}^{sp}_j, \hat{\phi}^{sp}_j)_{R} = \delta_{jj} \lambda_j \) in contrast to \( (\hat{\phi}^{sp}_j, \hat{\phi}^{sp}_j)_{R} = \delta_{ij} \).

Finally, we introduce Ritz isomorphisms \( D_S : S \rightarrow \mathbb{R}^{NS} \) and \( D_R : R \rightarrow \mathbb{R}^{NR} \) which map elements from \( S \) or \( R \) to a vector containing their FE coefficients in \( \mathbb{R}^{NS} \) or \( \mathbb{R}^{NR} \), respectively. For instance, \( D_S \) maps a function \( \xi = \sum_{i=1}^{N_S} \xi_i \psi_i |_{\Gamma_{out}} \in S \) to \( \xi \in \mathbb{R}^{NS} \). As a result we have the matrix of the transfer operator as \( \mathbf{T} = D_R TD_S^{-1} \).

3. Approximating the range of an operator by random sampling. In this section we present and analyze an algorithm which constructs a reduced space \( R^n \) that approximates the range of a finite dimensional linear operator \( T \) of rank \( N_T \) by iteratively enhancing the reduced space with applications of \( T \) to a random function. Although having the transfer operator (2.11) in mind we consider the general setting of a finite dimensional linear operator mapping between two finite dimensional Hilbert spaces \( S \) and \( R \). Note that in the context of localized MOR for inhomogeneous problems it is necessary to enhance \( R^n \) by the representation of the right-hand side and the boundary conditions.

The algorithm and parts of its analysis are an extension of results in randomized LA \[38\] to the setting of finite dimensional linear operators. In detail we first present an adaptive range finder algorithm in subsection 3.1 and discuss its computational complexity. This algorithm relies on a probabilistic a posteriori bound, which is a extension of a result in \[38\] and for which we prove as one new contribution its efficiency in subsection 3.3. Starting from results in randomized LA \[38\] we prove in subsection 3.2 that the reduced space \( R^n \) generated by the algorithm as presented in subsection 3.1 yields an approximation that converges with a nearly optimal rate.

3.1. An adaptive randomized range finder algorithm. We propose an adaptive randomized range approximation algorithm that constructs an approximation space \( R^n \) by iteratively extending its basis until a convergence criterion is satisfied. In each iteration, the basis is extended by the operator \( T \) applied to a random function.

The full algorithm is given in Algorithm 1 and has four input parameters, starting with the operator \( T \), whose range should be approximated. This could be represented by a matrix, but in the intended context it is usually an implicitly defined operator which is computationally expensive to evaluate. Only the evaluation of the operator on a vector is required. The second input parameter is the target accuracy \( \text{tol} \) such that \( \|T - P_{R^n}T\| \leq \text{tol} \). The third input parameter is the number of test vectors \( n_t \) to be used in the a posteriori error estimator which we will discuss shortly. A typical \( n_t \) could be 5, 10, or 20. The fourth input parameter is the maximum failure probability \( \varepsilon_{\text{algofail}} \) and the algorithm returns a space which has the required approximation properties with a probability greater than \( 1 - \varepsilon_{\text{algofail}} \).

The basis \( B \) of \( R^n \) is initialized as empty in line 2, test vectors are initialized as \( \hat{\phi}^{sp}_j \), \( \sigma_j \), and \( \delta_{ij} \) in line 3. Recall that \( TD_S^{-1} R \) is the operator \( T \) applied to a random normal vector. We use the term “random normal vector” to denote a vector whose entries are independent and identically distributed random variables with normal distribution. The main loop of the algorithm is terminated when the following a posteriori norm estimator applied to \( T - P_{R^n}T \) is smaller than \( \text{tol} \).
Algorithm 1: Adaptive Randomized Range Approximation

Function AdaptiveRandomizedRangeApproximation(T, tol, nt, εalgofail):

Input: Operator T,
       target accuracy tol,
       number of test vectors nt,
       maximum failure probability εalgofail

Output: space \( R^n \) with property \( P(\| T - P_{R^n} T \| \leq tol ) > (1 - εalgofail) \)

/* initialize basis */
1. \( B \leftarrow \emptyset \)
/* initialize test vectors */
2. \( M \leftarrow \{ T D_S^{-1} \xi_1, \ldots, T D_S^{-1} \xi_{nt} \} \)
/* determine error estimator factor */
3. \( ε_{testfail} \leftarrow ε_{algofail} / N_T \)
4. \( c_{est} \leftarrow \left( \sqrt{2 λ_{M_S}^{\min}} \frac{1}{\sqrt{2}} \right)^{-1} \)
/* basis generation loop */
5. while \( (\max_{t \in M} \| t \|_{R^n}) \cdot c_{est} > tol \) do
6. \( B \leftarrow B \cup (T D_S^{-1} \xi) \)
7. \( B \leftarrow \text{orthonormalize}(B) \)
8. \( M \leftarrow \{ t - P_{\text{span}(B)} t \mid t \in M \} \)
9. return \( R^n = \text{span}(B) \)

Definition 3.1 (A probabilistic a posteriori norm estimator). To estimate the operator norm of an operator \( O : S \rightarrow R \) of rank \( N_O \), we define the a posteriori norm estimator \( \Delta(O, n_t, ε_{testfail}) \) for \( n_t \) test vectors as

\[
\Delta(O, n_t, ε_{testfail}) := c_{est}(n_t, ε_{testfail}) \max_{i \in 1, \ldots, n_t} \| O D_S^{-1} \xi_i \|_{R^n}
\]

Here, \( c_{est}(n_t, ε_{testfail}) \) is defined as \( c_{est}(n_t, ε_{testfail}) := 1 / \left( \sqrt{2 λ_{M_S}^{\min}} \frac{1}{\sqrt{2}} \right)^{-1} \), \( \xi_i \) are random normal vectors, and \( λ_{M_S}^{\min} \) is the smallest eigenvalue of the matrix of the inner product in \( S \).

This error estimator \( \Delta(O, n_t, ε_{testfail}) \) is analyzed in detail in subsection 3.3. The constant \( c_{est}(n_t, ε_{testfail}) \), which appears in the error estimator, is calculated in line 4 and 5 using \( N_T \) — the rank of operator \( T \). In practice \( N_T \) is unknown and an upper bound for \( N_T \) such as \( \min(N_S, N_R) \) can be used instead. In line 6 the algorithm assesses if the convergence criterion is already satisfied. Note that the term \( (\max_{t \in M} \| t \|_{R^n}) \cdot c_{est}(n_t, ε_{testfail}) \) is the norm estimator (3.1) applied to \( T - P_{R^n} T \). The test vectors are reused for all iterations. The main loop of the algorithm consists of two parts. First, the basis is extended in line 7 and 8 by applying the operator \( T \) to a random normal vector and adding the result to the basis \( B \). Then the basis \( B \) is orthonormalized. The resulting basis vectors are denoted by \( \phi_i^{\text{rnd}} \). We emphasize that the orthonormalization is numerically challenging, as the basis functions are nearly linear dependent when \( R^n \) is already a good approximation of the range of \( T \). In the numerical experiments we use the numerically stable Gram-Schmidt with re-iteration from [15], which always succeeded to obtain an orthogonal set of vectors.
Instead of the Gram-Schmidt orthonormalization, one could apply an SVD to the matrix that contains the vectors in $B$ as columns after termination of Algorithm 1 to remove linear dependent vectors. In the case of almost linear dependent vectors, this could lead to slightly smaller basis sizes. Note that as we suggest to only remove the linear dependent vectors with the SVD the accuracy of the approximation is not compromised. Finally, the test vectors are updated in line 9.

In Algorithm 1, the smallest eigenvalue of matrix of the inner product in $S$, $\lambda_{min}^{M}$, or at least a lower bound for it, is required. The orthonormalization of $B$ in line 8 and the update of test vectors in line 9 use the inner product in $R$. These aspects should be taken into account when choosing the inner products in $S$ and $R$.

The presented algorithm has good performance properties for operators $T$ which are expensive to evaluate. To produce the space $R^n$ of dimension $n$, it evaluates the operator $n$ times to generate the basis and $n_t$ times to generate the test vectors, so in total $n + n_t$ times. In contrast, direct calculation of the optimal space, solving the eigenvalue problem (2.6), would require $N_S$ evaluations of the operator and solving a dense eigenproblem of dimension $N_S \times N_S$. Exploiting the low rank structure of $T$, one could calculate the eigenvectors of $T^*T$ using a Lanczos type algorithm as implemented in ARPACK [54], but this would require $O(n)$ evaluations of $T$ and $T^*$ in every iteration, potentially summing up to much more than $n + n_t$ evaluations, where the number of iterations is often not foreseeable.

### 3.2. A Probabilistic a Priori Error Bound

In this subsection we analyze the convergence behavior of Algorithm 1. In detail, we derive a probabilistic a priori error bound for the projection error $\|T - P_{R^n}T\|$ and its expected value. Recalling that the optimal convergence rate achieved by the optimal spaces from Theorem 2.2 is $\sqrt{\lambda_{n+1}} = \sigma_{n+1}$ we show that the reduced spaces constructed with Algorithm 1 yield an approximation that converges with a nearly optimal rate:

**Proposition 3.2.** Let $\lambda_{max}^{M_S}$, $\lambda_{min}^{M_S}$, $\lambda_{max}^{M_R}$, and $\lambda_{min}^{M_R}$ denote the largest and smallest eigenvalues of the inner product matrices $M_S$ and $M_R$, respectively and let $R^n$ be the outcome of Algorithm 1. Then, for $n \geq 4$ there holds

\[
E\|T - P_{R^n}T\| \leq \sqrt{\lambda_{max}^{M_S}} \min_{k+p=n, k \geq 2, p \geq 2} \left[ \left( 1 + \sqrt{\frac{k}{p-1}} \right) \sigma_{k+1} + \frac{\epsilon \sqrt{n}}{p} \left( \sum_{j \geq k} \sigma_j^2 \right)^{\frac{1}{2}} \right].
\]

Before addressing the proof of Proposition 3.2 we highlight that for operators with a fast decaying spectrum such as the transfer operator the last term in (3.2) behaves roughly as $(\epsilon \sqrt{k + p} \sigma_{k+1})/p$ and we therefore obtain an approximation that converges approximately as $\sqrt{n} \sigma_{n+1}$ and thus with a nearly optimal rate. Proposition 3.2 extends the results in Theorem 10.6 in [38] to the case of finite dimensional linear operators. The terms consisting of the square root of the conditions of the inner product matrices $M_S$ and $M_R$ in (3.2) are due to our generalization from the spectral matrix norm as considered in [38] to inner products associated with finite dimensional Hilbert spaces. We present a reformulation in the supplementary materials Proposition SM4.2 where the condition of $M_S$ does not appear. The occurrence of the remaining terms in (3.2) is discussed in section SM3 where we summarize the proof of Theorem 10.6 in [38], which read as follows:

**Theorem 3.3.** [38, Theorem 10.6] Let $T \in \mathbb{R}^{N_R \times N_S}$ and $P_{R^n,2}$ be the matrix of the orthogonal projection on $R^n$ in the euclidean inner product in $\mathbb{R}^{N_R}$ and $\|\cdot\|_2$
denote the spectral matrix norm. Then for \( n \geq 4 \) it holds

\[
E \left( \| T - P_{R^n,2} T \|_2 \right) \leq \min_{k \geq 2, p \geq 2} \left[ \left( 1 + \frac{k}{p} \right) \sigma_{k+1} + \frac{\sqrt{n}}{p} \left( \sum_{j>k} \sigma_j^2 \right)^{\frac{1}{2}} \right].
\]

To proceed with the proof of Proposition 3.2, we next bound \( \| T - P_{R^n,2} T \| \) by \( \| T - P_{R^n,2} T \|_2 \) times other terms in Lemma 3.4. Then we apply Theorem 3.3 to the matrix representation \( T \) of the operator \( T \) and finally bound the singular values \( \sigma_i \) of the matrix \( T \) by the singular values \( \sigma_i \) of the operator \( T \) in Lemma 3.5 below to conclude.

**Lemma 3.4.** There holds for some given reduced space \( R^n \)

\[
\| T - P_{R^n} T \| = \sup_{\xi \in S} \inf_{\xi \in R^n} \frac{\| T \xi - \xi \|_S}{\| \xi \|_S} \leq \sqrt{\frac{\lambda_{\max}^R}{\lambda_{\min}^R}} \| T - P_{R^n,2} T \|_2.
\]

**Proof.**

\[
\begin{align*}
&\sup_{\xi \in S} \inf_{\xi \in R^n} \frac{\| T \xi - \xi \|_S}{\| \xi \|_S} = \sup_{\xi \in S} \frac{\| T \xi - P_{R^n} T \xi \|_S}{\| \xi \|_S} \\
&= \sup_{\xi \in \mathbb{R}^N_S} \frac{\| (T \xi - P_{R^n,2} T \xi)^T M_{R^n} (T \xi - P_{R^n,2} T \xi) \|^{\frac{1}{2}}}{\| \xi \|_S} \\
&\leq \sup_{\xi \in \mathbb{R}^N_S} \frac{\| (T \xi - P_{R^n,2} T \xi)^T M_{S} (T \xi - P_{R^n,2} T \xi) \|^{\frac{1}{2}}}{\| \xi \|_S} \\
&\leq \sqrt{\frac{\lambda_{\max}^R}{\lambda_{\min}^R}} \sup_{\xi \in \mathbb{R}^N_S} \frac{\| T \xi - P_{R^n,2} T \xi \|_2}{\| \xi \|_2}.
\end{align*}
\]

}\]

**Lemma 3.5.** Let the singular values \( \sigma_j \) of the matrix \( T \) be sorted in non-increasing order, i.e. \( \sigma_1 \geq \ldots \geq \sigma_N \), and \( \sigma_j \) be the singular values of the operator \( T \), also sorted non-increasing. Then there holds \( \sigma_j \leq (\lambda_{\max}^S/\lambda_{\min}^S)^{1/2} \sigma_j \) for all \( j = 1, \ldots, N_T \).

**Proof.** For notational convenience we denote within this proof the \( j \)-th eigenvalue of a matrix \( A \) by \( \lambda_j(A) \). All singular values for \( j = 1, \ldots, N_T \) are different from zero. Therefore, there holds \( \sigma_j^2 = \lambda_j(T^T T) \) and \( \sigma_j^2 = \lambda_j(M_S^{-1} T M_R T) \). Recall that \( T \) is the matrix representation of \( T \) and note that \( M_S^{-1} T^T M_R \) is the matrix representation of the adjoint operator \( T^* \). The non-zero eigenvalues of a product of matrices \( AB \) are identical to the non-zero eigenvalues of the product \( BA \) (see e.g. [43, Theorem 1.3.22]), hence \( \lambda_j(M_S^{-1} T^T M_R T) = \lambda_j(T^T M_R T M_S^{-1}) \). We may then apply the Courant minimax principle to infer \( \lambda_j(T^T M_R T)(\lambda_{\max}^S)^{-1} \leq \lambda_j(M_S^{-1} T^T M_R T) \). Employing once again cyclic permutation and the Courant minimax principle yields

\[
\lambda_j(T^T T) \leq \lambda_j(T^T M_R T) \frac{1}{\lambda_{\min}^R} \leq \lambda_j(M_S^{-1} T^T M_R T) \frac{\lambda_{\max}^S}{\lambda_{\min}^S}
\]

and thus the claim. \( \square \)
Remark 3.6. The result of Algorithm 1, when interpreted as functions and not as coefficient vectors, is independent of the choice of the basis in \( \mathbb{R} \). Disregarding numerical errors, the result would be the same if the algorithm was executed in an orthonormal basis in \( \mathbb{R} \). Thus, we would expect Proposition 3.2 to hold also without the factor \( (\lambda_{\text{max}}/\lambda_{\text{min}})^{1/2} \).

3.3. Adaptive convergence criterion and a probabilistic a posteriori error bound. When approximating the range of an operator, usually its singular values are unknown. To construct a space with prescribed approximation quality, Algorithm 1 uses the probabilistic a posteriori error estimator defined in Definition 3.1, which is analyzed in this subsection.

Proposition 3.7 (Norm estimator failure probability). The norm estimator \( \Delta(O, n_t, \varepsilon_{\text{testfail}}) \) is an upper bound of the operator norm \( \|O\| \) with probability greater or equal than \( 1 - \varepsilon_{\text{testfail}} \).

Proposition 3.8 (Norm estimator effectivity). Let the effectivity \( \eta \) of the norm estimator \( \Delta(O, n_t, \varepsilon_{\text{testfail}}) \) be defined as

\[
\eta(O, n_t, \varepsilon_{\text{testfail}}) := \frac{\Delta(O, n_t, \varepsilon_{\text{testfail}})}{\|O\|}.
\]

Then, there holds

\[
P\left( \eta \leq c_{\text{eff}}(n_t, \varepsilon_{\text{testfail}}) \right) \geq 1 - \varepsilon_{\text{testfail}},
\]

where the constant \( c_{\text{eff}}(n_t, \varepsilon_{\text{testfail}}) \) is defined as

\[
c_{\text{eff}}(n_t, \varepsilon_{\text{testfail}}) := Q^{-1}\left( \frac{N_O}{2}, \frac{\varepsilon_{\text{testfail}}}{n_t} \right) \lambda_{\text{max}}^2 \lambda_{\text{min}}^{-2} (\text{erf}^{-1}(\frac{\varepsilon_{\text{testfail}}}{\sqrt{\varepsilon_{\text{testfail}}}}))^2 \]

and \( Q^{-1} \) is the inverse of the upper normalized incomplete gamma function; that means \( Q^{-1}(a, y) = x \) when \( Q(a, x) = y \).\(^5\)

The proofs of Propositions 3.7 and 3.8 follow at the end of this subsection.

In Proposition 3.7 we analyzed the probability for one estimate to fail. Based on that, we can analyze the algorithm failure probability. To quantify this probability, we first note that Algorithm 1 will terminate after at most \( N_T \) steps. Then, the approximation space \( \mathbb{R}^n \) has the same dimension as \( \text{range}(T) \) and as \( \mathbb{R}^n \subset \text{range}(T) \) we have \( \mathbb{R}^n = \text{range}(T) \) and thus \( \|T - P_{\mathbb{R}^n}T\| = 0 \). The a posteriori error estimator defined in Definition 3.1 is therefore executed at most \( N_T \) times. Each time, the probability for failure is given by Proposition 3.7 and with a union bound argument we may then infer that the failure probability for the whole algorithm is \( \varepsilon_{\text{algofail}} \leq N_T \varepsilon_{\text{testfail}} \).

To prove Propositions 3.7 and 3.8, it is central to analyze the distribution of the inner product \( (v, D_S^{-1}z)_S \) for any \( v \in S \) with \( \|v\|_S = 1 \) and a random normal vector \( z \).

\(^5\) Recall that the definition of the upper normalized incomplete gamma function is

\[
Q(a, x) = \int_0^x t^{a-1}e^{-t}dt = \frac{\Gamma(a, x)}{\Gamma(a)}.
\]
LEMMA 3.9 (Distribution of inner product). The inner product of a normed vector \( v \) in \( S \) with a random normal vector \( (v, D_S^{-1}r)_S \) is a Gaussian distributed random variable with mean zero and variance \( s^2 \), where \( \lambda_{\min}^S \leq s^2 \leq \lambda_{\max}^S \).

Proof. We use the spectral decomposition of the inner product matrix

\[
M_S = \sum_{i=1}^{N_S} m_{S,i} \lambda_i^S m_{S,i}^T
\]

with eigenvalues \( \lambda_i^S \) and eigenvectors \( m_{S,i} \). There holds

\[
(v, D_S^{-1}r)_S = \sum_{i=1}^{N_S} (D_S v)^T \lambda_i^S m_{S,i}^T m_{S,i} r.
\]

As \( m_{S,i} \) is normed with respect to the euclidean inner product, the term \( m_{S,i}^T r \) is a normal distributed random variable. Using the rules for addition and scalar multiplication of Gaussian random variables, one sees that the inner product \( (v, D_S^{-1}r)_S \) is a Gaussian random variable with variance \( s^2 = \sum_{i=1}^{N_S} ((D_S v)^T m_{S,i} \lambda_i^S)^2 \) The variance \( s^2 \) can easily be bounded as follows:

\[
s^2 = \sum_{i=1}^{N_S} ((D_S v)^T m_{S,i} \lambda_i^S)^2 \leq \sum_{i=1}^{N_S} ((D_S v)^T m_{S,i})^2 \lambda_i^S \max_i (\lambda_i^S) = \lambda_{\max}^S
\]

\[
s^2 = \sum_{i=1}^{N_S} ((D_S v)^T m_{S,i} \lambda_i^S)^2 \geq \sum_{i=1}^{N_S} ((D_S v)^T m_{S,i})^2 \lambda_i^S \min_i (\lambda_i^S) = \lambda_{\min}^S
\]

\[
\square
\]

Using this result, we can prove Propositions 3.7 and 3.8. Proof of Proposition 3.7:

Proof. We analyze the probability for the event that the norm estimator \( \Delta(O, n_t, \varepsilon_{\text{testfail}}) \) is smaller than the operator norm \( \lVert O \rVert \):

\[
P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}} < \lVert O \rVert)\right) = P\left(\epsilon_{\text{est}}(n_t, \varepsilon_{\text{testfail}}) \max_{i \in 1, \ldots, n_t} \lVert O D_S^{-1} \lambda_i \rVert_R < \lVert O \rVert\right).
\]

The probability that all test vector norms are smaller than a certain value is the product of the probabilities that each test vector is smaller than that value. So with a new random normal vector \( r \) it holds

\[
P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}} < \lVert O \rVert)\right) = P\left(\epsilon_{\text{est}}(n_t, \varepsilon_{\text{testfail}}) \lVert O D_S^{-1} r \rVert_R < \lVert O \rVert\right)^{n_t}.
\]

Using the singular value decomposition of the operator \( O \):

\[
O \varphi = \sum_i u_i \sigma_i (v_i, \varphi)_S
\]

we obtain

\[
P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}} < \lVert O \rVert)\right) \leq P\left(\epsilon_{\text{est}}(n_t, \varepsilon_{\text{testfail}}) \lVert u_1 \sigma_1 (v_1, D_S^{-1} r)_S \rVert_R < \lVert O \rVert\right)^{n_t}
\]

\[
= P\left(\epsilon_{\text{est}}(n_t, \varepsilon_{\text{testfail}}) \sigma_1 \lVert (v_1, D_S^{-1} r)_S \rVert < \lVert O \rVert\right)^{n_t}
\]

\[
= P\left(\epsilon_{\text{est}}(n_t, \varepsilon_{\text{testfail}}) \lVert (v_1, D_S^{-1} r)_S < 1\right)^{n_t}.
\]

The inner product \( \lVert (v_1, D_S^{-1} r)_S \rVert \) is a Gaussian distributed random variable with variance greater \( \lambda_{\min}^S \), so with a new normal distributed random variable \( r' \) it holds

\[
P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}} < \lVert O \rVert)\right) \leq P\left(\sqrt{\lambda_{\min}^S} \lVert r' \rVert < \sqrt{2} \lambda_{\min}^S \cdot \text{erf}^{-1}\left(\sqrt{\epsilon_{\text{testfail}}}\right)\right)^{n_t}
\]

\[
= \text{erf}\left(\sqrt{2} \text{erf}^{-1}\left(\sqrt{\epsilon_{\text{testfail}}}\right)\right)^{n_t} = \epsilon_{\text{testfail}}.
\]
Proof of Proposition 3.8:

Proof. The constant $c_{\text{est}}(n_t, \varepsilon_{\text{testfail}})$ is defined as in the proof of Proposition 3.7. To shorten notation, we write $c_{\text{est}}$ for $c_{\text{est}}(n_t, \varepsilon_{\text{testfail}})$ and $c_{\text{eff}}$ for $c_{\text{eff}}(n_t, \varepsilon_{\text{testfail}})$ within this proof. Invoking the definition of $\Delta(O, n_t, \varepsilon_{\text{testfail}})$ yields

$$P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}}) > c_{\text{eff}} \|O\|\right) = P\left(c_{\text{est}} \max_{i \in 1, \ldots, n_t} \|O D_{S_i}^{-1} r\|_R > c_{\text{eff}} \|O\|\right)$$

and by employing a new random normal vector $r$ we obtain

$$P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}}) > c_{\text{eff}} \|O\|\right) \leq n_t P\left(c_{\text{est}} \max_{i \in 1, \ldots, n_t} \|O D_{S_i}^{-1} r\|_R > c_{\text{eff}} \|O\|\right).$$

Using the singular value decomposition of the operator $O$: $O\varphi = \sum_i u_i \sigma_i(v_i, \varphi)_S$ results in

$$P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}}) > c_{\text{eff}} \|O\|\right) \leq n_t P\left(c_{\text{est}} \max_{i \in 1, \ldots, n_t} \|u_i \sigma_1(v_i, D_{S_i}^{-1} r)\|_R > c_{\text{eff}} \|O\|\right).$$

For a new random normal variables $r_i$ we have

$$P\left(\Delta(O, n_t, \varepsilon_{\text{testfail}}) > c_{\text{eff}} \|O\|\right) \leq n_t P\left(c_{\text{est}} \sigma_1 \sqrt{\lambda_{\text{max}}} \sqrt{\sum_i r_i^2} > c_{\text{eff}} \|O\|\right)$$

$$= n_t P\left(\sqrt{\sum_i r_i^2} > \frac{c_{\text{eff}}}{c_{\text{est}}} \sigma_1 \sqrt{\frac{M_s}{\lambda_{\text{max}}}} \|O\|\right) = n_t P\left(\sqrt{\sum_i r_i^2} > \frac{c_{\text{eff}}}{c_{\text{est}}} \sqrt{\frac{M_s}{\lambda_{\text{max}}}} \|O\|\right)$$

$$= n_t P\left(\sum_i r_i^2 > \frac{c_{\text{eff}}^2}{c_{\text{est}}^2} \frac{M_s}{\lambda_{\text{max}}^2} \|O\|^2\right) = \varepsilon_{\text{testfail}}.$$

4. Numerical experiments. In this section we demonstrate first that the reduced local spaces generated by Algorithm 1 yield an approximation that converges at a nearly optimal rate. Moreover, we validate the a priori error bound in (3.2), the a posteriori error estimator (3.1), and the effectivity (3.4). To this end, we consider four test cases, starting in subsection 4.1 with an example for which the singular values of the transfer operator are known. The main focus of this subsection is a thorough validation of the theoretical findings in section 3, including a comprehensive testing on how the results depend on various parameters such as the basis size $n$, the number of test vectors $n_t$, and the mesh size. In addition, CPU time measurements are given. The second numerical example in subsection 4.2 examines the behavior
of the proposed algorithm in the more challenging case of the Helmholtz equation. In subsection 4.3 we numerically analyze the theoretical results from section 3 for a transfer operator whose singular values decay rather slowly and discrete spaces with large $N$, $N_S$, and $N_R$. Furthermore, we demonstrate that Algorithm 1 is computationally efficient. Finally, we employ the GFEM to construct a global approximation from the local reduced spaces generated by Algorithm 1 in the fourth test case in subsection 4.4, demonstrating that the excellent local approximation capacities of the local reduced spaces carry over to the global approximation.

For the implementation of the first test case, no FEM software library was used. The implementation for the third test case is based on the finite element library libMesh [51]. For the second and fourth test case we used the software library pyMOR [68]. The complete source code for reproduction of all results shown in subsections 4.1, 4.2 and 4.4 is provided in [14].

### 4.1. Analytic interface problem.

To analyze the behavior of the proposed algorithm, we first apply it to an analytic problem where the singular values of the transfer operator are known. We refer to this numerical example as Example 1. We consider the problem $A = -\Delta, f = 0$, and assume that $\Omega = (-L,L) \times (0,W)$, $\Gamma_{\text{out}} = \{-L,L\} \times (0,W)$, and $\Gamma_{\text{in}} = \{0\} \times (0,W)$. Moreover, we prescribe homogeneous Neumann boundary conditions on $\partial\Omega \setminus \Gamma_{\text{out}}$ and arbitrary Dirichlet boundary conditions on $\Gamma_{\text{out}}$, see also Figure 2.1 (left). The analytic solution is further discussed in the supplementary materials section SM2. This example was introduced in [80, Remark 3.3]. We equip $S$ and $R$ with the $L^2$-inner product on the respective interfaces. Recall that the transfer operator maps the Dirichlet data to the inner interface, i.e. with $H$ as the space of all discrete solutions, we define

$$T(v|_{\Gamma_{\text{out}}}) := v|_{\Gamma_{\text{in}}} \quad \forall v \in H.$$  

6 The singular values of the transfer operator are $\sigma_i = 1/(\sqrt{2}\cosh((i-1)\pi L/W))$. For the experiments, we use $L = W = 1$, unless stated otherwise. We discretize the problem by meshing it with a regular mesh of squares of size $h \cdot h$, where $1/h$ ranges from 20 to 320 in the experiments. On each square, we use bilinear $Q_1$ ansatz functions, which results in $51,681$ DOFs, $N_S = 322$ and $N_R = 161$ for $1/h = 160$.

In Figure 4.1b the first five basis vectors as generated by Algorithm 1 in one particular run are shown side by side with the first five basis vectors of the optimal space, i.e. with $H$ as the space of all discrete solutions, we define

$$\sigma_i = 1/(\sqrt{2}\cosh((i-1)\pi L/W)).$$

For the experiments, we use $L = W = 1$, unless stated otherwise. We discretize the problem by meshing it with a regular mesh of squares of size $h \cdot h$, where $1/h$ ranges from 20 to 320 in the experiments. On each square, we use bilinear $Q_1$ ansatz functions, which results in $51,681$ DOFs, $N_S = 322$ and $N_R = 161$ for $1/h = 160$.

In Figure 4.1b the first five basis vectors as generated by Algorithm 1 in one particular run are shown side by side with the first five basis vectors of the optimal space, i.e. the optimal modes in Figure 4.1a. While not identical, the basis functions generated using the randomized approach are smooth and have strong similarity with the optimal ones. Unless stated otherwise, we present statistics over 100,000 evaluations, use a maximum failure probability of $\varepsilon_{\text{algofail}} = 10^{-15}$, and use $\min(N_S, N_R)$ as an upper bound for $N_T$.

We first quantify the approximation quality of the spaces $R^n$ in dependence of the basis size $n$, disregarding the adaptive nature of Algorithm 1. In Figure 4.2a, statistics over the achieved projection error $\|T - P_{R^n}T\|$ are shown along with the singular values $\sigma_{n+1}$ of the transfer operator $T$. $\sigma_{n+1}$ is a lower bound for the projection error and it is the projection error that is achieved using an optimal basis. It shows that while the algorithm most of the time produces a basis nearly as good as the optimal basis, sometimes it needs two or three basis vectors more. This is in line with the

$6$Thanks to the form of the $A$-harmonic functions (SM2.1) and the fact that $\Omega$ is symmetric with respect to the $x_2$-axis, we have that $u - (1/|\Gamma_{\text{out}}|) \int_{\Gamma_{\text{out}}} u = u - (1/|\Omega|) \int_{\Omega} u$ and therefore that the singular vectors and singular values of (2.4) equal the ones of (4.1) apart from the constant function, which has to be added for the former but not for the latter.
predictions by theory, see the discussion after Proposition 3.2. The mean value of
the projection error converges with the same rate as the a priori error bound given
in Proposition 3.2 with increasing basis size. The a priori error bound is consistently
around three orders of magnitude larger than the actual error, until the actual error
hits the numerical noise between $10^{-14}$ and $10^{-15}$, see Figure 4.2b. This is mainly
due to the fact that the singular values decay very fast for the present example and
an index shift in the singular values by $p \geq 2$ as required by the a priori error bound
(3.2) therefore results in a much smaller error than predicted by the a priori error
bound. Note that we have $(\lambda_{\max}^{M, R}/\lambda_{\min}^{M, R})^{1/2} \approx (\lambda_{\max}^{M, S}/\lambda_{\min}^{M, S})^{1/2} \approx 2$.

The adaptive behavior of Algorithm 1 is analyzed in Figure 4.3. Figure 4.3a shows
that for $n_t = 10$ the algorithm succeeded to generate a space with the re-
quested approximation quality every single time in the 100,000 test runs and most
of the time, the approximation quality is about one or two orders of magnitude bet-
ter than required. Figure 4.3b shows the influence of the number of test vectors $n_t$:
With a low number of test vectors like 3 or 5, the algorithm produces spaces with an
approximation quality much better than requested, which is unfavorable as the basis
sizes are larger than necessary. 10 or 20 test vectors seem to be a good compromise,
as enlarging $n_t$ to 40 or 80 results in only little improvements while increasing com-
putational cost. This different behavior of Algorithm 1 for various numbers of test vectors \( n_t \) is due to the scaling of the effectivity of the a posteriori error estimator \( \eta(T - P_{R^n}T, n_t, \epsilon_{testfail}) \) as defined in (3.4) in the number of test vectors \( n_t \). The median effectivity \( \eta(T - P_{R^n}T, n_t, \epsilon_{testfail}) \) is 29.2 for \( n_t = 10 \), 10.4 for \( n_t = 20 \), and 6.1 for \( n_t = 40 \). We may thus also conclude that the a posteriori error estimator (3.1) is a sharp bound for the present test case.

Analyzing the numerical effectivity of the a posteriori error estimator \( \eta(T - P_{R^n}T, n_t, \epsilon_{testfail}) \) and comparing it to its theoretical upper bound \( c_{\text{eff}}(n_t, \epsilon_{testfail}) \) in Figure 4.5a, it can be observed that the theoretical upper bound becomes a sharper bound with increasing number of test vectors \( n_t \). The reason is the decreasing test vector norm

\[
\max_{i=1,...,n_t} \frac{\| (T - P_{R^n}T)D_S^{-1} \Sigma_i \|_R}{\| T - P_{R^n}T \|_R},
\]

as shown in Figure 4.5b. The normalized test vector norm is bound from above by \( c_{\text{est}}(n_t, \epsilon_{testfail})^{-1}c_{\text{eff}}(n_t, \epsilon_{testfail}) \) and from below by \( c_{\text{est}}(n_t, \epsilon_{testfail})^{-1} \) with error probability \( \epsilon_{testfail} \).

The quality of the produced spaces \( R^n \) should be independent of the mesh size \( h \). Figure 4.4a confirms this. After a preasymptotic regime, the deviation \( \| T - P_{R^n}T \| \) is independent of the mesh size. In the preasymptotic regime, the finite element space is not capable of approximating the corresponding modes. But while the deviation \( \| T - P_{R^n}T \| \) is independent of the mesh size, the norm of the test vectors used in the a posteriori error estimator in Algorithm 1 is not (see Figure 4.4b). The maximum norm of test vectors scales with the deviation and with \( \sqrt{h} \). In the adaptive algorithm, the scaling with \( \sqrt{h} \) is compensated by the factor \( (\lambda_{min}^n)^{-1/2} \) in \( c_{\text{est}}(n_t, \epsilon_{testfail}) \). To analyze the behavior in \( h \), the geometry parameters were chosen as \( L = 0.5 \) and \( W = 1 \) to have a slower decay of the singular values of the transfer operator.

To examine CPU times we use Example 1 in a larger configuration with \( L = 1 \), \( W = 8 \) and \( 1/h = 200 \). This results in 638,799 unknowns, \( N_S = 3202 \), and \( N_R = 1601 \). The measured CPU times for a simple, single threaded implementation are given in Table 4.1. The transfer operator is implemented implicitly. Its matrix is not assembled. Instead, the corresponding problem is solved using the sparse direct solver SuperLU [56, 21] each time the operator is applied. For Algorithm 1, a target accuracy \( tol \) of \( 10^{-4} \), the number of testvectors \( n_t = 20 \), and a maximum failure probability \( \epsilon_{\text{algorithm fail}} = 10^{-15} \) is used. In one test run, it resulted in an approximation space \( R^n \) of dimension 39. It only evaluated the operator \( n + n_t = 59 \) times. Each operator evaluation was measured to take 0.301 seconds, so a runtime of approximately \( (n + n_t) * 0.301 \) s \( \approx 17.8 \) s is expected. The measured runtime of 20.4 seconds is slightly higher, due to the orthonormalization of the basis vectors and the projection of the test vectors.

CPU times for the calculation of the optimal space of same size are given for comparison. The “eigs” function in “scipy.sparse.linalg”, which is based on ARPACK, is used to find the eigensystem of \( TT^* \). However, the calculation using ARPACK is not adaptive. To employ ARPACK, the required number of vectors has to be known in advance, which is why we expect that in general, the comparison would be even more in favor of the adaptive randomized algorithm.

4.2. Helmholtz equation. In this subsection we analyze the behavior of the proposed algorithm in a numerical test case approximating the solution of the Helm-
Properties of transfer operator

unknowns of corresponding problem | 638,799
---|---
LU factorization time in s | 14.1
operator evaluation time in s | 0.301
adjoint operator evaluation time in s | 0.301

Properties of basis generation

| Algorithm 1 | Scipy/ARPACK |
|---|---|
| (resulting) basis size | 39 | 39 |
| operator evaluations | 59 | 79 |
| adjoint operator evaluations | 0 | 79 |
| execution time in s (w/o factorization) | 20.4 | 47.9 |

Table 4.1: CPU times for Example 1 with $L = 1$, $W = 8$ and $1/h = 200$. Single threaded performance.

(a) Quartiles for 10 test vectors. (b) Maximum error for given number of test vectors.

Figure 4.3: Projection error $\sup_{\zeta \in S} \inf_{\xi \in R^n} \frac{\|T\zeta - T\xi\|}{\|\xi\|} = \|T - P_{R^n}T\|$ over target projection error for Example 1. Meshsize $h = 1/160$.

(a) $h$ dependency of deviation (b) $h$ dependency of testvector norm

Figure 4.4: $h$ dependency for Example 1. Statistics over 10,000 samples.
holtz equation. The domain Ω, the boundaries Γ_{in} and Γ_{out} and the boundary conditions are the same as in subsection 4.1, only the operator \( A \) differs and is defined as \( A = -\Delta - \kappa^2 \) in this subsection. As for Example 1, it has 51,681 DOFs, \( N_S = 322 \) and \( N_R = 161 \) for \( 1/h = 160 \). We refer to this numerical example as Example 2. We assume the problem to be inf-sup stable and thus uniquely solvable, which is the case as long as it is not in a resonant configuration. A treatment of the resonant case is beyond the scope of this publication.

For \( \kappa = 0 \) we obtain Example 1. We observe that the singular values of the transfer operator first have a plateau and then decay exponentially, see Figure 4.6. The longer the plateau, the faster is the exponential decay. The length of the plateau is observed to be very close to the length of the inner interface divided by a half wavelength, i.e. \( 1/(\lambda/2) = \kappa/\pi \). Comparing this with the analysis of Finite Element methods for the Helmholtz equation (cf. [50]), one finds this similar to the “minimal resolution condition” \( 1/h \geq \sqrt{12}/\kappa \).

Algorithm 1 succeeds to generate reduced spaces \( R^n \) which achieve a projection error \( \| T - P_{R^n} \| \) which is close the the optimal projection error given by the singular values of the transfer operator. We show results for \( \kappa = 30 \) in Figure 4.7. Also in the adaptive case, we observe the expected behavior, see Figure 4.8a and Figure 4.8b. The plateaus which can be observed in Figure 4.8a are due to the very fast decay of the singular values. E.g. the first plateau is at an error of about \( 10^{-3} \), which is the
4.3. A transfer operator with slowly decaying singular values; application to linear elasticity. In this subsection we numerically analyze Algorithm 1 and the theoretical findings of section 3 for a numerical test case Example 3 where the singular values of the transfer operator exhibit a relatively slow decay and $N$, $N_S$, and $N_R$ are relatively large. Moreover, we shortly illustrate that Algorithm 1 is attractive from a computational viewpoint and with respect to memory requirement.

To that end let $\Omega_{in} = (-0.5, 0.5) \times (-0.5, 0.5) \times (-0.5, 0.5)$ be the subdomain on which we aim to construct a local approximation space, $\Omega = (-2,2) \times (-0.5, 0.5) \times (-2,2)$ the (oversampling) domain, and $\Gamma_{out} = \{-2,2\} \times (-0.5, 0.5) \times (-2,2)$ the target error usually achieved at a basis size of 10 (cf. Figure 4.7a). The next plateau at an error of about $10^{-7}$ corresponds to a basis size of 11.
$\sigma_k$ is the fourth-order stiffness tensor
\[ C_{ijkl} = \frac{\nu}{(1 + \nu)(1 - 2\nu)} \delta_{ij} \delta_{kl} + \frac{1}{2(1 + \nu)} (\delta_{ik} \delta_{jl} + \delta_{il} \delta_{jk}), \quad 1 \leq i, j, k, l \leq 3, \]

where $\delta_{ij}$ denotes the Kronecker delta, and we choose Poisson’s ratio $\nu = 0.3$. Moreover, $\varepsilon(u) = 0.5(\nabla u + (\nabla u)^T)$ is the infinitesimal strain tensor and the colon operator is defined as $C : \varepsilon(u) = \sum_{i,j,l=1}^{3} C_{ijkl}\varepsilon_{kl}(u)$.

For the FE discretization we use a regular mesh with hexahedral elements and a mesh size $h = 0.1$ in each space direction and a corresponding FE space $X$ with linear FE resulting in $\dim(X) = N = 55473$, $\dim(R) = N_R = 3987$, and $\dim(S) = N_S = 5280$. Note that although in theory we should subtract the orthogonal projection on the six rigid body motions from the FE basis functions, in actual practice we avoid that by subtracting the orthogonal projection from the $A$-harmonic extensions only. Finally, we equip the source space $S$ with the $L^2$-inner product and the range space $R$ with the energy inner product
\[ (w, v)_R := \int_{\Omega_n} \frac{\partial w^i}{\partial x_j} C_{ijkl} \frac{\partial v^k}{\partial x_l} dx. \]

Analyzing the convergence behavior of $\mathbb{E}(\|T - P_{R^{k+1}}T\|)$ for a growing number of randomly generated basis functions $k$ and a (fixed) oversampling parameter $p = 2$ in Figure 4.9a we observe that the local approximation spaces generated as proposed in section 3 yield an approximation that converges nearly with the optimal rate $\sigma_k$. Moreover, we see in Figure 4.9a that the a priori error bound as proposed in (3.2) reproduces the convergence behavior of $\mathbb{E}(\|T - P_{R^{k+1}}T\|)$ quite well, where the mean
of the deviation converges slightly faster than the a priori error bound. Furthermore, for the present test case the a priori error bound seems to behave like $\sqrt{K\sigma_{k+1}}$, arguing that the latter might be the dominating factor. Finally, we observe that the a priori bound is rather pessimistic as it overestimates $\mathbb{E}(\|T - P_{R^k}T\|)$ by a factor of more than 100. This is mainly due to the square root of the conditions of the inner product matrices which amount to $(\lambda_{\max}/\lambda_{\min})^{1/2} \approx 17.3197$ and $(\lambda_{\max}/\lambda_{\min})^{1/2} \approx 3.4404$.

If we consider a flatter domain $\hat{\Omega} = (-2,2) \times (-0.25,0.25) \times (-2,2)$ and a flatter subdomain $\tilde{\Omega}_{in} = (-0.5,0.5) \times (-0.25,0.25) \times (-0.5,0.5)$ instead, where $\tilde{\Gamma}_{out} = \{-2,2\} \times (-0.25,0.25) \times (-2,2) \cup (-2,2) \times (-0.25,0.25) \times \{-2,2\}$ and we still consider the same PDE and the same inner products as above, we observe in Figure 4.9b that until $k \approx 75$ the a priori bound reproduces the convergence behavior of $\mathbb{E}(\|T - P_{R^k}T\|)$ perfectly. We may thus conclude that the a priori bound in (3.2) seems to be sharp regarding the convergence behavior of $\mathbb{E}(\|T - P_{R^k}T\|)$ in the basis size $k$. The a priori estimates could be improved slightly by finding the optimal oversampling size $\rho$, which was fixed to its minimum value of 2 in this experiment. Expecially on the domain $\Omega$, where the singular values of the transfer operator show a slower decay, a larger oversampling would be beneficial. For the computations on $\hat{\Omega}$ we employed again a regular hexaedral mesh with $h = 0.1$ and linear FE with $N = 30258$, $N_R = 2172$, and $N_S = 2880$. Finally, for all results in this subsection we computed the statistics over 1000 samples. From now on all results are computed on $\Omega$.

Regarding the performance of Algorithm 1 we first observe in Figure 4.10a that the actual error $\|T - P_{R^t}T\|$ lies below the target tolerance $\text{tol}$ for all 1000 samples for $n_t = 10$, which holds also true for all other considered values of $n_t$. Here, we prescribe $\varepsilon_{\text{algofail}} = 10^{-10}$ and use 3993 as an upper bound for $N_T$ throughout this subsection. Compared with the performance of Algorithm 1 for Example 1 in Figure 4.3a the dispersion in Figure 4.10a is much smaller. This may be explained by the much faster decay of the singular values of $T$ and therefore $\|T - P_{R^t}T\|$ in subsection 4.1 compared with the present test case.

Similarly to Figure 4.3b and Figure 4.8b in subsection 4.1 and subsection 4.2 we see in Figure 4.10a that increasing the number of test vectors $n_t$ from 5 to 10 or from 10 to 20 increases the ratio between the median of the actual error $\|T - P_{R^t}T\|$ and

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Figure 4.10: Median of the projection error $\|T - P_{R^t}T\|$ for a decreasing target accuracy $\text{tol}$ for a varying number of test vectors $n_t$ and the minimal and maximal values for $n_t = 10$ (a). Behavior of the median of the effectivity $\eta(T - P_{R^t}T, n_t, 10^{-10})$ as defined in (3.4) for growing $n$ and various number of test vectors $n_t$ (b).
the target accuracy $\text{tol}$ significantly — for the former by more than one magnitude — while an increase from $n_t = 40$ to $n_t = 80$ has hardly any influence. This can be explained by the scaling of the effectiveness of the employed a posteriori error estimator defined in (3.4) which we will elaborate on shortly. Regarding the choice of $n_t$ it seems that for the present test case a value of about 20 is in the sweet spot.

In order to successfully apply the proposed algorithm in the context of a method, it has to be possible to define a transfer operator with quickly decaying spectrum. Moreover, allowing to bound the local approximation error in terms of the local error contributions, is sufficient (but not necessary) to yield a global error decaying as $\sqrt{n}\sigma_{n+1}$ or better. All of this is possible for the GFEM, which is why we employ this method in this subsection to build a global approximation from the local reduced spaces generated by Algorithm 1. We refer to this numerical example as Example 4. The convergence theory for the GFEM with randomized basis generation is given in the supplementary materials subsection SM5.2.

On $\Omega_{gl} = (0, 1)^2$ we consider the following PDE: Find $u_{gl} \in X_{gl} = \{v \in H^1(\Omega_{gl}) : v = 0 \text{ on } \Sigma_D\}$, such that

$$-\text{div}(k\nabla u_{gl}) = f \text{ in } X'_{gl},$$

where $k \in L^\infty(\Omega_{gl}), 0 < k_0 \leq k \leq k_1 < \infty$ and $f(\varphi) := \int_{\Omega_{gl}} \hat{f}\varphi \, dx$ for a source term $\hat{f} \in L^2(\Omega_{gl})$. We use the GFEM to compute an approximation of $u_{gl}$. To this end let $\{\omega_i\}_{i=1}^m$ be an open cover of $\Omega_{gl}$ such that $\Omega_{gl} = \bigcup_{i=1}^m \omega_i$. For each domain, we define a local space $X_i := \{v|_{\omega_i} \mid v \in X_{gl}\}$. We will construct local reduced spaces $R_i^n \subset X_i$ and the global GFEM space $X_{\text{GFEM}} := \bigoplus_{i=1,\ldots,m} \{q_i v_i \mid v_i \in R_i^n\}$, where $q_i$ is a suitably defined partition of unity (see the supplementary materials subsection SM5.2). The GFEM solution $u_{\text{GFEM}} \in X_{\text{GFEM}}$ is then defined as the solution of

$$-\text{div}(k\nabla u_{\text{GFEM}}) = f \text{ in } X'_{\text{GFEM}},$$

We construct reduced spaces $R_i^n$, each approximating the solution $u_{gl}$ on $\omega_i$. To this end, we introduce $\omega_i^\rho$, satisfying $\omega_i \subset \omega_i^\rho \subset \Omega_{gl}$ with $\text{dist}(\partial \omega_i^\rho \setminus \partial \Omega_{gl}, \partial \omega_i) \geq \rho > 0$, which denotes the oversampling domain used to construct the reduced space and thus
the interior, it holds differ for domains in the interior, at the boundary and in the corners. For domains in \( \omega \) due to the Dirichlet boundary. As local domains

\[ \text{Example 4.1} \]

We demonstrate the GFEM with randomized basis generation on two examples. In particular the solution of this example (Example 4.1 in the following) is more complex and features small details, while the domains

\[ \text{Example 4.2} \]

The spaces \( \text{Example 2} \); white equates to 1 and black to \( 10^2 \).

corresponds to \( \Omega \) in the remainder of this article. We denote its inner boundary \( \partial \omega_i \setminus \partial \Omega_{gl} \) by \( \Gamma_{out,i} \). Denoting the space of \( A \)-harmonic functions on \( \omega_i^* \) as \( \hat{H}_i \), the transfer operator is defined as

\[ T(w|_{\Gamma_{out,i}}) := (w - P_{\ker(A),\omega_i}(w))|_{\omega_i} \quad \text{for all } w \in \hat{H}_i. \]

The spaces \( S_i := \{ w|_{\Gamma_{out,i}} : w \in \hat{H}_i \} \) are equipped with the \( L^2 \)-inner product. In the range spaces \( R_i := \{ (w - P_{\ker(A),\omega_i})|_{\omega_i} : w \in \hat{H}_i \} \) we use the energy inner product. We apply Algorithm 1 to these transfer operators using \( n_t = 20 \) test vectors throughout this subsection and a global maximum failure probability of \( \varepsilon_{\text{fail}} = 10^{-15} \). For the full GFEM algorithm with all details, see the supplementary materials subsection SM5.1.

We demonstrate the GFEM with randomized basis generation on two examples. The first example (Example 4.1 in the following) is the Poisson problem \(-\Delta u = 1\), i.e. \( k \equiv 1 \) and \( \hat{f} \equiv 1 \). We consider this problem because the singular values and singular vectors of all transfer operators associated with subdomains \( \omega_i^* \) that do not lie on the boundary of \( \Omega_{gl} \) are the same and thus only the boundary has a (slight) influence. Therefore, we would expect that for this test case the convergence rate of the global error is similar to the convergence rate of \( \| T - P_{R_i} T \| \). The second example (Example 4.2 in the following) is more complex and features small details, high contrast, and high conductivity channels. In particular the solution of this example is non-smooth. For Example 4.2, we define a high conductivity region \( \Omega_{hc} := [(0.02, 0.1) \times (0.02, 0.98)] \cup [(0.9, 0.98) \times (0.02, 0.98)] \cup [(0.11, 0.89) \times (0.475, 0.485)] \cup [(0.1, 0.9) \times (0.495, 0.505)] \cup [(0.11, 0.89) \times (0.515, 0.525)] \) and define \( k(x) := 10^5 \) for \( x \in \Omega_{hc} \) and \( k(x) := 1 \) else. For the right hand side, we define a heating region \( \Omega_{heat} := (0.9, 0.98) \times (0.02, 0.98) \) and a cooling region \( \Omega_{cool} := (0.02, 0.1) \times (0.02, 0.98) \) and define the right hand side as

\[ \hat{f}(x) := 1 \quad \text{for } x \in \Omega_{heat}, \quad \hat{f}(x) := -1 \quad \text{for } x \in \Omega_{cool}, \quad \text{and } \hat{f}(x) := 0 \quad \text{else}. \]

For both examples, the domain is discretized using a regular mesh where the domain is partitioned into 200 \( \times \) 200 squares of size 0.005\(^2\), each of which is divided into four triangles. On this mesh, standard \( P1 \) basis functions are used, spanning the FEM space. It has 80401 degrees of freedom, of which 800 are constrained due to the Dirichlet boundary. As local domains \( \omega_i \) we use patches of size 0.2 \( \times \) 0.2 with an overlap of size 0.1. This accounts for 9 \( \times \) 9 = 81 subdomains \( \omega_i \). For the oversampling size we also use 0.1, so the domains \( \omega_i^* \) in the interior have size 0.4 \( \times \) 0.4 while the domains \( \omega_i^* \) at one boundary have size 0.4 \( \times \) 0.3 or 0.3 \( \times \) 0.4 and the domains \( \omega_i^* \) in the corners have size 0.3 \( \times \) 0.3. The dimension of the source spaces \( N_{S_i} \) and \( N_{R_i} \) differ for domains in the interior, at the boundary and in the corners. For domains in the interior, it holds \( N_{S_i} = 320 \) and \( N_{R_i} = 1681 \). All 81 transfer operators in the two

\[ \text{Figure 4.11: Coefficient field for GFEM Example 2; white equates to 1 and black to } 10^2. \]

\[ \text{Figure 4.12: The slowest SVD decay (svd), the maximum relative local error (local), and the relative global error (global) for the two GFEM examples (Example 4). Median values over 100 realizations.} \]
examples have an exponentially decaying spectrum. The slowest decaying spectrum is shown in Figure 4.12 along with the maximum local relative approximation error and the global relative approximation error. The maximum relative local approximation error follows the spectrum of the transfer operator closely for Example 4.1. For Example 4.2, the behavior is similar. As anticipated, this error decay propagates to the relative global approximation error, which flattens out due to numerical effects at about $10^{-13}$ for Example 4.1 and $10^{-10}$ for Example 4.2. To compute the spectrum of the transfer operators, the numerically more accurate eigenvalue problem presented in the supplementary materials section SM6 is used. To further examine the behavior of the local error $\min_{v_i \in R^n_{i}} \|k^{1/2} \nabla (u_{gl} - v_i)\|_{L^2(\omega_i)} / \|k^{1/2} \nabla u_{gl}\|_{L^2(\omega_i)}$ when using the adaptive algorithm, we construct local approximation spaces $R^n_{i}$ with the proposed adaptive range recovery and measure the relative local error. Statistics over 1,000 different realizations and all 81 local spaces show that results are more accurate than required by about 2.5 orders of magnitude for Example 4.1 and about 3.5 orders of magnitude for Example 4.2, see Figure 4.13. This discrepancy is in part caused by the fact that the adaptive range approximation generates spaces which are better than required, as was already discussed in subsections 4.1 and 4.3. This accounts for about 1 to 1.5 orders of magnitude. The other part is the pessimistic estimate for the local error given in the supplementary materials, Lemma SM5.2. The error decay propagates to the global relative error $(\|k^{1/2} \nabla (u_{gl} - u_{GFEM})\|_{L^2(\omega)} ) / (\|k^{1/2} \nabla u_{gl}\|_{L^2(\omega)})$. It is possible to choose a target error $tol_{GFEM}$ and choose all tolerances accordingly, so the resulting approximation will have at most this relative error: From $tol_{GFEM}$ as a target maximum for the global relative error, we calculate the maximum local relative error using Proposition SM5.1. Using this maximum local relative error, we calculate a limit for the operator norm $\|T_i - P_{R^n_{i}}\|$ using Lemma SM5.2. This limit for the operator norm is then used to steer the adaptive range finder algorithm, Algorithm 1. There are no unknown constants. For details, see the supplementary materials subsection SM5.2. The global relative error, shown in Figure 4.14, confirms this and is more accurate than required by about 4.5 orders of magnitude for Example 4.1 and about 6.5 orders of magnitude for Example 4.2. From the local error to the global error, we lose about 2 orders of magnitude for Example 4.1 and about 3 orders of magnitude for Example 4.2. This is due to the pessimistic estimate in Proposition SM5.1.
5. Conclusions. Recently, optimal local reduced spaces for localized MOR procedures have been proposed in [9, 80]. However, a straightforward FE approximation of those optimal local spaces is very expensive. In this article we have proposed an adaptive randomized range finder algorithm based on methods from randomized LA [38] that adaptively builds local reduced spaces for localized MOR procedures from local solutions of the PDE with random boundary conditions. Starting from results in randomized LA [38, 35, 36, 18] we have shown that the randomly generated local reduced spaces produce an approximation with a convergence rate that is only slightly worse than the optimal rate; the rate is deteriorated by about the square root of the basis size. Finally, the adaptive randomized range finder algorithm is steered by a probabilistic a posteriori error estimator for which we have demonstrated its efficiency.

The numerical experiments show that the local spaces constructed by the adaptive randomized range finder algorithm indeed converge with a nearly optimal rate. It can also be seen that the a priori error bound seems to be sharp in the sense that for some numerical experiments the projection error converges exactly as predicted by the a priori error bound. Moreover, we observed that after a preasymptotic regime the convergence behavior of the projection error is independent of the mesh size, indicating that it might be possible to generalize the results of the present paper to the continuous setting. This is the subject of future work. For the GFEM we exemplarily demonstrated in the numerical experiments that the excellent local approximation behavior of the randomly generated spaces carries over to the global level for an example with high conductivity channels. Regarding the probabilistic a posteriori error estimator the numerical experiments have showed also for a transfer operator with slowly decaying singular values and discrete source and range spaces of rather large dimensions that we only need 10 to 20 test vectors in order to obtain a sharp bound. Thanks to both the nearly optimal approximation capacities of the randomly generated local reduced spaces and the good effectivity of the probabilistic a posteriori error estimator also for few numbers of test vectors the adaptive randomized range finder algorithm requires only very few local solutions of the PDE in addition and therefore has a close to optimal computational complexity and is faster than the calculation of the optimal spaces using ARPACK on the corresponding eigenvalue problem.

The extension of the proposed method to transient and nonlinear problems is
the subject of future work. We note that although prescribing random boundary conditions and using the solution of the nonlinear PDE evaluated either on an interface or a subdomain might actually yield a reduced space with decent approximation properties, we believe that the corresponding numerical analysis might however be quite involved.

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