

PREPRINT 2005–06

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Chalmers Finite Element Center Chalmers University of Technology SE–412 96 Göteborg Sweden Göteborg, July 2005

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Printed in Sweden Chalmers University of Technology Göteborg, Sweden 2005

TOPOLOGICAL OPTIMIZATION OF THE EVALUATION OF FINITE ELEMENT MATRICES

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ABSTRACT. We present a topological framework for finding low-flop algorithms for evaluating element stiffness matrices associated with multilinear forms for finite element methods. This framework relies on phrasing the computation on each element as the contraction of each collection of reference element tensors with an element-specific geometric tensor. We then present a new concept of *complexity-reducing relations* that serve as distance relations between these reference element tensors. This notion sets up a graph-theoretic context in which we may find an optimized algorithm by computing a minimum spanning tree. We present experimental results for some common multilinear forms showing significant reductions in operation count and also discuss some efficient algorithms for building the graph we use for the optimization.

1. Introduction

Several ongoing projects have led to the development of tools for automating important aspects of the finite element method, with the potential for increasing code reliability and decreasing development time. By developing libraries for existing languages or new domain-specific languages, these software tools allow programmers to define variational forms and other parts of a finite element method with succinct, mathematical syntax. Existing C++ libraries for finite elements include DOLFIN [6, 7], Sundance [16, 17] and deal.II [2]. These rely on some combination of operator overloading and object orientation to present a high-level syntax to the user. Other projects have defined new languages with their own grammar and syntax, such as FreeFEM [20], GetDP [5], and Analysa [1]. Somewhere between these two approaches is the FEniCS Form Compiler, FFC [13, 15] developed primarily by the second author. This Python library relies on operator overloading to define variational forms, but rather than relying on the Python interpreter to evaluate the

Date: July 11, 2005.

Key words and phrases. finite element, variational form, optimized algorithm, minimum spanning tree. Robert C. Kirby, Department of Computer Science, University of Chicago, 1100 East 58th Street, Chicago, Illinois 60637, USA. Email: kirby@cs.uchicago.edu. This work was supported by the U.S. Department of Energy Early Career Principal Investigator Program under award number DE-FG02-04ER25650.

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forms, FFC generates low-level code that can be compiled into other platforms, especially DOLFIN.

While these tools are effective at exploiting modern software engineering to produce workable systems, we believe that additional mathematical insight will lead to even more powerful codes with more general approximating spaces and more powerful algorithms. For one, work by the first author [9, 10] shows how the Ciarlét definition of the finite element leads to a code for arbitrary order elements of general type. This code, FIAT, is used by FFC and is currently being integrated with Sundance. Second, the first three authors together with Knepley [11, 12], studied how to efficiently (in the sense of operation count) evaluate local stiffness matrices for finite element methods. All entries of the local stiffness matrix for an element e may be expressed as the contraction of some reference element tensor with a "geometric" tensor. To build the $n \times n$ stiffness matrix for one element, one must contract this tensor with n^2 reference element tensors. We saw that there are relations between many of these tensors (equality, colinearity, small Hamming distance) that, if exploited, lead to an algorithm with significantly fewer floating point operations.

In [11], we devised a crude algorithm that searched for and exploited these dependencies, generating simple Python code for evaluating the local stiffness matrix for the Laplacian on triangles given the geometric tensor. In this paper, we set the optimization process in a more abstract graph/topological context. In Section 2, we express the calculation for some finite element operators as tensor contractions. These ideas are more fully developed in [13], where the first two authors show how this tensor structure may be used to implement a compiler capable of generating the reference tensors and hence code for building the stiffness matrices for general multilinear forms. In Section 3, introduce the idea of complexity-reducing relations, which are a kind of distance relations among the tensors that serve to model the cost of complexity in computing the contractions. Using these ideas, we show how to derive an algorithm for performing the computation that is optimal in a certain sense. We demonstrate the reduction in operation count for several finite element operations in Section 4. After this, we show how this efficient algorithm may be derived more efficiently through some search procedures in Section 5 and make some conclusions and remarks about ongoing work in Section 6.

2. Finite element formulation

The finite element method is a general methodology for the discretization of differential equations. A linear (or linearized) differential equation for the unknown function u is expressed in the form of a canonical variational problem and the discrete approximation U of u is sought as the solution of a discrete version of the variational problem [4, 3]: Find $U \in V$ such that

(2.1)
$$a(v, U) = L(v) \quad \forall v \in \hat{V},$$

where (\hat{V}, V) is a pair of suitable discrete (typically piecewise polynomial) function spaces, $a: \hat{V} \times V \to \mathbb{R}$ a bilinear form and $L: \hat{V} \to \mathbb{R}$ a linear form.

The variational problem (2.1) corresponds to a linear system $A\xi = b$ for the expansion coefficients $\xi \in \mathbb{R}^M$ of the discrete function U in a basis $\{\varphi_i\}_{i=1}^M$ for V. If $\{\hat{\varphi}_i\}_{i=1}^M$ is the corresponding basis for \hat{V} , the entries of A and b are given by $A_{ij} = a(\hat{\varphi}_i, \varphi_j)$ and $b_i = L(\hat{\varphi}_i)$ respectively. When we consider consider general multilinear forms below, the multilinear form a is represented by a tensor A.

2.1. Multilinear forms. Let now $\{V_i\}_{i=1}^r$ be a given set of discrete function spaces defined on a triangulation $\mathcal{T} = \{e\}$ of $\Omega \subset \mathbb{R}^d$. We consider a general multilinear form a defined on the product space $V_1 \times V_2 \times \cdots \times V_r$:

$$(2.2) a: V_1 \times V_2 \times \cdots \times V_r \to \mathbb{R}.$$

Typically, r=1 (linear form) or r=2 (bilinear form), but forms of higher arity appear frequently in applications and include variable coefficient diffusion and advection of momentum in the incompressible Navier–Stokes equations.

Let $\{\varphi_i^1\}_{i=1}^{M_1}, \{\varphi_i^2\}_{i=1}^{M_2}, \dots, \{\varphi_i^r\}_{i=1}^{M_r}$ be bases of V_1, V_2, \dots, V_r and let $i = (i_1, i_2, \dots, i_r)$ be a multiindex. The multilinear form a then defines a rank r tensor given by

(2.3)
$$A_i = a(\varphi_{i_1}^1, \varphi_{i_2}^2, \dots, \varphi_{i_r}^r).$$

In the case of a bilinear form, the tensor A is a matrix (the stiffness matrix), and in the case of a linear form, the tensor A is a vector (the load vector).

To compute the tensor A by assembly, we need to compute the element tensor A^e on each element e of the triangulation \mathcal{T} of Ω [13]. Let $\{\varphi_i^{e,1}\}_{i=1}^{n_1}$ be the restriction to e of the subset of $\{\varphi_i^1\}_{i=1}^{M_1}$ supported on e and define the local bases on e for V_2, \ldots, V_r similarly. The rank r element tensor A^e is then given by

(2.4)
$$A_i^e = a_e(\varphi_{i_1}^{e,1}, \varphi_{i_2}^{e,2}, \dots, \varphi_{i_r}^{e,r}).$$

2.2. Evaluation by tensor representation. The element tensor A^e can be efficiently computed by representing A^e as a special tensor product. If the multilinear form a is given by an integral over the domain Ω , then each entry A^e_i of A^e is given by an integral over the element e. By a change of variables and a series of linear transformations, we may rewrite this integral as an integral over a reference element E. In particular, when the map from the reference element E is affine, the linear transformations of derivatives can be moved outside of the integral to obtain a representation of the element tensor A^e as a tensor product of a constant tensor A^0 and a tensor G_e that varies over the set of elements,

$$(2.5) A_i^e = A_{i\alpha}^0 G_e^\alpha,$$

or more generally a sum $A_i^e = A_{i\alpha}^{0,k} G_{e,k}^{\alpha}$ of such tensor products, where i and α are multiindices and we use the convention that repetition of an index means summation over that index. We refer to A^0 as the reference tensor and to G_e as the geometric tensor. The rank of the reference tensor is the sum of the rank r = |i| of the element tensor and the rank $|\alpha|$ of the geometric tensor G_e . As we shall see, the rank of the geometric tensor depends on the specific form. In [13], we present an algorithm that computes the tensor representation (2.5) for fairly general multilinear forms. This algorithm forms the foundation for the FEniCS Form Compiler, FFC [15].

As an example, we consider here the tensor representation of the element tensor A^e for Poisson's equation $-\Delta u(x) = f(x)$ with homogeneous Dirichlet boundary conditions on a domain Ω . The bilinear form a is here given by $a(v,u) = \int_{\Omega} \nabla v(x) \cdot \nabla u(x) dx$ and the linear form L is given by $L(v) = \int_{\Omega} v(x) f(x) dx$. By a change of variables using an affine map $F_e: E \to e$, we obtain the following representation of the element tensor A^e :

(2.6)
$$A_{i}^{e} = \int_{e} \nabla \varphi_{i_{1}}^{e,1}(x) \cdot \nabla \varphi_{i_{2}}^{e,2}(x) dx$$
$$= \det F_{e}' \frac{\partial X_{\alpha_{1}}}{\partial x_{\beta}} \frac{\partial X_{\alpha_{2}}}{\partial x_{\beta}} \int_{E} \frac{\partial \Phi_{i_{1}}^{1}(X)}{\partial X_{\alpha_{1}}} \frac{\partial \Phi_{i_{2}}^{2}(X)}{\partial X_{\alpha_{2}}} dX = A_{i\alpha}^{0} G_{e}^{\alpha},$$

where $A^0_{i\alpha} = \int_E \frac{\partial \Phi^1_{i_1}(X)}{\partial X_{\alpha_1}} \frac{\partial \Phi^2_{i_2}(X)}{\partial X_{\alpha_2}} \, \mathrm{d}X$, $G^\alpha_e = \det F'_e \frac{\partial X_{\alpha_1}}{\partial x_\beta} \frac{\partial X_{\alpha_2}}{\partial x_\beta}$, and $\Phi^j_{i_j} = \varphi^{e,j}_{i_j} \circ F_e$ for j=1,2. We see that the reference tensor A^0 is here a rank four tensor and the geometric tensor G_e is a rank two tensor. In [11], we saw that dependencies frequently occur between A^0_i and A^0_j for many multiindices i,j that can reduce the overall cost of computation.

3. Optimizing stiffness matrix evaluation

3.1. An abstract optimization problem. Since all of the computation to evaluate a local stiffness matrix for a multilinear form is tensor contraction, we may just as easily consider them as vectors and contraction as the Euclidean inner product. To formalize the optimization process then, we let $Y = \{y^i\}_{i=1}^n$ be a collection of n vectors in \mathbb{R}^m . In the most general case, this is a collection rather than a set, as some of the items in Y may be identical. Corresponding to Y, we must find a process for computing for arbitrary $g \in \mathbb{R}^m$ the collection of items $\{(y^i)^t g\}_{i=1}^n$. Throughout this paper, we will measure the cost of this as the total number of multiply-add pairs (MAPs) required to complete all the dot products. This cost is always bounded by nm, but we hope to improve on that. This could be alternatively formalized as building an abstract control-flow graph for performing the dot products that is equivalent to the naive process but contains a minimal number of nodes. Our techniques, however, rely on structure that is not apparent to traditional optimizing compilers, so we prefer the present formulation.

We seek out ways of optimizing the local matrix evaluation that rely on notions of distance between a few of the underlying vectors. The Euclidean metric is not helpful here; we focus on other, discrete measures of distance such that if y and z are close together, then y^tg is easy to compute once z^tg is known (and vice versa). Many of the dependencies we considered in [11] were between pairs of vectors — equality, colinearity, and Hamming distance. Here, we develop a theory for optimizing the evaluation of finite element matrices under binary relations between the members of the collection. This starts by introducing some notions of distance on the collection of vectors and finds an optimized computation with respect to those notions by means of a minimum spanning tree.

3.2. Complexity-reducing relations.

Definition 1. Let $\rho: Y \times Y \to [0, m]$ be symmetric. We say that ρ is complexity-reducing if for every $y, z \in Y$ with $\rho(y, z) \leq k < m$, $y^t g$ may be computed using the result $z^t g$ in no more than k multiply-add pairs.

The topological structure induced by complexity-reducing relations may not be a metric space, as we do not require a triangle inequality to hold. We shall remark on this further below.

Example 1. Let $e^+(y, z) = d(1 - \delta_{y,z})$, where $\delta_{y,z}$ is the standard Kronecker delta. Then, e^+ is seen to be complexity-reducing, for if $e^+(y, z) = 0$, then $y^t g = z^t g$ for all $g \in \mathbb{R}^m$ and hence the former requires no arithmetic once the latter is known. Similarly, we can let $e^-(u, v) = e^+(u, -v)$, for if u = -v, then computing $u^t g$ from $v^t g$ requires only a sign flip and no further floating point operations.

Example 2. Let

(3.1)
$$c(y,z) = \begin{cases} 0, & y = z \\ 1, & y = \alpha z \text{ for some } \alpha \in \mathbb{R}, \alpha \neq 0, 1 \\ m, & otherwise \end{cases}$$

Then c is complexity-reducing, for $y^tg = (\alpha z)^tg = \alpha(z^tg)$, so y^tg may be computed with one additional floating point operation once z^tg is known.

Example 3. Let $H^+(y, z)$ be the Hamming distance, the number entries in which y and z differ. Then H^+ is complexity-reducing. If $H^+(y, z) = k$, then y and z differ in k entries, so the difference y - z has only k nonzero entries. Hence, $(y - z)^t g$ costs k multiply-add pairs to compute, and we may write $y^t g = (y - z)^t g + z^t g$. By the same argument, we can let $H^-(y, z) = H^+(y, -z)$.

Theorem 1. Let ρ_1 and ρ_2 be complexity-reducing relations. Define

(3.2)
$$\rho(y,z) = \min(\rho_1(y,z), \rho_2(y,z)).$$

Then ρ is a complexity-reducing relation.

Proof. Pick $y, z \in Y$, let $1 \le i \le 2$ be such that $\rho(y, z) = \rho_i(y, z)$ and let $\rho_i(y, z) \equiv k$. But ρ_i is a complexity-reducing relation, so for any $g \in \mathbb{R}^m$, $y^t g$ may be computed in no more than $k = \rho(y, z)$ multiply-add pairs. Hence ρ is complexity-reducing.

This simple result means that we may consider any finite collection of complexity-reducing relations (e.g. colinearity together with Hamming distance) as if they were a single relation for the purpose of finding an optimized computation in the later discussion.

Definition 2. If ρ is a complexity-reducing relation defined as the minimum over a finite set of complexity-reducing relations, we say that it is composite. If it is not, we say that it is simple.

Remark 1. To see that not all complexity-reducing relations are metrics, it is easy to find a complexity-reducing relation that is the minimum over two metrics that violates the triangle inequality. To see this, let $\rho(y,z) = \min(H^+(y,z),c(y,z))$. It is not hard to show that H^+ and c are both metrics. If we take $y = (1,2,2)^t$ and $z = (0,4,4)^t$, then $\rho(y,z) = 3$ since the vectors are neither colinear nor share any common entries. However, if we let $x = (0,2,2)^t$, then $\rho(y,x) = 1$ since the vectors share two entries and $\rho(x,z) = 1$ by colinearity. Hence, $\rho(y,z) > \rho(y,x) + \rho(x,z)$, and the triangle inequality fails.

Remark 2. Later, we will put all of the vectors into a graph with weights given by the values of a complexity-reducing relation. Computing all-to-all shortest paths in this graph would provide a metric provided that $\rho(y, z) \neq 0$ whenever $y \neq z$. However, this is by no means necessary.

3.3. Finding an optimized computation. Having defined complexity-reducing relations and given several examples, we now show how they may be used to determine an optimized evaluation of the stiffness matrix. We shall work in the context of a single complexity-reducing relation ρ , which may be composite.

In order to compute $\{(y^i)^t g\}_{i=1}^n$, we would like to pick some $y^i \in Y$ and compute $(y^i)^t g$. Then, we want to pick some y^j that is very close to y^i under ρ and then compute $(y^j)^t g$. Then, we pick some y^k that is very close to either y^i or y^j and compute that dot product. So, this process continues until all the dot products have been computed. Moreover, since the vectors Y depend only on the variational form and finite element space and not the mesh or parameters, it is possible to do this search once offline and generate low-level code that will exploit these relations. We first formalize the notion of finding the optimal computation and then how the code may be generated.

We introduce a weighted, undirected graph G=(Y,E) where Y is our collection of vectors defined above. Our graph is completely connected; that is, every pair of vectors y^i, y^j are connected by an edge. The weight of this edge is defined to be $\rho(y^i, y^j)$. We may think of walking along the edge from y^i to y^j as using the dot product $(y^i)^t g$ to compute $(y^j)^t g$. If ρ is composite, it will be helpful to know later which underlying relation gave the minimum value used for ρ . So, suppose that $\rho(y,z) = \min\{\rho_i(y,z)\}_{i=1}^R$. For any fixed y,z, let $\varrho(y,z)$ be a in integer in [1,R] such that $\rho_{\varrho(y,z)}(y,z) = \rho(y,z)$. In addition to weights, we thus associate with each edge $\{y^i,y^j\}$ the entity $\varrho(y^i,y^j)$.

A standard graph-theoretic object called a minimum spanning tree is exactly what we need [14]. A spanning tree, which we shall denote (Y, E') is a subgraph that satisfies certain properties. First, it contains all of the n nodes of the original graph. Second, (Y, E') is connected. Finally, E' has n-1 edges, so that there are no cycles (thus it is a tree). Now, there are possibly many spanning trees for a given graph. Every spanning tree has a weight associated with it that is the sum of the weights of its edges. A minimum spanning tree is a spanning tree such that the sum of the edge weights is as small as possible. Minimum spanning tree algorithms start with a particular node of the graph, called the root. Regardless of which root is chosen, minimum spanning tree algorithms will generate trees with exactly the same sum of edge weights.

While technically a minimum spanning tree is undirected, we can think of it as being a directed graph with all edges going away from the root. Such a notion tells us how to compute all of the dot products with minimal operations with respect to ρ . We start with the root node, which we assume is y^0 , and compute $(y^0)^t g$. Then, for each of the children of y^0 in the tree, we compute the dot products with g using the result of $(y^0)^t g$. Then, we use the dot products of the children to compute the dot products of each of the children's children, and so on. This is just a breadth-first traversal of the tree. A depth-first traversal of the tree would also generate a correct algorithm, but it would likely require more motion of the computed results in and out of registers at run-time.

The total number of multiply-add pairs in this process is m for computing $(y^0)^t g$ plus the sum of the edge weights of (Y, E'). As the sum of edge weights is as small as it can be, we have a minimum-cost algorithm for computing $\{(y^i)^t g\}_{i=1}^n$ with respect to ρ for any $g \in \mathbb{R}^m$. On the other hand, it is not a true optimal cost as one could find a better ρ or else use relations between more than two vectors (say three coplanar vectors). One other variation is in the choice of root vector. If, for example some y^i has several elements that are zero, then it can be dotted with g with fewer than m multiply add pairs. Hence, we pick some $\bar{y} \in Y$ such that the number of nonzero entries is minimal to be the root. We summarize these results in a theorem:

Theorem 2. Let G = (Y, E) be defined as above and let $g \in \mathbb{R}^m$ be arbitrary. The total number of multiply-add pairs needed to compute $\{(y^i)^t g\}_{i=1}^n$ is no greater than m' + w, where m' is the minimum number of nonzero entries of a member of Y and W is the weight of a minimum spanning tree of G

The overhead of walking through the tree at runtime would likely outweigh the benefits of reducing the floating point cost. We can instead traverse the tree and generate low-level code for computing all of the dot products - this function takes as an argument the vector g and computes all of the dot products of Y with g. An example of such code was presented in [11].

3.4. Comparison to spectral elements. Our approach is remarkably different than the spectral element method. In spectral element methods, one typically works with tensor products of Lagrange polynomials over logically rectangular domains. Efficient algorithms for evaluating the stiffness matrix or its action follow naturally by working dimension-by-dimension. While such decompositions are possible for unstructured shapes [8], these are restricted to specialized polynomial bases. On the other hand, our approach is blind both to the element shape and kind of approximating spaces used. While spectral element techniques may ultimately prove more effective when available, our approach will enable some level of optimization in more general cases, such as Raviart-Thomas-Nedelec [21, 22, 18, 19] elements on tetrahedra.

4. Experimental results

Ä Here, we show that this optimization technique is successful at generating low-flop algorithms for computing the element stiffness matrices associated with some standard

TABLE 1. Element matrix indices and associated tensors (the slice A_i^0 for each fixed index i) displayed as vectors for the Laplacian on triangles with quadratic basis functions. All vectors are scaled by six so they appear as integers.

index		vec	tor		index		vec	tor		index		vec	tor	
(0, 0)	3	3	3	3	(2, 0)	0	0	1	1	(4, 0)	0	0	-4	-4
(0, 1)	1	0	1	0	(2, 1)	0	0	-1	0	(4, 1)	0	0	0	0
(0, 2)	0	1	0	1	(2, 2)	0	0	0	3	(4, 2)	0	-4	0	-4
(0, 3)	0	0	0	0	(2, 3)	0	0	4	0	(4, 3)	-8	-4	-4	0
(0, 4)	0	-4	0	-4	(2, 4)	0	0	-4	-4	(4, 4)	8	4	4	8
(0, 5)	-4	0	-4	0	(2, 5)	0	0	0	0	(4, 5)	0	4	4	0
(1, 0)	1	1	0	0	(3, 0)	0	0	0	0	(5, 0)	-4	-4	0	0
(1, 1)	3	0	0	0	(3, 1)	0	0	4	0	(5, 1)	-4	0	-4	0
(1, 2)	0	-1	0	0	(3, 2)	0	4	0	0	(5, 2)	0	0	0	0
(1, 3)	0	4	0	0	(3, 3)	8	4	4	8	(5, 3)	0	-4	-4	-8
(1, 4)	0	0	0	0	(3, 4)	-8	-4	-4	0	(5, 4)	0	4	4	0
(1, 5)	-4	-4	0	0	(3, 5)	0	-4	-4	-8	(5, 5)	8	4	4	8

variational forms. First, we consider the bilinear forms for the Laplacian and advection in one coordinate direction for tetrahedra. Second, we study the trilinear form for the weighted Laplacian. In all cases, we generated the element tensors using FFC, the FEniCS Form Compiler [13, 15], which in turn relies on FIAT [9, 10] to generate the finite element basis functions and integration rules. Throughout this section, we let d=2,3 refer to the spatial dimension of Ω .

4.1. **Laplacian.** We consider first the standard Laplacian operator

(4.1)
$$a(v,u) = \int_{\Omega} \nabla v(x) \cdot \nabla u(x) \, \mathrm{d}x.$$

We gave a tensor product representation of the local stiffness matrix in equation (2.6). The indices of the local stiffness matrix and the associated tensors are shown in Table 1.

Because the element stiffness matrix is symmetric, we only need to build the triangular part. Even without any optimization techniques, this naturally leads from computing $|P|^2$ contractions at a cost of d^2 multiply-add pairs each to $\binom{|P|+1}{2}$ contractions, where |P| is the dimension of the polynomial space P. Beyond this, symmetry opens up a further opportunity for optimization. For every element e, G_e is symmetric. The equality of its off-diagonal entries means that the contraction can be performed in $\binom{d+1}{2}$ rather than d^2 entries. This is readily illustrated in the two-dimensional case. We contract a symmetric

Table 2. Element matrix indices and associated tensors (the slice A_i^0 for each fixed index i) for the Laplacian on triangles with quadratic basis functions after transformation to make use of symmetry.

index	V	ecto	r
(0, 0)	3	6	3
(0, 1)	1	1	0
(0, 2)	0	1	1
(0, 3)	0	0	0
(0, 4)	0	-4	-4
(0, 5)	-4	-4	0
(1, 1)	3	0	0
(1, 2)	0	-1	0
(1, 3)	0	4	0
(1, 4)	0	0	0
(1, 5)	-4	-4	0

index	V	ecto	r
(2, 2)	0	0	3
(2, 3)	0	4	0
(2, 4)	0	-4	-4
(2, 5)	0	0	0
(3, 3)	8	8	8
(3, 4)	-8	-8	0
(3, 5)	0	-8	-8
(4, 4)	8	8	8
(4, 5)	0	8	0
(5, 5)	8	8	8

 2×2 tensor G with an arbitrary 2×2 tensor K:

(4.2)
$$G: K = \begin{pmatrix} G_{11} & G_{12} \\ G_{12} & G_{22} \end{pmatrix} : \begin{pmatrix} K_{11} & K_{12} \\ K_{21} & K_{22} \end{pmatrix}$$
$$= G_{11}K_{11} + G_{12}(K_{12} + K_{21}) + G_{22}K_{22}$$
$$= \tilde{G}^t \hat{K},$$

where
$$\tilde{G}^t = (G_{11}, G_{12}, G_{22})$$
 and $\hat{K}^t = (K_{11}, K_{12} + K_{21}, K_{22})$.

This simple calculation implies a linear transformation $\hat{\cdot}$ from $\mathbb{R}^{d\times d}$ into $\mathbb{R}^{\binom{d+1}{2}}$ obtained by taking the diagonal entries of the matrix together with the sum of the off-diagonal entries, that may be applied to each reference tensor, together with an associated mapping $\tilde{\cdot}$ on symmetric tensors that just takes the symmetric part and casts it as a vector. Hence, the overall cost of computing an element stiffness matrix before optimizations goes from $|P|^2 d^2$ to $\binom{|P|+1}{2} \binom{d+1}{2}$.

An interesting property of this transformation of the reference tensor is that it is contractive for the complexity-reducing relations we consider. The Hamming distance between two items under $\hat{\cdot}$ is bounded by the Hamming distance between the items. More precisely, $\rho(\hat{y},\hat{z}) \leq \rho(y,z)$. Furthermore, if items are colinear before transformation, their images will be as well. Hence, for the optimizations we consider, we will not destroy any dependencies. Moreover, the transformation may introduce additional dependencies. For example, before applying the transformation, entries (0,1) and (1,5) are not closely related by Hamming distance or colinearity, as seen in Table 1. However, after the transformation, we see that the same items in Table 2 are colinear. Other examples can be found readily.

We optimized the evaluation of the Laplacian for Lagrange finite elements of degrees one through three on triangles and tetrahedra using a composite complexity-reducing relation with H^+ , H^- , and c defined in Examples 2 and 3. We performed the optimization both

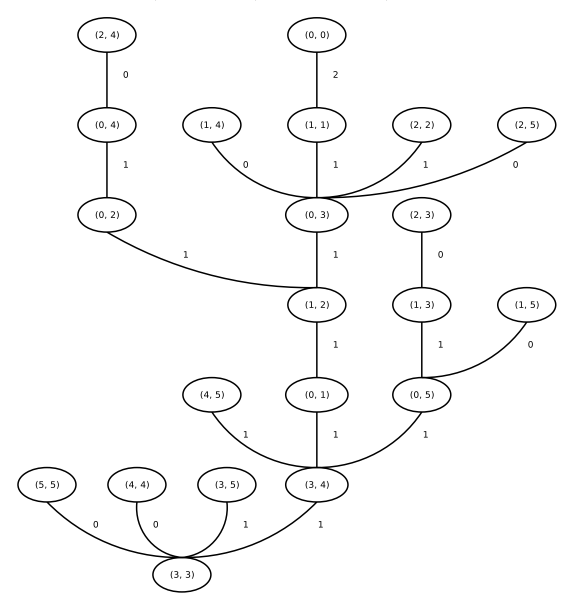


FIGURE 1. Minimum spanning tree for optimized computation of the Laplacian using quadratic elements on triangles. The node labeled (3,3) is the root, and the flow is from bottom to top.

with and without symmetry. The results are shown in Tables 3 and 4. Figure 1 shows a diagram of the minimum spanning tree computed by our code for the Laplacian on quadratic elements using symmetry. Each node of the graph is labeled with a pair (i, j) indicating the matrix entry (the vectors themselves are displayed in Table 2), and the edges are labeled with the associated weights. Simple inspection reveals that the sum of the edge weights is 14, which when added to 3 to compute the dot product for the root node, agrees with the entry for quadratics in Table 4.

Table 3. Number of multiply-add pairs in the optimized algorithm for computing the Laplacian element stiffness matrix on triangles and tetrahedra for Lagrange polynomials of degree one through three without using symmetry.

	tri	ang	les		tetrahedra						
degree	n	m	nm	MAPs		degree	n	m	nm	MAPs	
1	9	4	36	13		1	16	9	144	43	
2	36	4	144	25		2	100	9	900	205	
3	100	4	400	74		3	400	9	3600	864	

TABLE 4. Number of multiply-add pairs in the optimized algorithm for computing the Laplacian element stiffness matrix on triangles and tetrahedra for Lagrange polynomials of degree one through three using symmetry.

	tr	iang	gles			tet	rahe	edra	
degree	n	m	nm	MAPs	degree	n	m	nm	MAPs
1	6	3	18	9	1	10	6	60	27
2	21	3	63	17	2	55	6	330	101
3	55	3	165	46	3	210	6	1260	370

These techniques are successful in reducing the flop count, down to less than one operation per entry on triangles and less than two on tetrahedra for quadratic and cubic elements. We showed in [11] for low degree elements on triangles that going from standard numerical quadrature to tensor contractions led to a significant reduction in actual runtime for matrix assembly. From the tensor contractions, we got another good speedup by simply omitting multiplication by zeros. From this, we only gained a modest additional speedup by using our additional optimizations. However, this is most likely due to the relative costs of memory access and floating point operations. We have to load the geometric tensor from memory and we have to write every entry of the matrix to memory. Hence n+m in the tables gives a lower bound on memory access for computing the stiffness matrix. Our optimizations lead to algorithms for which there are a comparable number of arithmetic and memory operations. Hence, our optimization has succeeded in reducing the cost of computing the local stiffness matrix to a small increment to the cost of writing it to memory.

4.2. Advection in one coordinate direction. Now, we consider constant coefficient advection aligned with a coordinate direction

(4.3)
$$a(v,u) = \int_{\Omega} v(x) \frac{\partial u(x)}{\partial x_1} dx.$$

This is part of the operator associated with constant coefficient advection in some arbitrary direction — optimizing the other coordinate directions would give similar results.

TABLE 5. Number of multiply-add pairs in the optimized algorithm for computing the coordinate-advection element stiffness matrix on triangles and tetrahedra for Lagrange polynomials of degree one through three.

	tri	angl	les		tetrahedra						
degree	n	m	nm	MAPs		degree	n	m	nm	MAPs	
1	9	2	18	4		1	16	3	48	9	
2	36	2	72	22		2	100	3	300	35	
3	100	2	200	59		3	400	3	1200	189	

These results are shown in Table 5. Again, our optimization generates algorithms for which the predominant cost of computing the element matrix is writing it down, as there is significantly fewer than one floating point cycle per matrix entry in every case.

4.3. Weighted Laplacian. Our final operator is the variable coefficient Laplacian:

(4.4)
$$a_w(v,u) = \int_{\Omega} w(x) \nabla v(x) \cdot \nabla u(x) \, \mathrm{d}x.$$

This form may be viewed as a trilinear form a(w, v, u) in which w is the projection of the coefficient into the finite element space. For many problems, this can be performed without a loss in order of convergence. For nonlinear problems, we will have to reassemble this form at each nonlinear iteration, so it is an important step to optimize. The reference tensor is

(4.5)
$$A_{i\alpha}^{0} = \int_{E} \Phi_{\alpha_{1}}(X) \frac{\partial \Phi_{i_{1}}(X)}{\partial X_{\alpha_{2}}} \frac{\partial \Phi_{i_{2}}(X)}{\partial X_{\alpha_{2}}} dX$$

and the geometric tensor is

(4.6)
$$G_e^{\alpha} = w_{\alpha_1} \det F_e' \frac{\partial X_{\alpha_2}}{\partial x_{\beta}} \frac{\partial X_{\alpha_3}}{\partial x_{\beta}}.$$

Note that the geometric tensor is the outer product of the geometric tensor for the constant coefficient Laplacian, which we shall denote $(G^L)_e$ with the coefficients of the weight function. Unlike the constant coefficient case, the amount of arithmetic per entry in the element stiffness matrix grows with the polynomial degree.

We could simply proceed with the optimization as we did for other forms — the element tensor is just a collection of vectors that will have to be dotted into G_e for each element in the mesh, but now the dot products are more expensive since the vectors are longer. In this case, G_e must be explicitly formed for each element (this costs $|P|d^2$ once $(G^L)_e$ is formed). On the other hand, we could use the decomposition of G_e into $(G^L)_e$ and the coefficient vector w_k and do the contractions in stages. For example, we could organize the contraction as

(4.7)
$$A_i^e = \left(A_{i,(\alpha_1,\alpha_2,\alpha_3)}^0 \left(G^L \right)_e^{(\alpha_2,\alpha_3)} \right) w_{\alpha_1},$$

that is, for each of the $|P|^2$ entries of the stiffness matrix, we compute |P| contractions of $d \times d$ tensors with $(G^L)_e$. This is a similar optimization problem as the Laplacian, but with |P| times more elements to optimize over. After we do this set of computations, we must compute $|P|^2$ dot products with the coefficient vector w_k . Note that the contractions with $(G^L)_e$ may be optimized, but the resulting vectors to dot with w_k will not be known until run-time and must be computed at full cost. Hence, a lower bound for this approach is |P| multiply-add pairs per entry (assuming that the contractions with G^L were absolutely free). On the other hand, one could contract with the coefficient first to give an array of $|P|^2$ tensors of size $d \times d$, and then contract each of these with $(G^L)_e$. As before, the first step can be optimized, but the second step cannot.

In any of these cases, we may use the same transformations to exploit symmetry as we did in the constant coefficient case. Since $G_e^{\alpha_1,\alpha_2,\alpha_3} = G_e^{\alpha_1,\alpha_3,\alpha_2}$, we may view each slice A_i^0 of the reference tensor as an array of |P| tensors of size $d \times d$ and apply the transformation to each of these. If we fully form G_e , this reduces the cost from $|P|d^2$ to $|P|\binom{d+1}{2}$. In all of our experiments, we made use of this.

In Table 6, we see the cost of computing the weighted Laplacian by the first approach (optimizing directly the tensor product $A_i^e = A_{i\alpha}^0 G_e^{\alpha}$). While the optimizations are not as successful as for the constant coefficient operators, we still get reductions of 30%-50% in the operation counts.

When we perform the contraction in stages, we find more dependencies (for example, the slices of two of the tensors could be colinear although the entire tensors are not). We show the cost of performing the optimized stage for contracting with $(G^L)_e$ first in Table 7 and for contracting with w_k first in Table 8.

In order to get a fair comparison between these approaches, we must factor in the additional costs of building G_e or performing the second stage of contraction. Once $(G^L)_e$ is built and symmetrized, it costs an additional $|P|\binom{d+1}{2}$ multiply-add pairs to construct G_e . If we optimize the computation of contracting with $(G^L)_e$ first, we do not have to build G_e , but we must perform a dot product with w_k for each entry of the matrix. This costs |P| per contraction with $\binom{|P|+1}{2}$ entries in the matrix. If we optimize the contraction with each w_k first, then we have an additional $\binom{|P|+1}{2}$ contractions with $(G^L)_e$ at a cost of $\binom{d+1}{2}$ each. We expect that which of these will be most effective must be determined (automatically) on a case-by-case basis. Tables 10 and 9 show the comparisons for the first approach (labeled G_e), the second approach (labeled G_e) and the third approach (labeled G_e) by indicating the cost of the optimized computation plus the additional stages of computation. In most of these cases, contracting with the coefficient first leads to the lowest total cost.

5. Optimizing the optimization process

Since our graph (Y, E) is completely connected, we have $|E| = O(|Y|^2)$ and our optimization process requires complexity that is at least quadratic in the number of entries in the element stiffness matrix. In this section, we show how certain useful complexity-reducing relations may be evaluated over all of Y in better than quadratic time, then discuss how

Table 6. Number of multiply-add pairs in the optimized algorithm for computing the weighted Laplacian element stiffness matrix on triangles and tetrahedra for Lagrange polynomials of degree one through three using symmetry.

		t	riang	gles		tetrahedra						
ſ	degree	n	m	nm	MAPs	degree	n	m	nm	MAPs		
	1	6	9	54	27	1	10	24	240	108		
	2	21	18	378	218	2	55	60	3300	1650		
	3	55	30	1650	1110	3	210	120	25200	14334		

TABLE 7. Number of multiply-add pairs in the optimized algorithm for performing all of the contractions with $(G^L)_e$ in the weighted Laplacian on triangles and tetrahedra first, resulting in $\binom{|P|+1}{2}$ arrays of length |P| to contract with w_k .

		tr	iang	gles		tetrahedra						
	degree	n	m	nm	MAPs	degree	n	m	nm	MAPs		
Ī	1	18	3	54	9	1	40	6	240	27		
	2	126	3	378	115	2	550	6	3300	693		
	3	550	3	1650	683	3	4200	6	25200	7021		

TABLE 8. Number of multiply-add pairs in the optimized algorithm for performing all of the contractions with the coefficient in the weighted Laplacian on triangles and tetrahedra first, resulting in $\binom{|P|+1}{2}$ arrays of length $\binom{d+1}{2} = 6$ to contract with $(G^L)_e$.

		tr	iang	les		tetrahedra						
deg	ree	n	m	nm	MAPs	degree	n	m	nm	MAPs		
1		18	3	54	7	1	60	4	240	9		
2		63	6	378	138	2	330	10	3300	465		
3		165	10	1650	899	3	1260	20	25200	7728		

TABLE 9. Comparing the total number of multiply-add pairs for fully forming G_e , contracting with $(G^L)_e$ first, and contracting with w_k first on triangles.

		G_e			$(G^L)_e$ first		w_k first			
degree	MST	additional	total	MST	additional	total	MST	additional	total	
1	27	3*3	38	9	6*3	27	7	6*3	25	
2	218	3*6	236	115	21*6	241	138	21*3	201	
3	1110	3*10	1140	683	55*10	1233	899	55*3	1064	

TABLE 10. Comparing the total number of multiply-add pairs for fully forming G_e , contracting with $(G^L)_e$ first, and contracting with w_k first on tetrahedra.

		G_e			$(G^L)_e$ first		w_k first			
degree	MST	additional	total	MST	additional	total	MST	additional	total	
1	108	6*4	132	27	10*4	67	9	10*6	69	
2	1650	6*10	1710	693	55*10	1234	465	55*6	795	
3	14334	6*20	14454	7021	210*20	11221	7728	210*6	8988	

we may build a sparse graph (Y, E') with |E| = O(|Y|) that will admit a much more efficient optimization process. Even though this process must be run only once per form and element (say the Laplacian with quadratics on triangles), the quadratic algorithm can become very time consuming and challenge a single computer's resources for forms of high arity using high degree polynomials on tetrahedra.

5.1. **Search algorithms.** Before discussing how we may evaluate some of these complexity-reducing relations over the collection Y in better than quadratic time, we first describe some basic notation we will use for hash tables throughout this section and the next.

Hash tables are standard data structures [14] that associate each member of a set of keys to some value, possibly drawn from another set of objects. The important point about hash tables is that the basic operations of setting and getting values are expected to be independent of the number of entries in the table (expected constant time access). Many higher-level programming languages have library support or built-in features supporting hash tables (many implementations of the standard template library in C++ include hash tables, and scripting languages such as Python and Perl have them built in as primitive types).

We begin by establishing some notation for the basic operations we use. If a is a key of table T, then we find the value associated with a by T[a]. If there is no value associated with a (that is, if a is not a key of T, we may update T by adding a key a associated with value b by the notation $T[a] \leftarrow b$. We use the same notation to indicate setting a new value to an existing key.

As before, we label the vectors y^i for $1 \le i \le n$. We want to partition the labels into a set of subsets \mathcal{E} such that for each $E \in \mathcal{E}$, the vectors associated with each label in E are equal. Moreover, if two vectors are equal, then their labels must belong to the same E. This is easily accomplished by setting up a hash table whose keys are vectors and whose values are subsets of the integers $1 \le i \le n$. This process is described in Algorithm 1

Floating point arithmetic presents a slight challenge to hashing. Numbers which are close together (within some tolerance) that should be treated as equal must be rounded to so that they are indeed equal. Hashing relies on a function that maps items into a set of integers (the "hash code"). These functions are discontinuous and sensitive to small perturbations. For most numerical algorithms in floating point arithmetic, we may define equality to be "near equality", but hash tables require us to round or use rational arithmetic

Algorithm 1 Determining equality among vectors

```
E an empty table mapping vectors to subsets of \{i\}_{i=1}^n. for all 1 \leq i \leq n do
  if y^i is a key of E then
  E[y^i] \leftarrow E[v^i] \cup \{i\}
else if -y^i is a key of E then
  E[-y^i] \leftarrow E[-y^i] \cup \{i\}
else
  E[y^i] \leftarrow \{i\}
end if
end for
```

before any comparisons are made. We have successfully implemented our algorithms in both cases.

As an alternative to hashing, one could form a binary search tree or sort the vectors by a lexicographic ordering. These would rely on a more standard "close to equal" comparison operation, but only run in $O(mn \log{(mn)})$ time. So, for large enough data sets, hashing will be more efficient.

We may similarly partition the labels into a set of subsets C such that for each $C \in C$, the vectors associated with the labels in C are colinear. Similarly, if two vectors are colinear, then their labels must belong to the same C. This process may be performed by constructing the collection of unit vectors \hat{A} , with $\hat{y}_i = \frac{y_i}{\|y_i\|}$ for each $1 \le i \le n$ for some norm $\|\cdot\|$ and using Algorithm 1 on \hat{A} .

Finding vectors that are close together in Hamming distance is more subtle. At worst, the cost is $O(mn^2)$, as we have to compare every entry of every pair of vectors. However, we may do this in expected linear time with some assumptions about Y. We first describe the algorithm, then state the conditions under which the algorithm performs in worse than linear time.

Our vectors each have m components. We start by forming m empty hash tables. Each H_i will map numbers that appear in the i^{th} position of any vector to the labels of vectors that have that entry in that position. This is presented in Algorithm 2, in which y_j^i denotes the j^{th} entry of y^i .

This process runs in expected O(nm) time. From these tables, we can construct a table that gives the Hamming distance between any two vectors, as seen in Algorithm 3. This algorithm counts down from d each time it discovers an entry that two vectors share. Our algorithm reflects the symmetry of the Hamming distance.

On output, for $1 \le i < j \le n$, if D[i][j] has no entry, the distance between v^i and v_j is m. Otherwise, the table contains the Hamming distance between the two vectors.

Regarding complexity, there is a double loop over the entries of each H_i . Hence, the algorithm is quadratic in the maximum number of vectors that share the same entry at the same location. Presumably, this is considerably less than n on most data sets.

Algorithm 2 Mapping unique entries at each position to vectors containing them

```
for all 1 \leq i \leq d do H_i an empty table mapping numbers to sets of vector labels from \{i\}_{i=1}^n end for for all 1 \leq i \leq n do for all 1 \leq j \leq m do if y_j^i is a key of H_j then H_j[y_j^i] \leftarrow H_j[y_j^i] \cup \{i\} else H_j[y_j^i] := \{i\} end if end for end for
```

Algorithm 3 Computing Hamming distances efficiently

```
D an empty table
for all 1 \le i \le n do
   D[i] an empty table
end for
for all 1 \le i \le m do
  for all a in the keys of H_i do
     for all unique combinations k, \ell of labels in H_i[a] do
        \alpha := \min(k, \ell), \beta := \max(k, \ell)
        if D[\alpha] has a key \beta then
           D[\alpha][\beta] \leftarrow D[\alpha][\beta] - 1
        else
           D[\alpha][\beta] := m - 1
        end if
     end for
  end for
end for
```

5.2. Using a sparse graph. If we create a graph (Y, \tilde{E}) with significantly fewer edges than (Y, E), we may be able to get most of the reduction in operation count while having a more efficient optimization process. For example, we might choose to put a cutoff on ρ , only using edges that have a large enough complexity-reduction. So, we can define the set of edges being

(5.1)
$$E_k = \{ \{ y^i, y^j \} : \rho(y^i, y^j) \le k \}$$

For example, if we use Algorithms 2 and 3 to evaluate H^+ over all pairs from Y, then we are using $\rho = H^+$ and k = m - 1. Also, note that our structure D encodes a sparse graph. That is, the vectors of Y are the nodes of the graph, the neighbors of each y^i are

simply the keys of $D[y^i]$, and the edge weight between some y^i and neighbor y^j is $D[y^i][j^j]$. It is not hard to move from this table into whatever data structure is used for graphs.

Then, we could add colinearity or other complexity-reducing relations to this graph. If we use Algorithm 1 on the unit vectors to determine sets of colinear vectors, we can update the the graph by either adding edges or updating the edge weights for each pair of colinear vectors.

If $|E_k| = O(|Y|)$, then computing a minimum spanning tree will require only $O(n \log n)$ time rather than $O(n^2 \log n)$. However, there is no guarantee that (Y, E_k) will be a connected graph. Some vectors might not have close neighbors, or else some subgraphs do not connect with each other. An optimized algorithm can still be obtained by finding the connected components of the (Y, E_k) and finding a minimum spanning tree for each component. Then, the total cost of the computation is m times the number of connected components plus the sum of the weights of each of the minimum spanning trees.

6. Conclusion and ongoing work

We have developed a general optimization strategy for the evaluation of local stiffness matrices for finite elements. This is based on first formulating the computation as a sequence of tensor contractions, then introducing a new concept of complexity-reducing relations that allows us to set the optimization in a graph context. The optimization itself proceeds by computing a minimum spanning tree. These techniques worked very well at reducing the cost of evaluating finite element matrices for several forms using Lagrange elements of degrees one through three on triangles and tetrahedra. Finally, we discussed some efficient algorithms for detecting equality and colinearity and for evaluating the pairwise Hamming distance over the entire set of tensors.

In [11], we saw that frequently, some of the tensors will be linear combinations of two or more other tensors. However, both locating and making use of such relations in a more formal context has been difficult. We are working on geometric search algorithms to locate linear dependencies efficiently. However, once they are located, our optimization process must occur over a hypergraph rather than a graph. Finding a true minimum is also much more difficult, and we are working on heuristics that will allow us to combine these two approaches.

Finally, we plan to integrate our optimization strategy with FFC. While FFC currently generates very efficient code for evaluating variational forms, we will improve upon this generated code by piping the tensors through our optimization process before generating code to perform the contractions. This will lead to a domain-specific optimizing compiler for finite elements; by exploiting latent mathematical structure, we will automatically generate more efficient algorithms for finite elements than people write by hand.

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