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A Domain-Specific Compiler for Linear Algebra Operations

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ABSTRACT

We present a prototypical linear algebra compiler that automatically exploits domain-specific knowledge to generate high-performance algorithms. The input to the compiler is a target equation together with knowledge of both the structure of the problem and the properties of the operands. The output is a variety of high-performance algorithms to solve the target equation. Our approach consists in the decomposition of the input equation into a sequence of library-supported kernels. Since in general such a decomposition is not unique, our compiler returns not one but a number of algorithms. The potential of the compiler is shown by means of its application to a challenging equation arising within the *genome-wide association study*. As a result, the compiler produces multiple “best” algorithms that outperform the best existing libraries.

Categories and Subject Descriptors

D.1.2 [Programming Techniques]: Automatic Programming; G.4 [Mathematical software]: Algorithm Design and Analysis

General Terms

Algorithms, Experimentation, Performance

Keywords

Domain-Specific Compiler, Algorithm Generation, Numerical Linear Algebra, Pattern Matching, Rewrite Rules

1. INTRODUCTION

In the past 30 years, the development of linear algebra libraries has been tremendously successful, resulting in a variety of reliable and efficient computational kernels. Unfortunately these kernels are limited by a rigid interface that does not allow users to pass knowledge specific to the target problem. If available, such knowledge may lead to domain-specific algorithms that attain higher performance than any

traditional library [4]. The difficulty does not lay so much in creating flexible interfaces, but in developing algorithms capable of taking advantage of the extra information.

In this paper, we present preliminary work on a linear algebra compiler, written in Mathematica, that automatically exploits application-specific knowledge to generate high-performance algorithms. The compiler takes as input a target equation and information on the structure and properties of the operands, and returns as output algorithms that exploit the given information. In the same way that a traditional compiler breaks the program into assembly instructions directly supported by the processor, attempting different types of optimization, our linear algebra compiler breaks a target operation down to library-supported kernels, and generates not one but a family of viable algorithms. The decomposition process undergone by our compiler closely replicates the thinking process of a human expert.

We show the potential of the compiler by means of a challenging operation arising in computational biology: the *genome-wide association study* (GWAS), an ubiquitous tool in the fields of genomics and medical genetics [7, 8, 10]. As part of GWAS, one has to solve the following equation

$$\begin{cases} b_{ij} := (X_i^T M_j^{-1} X_i)^{-1} X_i^T M_j^{-1} y_j & \text{with } 1 \leq i \leq m \\ M_j := h_j \Phi + (1 - h_j)I & \text{and } 1 \leq j \leq t, \end{cases} \quad (1)$$

where X_i , M_j , and y_j are known quantities, and b_{ij} is sought after. The size and properties of the operands are as follows: $b_{ij} \in \mathcal{R}^p$, $X_i \in \mathcal{R}^{n \times p}$ is full rank, $M_j \in \mathcal{R}^{n \times n}$ is symmetric positive definite (SPD), $y_j \in \mathcal{R}^n$, $\Phi \in \mathcal{R}^{n \times n}$, and $h_j \in \mathcal{R}$; $n \approx 10^3$, $1 \leq p \leq 20$, $m \approx 10^{6-7}$, and t is either 1 or of the order of 10^5 .

At the core of GWAS lays a linear regression analysis with non-independent outcomes, carried out through the solution of a two-dimensional sequence of the Generalized Least-Squares problem (GLS)

$$b := (X^T M^{-1} X)^{-1} X^T M^{-1} y. \quad (2)$$

While GLS may be directly solved, for instance, by MATLAB, or may be reduced to a form accepted by LAPACK [1], none of these solutions can exploit the specific structure pertaining to GWAS. However, the nature of the problem, a sequence of correlated GLSs, allows multiple ways to reuse computation. Also, different sizes of the input operands demand different algorithms to attain high-performance in all possible scenarios. The application of our compiler to GWAS, Eq. 1, results in the automatic generation of dozens of algorithms, many of which outperform the current state of the art by a factor of four or more.

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The paper is organized as follows. Related work is briefly described in Section 2. Sections 3 and 4 uncover the principles and mechanisms upon which the compiler is built. In Section 5 we carefully describe the automatic generation of multiple algorithms, while in Section 6 we report on the performance of the generated algorithms through numerical experiments. We draw conclusions in Section 7.

2. RELATED WORK

A number of research projects concentrate their efforts on domain-specific languages and compilers. Among them, the SPIRAL project [9] and the Tensor Contraction Engine (TCE) [3], focused on signal processing transforms and tensor contractions, respectively. Centered on general dense linear algebra operations, one of the goals of the FLAME project is the systematic generation of algorithms. The FLAME methodology, based on the partitioning of the operands and the automatic identification of loop-invariants [6, 5], has been successfully applied to a number of operations, originating hundreds of high-performance algorithms.

The approach described in this paper is orthogonal to FLAME. No partitioning of the operands takes place. Instead, the main idea is the mapping of operations onto high-performance kernels from available libraries.

3. THE COMPILER PRINCIPLES

In this section we expose the human thinking process behind the generation of algorithms for a broad range of linear algebra equations. As an example, we derive an algorithm for the solution of the GLS problem, Eq. 2, as it would be done by an expert. Together with the derivation, we describe the rationale for every step of the algorithm. The exposed rationale highlights the key ideas on top of which we founded the design of our compiler.

Given Eq. 2, the **first concern is the inverse operator** applied to the expression $X^T M^{-1} X$. Since X is not square, the inverse cannot be distributed over the product and the expression needs to be processed first. The attention falls then on M^{-1} . The inversion of a matrix is costly and not recommended for numerical reasons; therefore, since M is a full matrix, we **factor** it. Given the structure of M (SPD), we choose a Cholesky factorization, resulting in

$$\begin{aligned} LL^T &= M \\ b &:= (X^T (LL^T)^{-1} X)^{-1} X^T (LL^T)^{-1} y, \end{aligned} \quad (3)$$

where L is square and lower triangular. As L is square, the inverse may now be distributed over the product LL^T , yielding $L^{-T} L^{-1}$. Next, we process $X^T L^{-T} L^{-1} X$; we observe that the quantity $L^{-1} X$ **appears multiple times**, and may be computed and reused to **save computation**:

$$\begin{aligned} W &:= L^{-1} X \\ b &:= (W^T W)^{-1} W^T L^{-1} y. \end{aligned} \quad (4)$$

At this point, since W is not square and the inverse cannot be distributed, there are two **alternatives**: 1) multiply out $W^T W$; or 2) factor W , for instance through a QR factorization. In this example, we choose the former:

$$\begin{aligned} S &:= W^T W \\ b &:= S^{-1} W^T L^{-1} y. \end{aligned} \quad (5)$$

One can prove that S is SPD, suggesting yet another factorization. We choose a Cholesky factorization and distribute the inverse over the product:

$$\begin{aligned} GG^T &= S \\ b &:= G^{-T} G^{-1} W^T L^{-1} y. \end{aligned} \quad (6)$$

Now that all the remaining inverses are applied to triangular matrices, we are left with a series of products to compute the final result. Since all operands are matrices except the vector y , we compute Eq. 6 from right to left to **minimize the number of flops**. The final algorithm is shown in Alg. 1.

Algorithm 1: Algorithm for the solution of the GLS problem as derived by a human expert

1	$LL^T = M$	(POTRF)
2	$W := L^{-1} X$	(TRSM)
3	$S := W^T W$	(SYRK)
4	$GG^T = S$	(POTRF)
5	$y := L^{-1} y$	(TRSV)
6	$b := W^T y$	(GEMV)
7	$b := G^{-1} b$	(TRSV)
8	$b := G^{-T} b$	(TRSV)

Three ideas stand out as the guiding principles for the thinking process:

- The first concern is to deal, whenever it is not applied to diagonal or triangular matrices, with the inverse operator. Two scenarios may arise: a) it is applied to a single operand, A^{-1} . In this case the operand is factored with a suitable factorization according to its structure; b) the inverse is applied to an expression. This case is handled by either computing the expression and reducing it to the first case, or factoring one of the matrices and analyzing the resulting scenario.
- When decomposing the equation, we give priority to a) common segments, i.e., common subexpressions, and b) segments that minimize the number of flops; this way we reduce the amount of computation performed.
- If multiple alternatives leading to viable algorithms arise, we explore all of them.

4. COMPILER OVERVIEW

Our compiler follows the above guiding principles to closely replicate the thinking process of a human expert. To support the application of these principles, the compiler incorporates a number of modules ranging from basic matrix algebra support to analysis of dependencies, including the identification of building blocks offered by available libraries. In the following, we describe the core modules.

Matrix algebra The compiler is written using Mathematica from scratch. We implement our own operators: addition (plus), negation (minus), multiplication (times), inversion (inv), and transposition (trans). Together with the operators, we define their precedence and properties, as commutativity, to support matrices as well as vectors and scalars. We also define a set

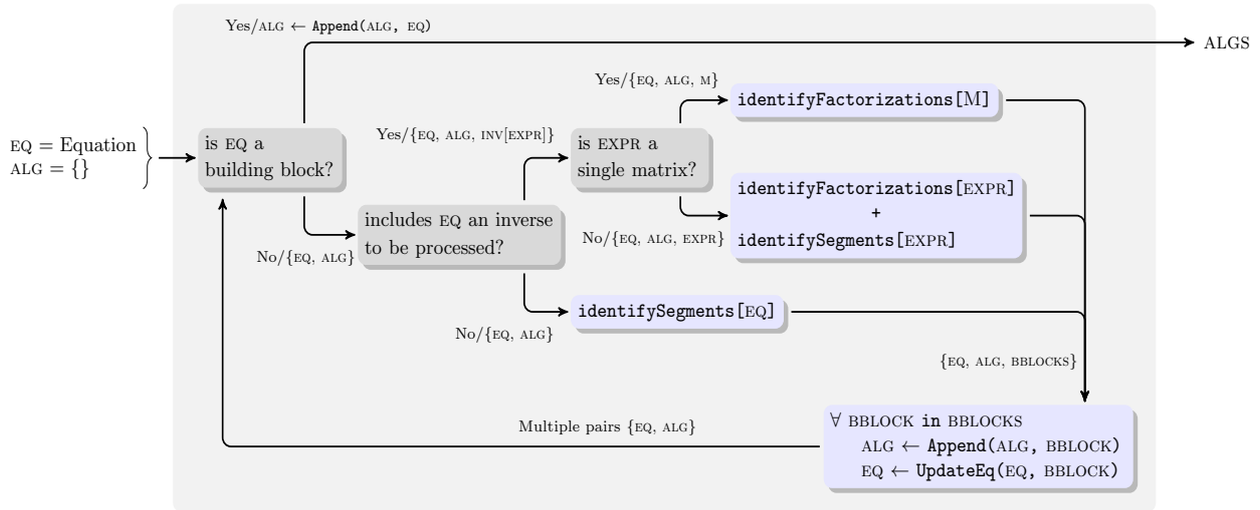


Figure 1: High-level description of the compiler’s *reasoning*.

of rewrite rules according to matrix algebra properties to freely manipulate expressions and simplify them, allowing the compiler to work on multiple equivalent representations.

Inference of properties In this module we define the set of supported matrix properties. As of now: identity, diagonal, triangular, symmetric, symmetric positive definite, and orthogonal. On top of these properties, we build an inference engine that, given the properties of the operands, is able to infer properties of complex expressions. This module is extensible and facilitates incorporating extra properties.

Building blocks interface This module contains an extensive list of patterns associated with the desired building blocks onto which the algorithms will be mapped. As with the properties module, if a new library is to be used, the list of accepted building blocks can be easily extended.

Analysis of dependencies When considering a sequence of problems, as in GWAS, this module analyzes the dependencies among operations and between operations and the dimensions of the sequence. Through this analysis, the compiler rearranges the operations in the algorithm, eliminating redundant computations.

To complete the description of our compiler, we include in Fig. 1 a high-level description of the algorithm behind the compiler’s *reasoning*. We emphasize that the algorithm does not rely on an exhaustive search; instead, the search space is constrained by the guiding principles listed in Sec. 3.

The process takes the equation EQ as input; initially, the algorithm ALG includes no building blocks, it is empty. If EQ corresponds to a supported building block, the building block is added to ALG, and the decomposition is complete. Otherwise, the compiler inspects EQ for inverses that need treatment. If such an inverse exists, two different cases may arise: 1) the inverse is applied to a single matrix M, then the compiler identifies a set of viable factorizations for M, based on its properties and structure; 2) the inverse is applied to an expression EXPR, then the compiler identifies both viable factorizations for the operands in EXPR, and segments

of EXPR that are directly mapped onto a building block. If there exist no inverse to process, the compiler identifies segments in EQ with a mapping onto a building block.

All three scenarios may return multiple alternatives. Each of the alternatives is studied, one at a time: the corresponding building block BBLOCK (either a factorization or a segment) is added to the sequence in ALG, and EQ is updated as shown several times in Sec. 3 (Eqs. 3, 4, 5, and 6). The *reasoning* process is repeated for every resulting pair {EQ, ALG} until the decomposition of every branch is complete. The result is the list of algorithms ALGS.

The compiler is still in its early stages and the code is not yet available for a general release. However, we include along the paper details on the input / output of the system, as well as screenshots of the actual working prototype.

5. COMPILER-GENERATED ALGORITHMS

We detail now the application of the process depicted in Fig. 1 to GWAS. Box 1 includes the input to the compiler: the target equation along with domain-specific knowledge arising from GWAS, e.g, operands’ shape and properties. As a result, dozens of algorithms are automatically generated; we report on three selected ones.

5.1 Algorithm 1

To ease the reader, we describe the process towards the generation of an algorithm similar to Alg. 1. The starting point is Eq. 1. Since X is not square, the inverse operator applied to $X^T(h\Phi + (1-h)I)^{-1}X$ cannot be distributed over the product; thus, the inner-most inverse is $(h\Phi + (1-h)I)^{-1}$. The inverse is applied to an expression, which is inspected for viable factorizations and segments. Among the identified alternatives are a) the factorization the operand Φ according to its properties, or b) the computation of the expression $h\Phi + (1-h)I$. Here we concentrate on the second case. The segment $h\Phi + (1-h)I$ is matched as the SCAL-ADD building block (scaling and addition of matrices); the operation is

```

equation = {
  equal[b,
    times[
      inv[times[
        trans[X],
        inv[plus[
          times[h, Phi],
          times[plus[1, minus[h]], id]
        ]],
      X]
    ],
  trans[X],
  inv[plus[
    times[h, Phi],
    times[plus[1, minus[h]], id]
  ]],
  y
]
];

operandProperties = {
  {X, {'Input', 'Matrix', 'FullRank'}},
  {y, {'Input', 'Vector'}},
  {Phi, {'Input', 'Matrix', 'Symmetric'}},
  {h, {'Input', 'Scalar'}},
  {b, {'Output', 'Vector'}}
};

expressionProperties = { inv[plus[ times[h, Phi],
  times[plus[1, minus[h]], id ]], 'SPD' };

sizeAssumptions = { rows[X] > cols[X] };

```

Box 1: Mathematica input to the compiler.

made explicit and replaced:

$$\begin{aligned}
M &:= h\Phi + (1-h)I \\
b &:= (X^T M^{-1} X)^{-1} X^T M^{-1} y.
\end{aligned} \tag{7}$$

Now, the inner-most inverse is applied to a single operand, M , and the compiler decides to factor it using multiple alternatives: Cholesky ($LL^T = M$), QR ($QR = M$), eigendecomposition ($ZWZ^T = M$), and SVD ($U\Sigma V^T = M$). All the alternatives are explored; we focus now on the Cholesky factorization (POTRF routine from LAPACK):

$$\begin{aligned}
LL^T &= M \\
b &:= (X^T L^{-T} L^{-1} X)^{-1} X^T L^{-T} L^{-1} y.
\end{aligned} \tag{8}$$

After M is factored and replaced by LL^T , the inference engine propagates a number of properties to L based on the properties of M and the factorization applied. Concretely, L is square, triangular and full-rank.

Next, since L is triangular, the inner-most inverse to be processed in Eq. 8 is $(X^T L^{-T} L^{-1} X)^{-1}$. In this case two routes are explored: either factor X (L is triangular and does not need further factorization), or map a segment of the expression onto a building block. We consider this second alternative. The compiler identifies the solution of a triangular system (TRSM routine from BLAS) as a common

segment appearing three times in Eq. 8, makes it explicit, and replaces it:

$$\begin{aligned}
W &:= L^{-1} X \\
b &:= (W^T W)^{-1} W^T L^{-1} y.
\end{aligned} \tag{9}$$

Since L is square and full-rank, and X is also full-rank, W inherits the shape of X and is labelled as full-rank. As W is not square, the inverse cannot be distributed over the product yet. Therefore, the compiler faces again two alternatives: either factoring W or multiplying $W^T W$. We proceed describing the latter scenario while the former is analyzed in Sec. 5.2. $W^T W$ is identified as a building block (SYRK routine of BLAS), and made explicit:

$$\begin{aligned}
S &:= W^T W \\
b &:= S^{-1} W^T L^{-1} y.
\end{aligned} \tag{10}$$

The inference engine plays an important role deducing properties of S . During the previous steps, the engine has inferred that W is full-rank and $\text{rows}[W] > \text{cols}[W]$; therefore the following rule states that W is SPD:

```

isSPDQ[ times[ trans[ A_?isFullRankQ, A_ ] ] ] /;
  Refine[ rows[A] > cols[A], And@@sizeAssumptions ]
:= True;

```

This knowledge is now used to determine possible factorizations for S . We concentrate on the Cholesky factorization:

$$\begin{aligned}
GG^T &= S \\
b &:= G^{-T} G^{-1} W^T L^{-1} y.
\end{aligned} \tag{11}$$

In Eq. 11, all inverses are applied to triangular matrices; therefore, no more treatment of inverses is needed. The compiler proceeds with the final decomposition of the remaining series of products. Since at every step the inference engine keeps track of the properties of the operands in the original equation as well as the intermediate temporary quantities, it knows that every operand in Eq. 11 are matrices except for the vector y . This knowledge is used to give matrix-vector products priority over matrix-matrix products, and Eq. 11 is decomposed accordingly. In case the compiler cannot find applicable heuristics to lead the decomposition, it explores the multiple viable mappings onto building blocks. The resulting algorithm, and the corresponding output from Mathematica, are assembled in Alg. 2, CHOL-GWAS.

Algorithm 2: CHOL-GWAS

<pre> 1 M := hPhi + (1-h)I (SCAL-ADD) 2 LL^T = M (POTRF) 3 W := L^{-1} X (TRSM) 4 S := W^T W (SYRK) 5 GG^T = S (POTRF) 6 y := L^{-1} y (TRSV) 7 b := W^T y (GEMV) 8 b := G^{-1} b (TRSV) 9 b := G^{-T} b (TRSV) </pre>	<pre> tmp1 = -(hid) + 1 id + h Phi L2 L2^T = tmp1 tmp5 = X^T L2^T tmp10 = tmp5 tmp5^T L3 L3^T = tmp10 tmp23 = L2^{-1} y tmp31 = tmp5 tmp23 tmp40 = L3^{-1} tmp31 tmp55 = L3^{-T} tmp40 b = tmp55 </pre>
--	---

5.2 Algorithm 2

In this subsection we display the capability of the compiler to analyze alternative paths, leading to multiple viable

algorithms. At the same time, we expose more examples of algebraic manipulation carried out by the compiler. The presented algorithm results from the alternative path arising in Eq. 10, the factorization of W . Since W is a full-rank column panel, the compiler analyzes the scenario where W is factored using a QR factorization (GEQRF routine in LAPACK):

$$\begin{aligned} QR &:= W \\ b &:= ((QR)^T QR)^{-1} (QR)^T L^{-1} y. \end{aligned} \quad (12)$$

At this point, the compiler exploits the capabilities of the *Matrix algebra* module to perform a series of simplifications on Eq. 12 (Box 2). First, it distributes the transpose operator over the product. Then, it applies the rule

```
times[ trans[ q_?isOrthonormalQ, q_ ] -> id,
```

included as part of the knowledge-base of the module. The rule states that the product $Q^T Q$, when Q is orthogonal with normalized columns, may be rewritten as the identity matrix. Next, since R is square, the inverse is distributed over the product. More mathematical knowledge allows the compiler to rewrite the product $R^{-T} R^T$ as the identity.

$$\begin{aligned} b &:= ((QR)^T QR)^{-1} (QR)^T L^{-1} y; \\ b &:= (R^T Q^T QR)^{-1} R^T Q^T L^{-1} y; \\ b &:= (R^T R)^{-1} R^T Q^T L^{-1} y; \\ b &:= R^{-1} R^{-T} R^T Q^T L^{-1} y; \\ b &:= R^{-1} Q^T L^{-1} y. \end{aligned} \quad (13)$$

Box 2: Simplifications performed by the *Matrix algebra* module.

In Eq. 13, the compiler does not need to process any more inverses; hence, the last step is to decompose the remaining computation into a sequence of products. Once more, y is the only non-matrix operand. Accordingly, the compiler decomposes the equation from right to left. The final algorithm is put together in Alg. 3, QR-GWAS.

Algorithm 3: QR-GWAS

1	$M := h\Phi + (1-h)I$	(SCAL-ADD)	tmp1 = - (h id) + 1 id + h Phi
2	$LL^T = M$	(POTRF)	L2 L2 ^T = tmp1
3	$W := L^{-1}X$	(TRSM)	tmp5 = x ^T L2 ^{-T}
4	$QR = W$	(GEQRF)	Q10 R10 = tmp5 ^T
5	$y := L^{-1}y$	(TRSV)	tmp16 = L2 ⁻¹ y
6	$b := Q^T y$	(GEMV)	tmp21 = Q10 ^T tmp16
7	$b := R^{-1}b$	(TRSV)	tmp29 = R10 ⁻¹ tmp21
			b = tmp29

5.3 Algorithm 3

This third algorithm exploits further knowledge from GWAS, concretely the structure of M , in a manner that may be overlooked even by human experts.

Again, the starting point is Eq. 1. The inner-most inverse is $(h\Phi + (1-h)I)^{-1}$. Instead of multiplying out the expression within the inverse operator, we now describe the alternative path also explored by the compiler: factoring

one of the matrices in the expression. We concentrate in the case where an eigendecomposition of Φ (SYEVD or SYEVR from LAPACK) is chosen:

$$\begin{aligned} ZWZ^T &= \Phi \\ b &:= (X^T (hZWZ^T + (1-h)I)^{-1} X)^{-1} \\ &\quad X^T (hZWZ^T + (1-h)I)^{-1} y \end{aligned} \quad (14)$$

where Z is a square, orthogonal matrix with normalized columns, and W is a square, diagonal matrix.

In this scenario, the *Matrix algebra* module is essential; it allows the compiler to work with alternative representations of Eq. 14. We already illustrated an example where the product $Q^T Q$, Q orthonormal, is replaced with the identity matrix. The freedom gained when defining its own operators, allows the compiler to perform also the opposite transformation, reusing Z as the orthonormal matrix in the rules:

```
id -> times[ Z, trans[ Z ] ];
id -> times[ trans[ Z ], Z ];
```

The resulting expression is

$$\begin{aligned} b &:= (X^T (hZWZ^T + (1-h)ZZ^T)^{-1} X)^{-1} \\ &\quad X^T (hZWZ^T + (1-h)ZZ^T)^{-1} y. \end{aligned} \quad (15)$$

The algebraic manipulation capabilities of the compiler lead to the derivation of further multiple equivalent representations of Eq. 15. We recall that, although we focus on a concrete branch of the derivation, the compiler analyzes the many alternatives. In the branch under study, the quantities Z and Z^T are grouped on the left- and right-hand sides of the inverse, respectively:

$$(X^T (Z(hW + (1-h)I)Z^T)^{-1} X)^{-1};$$

then, since both Z and $hW + (1-h)I$ are square, the inverse is distributed:

$$(X^T (Z^{-T} (hW + (1-h)I)^{-1} Z^{-1}) X)^{-1};$$

finally, by means of the rules:

```
inv[ q_?isOrthonormalQ ] -> trans[ q ];
inv[ trans[ q_?isOrthonormalQ ] ] -> q;
```

which state that the inverse of an orthonormal matrix is its transpose, the expression becomes:

$$(X^T Z (hW + (1-h)I)^{-1} Z^T X)^{-1}.$$

The resulting equation is

$$\begin{aligned} b &:= (X^T Z (hW + (1-h)I)^{-1} Z^T X)^{-1} \\ &\quad X^T Z (hW + (1-h)I)^{-1} Z^T y. \end{aligned} \quad (16)$$

The inner-most inverse in Eq. 16 is applied to a diagonal object (W is diagonal and h a scalar). No more factorizations are needed, $hW + (1-h)I$ is identified as a SCAL-ADD building block, and exposed:

$$\begin{aligned} D &:= hW + (1-h)I \\ b &:= (X^T Z D^{-1} Z^T X)^{-1} X^T Z D^{-1} Z^T y. \end{aligned} \quad (17)$$

D is a diagonal matrix; hence only the inverse applied to $X^T Z D^{-1} Z^T X$ remains to be processed. Among the alternative steps, we consider the mapping of the common segment

$X^T Z$, that appears three times, onto the GEMM building block (matrix-matrix product):

$$\begin{aligned} K &:= X^T Z \\ b &:= (KD^{-1}K^T)^{-1}KD^{-1}Z^T y. \end{aligned} \quad (18)$$

From this point on, the compiler proceeds as shown for the previous examples, and obtains, among others, Alg. 4, EIG-GWAS.

Algorithm 4: EIG-GWAS

<ol style="list-style-type: none"> 1 $ZWZ^T = \Phi$ (EIG) 2 $D := hW + (1-h)I$ (SCAL) 3 $K := X^T Z$ (GEMM) 4 $V := KD^{-1}$ (SCAL) 5 $S := VK^T$ (GEMM) 6 $QR = S$ (QR) 7 $y := Z^T y$ (GEMV) 8 $b := Vy$ (GEMV) 9 $b := Q^T b$ (GEMV) 10 $b := R^{-1}b$ (TRSV) 	<pre> z1 w1 z1^T = Phi tmp2 = - (hid) + 1 id + h w1 tmp7 = x^T z1 tmp13 = tmp7 tmp2⁻¹ tmp20 = tmp13 tmp7^T Q31 R31 = tmp20 tmp36 = z1^T y tmp51 = tmp13 tmp36 tmp66 = Q31^T tmp51 tmp76 = R31⁻¹ tmp66 b = tmp76 </pre>
--	--

At first sight, Alg. 4 might seem to be a suboptimal approach. However, as we show in Sec. 6, it is representative of a family of algorithms that play a crucial role when solving a certain sequence of GLS problems within GWAS.

So far, we have illustrated how our compiler, closely replicating the reasoning of a human expert, automatically generates algorithms for the solution of a single problem (GLS within GWAS). Next, we describe how the compiler proceeds when the target equation solves not a single instance of a problem but a sequence of correlated problems.

5.4 Sequences of problems

Often times an application has to solve not one single problem instance, but a sequence of them; this is the case in GWAS, where one needs to solve a two-dimensional sequence of correlated GLS problems. The naive approach to the solution of a n-dimensional sequence of problems is to wrap an algorithm to solve a single instance of the problem within a n-loop structure. However, this approach might result in a considerable amount of redundant computation. To overcome this limitation, our compiler, by means of the *Analysis of dependencies* module, takes into account the dependencies among the building blocks to reorganize the operations, and to eliminate redundant computation.

First, the compiler labels each input operand with the corresponding dependencies. Then, these dependencies are propagated to the temporary and output operands. Once every operand has been tagged, the compiler 1) creates one loop per dimension of the sequence, and 2) reorders the computation placing each operation in the outermost possible loop, thus eliminating redundant computation.

We use the example of the EIG-GWAS algorithm to illustrate the process. GWAS solves a two-dimensional sequence of problems denoted by the corresponding pair of iterators $\{i, j\}$ (Eq. 1). The extra input needed by the compiler is shown in Box 3. To keep track of the dependencies between dimensions and operands, the compiler attaches the corresponding iterator to each input operand. While Φ is constant, X , y and h are tagged as depending on i , j , and j , respectively, as shown in Alg. 5.

```

dependencies = {
  {X, {'i'}},
  {y, {'j'}},
  {Phi, {}},
  {h, {'j'}},
};

```

Box 3: Extra Mathematica input to the compiler for the analysis of dependencies.

Then, the dependencies are propagated from the input to the output and temporary operands. To this end, the compiler traverses the sequence of operations in Alg. 5 from top to bottom. For each of the operations, the left-hand side operand is labeled with the union of the dependencies of the right-hand side operands. For instance, since h depends on j , the temporary operand D in line 2 also depends on j , and is labelled accordingly (and so are all following occurrences of D). As a result of this process every operand in Alg. 6 is labelled with its dependencies.

Algorithm 5: EIG-GWAS labelled with the initial dependencies

- 1 $ZWZ^T = \Phi$
- 2 $D := h_j W + (1-h_j)I$
- 3 $K := X_i^T Z$
- 4 $V := KD^{-1}$
- 5 $S := VK^T$
- 6 $QR = S$
- 7 $y_j := Z^T y_j$
- 8 $b := Vy_j$
- 9 $b := Q^T b$
- 10 $b := R^{-1}b$

Algorithm 6: EIG-GWAS labelled with all the dependencies

<ol style="list-style-type: none"> 1 $ZWZ^T = \Phi$ 2 $D_j := h_j W + (1-h_j)I$ 3 $K_i := X_i^T Z$ 4 $V_{ij} := K_i D_j^{-1}$ 5 $S_{ij} := V_{ij} K_i^T$ 6 $Q_{ij} R_{ij} = S_{ij}$ 7 $y_j := Z^T y_j$ 8 $b_{ij} := V_{ij} y_j$ 9 $b_{ij} := Q_{ij}^T b_{ij}$ 10 $b_{ij} := R_{ij}^{-1} b_{ij}$

Next, the compiler builds a double loop structure to solve the sequence of problems. Both alternatives, $\forall i \forall j$ and $\forall j \forall i$, are generated and analyzed; the case $\forall j \forall i$ is shown in Alg. 7. A number of operations in Alg. 7, for instance lines 4 and 9, do not depend on i . This means that, for a fixed j , for each iteration of i these two operations unnecessarily recompute the same result m times. In a system that overlooks this situation, the generated algorithms will perform redundant computations.

On the contrary, our compiler identifies this situation and rearranges the operations pushing them to the outermost possible loop: lines 4 and 9 are shifted from the inner-most loop to the outer-most one. Most interestingly, line 3 is completely independent of the dimensions of the problems: Φ is constant along both dimensions, and so are its factors. Therefore, the computation may be performed only once before entering the loops, and reused for every iteration. We illustrate the resulting algorithm in Alg. 8.

Algorithm 7: EIG-GWAS wrapped with a double loop

```

1 for j in 1:t
2   for i in 1:m
3     ZWZT = Φ
4     Dj := hjW ...
5     Ki := XiTZ
6     Vij := KiDj-1
7     Sij := VijKiT
8     QijRij = Sij
9     yj := ZTyj
10    bij := Vijyj
11    bij := QijTbij
12    bij := Rij-1bij

```

Algorithm 8: Final version of EIG-GWAS

```

1 ZWZT = Φ
2 for j in 1:t
3   Dj := hjW ...
4   yj := ZTyj
5   for i in 1:m
6     Ki := XiTZ
7     Vij := KiDj-1
8     Sij := VijKiT
9     QijRij = Sij
10    bij := Vijyj
11    bij := QijTbij
12    bij := Rij-1bij

```

5.5 Cost analysis

We summarize now the computational cost of the three selected algorithms. Although the generation of the cost analysis is not yet automated, it is our top priority. Table 1 includes the cost of the three algorithms for one instance of GLS problem, as well as for the two most common sequences of GLS problems in GWAS (Eq. 1): a one-dimensional sequence where $t = 1$, and a two-dimensional sequence, $t \approx 10^5$.

While QR-GWAS and CHOL-GWAS share the same cost for both types of sequences, suggesting a very similar behavior in practice, the cost of EIG-GWAS differs in both cases. For the one-dimensional sequence the cost of EIG-GWAS is not only greater in theory, the practical constants associated to its terms increase the gap. On the contrary, for the two-dimensional sequence, the cost of EIG-GWAS is lower than the cost of the other two. This analysis suggests that QR-GWAS and CHOL-GWAS are better suited for the one-dimensional case, while EIG-GWAS is better suited for the two-dimensional one. In the next section we confirm these predictions through experimental results.

6. PERFORMANCE EXPERIMENTS

We turn now the attention to numerical results. In the experiments, we compare the algorithms automatically generated by our compiler with LAPACK and GenABEL [2], a widely used package for GWAS-like problems. We present results for the two most representative scenarios in GWAS: one-dimensional ($t = 1$), and two-dimensional ($t > 1$) sequences of GLS problems.

The experiments were performed on an 12-core Intel Xeon X5675 processor running at 3.06 GHz, with 96GB of memory. The algorithms were implemented in C, and linked to the multi-threaded GotoBLAS and the reference LAPACK libraries. The experiments were executed using 12 threads.

We first study the scenario $t = 1$. We compare the performance of QR-GWAS and CHOL-GWAS, with GenABEL’s GWFGLS, and GELS-GWAS, based on LAPACK’s GELS routine. The results are displayed in Fig. 2. As expected, QR-GWAS and CHOL-GWAS attain the same performance and overlap. Most interestingly, our algorithms clearly outperform GELS-GWAS and GWFGLS, obtaining speedups of 4 and 8, respectively.

Next, we present an even more interesting result. The current approach of all state of the art libraries to the case

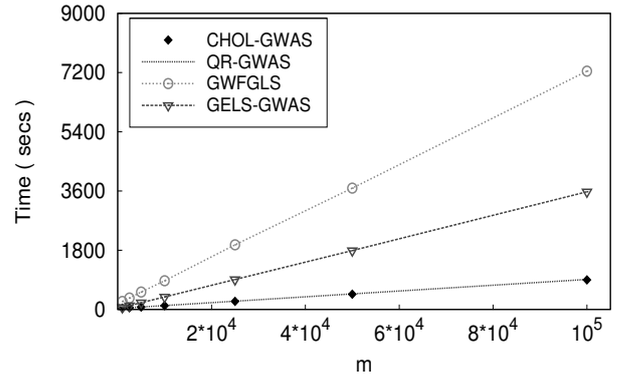


Figure 2: Timings for the one-dimensional sequence of GLS problems within GWAS. Problem sizes: $n = 10000$, $p = 4$, $t = 1$.

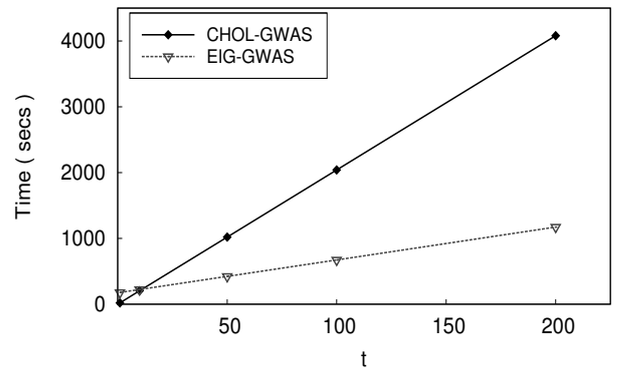


Figure 3: Timings for the two-dimensional sequence of GLS problems within GWAS. Problem sizes: $n = 5000$, $p = 4$, $m = 10^6$.

$t > 1$ is to repeat the experiment t times with the same algorithm used for $t = 1$. On the contrary, our compiler generates the EIG-GWAS algorithm, which particularly suits such scenario. Fig. 3, illustrates how EIG-GWAS outperforms the best algorithm for the case $t = 1$, CHOL-GWAS, by a factor of 4, and therefore outperforms GELS-GWAS and GWFGLS by a factor of 16 and 32 respectively.

The results remark two significant facts: 1) the exploitation of domain-specific knowledge may lead to improvements in state of the art algorithms; and 2) the library user may benefit from the existence of multiple algorithms, each matching a given scenario better than others. In the case of GWAS our compiler achieves both, thus enabling computational biologists to perform larger experiments in considerably less time.

7. CONCLUSIONS

We presented a linear algebra compiler that automatically exploits domain-specific knowledge to generate high-performance algorithms. Our linear algebra compiler mimics the reasoning of a human expert to, similar to a traditional compiler, decompose a target equation into a sequence of library-supported building blocks.

The compiler builds on a number of modules to support the replication of human reasoning. Among them, the *Ma-*

Scenario	QR-GWAS	CHOL-GWAS	EIG-GWAS
One instance	$O(n^3)$	$O(n^3)$	$O(n^3)$
1D sequence	$O(n^3 + mpn^2)$	$O(n^3 + mpn^2)$	$O(n^3 + mpn^2 + mp^2n)$
2D sequence	$O(tn^3 + mtpn^2)$	$O(tn^3 + mtpn^2)$	$O(n^3 + mpn^2 + mtp^2n)$

Table 1: Computational cost for the three selected algorithms generated by the compiler.

trix algebra module, which enables the compiler to freely manipulate and simplify algebraic expressions, and the *Properties inference* module, which is able to infer properties of complex expressions from the properties of the operands.

The potential of the compiler is shown by means of its application to the challenging *genome-wide association study* equation. Several of the dozens of algorithms produced by our compiler, when compared to state of the art ones, obtain n -fold speedups.

As future work we plan to develop two new modules for the compiler: a module for the generation of MATLAB and C code, and a module for the analysis of the computational cost of the algorithms, as a first approach towards a more robust metric to compare the performance of the generated algorithms.

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