A STENCIL SCALING APPROACH FOR ACCELERATING
MATRIX-FREE FINITE ELEMENT IMPLEMENTATIONS

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Abstract. We present a novel approach to fast on-the-fly low order finite element assembly
for scalar elliptic partial differential equations of Darcy type with variable coefficients optimized for
matrix-free implementations. Our approach introduces a new operator that is obtained by appro-
priately scaling the reference stiffness matrix from the constant coefficient case. Assuming sufficient
regularity, an a priori analysis shows that solutions obtained by this approach are unique and have
asymptotically optimal order convergence in the $H^1$- and the $L^2$-norms on hierarchical hybrid grids.
For the preasymptotic regime, we present a local modification that guarantees uniform ellipticity of
the operator. Cost considerations show that our novel approach requires roughly one-third of the
floating-point operations compared to a classical finite element assembly scheme employing nodal
integration. Our theoretical considerations are illustrated by numerical tests that confirm the expect-
tations with respect to accuracy and run-time. A large scale application with more than a hundred
billion ($1.6 \cdot 10^{11}$) degrees of freedom executed on 14310 compute cores demonstrates the efficiency
of the new scaling approach.

Key words. matrix-free, finite elements, variable coefficients, stencil scaling, variational crime
analysis, optimal order a priori estimates

AMS subject classifications. 65N15, 65N30, 65Y20

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1. Introduction. Traditional finite element implementations are based on com-
puting local element stiffness matrices, followed by a local-to-global assembly step,
resulting in a sparse matrix. However, the cost of storing the global stiffness matrix
is significant. Even for scalar equations and low order 3D tetrahedral elements, the
stiffness matrix has, on average, fifteen entries per row, and thus a standard sparse
matrix format will require thirty times as much storage for the matrix as for the solu-
tion vector. This limits the size of the problems that can be tackled and becomes the
dominating cost factor since the sparse matrix must be reread from memory repeat-
edly when iterative solvers are applied. On all current and future computing systems
memory throughput and memory access latency can determine the run-time more crit-
ically than the floating-point operations executed. Furthermore, energy consumption
has been identified as one of the fundamental roadblocks in exascale computing. In
this cost metric, memory access is again more expensive than computation. Against
the backdrop of this technological development, it has become mandatory to develop
numerical techniques that reduce memory traffic. In the context of partial differen-
tial equations (PDEs) this is leading to a renewed interest in so-called matrix-free
Matrix-free techniques are motivated by the observation that many iterative solvers, e.g., Richardson iteration or Krylov subspace methods, require only the result of multiplying the global system matrix with a vector, but not the matrix itself. The former can be computed by local operations in each element, which avoids setting up, storing, and loading the global stiffness matrix. One of the first papers in this direction is [10], which describes the so-called element-by-element (EBE) approach, in which the global matrix-vector product (MVP) is assembled from the contributions of MVPs of element matrices with local vectors of unknowns. The element matrices are either precomputed and stored or recomputed on the fly. Note that storing all element matrices, even for low order elements, has a higher memory footprint than the global matrix itself.

Consequently, the traditional EBE approach has not found wide application in unstructured mesh settings. However, it has been successfully applied in cases where the discretization is based on undeformed hexahedral elements with trilinear trial functions; see, e.g., [1, 7, 15, 37]. In such a setting, the element matrix is the same for all elements, which significantly reduces the storage requirements, and variable material parameters can be introduced by weighting local matrix entries in the same on-the-fly fashion as will be developed in this paper.

Matrix-free approaches for higher order elements, as described in, e.g., [9, 25, 28, 29, 32], differ from the classic EBE approach in that they do not set up the element matrix and consecutively multiply it with the local vector. Instead, they fuse the two steps by going back to numerical integration of the weak form itself. The process is accelerated by precomputing and storing certain quantities, such as, e.g., derivatives of basis functions at quadrature points within a reference element. These techniques in principle work for arbitrarily shaped elements and orders, although a significant reduction in complexity can be achieved for tensor-product elements.

However, these matrix-free approaches also have shortcomings. While the low order settings [1, 7, 15, 37] require structured hexahedral meshes, modern techniques for unstructured meshes only pay off for elements with tensor-product spaces with polynomial orders of at least two; see [25, 28].

In this paper we will present a novel matrix-free approach for low order finite elements designed for the hierarchical hybrid grids framework (HHG); see, e.g., [3, 4, 5]. HHG offers significantly more geometric flexibility than undeformed hexahedral meshes. It is based on two interleaved ideas. The first one is a special discretization of the problem domain. In HHG, the computational grid is created by way of a uniform refinement following the rules of [6], starting from a possibly unstructured simplicial macro-mesh. The resulting nested hierarchy of meshes allows for the implementation of powerful geometric multigrid solvers. The elements of the macro-mesh are called macro-elements, and the resulting subtriangulation reflects a uniform structure within these macro-elements. The second aspect is based on the fact that each row of the global finite element stiffness matrix can be considered as a difference stencil. This notion and point of view is classical on structured grids and recently has found renewed interest in the context of finite elements too; see, e.g., [14]. In combination with the HHG grid construction this implies that for linear simplicial elements one obtains stencils with identical structure for each inner node of a macro primitive.

However, it requires less memory than storage schemes for sparse direct solvers which reserve space for fill-in, the original competing scenario in [10].
We define macro primitives as the geometrical entities of the macro-mesh of different dimensions, i.e., vertex, edge, face, and tetrahedrons. If additionally the coefficients of the PDE are constant per macro-element, then also the stencil entries are exactly the same. Consequently, only a few different stencils (one per macro primitive) can occur and need to be stored. This leads to extremely efficient matrix-free techniques, as has been demonstrated, e.g., in [3, 18].

Let us now consider the setting of an elliptic PDE with piecewise smooth variable coefficients, assuming that the macro-mesh resolves jumps in the coefficients. In this case, a standard finite element formulation is based on quadrature formulas and introduces a variational crime. According to [11, 34], there is flexibility in how the integrals are approximated without degenerating the order of convergence. This was recently exploited in [2] with a method that approximates these integral values on the fly using suitable surrogate polynomials with respect to the macro-mesh. The resulting two-scale method is able to preserve the convergence order if the coarse and fine scales are related properly. Here we propose an alternative which is based on the fine scale.

For this article, we restrict ourselves to the lowest order case of conforming finite elements on simplicial meshes. Then the most popular quadrature formula is the one-point Gauss rule, which in the simplest case of \( \text{div}(k \nabla u) \) as PDE operator just weights the element-based reference stiffness matrix of the Laplacian by a factor of \( k(x_T) \), where \( x_T \) is the barycenter of the element \( T \). Alternatively, one can select a purely vertex-based quadrature formula. Here, the weighting of the element matrix is given by \( \sum_{i=1}^{d+1} k(x^i_T) / (d + 1) \), where \( d \) is the space dimension and \( x^i_T \) are the vertices of element \( T \). Using a vertex-based quadrature formula saves function evaluations and is thus attractive whenever the evaluation of the coefficient function is expensive, and it pays off to reuse once-computed values in several element stiffness matrices. Note that reusing barycentric data on general unstructured meshes will require nontrivial storage schemes.

In the case of variable coefficient functions, stencil entries can vary from one mesh node to another. The number of possibly different stencils within each macro-element becomes \( \frac{1}{2} 2^{d \ell} + \mathcal{O}(2^{(d-1)\ell}) \), where \( \ell \) is the number of uniform refinement steps for HHG. Now we can resort to two options: Either these stencils are computed once and then saved, effectively creating a sparse matrix data structure, or they are computed on the fly each time they are needed. Neither of these techniques is ideal for extreme scale computations. While for the first option \( \mathcal{O}(2^{d \ell}) \) extra memory is consumed and extensive memory traffic occurs, the second option requires recomputation of \( \mathcal{O}(2^{d \ell}) \) local contributions.

The efficiency of a numerical PDE solver can be analyzed following the textbook paradigm [8] that defines a work unit (WU) to be the cost of one application of the discrete operator for a given problem. With this definition, the analysis of iterative solvers can be conducted in terms of WU. Classical multigrid textbook efficiency is achieved when the solution is obtained in less than 10 WU. For devising an efficient method it is, however, equally critical to design algorithms that reduce the cost of a WU without sacrificing accuracy. Clearly, the real-life cost of a WU depends on the computer hardware and the efficiency of the implementation, as, e.g., analyzed for parallel supercomputers in [18]. On the other side, matrix-free techniques, such as the one proposed in this article, seek opportunities to reduce the cost of a WU by a clever rearrangement of the algorithms or by exploiting approximations where this is possible; see also, e.g., [2].

These preliminary considerations motivate our novel approach to reducing the
cost of a WU by recomputing the surrogate stencil entries for a matrix-free solver more efficiently. We find that these values can be assembled from a reference stencil of the constant coefficient case which is scaled appropriately using nodal values of the coefficient function. We will show that under suitable conditions, this technique does not sacrifice accuracy. However, we also demonstrate that the new method can reduce the cost of a WU considerably and in consequence helps reduce the time-to-solution.

The rest of this paper is structured as follows: In section 2, we define our new scaling approach. The variational crime is analyzed in section 3, where optimal order a priori results for the $L^2$- and $H^1$-norms are obtained. In section 4, we consider modifications in the preasymptotic regime to guarantee uniform ellipticity. Section 5 is devoted to the reproduction property and the primitive concept which allows for a fast on-the-fly reassembling in a matrix-free software framework. In section 6, we discuss the cost compared to a standard nodal-based elementwise assembling. Finally, in section 7 we perform numerically an accuracy study and a run-time comparison to illustrate the performance gain of the new scaling approach.

2. Problem setting and definition of the scaling approach. We consider a scalar elliptic PDE of Darcy type, i.e.,

$$-
\text{div}
K \nabla u = f \quad \text{in } \Omega, \quad \text{tr } u = 0 \quad \text{on } \partial \Omega,$$

where $\text{tr}$ stands for the boundary trace operator and $f \in L^2(\Omega)$. Here $\Omega \subset \mathbb{R}^d$, $d = 2, 3$, is a bounded polygonal/polyhedral domain, and $K$ denotes a uniformly positive and symmetric tensor with coefficients specified through a number of functions $k_m$, $m = 1, \ldots, M$, where $M \leq 3$ in 2D and $M \leq 6$ in 3D due to symmetry.

For the Darcy operator with a scalar uniform positive permeability, that is, $-\text{div}(k \nabla u)$, we can set $M = 1$ and $k_1 := k$. The above setting also covers blending finite elements approaches [19]. Here $K$ is related to the Jacobian of the blending function. For example, if the standard Laplacian model problem is considered on the physical domain $\Omega_{\text{phy}}$ but the actual assembly is carried out on a reference domain $\Omega := \Phi(\Omega_{\text{phy}})$, we have

$$a(v, w) = \int_{\Omega_{\text{phy}}} \nabla v_{\text{phy}} \cdot \nabla w_{\text{phy}} \, dx_{\text{phy}} = \int_{\Omega} \nabla v \cdot \frac{(D\Phi)(D\Phi)^\top}{|\det D\Phi|} \nabla w \, dx,$$

where $D\Phi$ is the Jacobian of the mapping $\Phi$, and $v_{\text{phy}} := v \circ \Phi$, $w_{\text{phy}} := w \circ \Phi$.

2.1. Definition of our scaling approach. The weak form associated with the PDE is defined in terms of the bilinear form $a(v, w) := \int_{\Omega} \nabla v \cdot K \nabla w \, dx$, and the weak solution $u \in V_0 := H^1_0(\Omega)$ satisfies

$$a(u, v) = (f, v), \quad v \in V_0.$$

This bilinear form can be affinely decomposed as

$$a(v, w) := \sum_{m=1}^{M} a_m(v, w), \quad a_m(v, w) := \int_{\Omega} k_m(x)(D_m v, D_m w) \, dx, \quad v, w \in V := H^1(\Omega),$$

where $D_m$ is a first order partial differential operator and $(\cdot, \cdot)$ stands for some suitable inner product. In the case of a scalar permeability we find $D_1 := \nabla$ and $(\cdot, \cdot)$ stands
for the scalar product in \( \mathbb{R}^d \). For (2.1) in 2D, however, one can, as one alternative, e.g., define

\[
\begin{align*}
k_1 &:= (K_{11} - K_{12}), & k_2 &:= K_{12}, & k_3 &:= (K_{22} - K_{11}), \\
D_1 &:= \nabla, & D_2 &:= \partial/\partial x + \partial/\partial y, & D_3 &:= \partial/\partial y,
\end{align*}
\]

where \( K = (K_{ij}) \) and the same scalar product \((\cdot,\cdot)\) as above. Note that this decomposition reduces to the one in case of a scalar permeability, i.e., for \( K = \text{diag}(k, k) \).

Let \( \mathcal{T}_H, H > 0 \) fixed, be a possibly unstructured simplicial triangulation resolving \( \Omega \). We call \( \mathcal{T}_H \) also macro-triangulation and denote its elements by \( T \). Using uniform mesh refinement, we obtain \( \mathcal{T}_{H/2} \) from \( \mathcal{T}_H \) by decomposing each element into \( 2^d \) subelements, \( h \in \{ H/2, H/4, \ldots \} \); see [6] for the 3D case. The elements of \( \mathcal{T}_H \) are denoted by \( T \). The macro-triangulation is then decomposed into the following geometrical primitives: elements, faces, edges, and vertices. Each of these geometric primitives acts as a container for a subset of unknowns associated with the refined triangulations. These sets of unknowns can be stored in array-like data structures, resulting in a contiguous memory layout that conforms inherently to the refinement hierarchy; see [5, 18]. In particular, the unknowns can be accessed without indirect addressing such that the overhead is reduced significantly when compared to conventional sparse matrix data structures. Associated with \( \mathcal{T}_H \) is the space \( V_h \subset V \) of piecewise linear finite elements. In \( V_h \), we do not include the homogeneous boundary conditions. We denote by \( \phi_i \in \chi_{V_h} \) the nodal basis functions associated to the ith mesh node. Node \( i \) is located at the vertex \( x_i \). For \( v_h := \sum_i \nu_i \phi_i \) and \( w_h := \sum_j \chi_j \phi_j \), we define our scaled discrete bilinear forms \( a_h(\cdot, \cdot) \) and \( a_m^h(\cdot, \cdot) \) by

\[
\begin{align*}
(2.3a) & \quad a_h(v_h, w_h) := \sum_{m=1}^M a_m^h(v_h, w_h), \\
(2.3b) & \quad a_m^h(v_h, w_h) := \frac{1}{4} \sum_{T \in \mathcal{T}_H} \sum_{i,j} (k_m|_T(x_i) + k_m|_T(x_j))(\nu_i - \nu_j)(\chi_j - \chi_i) \int_T (D_m \phi_i, D_m \phi_j) \, dx.
\end{align*}
\]

This definition is motivated by the fact that \( a_m(v_h, w_h) \) can be written as

\[
(2.4) \quad a_m(v_h, w_h) = \frac{1}{2} \sum_{T \in \mathcal{T}_H} \sum_{i,j} (\nu_i - \nu_j)(\chi_j - \chi_i) \int_T k_m(x)(D_m \phi_i, D_m \phi_j) \, dx.
\]

Here we have exploited symmetry and the row sum property. It is obvious that if \( k_m \) is a constant restricted to \( T \), we do obtain \( a_m^h(v_h, w_h) = a_m(v_h, w_h) \). In general, however, the definition of \( a_m(\cdot, \cdot) \) introduces a variational crime, and it does not even correspond to an elementwise local assembling based on a quadrature formula. We note that each node on \( \partial T \) is redundantly existent in the data structure and that we can easily account for jumps in the coefficient function when resolved by the macro-mesh elements \( T \).

Similar scaling techniques have been used in [39] for a generalized Stokes problem from geodynamics with coupled velocity components. However, for vectorial equations such a simple scaling does not asymptotically result in a physically correct solution. For the computation of integrals on triangles containing derivatives in the form of (2.2), cubature formulas of the form (2.3b) in combination with Euler–MacLaurin-type asymptotic expansions have been applied [31, Table 1].
Remark 2.1. At first glance the definition (2.3b) might not be more attractive than (2.4) regarding computational cost. In a matrix-free approach, however, where we have to reassemble the entries in each matrix call, (2.3b) turns out to be much more favorable. In order to see this, we have to recall that we work with hybrid hierarchical meshes. This means that for each inner node \( i \) in \( T \), we find the same entries in the sense that

\[
\int_T (D_m \phi_i, D_m \phi_j) \, dx = \int_T (D_m \phi_i, D_m \phi_{x_i + \delta x}) \, dx.
\]

Here we have identified the index notation with the vertex notation, and the vertex \( x_l \) is obtained from the vertex \( x_i \) by a shift of \( \delta x \), i.e., \( x_l = x_i + \delta x \). Consequently, the values of \( \int_T (D_m \phi_i, D_m \phi_j) \, dx \) do not have to be recomputed but can be efficiently stored.

For simplicity of notation, we shall restrict ourselves in the following to the case of the Darcy equation with a scalar uniformly positive definite permeability; i.e., let \( M = 1 \) and drop the index \( m \). However, the proofs in section 3 can be generalized to conceptually the same type of results as for \( M > 1 \). In subsection 7.3, we also show numerical results for \( M = 6 \) in 3D.

2.2. Stencil structure. We exploit the hierarchical grid structure to save a significant amount of memory compared to classical sparse matrix formats. Any direct neighbor \( x_j \in \mathcal{N}_T(x_i) \) can be described through a direction vector \( w_j \) such that \( x_j = x_i + w_j \). The regularity of the grid in the interior of a macro-element \( T \) implies that these vectors remain the same when we move from one node to another node. Additionally, for each neighbor \( x_j \in \mathcal{N}_T(x_i) \) \( \setminus \{x_i\} \) there is a mirrored neighbor \( x'_j \) of \( x_i \) reachable by \( w_j = -w_j' \); see Figure 2.1.

Let \( n_i = |\mathcal{N}_T(x_i)| \) denote the stencil size at mesh node \( x_i \). We define the stencil \( \mathbf{s}^T_{x_i} \in \mathbb{R}^{n_i} \) associated to the \( i \)th mesh node \( x_i \) restricted on \( T \) as

\[
(\mathbf{s}^T_{x_i})_j := \int_T (\nabla \phi_{x_i + w_j}, \nabla \phi_{x_i}) \, dx.
\]

The symmetry of the bilinear form yields

\[
(\mathbf{s}^T_{x_i})_j = (\mathbf{s}^T_{x_i + w_j})'_j.
\]

Fig. 2.1. From left to right: exemplary local indices \( j \) and their corresponding direction vectors of a 15 point stencil in 3D; six elements attached to one edge; four elements attached to one edge.
We recall that for each mesh node \( x_i \) we have \( n_i \leq 7 \) in 2D and \( n_i \leq 15 \) in 3D. Out of these entries only 3 in 2D and 7 in 3D have to be computed since the remaining ones follow from symmetry arguments and the observation that \( \sum_j (s^T_{x_i})_j = 0 \).

Due to the hierarchical hybrid grid structure, two stencils, \( s^T_{x_{i1}} \) and \( s^T_{x_{i2}} \), are exactly the same if \( x_{i1} \) and \( x_{i2} \) are two nodes belonging to the same primitive; i.e., we find only 15 different kinds of stencils per macro-element in 3D, one for each of its 15 primitives (4 vertices, 6 edges, 4 faces, 1 volume), and 7 in 2D. This observation allows for an extremely fast and memory-efficient on-the-fly (re)assembly of the entries of the stiffness matrix in stencil form. For each node \( x_i \) in the data structure, we save the nodal values of the coefficient function \( k \). With these considerations in mind, the bilinear form (2.3b) can be evaluated very efficiently and requires only a suitable scaling of the reference entries; see section 6 for detailed cost considerations.

### 3. Variational crime framework and a priori analysis
In order to obtain order \( h \) and \( h^2 \) a priori estimates of the modified finite element approximation in the \( H^1 \)- and \( L^2 \)-norms, respectively, we analyze the discrete bilinear form. From now on, we assume that \( k|_T \in W^{2,\infty}(T) \) for each \( T \in \mathcal{T}_h \). Moreover, we denote by \( \| \cdot \|_0 \) the \( L^2 \)-norm on \( \Omega \) and \( \| \cdot \|_{\infty} := \sup_{T \in \mathcal{T}_h} \| \cdot \|_{\infty,T} \) defines a broken \( L^\infty \)-norm. We recall that the coefficient function is only assumed to be elementwise smooth with respect to the macro-triangulation. Existence and uniqueness of a finite element solution \( u_h \in V_h \cap V_0 \) of

\[
a_h(u_h,v_h) = f(v_h), \quad v_h \in V_h \cap V_0,
\]

is given, provided that the following assumption (A1) holds true:

(A1) \( a_h(\cdot, \cdot) \) is uniformly coercive on \( V_h \cap V_0 \).

(A2) \( |a(v_h,w_h) - a_h(v_h,w_h)| \lesssim h \| \nabla k \|_\infty \| \nabla v_h \|_0 \| \nabla w_h \|_0, \quad v_h, w_h \in V_h \).

Here and in the following, the notation \( \lesssim \) is used as abbreviation for \( \leq C \), where \( C < \infty \) is independent of the mesh-size \( h \). The assumption (A2), if combined with Strang’s first lemma, yields that the finite element solution results in \( \mathcal{O}(h) \) a priori estimates with respect to the \( H^1 \)-norm; see, e.g., [11, 34]. We note that for \( h \) small enough, the uniform coercivity (A1) follows from the consistency assumption (A2), since for \( v_h \in V_h \)

\[
a_h(v_h,v_h) \geq a(v_h,v_h) - |a_h(v_h,v_h) - a(v_h,v_h)| \geq C(1 - ch)\| \nabla v_h \|_0^2.
\]

**Remark 3.1.** As is commonly done in the finite element analysis in unweighted Sobolev norms, we allow the generic constant \( C \) to be dependent on the global contrast of \( k \) defined by \( \sup_{\Omega} k / \inf_{\Omega} k \). Numerical results, however, show that the resulting bounds may be overly pessimistic for coefficients with large global variations. In [33] and the references therein, methods to improve the bounds in this case are presented. The examples show that the bounds may be improved significantly for coefficients with a global contrast in the magnitude of about \( 10^5 \). We are mainly interested in showing alternative assembly techniques to the standard finite element method and in comparing them to the well-established approaches in standard norms. Moreover, in our modification only the local variation of the coefficient \( k \) is important; therefore we shall not work out these subtleties here.

**3.1. Abstract framework for \( L^2 \)-norm estimates.** Since (A2) does not automatically guarantee optimal order \( L^2 \)-estimates, we employ duality arguments. To get a better feel for the required accuracy of \( a_h(\cdot, \cdot) \), we briefly recall the basic steps occurring in the proof of the upper bound. As is standard, we assume \( H^2 \)-regularity
of the primal and the dual problem. Restricting ourselves to the case of homogeneous Dirichlet boundaries, the dual PDE and boundary operators coincide with the primal ones. Let us denote by $P_h u$ the standard Galerkin approximation of $u$, i.e., the finite element solution obtained as the solution of a discrete problem using the bilinear form $a(\cdot, \cdot)$. It is well known that under the given assumptions $\|u - P_h u\|_0 = O(h^2)$. Now, to obtain an $L^2$-estimate for $u_h$, we consider the dual problem with $u_h - P_h u$ on the right-hand side. Let $w \in V_0$ be the solution of $a(v, w) = (u_h - P_h u, v)_0$ for $v \in V_0$. Due to the standard Galerkin orthogonality, we obtain

$$
(3.1) \quad \|u_h - P_h u\|_0^2 = a(u_h - P_h u, w) = a(u_h - P_h u, P_h w) = a(u_h, P_h w) - a_h(u_h, P_h w).
$$

This straightforward consideration shows us that compared to (A2), we need to make stronger assumptions on the mesh-dependent bilinear form $a_h(\cdot, \cdot)$. We define (A3) by

$$
(A3) \quad |a(v_h, w_h) - a_h(v_h, w_h)| \lesssim h^2 \|Hk\|_\infty \|\nabla v_h\|_0 \|\nabla w_h\|_0 + h \|\nabla k\|_\infty \|\nabla v_h\|_{0,S_h} \|\nabla w_h\|_{0,S_h},
$$

where $Hk$ denotes the Hessian of $k$ and $S_h := \cup_{T \in \mathcal{T}_h} S_h(T)$ with $S_h(T) := \{t \in \mathcal{T}_h; \partial T \cap \partial T' \neq \emptyset\}$; see Figure 3.1 for a 2D illustration. The seminorm $\|\cdot\|_{0,S_h}$ stands for the $L^2$-norm restricted to $S_h$.

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{fig3_1.png}
\caption{Elements in $S_h(T)$ (left) and $S_h$ (right) for $d = 2$.}
\end{figure}

**Lemma 3.2.** Let the problem under consideration be $H^2$-regular, let $h$ be sufficiently small, and let (A3) be satisfied. Then we obtain a unique solution and optimal order convergence in the $H^1$- and $L^2$-norms, i.e.,

$$
(3.2) \quad \|u_h - u\|_0 + h \|\nabla (u_h - u)\|_0 \lesssim h^2 (\|Hu\|_0 + \|Hk\|_\infty \|\nabla u\|_0 + \|\nabla k\|_\infty (\|\nabla u\|_0 + \|Hu\|_0)).
$$

**Proof.** Given that $h$ is small enough, (A1) follows from (A3). In terms of (3.1) and (A3), we get

$$
(3.3) \quad \|u_h - P_h u\|_0^2 \lesssim h^2 \|Hk\|_\infty \|\nabla u_h\|_0 \|\nabla P_h w\|_0 + h \|\nabla k\|_\infty \|\nabla u_h\|_{0,S_h} \|\nabla P_h w\|_{0,S_h}.
$$

The stability of the standard conforming Galerkin formulation yields $\|\nabla P_h w\|_0 \lesssim \|u_h - P_h u\|_0$. By definition (2.3), we find that the discrete bilinear form $a_h(\cdot, \cdot)$ is uniformly continuous for a coefficient function in $L^\infty(\Omega)$, and thus $\|\nabla u_h\|_0 \lesssim \|\nabla u\|_0$. To bound the two terms involving $\|\nabla \cdot\|_{0,S_h}$, we use the 1D Sobolev embedding $H^1((0,1)) \subset L^\infty((0,1))$ [36]. More precisely, for an element in $H^s(\Omega)$, $s > 0.5$, we have $\|v\|_{0,S_h} \lesssim \|v\|_{H^s(\Omega)}$; see [27]. Here we use $s = 1$, and in terms of the $H^2$-regularity assumption we obtain

$$
\|\nabla u_h\|_{0,S_h} \leq \|\nabla (u - u_h)\|_{0,S_h} + \|\nabla u\|_{0,S_h} \lesssim \|\nabla (u - u_h)\|_{0,S_h} + \sqrt{h}(\|\nabla u\|_0 + \|Hu\|_0),
$$

$$
\|\nabla P_h w\|_{0,S_h} \leq \|\nabla (w - P_h w)\|_{0,S_h} + \sqrt{h}\|u_h - P_h u\|_0 \lesssim \sqrt{h}\|u_h - P_h u\|_0.
$$
Using the above estimates, (3.3) reduces to
\[ \|u_h - P_h u\|_0 \lesssim h^2 (\|Hk\|_\infty \|\nabla u\|_0 + \|\nabla k\|_\infty (\|\nabla u\|_0 + \|Hu\|_0)) + h^{3/2} \|\nabla k\|_\infty \|\nabla (u - u_h)\|_0. \]

Applying the triangle inequality and using the approximation properties of the Galerkin finite element solution results in the extra term \( h^2 \|H u\|_0 \) in the upper bound for \( \|u - u_h\|_0 \). The bound for \( h\|\nabla(u_h - u)\|_0 \) follows by a standard inverse estimate for elements in \( V_h \) and the best approximation property of \( V_h \cap V_0 \). Since for \( h \) small enough it holds that \( c_1 \sqrt{h} \|\nabla k\|_\infty \leq 1/2 \), where \( c_1 < \infty \) is a suitably fixed positive constant, the upper bound (3.2) follows.

**3.2. Verification of the assumptions.** It is well known [11] that assumptions (A1)–(A3) are satisfied for the bilinear form
\[
(3.4) \quad \tilde{a}_h(u_h, v_h) := \sum_{t \in T_h} \frac{|t|}{d+1} \sum_{i=1}^{d+1} k_i(x_i^t) \nabla u_h|_t(x_i^t) \cdot \nabla v_h|_t(x_i^t) = \sum_{t \in T_h} \tilde{k}_i \int_t \nabla u_h \cdot \nabla v_h \, dx.
\]

Here \( x_i^t \) denotes the vertices of the \( d \)-dimensional simplex \( t \); i.e., we approximate the integral by a nodal quadrature rule and \( \tilde{k}_i := (\sum_{i=1}^{d+1} k_i(x_i^t))/(d+1) \). Thus, to verify the assumptions also for \( a_h(\cdot, \cdot) \), it is sufficient to consider \( a_h(v_h, w_h) - \tilde{a}_h(v_h, w_h) \) in more detail with \( a_h(\cdot, \cdot) \) given by (2.3). Let
\[
\hat{A}_t := \begin{pmatrix} a_{1,1}^t & a_{1,2}^t & \ldots & a_{1,d+1}^t \\ a_{1,2}^t & a_{2,2}^t & \ldots & a_{2,d+1}^t \\ \vdots & \vdots & \ddots & \vdots \\ a_{1,d+1}^t & a_{2,d+1}^t & \ldots & a_{d+1,d+1}^t \end{pmatrix}, \quad K_t := \begin{pmatrix} k_{1,1}^t & k_{1,2}^t & \ldots & k_{1,d+1}^t \\ k_{1,2}^t & k_{2,2}^t & \ldots & k_{2,d+1}^t \\ \vdots & \vdots & \ddots & \vdots \\ k_{1,d+1}^t & k_{2,d+1}^t & \ldots & k_{d+1,d+1}^t \end{pmatrix}
\]

be the local stiffness matrix associated with the nodal basis functions \( \phi_i^t \), i.e., \( a^t_{i,j} := \int_t \nabla \phi_i^t \cdot \nabla \phi_j^t \, dx \), and the local coefficient function with \( k^t_{i,j} := \frac{1}{2}(k_i(x_i^t) + k_j(x_j^t)) \), \( i \neq j \). The diagonal entries of \( K_t \) are defined differently as
\[
(3.5) \quad k^t_{i,i} := \frac{1}{a^t_{i,i}} \sum_{j \neq i} k^t_{i,j} a^t_{i,j}.
\]

We introduce the componentwise Hadamard product between two matrices as \((B \circ C)_{ij} := B_{ij} C_{ij}\) and define the rank-one matrix \( \tilde{K}_t \) by \( (\tilde{K}_t)_{ij} := \tilde{k}_i \).

Due to the symmetry of \( \hat{A}_t \) and the fact that the row sum of \( \hat{A}_t \) is equal to zero, we can rewrite the discrete bilinear forms. With \( v_h, w_h \in V_h \), we associate locally elements \( v_t, w_t \in \mathbb{R}^{d+1} \) with \( (v_t)_i := v_h(x_i^t), (w_t)_i := w_h(x_i^t) \). We recall that if \( v_h, w_h \in V_h \cap V_0 \) and \( x_i^t \) is a boundary node, then \( v_h(x_i^t) = 0 = w_h(x_i^t) \).

**Lemma 3.3.** The bilinear forms given by (2.3) and (3.4) have the algebraic form
\[
(3.6) \quad a_h(v_h, w_h) = \frac{1}{2} \sum_{t \in T_h} \sum_{i,j=1}^{d+1} ((v_t)_i - (v_t)_j)(K_t \circ \hat{A}_t)_{ij} ((w_t)_j - (w_t)_i), \quad v_h, w_h \in V_h,
\]
\[
(3.7) \quad \tilde{a}_h(v_h, w_h) = \frac{1}{2} \sum_{t \in T_h} \sum_{i,j=1}^{d+1} ((v_t)_i - (v_t)_j)(\tilde{K}_t \circ \hat{A}_t)_{ij} ((w_t)_j - (w_t)_i), \quad v_h, w_h \in V_h.
\]
Proof. We note that (3.5) yields that the row sum of $K_t \circ \hat{A}_t$ is equal to zero. Moreover, $K_t \circ \hat{A}_t$ is by construction symmetric. Introducing for $T \in \mathcal{T}_h$ the set $\mathcal{T}^{i,j:T}_h$ of all elements $T \ni t \in \mathcal{T}_h$ sharing the global nodes $i$ and $j$, we identify by $i_t$ the local index of the node $i$ associated with the element $t$, and by $j_t$ the local index of the global node $j$. Then the standard local-to-global assembling process yields that the right-hand side in (3.6) reads

$$\frac{1}{2} \sum_{T \in \mathcal{T}_h} \sum_{i,j} (\nu_i - \nu_j)(\chi_j - \chi_i) \sum_{t \in \mathcal{T}^{i,j:T}_h} k^t_{i,j} a^t_{i,j},$$

Comparing this result with the definition (2.3b), we find equality since the coefficient function is assumed to be smooth within each $T$. The proof of (3.7) follows with exactly the same arguments as the one for (3.6).

Although the proof of the previous lemma is straightforward, the implication of it for large scale simulations cannot be underestimated. In 2D, the number of different edge types per macro-element is three, while in 3D it is seven, assuming uniform refinement. All edges in 3D in the interior of a macro-element $T$ share only four or six elements; see Figure 2.1. We have three edge types that have four elements attached to them and four edge types with six adjacent elements.

The algebraic formulations (3.7) and (3.6) allow us to estimate the effects of the variational crime introduced by the stencil scaling approach.

Lemma 3.4. Assumptions (A2) and (A3) hold true.

Proof. The required $O(h)$ bound for (A2) is straightforward and also holds true for any unstructured mesh refinement strategy. Recalling that $K_t = \hat{K}_t$ if the coefficient function $k$ restricted to $t$ is a constant, (3.7) and (3.6) yield

$$|a_h(v_h, w_h) - \tilde{a}_h(v_h, w_h)| = \frac{1}{2} \sum_{t \in \mathcal{T}_h} \sum_{i,j=1}^{d+1} ((\nu_i - \nu_j)(\chi_j - \chi_i) (K_t - \hat{K}_t) \circ \hat{A}_t)_{ij} (w_j - w_i)_{ij},$$

$$\lesssim h^{2-d} \max_{t \in \mathcal{T}_h} \left( \lambda_{\max}(K_t - \hat{K}_t) \lambda_{\max}(\hat{A}_t) \right) \|\nabla v_h\|_0 \|\nabla w_h\|_0$$

$$\lesssim \max_{t \in \mathcal{T}_h} \left( \lambda_{\max}(K_t - \hat{K}_t) \right) \|\nabla v_h\|_0 \|\nabla w_h\|_0 \lesssim h^{2-d} \|\nabla v_h\|_0 \|\nabla w_h\|_0,$$

where $\lambda_{\max}()$ denotes the maximal eigenvalue of its argument.

To show (A3), we have to exploit the structure of the mesh $\mathcal{T}_h$. Let $\mathcal{E}_h^{e:T}$ be the set of all edges in $\hat{T}$, and $\mathcal{T}_h^{e:T}$ the subset of elements $t \subset T$ which share the edge $e$ having the two global nodes $i$ and $j$ as endpoints. As before, we identify the local indices of these endpoints by $i_t$ and $j_t$. We note that the two sets $\mathcal{E}_h^{e:T}$ and $\mathcal{E}_h^{e:T}$, $T \neq \hat{T}$, are not necessarily disjoint. Observing that each element $t \subset T$ is exactly contained in $\frac{1}{2}(d+1)$ elements of $\mathcal{T}_h^{e:T}$, we find

$$a_h(v_h, w_h) - \tilde{a}_h(v_h, w_h) = \frac{1}{d(d+1)} \sum_{t \in \mathcal{T}_h} \sum_{e \in \mathcal{E}_h^{e:T}} \sum_{t \in \mathcal{T}_h^{e:T}} ((\nu_i)_{i_t} - (\nu_j)_{j_t}) (K_t - \hat{K}_t) \circ \hat{A}_t_{ij} ((w_j)_{j_t} - (w_i)_{i_t}),$$

and thus it is sufficient to focus on the contributions resulting from $t \in (\mathcal{T}_h^{e:T})$. We consider two cases separately: First, we consider the case that the edge $e$ is part of
\( \partial T \) for at least one \( T \); then we directly find the upper bound
\[
h \| \nabla k \|_{L^\infty(T)} \| \nabla v_h \|_T \| \nabla w_h \|_T.
\]

Second, we consider the case that \( e \) is in the interior of one \( T \). Then for each element \( t \in T_h^{e,T} \) there exists exactly one \( t^m \in T_h^{e,T} \) such that \( t^m \) is obtained by point reflection at the midpoint of the edge \( e \); see Figure 3.2. In the following, we exploit that the midpoint of the edge is the barycenter of \( t \cup t^m \). Here the local indices \( i_t \) and \( j_t \) are associated with the global nodes \( i \) and \( j \), respectively. Without loss of generality, we assume a local renumbering such that \( i_t = 1 \), \( j_t = 2 \) and that \( x_{i_t}^{t_m} \) is the point reflected vertex of \( x_i \); see also Figure 3.2.

Let us next focus on
\[
E_{ij}^t(v_h, w_h) := \left| ((v_1)_1 - (v_2)_2)((K_1 - \tilde{K}_t) \circ \tilde{A}_t)_1,2((w_1)_2 - (w_2)_1)
+((v_1)_2 - (v_2)_1)((K_2 - \tilde{K}_t) \circ A_t)_2,1((w_1)_1 - (w_2)_2) \right|.
\]

Exploiting the fact that \( (\tilde{A}_t)_1,2 = (\tilde{A}_t)_2,1 \), we can bound \( E_{ij}^t(\cdot, \cdot) \) by the local \( H^1 \)-seminorms:
\[
E_{ij}^t(v_h, w_h) \lesssim \| \nabla v_h \|_{0,t;L^\infty} \| \nabla w_h \|_{0,t;L^\infty} \left( |k|_T(x_i) + k|_T(x_j) \right) - \frac{1}{d+1} \sum_{i=1}^{d+1} |(k|_T(x_i^1) + k|_T(x_i^m))|.
\]

A Taylor expansion of \( k \) in \( \frac{1}{2}(x_1^1 + x_1^m) = \frac{1}{2}(x_i + x_j) \) guarantees that the terms of zeroth and first order cancel out and only second order derivatives of \( k \) scaled with \( h^2 \) remain, i.e.,
\[
E_{ij}^t(v_h, w_h) \lesssim h^2 \| Hk \|_{L^\infty(t;L^\infty)} \| \nabla v_h \|_{0,t;L^\infty} \| \nabla w_h \|_{0,t;L^\infty}.
\]

Then the summation over all macro-elements, all edges, and all elements in the subsets \( T_h^{e,T} \) in combination with a finite covering argument [16] yields the upper bound of (A3).
4. Guaranteed uniform coercivity. While for our hierarchical hybrid mesh framework the assumptions (A2) and (A3) are satisfied, and thus asymptotically, i.e., for $h$ sufficiently small, (A1) is also satisfied, (A1) is not necessarily guaranteed for any given mesh $\mathcal{T}_h$.

**Lemma 4.1.** If the matrix representation of the discrete Laplace operator is an $M$-matrix, then the scaled bilinear form $a_h(\cdot, \cdot)$ is positive semidefinite on $V_h \times V_h$ for $k$ globally smooth.

**Proof.** Let $\hat{A}_h$ be the matrix representation of the discrete Laplace operator; then the $M$-matrix property guarantees that $(\hat{A}_h)_{ij} \leq 0$ for $i \neq j$. Taking into account that $\sum_{T \in \mathcal{T}_h} \hat{s}_{ij}^T = (\hat{A}_h)_{ij}$, definition (2.3) for the special case $M = 1$ and $k$ globally smooth yields

$$a_h(v_h, v_h) = -\frac{1}{4} \sum_{i,j} (k(x_i) + k(x_j))(v_i - v_j)^2 (\hat{A}_h)_{ij} \geq 0.$$

**Remark 4.2.** In 2D it is well known [12] that if all elements of the macro-mesh have no obtuse angle, then $A_h$ is an $M$-matrix and we are in the setting of Lemma 4.1.

4.1. Preasymptotic modification in 2D based on (A2). Here we work out the technical details of a modification in 2D that guarantees uniform ellipticity assuming that at least one macro-element $T$ has an obtuse angle. Our modification yields a linear condition on the local mesh-size depending on the discrete gradient of $k$. It only applies to selected stencil directions. In 2D our 7-point stencil associated with an interior fine grid node has exactly two positive off-center entries if the associated macro-element has an obtuse angle. We call the edges associated with a positive reference stencil entry to be of gray type. With each macro-element $T$, we associate the reference stiffness matrix $\hat{A}_T$. Without loss of generality, we assume that the local enumeration is done in such a way that the largest interior angle of the macro-element $T$ is located at the local node 3; i.e., if $T$ has an obtuse angle, then $a_{1,2}^T > 0$, $a_{1,3}^T < 0$, and $a_{2,3}^T < 0$, and otherwise $a_{1,2}^T \leq 0$, $1 \leq i < j \leq 3$. By $\lambda_{\min}^T$ we denote the smallest nondegenerated eigenvalue of the generalized eigenvalue problem

$$\hat{A}_T x := \begin{pmatrix} a_{1,1}^T & a_{1,2}^T & a_{1,3}^T \\ a_{2,1}^T & a_{2,2}^T & a_{2,3}^T \\ a_{3,1}^T & a_{3,2}^T & a_{3,3}^T \end{pmatrix} x = \lambda \begin{pmatrix} 2 & -1 & -1 \\ -1 & 2 & -1 \\ -1 & -1 & 2 \end{pmatrix} x, \quad x \in \mathbb{R}^3.$$

We note that both matrices in the eigenvalue problem are symmetric, are positive semidefinite, and have the same 1D kernel and thus $\lambda_{\min}^T > 0$.

Let $e$ be a gray type edge. For each such edge $e$, we possibly adapt our approach locally. We denote by $\omega_{e,T}$ the element patch of all elements $t \in \mathcal{T}_h$, such that $t \subset T$ and $e \subset \partial t$. Then we define

$$k_{e,\min} := \min_{e \in E_h^e} k_{\tilde{e}}, \quad k_{\tilde{e}} := \frac{1}{2}(k_{|_{T(x_1^e)}} + k_{|_{T(x_2^e)}}),$$

where $E_h^e$ is the set of all edges being in $\omega_{e,T}$, and $x_1^e$ and $x_2^e$ are the two endpoints of $\tilde{e}$. In the preasymptotic regime, i.e., if

$$(MA2) \quad (k_e - k_{e,\min}) a_{1,2}^T > k_{e,\min} \lambda_{\min}^T,$$
we replace the scaling factor \( k_e = \frac{1}{2}(k|_{T}(x_1^e) + k|_{T}(x_2^e)) \) in definition (2.3) by

\[
k^\text{mod}_{e} := k^\text{min}_{e} \left( 1 + \frac{\lambda_{T}^\text{min}}{a_{1,2}^0} \right) .
\]

Then it is obvious that \( k^\text{min}_{e} < k^\text{mod}_{e} < k_e \). We note that \( 2a_{1,2}^T \) is the value of the 7-point stencil associated with a gray edge and thus trivial to access.

**Lemma 4.3.** Let the bilinear form be modified according to (MA2); then it is uniformly positive definite on \( V_h \cap V_0 \times V_h \cap V_0 \) for all simplicial hierarchical meshes.

**Proof.** As it holds true for standard bilinear forms, our modified one can also be decomposed into element contributions. The local stiffness \( A_t \) matrix for \( t \subset T \) associated with (2.3) reads \( A_t = K_t \circ \tilde{A}_t = K_t \circ \hat{A}_t \) and can be rewritten in terms of \( k^t_{\text{min}} := \min_{1 \leq i < j \leq 3} k^t_{i,j} \) and \( \delta k^t_{i,j} := k^t_{i,j} - k^t_{\text{min}} \) as

\[
A_t = k^t_{\text{min}} \hat{A}_T + \delta k^t_{1,2} \begin{pmatrix}
-a_{1,2}^T & a_{1,2}^T & 0 \\
0 & -a_{1,2}^T & 0 \\
0 & 0 & 0
\end{pmatrix}
+ \delta k^t_{2,3} \begin{pmatrix}
0 & 0 & 0 \\
0 & -a_{2,3}^T & a_{2,3}^T \\
0 & a_{2,3}^T & -a_{2,3}^T
\end{pmatrix},
\]

(4.1)

where we use a consistent local node enumeration. We note that each \( \delta k^t_{i,j} \geq 0 \). Now we consider two cases separately.

First, let \( t \subset T \), and let \( T \) be a macro-element having no obtuse angle; then we find that all four matrices on the right-hand side of (4.1) are positive semidefinite. Thus, we obtain \( A_t \geq k^t_{\text{min}} \hat{A}_T \) and no modification is required.

Second, let \( t \subset T \), and let \( T \) be a macro-element having one obtuse angle. Then the second matrix on the right-hand side of (4.1) is negative semidefinite, while the three other ones are positive semidefinite. Now we find

\[
x^T A_t x \geq k^t_{\text{min}} x^T \hat{A}_T x + x^T \delta k^t_{1,2} \begin{pmatrix}
-a_{1,2}^T & a_{1,2}^T & 0 \\
0 & -a_{1,2}^T & 0 \\
0 & 0 & 0
\end{pmatrix} x
\]

\[
\geq \lambda_{T}^T k^t_{\text{min}} ((x_1 - x_2)^2 + (x_2 - x_3)^2) + \lambda_{T}^T k^t_{\text{min}} - a_{1,2}^T \delta k^t_{1,2} (x_1 - x_2)^2
\]

\[
\geq \lambda_{T}^T k^t_{\text{min}} ((x_1 - x_3)^2 + (x_2 - x_3)^2) + \lambda_{T}^T k^t_{\text{min}} - a_{1,2}^T \delta k^t_{1,2} (x_1 - x_2)^2,
\]

where \( e \) is the edge associated with the two local nodes 1 and 2. Provided (MA2) is not satisfied, we have \( x^T A_t x \geq \lambda_{T}^T k^t_{\text{min}} ((x_1 - x_3)^2 + (x_2 - x_3)^2) \geq \delta k^t_{1,2} \) with \( k^t_{\text{mod}} \). With this modification it is now obvious that the newly defined bilinear form is positive semidefinite on \( V_h \times V_h \) and moreover positive definite on \( V_h \cap V_0 \times V_h \cap V_0 \). The coercivity constant depends only on the shape regularity of the macro-mesh, \( \min k \), and the Poincaré–Friedrichs constant.

**Remark 4.4.** From the proof it is obvious that any other positive scaling factor less than or equal to \( k^t_{\text{mod}} \) also preserves the uniform ellipticity.

**Remark 4.5.** We can replace in the modification criterion the local condition (MA2) by

\[
l_T 2^{-(e+1)} \| \nabla k \|_{L^\infty(\omega_e; T)} a_{1,2}^T \geq \inf_{x \in \omega_e; T} k(x) \lambda_{T}^t \]

where
where $T_h$ is obtained by $\ell$ uniform refinement steps from $T_H$, and $l_T$ is the length of the second longest edge in $T$. Both these criteria allow a local marking of gray-type edges which have to be modified. To avoid computation of $k_{e\text{min}}$ each time it is needed, and thus further reducing computational cost, we set the scaling factor for all gray-type edges in a marked $T$ to

\[(4.2) \quad \frac{1}{2}(k_{\text{min}}(x_1^e) + k_{\text{min}}(x_2^e)), \quad k_{\text{min}}(x_i) := \min \{ k(x_j) \mid x_j \in \mathcal{N}_T(x_i) \},\]

where $\mathcal{N}_T(x_i)$ denotes the set of all mesh nodes that are connected to node $x_i$ via an edge and belonging to the macro-element $T$, including $x_i$ itself. The quantity $k_{\text{min}}$ can be precomputed for each node $x_i$ once at the beginning and stored as a node-based vector such as $k$ is. For nonlinear problems where $k$ depends on the solution itself, $k_{\text{min}}$ can be updated directly after the update of $k$.

The presented preasymptotic modification based on (A2) yields a condition on the local mesh-size and only affects edge types associated with a positive stencil entry. However, the proof of (A3) shows that a condition on the square of the mesh-size is basically sufficient to guarantee (A1). This observation allows us to design an alternative modification yielding a condition on the square of the local mesh-size and involving the Hessian of $k$, except for the elements which are in $S_h$. However, in contrast to the option discussed before, we possibly also have to alter entries which are associated with negative reference stencil entries, and therefore we do not discuss this case in detail. It is obvious that for piecewise smooth $k$ there exists an $\ell_0$ such that for all refinement levels $\ell \geq \ell_0$ no local modification has to be applied. This holds true for both types of modifications. Thus, all the a priori estimates also hold true for our modified versions. Since we are interested in piecewise moderate variations of $k$ and large scale computations, i.e., large $\ell$, we assume that we are already in the asymptotic regime, i.e., that no modification has to be applied for our 3D numerical test cases, and we do not work out the technical details for the modifications in 3D.

### 4.2. Numerical counterexample in the preasymptotic regime.

In the case where we are outside the setting of Lemma 4.1, it is easy to come up with an example where the scaled bilinear form is not positive semidefinite, even for a globally smooth $k$. For a given mesh-size $h$, one can always construct a $k$ with a variation large enough such that the scaled stiffness matrix has negative eigenvalues.

Here we consider a 2D setting on the unit square with an initial mesh which is not Delaunay; see the left part of Figure 4.1. For the coefficient function $k$, we use a sigmoid function defined as

\[k(x, y; m, \eta) = \frac{\eta}{1 + \exp(-m(y - x - 0.2))} + 1,\]

with $m = 50$, which yields a steep gradient. Further, we vary the magnitude of $k$ by setting $\eta \in \{1, 10, 100, 1000\}$.

Now, we assemble the global stiffness matrix and report in Table 4.1 for different choices of $\eta$ its minimal and maximal eigenvalues over a sequence of uniform refinement steps. We show the eigenvalues for our scaling approach (2.3) with and without modification (MA2) and for the standard nodal integration assembly (3.4).

We find that for $\eta \geq 100$ the stiffness matrix of the scaling approach has negative eigenvalues in the preasymptotic regime. In these cases three or four refinement steps are required, respectively, to enter the asymptotic regime. However, the positive definiteness of the matrix can be recovered on coarser resolutions if (MA2) is applied.
Application of (MA2): mesh with two critical macro-elements used for Table 4.1 (left) and mesh where all macro-elements have an obtuse angle (right).

Table 4.1

<table>
<thead>
<tr>
<th>Scaling approach</th>
<th>( \lambda_{\text{min}}^A )</th>
<th>( \lambda_{\text{max}}^A )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \eta ) ( \ell )</td>
<td>1 2 3 4 5</td>
<td>1 2 3 4 5</td>
</tr>
<tr>
<td>1</td>
<td>2.7 0.72 0.18 0.046 0.011 0.0029</td>
<td>9.1 13.3 15.1 17.4 19.8 21.3</td>
</tr>
<tr>
<td>10</td>
<td>2.8 0.89 0.24 0.063 0.016 0.0039</td>
<td>40.0 69.3 80.7 93.2 108 116</td>
</tr>
<tr>
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<td>353 633 739 853 985 1066</td>
</tr>
<tr>
<td>1000</td>
<td>-128 -148 -94.5 -3.0 0.019 0.0049</td>
<td>3486 6265 7317 8450 9759 10562</td>
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</tbody>
</table>

<table>
<thead>
<tr>
<th>Scaling approach modified with (MA2)</th>
<th>( \lambda_{\text{min}}^A )</th>
<th>( \lambda_{\text{max}}^A )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \eta ) ( \ell )</td>
<td>1 2 3 4 5</td>
<td>1 2 3 4 5</td>
</tr>
<tr>
<td>1</td>
<td>2.7 0.72 0.18 0.046 0.011 0.0029</td>
<td>9.1 13.3 15.1 17.4 19.8 21.3</td>
</tr>
<tr>
<td>10</td>
<td>3.8 1.1 0.26 0.063 0.016 0.0039</td>
<td>40.8 69.4 80.7 93.2 108 116</td>
</tr>
<tr>
<td>100</td>
<td>4.0 1.3 0.30 0.072 0.018 0.0045</td>
<td>364 634 739 853 985 1066</td>
</tr>
<tr>
<td>1000</td>
<td>4.4 1.3 0.33 0.078 0.019 0.0049</td>
<td>3598 6278 7321 8453 9759 10562</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Standard nodal integration</th>
<th>( \lambda_{\text{min}}^A )</th>
<th>( \lambda_{\text{max}}^A )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \eta ) ( \ell )</td>
<td>1 2 3 4 5</td>
<td>1 2 3 4 5</td>
</tr>
<tr>
<td>1</td>
<td>2.8 0.72 0.18 0.046 0.011 0.0029</td>
<td>9.0 12.8 14.7 17.1 19.7 21.2</td>
</tr>
<tr>
<td>10</td>
<td>5.7 1.3 0.27 0.065 0.016 0.0040</td>
<td>36.6 64.9 78.4 90.7 107 116</td>
</tr>
<tr>
<td>100</td>
<td>27.7 1.8 0.34 0.076 0.018 0.0045</td>
<td>316 590 717 829 977 1064</td>
</tr>
<tr>
<td>1000</td>
<td>247 1.9 0.38 0.084 0.020 0.0049</td>
<td>3116 5840 7104 8211 9682 10549</td>
</tr>
</tbody>
</table>

Asymptotically, the minimal or maximal eigenvalues of all three approaches tend to the same values.

In Figure 4.1, we illustrate the action of the modification and show how the region of elements where it has to be applied gets smaller and finally vanishes with increasing number of refinement steps. To further illustrate how the region that is affected by (MA2) changes, we show a second example where all initial elements have an obtuse angle. The underlying color bar represents the coefficient function \( k \) with \( \eta = 1000 \) for the first (left) example and \( \eta = 100 \) for the second (right). The location of the steepest gradient of \( k \) is marked by a dashed line. Note that the gradient of \( k \) is constant along lines parallel to the dashed one. For all edges that are modified by (MA2) the adjacent elements are shown. The color intensity refers to the refinement level and goes from bright (initial mesh) to dark.

4.3. The sign of the stencil entries in 3D. In contrast to the 2D setting, a macro-mesh with no obtuse angle does not yield that \( A_h \) is an \( M \)-matrix. Here
the uniform refinement rule yields that for each macro-element three subclasses of tetrahedra (gray, blue, green) exist. To each of these we associate one interior edge type (gray, blue, green) defined by not being parallel to any of the six edges of the respective tetrahedron type. Figure 4.2 shows the subclasses and as example the gray edge type. The coloring of the subclasses is up to now arbitrary. We always associate the gray color with the macro-element and call the associated interior edge a gray-type edge. The interior edges associated with the blue and green elements are called blue- and green-type edges, respectively. All other remaining edges are by notation red-type edges.

If the macro-element $T$ has no obtuse angle between two faces, then it follows from [26, 24] that the reference stencil entries, i.e., the entries associated with the Laplace operator, associated with gray-type edges have a positive sign; see Figure 4.3.
However, it can be shown by some simple geometrical considerations that not both, green- and blue-type edges, can have a positive sign. If one of these has a positive sign, we call this the blue- and the other one the green-type edge. Conversely, if both have a negative sign, the coloring is arbitrary. In case the macro-element $T$ has no obtuse angle, the sign associated with all red-type edges is automatically not positive. Thus, we find for such elements in our 15-point stencil either two or four positive off-diagonal entries. Then a modification similar to the one proposed for the 2D case can now be applied to the edges of gray and possibly blue type.

The situation can be drastically different if the macro-element $T$ has an obtuse angle between two faces. In that case, we find up to four edge directions which carry a positive sign in the stencil.

5. Reproduction property and primitive concept. As already mentioned, our goal is to reduce the cost of a WU and the run-times while preserving discretization errors that are qualitatively and quantitatively on the same level as for standard conforming finite elements. Here we focus on the 3D case, but similar results can be obtained for the 2D setting. The a priori bounds of Lemma 3.2 do not necessarily guarantee that for an affine coefficient function $k$ and affine solution $u$ the error is equal to zero. In the upper bound (3.2) a term of the form $\| \nabla k \|_\infty \| \nabla u \|_0$ remains. A closer look at the proof reveals that this nontrivial contribution can be traced back to the terms associated with nodes on the boundary of a macro-element. This observation motivates us to introduce a modification of our stencil scaling approach. As already mentioned, all nodes are grouped into primitives, and we have easy access to the elements of these primitives. Recall that $\tilde{a}_h(\cdot, \cdot)$ defined as in (3.4) is associated with the standard finite element approach with nodal quadrature. Let us by $W_V$, $W_E$, and $W_F$ denote the set of all nodes associated with the vertex, edge, and face primitives, respectively. Now we introduce a modified stencil scaling approach (cf. (2.3) and (3.4)):

\begin{equation}
(5.1) \quad a_{h,\mathcal{I}}(v_h, \phi_i) := \begin{cases} 
\tilde{a}_h(v_h, \phi_i), & i \in \mathcal{I}, \\
a_h(v_h, \phi_i), & i \notin \mathcal{I}.
\end{cases}
\end{equation}

Replacing $a_h(\cdot, \cdot)$ by $a_{h,\mathcal{I}}(\cdot, \cdot)$ with $\mathcal{I} \subset \mathcal{W} := W_V \cup W_E \cup W_F$ still yields that all node stencils associated with a node in a volume primitive are cheap to assemble. The number of node stencils which have to be more expensively assembled grow only at most with $4^d$, while the total number of nodes grows with $8^d$.

Remark 5.1. The modification (5.1) introduces an asymmetry in the definition of the stiffness matrix to which multigrid solvers are not sensitive. Moreover, the asymmetry tends asymptotically to zero with $O(h^2)$.

Lemma 5.2. Let $\mathcal{I} = \mathcal{W}$ or $\mathcal{I} = W_V \cup W_E$; then an affine solution can be reproduced if $k$ is affine, i.e., $u_h = u$.

Proof. For $i \notin \mathcal{W}$, associated with a node in the macro-element $T$ and $k$ affine, we have $k_{ij}(s^T_i) = (s^T_i)_j$, and thus the bilinear form $a_{h,\mathcal{I}}(\cdot, \cdot)$ is identical to $\tilde{a}_h(\cdot, \cdot)$ for $\mathcal{I} = \mathcal{W}$. Since then no variational crime occurs, we find $u_h = u$ for any affine solution $u$.

The case $\mathcal{I} = W_V \cup W_E$ is more involved. Here the bilinear forms $a_{h,\mathcal{I}}(\cdot, \cdot)$ and $\tilde{a}_h(\cdot, \cdot)$ are not identical. However, it can be shown that for any affine function $v_{aff}$ we find $a_{h,\mathcal{I}}(v_{aff}, \cdot) = \tilde{a}_h(v_{aff}, \cdot)$.
on $V_h \cap V_0$. To see this, we follow arguments similar to those in the proof of Lemma 3.4. This time we do not use a point reflected element $t^\infty$ but a shifted element $t^s$; see Figure 5.1. We note that $\nabla v_{\mathrm{aff}}$ is constant and

\begin{equation}
\frac{1}{2}(k(x_i) + k(x_i + w_j)) - \frac{1}{2}(k(x_i) + k(x_i + w_j')) = k(x_i) - k(x_i') .
\end{equation}

Here $x_i'$ and $x_i''$ stand for the barycenter of the elements $t$ and $t^s$, respectively. Two of the four vertices of element $t$ are the nodes $x_i$ and $x_i + w_j$. The shifted element $t^s$ is obtained from $t$ by a shift of $-w_j$.

This observation yields for a node $i \notin \mathcal{W}_V \cup \mathcal{W}_E$ that the sum over the nodes can be split into two parts,

\begin{align*}
a_{h,I}(v_{\mathrm{aff}}, \phi_i) &= a_h(v_{\mathrm{aff}}, \phi_i) = \frac{1}{2} \sum_{j \in \mathcal{W}} (k(x_j) + k(x_i))(v_{\mathrm{aff}}(x_i) - v_{\mathrm{aff}}(x_j))(\bar{s}_{x_j})_j \\
&\quad + \frac{1}{2} \sum_{j \notin \mathcal{W}} (k(x_j) + k(x_i))(v_{\mathrm{aff}}(x_i) - v_{\mathrm{aff}}(x_j))(\bar{s}_{x_j})_j .
\end{align*}

For the second sum on the right, we have already shown equality to the corresponding term in the bilinear form $\bar{a}_h(v_{\mathrm{aff}}, \phi_i)$. We recall that each node $j$ in $\mathcal{W}$ such that $x_i$ and $x_j$ form an edge has a point mirrored node $x'_j$, i.e., $x_i = 0.5(x_j + x_j')$. For the first term on the right, we find that for a node $j \in \mathcal{W}$ it holds that $(\bar{s}_{x_i})_j = (\bar{s}_{x_i'})_j$. In terms of $v_{\mathrm{aff}}(x_i + w_j) - v_{\mathrm{aff}}(x_i) = v_{\mathrm{aff}}(x_i) - v_{\mathrm{aff}}(x_i + w_j')$, the first summand on the right can thus be further simplified to

\begin{equation}
\frac{1}{2} \sum_{j=1}^{n_i} \left( \frac{1}{2}(k(x_i + w_j) + k(x_i)) - (k(x_i + w_j') + k(x_i)) \right)(v_{\mathrm{aff}}(x_i + w_j) - v_{\mathrm{aff}}(x_i))(\bar{s}_{x_j})_j .
\end{equation}

Together with (5.2), this yields the stated equality.

To illustrate Lemma 5.2, we consider a 3D example on the unit cube $\Omega = (0,1)^3$. We use as solution $u(x, y, z) = -7x + y + 3z$ and $k(x, y, z) = 2x + 3y + 5z + 1$ for the coefficient function. The standard finite element solution reproduces the exact solution up to machine precision. We test the influence of the stencil scaling approach on different

\begin{table}[h]
\centering
\begin{tabular}{|c|c|}
\hline
Used bilinear form to define FE approximation & Discretization error in discrete $L^2$-norm \\
\hline
$\bar{a}_h(\cdot, \cdot)$ standard FE & 3.85104e-15 \\
$a_{h,I}(\cdot, \cdot), I = \mathcal{W}$ & 2.10913e-15 \\
a_{h,I}(\cdot, \cdot), I = \mathcal{W}_V \cup \mathcal{W}_E & 1.75099e-15 \\
a_{h,I}(\cdot, \cdot), I = \mathcal{W}_V \cup \mathcal{W}_E & 6.82596e-04 \\
a_{h,I}(\cdot, \cdot), I = \mathcal{W}_V \cup \mathcal{W}_E & 9.19145e-04 \\
a_{h,I}(\cdot, \cdot), I = \mathcal{W}_V & 6.82032e-04 \\
a_{h,I}(\cdot, \cdot), I = \mathcal{W}_E & 9.17523e-07 \\
a_{h,I}(\cdot, \cdot)$ stencil scaling & 6.81942e-04 \\
\hline
\end{tabular}
\caption{Results for different choices of bilinear forms.}
\end{table}
sets of primitives. In the right of Figure 5.1, we report the discretization error in the discrete $L^2$-norm for different combinations. Note that the macro-triangulation of the unit cube consists of 12 tetrahedral elements and of one nonboundary macro-vertex at $(0.5, 0.5, 0.5)$. These considerations show that the choice $\mathcal{I} = \mathcal{W}_V \cup \mathcal{W}_E$ is quite attractive. The number of stencils which have to be expensively evaluated grows only with $2^\ell$, while we still can guarantee the reproduction property for an affine solution in the case of an affine coefficient function.

6. Cost of a work unit. The stencil scaling has been introduced as a means to reduce the cost of a WU and hence enable the design of less expensive and thus more efficient PDE solvers. We will first employ a cost metric based on operation count. While we are aware that real run-times will be influenced by many additional factors, including, e.g., the quality of the implementation, compiler settings, and various hardware details, the classic measure still provides useful insight. Another important aspect that we will address here is the question of memory accesses. A more technical hardware-aware performance analysis, as conducted in [18] to evaluate the efficiency on real computers, is beyond our current scope. Since asymptotically the contributions from the element primitives dominate the cost, we restrict ourselves to the study of stencils for nodes located in the interior of a macro-element $T$.

6.1. Cost for stencil scaling. Let $\tilde{s}_{ij}^T$ be the 15-point stencil associated with the Laplacian at an inner node $i$ of $T$ which is independent of the node location within $T$. Recall that the scaled stencil is given by

$$s_{ij}^T := \frac{1}{2} \left( \kappa(x_i) + \kappa(x_j) \right) \tilde{s}_{ij}^T \quad \forall j \in \mathcal{N}_T(i) \setminus \{i\}, \quad s_{ii}^T := - \sum_{j \in \mathcal{N}_T(i) \setminus \{i\}} s_{ij}^T,$$

where $\kappa(x_i)$ represents either the value of the parameter $k$ at node $x_i$ or a modified version of it as suggested in (4.2). Note that this definition is the direct translation of the approach described in section 2.1 from the bilinear form to the pure stencil. Assuming that the constant $1/2$ factor is incorporated into the stencil $\tilde{s}_{ij}^T$ at setup and taking into account the number of edges emanating from $x_i$, being 6 in 2D and 14 in 3D, we can compute the noncentral stencil entries with 12 and 28 operations, respectively. The computation of the central entry via the zero row sum property takes 5 and 13 additions, respectively. This is summarized in Table 6.1.

We next compare the cost for the stencil scaling approach to an on-the-fly computation based on classic FEM techniques which, however, exploits the advantages of hierarchical hybrid grids; see also [17].

Remark 6.1. The cost to approximate the stencil coefficients with the two-scale interpolation method of [2] amounts to $q$ operations per stencil entry when the polynomial degree is chosen as $q$. When the coefficients are locally smooth, $q = 2$ delivers good results in [2]. The cost of this method becomes $2 \times 7 = 14$ operations in 2D, and $2 \times 15 = 30$ operations in 3D, respectively. This is slightly cheaper than the stencil scaling of this paper, but it neglects the setup cost that is necessary to construct the polynomials and completely ignores the fact that for a fixed macro-mesh-size no asymptotic optimality can be achieved.

6.2. Cost of the on-the-fly computation for the classical FEM. Employing the bilinear form (3.4), we can write the stencil coefficients as

$$\tilde{s}_{ij}^T = \tilde{a}_h(\phi_j, \phi_i) = \sum_{t \in \mathcal{T}^T} \sum_{\ell = 1}^{d+1} \frac{k(x_{\ell}^t)}{(d+1)} \nabla\phi_j, \nabla\phi_i | t| = \sum_{t \in \mathcal{T}^T} (E^t)_{i,j} \sum_{\ell = 1}^{d+1} k(x_{\ell}^t),$$
where

\[(E^t)_{i,j} := \frac{1}{(d+1)} \int_t \nabla \phi_j \cdot \nabla \phi_i \, dx\]

denotes the local stiffness matrix including the volume averaging factor \(1/(d+1)\). The regularity of the mesh inside a macro-element implies that there exist only a fixed number of differently shaped elements \(t\). Thus, we can compute stencil entries on the fly from one (2D) or three (3D) precomputed stiffness matrices and the node-based coefficient values.\(^2\) In 2D, always six elements \(t\) are attached to an interior node. Summing nodal values of \(k\) for each element first would require 12 operations. Together with the fact that two elements are attached to each edge, computing all 6 noncentral stencil entries via (6.1) would then require a total of 30 operations. This number can be reduced by eliminating common subexpressions. We precompute the values

\[k(x_i) + k(j_2), \quad k(x_i) + k(j_4), \quad k(x_i) + k(j_6),\]

i.e., we sum \(k\) for the vertices at the ends of the edges marked as dashed in Figure 6.1. Then we obtain \(\sum_{t=1}^3 k(x_t)\) by adding the value of \(k\) at the third vertex of \(t\) to the precomputed expression. In this fashion, we can compute all six sums with 9 operations and obtain a total of 27 operations for the noncentral entries.

The situation in 3D is more complicated. This stems mainly from the fact that the number of elements sharing the edge from node \(i\) to node \(j\) is no longer a single value as in 2D. Instead we have in 3D that \(|\mathcal{T}_h^i,j| = 4\) for the gray-, blue-, and green-type edges that emanate from node \(i\), while \(|\mathcal{T}_h^i,j| = 6\) for the remaining 8 red-type edges. Thus, for the computation of the noncentral stencil entries we obtain the cost

\[6(4 + 3) + 8(6 + 5) + 40 = 170.\]

The summand 40 represents the operations required for summing the nodal \(k\) values employing again elimination of common subexpressions; see [17] for details. Assembling the central entry involves contributions from 6 elements in 2D and 24 in 3D. Thus, executing this via (6.1) cannot require less operations than using the row sum property.

\(^2\)Note that in our HHG implementation, we use 6 element matrices as this is advantageous with respect to local-to-global indexing; see [3] for details.
6.3. Cost of stencil application. The cost of the application of the assembled stencil is independent of the approach to compute the stencil. Using the first expression in

\[ s^T_i v = \sum_{j \in \mathcal{N}(i)} s^T_{ij} v(x_j) = \sum_{j \in \mathcal{N}(i) \setminus \{i\}} s^T_{ij}(v(x_j) - v(x_i)) \]

results in 13 operations in 2D and 29 in 3D. Exploiting the row sum property avoids computing the central entry, but increases the cost for the stencil application (cf. second expression), so that in total only a single operation is saved. Table 6.1 summarizes the results of this section. We observe that assembling the stencil and applying it once in the scaling approach requires in 2D only about \( \frac{2}{3} \) and in 3D about \( \frac{1}{3} \) of the operations compared to a classical on-the-fly variant.

<table>
<thead>
<tr>
<th>Approach</th>
<th>Dimension</th>
<th>Noncentral entries</th>
<th>Central entry</th>
<th>Assembly + Application</th>
</tr>
</thead>
<tbody>
<tr>
<td>stencil scaling</td>
<td>2D</td>
<td>6 add / 6 mult</td>
<td>5 add / 0 mult</td>
<td>17 add / 12 mult</td>
</tr>
<tr>
<td></td>
<td>3D</td>
<td>14 add / 14 mult</td>
<td>13 add / 0 mult</td>
<td>41 add / 28 mult</td>
</tr>
<tr>
<td>on-the-fly FEM</td>
<td>2D</td>
<td>15 add / 12 mult</td>
<td>5 add / 0 mult</td>
<td>26 add / 18 mult</td>
</tr>
<tr>
<td></td>
<td>3D</td>
<td>98 add / 72 mult</td>
<td>13 add / 0 mult</td>
<td>125 add / 86 mult</td>
</tr>
</tbody>
</table>

In 3D, the stencil scaling approach results in a total of 69 operations per node and is thus only 2.4 times more expensive than the constant-coefficient case treated in, e.g., [3]. The 69 operations save a factor of 3 compared to the 211 operations per node for the conventional, yet highly optimized, on-the-fly assembly and roughly an order of magnitude when compared with nonoptimized variants of on-the-fly-assembly techniques.

To which extent a factor of 3 savings in floating-point operations will be reflected in run-time (or other practical cost metrics, such as, e.g., energy consumption) depends on many details of the hardware and system software. For instance, we note that in the above algorithms, the addition and multiplication operations are not ideally balanced and do not always occur such that the fused multiply-add operations of a modern processor architecture can be used. Furthermore, advanced optimizing compilers will restructure the loops and will attempt to find a scheduling of the instruction stream that avoids dependencies. Thus, in effect, the number of operations executed on the processor may not be identical to the number of operations calculated from the abstract algorithm or the high-level source code. Nevertheless, though the operation count does not permit a precise prediction of the run-times, we will see in the following that our effort to reduce the number of operations pays off in terms of accelerated execution.

6.4. Memory accesses. One other key aspect governing the performance of any algorithm is the number of read and write operations it needs to perform and their pattern with respect to spatial and temporal locality. The influence of these properties results from the large disparity between peak floating-point performance of modern CPUs and the latency and bandwidth limitation of memory access. All modern architectures employ a hierarchy of caches [13, 21] that helps accelerate memory access but that also make an a priori prediction of run-times quite difficult.
As in the previous subsections, for the number of operations we will perform here a high-level analysis of memory traffic before presenting experimental results in section 7.4. As a baseline we are including in our consideration not only the on-the-fly and stencil scaling approaches, but also a stored stencil version. For the latter we assume that in a setup phase the stencil for each node was assembled, by whatever method, and then stored. For the stencil application the weights of the local stencil are then loaded from main memory and applied to the associated degrees of freedom. Since the stencil weights correspond to the nonzero matrix entries of the corresponding row, this would be the HHG analogue of performing a sparse matrix-vector multiplication with a matrix stored in a standard sparse storage format, such as, e.g., compressed row storage (CRS). Note, however, that due to the structuredness of the mesh inside a volume primitive, less organizational overhead and indirect memory accesses are required than for a sparse matrix format. Most importantly we neither need to store nor transfer over the memory system any information on the position of the nonzero matrix entries.

For this article we do not study the full details of algorithmic optimization for best usage of the memory subsystem. Instead, we are going to compare two idealized cases. These are the optimistic version, in which we assume perfect reuse of each data item loaded to the caches, and a pessimistic version, where there is no reuse at all. Any actual implementation will lie somewhere in between these two cases, with the closeness to one of them being determined by algorithmic properties and the quality of its implementation. Furthermore, we will only consider the 3D problem.

Common to all three approaches under consideration is that, in order to apply the local stencil and compute the residuum at a node, they need to load the DOFs at the node and its 14 neighbors and the value of the right-hand side at the node only. After the stencil application, the resulting nodal value must be written back. Thus, we are not going to inspect these parts of the stencil application and also neglect questions of write-back strategies for the caches.

With respect to the stencil weights the situation is, of course, a different one. Let us start with the stored stencil approach. We denote by $N$ the number of DOFs inside a single volume primitive. In 3D a scalar operator using our discretization is represented by a 15-point stencil. This structure is invariant inside the volume primitive due to the regular mesh structure. Hence, we obtain for the total number of data items to be loaded from memory $\mathcal{N}_{\text{stored}} = 15N$.

In the on-the-fly assembly we start by loading the 6 precomputed element matrices, which are in $\mathbb{R}^{4 \times 4}$ for the 3D case. Note that these are loaded only once when the first node of the volume primitive is treated and can stay in the L1 cache during the complete loop over the volume primitive as they only occupy 768 bytes. Assembling the local stencil from these matrices requires information on the coefficient function $k$. Assuming perfect reuse of the nodal $k$ information, i.e., each nodal value needs to be loaded only once, we obtain $\mathcal{N}_{\text{opt}} = 6 \times (4 \times 4) + N = 96 + N$. In the pessimistic case, where we assume absolutely no cache effects, we need to reload neighboring $k$ values each time we update another node. This gives us $15N$ load operations per node, resulting in $\mathcal{N}_{\text{pes}} = 96 + 15N$.

Finally we consider the stencil scaling approach. Here it is sufficient to load once the 14 noncentral weights of the reference stencil $\hat{s}$ for the volume primitive.\footnote{The current HHG implementation for technical reasons assembles the stencil from the precomputed element matrices also in this case, so the constant term in the memory access is the same 768 bytes as for the on-the-fly approach.}
Table 6.2
Cost of the three different approaches measured in total bytes loaded for computation of stencil weights, assuming the use of the IEEE binary64 data type.

<table>
<thead>
<tr>
<th>Approach</th>
<th>Optimistic</th>
<th>Pessimistic</th>
</tr>
</thead>
<tbody>
<tr>
<td>stored stencils</td>
<td>120N</td>
<td>120N</td>
</tr>
<tr>
<td>on-the-fly FEM</td>
<td>768 + 8N</td>
<td>768 + 120N</td>
</tr>
<tr>
<td>stencil scaling</td>
<td>112 + 8N</td>
<td>112 + 120N</td>
</tr>
</tbody>
</table>

These are 112 bytes, and the values can, as in the on-the-fly approach, remain in the L1 cache. The noncentral weights are then scaled depending on the neighboring $k$ values, while its central weight is derived using the zero sum property. This gives us $N_{scale}^{opt} = 14 + N$ and $N_{scale}^{pes} = 14 + 15N$.

Table 6.2 sums up our results. Comparing the entries for the stored stencil approach to the pessimistic bounds for the other two approaches, we find the same prefactors for $N$ plus small constant terms. One should note, however, that in the case of the stored stencils approach there will be no temporal cache effects when we proceed from one node to the next, since the stencil values/row entries cannot be reused. In the two other approaches, we expect to see positive cache effects due to temporal and spatial reuse of some values of the coefficient function $k$, i.e., results closer to $N_{opt}$. This can be seen in the comparisons in section 7.4.

7. Numerical accuracy study and run-time comparison. In this section, we provide different numerical results which illustrate the accuracy and run-time of the new scaling approach in comparison to the elementwise finite element assembling based on nodal integration within a matrix-free framework. We consider different cases such as scalar and tensorial coefficient functions $k$ and the scenario of a geometry mapping. Throughout this section, we denote the time-to-solution by tts and, by relative tts we always mean the ratio of the time-to-solution of the stencil scaling approach with respect to the nodal integration. From our theoretical considerations for one stencil application from Table 6.1, we expect a relative tts of roughly one-third. Further, we denote the estimated order of convergence by eoc and the asymptotic convergence rates of the multigrid solver by $\rho$ defined as $\rho = \left( \frac{r(i^*)}{r(5)} \right)^{1/(i^* - 5)}$, where $r(i)$ is the $L^2$ residual at iteration $i$, and $i^*$ is the final iteration of the solver. Each of the following 3D computations was conducted on SuperMUC Phase 2 using the Intel 17.0 compiler together with the Intel 2017 MPI library. For all runs, we specify the compiler flags `-O3 -march=native -xHost`. Note that the serial runs using only a single compute core are not limited to run on large machines like SuperMUC but can also be run on typical modern desktop workstations with enough memory. The peak memory usage by our largest serial run was at about 4.46 GiB.

7.1. A quantitative comparison in 2D for a scalar permeability. In this example, we consider as domain the unit-square $\Omega = (0, 1)^2$ and use a nonpolynomial manufactured solution

$$u(x, y) = \frac{x^4 y}{xy + 1}.$$  

We employ as coefficient function $k(x, y; m) = 2 + \sin(m \pi x) \sin(m \pi y)$ with $m \in \{2, 4, 8\}$. The right-hand side is computed by inserting the above definitions into the equation. This construction has the advantage that we can study the effect of the
Table 7.1

Results for a regular triangular mesh. Here, $L$ denotes the (uniform) refinement level and eoc the estimated order of convergence.

<table>
<thead>
<tr>
<th>$m$</th>
<th>Midpoint integration</th>
<th>Nodal integration</th>
<th>Stencil approach</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$L$</td>
<td>$|u - u_n^h|_0$</td>
<td>eoc</td>
</tr>
<tr>
<td>2</td>
<td>0</td>
<td>3.75e-03</td>
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<td>2.32e-04</td>
<td>1.98</td>
</tr>
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<td></td>
<td>3</td>
<td>5.97e-05</td>
<td>1.99</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>1.50e-05</td>
<td>1.99</td>
</tr>
</tbody>
</table>

magnitude of $\nabla k$ in a systematic fashion by adjusting $m$. We perform a study on a regular triangular mesh comparing errors of the discrete solutions obtained by two standard Galerkin finite element approaches and our proposed scaling approach. The error norms are approximated using a fifth order quadrature rule, while we use a second order scheme to evaluate the weak right-hand side. Results are listed in Table 7.1. Here, $u_n^h$ denotes the approximation obtained by employing a nodal quadrature rule, resulting in bilinear form (3.4), while $u_b^h$ uses a quadrature rule that evaluates $k$ at the triangle’s barycenter.

7.2. A quantitative comparison in 3D for a scalar permeability. As a second test, we consider a nonlinear solution similar to the 2D tests in the previous section,

$$u = \frac{x^3 y + z^2}{xyz + 1},$$

with a parameter-dependent scalar coefficient function $k(x, y, z; m) = \cos(m\pi xyz) + 2$ on the unit-cube $\Omega = (0, 1)^3$ discretized by six tetrahedra. Here, and in the second 3D example below, the weak right-hand side is computed by interpolating the right-hand side associated with our manufactured solution into our finite element ansatz space and subsequent multiplication with the mass matrix. The $L^2$ error is approximated by a discrete version, i.e., an appropriately scaled nodal $L^2$ error. While more advanced approaches could be used here, the ones chosen are completely sufficient for our purpose, which is to demonstrate that our stencil scaling approach behaves analogously to a classical finite element approach. We employ a multigrid solver with a V(3,3) cycle on a single compute core and stop after 10 multigrid iterations, i.e., $i^* = 10$, which is enough to reach the asymptotic regime. In Table 7.2, we report convergence of the discretization error for the three approaches, i.e., classical finite element with nodal
Table 7.2

3D results in the case of a scalar coefficient function with errors measured in the discrete $L^2$-norm.

<table>
<thead>
<tr>
<th>$L$</th>
<th>DOF</th>
<th>Nodal integration</th>
<th>Scale Vol+Face</th>
<th>Scale all</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Error</td>
<td>ecc</td>
<td>$\rho$</td>
</tr>
<tr>
<td>1</td>
<td>3.43e+02</td>
<td>2.46e-03</td>
<td>0.07</td>
<td>2.42e-03</td>
</tr>
<tr>
<td>2</td>
<td>3.38e+03</td>
<td>7.06e-04</td>
<td>1.80</td>
<td>5.97e-04</td>
</tr>
<tr>
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<td>1.80e-04</td>
<td>1.97</td>
<td>1.46e-04</td>
</tr>
<tr>
<td>4</td>
<td>2.50e+05</td>
<td>4.46e-05</td>
<td>2.01</td>
<td>3.59e-05</td>
</tr>
<tr>
<td>5</td>
<td>2.05e+06</td>
<td>1.11e-05</td>
<td>2.01</td>
<td>8.88e-06</td>
</tr>
<tr>
<td>6</td>
<td>1.66e+07</td>
<td>2.75e-06</td>
<td>2.01</td>
<td>2.21e-06</td>
</tr>
</tbody>
</table>

$m = 3$

<table>
<thead>
<tr>
<th>$L$</th>
<th>DOF</th>
<th>Nodal integration</th>
<th>Scale Vol+Face</th>
<th>Scale all</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Error</td>
<td>ecc</td>
<td>$\rho$</td>
</tr>
<tr>
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<td>0.07</td>
<td>5.43e-03</td>
</tr>
<tr>
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</tr>
<tr>
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<td>1.63</td>
<td>4.04e-04</td>
</tr>
<tr>
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<td>1.31e-04</td>
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<td>1.01e-04</td>
</tr>
<tr>
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<td>3.32e-05</td>
<td>1.98</td>
<td>2.52e-05</td>
</tr>
<tr>
<td>6</td>
<td>1.66e+07</td>
<td>8.30e-06</td>
<td>2.00</td>
<td>6.29e-06</td>
</tr>
</tbody>
</table>

$m = 8$

integration, stencil scaling on volumes and faces, but classical finite element assembly on edges and vertices, and stencil scaling on all primitives. The refinement is given by $L$, where $L = -2$ denotes the macro-mesh. We observe quadratic convergence of the discrete $L^2$ error for all three approaches and a relative tts of about 32% on level $L = 6$.

7.3. A quantitative comparison in 3D for a permeability tensor. As a third test, we consider a full symmetric and positive definite permeability tensor $K$ with nonlinear components. The off-diagonal components are negative or zero:

$$K = \begin{pmatrix}
  x^2 + 2y^2 + 3z^2 + 1 & -y^2 & -z^2 \\
  -y^2 & 2x^2 + 3y^2 + z^2 + 1 & -x^2 \\
  -z^2 & -x^2 & 3x^2 + y^2 + 2z^2 + 1
\end{pmatrix}.$$

The manufactured solution is set to

$$u = \frac{x^4y + 2z}{xyz + 1},$$

and we consider as domain the unit-cube discretized by twelve tetrahedra. We employ the same multigrid solver as in the previous subsection on a single compute core, but stop the iterations if the residual is reduced by a factor of $10^{-9}$. We denote the final iteration by $i^\ast$. The results for the classical finite element approach and the stencil scaling approach on volumes and faces with classical finite element assembly on edges and vertices are reported in Table 7.3. We observe quadratic convergence of the discrete $L^2$ error for both approaches and a relative tts of 31% on our finest level, $L = 6$.

Note that, as in Table 7.2, the relative tts exhibits a slightly nonmonotonic behavior. This stems from the fact that different types of primitives (e.g., faces and volume) have a different tts behavior for increasing $L$ and profit differently from the scaling approach. For large $L$, tts is dominated by the work performed on the volume DOFs, as is the relative tts between two approaches. A test with a single macro tetrahedron showed a monotonic behavior for the relative tts.
Table 7.3

3D results in the case of a tensorial permeability with discretization errors measured in the discrete $L^2$-norm.

<table>
<thead>
<tr>
<th>$L$</th>
<th>DOF</th>
<th>Nodal integration</th>
<th>Scale Vol+Face</th>
<th>Rel. tts</th>
</tr>
</thead>
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<td>8.55e+02</td>
<td>1.92e-03</td>
<td>0.07</td>
<td>2.21e-03</td>
</tr>
<tr>
<td>2</td>
<td>7.47e+03</td>
<td>4.40e-04</td>
<td>0.16</td>
<td>5.19e-04</td>
</tr>
<tr>
<td>3</td>
<td>6.26e+04</td>
<td>1.06e-04</td>
<td>0.24</td>
<td>1.27e-04</td>
</tr>
<tr>
<td>4</td>
<td>5.12e+05</td>
<td>2.00e-05</td>
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<td>4.15e+06</td>
<td>6.47e-06</td>
<td>0.35</td>
<td>7.86e-06</td>
</tr>
<tr>
<td>6</td>
<td>3.34e+07</td>
<td>1.61e-06</td>
<td>0.37</td>
<td>1.96e-06</td>
</tr>
</tbody>
</table>

7.4. Memory traffic and roofline analysis. In section 6.4 a theoretic assessment of memory accesses was presented. In order to verify these results we devised a benchmark to run on a single compute node of SuperMUC Phase 2. We compare our proposed scaling approach against the on-the-fly assembly and the stored stencil variant, described in section 6.4. Floating-point performance and memory traffic are measured using the Intel Advisor 2018 tool [23].

We start by giving a brief summary of the SuperMUC Phase 2 hardware details. Values were taken from [30]. One compute node consists of two Haswell Xeon E5-2697 v3 processors clocked at 2.6 GHz. Each CPU is equipped with 14 physical cores. Each core has a dedicated L1 (data) cache of size 32 kB and a dedicated L2 cache of size 256 kB. The theoretical bandwidths are 343 GB/s and 92 GB/s, respectively. The CPUs are running in cluster-on-die mode. Thus, each node represents four NUMA domains each consisting of 7 cores with a separate L3 cache of size 18 MB and a theoretical bandwidth of 39 GB/s. Note that the cache bandwidth scales linearly with the number of cores. Furthermore, each node provides 64 GB of shared memory with a theoretical bandwidth of 6.7 GB/s.

The benchmark computes the residual $r = f - A(k)u$ for a scalar operator $A(k)$. We only consider volume primitives and their associated DOFs in the benchmark. The residual computation is iterated 200 times to improve signal-to-noise ratio. The program is executed using 28 MPI processes, pinned to the 28 physical cores of a single node. This is essential to avoid overly optimistic bandwidth values when only a single core executes memory accesses. Measurements with the Intel Advisor are carried out solely on rank 0. Moreover, all measurements are restricted to the innermost loop, i.e., where the actual nodal updates take place, to obtain a clear picture. This restriction does not influence the results as the outer loops are identical in all three variants. We choose $L = 6$ as refinement level, which gives us $2.7 \cdot 10^6$ DOFs per MPI rank. The computation involves three scalar fields $u$, $k$, and $f$, which each require $\sim 22$ MB of storage. In Figure 7.1 (left) we present a roofline analysis (see [22, 38]) based on the measurements. The abscissa shows the arithmetic intensity, i.e., the number of FLOPs performed divided by the number of bytes loaded into CPU registers per nodal update. The ordinate gives the measured performance as FLOPs performed per second. The diagonal lines give the measured DRAM and cache bandwidths. The values for the caches are those reported by the Intel Advisor. Naturally these measured values are smaller than the theoretical ones given in the hardware description above.

In order to assess the practical DRAM bandwidth of the complete node we used the LIKWID tool (see [35]) and performed a memcopy benchmark with nontemporal stores. We found that the system can sustain a bandwidth of about 104 GB/s, which for perfect load-balancing on the nodes, as is the case in our benchmark, results in
about 3.7 GB/s per core. The maximum performance for double precision vectorized fused multiply-add operations is also reported by the Advisor tool (35.7 GFLOPs/s).

For all three kernels we observe a quite similar arithmetic intensity. The value for the stored stencil variant is close to what one would expect theoretically. In this case one needs to load 31 values of size 8 bytes (15 $u$ values, 1 $f$ value, and 15 stencil weights), write back one residual value of 8 bytes, and perform 30 FLOPs, which results in an intensity of around 0.12 FLOPs/Byte. Both the stencil scaling approach and the on-the-fly assembly show a slightly higher intensity, but the difference seems negligible.

Using FLOPs/s as a performance criterion, we see that the stored stencil approach achieves significantly smaller values than the two other approaches, which perform about a factor of 4 better. The limiting resource for both the on-the-fly assembly and the stencil scaling is the L3 bandwidth. This confirms our expectations from section 6.4 showing that values of $k$ are not evicted from the L3 cache and can be reused.

For the stored stencil approach the performance is somewhere in between the limits given by the L3 and DRAM bandwidths. Here, all stencil values have to be loaded from main memory, while values of $u$ may still be kept in the cache, as with the two other approaches. Due to overlapping of computation and load operations, the performance is still above the DRAM bandwidth limit, but clearly below what the L3 bandwidth would allow.

Further assessment requires a more detailed analysis like the execution-cache-memory model [20], which goes beyond the scope of this article.

From an application point of view, FLOPs/s is not the most relevant criterion, however. Of main importance to a user is overall run-time. In this respect the interesting measure is the number of stencil applications per second, or equivalently the number of DOFs updated per second (DOFs/s). The latter can be derived from the FLOPs/s value based on the number of operations required to (assemble and) apply a local stencil for the three different methods. These values and the derived DOFs/s are shown in the right part of Figure 7.1. The number of FLOPs required per update was computed from the number of performed FLOPs, as reported by the Intel Advisor, and the number of DOFs inside a volume primitive. Note that these values match very well our theoretical considerations from Table 6.1.

As the FLOPs/s value attained by the on-the-fly and stencil scaling approaches...
are almost identical, it is the reduced number of operations required in the stencil scaling variant which directly pays off. The three times lower FLOP count directly translates to a threefold increase in the DOFs/s and, thus, a similar reduction in run-time.

Furthermore, we observe that our stencil scaling version also gives a higher number of DOFs/s than the stored stencil approach. While this increase is not as dramatic compared to the on-the-fly assembly, we emphasize that the stored stencil approach within HHG does not require additional memory traffic for information on the matrix’s sparsity pattern or involve indirect accesses as in a classical CRS format. More importantly, for the largest simulation with \( L = 6 \), carried out in section 7.5 below, the stored stencil approach over all levels of the mesh hierarchy would require about 22 TB of storage. Together with the scalar fields \( u, f \) and the tensor \( K \) this would exceed the memory available on SuperMUC. Per core typically 2.1 GB are available to an application \[30\], which sums up to around 30 TB for the 14 310 cores used. The stencil scaling approach, on the other hand, only requires the storage of 120 bytes, i.e., a single 15-point stencil per primitive and level, for the operator construction. This results in less than 165 MB over all cores for the complete simulation. This value could even be further reduced to 26 MB by exploiting similarities of stencils on different levels and the zero sum property. However, this does not seem worth the effort.

7.5. Application to a blending setting and large scale results. To demonstrate the advantages of our novel scaling approach also for a more realistic scenario, we consider an example using a blending function, as mentioned in section 2. To this end, we consider a half cylinder mantle with inner radius \( r_1 = 0.8 \) and outer radius \( r_2 = 1.0 \), height \( z_1 = 4.0 \), and with an angular coordinate between 0 and \( \pi \) as our physical domain \( \Omega_{\text{phy}} \). The cylinder mantle is additionally warped inwards by \( w(z) = 0.2 \sin (z\pi/z_1) \) in an axial direction. The mapping \( \Phi : \Omega_{\text{phy}} \rightarrow \Omega \) is given by

\[
\Phi(x, y, z) = \left(\frac{\sqrt{x^2 + y^2 + w(z)}}{\arccos \left(\frac{z}{\sqrt{x^2 + y^2}}\right) \frac{\pi}{z}}\right),
\]

with the reference domain \( \Omega = (r_1, r_2) \times (0, \pi) \times (0, z_1) \). Using (2.1), it follows for the mapping tensor \( K \),

\[
K = \frac{(D\Phi)(D\Phi)^\top}{|\det D\Phi|} = \sqrt{x^2 + y^2} \begin{pmatrix} w'(z)^2 + 1 & 0 & w'(z) \\ 0 & 1/(x^2+y^2) & 0 \\ w'(z) & 0 & 1 \end{pmatrix}.
\]

Obviously, this tensor is symmetric and positive definite. In addition to the geometry blending, we use a variable material parameter \( a(x, y, z) = 1 + z \). On the reference domain \( \Omega \) this yields the PDE \( -\text{div} aK \nabla u = f \). As the analytic solution on the reference domain, we set

\[
u(\hat{x}, \hat{y}, \hat{z}) = \sin \left(\frac{\hat{x} - r_1}{r_2 - r_1} \pi\right) \cos (4\hat{y}) \exp \left(\frac{\hat{z}}{2}\right).
\]

The analytic solution mapped to the physical domain is illustrated in the right part of Figure 7.2.

For our numerical experiments, we employ a macro-mesh composed of 9540 hexahedral blocks, where each block is further split into six tetrahedral elements; see
Figure 7.2 (left). The resulting system is solved using 14,310 compute cores; i.e., we assign four macro-elements per core. For the largest run we have a system with $O(10^{11})$ DOF. We employ a multigrid solver with a V(3,3) cycle. The iteration is stopped when the residual has been reduced by a factor of $10^{-8}$. In Table 7.4, we report the resulting discretization error, the asymptotic multigrid convergence order $\rho$, and the time-to-solution.

These results demonstrate that the new scaling approach maintains the discretization error, as is expected on structured grids from our variational crime analysis, as well as the multigrid convergence rate. For small $L$ we observe that the improvement in time-to-solution is only modest as here the influence of vertex and edge primitives is more pronounced. But for increasing $L$ this influence decreases and for $L \geq 4$ the run-time as compared to the nodal integration approach is reduced to about 30%.

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