

A Domain-Specific Optimizing Compiler for Solving Partial Differential Equations

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Abstract—The abstract goes here.

I. INTRODUCTION

In many fields, like computational fluid dynamics, computational electromagnetics, and structural mechanics, programs model phenomena by means of partial differential equations (PDEs). Numerical techniques, like Finite Volume Method (FVM) and Finite Element Method (FEM), are widely employed in real-world applications to approximate solutions of PDEs. Unstructured meshes are often used to discretize the domain of a PDE, since their geometric flexibility allows solvers to be extremely effective. However, they are also one of the main responsible for performance degradation, due to the need for indirect accesses (e.g. $A[B[i]]$) to access mesh components. A typical problem, therefore, is that implementing a scientific application requires a great effort, other than domain knowledge, and even more time is needed to optimize their execution time.

An unstructured mesh discretizes the domain of the equation into two- or three-dimensional cells (or “elements”), within which the solution is sought. As the number of elements can be of the order of millions, a major issue is the time required to perform the numerical method, which can be hours or days. To address this issue, domain-specific languages (DSLs), which ease the programming burden and enable highly-efficient parallel execution, have been developed. The successful porting of Hydra, a CFD industrial application devised by Rolls Royce for turbomachinery design (based on FVM, roughly 50000 lines of code and mesh sizes that can be over 100 millions edges), to the OP2 DSL [2], demonstrates the effectiveness of the DSL approach to implementing PDEs solvers [?].

OP2, as well as Lizst [1], adopt a kernel-oriented programming model, in which the computation semantics is expressed through self-contained functions (or “kernels”). A kernel is applied to all elements in a set of mesh components (e.g. edges, vertices, elements), with an implicit synchronization between the application of two consecutive kernels. On commodity multi-cores, a kernel is executed sequentially by a thread, while parallelism is achieved partitioning the mesh and assigning each partition to a thread. Together with the indirections problem, kernel optimization is one of the major concerns in unstructured mesh applications. In this paper, we tackle this problem by proposing a domain-driven optimization strategy for a class of kernels used in FEM.

We focus on Finite Element Local Assembly (“assembly”, in the following), a fundamental step of FEM that covers an important fraction of the overall computation run-time, typically in the range 30%-60%. During the assembly phase, the solution of the PDE is approximated by executing a suitable kernel over all elements in the discretized domain. A kernel’s working set is usually small enough to fit the L1 cache; it might need L2 cache when (very) high-order methods are employed to improve the accuracy of the solution. However, we do not consider the latter case. An assembly kernel is characterized by the presence of an affine, usually non-perfect loop nest, where individual loops are rather small (the trip count rarely exceeds 60, with a minimum value of 3, depending on the order of the method). With such small kernels, our work focuses on aspects like minimization of floating-point operations, register allocation and instruction-level parallelism, especially in the form of SIMD vectorisation.

Our study is conducted in the context of Firedrake, a multilayered system for the automated solution of PDEs through FEM. PDEs are written in Firedrake using a domain-specific language called Unified Form Language (UFL). The high-level UFL notation is used by the Fenics Form Compiler (FFC) to produce assembly kernels. At the core of Firedrake [?] there is PyOP2 [2] - a python implementation of the OP2 abstraction - which triggers compilation of kernels using an available vendor compiler, and eventually manages parallel execution over the unstructured mesh. We exploit the domain knowledge and the structure inherent to FFC’s assembly kernels to develop an optimizing compiler for all problems that are expressible in Firedrake.

The compiler we have developed is integrated with PyOP2, and already used in the released version of Firedrake [?]. This allows us to evaluate the proposed code transformations in a range of real-world problems, employing different low-order methods (from $p = 1$ to $p = 4$; the higher, the more expensive is a kernel, but better is the accuracy), and varying the topology of the mesh (2D, 3D, 3D hybrid structured-unstructured, which in turn affect the kernel cost). Performance results show that compiler-generated code for non-trivial assembly kernels, if not adequately transformed, is sub-optimal. A cost-model-driven sequence of source-to-source code transformations, aimed at improving register allocation and register data locality, can result in performance improvements up to 33% over “softly-optimized” code (i.e. where only basic transformations are performed, such as auto-vectorizable loop-invariant code

```

1 void helmholtz(double A[3][3], double **coords) {
2   // K, det = Compute Jacobian (coords)
3
4   static const double W3[3] = {...}
5   static const double X_D10[3][3] = {...}
6   static const double X_D01[3][3] = {...}
7
8   for (int i = 0; i<3; i++)
9     for (int j = 0; j<3; j++)
10      for (int k = 0; k<3; k++)
11        A[j][k] += ((Y[i][k]*Y[i][j]+
12          +((K[1]*X_D10[i][k]+K[3]*X_D01[i][k])*
13            *(K[1]*X_D10[i][j]+K[3]*X_D01[i][j]))+
14            +((K[0]*X_D10[i][k]+K[2]*X_D01[i][k])*
15              *(K[0]*X_D10[i][j]+K[2]*X_D01[i][j])))*
16            *det*W3[i]);
17 }

```

Algorithm 1: Local assembly code generated by FFC for a Helmholtz problem (2D mesh, Lagrange $p = 1$ elements).

motion, padding and data alignment), and up to 77% over the original FFC kernels. The contribution of this paper is therefore twofold:

- An optimisation strategy for a class of kernels widely-used in scientific applications, namely Finite Element Local Assembly kernels. Our approach exploits domain knowledge to go beyond the limits of both vendor (e.g. *icc*) and research (e.g. those based on the polyhedral model) compilers.
- Automation of such code optimizations in Firedrake, a framework for the resolution of PDEs through FEMs. This enables performing an in-depth performance evaluation of the proposed optimization strategy in real-world simulations.

The paper is organized as follows. ...

II. BACKGROUND

Local assembly consists of evaluating so called element stiffness matrix and vector; in this work, we focus on computation of matrices, which is the costly part of the process. A stiffness matrix can be intuitively thought as an approximated representation of the PDE solution in a specific cell of the discretized domain. Quadrature algorithms (based on numerical integration), which follow on from the mathematical description of a stiffness matrix, are widely-used in scientific codes to implement assembly [?], [?]. The FFC compiler used by Firedrake and Fenics can generate assembly kernels using optimized quadrature schemes. It is worth noticing, however, that problems and considerations similar to that exposed below can be found also in non-FFC assembly codes.

The complexity of FFC-generated kernels depends on the mathematical problem being solved. In simpler cases, the loop nest is perfect, it has short trip counts (in the range 3-15), and the computation reduces to a summation of a few, small outer-product vector-operations. An example is provided in Figure 1, which shows an assembly kernel for a Helmholtz problem, using Lagrange basis functions on 2D elements with polynomial order $p = 1$. In other scenarios, for instance when solving a non-linear problem like Burgers (see Figure 2), the

```

1 void burgers(double A[12][12], double **c, double **w) {
2   // K, det = Compute Jacobian (c)
3
4   static const double W5[5] = {...}
5   static const double X1_D001[5][12] = {...}
6   static const double X2_D001[5][12] = {...}
7   //It follows 11 other basis functions definitions.
8   ...
9
10  for (int i = 0; i<5; i++) {
11    double F0 = 0.0;
12    //It follows 10 other declarations (F1, F2,...)
13    ...
14    for (int r = 0; r<12; r++) {
15      F0 += (w[r][0]*X1_D100[ip][r]);
16      //It follows 10 analogous statements (F1, F2, ...)
17    }
18    ...
19    for (int j = 0; j<12; j++)
20      for (int k = 0; k<12; k++)
21        A[j][k] += (.(K[5]*F9)+(K[8]*F10))*Y1[i][j])+
22          +(((K[0]*X1_D100[i][k])+(K[3]*X1_D010[i][k])+
23            +(K[6]*X1_D001[i][k]))*Y2[i][j]))*F11)+
24          +((K[2]*X2_D100[i][k])+...+(K[8]*X2_D001[i][k]))*
25            *(K[2]*X2_D100[i][j])+...+(K[8]*X2_D001[i][j])))+
26            + <a hundred of sum/muls go here>...)*
27            *det*W5[i]);
28  }
29 }

```

Algorithm 2: Local assembly code generated by FFC for a Burgers problem (3D mesh, Lagrange $p = 1$ elements).

number of arrays involved in the computation of A can be much larger: in this case, 14 unique arrays are accessed, and the same array can be referenced multiple times within the expression. Also, constants evaluated in outer loops (called F in the code), acting as scaling factors of arrays, may be required; trip counts can be larger (proportionally to the order of the method); arrays may be block-sparse. Despite such a big space of possible assembly implementations, it is still possible to identify a domain-specific structure common to all kernels, which can be turned into effective code transformations and simd-vectorization.

The class of kernels we are considering has, in particular, some peculiarities. 1) The computation of the Jacobian, which is the first step of the assembly, is independent of the loop nest; this is not true in general, since the unstructured mesh might have bended elements that require the Jacobian be re-computed every i iteration. 2) The stiffness matrix A is always the result of an outer-product computation along the j and k dimensions. 3) Memory accesses along the three loop dimensions are always 1-stride. 4) The j and k loops are interchangeable, whereas to permute i it might be necessary to precompute F values and to introduce temporaries. 5) Basis functions arrays (denoted by X, Y, \dots) are constants, and many sub-expressions on the right hand side of the stiffness matrix computation depend on just two loops (either $i-j$ or $i-k$). In Section III we show how to exploit these observations to define a set of systematic, composable optimizations.

One attractive possibility is to transform the loop nest into a sequence of calls to BLAS routines. There are several issues that make achieving this difficult and, potentially, not

effective. The first problem concerns automating identification and extraction of matrix-matrix multiplications from the code, which requires a non-trivial analysis of the memory access pattern. Related to this problem is the fact that, in general, many sub-expressions should be pre-computed (i.e. computed once before the loop nest), which leads to using temporaries and, possibly, to working sets that do not fit the L1 cache (for $p > 2$). In addition, using BLAS routines may impact data locality, since the same matrix can appear in multiple products, implying, at least, a certain degree of register spilling. Finally, it is known that BLAS routines perform far from peak performance when the dimension of the matrices is rather small [?]. Hand-made implementations of the simple Helmholtz problem ($p = 1, 2, 3, 4$) as a sequence of BLAS (*dgemm*) routines have demonstrated that performance is worse than that shown in Section V. A comprehensive study concerning BLAS optimization is therefore left as further work.

III. CODE TRANSFORMATIONS

1 The PyOP2 Compiler

Input: ast, wrapper, isa

Output: code

```
2 // Analyze ast and build optimization plan
3 it_space = analyze(ast)
4 if not it_space then
5     ast.apply_inter_kernel_vectorization(wrapper)
6     return ast.from_ast_to_c()
7 endif
8
9 // Optimize ast based on plan
10 ast.apply LICM()
11 if plan.sz_split then
12     ast.split(plan.sz_split)
13 endif
14 if plan.uaj_factor then
15     ast.vr_tile(plan.uaj_factor)
16 endif
17 ast.padding()
18 ast.data_align()
19 return ast.from_ast_to_c()
```

Algorithm 3: The PyOP2 compiler.

1 Algorithm: ApplyCostModel

Input: n_inner_arrays, n_regs

Output: uaj_factor, sz_split

```
2 sz_split = 0
3 // Compute splitting factor
4 while n_inner_arrays > n_regs do
5     n_inner_arrays = n_inner_arrays / 2
6     sz_split = sz_split + 1
7 endw
8 n_inner_regs = n_regs / 2
9 if n_inner_arrays > n_inner_regs or sz_split > 0 then
10     // Rely on autovectorization, as there might be not
11     // enough registers to improve performance by tiling
12     return <sz_split, 0>
13 endif
14 // Compute unroll-and-jam factor for vector-register
15 // tiling
16 n_regs_avail = n_regs - n_inner_arrays
17 uaj_factor = ⌈n_regs_avail / n_inner_arrays⌉
18 return <sz_split, uaj_factor>
```

Algorithm 4: Procedure to estimate the most suitable unroll-and-jam factor and/or split size.

IV. THE PYOP2 COMPILER

The compiler structure

The compiler cost model. Used to find out suitable unroll-and-jam factor and split size.

V. PERFORMANCE EVALUATION

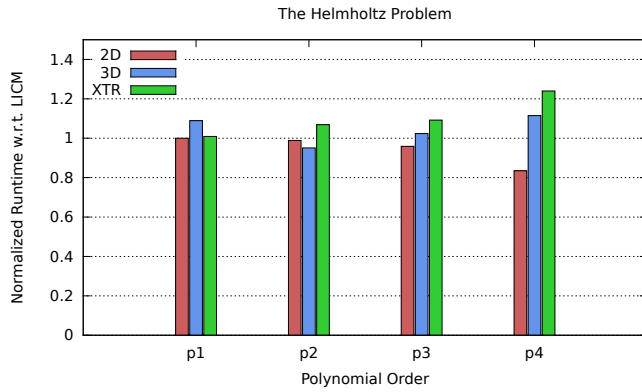


Fig. 1. Helmholtz

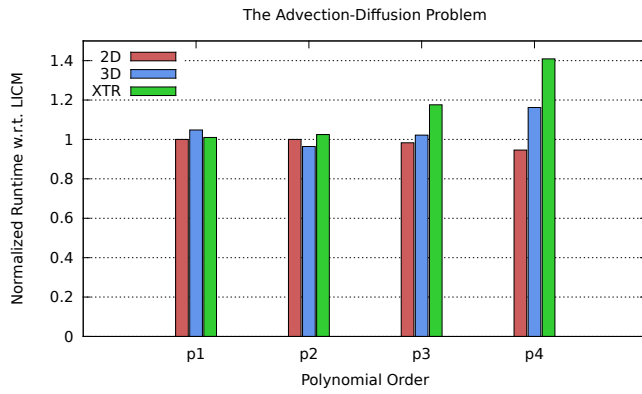


Fig. 2. Advection-Diffusion

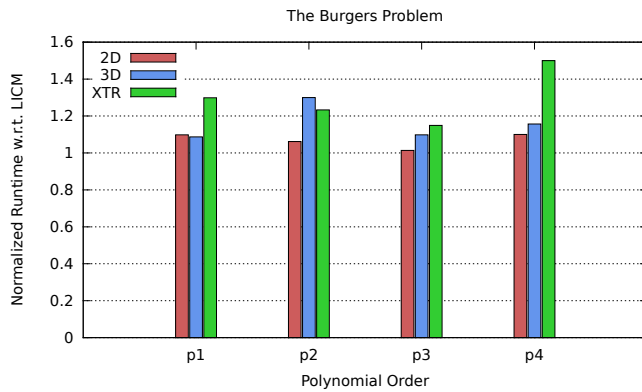


Fig. 3. Burgers

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VI. RELATED WORK

VII. CONCLUSIONS

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