# Imperial College London Department of Computing

# Automated Optimization of Numerical Methods for Partial Differential Equations

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# **Declaration**

I herewith certify that all material in this dissertation which is not my own work has been properly acknowledged.

Fabio Luporini

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# Chapter 1

## Introduction

#### 1.1 Thesis Statement

#### 1.2 Overview

In many fields, such as computational fluid dynamics, computational electromagnetics and structural mechanics, phenomena are modelled by partial differential equations (PDEs). Unstructured meshes, which allow an accurate representation of complex geometries, are often used to discretize their computational domain. Numerical techniques, like the finite volume method and the finite element method, approximate the solution of a PDE by applying suitable numerical operations, or kernels, to the various entities of the unstructured mesh, such as edges, vertices, or cells. On standard clusters of multicores, typically, a kernel is executed sequentially by a thread, while parallelism is achieved by partitioning the mesh and assigning each partition to a different node or thread. Such an execution model, with minor variations, is adopted, for instance, in Markall et al. [2013], Logg et al. [2012], AMCG [2010], DeVito et al. [2011], which are examples of frameworks specifically thought for writing numerical methods for PDEs.

The time required to execute these unstructured-mesh-based applications is a fundamental issue. An equation domain needs to be discretized into an extremely large number of cells to obtain a satisfactory approximation of the solution, possibly of the order of trillions (e.g. Rossinelli et al. [2013]), so applying numerical kernels all over the mesh is expensive. For

example, it is well-established that mesh resolution is crucial in the accuracy of numerical weather forecasts; however, operational centers have a strict time limit in which to produce a forecast - 60 minutes in the case of the UK Met Office - so, executing computation- and memory-efficient kernels has a direct scientific payoff in higher resolution, and therefore more accurate predictions. Motivated by this and analogous scenarios, this thesis studies, formalizes, and implements a number of code transformations to improve the performance of real-world scientific applications using numerical methods over unstructured meshes.

#### 1.3 Contributions

#### 1.4 Dissemination

The research exposed in this thesis has been disseminated in the scientific community through various channels:

- **Papers.** The following is the list of publications derived from the research activity (chronological order):
  - Strout, M.M.; Luporini, F.; Krieger, C.D.; Bertolli, C.; Bercea, G.-T.; Olschanowsky, C.; Ramanujam, J.; Kelly, P.H.J., "Generalizing Run-Time Tiling with the Loop Chain Abstraction," Parallel and Distributed Processing Symposium, 2014 IEEE 28th International, vol., no., pp.1136,1145, 19-23 May 2014
  - Fabio Luporini, Ana Lucia Varbanescu, Florian Rathgeber, Gheorghe-Teodor Bercea, J. Ramanujam, David A. Ham, and Paul H. J. Kelly. "Cross-loop optimization of arithmetic intensity for finite element local assembly". 2014. Submitted for publication.
  - Fabio Luporini, David A. Ham, Paul H. J. Kelly. "Optimizing Automated Finite Element Integration through Expression Rewriting and Code Specialization". 2014. To be written.
- Talks. Talks have been delivered at the following conferences/workshops:
  - "Generalised Sparse Tiling for Unstructured Mesh Computations in the OP2 Framework". Compilers for Parallel Computing, July

2013.

- 2. "COFFEE: an Optimizing Compiler for Fintie Element Local Assembly". FEniCS Workshop, July 2014.
- **Software.** The following software is released under open source licenses.
  - 1. COFFEE (COmpiler For Finit Element local assEmbly), the compiler described in Chapter 6.

, and the design of this software and results have been disseminated in the scientific community through publications.

## 1.5 Thesis Outline

# Chapter 2

# Background

#### 2.1 The Finite Element Method

Computational methods based upon finite elements are used to approximate the solution of partial differential equations (henceforth, PDEs) in a wide variety of domains. The mathematical abstraction used in finite element methods (or FEMs) is extremely powerful: not only does it help reasoning about the problem, but also provides systematic ways of deriving effective computer implementations. In ?, it is usefully suggested to consider an FEM as a black box that, given a differential equation, returns a discrete algorithm capable of approximating the equation solution. Unveiling the whole magic of such a black box is clearly out of the scope of this chapter. We rather limit ourselves to review the mathematical properties and the computational aspects that are essential for understanding the contributions in Chapters 4 and 5. The content and the notation used in this section are inspired from ???. For a complete treatment of the subject, the reader is invited to refer to ?.

#### 2.1.1 Variational Formulation

We consider the weak formulation of a linear variational problem

Find 
$$u \in U$$
 such that  $a(u,v) = L(v), \forall v \in V$  (2.1)

where a and L are, respectively, a bilinear form and a linear form. The term "variational" stems from the fact that the function v can vary arbitrarily. The unfamiliar reader may find this expression unusual. Understanding the actual meaning of this formulation is far beyond the goals of this review, since an entire course in functional analysis would be needed. Informally, we can think of V as a "very nice" space, in which functions have desirable properties. The underlying idea of the variational formulation is to "transfer" certain requirements (e.g., differentiability) from the unknown  $u \in U$  to  $v \in V$ .

The sets *U* and *V* and called, respectively, trial and test functions. The variational problem is discretized by using discrete test and trial spaces

Find 
$$u_h \in U_h \subset U$$
 such that  $a(u_h, v_h) = L(v_h), \forall v_h \in V_h \subset V$  (2.2)

Let  $\{\psi_i\}_{i=1}^N$  be the set of basis functions spanning  $U_h$  and let  $\{\phi_j\}_{j=1}^N$  be the set of basis functions spanning  $V_h$ . Then the unknown solution u can be approximated as a linear combination of the basis functions  $\{\psi_i\}_{i=1}^N$ ,

$$u_h = \sum_{j=1}^{N} U_j \psi_j. {(2.3)}$$

This allows us to rewrite 2.2 as:

$$\sum_{j=1}^{N} U_{j} a(\psi_{j}, \phi_{i}) = L(\phi_{i}), \ i = 1, 2, ..., N$$
(2.4)

From the solution of the following linear system we determine the set of degrees of freedom U to express  $u_h$ :

$$Au = b (2.5)$$

where clearly

$$A_{ij} = a(\phi_i(x), \phi_j(x))$$

$$b_i = L(\phi_i(x))$$
(2.6)

The matrix A and the vector b can be seen as the discrete operators arising from the bilinear form a and from the linear form L for the given choice of basis functions.

The variational formulation of a *non-linear variational problem* requires refinements that are out of the scope of this review. The interested reader should refer to ?.

#### 2.1.2 Finite Elements

In an FEM the domain  $\Omega$  of the PDE is partitioned into a finite set of disjoint cells  $\{K\}$ ; that is,  $\bigcup K = \Omega$  and  $\bigcap K = \emptyset$ . This forms a *mesh*. A finite element is usually defined (see for example ?) as a triple  $\langle K, \mathcal{P}_K, \mathcal{L}_K \rangle$ , where:

- *K* is a cell in the mesh with non-empty interior and piecewise smooth boundary;
- $\mathcal{P}_K$  is a finite dimensional "local" function space of dimension  $n_K$ ;
- the set of degrees of freedom  $\mathcal{L}_K$  is a basis  $\{l_1^K, l_2^K, ..., l_{n_K}^K\}$  for  $\mathcal{P}'_K$ , the dual space of  $\mathcal{P}_K$ .

This definition allows to impose constraints on the set of basis functions  $\{\phi_1^K, \phi_2^K, ..., \phi_{n_K}^K\}$  spanning  $\mathcal{P}_K$ . For instance, to enforce a *nodal basis* for  $\mathcal{P}_K$  – a particularly useful property for expressing solutions in  $U_h$  – we can impose that the relationship

$$l_i^K(\phi_i^K) = \delta_{ij}, \ i, j = 1, 2, ..., n_K$$
 (2.7)

must be satisfied ( $\delta_{ij}$  is the Kronecker delta). This allows to express any  $v \in \mathcal{P}_K$  as

$$v = \sum_{i=1}^{n_K} l_i^K(v) \phi_i^K.$$
 (2.8)

Each linear functional in  $\mathcal{L}_K$  is used to evaluate one degree of freedom of v in terms of the chosen nodal basis. In other words, we can refer to both the coefficients U introduced in the previous section and  $\mathcal{L}_K$  as the degrees of freedom.

**Example:** the triangular Lagrange element As an example<sup>1</sup>, consider a triangular cell K and let  $\mathcal{P}_K$  be the space of polynomials of order 1 on K.  $\mathcal{L}_K$  may be the set of bounded linear functionals representing point evaluation at the vertices  $x^i$  (i = 1, 2, 3) of K such that

$$l_i^K : \mathcal{P}^K \to \mathbb{R}$$

$$l_i^K(v) = v(x^i)$$
(2.9)

Since if v is zero at each vertex then v must be zero everywhere,  $\mathcal{L}_K$  really is a basis for  $\mathcal{P}'_K$ , so what we have defined is indeed a finite element. In particular, if we take  $x^1 = (0,0)$ ,  $x^2 = (1,0)$ ,  $x^3 = (0,1)$ , we have that the nodal basis is given by:

$$\phi_1(x) = 1 - x_1 - x_2, \ \phi_2(x) = x_1, \ \phi_3(x) = x_2.$$
 (2.10)

#### 2.1.3 Global Discretization

A *local-to-global mapping* allows to patch together the finite elements to form a global function space, for instance the set of trial functions  $U_h = \text{span}\{\psi_i\}_{i=1}^N$  introduced in Section 2.1.1. A local-to-global mapping is a function

$$\iota_K : [1, n_K] \to [1, N]$$
 (2.11)

that maps the local degrees of freedom  $\mathcal{L}_K$  to global degrees of freedom  $\mathcal{L}$ . The mappings  $\iota_K$ , together with the choice of  $\mathcal{L}_K$ , determine the continuity of a function space or, in simpler words, the continuity of a function throughout the domain  $\Omega$ . The reader is invited to refer to ? for a comprehensive yet simple description of this step.

One of the crucial aspects of an FEM is that global function spaces are often defined in terms of a *reference finite element*  $\langle \hat{K}, \hat{P}, \hat{\mathcal{L}} \rangle$  and a set of invertible mappings  $\{\mathcal{G}_K\}_K$  from  $\hat{K}$  to each cell in the mesh such that  $K = \mathcal{G}_K(\hat{K})$ . This situation is illustrated in Figure ??.

For each K,  $\mathcal{G}_K$  also allows to generate  $\mathcal{P}_K$  and  $\mathcal{L}_K$ . The complexity of this process depends on the mapping itself. In the simplest case, the

<sup>&</sup>lt;sup>1</sup>The example is extracted from ?

mapping is affine; that is, expressible as  $G_K(\hat{x}) = A_K \hat{x} + b_K$ , where  $A_K$  and  $b_K$  are, respectively, some matrix and vector.

#### 2.1.4 Assembly

The assembly of an FEM is the phase in which the matrix A and the vector b from 2.6 are constructed. This is accomplished by adding the contributions from each K to A and b. Let us consider the bilinear form a, and assume this is an integral over  $\Omega$ . Thanks to the linearity of the operator, we can express a as

$$a = \sum_{K} a_{K} \tag{2.12}$$

where  $a_K$  is an element bilinear form. We can then define the local element matrix

$$A_i^K = a_K(\psi_{i_1}^K, \phi_{i_2}^K) \tag{2.13}$$

where  $i \in \mathcal{I}_K$  is the index set on  $A_i^K$ . That is,  $\mathcal{I}_K = \{(1,1), (1,2), ..., (n_U, n_V)\}$ , with  $n_U$  and  $n_V$  representing the number of degrees of freedom for the local trial functions  $\psi^K \in U_h^K$  and the local test functions  $\phi^K \in V_h^K$ . The element matrix  $A^K$  is therefore a (typically dense) matrix of dimension  $n_U \times n_V$ .

Now let  $\iota_K^U$  and  $\iota_K^V$  be the local-to-global mappings for the local discrete function spaces  $U_h^K$  and  $V_h^K$ . We can define, for each K, the collective local-to-global mapping  $\iota_K : \mathcal{I}_K \to \mathcal{I}$  such that

$$\iota_K(i) = (\iota_K^U(i_1), \iota_K^V(i_2)) \ \forall i \in \mathcal{I}_K.$$
(2.14)

This simply maps a pair of local degrees of freedom to a pair of global degrees of freedom. Let also  $\mathcal{T}$  be the subset of the cells in the mesh in which  $\psi_{i_1}$  and  $\phi_{i_2}$  are both non-zero. Note that here we are talking about the global functions whose restrictions to K gives  $\psi_{i_1}^K$  and  $\phi_{i_2}^K$ . By construction,  $\iota_K$  is invertible if  $K \in \mathcal{T}$ . At this point, we have all the ingredients to compute A as the sum of local contributions from the cells

in the mesh:

$$A_{i} = \sum_{K \in \mathcal{T}} a_{K}(\psi_{i_{1}}, \phi_{i_{2}})$$

$$= \sum_{K \in \mathcal{T}} a_{K}(\psi_{(i_{K}^{U})^{-1}(i_{1})}^{K}, \phi_{(i_{K}^{V})^{-1}(i_{2})}^{K}) = \sum_{K \in \mathcal{T}A_{i_{r}}^{K}(i)}$$
(2.15)

Similar conclusions may be drawn for the linear form L. We observe that this computation can be implemented as a single iteration over all cells in the mesh. On each cell,  $A^K$  is computed and added to A using the inverse mappings. This approach is particularly efficient because it only evaluates the non-zero entries in the sparse matrix A. Other more trivial implementations of the assembly phase are possible, although rarely used in practice because ineffective.

We conclude with a clarification concerning the terminology. The assembly process is often naturally described as a two-step procedure: *local assembly* and *global assembly*. The former consists of computing the contributions of each single element (i.e., the element matrices  $A^K$ ); the latter represents the "coupling" of all  $A^K$  into A. As we shall see, one of the main subjects of this thesis is the computational optimization of the local assembly phase.

#### 2.1.5 Local Assembly Example: from Math to Code

Because of its relevance in this thesis, we illustrate local assembly in a concrete example, the evaluation of the element matrix for a Laplacian operator.

#### Specification of the Laplacian operator

Consider the weighted Laplace equation

$$-\nabla \cdot (w\nabla u) = 0 \tag{2.16}$$

in which u is unknown, while w is prescribed. The bilinear form asso-

ciated with the weak variational form of the equation is:

$$a(v,u) = \int_{\Omega} w \nabla v \cdot \nabla u \, dx \tag{2.17}$$

The domain  $\Omega$  of the equation is partitioned into a set of cells (elements) T such that  $\bigcup T = \Omega$  and  $\bigcap T = \emptyset$ . Assuming for simplicity that the sets of trial and test functions are the same and by defining  $\{\phi_i^K\}$  as the set of local basis functions spanning U and V on the element K, we can express the local element matrix as

$$A_{ij}^K = \int_K w \nabla \phi_i^K \cdot \nabla \phi_j^K \, \mathrm{d}x \tag{2.18}$$

The element vector *L* can be determined in an analogous way.

In this example, the element tensors are expressed as a single integral over the cell domain. In general, they are expressed as a sum of integrals over *K*, each integral being the product of derivatives of functions from sets of discrete spaces and, possibly, functions of some spatially varying coefficients. Such an integral is often called *monomial*.

#### Quadrature Mode

A quadrature scheme is typically used to numerically evaluate  $A_{ij}^K$ . It consists of evaluating an integral at a set of given *quadrature points*, each point multiplied with some suitable *quadrature weight*. By mapping the computation to a reference element as explained in Section 2.1.3 and using the same approach as in Section 2.1.4, a quadrature scheme for the Laplacian operator on K is as follows

$$A^{K} = \int_{K} w \nabla \phi_{i}^{K} \cdot \nabla \phi_{j}^{K}$$

$$\approx w \sum_{k=1}^{N_{q}} W_{k} \nabla \phi_{i}^{K}(\mathbf{x}^{k}) \cdot \nabla \phi_{i}^{K}(\mathbf{x}^{k}) |\det \mathcal{G}_{K}'(\mathbf{x}^{k})|,$$
(2.19)

where  $\{x^1, x^2, ..., x^{N_q} \text{ is the set of } N_q \text{ quadrature points, and } \{W_1, W_2, ..., W_{N_q}\}$  the corresponding set of quadrature weights scaled such that  $\sum_{k=1}^{N_q} W_k = |\hat{K}|$ .

To compute a local basis function  $\phi_i^K$  from a reference element basis

function  $\Phi_i$  we exploit the invertible map  $\mathcal{G}_K^{-1}$ , which allows us to write  $\phi_i^K = \Phi_i \circ \mathcal{G}_K^{-1}$ . To evaluate the gradient of a basis function  $\phi_i^K$  at a quadrature point  $x^k$ , with  $x^k = \mathcal{G}_K(X^k)$  and  $X^k \in \hat{K}$ , we therefore have to compute a matrix-vector product

$$\nabla_{\mathbf{x}} \phi_i^K(\mathbf{x}^k) = ((\mathcal{G}_K')^{-1})^T(\mathbf{x}^k) \nabla_X \Phi_i(\mathbf{X}^k). \tag{2.20}$$

The term  $\mathcal{G}_K'$ )<sup>-1</sup> represents the inverse of the Jacobian matrix originating from the change of coordinates. The resulting scalar-valued expression for each entry  $A_{ii}^K$ , assuming  $\Omega$  to be a domain of dimension d, reads as:

$$A_{ij}^{K} = \sum_{k=1}^{N_q} \sum_{\alpha_3=1}^{n} \phi_{\alpha_3}(\mathbf{X}^k) w_{\alpha_3} \sum_{\alpha_1=1}^{d} \sum_{\alpha_2=1}^{d} \sum_{\beta=1}^{d} \frac{\partial X_{\alpha_1}}{\partial x_{\beta}} \frac{\partial \phi_i^K(\mathbf{X}^k)}{\partial X_{\alpha_1}} \frac{\partial X_{\alpha_2}}{\partial x_{\beta}} \frac{\partial \phi_j^K(\mathbf{X}^k)}{\partial X_{\alpha_2}} \det \mathcal{G}_K' W^k.$$
(2.21)

#### **Tensor Contraction Mode**

Consider the case in which  $\mathcal{G}_K : \hat{K} \to K$  is an affine mapping. Starting from Equation 2.21 and exploiting linearity, associativity and distributivity of the involved mathematical operators, we can rewrite 2.21 as

$$A_{ij}^{K} = \sum_{\alpha_{1}=1}^{d} \sum_{\alpha_{2}=1}^{d} \sum_{\alpha_{3}=1}^{n} \det \mathcal{G}_{K}' w_{\alpha_{3}} \sum_{\beta=1}^{d} \frac{X_{\alpha_{1}}}{\partial x_{\beta}} \frac{\partial X_{\alpha_{2}}}{\partial x_{\beta}} \int_{K_{0}} \phi_{\alpha_{3}} \frac{\partial \phi_{i_{1}}}{\partial X_{\alpha_{1}}} \frac{\partial \phi_{i_{2}}}{\partial X_{\alpha_{2}}} dX.$$
 (2.22)

A generalization of this transformation has been proposed in ?. Because of only involving reference element terms, the integral in the equation can be pre-evaluated and stored in temporary variables. The evaluation of the local tensor can then be abstracted as

$$A_{ij}^{K} = \sum_{\alpha} A_{i_1 i_2 \alpha}^{0} G_K^{\alpha}$$
 (2.23)

in which the pre-evaluated "reference tensor"  $A_{i_1i_2\alpha}$  and the cell-dependent "geometry tensor"  $G_K^{\alpha}$  are exposed.

#### Qualitative comparison

Depending on form and discretization, the relative performance of the two modes, in terms of the operation count, can vary quite dramatically. The presence of derivatives or coefficient functions in the input form increases the rank of the geometry tensor, making the traditional quadrature mode preferable for "complex" forms. On the other hand, speed-ups from adopting tensor mode can be significant in a wide class of forms in which the geometry tensor remains "sufficiently small". The discretization, particularly the relative polynomial order of trial, test, and coefficient functions, also plays a key role in the resulting operation count.

These two modes have been implemented in the FEniCS Form Compiler?, which we review in later sections. In this compiler, a heuristic is used to choose the most suitable mode for a given form. It consists of analysing each monomial in the form, counting the number of derivatives and coefficient functions, and checking if this number is greater than a constant found empirically (Logg et al. [2012]). In Chapter 4, we will describe a code generation system that goes beyond the dichotomy between quadrature and tensor modes, relying on cost models comparing the impact of different rewrite operators (e.g., expansion of products, factorization).

#### Code Examples

A possible C implementation of Equation 2.21 is illustrated in Listing 1. We assume a domain of dimension d=2 and polynomial order 1 Lagrange elements. The values at the quadrature points of the derivatives of the basis functions are pre-tabulated in the B and D arrays (representing, respectively, the derivatives with respect to the coordinates x and y). The values at the quadrature points of the basis functions spanning the field w are pre-tabulated in the array C. Pre-tabulation, which is made possible by mapping the computation to a reference element, is of fundamental importance to speed-up the local assembly phase. The summation over the  $N_q=6$  quadrature points is implemented by the i loop. The r loop implements the summation over  $\alpha_3$  for discretization of the weight w. Given their small range, the summations over the spatial dimensions  $\alpha_1$ ,  $\alpha_2$  and  $\beta$  have all been expanded in the "assembly expression" that evaluates the

local element matrix *A*. The *K* array includes the four components of the inverse of the Jacobian matrix for the change of coordinates.

**LISTING 1:** A possible implementation of Equation 2.21 assuming a 2D triangular mesh and polynomial order 1 Lagrange basis functions.

```
void weighted_laplace(double A[3][3], double **coords, double w[3])
     // Compute Jacobian
     double J[4];
4
     compute_jacobian_triangle_2d(J, coords);
     // Compute Jacobian inverse and determinant
     double K[4];
     double detJ;
     compute_jacobian_inverse_triangle_2d(K, detJ, J);
10
     const double det = fabs(detJ);
11
12
13
     // Quadrature weights
     static const double W[6] = 0.5;
14
15
     // Basis functions
16
     static const double B[6][3] = \{\{...\}\}; static const double C[6][3] = \{\{...\}\}; static const double D[6][3] = \{\{...\}\};
17
18
19
20
21
     for (int i = 0; i < 6; ++i) {
22
        double f0 = 0.0;
        for (int r = 0; r < 3; ++r) {
23
          f0 += (w[r] * C[i][r]);
24
25
        for (int j = 0; j < 3; ++j) {
26
          for (int k = 0; k < 3; ++k) {
27
            A[j][k] += ((((K[1]*B[i][k]) + (K[3]*D[i][k])) *
28
29
                            ((K[1]*B[i][j]) + (K[3]*D[i][j]))) +
                          (((K[0]*B[i][k]) + (K[2]*D[i][k])) *
30
                            ((K[0]*B[i][j]) + (K[2]*D[i][j])))*det*W[i]*f0);
31
32
        }
33
34
     }
35
```

The evaluation of integrals becomes more computationally expensive if the complexity of a variational form grows, in terms of number of coefficients and tensor algebra or differential operators employed. It is not pathological a scenario in which the computation of a local element tensor requires more than hundreds or even thousands of floating point operations. An excerpt from one such example is shown in Listing 2: here, in the main expression, 14 unique arrays are accessed (with the same array referenced multiple times within the expression) along with several other constants. The loop trip counts are also larger due to the different domain discretization employed. As we shall see, automated code generation for

**LISTING 2:** Local assembly implementation for a Burgers problem on a 3D mesh using polynomial order q = 1 Lagrange basis functions.

```
void burgers(double A[12][12], double **coords, double **w)
2
     // Compute Jacobian
     double J[9];
     compute_jacobian_triangle_3d(J, coords);
     // Compute Jacobian inverse and determinant
     double K[9];
     double detJ;
     compute_jacobian_inverse_triangle_3d(K, detJ, J);
10
     const double det = fabs(detJ);
12
13
     // Quadrature weights
     static const double W[5] = {\ldots}
14
15
     // Basis functions
    static const double B[5][12] = \{\{...\}\}
static const double C[5][12] = \{\{...\}\}
17
18
     //11 other basis functions definitions
20
     for (int i = 0; i < 5; i++) {
21
       double f0 = 0.0;
       //10 other declarations (f1, f2,...)
23
24
       for (int r = 0; r < 12; r++) {
25
         f0 += (w[r][0]*C[i][r]);
26
         //10 analogous statements (f1, f2, ...)
27
28
29
30
       for (int j = 0; j < 12; j++) {
         for (int k = 0; k<12; k++) {
31
32
           A[j][k] += ..
      + ((K[5]*F9) + (K[8]*F10))*Y1[i][j]) + ... +
33
      + \ (((K[0]*C[i][k]) \ + \ (K[3]*D[i][k]) \ + \ (K[6]*E[i][k]))*Y2[i][j]))*f11) \ +
34
      + (((K[2]*C[i][k]) + (K[5]*D[i][k]) + (K[8]*E[i][k]))*((K[2]*E[i][j]) + ...))) +
36
      + <roughly a hundred sum/muls go here>)..)*det*W[i];
37
38
39
40
```

finite element operators has had two main objectives:

- relieving the implementation burden when it comes to translate into code non-trivial operators, a tedious, error-prone task;
- facilitating the introduction of techniques to reduce the operation count and powerful low-level optimizations.

The second point, as already anticipated, is one of the topics treated in this thesis.

#### 2.1.6 Linear Solvers

The last step of an FEM is the resolution of the linear system (2.5) arising from the variational form of the input problem. This and the assembly of *A* and *b* are the most expensive phases of an FEM. There is a whole science concerning the efficient resolution of linear systems. Among the most effective solvers are the family of *Krylov-type iteration methods*, such *conjugate gradient* for symmetric positive-definite matrices and *generalized minimal residual* (GMRES), which does not require *A* in explicit form. *Multi-grid* methods are also widely used, whereas *direct methods* computing an LU factorization using *Gaussian elimination* have limited applicability.

The resolution of linear systems is not one of the topics of this thesis, so the interested reader is invited to refer to ?. It is however important to keep in mind that this phase usually has a significant impact on the execution time of an FEM: the performance optimization of the assembly phase has marginal impact if the method is solver-dominated.

## 2.2 Abstractions in Computational Science

This thesis centres on performance optimizations for scientific codes targeting different layers of abstraction. In this section, we dive into such abstractions and review the state-of-the-art on established software. The objective is to provide a foundation upon which motivating and explaining the contributions in Chapters 3, 4 and 5.

#### 2.2.1 Automating the Finite Element Method

The need for rapid implementation of high performance, robust, and portable scientific simulations has led to approaches based on automated code generation. This has been proven extremely successful in the context of the FEniCS (Logg et al. [2012]) and Firedrake (?) projects, which target the finite element method. In these frameworks, the weak variational form of a problem is expressed at high level by means of a domain-specific language. The mathematical specification is then manipulated by a form compiler that generates a representation of the assembly operators. The representation is transformed for performance optimization and subsequently translated into C code. The generated code is compiled and executed over the mesh. This entire process occurs at run-time because both FEniCS and Firedrake are implemented in Python and rely on just-in-time code generation and compilation. When the operators are assembled, a linear system needs be solved. For this, the PETSc library? is employed. In the following, we describe the aspects of this tool-chain that are relevant for the following chapters. We focus on Firedrake, rather than FEniCS, because all algorithms and techniques developed in this thesis have been integrated with this framework.

#### **Problem Specification**

Firedrake uses a mathematical language called UFL, the *Unified Form Language*? UFL is unrelated to meshes, function spaces, and solvers. It only concerns with the variational formulation of a problem, by providing different kinds of operators – like grad (gradient) and inner (inner product), but also algebraic operators (e.g., transpose, inverse) and elementary functions (e.g., abs, sqrt). Before emitting a representation suitable for the underlying compiler, UFL analyzes the form to collect useful information and applies some preliminary transformations, including automatic differentiation.

A UFL representation of the weighted Laplace operator shown in (2.17) is given in Listing 3. When constructing a finite element, three pieces of information are specified: *family*, *cell*, and *polynomial degree*. The family allows varying the type of the element. UFL supports many types, ranging from the most standard *Lagrange* element to *Discontinuous Lagrange* 

**LISTING 3:** UFL specification of the weighted Laplace operator defined in (2.17). In orange the keywords of the language.

```
1 element = FiniteElement ('Lagrange', triangle, 1)
2
3 u = TrialFunction (element)
4 v = TestFunction (element)
5 w = Coefficient (element)
6
7 a = w*dot (grad(v), grad(u))*dx
```

and even mixed elements such as H(div) and H(curl). This allows solving problems with the most disparate requirements on continuity of the functions. A thorough description is provided in ?. The cell represents the shape of the reference element: possible values include triangle, quadrilateral and tetrahedron. The polynomial order drives the number of degrees of freedom in an element. Functions can also be vector-valued, in which case one must use the special construct VectorElement in place of FiniteElement.

We are not interested in a complete review of UFL, which would also require a knowledge of the finite element method that goes far beyond the tool-kit we need in this thesis. It is however worth appreciating the power, versatility and depth of this language, since it opened a new era in the development of computational methods.

#### Form Compilers

The UFL specification of a variational form is eventually passed to a form compiler, whose objective is to construct a representation of the assembly operators. The *FEniCS Form Compiler*, or FFC, was originally adapted and used in Firedrake for this task. FFC, which supports both quadrature and tensor modes (see Section 2.1.5), used to directly emit C code. It was later modified (one of the initial steps of this thesis) to rather output abstract syntax trees, a structure allowing further manipulation in the lower layers of the stack. More recently, FFC has been supplanted by a new system, the *Two-Stage Form Compiler*, or TSFC. Just like FFC, TSFC emits abstract syntax trees (ASTs). The optimizations described in Chapters 4 and 5 are implemented by manipulating the AST representation in a lower-level compiler, COFFEE (outlined in Chapter 6).

TSFC has two main features:

- It is a *structure-preserving compiler* in that it keeps intact the structure of algebraic operations (e.g., index sums, inner products), rather than committing to a specific implementation. This lets the lower-level compiler to explore the space of all possible transformations.
- As opposed to FFC, it supports the compilation of complicated forms
  making extensive use of tensor algebra. TSFC can efficiently identify
  repeated sub-expressions and assign them to temporary variables, thus
  drastically reducing the code generation time.

We observe that in Firedrake there is a neat separation of concerns:

- UFL is the mathematical language that allows expressing finite element problems.
- TSFC progressively abstracts away the domain knowledge and transforms the input into an AST. Some nodes in the ASTs are decorated to keep track of properties (e.g., linearity of an expression in a given set of symbols) exploitable for later optimization.
- COFFEE uses several rewrite operators for reducing the operation count of the operators returned by TSFC.

The COFFEE layer, design and optimization algorithms constitute one of the main contributions of this thesis. Chapters 4, 5 and 6 are entirely devoted to these topics.

#### Iteration over the Mesh

Finite element problems require the application of computational *kernels* over the discretized equation domain. In Firedrake, this is accomplished through *PyOP2* Markall et al. [2013], a domain specific language embedded in Python relying on just-in-time compilation and execution.

Typical kernels are the assembly operators presented in Section 2.1.4, usually the most expensive from a computational viewpoint. Other relevant examples are the application of boundary conditions as well as the interpolation and projection of fields. The fact that a kernel is a local operation – its application on an element of the mesh is independent of the application on any another element – suits naturally parallel execution. This does not mean, however, that parallelizing a finite element computa-

tion is straightforward. As clarified in Section 2.2.2, a kernel can update a field either directly or indirectly. In the latter case, a subset of values, for instance the degrees of freedom at the boundary between two elements, may need be incremented by two different processes/threads, which requires non-trivial coordination.

PyOP2 supports the parallel application of kernels over unstructured meshes, a key requirement for FEMs. It also provides global data structures such as sparse matrices, which are essential when it comes to solve linear systems as 2.5.

This abstraction implements another separation of concerns. The kernels, which encode the numerics of an FEM, are produced at layers above PyOP2, usually in a completely automated fashion by Firedrake and the form compilers themselves. On the other hand, the parallel execution is completely masked by PyOP2.

The PyOP2 layer, and its relationship with the OP2 library Giles et al. [2011], are documented in mode detail in Section 2.2.2.

#### **Unstructured Meshes**

PyOP2 only works with sets and fixed-arity maps. Possible sets are topological entities such as cells or degrees of freedom. A map is an object describing the adjacency relationship between the elements of two distinct sets (e.g., a map from cells to degrees of freedom). PyOP2 has therefore no concept of what a mesh is or how it can be constructed. This task, in Firedrake, is carried out by another software module, PETSc's *DMPlex*?. The maps required by PyOP2 are then directly derived from DMPlex.

DMPlex is a data management abstraction representing unstructured mesh data through direct acyclic graphs. DMPlex relieves PyOP2 from the duty of handling two complex operations: partitioning for distributed parallel execution and reordering for efficient memory accesses. Moreover, the flexibility of the abstraction allows implementing techniques for communication-computation overlap, for instance via redundant computation along partition boundaries.

In this thesis, we will exploit the versatility of DMPlex for implementing automated loop fusion (Chapter 3).

#### Solving systems of linear equations

As we have seen, one of the key design principles characterizing Firedrake is exploiting successful, available software to carry out specific tasks of an FEM. This philosophy is also applied for what concerns the resolution of systems of linear equation, for which the *Portable, Extensible Toolkit for Scientific Computation* library?, or simply PETSc, is employed.

PETSc is entirely implemented in C, although its Python interface *petsc4py* makes it easily usable by frameworks such as Firedrake. It provides a wide range of algorithms for solving linear systems as well as a considerable number of options to drive the solvers. All algorithms are parallelized for distributed-memory execution through MPI. All these aspects probably make PETSc the most powerful tool when it comes to deal with expensive linear algebra operations.

Similarly to Firedrake, PETSc never attempts to reinvent science: many functionalities are implemented on top of existing libraries (e.g., BLAS) or offered via third-party implementations through suitable wrappers.

#### 2.2.2 The PyOP2 and OP2 Libraries

PyOP2 is inspired from and shares many ideas with OP2<sup>2</sup> Giles et al. [2011], although it differs in a few yet significant ways. In this section, we first describe the main features of the abstraction and the implementation that are in common; we then conclude explaining how PyOP2 detaches itself from OP2.

#### Programming Model

OP2 offers abstractions for modeling an unstructured mesh, in terms of *sets* (e.g. vertices, edges), and mapping between them (e.g. mapping between edges and vertices) that express the mesh connectivity. An OP2 dataset associates data to each element of a mesh set (e.g. 3D coordinates).

OP2 programs, such as the one in Listing ??, are expressed as sequences of parallel loops, each loop applying a computational kernel to every element in a mesh set. These *kernels* can access data associated to either the

<sup>&</sup>lt;sup>2</sup>The name OP2 indicates that this is the second software engineering iteration of the OPlus library, or Oxford Parallel Library.

#### LISTING 4: Section of a toy OP2 program.

```
void kernel1 (double * x, double * tmp1, double * tmp2) {
     *tmp1 += *x;
2
     *tmp2 += *x;
3
4 }
5
6 // loop over edges
7 op_par_loop (edges, kernel1,
               op_arg_dat (x, -1, OP_ID, OP_READ).
               op_arg_dat (temp, 0, edges2vertices, OP_INC),
               op_arg_dat (temp, 1, edges2vertices, OP_INC))
10
11
12 // loop over cells
13 op_par_loop (cells, kernel2,
               op_arg_dat (temp, 0, cells2vertices, OP_INC),
               op_arg_dat (temp, 1, cells2vertices, OP_INC),
15
               op_arg_dat (temp, 2, cells2vertices, OP_INC),
16
               op_arg_dat (res, -1, OP_ID, OP_READ))
17
18
19 // loop over edges
20 op_par_loop (edges, kernel3,
               op_arg_dat (temp, 0, edges2vertices, OP_INC),
21
               op_arg_dat (temp, 1, edges2vertices, OP_INC))
```

loop iteration set (direct access) or to other sets (indirect access) through mappings (or OP2 maps). As an example, a parallel loop iterates over all edges of the mesh and the kernel increments a dataset associated to vertices (i.e. each edge updates its two vertices). OP2 maps are implemented as arrays of indices.

The running example including three parallel loops. The first loop iterates over edges, as indicated by the first parameter of the *op\_par\_loop* function. In OP2, an iteration space (edges, in this case) is expressed as an op\_set structured type, whose declaration is omitted in the example. The op\_par\_loop applies a kernel "kernel1" to every element in the indicated iteration space, i.e. to each edge. The kernel of this example reads data associated to an edge and increments the two adjacent vertices with the read value. This is indicated by the OP\_READ and OP\_INC modes in the access descriptors.

The access descriptors (or op\_arg\_dat) are used to indicate which datasets are being accessed when iterating on an OP2 set. When the access descriptor indicates OP\_ID for a data array, then that data array is being accessed with the identity function. The second and third op\_args in the example use indirections. Assuming that *temp* is a dataset associated to vertices, we need to tell OP2 what vertex data should be passed as second and third

parameters (i.e. what fields of *temp* are required for each invocation). The index array *edges2vertices* maps each edge index into the indices of its two incident vertices with the integer values of 0 and 1 indicating which of the two vertices should be considered.

#### **Execution Model for Shared-Memory Parallelism**

The execution order of parallel loop iterations does not influence the result. The best ordering scheme, therefore, can be chosen by the run-time implementation. For shared-memory-parallelism, in case of indirect increments/reductions (as in the example), OP2 guarantees data race avoidance by scheduling serially (in any order) iterations incrementing the same variable.

Data races are avoided by partitioning the vertices in the underlying mesh. Partitions that share an iteration set element (e.g. edge, cell) access shared vertex data and, therefore, are considered conflicting partitions. The partition conflict graph is eventually colored to enforce serialisation of increments to shared data. Partitions assigned the same color can be executed in parallel by different threads. On the other hand, partitions of different colors are scheduled serially, to prevent race conditions. The execution of elements inside a partition is serialized by scheduling them to a single thread.

Partitioning is performed on a per-loop basis. Each loop iteration set is partitioned by creating a dependency list between iterations using one of the maps used to access data in the loop. The built adjacency list can optionally be passed to a partitioning algorithm (METIS and PT-Scotch have been demonstrated for use in OP2) to maximize iteration locality according to their data access pattern. In other words, iterations accessing the same data will be more likely to be assigned to the same partition rather than independent iterations.

#### **Execution Model for Distributed-Memory Parallelism**

Distributed-memory parallelism is conceptually simpler than shared-memory parallelism. During the OP2 initialization phase, sets, maps, and datasets are partitioned and then distributed to different processes. For executing a parallel loop, MPI messages are usually exchanged to refresh the out-

of-date values along the partition boundaries. In overlap, each process computes its own portion of "local" iterations. Once both phases have finished, a process can correctly execute the remaining boundary iterations.

To implement this parallelization scheme, the iterations of each locally stored OP2 set are divided into four contiguous regions:

**Core** Iterations owned that can be processed without reading halo data.

Owned Iterations owned that can be processed only by reading halo data.

**Exec halo** Off-process iterations that are redundant executed because they indirectly increment owned iterations.

**Non-exec halo** Off-process iterations that only need be read to compute the exec halo iterations.

This situation is depicted in Figure ??. Clearly, a good partitioning maximizes the ratio between the sizes of the core and non-core regions.

#### **PyOP2 Features**

PyOP2 distinguishes itself from OP2 in a number of different ways.

...

#### 2.2.3 Stencil Languages

...

## 2.3 Compilers and Libraries for Loop Optimization

#### 2.3.1 Loop Optimization - Static Analysis

Polyhedral compilers ... scheduling functions Mention their unsuitability for tiling unstructured meshes... (use email I sent listing all issues...)

#### 2.3.2 Loop Optimization - Dynamic Analysis

- Inspector/Executor schemes
- Space filling curves for unstructured meshes
- sparse tiling = loop fusion + loop tiling

2.3.3 On the Dichotomy between Loop Tiling and Loop Fusion	2.3.3	On the	Dichotomy	between	Loop	Tiling	and	Loop	Fusio	01
--	-------	--------	-----------	---------	------	--------	-----	------	-------	----

• Tiling is for a nested loop!
• tiles are atomic blabla
• split vs overlapped
Split tiling
Overlapped tiling
Fusion vs Tiling
so time tiling is just fusion across the time stepping loop, then tiling
2.4 Domain Specific Optimization
Tensor Contraction Engine
LGen
Why potentially useful
Halide
Spiral
opiiui –
<b></b>
2.5 State-of-the-art Hardware Architectures
2.0 State of the art Haraware Michilectures

#### 2.5.1 SIMD Vectorization

...

### 2.5.2 Terminology

Memory pressure, Register pressure

**Arithmetic intensity** 

**Operational intensity** Maybe add a subsection of the roofline model?

Flops

Access function (for array)

General-purpose compiler

Communication vs computation

**CPU- vs Memory-bound** 

Local vs Global reduction

# Chapter 3

# Automated Loop Fusion and Tiling for Irregular Computations

#### 3.1 Motivation

Many numerical methods for partial differential equations (PDEs) are structured as sequences of parallel loops. This exposes parallelism well, but does not convert data reuse between loops into data locality, since large datasets are usually accessed. In Section 2.3.1 we have explained that loop fusion and loop tiling may be used to retain some of this potential data locality. As we elaborate in the upcoming sections, however, it is extremely challenging to implement these optimizations in most real-world applications.

Our focus is on unstructured mesh PDE solvers, like those based on the finite volume or the finite element methods. Here, the loop-to-loop dependence structure is data-dependent due to indirect references such as A[map[i]]; the map array stores connectivity information, for example from elements in the mesh to degrees of freedom. A similar pattern occurs in molecular dynamics simulations and graph processing, so both the theory and the tools that we will develop in this chapter are generalizable to these domains.

Because of the irregular memory access pattern, our approach to loop transformation is based on dynamic analysis, particularly on *inspector/ex-*

ecutor schemes. Our hypothesis, backed by the studies reviewed in Section 2.3.2, is that dynamic loop optimization can improve the performance of a class of real-world unstructured mesh applications. Among the possible dynamic loop optimizations, we target *sparse tiling*. We recall from Section 2.3.3 that sparse tiling aims to exploit data reuse across consecutive loops by composing three transformations: loop fusion, loop tiling, and automatic parallelization.

The three main issues that we tackle in this chapter are:

- Previous approaches to sparse tiling were all based upon "ad-hoc" inspector/executor strategies; that is, developed "by hand", per application. We seek for a general technique, applicable to arbitrary computations on unstructured meshes.
- Automation is more than a desired feature because application specialists tend to avoid complicated optimizations that harm source code comprehensibility. We therefore aim for a fully-automated framework, based upon a mixed compiler/library approach.
- Very few studies have tackled fusion when loops are interleaved by routines for message passing. We are aware of none for the case in which the memory accesses pattern is irregular, as in unstructured mesh applications. We describe and implement a technique that solves this problem. This is a fundamental contribution because most scientific simulations run on multi-node architectures.

## 3.2 Open Problems, Questions, Hypotheses

Loop fusion and loop tiling have been widely studied in the literature. However, it is not clear to what extent they are actually employed in real-world applications. Most studies centre their experimentation on relatively simple benchmarks and single-node performance. This unfortunately does not expose the complexity and the limitations of scientific codes. On the other hand, it has repeatedly been shown that applying these transformations to "simple" memory-bound loop nests can result in considerable speed-ups. Some of the most notable examples are stencil codes arising in finite difference methods [Zumbusch, 2013, Holewinski et al., 2012, Bondhugula et al., 2014], linear algebra routines [Buttari

et al., 2008, 2009], and image processing kernels [Ragan-Kelley et al., 2013]. Since numerical methods for partial differential equations (PDEs) are often structured as sequences of parallelizable loops, or "sweeps", over the discretized equation domain, the following questions arise naturally:

**Applicability** Can we adopt sparse tiling (i.e., fusion and tiling) in real-life numerical methods for solving PDEs? If so, what kind of gain in performance should we expect?

**Lack of evidence** Why, despite decades of research, loop transformations are rarely used in scientific (production) codes?

**Challenges** What are the theoretical and technical challenges that we have to overcome to automate sparse tiling, thus making it available to wider audiences?

We tackle these questions in the following context:

**Irregular codes** Unstructured meshes are often used to discretize the computational domain, since they allow for an accurate representation of complex geometries. Their connectivity is stored by means of adjacency lists (or equivalent data structure). This leads to indirect memory accesses (e.g., A[B[i]]) within loop nests. Indirections break static analysis, thus making compiler-based approaches to loop transformation (e.g., polyhedral optimization) unsuitable for our context. Runtime data dependence analysis enables dynamic loop optimization, although at the price of additional overhead.

**Realistic datasets** Complex simulations usually operate on at least gigabytes of data, requiring multi-node execution. Any loop transformation we consider will have to co-exists with message passing for distributed-memory parallelism.

**Automation, but no legacy code** Sparse tiling is an "extreme optimization". It requires a great deal of effort to be implemented because it imposes a complete restructuring of the core of the application. Similarly to many other low level transformations, it also tends to render the source code impenetrable, increasing the maintenance and the extension costs. We therefore aim for a fully automated system based on DSLs, which abstracts sparse tiling through a simple interface (i.e., a simple construct that lets the users tell the compiler "apply sparse tiling to the following sequence of loops") and a tiny set of parameters for performance tuning

(e.g., the tile size). We are not interested in supporting legacy code, where the key computational aspects (e.g., mesh iteration, distributed-memory parallelism) are usually hidden for software modularity, thus making automation almost impossible.

Based upon these observations and requirements, we decompose our problem into two tasks:

- a library for writing inspector/executor schemes (Salz et al. [1991]) to enable sparse tiling in arbitrary computations on unstructured meshes;
- 2. integration of the library with a multilayer framework based on DSLs and runtime code generation.

Before addressing these two tasks, respectively in Sections 3.5-3.7 and Section 3.8, we elaborate on the theoretical and technical challenges that arise when applying loop fusion and tiling (Section 3.3), and review the related work (Section 3.4).

# 3.3 Applying Loop Fusion and Tiling is More Difficult than Commonly Thought

We show in Listing 5 the "skeleton" of a typical PDE solver on an unstructured mesh. This will be useful throughout the analysis presented in this section.

We identify three classes of problems that are neglected, or at least treated with scarce emphasis, in the relevant literature.

**Theoretical questions** We first wonder about the effectiveness of fusion and tiling in unstructured mesh applications.

**Computational Boundedness** Computational methods for PDEs are structured as sequences of loop nests, each loop nest characterized by its own operational intensity. In the same application, some loop nests may be memory-bound, while others CPU-bound. This clearly depends on several parameters of the numerical method, including the arithmetic complexity of the operators and the discretization employed (e.g., polynomial order of function spaces). Obviously, if most loop nests in a code are CPU-bound, the benefits of spars tiling on data locality will be marginal. Before even thinking about aggressive optimizations such as fusion and tilling,

**LISTING 5:** The "bare" structure of a numerical method for solving a partial differential equation. Three parallelizable sweeps over sets of mesh components – cells, nodes, boundary nodes – are executed within a time-stepping loop. In the cells loop, we show the invocation of a kernel: first, the memory indirections are resolved; the kernel, which receives data that is now contiguous in memory (this hopefully increases the chances of vectorisation), is then invoked; finally, the computed values are "scattered" back to memory. Distributed-memory parallelism is achieved through MPI; messages are exchanged between processes (MPI\_Comm (...)) between different mesh sweeps. Additional computation (Calc (...)) could also be present (e.g., sparse linear algebra operations, as in the finite element method; checkpointing for fault tolerance).

```
// Time-stepping loop (T = total number of iterations)
2 for t = 0 to T
     // 1st sweep over the {\it C} cells of the mesh
    for i = 0 to C {
       buffer_0 = gather_data ( A[f(map[i])], ... )
       kernel_1( buffer_0, buffer_1, ... );
       scatter_data ( buffer_0, f(map[i]) )
    Calc (...);
10
    MPI_Comm (...);
11
     // 2nd sweep over the N nodes of the mesh
12
13
    for i = 0 to N {
      ... // Similar to sweep 1
14
15
     // Boundary conditions: sweep over the {\it BV} boundary nodes
16
     for i = 0 to BV {
17
       \dots // Similar to sweep 1
18
19
20
21
    Calc (...);
22
    MPI_Comm (...);
23
```

it is fundamental to understand the limiting factors of an application. In essence, two questions need be answered: (i) what fraction of the execution time is due to memory-bound loop nests; (ii) can CPU-boundedness be relieved by applying other optimizations (e.g., vectorization).

Loop Tiling vs Space Filling Curves Loop tiling and Space Filling Curves (SFCs) are two solutions produced by different communities to the same problem: improving the performance of mesh-based computations by making a better use of memory. Our view is that both tiling and SFCs are possible instances of the scheduling function of a given loop nest (see Section 2.3.1). We could not find studies comparing the performance of the two approaches. This seems to denote a lack of communication be-

tween communities that have tackled similar problems for years.

**Practical issues** Several studies on fusion and tiling for structured mesh applications have addressed key problems such as automation (e.g., polyhedral compilers), time-loop tiling (i.e., time skewing), exotic tile shapes for communication minimization (e.g., diamond tiling). However, the following issues were rarely given the right importance.

**Unstructured meshes** Although ad-hoc inspector-executor strategies for some proxy applications had previously been developed, a general technique to fuse and tile arbitrary computations on unstructured meshes has been missing until this thesis<sup>1</sup>. As already explained, the main problem with unstructured meshes is the presence of indirect memory accesses, which essentially inhibit the static analysis needed by any loop transformation.

**Time tiling and MPI** We reiterate the fact that real-world computations almost always run on distributed-memory architectures. This is evident in Listing 5, where MPI calls appear between consecutive mesh sweeps. Distributed-memory parallelism poses a big challenge for time tiling, because tiles at the partition boundary need special handling.

**Time tiling and extra code** The Comp(...) function in Listing 5 denotes the possibility that additional computation is performed between consecutive sweeps. This could be, for instance, check-pointing or I/O. Also, conditional execution of loops (e.g., through if-then-else) breaks time tiling.

**Legacy code is usually impenetrable** Loop transformation opportunities are often hidden in existing scientific codes. As explained in Strout [2013], common problems are: 1) potentially fusible or tilable loop nests are separated for code modularity; 2) handling of boundary conditions; 3) source code not amenable for data dependency analysis (e.g., extensive use of pointers, function calls).

**Limitations inherent in the numerical method** Two loops cannot be fused if they are separated by a global synchronization point. This is often a global reduction, either explicit (e.g., the first loop updates a global variable that is read by the second loop) or implicit (i.e., within an external function

<sup>&</sup>lt;sup>1</sup>We reinforce once more that the generalized sparse tiling algorithm is the result of a joint collaboration among the authors of Strout et al. [2014].

invoked between the two loops, like in many iterative solvers for linear systems). By limiting the applicability of many loop optimizations, global synchronization points pose great challenges and research questions. If strong scaling is the primary goal and memory-boundedness is the key limiting factor, then interesting questions are: (i) can the numerical method be reformulated to relieve the constraints on low level optimization (which requires a joint effort between application and performance specialists); (ii) can the tools be made more sophisticated to work around these problems; (iii) will the effort be rewarded by significant performance improvements.

All these issues and questions will be progressively addressed in the upcoming sections.

#### 3.4 Related Work

#### **Loop Chain**

The data dependence analysis that we develop in this chapter is based on an abstraction called *loop chain*, which was originally presented in Krieger et al. [2013]. This abstraction is sufficiently general to capture data dependency in programs structured as arbitrary sequences of loops. We will detail our loop chain abstraction in Section 3.5.

#### Inspector/Executor and Sparse Tiling

The loop chain abstraction provides sufficient information to create an inspector/executor scheme for an arbitrary unstructured mesh application. Inspector/executor strategies were first formalized by Salz et al. [1991]. They have been used to fuse and tile loops for exploiting data reuse and providing parallelism in several studies [Douglas et al., 2000, Strout et al., 2002, Demmel et al., 2008, Krieger and Strout, 2012].

Sparse tiling, which we introduced in Section 2.3.2, is the most notable technique based upon inspection/execution. The term was coined by Strout et al. [2002, 2004] in the context of the Gauss-Seidel algorithm and in Strout et al. [2003] in the context of the moldyn benchmark. However, the technique was initially proposed by Douglas et al. [2000] to parallelize computations over unstructured meshes, taking the name of *unstruc*-

tured cache blocking. The mesh was initially partitioned; the partitioning represented the tiling in the first sweep over the mesh. Tiles would then shrink by one layer of vertices for each iteration of the loop. This shrinking represented what parts of the mesh could be update in later iterations of the outer loop without communicating with the processes executing other tiles. The unstructured cache blocking technique also needed to execute a serial clean-up tile at the end of the computation. Adams and Demmel [1999] also developed an algorithm very similar to sparse tiling, to parallelize Gauss-Seidel computations. The main difference between Strout et al. [2002, 2004] and Douglas et al. [2000] was that in the former work the tiles fully covered the iteration space, so a sequential clean-up phase at the end could be avoided.

We reiterate the fact that all these approaches were either specific to individual benchmarks or general but not scheduling across loops (i.e., loop fusion). Filling this gap is one of the contributions of this chapter.

## Automated Code Generation and Optimization for Mesh-Based Computations

The automated code generation technique presented in Ravishankar et al. [2012] examines the data affinity among loops and partitions the execution with the goal of minimizing communication between processes, while maintaining load balancing. This technique supports unstructured mesh applications (being based on an inspector/executor strategy) and targets distributed memory systems, although it does not exploit the loop chain abstraction and abstracts from loop optimization.

Automated code generation techniques, such as those based on polyhedral compilers (reviewed in Section 2.3.1), have been applied to structured mesh benchmarks or proxy applications. Notable examples are Bondhugula et al. [2008], Grosser et al. [2012], Klöckner [2014]. There has been very little effort in providing evidence that these tools can be effective in real-world applications. Time-loop diamond tiling was applied in ?? to a proxy code, but experimentation was limited to a single-node.

#### Overlapped Tiling

In structured codes, Using multiple layers of halo, or "ghost", elements is a common optimization in structured codes Bassetti et al. [1998]. Overlapped tilling (see Section 2.3.3) exploits the same principle to reduce communication at the expense of performing redundant computation along the boundary Zhou et al. [2012]. Several studies centered on overlapped tiling within single regular loop nests (mostly stencil-based computations), for example Meng and Skadron [2009], Krishnamoorthy et al. [2007], Chen et al. [2002]. Techniques known as "communication avoiding" [Demmel et al., 2008, Mohiyuddin et al., 2009] also fall in this broader category. To the best of our knowledge, overlapped tiling for unstructured grids by automated code generation has been studied only analytically in Giles et al. [2012].

### 3.5 Generalized Inspector/Executor Schemes

In this section we explain how to build inspector/executor schemes for arbitrary sequences of loops.

#### 3.5.1 Relationship between Loop Chain and Inspector

The *loop chain* is an abstraction introduced in Krieger et al. [2013]. Informally, a loop chain is a sequence of loops with no global synchronization points, with attached some extra information to enable run-time data dependence analysis.

We recall from Section 3.3 that the indirect memory accesses inhibit static optimization for data locality in unstructured mesh applications. The idea to work around this issue is to exploit the data dependence information carried by a loop chain to replace compile-time with run-time analysis. The source code must be modified adding a description of the loop chain and the inspector, as well as by replacing a sequence of loops with a new piece of code, the executor. At run-time, the inspector exploits the data dependence information in the loop chain and builds a "scheduling", which is eventually used as input to the executor.

Before diving into the description of the loop chain abstraction, it is worth observing the following:

- The execution of the inspection phase introduces an overhead. In many scientific computations, however, the data dependence pattern is static; or, more informally, "the mesh does not change over time". This means that the inspection cost can be amortized over multiple iterations of the time loop. If instead the mesh changes over time (e.g., in case of adaptive mesh refinement), the inspection needs be executed again. We have spent a considerable amount of time in designing and implementing a highly optimized inspector algorithm; this hopefully will make the overhead negligible even in the unfortunate case of frequent changes to the data dependence pattern.
- We have explained that the loop chain must be somewhat provided. One possibility is that application specialists manually change their programs by constructing the loop chain and the inspector/executor scheme through library calls. As already observed, this is clearly challenging. Another possibility is that the loop chain is constructed at a higher layer of abstraction, in which case the optimization process is fully automated. The tools we have built enable both approaches.

Further details on these two points will be provided in the later sections.

#### 3.5.2 The Loop Chain Abstraction

In Krieger et al. [2013], a loop chain L is defined as follows:

- L consists of n loops,  $L_0, L_1, ..., L_{n-1}$ . There are no global synchronization points between the loops. The execution order of a loop's iterations does not influence the result. This means that given  $L_i$ , we can choose an arbitrary scheduling of iterations because there will not be loop-carried dependencies.
- $\mathbb{D}$  is a set of disjoint m data spaces,  $D_0, D_1, ..., D_{m-1}$ . Each loop accesses (reads from, writes to) a certain subset of these data spaces. An access can be either direct (e.g., A[i]) or indirect (e.g., A[map(i)]).
- $R_{L_l \to D_d}(\vec{i})$  and  $W_{L_l \to D_d}(\vec{i})$ , where the R and W access relations are defined over for each data space  $D_d \in D$ . They indicate which data locations in data space  $D_d$  an iteration  $i \in L_i$  reads from and writes to respectively (e.g., if we have  $A[B(i)] = \ldots$  in loop  $L_j$ , the access relation  $B_{L_i \to A}(\vec{i})$  will be available).

## 3.5.3 The Loop Chain Abstraction Revisited for Unstructured Mesh Computations

Because of our focus on unstructured mesh computations, and inspired by the programming and execution models of OP2 (see Section 2.2.2), the definition of a loop chain L is changed as follows:

- L consists of n loops,  $L_0, L_1, ..., L_{n-1}$ . There are no global synchronization points between the loops. The execution order of a loop's iterations does not influence the result. This means that given  $L_i$ , we can choose an arbitrary scheduling of iterations because there will not be loop-carried dependencies.
- S is a set of disjoint m sets,  $S_0, S_1, ..., S_{m-1}$ . Possible sets are the cells in the mesh or the degrees of freedom associated to a certain function. Sets are used to define both iteration spaces and datasets. In the former case, a set element is an iteration; in the latter case, a set element is logically associated with a data value (not necessarily a scalar).

A set S is logically split into three contiguous regions: core ( $S^c$ ), boundary ( $S^b$ ), and non-exec ( $S^{ne}$ ). Given a process P and a set S, we have that:

 $S^c$  The iterations of S that exclusively belong to P.

 $S^b$  The boundary region can be seen as the union of two sub-regions, owned ( $S^{owned}$ ) and exec ( $S^{exec}$ ).  $S^{owned}$  are iterations that belong to P which will be redundantly executed by some other processes;  $S^{exec}$  are iterations from other processes which are redundantly executed by P. We will see that redundant computation is exploited to preserve atomic execution of a tile – the property that enables execution without the need for synchronization.

 $S^{ne}$  These are iterations of other processes that are communicated to P because they need be indirectly read to correctly compute  $S^b$ .

A set is uniquely identified by a name and the sizes of its three regions. For example, the notation S = (vertices, C, B, N) defines the vertices set, which comprises C elements in the core region (iterations 0 to C - 1), B elements in the boundary region (iterations C to C + B - 1), and N elements in the non-exec region (iterations C + B to

$$C + B + N - 1$$
).

- The *depth* is an integer indicating the extent of the boundary region of a set. This constant is the same for all sets.
- M is a set of k maps,  $M_0, M_1, ..., M_{k-1}$ . A map of arity a is a vector-valued function  $M: S_i \to S_j^0 \times S_j^1 \times ... \times S_j^{a-1}$  that connects each element of  $S_i$  to one or more elements in  $S_j$ . For example, if a triangular cell c is connected to three vertices  $v_0, v_1, v_2$ , we have  $M(c) = [v_0, v_1, v_2]$ .
- A loop  $L_i$  over the iteration space S is associated with d descriptors,  $D_0, D_1, ..., D_{d-1}$ . A descriptor D is a 2-tuple  $D = \langle M, \bmod \rangle$ . M is either a map from  $S_j$  to some other sets or the special placeholder  $\bot$ , which indicates that the access is direct to some data associated with  $S_j$ . mode is one of [r, w, i], meaning that a memory access is respectively of type read, write or increment.

With respect to the original definition, one of the most important differences is the lack of data spaces. In unstructured mesh applications, a loop tends to access multiple data spaces associated with the same set. The idea, therefore, is to rather rely on sets to perform data dependency analysis. This can significantly improve the inspection cost, because typically  $|S| << |\mathbb{D}|$ .

The second fundamental difference is the characterization of sets into the three regions core, boundary and non-exec. This separation is essential for distributed-memory parallelism (we simply have that both  $S^b$  and  $S^{ne}$  contain no elements in the special case of execution entirely based on shared-memory parallelism). The "extent" of the boundary regions is captured by the *depth* of the loop chain; the actual role of this parameter will become clear in Section 4.

### 3.6 Loop Chain and Inspection Examples

Using the example in Listing 6 – a plain C implementation of the unstructured mesh OP2 program illustrated in Listing 4 – we describe the actions performed by a sparse tiling inspector. The inspector takes as input a loop chain, as illustrated in Listing 7. We will show two variants of inspection, for shared-memory and distributed-memory parallelism. Which variant is executed depends on the value of the variable mode at line 18 in Listing 7.

**LISTING 6:** Section of a toy program that is used as a running example to illustrate the loop chain abstraction and show how the tiling algorithm works.

```
for t = 0 to T {
     // Loop over edges
     for e = 0 to E 
       x = X[e];
      tmp_0 = tmp + edges2vertices[e + 0];
      tmp_1 = tmp + edges2vertices[e + 1];
       kernel1 (x, tmp_0, tmp_1);
8
     // Loop over cells
10
    for c = 0 to C
11
      res = R[C];
13
      tmp_0 = tmp + cells2vertices[c + 0];
14
      tmp_1 = tmp + cells2vertices[c + 1];
15
      tmp_2 = tmp + cells2vertices[c + 2];
16
      kernel2 (res, tmp_0, tmp_1, tmp_2);
17
18
     // Loop over edges
19
20
     for e = 0 to E
      tmp_0 = tmp + edges2vertices[e + 0];
21
22
      tmp_1 = tmp + edges2vertices[e + 1];
      kernel3 (tmp_0, tmp_1);
23
24
25
```

#### Overview

The inspector starts with partitioning the iteration space of a *seed loop*, for example  $L_0$ . Partitions are used to initialize tiles: the iterations of  $L_0$  falling in  $P_i$  – or, in other words, the edges in partition  $P_i$  – are assigned to the tile  $T_i$ . Figure 3.1 displays the situation after the initial partitioning of  $L_0$  for a given input mesh. There are four partitions, two of which are not connected through any edge or cell. These four partitions correspond to four tiles,  $[T_0, T_1, T_2, T_3]$ .

In the later stages of the inspection,  $T_i$  is populated with iterations from  $L_1$  and  $L_2$ . The challenge in scheduling iterations from  $L_j$  to  $T_i$  is to guarantee that all data dependencies – read after write, write after read, write after write – are honored. In the next two sections, we use the running example to illustrate how iterations are assigned to tiles to enable shared-memory and distributed-memory parallelism.

The result of the inspection is eventually passed to the executor. The inspection carries sufficient information for a parallel computation of different tiles. A given tile  $T_i$  is always executed by a single thread/pro-

#### **LISTING 7:** Building the loop chain for inspection.

```
nspector = init_inspector (...);

// Three sets, edges, cells, and vertices
Let = set (inspector, core_edges, boundary_edges, nonexec_edges, ...);
C = set (inspector, core_cells, boundary_cells, nonexec_cells, ...);
V = set (inspector, core_vertices, boundary_vertices, nonexec_vertices, ...);

// Two maps: from edges to vertices and from cells to vertices
e2vMap = map (inspector, E, V, edges2vertices, ...);
c2vMap = map (inspector, C, V, cells2vertices, ...);

// The loop chain comprises three loops; each loop has a set of descriptors
loop (inspector, E, {\perp , READ}, {e2vMap, INC});
loop (inspector, C, {\perp , READ}, {c2vMap, INC});
loop (inspector, E, {e2vMap, INC});
// Now can run the inspector
inspection = run_inspection (mode, inspector, tile_size, ...)
return inspection;
```

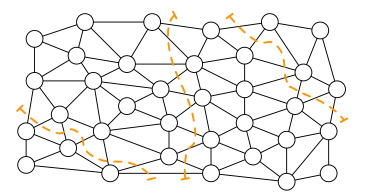


Figure 3.1: Partitioning of the seed loop.

cess "atomically"; that is, without ever communicating with other threads/processes. When executing  $T_i$ , first all iterations from  $L_0$  are executed, then all iterations from  $L_1$  and finally those from  $L_2$ .

#### **Inspection for Shared-Memory Parallelism**

Similarly to OP2, to achieve shared-memory parallelism we use coloring. Two tiles that are given the same color can be executed in parallel by different threads. Two tiles can have the same color if they are not connected, because this ensures the absence of race conditions through indirect memory accesses during parallel execution. In the example we can use three colors: red (R), green (G), and blue (B).  $T_0$  and  $T_3$  are not connected, so they are assigned the same color. The colored tiles are shown in Figure 3.2. In the following, with the notation  $T_i^c$  we indicate that the i-th tile has color c.

To populate the tiles with iterations from  $L_1$  and  $L_2$ , we first have to establish a total ordering for the execution of partitions with different colors. Here, we assume the following order: green (G), blue (B), and red (R). This means, for instance, that *all iterations* assigned to  $T_1^B$  must be executed *before all iterations* assigned to  $T_2^R$ . By "all iterations" we mean the iterations from  $L_0$  (determined by the seed partitioning) as well as the iterations that will later be assigned from  $L_1$  and  $L_2$ . We assign integer positive numbers to colors to reflect their ordering, where a smaller number means higher execution priority. In this case, we can assign 0 to green, 1 to blue, and 2 to red.

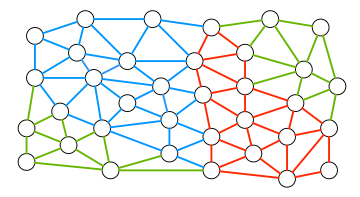


Figure 3.2: Tiling  $L_0$ .

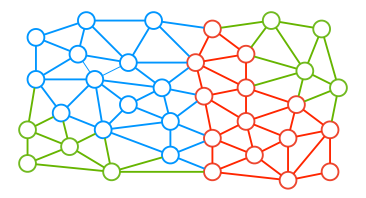


Figure 3.3: Projection of  $L_0$  to  $L_1$ .

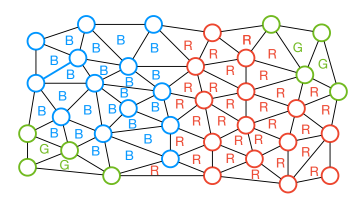


Figure 3.4: Tiling  $L_1$ .

To schedule the iterations of  $L_1$  to  $[T_0^G, T_1^B, T_2^R, T_3^G]$ , we first need to compute a *projection* of any write or local reduction performed within  $L_0$ . A projection for  $L_0$  is a function  $\phi: V \to \mathbb{T}$  mapping a vertex indirectly incremented during the execution of  $L_0$  (see Listing 6) to a tile. Consider the vertex  $v_0$  in Figure 3.3.  $v_0$  has 7 incident edges, 2 of which belong to  $T_0^G$ , while the remaining 5 to  $T_1^B$ . Since we established that  $G \prec B$ ,  $v_0$  can only be read after  $T_1^B$  has finished executing the iterations from  $L_0$  (i.e., the 5 incident blue edges). We express this condition by setting  $\phi(v_0) = T_1^B$ . Observe that we can compute  $\phi$  by iterating over V and, for each vertex, applying the maximum function (MAX) to the color of the adjacent edges.

We now use  $\phi$  to schedule  $L_1$ , a loop over cells, to the tiles. Consider again  $v_0$  and the adjacent cells  $[c_0, c_1, c_2]$ . These three cells have in common the fact that they are adjacent to both green and blue vertices. For  $c_1$ , and similarly with the other cells, we compute  $MAX(\phi(v_0), \phi(v_1), \phi(v_2)) =$ 

MAX(B,G,G) = B = 1. This establishes that  $c_1$  must be assigned to  $T_1^B$ , because otherwise ( $c_1$  assigned instead to  $T_0^G$ ) a read to  $v_0$  would occur before the last increment from  $T_1^B$  took place. We indeed reiterate once more that the execution order is "all iterations from [ $L_0, L_1, L_2$ ] in the green tiles before all iterations from [ $L_0, L_1, L_2$ ] in the blue tiles". The result of scheduling  $L_1$  to tiles is displayed in Figure 3.4.

To schedule  $L_2$  to  $[T_0^G, T_1^B, T_2^R, T_3^G]$  we employ a similar process. Vertices are again written by  $L_1$ , so a new projection over V will be computed and will be used in place of  $\phi$  to schedule the edges in  $L_2$ . Figure 3.5 shows the result of this last phase.

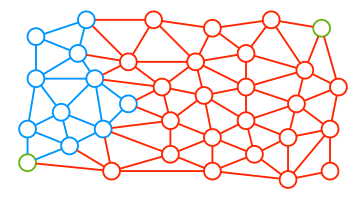


Figure 3.5: Tiling  $L_2$ .

#### **Conflicting Colors**

It is worth noting how  $T_2^R$  "consumed" the frontier elements of all other tiles every time a new loop was scheduled. Tiling a loop chain consisting of k loops had the effect of expanding the frontier of a tile of at most k vertices. With this in mind, we re-inspect the loop chain of the running example, although this time with different execution order – blue (B), red (R), and green (G) – and initial partitioning. We recall that the total ordering of colors is indeed arbitrary. By applying the same inspection procedure that we just described,  $T_0^G$  and  $T_3^G$  will eventually become connected (see Figure 3.6), thus violating the precondition "tiles/partitions with the same color can run in parallel". Race conditions during the execution of iterations belonging to  $L_2$  are now theoretically possible. We will solve this problem in Section 3.7.2.

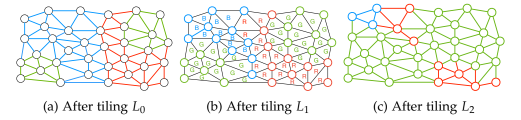


Figure 3.6: Tiling the program in Listing 6 for shared memory parallelism can lead to conflicts. Here, the two green tiles eventually become adjacent, exposing potential race conditions.

#### Inspection for Distributed-Memory Parallelism

If parallelism is achieved through message-passing, the mesh and its datasets are distributed to different processes. Similarly to the example in Listing 5, MPI calls now separate the execution of two consecutive loops in the chain. This means that our inspector/executor scheme will have to take extra care of data dependencies along the mesh partition boundaries.

From Section 3.5.3 we know that all sets are logically split into three regions, *core*, *boundary*, and *non-exec*. The boundary region includes all iterations that cannot be executed until the MPI communications have successfully been accomplished. With this in mind, we take again  $L_0$  as seed loop and make the following adjustments to the procedure illustrated in the previous section:

- 1. We take the core region of  $L_0$  and partition it into tiles. Unless we aim for a mixed distributed/shared-memory parallelization scheme, there is no need to reassign the same color to unconnected tiles since we now have a single process executing sequentially a group of tiles . We can assign colors increasingly:  $T_i$  is given color i. This allows us to uniquely identify a tile's color with its index, i. As long as tiles with contiguous ID are also physically contiguous in the mesh, this assignment retains spatial locality when "jumping" from executing  $T_i$  to  $T_{i+1}$ .
- 2. We replicate for the boundary region of  $L_0$  what we just did for the core. By neatly separating the core and boundary regions, we prevent tiles from crossing the two regions by construction. Further, all tiles within boundary have greater color than those in core, which poses a constraint on the execution order: no boundary tiles can be

executed until all core tiles have been scheduled.

3. We map the whole non-exec region of  $L_0$  to a single special tile,  $T_{ne}$ . This tile has highest color and will actually never be executed. Its role will be explained in later sections.

In Figure 3.7, the same mesh of Figure 3.1 has been distributed to two processes and the output of the initialization phase is displayed. The inspection then proceeds as in the case of shared memory parallelism. The application of the MAX function when scheduling iterations from  $L_1$  and  $L_2$  to tiles makes higher color tiles (i.e., those that will be scheduled later at execution time) "grow over" lower color ones. The whole boundary region "expands" over core, and so does  $T_{ne}$  over boundary (see Figure ??).

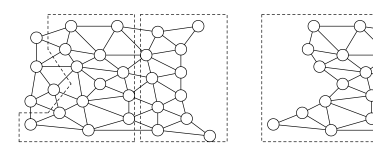


Figure 3.7: Seed partitioning for distributed memory execution.

On each process, the executor starts with triggering the MPI communications required for the execution of (all tiles over) the boundary regions; it proceeds to scheduling the core tiles, thus overlapping communication with computation; eventually, as soon as all core tiles have been executed and the MPI communications been accomplished, it computes the boundary tiles.

Finally, we observe that:

**Correctness** This inspector/executor scheme relies on redundant computation along the boundaries of mesh partitions. The depth of the boundary region – a parameter of the loop chain, as explained in Section 3.5.3 – grows proportionally with the number of loops that are fused. Roughly speaking, if the loop chain consists of n loops, then the boundary region needs n-1 extra layers of elements. In Figure 3.7, the elements [...] belong to  $R_1$ , although they also need be executed by  $R_0$  in order for [...] to be correctly computed when  $T_x$  executes iterations from  $L_2$ . This is a conceptually simple expedient, although it poses a big challenge on software

engineering.

**Efficiency** The underlying hypothesis is that the increase in data locality achieved through sparse tiling will outweigh the overhead introduced by the redundant computation. This is based on the consideration that, in real applications, the core region tends to be way larger than the boundary one. In addition, not all iterations along the boundary region need be redundantly executed at every loop. For example, if we consider Figure ??, we see that the strip of elements [...] is not relevant any more for the correct computation of [...]. The non-exec tile  $T_{ne}$ , which we recall is not scheduled by the executor, is deliberately given the highest color to grow over the boundary region: this will leave dirty values in some datasets until the next round of MPI communications, but will not hurt correctness.

#### 3.7 Formalization

#### 3.7.1 Data Dependency Analysis for Loop Chains

As with all loop optimizations that reschedule the iterations in a sequence of loops, any sparse tiling must satisfy the data dependencies. The loop chain abstraction, which we have described in Section 3.5, provides enough information to construct an inspector which analyzes all of the dependencies in a computation and builds a legal sparse tiling. We recall that one of the main assumptions in a loop chain is that each loop is fully parallel or, equivalently, that there are no loop carried dependencies.

The descriptors in the loop chain abstraction enable a general derivation of the storage-related dependencies between loops in a loop chain. The storage related dependencies between loops can be described as either flow (read after write), anti (write after read), or output (write after write) dependencies. In the following, assume that loop  $L_x$ , having iteration space  $S_x$ , always comes before loop  $L_y$ , having iteration space  $S_y$ , in the loop chain. Let us identify a descriptor of a loop L with  $m_{S_i \to S_j}^{mode}$ : this simply indicates that the loop  $L_i$  has iteration space  $S_i$  and uses a map m to write/read/increment elements (respectively, mode  $\in \{w, r, i\}$ ) in the space  $S_i$ .

The flow dependencies can then be enumerated by considering pairs of

points  $(\vec{i} \text{ and } \vec{j})$  in the iteration spaces of the two loops  $L_x$  and  $L_y$ :

$$\{\vec{i} \to \vec{j} \mid \vec{i} \in S_x \land \vec{j} \in S_y \land m^w_{S_x \to S_z}(\vec{i}) \cap m^r_{S_y \to S_z}(\vec{j}) \neq \emptyset\}.$$

Anti and output dependencies are defined in a similar way. The anti dependencies for all pairs of loops  $L_x$  and  $L_y$  are:

$$\{\vec{i} \to \vec{j} \mid \vec{i} \in S_x \land \vec{j} \in S_y \land m^r_{S_x \to S_z}(\vec{i}) \cap m^w_{S_y \to S_z}(\vec{j}) \neq \emptyset\}.$$

While the output dependencies between loops  $L_x$  and  $L_y$  are:

$$\{\vec{i} \to \vec{j} \mid \vec{i} \in S_x \land \vec{j} \in S_y \land m^w_{S_x \to S_z}(\vec{i}) \cap m^w_{S_y \to S_z}(\vec{j}) \neq \emptyset\}.$$

In essence, there is a storage-related data dependence between two iterations from different loops (and therefore between the tiles they are placed in) when one of those iterations writes to a data element and the other iteration reads from or writes to the same data element.

There are local reductions, or "reduction dependencies" between two or more iterations of the same loop when those iterations "increment" the same location(s); that is, when they read, modify with a commutative and associative operator, and write to the same location(s). The reduction dependencies in  $L_x$  are:

$$\{\vec{i} \to \vec{j} \mid \vec{i} \in S_x \land \vec{j} \in S_x \land m^i_{S_x \to S_z}(\vec{i}) \cap m^i_{S_x \to S_z}(\vec{j}) \neq \emptyset\}.$$

The reduction dependencies between two iterations within the same loop indicates that those two iterations must be executed atomically with respect to each other.

As seen in the example in Section 19, our inspector algorithm handles data dependencies, including those between non-adjacent loops, by tracking *projections*. In the next section we explain how projections are constructed and used.

#### 3.7.2 The Generalized Sparse Tiling Inspector

The pseudo-code for the generalized sparse tiling inspector is showed in Algorithm 1. Given a loop chain and an average tile size, the algorithm produces a schedule suitable for mixed distributed/shared memory par-

Symbol	Meaning
$\overline{\mathbb{L}}$	The loop chain
$L_i$	The $i$ -th loop in $\mathbb L$
$S_i$	The iteration space of $L_i$
$S_i^c, S_i^b, S_i^n$	the core, boundary, and non-exec regions of $S_i$
S	A generic set in L
$S_{in}$ , $S_{out}$	The input and output sets of a map
D	A descriptor of a loop
r, w, i	Possible values for D.mode
${\mathbb T}$	The set of all tiles
$\mathbb{T}[i]$	Accessing the <i>i</i> -th tile
$T_i^c$ , $T_i^b$	the <i>i</i> -th tile over the core and boundary regions
$\phi_S$	A projection $\phi_S:S o \mathbb{T}$
Φ	The set of all available projections
$\sigma_i$	The tiling function $\sigma_i : S_i \to \mathbb{T}$ for $L_i$

Table 3.1: Summary of the notation used throughout the section.

allelism. In the following, we elaborate on the main steps of the algorithm. The notation used throughout the section is summarized in Table 3.1.

**Choice of the seed loop** The seed loop  $L_{seed}$  is used to initialize the tiles. Theoretically, any loop in the chain can be chosen as seed. Supporting distributed memory parallelism, however, is cumbersome if  $L_{seed} \neq L_0$ . This is because more general partitioning and coloring schemes would be needed to ensure that no iterations in any  $S_i^b$  are assigned to a core tile. A constraint of our inspector algorithm in the case of distributed memory parallelism is that  $L_{seed} = L_0$ .

In the special case in which there is no need to distinguish between core and boundary tiles (because a program is executed on a single shared-memory system), the seed loop can be chosen arbitrarily. If we however pick  $L_{seed}$  in the middle of the loop chain ( $L_0 \prec ... \prec L_{seed} \prec ...$ ), a mechanism for constructing tiles in the reverse direction ("backwards"), from  $L_{seed}$  towards  $L_0$ , is necessary. In Strout et al. [2014], we propose two "symmetric" algorithms to solve this problem, forward tiling and backward tiling, with the latter using the MIN function in place of MAX when computing projections. For ease of exposition, and since in the fundamental case of distributed memory parallelism we impose  $L_{seed} = L_0$ , we here neglect

#### **ALGORITHM 1:** The inspection algorithm

```
Input: The loop chain \mathbb{L} = [L_0, L_1, ..., L_{n-1}], a tile size ts Output: A set of tiles \mathbb{T}, populated with iterations from \mathbb{L}
```

```
// Initialization
 1 seed \leftarrow 0;
 _{2}\ \Phi\leftarrow\varnothing;
 з C \leftarrow \perp;
    // Creation of tiles
 4 \sigma_{seed}, \mathbb{T} \leftarrow \text{partition}(S_{seed}, ts);
 5 seed_map \leftarrow find_map(S_{seed}, \mathbb{L});
 6 conflicts ← false;
    // Schedule loops to tiles
 7 do
         color(\mathbb{T}, seed\_map);
         for i = 1 to n - 1 do
10
              project(L_{i-1}, \sigma_{i-1}, \Phi, C);
11
              \sigma_i \leftarrow \text{tile}(L_i, \Phi);
              assign(\sigma_i, \mathbb{T});
12
         end for
13
         if found_conflicts(C) then
14
15
              conflicts \leftarrow true;
              add_fake_connection(seed_map, C);
16
        end if
17
18 while conflicts;
    // Inspection successful, create local maps and return
   create_local_maps(\mathbb{T});
20 return T
```

this distinction<sup>2</sup>.

**Tiles initialization** The algorithm starts with partitioning  $S_{seed}^c$  into m subsets  $\{P_0, P_1, ..., P_{m-1}\}$  such that  $P_i \cap P_j = \emptyset$  and  $\bigcup_{i=0}^{m-1} P_i = S_{seed}^c$ . The partitioning is sequential:  $S_{seed}$  is "chunked" every ts iterations, with  $P_{m-1}$  that may be smaller than all other partitions. Similarly, we partition  $S_{seed}^b$  into k subsets.

<sup>&</sup>lt;sup>2</sup>The algorithm that is implemented in the SLOPE library, however, is more general and supports backwards tiling for the case in which distributed memory parallelism is not required.

Field	Possible values		
region	core, boundary, non-exec		
color	an integer representing the execution priority		
iterations lists local maps	one list of iterations (integers) $T_i^{L_j}$ for each $L_j \in \mathbb{L}$ one list of local maps for each $L_j \in \mathbb{L}$ ; one local		
iocai maps	map for each map used in $L_j \in \mathbb{L}$ , one local		

Table 3.2: The tile data structure.

We then create m + k + 1 tiles,  $\mathbb{T} = \{T_0^c, ..., T_{m-1}^c, T_m^b, ..., T_{m+k-1}^b, T_{m+k}^n\}$ , one for each partition and one extra tile for  $S_{seed}^n$ .

A tile  $T_i$  has four fields, as summarized in Table 3.2. The region is used by the executor to schedule tiles in a certain order. The value of this field is determined right after the seed partitioning, since a tile is exclusively in one of  $\{S_{seed}^c, S_{seed}^b, S_{seed}^n, S$ 

Each tile needs be assigned a color, which gives it a scheduling priority. If shared memory parallelism is requested, adjacent tiles are given different colors (the adjacency relation is determined through the maps available in  $\mathbb{L}$ ). Otherwise, the colors are assigned to the tiles in increasing order (i.e.,  $T_i$  is given color i). The boundary tiles are always given colors higher than that of core tiles; the non-exect le has highest color. In essence, this will allow the executor to schedule all core tiles first and all boundary tiles afterwards.

Construction of tiles by tracking data dependencies To schedule a loop to tiles we use projections. A projection is a function  $\phi_S : S \to \mathbb{T}$ . Projections are computed and/or updated after tiling a loop. Initially, the set  $\Phi$  of all projections is empty. Starting with the seed tiling  $\sigma_{seed} : S_{seed} \to \mathbb{T}$ , we iteratively update  $\Phi$  and build tiling functions for all other loops  $[L_1, L_2, ..., L_{n-1}]$ .

#### **ALGORITHM 2:** Projection of a tiled loop

**Input**: A loop  $L_i$ , a tiling function  $\sigma_i$ , the projections set Φ, the conflicts matrix C **Result**: Update Φ and C

```
1 foreach D in L_i.descriptors do
        if D.mode == r then
2
          skip;
3
        end if
4
        if D.map == \perp then
5
              \Phi = \Phi \cup \sigma_i;
6
7
        else
              inverse\_map \leftarrow map\_invert(D.map);
8
              S_i, S_i, values, offsets \leftarrow inverse_map;
10
              \phi_{S_i} \leftarrow \perp;
11
              for e = 0 to S_i.size do
                   for k = offsets[e] to offsets[e+1] do
12
                        adjacent_tile = T[values[k]];
13
                        \max\_color \leftarrow MAX(\phi_{S_i}[e].color, adjacent\_tile.color);
                        if max\_color \neq \phi_{S_i}[e].color then
15
                             \phi_{S_i}[e] \leftarrow \text{adjacent\_tile};
16
17
                        end if
                   end for
18
              end for
19
              update(C, \mathbb{T}, \phi_{S_i});
20
              \Phi = \Phi \cup \phi_{S_i};
21
        end if
23 end foreach
```

**Projecting tiled sets** Algorithm 2 takes as input  $\sigma_{i-1}$  and (the descriptors of)  $L_{i-1}$  to update  $\Phi$ . Further, the conflicts matrix  $C \in \mathbb{N}^{m \times m}$ , indicating whether two different tiles having the same color will become adjacent after tiling  $L_{i-1}$ , is updated.

A projection tells what tile a set element logically belongs to at a given point of the tiling process. A new projection  $\phi_S$  is needed if the elements of S are written by  $L_i$ . Let us consider the non-trivial case in which writes or increments occur indirectly through a map  $M: S_i \to S_j^0 \times S_j^1 \times ... \times S_j^{a-1}$ . To compute  $\phi_{S_j}$ , we first determine the inverse map (an example is shown in Figure ??). Then, we iterate over all elements of  $S_i$  that indirectly access elements in  $S_j$ . In particular, given  $e \in S_j$ , we determine the last tile that writes to e, say  $T_{last}$ , through the application of the MAX function to tile colors. We then simply set  $\phi_{S_j}[e] = T_{last}$ .

#### **ALGORITHM 3:** Building a tiling function

**Input**: A loop  $L_i$  (with iteration space  $S_i$ ), the projections set  $\Phi$  **Output**: The tiling function  $\sigma_i$ 

```
1 \sigma_i \leftarrow \perp;
 2 foreach D in L<sub>i</sub>.descriptors do
         if D.map == \perp then
               \sigma_i \leftarrow \Phi[S_i];
 4
 5
          else
               arity \leftarrow D.map.arity;
 6
                \phi_S \leftarrow \Phi[D.\text{map.}S_i];
 7
                for e = 0 to S_i.size do
 8
                     \sigma_i[e] \leftarrow T_{\perp};
                     for k = 0 to arity do
10
11
                           adjacent_tile \leftarrow \phi_S[D.\text{map.values}[e^*\text{arity} + k]];
                           \max\_color \leftarrow MAX(\sigma_i[e].color, adjacent\_tile.color);
12
                           if max\_color \neq \sigma_i[e].color then
13
                               \sigma_i[e] \leftarrow \text{adjacent\_tile};
14
15
                           end if
                     end for
16
               end for
17
          end if
18
19 end foreach
   return \sigma_i
```

**Scheduling loops** Using  $\Phi$ , we compute  $\sigma_i$  as described in Algorithm 3. The algorithm is similar to the projection of a tiled loop, with the main difference being that now we use  $\Phi$  to schedule iterations correctly. Finally,  $\sigma_i$  is inverted and the iterations added to the corresponding iteration lists  $T_j^{L_i}$  of all  $T_j \in \mathbb{T}$ .

**Detection of conflicts** If C indicates the presence of at least one conflict, say between  $T_i$  and  $T_j$ , we add a "fake connection" between  $T_i$  and  $T_j$  and loop back to the coloring stage.  $T_i$  and  $T_j$  are now connected, so they will receive different colors.

#### 3.7.3 The Generalized Sparse Tiling Executor

The sparse tiling executor is illustrated in Algorithm 4. It consists of four main phases: triggering of MPI communications; execution of core tiles (in overlap with communication); waiting for the termination of all MPI

#### **ALGORITHM 4:** The executor algorithm

```
Input: A set of tiles \mathbb{T}
   Result: Execute the loop chain, thus modifying the data sets written or
            incremented within L
1 \mathbb{T}^c, \mathbb{T}^b \leftarrow \text{group\_tiles\_byregion}(\mathbb{T});
2 start_MPI_communications();
3 foreach available color c do
       foreach T \in \mathbb{T}^c s.t. T.color == c do
        execute_tile(T);
       end foreach
7 end foreach
8 end_MPI_communications();
  foreach available color c do
       foreach T \in \mathbb{T}^b s.t. T.color == c do
        execute_tile(T);
11
       end foreach
12
13 end foreach
```

communications; execution of boundary tiles.

As seen in the example in Section 19 and based on the explanation in Section 3.7.2, we know that the core tiles do not require any off-process information to execute as long as the boundary regions are (i) up-to-date and (ii) "sufficiently deep". If both conditions hold, the execution is semantically correct and it is safe to overlap computation of core tiles with the communication of boundary data.

The depth of the boundary region In  $\mathbb{L}$  we have n loops. A tile "crosses" all of these loops and is executed atomically; that is, once it starts executing its iterations, it reaches the end without the need for synchronization with other processes. With this execution model, the boundary region must include a sufficient amount of off-process iterations for a correct computation of the tiles along the border with the core region.

In the extreme case n = 1, a single "strip" of iterations belonging to adjacent processes need be redundantly executed. As n increases, more off-process data is required. If n = 3, as in the example in Figure ??,

Phase	Cost shared memory	Cost distributed memory	
Partitioning	N	N	
Coloring	RK	N	
Projection	$R(nMNK^2)$	$nMNK^2$	
Tiling	R(nMNK)	nMNK	
Local maps	nM	nM	

Table 3.3: Worst-case costs of inspection.

we need three "strips" of off-process iterations to be computed in order for the datasets over X to be correctly updated when executing iterations belonging to  $L_3$ .

The *depth* is a parameter provided through the loop chain abstraction that informs the inspector about the extent of the boundary region. For correctness, we cannot tile more than *depth* loops, so if n > depth the loop chain must first be split into smaller sequences of loops, to be individually inspected and executed.

#### 3.7.4 Computational Complexity of Inspection

Let N be the maximum size of a set in  $\mathbb{L} = [L_0, L_1, ..., L_{n-1}]$  and let M be the maximum number of sets accessed in a loop. If a is the maximum arity of a map, then K = aN is the maximum cost for iterating over a map. K is also the worst-case cost for inverting a map. Let p < 1 be the probability that a conflict arises during inspection in the case of shared memory parallelism; thus, the expected number of inspection rounds is  $K = \frac{1}{1-p}$ . Hence, the worst-case computational costs of the main inspection phases are as in Table 3.3.

### 3.8 Implementation

The implementation of generalized sparse tiling is distributed over three software modules.

**Firedrake** This is the framework for the automated solution of finite element methods described in Section ??.

**PyOP2** Firedrake produces numerical kernels to be applied over sets of mesh components. The parallel iteration over the mesh is handled by

PyOP2 (see Section 2.2.2).

**SLOPE** A library for writing generalized inspector/executor schemes, with primary focus on sparse tiling. PyOP2 uses SLOPE for tiling loop chains. The interaction among these modules is illustrated in Figure ??.

There are several reasons behind the choice of this three-layer structuring:

**Simpler analysis** The abstractions used in Firedrake and PyOP2 simplify the analysis required for automation. For example, from the parallel loop construct in PyOP2 we can derive sufficient information to build a loop chain (see Section 3.5.3).

**User base** We wish generalized sparse tiling to be used in real scientific software, so we targeted a system with a user base in steady expansion. This increased the implementation effort, but it has also made sparse tiling promptly available to a wide audience.

**Flexibility** Adopting sparse tiling in a Firedrake program is the simplest option, because the generation of inspector/executor schemes is fully automated. If we instead have a PyOP2 program, the generation of inspector/executor schemes is only partly automated, since the separation of a set into the core, boundary and non-exec regions for distributed-memory parallelism must be coded explicitly. Users can also write an inspector-executor scheme from scratch, in which case the entire loop chain must be provided through direct calls to the SLOPE library (an example was provided in Listing 7). This is clearly the most flexible yet complicated option.

In the next sections, we describe the interplay between these three frameworks and how this leads to automation.

#### 3.8.1 SLOPE: a Library for Tiling Irregular Computations

SLOPE is an open source software that provides a set of functions to build a loop chain and to write inspector/executor schemes for sparse tiling<sup>3</sup>.

The loop chain abstraction implemented by SLOPE is the one described in Section 3.5.3. In essence, a loop chain comprises some sets (including the separation into core, boundary, and non-exec regions), maps between

<sup>&</sup>lt;sup>3</sup>SLOPE is freely available at https://github.com/coneoproject/SLOPE

sets, and a sequence of loops. Each loop has one or more descriptors specifying what and how dataset are accessed. The example in Listing ?? illustrates the interface exposed by SLOPE for constructing loop chains.

SLOPE implements all of the algorithms in Section 3.7.2. It also provides features to verify the effectiveness and the correctness of the transformations:

**VTK file generator** For each tiled loop, a file showing the mesh and the repartition into colored tiles is generated. The file is suitable for visualization in Paraview ?.

The summary routine The inspector returns useful information concerning the tiling process, including: the number and the average size of tiles, the total number of colors used (which can give an indication of how effective a shared-memory parallelization will be), times for the most critical inspection phases.

In the case of shared-memory parallelism, the following sections of code are parallelized through OpenMP:

- The projection and tiling algorithms; in particular, the loop at line 11 of Algorithm 2 and the loop at line 8 of Algorithm 3).
- The execution of tiles having the same color; that is, the loops at lines 4 and 10 of Algorithm 4.

## 3.8.2 PyOP2: a Runtime Library for Mesh Iteration based on Lazy Evaluation

PyOP2 has been described in Section 2.2.2. This section focuses on three complementary aspects: (i) the interface with the user for identifying sequences of "chainable" loops; (ii) the lazy evaluation mechanism that allows building loop chains; (iii) the interface with SLOPE to run an inspector/executor scheme.

To trigger the creation of a loop chain, PyOP2 provides the *loop\_chain* construct. The use of this interface is exemplified in Listing ??. The *loop\_chain* interface is exposed to both users and higher layers of abstraction. Thus, in Firedrake we define a loop chain in a way analogous to PyOP2's, with the only difference being that loops are not visible in the source code because they are automatically generated from the mathemat-

ical notation.

The second fundamental aspect is that PyOP2 implements lazy evaluation of parallel loops. The parallel loops encountered or generated by Firedrake are first pushed into a queue. The sequence of parallel loops in the queue is called trace. If a field f needs be read, for example because a user wants to inspect its values or a global linear algebra operation is about to be performed, the trace is traversed – from the most recent parallel loop to the oldest one – and a new sub-trace produced. The sub-trace, which includes all parallel loops that need be executed for a correct update of f, is finally executed.

All loops in the trace that are created within a *loop\_chain* scope are sparse tiled through SLOPE. In detail, the interaction between PyOP2 and SLOPE is as follows:

- 1. As seen in the examples in Listing ??, the  $loop\_chain$  defines a new scope. As this scope is entered, a stamp  $s_1$  of the loop chain is generated. This happens "behind the scenes", because the  $loop\_chain$  is a Python context manager, which executes pre-specified routines prior and after the execution of the body. As the  $loop\_chain$  is exited, a new stamp  $s_2$  of the trace is computed, and all parallel loops generated between  $s_1$  and  $s_2$  are placed into a list for pre-processing.
- 2. The pre-processing consists of two main steps: "simple" fusion of all consecutive parallel loops iterating over the same iteration space that do not present read-after-write or write-after-read dependencies through indirections/maps; generation of a loop chain suitable for SLOPE.
- 3. SLOPE provides a thin Python interface to ease integration with other frameworks. PyOP2 inspects the sequence of loops and translates all relevant data structures (sets, maps, loops) into a format suitable for SLOPE. Eventually, a string of C code implementing an inspector for the loops in the *loop\_chain* is returned. PyOP2 just-intime compiles this code and runs it, obtaining the *inspection* data structure.
- 4. A "software cache" mapping *loop\_chains* to *inspection* data structures is used. This whole process is therefore executed only once for each unique *loop\_chain* encountered in the code.

5. The executor is built analogously, and invoked each time a *loop\_chain* is exited.

#### 3.8.3 Firedrake/DMPlex: the S-depth mechanism for MPI

Firedrake uses DMPlex ? to handle meshes. In particular, DMPlex is responsible for partitioning, distributing over multiple processes, and locally reordering a mesh. The MPI parallelization, therefore, is completely handled in Firedrake.

During the start-up phase, each MPI process receives a contiguous partition of the original mesh from DMPlex. It then creates various PyOP2 sets, representing either mesh components (e.g., cells, vertices) or function spaces. As seen in Section 2.2.2, PyOP2 sets distinguish between different regions: core, owned, exec, and non-exec. Firedrake is directly responsible for setting these regions and creating data structures enabling message passing among different processes.

To support our loop chain abstraction, Firedrake must be able to allocate arbitrarily deep halo regions; both Firedrake and DMPlex have been extended with this feature<sup>4</sup>. A parameter called *S-depth* (for historical reasons, see ?), used for initializing a mesh, specifies the extent of the halo regions. A value of *S-depth* equal to *n* indicates the presence of *n* strips of off-process data elements in each set. The default value for *S-depth* is 1, which corresponds to the standard execution model of Firedrake/PyOP2. Even in absence of sparse tiling, this value enables overlap of computation with communication, at the price of a small amount of redundant computation along partition boundaries.

#### 3.9 Performance Evaluation

The experimentation was divided into two phases:

 Generalized sparse tiling was initially applied to two benchmarks: a sparse Jacobi kernel and a proxy unstructured mesh application, originally developed as a demo for the OP2 framework. This was especially useful to identify strengths and limitations of the technique.

<sup>&</sup>lt;sup>4</sup>Michael Lange carried out the bulk of the implementation.

Matrix name	Execution time reduction	Speed-up
ldoor	40.34	12.11
pwtk	38.42	11.98
thermal2	25.78	11.08
xenon2	20.15	9.53
audikw_1	13.42	8.70
nd24k	-151.72	3.06

Table 3.4: Execution time reductions over the original implementation (in percentage) and speed-ups over the single-threaded tiled implementation for the sparse Jacobi solver with 15 threads.

2. Then, it was used in an application developed in Firedrake, Seigen, an Elastic Wave Equation Solver for Seismological Problems<sup>5</sup>. The aim of this phase was to evaluate the performance improvements in a real-world simulation.

#### 3.9.1 Benchmarks

In this section, our objective is to explore the impact of generalized sparse tiling on performance, to characterize the circumstances where the approach is profitable, to identify the major limitations to a widespread adoption of the technique, and to justify some of the design choices made in the previous sections.

All benchmarks are written in C. The sparse tiled implementations only support shared memory parallelism via OpenMP. A previous version of the inspector algorithms presented in this chapter, described in Strout et al. [2014], was used. As detailed next, one of the major drawbacks of this older inspector was its cost, which grew very rapidly with the number of loops in the chain and the number of distinct datasets accessed.

In all the experiments presented in this section, the optimal tile size (i.e. the one leading to the best execution time) was determined empirically, for each combination of architecture and application.

#### Sparse Jacobi

The first experiment was the full sparse tiling of a Jacobi sparse matrix solver<sup>6</sup>. Given a sparse matrix A, and a vector  $\vec{f}$ , related by  $A\vec{u} = \vec{f}$ , each iteration of the sparse Jacobi method produces an approximation to the unknown vector  $\vec{u}$ . In our experiments, the Jacobi convergence iteration loop is unrolled by a factor of two and the resulting two loops are chained together (1000 iterations of the loop chain was executed). Using a ping-pong strategy, each loop reads from one copy of the  $\vec{u}$  vector and writes to the other copy. This experiment was run on an Intel Westmere (dual-socket 8-core Intel Xeon E7-4830 2.13 GHz, 24MB shared L3 cache per socket). The code was compiled using gcc-4.7.0 with options -03 -fopenmp.

The Jacobi recurrence equation includes a sparse matrix vector multiplication and is representative of a broad class of sparse linear algebra applications. It is also an effective test-bed because different data dependency patterns can be examined simply by using different input matrices. In these experiments, a set of 6 input matrices, drawn from the University of Florida Sparse Matrix Collection Davis and Hu [2011], was used. The matrices were selected so that they would vary in overall data footprint, from 45 MB to 892 MB, and in percentage of non-zeros, from very sparse at 0.0006% to much more dense at 0.5539% non-zeros.

Table 3.4 compares the performance of the tiled Jacobi solver to that of a simple blocked version. Both codes use OpenMP parallel for directives to achieve parallelism. The execution time reduction varied from 13% to 47% with the exception of the *nd24k* matrix, which showed as much as a 1.52x slowdown when sparse tiled. This matrix is highly connected, thus limiting the number of tiles that can be scheduled in parallel. The greater parallelism available under a blocked approach provides more benefit in this case than the performance improvements due to improved locality from full sparse tiling. Overall, speed-ups of between 8 and 12 times over the single-threaded tiled implementation were observed when using 15 threads; a clear outlier is again the *nd24k* matrix that did not scale past 3.2

<sup>&</sup>lt;sup>5</sup>Seigen has been developed by Christian Jacobs.

<sup>&</sup>lt;sup>6</sup>This section is partly extracted from Strout et al. [2014], and the results were obtained from experiments conducted by Christopher D. Krieger, Catherine Olschanowsky, and Michelle Mills Strout

Implementation	Execution time	Speed-up
Westmere omp	36.87	6.43
Westmere mpi	31.0	7.66
Westmere tiled	26.49	8.96
Sandy Bridge omp	30.01	6.65
Sandy Bridge mpi	24.42	8.17
Sandy Bridge tiled	20.63	9.67

Table 3.5: Execution time (in seconds) and speed-ups over the slowest single-threaded implementation for the Airfoil benchmark. The values are obtained from simulations with fully-loaded machines (16 and 24 threads/processes on the Sandy Bridge and the Westmere architectures, respectively).

times the single thread performance.

The values in Table 3.4 do not include the inspection time necessary to full sparse tile the loop chain. To break even when this cost is considered, the inspector time must be amortized over between 1000 and 3000 iterations of the executor, depending on the specific matrix being solved. We will further elaborate on this aspect in Section 3.9.1.

#### Airfoil

Airfoil is a representative unstructured mesh application? Three implementations of Airfoil, *omp*, *mpi* and *tiled*, were compared on two shared-memory machines, an Intel Westmere (dual-socket 6-core Intel Xeon X5650 2.66 GHz, 12MB of shared L3 cache per socket) and an Intel Sandy Bridge (dual-socket 8-core Intel Xeon E5-2680 2.00Ghz, 20MB of shared L3 cache per socket). The code was compiled using the Intel icc 2013 compiler with optimizations enabled (-03, -xSSE4.2/-xAVX).

The Airfoil code consists of a main time loop with 2000 iterations. This loop contains a sequence of four parallel loops that carry out the computation. In this sequence, the first two loops, called *adt-calc* and *res-calc*, constitute the bulk of the computation. *Adt-calc* iterates over cells, reads from adjacent vertices and write to a local dataset, whereas *res-calc* iterates over edges and exploits indirect mappings to vertices and cells for incrementing indirect datasets associated to cells. These loops share datasets associated with cells and vertices. Datasets are composed of doubles.

In the *omp* and *mpi* implementations of Airfoil, the OpenMP and the MPI back-ends of OP2, were used. The effectiveness of these parallelization schemes has been demonstrated in Giles et al. [2011]. The *tiled* implementation uses an early version of the SLOPE library (the differences with the inspector algorithms shown in Section 3.7.2 are discussed later) for tiling a loop chain. We manually unrolled the time loop by a factor of two to be able to tile over 6 loops in total.

Table 3.5 shows the scalability and runtime reduction achieved by sparse tiling the loop chain on the Westmere and Sandy Bridge architectures. The input unstructured mesh was composed of 1.5 million edges. It is worth noticing that both the *omp* and *tiled* versions suffer from the well-known NUMA effect as threads are always equally spread across the two sockets. Nevertheless, compared to *mpi*, the *tiled* version exhibits a peak runtime reduction of 15% on the Westmere and of 16% on the Sandy Bridge.

Results shown for *tiled* do not include, however, the inspector overhead. By also including it, the aforementioned improvements over *mpi* reduce to roughly 10% on both platforms. Similarly to the sparse Jacobi solver, the slow-downs when including the inspection overhead are significant.

#### Observations

One of the most important outcomes of this first set of experiments concerns the inspection cost, which we found to significantly impact the overall execution time. To effectively support real applications like Seigen, typically characterized by a number of parallel loops larger than that of these benchmarks, we had to rework the original inspector algorithms into the versions described in Section 3.7.2. The most notable differences are: (i) data dependency analysis abstracted to the level of sets, rather than datasets; (ii) optimistic coloring with backtracking in case of conflicts; (iii) fully parallel projection and tiling phases, through the use of inverse maps.

A second observation is about the importance of automation. Writing the inspector as a sequence of calls to SLOPE is relatively simple, although tedious and error-prone. Much more complicated is integrating the executor, because this essentially means rewriting entire sequences of loops – a severe limitation when using SLOPE in plain C (perhaps legacy) code.

These considerations on the implementation complexity led to the multilayer framework detailed in Section 3.8 for automating the generation of inspector/executor schemes.

The Airfoil experiments highlighted that shared memory parallelism over multi-socket architectures is considerably affected by the NUMA issue. The difference in execution time between the OP2 OpenMP and MPI versions is indeed remarkable. As discussed in ?, the irregular nature of the computation makes it hard to find systematic solutions to the NUMA problem when relying on OpenMP. The MPI and the hybrid OpenMP/MPI execution models are simply better candidates for unstructured mesh computations. The sparse tiled implementation, which in these experiments is also purely based on OpenMP, suffers from the NUMA issue as well, although it manages to outperform the OP2 MPI version thanks to the relieved memory pressure. We wondered, however, what the gain would be if we managed to integrate MPI with sparse tiling. This led to the definition of more general loop chain abstraction (Section 3.5.3) and inspector algorithms (Section 3.7.2), for supporting MPI-based executors.

### 3.9.2 Seigen: an Elastic Wave Equation Solver for Seismological Problems

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#### 3.10 Conclusions and Future Work

# Chapter 4

# On Optimality of Finite Element Integration

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## Chapter 5

# Cross-loop Optimization of Arithmetic Intensity for Finite Element Integration

#### 5.1 Recapitulation and Objectives

In Chapter 4, we have developed a method to minimize the operation count of finite element operators, or "assembly kernels". This chapter focuses on the same class of kernels, but tackles an orthogonal issue: the low level optimization of the generated code. We will abstract from the mathematical structure inherent in the expressions and concentrate on the aspects impacting the computational efficiency.

We know that an assembly kernel is characterized by the presence of an affine, often non-perfect loop nest, in which individual loops are rather small: their trip count rarely exceeds 30, and may be as low as 3 for low order methods. In the innermost loop, a problem-specific, compute intensive expression evaluates a two dimensional array, representing the result of local assembly in an element of the discretized domain. With such a kernel structure, we focus on aspects like register locality and SIMD vectorization.

We aim to maximize our impact on the platforms that are realistically used for finite element applications, so we target conventional CPU architectures rather than GPUs. The key limiting factor to the execution on GPUs is the stringent memory requirements. Only relatively small prob-

lems fit in a GPU memory, and support for distributed GPU execution in general purpose finite element frameworks is minimal. There has been some research on adapting local assembly to GPUs, although it differs from ours in several ways, including: (i) not relying on automated code generation from a domain-specific language (explained next), (ii) testing only very low order methods, (iii) not optimizing for cross-loop arithmetic intensity (the goal is rather effective multi-thread parallelization). In addition, our code transformations would drastically impact the GPU parallelization strategy, for example by increasing a thread's working set. For all these reasons, a study on extending the research to GPU architectures is beyond the scope of this work. In Section 5.6, however, we provide some intuitions about this research direction.

Achieving high-performance on CPUs is non-trivial. The complexity of the mathematical expressions, which we know to be often characterized by a large number of operations on constants and small vectors, makes it hard to determine a single or specific sequence of transformations that is successfully applicable to all problems. Loop trip counts are typically small and can vary significantly, which further exacerbates the issue. The complexity of the memory access pattern also depends on the kernel, specifically on the function spaces employed by the method, ranging from unitstride (e.g., A[i], A[i+1], A[i+2], A[i+3], ...) to random-stride (e.g., A[i], A[i+1], A[i+2], A[i+N], A[i+N+1], ...). We will show that traditional vendor compilers, such as GNU's and Intel's, fail at maximizing the efficiency of the generated code because of such a particular structure. Polyhedralmodel-based source-to-source compilers, for instance Bondhugula et al. [2008], can apply aggressive loop optimizations, such as tiling, but these are not particularly helpful in our context since they mostly focus on cache locality.

Like in Chapter 4, we focus on optimizing the performance of assembly kernels produced through automated code generation, so we seek transformations that are generally applicable and effective. In particular, we will study the following transformations:

**Padding and data alignment** SIMD vectorization is more effective when the CPU registers are packed (unpacked) by means of aligned load (store) instructions. Data alignment is achieved through array padding, a conceptually simple yet powerful transformation that can result in dramatic

reductions in execution time. We will see that the complexity of the transformation increases if non unite-stride memory accesses are present.

**Vector-register tiling** Blocking at the level of vector registers aims to improve data locality. This transformation exploits the peculiar memory access pattern inherent in finite element operators (i.e., inner products involving test and trial functions).

**Expression splitting** Complex expressions are often characterized by high register pressure (i.e., the lack of available registers inducing the compiler to "spill" data from registers to cache). This happens, for example, when the number of arrays (e.g., basis functions, temporaries introduced by generalized code motion, temporaries produced by pre-evaluation) and constants is large compared to the number of available registers (typically 16 on state-of-the-art CPUs, 32 on future generations). This transformation exploits the associativity of addition to distribute, or "split", an expression into multiple sub-expressions; each sub-expression is then computed in a separate loop nest.

We will also provide insights into the effects of more "traditional" compiler optimizations, such as loop unroll, loop interchange, loop fusion and vector promotion.

To summarize, the contributions of this chapter are as follows:

- A number of low level transformations for optimizing the performance of assembly kernels. Some of these transformations are directly inspired by the structure of assembly kernels.
- Extensive experimentation using a set of real-world forms commonly arising in finite element methods.
- A discussion concerning the generality of the transformations and their applicability to different domains.

#### 5.2 Low-level Optimization

#### 5.2.1 Padding and Data Alignment

The absence of stencils renders the local element matrix computation easily auto-vectorizable by a general-purpose compiler. Nevertheless, auto-vectorization is not efficient if data are not aligned to cache-line bound-

aries and if the length of the innermost loop is not a multiple of the vector length VL, especially when the loops are small as in local assembly.

Data alignment is enforced in two steps. Firstly, all arrays (but the element matrix, for reasons discussed shortly) are padded by rounding the innermost dimension to the nearest multiple of VL. For instance, assume the original size of a basis function array is  $3\times3$  and VL = 4 (e.g. AVX processor, with 32-byte long vector registers and 8-byte double-precision floats). In this case, a padded version of the array will have size  $3\times4$ . Secondly, their base address is enforced to multiples of VL by means of special attributes. The compiler is explicitly told about data alignment using suitable pragmas; for example, in the case of the Intel compiler, the annotation #pragma vector aligned is added before the loop (as shown in later figures) to inform that all of the memory accesses in the loop body will be properly aligned. This allows the compiler to issue aligned load and store instructions, which are notably faster than unaligned ones.

In our computational model, the element matrix is one of the kernel's input parameters, so it needs special handling when padding (the signature of the kernel must not be changed, otherwise the abstraction would be broken). We create a "shadow" copy of the element matrix, padded, aligned, and initialized to 0. The shadow element matrix is used in place of the original element matrix. Right before returning to the caller, a loop nest copies, discarding the padded region, the shadow matrix back into the input buffer.

Array padding also allows to safely round the loop trip count to the nearest multiple of VL. This avoids the introduction of a remainder (scalar) loop from the compiler, which would render vectorization less efficient. These extra iterations only write to the padded region of the element matrix, and therefore have no side effects on the final result.

Listing 1 illustrates the effect of padding and data alignment on top of generalized code motion applied to the weighted Laplace operator presented in Listing 1.

#### 5.2.2 Expression Splitting

In complex kernels, like Burgers in Listing 2, and on certain architectures, achieving effective register allocation can be challenging. If the number

**LISTING 1:** The assembly kernel for the weighted Laplace operator in Listing 1 after application of padding and data alignment (on top of generalized code motion). An AVX architecture, which implies VL = 4, is assumed.

```
void weighted_laplace(double A[3][3], double **coords, double w[3]) {
     #define ALIGN __attribute__((aligned(32)))
     // K, det = Compute Jacobian (coords)
     // Quadrature weights
     static const double W[6] ALIGN = 0.5;
     // Basis functions
     static const double B[6][4] ALIGN = \{\{...\}\};
     static const double C[6][3] ALIGN = \{\{...\}\}\; static const double D[6][4] ALIGN = \{\{...\}\}\;
10
11
12
     // Padded buffer
13
     double A[3][4] ALIGN = {{0.0}};
14
15
     for (int i = 0; i < 6; i++) {
16
       double f0 = 0.0;
17
18
       for (int r = 0; r < 3; ++r) {
         f0 += (w[r] * C[i][r]);
19
20
21
       double T_0[4] ALIGN;
       double T_1[4] ALIGN;
22
23
       #pragma vector aligned
       for (int k = 0; k < 4; r++) {
24
         T_0[r] = ((K[1]*B[i][k])+(K[3]*D[i][k]));
25
26
         T_{-1}[r] = ((K[0]*B[i][k])+(K[2]*D[i][k]));
27
       for (int j = 0; j < 3; j++) {
28
         #pragma vector aligned
         for (int k = 0; k < 4; k++) {
30
            A[j][k] += (T_0[k]*T_0[j] + T_1[k]*T_1[j])*det*W[i]*f0);
31
32
33
34
35
36 for (int j = 0; j < 3; j++) {
     for (int k = 0; k < 3; k++) {
       A[j][k] = A[j][k];
38
39
40
```

of variables independent of the innermost-loop dimension is close to or greater than the number of available CPU registers, poor register reuse is likely. This usually happens when the number of basis function arrays, temporaries introduced by either generalized code motion or preevaluation, and problem constants is large. For example, applying code motion to the Burgers example on a 3D mesh requires 24 temporaries for the ijk loop order. This can make hoisting of the invariant loads out of the k loop inefficient on architectures with a relatively low number of registers. One potential solution to this problem consists of suitably "splitting" the computation of the element matrix *A* into multiple sub-expressions. An example of this idea is given in Listing 2. The transformation can be regarded as a special case of classic loop fission, in which associativity of the sum is exploited to distribute the expression across multiple loops. To the best of our knowledge, expression splitting is not supported by available compilers.

**LISTING 2:** The assembly kernel for the weighted Laplace operator in Listing 1 after application of expression splitting (on top of generalized code motion). In this example, the split factor is 2.

```
void weighted_laplace(double A[3][3], double **coords, double w[3]) {

// Omitting redundant code

for (int j = 0; j<3; j++) {

for (int k = 0; k<3; k++) {

    A[j][k] += (T_0[k]*T_0[j])*det*W[i]*f0;

}

for (int j = 0; j<3; j++) {

    for (int j = 0; j<3; j++) {

        for (int k = 0; k<3; k++) {

            A[j][k] += (T_1[k]*T_1[j])*det*W[i]*f0;

        }

        A[j][k] += (T_1[k]*T_1[j])*det*W[i]*f0;

        }

    }

</pre>
```

Splitting an expression (henceforth *split*) has, however, several drawbacks. Firstly, it increases the number of accesses to A in proportion to the "split factor", which is the number of sub-expressions produced. Also, depending on how splitting is done, it can lead to redundant computation. For example, the number of times the product det\*W3[i] is performed is proportional to the number of sub-expressions, as shown in the code snippet. Further, it increases loop overhead, for example through additional branch instructions. Finally, it might affect register locality: for instance,

the same array could be accessed in different sub-expressions, requiring a proportional number of loads be performed; this is not the case of the running example, though. Nevertheless, the performance gain from improved register reuse can still be greater if suitable heuristics are used. Our approach consists of traversing the expression tree and recursively splitting it into multiple sub-expressions as long as the number of variables independent of the innermost loop exceeds a certain threshold. This is elaborated in the next sections, and validated against empirical search in Section 5.3.2.

#### 5.2.3 Model-driven Vector-register Tiling

One notable problem of assembly kernels concerns register allocation and register locality. The critical situation occurs when the loop trip counts and the variables accessed are such that the vector-register pressure is high. Since the kernel's working set is expected to fit the L1 cache, it is particularly important to optimize register management. Standard optimizations, such as loop interchange, unroll, and unroll-and-jam, can be employed to deal with this problem. Tiling at the level of vector registers represents another opportunity. Based on the observation that the evaluation of the element matrix can be reduced to a summation of outer products along the j and k dimensions, a model-driven vector-register tiling strategy can be implemented. If we consider the codes in the various listings and we focus on the body of the test and trial functions loops (j and k), the computation of the element matrix is abstractly expressible as

$$A_{jk} = \sum_{\substack{x \in B' \subseteq B \\ y \in B'' \subseteq B}} x_j \cdot y_k \qquad j, k = 0, ..., 2$$
 (5.1)

where B is the set of all basis functions or temporary variables accessed in the kernel, whereas B' and B'' are generic problem-dependent subsets. Regardless of the specific input problem, by abstracting from the presence of all variables independent of both j and k, the element matrix computation is always reducible to this form. Figure 5.1 illustrates how we can evaluate 16 entries (j, k = 0, ..., 3) of the element matrix using just 2 vector registers, which represent a 4×4 tile, assuming |B'| = |B''| = 1. Values in a register are shuffled each time a product is performed. Standard compiler

**LISTING 3:** The assembly kernel for the weighted Laplace operator in Listing 1 after application of vector-register tiling (on top of generalized code motion, padding, and data alignment). In this example, the unroll-and-jam factor is 1.

```
void weighted_laplace(double A[3][3], double **coords, double w[3]) {
    // Omitting redundant code
     // Padded buffer (note: both rows and columns)
     double A[4][4] ALIGN = {{0.0}};
     for (int i = 0; i < 3; i++) {
       // Omitting redundant code
       // ...
       for (int j = 0; j < 4; j += 4)
10
11
         for (int k = 0; k < 4; k += 4) {
           // Sequence of LOAD and SET intrinsics
12
13
           // Compute _A[0][0], _A[1][1], _A[2][2], _A[3][3]
           // One _mm256_permute_pd per k-loop LOAD
14
           // Compute A[0][1], A[1][0], A[2][3], A[3][2]
15
           // One _mm256_permute2f128_pd per k-loop LOAD
17
18
19
       // Scalar remainder loop (not necessary in this example)
20
     // Restore the storage layout
21
     for (int j = 0; j < 4; j += 4) {
       for (int k = 0; k < 4; k += 4) {
23
24
         _{-m256d} r0 = _{mm256\_load\_pd} (&_{A[j+0][k]);}
         // LOAD _A[j+1][k], _A[j+2][k], _A[j+3][k]
25
26
         r4 = _mm256\_unpackhi\_pd (r1, r0);
27
         r5 = _mm256\_unpacklo\_pd (r0, r1);
         r6 = _mm256_unpackhi_pd (r2, r3);
28
29
         r7 = _mm256\_unpacklo\_pd (r3, r2);
30
         r0 = _mm256_permute2f128_pd (r5, r7, 32);
         r1 = _mm256_permute2f128_pd (r4, r6, 32);
31
         r2 = _mm256_permute2f128_pd (r7, r5, 49);
         r3 = _mm256_permute2f128_pd (r6, r4, 49);
33
         _mm256_store_pd (&_A[j+0][k], r0);
34
35
         // STORE _A[j+1][k], _A[j+2][k], _A[j+3][k]
36
37
38 }
39
```

auto-vectorization for both GNU and Intel compilers, instead, executes 4 broadcast operations (i.e., "splat" of a value over all of the register locations) along the outer dimension to perform the calculation. In addition to incurring a larger number of cache accesses, it needs to keep between f=1 and f=3 extra registers to perform the same 16 evaluations when unroll-and-jam is used, with f being the unroll-and-jam factor.

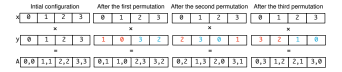


Figure 5.1: Outer-product vectorization by permuting values in a vector register.

The storage layout of *A*, however, is incorrect after the application of this outer-product-based vectorization (*op-vect*, in the following). It can be efficiently restored with a sequence of vector shuffles following the pattern highlighted in Figure 5.2, executed once outside of the ijk loop nest. The pseudo-code for the weighted Laplace assembly kernel using *op-vect* is shown in Listing 3.

Initial configuration			 After the first set of shuffles			After the second set of shuffles			Layout restored						
(0, 0)	(1, 1)	(2, 2)	(3, 3)	(0, 0)	(1, 1)	(2, 2)	(3, 3)	(0, 0)	(0, 1)	(2, 2)	(2, 3)	(0, 0)	(0, 1)	(0, 2)	(0, 3)
(0, 1)	(1, 0)	(2, 3)	(3, 2)	(0, 1)	(1, 0)	(2, 3)	(3, 2)	(1, 0)	(1, 1)	(3, 2)	(3, 3)	(1, 0)	(1, 1)	(1, 2)	(1, 3)
(0, 2)	(1, 3)	(2, 0)	(3, 1)	(0, 2)	(1, 3)	(2, 0)	(3, 1)	(0, 2)	(0, 3)	(2, 0)	(2, 1)	(2, 0)	(2, 1)	(2, 2)	(2, 3)
(0, 3)	(1, 2)	(2, 1)	(3, 0)	(0, 3)	(1, 2)	(2, 1)	(3, 0)	(1, 2)	(1, 3)	(3, 0)	(3, 1)	(3, 0)	(3, 1)	(3, 2)	(3, 3)

Figure 5.2: Restoring the storage layout after *op-vect*. The figure shows how  $4\times4$  elements in the top-left block of the element matrix A can be moved to their correct positions. Each rotation, represented by a group of three same-colored arrows, is implemented by a single shuffle intrinsic.

#### 5.3 Experiments

#### 5.3.1 Setup

The objective is to evaluate the impact of the code transformations presented in the previous sections in three representative PDEs, which we refer to as (i) Helmholtz, (ii) Diffusion, and (iii) Burgers.

The three chosen equations are *real-life kernels* and comprise the core differential operators in some of the most frequently encountered finite element problems in scientific computing. This is of crucial importance because distinct problems, possibly arising in completely different fields, may employ (subsets of) the same differential operators of our benchmarks, which implies similarities and redundant patterns in the generated code. Consequently, the proposed code transformations have a domain of applicability that goes far beyond that of the three analyzed equations.

The Helmholtz and Diffusion kernels are archetypal second order elliptic operators. They are complete and unsimplified examples of the operators used to model diffusion and viscosity in fluids, and for imposing pressure in compressible fluids. As such, they are both extensively used in climate and ocean modeling. Very similar operators, for which the same optimisations are expected to be equally effective, apply to elasticity problems, which are at the base of computational structural mechanics. The Burgers kernel is a typical example of a first order hyperbolic conservation law, which occurs in real applications whenever a quantity is transported by a fluid (the momentum itself, in our case). We chose this particular kernel since it applies to a vector-valued quantity, while the elliptic operators apply to scalar quantities; this impacts the generated code, as explained next. The operators we have selected are characteristic of both the second and first order operators that dominate fluids and solids simulations.

The benchmarks were written in UFL (code available at [Luporini, 2014b]) and executed over real unstructured meshes through Firedrake. The Helmholtz code has already been shown in Listing ??. The Diffusion equation uses the same differential operators as Helmholtz. In the Diffusion kernel code, the main differences with respect to Helmholtz are the absence of the *Y* array and the presence of additional constants for computing the element matrix. Burgers is a non-linear problem employing differential operators different from those of Helmholtz and relying on vector-valued quantities, which has a major impact on the generated assembly code (see Listing 2), where a larger number of basis function arrays (*X*1, *X*2, ...) and constants (*F*0, *F*1, ..., *K*0, *K*1,...) are generated.

These problems were studied varying both the shape of mesh elements and the polynomial order q of the method, whereas the element family, Lagrange, is fixed. As might be expected, the larger the element shape and

q, the larger the iteration space. Triangles, tetrahedra, and prisms were tested as element shape. For instance, in the case of Helmholtz with q=1, the size of the j and k loops for the three element shapes is, respectively, 3, 4, and 6. Moving to bigger shapes has the effect of increasing the number of basis function arrays, since, intuitively, the behaviour of the equation has to be approximated also along a third axis. On the other hand, the polynomial order affects only the problem size (the three loops i, j, and k, and, as a consequence, the size of X and Y arrays). A range of polynomial orders from q=1 to q=4 were tested; higher polynomial orders are excluded from the study because of current Firedrake limitations. In all these cases, the size of the element matrix rarely exceeds  $30 \times 30$ , with a peak of  $105 \times 105$  in Burgers with prisms and q=4.

#### 5.3.2 Impact of Transformations

Experiments were run on a single core of an Intel architecture, a Sandy Bridge I7-2600 CPU running at 3.4 GHz, with 32KB of L1 cache and 256KB of L2 cache). The icc 14.1 compiler was used. On the Sandy Bridge, the compilation flags used were -02 and -xAVX for auto-vectorization (other optimization levels were tried, but they generally resulted in higher execution times).

The speed-ups achieved by applying the transformations on top of the original assembly kernel code are shown in Figure 5.3. This figure is a three-dimensional plot: element shape and equation vary along the outermost axes, whereas *q* varies within each sub-plot. In the next sections, we will refer to this figure and elaborate on the impact of the individual transformations. We shorten generalized loop-invariant code motion as *licm*; padding and data alignment as *ap*; outer-product vectorization as *op-vect*; expression splitting as *split*.

#### Impact of Generalized Loop-invariant Code Motion

In general, the speed-ups achieved by *licm* are notable. The main reasons were anticipated in Section ??: in the original code, 1) sub-expressions invariant to outer loops are not automatically hoisted, while 2) sub-expressions invariant to the innermost loop are hoisted, but their execution is not autovectorized. These observations come from inspection of assembly code

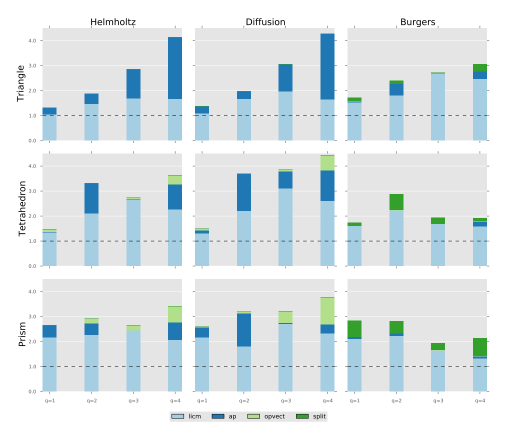


Figure 5.3: Performance improvement due to generalized loop-invariant code motion (*licm*), data alignment and padding (*ap*), outerproduct vectorization (*op-vect*), and expression splitting (*split*) over the original non-optimized code. In each plot, the horizontal axis reports speed ups, whereas the polynomial order *q* of the method varies along the vertical axis.

#### generated by the compiler.

The gain tends to grow with the computational cost of the kernels: bigger loop nests (i.e., larger element shapes and polynomial orders) usually benefit from the reduction in redundant computation, even though extra memory for the temporary arrays is required. Some discrepancies to this trend are due to a less effective auto-vectorization. For instance, on the Sandy Bridge, the improvement at q=3 is larger than that at q=4 because, in the latter case, the size of the innermost loop is not a multiple of the vector length, and a remainder scalar loop is introduced at compile time. Since the loop nest is small, the cost of executing the extra scalar

iterations can have a significant impact.

#### Impact of Padding and Data Alignment

Padding, which avoids the introduction of a remainder loop as described in Section 5.2.1, as well as data alignment, enhance the quality of autovectorization. Occasionally the impact of *ap* is marginal. These may be due to two reasons: (i) the non-padded element matrix size is already a multiple of the vector length; (ii) the number of aligned temporaries introduced by *licm* is so large to induce cache associativity conflicts (e.g. Burgers equation).

#### Impact of Vector-register Tiling

In this section, we evaluate the impact of vector-register tiling. *op-vect* requires the unroll-and-jam factor to be explicitly set. Here, we report the best speed-up obtained after all feasible unroll-and-jam factors were tried.

The rationale behind these results is that the effect of *op-vect* is significant in problems in which the assembly loop nest is relatively big. When the loops are short, since the number of arrays accessed at every loop iteration is rather small (between 4 and 8 temporaries, plus the element matrix itself), there is no need for vector-register tiling; extensive unrolling is sufficient to improve register re-use and, therefore, to maximize the performance. However, as the iteration space becomes larger, *op-vect* leads to improvements up to  $1.4\times$  (Diffusion, prismatic mesh, q=4 - increasing the overall speed up from  $2.69\times$  to  $3.87\times$ ).

Using the Intel Architecture Code Analyzer tool Intel Corporation [2012], we confirmed that speed ups are a consequence of increased register reuse. In Helmholtz q=4, for example, the tool showed that when using *op-vect* the number of clock cycles to execute one iteration of the j loop decreases by roughly 17%, and that this is a result of the relieved pressure on both of the data (cache) ports available in the core.

The performance of individual kernels in terms of floating-point operations per second was also measured. The theoretical peak on a single core, with the Intel Turbo Boost technology activated, is 30.4 GFlop/s. In the case of Diffusion using a prismatic mesh and q=4, we achieved a maximum of 21.9 GFlop/s with op-vect enabled, whereas 16.4 GFlop/s was

obtained when only *licm-ap* is used. This result is in line with the expectations: analysis of assembly code showed that, in the jk loop nest, which in this problem represents the bulk of the computation, 73% of instructions are actually floating-point operations.

Application of *op-vect* to the Burgers problem induces significant slow-downs due to the large number of temporary arrays that need to be tiled, which exceeds the available logical registers on the underlying architecture. Expression splitting can be used in combination with *op-vect* to alleviate this issue; this is discussed in the next section.

#### Impact of Expression Splitting

Expression splitting relieves the register pressure when the element matrix evaluation needs to read from a large number of basis function arrays. As detailed in Section 5.2.2, the price to pay for this optimization is an increased number of accesses to the element matrix and, potentially, redundant computation.

For the Helmholtz and Diffusion kernels, in which only between 4 and 8 temporaries are read at every loop iteration, split tends to slow down the computation, because of the aforementioned drawbacks. Slow downs up to  $1.4\times$  were observed.

In the Burgers kernels, between 12 and 24 temporaries are accessed at every loop iteration, so *split* plays a key role since the number of available logical registers on the Sandy Bridge architecture is only 16. In almost all cases, a split factor of 1, meaning that the original expression was divided into two parts, ensured close-to-peak perforance. The transformation negligibly affected register locality, so speed ups up to  $1.5\times$  were observed. For instance, when q=4 and a prismatic mesh is employed, the overall performance improvement increases from  $1.44\times$  to  $2.11\times$ .

The performance of the Burgers kernel on a prismatic mesh was 20.0 GFlop/s from q = 1 to q = 3, while it was 21.3 GFlop/s in the case of q = 4. These values are notably close to the peak performance of 30.4 GFlop/s. Disabling *split* makes the performance drop to 17.0 GFlop/s for q = 1,2,18.2 GFlop/s for q = 3, and 14.3 GFlop/s for q = 4. These values are in line with the speed-ups shown in Figure 5.3.

The split transformation was also tried in combination with op-vect (split-

op-vect). Despite improvements up to 1.22×, split-op-vect never outperforms split. This is motivated by two factors: for small split factors, such as 1 and 2, the data space to be tiled is still too big, and register spilling affects run-time; for higher ones, sub-expressions become so small that, as explained in Section 5.3.2, extensive unrolling already allows to achieve a certain degree of register re-use.

# 5.4 Experience with Traditional Compiler Optimizations

#### 5.4.1 Loop Interchange

All loops are interchangeable, provided that temporaries are introduced if the nest is not perfect. For the employed storage layout, the loop permutations ijk and ikj are likely to maximize the performance. Conceptually, this is motivated by the fact that if the i loop were in an inner position, then a significantly higher number of load instructions would be required at every iteration. We tested this hypothesis in manually crafted kernels. We found that the performance loss is greater than the gain due to the possibility of accumulating increments in a register, rather than memory, along the i loop. The choice between ijk and ikj depends on the number of load instructions that can be hoisted out of the innermost dimension. A good heuristics it to choose as outermost the loop along which the number of invariant loads is smaller so that more registers remain available to carry out the computation of the local element matrix.

Our experience with the Intel's and GNU's compilers is controversial: if, from one hand, the former applies this transformation following a reasonable cost model, the latter results in general more conservative, even at highest optimization level. This behaviour was verified in different variational forms (by looking at assembly code and compiler reports), including the complex hyperelastic model analyzed in Chapter 4.

#### 5.4.2 Loop Unroll

Loop unroll (or unroll-and-jam of outer loops) is fundamental to the exposure of instruction-level parallelism, and tuning unroll factors is partic-

ularly important.

We first observe that manual full (or extensive) unrolling is unlikely to be effective for two reasons. Firstly, the ijk loop nest would need to be small enough such that the unrolled instructions do not exceed the instruction cache, which is rarely the case: it is true that in a local assembly kernel the minimum size of the ijk loop nest is  $3\times3\times3$  (triangular mesh and polynomial order 1), but this increases rapidly with the polynomial order of the method and the discretization employed (e.g. tetrahedral meshes imply larger loop nests than triangular ones), so sizes greater than  $10\times10\times10$ , for which extensive unrolling would already be harmful, are in practice very common. Secondly, manual unrolling is dangerous because it may compromise compiler auto-vectorization by either removing loops (most compilers search for vectorizable loops) or losing spatial locality within a vector register.

By comparison to implementations with manually-unrolled loops, we noticed that recent versions of compilers like GNU's and Intel's estimate close-to-optimal unroll factors when the loops are affine and their bounds are relatively small and known at compile-time, which is the case of our kernels. Our choice, therefore, is to leave the back-end compiler in charge of selecting unroll factors.

#### 5.4.3 Vector promotion

Vector promotion is a transformation that "trades" space in exchange of a parallel dimension (a "clone" of the integration loop), thus promoting SIMD vectorization at the level of an outer loop.

Consider Listing 4. The evaluation of the coefficient w at each quadrature point can be vectorized by "promoting" f from a scalar to a vector of size 3. Any other sub-expression hoisted at the level of the integration loop (as described in Chapter 4) can be transformed in a similar way. The impact of this optimization obviously increases with the number of operations involving coefficients. At the same time, the allocation of extra memory may lead to the same issues described in Section ??. Loop tiling could be used to counteract this negative effect, although this would significantly increase the implementation complexity.

We have not seen this transformation being applied by neither the GNU's

**LISTING 4:** The assembly kernel for the weighted Laplace operator in Listing 1 after application of vector promotion (on top of generalized code motion).

```
void weighted_laplace(double A[3][3], double **coords, double w[3]) {
     // Omitting redundant code
     double f0[3] = \{0.0\};
     for (int i = 0; i < 6; i + +) {
       for (int r = 0; r < 3; ++r) {
         f0[i] += (w[r] * C[i][r]);
8
     for (int i = 0; i < 6; i++) {
10
       double T_0[3] ALIGN;
11
       double T_1[3] ALIGN;
12
13
       for (int k = 0; k < 3; r++) {
         T_0[r] = ((K[1]*B[i][k])+(K[3]*D[i][k]));
14
15
         T_{-1}[r] = ((K[0]*B[i][k])+(K[2]*D[i][k]));
16
       for (int j = 0; j < 3; j++) {
17
         for (int k = 0; k < 3; k++) {
18
           A[j][k] += (T_0[k]*T_0[j] + T_1[k]*T_1[j])*det*W[i]*f0[i]);
19
20
21
       }
22
     }
23
```

nor the Intel's compilers. In our experience – and in absence of loop tiling – the impact on execution time is difficult to predict. This transformation requires further investigation. Despite being fully implemented in COFFEE, it is therefore not applied in the default optimization process.

#### 5.4.4 Loop Fusion

Loop fusion is a well-known compiler transformation that consists of merging a sequence of loops into a single one. This optimization can be applied by most general-purpose compilers. What we cannot expect these compilers to do, however, is identifying common sub-expressions across the fused loops – an optimization of domain-specific nature.

In assembly kernels arising from bilinear forms, test and trial functions may belong to the same function space. More interestingly, the same operators could be applied to both sets of functions. This would result in both linear loops having the same iteration space and common sub-expressions arising across them. To avoid this kind of redundant computation and simultaneously enforcing fusion, we implemented in COFFEE a specialized version of loop fusion. In our experiments, this optimization always re-

sulted in relatively small performance improvements, ranging between 2% and 8%. Therefore, it is automatically enabled in the default optimization process.

#### 5.5 Related Work

The code transformations presented are inspired by standard compilers optimizations and exploit several domain properties. Our loop-invariant code motion technique individuates invariant sub-expressions and redundant computation by analyzing all loops in an iteration space, which is a generalization of the algorithms often implemented by general-purpose compilers. Expression splitting is an abstract variant of loop fission based on properties of arithmetic operators. The outer-product vectorization is an implementation of tiling at the level of vector registers; tiling, or "loop blocking", is commonly used to improve data locality (especially for caches). Padding has been used to achieve data alignment and to improve the effectiveness of vectorization. A standard reference for the compilation techniques re-adapted in this work is [Aho et al., 2007].

Our compiler-based optimization approach is made possible by the top-level DSL, which enables automated code generation. DSLs have been proven successful in auto-generating optimized code for other domains: Spiral [Püschel et al., 2005] for digital signal processing numerical algorithms, [Spampinato and Püschel, 2014] for dense linear algebra, or Pochoir [Tang et al., 2011] and SDSL [Henretty et al., 2013] for image processing and finite difference stencils. Similarly, PyOP2 is used by Firedrake to express iteration over unstructured meshes in scientific codes. COFFEE improves automated code generation in Firedrake.

Many code generators, like those based on the Polyhedral model [Bondhugula et al., 2008] and those driven by domain-knowledge [Stock et al., 2011], make use of cost models. The alternative of using auto-tuning to select the best implementation for a given problem on a certain platform has been adopted by nek5000 [Shin et al., 2010] for small matrix-matrix multiplies, the ATLAS library [Whaley and Dongarra, 1998], and FFTW [Frigo and Johnson, 2005] for fast fourier transforms. In both cases, pruning the implementation space is fundamental to mitigate complexity and overhead. Likewise, COFFEE uses heuristics and a model-driven auto-tuning

#### 5.6 Applicability to Other Domains

We have demonstrated that our cross-loop optimizations for arithmetic intensity are effective in the context of automated code generation for finite element integration. In this section, we discuss their applicability in other computational domains and, in general, their integrability within a general-purpose compiler.

There are neither conceptual nor technical reasons which prevent our transformations from being used in other (general-purpose, research, ...) compilers. It is challenging, however, to assess the potential of the presented optimizations is another computational domains, and to what extent they would be helpful for improving the full application performance. To answer these questions, we first need to go back to the origins of our study. The starting point of our work was the mathematical formulation of a finite element operator, expressible as follows

$$\forall_{i,j} \quad A_{ij}^K = \sum_{q=1}^{n_1} \sum_{k=1}^{n_2} \alpha_{k,q}(a',b',c',...) \beta_{q,i,j}(a,b,c,d,...) \gamma_q(w_K, z_K)$$
(5.2)

The expression represents the numerical evaluation of an integral at  $n_1$  points in the mesh element K computing the local element matrix A. Functions  $\alpha$ ,  $\beta$  and  $\gamma$  are problem-specific and can be intricately complex, involving for example the evaluation of derivatives. We can however abstract from the inherent structure of  $\alpha$ ,  $\beta$  and  $\gamma$  to highlight a number of aspects

• Optimizing mathematical expressions. Expression manipulation (e.g. simplification, decomposition into sub-expressions) opens multiple semantically equivalent code generation opportunities, characterized by different trade-offs in parallelism, redundant computation, and data locality. The basic idea is to exploit properties of arithmetic operators, such as associativity and commutativity, to re-schedule the computation suitably for the underlying architecture. Loop-invariant code motion and expression splitting follow this principle, so they can be re-adapted or extended to any domains involving numerical evaluation of complex mathematical expressions (e.g. electronic structure calculations in

physics and quantum chemistry relying on tensor contractions Hartono et al. [2009]). In this context, we highlight three notable points.

- 1. In Equation (5.2), the summations correspond to reduction loops, whereas loops over indices *i* and *j* are fully parallel. Throughout the paper we assumed that a kernel will be executed by a single thread, which is likely to be the best strategy for standard multi-core CPUs. On the other hand, we note that for certain architectures (for example GPUs) this could be prohibitive due to memory requirements. Intra-kernel parallelization is one possible solution: a domain-specific compiler could map mathematical quantifiers and operators to different parallelization schemes and generate distinct variants of multi-threaded kernel code. Based on our experience, we believe this is the right approach to achieve performance portability.
- 2. The various sub-expressions in  $\beta$  only depend on (i.e. iterate along) a subset of the enclosing loops. In addition, some of these sub-expressions might reduce to the same values as iterating along certain iteration spaces. This code structure motivated the generalized loop-invariant code motion technique. The intuition is that whenever sub-expressions invariant with respect to different sets of affine loops can be identified, the question of whether, where and how to hoist them, while minimizing redundant computation, arises. Pre-computation of invariant terms also increases memory requirements due to the need for temporary arrays, so it is possible that for certain architectures the transformation could actually cause slowdowns (e.g. whenever the available per-core memory is small).
- 3. Associative arithmetic operators are the prerequisite for expression splitting. In essence, this transformation concerns resource-aware execution. In our context, expression splitting has successfully been applied to improve register pressure. However, the underlying idea of re-scheduling (re-associating) operations to optimize for some generic parameters is far more general. It could be used, for example, as a starting point to perform kernel fission; that is, splitting a kernel into multiple parts, each part characterized by

less stringent memory requirements (a variant of this idea for non-affine loops in unstructured mesh applications has been adopted in [Bertolli et al., 2013]). In Equation (5.2), for instance, not only can any of the functions  $\alpha$ ,  $\beta$  and  $\gamma$  be split (assuming they include associative operators), but  $\alpha$  could be completely extracted and evaluated in a separate kernel. This would reduce the working set size of each of the kernel functions, an option which is particularly attractive for many-core architectures in which the available per-core memory is much smaller than that in traditional CPUs.

- Code generation and applicability of the transformations. All array sizes and loop bounds, for example *n*1 and *n*2 in Equation 5.2, are known at code generation time. This means that "good" code can be generated. For example, loop bounds can be made explicit, arrays can be statically initialized, and pointer aliasing is easily avoidable. Further, all of these factors contribute to the applicability and the effectiveness of some of our code transformations. For instance, knowing loop bounds allows both generation of correct code when applying vector-register tiling and discovery of redundant computation opportunities. Padding and data alignment are special cases, since they could be performed at run-time if some values were not known at code generation time. Theoretically, they could also be automated by a general-purpose compiler through profile-guided optimization, provided that some sort of dataflow analysis is performed to ensure that the extra loop iterations over the padded region do not affect the numerical results.
- Multi-loop vectorization. Compiler auto-vectorization has become increasingly effective in a variety of codes. However, to the best of our knowledge, multi-loop vectorization involving the loading and storing of data along a subset of the loops characterizing the iteration space (rather than just along the innermost loop), is not supported by available general-purpose and polyhedral compilers. The outer-product vectorization technique presented in this paper shows that two-loop vectorization can outperform standard auto-vectorization. In addition, we expect the performance gain to scale with the number of vectorized loops and the vector length (as demonstrated in the Xeon Phi experiments). Although the automation of multi-loop vectorization in a

general-purpose compiler is far from straightforward, especially if stencils are present, we believe that this could be more easily achieved in specific domains. The intuition is to map the memory access pattern onto vector registers, and then to exploit in-register shuffling to minimize the traffic between memory and processor. By demonstrating the effectiveness of multi-loop vectorization in a real scenario, our research represents an incentive for studying this technique in a broader and systematic way.

#### 5.7 Conclusion

In this chapter, we have presented the study and systematic performance evaluation of a class of composable cross-loop optimizations for improving arithmetic intensity in finite element local assembly kernels. In the context of automated code generation for finite element local assembly, COFFEE is the first compiler capable of introducing low-level optimizations to simultaneously maximize register locality and SIMD vectorization. Assembly kernels have particular characteristics. Their iteration space is usually very small, with the size depending on aspects like the degree of accuracy one wants to reach (polynomial order of the method) and the mesh discretization employed. The data space, in terms of number of arrays and scalars required to evaluate the element matrix, grows proportionally with the complexity of the finite element problem. The various optimizations overcome limitations of current vendor and research compilers. The exploitation of domain knowledge allows some of them to be particularly effective, as demonstrated by our experiments on a state-ofthe-art Intel platform. The generality and the applicability of the proposed code transformations to other domains has also been discussed.

## Chapter 6

# **COFFEE: a Compiler for Fast Expression Evaluation**

#### 6.1 Overview

Sharing elimination and pre-evaluation, which we presented in Chapter 4, as well as the low level optimizations discussed in Chapter 5, have been implemented in COFFEE<sup>1</sup>, a mature, platform-agnostic compiler. COFFEE has fully been integrated with Firedrake, the framework based on the finite element method introduced in Section ??. The code, which comprises more than 5000 lines of Python, is available at [Luporini, 2014a].

Firedrake users employ the Unified Form Language to express problems in a notation resembling mathematical equations. At run-time, the high-level specification is translated by a form compiler, the Two-Stage Form Compiler (TSFC) ?, into one or more abstract syntax trees (ASTs) representing assembly kernels. ASTs are then passed to COFFEE for optimization. The output of COFFEE, C code, is eventually provided to PyOP2 [Markall et al., 2013], where just-in-time compilation and execution over the discretized domain take place. The flow and the compiler structure are outlined in Figure 6.1.

<sup>&</sup>lt;sup>1</sup>COFFEE is the acronym for COmpiler For Fast Expression Evaluation.

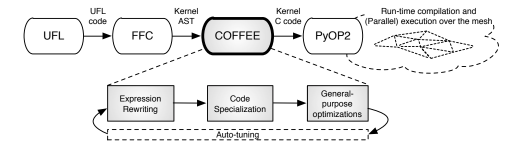


Figure 6.1: High-level view of Firedrake. COFFEE is at the core, receiving ASTs from a modified version of the FEniCS Form Compiler and producing optimized C code kernels.

#### 6.2 The Compilation Pipeline

Similarly to general-purpose compilers, COFFEE provides different optimization levels, namely 00, 01, 02 and 03. Apart from 00, which does not transform the code received from the form compiler (useful for debugging purposes), all optimization levels apply ordered sequences of optimizations. In essence, the higher the optimization level, the more aggressive (and potentially slower) is the transformation process. In the following, when describing aspects of the optimization process common to 01, 02 and 03, we will use the generic notation 0x ( $x \in \{1,2,3\}$ ).

The optimization level 0x can logically be split into three phases:

**Expression rewriting** Any transformation changing the structure of the expressions in the assembly kernel belongs to this class. For example, a high level optimization (sharing elimination, pre-evaluation) or, more in general, any rewrite operator (described later in Section 6.4) such as generalized code motion or factorization.

**Handling of block-sparse tables** Explained in Section ??, this phase consists of restructuring the iteration spaces searching for a trade-off between the avoidance of useless operations involving blocks of zeros in basis function tables and the effectiveness of low level optimization.

**Code Specialization** The class of low level optimizations. The primary focus of this thesis has been code specialization for conventional CPUs, although a generalization to other platforms is possible. In this phase, a

specific combination of the transformations presented in Chapter 6 is applied.

These three phases are totally ordered. Expression rewriting introduces temporaries and creates loops. All loops, including those produced by expression rewriting, and the statements therein are potentially transformed in the subsequent phase, by adjusting bounds and introducing memory offsets, respectively. The output of the first two phases is finally processed for padding and data alignment, vector-register tiling and vector promotion.

**Phase 1: analysis** During the analysis phase, an AST is visited and several kinds of information are collected. In particular, COFFEE searches for expression rewriting candidates. These are represented by special nodes in the AST, which we refer to as "expression nodes". In plain C, we could think of an expression node as a statement preceded by a directive such as #pragma coffee expression; the purpose of the directive would be to trigger COFFEE's 0x. This is for example similar to the way loops are parallelized through OpenMP. If at least one expression node is found, we proceed to the next phase, otherwise the AST is unparsed and C code returned.

Phase 2: checking legality In addition to 0x, users can craft their own custom optimization pipelines by composing the individual transformations available in COFFEE. However, since some of the low level transformations are inherently not composable (e.g., loop unrolling with vector-register tiling), the compiler always checks the legality of the transformation sequence.

**Phase 3: AST transformation** If the sequence of optimizations is legal, the AST is processed. In particular:

**01** At lowest optimization level, expression rewriting reduces to generalized code motion, while only padding and data alignment are applied among the low level optimizations.

**02** With respect to 01, there is only one yet fundamental change: expression rewriting now performs sharing elimination (i.e., Algorithm ??).

**03** Algorithm **??**, which coordinates sharing elimination and pre-evaluation, is applied. This is followed by handling block-sparse tables, and finally by padding and data alignment.

**Phase 4: code generation** Once all optimizations have been applied, the AST is visited one last time and a C representation (a string) is returned.

#### 6.3 Plugging COFFEE into Firedrake

#### 6.3.1 Abstract Syntax Trees

In this section, we highlight peculiarities of the hierarchy of AST nodes.

**Special nodes** Firstly, we observe that some nodes have special semantics. The expression nodes described in the previous section is one such example. A whole sub-hierarchy of LinAlg nodes is available, with objects such as Invert and Determinant representing basic linear algebra operation. Code generation for these objects can be specialized based upon the underlying architecture and the size of the involved tensors. For instance, a manually-optimized loop nest may be preferred to a BLAS function when the tensors are small<sup>2</sup>. Another special type of node is ArrayInit, used for static initialization of arrays. An ArrayInit wraps an N-dimensional Numpy array? and provides a simple interface to obtain information useful for optimization, like the sparsity pattern of the array.

**Symbols** A Symbol represents a variable in the code. The *rank* of a Symbol captures the dimensionality of a variable, with a rank equal to N indicating that a variable is an N-D array (N=0 implies that the variable is a scalar). The rank is implemented as an N-tuple, each entry being either an integer or a string representing a loop dimension. The *offset* of a Symbol is again an N-tuple where each element is a 2-tuple. For each entry r in the rank, there is a corresponding entry < scale, stride> in the offset. Rank and offset are used as in Figure 6.2 to access specific memory locations. By clearly identifying rank and offset of a Symbol - rather

<sup>&</sup>lt;sup>2</sup>It is well-known that BLAS libraries are highly optimized for big tensors, while their performance tends to be sub-optimal with small tensors, which are very common in assembly kernels.

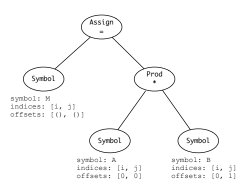


Figure 6.2: AST representation of a C assignment in COFFEE.

than storing a generic expression – the complexity of the data dependency analysis required by the rewrite operators is greatly reduced. The underlying assumption, however, is that all symbols in the kernel (at least those relevant for optimization) have access functions (see Section 2.5.2) that are affine in the loop indices. As motivated in Chapter 4, this is definitely the case for the class of kernels in which we are interested.

**Building an AST** Rather than using a parser, COFFEE exposes to the user the whole hierarchy of nodes for explicitly building ASTs. This is because the compiler is meant to be used as an intermediate step in a multilayer framework based on DSLs. To ease the construction of ASTs (especially nested loops), a set of utility functions is provided. We will elaborate on these aspects in the next section.

#### 6.3.2 Integration with Form Compilers

So far, COFFEE has been integrated with two form compilers: the FEniCS Form Compiler (FFC) and the Two-Stage Form Compiler (TSFC)<sup>3</sup>. These form compilers have their own internal representation of an assembly kernel; the objective is to turn such a representation into an AST suitable for COFFEE. We here describe how we achieved this in the case of FFC.

<sup>&</sup>lt;sup>3</sup>The generation of ASTs in TSFC has been written by Myklos Homolya.

The key idea is to enrich the FFC's intermediate representation at construction time; that is, when the UFL specification of a form is translated. We made the following changes.

- The mathematical expression evaluating the element tensor is represented as a tree data structure, or "FFC-AST". A limitation of an FFC-AST was that its nodes symbols or arithmetic operations were not bound to loops. For instance, the FFC-AST node corresponding to the symbol A[i][j] did not separate the variable name A from the loop indices i and j. We have therefore enriched FFC-AST symbols with additional fields to capture these information.
- Basis functions in an FFC-AST are added a new field storing the dimensionality of their function space. This information is used to enrich ArrayInit objects with the sparsity pattern of the values they are representing (recall that the tabulation of vector-valued basis functions is characterized by the presence of zero-valued blocks).

The improved FFC-AST is intercepted prior to code generation (the last phase in the original FFC, which outputs C code directly) and forwarded to a new module, where a COFFEE AST is finally built. In this module:

- the template originally used by FFC for code generation (i.e., the parts
  of an assembly kernels that are immutable across different forms) is
  changed in favour of "static" pieces of AST (kernel signature, loop nests,
  etc).
- the FFC-AST is visited and translated into a COFFEE AST by a suitable AST-to-AST converter routine.

The Two-Stage Form Compiler was originally conceived to produce ASTs for COFFEE, so no particular changes to its intermediate representation were needed.

#### 6.4 Rewrite Operators

COFFEE implements sharing elimination and pre-evaluation by composing "building-block" operators, or "rewrite operators". This has several advantages. Firstly, extendibility: novel transformations – for instance,

sum-factorization in spectral methods – could be expressed using the existing operators, or with small effort building on what is already available. Secondly, generality: COFFEE can be seen as a lightweight, low level computer algebra system, not necessarily tied to finite element integration. Thirdly, robustness: the same operators are exploited, and therefore stressed, by different optimization pipelines. The rewrite operators, whose implementation is based on manipulation of the kernel's AST, essentially compose the COFFEE language.

The most important rewrite operators in COFFEE are:

**Generalized code motion** It pre-computes the values taken by a sub-expression along an invariant dimension. This is implemented by introducing a temporary array per invariant sub-expression and by adding a new "clone" loop to the nest (Several examples, e.g. Figure ??, have been provided throughout the thesis). At the price of some extra memory for storing temporaries, all lifted terms are now amenable to auto-vectorization.

**Expansion** This transformation consists of expanding (i.e., distributing) a product between two generic sub-expressions. Expansion has several effects, the most important ones being exposing factorization opportunities and increasing the operation count. It can also help relieving the register pressure within a loop, by allowing further code motion.

**Factorization** Collecting, or factorizing, symbols reduces the number of multiplications and potentially exposes, as illustrated through sharing elimination, code motion opportunities.

**Symbolic evaluation** This operator evaluates sub-expressions that only involve statically initialized, read-only arrays (e.g., basis function tables). The result is stored into a new array, and the AST modified accordingly All these operators are used by both sharing elimination and pre-evaluation (apart from symbolic evaluation, only employed by pre-evaluation).

The rewrite operators accept a number of options to drive the transformation process. With code motion, for example, we can specify what kind of sub-expressions should be hoisted (by indicating the expected invariant loops) or the amount of memory that is spendable in temporaries. Factorization can be either "explicit", by providing a list of symbols to be factorized or a loop dimension along which searching for factorizable symbols, or "heuristic", with the algorithm searching for the groups of

#### 6.5 Features of the Implementation

Rather than providing the pseudo-code and an explanation for each of the algorithms implemented in COFFEE – a mere exercise of scarce interest for the reader, given that the implementation is open-source and well-documented – this section focuses on the structure of the compiler and its "toolkit" for implementing or extending rewrite operators.

#### 6.5.1 Tree Visitor Pattern

The need for a generic infrastructure for traversing ASTs has grown rapidly, together with the complexity of the compiler. In the early stages of COF-FEE, any time that a new transformation (e.g., a rewrite operator) or data collector (e.g., for dependence analysis) were required, the full AST traversal had to be (re-)implemented. In addition, the lack of a common interface for tree traversals made the code more difficult to understand and to extend. This led to the introduction of a tree visitor design pattern<sup>4</sup>, whose aim is to decouple the algorithms from the data structure on which they are applied ?.

Consider, without loss of generality, an algorithm that needs to perform special actions (e.g., collecting loop dependence information) any time a Symbol or a ForLoop nodes are encountered. Then, a tree visitor will only need to implement three methods, namely visit\_Symbol and visit\_ForLoop – the actual handlers – as well as visit\_Node, which implements the "fallback" action for all other node types (typically, just a propagation of the visit).

Tree visitors exploit the hierarchy of AST nodes by always dispatching to the most specialized handler. For example, symbols are simultaneously of type Symbol and Expression, but if a Symbol is encountered and visit\_Symbol is implemented, then visit\_Symbol is executed, whereas visit\_Expression (if any) is ignored.

Most of the algorithms in COFFEE exploit the tree visitor pattern; a few,

<sup>&</sup>lt;sup>4</sup>The tree visitor infrastructure was mainly developed by Lawrence Mitchell, and was inspired by that adopted in UFL, the language used to specify forms in Firedrake.

the "oldest" ones, still do not, due to the lack of time for porting to the new infrastructure.

#### 6.5.2 Flexible Code Motion

Code motion consists of lifting, or hoisting, a (sub-)expression out of one or more loops. This rewrite operator is used in many different contexts: as a stand-alone transformation (optimization level 01); in multiple steps during sharing elimination; in pre-evaluation.

When applying the operator, several pieces of information must be known:

- 1. What sub-expression should be hoisted; for instance, should they be constant in the whole loop nest or invariant in at most one of the linear loops.
- 2. Where to hoist it; that is, how many loops is the operator allowed to cross.
- 3. How much memory are we allowed to use for a temporary.
- 4. If a common sub-expression had already been hoisted.

The code motion operator is flexible and let the caller (i.e., a higher-level transformation) drive the hoisting process by specifying how to behave with respect to the aforementioned points.

COFFEE must therefore track all of the hoisted sub-expressions for later retrieval. A dictionary mapping each of the temporaries introduced to a tuple of metadata is employed. For a temporary t, the dictionary records:

- A reference to the hoisted expression e assigned to t.
- A reference to the loop in which e is lifted.
- A reference to the declaration of t.

This dictionary belongs to the "global state" of COFFEE. It is updated each time the code motion operator is invoked, and read by other transformations (e.g., by all of the lower level optimizations).

The code motion operator "silently" applies common sub-expression elimination. A look-up in the dictionary tells whether a hoistable subexpression e has been assigned to a temporary t by a prior call to the operator; in such a case, e is straightforwardly replaced with t, that is, no further temporaries are introduced.

#### 6.5.3 Tracking Data Dependency

Data dependency analysis is necessary to ensure the legality of some transformations. For example:

- When lifting a sub-expression e, we may want to hoist "as far as possible" in the loop nest (possibly even outside of it); that is, right after the last write to a variable read in e.
- When expanding a product, some terms may be aggregated with previously hoisted sub-expressions. This would avoid introducing extra temporaries and increasing the register pressure. For example, if we have (a + b)\*c and both a and b are temporaries created by code motion, we could expand the product and aggregate c with the sub-expressions stored by a and b. Obviously, this is as long as neither a nor b are accessed in other sub-expressions.
- For loop fusion (see Section 5.4.4).

In a similar way to general-purpose compilers, COFFEE uses a dependency graph for tracking data dependencies. The dependency graph has as many vertices as symbols in the code; a direct edge from A to B indicates that symbol B depends on (i.e., is going to read) symbol A. Since COFFEE relies on *static single assignment* – a property that ensures that variables are assigned exactly once – such a minimalistic data structure suffices for data dependence analysis.

#### 6.5.4 Minimizing Temporaries

Both code motion operator (Section 6.5.2) and common sub-expression elimination induced by loop fusion (Section 5.4.4) impact the number of temporaries in the assembly kernel. At the end of expression rewriting, a routine in COFFEE attempts to remove all of the unnecessary temporaries. This makes the code more readable and, potentially, relieves the register pressure.

The main rule for removing a temporary t storing an expression e is that if t is accessed only in a single statement s, then e is inlined into s and t

is removed. Secondly, if some of the transformations in the optimization pipeline reduced e to a symbol, then any appearance of t is also replaced by e.

# Chapter 7

# **Conclusions**

...

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