

## ADJUSTING A PROGRAM TRANSFORMATION FOR LEGALITY

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### ABSTRACT

Program transformations are one of the most valuable compiler techniques to improve parallelism or data locality. However, restructuring compilers have a hard time coping with data dependences. A typical solution is to focus on program parts where the dependences are simple enough to enable any transformation. For more complex problems is only addressed the question of checking whether a transformation is legal or not. In this paper we propose to go further. Starting from a transformation with no guarantee on legality, we show how we can correct it for dependence satisfaction. Two directions are explored: first when transformation properties can be explicitly expressed and second when they are implicit as in the data locality transformation case. Generating code having the best properties is a direct application of this result.

*Keywords:* Program transformations, legality, dependences, polyhedral model, locality.

### 1. Introduction

The task of optimizing compute-bound programs is crucial for present day supercomputers if we notice that most of these machines run at a few percent of their peak performance. The problem can be stated as a combinatorial optimization problem, but due to the complexity of real-life programs and computers, this approach is not practical. Most of the time, we start from a first implementation, and try to improve its performance by successive transformations. Beside improving the performances, a transformation must be legal, i.e. must not change the final results of the program. This is usually enforced by using only transformations that respect dependences [22]. While selecting an optimizing transformation is not too difficult for an experienced programmer, adjusting this transformation for legality is a tedious and error-prone process.

To bypass the dependence problem, most of the existing methods apply only to perfect loop nests in which dependences are non-existent or have a special form (fully permutable loop nests) [27]. To enlarge their application domain some preprocessing, e.g. *loop skewing* or *code sinking*, may be applied [27,1,14]. More ambitious techniques do not lay down any requirement on dependences, but are limited to propose *solution candidates* then to *check* them for legality [17,8]. If the candidate is proved to violate dependences, then the proposed transformation is discarded and another candidate, perhaps having less interesting properties is studied. In this paper, we present a method that goes beyond checking by adjusting – if possible – a transformation for dependence satisfaction, without modifying its optimizing properties. This technique can be used to correct a transformation candidate as well as to replace preprocessing. The technique has been designed in the context of locality-improving transformations, but can be applied in many other cases. On the other hand, only transformations which can be represented as affine transformations in iteration space can be corrected in this way.

This paper is organized as follows. In section 2 we outline the background of this work. Section 3 deals with the transformations in the polyhedral model and focuses on their dependences constraints. Section 4 shows how it is possible to correct a transformation for legality. Section 5 compares our proposal to previous work in the field of locality enhancement then section 6 concludes and discusses future work.

## 2. Background and Notations

A loop in an imperative language like C or FORTRAN can be represented using a  $n$ -entry column vector called its *iteration vector*:  $\vec{x} = (i_1, i_2 \dots i_n)^T$ , where  $i_k$  is the  $k^{th}$  loop index and  $n$  is the innermost loop. The surrounding loops and conditionals of a statement define its *iteration domain*. The statement is executed once for each element of the iteration domain. When loop bounds and conditionals only depend on surrounding loop counters, formal parameters and constants, the iteration domain can be specified by a set of linear inequalities defining a polyhedron [18]. The term *polyhedron* will be used in a broad sense to denote a *convex set of points in a lattice* (also called  $\mathbb{Z}$ -polyhedron or lattice-polyhedron), i.e. a set of points in a  $\mathbb{Z}$  vector space bounded by affine inequalities [24]. A maximal set of consecutive statements in a program with such polyhedral iteration domains is called a *static control part* (SCoP) [7]. Figure 1 illustrates the correspondence between static control and polyhedral domains. Each integral point of the polyhedron corresponds to an *operation*, i.e. an instance of the statement. The notation  $S(\vec{x})$  refers to the instance of the statement  $S$  with iteration vector  $\vec{x}$ . The execution of the operations follows *lexicographic order*. This means in a  $n$ -dimensional polyhedron, the operation corresponding to the integral point defined by the coordinates  $(a_1 \dots a_n)$  is executed before those corresponding to the coordinates  $(b_1 \dots b_n)$  iff

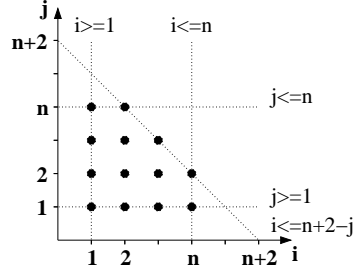
$$\exists i, 1 \leq i < n, (a_1 \dots a_i) = (b_1 \dots b_i) \wedge a_{i+1} < b_{i+1}.$$

We will use  $\ll$  and  $\leq$  for the strict and non strict lexicographic order, respectively.

```

do i = 1, n
  do j = 1, n
    if (i <= n+2-j)
S1:      B[i+j][2*i+1] = ...

```



$$\begin{bmatrix} 1 & 0 \\ -1 & 0 \\ 0 & 1 \\ 0 & -1 \\ -1 & -1 \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix} + \begin{pmatrix} -1 \\ n \\ -1 \\ n \\ n+2 \end{pmatrix} \geq \vec{0}$$

(a) surrounding control of  $S_1$ (b) iteration domain of  $S_1$ 

Figure 1: Static control and corresponding iteration domain

Each statement may include one or several *references* to arrays (scalars are zero-dimensional arrays). When the subscript function  $f(\vec{x})$  of a reference is affine, we can write it  $f(\vec{x}) = F\vec{x} + \vec{a}$  where  $F$  is called the *subscript matrix* and  $\vec{a}$  is a constant vector. For instance, the reference to the array  $B$  in figure 1(a) is  $B[f(\vec{x})]$  with  $f\left(\begin{smallmatrix} i \\ j \end{smallmatrix}\right) = \begin{bmatrix} 1 & 1 \\ 2 & 0 \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix} + \begin{pmatrix} 0 \\ 1 \end{pmatrix}$ .

In this paper, matrices are always denoted by capital letters, vectors and functions in vector spaces are not. When an element is statement-specific, it is subscripted like  $A_S$ ; the subscript may be omitted when it is clear from the context.

### 3. Affine Transformations

#### 3.1. Formulation

The goal of a transformation is to modify the original execution order of the operations. A convenient way to express the new order is to give for each operation an execution date. However, defining all the execution dates separately would usually require very large scheduling systems. Thus optimizing compilers build schedules at the statement level by finding a function specifying an execution time for each instance of the corresponding statement. These functions are chosen affine for multiple reasons: this is the only case where we are able to decide exactly the transformation legality and where we know how to generate the target code. Thus, scheduling functions have the following shape:

$$\theta_S(\vec{x}_S) = T_S \vec{x}_S + \vec{t}_S, \quad (1)$$

where  $\vec{x}_S$  is the iteration vector,  $T_S$  is a constant transformation matrix and  $\vec{t}_S$  is

a constant vector (possibly including structure parameters).

It has been extensively shown that linear transformations can express most of the useful transformations. In particular, loop transformations (such as loop reversal, permutation or skewing) can be modeled as a simple particular case called unimodular transformations (the  $T_S$  matrix has to be square and has determinant  $\pm 1$ ) [5,25]. Complex transformations such as tiling [26] can be achieved using linear transformations as well [28]. These transformations modify the source polyhedra into target polyhedra containing the same points, but with a new lexicographic order. Considering an original polyhedron defined by the system of affine constraints  $A\vec{x} + \vec{c} \geq \vec{0}$  and the transformation function  $\theta$  leading to the target index  $\vec{y} = T\vec{x}$ , we deduce that the transformed polyhedron can be defined by  $(AT^{-1})\vec{y} + \vec{c} \geq \vec{0}$  (there exists more convenient way to describe the target polyhedron as discussed in [6]). For instance, let us consider the polyhedron in figure 2(a) and the transformation function  $\theta\left(\begin{smallmatrix} i \\ j \end{smallmatrix}\right) = \begin{bmatrix} 1 & 1 \\ 0 & 1 \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix}$ . The corresponding transformation is a well known *iteration space skewing* and the resulting polyhedron is shown in figure 2(c).

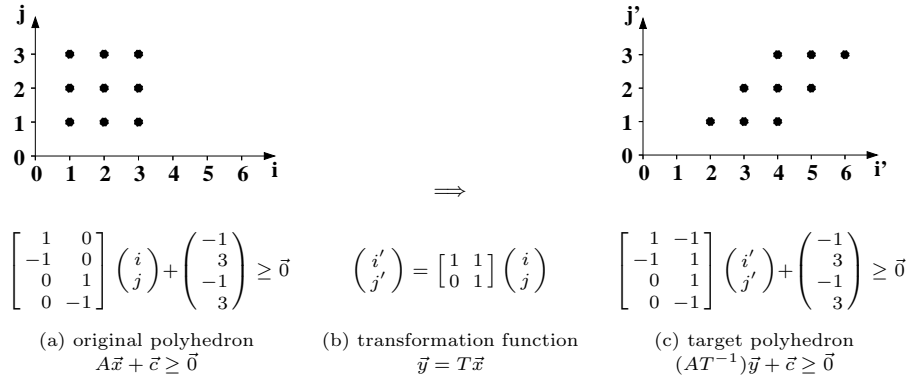


Figure 2: A skewing transformation

### 3.2. Legality

In general, applying an arbitrary transformation to a program will change its semantics.

Two operations are said to be *in dependence* if they share a variable (memory cell) and at least one the operations modifies it. This definition was suggested by Bernstein [9] and is the most widely used one in works about program transformation. It is a sufficient condition for parallelism, but is by no means necessary, as the well known case of *reductions* shows.

Many tests have been designed for dependence checking. Most of these are based on sufficient conditions for independence. They give an approximate, conservative answer. The best known examples are the GCD-test [3], and the Banerjee test [4]. On the other hand, one can use classical algorithms from Linear Integer program-

ming to get exact answers, as in the Omega-test [23] and the Simplex-Gomory test [12]. In the same way many dependence representations are possible, from the simplest ones as *dependence levels* [2] to the most precise as *dependence polyhedra* [16]. We chose in this paper to use the most precise representation of dependences: the dependence polyhedra. However, many authors have noticed that approximate dependences are special cases of dependence polyhedra. Hence, our method applies whatever representation is chosen, provided the approximation is conservative.

In section 3.2.1, we recall how dependences in a SCoP can be expressed exactly using linear (in)equalities. Then we show in section 3.2.2 how to build the legal transformation space.

### 3.2.1. Dependence Graph

A convenient way to represent the scheduling constraints is the *dependence graph*. In this directed graph, each program statement is represented using a unique vertex, and the existing dependence relations are represented using edges. Each vertex is labelled with the iteration domain of the corresponding statement and the edges with the dependence polyhedron describing the dependence.

The dependence relation can be defined in the following way:

**Definition 1** A statement  $R$  **depends** on a statement  $S$  (written  $S\delta R$ ) if there exists an operation  $S(\vec{x}_1)$ , an operation  $R(\vec{x}_2)$  and a memory location  $m$  such that:

1.  $S(\vec{x}_1)$  and  $R(\vec{x}_2)$  refer the same memory location  $m$ , and at least one of them writes to that location;
2.  $\vec{x}_1$  and  $\vec{x}_2$  respectively belong to the iteration domain of  $S$  and  $R$ ;
3. in the original sequential order,  $S(\vec{x}_1)$  is executed before  $R(\vec{x}_2)$ .

From this definition follows the description of the *dependence polyhedron* by affine (in)equalities. The constraints systems have the following components:

1. *Same memory location*: assuming that  $m$  is an array location, this constraint is the equality of the subscript functions of a pair of references to the same array:  $F_S\vec{x}_S + \vec{a}_S = F_R\vec{x}_R + \vec{a}_R$ .
2. *Iteration domains*: both  $S$  and  $R$  iteration domains can be described using affine inequalities, respectively  $A_S\vec{x}_S + \vec{c}_S \geq \vec{0}$  and  $A_R\vec{x}_R + \vec{c}_R \geq \vec{0}$ .
3. *Precedence order*: this constraint can be separated into a disjunction of as many parts as there are common loops to both  $S$  and  $R$ . Each case corresponds to a common loop depth and is called a *dependence level*. For each dependence level  $l$ , the precedence constraints are the equality of the loop index variables at depth lesser to  $l$ :  $x_{R,i} = x_{S,i}$  for  $i < l$  and  $x_{R,l} > x_{S,l}$  if  $l$  is less than the common nesting level. Otherwise, there are no additional constraints and the dependence only exists if  $S$  is textually before  $R$ . Such constraints can be written using linear inequalities:  $P_S\vec{x}_S - P_R\vec{x}_R + \vec{b} \geq \vec{0}$ .

Thus, the dependence polyhedron for  $S\delta R$  at a given level  $l$  and for a given pair of references  $p$  can be described using the following system of (in)equalities:

$$\mathcal{D}_{S\delta R,l,p} : D \begin{pmatrix} \vec{x}_S \\ \vec{x}_R \end{pmatrix} + \vec{d} = \begin{bmatrix} F_S & -F_R \\ A_S & 0 \\ 0 & A_R \\ P_S & -P_R \end{bmatrix} \begin{pmatrix} \vec{x}_S \\ \vec{x}_R \end{pmatrix} + \begin{pmatrix} \vec{a}_S - \vec{a}_R \\ \vec{c}_S \\ \vec{c}_R \\ \vec{b} \end{pmatrix} \stackrel{=}{\geq} \vec{0} \quad (2)$$

There is a dependence  $S\delta R$  if there exists an integral point inside  $\mathcal{D}_{S\delta R,l,p}$ . This can be easily checked with some linear integer programming tool like PipLib<sup>a</sup> [11]. If this polyhedron is not empty, there is an edge in the dependence graph from the vertex corresponding to  $S$  up to the one corresponding to  $R$ , labelled with  $\mathcal{D}_{S\delta R,l,p}$ . For the sake of simplicity we will ignore subscripts  $l$  and  $p$  and refer in the following to  $\mathcal{D}_{S\delta R}$  as the only dependence polyhedron describing  $S\delta R$ .

### 3.2.2. Legal Transformation Space

Considering the transformations as scheduling functions, the time interval in the target program between the executions of two operations  $R(\vec{x}_R)$  and  $S(\vec{x}_S)$  is

$$\Delta_{R,S} \begin{pmatrix} \vec{x}_S \\ \vec{x}_R \end{pmatrix} = \theta_R(\vec{x}_R) - \theta_S(\vec{x}_S). \quad (3)$$

If there exists a dependence  $S\delta R$ , i.e. if  $\mathcal{D}_{S\delta R}$  is not empty, then  $\Delta_{R,S} \begin{pmatrix} \vec{x}_S \\ \vec{x}_R \end{pmatrix}$  must be lexicopositive in  $\mathcal{D}_{S\delta R}$  (intuitively, the time interval between two operations  $R(\vec{x}_R)$  and  $S(\vec{x}_S)$  such that  $R(\vec{x}_R)$  depends on  $S(\vec{x}_S)$  must be at least  $(0, \dots, 0, 1)^T$ , the smallest time interval: this guarantees that the operation  $R(\vec{x}_R)$  is executed after  $S(\vec{x}_S)$  in the target program). This condition represents as many constraints as there are points in  $\Delta_{R,S}$ . Fortunately, all these constraints can be compacted in a small set of affine constraints with the help of Farkas Lemma [13].

**Lemma 1** (Affine form of Farkas Lemma [24]) *Let  $\mathcal{D}$  be a nonempty polyhedron defined by the inequalities  $A\vec{x} + \vec{b} \geq \vec{0}$ . Then any affine function  $f(\vec{x})$  is nonnegative everywhere in  $\mathcal{D}$  iff it is a positive affine combination:*

$$f(\vec{x}) = \lambda_0 + \vec{\lambda}^T (A\vec{x} + \vec{b}), \text{ with } \lambda_0 \geq 0 \text{ and } \vec{\lambda}^T \geq \vec{0}.$$

$\lambda_0$  and  $\vec{\lambda}^T$  are called Farkas multipliers.

$\Delta_{R,S}$  is a vector. For it to be lexicopositive, some of its components must be constrained to be either non negative or strictly positive. Let us apply Farkas Lemma to one of the constrained components. We can find a non-negative scalar  $\lambda_0$  and a non-negative vector  $\vec{\lambda}^T$  such that:

$$T_{R,\bullet}\vec{x}_R + t_R - (T_{S,\bullet}\vec{x}_S + t_S) - \delta = \lambda_0 + \vec{\lambda}^T \left( D \begin{pmatrix} \vec{x}_S \\ \vec{x}_R \end{pmatrix} + \vec{d} \right) \quad (4)$$

In this formula,  $T_{R,\bullet}$  and  $T_{S,\bullet}$  are corresponding rows in the  $T_R$  and  $T_S$  matrices, and  $\delta$  is zero or one according to the position of the rows.

<sup>a</sup>PipLib is freely available at <http://www.prism.uvsq.fr/~cedb>

This formula can be split in as many equalities as there are independent variables ( $\vec{x}_S$  and  $\vec{x}_R$  components and parameters) by equating their coefficients in both sides of the formula. The Farkas multipliers can be eliminated by using the Fourier-Motzkin projection algorithm [24]. The result is a system of affine constraints on the coefficients of the transformation (the elements of  $T_{R,\bullet}$  and  $T_{S,\bullet}$ ). The important point is that this system is the same for all rows of the scheduling matrices and depends only on the dependence to be satisfied. Furthermore, it depends linearly on the value of  $\delta$ . These systems completely characterize the legal transformations of a program, and can be computed once and for all as soon as the dependences are known.

#### 4. Correcting Transformations

Both optimizing compilers and programmers have a tendency to think of performances first and to check legality afterward. The basic framework is first to find the best transformation (e.g. in the case of data locality improvement, which references carry the most reuse and necessitate new access patterns, which rank constraints should be respected by the corresponding transformation functions, etc.), then to *check* if a candidate transformation is legal or not<sup>b</sup>. If the check fails, build and test another candidate, and so on. The major advantage of such a framework is to focus firstly on the most interesting properties, and the main drawback is to forsake these properties if a legal transformation is not directly found after a simple check of a candidate solution. In this section we will show how it is possible to correct a candidate transformation for dependences, firstly when it can be described using explicit constraints as discussed in section 4.1. Then in section 4.2 we study the special case of data locality improvement where the transformation properties are hidden.

##### 4.1. Transformations With Explicit Properties

Experts or optimizing compilers have a wide choice of optimizing transformations for a given program. Each transformation has a more or less precise cost model which helps in deciding whether to apply the transformation or not. In the polyhedral framework, many transformations are related to well chosen scheduling functions [5,13,10]. For instance, generalized loop interchange is associated to schedules whose matrix is a permutation matrix [5]. Trying to use these transformations as they are may result in a negative dependence test. Let us consider the code in Figure 3. An expert or an optimizing compiler may decide that moving the  $i$ -loop innermost would result in better locality. But because of complex dependences, using directly the loop interchange transformation  $\theta_S \begin{pmatrix} i \\ j \end{pmatrix} = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix}$  is not legal. This will lead usually to rejecting the transformation.

<sup>b</sup>This can be done easily by instantiating the transformation functions in the space of all affine transformation as defined in section 3.2, then checking whether it belong to the legal subset using any linear algebra tool.

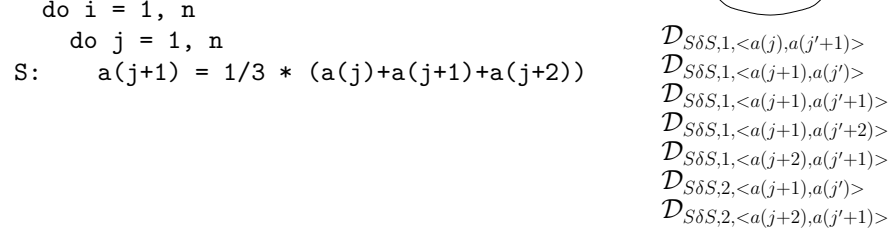


Figure 3: Original Hyperbolic-PDE program and dependence graph

The polyhedral model allow more flexibility when defining such transformations. Moreover, one can work with an incompletely specified transformations and use the legality constraints as a way of solving for the missing coefficients. The method consists in stating the constraints the transformation has to satisfy, then solving these constraints and the legality constraints (4), using a linear algebra tool as PipLib. If the system does not have a solution, we conclude that there is no legal instance of the proposed transformation. For example, “innermosting” the  $i$ -loop in the code in Figure 3 means that we are looking for a transformation function  $\theta_S \begin{pmatrix} i \\ j \end{pmatrix} = \begin{bmatrix} T_{1,1} & T_{1,2} \\ T_{2,1} & T_{2,2} \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix} + \begin{pmatrix} t_1 \\ t_2 \end{pmatrix}$  with as only constraints  $T_{2,1} = 1$  and  $T_{2,2} = 0$ . By solving the system we find the solution  $\theta_S \begin{pmatrix} i \\ j \end{pmatrix} = \begin{bmatrix} 1 & 1 \\ 1 & 0 \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix}$ . Expressed using classical transformation techniques, it is a combination of loop skewing and loop interchange. It leads to the target program in Figure 4 and as expected to a better cache behavior (on a i386 1GHz system with 128KB L1 cache memory and  $n=33000$  the number of cache misses of the original program is 68M but 43M for the target one).

```

do i' = 2, 2*n
  do j' = max(i'-n,1), min(i'-1,n)
    j = i'-j' ;
    i = i' ;
S:    a(j+1) = 1/3 * (a(j)+a(j+1)+a(j+2))

```

Figure 4: Final Hyperbolic-PDE program

#### 4.2. Transformations With Implicit Properties: Data Locality

Cache are used in most computer systems to compensate for the mismatch



between processor and memory performance (at the time of writing, factors of 10 to 100 are commonplace). Caches work by exploiting locality, i.e. the fact that accesses to each memory cell and its close neighbors have a tendency to cluster in the program code. While this is found to work very well for ordinary programs, it fails for compute-bound codes where large datasets are accessed according to very regular patterns. The basic framework for increasing cache hit rates is to move references to a given memory cell (or cache line) to neighboring iterations of some innermost loop. This reduces the elapsed time between two accesses and hence decreases the probability that the cell has been evicted from the cache. Such a transformation usually changes the execution order of the program, hence it must be checked for legality before being applied.

Another way of expressing the same intuition is to assign an execution date (a schedule) to each operation, and to take care that the date of accesses to the same memory cell are “almost” equal. This is usually obtained by requiring that the outer components of the schedule are equal. The method continues by applying a completion procedure to achieve an invertible transformation function (see [27] for references).

For instance, let us consider self-temporal locality and a reference  $B[f(\vec{x})]$  to an array  $B$  with the affine subscript function  $f(\vec{x}) = F\vec{x} + \vec{a}$ . Two instances of this reference,  $B[f(\vec{x}_1)]$  and  $B[f(\vec{x}_2)]$  refers the same memory location iff  $f(\vec{x}_1) = f(\vec{x}_2)$ , that is when  $F\vec{x}_1 + \vec{a} = F\vec{x}_2 + \vec{a}$ , then iff  $F\vec{x}_r = \vec{0}$  with  $\vec{x}_r = \vec{x}_1 - \vec{x}_2$ . Thus there is self-temporal reuse when  $\vec{x}_r \in \ker F$ . The basis vectors of  $\ker F$  give the reuse directions for the reference  $B[f(\vec{x})]$ ; if  $\ker F$  is trivial, there is no self-temporal reuse for the corresponding reference. Reuse can be exploited if the transformed iteration order follows one of the reuse directions. Then we have to find a vector orthogonal to the chosen reuse direction to be the first part of the transformation matrix  $T$ . If this partial transformation does not violate dependences, we have many choices for the completion procedure in order for the transformation function to be one-to-one either by considering artificial dependences [20,15] or not [6]. As an example, consider the following pseudo-code:

```

do i = 1, n
  do j = 1, n
S1:    ... B[j] ...

```

the subscript function of the reference  $B[j]$  is  $f\left(\begin{smallmatrix} i \\ j \end{smallmatrix}\right) = \begin{bmatrix} 0 & 1 \\ 0 & 0 \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix}$ , the kernel of the subscript matrix is then  $\ker F = \text{span}\{(1,0)\}$ . Thus there is reuse generated by the reference  $B[j]$ , and we can exploit it thanks to a transformation matrix built with an orthogonal vector to the reuse direction, e.g.  $[0 \ 1]$  and its completion to a unimodular transformation matrix as described in [15]:  $T = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix}$ . The transformation function would be  $\theta\left(\begin{smallmatrix} i \\ j \end{smallmatrix}\right) = \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} \begin{pmatrix} i \\ j \end{pmatrix}$ , i.e. a loop interchange (the reader may care to verify that this solution does exploit the reuse of the reference

$B[j]$ ). It is easy to generalize this method to several references by considering not only a reuse direction vector, but a reuse direction space (built with one basis vector per reference). It appears that there are a lot of degrees of freedom when looking for a transformation improving self-temporal locality, since it is possible to choose the reuse direction space, the completion method and the constant vector of the transformation function.

Let us consider self-temporal locality and a transformation candidate before completion  $\theta_{S_c}(\vec{x}_S) = T_{S_c}\vec{x}_S$ . This function has the property that, modified in the following way:

$$\theta_S(\vec{x}_S) = C_S T_{S_c} \vec{x}_S + \vec{t}_S, \quad (5)$$

where  $C_S$  is an invertible matrix and  $\vec{t}_S$  is a constant vector, the locality properties are left unmodified for each time step. Intuitively, if  $\theta_{S_c}$  gives the same execution date for  $\vec{x}_1$  and  $\vec{x}_2$ , then the transformed function  $\theta_S$  does it as well. In the same way if the dates are different with  $\theta_{S_c}$ , then the transformed function  $\theta_S$  returns different dates. But while the values of  $C_S$  and  $\vec{t}_S$  do not change the self-temporal locality properties<sup>c</sup>, they can change the transformation from an illegal to a legal one.

It is clearly not possible to check all these transformations for legality. In the following we study another way: we show how to find, when possible, the unknown components  $C_S T_{S_c}$  and  $\vec{t}_S$  of formula 5 in order to construct a legal transformation.

Correcting formulae similar to (5) and having the same type of degrees of freedom can be used to achieve every type of locality (*self* or *group - temporal* or *spatial*) [25,8]. The challenge is, considering the candidate transformation matrices  $T_{S_c}$ , to find the *corrected matrices*  $C_S T_{S_c}$  and the constant vectors  $\vec{t}_S$  in order for the transformation system to be legal for dependences.

This problem can be solved in an iterative way, each dimension being considered as a stand-alone transformation. Each row of  $C_S T_{S_c}$  is a linear combination of the rows of  $T_{S_c}$ . Thus, the unknown in the  $i^{th}$  algorithm iteration are, for each statement, the linear combination coefficients building the  $i^{th}$  row of  $C_S T_{S_c}$  from  $T_{S_c}$  and the constant factor of the corresponding  $\vec{t}_S$  entry. After each iteration, we have to update the dependence graph since, by a property of lexicographic order, there is no need to consider the already satisfied dependences. Thus, to find a solution is easier as the algorithm iterates. The algorithm is shown in figure 5.

Let us illustrate how the algorithm works using the example in figure 6. Suppose that an optimizing compiler would like to exploit the data reuse generated by the references to the array  $A$  of the program in figure 6(a) and that it suggests the transformation candidates in figure 6(b). As shown by the graph describing the resulting operation execution order, where each arrow represents a dependence relation and each backward arrow is a dependence violation, the transformation system is not

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<sup>c</sup>This amount to noticing that the amount of locality in the transformed program is linked to the *rank* of  $T_{S_c}$ . For a more formal discussion, see [8].

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**Correction Algorithm:** adjust a transformation system to respect dependences

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**Input:** a dependence graph DG, the transformation candidates  $\theta_{S_c}(\vec{x}_S) = T_{S_c}\vec{x}_S$ .

**Output:** the legal transformations  $\theta_S(\vec{x}_S) = C_S T_{S_c} \vec{x}_S + \vec{t}_S$ .

1. for dimension  $i = 1$  to maximum dimension of  $T_{S_c}$ 
    - (a) build the legal transformation space with:
      - for each edge in DG, the constraints of (4) for the  $i^{th}$  row of  $T_{R_c}$  and  $T_{S_c}$
      - the constraints equating the  $i^{th}$  row entries of each  $C_S T_{S_c}$  with a linear combination of  $T_{S_c}$  entries whose coefficients are unknown
    - (b) for each statement, remove from the solution space the trivial solution where  $\forall j \geq i$  the linear combination coefficient of the  $j^{th}$  row of  $T_{S_c}$  is null
    - (c) if the solution space is empty, return  $\emptyset$ , else
      - i. pick the solution giving for each statement the minimum values for the entries of the  $i^{th}$  row of  $C_S T_{S_c}$  and the  $i^{th}$  element of  $\vec{t}_S$
      - ii. update DG: for each edge in DG, add to the dependence polyhedron the constraint equating the  $i^{th}$  dimension of  $C_S T_{S_c} \vec{x}_S + \vec{t}_S$  of the statements labelling the source and destination vertices (this may empty the polyhedron for integral solutions)
      - iii. if every dependence polyhedra in DG are empty, goto 2
      - iv. for each statement, update the candidate transformation  $T_{S_c}$ :
        - replace a row such that the corresponding linear combination coefficient is not null with the  $i^{th}$  row
        - replace the  $i^{th}$  row with the  $i^{th}$  row of  $C_S T_{S_c}$
  2. return the transformation functions  $\theta_S(\vec{x}_S) = C_S T_{S_c} \vec{x}_S + \vec{t}_S$ .
- 

Figure 5: Algorithm to correct the transformation functions

legal. The correction algorithm modifies successively each transformation dimension. Each stand-alone transformation splits up the operations into sets such that there are no backward arrows between sets. The algorithm stops when there are no more backward arrows or when every dimension has been corrected. Then any polyhedral code generator, like CLooG<sup>d</sup> [6], can generate the target code. Choosing transformation coefficients as small as possible (step 1(c)i) is a heuristic helping code generators to avoid control overhead.

The correctness of the algorithm comes from two properties: (1) the target

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<sup>d</sup>CLooG is freely available at <http://www.prism.uvsq.fr/~cedb>

transformations are legal, (2) the  $C_S$  matrices are invertible. The legality is achieved because each transformation part is chosen in the legal transformation space (step 1a). The second property follows from the updating policy (step 1(c)iv): at start the  $C_S$  matrices are identities. During each iteration, we exchange their rows, multiply some rows by non null constants (as guaranteed by step 1b) and add to these rows a linear combination of the other rows. Each of these transformations does not modify the invertibility property.

## 5. Related Work

Since they cannot deal with (complex) dependences, the earliest works on locality improvement discuss *enabling transformations* to modify the program in such a way that the proposed method can apply. Wolf and Lam [25] proposed in their seminal *data locality optimizing algorithm* to use *skewing* and *reversal* to enable *tiling* as in previous works on automatic parallelization. McKinley et al. [21] proposed a technique based on a detailed cost model that drives the use of *fusion* and *distribution* mainly to enable loop *permutation*. Such methods are limited by the set of directives they use (like *fuse* or *skew*) and because they have to apply them in a definite order. We claim that proposing (and correcting) scheduling functions is more complete and has better compositionality properties.

A significant step on preprocessing techniques to produce fully permutable loop nests has been achieved by Ahmed et al. [1]. They use Farkas Lemma to find a valid *code sinking*-like transformation if it exists. But this transformation is still independent from the optimization itself and it is limited to produce a fully permutable loop nest. The method proposed in this paper may find solutions even when it is not possible to extract such a loop nest.

The method of Griehl et al. [14] is quite different. Their aim is to minimize the amount of communication in a distributed program, which is indeed a kind of locality optimization. They first take care of dependences by finding a legal space-time transformation (i.e. a schedule and a placement) and then tile in space-time to achieve the optimal granularity. Adapting these ideas to cache optimization seems by no mean obvious, although it is an interesting subject for further research.

Reasoning directly on scheduling functions, Li and Pingali proposed a completion algorithm to build a non-unimodular transformation function from a partial matrix, such that starting from a legal transformation, the completed transformation stay legal for dependences [20]. In the same spirit, Griehl et al. [15] extended an arbitrary matrix describing a legal transformation to a square invertible matrix. In contrast, we show in this paper how to find the valid functions before completion.

## 6. Conclusion and Future Work

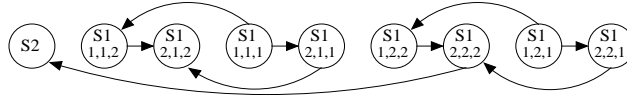
In this paper we presented a general method correcting a program transformation for legality with no consequence on its properties. It can be applied either

```

do i = 1, n
  do j = 1, n
    do k = 1, n
S1:      A(j,k) = A(j,k) + B(i,j,k) / A(j,k-1)
S2:      c = A(n,n) + 1
    
```

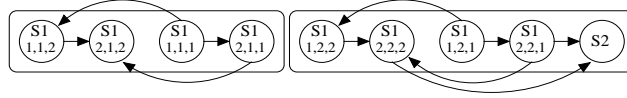
(a) Original program

$$\theta_{S1c} \begin{pmatrix} i \\ j \\ k \end{pmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & -1 \\ 0 & 0 & 0 \end{bmatrix} \begin{bmatrix} i \\ j \\ k \end{bmatrix} + \begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}; \theta_{S2c} = \begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}$$



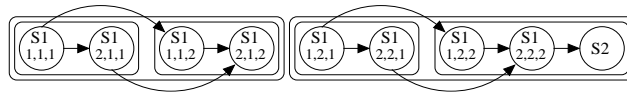
(b) Transformation function candidates

$$\theta_{S1c} \begin{pmatrix} i \\ j \\ k \end{pmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & -1 \\ 0 & 0 & 0 \end{bmatrix} \begin{bmatrix} i \\ j \\ k \end{bmatrix} + \begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}; \theta_{S2c} = \begin{pmatrix} n \\ 0 \\ 0 \end{pmatrix}$$



(c) First correction iteration

$$\theta_{S1c} \begin{pmatrix} i \\ j \\ k \end{pmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & -1 & 0 \\ 0 & 0 & 1 \end{bmatrix} \begin{bmatrix} 0 & 1 & 0 \\ 0 & 0 & -1 \\ 0 & 0 & 0 \end{bmatrix} \begin{bmatrix} i \\ j \\ k \end{bmatrix} + \begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix}; \theta_{S2c} = \begin{pmatrix} n \\ n \\ 0 \end{pmatrix}$$



(d) Second and last correction iteration

```

do j = 1, n
  do k = 1, n
    do i = 1, n
S1:      A(j,k) = A(j,k) + B(i,j,k) / A(j,k-1)
S2:      c = A(n,n) + 1
    
```

(e) Target program

Figure 6: Iterative transformation correction principle ( $n = 2$  for graphs)

when the properties can be explicitly expressed as affine constraints, either when they are carried implicitly as data locality properties. It has been implemented in the Chunky prototype [8], advantageously replacing usual *enabling* preprocessing techniques and saving a significant amount of interesting transformations from being ignored. It could be used combined with a wide range of existing optimizing techniques and in particular for data locality improvement methods, for the single processor case as well as for parallel systems using space-time mappings [19].

Further implementation work is necessary to handle real-life benchmarks in our prototype and to provide full statistics on corrected transformations. Moreover, the question of scalability is left open since, for several tenth of deeply nested statements, the number of unknown in the constraint systems can become embarrassingly large. Splitting up the problem according to the dependence graph is a solution under investigation.

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