On the Synthesis of Integral and Dynamic Recurrences

by

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To Jon, with Love.
Abstract

Synthesis techniques for regular arrays provide a disciplined and well-founded approach to the design of classes of parallel algorithms. The design process is guided by a methodology which is based upon a formal notation and transformations.

The mathematical model underlying synthesis techniques is that of affine Euclidean geometry with embedded lattice spaces. Because of this model, computationally powerful methods are provided as an effective way of engineering regular arrays. However, at present the applicability of such methods is limited to so-called affine problems.

The work presented in this thesis aims at widening the applicability of standard synthesis methods to more general classes of problems. The major contributions of this thesis are the characterisation of classes of integral and dynamic problems, and the provision of techniques for their systematic treatment within the framework of established synthesis methods. The basic idea is the transformation of the initial algorithm specification into a specification with data dependencies of increased regularity, so that corresponding regular arrays can be obtained by a direct application of the standard mapping techniques.

We will complement the formal development of the techniques with the illustration of a number of case studies from the literature.
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Any remaining errors or omissions are the author's sole responsibility.

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Chapter 1

Introduction

Parallel computing (see, e.g., [HwBr85, Kri89]) is a fast growing branch of computing science. Many of today's computer applications require a great computing power at very high speed. Higher performance for these applications can be achieved through parallel processing, that is by allowing the concurrent and cooperative execution of their computations. Parallel computing has become a necessary and effective alternative to building faster sequential computers: - necessary as hardware components are approaching their technological limits as for the level of circuit integration and the speed of signal propagation; - effective as significant "speed-ups" have been obtained for many applications of interest. Also, technological advances have made it possible for both commercial and research parallel computers and systems to be widely available at a relatively low cost. Research in parallel computing spans from architectural and algorithmic issues, to programming languages and compiler technology, to theoretical models and complexity theory (see, e.g., [Kri89] for an overview).

A substantial part of the research in parallel computing has been devoted to the study of parallel machines which are *algorithm specific* [Sn-et-al85]. Algorithm specific machines are machines whose architectures are specialised to provide efficient solutions for classes of problems which share a common solution method. Hence they provide a trade-off in which improved performance is gained at the expense of generality. This thesis deals with a particular aspect of algorithm specific computing, that of the synthesis of *regular (processor) arrays*.

1.1 Algorithm Specialisation

The basic objective of algorithm specialisation is the definition of a (parallel) machine for the efficient execution of an algorithm. Efficiency is obtained by an optimal exploitation of the
structural properties of the algorithm within a number of design constraints. The structural properties of an algorithm are those related, for instance, to its sub-tasks, their generation and data communications. Architectural specialisation is based on such properties. Design constraints are those related, for instance, to processing speed, physical size, accuracy of the results, etc.. Design constraints can be typical of the application area of an algorithm, or imposed by the available physical resources.

A methodological approach to algorithm specialisation considers the definition of an algorithm specific machine as a design process in which methods are provided for the formal description and transformation of an algorithm, and optimisation strategies for the optimal exploitation of its structural properties within a given set of constraints.

We may classify methods for algorithm specialisation into two broad categories. On the one hand, we find general design strategies, including divide and conquer, branch and bound, dynamic programming, search and traversal methods, backtracking, etc.\(^1\) [Kri89]. Given a problem, the designer adopts a number of these strategies until a satisfactory algorithm specific machine is defined. In general, this approach to design requires a high degree of expertise from the designer who needs to have a deep knowledge of the various design strategies in order to apply them optimally for the particular problem. Usually, little automatic support can be provided because of the ad hoc transformations which are involved. These design strategies, however, have the benefit of being generally applicable and likely to produce optimal solutions given the problem requirements.

On the other hand, we find methods which focus on a particular model of parallel computation and try and identify those algorithms which can be efficiently executed within such a model. This viewpoint has led to the development of parallel compilation techniques and systematic synthesis of regular arrays. The emphasis, in this case, is on the syntactic characterisation of algorithms and their systematic manipulation by formal transformations. In this approach a prominent rôle is given to the development of automatic support, so that the transformations are not only systematic, but also largely mechanised. Because of the relevance given to tool support, in the design process a lower level of expertise is expected from the designer. However, there is a loss of generality, as only algorithms which conform to a particular syntax can be treated. Also, sub-optimal solutions are likely to be derived, as optimisation strategies are formulated in general terms for classes of problems instead of being targeted to particular

\(^1\)Most of these strategies were actually developed independently and prior to parallel computing, motivated by the desire of devising optimal algorithms for applications of interest.
applications.

In this thesis we take this second approach and aim at developing design methods for the systematic synthesis of regular arrays.

1.2 Regular Array Synthesis

The type of parallelism we address in this work is that characteristic of regular (processor) arrays, i.e., synchronous regularly connected networks of processors. We will refer to this form of parallelism as regular parallelism. Regular parallelism is synchronous and deterministic in that data are supposed to be transferred through the processor network at regular, consecutive and specified instants of time. Regular parallelism is massive in that the number of processors in the network is assumed to be of the same order of the size parameters of the problem, with each processor performing simple operations, corresponding to the basic computations of the algorithm.

Historically, the development of regular arrays and their synthesis techniques is related to the advent of VLSI design and fabrication techniques, and was initiated, in the late 70s, by Kung and Leiserson [KuLe80] with the introduction of systolic arrays. Regular arrays have developed from systolic arrays by relaxing some of the initial constraints on the topology of the network and the complexity of the processing elements (such as strictly neighbour connections, or bit and word level operations). Also, in time regular arrays have evolved from being seen as particular types of hardware components to being considered as special types of synchronous parallel programs, hence they have been adopted as a model of parallel computation. (A brief survey of the development of regular array synthesis with the relevant references is given in Chapter 2.)

This development of the subject has brought regular array synthesis closer to parallel compilation techniques for Fortran-like programs, and, in particular, the automatic parallelisation of nested for-loops [Wol89]. Besides, together with the traditional exploitation of parallelism for vector processors and shared memory machines, parallel compilation techniques have also evolved to include techniques for distributed memory architectures [OBo93]. The combined effect of these developments have resulted in an even stronger bond between parallel compilation and regular array synthesis [Me-et-al95b].
1.2.1 Algorithm Specification

Synthesis methods for regular arrays are based on the systematic manipulation of algorithm descriptions or specifications. A specification should conform to a formal syntax and corresponds to a functional description of the algorithm. Restricted forms of imperative nested for-loops or recurrence equations are usually admitted as a specification (no standard syntax exists in the literature). For example, an algorithm which computes the first \( n + 1 \) entries of the Fibonacci sequence could be specified as the code segment:

\[
F(0) := 1; \\
F(1) := 1; \\
\text{for } i := 2 \text{ to } n \text{ do} \\
F(i) := F(i - 1) + F(i - 2);
\]

or the recurrence equation:

\[
F(i) = \begin{cases} 
1 & i = 0, 1 \\
F(i - 1) + F(i - 2) & i = 2, \ldots, n
\end{cases}
\]

The specification does not contain any explicit directive for the parallel execution of the computations of the algorithm, because, in principle, the specifier should not be concerned with the model of computation adopted.

Although, it could be argued that the use of imperative code may imply a degree of awareness of some model of computation by the algorithm designer, its traditional use is for syntactic description only. Indeed, the subset of imperative code used corresponds to so-called single assignment code. To a certain extent, the use of single assignment code allows the specifier to adopt a (perhaps more familiar) imperative programming style, while a functional interpretation of the specification is assumed by the method. (The reader interested in the debate imperative vs. functional is referred to the famous article by Backus in [Bac78].)

Our main concern in this thesis is the development of methods for the systematic synthesis of regular arrays, and a purely functional approach to the specification of algorithms will be adopted. We will, however, use nested for-loops in examples and illustrations, so that the reader less accustomed to mathematical notations may gain an intuition of the effects of the specifications.
formal transformations. Work on how to convert imperative nested for-loops into a functional
description exists in the literature. See, for instance, [BuDe88, Lis89].

1.2.2 Parallelism and Data Dependence Relations

While there are no directives for concurrency or data communication, the algorithm specification contains a description of the data dependence relations between the computations of the specified algorithm.

A data dependence relation introduces a sequentiality constraint by expressing that the execution of some computations relies on data generated by other computations. For instance, if we consider the specification of the Fibonacci sequence of the previous section, the evaluation of \( F_i \) at each \( i \) depends on the evaluations of \( F \) both at \( i - 1 \) and \( i - 2 \). Hence, the entries of \( F \) have to be computed sequentially for \( i \) from 2 to \( n \).

Fortunately, strictly sequential computations do not characterise all algorithms. Indeed, the source of the exploitable parallelism of a specification is represented by the sets of its computations which are not related under any data dependence relation. For instance, consider the following segment of code, which computes a vector \( C \) whose entries are the sums of two adjacent entries of a given vector \( A \):

\[
\text{for } i := 2 \text{ to } n \text{ do } \\
\quad C[i] := A[i] + A[i - 1];
\]

The corresponding equation is:

\[
C(i) = A(i) + A(i - 1) \quad i = 2, \ldots, n
\]

Data dependence relations can be established between \( C \) and \( A \). In particular, for all \( i \), the evaluation of \( C \) at \( i \) depends on the values of \( A \) both at \( i \) and \( i - 1 \). On the other hand, there is no data dependence between the computations of \( C \) for different values of the index \( i \). Therefore, the entries of \( C \) may be computed in parallel.

That the ordering of the operations of a program is based on the needs of data (instead of being specified by the programmer) is the basic principle of data flow computing [Ada68, Ada70, DeWe77, Den80]. The analysis of the data dependencies of a program has a prominent rôle in regular array synthesis, and more generally in the exploitation of algorithm parallelism and parallel compilation techniques, and a number of tools have been developed for the representation of data dependence relations (mainly based on graph theory [Car79]). A basic
tool, which we will use extensively in this work, is the so-called data dependence graph, which provides a graphical representation of the data dependence relations of an algorithm. Nodes in a data dependence graph represent computations, with arcs representing their data inter-dependencies.

1.2.3 Algorithm Specialisation and Space-Time Mapping

By decomposing an algorithm into basic computations and their inter-dependencies, the maximal inherent parallelism of the algorithm is uncovered. In particular, if the corresponding data dependence graph defines a partial order, its sets of incomparable nodes correspond to computations that can be executed in parallel, while its arcs indicate the data communications required between computations. In being a graphical representation, the data dependence graph allows for structural and topological considerations. In particular, a processor network whose topology matches the data dependence graph may provide a specialised parallel machine for the execution of the algorithm. The correspondence couples processors of the network with nodes of the data dependence graph (computations) and communication links with arcs of the data dependence graph (data communications).

More complex correspondences can be established between a data dependence graph and a network of processors, for instance by associating several computations to the same processor. In this case, care should be taken that the partial order induced by the data dependencies is reflected by the ordering of execution of the computations in the processor network.

Therefore, in regular array synthesis we can consider algorithmic specialisation as a mapping of the data dependence graph of the algorithm onto a processor network. This mapping is usually known as a space-time mapping, as it can be seen as characterised both by a spatial and a temporal component. The spatial component defines a correspondence between computations and processors. The temporal component indicates the ordering of the computations at each processor.

1.2.4 The Rôle of Regularity

A space-time mapping is straightforward only if data dependence graph and processor network share a similar topology. When the target processor network is a regular array, we are left with the problem of characterising regular data dependence graphs and the type of algorithms to which they correspond. In other words, we need to identify those properties of an algorithm which yield regular data dependence graphs and, hence, regular array designs.
Regular array synthesis has developed from the realisation that particular forms of nested for-loops and recurrence equations correspond to regular data dependence graphs. They are characteristic of so-called uniform problems [Ka-et-al67]. The term uniform refers to characteristics of the data dependencies of the problem. Our description of the Fibonacci sequence is an example of a uniform algorithm. In particular, for all $i$, the computations of $F$ are characterised by the data dependence of $F(i)$ on $F(i - 1)$ and $F(i - 2)$. In terms of data dependence graph, the same arcs, from $i - 1$ to $i$ and from $i - 2$ to $i$, are uniformly replicated at each node of the graph (which is then highly regular). Uniform problems can be easily recognised by considering the form of the index expressions of their variables. If the indices are seen as representing the axes of a Euclidean space, uniform index expressions correspond to translations in that space.

From a designer's point of view, uniform recurrences offer a limited abstraction power for the specification of algorithms. The expression of a generic algorithm as uniform nested for-loops or uniform recurrences requires considerable effort and expertise from the designer, often involving several manipulations of (from a human's point of view) more natural expressions of the algorithm. It also implies a high level of awareness of the underlying model of computation.

In order to increase the abstraction power of the languages for algorithm specification, so-called affine problems have been considered [Mol83, RaFu90]. This type of specifications are characterised by index expressions which are affine expressions of the indices, that is the combination of linear expressions and translations [Ner63, Roc70, Sch86]. An example of an affine specification is the following set of nested for-loops for the computation of the product of two $n \times n$ matrices $A$ and $B$:

```plaintext
for i := 1 to n do
  for j := 1 to n do
    begin
      C(i, j, 0) := 0;
      for k := 1 to n do
        C(i, j, k) := C(i, j, k - 1) + A(i, k) * B(k, j);
    end;
```
The recurrence formulation of the same algorithm is:

\[
C(i, j, k) = \begin{cases} 
0 & i, j = 1, \ldots, n, k = 0 \\
C(i, j, k-1) + A(i, k) \cdot B(k, j) & i, j, k = 1, \ldots, n 
\end{cases}
\]

Note that the single assignment form of the code requires a third index for the accumulation variable \(C\). The entries of the result matrix correspond to the values \(C(i, j, n)\) for all combinations of \(i\) and \(j\). Each computation \(C(i, j, k)\) depends on the computation of the values \(C(i, j, k - 1), A(i, k)\) and \(B(k, j)\), and the relation between each pair of index expressions can be described as an affine mapping.

As uniform index expressions define translations and affine index expressions combine translations and linear transformations, then a uniform data dependence relation is a particular case of affine data dependence relation. Hence, the power of abstraction of affine specifications is higher than that of uniform specifications. The gain in abstraction is, however, counterbalanced by a loss of regularity of the data dependencies.

### 1.2.5 Enforcing Regularity

The increase of abstraction power from uniform to affine can be usefully exploited only if it is accompanied by the provision of systematic transformations of an affine description into a uniform algorithm. These transformations have, indeed, been the object of study by several authors and the problem of making an affine problem uniform is now well-understood [FoMo84, RaFu87, Raj89, WoDe92, QuVa89]. As the effect of these transformations is that of enforcing regularity, we call them *regularising transformations*. (This is not a standard terminology. Other names which have appeared in the literature include *uniformisation* and *localisation*. We feel, however, that regularity carries a more general meaning than the others – in fact, uniform does not necessarily mean local, while locality does not imply uniformity.)

With the introduction of regularising transformations, algorithm specialisation through regular array synthesis can be logically outlined as in the diagram of Fig. 1.1. Initially, a non-regular specification is provided. This is transformed into a regular algorithm (transformation \(\tau_1\)) by the application of regularising transformations, and subsequently mapped onto a regular array (transformation \(\tau_2\)) through a mapping of its regular data dependence graph.

Although in the diagram regularisation has been represented as a mapping from non-regular to regular algorithms, it should be represented more faithfully as a transformational process iterating on non-regular specifications, and which terminates with a regular algorithm. This is
because, in general, regularisation techniques apply to single non-uniform data dependencies, rather than to a specification as a whole. For instance, the matrix product example of the previous section would be transformed by classic regularisation techniques into a uniform algorithm by transforming separately the data dependence relations between $C(i, j, k)$ and $A(i, k)$ and between $C(i, j, k)$ and $B(k, j)$ (the relation between $C(i, j, k)$ and $C(i, j, k - 1)$ is already uniform).

The fact that regularising transformations apply to single data dependence relations has a significant impact on the simplicity of their definition and software implementation. Indeed, regularising transformations have also to be consistent with optimisation strategies which apply to the problem as a whole.

The balance between scope of application, complexity and optimality of the transformations plays an important rôle in the design process. While fully automated optimisation is not supported in regular array synthesis as yet (mainly because an exhaustive search of the solution space of the problem would not be computationally feasible), a possibility is offered to the designer to choose an optimisation strategy and perform some basic transformations of the specification automatically. The development of optimising compilers for regular arrays is, however, an active area of research and likely to become more and more relevant as realistically computational approaches to optimisation are developed.

### 1.3 From Affine to Integral Problems

Although affine data dependencies characterise a large number of problems, they still impose severe restrictions on the specification of algorithms. For instance, the nested for-loops shown below (in which each entry of variable $P$ accumulates the values of $m$ entries of $Q$, as specified by the index expressions, with $Q$ suitably initialised) cannot be treated in a systematic way under current synthesis methods because they do not apply to data dependencies which are not expressed as affine index expressions:

```plaintext
def i := 1 to n do
    begin
```
In the example, the value $P(i, j)$ is data dependent on the value $Q(i * j)$ (as well as $P(i, j - 1)$), where $i * j$ is not an affine expression.

The limitations of the present techniques are mainly due to the choice of the underlying mathematical model for regular array synthesis, that of linear algebra and affine geometry [Ner63, Roc70, Sch86]. Linear and affine transformations as well as convex polyhedral sets are the key concepts which are exploited by synthesis techniques both from a theoretical and an applicative point of view. Hence data dependencies which are defined through arbitrary index expressions fall outside the scope of the established synthesis techniques.

That synthesis techniques are not applicable does not imply that regular parallel solutions are not possible for non-affine problems. It simply means that the techniques are not powerful enough to provide such solutions systematically. Indeed, \textit{ad hoc} regular arrays for non-affine problems have been proposed in the literature (see, e.g., [Eva91]).

One of the main objectives of the thesis is the characterisation of classes of non-affine problems and the provision of regularising techniques for their systematic transformation into uniform specifications. The characterisation that we will provide yields the definition of \textit{integral problems} as problems whose index expressions are general integer mappings of the indices.

1.4 From Static to Dynamic Problems

Both affine and integral problems share the property that their data dependencies are entirely specified by the algorithm description. Hence, a \textit{static} or \textit{compile-time} analysis of those data dependencies is possible, as well as their representation as a data dependence graph. There exist problems, however, which do not share this property. For instance, consider the code segment below (obtained through a small modification of the code segment of the previous section):

\begin{verbatim}
for i := 1 to n do
    read(G(i));
end;
\end{verbatim}
begin
  \( P(i, 0) := 0; \)
  for \( j := 1 \) to \( m \) do
  \[ P(i, j) := P(i, j - 1) + Q(G(i)); \]
end;

In this case, the computation of \( P(i, j) \) depends on the value \( P(i, j - 1) \) as well as on the value of \( Q \) at \( G(i) \). However, this index expression cannot be resolved until the program is executed and a value is provided for \( G(i) \). Moreover, such a value may be different at each execution of the algorithm, as different inputs may be provided. (In this case we also need to ensure that \( G \) is well-defined and the value provided generates a valid index expression for \( Q \).)

A data dependence of this type is called \textit{dynamic} or \textit{run-time} as, in general, its complete representation and analysis is not feasible before the algorithm is executed. The main problem with dynamic data dependencies is that, in general, they do not allow one to derive array designs statically. Therefore, in principle, dynamic problems and regular array synthesis are, at a simple level, incompatible.

However, a second and surprising result of this thesis is to show that for restricted classes of dynamic problems the provision of a regular array solution in a systematic fashion and at compile time is still feasible. The approach we take will require some boundedness assumptions on the specification. For instance, if for the above segment of code we assume that the input values of \( G(i) \) are contained in a finite range, we can define a regular array which will execute the algorithm for all the inputs in that range.

1.5 Outline of the Thesis

This thesis is organised as follows. Chapter 2 describes the design process in regular array synthesis, and introduces some basic definitions and properties. Technical issues related to the regularisation of non-uniform data dependencies are discussed at length as the basis for the development of the following chapters. A brief survey of the most significant contributions to regular array synthesis is given.

Chapter 3 introduces integral problems and their regularisation. The relation between integral and affine problems is explored. The advantages and limitations of the approach are discussed.
The step from static to dynamic problems is formalised in Chapter 4, which also introduces a subclass of dynamic problems for which we provide systematic regularisation techniques.

A number of case studies are presented in Chapter 5 to illustrate the application of our techniques to well-known problems from the literature. Integral and dynamic problems which fall outside the scope of our techniques are also illustrated for comparison, and ad hoc solutions provided.

Chapter 6 outlines some possible developments of the work and draws a number of conclusions.
Chapter 2

Regular Array Synthesis

Regular array synthesis is the process of transforming the functional description of an algorithm into its implementation as a regular array. We call this process the design process of the algorithm. Its basic steps, illustrated in Fig. 2.1, include specification, analysis, regularisation, space-time mapping and implementation.

![Fig. 2.1. Design process.](image)

The specification of the algorithm denotes the first stage of the process, in which a functional description of the algorithm is provided as recurrence equations (or some equivalent, such as single assignment code). The design process terminates with the implementation of the algorithm either as parallel code for a target machine or as specialised hardware. The remaining design steps (contained within a dotted box in the figure) constitute the core of the synthesis method, in which a regular spreading of the computations of the algorithm in space and time is performed, so that the data dependence relations between computations are exposed and

---

1 This is a simplified view of the design process. A more complete description is given in Section 2.4, including classes of transformations which are not considered in this thesis.
the maximal inherent parallelism of the algorithm can be exploited. A spatial distribution is obtained by representing the computations as the nodes of a graph. A temporal distribution follows from the partial order induced by the data dependence relations, and the assumption that a constant time cycle is associated with the execution of each computation. Regularity is enforced through systematic transformations of the specification.

In the figure, the arrows indicate the logical flow in the design process. That all arrows in the synthesis core are bi-directional follows from the fact that several iterations of the design steps may be necessary before the algorithm reaches a form which satisfies optimality, design and correctness requirements.

This chapter contains some preparatory work which is necessary to the development of the remainder of the thesis, and reviews some basic concepts of regular array synthesis which are of particular significance for our approach. The chapter is organised as follows. Section 2.1 addresses the basic definitions for the specification, analysis and space-time mapping of regular parallel algorithms. In the discussion, a generic index space is assumed for the representation of the computations of the algorithm, and the theory is developed at an abstract level. These basic concepts are made more specific in Section 2.2, where the computation space is assumed to be a Euclidean (lattice) space. The advantages of this embedding are highlighted. Proofs of relevant properties can be found in Appendix B. The main issues regarding regularising transformations are discussed in Section 2.3, laying the basis for the development of the following chapters. A brief survey of the major contributions in the literature to regular array synthesis is given in Section 2.4. We summarise the main issues of the chapter in Section 2.5.

2.1 Basic Design Steps

In this section, the basic concepts relative to the specification, analysis and space-time mapping of algorithms are introduced. An index space with no particular structure (apart from being a set) is assumed for representing the computations of an algorithm. We call it computation space, denoted by $\mathcal{CS}$. We also assume the existence of a set $\mathcal{Var}$ of variable names, from which new variables can always be selected, and a set of data values $\mathcal{Val}$, from which variables are assigned and which contains a special undefined value $\bot$. Variables will be indexed by computation points. In particular, if $V$ is a variable and $c \in \mathcal{CS}$, then $V(c)$ will be called an instance of $V$ (at $c$). Also, we will call an index mapping a mapping from $\mathcal{CS}$ to itself.
2.1.1 Algorithm Specification

In our context, an algorithm is specified as a system of recurrence equations, defining the computations and data dependencies of the algorithm. Each recurrence equation expresses the evaluation of a function (the applied function) on a set of computation points (the computation domain). The equations are recurrent in that the evaluation of a function may depend on other evaluations of the same function. Also, the equations constitute a system because, in general, the evaluation of a function depends on the evaluation of other functions of the system (so-called simultaneous recursion).

The result of evaluating a function on each point of a domain as prescribed by a recurrence equation, is recorded by an indexed variable, called the result (variable) of the recurrence equation. The indexing corresponds to the computation point. The definition of the applied function by a recurrence equation is obtained by specifying a number of indexed variables (arguments) to which the function is applied. The indexed variables provide a tabulation of the functions computed by the algorithm. The index expressions of the result and argument variables of a recurrence equation establish the data dependence relations existing between computations.

We distinguish two types of equations: recurrence equations, which define indexed variables in terms of functions applied to a number of argument indexed variables; and input equations, which define variables by the assignment of input values.

**Definition 2.1.1** [Recurrence and Input Equations] A recurrence equation $E$ (with $m$ arguments) is defined as a 5-tuple $(D_E, \cdot E, E^*, f_E, I_M E)$, where: $D_E \subseteq CS$ is the computation domain of the equation; $\cdot E \in Var$ is the result of the equation; $E^* \in Var^m$ is the $m$-tuple of its arguments; $f_E : Val^m \rightarrow Val$ the applied function; $I_M E \in [CS \rightarrow CS]^m$ the $m$-tuple of its index mappings.

An input equation $E$ is defined as a 3-tuple $(D_E, \cdot E, f_E)$, where: $D_E \subseteq CS$ and $\cdot E \in Var$ are as above; and $f_E : CS \rightarrow Val$ is the applied function.

The applied function $f_E$ depends strictly on its arguments (see [Ka-et-al67]) and has constant complexity. The notation $\cdot E$ and $E^*$ (notation reminiscent of the pre- and post-sets in Petri net theory [Br-et-al87]) simply indicates the left- and right-hand sides of the equation $E$. Given a recurrence equation $E$, the order of $E$ is equal to the number of its arguments (i.e.,...
\(|E^*| = m\) in the above definition). The order of an input equation is zero. Given an equation \(E\), we say that \(E\) defines variable \(E\) on each point of the domain \(D_E\).

In the literature a more usual notation for a recurrence equation is (similar to) the following\(^2\):

\[
c \in D : \quad U(c) = f(V_1(I_1(c)), \ldots, V_m(I_m(c))).
\]

where \(D\) is the domain of the equation, \(U\) the result, \(f\) the applied function, and for all \(i, V_i\) are the arguments and \(I_i\) the index mappings. The correspondence between the two notations is straightforward, as illustrated in the following example.

**Example 2.1.2** Consider the specification of the Fibonacci sequence given in Chapter 1. For convenience, we recall it here, both as imperative code:

\[
F(0) := 1;
F(1) := 1;
for \(i := 2\) to \(n\) do
\]

\[
F(i) := F(i - 1) + F(i - 2);
\]

and as the recurrence equation:

\[
F(i) = \begin{cases}
1 & i = 0, 1 \\
F(i - 1) + F(i - 2) & i = 2, \ldots, n
\end{cases}
\]

In our notation, the specification can be expressed as:

\[
E_1 = (D_1, F, in_F)
\]

\[
E_2 = (D_2, F, (F, F), +, (I_1, I_2))
\]

where \(D_1 = \{0, 1\}, D_2 = \{2, \ldots, n\}, in_F(i) = 1, + (a, b) = a + b, I_1(i) = i - 1\) and \(I_2(i) = i - 2\).

In the more traditional notation the specification might be:

\[
i \in D_1 : \quad F(i) = in_F(i)
\]

\[
i \in D_2 : \quad F(i) = +(F(I_1(i)), F(I_2(i)))
\]

\[\Box\ 2.1.2\]

The choice of an algebraic notation as given above is not just a matter of personal taste, but will prove to be convenient for the expression of the definitions and properties necessary in the development of regularisation techniques for dynamic problems (see Chapter 4).

\(^2\text{Many variants of this notation exists. See, for instance, [Ka-et-al67, Rao85, QuVa89, Raj89, QuRo91].}\)
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An algorithm is specified as a system of recurrence and input equations. In such a system, each variable is assumed to be assigned exactly once at each computation point. This is enforced by assuming that equations of the system with the same variable as result have disjoint domains. Note that the same property could be enforced in other ways, for instance by assuming that if the domains of such equations intersect, then the applied functions evaluate to the same value for each point of the intersection. Although, the former condition is more restrictive, it is also easier to enforce and verify, hence it is the one usually adopted.

Definition 2.1.3 [System of Equations] A system of equations $S$ is defined as a set of equations $\{E_1, \ldots, E_r\}$, where for all $E_i, E_j \in S$, with $i \neq j$, for $i, j = 1, \ldots, r$, $E_i = E_j$ implies $D_{E_i} \cap D_{E_j} = \emptyset$.

Given a system of equations $S$, we define its domain as the union of the domains of its equations, i.e., as the set $D_S = \bigcup_{E \in S} D_E$.

Example 2.1.4 [System of Equations] Consider the equations of Example 2.1.2 for the computation of the Fibonacci sequence. Equation $E_1$ is an input equation, while $E_2$ is a recurrence equation. Their domains are disjoint sets, so that $S = \{E_1, E_2\}$ is a system of equations.

The set of variables of an equation $E$ is the set $\text{Var}_E = \{*E\} \cup S(E^*)$ (see Appendix A, for a definition of the support set $S(u)$ of a tuple $u$), for a recurrence equation, and $\text{Var}_E = \{*E\}$, for an input equation. A system of equations $S$ has set of variables $\text{Var}_S = \bigcup_{E \in S} \text{Var}_E$. Moreover, given a variable $V \in \text{Var}_S$, we can identify the set of equations defining $V$, as $\text{Def}_E V = \{E \in S \mid *E = V\}$, as well as the set of computation points on which $V$ is defined, as $\text{Def}_D V = \bigcup_{E \in \text{Def}_E V} D_E$. In the following, we assume that an indexed variable has undefined value outside its definition domain, i.e., $V(c) = \perp$ for all $c \notin \text{Def}_D V$.

2.1.2 Analysis of the Data Dependencies

The process of extracting parallelism from an algorithm specification is based on the analysis of the data dependencies existing between its computations. To this end, several notions of dependence relation have been developed either among variables or their instances [Ka-et-al67, Rao85, SYKun88].

A basic dependence relation is the so-called data dependence relation (or simply data dependence). This relation is defined between variable instances and is based on index mappings.
A recurrence equation implicitly defines a number of data dependencies equal to the order of the equation. Each of them involves the result of the equation and one of the arguments. If \( D \) is the domain of the equation, \( U \) its result, \( V \) any such argument and \( I \) the corresponding index mapping, a data dependence relation between \( U \) and \( V \) is established as follows: for each point \( c \) of the domain \( D \), the computation of \( U(c) \) depends on the value of \( V \) computed at the point \( I(c) \) of the computation space, hence the value \( V(I(c)) \) has to be available for the computation of \( U(c) \) to be possible.

Abstracting away from particular equations, a data dependence can be represented as a 4-tuple \( \mathcal{D} = (D, U, V, I) \), where \( D \subseteq CS, U, V \in Var \) and \( I : CS \rightarrow CS \), expressing that for all \( c \in D \), \( U(c) \) is data dependent on \( V(I(c)) \).

Let \( E \) be a recurrence equation of order \( m \). Then, the \( i^{th} \) data dependence generated by \( E \) is \( \mathcal{D}E_i = (DE, p(E), pr(I, ME)) \) (where \( pr \) is a projection – see Appendix A), and the set of data dependencies of \( E \) is \( \mathcal{D}E = \bigcup_{i=1}^{m} \{\mathcal{D}E_i\} \). As the order of an input equation is 0, its set of data dependencies is empty. Finally, the set of data dependencies of a system of equations \( S \) is \( \mathcal{D}S = \bigcup_{E \in S} \mathcal{D}E \).

**Example 2.1.5 [Data Dependencies]** Consider the system \( S \) of Example 2.1.4. Its data dependencies are:

\[
\begin{align*}
\mathcal{D}D_1 &= (D_2, F, F, I_1) \\
\mathcal{D}D_2 &= (D_2, F, F, I_2).
\end{align*}
\]

Graphs are the usual tools for the representation and analysis of data dependencies. The following three types of graphs are commonly used in regular array synthesis\(^3\). A fourth type of graph, the so-called signal flow graph, which also plays a very important rôle in synthesis, will be introduced in Section 2.1.3 for the description of regular array designs.

**Data Dependence Graph**

A data dependence graph is the natural graph representation of a data dependence. Its nodes are computation points and its arcs relate computation points under the index mapping of the data dependence.

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\(^3\)Many variants of these graphs are known from the literature. According to Feautrier [Feaut92a]: "dependence graphs are used in every form of parallel programming [...]. There are nearly as many dependence graphs as there are workers in the field".
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Fig. 2.2. System S: a) data dependence graph; b) complete data dependence graph; c) reduced dependence graph.

Definition 2.1.6 [Data Dependence Graph] Let $DD = (D, U, V, I)$ be a data dependence. Define its data dependence graph $DDG$ as the graph $(N, A)$, where:

- $N = D \cup I(D)$; and
- $A = \{(I(c), c) | c \in D\}$.

$\blacksquare$ 2.1.6

The data dependence graph of an equation or a system of equations can be defined as the union of the graphs of their data dependencies, with the data dependence graph of an input equation being the empty graph (as an input equation defines no data dependencies).

Example 2.1.7 [Data Dependence Graph] Consider the system S of Example 2.1.4. Its data dependence graph, illustrated in Fig. 2.2 a), is $DDG = (N, A)$, where:

$$\begin{align*}
N &= \{i, i-1, i-2 | i = 2, \ldots, n\} \\
A &= \{(i-1, i), (i-2, i) | i = 2, \ldots, n\}.
\end{align*}$$

$\blacksquare$ 2.1.7

Complete Data Dependence Graph

In a data dependence graph variable names are abstracted away: the nodes represent computation points, and distinct variable instances at the same computation point are not distinguished in the graph. When a distinction between variable instances is important, the following type of graph is used$^4$:

$^4$The notation $(U,c)$, where $U$ is a variable name and $c$ a computation point, is used to distinguish a node of the complete data dependence graph from $U(c)$, which represents the instance of $U$ at $c$. 
Definition 2.1.8 [Complete Data Dependence Graph] Let \( \mathcal{DD} = (D, U, V, I) \) be a data dependence. Define its complete data dependence graph \( \mathcal{CDDG} \) as the graph \( (\mathcal{N}, \mathcal{A}) \), where:

- \( \mathcal{N} = \{(U, c) \mid c \in D\} \cup \{(V, d) \mid d \in I(D)\} \); and
- \( \mathcal{A} = \{(V, I(c)), (U, c) \mid c \in D\} \).

The complete data dependence graph of an equation may be defined as the union of the complete dependence graphs of its data dependencies. Similarly, the complete data dependence graph of a system of equations may be defined as the union of the complete data dependence graphs of its equations.

Example 2.1.9 [Complete Data Dependence Graph] Consider the system \( S \) of Example 2.1.4. Its complete data dependence graph, illustrated in Fig. 2.2 b), is \( \mathcal{CDDG} = (\mathcal{N}, \mathcal{A}) \), where:

\[
\mathcal{N} = \{(F, i), (F, i-1), (F, i-2) \mid i = 2, \ldots, n\}
\]
\[
\mathcal{A} = \{((F, i-1), (F, i)), ((F, i-2), (F, i)) \mid i = 2, \ldots, n\}
\]

Reduced Dependence Graph

Data dependence graph and complete data dependence graph allow a fine-grain analysis of the algorithm at the level of its basic computations. The reduced dependence graph, instead, allows a coarser-grain dependence analysis, in which the inter-dependencies between variables (hence, computed functions) are exposed.

Definition 2.1.10 [Reduced Dependence Graph] Let \( S \) be a system of equations. Define its reduced dependence graph \( \mathcal{RDG} \) as the graph \( (\mathcal{N}, \mathcal{A}) \), where:

- \( \mathcal{N} = \text{Var}_S \); and
- \( \mathcal{A} = \{(U, V) \mid \exists D, I \text{ such that } (D, U, V, I) \in \mathcal{DDS}\} \). 

\begin{itemize}
  \item 2.1.8
  \item 2.1.9
  \item 2.1.10
\end{itemize}
Example 2.1.11 [Reduced Dependence Graph] Consider the system $S$ of Example 2.1.4. Its reduced dependence graph, illustrated in Fig. 2.2 c), is $RDG_S = (\mathcal{N}_S, \mathcal{A}_S)$, where $\mathcal{N}_S = \{F\}$ and $\mathcal{A}_S = \{(F, F)\}$.

2.1.3 Space-Time Mapping

A (processor) array design for a specification is obtained by mapping its data dependence graph onto a graph describing the design. The mapping is realised by identifying a timing function (or scheduling) and an allocation function (or placement) of the computations.

A timing and allocation pair are usually selected on the basis of minimising some parameters, such as the number of execution steps of the algorithm, the number of processing elements of the network, their connections and/or memory requirements, etc. The selection of an optimal space-time mapping is of fundamental importance in the synthesis process, and has been extensively studied (see, e.g., [Da-et-al91, ShF092, DaRo94]). Whilst in this section we only recall some of the basic requirements and properties of a space-time mapping, in the following section we will discuss how optimal timing functions can be systematically derived for particular forms of recurrence equations embedded in Euclidean lattice spaces.

Timing Function

A timing function associates an execution time with each computation point of a system of equations. As we consider discrete schedulings only, we may assume that time coincides with the integers, so that a timing function will be a partial mapping from $CS$ to $\mathbb{Z}$. In particular, a timing function for a system of equations is a mapping from $CS$ to $\mathbb{Z}$ which is defined (at least) at each computation point of the system. A valid timing function for a system of equations is a timing function which preserves the ordering of the computations induced by their data dependencies$^5$:

Definition 2.1.12 [Valid Timing Function for a Data Dependence Graph] Let $DDG = (\mathcal{N}, \mathcal{A})$ be a data dependence graph and $t$ a timing function for $DDG$. Then $t$ is a valid timing function for $DDG$ if and only if for all $(c, c') \in \mathcal{A}$, $t(c) < t(c')$.

$^5$Feautrier, in [Fea92a], uses the name causality to indicate a condition similar to the one we use in the definition of valid timing function.
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Given a timing function \( t \), an infinite family of timing functions can be derived from \( t \) by introducing integral delays\(^6\). In fact, if \( t \) is a valid timing function for a data dependence graph \( \mathcal{D} \mathcal{G} \), then for all \( c \in CS \), the mapping defined by \( t(\theta(c)) = t(c) + 3 \), with \( 3 \in \mathbb{Z} \), is a valid timing function for \( \mathcal{D} \mathcal{G} \). The proof is trivial, and rests on the observation that the ordering defined by a valid timing function among the nodes of a data dependence graph is invariant under the addition of constants. We will use this invariant to restrict ourselves to valid timing functions which are non-negative.

For implementation on finite array architectures, the following two properties of a timing function are required: that only finite sets of computations are scheduled for parallel execution; and that there is a starting time for the execution of the algorithm. For specifications with a finite computation set any valid timing function satisfies these requirements.

**Definition 2.1.13 [Finite and Bounded Timing Function]** Let \( \mathcal{D} \mathcal{G} = (\mathcal{N}, \mathcal{A}) \) be a data dependence graph and \( t \) a valid timing function for \( \mathcal{D} \mathcal{G} \). Then:

- \( t \) is finite if and only if for all \( \tau \in \text{range}_t(\mathcal{N}) \), the set \( \{ c \in \mathcal{N} \mid t(c) = \tau \} \) is finite;
- \( t \) is bounded (below) if and only if there exists \( \bar{\tau} \in \text{range}_t(\mathcal{N}) \) such that for all \( \tau \in \text{range}_t(\mathcal{N}), \bar{\tau} \leq \tau \).

From the previous discussion, it follows that, given a valid bounded timing function \( t \) with lower bound \( \bar{\tau} \), it is always possible to derive a non-negative valid bounded timing function from \( t \), for instance, by adding a delay equal to \( \bar{\tau} \). Without loss of generality, in the following we will restrict ourselves to non-negative timing functions, unless otherwise specified.

The above definitions naturally extend from a data dependence graph \( \mathcal{D} \mathcal{G} \) to, respectively, the data dependence \( \mathcal{D} \), equation \( \mathcal{E} \) or system of equations \( \mathcal{S} \), of which \( \mathcal{D} \mathcal{G} \) is the data dependence graph. Hence, for instance, we may refer to a valid timing function of a system of equations, by which we mean a valid timing function of its data dependence graph.

**Example 2.1.14 [Timing Function]** Consider the data dependence graph in Fig. 2.3 a). A valid timing function is defined by: \( t(i) = 0 \), for \( i = 1, \ldots, 8 \); \( t(i) = 1 \), for \( i = 9, \ldots, 12 \); \( t(i) = 2 \), for \( i = 13, 14 \); and \( t(15) = 3 \).

All timing functions will be considered valid from hereon.

---

\(^6\)Indeed, this is not the only operation which preserves timing functions. It is, however, the only operation we will consider in this work.
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Fig. 2.3. a) Data dependence graph; b) signal flow graph.

Allocation Function

An allocation function determines the distribution of the computations of a system of equations among a set of processing elements.

We assume the existence of an index space, which we call the processor space and denote by $\mathcal{PS}$, as a convenient abstraction in which the placement of computations onto processing elements can be represented (in the same way the space $\mathcal{CS}$ provides a convenient abstraction for the representation of computations). No particular structure is assumed of $\mathcal{PS}$, other than being a set.

An allocation function can be represented as a partial mapping from $\mathcal{CS}$ to $\mathcal{PS}$ (as for timing functions, an allocation function is required to be defined at least on the computation points of a system of equations).

We term a timing and an allocation functions as compatible\(^7\) if and only if computations which are scheduled at the same time are not allocated to the same processing element, i.e., potentially parallel computations are not forced to be executed sequentially. This implies that the maximal parallelism of the specification is exploited, and no sequentiality is enforced which is not prescribed by the partial order defined by the data dependencies of the algorithm\(^8\).

The property of compatibility may be expressed more formally as follows:

**Definition 2.1.15** [Compatibility] Let $\mathcal{DDG} = (\mathcal{N}, \mathcal{A})$ be a data dependence graph and let $t$ be a timing function and $a$ an allocation function for $\mathcal{DDG}$. Then $t$ and $a$ are compatible if and only if for all $c, c' \in \mathcal{N}$, with $c \neq c'$, $t(c) = t(c')$ implies $a(c) \neq a(c')$.

Note that given a valid timing function $t$ for, and the $\mathcal{DDG}$ of, a system, any allocation function which is injective on the nodes of $\mathcal{DDG}$ is trivially compatible with $t$. However, as

---

\(^7\)Another term used in the literature is conflict-free [Fes92a].

\(^8\)Note that with this notion of compatibility, design constraints related to the implementation of the algorithm, such as a limited number of processing elements, are not taken into consideration. See also the discussion on partitioning techniques in Sections 2.2.4 and 2.4.
we will see in the following section, such allocation functions correspond to highly inefficient array designs.

**Example 2.1.16 [Allocation Function]** Consider the data dependence graph in Fig. 2.3 a) with valid timing function \( t \) as given in Example 2.1.14. An allocation function \( a \) compatible with \( t \) is defined as: \( a(i) = p_i \), for \( i = 1, \ldots, 9 \); \( a(10) = a(13) = p_{10} \); and \( a(11) = a(14) = a(15) = p_{11} \); and \( a(12) = p_{12} \), where \( p_i \), for \( i = 1, \ldots, 12 \), denote distinct processors in \( \mathcal{PS} \). In Fig. 2.3 a), dotted lines surround computation points which are allocated to the same processing element under \( a \).

A non-compatible allocation function is, for instance, \( a' \) defined as: \( a'(i) = p_i \), for \( i = 1, \ldots, 8 \); \( a'(9) = a'(13) = p_9 \), \( a'(10) = a'(11) = p_{10} \); and \( a'(12) = a'(14) = a'(15) = p_{11} \). Under \( a' \), computations 10 and 11 would contend the use of processor \( p_{10} \) at time \( t = 2 \).

A trivially compatible allocation function is \( a'' \) defined as \( a''(i) = p_i \), for \( i = 1, \ldots, 15 \), with \( p_i \), distinct processors in \( \mathcal{PS} \), for \( i = 1, \ldots, 15 \).

In the above presentation, we have assumed that a timing function is selected first, followed by the choice of a compatible allocation function. In this approach, the emphasis is on the optimality of the scheduling as a timing function which guarantees the minimum number of computation steps. Alternatively, an allocation function may be selected first [ClMo93]. In this case, the optimality of a placement can be expressed in terms of a number of design parameters, such as the locality of the connections of the network or minimising the memory requirements for its processing elements.

**Signal Flow Graph**

A (valid) timing function for a system of equations together with a compatible allocation function determine an array design for a system of equations. Such a design is represented by a labelled graph known as a signal flow graph\(^9\). Its nodes represent processing elements, its edges communication links and their labels communication delays.

**Definition 2.1.17 [Signal Flow Graph under \( t \) and \( a \)]** Let \( S \) be a system of equations and \( \mathcal{DDG} = (N, A) \) its data dependence graph. Let \( t \) be a valid timing function for \( S \) and \( a \) a

\(^9\)Signal flow graphs are well-known design tools for digital signal processing applications. The type of signal flow graph addressed in this work is only a particular type of signal flow graph, and more general definitions may be found, for instance, in [CrRa83, SYKun88].
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compatible allocation function. Define the signal flow graph $G^{t,a}$ of $S$ under $t$ and $a$, as the
labelled graph $(N^{t,a}, A^{t,a}, \ell)$ such that:

- $N^{t,a} = \{a(c) | c \in N\};$
- $A^{t,a} = \{(a(c), a(c')) | (c, c') \in A\};$ and
- $\ell : A^{t,a} \rightarrow \mathbb{N}$ such that $\forall (c, c') \in A, \ell((a(c), a(c'))) = t(c') - t(c).$

\[2.1.17\]

Given a timing function for, and the data dependence graph of, a system of equations, an
allocation function which is trivially compatible (i.e., an allocation which is just an injective
mapping on the nodes of the graph – see Section 2.1.3) generates a highly inefficient array
design whose processing elements are active exactly once.

Example 2.1.18 [Signal Flow Graph] Consider the data dependence graph in Fig. 2.3 a),
the timing function $t$, given in Example 2.1.14, and the compatible allocation function $a$, given in Example 2.1.16. The corresponding signal flow graph is illustrated in Fig. 2.3 b).
All the arcs have a unit label, which we have omitted in the figure. In this (and in the
general) case, $a$ associates more than one computation to some of the nodes of the signal flow
graph, and the correct order of execution of such computations is determined by the timing
function $t$.

\[2.1.18\]

Note that a signal flow graph provides an abstraction for the algorithm implementation. Such
an implementation can be realised in a number of ways, including custom VLSI circuits whose
topology matches that of the signal flow graph, or parallel code to be executed on a general
purpose parallel machine.

Timing Function and Cyclic Data Dependence Graph

Definition 2.1.12 precludes the definition of a valid timing function for a data dependence
graph containing cycles (given a data dependence graph $\mathcal{DG}$ and a path $n_1, \ldots, n_p$ of $\mathcal{DG}$,
where $n_1 = n_p = n$, then for a function $t$ to be a valid timing function of $\mathcal{DG}$ the definition
implies that $t(n_1) < t(n_p)$, i.e., $t(n) < t(n)$. As $t$ is a function to the integers, this can never
be the case). As a consequence only acyclic data dependence graphs admit valid timing
functions.
Requiring that the data dependence graph be acyclic is too strong a restriction: there are many examples of recurrences with cyclic data dependence graphs which are (intuitively) computable and for which we would like to define a timing function. For instance:

Example 2.1.19 The system $S = \{E_1, \ldots, E_5\}$ of equations below computes the sum $a + b$, where $a$ and $b$ are input values (at points 1 and $n$, respectively). The sum is computed at each point of the sub-domain $D_2$ of $S$ (we have intentionally defined $S$ so that its data dependence graph is cyclic, although simpler definitions no doubt exist). The equations are:

\[
\begin{align*}
E_1 &= (D_1, A, in_a) \\
E_2 &= (D_2, A, A, id, I_1) \\
E_3 &= (D_3, B, in_b) \\
E_4 &= (D_2, B, B, id, I_2) \\
E_5 &= (D_2, F, (A, B), +, (I, I)),
\end{align*}
\]

with: domains $D_1 = \{1\}$, $D_2 = \{2, \ldots, n - 1\}$, $D_3 = \{n\}$; index mappings $I(i) = i$, $I_1(i) = i - 1$, $I_2(i) = i + 1$; computed functions $id(a) = a$, $in_a(i) = a$, $in_b(i) = b$, with $a, b \in \mathbb{R}$, and $+(a, b) = a + b$. The addition $a + b$ at each point of $D_2$ is realised by the computation of variable $F$. Variables $A$ and $B$ transfer, respectively, $a$ and $b$ among the computation points. The data dependence graph and complete data dependence graph of $S$ are illustrated in Fig. 2.4 a) and b), respectively. We note that the complete data dependence graph is acyclic. 

The reason why cycles may appear in a data dependence graph even when the complete data dependence graph is acyclic is that a data dependence graph abstracts away from variable names, i.e., does not distinguish between variables, so that variable instances with different names at the same computation point are collapsed onto the same node of the graph. Such cycles can be eliminated if the specification is modified, for instance, to redistribute its
computations in the space, so that a distinct computation point is associated with each node of the acyclic complete data dependence graph.

In the following, without loss of generality and unless otherwise indicated, we will always assume that data dependence graphs are acyclic.

### 2.2 Euclidean Synthesis

Although in the previous discussion no assumption was made of the nature and properties of the computation space $CS$ and processor space $PS$, regular array synthesis techniques have developed mainly for computation and processor spaces represented by multi-dimensional Euclidean spaces. This geometric embedding of regular array synthesis has the following important advantages. From a theoretical point of view, linear algebra and affine geometry provide for rich mathematical models. From an applicative point of view, their constructive and algorithmic properties may be exploited for the development of automatic support.

The central idea of Euclidean regular array synthesis is the embedding of the data dependence graph of a specification into convex regions of the space. Of such convex regions only lattice points are considered. Therefore, the actual representation of the graph is in a lattice space. However, properties of the embedding linear and affine spaces can be exploited, such as polyhedral convexity, which allows for a finite representation of the computation domains, and linear and affine transformations, which, by preserving polyhedral convexity, allow for a finite manipulation of the domains.

The choice of linear and affine (lattice) spaces for the representation of algorithm specifications is not arbitrary, but comes from the realisation that multi-dimensional lattice spaces offer a natural interpretation of the index spaces defined by sets of nested for-loops, which, in turn, are widely used in the sequential coding of large classes of numerical algorithms [Lam74]. Moreover, for those algorithms, loop bounds are often given as affine expressions of the indices, which can be interpreted as the definition of the boundaries of polyhedral convex sets in the embedding Euclidean space.

Finally, another major benefit of this representation is the possibility of defining systematic techniques for generating optimal space-time mappings, by relating the problem specification to the formulation of (integer) linear programming problems, for which computationally effective solution methods are well-developed [Sch86].

In the remainder of this section we will discuss the basic concepts of Euclidean synthesis
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and the specialisation of regular array synthesis concepts arising from this embedding strategy. From now on, computation and processor spaces are assumed to be multi-dimensional Euclidean spaces and computation domains to be (convex) polyhedra.

2.2.1 Representation and Mechanisation

For a specification to be formally represented and manipulated, it has to be expressible as a term according to some syntactic rules. Euclidean synthesis techniques account for this requirement. In particular computation domains, index mappings, and timing and allocation functions admit term representations, such as linear and affine expressions.

As computation domains are assumed to be polyhedral convex sets, by definition, they admit a finite representation either as the intersections of a finite set of half-spaces or as the convex closure of a finite number of points and directions (see also Appendix E).

Index mappings, in their most general form, are assumed to be affine transformations. Hence they can be represented as a combination of a linear transformation (as a matrix-vector product) and a translation vector. A similar form is provided for timing and allocation functions, which are assumed to be affine and linear transformations, respectively. This form and related properties are discussed in the remainder of this section.

2.2.2 Normalisation of Index Expressions

In Euclidean synthesis the computation domain is assumed to be a multi-dimensional lattice space. Let $\mathbb{Z}^n$ denote such a space for some $n \in \mathbb{N}$. Then an index mapping in $\mathbb{Z}^n$ is a mapping $I$ from $\mathbb{Z}^n$ to $\mathbb{Z}^n$.

In general, however, the initial specification of an algorithm may contain a more general form of index mapping as a mapping $I : \mathbb{Z}^n \rightarrow \mathbb{Z}^l$, where $l \leq n$. Such mappings need to be normalised [QuVa89] before synthesis techniques can be applied. For example, let us consider the specification of the matrix product of Chapter 1 (recalled here, for convenience):

\begin{verbatim}
for i := 1 to n do
for j := 1 to n do
begin
    C(i, j, 0) := 0;
    for k := 1 to n do
\end{verbatim}
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\[ C(i, j, k) := C(i, j, k - 1) + A(i, k) \cdot B(k, j); \]

end;

Its computation space can be assumed to be a 3-dimensional space in which each value \( C(i, j, k) \) is computed for all \( i, j, k \) from 1 to \( n \). Note that the number of dimensions of the space corresponds to the number of nested for-loops. Variables \( A \) and \( B \), however, are indexed by two indices only. Hence, they need to be normalised in \( \mathbb{Z}^3 \).

Normalisation usually consists of "padding" the index expressions of variables which are not fully indexed, with extra indices up to the number \( n \) of dimensions of the space. Geometrically, this corresponds to positioning the computations of such variables in particular sub-spaces of the \( n \)-dimensional space. In principle, any arbitrary choice of the extra indices can be made. However, because of the relation between index expressions and data dependencies, this choice has an impact on the regularity of the specification. When the variables to be fully indexed correspond to input or output data of the algorithm (as in our example), it is common practice to position their evaluation on some boundary of the computation domains. This, in general, translates into array designs where input and output operations are confined to border processing elements.

In our examples, we could pad the index expressions of \( A \) and \( B \) with extra null entries and obtain:

\[
\text{for } i := 1 \text{ to } n \text{ do }
\text{for } j := 1 \text{ to } n \text{ do }
\begin{align*}
& \text{begin} \\
& \quad C(i, j, 0) := 0; \\
& \quad \text{for } k := 1 \text{ to } n \text{ do} \\
& \quad \quad C(i, j, k) := C(i, j, k - 1) + A(i, k, 0) \cdot B(k, j, 0); \\
& \text{end;}
\end{align*}
\]

If we interpret the overall computation domain as the hypercube \( D = \{ (i, j, k) \mid 1 \leq i, j \leq n, 0 \leq k \leq n \} \) (which corresponds to the evaluation of all index expressions in the nested for-loops, for \( i, j \) and \( k \) ranging between their bounds), then the evaluation of \( A \) and \( B \) occurs on the boundary \( D' = \{ (i, j, k) \in D \mid k = 0 \} \).

This type of normalisation is generally applicable [QuVa89] and, in the following, we always
assume that all variables are normalised.

2.2.3 Uniform and Affine Data Dependencies

An affine index mapping is an index mapping in $\mathbb{Z}^n$ which defines an affine transformation. An affine index mapping has the following form:

**Definition 2.2.1 [Affine Index Mapping]** Let $\mathcal{I}$ be an index mapping. $\mathcal{I}$ is affine if for all $z \in \mathbb{Z}^n$, $\mathcal{I}(z) = A \cdot z + b$, with $A \in \mathbb{Z}^{n \times n}$ and $b \in \mathbb{Z}^n$. In addition, $\mathcal{I}$ is uniform if $A = I_n$.

Together with index mappings, so-called dependence mappings are commonly used in the analysis of the data dependencies. Dependence mappings are specific to Euclidean synthesis in that their definition relies upon the existence of arithmetic operations in the computation space. A dependence mapping is derived from an index mapping as follows:

**Definition 2.2.2 [Affine Dependence Mapping]** Let $\mathcal{I}$ be an affine index mapping. Its dependence mapping $\Theta_{\mathcal{I}}$ is the mapping defined as $\Theta_{\mathcal{I}}(z) = z - I(z)$, for all $z \in \mathbb{Z}^n$.

For all $z$, $\Theta_{\mathcal{I}}(z)$ is a vector, called the *data dependence vector* determined by $\mathcal{I}$ at $z$. Note that as $\mathcal{I}$ is affine, $\Theta_{\mathcal{I}}$ is also affine. Affine index mappings are used in the definition of affine data dependencies:

**Definition 2.2.3 [Affine Data Dependence]** Let $\mathcal{D} = (D, U, V, \mathcal{I})$ be a data dependence. $\mathcal{D}$ is affine if $\mathcal{I}$ is an affine index mapping.

Given an affine data dependence $\mathcal{D} = (D, U, V, \mathcal{I})$, with $\Theta_{\mathcal{I}}$ the dependence mapping defined by $\mathcal{I}$, the image of $D$ under $\Theta_{\mathcal{I}}$ is called the *dependence domain* of $\mathcal{D}$:

**Definition 2.2.4 [Dependence Domain]** Let $\mathcal{D} = (D, U, V, \mathcal{I})$ be an affine data dependence and $\Theta_{\mathcal{I}}$ the dependence mapping defined by $\mathcal{I}$. The dependence domain $\Omega_{\mathcal{I}}$ of $\mathcal{D}$ is $\Omega_{\mathcal{I}} = \Theta_{\mathcal{I}}(D)$.

As $D$ is a convex polyhedron and $\Theta_{\mathcal{I}}$ is affine, $\Omega_{\mathcal{I}}$ is also a convex polyhedron (see Appendix E).
An important sub-class of affine data dependencies is represented by uniform data dependencies. A uniform data dependence is characterised by a constant vector replicated for each point of the domain. Because of their regularity, uniform data dependencies generate regular array designs under a space-time mapping. Uniformity depends both on the index mapping and the domain of the data dependence:

**Definition 2.2.5 [Uniform Data Dependence]** Let $\mathcal{D} = (D, U, V, I)$ be an affine data dependence. $\mathcal{D}$ is uniform if and only if there exists an integer vector $c \in \mathbb{Z}^n$ such that, for all $z \in D$, $\Theta_I(z) = c$. \hfill 2.2.5

From this definition it follows that the dependence domain of a uniform data dependence reduces to a singleton set.

**Example 2.2.6 [Uniform and Non-Uniform Data Dependencies]** Consider the index mapping

$$I \begin{pmatrix} i \\ j \\ k \end{pmatrix} = \begin{pmatrix} k \\ j - 1 \\ k \end{pmatrix} = \begin{bmatrix} 0 & 0 & 1 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix} \begin{pmatrix} i \\ j \\ k \end{pmatrix} + \begin{pmatrix} 0 \\ -1 \\ 0 \end{pmatrix},$$

in $\mathbb{Z}^3$, and its dependence mapping $\Theta_I(i, j, k) = (i - k, 1, 0)$. Let $\mathcal{D} = (D, U, V, I)$ and $\mathcal{D}' = (D', U, V, I)$ be data dependencies with domains, respectively:

$$D = \{(i, j, k) | 1 \leq i, j \leq m, 1 \leq k \leq i\}$$

$$D' = \{(i, j, k) | 1 \leq i, j \leq m, i = k\},$$

where $m$ is a constant in $\mathbb{N}$. Domains $D$ and $D'$ are illustrated in Fig. 2.5 a) and b), respectively. A section (for $k = 1$) of the data dependence graph relative to $\mathcal{D}$ is given in Fig. 2.5 c), while the data dependence graph of $\mathcal{D}'$ is illustrated in part d) of the figure. Note that $\mathcal{D}$ is not uniform. For instance, by assuming $m \geq 3$ and considering the points $(2, 2, 1)$ and $(3, 2, 1)$ of $D$, then $\Theta(2, 2, 1) = (1, 1, 0) \neq \Theta(3, 2, 1) = (2, 1, 0)$. However, $\mathcal{D}'$ is uniform as for all $(i, j, k) \in D'$, $\Theta_I(i, j, k) = (0, 1, 0)$. \hfill 2.2.6

It can be proved (see Appendix B) that given an affine data dependence $\mathcal{D} = (D, U, V, I)$, with $I(z) = A \cdot z + b$, for all $z \in D$, $\mathcal{D}$ is uniform if and only if $\text{lin}(D) \subseteq \text{null}(I_n - A)$ (this is trivially true if $I$ is a uniform index mapping, as $A = I_n$). Moreover, if $\mathcal{D}$ is uniform then $I$ can always be replaced by a uniform index mapping.
Example 2.2.7 In Example 2.2.6, \( \text{lin}(D) = \mathbb{Z}^3 \), \( \text{lin}(D') = \{(1,0,1),(0,1,0)\} \), \( A = \begin{bmatrix} 0 & 0 & 1 \\ 0 & 1 & 0 \\ 0 & 0 & 1 \end{bmatrix} \), and \( \text{null}(I_3 - A) = \{(1,0,1),(0,1,0)\} \). As \( \text{lin}(D) \not\subseteq \text{null}(I_3 - A) \), \( DD \) is not uniform. As \( \text{lin}(D') \subseteq \text{null}(I_3 - A) \) (in particular they are equal), \( DD' \) is uniform. Note that, in \( DD' \), \( I \) can be replaced by \( I'(i,j,k) = (i,j,k) + (0,-1,0) \).  

\[ 2.2.7 \]

2.2.4 Affine Space-Time Mapping

An affine timing function is a timing function which is also an affine transformation:

Definition 2.2.8 [Affine Timing Function] Let \( t : \mathbb{Z}^n \to \mathbb{Z} \) be a timing function. \( t \) is affine if and only if, for all \( z \in \mathbb{Z}^n \), \( t(z) = \lambda \cdot z + \mu \), where \( \lambda \) is a non-null vector in \( \mathbb{Z}^n \) and \( \mu \in \mathbb{Z} \).

\[ 2.2.8 \]

The vector \( \lambda \) may be regarded as the normal vector to a family of parallel hyperplanes which it defines. There exists one such hyperplane for each (instant) \( i \in \mathbb{Z} \), called an *isochronous* or *equitemporal* hyperplane. Intuitively, all the points on such a hyperplane are assigned by \( t \) the same instant of time, and so should be computed on different processors.

A linear allocation function is an allocation function which is defined as a linear transformation from the computation space to the processor space. The processor space is usually assumed to have dimensionality no greater than the computation space, as, in general, in order to increase the efficiency of the array, several (sequential) computations are allocated
Fig. 2.6. a) Compatible and b) non-compatible affine scheduling and linear allocation.

to the same processor. Hence, in general, the image of an allocation function has a lower dimensionality than its application domain. Computationally, a mapping from $\mathbb{Z}^n$ onto a space of lower dimension can be conveniently realised as a linear projection.

**Definition 2.2.9 [Linear Allocation Function]** Let $a : \mathbb{Z}^n \rightarrow \mathbb{Z}^l$ be an allocation function. $a$ is linear if and only if, for all $z \in \mathbb{Z}^n$, $a(z) = \sigma \cdot z$, where $\sigma$ is a non-null matrix in $\mathbb{Z}^{l \times n}$ and $l \leq n$. 

The simplest type of linear allocation functions are projections from $\mathbb{Z}^n$ to $\mathbb{Z}^{n-1}$. These projections can be expressed by corresponding projection vectors:

**Definition 2.2.10 [Projection Vector]** Let $a : \mathbb{Z}^n \rightarrow \mathbb{Z}^{n-1}$ be a linear allocation function, with $a(z) = \sigma \cdot z$ and $\sigma \in \mathbb{Z}^{(n-1) \times n}$, for all $z \in \mathbb{Z}^n$. A projection vector for $a$ is a non-null vector $u \in \text{null}(\sigma)$. 

For affine timing and allocation functions $t$ and $a$, respectively, their compatibility can be verified as follows. If $t(z) = \lambda \cdot z + \mu$, and $a$ has projection vector $u$, then $t$ and $a$ are compatible if and only if $\lambda \cdot u \neq 0$ (see Appendix B). That the dot product of $\lambda$ and $u$ is not zero, implies that points on an equitemporal hyperplane defined by $\lambda$ are not projected onto the same processor, hence their computations can proceed in parallel. An intuitive picture of this property is given in Fig. 2.6 for $\lambda$ and $u$ in $\mathbb{Z}^2$.

An affine timing function $t$, defined as $t(z) = \lambda \cdot z + \mu$, together with a compatible linear allocation function $a$, defined as $a(z) = \sigma \cdot z$, can be regarded as a space-time mapping from
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$Z^n$ to itself. This mapping is usually represented in its matrix form as:

$$[t, a](z) = \begin{bmatrix} \lambda \\ \sigma \end{bmatrix}.$$  

An affine space-time mapping $[t, a]$ defines a signal flow graph according to Definition 2.1.17. If $S$ is a system of equations with dependence domain $\Omega$, the arcs of the signal flow graph under $[t, a]$ are the images of the vectors in $\Omega$. In particular, each data dependence vector $d$ in $\Omega$ corresponds to the arc $\sigma \cdot d$ of the signal flow graph, with label $\lambda \cdot d$.

2.2.5 Linear Optimisation and Affine Scheduling

One of the main advantages of Euclidean synthesis is the possibility of defining semi-automatic techniques for the scheduling of specifications containing affine data dependencies. Such techniques are based on the formulation of an integer linear programming problem, whose solution is an optimal affine timing function for the specification. These techniques were first introduced by Quinton in [Qui84] for uniform problems, and subsequently extended to affine problems in [RaFu87, RaFu89, QuVa89].

If $DD$ is an affine data dependence, the optimisation problem which defines an optimal affine timing function for $DD$ is defined both on the computation domain and the dependence domain of $DD$. The fundamental result is expressed by the following theorem, which is based on the formulation given in [QuVa89]:

**Theorem 2.2.11** Let $DD = (D, U, V, I)$ be an affine data dependence, with $\Omega_T$ its dependence domain. Let $t$ be an affine timing function, such that for all $z \in Z^n$, $t(z) = \lambda \cdot z + \mu$, where $\lambda \in Z^n$ and $\mu \in Z$. Then $t$ is a non-negative, finite, bounded valid timing function for $DD$ if and only if:

i) for all $v \in \text{vert}(\Omega_T)$, $\lambda \cdot v > 0$;

ii) for all $r \in \text{ray}(\Omega_T)$, $\lambda \cdot r \geq 0$;

iii) for all $v \in \text{vert}(D)$, $\lambda \cdot v + \mu \geq 0$;

iv) for all $r \in \text{ray}(D)$, $\lambda \cdot r > 0$.

\(^{10}\)More general types of affine space-time mappings have been considered in the literature for partitioning techniques (see the discussion in Section 2.4). In such cases, both scheduling and allocation functions are seen as multi-dimensional mappings, with the dimensions of their ranges summing up to the dimension of the computation space. As we do not address partitioning in this work, and for ease of presentation, we do not consider this type of mapping formally.
A proof of Theorem 2.2.11 is given in Appendix B. Condition ii) of the theorem implies that if $\Omega_T$ contains a line with direction $l \in \mathbb{Z}^n \ (l \neq 0)$, then $^\lambda \cdot l = 0$. The following correspondence exists between the conditions of Theorem 2.2.11 and the properties of $t$:

- Conditions i) and ii) correspond to the validity of $t$. As polyhedral convex sets are finitely generated, i) and ii) imply that for all $z \in D$, $\lambda \cdot \Theta_T(z) > 0$, which in turn, implies that for all $z \in D$, $t(I(z)) < t(z)$. Hence, $t(n) < t(n')$ for all pairs of nodes $(n, n')$ of the data dependence graph of $DD$. As a corollary, the validity of $t$ can be expressed by the condition $\lambda \cdot \Theta(z) > 0$, for all $z \in D$. Also, if there exists a valid affine timing function, then $\Omega_T$ does not contain two non-null vectors $d, d'$ such that $d' = -cd$, with $c > 0$.

- Conditions iii) and iv) correspond to the non-negativity of $t$ over $D$. This comes from the property of finite generation of polyhedral sets applied to the domain $D$; and

- Condition iv) corresponds to the finiteness and boundedness of $t$. In particular, $t$ is finite if and only if $\lambda \cdot r \neq 0$, for all $r \in ray(D)$ (see Appendix B).

**Example 2.2.12** Consider the affine data dependence $DD = (D, U, V, I)$, with $D = \{(i, j) \mid 1 \leq j \leq n, i \in \mathbb{Z}\}$, where $n$ is a constant in $\mathbb{N}$, and $I(i, j) = (2i, j - 1)$. The corresponding data dependence graph is depicted in Fig. 2.7 a). Note that $D$ is finitely generated, for instance, by the points $(0, 1)$ and $(0, n)$ and directions $(1, 0)$ and $(-1, 0)$. The dependence mapping is $\Theta_T(i, j) = (-i, 1)$ so that $\Omega_T$ is finitely generated by the point $(0, 1)$ and the directions $(1, 0)$ and $(-1, 0)$ (see Fig. 2.7 b)).

A valid affine timing function for $DD$ is that given by $\lambda = (0, 1)$, which satisfies conditions i) and ii) of Theorem 2.2.11. Note, however, that any affine timing function defined by $\lambda$ is not finite, as $\lambda$ does not satisfy the inequality $\lambda \cdot r \neq 0$ for all $r \in ray(D)$. Indeed, for this data dependence there is no valid and finite affine timing function as the constraints for the two properties are in contradiction (as the infinite directions of $D$ and $\Omega_T$ are the same). However, if the domain $D$ were bounded, a valid finite timing function might exist. Consider the data dependence $DD' = (D', U, V, I)$, with $D' = \{(i, j) \mid 1 \leq i \leq m, 1 \leq j \leq n\}$, where $n$ and $m$ are constant in $\mathbb{N}$. The corresponding data dependence graph is given in Fig. 2.7 c). Note

---

$^\lambda$If $l$ is the direction of a line of $\Omega_T$, then there exist two rays in $\Omega_T$ with direction $l$ and $-l$, respectively. From condition ii), it follows that $\lambda \cdot l = 0$. 


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Fig. 2.7. Data dependence $DD$: a) data dependence graph; b) dependence domain. Data dependence $DD'$: c) data dependence graph; d) dependence domain.

that $D'$ is finitely generated by its vertices $(-m, 1), (-m, n), (m, 1)$ and $(m, n)$, and $\Omega'_T$, by the vertices $(-m, 1)$ and $(m, 1)$ (see Fig. 2.7 d)). Then $\lambda = (0, 1)$ satisfies all the conditions of Theorem 2.2.11.

For a uniform data dependence, the dependence domain reduces to a singleton set and conditions i) and ii) of Theorem 2.2.11 reduce to the simpler condition i) of the following corollary (corresponding to a result in [Qui84]):

**Corollary 2.2.13** Let $DD = (D, U, V, I)$ be a uniform data dependence, with index mapping $I(z) = z + b$, for all $z \in \mathbb{Z}^n$, and $b \in \mathbb{Z}^n$. Let $t$ be an affine timing function, such that for all $z \in \mathbb{Z}^n$, $t(z) = \lambda \cdot z + \mu$, with $\lambda \in \mathbb{Z}^n$ and $\mu \in \mathbb{Z}$. Then $t$ is a non-negative, finite, bounded valid timing function for $DD$ if and only if:

i) $\lambda \cdot b < 0$;

ii) for all $v \in \text{vert}(D)$: $\lambda \cdot v + \mu \geq 0$;

iii) for all $r \in \text{ray}(D)$: $\lambda \cdot r > 0$. 

2.2.12
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Note that the above results refer to a single data dependence. In general, as algorithm specifications are systems of recurrence equations, several data dependencies will have to be considered at the same time. The results extend naturally to a system of equations by considering as domain and dependence domain of the system, the smallest polyhedral convex sets containing the union of, respectively, the domains and dependence domains of its equations.

2.2.6 Affine Scheduling and Dependence Cone

Given a data dependence, its dependence cone is defined as the smallest polyhedral convex cone containing its dependence domain:

Definition 2.2.14 [Dependence Cone] Let \( DD = (D, U, V, I) \) be an affine data dependence with dependence domain \( \Omega_I \). The dependence cone \( \Theta_I^\ast \) of \( DD \) is \( \Theta_I^\ast = cone(\Omega_I) \).

In the previous section we have established necessary and sufficient conditions to the existence of a valid affine timing function based on the generators of the dependence domain of an affine data dependence. Here we discuss a sufficient condition for the existence of a valid affine timing function which is based on its dependence cone.

The existence of a valid affine timing function may be related to the pointedness of \( \Theta_I^\ast \). If \( \Theta_I^\ast \) is pointed (see Appendix E), then there exists a hyperplane with a non-null, integer normal vector \( \lambda \) such that: \( \Theta_I^\ast \) is entirely contained in one of the half-spaces defined by the hyperplane; and \( \lambda \cdot z > 0 \) for all \( z \) in \( \Theta_I^\ast \). As, by definition, \( \Omega_I \) is contained in \( \Theta_I^\ast \), then, in particular, \( \lambda \cdot z > 0 \) for all \( z \) in \( \Omega_I \). Therefore, \( \lambda \) defines a (family of) valid affine timing function(s) for the data dependence \( DD \). The formal result is stated in the following proposition (a proof of which may be found in Appendix B).

Proposition 2.2.15 Let \( DD = (D, U, V, I) \) be a data dependence and \( \Theta_I^\ast \) its dependence cone. If \( \Theta_I^\ast \) is pointed then there exists a valid affine timing function for \( DD \).

Note that pointedness of \( \Theta_I^\ast \) gives a sufficient condition for the existence of a valid timing function. That it is not necessary can be seen by considering, for instance, the data dependence of Example 2.2.12, which admits a valid (but not finite) affine timing function even
though its dependence cone is not pointed (such a cone is $\Omega_i^* = \text{cone}\{(-1,0),(0,1),(1,0)\}$, corresponding to the dependence domain in Fig. 2.7 b)).

As we will see, the main advantage of this condition is that it can be exploited even in those cases when the dependence domain is not a convex polyhedron. It is this weaker condition which will be used in the remainder of this work.

2.3 Regularisation

One of the requirements of regular array design is that the processing elements are locally and uniformly connected. This was originally motivated by technological consideration. If the design is implemented as a VLSI circuit, long wiring is too expensive and introduces undesirable delays in signal propagation. On the other hand, a regular communication topology guarantees simple and cost-effective layouts. Although these requirements can be partially relaxed if the regular array is implemented in software, regularity remains one the basic feature of regular arrays as a model of computation.

Because of the relationships between data dependence structure of the specification and communication structure of the design, notions of uniformity and locality can be formulated on data dependencies. We have already formally defined what we intend by a uniform data dependence, and we can think of a local data dependence as a data dependence under which only (nearest) neighbour computation points are related. The lack of regularity of a data dependence has to be addressed before the mapping onto a regular array design is feasible. We use the term regularisation to refer to the set of techniques which can be used to improve the uniformity and locality of data dependencies.

As discussed in Chapter 1, regular array synthesis is based on the specification of algorithms as systems of equations whose data dependencies conform to a particular syntax, and, historically, recurrences with uniform data dependencies were the first class of recurrences to be considered, for their natural correspondence with regular arrays. In fact, any regular array can be expressed as a system of uniform recurrences (as proved by Rao in [Rao85]) and uniform recurrences always map to regular array designs under linear space-time mappings. Also, we have already mentioned that in order to facilitate the specification of algorithms, more general classes of recurrences have been introduced together with regularising transformations for their manipulation into systems of uniform recurrences. These new recurrences are those characterised by affine data dependencies, and constitute the most general type of
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data dependencies which is considered in regular array synthesis at present.

The main goal of this thesis is the extension of synthesis techniques to more general classes of problems. This will be formally addressed in the following chapters. In this section, however, we intend to discuss in general terms the basic issues and properties which are involved in the regularisation of non-uniform data dependencies. We will capitalise on these properties in the development of the regularisation techniques of the following chapters.

2.3.1 Decomposition and Uniformisation

Regularisation involves replacing a non-uniform data dependence with a set of data dependencies which exhibit improved uniformity and locality. Although regularisation mainly refers to transformations of the data dependence graph, it can be described in terms of pipelining and routing of the data flow between computation points. This terminology can be justified if we think of a data dependence graph as a signal flow graph (this is always possible - consider the trivial allocation function in Section 2.1.3). Also, regularisation has the ultimate goal of producing specifications for which a regular array design is guaranteed under a space-time mapping. Hence, ultimately the application of regularisation techniques is reflected in the improved uniformity and locality of the connection topology of the design. Early forms of regularisation were actually defined directly on the signal flow graph rather than the data dependence graph (see, e.g., the work on retiming in [KuLe80, LeSa83, SYKun88]). The appeal of this terminology is that of providing an intuitive picture of the effect of the transformations, which certainly helps in the understanding of the mathematics.

In general, we associate the term pipelining with a particular direction vector in the lattice space, and we expect the data to be propagated in a regular fashion through the computation points in that direction. On the other hand, we associate the term routing with a set of direction vectors in the lattice space, used to define a sequence of directions used for the propagation of the data among computations. Such directions contribute to the definition of routing paths on the nodes of the dependence graph. A regular routing scheme is one such that the propagation of the data flow in each of the routing directions is pipelined.

Given a non-uniform data dependence, its regularisation may be achieved by selecting a number of routing directions in the lattice space and pipelining the data flow in each of those directions. We illustrate these two activities on the data dependence graph depicted in Fig. 2.8 a).
Fig. 2.8. Regularisation: a) a generic data dependence; b) and c) possible decompositions; d) a possible uniformisation.

The basic requirements for the choice of routing directions, is that each data dependence vector can be expressed as an integer combination of those directions. For instance, any unimodular basis (see Appendix F) of the space will meet this requirement. The integer combination can be used for the definition of a routing path which replaces the dependence vector, simply by fixing an ordering of the routing vectors and considering the summands of the combination in the sequence established by the ordering. In our example, let us choose as routing directions the vectors (1,1) and (1,0). These vectors constitute a unimodular basis in $\mathbb{Z}^2$, therefore any data dependence vector in Fig. 2.8 a) can be rewritten as a combination of (1,1) and (1,0) with suitable integer coefficients. By considering (1,1) and (1,0) in this order, the data dependence vectors of Fig. 2.8 a) may be replaced by the routing paths of Fig. 2.8 b), where solid lines correspond to the direction (1,1) and dashed lines to (1,0). Reversing the ordering of the routing directions produces the paths of Fig. 2.8 c). Note that one of the effects of this substitution is that more points of the lattice space are involved in the transfer of the data among computations. Hence, from a point-to-point connection represented by a data dependence vector, we define a path of lattice points through which the data is transferred. The end points of the path are the pair of points initially related by the dependence vector.

In the following chapters, we will use the term *decomposition* for the techniques which allow the definition of these routing paths for various classes of non-uniform data dependencies. In general, decomposition will also allow us to separate and consider singularly the sub-
paths generated in each of the routing directions (hence the name we have chosen for the transformation).

Once the routing paths have been established by a selection of the routing directions and their ordering, the data flow can be pipelined along the paths. In our example, the routing paths of Fig. 2.8 b) are transformed into those of Fig. 2.8 d), by pipelining the data according to the direction of \((1,0)\). Note that the result of this transformation is a set of more dense routing paths, which involve nearest neighbour points in each of the routing directions. The name which we have chosen for this transformation is uniformisation, as, in general, it corresponds to the definition of a set of uniform data dependencies.

2.3.2 Data Conflicts and Data Broadcasts

A data conflict is an attempt by distinct pieces of data to share the same communication channel during the same time cycle. A data conflict is avoided by providing distinct communication channels for each of these pieces of data. If the array design is realised in hardware, because of the cost involved, the amount of data conflicts needs to be contained. Also, if the level of data conflicts increases with the size of the problem, the array design is not scalable. This is a strong limitation for hardware implementation as it would involve the replacement of the whole piece of hardware every time a problem of larger size needs to be solved. Therefore, array designs characterised by data conflicts which are dependent on the size of the problem should be avoided.

Data conflicts may be generated by overloading the dependence graph of a specification. The nodes of a dependence graph become overloaded when the pipelining of the data flow according to a certain direction produces several routing paths which cross the same computation points. Then distinct communication channels at the corresponding processing elements need to be provided in order to avoid conflicts. If the level of overloading is dependent on the problem size, the level of conflicts exhibits the same dependence. Therefore, a major constraint in the selection of routing directions is to avoid directions of the lattice space which yield problem size dependent overloading of the data dependence graph. Let us consider once again the data dependence graph of Fig. 2.8 a). The choice we made of routing directions and their ordering resulted in the graphs in Fig. 2.8 b) and d). Note that some of the nodes of the graph in Fig. 2.8 d) are overloaded (those labelled as A and B).

The appearance of overloading can be related to the presence of data broadcasts. In terms of array design, a data broadcast indicates that the same data is shared by several processing
elements during the same time cycle, and therefore has to be transferred from some source processing element to all the recipients with a direct communication. Data broadcasts usually involve non-local communication between processing elements and in regular array design are always replaced by a pipelining of the data among neighbour recipients. In a specification, a data broadcast corresponds to a one-to-many data dependence relation, that is several nodes of the dependence graph depend on the same node. In Fig. 2.8 a) several data broadcasts are present. When compared with Fig. 2.8 d) the overloading appears exactly on those paths replacing data broadcasts. A more precise way of defining a data broadcast is as the data dependence relation defined by a non-injective index mapping (trivially, if several points depend on the same point, that point is their image under the index mapping, and the mapping is not injective). Therefore, overloading the nodes of the graph can be avoided by avoiding the definition of data broadcasts.

It is this important principle which is exploited in the definition of the regularisation techniques of the next chapters: routing directions are selected such that their routing paths correspond to data dependencies defined by injective index mappings. Indeed the question to address is whether this selection is always possible. Fortunately, in general the answer is affirmative. However, it may require an increase in the number of dimensions of the specification. For instance, let us consider, once again, the graph of Fig. 2.8 b). Overloading can be avoided by adding a third dimension to the problem and producing the graph of Fig. 2.9. In the figure, white nodes denote the extra routing points in the 3-dimensional space. Note that not only an increase of dimensionality is required, but also new routing directions are used (the order of the paths, however, remains unchanged).
2.3.3 Regularisation and Dependence Cone

Because of the relation between affine timing functions and pointed dependence cones, a condition for the preservation of affine scheduling by regularisation may be formulated in terms of the preservation of pointedness of the dependence cones. In particular, the routing directions need to be chosen as vectors of the lattice space which, together with the original dependence cone of the specification, generate a pointed polyhedral convex cone. This is possible, in general, but may require an increase of the dimensionality of the space. The formulation of this condition for various types of data dependencies will be provided in the following chapters.

2.3.4 Substitution of a Data Dependence

The definition of a routing scheme which replaces a data dependence finds a mathematical expression in the function composition of index mappings. It is not limited, however, to such a composition as the computation domains and variables of the data dependencies have also to be taken into account. The definition of a routing scheme may be described as follows.

Let $\mathcal{DD} = (D, U, V, I)$ be a data dependence describing the dependence relation between the instances of variables $U$ and $V$ on the domain $D$. In particular, for all $z \in D$, $U(z)$ is data dependent on $V(I(z))$. In other words, the value of $V$ at the computation point $I(z)$ has to be transferred to $z$ in order to enable the computation of $U$. $\mathcal{DD}$ implies that there exists a direct data communication from $I(z)$ to $z$. A routing scheme for $\mathcal{DD}$ defines a communication path from $I(z)$ to $z$ via a number of intermediate points of the lattice space. The routing scheme has to guarantee the following two requirements: that for each $z \in D$ the communication path from $I(z)$ to $z$ is finite; and that the value $V(I(z))$ is transferred among neighbour points on the communication path from $I(z)$ to $z$ via a set of routing variables.

In order to define the routing scheme, a system of recurrence equations is introduced which defines the routing variables, and the original data dependence is replaced by a new data dependence which is defined on such variables. Let us illustrate the transformation in the following example.

**Example 2.3.1** Consider the data dependence $\mathcal{DD} = (D, U, V, I)$, such that $D = \{(i, j) | i \geq 1, 1 \leq j \leq m\}$, for some $m$ in $\mathbb{N}$, and $I(i, j) = (0, j)$. $I$ defines the data broadcasts sketched in Fig. 2.10 a). For each point of $D$, we want to replace any such data broadcast with a pipelined propagation as illustrated in Fig. 2.10 b). This is achieved by replacing $\mathcal{DD}$
by the data dependence
\[ DD' = (D, U, R, I_0) \]
where \( I_0(i, j) = (i, j) \) and \( R \) is the routing variable defined by the equations:
\[
E_1 = (D_1, R, R, id, I_1) \\
E_2 = (D_2, R, V, id, I_1)
\]
with \( I_1(i, j) = (i - 1, j) \), \( id(a) = a \), and
\[
D_1 = \{ (i, j) \mid i \geq 2, 1 \leq j \leq m \} \\
D_2 = \{ (i, j) \mid i = 1, 1 \leq j \leq m \}.
\]
By induction, it is easy to show that, for all \((i, j) \in D, I_1^{i} \circ I_0(i, j) = I(i, j)\) (where \( I_1^{i} \) denotes the composition \( I_1 \circ I_1 \circ \ldots \circ I_1 \), \( i \) times – see Appendix A), and
\[
U(i, j) = R(I_0(i, j)) = R(I_1^{i-1} \circ I_0(i, j)) = V(I_1^{i} \circ I_0(i, j)) = V(I(i, j)).
\]

### 2.4 A Brief Survey

The basic ideas behind regular array design can be related (see [Meg92]) to the theory of cellular automata, whose foundations were established by John von Neumann [VonN66] in the early 60s. Cellular automata were initially developed for the study of the evolution of biological systems, although several applications in mathematical and physical sciences have successively been developed. Cellular automata deal with large (possibly infinite) collections of interconnected finite state automata and, hence provide a framework for the investigation of systems characterised by homogeneous and scalable components. (An overview of the theory and application of cellular automata can be found in [Wol86].)
The formal development of regular array synthesis began in the late 60s with the work by Karp, Miller and Winograd [Ka-et-al67], who proposed the idea of expressing (classes of) iterative algorithms as systems of uniform recurrence equations (UREs). In their work, the computability of the algorithm was related to properties of the (reduced) dependence graph of the equations, and necessary and sufficient conditions for the existence of a scheduling were stated.

While the work by Karp et al. is based on a functional representation of an algorithm, the work by Lamport [Lam74], may be considered as its imperative counterpart. His work is based on the observation that for large classes of (numerical) algorithms, most of the computation is devoted to the execution of loops, in particular FORTRAN-like for-loops. Under a number of assumptions on the form of the loops (such as the absence of input/output operations or of control transferred outside the loop), Lamport proposed a systematic rewriting of sequential nested for-loops into concurrent loops, based on the analysis of the data dependencies. He also introduced the notion of equitemporal hyperplanes for the scheduling of the computations, a notion which has been extensively exploited in regular array synthesis. His work has been also most influential in the subsequent development of parallel compilers.

In the early 70s, data flow computing [Ada68, Ada70, DeWe77, Den80] was defined for the maximal exploitation of parallelism as an alternative to conventional control flow computers. The essential idea of data flow computing is that of enabling the execution of an instruction as soon as its operands become available. In other words, computations are driven by data availability rather than explicit control.

In the early 80s, systolic arrays were introduced by H.T.Kung and C.E.Leiserson [KuLe80]. The term array indicates their structural affinity with array processors, while the term systolic describes their behaviour using the human circulatory system as a metaphor (signals are rhythmically "pumped" among processing elements). Systolic arrays are structurally simple, regular and modular and, typically, are realised through replication and local interconnection of simple processing elements which perform basic operations. Multiprocessing and pipelining are principles of systolic behaviour ensuring high performance with low memory and input/output bandwidth (the same data can be propagated among neighbouring processing elements to be, thus, reused by the recipients). Also, parallelism is synchronous and decentralised: computations occur in lockstep, with signals representing both data and control information. Although systolic arrays were initially designed for hardware implementation as VLSI circuits, the principles of systolic computation are now considered more
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generally as a paradigm for parallel processing, and in synthesis methods they are retained as a computational model.

Asynchronous regular arrays, so-called wavefront arrays, were introduced by S.Y. Kung et al. in the early 80s [Ku-et-al81, Ku-et-al82]. Wavefront arrays are based on the principles of data flow computing. The name wavefront is evocative of the way their computations proceed, which resembles a wave propagation. A major advantage of wavefront arrays over systolic arrays is that they do not require global synchronisation.

An automata and complexity theory for systolic computing, known as systolic automata theory, developed through the 80s and early 90s [Cu-et-al83, Cu-et-al84, Gru84, FaNa88. Gru90], mainly dealing with a restricted class of problems which have systolic solutions as linear array, trellis or tree-like forms.

From an applicative point of view, a vast literature exists on systolic arrays for the solution of numerical and non-numerical problems. Systolic arrays have become a popular form of parallel computing, and collections of systolic algorithms can be found, for example, in [QuRo91, Eva91, Meg92].

Initially, array designs were derived manually and in an ad hoc fashion. Attempts at coding the expertise acquired in the development of systolic algorithms into a methodology with mechanised support started in the middle of the 80s. Between 1983 and 1985 a number of independent contributions defined the foundations of regular array synthesis. Moldovan [Mol83] introduced the notion of space-time mapping of the data dependence graph as a pair of linear functions. He also defined the concept of dependence mapping (rather than simply a dependence vector as it is for UREs), hence defining a new type of recurrence equations for the specification of algorithms. We refer to such recurrences as affine recurrence equations (AREs), a name introduced later by Rajopadhye and Fujimoto [RaFu90]. Quinton [Qui84] proposed the detection of a scheduling for system of UREs as the solution of an (integer) linear optimisation problem. Cappello and Steiglitz [CaSt84] focused on the geometric representation of an algorithm and its data dependencies as a graph in a Euclidean lattice space together with sets of linear transformations of such a representation. They showed that this geometric framework is a powerful unification tool, as several array designs for the same algorithm correspond to the mapping of data dependence graphs which are linear transforms of the same specification. Miranker and Winkler's contribution [MiWi84] can be seen as emphasising that the mapping problem reduces to a mapping between graphs: the space-time representation of the algorithm and its data flow representation.
Between 1985 and the early 90s, several contributions followed these seminal works on methodology. It would be impractical to enter the details of each of them. To summarise, even though there were no major revisions to the methodology, particular aspects of regular array synthesis were addressed, resulting in the refinement of the design process to the structure illustrated in Fig. 2.11. This diagram is a more detailed version of that of Fig. 2.1 at the beginning of this chapter, with more steps included in the core of the design process. In the following we provide a brief review of some of the major contributions.

The analysis step, together with the analysis of the data dependencies, also addresses the computability of the specification, where a computable specification is a specification which admits a valid timing function. The computability of uniform and affine recurrence equations has been investigated widely [Ka-et-al67, Rao85, Delp86, Delp87, Ra-Ka88, SaQu90]. It has been established that if the computation domains are unbounded, the computability of a system of recurrence equations can be reduced to the halting problem [BoJe89], and so is undecidable. On the other hand, for uniform recurrence equations with a finite computation set, computability is decidable and corresponds to checking whether the (complete) data dependence graph is acyclic.

A correspondence between systolic arrays and algorithms specified by recurrence equations was formally established by Rao in his doctoral thesis in 1985 [Rao85]. In particular he showed that if an algorithm can be implemented as a systolic array then it can be expressed as a Regular Iterative Algorithm (RIA), an extension of UREs with finite conditional branches in
CHAPTER 2. REGULAR ARRAY SYNTHESIS

each recurrence equation. Li and Wah [LiWa85] treated the derivation of a systolic design from a specification as an optimisation problem, based on a set of design parameters such as the velocity of data flows, the spatial distribution of data, or the periods of computation. Similarly, M. Chen [Che86] treated the design process as an optimisation problem applied to algorithms specified in the parallel programming language Crystal [Che86b]. Her work included an initial investigation into aspects of the design such as regularisation, synthesis of control signals and mapping to fixed-size architectures. Delosme and Ipsen [DeIp86] concentrated on computability issues and extended the work of Karp et al. [Ka-et-al67] to systems of AREs. Based on the work by Delosme and Ipsen, Yaacoby and Cappello [YaCa88] provided necessary and sufficient conditions for the existence of an affine scheduling and a procedure to construct a scheduling vector. Similarly, Rajopadhye and Fujimoto [RaFu90] extended the work of Quinton [Qui84] to AREs, defining a linear optimisation problem for the automatic derivation of an affine scheduling.

Early work on regularisation includes the retiming of a signal flow graph, for example, by Leiserson and Saxe [LeSa83]. Subsequent regularisation techniques were defined by several authors, mainly for the removal of data broadcast in AREs. As data broadcasts correspond to non-injective index mappings, for affine data dependencies, they can be related to rank deficient matrices (defining the linear part of the index mapping), and simple forms of regularisation can be provided based on the selection of pipelining directions in the null space of such matrices. This approach was taken by Fortes and Moldovan [FoMo84] and Rajopadhye and Fujimoto [RaFu87, Raj89]. Wong and Delosme [WoDe92] proposed more general forms of regularisation of data broadcasts based on the selection of routing vectors as the elements of canonical and non-canonical basis of the lattice space. Regularisation techniques for more general forms of non-uniform AREs (i.e., not limited to data broadcasts) were proposed by Quinton and Van Dongen [QuVa89], via a combination of pipelining and routing. They also defined a new class of recurrences, generalising AREs with linear size parameters. Our work in this thesis on the regularisation of non-affine types of data dependencies has developed from their approach.

The systematic derivation of control signals from conditional expressions was first addressed by M. Chen [Che86] in 1986. In this work she proposed methods to replace conditionals of a recurrence equation by control signals pipelined from the boundary of the array. Rajopadhye and Fujimoto [RaFu87] also proposed a systematic pipelining of conditional expressions. However, they considered a more restricted class of conditional expressions which arise be-
cause of the application of regularisation techniques to AREs. The characterisation of such conditionals resulted in the definition of the class of so-called conditional uniform recurrence equations (CUREs). Teich and Thiele in [TeTh91] adopted an approach similar to Chen’s for what they defined as piecewise regular iterative algorithms (an extension of Rao’s RIAs with affine data dependencies). Their formalism is based on Chandry and Misra’s UNITY [ChMi88]. Finally, Xue [XuLe92, Xue92] proposed a more general method that also extends to space-time mappings of a dependence graph onto fixed-size array designs.

Regular array synthesis aims at producing optimal array designs based on the assumption that unbounded computational resources are available. For more realistic situations in which this assumption cannot be made, partitioning techniques have been developed which allows the mapping of, possibly, multi-dimensional algorithms onto fixed-size lower-dimensional array designs. The development of partitioning techniques is not confined to regular array design, but is part of the more general problem of parallel code generation from sequential code (see, e.g., [IrTr88, AnIr91, Fea92b]). The main goal of partitioning methods is to operate a compression of the design. This can be achieved either at the data dependence level, by defining clusters of computations to allocate onto the same processor, or at the signal flow graph level, by merging clusters of cells into a single (super-) processor. Partitioning techniques are usually related to the use of multi-dimensional schedules (see, e.g., [Fea92b]), i.e., the timing function is a mapping between multi-dimensional lattice spaces. Work on partitioning for regular arrays started in 1986 with a contribution by Moldovan and Fortes [MoFo86]. Partitioning techniques were classified by Darte [Dar91] into Locally Parallel Globally Sequential (LPGS) and Locally Sequential Globally Parallel (LSGP). In LPGS partitioning, each partition is a block of parallel computations, while the blocks are processed sequentially. This class includes the work by Moldovan and Fortes [MoFo86] and Bu et al. [Bu-et-al90]. In LSGP partitioning, each partition is a sequential block of computations and blocks are executed in parallel. Darte’s work [Dar91] belongs to this class. Independent partitioning was proposed by Shang and Fortes [ShFo92b], where independence means that no communication is needed between the blocks of the partition, while X. Chen and Megson [MeCh94] related partitioning to code generation for parallel platforms (in particular, transputers), by exploiting the idea of canonical dependencies derived from a positive expressive basis.

Verification refers to the formal proof of some correctness properties of a regular array design. Early work on verification, undertaken since 1983, includes: the approach by M. Chen [Che83], based on systems of recurrence equations and fixed point induction; the algebraic
approach by Kung and Lin [KuLi84]; and the approach based on the solution of systems of differential equations by Melhem and Rheinboldt [MeRh84]. In 1986, Hennessy [Hen86] used process algebras for the specification of a systolic circuit and fixed point induction to derive an implementation from the specification. With work started in 1988, Thompson and Tucker [ThTu88, ThTu91, ThTu94] have developed formal specification and verification techniques for *Synchronous Concurrent Algorithms (SCA)*, of which systolic arrays are a particular case. Their method is based on many-sorted universal algebras, primitive recursion and equational logic. Work towards automatic verification systems for systolic arrays includes: the approach by Abdulla [Abd90, Abd92], which addresses a general model for the description and verification of systolic circuits over arbitrary algebras (he provides completely automatic verification for subclasses of systolic circuits [Abd92]); and the work by Ling and Bayoumi [LiBa94], which defines a systolic temporal arithmetic (based on temporal logic) for the specification and verification of systolic designs at the array level (for which they provide a Prolog-based verifier).

Finally, as the automation of the design process is one of the basic objectives of regular array synthesis, a number of support software tools and environments have been developed. Among the major contributions in the form of Computer-Aided Design (CAD) tools, we may recall DIASTOL [Ga-et-al87] and its more complete version Alpha du Centaur (AdC) [Ga-et-al88], ADVIS (Automatic Design of VLSI Systems) [Mo187], PRESAGE, [VanD88], DECOMP [VeCr91] and SADE (Systolic Array Design Environment) [MeCo91]. In general, these tools support (some of) the basic steps of regular array synthesis, such as the generation of the data dependence graph and its (optimal) space-time mapping. Languages for the initial specification of an algorithm vary from case to case, and include specialised languages based on recurrence equations or restricted forms of nested for-loops. Early tools accept as inputs only uniform problems, while more recent developments can be used also for the synthesis of affine problems. Often graphical interfaces have been developed for the representation of the data dependence graph as well as the animation of the array design through snapshots of computations. More sophisticated tools also include parallel code generation. Compilers for systolic and regular programs have also been developed. The first systolic compiler was developed by H.T.Kung *et al.* in the early 80s, for the CMU Warp machine [KuWe85, An-et-al87], a systolic array computer of linearly interconnected programmable cells. A more general approach to systolic compilation is due to Lengauer *et al.* [HuLe87], in which, from a formally specified program, traces (of operations) are extracted, transformed into
parallel traces, and a corresponding systolic design is derived. The design can then be either implemented in hardware or corresponding parallel code can be generated for a target machine [Le-et-al91].

2.5 Summary

In this chapter we have introduced some of the basic concepts of regular array synthesis. In particular, the design process was outlined and the basic notions of recurrence and input equation, equation system, data dependence and relative graphical representations were formally discussed. We have explained how a regular array can be derived from an algorithm specification through a space-time mapping, that is a scheduling of the computations and their placement onto processing elements. Formal properties of such a space-time mapping were stated such as the validity, finiteness and boundedness of the scheduling, and the compatibility of scheduling and placement.

We have discussed the advantages of developing synthesis methods in the framework of Euclidean geometry, both theoretically and in the practical development of systematic transformations. In this framework, powerful techniques can be developed by restricting ourselves to affine data dependencies and affine timing and allocation functions, and exploiting the basic properties of linear and affine spaces. Among the major advantages is the possibility for the systematic derivation of optimal affine timing functions. Central to Euclidean synthesis methods is the concept of dependence cone, as a polyhedral convex cone embedding all possible data dependence vectors of an algorithm specification. Its pointedness can be related to the existence of linear schedulings and their preservation by transformations of the specification.

The rôle of regularisation in the design process was discussed at length, the development of regularisation techniques being the main theme of this thesis. Routing and pipelining were outlined, on a small example, as basic transformations for regularisation. We have explained the relation between data conflicts and data broadcasts, and how they can be detected and avoided in regularisation. The relation between regularisation directions and dependence cones was discussed together with guidelines for the preservation of affine timing functions.

From the discussion, a regularisation scheme has emerged, which will be used in the development of the following chapters. It consists of a syntactic characterisation of classes of non-uniform data dependencies, and their systematic substitution with new data dependen-
cies which exhibit improved locality and uniformity, the target being specifications as systems of uniform recurrence equations. The new data dependencies are defined by selecting regularisation directions in the embedding Euclidean space and defining regular routing systems for transferring data among computation points. Conditions have to be provided for the preservation of affine scheduleings by relating regularisation directions and (the generators of) data dependence cones. Conflict-freeness has to be guaranteed.

The last part of the chapter was devoted to a brief survey of the major contributions in regular array synthesis.
Chapter 3

Integral Recurrence Equations

In this chapter we introduce integral recurrence equations and their systematic regularisation. We will base the definition of integral recurrence on the notion of integral index mapping, as an index mapping in $\mathbb{Z}^n$ which is not required to define an affine transformation. The syntactic form of an integral index mapping is an integer combination of a finite set of directions of the lattice space, in which the coefficients are functions from $\mathbb{Z}^n$ to $\mathbb{Z}$. With this form we will be able to establish an explicit relation between the index mapping and a finite set of vectors of the space, which can be exploited for regularisation purposes. Based on this syntactic form we will show that affine data dependencies are particular types of integral data dependencies.

The regularisation techniques that we will define will allow us to transform integral specifications systematically into systems of uniform recurrence equations. This fact implies that ordinary mapping techniques can subsequently be applied for the derivation of regular array designs. Regularisation directions will be chosen among the direction vectors defining the index mappings. Based on the same vectors, conditions will be given for the existence of affine timing functions and their preservation through regularisation.

The main difficulty in the definition of regularisation techniques for integral data dependencies stems from the necessity of reconciling the existence of polyhedral convex sets and the application of non-affine transformations. Because of the role of convexity in Euclidean synthesis, we need to guarantee that all computation domains are (convex) polyhedra. However, non-affine index mappings, in general, do not preserve convexity. The solution we have chosen is that of enforcing convexity, at the expense of an increased complexity of the regularisation techniques. In particular, we will consider enlarged polyhedral convex domains and define control variables\(^1\) in order to identify non-convex subsets of computation points in those do-

\(^1\)A control variable is an ordinary variable of the specification, whose values are interpreted as control signals rather than data. As in a regular array, control signals (other than clock synchronisation) are entirely
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mains. Hence, the solution we will adopt introduces some control overhead, but it allows us to recover convexity and remain within the domain of classic regular array synthesis. As we will see in the next chapter, the use of control variables also accounts for a degree of reconfigurability of the array design, which we will exploit for particular classes of dynamic data dependencies.

This chapter is organised as follows. In Section 3.1 we define integral index mappings and data dependencies, and discuss their relation with the affine and uniform cases. Section 3.2 is devoted to regularisation techniques. In particular, a condition for the injectivity of an integral index mapping is established, and used for the development of regularisation techniques which guarantee that no size dependent overloading of the dependence graph is generated. The regularisation techniques developed have the form of decomposition and uniformisation techniques. A parametric version of uniformisation is also defined, where the parameter is seen as an upper bound on the level of overloading allowed, hence the level of conflicts and corresponding physical resources of the array design can be controlled to some extent. Parametric uniformisation allows for compact designs at the expense of increased communication resources. That the regularisation techniques preserve affine scheduling is formally addressed in Section 3.3. Toy examples are used in the chapter to clarify the basic results. More interesting applications of the techniques will be found in Chapter 5.

3.1 Integral Data Dependencies

We define an integral index mapping as an index mapping in $\mathbb{Z}^n$, which can be expressed as an integer combination of a finite set of direction vectors in the lattice space. In addition, when only one direction vector is used, we call the index mapping atomic integral (or simply atomic).

Definition 3.1.1 [Integral Index Mapping] Let $I$ be an index mapping. $I$ is integral if for all $z \in \mathbb{Z}^n$, $I(z) = z + \sum_{j=1}^{m} g_j(z)d_j$, where, for $j = 1, \ldots, m$, $g_j : \mathbb{Z}^n \rightarrow \mathbb{Z}$ and $d_j$ is a non-null vector in $\mathbb{Z}^n$. In addition, $I$ is atomic integral if $m = 1$.

We call the vectors $d_j$ the generators of $I$, and the integer functions $g_j$ its coefficients. The form of an integral index mapping of Definition 3.1.1 is not unique, as illustrated in the following example.
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Example 3.1.2  Consider the index mapping $I(i, j, k) = (i - 2^j, j - 1, k - 1)$. $I$ may be expressed, for instance, by any of the two integral forms:

$$I(i, j, k) = (i, j, k) + 2^j (-1, 0, 0) + 1 (0, -1, 0) + 1 (0, 0, -1)$$

In theory, any number of generators and corresponding coefficients can be chosen to express an integral index mapping. In practice, however, only expressions with a number of generators less or equal to the number of dimensions of the lattice space are actually considered. Indeed, this is always possible (trivially, by considering the elements of the standard basis of the space). As we will see, the choice of generators has an impact on the regularisation of the corresponding data dependencies, hence on the complexity of the resulting array design. We will consider this issue in greater detail in Section 3.2.1.

Integral dependence mappings may be defined, similarly to the affine case, as follows:

Definition 3.1.3 [Integral Dependence Mapping] Let $I$ be an integral index mapping. Its dependence mapping $\Theta_I$ is the mapping defined as $\Theta_I(z) = z - I(z)$, for all $z \in \mathbb{Z}^n$.

If $I$ is an integral index mapping defined as $I(z) = z + \sum_{j=1}^m g_j(z)d_j$, its dependence mapping is $\Theta_I(z) = - \sum_{j=1}^m g_j(z)d_j = \sum_{j=1}^m g_j(z)(-d_j)$. We call the vectors $-d_j$ the generators of $\Theta_I$ and the integer functions $g_j$ its coefficients. Note that, in general, $\Theta_I$ is not affine. Integral index mappings characterise integral data dependencies:

Definition 3.1.4 [Integral Data Dependence] Let $\mathcal{D} = (D, U, V, I)$ be a data dependence. $\mathcal{D}$ is integral if $I$ is an integral index mapping. In addition, $\mathcal{D}$ is atomic integral if $I$ is atomic integral.

The relation between atomic integral and integral data dependencies is similar to that between uniform and affine data dependencies. In particular, atomic integral data dependencies are simple forms of integral data dependencies, and for regularisation purposes we will aim at substituting a generic integral data dependence with a set of corresponding atomic integral data dependencies.

The dependence domain and cone of an integral data dependence can also be defined similarly to the affine case.
Definition 3.1.5 [Integral Dependence Domain] Let $D \subseteq (D, U, V, I)$ be an integral data dependence and $\Theta_T$ the dependence mapping defined by $I$. The dependence domain $\Omega_T$ of $D \subseteq$ is $\Omega_T = \Theta_T(D)$.  

As, in general, $\Theta_T$ is not affine, the dependence domain $\Omega_T$ is not a convex polyhedral set.

Definition 3.1.6 [Integral Dependence Cone] Let $D \subseteq (D, U, V, I)$ be an integral data dependence with dependence domain $\Omega_T$. The dependence cone $\Theta_T^*$ of $D \subseteq$ is $\Theta_T^* = \text{cone}(\Omega_T)$.  

Because of the relation between pointed dependence cones and affine scheduling discussed in Section 2.2.6, in the following we always assume that $\Theta_T^*$ is pointed.

Example 3.1.7 Consider the data dependence $D \subseteq (D, U, V, I)$ with domain $D = \{(i, j, k) \mid 1 \leq i, j \leq p, k = 1\}$, for some $p \in \mathbb{N}$, and index mapping $I(i, j, k) = (i, j, k) + 2^i(-1, 0, 0) + (0, -1, -1)$. Its dependence mapping is $\Theta_T(i, j, k) = 2^i(1, 0, 0) + (0, 1, 1)$. Its dependence domain is $\Omega_T = \{(2^i, 1, 1) \mid 1 \leq j \leq p\}$ and its dependence cone $\Theta_T^*$ is pointed (see Fig. 3.1).  

3.1.1 Integral vs. Affine Recurrences

Because of the generality of our definition of an integral index mapping, it is easy to show that affine index mappings constitute a particular type of integral index mappings.

Let us consider an affine index mapping $I(z) = Az + b$, with $A \in \mathbb{Z}^{n \times n}$ and $b \in \mathbb{Z}^n$. Then $I$ can be expressed as the integral index mapping below. where, for $j = 1, \ldots, n$: each coefficient $g_j$ is defined by the vector expression $g_j(z) = (A - I_n)_{j \cdot z} + b_j$, with $(A - I_n)_{j \cdot}$ denoting the $j^{th}$ row of the matrix $A - I_n$, and $b_j$ the $j^{th}$ component of $b$; and each generator $e_j$ is the $j^{th}$
Fig. 3.2. Inclusions among the classes of uniform (UREs), affine (AREs), atomic integral (AIREs) and integral (IREs) recurrence equations.

vector of the standard basis of $\mathbb{Z}^n$. The mapping is:

$$I(z) = Az + b = z + (A - I_n)z + b = z + \sum_{j=1}^{n} g_j(z)e_j.$$ 

Trivially, all uniform index mappings are atomic integral. In fact, a uniform index mapping has a generic form $I(z) = z + b$, which can be seen as an integral form with a single generator $b$ and, as coefficient, the constant function $g(z) = 1$.

Given the above relations, a taxonomy of the different classes of recurrences\textsuperscript{2} can be defined as illustrated in Fig. 3.2. In particular, affine recurrences (AREs) constitute a proper subclass of integral recurrences (IREs), while uniform recurrences (UREs) are in particular atomic integral (AIREs).

Therefore, integral recurrence equations can be used as more general forms of specification than those obtained with affine recurrences only. For specification in the intersection of the two classes, the convenience of one formalism over the other needs to be evaluated case by case. In particular, a trade-off between ease of expression and complexity of the required regularisation should be considered. We will return to this point in the discussion at the end of the chapter.

### 3.2 Regularisation

The regularisation of an integral data dependence consists of two main steps:

- the substitution of a generic integral data dependence by a set of atomic integral data dependencies; and

\textsuperscript{2}It is custom to name recurrence equations after the (most general) type of their data dependencies.
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- the substitution of each atomic integral data dependence by a set of uniform data dependencies.

Routing directions are chosen among the generators of the data dependence mapping, which guarantee the definition of injective index mappings, hence no overloaded data dependence graphs are generated. The major difference with respect to regularisation techniques for the affine case is the necessity of using control variables to reconcile the preservation of convexity with the application of non-affine transformations.

3.2.1 Regularisation Directions

Let \( \mathcal{D} = (D, U, V, I) \) be an integral data dependence, with pointed dependence cone \( \Theta_I \). We intend to use a set of non-null generators of \( I \) as regularisation direction vectors. To do so, we want to express \( I \) so that the generators of \( I \) form a pointed cone \( C' \) containing \( \Theta_I \).

Such a representation is particularly convenient because:

- it allows us to formulate conditions for the existence and preservation of affine scheduling through regularisation;

- it simplifies the definition of regularisation techniques, as the coefficients define non-negative integer functions on \( D \).

Such a form always exists, as can be proved by using a result due to Quinton and Van Dongen [QuVa89]. The result is contained in the following proposition (a proof of which is sketched in Appendix B):

**Proposition 3.2.1 [QuVa89]** Let \( C \) be a pointed polyhedral convex cone of full dimension in \( \mathbb{Q}^n \). Then there exists a pointed polyhedral convex cone \( C' \) such that \( C' \) contains \( C \) and its extremal rays constitute a unimodular basis of \( \mathbb{Z}^n \).

We illustrate the result in the following example.

**Example 3.2.2** Consider the data dependence \( \mathcal{D} = (D, U, V, I) \) with domain \( D = \{(i, j) \mid -2 \leq i \leq 2, 1 \leq j \leq 2 \} \) and index mapping \( I(i, j) = (i, j) + 6(-1, -1) + i^3(-1, 0) \).

The dependence mapping is \( \Theta_I(i, j) = 6(1, 1) + i^3(1, 0) \) and \( \Theta_I \) is illustrated in Fig. 3.3 a).

The dependence cone is pointed (intuitively, it does not contain any line). However, the coefficient \( i^3 \) does not define a non-negative integer function over \( D \).
Fig. 3.3. a) Data dependence cone $\Theta^*_T$; b) Pointed cone $C$ containing $\Theta^*_T$.

Consider the pointed cone $C = \text{cone} \{(-1,1),(1,0)\}$. $C$ contains $\Theta^*_T$ (see Fig. 3.3 b)) and its generators constitute a unimodular basis (i.e., they integrally span $\mathbb{Z}^2$ – see Appendix F, where unimodularity is defined). The correspondence between the generators of $\Theta_T$ and those of $C$ is the following:

\[(1,1) = 1(-1,1) + 2(1,0)\]
\[(1,0) = 0(-1,1) + 1(1,0).\]

Therefore we can rewrite $\Theta_T$ and $I$ as follows:

\[\Theta_T(i,j) = 6[1(-1,1) + 2(1,0)] + i^3[0(-1,1) + 1(1,0)]\]
\[= 6(-1,1) + (12 + i^3)(1,0)\]
\[I(i,j) = (i,j) + 6(1,-1) + (12 + i^3)(-1,0).\]

With this rewriting, the coefficients of $I$ are non-negative over $D$.  

Note that if $\Theta^*_T$ is pointed and the coefficients of $\Theta_T$ are non-negative integer functions on $D$, the cone $C$ defined by the generators of $\Theta_T$ contains the dependence cone $\Theta^*_T$. However, $C$ is not guaranteed to be pointed, as shown in the following example.

**Example 3.2.3** Consider the data dependence $D = (D,U,V,I)$ with domain $D = \{(i,j) \mid 1 \leq i, j \leq m\}$, where $m \in \mathbb{N}$, and index mapping $I(i,j) = (i,j) + 6(1,0) + 4(-1,0)$. The dependence cone $\Theta^*_T$ is pointed. However, $C = \text{cone} \{(1,0),(-1,0)\}$ contains a line (see Fig. 3.4). Indeed, a straightforward rewriting of $I$ exists such that $\Theta^*_T$ and $C$ coincide. The rewriting is $I(i,j) = (i,j) + (6 - 4)(1,0) = (i,j) + 2(1.0).$  

In the following, we always assume that given an integral data dependence $D = (D,U,V,I)$, the generators of $I$ are non-null vectors, its coefficients define non-negative integer functions on $D$, and the cone $C$ defined by the generators of $\Theta_T$ is pointed. We will also refer to the cone $C$ as the embedding dependence cone.
3.2.2 Injectivity of an Atomic Integral Index Mapping

From the discussion in Section 2.3, we know that a data broadcast corresponds to a non-injective index mapping. A sufficient condition for the injectivity of an atomic integral index mapping $\mathcal{I}$ on a domain $D$ can be established by considering the relation between the generator of $\mathcal{I}$ and the direction $\text{lin}(D)$ of the domain $D$ (for a definition of $\text{lin}(D)$, see Appendix F). The result is contained in the following proposition:

**Proposition 3.2.4** Consider an atomic integral index mapping $\mathcal{I}$ and a domain $D \subseteq \mathbb{Z}^n$, such that for all $z \in D$, $\mathcal{I}(z) = z + g(z)d$ and $g(z) \geq 0$. If $d \notin \text{lin}(D)$ then $\mathcal{I}$ is injective over $D$.

**Proof:** From linear algebra, if $z, z' \in D$ then $z - z' \in \text{lin}(D)$. Assume that there exist $z, z' \in D$, such that $z \neq z'$ and $\mathcal{I}(z) = \mathcal{I}(z')$. We want to prove that this assumption always implies a contradiction with respect to the hypotheses of the proposition and therefore for all $z, z' \in D$, $z \neq z'$ implies $\mathcal{I}(z) \neq \mathcal{I}(z')$. There are only two possibilities, both leading to a contradiction. If $g(z) = g(z') = c$, then $\mathcal{I}(z) = \mathcal{I}(z')$ implies $z + cd = z' + cd$, i.e., $z = z'$. Otherwise, if $g(z) \neq g(z')$, then $g(z') - g(z) = c \neq 0$ and $\mathcal{I}(z) = \mathcal{I}(z')$ implies $z - z' = cd$, i.e., $d \in \text{lin}(D)$. \[3.2.4\]

**Example 3.2.5** Consider a data dependence $\mathcal{D}D = (D, U, V, I)$ with domain $D = \{(i, j) | i \geq 0, i = j\}$ and integral index mapping $I(i, j) = (i, j) + 2(1, -1)$. The direction of $D$ is $\text{lin}(D) = \{(1, 1)\}$ and $d = (1, -1) \notin \text{lin}(D)$. Hence, according to Proposition 3.2.4, $\mathcal{I}$ is injective over $D$. Some of the corresponding data dependence vectors are sketched in Fig. 3.5. \[3.2.5\]

Note that the condition is only sufficient and there may exist atomic index mappings which are injective regardless of the geometric relation between their generators and the domain. This is illustrated in the following example.
Example 3.2.6  Consider the atomic index mapping $I(i, j) = (i, j) + j(1, 0)$ and the domain $D = \{(i, j) | 1 \leq i, j \leq p\}$, for some $p > 1$. As $D$ is of full dimension in $\mathbb{Z}^2$, then $d \in \text{lin}(D)$. However, $I$ is injective on $D$. In particular $I$ defines an injective mapping on $\mathbb{Z}^2$.

Given an atomic integral index mapping which is injective according to Proposition 3.2.4, we want to define a corresponding inverse mapping. Inverse mappings will be needed in the definition of the decomposition techniques for integral index mappings in Section 3.2.5.

An inverse of an atomic integral index mapping which satisfies the condition of Proposition 3.2.4, can be obtained in a systematic way, based only on geometric properties. This definition is given in Proposition 3.2.7 below, and is based on the following:

- the choice of a hyperplane $[\pi : \theta]$, with $\pi \neq 0$, containing $D$. By definition, $\pi$ has to be a vector orthogonal to all vectors in $D$. Hence $\pi$ has to be chosen in the space $D^\perp$ (the orthogonal complement of $D$ - see Appendix F for a definition). We also require that $\pi$ is in the space generated by $D$ and $d$. This choice will allow us to define a measure of the distance from $D$ of a point in the direction of $d$. Formally, this choice is expressed by the condition $\pi \in \text{lin}(P) \cap D^\perp$ of the proposition, where $P$ is the polyhedron generated by $D$ and $d$ (see also Appendix E). The effect of this choice is illustrated in Fig. 3.6 a) in $\mathbb{Z}^2$, for a 1-dimensional domain $D$. Fig. 3.6 b) illustrates a (non-admissible) choice of $\pi$ outside the polyhedron $P$.

- the scalar product $\eta = \pi \cdot d$, which represents the projection of $d$ along the direction of $\pi$. The integer $\eta$ may be seen as establishing a “distance” between parallel hyperplanes with normal vector $\pi$ intersecting the polyhedron $P$ (see Fig. 3.6 c)).

- a linear transformation $l$, defined so that for all $z \in D$, $l(z) = g(z)$. For each point $z \in D$, $l(z)$ defines the “distance” (as a multiple of $\eta$) between the parallel hyperplanes
**Proposition 3.2.7** Consider an atomic integral index mapping $\mathcal{I}$ and a domain $D \subseteq \mathbb{Z}^n$. such that for all $z \in D$, $\mathcal{I}(z) = z + g(z)d$ and $g(z) \geq 0$. Let $d \notin \text{lin}(D)$ and $P$ the convex polyhedron generated by $D$ and $d$. Consider $\pi \in \text{lin}(P) \cap D^\perp$, with $\pi \neq 0$, and the hyperplane $[\pi : \theta]$ containing the domain $D$.

Then the mapping $\mathcal{I}'(z) = z + l(z)(-d)$, where $l(z) = (\pi \cdot z - \theta)/\eta$ and $\eta = \pi \cdot d$, defines an inverse of $\mathcal{I}$ over $D$.

**Proof:** By definition, for all $z \in D$, $\pi \cdot z = \theta$. Let $z \in D$, then:

$$l(\mathcal{I}(z)) = l(z + g(z)d) = (\pi \cdot (z + g(z)d) - \theta)/\eta$$

$$= (\pi \cdot z + g(z)\pi \cdot d - \theta)/\eta = (\theta + g(z)\eta - \theta)/\eta = g(z).$$

Therefore, for all $z \in D$,

$$I' \circ \mathcal{I}(z) = \mathcal{I}'(\mathcal{I}(z)) = \mathcal{I}(z) + l(\mathcal{I}(z))(-d) = z + g(z)d - g(z)d = z.$$ 

**Example 3.2.8** Let us consider the atomic integral index mapping of Example 3.2.5, and apply Proposition 3.2.7. Let $\pi = (1, -1)$, $\theta = 0$ and $\eta = \pi \cdot d = (1, -1) \cdot (1, -1) = 2$. Then $l(i, j) = (i - j)/2$ and $I'(i, j) = ((i + j)/2, (i + j)/2)$. For instance, $I' \circ I(3, 3) = I'(11, -5) = (6/2, 6/2) = (3, 3)$. 

---

Fig. 3.6. Inverse of an injective atomic integral index mapping: a) admissible choice of $\pi$; b) non-admissible choice of $\pi$; c) intuitive meaning of $\eta$; d) intuitive meaning of $l$.

$[\pi : \theta]$ and $[\pi : \theta + g(z)\eta]$ (see Fig. 3.6 d)).
Proposition 3.2.7 provides a way of defining an inverse for an atomic index mapping (which satisfies the conditions of the proposition) that can be easily mechanised. It may be the case that an inverse for the mapping can be provided by the designer by other means. This happens, in particular, for injective linear mappings, as shown in the following example.

**Example 3.2.9** Consider the index mapping $I(i, j) = (i + j, j)$ of Example 3.2.6. $I$ defines an injective linear mapping in $\mathbb{Z}^n$. In its matrix form, $I$ can be expresses as:

$$I\begin{pmatrix} i \\ j \end{pmatrix} = \begin{bmatrix} 1 & 1 \\ 0 & 1 \end{bmatrix} \cdot \begin{pmatrix} i \\ j \end{pmatrix}$$

An inverse for $I$ can be determine by computing the inverse of the matrix

$$A = \begin{bmatrix} 1 & 1 \\ 0 & 1 \end{bmatrix}$$

that is the matrix

$$A^{-1} = \begin{bmatrix} 1 & -1 \\ 0 & 1 \end{bmatrix}$$

Therefore, the mapping $I'(i, j, k) = (i - j, j)$ defines an inverse of $I$.

From the above discussion, it follows that we may consider two ways of deciding whether an atomic integral index mapping is injective on a certain domain, as well as defining a corresponding inverse: automatically, by considering geometric properties only, or in an *ad hoc* fashion by direct intervention of the designer. The two approaches are complementary and should both be accounted for by the method. In Section 3.2.5 we will take advantage of both for the definition of decomposition techniques.

### 3.2.3 Uniformisation

Given an atomic integral data dependence $DD$, uniformisation defines a substitution of $DD$ with a uniform routing system defined by a set of uniform recurrence equations.

Let $I$ denote the atomic integral index mapping of $DD$ and $g$ its coefficient. Uniformisation can be applied only if $g$ admits an upper bound on the domain $D$, i.e., there exists $m \in \mathbb{N}$ such that for all $z \in D$, $g(z) \leq m$. Technically, it is this condition which allows us to define and initialise the control variables which control the uniform routing of the data. Although this is a restriction to the application of the technique, we can justify it in terms of requirements for a realistic implementation. In fact, if $g$ is not bounded over $D$, the Euclidean distance between pairs of data dependent points increases without a bound, and so does the order of the routing paths which are defined between the two points. Because
of the relation between data dependence graph and data flow graph, this fact results in unbounded resources (either memory or number of processing elements and connections) of any regular array implementation of the specification.

The following propositions define two uniformisation techniques with the guarantee that no problem size dependent overloading of the data dependence graph is generated. The difference between the two techniques is the geometric relation between the direction vector $d$ and the domain $D$ of the data dependence. In particular, if $d \not\in \text{lin}(D)$, because of Proposition 3.2.4, $I$ is injective and $d$ can be used as the direction for pipelining the data. Otherwise, new pipelining directions need to be found outside $\text{lin}(D)$, which guarantee the injectivity of the corresponding index mappings.

Control variables are used in the definition of the routing schemes. The need for control variables comes from the contrasting necessity of obtaining polyhedral convex regions even when non-affine mappings are applied. As, in general, $I$ is not linear, the image of $D$ under $I$ is not a polyhedral convex set. However, polyhedral convex routing domains are enforced, which embed the actual routing paths of the data. The points needed for routing constitute non-convex sub-sets of such domains. The identification of the required points is achieved via conditional expressions based on the integer coefficient $g$.

The content of Proposition 3.2.10 is illustrated in Fig. 3.7, for a 2-dimensional case. The aim is to transform the data dependence graph in Fig. 3.7 a) into that of Fig. 3.7 d). A new domain $D_1$ is defined from $D$ and the maximum value $\tilde{g}$ that the coefficient $g$ assumes on $D$ (see Fig. 3.7 b)). The choice of $\pi$ and the hyperplane $[\pi : \theta]$ containing $D$, already discussed.
in Section 3.2.2, guarantees that no data broadcast is generated. In Fig. 3.7 d) solid arrows correspond to the routing paths which substitute the data dependence vectors of Fig. 3.7 a). The dashed arrows result because of the uniformisation on the enlarged domains and do not contribute to the actual routing. The computations on the routing paths are identified by the evaluation of control variables \( \alpha \) and \( \gamma \). The definition of \( \alpha \) and \( \gamma \) is based on the coefficient \( g \) of the index mapping, and their use is illustrated in Fig. 3.7 c). In the figure, three points of the domain \( D_1 \) are emphasised: point \( z \) needs to receive the data from \( z + g(z)d \); variables \( \alpha \) and \( \gamma \) are initialised at \( z + \tilde{g}d \). Note that the three points lie on the line \( z + ld \), for \( l \in \mathbb{R} \). At \( z + \tilde{g}d \), variable \( \alpha \) is initialised to the value \( g(z) \) and \( \gamma \) to \( \tilde{g} \). Hence, initially, \( \gamma \geq \alpha \). The tags \( \alpha \) and \( \gamma \) are then pipelined according to the direction of \( -d \). At each step, the value of \( \gamma \) is decreased by 1. Also, at each step the values of \( \alpha \) and \( \gamma \) are compared, and when they become equal (i.e., both equal to \( g(z) \)), the data is collected by a routing variable \( R \) and subsequently pipelined to \( z \) along \( -d \).

**Proposition 3.2.10 [Uniformisation 1]** Let \( DD = (D, U, V, I) \) be an atomic integral data dependence, with \( I(z) = z + g(z)d \). Let \( g \) be non-negative and bounded over \( D \), and \( \tilde{g} \) the least upper bound of \( g \). Let \( d \notin \text{lin}(D) \) and \( P \) be the convex polyhedron generated by \( D \) and \( d \). Consider \( \pi \in \text{lin}(P) \cap D^\perp \), with \( \pi \neq 0 \), and the hyperplane \( [\pi : \theta] \) containing the domain \( D \).

Then \( DD \) can be substituted by the uniform data dependence

\[
DD' = (D, U, R, I_0)
\]

and the system of equations:

\[\begin{align*}
E_1 &= (D_1, R, (R, V, \alpha, \gamma), f, (I_1, I_0, I_0, I_0)) \\
E_2 &= (D_{1,1}, \alpha, \alpha, id, I_1) \\
E_3 &= (D_{1,2}, \alpha, in_\alpha) \\
E_4 &= (D_{1,1}, \gamma, \gamma, \text{dec}, I_1) \\
E_5 &= (D_{1,2}, \gamma, in_\gamma)
\end{align*}\]

where:

- the index mappings are:

\[
\begin{align*}
I_0(z) &= z \\
I_1(z) &= z + d
\end{align*}
\]
- $R, \alpha$ and $\gamma$ are new variables;

- the applied functions are:

\[
\begin{align*}
in_\alpha(z) & = g(z - \bar{g}d) \\
in_\gamma(z) & = \bar{g} \\
id(a) & = a \\
dec(a) & = a - 1 \\
f(a, b, c, d) & = \begin{cases} 
  a & c \neq d \\
  b & \text{otherwise}
\end{cases}
\end{align*}
\]

- the new domains are:

\[
\begin{align*}
D_1 & = \{z + ld \mid z \in D, 0 \leq l \leq \bar{g}\} \\
D_{1,1} & = \{z + ld \mid z \in D, 0 \leq l < \bar{g}\} \\
D_{1,2} & = \{z + \bar{g}d \mid z \in D\}
\end{align*}
\]

**Proof:** For all $z \in D$, let $\text{segm}(z) = \{z + ld \mid 0 \leq l \leq \bar{g}\}$. By definition, for all $z' \in \text{segm}(z)$, $\alpha(z') = g(z)$. In fact, for $z \in D$,

\[
\alpha(z) = \alpha(z + d)
\]

\[
= \ldots = \alpha(z + \bar{g}d) = g(z + \bar{g}d - \bar{g}d) = g(z).
\]

Also for all $z' = z + ld \in \text{segm}(z)$, $\gamma(z') = l$. In fact, for $z \in D$,

\[
\gamma(z) = \gamma(z + d) - 1
\]

\[
= \ldots = \gamma(z + \bar{g}d) - \bar{g} = \bar{g} - \bar{g} = 0.
\]

Therefore, for all $z' = z + ld \in \text{segm}(z)$, $\alpha(z') = \gamma(z')$ if and only if $l = g(z)$.

Hence, for all $z \in D$,

\[
U(z) = R(\mathcal{I}_0(z)) = R(\mathcal{I}_1 \circ \mathcal{I}_0(z))
\]

\[
= \ldots = R(\mathcal{I}_1^{g(z)} \circ \mathcal{I}_0(z)) = V(\mathcal{I}_0 \circ \mathcal{I}_1^{g(z)} \circ \mathcal{I}_0(z))
\]

\[
= V(z + g(z)d) = V(\mathcal{I}(z)).
\]

\[\blacksquare\]
Corollary 3.2.11 Let $DD = (D, U, V, I)$ be an atomic integral data dependence, with $I(x) = x + g(x)d$. Let $g$ be non-negative and bounded over $D$, and $\tilde{g}$ the least upper bound of $g$.

If $\tilde{g} = 0$, then $DD$ can be substituted by the uniform data dependence

$$DD' = (D, U, V, I_0)$$

where $I_0(x) = x$.  

Example 3.2.12 Consider the atomic integral data dependence $DD = (D, U, V, I)$ with index mapping $I(i, j) = (i, j) + g(i, j)(1, 0)$, where $g(i, j) = (i^2 + j^2)mod 6$, and domain $D = \{(i, j) \mid 1 \leq j \leq 5, i = 11 - 2j\}$. $lin(D) = \langle(2, -1)\rangle$ and $d = (1, 0) \not\in lin(D)$. $DD$ is illustrated in Fig. 3.8 a).

The space generated by $D$ and $d$ is the whole $Z^2$, and $\pi$ can be any vector in $D^1$. For instance, the vector $\pi = (1, 2)$ (which is orthogonal to the generator $(2, -1)$ of $lin(D)$) satisfies the conditions of Proposition 3.2.10, and $[(1, 2) : 11]$ is a hyperplane containing $D$. Also, for all $(i, j) \in D$, $g(i, j) \leq 5$. If we apply Proposition 3.2.10 we obtain the new data dependence

$$DD' = (D, U, R, I_0)$$

and the system of equations:

$$E_1 = (D_1, R, (R, V, \alpha, \gamma), f, (I_1, I_0, I_0, I_0))$$
$$E_2 = (D_{11}, \alpha, \alpha, id, I_1)$$
$$E_3 = (D_{12}, \alpha, in_{\alpha})$$
$$E_4 = (D_{11}, \gamma, \gamma, dec, I_1)$$
$$E_5 = (D_{12}, \gamma, in_{\gamma})$$

where:

Fig. 3.8. Uniformisation: a) domain $D$ and corresponding data dependence vectors; b) domain $D_1$ and data dependence vectors after uniformisation.
- the index mappings are:

\[
\begin{align*}
I_0(i, j) &= (i, j) \\
I_1(i, j) &= (i, j) + (1, 0)
\end{align*}
\]

- the applied functions are:

\[
\begin{align*}
in_\alpha(i, j) &= g(i - 5, j) \\
in_{\gamma}(z) &= z \\
id(a) &= a \\
dec(a) &= a - 1 \\
f(a, b, c, d) &= \begin{cases} a & \text{if } c \neq d \\ b & \text{otherwise} \end{cases}
\end{align*}
\]

- the new domains are:

\[
\begin{align*}
D_1 &= \{(i, j) \mid 1 \leq j \leq 5, 11 - 2j \leq i \leq 16 - 2j\} \\
D_{1,1} &= \{(i, j) \in D_1 \mid i + 2j < 16\} \\
D_{1,2} &= \{(i, j) \in D_1 \mid i + 2j = 16\}
\end{align*}
\]

The resulting data dependence graph is illustrated in Fig. 3.8 b). The correct routing of the data is achieved by evaluating the control variables $\alpha$ and $\gamma$. For simplicity, in the figure, the propagation of the control variables has been omitted.

The second uniformisation technique applies when the generator $d$ of the index mapping $I$ cannot be used as a regularisation direction. Fig. 3.9 illustrates the result in a 3-dimensional space. In this case we assume that $d$ is contained in $\text{lin}(D)$ (in Fig. 3.9 a), $\text{lin}(D)$ is the space generated by $(1,0,0)$ and $(0,1,0))$. A vector $\tau$ is chosen in $D_1^\perp$ (in the figure, $\tau = (0,0,1)$). Three regularisation vectors are involved: $d$ itself, together with $\bar{d} = d + \pi$ and $\bar{d} = d - \pi$ (see Fig. 3.9 d)). Also, three control variables, $\alpha, \beta$ and $\gamma$, are used. The effect of uniformisation is illustrated in Fig. 3.9 b) and c) for $g(z)$ even and odd, respectively. The data dependence vector between $z$ and $z + g(z)d$ is replaced by a routing path of order $g(z)$. This path is the result of:

- a sub-path of order $\lfloor g(z)/2 \rfloor$ according to the direction of $\bar{d}$. On this sub-path the data is pipelined by a routing variable $R^2$ (R2 in the figure):
- a sub-path of order \( g(z) \mod 2 \) according to the direction of \( d \). On this sub-path the data is pipelined by a routing variable \( R_1 \) (R1 in the figure). This is necessary only if \( g(z) \) is odd, as in Fig. 3.9 c);

- a sub-path of order \( \lfloor g(z)/2 \rfloor \) according to the direction of \( \bar{d} \). On this sub-path the data is also pipelined by the routing variable \( R_1 \).

The control variables \( \alpha, \beta \) and \( \gamma \) are initialised at \( z + \bar{g} \bar{d} \), where \( \bar{g} = \lfloor m/2 \rfloor \) and \( m \) is the least upper bound of \( g \) on \( D \). Variable \( \alpha \) is initialised to \( \lfloor g(z)/2 \rfloor \), \( \gamma \) to \( \bar{g} \), and \( \beta \) to \( g(z) \mod 2 \).

The values of \( \alpha, \beta \) and \( \gamma \) are pipelined according to the direction of \( -\bar{d} \), with the value of \( \gamma \) decreased by one at each step. Also, the values of \( \alpha \) and \( \gamma \) are compared at each step; when they become equal (i.e., both equal to \( \lfloor g(z)/2 \rfloor \)), the value of \( \beta \) is considered to determine whether \( g(z) \) is either even or odd, and the data is transferred accordingly from \( R_2 \) to \( R_1 \).

**Proposition 3.2.13 [Uniformisation 2]** Let \( DD = (D, U, V, I) \) be an atomic finitely generated dynamic data dependence, with \( I(z) = z + g(z)d \). Let \( g \) be non-negative and bounded over \( D \), \( m \) the least upper bound of \( g \) and \( \bar{g} = \lfloor m/2 \rfloor \). Let \( d \in \text{lin}(D) \) and \( \text{dim}(D) < n \).

Consider \( \pi \in D^\perp \), with \( \pi \neq 0 \), the hyperplane \( [\pi : \theta] \) containing the domain \( D \). and let \( \bar{d} = d + \pi \) and \( \bar{d} = d - \pi \).
Then $DD$ can be substituted by the uniform data dependence

$$DD' = (D, U, R^1, I_0)$$

and the system of equations:

$$E_1 = (D_1, R^1, (R^1, R^2, R^2, \alpha, \beta, \gamma), f, (I_{1,0}, I_{2,1}, I_{0}, I_{0}, I_{0}))$$

$$E_2 = (D_{2,1}, R^2, R^2, id, I_3)$$

$$E_3 = (D_{2,2}, R^2, V, id, I_0)$$

$$E_4 = (D_{1,1}, \alpha, \alpha, id, I_1)$$

$$E_5 = (D_{1,2}, \alpha, in_\alpha)$$

$$E_6 = (D_{1,1}, \beta, \beta, id, I_1)$$

$$E_7 = (D_{1,2}, \beta, in_\beta)$$

$$E_8 = (D_{1,1}, \gamma, \gamma, dec, I_1)$$

$$E_9 = (D_{1,2}, \gamma, in_\gamma)$$

where:

- the index mappings are:

  $$I_0(z) = z$$

  $$I_1(z) = z + \hat{d}$$

  $$I_{2,0}(z) = z$$

  $$I_{2,1}(z) = z + d$$

  $$I_3(z) = z + \hat{d}$$

- $R^1, R^2, \alpha, \beta$ and $\gamma$ are new variables;

- the applied functions are:

  $$in_\alpha(z) = \lfloor g(z - \hat{g}\hat{d})/2 \rfloor$$

  $$in_\beta(z) = g(z - \hat{g}\hat{d}) \mod 2$$

  $$in_\gamma(z) = \hat{g}$$

  $$id(a) = a$$

  $$dec(a) = a - 1$$

  $$f(a, b, c, d, e, f) = \begin{cases} 
    a & d \neq f \\
    b & d = f, e = 0 \\
    c & d = f, e = 1
  \end{cases}$$
the new domains are:

\[
D_1 = \{ z + l \hat{d} \mid z \in D, 0 \leq l \leq \bar{g} \}
\]

\[
D_{1,1} = \{ z + l \hat{d} \mid z \in D, 0 \leq l < \bar{g} \}
\]

\[
D_{1,2} = \{ z + \bar{g} \hat{d} \mid z \in D \}
\]

\[
D_2 = \{ z + l_1 d + l_2 \hat{d} \mid z \in D_1, 0 \leq l_1 \leq 1, 0 \leq l_2 \leq \bar{g} \} \cap \{ z \in \mathbb{Z^n} \mid \pi \cdot z \geq \theta \}
\]

\[
D_{2,1} = \{ z \in D_2 \mid \pi \cdot z > \theta \}
\]

\[
D_{2,2} = \{ z \in D_2 \mid \pi \cdot z = \theta \}.
\]

**Proof:** For all \( z \in D \), let \( \text{segm}(z) = \{ z + ld \mid 0 \leq l \leq \bar{g} \} \). By definition, for all \( z' \in \text{segm}(z) \),

\[
\alpha(z') = \lfloor g(z)/2 \rfloor.
\]

In fact, for \( z \in D \),

\[
\alpha(z) = \alpha(z + \hat{d}) = \ldots = \alpha(z + \bar{g} \hat{d}) = \lfloor g(z + \bar{g} \hat{d} - \bar{g} \hat{d})/2 \rfloor = \lfloor g(z)/2 \rfloor.
\]

Also, for all \( z' \in \text{segm}(z) \), \( \beta(z') = g(z) \mod 2 \). In fact, for \( z \in D \),

\[
\beta(z) = \beta(z + \hat{d}) = \ldots = \beta(z + \bar{g} \hat{d}) = g(z + \bar{g} \hat{d} - \bar{g} \hat{d}) \mod 2 = g(z) \mod 2.
\]

Finally, for all \( z' = z + l \hat{d} \in \text{segm}(z) \), \( \gamma(z') = l \). In fact, for \( z \in D \),

\[
\gamma(z) = \gamma(z + \hat{d}) - 1 = \ldots = \gamma(z + \bar{g} \hat{d}) - \bar{g} = \bar{g} - \bar{g} = 0.
\]

Therefore, for all \( z' = z + ld \in \text{segm}(z) \), \( \alpha(z') = \gamma(z') \) if and only if \( l = \lfloor g(z)/2 \rfloor \).

We observe that, for all \( c \in \mathbb{Z} \),

\[
\lfloor c/2 \rfloor = (c - c \mod 2)/2
\]

\[
c = 2\lfloor c/2 \rfloor + c \mod 2.
\]

Hence, for all \( z \in D \),

\[
U(z) = R^1(I_0(z)) = R^1(I_1 \circ I_0(z))
\]

\[
= \ldots = R^1(I_1^{[g(z)/2]} \circ I_0(z))
\]

\[
= R^2(I_2 \circ I_0(z) \mod 2 \circ I_1^{[g(z)/2]} \circ I_0(z))
\]

\[
= R^2(I_3 \circ I_2 \circ I_0(z) \mod 2 \circ I_1^{[g(z)/2]} \circ I_0(z)).
\]
Because of the condition $\dim(D) < n$, the application of Proposition 3.2.13 may require an increase in the number of dimensions of the lattice space and to reindex the data dependence accordingly. Such a reindexing is a particular case of normalisation as described in Section 2.2.2 and is realised by "padding" the index expressions with extra indices. Note that this normalisation should not alter the dimensionality of the domains (it simply produces an embedding of the domains in a space of higher dimension).

**Corollary 3.2.14** Let $\mathcal{DD} = (D, U, V, I)$ be an atomic integral data dependence, with $I(z) = z + g(z)d$. Let $g$ be non-negative and bounded over $D$, $m$ the least upper bound of $g$, and $\bar{g} = \lfloor m/2 \rfloor$.

If $\bar{g} = 0$ then $\mathcal{DD}$ can be substituted by the uniform data dependence:

$$\mathcal{DD}' = (D, U, R, I_0)$$

and the equations:

$$E_1 = (D, R, (V, V, \beta), f, (I_0, I_1, I_0))$$

$$E_2 = (D, \beta, in_\beta)$$

where:

- the index mappings are:

$$I_0(z) = z$$

$$I_1(z) = z + d$$

- $R$ and $\beta$ are new variables;
Fig. 3.10. Uniformisation: a) domain $D$ in $\mathbb{Z}^2$ and corresponding data dependence vectors; b) domain $D$ in $\mathbb{Z}^3$; c) domain $D_1$; d) domain $D_2$.

- the applied functions are:

$$
in_g(z) = g(z) \mod 2$$
$$f(a,b,c) = \begin{cases} 
a & c = 0 
b & c = 1 \end{cases}
$$

Example 3.2.15 Consider the atomic integral data dependence $\mathcal{DD} = (D,U,V,I)$ with index mapping $I(i,j) = (i,j) + g(i,j)(1,0)$, where $g(i,j) = (i^2 + j^2) \mod 6$, and domain $D = \{(i,j) | 1 \leq j \leq 5, 11 - 2j \leq i \leq 16 - 2j\}$. The domain $D$ is illustrated in Fig. 3.10 a). $\text{lin}(D) = \mathbb{Z}^2$ and $d = (1,0) \in \text{lin}(D)$. As $D$ is of full dimension in $\mathbb{Z}^2$, we need to reindex the data dependence in $\mathbb{Z}^3$, before applying Proposition 3.2.13 (Fig. 3.10 b) illustrates $D$ in $\mathbb{Z}^3$). Any non-null vector in $D^\perp$ can be chosen as $\pi$. For instance, vector $\pi = (0,0,1)$ satisfies the conditions of Proposition 3.2.13 and $[(0,0,1):0]$ is a hyperplane containing $D$. Let $\tilde{d} = d + \pi = (1,0,1)$ and $\tilde{d} = d - \pi = (1,0,-1)$. For all $(i,j,k) \in D$, $g(i,j,k) \leq 5$, hence we can assume $\bar{g} = 2$. By applying Proposition 3.2.13, we obtain the data dependence

$$\mathcal{DD}' = (D,U,R^1,I_0)$$

and the system of equations:

$$E_1 = (D_1,R^1,(R^1,R^2,R^2,\alpha,\beta,\gamma),f,(I_1,I_{2,0},I_{2,1},I_0,I_0))$$
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\[ E_2 = (D_{2,1}, R_1, R_2, id, I_3) \]
\[ E_3 = (D_{2,2}, R_1, V, id, I_0) \]
\[ E_4 = (D_1, \alpha, \alpha, id, I_1) \]
\[ E_5 = (D_1, \alpha, in_\alpha) \]
\[ E_6 = (D_1, 2, R_1, id, I_1) \]
\[ E_7 = (D_{1,2}, \beta, V, id, I_1) \]
\[ E_8 = (D_1, \gamma, \gamma, dec, I_1) \]
\[ E_9 = (D_{1,2}, \gamma, in_\gamma) \]

where:

- the index mappings are:

\[ I_0(i,j,k) = (i,j,k) \]
\[ I_1(i,j,k) = (i,j,k) + (1,0,1) \]
\[ I_2,0(i,j,k) = (i,j,k) \]
\[ I_2,1(i,j,k) = (i,j,k) + (1,0,0) \]
\[ I_3(i,j,k) = (i,j,k) + (1,0,-1) \]

- the applied functions are:

\[ in_\alpha(i,j,k) = \lfloor g(i - 2, j, k - 2)/2 \rfloor \]
\[ in_\beta(i,j,k) = g(i - 2, j, k - 2) mod 2 \]
\[ in_\gamma(i,j,k) = 2 \]
\[ id(a) = a \]
\[ dec(a) = a - 1 \]
\[ f(a,b,c,d,e,f) = \begin{cases} 
  a & d \neq f \\
  b & d = f, e = 0 \\
  c & d = f, e = 1 
\end{cases} \]

- the new domains are:

\[ D_1 = \{(i,j,k) \mid 1 \leq j \leq 5, 0 \leq k \leq 2, 11 + 2j + k \leq i \leq 16 - 2j + k \} \]
\[ D_{1,1} = \{(i,j,k) \in D_1 \mid k < 2 \} \]
\[ D_{1,2} = \{(i,j,k) \in D_1 \mid k = 2 \} \]
\[ D_2 = \{(i,j,k) \mid 1 \leq j \leq 5, 0 \leq k \leq 2, 11 + 2j + k \leq i \leq 21 - 2j - k \} \]
The routing domain $D_1$ and $D_2$ are sketched in Fig. 3.10 c) and d), respectively, while sections of the data dependence graph in $D_1$ and $D_2$ are given in Fig. 3.11 a) and b), respectively. As shown in the figure, the data dependence graph is uniform and its nodes are locally connected.

### 3.2.4 Parametric Uniformisation

While uniformisation according to Proposition 3.2.10 defines data pipelining in the existing computation space of the problem, the technique defined by Proposition 3.2.13, in general, adds new dimensions to the specification and artificially introduces new domains for the sole purpose of routing the data. Because of the correspondence between computation points and processing elements under space-time mappings, a correspondent increase in both latency and number of processors will characterise the resulting array design. In this section we introduce a parametric version of the technique which allows us to control the amount of routing overhead to a certain extent.

The size of the routing domains defined through the uniformisation technique depends on the quantity $\bar{g} = \lfloor m/2 \rfloor$, where $m$ is the least upper bound of the values of the coefficient $g$ on the domain $D$. In this section we show that by allowing a limited amount of overloading of the data dependence graph, the size of the routing domains can be reduced. In particular, we introduce a parameter $p$ as a small positive integer, in order to represent an upper bound to the amount of overloading allowed in the specification. Then we define new routing domains whose size depends on the quantity $\bar{g} = \lfloor m/(p+1) \rfloor$, and new routing paths in these smaller domains. For each $z$ of the domain, the order of the corresponding routing path remains equal to $g(z)$. However, the shape changes according to the parameter $p$. Fig. 3.12 may
help to illustrate how the technique works. Fig. 3.12 a) and b) illustrate the routing path (for \( g(z) \) odd and even, respectively) corresponding to the data dependence vector between \( z \) and \( z + g(z)d \), as defined in the previous section. For the same data dependence vector, Fig. 3.12 c) and d) and Fig. 3.12 f) and g) illustrate the corresponding routing paths for values of the parameter \( p = 2 \) and \( p = 3 \), respectively. Note that as \( p \) increases, \( \bar{g} \) decreases, and the hyperplanes \([\pi : \theta]\) and \([\pi : \bar{g}\eta + \theta]\) become closer together. Control variables \( \alpha, \beta \) and \( \gamma \) are still initialised at the point \( z + \bar{g}d \). However, their initialisation values become \([g(z)/(p+1)], g(z) \mod (p+1)\) and \( \bar{g} \), respectively. As before, the control variables are pipelined according to the direction of \(-\hat{d}\). At each step the value of \( \gamma \) is decreased by one, and \( \alpha \) and \( \gamma \) are compared. When their values are equal, the value of \( \beta \) determines at which lattice point the data has to be transferred between \( R^2 \) and \( R^1 \). Note that three variables \( R^1, R^2 \) and \( R^3 \) are used to route the data. The figure illustrates the correspondence between the routing variables and the routing directions \( d, \hat{d} \) and \( \bar{d} \).

**Proposition 3.2.16 [Parametric Uniformisation]** Let \( DD = (D, U, V, I) \) be an atomic integral data dependence, with \( I(z) = z + g(z)d \). Let \( g \) be non-negative and bounded over \( D \), and \( m \) the least upper bound of \( g \). Let \( p \in \mathbb{N}^+ \) and \( \bar{g} = [m/(p+1)] \). Let \( d \in lin(D) \) and \( dim(D) < n \). Consider \( \pi \in D^\perp \), with \( \pi \neq 0 \), the hyperplane \([\pi : \theta]\) containing the domain \( D \). and let \( \hat{d} = d + \pi \) and \( \bar{d} = d - \pi \).
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Then $\mathcal{DD}$ can be substituted by the uniform data dependence

$$\mathcal{DD}' = (D, U, R^1, I_0)$$

and the system of equations:

$$E_1 = (D_1, R^1, (R^1, R^2, \ldots, R^2, \alpha, \beta, \gamma), f, (I_1, I_{2,0}, \ldots, I_{2,p}, I_0, I_0, I_0))$$

$$E_2 = (D_{2,1}, R^2, R_3, id, I_3)$$

$$E_3 = (D_{2,2}, R^2, V, id, I_0)$$

$$E_4 = (D_{2,3}, R^3, R^2, id, I_4)$$

$$E_5 = (D_{1,1}, \alpha, \alpha, id, I_1)$$

$$E_6 = (D_{1,2}, \alpha, in_\alpha)$$

$$E_7 = (D_{1,1}, \beta, \beta, id, I_1)$$

$$E_8 = (D_{1,2}, \beta, in_\beta)$$

$$E_9 = (D_{1,1}, \gamma, \gamma, dec, I_1)$$

$$E_{10} = (D_{1,2}, \gamma, in_\gamma)$$

where:

- the index mappings are:

  $$I_0(z) = z$$

  $$I_1(z) = z + \hat{d}$$

  $$I_{2,0}(z) = z$$

  $$I_{2,1}(z) = z + d$$

  ...\n
  $$I_{2,p}(z) = z + pd$$

  $$I_3(z) = z + \hat{d}$$

  $$I_4(z) = z + (p-1)d$$

- $R^1, R^2, R^3, \alpha, \beta$ and $\gamma$ are new variables;

- the applied functions are:

  $$in_\alpha(z) = |g(z - \hat{g}\hat{d})/(p + 1)|$$

  $$in_\beta(z) = g(z - \hat{g}\hat{d}) \mod (p + 1)$$
\[ \text{in}_\gamma(z) = \tilde{g} \]
\[ \text{id}(a) = a \]
\[ \text{dec}(a) = a - 1 \]
\[
\begin{cases} 
  a & c \neq e \\
  b_0 & c = e, d = 0 \\
  \ldots & \\
  b_p & c = e, d = p
\end{cases}
\]

- the new domains are:

\[
\begin{align*}
  D_1 &= \{ z + l \hat{d} \mid z \in D, 0 \leq l \leq \tilde{g} \} \\
  D_{1,1} &= \{ z + l \hat{d} \mid z \in D, 0 \leq l < \tilde{g} \} \\
  D_{1,2} &= \{ z + \tilde{g} \hat{d} \mid z \in D \} \\
  D_2 &= \{ z + (l_1 + l_2)d + l_3 \hat{d} \mid z \in D, 0 \leq l_1 \leq p, 0 \leq l_2 \leq (p - 1)\tilde{g}, 0 \leq l_3 \leq \tilde{g} \} \\
  \cap \{ z \in \mathbb{Z}^n \mid \pi \cdot z \geq \theta \} \\
  D_{2,1} &= \{ z \in D_2 \mid \pi \cdot z > \theta \} \\
  D_{2,2} &= \{ z \in D_2 \mid \pi \cdot z = \theta \} \\
  D_{2,3} &= \{ z \in D_2 \mid \pi \cdot z < \theta + \tilde{g} \eta \}.
\end{align*}
\]

**Proof:** For all \( z \in D \), let \( \text{segm}(z) = \{ z + l \hat{d} \mid 0 \leq l \leq \tilde{g} \} \). By definition, for all \( z' \in \text{segm}(z) \),

\[ \alpha(z') = \lfloor g(z)/(p+1) \rfloor. \]

In fact, for \( z \in D \),

\[
\begin{align*}
  \alpha(z) &= \alpha(z + \hat{d}) = \ldots = \alpha(z + \tilde{g} \hat{d}) \\
  &= \lfloor g(z + \tilde{g} \hat{d} - \tilde{g} \hat{d})/(p+1) \rfloor = \lfloor g(z)/(p+1) \rfloor.
\end{align*}
\]

Also, for all \( z' = z + l \hat{d} \) in the segment \( \text{segm}(z) \), with \( z \in D \), \( \beta(z') = g(z) \mod (p+1) \).

In fact, for \( z \in D \),

\[
\begin{align*}
  \beta(z) &= \beta(z + \hat{d}) = \ldots = \beta(z + \tilde{g} \hat{d}) \\
  &= g(z + \tilde{g} \hat{d} - \tilde{g} \hat{d}) \mod (p+1) = g(z) \mod (p+1).
\end{align*}
\]

Finally, for all \( z' = z + l \hat{d} \in \text{segm}(z) \), \( \gamma(z') = l \). In fact, for \( z \in D \),

\[
\begin{align*}
  \gamma(z) &= \gamma(z + \hat{d}) - 1 \\
  &= \ldots = \gamma(z + \tilde{g} \hat{d}) - \tilde{g} = \tilde{g} - \tilde{g} = 0.
\end{align*}
\]

Therefore, for all \( z' = z + l \hat{d} \in \text{segm}(z) \), \( \alpha(z') = \gamma(z') \) if and only if \( l = \lfloor g(z)/(p+1) \rfloor \).
We observe that, for all \( c \in \mathbb{Z} \),
\[
\left\lfloor \frac{c}{p+1} \right\rfloor = \frac{c - c \mod (p+1)}{p+1}
\]
\[
c = (p+1)\left\lfloor \frac{c}{p+1} \right\rfloor + c \mod (p+1).
\]

Hence, for all \( z \in D \),
\[
U(z) = R^1(I_0(z)) = R^1(I_1 \circ I_0(z))
\]
\[
= \ldots = R^1(I_1^{[g(z)/(p+1)]} \circ I_0(z))
\]
\[
= R^2(I_{2, (g(z) \mod (p+1))} \circ I_1^{[g(z)/(p+1)]} \circ I_0(z))
\]
\[
= R^3(I_3 \circ I_{2, (g(z) \mod (p+1))} \circ I_1^{[g(z)/(p+1)]} \circ I_0(z))
\]
\[
= R^2(I_4 \circ I_3 \circ I_{2, (g(z) \mod (p+1))} \circ I_1^{[g(z)/(p+1)]} \circ I_0(z))
\]
\[
= \ldots
\]
\[
= R^2(I_4^{[g(z)/(p+1)]} \circ I_3^{[g(z)/(p+1)]} \circ I_{2, (g(z) \mod (p+1))} \circ I_1^{[g(z)/(p+1)]} \circ I_0(z))
\]
\[
= V(I_0 \circ I_{4}^{[g(z)/(p+1)]} \circ I_{3}^{[g(z)/(p+1)]} \circ I_{2, (g(z) \mod (p+1))} \circ I_1^{[g(z)/(p+1)]} \circ I_0(z))
\]
\[
= V(z + [g(z)/(p+1)]d + (g(z) \mod (p+1))d + [g(z)/(p+1)]d +
\]
\[
[g(z)/(p+1)](p-1)d
\]
\[
= V(z + (p+1)[g(z)/(p+1)] + g(z) \mod (p+1))d)
\]
\[
= V(z + g(z)d) = V(I(z)).
\]

\[ \blacksquare \]

**Corollary 3.2.17** Let \( \mathcal{DD} = (D, U, V, I) \) be an integral data dependence, with \( I(z) = z + g(z)d \). Let \( g \) be non-negative and bounded over \( D \), and \( m \) the least upper bound of \( g \).

Let \( p \in \mathbb{N}^+ \) and \( \tilde{g} = \lceil m/(p+1) \rceil \).

If \( \tilde{g} = 0 \) then \( \mathcal{DD} \) can be substituted by the uniform data dependence:

\[
\mathcal{DD}' = (D, U, R, I_0)
\]

and the equations:

\[
E_1 = (D, R, (V, \ldots, V, \beta). f, (I_{1,0}, \ldots, I_{1,p}, I_0))
\]
\[
E_2 = (D, \beta, \text{in}_\beta)
\]

where:
- the index mappings are:

\[
\begin{align*}
I_0(z) &= z \\
I_{1,0}(z) &= z \\
I_{1,1}(z) &= z + d \\
\ldots \\
I_{1,p}(z) &= z + pd
\end{align*}
\]

- \( R \) and \( \beta \) are new variables;

- the applied functions are:

\[
\begin{align*}
in_\beta(z) &= g(z) \mod (p + 1) \\
f(a_0, \ldots, a_p, b) &= \begin{cases} 
  a_0 & b = 0 \\
  a_1 & b = 1 \\
  \ldots & \\
  a_p & b = p 
\end{cases}
\]

Note that uniformisation according to Proposition 3.2.13 corresponds to parametric uniformisation with parameter \( p = 1 \).

Example 3.2.18 Let us apply parametric uniformisation to the atomic integral data dependence of Example 3.2.15, with the same choice of routing directions, i.e., \( d = (1,0,0) \), \( \hat{d} = (1,0,1) \) and \( \dd = (1,0,-1) \).

Let \( \bar{g} = [5/(p+1)] \), with \( p \in \mathbb{N}^+ \). The application of parametric uniformisation according to Proposition 3.2.16 yields the data dependence

\[
\mathcal{DD}' = (D, U, R^1, I_0)
\]

and the system of equations:

\[
\begin{align*}
\mathbf{E}_1 &= (D_1, R^1, (R^1, R^2, \ldots, R^2, \alpha, \beta, \gamma), f, (I_{1,0}, I_{2,0}, \ldots, I_{2,p}, I_0, I_0, I_0)) \\
\mathbf{E}_2 &= (D_{2,1}, R^2, R^3, id, I_3) \\
\mathbf{E}_3 &= (D_{2,2}, R^2, V, id, I_0) \\
\mathbf{E}_4 &= (D_{2,3}, R^3, R^2, id, I_4) \\
\mathbf{E}_5 &= (D_{1,1}, \alpha, \alpha, id, I_1) \\
\mathbf{E}_6 &= (D_{1,2}, \alpha, in_\alpha)
\end{align*}
\]
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\[ \mathbf{E}_7 = (D_{1,1}, \beta, \beta, \text{id}, I_1) \]
\[ \mathbf{E}_8 = (D_{1,2}, \beta, \text{in}_\beta) \]
\[ \mathbf{E}_9 = (D_{1,1}, \gamma, \gamma, \text{dec}, I_1) \]
\[ \mathbf{E}_{10} = (D_{1,2}, \gamma, \text{in}_\gamma) \]

where:

- the index mappings are:
  \[ I_0(i, j, k) = (i, j, k) \]
  \[ I_3(i, j, k) = (i, j, k) + (1, 0, 1) \]
  \[ I_{2,0}(i, j, k) = (i, j, k) \]
  \[ I_{2,1}(i, j, k) = (i, j, k) + (1, 0, 0) \]
  \[ \ldots \]
  \[ I_{2,p}(i, j, k) = (i, j, k) + (p, 0, 0) \]
  \[ I_3(i, j, k) = (i, j, k) + (1, 0, -1) \]
  \[ I_4(i, j, k) = (i, j, k) + (p - 1, 0, 0) \]

- the applied functions are:
  \[ \text{in}_\alpha(i, j, k) = \lfloor g(i - \bar{g}, j, k - \bar{g})/(p + 1) \rfloor \]
  \[ \text{in}_\beta(i, j, k) = g(i - \bar{g}, j, k - \bar{g}) \mod (p + 1) \]
  \[ \text{in}_\gamma(i, j, k) = \bar{g} \]
  \[ \text{id}(a) = a \]
  \[ \text{dec}(a) = a - 1 \]
  \[ f(a, b_0, \ldots, b_p, c, d, e) = \begin{cases} a & c \neq e \\ b_0 & c = e, d = 0 \\ \ldots & c = e, d = p \end{cases} \]

- the new domains are:
  \[ D_1 = \{(i, j, k) | 1 \leq j \leq 5, 0 \leq k \leq \bar{g}, 11 - 2j + k \leq i \leq 16 - 2j + k\} \]
  \[ D_{1,1} = \{(i, j, k) \in D_1 | k < \bar{g}\} \]
  \[ D_{1,2} = \{(i, j, k) \in D_1 | k = \bar{g}\} \]
  \[ D_2 = \{(i, j, k) | 1 \leq j \leq 5, 0 \leq k \leq \bar{g}, 11 - 2j + k \leq i \leq 21 - 2j - k\} \]
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Fig. 3.13. Data dependence vectors after parametric uniformisation: a) and b) sections of $D_1$ and $D_2$ for $p = 2$; c) and d) the same sections for $p = 3$.

\begin{align*}
D_{2,1} &= \{(i,j,k) \in D_2 \mid k > 0\} \\
D_{2,2} &= \{(i,j,k) \in D_2 \mid k = 0\} \\
D_{2,3} &= \{(i,j,k) \in D_2 \mid k < \bar{g}\}.
\end{align*}

The routing domains $D_1$ and $D_2$ are similar to those sketched in Fig. 3.10 (as the same regularisation vectors are considered). A section of the data dependence graph in $D_1$ and $D_2$ is given in Fig. 3.13 a), b) and Fig. 3.13 c), d) for values of the parameter $p = 2$ and $p = 3$, respectively.

3.2.5 Decomposition

While uniformisation replaces an atomic integral data dependence with a system of uniform recurrences, decomposition allows us to substitute an integral data dependence with a set of atomic integral data dependencies. This is obtained by decomposing all data dependence vectors into their components along the generators of the integral dependence mapping.

Given an integral data dependence $\mathcal{D} = (D, U, V, I)$, whose integral index mapping $I$ has $m$ generators, where $m > 1$, the application of a decomposition technique aims at generating a set of integral data dependencies such that:

- each of the new index mappings is integral with less than $m$ generators; and

- the composition of the new index mappings is equal to $I$ for each point of the domain $D$. 

\[ \blacksquare \]
A recursive application of the techniques yields a system of atomic integral data dependencies. Decomposition is subject to the same constraint as uniformisation, namely that the coefficients of the index mapping define bounded integer functions on the computation domain. Also, similar to uniformisation, two decomposition techniques can be defined according to the relation between the generators of the index mapping and the domain of the data dependence.

In the first case, illustrated in Fig. 3.14 (in 2 dimensions), the application of the technique consists of:

- the selection of a generator \( d \) of \( I \) such that \( d \notin \text{lin}(D) \). Let \( g \) be the coefficient of \( I \) relative to \( d \);

- the definition of the atomic integral index mapping \( I_0(z) = z + g(z)d \). Because of Proposition 3.2.4, \( I_0 \) is guaranteed to be injective over \( D \) and an inverse \( I_0^{-1} \) can be defined according to Proposition 3.2.7;

- the definition of a new integral index mapping \( I_1 \), based on the remaining generators of \( I \), and having as coefficients the composition of \( I_0^{-1} \) with the relative coefficients of \( I \).

Suitable routing domains and an auxiliary variable are introduced. Fig. 3.14 a) and b) illustrate the decomposition technique for \( I(z) \), respectively, inside and outside \( \text{lin}(D) \).

**Proposition 3.2.19** [Decomposition 1] Let \( \mathcal{DD} = (D, U, V, I) \) be an integral data dependence, with \( I(z) = z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j \). Let \( g \) be non-negative and bounded over \( D \), and \( \bar{g} \) the least upper bound of \( g \). Let \( d \notin \text{lin}(D) \) and \( P \) be the convex polyhedron generated by \( D \) and \( d \). Consider \( \pi \in \text{lin}(P) \cap D^\perp \), with \( \pi \neq 0 \), and the hyperplane \( [\pi : \theta] \) containing the domain \( D \).

Then \( \mathcal{DD} \) can be substituted by the atomic integral data dependence

\[
\mathcal{DD}' = (D, U, R, I_0)
\]
and the equation

\[ E = (D_1, R, V, id, I_1) \]

where:

- the index mappings are:

\[
\begin{align*}
I_0(z) &= z + g(z)d \\
I_1(z) &= z + \sum_{j=1}^{m} g_j(z)d_j
\end{align*}
\]

with \( g_j(z) = g_j(I_0^{-1}(z)) \), \( I_0^{-1}(z) = z + l(z)(-d) \), \( l(z) = (\pi \cdot z - \theta)/\eta \) and \( \eta = \pi \cdot d \);

- \( R \) is a new variable;

- the applied function is \( id(a) = a \);

- the new domain is \( D_1 = \{ z + ld \mid z \in D, 0 \leq l \leq \tilde{g}(z) \} \).

**Proof:** \( I_0^{-1} \) defines an inverse of \( I_0 \) on \( D \), according to Proposition 3.2.7. Hence, for all \( z \in D, I_0^{-1} \circ I_0(z) = z \).

Therefore, for all \( z \in D \),

\[
\begin{align*}
I_1 \circ I_0(z) &= I_0(z) + \sum_{j=1}^{m} g_j(I_0(z))d_j \\
&= I_0(z) + \sum_{j=1}^{m} g_j(I_0^{-1}(z))d_j \\
&= z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j = I(z)
\end{align*}
\]

Finally, for all \( z \in D \),

\[
U(z) = R(I_0(z)) = V(I_1 \circ I_0(z)) = V(I(z)).
\]

\( \blacksquare \) 3.2.19
Example 3.2.20  Consider the integral data dependence $\mathcal{D} = (D, U, V, I)$ with index mapping $I(i, j) = (i, j) + g_1(i, j)(1, 0) + g_2(i, j)(0, 1)$, where $g_1(i, j) = (i^2 + j^2) \mod 6$ and $g_2(i, j) = 2^j$, and domain $D = \{(i, j) | 1 \leq j \leq 5, i = 11 - 2j\}$. $\text{lin}(D) = \langle(2, -1)\rangle$ and $d = (1, 0) \not\in \text{lin}(D)$.

We can choose $\pi$ as any vector in $D^\perp$. In particular $\pi = (1, 2)$ (which is orthogonal to the generator $(2, -1)$ of $\text{lin}(D)$) satisfies the conditions of Proposition 3.2.19. Then, $[(1, 2): 11]$ is a hyperplane containing $D$, $\eta = (1, 2) \cdot (1, 0) = 1$ and $l(i, j) = i + 2j - 11$. Also, for all $(i, j) \in D$, $g_1(i, j) \leq 5$. If we apply Proposition 3.2.19, we obtain the data dependence

$$\mathcal{D}' = (D, U, R, I_0)$$

and the equation

$$E = (D_1, R, V, id, I_1)$$

where $id(a) = a$, the index mappings are:

$$I_0(i, j) = (i, j) + g_1(i, j)(1, 0)$$

$$I_1(i, j) = (i, j) + g_2'(i, j)(0, 1)$$

and the new domain is $D_1 = \{(i, j) | 1 \leq j \leq 5, 11 - 2j \leq i \leq 16 - 2j\}$. Note the new coefficient $g_2'$, which is defined, according to the proposition, as the composition of $g_2$ and $I_0^{-1}$, that is:

$$I_0^{-1}(i, j) = (i, j) + l(i, j)(-1, 0)$$

$$g_2'(i, j) = g_2(I_0^{-1}(i, j)) = 2^j$$

$\blacksquare$ 3.2.20
The second decomposition technique is similar to the first one, and corresponds to a situation in which the selected generator \( d \) of \( I \) is contained in the direction of the domain \( D \), i.e., \( d \in \text{lin}(D) \). As for uniformisation, a vector \( \pi \) needs to be chosen so that routing directions outside \( \text{lin}(D) \) can be considered. The choice of \( \pi \) is that of a vector orthogonal to both \( \text{lin}(D) \) and the space spanned by the generators of \( I \). The resulting decomposition is illustrated (in 3 dimensions) in Fig. 3.15 a) and b) for \( I(z) \) inside and outside \( \text{lin}(D) \), respectively. Note that this choice of \( \pi \) implies that even for a 1-dimensional domain \( D \), the application of the technique may require a 3-dimensional space (see Fig. 3.15 b)). The direction vectors \( \hat{d} = d + \pi \) and \(-\pi\) are considered as the generators of the new integral index mappings. Let \( g \) be the coefficient of \( I \) relative to \( d \). The technique consists of:

- the definition of the atomic integral index mapping \( I_0 = z + g(z)d \). Because of Proposition 3.2.4 and the choice of \( \pi \), \( \hat{d} \) is not in \( \text{lin}(D) \), the mapping \( I_0 \) is guaranteed to be injective over \( D \), and an inverse \( I_0^{-1} \) can be defined according to Proposition 3.2.7;

- the definition of an integral index mapping \( I_1 \) based on the remaining generators of \( I \), and having as coefficients the composition of \( I_0^{-1} \) with the relative coefficients of \( I \); and

- the definition of a third atomic integral index mapping \( I_2 \) with generator \(-\pi\).

**Proposition 3.2.21** [Decomposition 2] Let \( DD = (D, U, V, I) \) be an integral data dependence, with \( I(z) = z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j \). Let \( g, g_j \) be non-negative and bounded over \( D \) and let \( \bar{g}, \bar{g}_j \) be the least upper bounds of \( g, g_j \), respectively. Let \( d \in \text{lin}(D) \), \( P \) be the convex polyhedron generated by \( D \) and \( \{d_1, \ldots, d_m\} \), and \( \dim(P) < n \). Consider \( \pi \in P^\perp \), with \( \pi \neq 0 \), the hyperplane \([\pi : \theta]\) containing the domain \( D \), and let \( \hat{d} = d + \pi \).

Then \( DD \) can be substituted by the atomic integral data dependence

\[
DD' = (D, U, R^1, I_0)
\]

and the equations:

\[
E_1 = (D_1, R^1, R^2, id, I_1)
\]

\[
E_2 = (D_2, R^2, V, id, I_2)
\]

where:
- the index mappings are:

\[
\begin{align*}
I_0(z) &= z + g(z)d \\
I_1(z) &= z + \sum_{j=1}^{m} g_j(z)d_j \\
I_2(z) &= z + g'(z)(-\pi)
\end{align*}
\]

with \( g_j^j(z) = g_j(I_0^{-1}(z)) \), \( I_0^{-1}(z) = z + l(z)(-\hat{d}) \), \( g'(z) = l(z) \), \( l(z) = (\pi \cdot z - \theta)/\eta \) and \( \eta = \pi \cdot \hat{d} \);

- \( R^1, R^2 \) are new variables;

- the applied function is \( id(a) = a \);

- the new domains are:

\[
\begin{align*}
D_1 &= \{ z + l\hat{d} \mid z \in D, 0 \leq l \leq \hat{g} \} \\
D_2 &= \{ z + \sum_{j=1}^{m} l_jd_j \mid z \in D_1, 0 \leq l_j \leq \hat{g}_j \}.
\end{align*}
\]

PROOF: As \( \pi \notin \text{lin}(D) \), then \( \hat{d} = d + \pi \notin \text{lin}(D) \) and \( I_0 \) is injective over \( D \). Hence, \( I_0^{-1} \) defines an inverse of \( I_0 \) on \( D \), according to Proposition 3.2.7. Therefore, for all \( z \in D \), \( I_0^{-1} \circ I_0(z) = z \).

For all \( z \in D \),

\[
I_1 \circ I_0(z) = \\
= I_0(z) + \sum_{j=1}^{m} g_j(I_0(z))d_j \\
= I_0(z) + \sum_{j=1}^{m} g_j(I_0^{-1} \circ I_0(z))d_j \\
= z + g(z)\hat{d} + \sum_{j=1}^{m} g_j(z)d_j.
\]

Also, as \( \pi \in P^\perp \), then \( \pi \cdot d_j = 0 \) and \( \pi \cdot (\sum_{j=1}^{m} g_j(z)d_j) = 0 \). Hence, for all \( z \in D \),

\[
g'(I_1 \circ I_0(z)) = l(I_1 \circ I_0(z)) \\
= (\pi \cdot (I_1 \circ I_0(z)) - \theta)/\eta \\
= (\pi \cdot (z + g(z)\hat{d} + \sum_{j=1}^{m} g_j(z)d_j) - \theta)/\eta \\
= (\pi \cdot z + g(z)\pi \cdot \hat{d} + \pi \cdot (\sum_{j=1}^{m} g_j(z)d_j) - \theta)/\eta \\
= (\theta + g(z)\eta + 0 - \theta)/\eta = g(z).
\]


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Therefore, for all \( z \in D \),
\[
I_2 \circ I_1 \circ I_0(z) = I_2(I_1 \circ I_0(z)) \\
= I_1 \circ I_0(z) + g'(I_1 \circ I_0(z))(-\pi) \\
= I_1 \circ I_0(z) + g(z)(-\pi) \\
= z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j + g(z)(-\pi) \\
= z + g(z)(\pi + d - \pi) + \sum_{j=1}^{m} g_j(z)d_j \\
= z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j = I(z).
\]

Finally, for all \( z \in D \),
\[
U(z) = R^1(I_0(z)) = R^2(I_1 \circ I_0(z)) \\
= V(I_2 \circ I_1 \circ I_0(z)) = V(I(z)).
\]

Condition \( \text{dim}(P) < n \) above implies that it may be necessary to increase the number of dimensions of the space prior to the application of Proposition 3.2.21.

Example 3.2.22 Consider the integral data dependence \( \mathcal{D} = (D, U, V, I) \) with index mapping \( I(i, j) = (i, j) + g_1(i, j)(1, 0) + g_2(i, j)(0, 1) \), where \( g_1(i, j) = (i^2 + j^2) \mod 6 \) and \( g_2(i, j) = 2^i \), and domain \( D = \{(i, j) \mid 1 \leq j \leq 5, 11 - 2j \leq i \leq 16 - 2j \} \). \( \text{lin}(D) = \mathbb{Z}^2 \) and \( d = (1, 0) \in \text{lin}(D) \). As the polyhedron \( P \) generated by \( D \) and \( d_2 = (0, 1) \) is of full dimension in \( \mathbb{Z}^2 \), we need to reindex the data dependence in \( \mathbb{Z}^3 \), before applying Proposition 3.2.21.

Any non-null vector in \( D^\perp \) can be chosen as \( \pi \). For instance, vector \( \pi = (0, 0, 1) \) satisfies the conditions of Proposition 3.2.21 and \( [0, 0, 1 : 0] \) is a hyperplane containing \( P \). Let \( d = d + \pi = (1, 0, 1) \), \( \eta = (0, 0, 1) \cdot (1, 0, 1) = 1 \) and \( l(i, j, k) = k \). Note that, for all \( (i, j, k) \in D, g_1(i, j, k) \leq 5, \) and \( g_2(i, j, k) \leq 2^6 \). If we apply Proposition 3.2.21 we obtain the data dependence

\[
\mathcal{D}' = (D, U, R^1, I_0)
\]

and the equations:

\[ E_1 = (D_1, R^1, R^2, id, I_1) \]
\[ E_2 = (D_2, R^2, V, id, I_2) \]

where:
- the index mappings are:

\[ I_0(i,j,k) = (i,j,k) + g_1(i,j,k)(1,0,1) \]
\[ I_1(i,j,k) = (i,j,k) + g'_2(i,j,k)(0,1,0) \]
\[ I_2(i,j,k) = (i,j,k) + g'(i,j,k)(0,0,-1) \]

- the new domains are:

\[ D_1 = \{(i,j,k) | 1 \leq j \leq 5, 0 \leq k \leq 5, 11 - 2j + k \leq i \leq 16 - 2j + k \} \]
\[ D_2 = \{(i,j,k) | 16 \leq i + 2j - k \leq 16 + 2^6, 1 \leq 1i - k \leq 14, 1 \leq j \leq 5 + 2^5 \} \]

Note the new coefficients \( g'_2 \) and \( g' \) defined, according to Proposition 3.2.21, as:

\[ g'_2(i,j,k) = g_2(I_0^{-1}(i,j,k)) = 2^j \]
\[ g'(i,j,k) = l(i,j,k) = k \]

where \( I_0^{-1}(i,j,k) = (i,j,k) + l(i,j,k)(-1,0,-1) = (i - k, j, 0) \).

The previous results provide means of defining decomposition techniques for integral data dependencies which can be automated. The decomposition procedure is based on the possibility of defining, at each step, an inverse of an atomic integral index mapping. This definition relies upon the results in Propositions 3.2.4 and 3.2.7, and, in general, may require an increase in the dimensionality of the computation space. A more economic decomposition step may be defined if such an inverse is known independently from geometric considerations. In such a case, a decomposition technique may be defined as follows:

**Proposition 3.2.23 [Decomposition 3]** Let \( \mathcal{DD} = (D, U, V, \mathcal{I}) \) be an integral data dependence, with \( \mathcal{I}(z) = z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j \). Let \( g \) be non-negative and bounded over \( D \), and \( \bar{g} \) the least upper bound of \( g \). Let \( I_0(z) = z + g(z)d \) be an injective mapping over \( D \) with inverse \( I_0^{-1} \).

Then \( \mathcal{DD} \) can be substituted by the atomic integral data dependence

\[ \mathcal{DD}' = (D, U, R, I_0) \]

and the equation

\[ E = (D_1, R, V, id, I_1) \]

where:
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- the index mappings are:

\[ I_0(z) = z + g(z)d \]
\[ I_1(z) = z + \sum_{j=1}^{m} g'_j(z)d_j \]

with \( g'_j(z) = g_j(I_0^{-1}(z)) \);

- \( R \) is a new variable;

- the applied function is \( id(a) = a \);

- the new domain is \( D_1 = \{ z + ld \mid z \in D, 0 \leq l \leq \bar{g} \} \).

**Proof:** As \( I_0^{-1} \) is an inverse of \( I_0 \) on \( D \), then for all \( z \in D \), \( I_0^{-1} \circ I_0(z) = z \).

Therefore, for all \( z \in D \),

\[ I_1 \circ I_0(z) = I_0(z) + \sum_{j=1}^{m} g'_j((I_0(z))d_j \]
\[ = I_0(z) + \sum_{j=1}^{m} g_j(I_0^{-1} \circ I_0(z))d_j \]
\[ = z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j = I(z). \]

Finally, for all \( z \in D \),

\[ U(z) = R(I_0(z)) \]
\[ = V(I_1 \circ I_0(z)) = V(I(z)). \]

3.2.23

**Example 3.2.24** Consider the integral data dependence \( DD = (D, U, V, I) \) with index mapping \( I(i, j) = (i, j) + (1, 1) + g(i, j)(1, 0) \), where \( g(i, j) = (i^2 + j^2) \mod 6 \), and domain \( D = \{ (i, j) \mid 1 \leq j \leq 5, 11 - 2j \leq i \leq 16 - 2j \} \).

The index mapping \( I_0(i, j) = (i, j) + (1, 1) \) admits the inverse \( I_0^{-1}(i, j) = (i, j) + (-1, -1) \). Hence, \( DD \) can be substituted by the data dependence

\[ DD' = (D, U, R, I_0) \]
and the equation
\[ E = (D_1, R, V, id, I_1) \]
where \( I_1(i, j) = (i, j) + g'(i, j)(1, 0), g'(i, j) = g(I_0^{-1}(i, j)) = ((i - 1)^2 + (j - 1)^2) \mod 6 \) and \( D_1 = \{(i + 1, j + 1) \mid (i, j) \in D\} \).

Note that as \( \text{lin}(D) = \mathbb{Z}^2 \) and \( d = (1, 0) \in \text{lin}(D) \), the application of Proposition 3.2.21 would require an increase in the dimensionality of the space. \( \blacksquare \) 3.2.24

From the previous results, we obtain the corollary:

Corollary 3.2.25 Let \( \mathcal{D}D = (D, U, V, I) \) be an integral data dependence, with \( I(z) = z + g(z)d + \sum_{j=1}^{m} g_j(z)d_j \). Let \( g \) be non-negative and bounded over \( D \), and \( \bar{g} \) the least upper bound of \( g \).

If \( \bar{g} = 0 \), then \( \mathcal{D}D \) can be substituted by the integral data dependence
\[ \mathcal{D}D_0 = (D, U, V, I_0) \]
where \( I_0(z) = z + \sum_{j=1}^{m} g_j(z)d_j \). \( \blacksquare \) 3.2.25

### 3.3 Regularisation and Affine Scheduling

Form the discussion in Section 2.2.6, a pointed dependence cone implies the existence of valid affine timing functions. The preservation of such functions by regularisation can be guaranteed if the techniques realise transformations of pointed dependence cones.

The following two propositions provide the basic results to prove that integral regularisation techniques preserve affine scheduling. The results are illustrated in Fig. 3.16 for a 3-dimensional space. Parts a), b) and c) of the figure illustrate Proposition 3.3.1, while parts d), e) and f) correspond to Proposition 3.3.2.

**Proposition 3.3.1** Let \( C \) be a pointed polyhedral convex cone with generators \( r_1, \ldots, r_p \). Let \( \rho \in C^\perp \) and \( r \in C \), with \( \rho \neq 0 \) and \( r \neq 0 \). Define \( \bar{r} = r + \rho \) and \( \bar{r} = r - \rho \). Then the cone \( C' \) generated by \( r_1, \ldots, r_p, \bar{r}, \bar{r} \) is pointed.

**Proof:** The result is based on the separation theorem for pointed cones (see Appendix E). As \( C \) is pointed, there exists \( \lambda \in \text{lin}(C) \) such that \( \lambda \cdot c > 0 \) for all \( c \in C \), with \( c \neq 0 \). In particular, \( \lambda \cdot r > 0 \) and \( \lambda \cdot r_j > 0 \), for all \( j = 1, \ldots, p \). As \( r \in C^\perp \), then \( \lambda \cdot \rho = 0 \).
We prove that \( \lambda \cdot c > 0 \), for all \( c \in C' \), with \( c \neq 0 \). Trivially the property is true for each of the generators \( r_j \) of \( C \). Besides:

\[
\begin{align*}
\lambda \cdot \tilde{r} &= \lambda \cdot r + \lambda \cdot \rho = \lambda \cdot r > 0 \\
\lambda \cdot \tilde{r} &= \lambda \cdot r - \lambda \cdot \rho = \lambda \cdot r > 0
\end{align*}
\]

Therefore, \( C' \) is pointed.

**Proposition 3.3.2** Let \( C \) be a pointed polyhedral convex cone with generators \( r_1, \ldots, r_p \).

Let \( \rho \in C^\perp \) and \( r \in C \), with \( \rho \neq 0 \) and \( r \neq 0 \). Define \( \tilde{r} = r + \rho \). Then the cone \( C' \) generated by \( r_1, \ldots, r_p, \tilde{r}, -\rho \) is pointed.

**Proof:** The result is based on the separation theorem for pointed cone (see Appendix E).

As \( C \) is pointed, there exists \( \lambda \in \text{lin}(C) \) such that \( \lambda \cdot c > 0 \), for all \( c \in C \), with \( c \neq 0 \). In particular, \( \lambda \cdot r > 0 \) and \( \lambda \cdot r_j > 0 \), for all \( j = 1, \ldots, p \). As \( \rho \in C^\perp \), then \( \lambda \cdot \rho = 0 \).

Let \( \lambda' = a\lambda - \rho \), with \( a > |\rho|^2/\lambda \cdot r > 0 \). We prove that \( \lambda' \cdot c > 0 \) for all \( c \in C' \), with \( c \neq 0 \). In fact, for all \( c \in C \), with \( c \neq 0 \), \( \lambda' \cdot c = a\lambda \cdot c - \rho \cdot c = a\lambda \cdot c > 0 \).

Hence, \( \lambda' \cdot r_j > 0 \) for all \( j = 1, \ldots, p \). Besides:

\[
\begin{align*}
\lambda' \cdot \tilde{r} &= \lambda' \cdot (r + \rho) = a\lambda \cdot (r + \rho) - \rho \cdot (r + \rho) \\
&= a\lambda \cdot r + a\lambda \cdot \rho - \rho \cdot r - \rho \cdot \rho = a\lambda \cdot r - |\rho|^2 > 0 \\
\lambda' \cdot (-\rho) &= -a\lambda \cdot \rho + \rho \cdot \rho = |\rho|^2 > 0
\end{align*}
\]
CHAPTER 3. INTEGRAL RECURRENCE EQUATIONS

3.3.2

Note that if \( C \) is a dependence cone and \( \lambda \) corresponds to a valid affine timing function for \( C \), the proofs of the above propositions indicate how to determine a vector \( \lambda' \) from \( \lambda \), which defines a valid affine timing function for the dependence cone \( C' \). In particular, according to Proposition 3.3.1, \( \lambda' \) is equal to \( \lambda \), while according to Proposition 3.3.2, \( \lambda' \) is equal to \( a\lambda - \rho \), with \( a > |\rho|^2/(\lambda \cdot r) \).

Example 3.3.3 Consider the pointed cone \( C \) of Fig. 3.17 a), generated by \( r_1 = (1, 2, 0) \), \( r_2 = (3, 1, 0) \), and the non-null vector \( \lambda = (0, 1, 0) \). Note that \( \lambda \cdot r_i > 0 \), for \( i = 1, 2 \). Consider the vector \( r = r_1 + r_2 = (4, 3, 0) \) in \( C \) (see Fig. 3.17 b)) and the vector \( \rho = (0, 0, 1) \) in \( C^\perp \).

Let us first consider the cone \( C' \) (see Fig. 3.17 c)) generated by \( r_1, r_2, \hat{r} \) and \( \hat{r} \), where \( \hat{r} = r + \rho = (4, 3, 1) \) and \( \hat{r} = r - \rho = (4, 3, -1) \). Then \( C' \) is pointed and \( \lambda \cdot c > 0 \) for all \( c \in C' \). This fact can be verified by considering the generators of \( C' \), i.e., :

\[
\begin{align*}
\lambda \cdot r_1 &= 2 > 0 \\
\lambda \cdot r_2 &= 1 > 0 \\
\lambda \cdot \hat{r} &= 3 > 0 \\
\lambda \cdot \hat{r} &= 3 > 0
\end{align*}
\]

Let us now consider the cone \( C'' \) (see Fig. 3.17 d)) generated by \( r_1, r_2, \hat{r} \) and \( -\rho \), where \( \hat{r} = r + \rho = (4, 3, 1) \) and \( -\rho = (0, 0, -1) \). Then \( C'' \) is pointed and any \( \lambda' = a\lambda - \rho \), with
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<table>
<thead>
<tr>
<th>Technique</th>
<th>Generators of $C$</th>
<th>Generators of $C'$</th>
<th>Proposition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unif. (3.2.10)</td>
<td>$r$</td>
<td>$r$</td>
<td>—</td>
</tr>
<tr>
<td>Unif. (3.2.13)</td>
<td>$r$</td>
<td>$r, \hat{r}, \tilde{r}$</td>
<td>3.3.1</td>
</tr>
<tr>
<td>Par. Unif. (3.2.16)</td>
<td>$r$</td>
<td>$r, \hat{r}, \tilde{r}$</td>
<td>3.3.1</td>
</tr>
<tr>
<td>Decomp. (3.2.19)</td>
<td>$r, r_1, \ldots, r_m$</td>
<td>$r, r_1, \ldots, r_m$</td>
<td>—</td>
</tr>
<tr>
<td>Decomp. (3.2.21)</td>
<td>$r, r_1, \ldots, r_m$</td>
<td>$r, r_1, \ldots, r_m, \hat{r}, \rho$</td>
<td>3.3.2</td>
</tr>
<tr>
<td>Decomp. (3.2.23)</td>
<td>$r, r_1, \ldots, r_m$</td>
<td>$r, r_1, \ldots, r_m$</td>
<td>—</td>
</tr>
</tbody>
</table>

Table 3.1. Regularisation and the preservation of affine scheduling.

$a > 1/3$, is such that $\lambda' c > 0$ for all $c \in C''$. For instance, let $a = 1$ and $\lambda' = \lambda - \rho = (0.1, -1)$.

Then:

$$
\begin{align*}
\lambda' \cdot r_1 &= 2 > 0 \\
\lambda' \cdot r_2 &= 1 > 0 \\
\lambda' \cdot \hat{r} &= 2 > 0 \\
\lambda' \cdot (-\rho) &= 1 > 0 
\end{align*}
$$

Note that $\lambda$ does not have the same property as, for instance, $\lambda \cdot (-\rho) = 0$. ■ 3.3.3

Both the uniformisation and decomposition techniques which we have introduced for integral data dependencies, define particular cases to which the above propositions apply. We have summarised the various techniques in Table 3.1. The first column of the table indicates the technique with, in brackets, the reference number of the corresponding formal result. The generators of the (embedding dependence) cone before ($C$) and after ($C'$) the application of the technique are given in the second and third columns, respectively. In particular, in such columns $r = -d$ and $r_j = -d_j$, for all $j = 1, \ldots, m$, denote the generators of the corresponding dependence cone, while $\hat{r} = -\hat{d}, \tilde{r} = -\tilde{d}$ and $\rho = -\pi$ the new generators introduced by the various techniques. The last column indicates which of the above proposition applies. A line — indicates that no result applies and corresponds to a case in which the dependence cone is not modified by the regularisation technique.

From the above results, it follows that the integral regularisation techniques which we have defined preserve the affine scheduling of a data dependence. In order to guarantee a similar result more globally for the whole specification, we need to consider all its data dependencies at the same time. Fortunately, this is not a major problem. In fact, we may define the dependence cone of a system of equations $S$ as the smallest polyhedral convex cone which
Table 3.2. Summary of regularisation techniques.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Condition</th>
<th>Extra Dimensions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unif. (3.2.10)</td>
<td>( d \notin \text{lin}(D) )</td>
<td>no</td>
</tr>
<tr>
<td>Unif. (3.2.13)</td>
<td>( d \in \text{lin}(D) )</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td>( \text{dim}(D) &lt; n )</td>
<td></td>
</tr>
<tr>
<td>Par. Unif. (3.2.16)</td>
<td>( d \in \text{lin}(D) )</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td>( \text{dim}(D) &lt; n )</td>
<td></td>
</tr>
<tr>
<td>Decomp. (3.2.19)</td>
<td>( d \notin \text{lin}(D) )</td>
<td>no</td>
</tr>
<tr>
<td>Decomp. (3.2.21)</td>
<td>( d \in \text{lin}(D) )</td>
<td>yes</td>
</tr>
<tr>
<td></td>
<td>( \text{dim}(P) &lt; n )</td>
<td></td>
</tr>
<tr>
<td>Decomp. (3.2.23)</td>
<td>known inverse</td>
<td>no</td>
</tr>
</tbody>
</table>

contains all the dependence cones of \( S \). Let \( C_S \) denote such a cone. Then, by definition, for each data dependence \( \mathcal{D} \) of \( S \), the corresponding cone \( C \) is contained in \( C_S \). Hence, a choice of \( \pi \) in \( (C_S)^\perp \) guarantees both that \( \pi \in C^\perp \) and that affine schedulings are preserved for the system \( S \).

3.4 Summary

In this chapter we have discussed how integral data dependencies can be defined and made uniform in the context of classic regular array synthesis. The key issue was the relationship established between the integral index mapping and sets of direction vectors of the lattice space. This relationship has allowed us both to relate integral data dependencies to affine data dependencies, and to generalise forms of regularisation techniques from the affine to the integral case.

The notion of atomic integral data dependence was formulated as an integral data dependence with a particular role in regularisation. Based on such a notion, we have developed both decomposition and uniformisation techniques. We have summarised those techniques in Table 3.2. In the table, the first column indicates the technique and in brackets the reference number of the corresponding formal result. The second column indicates the basic conditions for its applicability. In particular, \( d \) denotes a generator of the integral index mapping, \( D \) the domain of the data dependence, and \( P \) the polyhedron generated by \( d \) and \( D \). The last column indicates whether the application of the technique may require an increase in the number of dimensions of the space.

The techniques which we have developed are amenable to semi-automatic support. Designer's
intervention is required, typically for the choice of the regularisation directions (the generators of the integral index mapping and their ordering) as well as the hyperplane on which pipelining and routing schemes are based (the vector $\pi$ in the formal propositions). The details of the transformations are, however, accounted for by the techniques. Also, the conditions for the existence and preservation of affine scheduling which we have developed are not just of theoretical relevance, but provide guidelines to the designer in the choice of the regularisation directions.

As affine data dependencies are particular cases of integral data dependencies, several techniques are available for their regularisation, including those specifically developed in the literature for them, and those we have developed in this work for the integral case. In general, the designer has to evaluate which technique should be adopted on the basis of the requirements of each specific problem. As a general rule, however, the more specialised is a technique, the simpler and more effective it is likely to be. For instance, integral techniques always introduce control overhead as they are tailored to situations in which convexity needs to be recovered. Indeed, in the affine case convexity comes for free. Also, the removal of affine broadcasts can be achieved very effectively through a technique (known in the literature as pipelining [QuVa89]) which selects regularisation directions in the null space of the linear part of the affine transformation defined by an index mapping. A similar technique cannot be adopted for integral mappings, as linearity is not one of their general properties. On the other hand, the regularisation schemes that we have developed have a property of reconfigurability, achieved by the initialisation of the control variables, which is not characteristics of any affine regularisation technique. Our techniques are also, in general, more detailed than the affine techniques presented in [QuVa89] (on which our work in based), which tend to state the applicability conditions without providing all the details for their practical implementation.
Chapter 4

Dynamic Recurrence Equations

Both affine and integral data dependencies are static in that they are completely determined when the algorithm is specified. There exist algorithms, however, whose data dependence relations do not share this characteristic. For example, their definition may rely upon values which are provided or computed only when the algorithm is executed. We call such data dependence relations dynamic. In this chapter we aim at introducing the concept of dynamic data dependence in the context of regular array synthesis and identifying classes of dynamic data dependencies which are amenable to a systematic transformation into regular arrays.

The development of synthesis techniques for dynamic problems constitutes an entirely new chapter in regular array design. Researchers in the field have so far avoided consideration of dynamic problems, mainly on the belief that dynamic dependencies and the static topology of regular arrays cannot be reconciled. However, examples of ad hoc regular arrays for problems which may be considered as dynamic exist in the literature (one of these problems, that of Gaussian elimination with pivoting, will be considered in Chapter 5). Therefore, it appears that regular array design is feasible at least for restricted classes of dynamic problems.

One of the basic issues we need to address in the synthesis of dynamic problems is the type of dynamic data dependence relation that we want to consider. The notion we adopt in this chapter is that of a data dependence relation which is dynamic with respect to the inputs of the algorithm. More precisely, the data dependence relation between two variables of a specification changes (i.e., different pairs of their instances may be related under the data dependence) at each execution of the algorithm on the basis of the input values provided. In formalising this notion in the context of regular array synthesis, a major difficulty to overcome is the lack, in the classical framework, of some of the necessary basic concepts, primarily a formal notion of input. Hence, the first part of this chapter will be devoted to introducing
some of these basic notions. In particular, formal definitions of input and indexed variable will be provided in Sections 4.1, while notions of dynamic index mapping, dynamic data dependence and dependence graph will be developed in Section 4.2.

Once the basic notions have been established, we proceed to identify classes of dynamic problems which can be reduced systematically to uniform problems. The problems we consider may be regarded as a dynamic generalisation of integral problems and possibilities for their regularisation stem from the use of control variables in the definition of the uniformisation techniques, which can be dynamically reconfigured at each execution of the algorithm. Reconfigurable control variables enable us to reconcile the existence of dynamic data dependencies in the algorithm with its execution on a regular array of static topology. Namely, the connections between processing elements are not altered, while software control provides for a flexible routing of the data through the network. This type of dynamic problems and their regularisation will be considered in the second part of the chapter, including Sections 4.3, 4.4 and 4.5. We will draw some conclusions in Section 4.6.

4.1 Inputs and Indexed Variables

As mentioned above, we consider data dependencies which are dynamic with respect to the inputs of a system of equations. For their characterisation we need first to define more precisely what an input is. The basic concepts of regular array synthesis introduced in Chapter 2 do not account for a such a definition, because inputs do not play an essential part in the analysis and transformation of static data dependencies (hence their explicit treatment can be avoided).

By recalling the notation of Chapter 2, we will denote by $CS$ the computation space and $Val$ the set of data values, containing the undefined value $\perp$.

Given a system of equations $S$, we define an input of $S$ as an assignment of values to some of the variables of $S$ at points of the computation space. As may be expected, such an assignment is based on the input equations of $S$. More precisely:

**Definition 4.1.1 [Inputs]** Let $S$ be a system of equations with input equations $E_1, \ldots, E_r$, where $E_i = (D_i, U_i, \text{ini})$, for $i = 1, \ldots, r$. Let $V = \{U_1, \ldots, U_r\}$ and $D = \bigcup_{i=1}^r D_i$. An input of $S$ is defined as a mapping $\text{in} : V \times D \to Val$, where:

$$\text{in}(U_i, c) = \begin{cases} \text{ini}(c) & c \in D_i \\ \perp & \text{otherwise} \end{cases}$$
We denote the set of inputs to $S$ by $\text{Inputs}_S$.

**Example 4.1.2** Consider the following system of equations (which generalises that in Example 2.1.2 for the Fibonacci sequence):

$$E_1 = (D_1, F, in_F)$$
$$E_2 = (D_2, F, (F, F) \cdot (I_1, I_2)).$$

with domains $D_1 = \{0, 1\}$, $D_2 = \{2, \ldots, n\}$, for some integer $n > 2$; index mappings $I_1(i) = i - 1$, $I_2(i) = i - 2$; and applied functions $+(a, b) = a + b$, $in_F(i) = f_i$, with $f_i \in \mathbb{Z}$. The difference with respect to Example 2.1.2 is the definition of $in_F$ that, here, is not a constant function.

$E_1$ is the only input equation of the system. The inputs of $S$ are functions of the type $in : \{F\} \times D_1 \to \mathbb{Z} \cup \{\bot\}$, where:

$$in(F, i) = \begin{cases} 
in_F(i) & i \in D_1 \\ \bot & \text{otherwise} \end{cases}$$

The definition of inputs above allows us to define more precisely the indexed variables of a system of equations. In particular, while in Chapter 2 we informally associated indexed variables with the (tabulated) functions computed by the algorithm, we can now define an indexed variable as a mapping which associates a (tabulated) function with each input of the algorithm. The definition is the following:

**Definition 4.1.3** [Indexed Variable] Let $S$ be a system of equations and $\text{Inputs}_S$ its set of inputs. A variable $V$ of $S$, with definition domain $Def_DV$, is a mapping from $\text{Inputs}_S$ to $[Def_DV \to Val]$.

**Example 4.1.4** Let us consider the system of equations in Example 4.1.2. For $in_F(i) = 1$ on $D_1$, variable $F$ represents the first $n + 1$ entries of the Fibonacci sequence. For $in_F(i) = 0$, $F$ is the zero function. For $in_F(i) = 2$, $F$ represents the first $n + 1$ entries of the sequence $2, 2, 4, 6, 10, \ldots$. Indeed infinitely many functions may be obtained.
In the following, given a variable \( V \), we adopt the notation \( V[\cdot] \), where [\( \cdot \)] is a place holder for inputs. Then, for all \( in \in Input_S \), \( V[in] \) defines a function from \( \text{Def}Dv \) to \( Val \).

Once a formal definition of indexed variable has been provided, a number of properties for those variables can be given. The following properties characterise indexed variables which assume integer values. They will be used in the second part of this chapter for the definition of a class of dynamic data dependencies in the context of Euclidean synthesis methods. For ease of presentation, we prefer to introduce them here rather than later in the chapter.

**Definition 4.1.5 [Integer-valued Indexed Variable]** Let \( S \) be a system of equations, \( Input_S \) its set of inputs, and \( V \) a variable of \( S \) with definition domain \( \text{Def}Dv \). \( V \) is integer-valued if \( V \) defines a mapping from \( Input_S \) to \( [\text{Def}Dv \rightarrow \mathbb{Z}] \). In addition, let \( D \subseteq \text{Def}Dv \). Then \( V \) is:

- **non-negative over** \( D \), if for all \( in \in Input_S \), \( V[in](c) \geq 0 \) for all \( c \in D \);
- **bounded (above) over** \( D \), if there exists \( m \in \mathbb{Z} \), such that for all \( in \in Input_S \), \( V[in](c) \leq m \) for all \( c \in D \).

We assume an integer-valued indexed variable \( V \) equal to 0 outside its definition domain, i.e., for all \( in \in Input_S \), \( V[in](c) = 0 \) for \( c \notin \text{Def}Dv \). This assumption replaces that made in Chapter 2 that a variable is undefined (equal to \( \perp \)) outside its definition domain. Although there is a small change of semantics (we cannot distinguish when the variable is defined and equal to 0 from when it is undefined), this assumption simplifies the definitions of the regularisation techniques of the following sections.

**4.1.1 Implicit Quantification**

As index variables are defined in terms of inputs, all definitions which involve indexed variables are implicitly universally quantified over those inputs. For instance, a recurrence equation

\[
E = (D, U, (V_1, \ldots, V_m), f, (I_1, \ldots, I_m))
\]

in a system \( S \) can be seen as a short-hand for the expression: for all \( in \in Input_S \),

\[
E[in] = (D, U[in], (V_1[in], \ldots, V_m[in]), f, (I_1, \ldots, I_m)).
\]
This implicit quantification is not usually addressed in the synthesis methods (indeed, we have
ignored it in the previous chapters) as only static elements of the system are manipulated. In
particular, the emphasis is on the analysis and transformation of data dependence relations
between computation points, and those relations are defined as static (both computation
points and index mappings do not depend on the inputs of a system of equations). However,
the shift from static to dynamic data dependencies addressed in the following sections, will
require us to consider the inputs of the system explicitly. In the following, we will retain the
implicit quantification whenever possible.

4.2 Dynamic Data Dependencies

Our concept of dynamic data dependencies is based on the idea of dynamic index mapping
as an index mapping which is dynamic with respect to the inputs of an algorithm. That
an index mapping becomes dynamic may appear as a simple extension of the static notion.
It has, however, far reaching repercussion on the whole formalism, as index mappings are
among the basic elements of synthesis methods. In particular, data dependence relations and
their graphical representation as dependence graphs are affected, in that a new dependence
relation and graph are obtained for each input. Formally, we define a dynamic index mapping
as follows:

Definition 4.2.1 [Dynamic Index Mapping] Let $S$ be a system of equations and $\text{Input}_S$ its
set of inputs. A dynamic index mapping $\mathcal{I}$ is a mapping from $\text{Input}_S$ to $[CS \rightarrow CS]$.

Definition 4.2.1 implies that for all $in \in \text{Input}_S$, $\mathcal{I}[in]$ is an index mapping. Note that
the definition assumes the existence of a system of equations upon whose inputs the index
mapping is defined. For convenience, in the following we will omit the explicit reference to
this system, and will denote the set of its inputs simply by Input, without decoration.

Note that a (static) index mapping can be interpreted as a dynamic index mapping with a
singleton set as its range, i.e., the dynamic index mapping always associates the same (static)
index mapping with all the inputs to the system.

Example 4.2.2 Consider the following single assignment code segment:

$$
\text{for } i := 1 \text{ to } 3 \text{ do}
$$
CHAPTER 4. DYNAMIC RECURRENCE EQUATIONS

\[ \text{read}(Y(i)); \]
\[ \text{for } i := 4 \text{ to } n \text{ do} \]
\[ \quad \text{begin} \]
\[ \quad \text{read}(G(i)); \]
\[ \quad \text{read}(X(i)); \]
\[ \quad I(i) := X(i) - (G(i) \mod 4); \]
\[ \quad Y(i) := \text{sqr}(Y(i - I(i))); \]
\[ \text{end;} \]

The code corresponds to the system \( S \) of equations:

\[ E_1 = (D_1, Y, \text{iny}) \]
\[ E_2 = (D_2, G, \text{inG}) \]
\[ E_3 = (D_2, X, \text{inX}) \]
\[ E_4 = (D_2, I, (X, G), h, (I_1, I_1)) \]
\[ E_5 = (D_2, Y, \text{sqr}, I_2) \]

with domains \( D_1 = \{0, \ldots , 3\} \) and \( D_2 = \{4, \ldots , n\} \), index mappings \( I_1(i) = i, I_2(i) = i - I(i) \), and applied functions \( h(a, b) = a - (b \mod 4), \text{sqr}(a) = a^2, \text{iny}(i) = y_i, \) with \( y_i \in \mathbb{R} \), \( \text{inG}(i) = g_i, \) with \( g_i \in \mathbb{Z} \), and \( \text{inX}(i) = i \).

The mapping \( I_2(i) = i - I(i) \) is a dynamic index mapping. In fact, \( I_2 \) is defined in terms of the indexed variable \( I \) that, by definition, depends on the inputs of \( S \).

Dynamic data dependencies and data dependence graphs can be defined on the basis of dynamic index mappings. The main difference from the static case is that for each input a new relation and corresponding graph are defined, based on the corresponding index mapping. The reader may compare the definitions below with those in Chapter 2.

**Definition 4.2.3 [Dynamic Data Dependence]** Let \( I \) be a dynamic index mapping. A dynamic data dependence \( DD \) is a 4-tuple \((D, U, V, I)\), such that for all \( in \in \text{Input} \), \( DD[\text{in}] = (D, U, V, I[\text{in}]) \) is a data dependence.
Definition 4.2.4 [Dynamic Data Dependence Graph] Let $\mathcal{DD} = (D, U, V, I)$ be a dynamic data dependence. Its dynamic data dependence graph $\mathcal{DDG}$ is a 2-tuple $(V, A)$, such that for all $in \in Input$, $(\mathcal{N}[in], \mathcal{A}[in])$ is the data dependence graph defined by:

- $\mathcal{N}[in] = D \cup I[in](D)$; and
- $\mathcal{A}[in] = \{(I[in](d), d) | d \in D\}$.

Example 4.2.5 Consider the system $S$ of Example 4.2.2. Its data dependencies are:

$$
\mathcal{DD}_1 = (D_2, I, X, I_1)
$$
$$
\mathcal{DD}_2 = (D_2, I, G, I_1)
$$
$$
\mathcal{DD}_3 = (D_2, Y, Y, I_2)
$$

with $I_1(i) = i$ and $I_2(i) = i - I(i)$. The data dependencies $\mathcal{DD}_1$ and $\mathcal{DD}_2$ are (static) uniform, while $\mathcal{DD}_3$ is dynamic. The data dependence graph defined by $\mathcal{DD}_3$ is also dynamic. Fig. 4.1 a) and b) illustrate the dynamic data dependence graph for two inputs to $S$ (and for $n = 11$). The related tables give the corresponding values of the variables of the system on their domains.

4.3 Dynamic Data Dependencies in Euclidean Synthesis

In this section we introduce a class of dynamic data dependencies which are amenable to systematic transformation into regular arrays in the context of Euclidean synthesis methods.

The benefits of adopting Euclidean geometry and linear algebra as the mathematical framework for synthesis methods, both from a theoretical and applicative point of view, have already been discussed at length in Chapter 2. We have also already mentioned the necessity of providing an explicit syntactic characterisation for the formal manipulation of algorithm specifications in such a context (see Section 2.2.1).

In the following we will provide a syntactic characterisation of a class of dynamic data dependencies, which we see as a natural dynamic generalisation of integral data dependencies. The generalisation is provided by replacing integer functions with integer-valued variables as
Fig. 4.1. Values of the variables of $S$ for two inputs to the system and relative data dependence graphs.
the coefficients of the index mappings. Such a replacement introduces a form of dynamicity which depends on the inputs of the system of equations.

The main benefit of this approach is that regularisation techniques can be developed from those of integral data dependencies. In particular, we will show that the reconfigurability characteristics of the integral routing scheme that we have defined, provides the necessary flexibility for the treatment of this particular type of dynamic problems. There is, however, a restriction to their applicability, which is formally captured by the notion of separability between variables. Separability indicates that all computations necessary to resolve the dynamic data dependence relations can be separated and carried out prior to any other computation of the algorithm. Intuitively, if an algorithm is specified as a sequence of nested for-loops, separability allows us to reorganise the nested for-loops so that the control variables which define the dynamic routing of the data are computed initially, while all remaining nested for-loops follow. This type of restructuring of the computations implies that a further updating of the control signals is not possible. That is, once a dynamic data dependence relation has been established, it becomes static for the rest of the execution of the algorithm. From a mathematical point of view, the reorganisation of the computations is achieved through geometric translations of computation domains and a redefinition of the relative equations. That separability constitute a restriction for the regularisation of dynamic problems will become clear also from the case studies of Chapter 5.

For presentation purposes, the remainder of the chapter is structured similar to Chapter 3 for integral recurrences. Such an organisation facilitates the comparison between the two approaches. The main differences to keep in mind concern the notion of separability and the corresponding reorganisation of the computations. Other minor differences will be enlightened in the discussion.

4.3.1 Finitely Generated Index Mapping

A natural dynamic generalisation of an integral index mapping is the following (the reader may compare this definition with Definition 3.1.1 in Chapter 3), which replaces integer functions with integer-values variables as the coefficients of the index mapping:

**Definition 4.3.1** [Finitely Generated Index Mapping] Let $I$ be a dynamic index mapping. $I$ is finitely generated if for all $in \in Input$ and for all $z \in \mathbb{Z}^n$, $I[\text{in}](z) = z + \sum_{j=1}^{m} G_j[\text{in}](z)d_j,$ where, for $j = 1, \ldots, m$, $G_j$ is an integer-valued indexed variable and $d_j$ is a non-null vector.
in $\mathbb{Z}^n$. In addition, $I$ is atomic finitely generated if $m = 1$. \[4.3.1\]

Variables $G_j$ are called the coefficients of the index mapping and vectors $d_j$ its generators. Following the convention in Section 4.1.1, in the following, when possible, we will express such an index mapping $I$ more succinctly as $I(z) = z + \sum_{j=1}^{m} G_j(z) d_j$, i.e., we will leave the quantification over the inputs implicit.

Example 4.3.2 Consider the system of equations in Example 4.2.2. The index mapping $I_2$, defined as $I_2(i) = i - I(i)$, is finitely generated and, in particular, is atomic. The generator is the vector $(-1)$ with coefficient the integer-valued variable $I$.

Similar to the integral case we may define:

Definition 4.3.3 [Finitely Generated Dependence Mapping] Let $I$ be a finitely generated index mapping. Its dependence mapping $\Theta_I$ is such that, for all $i \in Input$ and for all $z \in \mathbb{Z}^n$, $\Theta_I[iin](z) = z - I[iin](z)$. \[4.3.3\]

If $I$ is expressed as $I(z) = z + \sum_{j=1}^{m} G_j(z) d_j$, its dependence mapping can be expressed as $\Theta_I(z) = - \sum_{j=1}^{m} G_j(z) d_j = \sum_{j=1}^{m} G_j(z) (-d_j)$. We call the vectors $-d_j$ the generators of $\Theta_I$ and the variables $G_j$ its coefficients.

4.3.2 Finitely Generated Data Dependence

Finitely generated index mappings characterise finitely generated dynamic data dependencies:

Definition 4.3.4 [Finitely Generated Data Dependence] Let $D = (D, U, V, I)$ be a dynamic data dependence. $D$ is finitely generated if $I$ is a finitely generated index mapping. In addition, $D$ is atomic finitely generated if $I$ is atomic finitely generated. \[4.3.4\]

The dependence domain and dependence cone of a finitely generated data dependence can be defined accordingly:

Definition 4.3.5 [Finitely Generated Dependence Domain] Let $D = (D, U, V, I)$ be a finitely generated data dependence and $\Theta_I$ the dependence mapping defined by $I$. The dependence domain $\Omega_I$ of $D$ is such that for all $iin \in Input$, $\Omega_I[iin] = \Theta_I[iin](D)$. \[4.3.5\]
Definition 4.3.6 [Finitely Generated Dependence Cone] Let $\mathcal{D} = (D, U, V, I)$ be a finitely generated data dependence with dependence domain $\Omega_I$. The dependence cone $\Theta^I_I$ of $\mathcal{D}$ is such that for all $in \in Input$, $\Theta^I_I[in] = cone(\Omega_I[in])$. 

Example 4.3.7 Consider the finitely generated data dependence $\mathcal{D} = (D, U, V, I)$ with index mapping $I(i,j) = (i,j) + G_1(i,j)(1,1) + G_2(i,j)(1,0)$ and domain $D = \{(i,j) \mid 1 \leq i,j \leq n\}$, for some $n \in N$. Assume that the coefficients of $I$ are defined by the input equations:

$$E_1 = (D, G_1, in_{G_1})$$
$$E_2 = (D, G_2, in_{G_2})$$

with $in_{G_1}(i,j) \in \mathbb{Z}$ and $in_{G_2}(i,j) = i + j$, for all $(i,j) \in D$.

The points of the dependence domains and the dependence cones of $\mathcal{D}$ for two inputs are illustrated in Fig. 4.2 (for $n = 3$). The values of the variables and of the index and dependence mappings for each point of $D$ are given in the related tables.

4.3.3 Dependence Cone and Pointedness

From a previous discussion (see Sections 2.2.6), we know that pointed dependence cones guarantee the existence of an affine scheduling. For an integral data dependencies this property was exploited as a guideline in the choice of the generators of the index mapping, and sufficient conditions were given for an integral dependence cone to be pointed and integral regularisation techniques to realise transformations of pointed dependence cones (see Sections 3.2.1 and 3.3).

In this section we investigate the possibility of formulating similar conditions for finitely generated data dependencies. The main difficulty here is that a dynamic dependence cone actually defines a family of dependence cones, one for each input, and that each such cone is known only when the algorithm is executed. Hence, in general, we cannot decide at compile time if all the cones of the family will be pointed. However, we may be able to guarantee that this is the case if we can make a number of assumptions on its finitely generated index mapping. The result is contained in the following proposition:
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\begin{align*}
(i, j) & \quad G_1(i, j) & \quad G_2(i, j) & \quad I(i, j) & \quad \Theta_I(i, j) \\
(1, 1) & 3 & 2 & (6, 4) & (-5.3) \\
(1, 2) & 2 & 3 & (6, 4) & (-5.2) \\
(1, 3) & 5 & 4 & (10, 8) & (-9.5) \\
(2, 1) & 1 & 3 & (6, 2) & (-4.1) \\
(2, 2) & 1 & 4 & (7, 3) & (-5.1) \\
(2, 3) & 2 & 5 & (9, 5) & (-7.2) \\
(3, 1) & 3 & 4 & (10, 4) & (-7.3) \\
(3, 2) & 2 & 5 & (10, 4) & (-7.2) \\
(3, 3) & 1 & 6 & (10, 4) & (-7.1)
\end{align*}

\begin{align*}
(i, j) & \quad G_1(i, j) & \quad G_2(i, j) & \quad I(i, j) & \quad \Theta_I(i, j) \\
(1, 1) & 2 & 2 & (5, 3) & (-4.2) \\
(1, 2) & 3 & 3 & (7, 5) & (-6.3) \\
(1, 3) & 2 & 4 & (7, 5) & (-6.2) \\
(2, 1) & 4 & 3 & (9, 5) & (-7.4) \\
(2, 2) & 1 & 4 & (7, 3) & (-5.1) \\
(2, 3) & 1 & 5 & (8, 4) & (-6.1) \\
(3, 1) & 5 & 4 & (11, 6) & (-9.5) \\
(3, 2) & 1 & 5 & (9, 3) & (-6.1) \\
(3, 3) & 2 & 6 & (11, 5) & (-8.2)
\end{align*}

Fig. 4.2. Dependence domains and cones for two inputs of $G_1$ and $G_2$. 
Proposition 4.3.8 Let \( DD = (D, U, V, I) \) be a finitely generated data dependence with index mapping \( I(z) = z + \sum_{j=1}^{m} G_j(z)d_j \). If:

- the coefficients \( G_j \) are non-negative over \( D \); and

- the embedding dependence cone \( C = \text{cone}(\{ -d_1, \ldots, -d_m \}) \) is pointed,

then, for all \( in \in \text{Input} \), \( \Theta^*[in] \) is pointed and contained in \( C \).

**Proof:** The result follows from the definition of \( C \). In fact, by definition \( C \) contains all points \( c \) in \( \mathbb{Z}^n \) such that \( c = \sum_{j=1}^{m} a_j(-d_j) \), with \( a_j \geq 0 \). For all inputs \( in \), \( \Omega_T[in] \) is defined as the set \( \{ \sum_{j=1}^{m} G_j[in](z)(-d_j) \mid z \in D \} \). If for all \( j \), \( G_j[in](z) \geq 0 \) then \( \Omega_T[in] \subseteq C \) and \( \Theta^*[in] \subseteq C \). Furthermore, because of the inclusion \( \Theta^*[in] \subseteq C \), pointed implies \( \Theta^*[in] \) pointed.

From Proposition 4.3.8, it follows that if \( \lambda \) is a non-null vector in \( \mathbb{Z}^n \), such that \( \lambda \cdot z > 0 \) for each element \( z \) of \( C \), then \( \lambda \cdot z > 0 \) for all \( z \in \Theta^*[in] \) and for all inputs \( in \). Therefore any affine timing function defined by \( \lambda \) is valid for all data dependence cones \( \Theta^*[in] \).

In the following we always assume that the conditions of Proposition 4.3.8 are satisfied. Note that if this assumption is not fulfilled, we can guarantee neither the existence of a timing function for all inputs, nor that our regularisation techniques will preserve affine scheduling.

**Example 4.3.9** Consider the data dependence of Example 4.3.7 and its dependence cones in Fig. 4.2 for two inputs of \( G_1 \) and \( G_2 \). The cone \( C = \text{cone}(\{(-1, -1), (-1, 0)\}) \), illustrated in Fig. 4.3, is pointed and contains both dependence cones.

**4.3.4 Finitely Generated vs. Integral Recurrences**

Given a finitely generated data dependence \( DD \), for all \( in \in \text{Input} \), \( DD[in] \) defines an integral data dependence. Vice versa, any integral data dependence may be regarded as a finitely
generated data dependence. This can be achieved, for instance, by replacing the coefficients of its index mapping with integer-valued variables, as explained below. Let \(DD = (D, U, V, I)\) be an integral data dependence, with index mapping \(I(z) = z + \sum_{j=1}^{m} g_j(z)d_j\). Then \(DD\) can be transformed in a dynamic data dependence by redefining its index mapping as \(I(z) = z + \sum_{j=1}^{m} G_j(z)d_j\), where each coefficient \(G_j\), for \(i = 1, \ldots, m\), is an integer-valued variable defined by the input equation \(E_j = (D, G_j, g_j)\). Note that this transformation is not just a simple syntactic manipulation of the index mapping (as, for instance, that we used to demonstrate that an affine index mapping is also integral – see Section 3.1.1). The change from static to dynamic is a semantic transformation, as the new interpretation modifies the possible models of the specification. Also, the transformation is not unique. Here we have chosen the simplest transformation we could think of. The reader may experiment other ways of interpreting an integral data dependence as a dynamic data dependence.

Note that, although theoretically an interpretation of a static data dependence as a dynamic data dependence is possible, from a practical point of view there is little convenience in doing so, as, in general, dynamic data dependencies require more complex (and often less efficient) forms of regularisation.

By recalling the taxonomy in Section 3.1.1, we may represent the relations between the various classes of recurrences which we have considered in this thesis, as illustrated in Fig. 4.4. The abbreviations used in the figure are: UREs, for uniform recurrences; AREs, for affine; AIREs and IREs, for atomic integral and integral recurrences; and FGREs, for finitely generated recurrences.

4.3.5 Extended Dependence Graph

By introducing variables in the index mappings, extra dependencies among the variables of a system of equations are generated. In fact, given a finitely generated data dependence \(DD = (D, U, V, I)\) with index mapping \(I(z) = z + \sum_{j=1}^{m} G_j(z)d_j\), for all \(z \in \mathbb{Z}^n\), the computation of
$U$ depends on the values of $G_j$ as well as the values of $V$. Let us define the set of variables of $I$ as $\text{Var}_I = \{G_1, \ldots, G_m\}$. For an (uninterpreted) static index mapping $I$ we assume $\text{Var}_I = \emptyset$. We define an extended version of the reduced dependence graph of a system of equation as follows:

**Definition 4.3.10 [Extended Dependence Graph]** Let $S$ be a system of equations. Define its extended dependence graph $\mathcal{EDG}$ as the graph $(\mathcal{N}, \mathcal{A})$, where:

- $\mathcal{N} = \text{Var}_S$; and
- $\mathcal{A} = \{(U, V) | \exists D, I \text{ such that } (D, U, V, I) \in \mathcal{DD}_S\}$
  $\cup \{(U, G) | \exists D, V, I \text{ such that } (D, U, V, I) \in \mathcal{DD}_S \text{ and } G \in \text{Var}_I\}$.

If no finitely generated index mapping is present in the system, its extended dependence graph is equal to its reduced dependence graph.

Given a system of equations $S$ and a variable $V$ of $S$, based on the notion of extended dependence graph, we can define the subsystem of equations defining $V$. In particular, let $V^A$ denote the set of the nodes of the graph which are accessible from $V$ (see Appendix D for a definition). Then the subsystem of equations defining $V$ is $\text{Def}_{SV} = \cup_{U \in V^A} \text{Def}_U$. Similarly, for a set of variables $V$, the subsystem of equations defining $V$ is $\text{Def}_{SV} = \cup_{V \in V} \text{Def}_V$.

### 4.3.6 Separability

Separability is a property defined between the variables of a system of equations which allows the identification of separable sets of computations. The notion of separability is based on the dependence relations existing among variables, instead of their instances, i.e., its formulation is based on the extended dependence graph of the system. The definition of separability relies upon the strong connectivity relation of graphs (see Appendix D for a formal definition). We define:

**Definition 4.3.11 [Separability]** Let $S$ be a system of equations, $\mathcal{EDG}$ its extended dependence graph and $U, V \in \text{Var}_S$. $U$ and $V$ are separable in $S$ if and only if $U$ and $V$ are not strongly connected in $\mathcal{EDG}$.
Example 4.3.12 Consider the system of equations of Example 4.2.2. Its extended dependence graph is given in Fig. 4.5 a). The equivalence classes of the nodes of the graph under strong connectivity are $\{[G], [Y], [I], [X]\}$. Therefore, all variables are pairwise separable in $S$.

In the following, given a finitely generated data dependence $\mathcal{DD} = (D, U, V, I)$, we will be interested in the separability of variable $U$ from the coefficients of the index mapping $I$. If these variables are separable, it will be possible to reorganise their computations in the lattice space and to define, similar to the integral case, a uniform and acyclic routing of the data which substitutes the data dependence $\mathcal{DD}$. Note that when an integral data dependence is regarded as finitely generated (with the interpretation of Section 4.3.4), this separation is always possible, as the coefficients of its index mapping are defined by input equations.

We will not be able to localise finitely generated data dependencies for which this separation is not possible. One such case is illustrated in the following example. The separability of the variables (or the lack of it) is the main restriction to the application of the regularisation techniques developed in this chapter.

Example 4.3.13 Consider the following system of equations $S'$, obtained by modifying that in Example 4.2.2:

\[
\begin{align*}
E_1 &= (D_1, Y, in_Y) \\
E_2 &= (D_2, G, Y, mod_4, I_1) \\
E_3 &= (D_2, X, in_X) \\
E_4 &= (D_2, I_1, (X, G), sub, (I_2, I_2)) \\
E_5 &= (D_2, Y, Y, inc, I_3)
\end{align*}
\]
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with domains \( D_1 = \{0, \ldots, 3\} \), \( D_2 = \{4, \ldots, n\} \), index mappings \( I_1(i) = i - 1 \), \( I_2(i) = i \), \( I_3(i) = i - I(i) \), and applied functions \( \text{mod}_4(a) = a \mod 4 \), \( \text{sub}(a, b) = a - b \), \( \text{inc}(a) = a + 1 \), and, on their domains, \( \text{in}_X(i) = i \) and \( \text{in}_Y(i) = y_i \in \mathbb{Z} \).

The extended dependence graph of the system is given in Fig. 4.5 b), and the corresponding strongly connected equivalence classes are \( \{[G, Y, I], [X]\} \). Note that \( Y \) and \( I \) are not separable in \( S' \).

4.4 Regularisation

The regularisation of finitely generated data dependencies aims at replacing a dynamic specification with a static uniform specification. Once this specification is obtained, classical mapping techniques apply for the derivation of an array design. The transformation from dynamic to static relies upon the definition of control variables which allow us to define a reconfigurable routing scheme for the data without having to alter the topology of the array design.

Regularisation techniques for finitely generated data dependencies extend those for integral data dependencies and have the form of uniformisation and decomposition. The main difference from the integral case stems from the necessity of redefining subsystems of computations of the specification, due to the definition of finitely generated index mappings, in which the coefficients are variables of the system. This characteristics of finitely generated index mappings implies that the corresponding dependence relations can be established only once such coefficients have been computed by the algorithm. Reorganising the computations guarantees that the coefficients are computed prior to their use. The way the computations are reorganised in the computation space of the algorithm is critical for the synchronisation of the data and the possibility of providing affine scheduling. In particular, as an effect of the transformation, new uniform dependence relations are introduced corresponding to acyclic data dependence graphs.

4.4.1 Translation of a System of Equations

We call translation of a system of equations the technique which allows us to reorganise the computations of a specification in its computation space. As the name indicates, the technique is based on a geometric translation (that is a simple type of affine transformation).

Systems of equations are translated in the lattice space by translating their computation
domains along some direction vector of the space. The technique is defined here with respect to an arbitrary direction vector. In the regularisation techniques of the following section, such a direction vector will correspond to one which is subsequently used for the uniform propagation of the data.

Technically, the translation of a system of equations is a combination of the translation of its domains together with a composition of index mappings which replicates on the translated domains the same data dependence relations of the original domains. In particular, if a computation point \( x \) in the original domain is data dependent on a point \( y \) under an index mapping \( \mathcal{I} \), a similar data dependence relation is established between the translated images of \( x \) and \( y \).

As a translation \( tr \) is just a mapping in \( \mathbb{Z}^n \), the translation of a system of equations can be defined by "lifting" \( tr \) from computation points to all the components of the system which are affected by the translation. This lifting is defined in Definition 4.4.1.

Usually we will have to combine a translation of the equations with a renaming of their variables. In fact, in general, the original system of equations is not removed from the algorithm specification, as there might exist other data dependencies in the specification relying on the same equations (this is a side-effect of applying regularisation techniques locally to each data dependence). Indeed all redundant equations may be eliminated once all the necessary regularisations have been completed.

**Definition 4.4.1 [Liftings]** Let \( tr : \mathbb{Z}^n \rightarrow \mathbb{Z}^n \) be a translation and \( tr^{-1} \) denote its inverse. Define the following liftings of \( tr \):

- Let \( D \subseteq \mathbb{Z}^n \). Define \( tr(D) = \{ tr(d) \mid d \in D \} \);

- Let \( \mathcal{I} \in [\text{Input} \rightarrow [\mathbb{Z}^n \rightarrow \mathbb{Z}^n]] \). Define \( tr(\mathcal{I}) \) such that for all \( in \in \text{Input} \), \( tr(\mathcal{I})[in] = tr \circ \mathcal{I}[in] \circ tr^{-1} \);

- Let \( \mathcal{I} \mathcal{M} \in [\text{Input} \rightarrow [\mathbb{Z}^n \rightarrow \mathbb{Z}^n]]^m \). Define \( tr(\mathcal{I} \mathcal{M}) = (tr(pr_1(\mathcal{I} \mathcal{M})), \ldots, tr(pr_m(\mathcal{I} \mathcal{M}))) \);

- Let \( E = (D_E, \star_E, E^*, f_E, \mathcal{I} \mathcal{M}_E) \). Define \( tr(E) = (tr(D_E), \star_E, E^*, f_E, tr(\mathcal{I} \mathcal{M}_E)) \);

- Let \( E = (D_E, \star_E, f_E) \). Define \( tr(E) = (tr(D_E), \star_E, f_E \circ tr^{-1}) \);

- Let \( S = \{ E_1, \ldots, E_s \} \). Define \( tr(S) = \{ tr(E_1), \ldots, tr(E_s) \} \).

Let \( ren : \text{Var} \rightarrow \text{Var} \) be a variable renaming. Define the following liftings of \( r \):
- Let \( V = (V_1, \ldots, V_m) \in \text{Var}^m \). Define \( \text{ren}(V) = (\text{ren}(V_1), \ldots, \text{ren}(V_m)) \).

- Let \( E = (D_E, E^*, f_E, \text{IM}_E) \). Define \( \text{ren}(E) = (D_E, \text{ren}(E^*), \text{ren}(E^*), f_E, \text{IM}_E) \).

- Let \( E = (D_E, E^*, f_E) \). Define \( \text{ren}(E) = (D_E, \text{ren}(E^*), f_E) \).

- Let \( S = \{E_1, \ldots, E_s\} \). Define \( \text{ren}(S) = \{\text{ren}(E_1), \ldots, \text{ren}(E_s)\} \).

By combining the renaming of the variables of a system of equations \( S \) with a translation of its equations, we obtain a new system of equations, which is an image of \( S \) with renamed variables and computations positioned differently from the computations of \( S \) in the lattice space.

**Definition 4.4.2** [Translated Image of a System of Equations] Let \( S \) be a system of equations. Let \( tr: \mathbb{Z}^n \to \mathbb{Z}^n \) be a translation and \( \text{ren}: \text{Var} \to \text{Var} \) an injective renaming of the variables of \( S \) with new names. Define the translated image of \( S \) according to \( tr \) and \( \text{ren} \) as \( S^{tr,\text{ren}} = \text{ren} \circ tr(S) \).

**Example 4.4.3** Consider the system \( S \) of equations of Example 4.2.2, and its sub-system \( \text{Def} S_I = \{E_2, E_3, E_4\} \), defining variable \( I \). The data dependence graph and reduced dependence graphs of \( \text{Def} S_I \) are illustrated in Fig. 4.6 a) and b), respectively. Let \( tr \) be the translation \( tr(i) = i - 5 \) and \( \text{ren} \) a renaming such that \( \text{ren}(I) = C, \text{ren}(G) = B \) and \( \text{ren}(X) = A \). Then \( (\text{Def} S_I)^{tr,\text{ren}} \) is the following system of equations (whose data dependence graph and reduced dependence graphs are illustrated in Fig. 4.6 c) and d), respectively):

\[
\begin{align*}
E'_2 &= (D_2^{tr}, B, \text{in}_B) \\
E'_3 &= (D_3^{tr}, A, \text{in}_A) \\
E'_4 &= (D_4^{tr}, C, (B, A), h, (I'_1, I'_1))
\end{align*}
\]

where \( D_2^{tr} = \{-1, \ldots, n-5\} \), \( \text{in}_B(i) = \text{in}_G(i+5), \text{in}_A(i) = \text{in}_X(i+5), h(a, b) = a - (b \mod 4) \), and \( I'_1(i) = I_1(i+5) - 5 = i \).

**4.4.2 Regularisation Directions**

Similar to the integral case, given a finitely generated data dependence, we will choose its regularisation directions among the generators of its index mapping. Because of the assumptions
made in Section 4.3.3, that the coefficients of the index mapping are non-negative variables and the cone defined by the generators of the dependence mapping is pointed, this choice will guarantee the preservation of affine scheduling. We will address this issue in Section 4.5.

4.4.3 Injectivity of an Atomic Finitely Generated Index Mapping

We can avoid data broadcasts by restricting ourselves to injective index mappings (see Section 2.3.2). Given an atomic finitely generated index mapping $I$ and a domain $D$ in $\mathbb{Z}^n$, the injectivity of the index mappings $I[in]$, for all inputs $in$, can be established by considering the geometric relation between $D$ and the generator of $I$. Indeed, this fact generalises the similar result for atomic integral index mappings (Proposition 3.2.4).

**Proposition 4.4.4** Consider an atomic finitely generated index mapping $I$ and a domain $D \subseteq \mathbb{Z}^n$, such that for all $z \in D$, $I(z) = z + G(z)d$. If $d \not\in \text{lin}(D)$ then for all $in \in Input$, $I[in]$ is injective over $D$.

**Proof:** For $in \in Input$, $I[in]$ is the integral index mapping defined by $I[in](z) = z + G[in](z)d$. From linear algebra, if $z, z' \in D$ then $z - z' \in \text{lin}(D)$. Assume that there exist $z, z' \in D$, such that $z \neq z'$ and $I[in](z) = I[in](z')$. We want to prove that this assumption always implies a contradiction with respect to the hypotheses of the proposition and therefore for all $z, z' \in D$, $z \neq z'$ implies $I[in](z) \neq I[in](z')$. There are only two possibilities, both leading to a contradiction. If $G[in](z) = G[in](z') = c$, then $I[in](z) = I[in](z')$ implies $z + cd = z' + cd$, i.e., $z = z'$. Otherwise, if
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Fig. 4.7. Injectivity of an atomic finitely generated index mapping.

\[ G[\text{in}](z) \neq G[\text{in}](z') \text{, then } G[\text{in}](z') - G[\text{in}](z) = c \neq 0 \text{ and } I[\text{in}](z) = I[\text{in}](z') \]

implies \( z - z' = cd \), i.e., \( d \in \text{lin}(D) \).

**Example 4.4.5** Consider the data dependence \( \mathcal{D} = (D, U, V, I) \), with index mapping \( I(i, j) = (i, j) + G(i, j)(1, 1) \), domain \( D = \{(i, j) \mid 1 \leq i \leq n, j = 2\} \), and \( G \) defined by the equation \( E = (D, G, \text{in}G) \), with \( \text{in}G(i, j) \in \mathbb{Z} \) for all \((i, j) \in D\).

The generator of \( I \) is \( d = (1, 1) \), \( \text{lin}(D) = \{(1, 0)\} \) and \( d \not\in \text{lin}(D) \) (see Fig. 4.7 a)). Hence, according to Proposition 4.4.4, all index mappings \( I[\text{in}] \), for all inputs, are injective on \( D \).

In fact, for all inputs \( \text{in} \), \( (i, j) + G[\text{in}](i, j)d \) belongs to the line \( \{(i, j) + ld \mid l \in \mathbb{R}\} \) (see Fig. 4.7 b)). In addition, such lines are parallel for all points in \( D \). This is illustrated in Fig. 4.7 c) and d) for two possible inputs of \( G \).

**4.4.4 Uniformisation**

Uniformisation substitutes an atomic finitely generated data dependence with a system of uniform data dependencies. Let \( \mathcal{D} = (D, U, V, I) \) be an atomic finitely generated data dependence, with \( I(z) = z + G(z)d \). The application of uniformisation techniques is subject to the following constraints:

- that the coefficient \( G \) of the index mapping \( I \) is bounded over \( D \). This restriction also applies to the integral case and its justification was discussed in Section 3.2.3;

- that variables \( G \) and \( U \) are separable. This property allows us to translate in the lattice space, the system of equations defining the variable \( G \), so that the computations of \( G \)
can be performed prior to those of $U$. The property also guarantees that no cycles are generated in the data dependence graph for effect of the translation.

Note that we are implicitly assuming that the index mapping has been specified correctly, hence that $G$ is defined at each point of $D$. If this is not the case, the assumption we made in Section 4.1 on the default value of an integer-valued indexed variable implies that our techniques are still well-defined. This fact should be taken into account if verification issues are considered. To this end, a different choice of default value, which allows to identify a condition of error, may be more appropriate.

As for integral uniformisation, in the finitely generated case two uniformisation techniques are defined based on the geometric relation between the direction vector $d$ and the domain $D$ of the data dependence.

The first technique is illustrated in Fig. 4.8 for a 2-dimensional case. In the figure, $G$ denotes the upper bound of $G$ on $D$. Routing directions and domains are chosen similar to the integral case, and two control variables, $\alpha$ and $\gamma$, are required. In the figure, the vector $\pi$ defines the hyperplane $[\pi: \theta]$ containing $D$ and $\eta = \pi \cdot d$. For each $z$ in $D$, $\alpha$ and $\gamma$ are initialised at the point $z + \tilde{G}d$. In contrast to the integral case, their initialisation requires the redefinition of the computations of $G$. For simplicity, let us assume that $D$ is the definition domain of $G$. Then its computations are redefined on the translated domain $D^{tr}$, shown in Fig. 4.8 b). A new variable $G^{tr}$ is used such that for each $z$ in $D$, $G^{tr}(z + \tilde{G}d)$ is defined to be the same as $G(z)$. The initialisation of $\alpha$ and $\gamma$ is obtained through variable $G^{tr}$ instead of $G$. The
Fig. 4.9. The need for the redefinition of the coefficient $G$.

Routing scheme is developed as for integral data dependencies. Fig. 4.8 c) and d) illustrate the routing scheme for two possible values of $G(z)$. That the scheme works for all inputs follows from the assumption that $G$ is a non-negative bounded variable on $D$. In fact, for all inputs $in$ and for all $z \in D$, $z + G[in](z)d$ lies on the segment $\{z + ld \mid 0 \leq l \leq \hat{G}\}$. On this segment the dynamic reconfiguration of the control variables $\alpha$ and $\gamma$ guarantees that, for all inputs, the right data is transferred. The redefinition of variable $G$ is necessary to avoid cycles in the resulting data dependence graph. If $G$ were not redefined as $G^{tr}$, the resulting data dependence graph would resemble the cyclic graph sketched in Fig. 4.9.

**Proposition 4.4.6 [Uniformisation 1]** Let $D \subseteq (D, U, V, I)$ be an atomic finitely generated data dependence, with $I(z) = z + G(z)d$. Let $G$ be non-negative and bounded over $D$ and let $\hat{G}$ be the least upper bound of $G$. Let $d \notin \text{lin}(D)$ and $P$ be the convex polyhedron generated by $D$ and $d$. Consider $\pi \in \text{lin}(P) \cap D^\perp$, with $\pi \neq 0$, and the hyperplane $[\pi : \theta]$ containing the domain $D$. Let $U$ and $G$ be separable. Consider the system of equations $(DefS_G)^{tr, ren}$, defined by the renaming $ren(G) = G^{tr}$ and the translation $tr(z) = z + \hat{G}d$.

Then $D \subseteq$ can be substituted by the uniform data dependence

$$D \subseteq' = (D, U, R, I_0)$$

and the system of equations:

$E_1 = (D_1, R, (R, V, \alpha, \gamma), f, (I_1, I_0, I_0, I_0))$

$E_2 = (D, U, \alpha, \alpha, id, I_1)$

$E_3 = (D, U, G^{tr}, id, I_0)$

$E_4 = (D_1, U, \gamma, \gamma, dec, I_1)$

$E_5 = (D_1, U, \gamma, in, \gamma)$

where:
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- the index mappings are:

\[ I_0(z) = z \]
\[ I_1(z) = z + d \]

- \( R, \alpha \) and \( \gamma \) are new variables;

- the applied functions are:

\[ i_{in}(z) = \tilde{G} \]
\[ id(a) = a \]
\[ dec(a) = a - 1 \]
\[ f(a, b, c, d) = \begin{cases} a & c \neq d \\ b & \text{otherwise} \end{cases} \]

- the new domains are:

\[ D_1 = \{ z + ld \mid z \in D, 0 \leq l \leq \tilde{G} \} \]
\[ D_{1,1} = \{ z + ld \mid z \in D, 0 \leq l < \tilde{G} \} \]
\[ D_{1,2} = \{ z + Gd \mid z \in D \} \]

PROOF: The proof relies upon the fact that, for all \( z \in D \) and for all inputs \( in, G[in](z) \in \{0, \ldots, \tilde{G}\} \).

For all \( z \in D \), let \( \text{segm}(z) = \{ z + ld \mid 0 \leq l \leq \tilde{G} \} \). By definition, for all \( z' \in \text{segm}(z) \),
\[ \alpha(z') = G(z). \]
In fact, for \( z \in D \),
\[ \alpha(z) = \alpha(z + d) \]
\[ = \ldots = \alpha(z + \tilde{G}d) = G^{\tilde{G}}(z + \tilde{G}d) = G(z). \]

Also for all \( z' = z + ld \in \text{segm}(z) \), \( \gamma(z') = l \). In fact, for \( z \in D \),
\[ \gamma(z) = \gamma(z + d) - 1 \]
\[ = \ldots = \gamma(z + \tilde{G}d) - \tilde{G} = \tilde{G} - \tilde{G} = 0. \]

Therefore, for all \( z' = z + ld \in \text{segm}(z) \), \( \alpha(z') = \gamma(z') \) if and only if \( l = G(z) \).

Hence, for all \( z \in D \),
\[ U(z) = R(I_0(z)) = R(I_1 \circ I_0(z)) \]
\[ = \ldots = R(I_1^{G(z)} \circ I_0(z)) = V(I_0 \circ I_1^{G(z)} \circ I_0(z)) \]
\[ = V(z + G(z)d) = V(I(z)). \]
Corollary 4.4.7 Let $\mathcal{D} = (D, U, V, I)$ be an atomic finitely generated data dependence, with $I(z) = z + G(z)d$. Let $G$ be non-negative and bounded over $D$ and $\tilde{G}$ the least upper bound of $G$.

If $\tilde{G} = 0$, then $\mathcal{D}$ can be substituted by the following uniform data dependence

$$\mathcal{D}_0 = (D, U, V, I_0)$$

where $I_0(z) = z$.

The second uniformisation technique consists of the combination of the corresponding integral techniques with the redefinition of the computations of the coefficient $G$ of the index mapping. Once again, for simplicity, let us assume that $D$ is the definition domain of $G$. The technique is illustrated in Fig. 4.10 for a 3-dimensional case. In the figure $D^{tr}$ indicates the translated domain and $G^{tr}$ the new variable defined as $G$ on $D^{tr}$. Fig. 4.10 c) and d) illustrate the routing scheme for two possible values of $G(z)$. Regularisation directions and domains, as well as routing and control variables are defined as for integral uniformisation.

Proposition 4.4.8 [Uniformisation 2] Let $\mathcal{D} = (D, U, V, I)$ be an atomic finitely generated data dependence, with $I(z) = z + G(z)d$. Let $G$ be non-negative and bounded over $D$, $m$ the least upper bound of $G$ and $\tilde{G} = [m/2]$. Let $d \in \text{lin}(D)$ and $\text{dim}(D) < n$. Consider $\pi \in D^d$, with $\pi \neq 0$, the hyperplane $[\pi : \theta]$ containing the domain $D$. and let $\hat{d} = d + \pi$ and $\check{d} = d - \pi$. Let $U$ and $G$ be separable. Consider the system of equations $(D \epsilon f S_G)^{tr, ren}$, defined by the renaming $\text{ren}(G) = G^{tr}$ and the translation $\text{tr}(z) = z + \tilde{G}\hat{d}$.
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Then $\mathcal{DD}$ can be substituted by the uniform data dependence

$$\mathcal{DD}' = (D, U, R^1, I_0)$$

and the system of equations:

$$
\begin{align*}
E_1 &= (D_1, R^1, (R^1, R^2, R^2, \alpha, \beta, \gamma).f.(I_1, I_{2.0}, I_{2.1}, I_0, I_0, I_0)) \\
E_2 &= (D_{2.1}, R^2, R^2, id, I_3) \\
E_3 &= (D_{2.2}, R^2, V, id, I_0) \\
E_4 &= (D_{1.1}, \alpha, \alpha, id, I_1) \\
E_5 &= (D_{1.2}, \alpha, G^{tr}, half\_floor, I_0) \\
E_6 &= (D_{1.1}, \beta, \beta, id, I_1) \\
E_7 &= (D_{1.2}, \beta, G^{tr}, mod_2, I_0) \\
E_8 &= (D_{1.1}, \gamma, \gamma, dec, I_1) \\
E_9 &= (D_{1.2}, \gamma, in_{\gamma})
\end{align*}
$$

where:

- the index mappings are:

$$
\begin{align*}
I_0(z) &= z \\
I_1(z) &= z + \dd \\
I_{2.0}(z) &= z \\
I_{2.1}(z) &= z + d \\
I_3(z) &= z + \dd
\end{align*}
$$

- $R^1, R^2, \alpha, \beta$ and $\gamma$ are new variables;

- the applied functions are:

$$
\begin{align*}
in_{\gamma}(z) &= \hat{G} \\
id(a) &= a \\
\text{half\_floor}(a) &= \lfloor a/2 \rfloor \\
\text{mod}_2(a) &= a \mod 2 \\
\text{dec}(a) &= a - 1 \\
f(a, b, c, d, e, f) &= \begin{cases} 
a & \quad d \neq f \\
b & \quad d = f, e = 0 \\
c & \quad d = f, e = 1 \end{cases}
\end{align*}
$$
- the new domains are:

\[
D_1 = \{z + l\hat{d} \mid z \in D, 0 \leq l \leq \hat{G}\}
\]

\[
D_{1,1} = \{z + l\hat{d} \mid z \in D, 0 \leq l < \hat{G}\}
\]

\[
D_{1,2} = \{z + \hat{G}\hat{d} \mid z \in D\}
\]

\[
D_2 = \{z + l_1d + l_2\hat{d} \mid z \in D_1, 0 \leq l_1 \leq 1, 0 \leq l_2 \leq \hat{G}\} \cap \{z \in \mathbb{Z}^n \mid \pi \cdot z \geq \theta\}
\]

\[
D_{2,1} = \{z \in D_2 \mid \pi \cdot z > \theta\}
\]

\[
D_{2,2} = \{z \in D_2 \mid \pi \cdot z = \theta\}
\]

**Proof:** The proof relies upon the fact that, for all \( z \in D \) and for all inputs \( \text{in} \), \( G[\text{in}](z) \in \{0, \ldots, \hat{G}\} \).

For all \( z \in D \), let \( \text{segm}(z) = \{z + ld \mid 0 \leq l \leq \hat{G}\} \). By definition, for all \( z' \in \text{segm}(z) \), \( \alpha(z') = [G(z)/2] \). In fact, for \( z \in D \),

\[
\alpha(z) = \alpha(z + \hat{d}) = \ldots = \alpha(z + \hat{G}\hat{d})
\]

\[
= \text{half}_\text{floor}(G_{tr}(z + \hat{G}\hat{d})) = \text{half}_\text{floor}(G(z)) = [G(z)/2].
\]

Also, for all \( z' \in \text{segm}(z) \), \( \beta(z') = G(z) \mod 2 \). In fact, for \( z \in D \),

\[
\beta(z) = \beta(z + \hat{d}) = \ldots = \beta(z + \hat{G}\hat{d})
\]

\[
= \text{mod}_2(G_{tr}(z + \hat{G}\hat{d})) = \text{mod}_2(G(z)) = G(z) \mod 2.
\]

Finally, for all \( z' = z + l\hat{d} \in \text{segm}(z) \), \( \gamma(z') = l \). In fact, for \( z \in D \),

\[
\gamma(z) = \gamma(z + \hat{d}) - 1
\]

\[
= \ldots = \gamma(z + \hat{G}\hat{d}) - \hat{G} = \hat{G} - \hat{G} = 0.
\]

Therefore, for all \( z' = z + ld \in \text{segm}(z) \), \( \alpha(z') = \gamma(z') \) if and only if \( l = [G(z)/2] \).

We observe that, for all \( c \in \mathbb{Z} \),

\[
|c/2| = (c - c \mod 2)/2
\]

\[
c = 2|c/2| + c \mod 2.
\]

Hence, for all \( z \in D \),

\[
U(z) = R^1(I_0(z)) = R^1(I_1 \circ I_0(z))
\]

\[
= \ldots = R^1(I_1^{[G(z)/2]} \circ I_0(z))
\]
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\[ R^2(I_2, G(z) \mod 2 \circ I_1^{[G(z)/2]} \circ I_0(z)) = R^2(I_3 \circ I_2, G(z) \mod 2 \circ I_1^{[G(z)/2]} \circ I_0(z)) = \ldots = R^2(I_3^{[G(z)/2]} \circ I_2, G(z) \mod 2 \circ I_1^{[G(z)/2]} \circ I_0(z)) = V(I_0 \circ I_3^{[G(z)/2]} \circ I_2, G(z) \mod 2 \circ I_1^{[G(z)/2]} \circ I_0(z)) = V(z + \lfloor G(z)/2 \rfloor d + \lfloor G(z) \mod 2 \rfloor d) = V(z + 2 \lfloor G(z)/2 \rfloor d + \lfloor G(z) \mod 2 \rfloor d) = V(z + (2 \lfloor G(z)/2 \rfloor + G(z) \mod 2) d) = V(z + G(z) d) = V(I(z)). \]

Corollary 4.4.9 Let \( \mathcal{D} = (D, U, V, I) \) be an atomic finitely generated data dependence, with \( I(z) = z + G(z) d \). Let \( G \) be non-negative and bounded over \( D \), \( m \) the least upper bound of \( G \), and \( \tilde{G} = \lfloor m/2 \rfloor \).

If \( \tilde{G} = 0 \) then \( \mathcal{D} \) can be substituted by the uniform data dependence:

\[ \mathcal{D}' = (D, U, R, I_0) \]

and the equations:

\[ E_1 = (D, R, (V, V, \beta), f, (I_0, I_1, I_0)) \]
\[ E_2 = (D, \beta, G, \text{mod}_2, I_0) \]

where:

- the index mappings are:
  \[ I_0(z) = z \]
  \[ I_1(z) = z + d \]

- \( R \) and \( \beta \) are new variables;

- the applied functions are:
  \[ \text{mod}_2(a) = a \mod 2 \]
  \[ f(a, b, c) = \begin{cases} a & c = 0 \\ b & c = 1 \end{cases} \]

\[ \square 4.4.9 \]
Example 4.4.10  Consider the system S of equations of Example 4.2.2. As $Y$ and $I$ are separable, we can apply uniformisation to the data dependence $\mathcal{DD} = (D_2, Y, Y, I_2)$, with $I_2(i) = i - I(i)$ and $D_2 = \{4, \ldots, n\}$.

As $\text{lin}(D) = \mathbb{Z}$, $d = (1) \in \text{lin}(D)$ and $\text{dim}(D) = 1$ we need to reindex (see Section 2.2.2) the equations in $\mathbb{Z}^2$ (so that $\text{dim}(D) < 2$). Let $\pi = (0,1)$, $\hat{d} = d + \pi = (-1,1)$ and $\hat{d} = d - \pi = (-1,-1)$. Note that $I(i) \leq n$ for all $i \in \{4, \ldots, n\}$. Consider the translation $\text{tr}(i,j) = (i,j) + \lfloor n/2 \rfloor \hat{d} = (i - \lfloor n/2 \rfloor, j + \lfloor n/2 \rfloor)$, and the variable renaming $\text{ren}$ such that $\text{ren}(G) = G^\text{tr}$, $\text{ren}(X) = X^\text{tr}$ and $\text{ren}(I) = I^\text{tr}$.

The application of Proposition 4.4.8 and the generation of the corresponding equations produces the following system:

\[
\begin{align*}
\text{(inputs of } Y) \\
E_1 &= (D_0, Y, inY) \\
\text{(translated sub-system of equations)} \\
E_2 &= (D^\text{tr}, G^\text{tr}, inG) \\
E_3 &= (D^\text{tr}, X^\text{tr}, inX) \\
E_4 &= (D^\text{tr}, I^\text{tr}, (X^\text{tr}, G^\text{tr}), h, (I_1, I_1)) \\
\text{(computations of } Y) \\
E_5 &= (D, Y, R^1, sqr, I_1) \\
\text{(routing variables } R^1, R^2) \\
E_6 &= (D_1, R^1, (R^1, R^2, R^2, \alpha, \beta, \gamma), f, (I_2, I_2, I_3, I_1, I_1, I_1)) \\
E_7 &= (D_2, R^2, R^2, id, I_4) \\
E_8 &= (D_2, R^2, Y, id, I_1) \\
\text{(control signals } \alpha, \beta, \gamma) \\
E_9 &= (D_1, \alpha, \alpha, id, I_2) \\
E_{10} &= (D_1, \alpha, I^\text{tr}, \text{half-floor}, I_1) \\
E_{11} &= (D_1, \beta, \beta, id, I_2) \\
E_{12} &= (D_2, \beta, I^\text{tr}, \text{mod}_2, I_1) \\
E_{13} &= (D_1, \gamma, \gamma, \text{dec}, I_2) \\
E_{14} &= (D_2, \gamma, in_{\gamma})
\end{align*}
\]

where:
the index mappings are:

\[ I_1(i, j) = (i, j) \]
\[ I_2(i, j) = (i, j) + \hat{d} = (i - 1, j + 1) \]
\[ I_3(i, j) = (i, j) + d = (i - 1, j) \]
\[ I_4(i, j) = (i, j) + \hat{d} = (i - 1, j - 1) \]

- the applied functions are \( \text{half\_floor}(a) = \lfloor a/2 \rfloor \), \( \text{mod}_2(a) = a \mod 2 \), \( \text{id}(a) = a \), \( \text{dec}(a) = a - 1 \), \( \text{in}_n(i, j) = \lfloor n/2 \rfloor \), \( h(a, b) = a - (b \mod 4) \), and

\[
f(a, b, c, \alpha, \beta, \gamma) = \begin{cases} 
  a, & \alpha \neq \gamma \\
  b, & \alpha = \gamma, \beta = 0 \\
  c, & \alpha = \gamma, \beta = 1
\end{cases}
\]

- the domains are:

\[ D_0 = \{(i, j) \mid 0 \leq i \leq 3, j = 0\} \]
\[ D = \{(i, j) \mid 4 \leq i \leq n, j = 0\} \]
\[ D^{tr} = \{(i, j) \mid 4 - \lfloor n/2 \rfloor \leq i \leq n - \lfloor n/2 \rfloor, j = \lfloor n/2 \rfloor\} \]
\[ D_1 = \{(i, j) \mid 4 - j \leq i \leq n - j, 0 \leq j \leq \lfloor n/2 \rfloor\} \]
\[ D_{1,1} = \{(i, j) \mid 4 - j \leq i \leq n - j, 0 \leq j \leq \lfloor n/2 \rfloor - 1\} \]
\[ D_{1,2} = \{(i, j) \mid 4 - \lfloor n/2 \rfloor \leq i \leq n - \lfloor n/2 \rfloor, j = \lfloor n/2 \rfloor\} \]
\[ D_2 = \{(i, j) \mid 2 - 2\lfloor n/2 \rfloor + j \leq i \leq n - j, 0 \leq j \leq \lfloor n/2 \rfloor\} \]
\[ D_{2,1} = \{(i, j) \in D_2 \mid j > 0\} \]
\[ D_{2,2} = \{(i, j) \in D_2 \mid j = 0\} \]

The resulting domains, data dependence graph and routing direction vectors are illustrated in Fig. 4.11 a), b) and c), respectively. A possible corresponding code segment is the following (that the reader may compare with that initially given in Example 4.2.2):

\[ n' := \text{floor}(n/2); \]

(* separated code segment *)

for \( i := 4 - n' \) to \( n - n' \) do

begin

\[ \text{read}(G^{tr}(i)); \]

\[ X^{tr}(i) := i + n'; \]
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\[ I^{tr}(i) := X^{tr}(i) - G^{tr}(i) \mod 4; \]
end;

(* control signals *)

for \( i := 4 - n' \) to \( n - n' \) do
begin
\[ \alpha(i, n') := \text{floor}(I^{tr}(i)/2); \]
\[ \beta(i, n') := I^{tr}(i) \mod 2; \]
\[ \gamma(i, n') := n'; \]

for \( j := 1 \) to \( n \) do
begin
\[ \alpha(i + j, n' - j) := \alpha(i + (j - 1), n' - (j - 1)); \]
\[ \beta(i + j, n' - j) := \beta(i + (j - 1), n' - (j - 1)); \]
\[ \gamma(i + j, n' - j) := \gamma(i + (j - 1), n' - (j - 1)) - 1; \]
end;
end;

(* inputs of \( Y \) and their routing *)

for \( i := 0 \) to \( 3 \) do

Fig. 4.11. a) Domains after uniformisation; b) Data dependence graph; c) routing direction vectors.
begin
read(Y(i));
R2(i, 0) := Y(i);
for j := 1 to floor(I(i)/2) do
    R2(i + j, j) := R2(i + j - 1, j - 1);
end;
(* computations of Y and their routing *)
for i := 4 to n do
    begin
        R1(i - n' - 1, n' + 1) := ⊥;
        for j := 0 to n' do
            if (α(i - n' + j, n' - j) ≠ γ(i - n' + j, n' - j))
                then R1(i - n' + j, n' - j) := R1(i - n' + j - 1, n' - j + 1);
            else if (β(i - n' + j, n' - j) = 1)
                then R1(i - n' + j, n' - j) := R2(i - n' + j - 1, n' - j);
                else R1(i - n' + j, n' - j) := R2(i - n' + j, n' - j);
        Y(i) := R1(i, 0);
        R2(i, 0) := Y(i);
        for j := 1 to n' do
            R2(i + j, j) := R2(i + j - 1, j - 1);
    end;

4.4.5 Parametric Uniformisation

Parametric uniformisation techniques can be used to reduce routing overhead by allowing a restricted amount of overloading of the data dependence graph (see Sections 2.3.2 and 3.2.4). A technique similar to that for atomic integral data dependencies can be developed for atomic finitely generated data dependencies, the main difference being the separation of the computations of the coefficient of the index mapping. The reader is referred to the discussion in Section 3.2.4 for a description of the technique.
**Proposition 4.4.11 [Parametric Uniformisation]** Let $\mathcal{DD} = (D, U, V, I)$ be an atomic finitely generated data dependence, with $I(z) = z + G(z)d$. Let $G$ be non-negative and bounded over $D$ and $m$ the least upper bound of $G$. Let $p \in \mathbb{N}^+$ and $\tilde{G} = \lfloor m/(p + 1) \rfloor$. Let $d \in \text{lin}(D)$ and $\dim(D) < n$. Consider $\pi \in D^\perp$, with $\pi \neq 0$, the hyperplane $[\pi : \theta]$ containing the domain $D$, and let $\tilde{d} = d + \pi$ and $\tilde{\pi} = d - \pi$. Let $U$ and $G$ be separable. Consider the system of equations $(DefS_G)^{tr, ren}$, defined by the renaming $ren(G) = G^{tr}$ and the translation $tr(z) = z + \tilde{G}\tilde{d}$.

Then $\mathcal{DD}$ can be substituted by the uniform data dependence

$$\mathcal{DD}' = (D, U, R^1, I_0)$$

and the system of equations:

$$\begin{align*}
E_1 &= (D_1, R^1, (R^1, R^2, \ldots, R^2, \alpha, \beta, \gamma), f.(I_1, I_{2.0}, \ldots, I_{2.p}, I_0, I_0, I_0)) \\
E_2 &= (D_{2.1}, R^2, R^3, id, I_3) \\
E_3 &= (D_{2.2}, R^2, V, id, I_0) \\
E_4 &= (D_{2.3}, R^3, R^2, id, I_4) \\
E_5 &= (D_{1.1}, \alpha, \alpha, id, I_1) \\
E_6 &= (D_{1.2}, \alpha, G^{tr}, (p + 1) \cdot \text{floor}, I_0) \\
E_7 &= (D_{1.1}, \beta, \beta, id, I_1) \\
E_8 &= (D_{1.2}, \beta, G^{tr}, \text{mod}_{p+1}, I_0) \\
E_9 &= (D_{1.1}, \gamma, \gamma, \text{dec}, I_1) \\
E_{10} &= (D_{1.2}, \gamma, \text{in}_n) \\
\end{align*}$$

where:

- the index mappings are:

$$\begin{align*}
I_0(z) &= z \\
I_1(z) &= z + \tilde{d} \\
I_{2.0}(z) &= z \\
I_{2.1}(z) &= z + d \\
\ldots \\
I_{2.p}(z) &= z + pd \\
I_3(z) &= z + \tilde{d}
\end{align*}$$
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$$I_4(z) = z + (p - 1)d$$

- $R_1, R_2, R_3, \alpha, \beta$ and $\gamma$ are new variables;

- the applied functions are:

$$in_{\gamma}(z) = \tilde{G}$$
$$id(a) = a$$
$$\ (p + 1).floor(a) = \lfloor a/(p + 1) \rfloor$$
$$mod_{p+1}(a) = a \ mod \ (p + 1)$$
$$dec(a) = a - 1$$
$$f(a, b_0, \ldots, b_p, c, d, e) = \begin{cases} a & c \neq e \\ b_0 & c = e, d = 0 \\ \ldots & \\ b_p & c = e, d = p \end{cases}$$

- the new domains are:

$$D_1 = \{ z + l d \mid z \in D, 0 \leq l \leq \tilde{G} \}$$
$$D_{1,1} = \{ z + l d \mid z \in D, 0 \leq l < \tilde{G} \}$$
$$D_{1,2} = \{ z + \tilde{G}d \mid z \in D \}$$
$$D_2 = \{ z + (l_1 + l_2)d + l_3 \tilde{d} \mid z \in D_1, 0 \leq l_1 \leq p, 0 \leq l_2 \leq (p - 1)\tilde{G}, 0 \leq l_3 \leq \tilde{G} \} \cap \{ z \in \mathbb{Z}^n \mid \pi \cdot z \geq \theta \}$$
$$D_{2,1} = \{ z \in D_2 \mid \pi \cdot z > \theta \}$$
$$D_{2,2} = \{ z \in D_2 \mid \pi \cdot z = \theta \}$$
$$D_{2,3} = \{ z \in D_2 \mid \pi \cdot z < \theta + \tilde{G}\eta \}.$$

PROOF: The proof relies upon the fact that for all $z \in D$ and for all inputs in, $G[\text{in}](z) \in \{0, \ldots, \tilde{G}\}$.

For all $z \in D$, let $\text{segm}(z) = \{ z + ld \mid 0 \leq l \leq \tilde{G} \}$. By definition, for all $z' \in \text{segm}(z)$.

$$\alpha(z') = \lfloor G(z)/(p + 1) \rfloor.$$ In fact, for $z \in D$,

$$\alpha(z) = \alpha(z + \tilde{d}) = \ldots = \alpha(z + \tilde{G}d)$$
$$= (p + 1).floor(G^{tr}(z + \tilde{G}d)) = (p + 1).floor(G(z)) = \lfloor G(z)/(p + 1) \rfloor.$$

Also, for all $z' = z + ld$ in the segment $\text{segm}(z)$, with $z \in D$, $\beta(z') = G(z) \ mod \ (p+1)$.

In fact, for $z \in D$,

$$\beta(z) = \beta(z + \tilde{d}) = \ldots = \beta(z + \tilde{G}d)$$
= \text{mod}_{p+1}(G''(z + \tilde{G}d)) = \text{mod}_{p+1}(G(z)) = G(z) \mod (p+1).

Finally, for all $z' = z + l\tilde{d} \in \text{segm}(z)$. $\gamma(z') = l$. In fact, for $z \in D$.

\[
\gamma(z) = \gamma(z + \tilde{d}) - 1
\]
\[
= \ldots = \gamma(z + \tilde{G}\tilde{d}) - \tilde{G} = \tilde{G} - \tilde{G} = 0.
\]

Therefore, for all $z' = z + ld \in \text{segm}(z)$. $\alpha(z') = \gamma(z')$ if and only if $l = [G(z)/(p + 1)]$.

We observe that, for all $c \in \mathbb{Z}$,

\[
[c/(p+1)] = (c - c \mod (p + 1))/(p + 1)
\]
\[
c = (p + 1)[c/(p + 1)] + c \mod (p + 1).
\]

Hence, for all $z \in D$,

\[
U(z) = R^1(I_0(z)) = R^1(I_1 \circ I_0(z))
\]
\[
= \ldots = R^1(I_1^{[G(z)/(p+1)]} \circ I_0(z))
\]
\[
= R^2(I_2,(G(z) \mod (p+1)) \circ I_1^{[G(z)/(p+1)]} \circ I_0(z))
\]
\[
= R^3(I_3 \circ I_2,(G(z) \mod (p+1)) \circ I_1^{[G(z)/(p+1)]} \circ I_0(z))
\]
\[
= R^2(I_4 \circ I_3 \circ I_2,(G(z) \mod (p+1)) \circ I_1^{[G(z)/(p+1)]} \circ I_0(z))
\]
\[
= \ldots
\]
\[
= R^2(I_4^{[G(z)/(p+1)]} \circ I_3^{[G(z)/(p+1)]} \circ I_2,(G(z) \mod (p+1)) \circ I_1^{[G(z)/(p+1)]} \circ I_0(z))
\]
\[
= V(I_0 \circ I_4^{[G(z)/(p+1)]} \circ I_3^{[G(z)/(p+1)]} \circ I_2,(G(z) \mod (p+1)) \circ I_1^{[G(z)/(p+1)]} \circ I_0(z))
\]
\[
= V(z + [G(z)/(p+1)]\tilde{d} + (G(z) \mod (p + 1))d + [G(z)/(p + 1)]\tilde{d} +
\]
\[
[G(z)/(p+1)](p-1)d
\]
\[
= V(z + ((p + 1)[G(z)/(p+1)] + G(z) \mod (p + 1))d
\]
\[
= V(z + G(z)d) = V(I(z)).
\]

\[\blacksquare\] 4.4.11

**Corollary 4.4.12** Let $DD = (D, U, V, I)$ be an atomic finitely generated data dependence, with $I(z) = z + G(z)d$. Let $G$ be non-negative and bounded over $D$ and $m$ the least upper bound of $G$. Let $p \in \mathbb{N}^+$ and $\bar{G} = [m/(p + 1)]$. 

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If $\bar{G} = 0$ then $DD$ can be substituted by the uniform data dependence

$$DD' = (D, U, R, I_0)$$

and the equations:

$$E_1 = (D, R, (V, \ldots, V, \beta), f, (I_{1,0}, \ldots, I_{1,p}, I_0))$$

$$E_2 = (D, \beta, G, \text{mod}_{p+1}, I_0)$$

where:

- the index mappings are:

  $$I_0(z) = z$$
  $$I_{1,0}(z) = z$$
  $$I_{1,1}(z) = z + d$$
  $$\ldots$$
  $$I_{1,p}(z) = z + pd$$

- $R$ and $\beta$ are new variables;

- the applied functions are:

  $$\text{mod}_{p+1}(a) = a \mod (p+1)$$
  $$f(a_0, \ldots, a_p, b) = \begin{cases} a_0 & b = 0 \\ a_1 & b = 1 \\ \ldots \\ a_p & b = p \end{cases}$$

4.4.6 Decomposition

Decomposition techniques allow us to substitute a finitely generated data dependence with a number of generators $m > 1$, with a corresponding finite set of atomic finitely generated data dependencies. Once again, the techniques derive from those for integral data dependencies by taking the separation of the coefficients of the index mappings into account. Note, however, that because of the translation of their subsystems of equations, no inverse mappings are needed in the transformations. In particular, let $DD = (D, U, V, T)$ be a finitely generated data dependence with index mapping $I(z) = z + G(z)d + \sum_{j=1}^{m} G_j(z)d_j$. Two decomposition techniques are developed.
The first technique applies when \( d \not\in \text{lin}(D) \) and is illustrated in Fig. 4.12 (in 2 dimensions). For all \( z \) in \( D \), \( I(z) \) and \( z + G(z)d \) assume different values for different inputs. Two of these values are illustrated in parts a) and b) of the figure. However, for all inputs, \( z + G(z)d \) lies on the segment \( \{ z + ld \mid 0 \leq l \leq \bar{G} \} \), where \( \bar{G} \) is an upper bound of \( G \) on \( D \). In Fig. 4.12 a) and b) the segment is represented as a dashed line through \( z \). The expression \( z + G(z)d \) defines the atomic finitely generated index mapping \( I_0 \) of the decomposition. New variables \( G_y \) are defined such that for all \( z \in D \), \( G(z)d \) assumes the same value as \( G_j(z) \). Their values are pipelined from \( z + \bar{G}d \) to \( z \) along the direction of \(-d\), and used in the definition of the finitely generated index mapping \( I_1 \) of the decomposition (see Fig. 4.12 c) and d)). Hence, an inverse of \( I_0 \) is not necessary for the definition of \( I_1 \) (as it is for integral decomposition — see Proposition 3.2.19).

**Proposition 4.4.13 [Decomposition 1]** Let \( DD = (D, U, V, I) \) be a finitely generated data dependence, with \( I(z) = z + G(z)d + \sum_{j=1}^{m} G_j(z)d_j \). Let \( G \) be non-negative and bounded over \( D \), and \( \bar{G} \) the least upper bound of \( G \). Let \( d \not\in \text{lin}(D) \) and \( P \) be the convex polyhedron generated by \( D \) and \( d \). Consider \( \pi \in \text{lin}(P) \cap D^\perp \), with \( \pi \neq 0 \), and the hyperplane \([\pi : \theta]\) containing the domain \( D \). Let \( U \) and \( G_j \) be separable, for all \( j \). Consider the system of equations \((\text{defS}_{G_1,\ldots,G_m})^{tr,\text{ren}}\), defined by the renaming \( \text{ren}(G_j) = G_j^{tr} \) and the translation \( \text{tr}(z) = z + \bar{G}d \).

Then \( DD \) can be substituted by the atomic finitely generated data dependence

\[
DD' = (D, I', R, I_0)
\]
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and the equations:

\[ E_1 = (D_1, R, V, id, I_1) \]
\[ E_{11} = (D_{1,1}, G'_j, G'_1, id, I_{2}) \]
\[ E_{12} = (D_{1,2}, G'_j, G'_{tr}, id, I_{3}) \]
\[ \ldots \]
\[ E_{m1} = (D_{1,1}, G'_j, G'_m, id, I_{2}) \]
\[ E_{m2} = (D_{1,2}, G'_j, G'_{tr}, id, I_{3}) \]

where:

- the index mappings are:
  \[ I_0(z) = z + G(z)d \]
  \[ I_1(z) = z + \sum_{j=1}^{m} G'_j(z)d_j \]
  \[ I_2(z) = z + d \]
  \[ I_3(z) = z \]

- \( R \) and \( G'_j \), for \( j = 1, \ldots, m \), are new variables;

- the applied function is \( id(a) = a \);

- the new domains are:
  \[ D_1 = \{ z + ld \mid z \in D, 0 \leq l \leq \bar{G} \} \]
  \[ D_{1,1} = \{ z + ld \mid z \in D, 0 \leq l < \bar{G} \} \]
  \[ D_{1,2} = \{ z + \bar{G}d \mid z \in D \} \]

**PROOF:** In the proof, we use the condition \( 0 \leq G(z) \leq \bar{G} \) which is satisfied for all inputs.

For all \( z \in D \), let \( \text{segm}(z) = \{ z + ld \mid 0 \leq l \leq \bar{G} \} \). By definition, for all \( z' \in \text{segm}(z) \),
\( G'_j(z') = G_j(z) \). In fact, for all \( z \in D \),
\[ G'_j(z) = G'_j(z + d) = \ldots = G'_j(z + \bar{G}d) = G'_j(z + \bar{G}d) = G_j(z). \]

Therefore, for all \( z \in D \),
\[ I_1 \circ I_0(z) = z + G(z)d + \sum_{j=1}^{m} G'_j(z + G(z)d)d_j \]
\[ = \sum_{0 \leq G(z) \leq \bar{G}} z + G(z)d + \sum_{j=1}^{m} G_j(z)d_j = I(z). \]
Finally, for all \( z \in D \),

\[
U(z) = R(I_0(z)) = V(I_1 \circ I_0(z)) = V(I(z)).
\]

The second technique extends the scheme discussed for the previous technique to the case when the generator \( d \) is contained in \( \text{lin}(D) \). It combines integral decomposition (see Proposition 3.2.21) with the redefinition of the coefficients \( G_1, \ldots, G_m \). The technique is illustrated in Fig. 4.12 (in 3-dimensions) for two possible values of \( I(z) \) (\( I[z] \) in part a) and \( I[z'][z] \) in part b).

**Proposition 4.4.14** [Decomposition 2] Let \( DD = (D, U, V, I) \) be a finitely generated data dependence, with \( I(z) = z + G(z)d + \sum_{j=1}^{m} G_j(z)d_j \). Let \( G, G_j \) be non-negative and bounded over \( D \) and let \( \bar{G}, \bar{G}_j \) be the least upper bounds of \( G, G_j \), respectively. Let \( d \in \text{lin}(D), P \) be the convex polyhedron generated by \( D \) and \( \{d_1, \ldots, d_m\} \) and \( \text{dim}(P) < n \). Consider \( \pi \in P_+^{\perp} \) with \( \pi \neq 0 \), and the hyperplane \( [\pi : 0] \) containing the domain \( D \), and let \( \bar{d} = d + \pi \). Let \( U \) and \( G_j \) be separable, for all \( j \). Consider the system of equations \((DefS_{(G_1, \ldots, G_m)})^{\text{tr}, \text{ren}}, \) defined by the renaming \( \text{ren}(G_j) = G_j^{\text{tr}} \) and the translation \( \text{tr}(z) = z + G\bar{d} \).
Then $DD$ can be substituted by the data dependence

$$DD' = (D, U, R^1, I_0)$$

and the equations:

$$E_1 = (D_1, R^1, R^2, id, I_1)$$
$$E_2 = (D_2, R^2, V, id, I_2)$$
$$E_{11} = (D_{1,1}, G'_j, G'_1, id, I_3)$$
$$E_{12} = (D_{1,2}, G'_j, G'_{1r}, id, I_4)$$

$$E_{m_1} = (D_{1,1}, G'_j, G'_{m_1}, id, I_3)$$
$$E_{m_2} = (D_{1,2}, G'_j, G'_{m_2}, id, I_4)$$
$$E_3 = (D_2, G', in_{GR})$$

where:

- the index mappings are:

  $$I_0(z) = z + G(z)\hat{d}$$
  $$I_1(z) = z + \sum_{j=1}^{m} G'_j(z)d_j$$
  $$I_2(z) = z + G'(z)(-\pi)$$
  $$I_3(z) = z + \hat{d}$$
  $$I_4(z) = z$$

- $R^1, R^2, G'$ and $G'_j$, for $j = 1, \ldots, m$, are new variables;

- the applied functions are:

  $$in_{GR}(z) = (\pi \cdot z - \theta)/(\pi \cdot \hat{d})$$
  $$id(a) = a$$

- the new domains are:

  $$D_1 = \{ z + l\hat{d} | z \in D, 0 \leq l \leq \bar{G} \}$$
  $$D_{1,1} = \{ z + l\hat{d} | z \in D, 0 \leq l < \bar{G} \}$$
  $$D_{1,2} = \{ z + \bar{G}\hat{d} | z \in D \}$$
  $$D_2 = \{ z + \sum_{j=1}^{m} l_jd_j | z \in D_1, 0 \leq l_j \leq \bar{G}_j \}$$
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PROOF: In the proof, we use the condition $0 \leq G(z) \leq \bar{G}$, which is satisfied for all inputs.

For all $z \in D$, let $\text{segm}(z) = \{z + ld \mid 0 \leq l \leq \bar{G}\}$. By definition, for all $z' \in \text{segm}(z)$, $G_j(z') = G_j(z)$. In fact, for all $z \in D$,

$$G_j'(z) = G_j'(z + \hat{d}) = \ldots = G_j'(z + \bar{G}\hat{d}) = G_j^{tr}(z + \bar{G}\hat{d}) = G_j(z).$$

Therefore, for all $z \in D$,

$$I_1 \circ I_0(z) = z + G(z)\hat{d} + \sum_{j=1}^{m} G_j'(z + G(z)\hat{d})d_j$$

$$= [0 \leq G(z) \leq G] \quad z + G(z)\hat{d} + \sum_{j=1}^{m} G_j(z)d_j.$$

Also, as $\pi \in P^\perp$, then $\pi \cdot d_j = 0$ and $\pi \cdot (\sum_{j=1}^{m} G_j(z)d_j) = 0$. Hence, for all $z \in D$,

$$G'(I_1 \circ I_0(z)) = \text{in}G'(I_1 \circ I_0(z))$$

$$= (\pi \cdot (I_1 \circ I_0(z)) - \theta)/(\pi \cdot \hat{d})$$

$$= (\pi \cdot (z + G(z)\hat{d} + \sum_{j=1}^{m} G_j(z)d_j) - \theta)/(\pi \cdot \hat{d})$$

$$= (\pi \cdot z + G(z)\pi \cdot \hat{d} + \pi \cdot (\sum_{j=1}^{m} G_j(z)d_j) - \theta)/(\pi \cdot \hat{d})$$

$$= (\theta + G(z)\pi \cdot \hat{d} + 0 - \theta)/(\pi \cdot \hat{d}) = G(z).$$

Therefore, for all $z \in D$,

$$I_2 \circ I_1 \circ I_0(z) = I_2(I_1 \circ I_0(z))$$

$$= I_1 \circ I_0(z) + G'(I_1 \circ I_0(z))(-\pi)$$

$$= I_1 \circ I_0(z) + G(z)(-\pi)$$

$$= z + G(z)\hat{d} + \sum_{j=1}^{m} G_j(z)d_j + G(z)(-\pi)$$

$$= z + G(z)(\pi + d - \pi) + \sum_{j=1}^{m} G_j(z)d_j$$

$$= z + G(z)\hat{d} + \sum_{j=1}^{m} G_j(z)d_j = I(z).$$

Finally, for all $z \in D$,

$$U(z) = R^1(I_0(z)) = R^2(I_1 \circ I_0(z))$$

$$= V(I_2 \circ I_1 \circ I_0(z)) = V(I(z)).$$

From the previous to results, we obtain the corollary:
Corollary 4.4.15  Let $\mathcal{DD} = (D, U, V, I)$ be a finitely generated data dependence, with $I(z) = z + G(z)d + \sum_{j=1}^{m} G_j(z)d_j$. Let $G$ be non-negative and bounded over $D$ and $\dot{G}$ the least upper bound of $G$.

If $\dot{G} = 0$, then $\mathcal{DD}$ can be substituted by the data dependence:

$$\mathcal{DD}_0 = (D, U, V, I_0)$$

where $I_0(z) = z + \sum_{j=1}^{m} G_j(z)d_j$.  \hfill 4.4.15

### 4.5 Regularisation and Affine Scheduling

The preservation of affine scheduling by the regularisation techniques of the previous section can be proved by applying Propositions 3.3.1 and 3.3.2. The results are summarised in Table 4.1, where: the first column indicates each regularisation technique with, in brackets, the reference number of the corresponding formal statement; the generators of the embedding dependence cone before $(C)$ and after $(C')$ the application of each technique are given in the second and third columns, respectively. In particular, in such columns $r = -d$ and $r_j = -d_j$, for all $j = 1, \ldots, m$, denote the generators of the corresponding dependence cone, while $\dot{r} = -\dot{d}, \ddot{r} = -\ddot{d}$ and $\rho = -\pi$ the new generators introduced by the various techniques; the last column indicates which of Propositions 3.3.1 and 3.3.2 applies. A line $-$ indicates that no proposition applies and corresponds to a case in which the dependence cone is not modified by the regularisation technique.

Note that Propositions 3.3.1 and 3.3.2 apply to the dynamic case because of the assumption that the coefficients of a finitely generated index mapping are non-negative and bounded variables. In turn, this condition guarantees that for all inputs, the corresponding integral index mappings are characterised by coefficients which are also non-negative and bounded.
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### Table 4.2. Summary of regularisation techniques.

<table>
<thead>
<tr>
<th>Technique</th>
<th>Condition</th>
<th>Extra Dimensions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unif. (4.4.6)</td>
<td>$d \notin \text{lin}(D)$</td>
<td>no</td>
</tr>
<tr>
<td>Unif. (4.4.8)</td>
<td>$d \in \text{lin}(D)$, $\dim(D) &lt; n$</td>
<td>yes</td>
</tr>
<tr>
<td>Par. Unif. (4.4.11)</td>
<td>$d \in \text{lin}(D)$, $\dim(D) &lt; n$</td>
<td>yes</td>
</tr>
<tr>
<td>Decomp. (4.4.13)</td>
<td>$d \notin \text{lin}(D)$</td>
<td>no</td>
</tr>
<tr>
<td>Decomp. (4.4.14)</td>
<td>$d \in \text{lin}(D)$, $\dim(P) &lt; n$</td>
<td>yes</td>
</tr>
</tbody>
</table>

In order to guarantee the preservation of an affine scheduling for the whole specification we can consider (as we did for integral problems in Section 3.3) the dependence cone of the corresponding system of equations. Let $S$ be such a system, and $C_S$ its dependence cone. Then, when a regularisation technique is applied, the normal vector ($\pi$ in the formal statements) to the hyperplane at the basis of the transformation can be chosen in the space $(C_S)^\perp$. This choice guarantees both that $\pi \in C^\perp$ and that affine schedulings are preserved for the system $S$.

### 4.6 Summary

The main objective of this chapter was to demonstrate that classes of dynamic problems can be transformed systematically into regular arrays in the context of classic synthesis methods. We have reached our objective by characterising a type of dynamic data dependencies (those which vary with the inputs of an algorithm) and identifying a subclass which can be represented and treated within Euclidean synthesis.

Both the formalisation and treatment of this type of problems was based on a generalisation of integral index mappings, in which indexed variables replace integer functions as the coefficients of the mappings. The implications of this substitution were enlightened. In particular the notions of extended dependence graph and variable separability were developed, with variable separability as the primary condition for the application of the regularisation techniques.

The regularisation techniques that we have developed are summarised in Table 4.2. In the table, the first column indicates the technique and in brackets the reference number of the corresponding formal result. The second column indicates the basic conditions for its appli-
cability. In particular, $d$ denotes a generator of the index mapping, $D$ the domain of the data dependence, and $P$ the polyhedron generated by $d$ and $D$. The last column indicates whether the application of the technique may require an increase in the number of dimensions of the space.

A major difference with respect to integral techniques concerns the necessity of redefining the subsystems of equations relative to the coefficients of the index mappings in the computation space of the problem. This redefinition is necessary to reconfigure the data routing scheme through the initialisation of its control variables. This also limits the applicability of the techniques, and many interesting problems from the literature cannot be treated with our methods. One of these problems, that of Gaussian elimination with pivoting, will be discussed in the next chapter.
Chapter 5

Case Studies

In this chapter we will consider a number of problems from the literature in order to illustrate the application, and limitations, of the techniques which we have developed in the previous chapters.

The first problem considered is a cyclic reduction algorithm for the solution of tridiagonal systems [Mod88, LoZa94]. A tridiagonal system is a system of linear equations whose matrix of coefficients is tridiagonal, i.e., the non-null entries of the matrix are all concentrated around the main diagonal in a band of width equal to 3. The solution of tridiagonal systems is at the base of several numerical applications, such as the solution of systems of partial differential equations [BuFa93]. Cyclic reduction algorithms for the solution of tridiagonal systems are known from the literature [Mod88, LoZa94]. In this chapter, we will show how the problem can be synthesised with our techniques through its specification as a system of integral recurrence equations.

The difficulties posed by integral data dependencies with unbounded coefficients will be discussed by considering the example of a decimation filter. Decimation filters are used for the conversion of the sampling rates of digital signals and constitute significant applications in digital signal processing [CrRa83]. We will show that the problem can be specified as a system of integral recurrence equations (the problem is, in particular, affine). However, because of the unbounded coefficients of its data dependencies, our techniques are not applicable. Instead, a simple reformulation of the problem will be given directly as a system of uniform recurrence equations. We will also show that not even affine regularisation techniques apply to the problem in its first specification.

As an example of dynamic problem we will consider the well-studied knapsack problem [MaTo90, Hu82], an NP-hard combinatorial optimisation problem both of theoretical and
practical interest. We will show that a treatment of the knapsack problem as a dynamic finitely generated problem is possible and that our techniques can be applied for the systematic derivation of corresponding regular array designs.

Finally, we will consider the technique of Gaussian elimination with partial pivoting [Mod88] [BaEl88, Ho-et-al89, ElBa90, Meg90] for the reduction of a dense matrix to a triangular form. The addition of pivoting to the elimination process increases its numerical stability and avoids breakdown due to division by zero. Pivoting, however, introduces dynamic data dependencies which cannot be handled by classic synthesis methods. When formulated as a dynamic problem, Gaussian elimination with pivoting is characterised by computations which cannot be separated. Therefore our techniques are not applicable. Instead, a static specification of the problem will be given, corresponding to regular array designs which have appeared in the literature [Meg90].

We have summarised the case studies in Table 5.1. The table indicates the problem, the type of the corresponding specification, whether our regularisation techniques apply and, if not, the condition that prevents their application.

### 5.1 Cyclic Reduction

The first problem that we consider is an algorithm for the solution of tridiagonal systems of equations through cyclic reduction. The following formulation of the problem is based on [Sa-et-al93]. Let $N = 2^r - 1$, for some integer $r$. A tridiagonal set of $N$ irreducible linear equations is:

\[
\begin{align*}
  a_{i-1} s_{i-2} + b_{i-1} x_{i-1} + c_{i-1} x_i &= d_{i-1} \\
  a_i s_i + b_i x_i + c_i x_{i+1} &= d_i \\
  a_{i+1} s_i + b_{i+1} x_{i+1} + c_{i+1} x_{i+2} &= d_{i+1}
\end{align*}
\]
for \( i = 2, 4, 6, \ldots, N - 1 \), with non-null coefficients and boundary values \( x_0 = x_{N+1} = 0 \).

The solution of this set of equations with cyclic reduction consists of a so-called reduction phase followed by a backsubstitution phase. The reduction phase aims at reducing the system to a single equation in a single variable, whose value can be determined with a single arithmetic operation. The backsubstitution phase recursively substitutes the variable values already determined in the sets of equations generated in the reduction phase, until all variable values are determined.

The reduction phase includes \( r - 1 \) reduction steps. Let \( j \) denote the current step, with \( j = 1, 2, \ldots, r - 1 \), and let \( g(j) = 2^j \). Each reduction step \( j \) consists of the computations of the following coefficients, for \( i = 2g(j), 4g(j), \ldots, 2r - 2g(j) \):

\[
\begin{align*}
a_i^j &= \frac{-a_i^{j-1} a_i^{j-1}}{b_i^{j-1}} \\
b_i^j &= b_i^{j-1} + \frac{-a_i^{j-1} c_i^{j-1}}{b_i^{j-1}} + \frac{-c_i^{j-1} a_i^{j-1}}{b_i^{j-1}} \\
c_i^j &= \frac{-c_i^{j-1} c_i^{j-1}}{b_i^{j-1}} \\
d_i^j &= d_i^{j-1} + \frac{-a_i^{j-1} d_i^{j-1}}{b_i^{j-1}} + \frac{-c_i^{j-1} d_i^{j-1}}{b_i^{j-1}}
\end{align*}
\]

The initial values are \( a_0^0 = a_i, b_0^0 = b_i, c_0^0 = c_i \) and \( d_0^0 = d_i \).

At the end of the reduction phase, the remaining equation is:

\[
a_{2g(r-1)}^{r-1} x_0 + b_{2g(r-1)}^{r-1} x_{2g(r-1)} + c_{2g(r-1)}^{r-1} x_{4g(r-1)} = d_{2g(r-1)}^{r-1}.
\]

Note that \( 4g(r-1) = 4 \times 2^{r-2} = 2^r = N + 1 \), hence \( x_{4g(r-1)} = x_{N+1} \). Therefore, because of the boundary conditions \( x_0 = x_{N+1} = 0 \), the value of variable \( x_{2g(r-1)} \) can be obtained from the above equation in a single operation.

The other values of \( x \) can be obtained in \( r - 1 \) backsubstitution steps. Once again, let \( j \) denote the current step, with \( j = r - 1, r - 2, \ldots, 1 \), and let \( g'(j) = 2^j \). Then, at each step \( j \), and for \( i = g'(j), 3g'(j), \ldots, 2^r - g'(j) \), \( x_i \) can be determined as:

\[
x_i = \frac{d_i^j - a_i^j x_i-g'(j) - c_i^j x_{i+g'(j)}}{b_i^j}
\]

Note that the two phases are strictly sequential and can be treated separately.
5.1.1 Specification

Let us specify the two phases of the algorithm separately.

Reduction Phase

The formulation of the reduction phase in the previous section was that of a system of recurrence equations, with coefficients indexed by two indices $i$ and $j$. We can specify this phase in $\mathbb{Z}^2$ by introducing the variables $A, B, C$ and $D$, corresponding to the coefficients of the equations.

The main difficulty with this specification is that the computation points do not constitute convex polyhedral domains. We solve the problem by embedding the computation points in a convex polyhedral region $D_1$ and defining a control variable $T$ to identify these computation points in $D_1$. The domain $D_1$ is defined as:

$$D_1 = \{(i, j) | 1 \leq j \leq r - 1, 2 \leq i \leq 2^r - 2\}$$

while the definition of $T$ is based on the function:

$$f(i, j) = \begin{cases} 1 & i/2^{j-1} \in \{2, 4, 6, \ldots, 2^r - 2\} \\ 0 & \text{otherwise} \end{cases}$$

We make variables $A, B, C$ and $D$ undefined (equal to $\perp$) at each $(i, j) \in D_1$ such that $f(i, j) = 0$. The specification is:

\begin{align*}
E_1 &= (D_0, A, in_A) \\
E_2 &= (D_1, A, (T, A, A, B), f_1, (I_0, I_1, I_2, I_3)) \\
E_3 &= (D_0, B, in_B) \\
E_4 &= (D_1, B, (T, B, A, C, B, C, A, B), f_2, (I_0, I_1, I_2, I_3, I_4, I_5)) \\
E_5 &= (D_0, C, in_C) \\
E_6 &= (D_1, C, (T, C, C, C, B), f_1, (I_0, I_1, I_2, I_3)) \\
E_7 &= (D_0, D, in_D)
\end{align*}
\[ E_8 = (D_1, D, (T, D, A, D, B, C, D, B), f_2, (I_0, I_1, I_2, I_2, I_1, I_3, I_3)) \]

(control variable \( T \))

\[ E_9 = (D_1, T, f) \]

with:

- domains:

\[
D_0 = \{(i, j) | j = 0, 1 \leq i \leq 2^r - 1\}
\]

\[
D_1 = \{(i, j) | 1 \leq j \leq r - 1, 2 \leq i \leq 2^r - 2\}
\]

- index mappings:

\[
I_0(i, j) = (i, j)
\]

\[
I_1(i, j) = (i, j - 1)
\]

\[
I_2(i, j) = (i - g(i, j), j - 1)
\]

\[
I_3(i, j) = (i + g(i, j), j - 1)
\]

where \( g(i, j) = 2^{j-1} \); and

- applied functions:

\[
in_A(i, j) = a_i
\]

\[
in_B(i, j) = b_i
\]

\[
in_C(i, j) = c_i
\]

\[
in_D(i, j) = d_i
\]

\[
f_1(t, a, b, c) = \begin{cases} -a * b / c & t = 1 \\ 1 & \text{otherwise} \end{cases}
\]

\[
f_2(t, a, b, c, d, e, f, g) = \begin{cases} a * b * c / d - e * f / g & t = 1 \\ 1 & \text{otherwise} \end{cases}
\]

\[
f(i, j) = \begin{cases} 1 / 2^{j-1} \in \{2, 4, 6, \ldots, 2^r - 2\} \\ 0 & \text{otherwise} \end{cases}
\]

A picture of the computation points of the phase is given in Fig. 5.1 a), for \( r = 4 \).

**Backsubstitution Phase**

The backsubstitution phase of the algorithm can also be formulated as a system of recurrences in \( \mathbb{Z}^2 \). To this end we introduce a variable \( X \), corresponding to \( x \), which we full index in \( \mathbb{Z}^2 \).
(in the previous section \( x \) is indexed by a single index \( i \)). Let us consider the domains:

\[
D'_0 = \{ (i,j) \mid j = r, 0 \leq i \leq 2^r \} \\
D'_1 = \{ (i,j) \mid 0 \leq j \leq r-1, 0 \leq i \leq 2^r \}
\]

and the control variable \( T' \) defined on \( D'_1 \) by the function:

\[
f'(i,j) = \begin{cases} 
2^j / i & i \in \{1, 3, 7, \ldots, 2^r - 1\} \\
0 & \text{otherwise}
\end{cases}
\]

Then the backsubstitution phase may be specified as:

\[
\begin{align*}
E'_1 &= (D'_0, X, \text{in}_x) \\
E'_2 &= (D'_1, X, (T', X, D, A, X, C, X, B), f, (I_0, I'_1, I_0, I'_0, I'_2, I_0, I'_3, I_0)) \\
E'_3 &= (D'_1, T', f')
\end{align*}
\]

with index mappings:

\[
\begin{align*}
I'_0(i,j) &= (i,j) \\
I'_1(i,j) &= (i,j+1) \\
I'_2(i,j) &= (i-g'(i,j), j+1) \\
I'_3(i,j) &= (i+g'(i,j), j+1)
\end{align*}
\]

where \( g'(i,j) = 2^j \), and applied functions:

\[
\text{in}_x(i,j) = 0
\]
Fig. 5.2. Backsubstitution phase: a) computation points; b) data dependence graph.

\[ f(t', a, b, c, d, e, f, g) = \begin{cases} \frac{(b - c + d - e + f)}{g} & t' = 1 \\ a & \text{otherwise} \end{cases} \]

\[ f'(i, j) = \begin{cases} 1 & 2^j/i \in \{1, 3, 7, \ldots, 2^r - 1\} \\ 0 & \text{otherwise} \end{cases} \]

The values \( x_i \) correspond to the values \( X(i, 0) \), for all \( i = 0, 1, \ldots, 2^r \). The computation points of this phase are given in Fig. 5.2 a), for \( r = 4 \). In the figure, black nodes indicate points at which values \( x_i \) are actually computed, while white nodes the points at which already computed \( x_i \) are pipelined. Note the similarity existing between the two phases, as illustrated in Fig. 5.1 and Fig. 5.2.

5.1.2 Analysis of the Data Dependencies

Although several variables are involved, and the overall specification may appear rather complicated, only a few types of data dependencies characterise all the computations of the two phases. Besides, the two phases and their data dependencies are very similar. Therefore, for brevity, in the following discussion we restrict ourselves to the reduction phase of the algorithm.

Let \( U, V \) represent any of the variables \( A, B, C, D \) or \( T \). All the data dependencies of the phase reduce to one of the following cases:

\[ \mathcal{DD}_0 = (D_1, U, V, I_0) \]
\[ \mathcal{DD}_1 = (D_1, U, V, I_1) \]
\[ \mathcal{DD}_2 = (D_1, U, V, I_2) \]
\[ \mathcal{DD}_3 = (D_1, U, V, I_3) \]
with index mappings:

\[ I_0(i, j) = (i, j) \]
\[ I_1(i, j) = (i, j - 1) \]
\[ I_2(i, j) = (i - g(i, j), j - 1) \]
\[ I_3(i, j) = (i + g(i, j), j - 1) \]

where \( g(i, j) = 2^j - 1 \). The data dependence graph is illustrated in Fig. 5.1 b). The dependence cones \( \Theta^*_I, \Theta^*_2, \) and \( \Theta^*_3 \) are illustrated in Fig. 5.3 a), b) and c), respectively, and the overall dependence cone \( \Theta^* \) in Fig. 5.4 a).

The above index mappings are all integral and, in particular, \( I_0 \) and \( I_1 \) are uniform. An explicit integral form for the mappings is the following:

\[ I_0(i, j) = (i, j) \]
\[ I_1(i, j) = (i, j) + (0, -1) \]
\[ I_2(i, j) = (i, j) + (-1, -1) + (g(i, j) - 1)(-1, 0) \]
\[ I_3(i, j) = (i, j) + 2^{-i-1}, -1) + (2^{-i-1} - g(i, j))(-1, 0) \]

The generators of the above index mappings have been chosen so that the cone \( C \) that they span (see Fig. 5.4 b)) contains the data dependence cone \( \Theta^* \), is pointed and has as
extremal rays the vectors of the unimodular basis \{((2^{r-1} - 1), (1, 0))\} (see also the discussion in Section 3.2.1). Moreover, the corresponding coefficients define non-negative and bounded integer functions over \( D_1 \). In particular, as for all \((i,j) \in D_1, 1 \leq g(i,j) = 2^{j-1} \leq 2^{r-2}\), then:

\[
0 \leq g(i,j) - 1 \leq 2^{r-2} - 1
\]

\[
2^{r-1} - 1 \geq 2^{r-1} - g(i,j) \geq 2^{r-2}
\]

Although all the conditions are met for the application of integral regularisation techniques, array designs derived from this specification would be non-scalable and exhibit non-local (albeit regular) connections. To see that this is the case, let us consider the dependence cone more closely. As the size parameter \( r \) of the problem increases, the dependence cone \( \Theta^* \) tends toward a non-pointed dependence cone. This is illustrated in Fig. 5.5 a), for \( r = 3 \) and \( r = 4 \). In addition, any possible decomposition of the index mappings is based on generators which are also dependent on the size parameter \( r \), namely the vector \((-2^{r-1}, 1)\) (Fig. 5.5 b) shows how the cone \( C \) varies with \( r \). By stepping ahead a little, Fig. 5.6 shows the data dependence graph that would result from a decomposition of the data dependencies according to the above formulation. Parts a) and b) of the figure illustrate the decomposition of \( DD_2 \), while parts c) and d) that of \( DD_3 \). The fact that vector \((-2^{r-1}, 1)\) depends on \( r \) implies that the data dependence graph corresponding to \( DD_3 \) (hence the resulting signal flow graph under a space-time mapping) is not scalable, that is as \( r \) varies a different connection topology is required. If the array design is realised in hardware, this fact implies that an entirely new component has to be built any time we want to solve a problem of different size. On the contrary, a scalable design allows one simply to update an existing component, typically by adding or removing processing elements, when a problem of larger or smaller size needs to be addressed. Although scalability is a design constraint which is mainly relevant to hardware implementation, it is good design practice to try and generate fully scalable regular array designs. From a theoretically point of view, the notion of scalability can be captured by a notion of uniformity of the data dependencies with respect to the size parameters of the problem. This notion was developed by Quinton and Van Dongen in [QuVa89]. Fig. 5.7 a) and b) show the non-scaling data dependence graphs of \( DD_3 \) for values of the size parameter \( r = 3 \) and \( r = 4 \), respectively. The reader may notice how the data dependence vectors change non uniformly as the size parameter increases, by comparing the sub-graph in b) corresponding to the graph in a).
Fig. 5.5. Reduction phase: a) $\Omega^*$ for $r = 3$ and $r = 4$; b) $C'$ for $r = 3$ and $r = 4$.

Fig. 5.6. Reduction phase: a) and b) decomposition of $\mathcal{D}_2$; c) and d) decomposition of $\mathcal{D}_3$. 
5.1.3 New Specification

We want to reformulate the specification so that the data dependence graph for its reduction phase assumes the form of Fig. 5.8. As we will see this new specification will result in scalable regular array designs. The transformation we are looking for is defined by the mapping \( T(i, j) = (i + 2^j - 1, j) \) in \( \mathbb{Z}^2 \). The computation domain \( D_1 \) becomes

\[
D_1 = \{(i, j) \mid 1 \leq j \leq r - 1, 3 \leq i \leq 2^r - 1\}
\]

and the actual computations in \( D_1 \) are identified by the guard:

\[
f(i, j) = \begin{cases} 
1 & (i - 2^j + 1)/2^{j-1} \in \{2, 4, 6, \ldots, 2^r - 2\} \\
0 & \text{otherwise}
\end{cases}
\]

The new index mappings are:

\[
\begin{align*}
\mathcal{I}_0(i, j) &= (i, j) \\
\mathcal{I}_1(i, j) &= (i - g(i, j) \cdot j - 1) \\
\mathcal{I}_2(i, j) &= (i - g'(i, j), j - 1) \\
\mathcal{I}_3(i, j) &= (i, j - 1)
\end{align*}
\]
where \( g(i, j) = 2^{j-1} \) and \( g'(i, j) = 2^j \). The input domain \( D_0 \) and the applied functions are the same as in the previous specification.

A similar, transformation applied to the backsubstitution phase, produces the data dependence graph of Fig. 5.9.

5.1.4 Analysis of the New Data Dependencies

Once again let us restrict ourselves to the reduction phase of the algorithm, and let \( I, V \) represent any of the variables \( A, B, C, D \) or \( T \). All data dependencies reduce to one of the following cases:

\[
\begin{align*}
\mathcal{D}D_0 & = (D_1, U, V, I_0) \\
\mathcal{D}D_1 & = (D_1, U, V, I_1) \\
\mathcal{D}D_2 & = (D_1, U, V, I_2) \\
\mathcal{D}D_3 & = (D_1, U, V, I_3)
\end{align*}
\]

with index mappings:

\[
\begin{align*}
I_0(i, j) & = (i, j) \\
I_1(i, j) & = (i - g(i, j), j - 1) \\
I_2(i, j) & = (i - g'(i, j), j - 1) \\
I_3(i, j) & = (i, j - 1)
\end{align*}
\]

where \( g(i, j) = 2^{j-1} \) and \( g'(i, j) = 2^j \). The dependence cones relative to \( I_1, I_2, \) and \( I_3 \) are sketched in Fig. 5.10 a), b) and c), respectively, while the overall dependence cone \( \Theta^* \) is illustrated in Fig. 5.11 a).

Explicit integral forms of the index mappings are:

\[
I_0(i, j) \quad (i, j)
\]
5.1.3 The cone \( C \) generated by \((0,1), (1,0)\) and \((1,1)\) is pointed and contains \( \Theta^* \) (see Fig. 5.11 b)). Also, all the coefficients of the mappings define non-negative and bounded integer functions over \( D_1 \). In particular, for all \((i,j) \in D_1\), 

\[
1 \leq g(i,j) = 2^{j-1} \leq 2^{r-2}, \quad 2 \leq g'(i,j) = 2^j \leq 2^{r-1},
\]

and:

\[
0 \leq g(i,j) - 1 \leq 2^{r-2} - 1 \\
1 \leq g'(i,j) - 1 \leq 2^{r-1} - 1
\]

5.1.5 Regularisation

Both integral data dependencies \( \mathcal{DD}_1 \) and \( \mathcal{DD}_2 \) are not atomic. Therefore their decomposition is necessary before uniformisation may be applied. We present the two transformations separately.

**Decomposition**

Both data dependencies admit a simple decomposition, as their first component defines a uniform (hence, linear and injective) index mapping for which an inverse can be determined directly.

![Fig. 5.10. Dependence cones relative to: a) \( I_1 \); b) \( I_2 \); c) \( I_3 \).](image1)

![Fig. 5.11. Reduction phase (new specification): a) \( \Theta^* \); b) \( C \).](image2)
We substitute $\mathcal{DD}_1$ with the atomic integral data dependencies:

$$
\begin{align*}
\mathcal{DD}_4' &= (D_1, U, P, I_1') \\
\mathcal{DD}_4 &= (D_2, P, V, I_4)
\end{align*}
$$

where $P$ is a new variable, the index mappings are:

$$
\begin{align*}
I_1'(i, j) &= (i, j) + (-1, -1) \\
I_4(i, j) &= (i, j) + (g'(i, j) - 1)(-1, 0)
\end{align*}
$$

with $g'(i, j) = 2^i$, and the new domain is:

$$
D_2 = \{(i-1, j-1) \mid (i, j) \in D_1\} = \{(i, j) \mid 0 \leq j \leq \tau - 2, 2 \leq i \leq 2^\tau - 2\}.
$$

Similarly, $\mathcal{DD}_2$ can be substituted by:

$$
\begin{align*}
\mathcal{DD}_2' &= (D_1, U, Q, I_1') \\
\mathcal{DD}_5 &= (D_2, Q, V, I_5)
\end{align*}
$$

with index mappings:

$$
\begin{align*}
I_1'(i, j) &= (i, j) + (-1, -1) \\
I_5(i, j) &= (i, j) + (g''(i, j) - 1)(-1, 0)
\end{align*}
$$

where $g''(i, j) = 2^{i+1}$.

The effects of these substitutions are illustrated in Fig. 5.12 a) and b) for the components of $\mathcal{DD}_1$ and Fig. 5.12 c) and d) for the components of $\mathcal{DD}_2$.

**Uniformisation**

We can now proceed to the uniformisation of $\mathcal{DD}_4$ and $\mathcal{DD}_5$, the only remaining non-uniform data dependencies.

The two cases are similar. In particular, by adopting the notation of Section 3.2.3, in both cases $d = (-1, 0)$. Also $\text{lin}(D_2) = \mathbb{Z}^2$ and $d$ is contained in this space. Hence, in both cases. a reindexing in $\mathbb{Z}^3$ is required. We choose a new index $k$ and a system of axes $i, j, k$ in this order. With this choice, a hyperplane $[\pi : \theta]$ containing the domain $D_2$ is determined by the vector $\pi = (0, 0, 1)$ and the coefficient $\theta = 0$. Hence, the uniformisation directions are $\hat{d} = d + \pi = (-1, 0, 1)$ and $\hat{d} = d - \pi = (-1, 0, -1)$. 

Fig. 5.12. Decomposition of: a) and b) $\mathcal{D}_2$; a) and c) $\mathcal{D}_2$. 
The differences between the two cases concern: the routing domains (as the upper bounds of the coefficients of the index mappings are different); and the values of the control variables (as the coefficients of the index mappings define different functions on $D_2$).

The data dependence $DD_4$ can be replaced by:

$$DD_4' = (D_2, P, R^1, I_0)$$

where routing and control variables are defined by the equations:

$$
\begin{align*}
E_{10} &= (D_{2,1}, R^1, (\alpha, \beta, \gamma, R^1, R^2), \delta, (I_0, I_0, I_0, I_0, I_0)) \\
E_{11} &= (D_{2,1}, R^2, R^2, id, I_7) \\
E_{12} &= (D_{2,2}, R^2, V, id, I_0) \\
E_{13} &= (D_{2,1}, \alpha, \alpha, id, I_4') \\
E_{14} &= (D_{2,2}, \alpha, in_\alpha) \\
E_{15} &= (D_{2,1}, \beta, \beta, id, I_4') \\
E_{16} &= (D_{2,2}, \beta, in_\beta) \\
E_{17} &= (D_{2,1}, \gamma, \gamma, dec, I_4') \\
E_{18} &= (D_{2,2}, \gamma, in_\gamma)
\end{align*}
$$

with:

- index mappings:

$$
\begin{align*}
I_0(i, j, k) &= (i, j, k) \\
I_4'(i, j, k) &= (i, j, k) + (-1, 0, 1) \\
I_6(i, j, k) &= (i, j, k) + (-1, 0, -1) \\
I_7(i, j, k) &= (i, j, k) + (-1, 0, 0)
\end{align*}
$$

- applied functions (where $\bar{g'} = [(2^r-2) - 1]/2$):

$$
\begin{align*}
in_\alpha(i, j, k) &= [(g'(i + \bar{g'}, j, k - \bar{g'}) - 1)/2] \\
in_\beta(i, j, k) &= g'(i + \bar{g'}, j, k - \bar{g'}) \mod 2 \\
in_\gamma(i, j, k) &= \bar{g'}
\end{align*}
$$
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\[\text{id}(a) = \ a\]
\[\text{dec}(a) = \ a - 1\]
\[f_3(\alpha, \beta, \gamma, a, b, c) = \begin{cases} a & \alpha \neq \gamma \\ b & \alpha = \gamma, \beta = 3 \\ c & \alpha = \gamma, \beta = 1 \end{cases}\]

-Domains:

\[D_{2_1} = \{(i, j, k) \mid 1 - k \leq i \leq 2^r - 2 - k, 0 \leq k \leq \bar{g}', 0 \leq j \leq r - 2\}\]
\[D_{2_1,1} = \{(i, j, k) \in D_{2_1} \mid k < \bar{g}'\}\]
\[D_{2_1,2} = \{(i, j, k) \in D_{2_1} \mid k = \bar{g}'\}\]
\[D_{2_2} = \{(i, j, k) \mid k - 2\bar{g}' \leq i \leq 2^r - 2 - k, 0 \leq k \leq \bar{g}', 0 \leq j \leq r - 2\}\]
\[D_{2_2,1} = \{(i, j, k) \in D_{2_2} \mid k > 0\}\]
\[D_{2_2,2} = \{(i, j, k) \in D_{2_2} \mid k = 0\}\]

Similarly, the uniformisation of \(D_5\) yields the data dependence:

\[DD'_5 = (D_2, Q, S^1, I_0)\]

and the equations:

\[(\text{routing variables } S^1, S^2)\]
\[E_{19} = (D_{2_3}, S^1, (\xi, \phi, \psi, S^1, S^2, S^2), f_3, (I_0, I_0, I_0, I_4, I_0, I_0))\]
\[E_{20} = (D_{2_4}, S^2, S^2, id, I_7)\]
\[E_{21} = (D_{2_4}, S^2, V, id, I_0)\]

\[(\text{control variables } \xi, \phi \text{ and } \psi)\]
\[E_{22} = (D_{2_3}, \xi, \xi, id, I_4')\]
\[E_{23} = (D_{2_3}, \xi, \text{in}_\xi)\]
\[E_{24} = (D_{2_3}, \phi, \phi, id, I_4')\]
\[E_{25} = (D_{2_3}, \phi, \text{in}_\phi)\]
\[E_{26} = (D_{2_3}, \psi, \psi, \text{dec}, I_4')\]
\[E_{27} = (D_{2_3}, \psi, \text{in}_\psi)\]

where:

- the index mappings are:

\[I_0(i, j, k) = (i, j, k)\]
\[ T_4(i, j, k) = (i, j, k) + (-1, 0, 1) \]
\[ T_6(i, j, k) = (i, j, k) + (-1, 0, 0) \]
\[ T_7(i, j, k) = (i, j, k) + (-1, 0, -1) \]

- the applied functions are (where \( \bar{g}'' = \lfloor (2^{r-1} - 1)/2 \rfloor \)):

\[ \text{in}_\xi(i, j, k) = [(g''(i + \bar{g}'', j, k - \bar{g}'') - 1)/2] \]
\[ \text{in}_\phi(i, j, k) = (g''(i + \bar{g}'', j, k - \bar{g}'') - 1) \mod 2 \]
\[ \text{in}_\psi(i, j, k) = \bar{g}'' \]
\[ \text{id}(a) = a \]
\[ \text{dec}(a) = a - 1 \]
\[ f_3(\xi, \phi, \psi, a, b, c) = \begin{cases} 
  a & \xi \neq \psi \\
  b & \xi = \psi, \phi = 0 \\
  c & \xi = \psi, \phi = 1 
\end{cases} \]

- the domains are:

\[ D_{23} = \{(i, j, k) \mid 1 - k \leq i \leq 2^r - 2 - k, 0 \leq k \leq \bar{g}'', 0 \leq j \leq r - 2\} \]
\[ D_{23,1} = \{(i, j, k) \in D_{23} \mid k < \bar{g}''\} \]
\[ D_{23,2} = \{(i, j, k) \in D_{23} \mid k = \bar{g}''\} \]
\[ D_{24} = \{(i, j, k) \mid k - 2g'' \leq i \leq 2^r - 2 - k, 0 \leq k \leq g'', 0 \leq j \leq r - 2\} \]
\[ D_{24,1} = \{(i, j, k) \in D_{24} \mid k > 0\} \]
\[ D_{24,2} = \{(i, j, k) \in D_{24} \mid k = 0\}. \]

The data dependence graphs corresponding to \( DD_4 \) and \( DD_5 \) are sketched in Fig. 5.13 a) and b), respectively, where only the routing paths of interest are illustrated.

### 5.1.6 Space-Time Mapping

A uniform system of equations is obtained by applying regularisation techniques to all the data dependencies of the specification as explained in the previous section. The resulting uniform specification (which we have coded in Mathematica in Appendix C) includes 35 variables, which we have summarised in Table 5.2.

The uniform data dependencies of the specification are summarised in Tables 5.3-5.7, where:

- the first column indicates the pair of variables which are related;
- the second and third
Fig. 5.13. Uniformisation of: a) $DD_4$; b) $DD_5$.

<table>
<thead>
<tr>
<th>Type</th>
<th>Name</th>
<th>Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Original variables</td>
<td>$A, B, C, D$</td>
<td>4</td>
</tr>
<tr>
<td>Original control variables</td>
<td>$T$</td>
<td>1</td>
</tr>
<tr>
<td>Decomposition variables</td>
<td>$P_A, P_B, P_C, P_D$</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>$Q_A, Q_B, Q_C, Q_D$</td>
<td></td>
</tr>
<tr>
<td>Uniformisation variables</td>
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<td>16</td>
</tr>
<tr>
<td></td>
<td>$R_A^2, R_B^2, R_C^2, R_D^2$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$S_A^1, S_B^1, S_C^1, S_D^1$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$S_A^2, S_B^2, S_C^2, S_D^2$</td>
<td></td>
</tr>
<tr>
<td>Uniformisation control variables</td>
<td>$\alpha, \beta, \gamma$</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td>$\xi, \psi, \phi$</td>
<td></td>
</tr>
</tbody>
</table>

Table 5.2. Variables in the final specification.
Table 5.3. Data dependencies relative to variable A.

<table>
<thead>
<tr>
<th>Pair</th>
<th>Decomp.</th>
<th>Unifor.</th>
<th>DDVector</th>
</tr>
</thead>
<tbody>
<tr>
<td>A, T</td>
<td>-</td>
<td>-</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td>A, A</td>
<td>-</td>
<td>-</td>
<td>(0,1,0)</td>
</tr>
<tr>
<td>A, A</td>
<td>A, P_A</td>
<td>P_A, A</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P_A, R_1</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_1, P_A</td>
<td>(1,0,-1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_1, R_A</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_A, R_A</td>
<td>(1,0,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_A, R_A</td>
<td>(1,0,1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_A, A</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td>A, B</td>
<td>A, P_B</td>
<td>P_B, A</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>P_B, R_1</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_1, R_B</td>
<td>(1,0,-1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_B, R_B</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_B, R_B</td>
<td>(1,0,0)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_B, R_B</td>
<td>(1,0,1)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>R_B, B</td>
<td>(0,0,0)</td>
</tr>
</tbody>
</table>

Fig. 5.14. Dependence cone $\Theta^*$ after uniformisation.

columns, respectively, the new data dependencies introduced by decomposition and uniformisation (when applicable, a dash, - , otherwise); and the last column the corresponding (uniform) data dependence vector.

By considering all data dependence vectors, the resulting dependence cone is

$$\Theta^* = \text{cone}(\{(0,1,0),(1,1,0),(1,0,0),(1,0,-1),(1,0,1)\}),$$

with extremal rays $(0,1,0),(1,0,-1)$ and $(1,0,1)$. The cone is illustrated in Fig. 5.14.

Such a cone is pointed and, according to the conditions discussed in Section 2.2.6, an affine timing function for the specification is determined by any vector $\lambda = (\lambda_1, \lambda_2, \lambda_3)$ in $ \mathbb{Z}^3$, such that:

$$\lambda_2 > 0$$
**Pair** | **Decomp.** | **Unifor.** | **DDVector**
--- | --- | --- | ---
*B, T* | – | – | (0,0,0)
*B, B* | – | – | (0,1,0)
*B, A* | – | – | (0,1,0)
*B, C* | B, P<sub>C</sub> | P<sub>C</sub>, C | (1,1,0)
*B, B* | B, P<sub>B</sub> | – | (1,1,0)
*B, C* | – | – | (0,1,0)
*B, A* | B, P<sub>A</sub> | – | (1,1,0)

Table 5.4. Data dependencies relative to variable B.

**Pair** | **Decomp.** | **Unifor.** | **DDVector**
--- | --- | --- | ---
*C, T* | – | – | (0,0,0)
*C, C* | – | – | (0,1,0)
*C, C* | C, Q<sub>C</sub> | Q<sub>C</sub>, C | (1,1,0)
*C, B* | C, Q<sub>B</sub> | – | (1,1,0)

Table 5.5. Data dependencies relative to variable C.
<table>
<thead>
<tr>
<th>Pair</th>
<th>Decomp.</th>
<th>Unifor.</th>
<th>DDVector</th>
</tr>
</thead>
<tbody>
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<td>-</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td>D,D</td>
<td>-</td>
<td>-</td>
<td>(0,1,0)</td>
</tr>
<tr>
<td>D,A</td>
<td>-</td>
<td>-</td>
<td>(0,1,0)</td>
</tr>
<tr>
<td>D,D</td>
<td>D,P_D</td>
<td>P_D,D</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td></td>
<td>P_D,D</td>
<td>P_D,R_D^1</td>
<td>(0,0,0)</td>
</tr>
<tr>
<td></td>
<td>R_D^1,R_D^1</td>
<td>(1,0,-1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>R_D^2,R_D^1</td>
<td>(0,0,0)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>R_D^1,R_D^2</td>
<td>(1,0,0)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>R_D^2,R_D^2</td>
<td>(1,0,1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>R_D^3,D_D</td>
<td>(0,0,0)</td>
<td></td>
</tr>
<tr>
<td>D,B</td>
<td>D,P_B</td>
<td>P_B,D</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td></td>
<td>P_B,B</td>
<td>-</td>
<td>(1,1,0)</td>
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<td>-</td>
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<tr>
<td>B,A</td>
<td>B,P_A</td>
<td>P_A,A</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td></td>
<td>-</td>
<td>-</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td>D,D</td>
<td>D,Q_D</td>
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<tr>
<td></td>
<td>Q_D,D</td>
<td>Q_D,S_D^1</td>
<td>(1,1,0)</td>
</tr>
<tr>
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<td>S_D^1,S_D^1</td>
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<td></td>
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<tr>
<td></td>
<td>S_D^2,S_D^2</td>
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<tr>
<td></td>
<td>S_D^1,S_D^2</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>S_D^2,S_D^1</td>
<td>(1,0,0)</td>
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<tr>
<td></td>
<td>S_D^3,D_D</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>S_D^2,D_D</td>
<td>(0,0,0)</td>
<td></td>
</tr>
<tr>
<td>D,B</td>
<td>D,Q_B</td>
<td>Q_B,D</td>
<td>(1,1,0)</td>
</tr>
<tr>
<td></td>
<td>Q_B,B</td>
<td>-</td>
<td>(1,1,0)</td>
</tr>
</tbody>
</table>

Table 5.6. Data dependencies relative to variable D.

<table>
<thead>
<tr>
<th>Pair</th>
<th>Decomp.</th>
<th>Unifor.</th>
<th>DDVector</th>
</tr>
</thead>
<tbody>
<tr>
<td>α,α</td>
<td>-</td>
<td>-</td>
<td>(1,0,-1)</td>
</tr>
<tr>
<td>β,β</td>
<td>-</td>
<td>-</td>
<td>(1,0,-1)</td>
</tr>
<tr>
<td>γ,γ</td>
<td>-</td>
<td>-</td>
<td>(1,0,-1)</td>
</tr>
<tr>
<td>ξ,ξ</td>
<td>-</td>
<td>-</td>
<td>(1,0,-1)</td>
</tr>
<tr>
<td>φ,φ</td>
<td>-</td>
<td>-</td>
<td>(1,0,-1)</td>
</tr>
<tr>
<td>ψ,ψ</td>
<td>-</td>
<td>-</td>
<td>(1,0,-1)</td>
</tr>
</tbody>
</table>

Table 5.7. Data dependencies relative to the routing control variables.
Note that, according to our discussion in Section 2.2.5, further inequalities may be derived by considering the properties of time optimality and non-negativeness of the corresponding timing functions. In particular, such inequalities can be obtained by considering the generators (vertices and rays) of the computation domain of the specification, which is the smallest convex polyhedral set containing all computation domains of the system. The resulting inequalities define an integer linear programming problem whose solution provides an optimal affine scheduling. For brevity, here (and in the following examples) we omit the complete formulation of the problem. The reader may be convinced that the vector $\lambda = (1, 1, 0)$ provides such an optimal solution, corresponding to the affine timing function $t(i, j, k) = i + j$. To make $t$ non-negative, a suitable delay can be added so that $t$ associates an initial time $0$ with the first set of computations of the algorithm. With this scheduling the algorithm executes in $O(N + 2\lfloor N/2 \rfloor + 2)$ time steps, where $N = 2^r - 1$.

A possible projection vector is any $u = (u_1, u_2, u_3)$ in $\mathbb{Z}^3$ such that $\lambda \cdot u \neq 0$, i.e., $u_1 + u_2 \neq 0$. For instance, $u = (1, 0, 0)$ satisfies the requirement and corresponds to the allocation function $a(i, j, k) = (j, k)$. The effect of the space-time mapping $[t, a]$ on the (uniform) data dependence vectors of the specification is summarised in Table 5.8, and the image of the data dependence graph under $[t, a]$ is the signal flow graph of Fig. 5.15 a). In the figure, two types of nodes are indicated, which correspond to two basic types of processing elements (see later on). For simplicity, in the figure, unit arc labels have been omitted. Note that here we have considered only a single instance of each data dependence vector. Hence, this signal flow graph is a simplified version of the actual signal flow graph of the specification, in which each arc should be replicated for all the corresponding data dependencies of the system.

The signal flow graph together with the information provided by the recurrences are used for a detailed description of the array design. In particular, a processing element is associated with each node of the signal flow graph and a communication channel or a memory cell with each of its arcs. The operations at each processing element are specified by the applied functions of the recurrences whose computation points are mapped onto that processing element by the allocation function. The association of variables and communication channels or memory cells is also determined by the allocation function though the mapping of the corresponding data dependence vectors. A description of the two basic cells of the regular array relative to the signal flow graph in Fig. 5.15 a) is given in Fig. 5.16 a). In the figure we have
Table 5.8. Transformation of the data dependence vectors under the space-time mappings \([t, a]\), \([t, a']\), and \([t, a'']\).

<table>
<thead>
<tr>
<th>(d)</th>
<th>(\lambda \cdot d)</th>
<th>(a(d))</th>
<th>(a'(d))</th>
<th>(a''(d))</th>
</tr>
</thead>
<tbody>
<tr>
<td>((0,1,0))</td>
<td>1</td>
<td>((1,0))</td>
<td>((-1,0))</td>
<td>((0,0))</td>
</tr>
<tr>
<td>((1,1,0))</td>
<td>2</td>
<td>((1,0))</td>
<td>((0,0))</td>
<td>((1,0))</td>
</tr>
<tr>
<td>((1,0,0))</td>
<td>1</td>
<td>((0,0))</td>
<td>((1,0))</td>
<td>((1,0))</td>
</tr>
<tr>
<td>((1,0,-1))</td>
<td>1</td>
<td>((0,-1))</td>
<td>((1,-1))</td>
<td>((1,-1))</td>
</tr>
<tr>
<td>((1,0,1))</td>
<td>1</td>
<td>((0,1))</td>
<td>((1,1))</td>
<td>((1,1))</td>
</tr>
</tbody>
</table>

assumed that: \(U, V \in \{A, B, C, D\}\); \(P, Q \in \{P_A, Q_A, P_B, Q_B, P_C, Q_C, P_D, Q_D\}\); \(R^1, S^1 \in \{R^1_A, S^1_A, R^1_B, S^1_B, R^1_C, S^1_C, R^1_D, S^1_D\}\); and \(R^2, S^2 \in \{R^2_A, S^2_A, R^2_B, S^2_B, R^2_C, S^2_C, R^2_D, S^2_D\}\). Dotted arrows correspond to control signals. Note that the loops of the signal flow graph correspond to memory cells in the processing elements. A slash, \(/\), indicates an alternative (between variables of the two routing schemes of Section 5.1.5, while a single quote, \(\-'\), denotes an output signal. Note that the description of the processing elements is only partial, as a complete description should include signals corresponding to all 35 variables of the specification.

Other compatible projection vectors for the given timing function \(t\) are, for instance, \(u' = (1,1,0)\) and \(u'' = (0,1,0)\), corresponding to the allocation functions \(a'(i,j,k) = (i-j,k)\) and \(a''(i,j,k) = (i,k)\), respectively. The effect of the resulting space-time mappings on the data dependence vectors are also given in Table 5.8, while Fig. 5.15 b) and c) illustrate the corresponding signal flow graphs, and Fig. 5.16 b) and c) the corresponding basic processing elements.

### 5.2 N Points FIR Filter for \(M\)-to-1 Decimation

One of the most fundamental concepts of digital signal processing [CrRa83] is the idea of sampling a continuous process to obtain a set of numbers which, in some sense, is represen-
CHAPTER 5. CASE STUDIES

Fig. 5.16. Basic processing elements corresponding to: a) \([t, a]\); b) \([t, a']\); c) \([t, a'']\).

tative of the characteristics of the process being sampled. Let \(x_C(t)\) denote a continuous function of the process being sampled, where \(t\) is a continuous variable (typically time) and \(-\infty < t < \infty\). A set of samples of \(x_C\) can be defined by another function \(x_D(n)\), with \(-\infty < n < \infty\), where the correspondence between \(t\) and \(n\) can be expressed by an equation \(n = q(t)\), for some function \(q\) specified by the sampling process.

A common form of sampling, called uniform (periodic) sampling, is one in which \(q(t) = t/T = n\), where \(n\) is an integer. That is, the samples \(x_D(n)\) are uniformly spaced (occurring \(T\) apart) in the dimension \(t\). \(T\) is called the sampling period.

In some cases the input signal may already be sampled at some predetermined sampling period \(T\) and the goal is to convert this sampled signal into a new sampled signal with a different sampling period \(T'\). If \(T' > T\) this digital conversion is called decimation. In particular, an \(M\)-to-1 decimator, for some natural number \(M\), is a decimator which discards \(M - 1\) every \(M\) (output) samples.

Decimation is usually obtained by filtering the input digital signal. A Finite Impulse Response (FIR) filter is a digital filter whose impulse response \(h(k)\) is of finite duration, i.e., it is zero outside a finite interval of samples \(k\). In particular, the direct form of an \(N\) points FIR filter is the convolution

\[
y(m) = \sum_{n=0}^{N-1} h(n)x(m - n)
\]

where the filter response \(h\) is assumed to be 0 for \(n < 0\) and \(n > N - 1\).

By combining the above definitions, the direct form of an \(N\) points FIR filter for \(M\)-to-1
decimation corresponds to the convolution:

\[ y(m) = \sum_{n=0}^{N-1} h(n)x(Mm - n) \]

for \( m \geq 0 \).

### 5.2.1 Specification

We want to specify the problem as a system of integral recurrence equations. The first step is to transform equation

\[ y(m) = \sum_{n=0}^{N-1} h(n)x(Mm - n) \]

by exploiting the associativity of addition and introducing a variable \( Y \) which accumulates the partial sums. Then each \( y(m) \) can be computed by the system of equations:

\[
\begin{align*}
Y(m,0) &= 0 \\
Y(m,1) &= Y(m,0) + h(0)x(Mm) \\
Y(m,2) &= Y(m,1) + h(1)x(Mm - 1) \\
&\quad \ldots \\
Y(m,N) &= Y(m,N - 1) + h(N - 1)x(Mm - N + 1)
\end{align*}
\]

That is, more concisely:

\[
\begin{align*}
Y(m,0) &= 0 \\
Y(m,j) &= Y(m,j - 1) + h(j - 1)x(Mm - j + 1)
\end{align*}
\]

for \( j = 1, \ldots, N \).

By introducing two new variables \( H \) and \( X \), corresponding to \( h \) and \( x \), respectively, we can specify the problem in 2 dimensions as follows. An index \( i \) is used which corresponds to \( m \). For simplicity, let us assume that the new digital signal \( y \) is sampled starting at \( i = 1 \), so that \( y \) is computed for \( i \geq 1 \). Variable \( X \) is initialised to the values of \( x \) for all \( i \), while variable \( H \) is initialised to the values of \( h \). As those values are used in each convolution, variable \( H \) also pipelines the values through the computation space. The system of equations is the following:

\[
\begin{align*}
E_1 &= (D_1, X, in_X) \\
E_2 &= (D_2, Y, in_Y)
\end{align*}
\]
\[ E_3 = (D_3, Y, (Y, H, X), f(I_1, I_2, I_3)) \]
\[ E_4 = (D_4, H, \text{id}_H) \]
\[ E_5 = (D_3, H, H, \text{id}, I_2) \]

where:

- the index mappings are:
  \[ I_1(i, j) = (i, j - 1) \]
  \[ I_2(i, j) = (i - 1, j) \]
  \[ I_3(i, j) = (Mi - j + 1, 0) \]

- the applied functions are:
  \[ f(a, b, c) = a + bc \]
  \[ \text{id}(a) = a \]
  \[ \text{inx}(i, j) = x(i) \]
  \[ \text{iny}(i, j) = 0 \]
  \[ \text{in}H(i, j) = h(j - 1) \]

- the domains are:
  \[ D_1 = \{(i, j) | i \geq -(N - 1), j = 0\} \]
  \[ D_2 = \{(i, j) | i \geq 1, j = 0\} \]
  \[ D_3 = \{(i, j) | i \geq 1, 1 \leq j \leq N\} \]
  \[ D_4 = \{(i, j) | i = 0, 1 \leq j \leq N\} \]

### 5.2.2 Analysis

The data dependencies of the system are the following:

\[ DD_1 = (D_3, Y, Y, I_1) \]
\[ DD_2 = (D_3, Y, H, I_2) \]
\[ DD_3 = (D_3, Y, X, I_3) \]
\[ DD_4 = (D_3, H, H, I_2) \]
where the index mappings are integral and can be expressed, for instance, as:

\[
I_1(i,j) = (i,j) + (0, -1)
\]

\[
I_2(i,j) = (i,j) + (-1, 0)
\]

\[
I_3(i,j) = (i,j) + ((M - 1)i + 1, 0) + j(-1, -1)
\]

In particular, \(I_1\) and \(I_2\) are uniform. The form we have chosen for \(I_3\) guarantees that its coefficients define non-negative integer functions on \(D_3\). Let \(g_1(i,j) = (M - 1)i + 1\) and \(g_2(i,j) = j\) be such coefficients.

The corresponding data dependence graph is sketched in Fig. 5.17, where, for clarity, we have separated the uniform data dependence vectors from those relative to \(DD_3\) (parts a) and b) of the figure, respectively). In the figure we have assumed \(M = 3\).

A simple analysis of \(g_1\) shows that the coefficient is not bounded on \(D_3\). In fact, its value grows linearly in \(i\). Therefore integral regularisation techniques are not applicable.

Although we have been treating \(DD_3\) as an integral data dependence, \(DD_3\) is in particular an affine data dependence. In fact, its index mapping can be written in matrix form as:

\[
I_3 \left( \begin{array}{c} i \\ j \end{array} \right) = \left[ \begin{array}{cc} M & -1 \\ 0 & 0 \end{array} \right] \left( \begin{array}{c} i \\ j \end{array} \right) + \left( \begin{array}{c} 1 \\ 0 \end{array} \right)
\]

As the matrix \(A = \left[ \begin{array}{cc} M & -1 \\ 0 & 0 \end{array} \right]\) is rank deficient, a well known regularisation technique (known in the literature as pipelining – see [FoMo84, RaFu87, Raj89, QuVa89]) can be applied. According to this technique, a regularisation vector can be chosen as a non-null vector in \(null(A) \cap lin(D_3)\) (see Appendix F for a definition of \(null(A)\)). \(D_3\) is of full dimension, hence we can restrict ourselves to \(null(A)\). The space \(null(A)\) is spanned, for instance, by the vector \(d = (1, M)\). As \(d\) is not a null vector, it can be chosen as a pipelining vector. The technique prescribes to partition the domain \(D_3\) in two non-empty sub-sets defined as:

\[
D_{3_1} = \{(i,j) \in D_3 | (i,j) + d \in D_3\}
\]
\[ D_{3_2} = \{(i, j) \in D_3 \mid (i, j) + d \notin D_3\} \]

Intuitively, \( D_{3_1} \) corresponds to computation points inside \( D_3 \), while \( D_{3_2} \) to points on (or around) some boundary of \( D_3 \). The result is obtained by pipelining the data among neighbour points in \( D_{3_1} \) according to the direction of \( d \), while possible residual non-uniform data dependence vectors are confined to \( D_{3_2} \).

Unfortunately, as \( d = (1, M) \) depends on the problem parameter \( M \), the applicability of the technique also depends on the values of \( M \). In particular, for \( M \geq N \), the subset \( D_{3_1} \) reduces to the empty set, and the technique is ineffective. Note that a different choice of pipelining vector would not solve the problem as any pipelining vector for the problem is an integer multiple of \( d \).

### 5.2.3 Problem Revisited

Let us consider the original equation:

\[ y(m) = \sum_{n=0}^{N-1} h(n)x(Mm - n) \]

For each \( m \), \( y(m) \) is a convolution of \( N \) terms, involving the filter responses \( h(0), h(1), \ldots, h(N-1) \) and the input samples \( x(Mm), x(Mm-1), \ldots, x(Mm-N+1) \). In other words, each \( y(m) \) is just a convolution of \( N \) input samples. However, different from ordinary convolution, the selection of such input samples depends on the decimation rate \( M \).

A straightforward formulation of the problem can be obtained by computing ordinary convolution and then decimating the outputs according to \( M \). This can be obtained through the following equations, where \( y' \) computes all convolutions, while \( y \) is assigned the decimated samples only:

\[
\begin{align*}
y'(m) &= \sum_{n=0}^{N-1} h(n)x(m - n) \\
y(k) &= y'(k)
\end{align*}
\]

for \( m \geq 0 \) and \( k = 0, M, 2M, 3M, \ldots \).

The convolution problem is a very well known problem in regular array synthesis (see, e.g., [QuRo91, Meg92]). The decimation is easy to specify with the help of control signals: it simply amounts to locate a non-convex sub-domain of a convex polyhedral domain. We have encountered this problem earlier in this chapter for cyclic reduction (see Section 5.1.1). In this case, we may define a control variable \( C \) which assumes value equal to 1 on those \( k \) of the
domains which are integer multiples of $M$. A possible specification is the following system of equations (whose uniform data dependence graph is illustrated in Fig. 5.18):

\[
\begin{align*}
\text{(variable } X) & \quad \mathbb{E}_1 = (D_1, X, \text{in}_X) \\
\text{(variable } \bar{Y}) & \quad \mathbb{E}_3 = (D_3, \bar{Y}, \text{in}_\bar{Y}) \\
\text{(variable } H) & \quad \mathbb{E}_5 = (D_4, H, \text{in}_H) \\
\text{(control variable } C) & \quad \mathbb{E}_7 = (D_5, C, \text{in}_C) \\
\text{(variable } Y) & \quad \mathbb{E}_8 = (D_5, Y, (\bar{Y}, C), f_2, (I_1, I_0))
\end{align*}
\]

where:

- the index mappings are:

\[
\begin{align*}
I_0(i, j) &= (i, j) \\
I_1(i, j) &= (i, j - 1) \\
I_2(i, j) &= (i - 1, j - 1) \\
I_3(i, j) &= (i - 1, j)
\end{align*}
\]
Chapter 5. Case Studies

5.2.4 Space-Time Mapping

All data dependencies are uniform and the corresponding dependence cone is

$$\Theta^* = \text{cone}((0,1),(1,1),(1,0)),$$

with extremal rays $(0,1)$ and $(1,0)$. The dependence cone is illustrated in Fig. 5.19 a).

As the dependence cone is pointed, a valid affine timing function is determined by any

$$\lambda = (\lambda_1, \lambda_2) \in \mathbb{Z}^2$$

such that the following system of inequalities is satisfied:

$$\lambda_1 > 0$$

$$\lambda_2 > 0.$$
The vector $\lambda = (1, 1)$ is a possible choice. A non-negative timing function determined by $\lambda$ is the function $t(i, j) = i + j$.

A compatible linear allocation function is a projection according to any non-null vector $u = (u_1, u_2) \in \mathbb{Z}^2$, such that $\lambda \cdot u \neq 0$, i.e., $u \in \mathbb{Z}^2$ such that $u_1 + u_2 \neq 0$. However, a finite array design can be obtained only if the projection is done in the same direction of the ray of the domain $D_2$, that is for a projection vector $u = (1, 0)$, which corresponds to the allocation function $a(i, j) = j$.

The transformation of the dependence vectors of the system under the space-time mapping $[t, a]$ is summarised in Table 5.9. The resulting signal flow graph is illustrated in Fig. 5.19 b). For simplicity arcs with a unit delay are not labelled. A description of the corresponding basic processing element and its operations is given in Fig. 5.20.

### Table 5.9. Transformation of data dependence vectors under the space-time mapping $[t, a]$.

<table>
<thead>
<tr>
<th>$d$</th>
<th>$\lambda \cdot d$</th>
<th>$a(d)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1, 0)</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>(1, 1)</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>(0, 1)</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

**Fig. 5.20. Processing element.**

5.3 Knapsack Problem

As an example of dynamic problem, in this section we consider the knapsack problem. The knapsack problem can be formulated either as a linear optimisation problem or as a dynamic programming problem (see, e.g., [Hu82]). The latter formulation is based on recursive functions, and is the formulation we have chosen in this section.

The problem can be described as follows. Let us consider a knapsack of finite capacity $c$, together with objects of $n$ different types, each type $i$, for $i = 1, \ldots, n$, characterised by a weight $w_i$ and a value $v_i$. We assume that both the capacity of the knapsack and the weights of the objects are integers, and that, for all $i$, $0 < w_i \leq c$. We also assume that there exists an unbounded number of objects of each type. The knapsack problem consists of determining the optimal (i.e., the most valuable) selection of the given objects, which can be carried in
the knapsack without exceeding its capacity.

A possible dynamic programming formulation of the algorithm is the following\(^1\). Let \(f_i(j)\) represent the maximum value which can be carried in a knapsack of capacity \(j\) by selecting objects of types from 1 to \(i\). \(f_i(j)\) may be defined as follows:

\[
\begin{align*}
\text{for } i = 1, \ldots, n, \quad j = 1, \ldots, c: & \quad f_i(j) = \max(f_{i-1}(j), f_i(j - w_i) + v_i) \\
\text{for } i = 0, \quad j = 0, \ldots, c: & \quad f_i(j) = 0 \\
\text{for } i = 0, \ldots, n, \quad j = 0: & \quad f_i(j) = 0 \\
\text{for } i = 1, \ldots, n, \quad j < 0: & \quad f_i(j) = -\infty
\end{align*}
\]

The boundary conditions express that \(f_i(j)\) is 0 if either the capacity of the knapsack is equal to 0 or there are no objects which can be selected; and \(f_i(j)\) is \(-\infty\) if the capacity of the knapsack is negative\(^2\). The solution to the knapsack problem is the value \(f_n(c)\).

Note that these equations only compute the optimal value of the objects which can be carried in the knapsack. In the literature this is known as the forward phase of the algorithm [Hu82]. The corresponding combination of objects can be determined subsequently as a backward substitution process on the sub-optimal values of \(f\). This is known as the backward phase of the algorithm. This phase is essentially sequential and will not be considered here. (An efficient algorithm for the backward phase is given in [Hu82].)

5.3.1 Specification

In order to specify the algorithm as a system of recurrence equations, we make the following observations:

- two indices, \(i\) and \(j\), are sufficient to the expression of the recurrences. Hence, we adopt \(\mathbb{Z}^2\) as the computation space and associate \(i\) to the types of the objects and \(j\) to the capacity of the knapsack;

- a variable \(F\), corresponding to \(f\), is needed to compute the optimal solution, together with two variables, \(W\) and \(V\) respectively, for the weights and values of the objects;

- the computation domain of \(F\) is the rectangular region determined by the points \((i, j)\), such that \(i = 1, \ldots, n\) and \(j = 1, \ldots, c\). Variables \(W\) and \(V\) are initialised on the boundary of this domain and their values are subsequently pipelined through the region.

---

\(^1\)This formulation is based on [Hu82].

\(^2\)We have used the symbol \(\infty\) to indicate the largest integer representable in a machine.
A possible system of equations is the following:

(variable $F$)

$E_1 = (D_1, F, \text{in}_F^1)$

$E_2 = (D_2, F, \text{in}_F^1)$

$E_3 = (D_3, F, \text{in}_F^2)$

$E_4 = (D_4, F, (F, F, V), f, (I_1, I_2, I_3))$

(variable $V$)

$E_5 = (D_2, V, \text{inv}_V)$

$E_6 = (D_4, V, V, \text{id}, I_3)$

(variable $W$)

$E_7 = (D_2, W, \text{in}_W)$

$E_8 = (D_4, W, W, \text{id}, I_3)$

where:

- index mappings:

  $I_1(i, j) = (i - 1, j)$

  $I_2(i, j) = (i, j - W(i, j))$

  $I_3(i, j) = (i, j - 1)$

- applied functions:

  $\text{in}_F^1(i, j) = 0$

  $\text{in}_F^2(i, j) = -\infty$

  $\text{inv}_V(i, j) = v_i$

  $\text{in}_W(i, j) = w_i$

  $f(a, b, c) = \text{max}(a, b + c)$

where $w_i$ is an integer, with $0 < w_i \leq c$, and $v_i \geq 0$;

- domains:

  $D_1 = \{(i, j) | i = 0, 1 \leq j \leq c\}$

  $D_2 = \{(i, j) | 1 \leq i \leq n, j = 0\}$
\[ D_3 = \{(i,j) | 1 \leq i \leq n, j < 0\} \]
\[ D_4 = \{(i,j) | 1 \leq i \leq n, 1 \leq j \leq c\}. \]

5.3.2 Analysis

The data dependencies of the specification are the following:

\[ \mathcal{DD}_1 = (D_4, F, F, I_1) \]
\[ \mathcal{DD}_2 = (D_4, F, F, I_2) \]
\[ \mathcal{DD}_3 = (D_4, F, V, I_3) \]
\[ \mathcal{DD}_4 = (D_4, V, V, I_3) \]
\[ \mathcal{DD}_5 = (D_4, W, W, I_3) \]

with index mappings:

\[ I_1(i,j) = (i-1, j) \]
\[ I_2(i,j) = (i, j-W(i,j)) \]
\[ I_3(i,j) = (i, j-1) \]

Data dependencies \( \mathcal{DD}_1, \mathcal{DD}_3, \mathcal{DD}_4 \) and \( \mathcal{DD}_5 \) are static and uniform, while data dependence \( \mathcal{DD}_2 \) is dynamic and atomic finitely generated. An explicit form for the index mapping is:

\[ I_1(i,j) = (i, j) + (-1, 0) \]
\[ I_2(i,j) = (i, j) + W(i,j)(0, -1) \]
\[ I_3(i,j) = (i, j) + (0, -1) \]

where \( 0 < W(i,j) \leq c \), for all \((i, j)\). As \( \mathcal{DD}_2 \) is dynamic, for each configuration of weights in input we obtain a different data dependence graph. For \( c = 8 \) and \( n = 3 \), two instances of the dynamic data dependence graph are given in Fig. 5.21 a) and b), corresponding to the two distributions of weights \( w_1 = 5, w_2 = 4, w_3 = 2 \) and \( w'_1 = 4, w'_2 = 1, w'_3 = 3 \), respectively. Note that only the sub-graph relative to \( \mathcal{DD}_2 \) varies with the inputs.

By considering the generators of all index mappings, we obtain the embedding dependence cone \( C = cone(\{(1, 0), (0, 1)\}) \), illustrated in Fig. 5.22 a). Note that \( C \) is pointed and, because of the condition \( 0 < W(i,j) \leq c \) contains the dependence cone of the specification for all inputs (see also the discussion in Section 4.3.3).

The separability of \( F \) and \( W \), which is necessary for uniformisation, can be verified on the extended dependence graph of the system. This graph is illustrated in Fig. 5.22 b). and is defined.
Fig. 5.21. Two instances of the dynamic dependence graph of the knapsack problem, for $c = 8$, $n = 3$, and weight distributions, respectively: a) $w_1 = 5, w_2 = 4, w_3 = 2$; b) $w'_1 = 4, w'_2 = 1, w'_3 = 3$.

Fig. 5.22. a) Cone $C$; b) Extended dependence graph.

as the graph $\mathcal{EDG} = (\mathcal{N}, \mathcal{A})$, where $\mathcal{N} = \{F, V, W\}$ and $\mathcal{A} = \{(F, F), (F, V), (F, W), (V, V), (W, W)\}$. As is clear, variables $F$ and $W$ are separable in $\mathcal{EDG}$.

5.3.3 Regularisation

We want to make the data dependence $\mathcal{DD}_2$ uniform, by applying parametric uniformisation according to Proposition 4.4.11. Let $p \in \mathbb{N}^+$. We note that:

- for all $(i, j), 0 < W(i, j) \leq c$, then $W = \lfloor c/(p + 1) \rfloor$.

- $\text{lin}(D) = \mathbb{Z}^2$ and $d = (0, -1) \in \text{lin}(D)$, hence we need to reindex the system in $\mathbb{Z}^3$. Let $k$ indicate the index of the added axis. We choose the new system of axes characterised by the indices $i, j, k$ in this order.
- $\pi$ can be chosen as the vector $\pi = (0, 0, 1)$. As is clear, $\pi$ is in $D_3^4$ and $D_4$ is contained in the hyperplane $[\pi : 0]$. Then $\tilde{d} = d + \pi = (0, -1.1)$ and $\tilde{d} = d - \pi = (0, -1, -1)$.

- $tr$ is the translation defined as $tr(i, j, k) = (i, j, k) + (0, -\tilde{W}, \tilde{W}) = (i, j - \tilde{W}, k + \tilde{W})$, and $ren$ renames $W$ as $W'$. The system of equations defining $W$ is $DefS_W = \{E_7, E_8\}$, and its image under $tr$ and $ren$ is $(DefS_W)^{tr, ren}$ defined as:

\[
E_7' = (D_2^{tr}, W^{tr}, inv_{W^{tr}}) \\
E_8' = (D_4^{tr}, W^{tr}, W^{tr}, id, T_3^{tr})
\]

with domains:

\[
D_2^{tr} = \{(i, j, k) \mid 1 \leq i \leq n, j = -\tilde{W}, k = \tilde{W}\} \\
D_4^{tr} = \{(i, j, k) \mid 1 \leq i \leq n, 1 - \tilde{W} \leq j \leq c - \tilde{W}, k = \tilde{W}\}
\]

and index mapping:

\[
T_3^{tr}(i, j, k) = t \circ T_3 \circ t^{-1}(i, j, k) = (i, j, k) + (0, -1, 0).
\]

The parametric uniformisation of $DD_2$ produces the system of equations:

(variable $F$)

\[
E_1 = (D_1, F, inv_1^F) \\
E_2 = (D_2, F, inv_2^F) \\
E_3 = (D_3, F, inv_3^F) \\
E_4 = (D_4, F, (F, R^1, V), f, (T_1, T_2, T_3))
\]

(variable $V$)

\[
E_5 = (D_2, V, inv) \\
E_6 = (D_4, V, V, id, T_3)
\]

(variable $W^{tr}$)

\[
E_7 = (D_2^{tr}, W^{tr}, inv_{W^{tr}}) \\
E_8 = (D_4^{tr}, W^{tr}, W^{tr}, id, T_3^{tr})
\]

(routing variables $R^1, R^2$ and $R^3$)

\[
E_9 = (D_4, R^1, (\alpha, \beta, \gamma, R^1, R^2, \ldots, R^2), f', (T_2, T_2, T_2, T_4, T_5, 0, \ldots, T_{5,p}))
\]
\[ E_{10} = (D_{42,1}, R^2, R^3, id, I_6) \]
\[ E_{11} = (D_{42,2}, R^2, F, id, I_2) \]
\[ E_{12} = (D_{42,3}, R^3, R^2, id, I_7) \]

(control variables \( \alpha, \beta \) and \( \gamma \))

\[ E_{13} = (D_{41,1}, \alpha, \alpha, id, I_4) \]
\[ E_{14} = (D_{41,2}, \alpha, W^{tr}, (p + 1) - \text{floor}, I_2) \]
\[ E_{15} = (D_{41,1}, \beta, \beta, id, I_4) \]
\[ E_{16} = (D_{41,2}, \beta, W^{tr}, \text{mod}_{p+1}, I_2) \]
\[ E_{17} = (D_{41,1}, \gamma, \gamma, \text{dec}, I_4) \]
\[ E_{18} = (D_{41,2}, \gamma, \text{in}_\gamma) \]

with:

- index mappings:

\[ I_1(i, j, k) = (i, j, k) + (-1, 0, 0) \]
\[ I_2(i, j, k) = (i, j, k) \]
\[ I_3(i, j, k) = (i, j, k) + (0, -1, 0) \]
\[ I_5^{tr}(i, j, k) = (i, j, k) + (0, -1, 0) \]
\[ I_4(i, j, k) = (i, j, k) + (0, -1, 1) \]
\[ I_{5,0}(i, j, k) = (i, j, k) \]
\[ I_{5,1}(i, j, k) = i, j, k + (0, -1, 0) \]

\[ \ldots \]
\[ I_{5,p}(i, j, k) = (i, j, k) + (0, -p, 0) \]
\[ I_6(i, j, k) = (i, j, k) + (0, -1, -1) \]
\[ I_7(i, j, k) = (i, j, k) + (0, 1 - p, 0) \]

- applied functions:

\[ \text{in}^1_P(i, j, k) = 0 \]
\[ \text{in}^2_P(i, j, k) = -\infty \]
\[ \text{in}_V(i, j, k) = v_i \]
\[ \text{in}^*_W(i, j, k) = w_i \]
in_{\gamma}(i, j, k) = \bar{W} \\
(p + 1)_{\text{floor}}(a) = \lfloor a/(p + 1) \rfloor \\
mod_{p+1}(a) = a \pmod{(p + 1)} \\
id(a) = a \\
dec(a) = a - 1 \\
f(a, b, c) = \max(a, b + c) \\
f'(\alpha, \beta, \gamma, a_1, a_2, \ldots, a_p, \ldots) = \left\{ \begin{array}{ll}
a_1 & \alpha \neq \gamma \\
a_2, 0 & \alpha = \gamma, \beta = 0 \\
\ldots & \\
a_2, p & \alpha = \gamma, \beta = p \end{array} \right. \\
- \text{ domains:} \\
D_1 = \{(i, j, k) \mid i = 0, 1 \leq j \leq c, k = 0\} \\
D_2 = \{(i, j, k) \mid 1 \leq i \leq n, j = 0, k = 0\} \\
D_3 = \{(i, j, k) \mid 1 \leq i \leq n, j < 0, k = 0\} \\
D_4 = \{(i, j, k) \mid 1 \leq i \leq n, 1 \leq j \leq c, k = 0\} \\
D_{41,1} = \{(i, j, k) \mid 1 \leq i \leq n, 0 \leq k \leq \bar{W}, 1 - k \leq j \leq c - k\} \\
D_{41,2} = \{(i, j, k) \mid 1 \leq i \leq n, 0 \leq k \leq \bar{W}, k - W \leq j \leq c - k\} \\
D_{42,1} = \{(i, j, k) \mid 1 \leq i \leq n, 0 \leq k \leq \bar{W}, k - 2W \leq j \leq c - k\} \\
D_{42,2} = \{(i, j, k) \mid 1 \leq i \leq n, 0 \leq k \leq \bar{W}, k - W \leq j \leq c - k\} \\
D_{42,3} = \{(i, j, k) \mid 1 \leq i \leq n, 0 \leq k \leq \bar{W}, k - 2W \leq j \leq c - k\} \\
\text{Domains } D_4, D_{41} \text{ and } D_{42} \text{ are sketched in Fig. 5.23, a), b) and c), respectively, while Fig. 5.24 a) and b) illustrate sections of the data dependence graphs in } D_{41} \text{ and } D_{42} \text{ for values of the parameter } p = 1, 2, \text{ respectively (in the figure, we assume } c = 8). \text{ Only the routing variables } R^1, R^2 \text{ and } R^3 \text{ are considered.} \\

5.3.4 Space-Time Mapping

After regularisation, all data dependencies are uniform. They are summarised in Table 5.10 (with similar conventions to those in Section 5.1.6). The corresponding dependence cone is

$$\Theta^* = cone(\{(0, 1, 0), (0, 2, 0), \ldots, (0, p, 0), (0, 1, -1), (0, 1, 1), (1, 0, 0)\}).$$
Fig. 5.23. Domains: a) $D_4$; b) $D_{41}$; c) $D_{42}$.

Fig. 5.24. Sections of the routing domains $D_{41}$ and $D_{42}$ for values of the parameter: a) $p = 1$; b) $p = 2$. 
**Table 5.10. Summary of the data dependencies.**

<table>
<thead>
<tr>
<th>Pair</th>
<th>Unifor.</th>
<th>DDVector</th>
</tr>
</thead>
<tbody>
<tr>
<td>F, F</td>
<td>–</td>
<td>(1, 0, 0)</td>
</tr>
<tr>
<td>F, F</td>
<td>F, R₁</td>
<td>(0, 0, 0)</td>
</tr>
<tr>
<td></td>
<td>R₁, R₁</td>
<td>(0, 1, −1)</td>
</tr>
<tr>
<td></td>
<td>R₁, R₂</td>
<td>(0, 0, 0)</td>
</tr>
<tr>
<td></td>
<td>R₂, R₂</td>
<td>(0, 1, 0)</td>
</tr>
<tr>
<td></td>
<td>…</td>
<td></td>
</tr>
<tr>
<td></td>
<td>R₁, R₂</td>
<td>(0, p, 0)</td>
</tr>
<tr>
<td></td>
<td>R₂, R₃</td>
<td>(0, 1, 1)</td>
</tr>
<tr>
<td></td>
<td>R₃, R₂</td>
<td>(0, p − 1.0)</td>
</tr>
<tr>
<td></td>
<td>R₂, F</td>
<td>(0, 0, 0)</td>
</tr>
<tr>
<td>F, V</td>
<td>–</td>
<td>(0, 1, 0)</td>
</tr>
<tr>
<td>V, V</td>
<td>–</td>
<td>(0, 1, 0)</td>
</tr>
<tr>
<td>Wᵀᵣ, Wᵀᵣ</td>
<td>–</td>
<td>(0, 1, 0)</td>
</tr>
<tr>
<td>α, α</td>
<td>–</td>
<td>(0, 1, −1)</td>
</tr>
<tr>
<td>β, β</td>
<td>–</td>
<td>(0, 1, −1)</td>
</tr>
<tr>
<td>γ, γ</td>
<td>–</td>
<td>(0, 1, −1)</td>
</tr>
</tbody>
</table>

Fig. 5.25. Dependence cone after uniformisation.

with extremal rays (0, 1, −1), (0, 1, 1) and (1, 0, 0). The dependence cone, illustrated in Fig. 5.25, is pointed.

A valid affine timing function is determined by any \( \lambda = (\lambda_1, \lambda_2, \lambda_3) \in \mathbb{Z}^3 \) such that the following system of inequalities is satisfied:

\[
\begin{align*}
\lambda_2 - \lambda_3 &> 0 \\
\lambda_2 + \lambda_3 &> 0 \\
\lambda_1 &> 0
\end{align*}
\]

The vector \( \lambda = (1, 1, 0) \) is a possible choice, corresponding to the timing function \( t(i, j, k) = i + j \). With this timing function, the algorithm is executed in \( O(c + 2\lceil c/2 \rceil + n) \) steps.
<table>
<thead>
<tr>
<th>$d$</th>
<th>$\lambda \cdot d$</th>
<th>$a(d)$</th>
<th>$a'(d)$</th>
<th>$a''(d)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$(1,0,0)$</td>
<td>1</td>
<td>$(1,0)$</td>
<td>$(1,0)$</td>
<td>$(0,0)$</td>
</tr>
<tr>
<td>$(0,1,0)$</td>
<td>1</td>
<td>$(0,0)$</td>
<td>$(-1,0)$</td>
<td>$(1,0)$</td>
</tr>
<tr>
<td>$(0,2,0)$</td>
<td>2</td>
<td>$(0,0)$</td>
<td>$(-2,0)$</td>
<td>$(2,0)$</td>
</tr>
<tr>
<td>$(0,p,0)$</td>
<td>$p$</td>
<td>$(0,0)$</td>
<td>$(-p,0)$</td>
<td>$(p,0)$</td>
</tr>
<tr>
<td>$(0,1,-1)$</td>
<td>1</td>
<td>$(0,-1)$</td>
<td>$(-1,-1)$</td>
<td>$(1,-1)$</td>
</tr>
<tr>
<td>$(0,1,1)$</td>
<td>1</td>
<td>$(0,1)$</td>
<td>$(-1,1)$</td>
<td>$(1,1)$</td>
</tr>
</tbody>
</table>

Table 5.11. Transformation of data dependence vectors under the space-time mappings $[t,a]$, $[t,a']$, and $[t,a'']$.

A compatible linear allocation function is a projection according to any non-null vector $u = (u_1, u_2, u_3) \in \mathbb{Z}^3$, such that $\lambda \cdot u \neq 0$, i.e., $u \in \mathbb{Z}^3$ such that $u_1 + u_2 \neq 0$. A possible projection vector is $u = (0,1,0)$, corresponding to the allocation function $a(i,j,k) = (i,k)$. The transformation of the dependence vectors of the system under the space-time mapping $[t,a]$ is summarised in Table 5.11.

The resulting signal flow graph is illustrated in Fig. 5.26 a), where the left-hand side illustrates the nodes and the non-parametric communication channels, while the right-hand side details the parametric communication channels at each node. For simplicity arcs with a unit delay are not labelled. Note that there are three types of nodes in the graph (illustrated as white, black and dotted nodes).

Under $[t,a]$, the $p$ parametric communication links may be implemented as $p$ locations of the local RAM at each processing element of the array. The three types of cells are described in Fig. 5.27 a).

Different compatible placements are determined, for instance, by the projection vectors $u' = (1,1,0)$ and $u'' = (1,0,0)$, corresponding to the allocation functions $a'(i,j,k) = (i-j,k)$ and $a''(i,j,k) = (j,k)$, respectively. The transformation of the dependence vectors under $[t,a']$ and $[t,a'']$ are also given in Table 5.11. The resulting signal flow graphs are illustrated in Fig. 5.26 b) and c), while the corresponding processing elements are sketched in Fig. 5.27 b) and c), respectively.

### 5.4 Gaussian Elimination with Partial Pivoting

Gaussian elimination reduces an $n \times n$ matrix $A = [a_{ij}]$ to a triangular form in $n-1$ iterations. At each iteration the elements, under the main diagonal, of one of the columns of the matrix
Fig. 5.26. Signal flow graphs corresponding to: a) \([t, a]\); b) \([t, a']\); c) \([t, a'']\).

Fig. 5.27. Processing elements corresponding to: a) \([t, a]\); b) \([t, a']\); c) \([t, a'']\).
Fig. 5.28. Pivot operation: the black circle indicates the pivot; the white circle indicates the entry to be updated; and the other circles indicate the corresponding row and column coefficients.

are nullified by a series of elementary row operations\(^3\). The columns are processed from left to right. At each iteration \(t\), for \(1 \leq t \leq n - 1\), a new matrix is generated as follows. The element \(a_{t,t}\) of the current matrix is selected and used to update each \(a_{i,j}\), with \(i > t\) and \(j \geq t\), by applying the operation \(a_{i,j} - a_{t,t} \cdot a_{t,j}/a_{t,t}\).

When Gaussian elimination is combined with partial pivoting, at iteration \(t\), an element of the subcolumn \([a_{t,t}, a_{t+1,t}, \ldots, a_{n,t}]\) with maximum modulo is chosen as the pivot at that iteration and used to update the current matrix. Let \(a_{p,t}\) denote the pivot at iteration \(t\). The new matrix is updated by applying the operation \(a_{i,j} - a_{t,t} \cdot a_{p,j}/a_{p,t}\), for all \(i, j\) such that \(i \geq t, i \neq p\) and \(j \geq t\). (By generalising, Gaussian elimination without pivoting may be seen as a form of pivoting in which at each iteration \(t\), \(a_{p,t} = a_{t,t}\).)

A way of exploiting parallelism in Gaussian elimination with pivoting is to break the computation of the operation which updates the elements of the matrix into sub-computations, which can be executed in a distributed fashion. Let us consider the operation \(a_{i,j} - a_{t,t} \cdot a_{p,j}/a_{p,t}\). The quotient \(a_{i,t}/a_{p,t}\) is used to update all the elements of the \(i^{th}\) row of the matrix, while \(a_{p,j}\) is applied to all the elements of the \(j^{th}\) column. This is illustrated in Fig. 5.28. These values can be treated as coefficients, which can be propagated among computation points and used for the parallel updating of the matrix. In the following we will call them row and column coefficients, respectively.

In Gaussian elimination without pivoting, at each iteration a particular element of the matrix is selected according to its position in the matrix (namely, \(a_{t,t}\) at iteration \(t\)). When pivoting is present, the pivot is selected according to its absolute value as an entry of the current matrix, and this value is known at run-time only, i.e., when the algorithm is executed on

\(^3\)Elementary row operations correspond to changes of basis of a vector space [Ner63]. They include: swapping rows, multiplying a row by a non-zero scalar; and subtracting from one row a multiple of another row.
actual data. It is this characteristics which makes Gaussian elimination with pivoting a dynamic problem.

5.4.1 Dynamic Formulation

In order to specify Gaussian elimination with pivoting as a dynamic problem, let us consider the basic operation

\[ a_{i,j} = a_{i,t} \cdot a_{p,j}/a_{p,t} \]

more closely. The index \( p \) indicating the row of the pivot is actually a function of the iteration index \( t \), and should be indicated more precisely as \( p(t) \). Therefore, both \( a \) and \( p \) can be seen as variables to be computed by the algorithm. Then at each iteration \( t \), for \( t = 1, \ldots, n \), the algorithm computes the values:

\[ p(t) = \bigcap \{a_{i,t}^{-1} \mid t \leq i \leq n\} \]

\[ a_{i,j}^t = a_{i,j}^{t-1} - a_{i,t}^{t-1} \cdot a_{p(t),j}^{t-1}/a_{p(t),t}^{t-1} \]

for all \( i, j \) such that \( i \geq t \) and \( i \neq p(t) \). The operation \( \bigcap \) compares the entries of the \( t \)th column of the current matrix to find the pivot, and once found it, returns the corresponding row index. The initial values are \( a_{i,j}^0 = a_{i,j} \) for all \( i, j \).

If we had to specify the problem more formally as a system of dynamic recurrence equations we would soon realise that the techniques we have developed are not powerful enough to deal with this problem, the reason being that the computations of \( p(t) \) and the updating of the entries \( a_{i,j}^t \) of the matrix are interleaved at each iteration. This fact introduces a mutual dependence between the computations of the pivot and the new entries of the current matrix so that the variables corresponding to \( p \) and \( a \) are not separable.

5.4.2 Static Formulation

Regular array designs for Gaussian elimination with partial pivoting have been proposed in the literature, which are based on a static formulation of the problem [BaEl88, Meg90, ElBa90]. This is obtained if, at each iteration, the elements of the current matrix are rearranged before the new entries are computed. In this way the pivot always assumes a predetermined position.

Each iteration of the new algorithm includes the following operations: finding the pivot and rearranging the elements of the matrix; determining and propagating the row and column coefficients; and computing the new entries of the matrix. We note that:
- at each iteration of the algorithm the same sequence of operations is carried out on the current matrix: hence the specification of the algorithm can be simplified by concentrating on the single iterations (otherwise we would need to consider a specification in 4 dimensions);

- the size of the current matrix decreases at each iteration: hence the specification of the iterations can be induced from the specification of the first iteration by a restriction to appropriate submatrices;

- the output matrix of each iteration represents the input matrix of the following iteration: hence a strong sequentiality constraint exists between consecutive iterations, and the overall design of the algorithm can be obtained as a sequential composition of the specifications of the single iterations. Indeed pipelining techniques should be used to overlap some of the operations of consecutive iterations.

Preliminary Specification

At each iteration of the algorithm, the pivot is chosen as the element with maximum modulo in a set of entries of the matrix. In this section we illustrate a technique for the selection of the pivot, which is used in the specifications of the following sections. The technique is a sorting of the elements according to their absolute value.

Let us consider \( n \) numbers, \( a_1, \ldots, a_n \). Sorting the \( n \) numbers \( a_1, \ldots, a_n \) according to their absolute value can be realised as follows. We introduce two variables \( V, H \), with \( V \) initialised with the \( n \) given numbers (plus \( n - 1 \) zero entries) and \( H \) with all its \( n - 1 \) entries equal to \( \infty \). The computations of \( V \) and \( H \) in a two dimensional space are illustrated by the data dependence graph in Fig. 5.29 (in the figure we have assumed \( n = 4 \)). At each point \((i, j)\) the pair of values \( V(i - 1, j) \) and \( H(i, j - 1) \) are compared. The element with greater absolute value is stored in \( V(i, j) \), while the other element in \( H(i, j) \). The recurrences are:

\[
\begin{align*}
\text{(variable } V) \\
E_1 &= (D_1, V, inv_1) \\
E_2 &= (D_2, V, inv_2) \\
E_3 &= (D_4, V, (V, H), |max|, (I_1, I_2)) \\
&\quad \text{(variable } H) \\
E_4 &= (D_3, H, in_H)
\end{align*}
\]
Fig. 5.29. Sorting: a) data dependence graph; b) domains.

\[ E_5 = (D_4, H, (V, H), |\text{min}|, (I_1, I_2)) \]

where:

- the index mappings are:
  \[ I_1(i, j) = (i - 1, j) \]
  \[ I_2(i, j) = (i, j - 1) \]

- the applied functions are:
  \[ \text{inv}_1(i, j) = a_j \]
  \[ \text{inv}_2(i, j) = 0 \]
  \[ \text{in}_H(i, j) = \infty \]
  \[ |\text{max}|(a, b) = \begin{cases} a & a \geq b \\ b & \text{otherwise} \end{cases} \]
  \[ |\text{min}|(a, b) = \begin{cases} b & a \geq b \\ a & \text{otherwise} \end{cases} \]

- the domains are:
  \[ D_1 = \{(i, j) | i = 0, 1 \leq j \leq n\} \]
  \[ D_2 = \{(i, j) | 0 \leq i \leq n - 2, j = i + n + 1\} \]
  \[ D_3 = \{(i, j) | 1 \leq i \leq n - 1, j = i - 1\} \]
  \[ D_4 = \{(i, j) | 1 \leq i \leq n - 1, i \leq j \leq i + n\} \]
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Specification

At each iteration $t$, the subcolumn $[a_{t,t}, a_{t+1,t}, \ldots a_{n,t}]$ of the matrix is scanned to identify the pivot. Every time two elements of the column are swapped, their corresponding rows are also exchanged. This provides the necessary rearrangement of the elements of the matrix before updating its entries. Once all the rows of the matrix have been rearranged, the row and column coefficients are determined and the new matrix computed.

Let us consider the first iteration of the algorithm, i.e., $t = 1$. The iteration can be represented in a 3-dimensional space, in which we assume that $A$ is input on the $(k,j)$-plane (the index $k$ corresponds to columns of $A$). For $k = 1$, the elements of the first column of $A$ are processed, looking for the pivot, as explained in the previous section. When the values of $H(i,j-1,1)$ and $V(i-1,j,1)$ are swapped, a control signal (variable $C$ below) is issued and propagated upwards (direction of $(0,0,1)$). In this way, for $k > 1$, the corresponding exchange of $H(i,j-1,k)$ and $V(i-1,j,k)$ is performed. The first phase of the iteration can be specified as follows:

\begin{align*}
E_1 &= (D_1, V, invV_1) \\
E_2 &= (D_2, V, invV_2) \\
E_3 &= (D_{4,1}, V, (V,H), |max|, (I_1,I_2)) \\
E_4 &= (D_{4,2}, V, (V,H,C), swapV, (I_1,I_2,I_3)) \\
E_5 &= (D_3, H, inH) \\
E_6 &= (D_{4,1}, H, (V,H), |min|, (I_1,I_2)) \\
E_7 &= (D_{4,2}, H, (V,H,C), swapH, (I_1,I_2,I_3)) \\
E_8 &= (D_{4,1}, C, (V,H), |cmp|, (I_1,I_2)) \\
E_9 &= (D_{4,2}, C, C, id, I_3)
\end{align*}

where:

- the index mappings are:

$$I_1(i,j,k) = (i-1,j,k)$$
\[ I_2(i, j, k) = (i, j - 1, k) \]
\[ I_3(i, j, k) = (i, j, k - 1) \]

- the applied functions are:

\[ \text{inV}_1(i, j, k) = a_{j,k} \]
\[ \text{inV}_2(i, j, k) = 0 \]
\[ \text{inH}(i, j, k) = \infty \]
\[ \text{id}(a) = a \]
\[ \text{swapv}(a, b) = b \]
\[ \text{swapH}(a, b) = a \]
\[ |\text{max}|(a, b) = \begin{cases} a & a \geq b \\ b & \text{otherwise} \end{cases} \]
\[ |\text{min}|(a, b) = \begin{cases} b & a \geq b \\ a & \text{otherwise} \end{cases} \]
\[ |\text{cmp}|(a, b) = \begin{cases} 0 & a \geq b \\ 1 & \text{otherwise} \end{cases} \]

- the domains are:

\[ D_1 = \{(i, j, k) \mid i = 0, 1 \leq j \leq n, 1 \leq k \leq n\} \]
\[ D_2 = \{(i, j, k) \mid 0 \leq i \leq n - 2, j = i + n + 1, 1 \leq k \leq n\} \]
\[ D_3 = \{(i, j, k) \mid 1 \leq i \leq n - 1, j = i - 1, 1 \leq k \leq n\} \]
\[ D_{4,1} = \{(i, j, k) \mid 1 \leq i \leq n - 1, i \leq j \leq i + n, k = 1\} \]
\[ D_{4,2} = \{(i, j, k) \mid 1 \leq i \leq n - 1, i \leq j \leq i + n, 2 \leq k \leq n\} \]

\(|\text{cmp}|\) compares the absolute values of its arguments and sets the signal \(C\) to 1 if an exchange is necessary (0 otherwise). \(\text{swapH}\) and \(\text{swapv}\) swap the values of \(H\) and \(V\) if the signal \(C\) is set to 1 (they pipeline \(H\) and \(V\), respectively, otherwise). At the end of this phase, \(V(n - 1, j, 1)\), for \(n \leq j \leq 2n - 1\), contain the elements of the pivot column, sorted in decreasing absolute value, and \(V(n - 1, n, 1)\) is the pivot. These elements are used to compute the row coefficients, which are subsequently pipelined upwards (direction of \((0, 0, 1)\)). The pivot row is contained in \(V(n - 1, n, k)\), for \(1 \leq k \leq n\). These elements represent the column coefficients, which are propagated horizontally (direction of \((0, 1, 0)\)). The computation of the new matrix is specified by:

\[(\text{variable } V)\]
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\[ E_{10} = (D_{5,1}, V, inv_3) \]
\[ E_{11} = (D_{5,2}, V, (V, H, RC).update, (I_1, I_2, I_3)) \]
\[ E_{12} = (D_6, V, V, id, I_1) \]
\[ \text{(variable H - column coefficients)} \]
\[ E_{13} = (D_6, H, V, id, I_1) \]
\[ E_{14} = (D_7, H, H, id, I_2) \]
\[ \text{(variable RC - row coefficients)} \]
\[ E_{15} = (D_{5,1}, RC, (V, H), div, (I_1, I_2)) \]
\[ E_{16} = (D_{5,2}, RC, RC, id, I_3) \]

where:

- the index mappings are as above;
- the applied functions are:
  \[ inv_3(i, j, k) = 0 \]
  \[ id(a) = a \]
  \[ div(a, b) = a/b \]
  \[ update(a, b, c) = a - b \times c \]
- the domains are:
  \[ D_{5,1} = \{(i, j, k) | i = n, n + 1 \leq j \leq 2n - 1, k = 1\} \]
  \[ D_{5,2} = \{(i, j, k) | i = n, n + 1 \leq j \leq 2n - 1.2 \leq k \leq n\} \]
  \[ D_6 = \{(i, j, k) | i = n, j = n, 1 \leq k \leq n\} \]
  \[ D_7 = \{(i, j, k) | i = n, n + 1 \leq j \leq 2n - 1, 1 \leq k \leq n\}. \]

The data dependence graph is given in Fig. 5.30 a) and b), and for \( k = 1 \) in Fig. 5.31. The iteration itself is illustrated in Fig. 5.32.

5.4.3 Space-Time Mapping

At each iteration \( t \) for \( 1 \leq t \leq n - 1 \), a submatrix of \((n+1-t) \times (n+1-t)\) elements is considered. as the remaining \( t - 1 \) rows and columns of the current matrix are already in their final format. On each of these submatrices the same sequence of operations is performed. Therefore the
CHAPTER 5. CASE STUDIES

Fig. 5.30. Data dependence graph: a) finding the pivot \((i = c \text{ and } 1 \leq c \leq n - 1)\); b) computing the new matrix \((i = n)\).

Fig. 5.31. Iteration for \(k = 1\): a) data dependence graph; b) domains.

Fig. 5.32. Phases of the iteration: a) finding the pivot; b) computing the new matrix; c) composition of the phases.
specification given in the previous section can be used as the specification of each iteration of the algorithm (indeed the computation domains have to be "resized" accordingly at each iteration). In order to obtain an array design for the algorithm, we first apply a space-time mapping to each iteration and compose the mapped iterations subsequently. Indeed, the combinations of different mappings and compositions of the iterations produce several distinct array designs for the algorithm.

The data dependencies of the specification are uniform and the corresponding dependence cone is

$$\Theta^* = \text{cone}(\{(1, 0, 0), (0, 1, 0), (0, 0, 1)\}).$$

The dependence cone, illustrated in Fig. 5.33, is pointed.

A valid affine timing function is determined by any $$\lambda = (\lambda_1, \lambda_2, \lambda_3) \in \mathbb{Z}^3$$ such that the following system of inequalities is satisfied:

$$\lambda_1 > 0$$
$$\lambda_2 > 0$$
$$\lambda_3 > 0.$$

The vector $$\lambda = (1, 1, 1)$$ is a possible choice, determining the timing function $$t(i, j, k) = i + j + k.$$ With this timing function, a single iteration on an $$n \times n$$ matrix requires $$O(4n)$$ steps.

A compatible allocation is determined by any projection vector $$u = (u_1, u_2, u_3) \in \mathbb{Z}^3$$ such that $$\lambda \cdot u \neq 0,$$ that is $$u_1 + u_2 + u_3 \neq 0.$$ For instance, $$u = (0, 1, 0)$$ satisfies the condition and determines the linear scheduling $$a(i, j, k) = (i, k).$$ The corresponding projection, summarised in Table 5.12, produces the signal flow graph of Fig. 5.34. There are four types of nodes in the graph: type a) (represented as white nodes) corresponds to the computations to determine the pivot; type b) (black nodes) corresponds to the computations to rearrange the entries of
### Table 5.12. Transformation of data dependence vectors under the space-time mapping $[t, a]$.

<table>
<thead>
<tr>
<th>$d$</th>
<th>$\lambda \cdot d$</th>
<th>$a(d)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$(1, 0, 0)$</td>
<td>1</td>
<td>$(1, 0)$</td>
</tr>
<tr>
<td>$(0, 1, 0)$</td>
<td>1</td>
<td>$(0, 0)$</td>
</tr>
<tr>
<td>$(0, 0, 1)$</td>
<td>1</td>
<td>$(0, 1)$</td>
</tr>
</tbody>
</table>

Fig. 5.34. Signal flow graph of the iteration.

Fig. 5.35. Processing elements.
the current matrix; type c) (crossed white node at the bottom left-hand corner) corresponds to the computations to determine the row coefficients; and type d) (white dotted nodes) corresponds to the computations to update the entries of the current matrix. A description of the corresponding processing elements and their operations is given in Fig. 5.35. Note that signal flow graph and processing elements refer to a single iteration.

A sequential composition of the iterations produces the signal flow graph in Fig. 5.36 a) (where we have omitted the loops at each node and different types of nodes are not distinguished). As the iterations of the algorithm are strictly sequential, a more compact graph (which yields a more efficient use of the processing elements) can be obtained by superimposing the iterations. The resulting signal flow graph is given in Fig. 5.36 b). Regular array designs corresponding to both these graphs were proposed by Megson in [Meg90]. These designs compute the algorithm in, respectively, $O(2n^2 + n)$ and $O(n^2 + 3n)$ time steps. Note that, as data are pipelined between consecutive iterations, some of the computation of the next iteration can begin before those of the current iteration have all been completed.

Finally, note that both designs require the specification of control signals (which for brevity we have omitted) in order to tag the beginning of each new iteration.

### 5.5 Summary

We have presented a number of case studies for the illustration of the techniques developed in the previous chapters of this thesis, and shown that some interesting problems from the literature, such as cyclic reduction techniques for the solution of tridiagonal systems or the knapsack problem, can be treated systematically with our methods.

From the work presented in this chapter we can make a number of observations. While, in general, some expertise and problem-specific knowledge is required from the designer both
for the initial specification of an algorithm and the application of the subsequent transformations, synthesis methods provide guidelines to support the designer in his/her task as well as the mechanisation of some of the design steps. These guidelines include, for instance, conditions related to the scalability of the array design (i.e., the uniformity of the regularisation directions with respect to the size parameters of the problem) and the existence of an affine scheduling (i.e., guaranteeing that the dependence cone is pointed). For example, in the case of cyclic reduction for the solution of tridiagonal systems, we could identify the lack of scalability of our specification in the early phases of the design, by analysing the generators of the dependence cone. On the other hand, little support is provided by synthesis methods to indicate how a "better" specification can be chosen. For instance, in the same example, the transformation $T$ at the basis of the new specification was chosen arbitrarily, based on our knowledge of the regularisation techniques and computational model.

Characterising what makes a specification particularly suitable for regular array synthesis is not a trivial problem, and very little work exists in the literature on the subject (the problem is mentioned in [LeXu91]). It is, however, an important issue as sometimes reformulating a problem is necessary (for instance, because the synthesis method breaks down). This was the case in our example of an $M$-to-1 decimator, where neither integral nor affine regularisation techniques could be applied successfully. Note that the necessary reformulation was not just a simple syntactic manipulation of the recurrences, but was based on a totally different view of the algorithm.

Changing the specification was also necessary in our example of Gaussian elimination with pivoting, where we replaced a dynamic algorithm with a static specification. We have already discussed (in Chapter 4) how a transformation from dynamic to static is necessary for the synthesis of dynamic problems as regular arrays. However, the type of transformation that we have applied in the example appears to be more complex than those we have defined through our regularisation techniques in Chapter 4. In particular, the transformation of the example was applied to the algorithm as a whole rather than its single dynamic data dependencies (as we do with our method). The relation between complexity and scope of application of synthesis transformations was discussed in Chapter 1, where we stressed how synthesis methods have developed by favouring simple locally applicable transformations. The limitations of this approach appear to be critical with dynamic problems, where more global (hence complex) transformations seem to be required (even our techniques require the redefinition of sub-systems of equations in order to make a dynamic data dependence
uniform).

The two dynamic examples, knapsack problem and Gaussian elimination with pivoting, have been useful to stress the type of problems characterised by finitely generated dynamic recurrences. In the knapsack problem the dynamicity is resolved as soon as the weight distribution is known. The control variables are assigned once and maintain their values throughout the algorithm. On the other hand, Gaussian elimination requires to compute a new pivot at each iteration, hence (the same) dynamic data dependence relations have to be resolved several times throughout the algorithm.

We may conclude that finitely generated recurrences characterise a restricted class of dynamic problems, and that a more general treatment of dynamic problems requires the consideration of specifications at a more global (and abstract) level. We will return to these points in the following final chapter.
Chapter 6

Conclusions

The focus of this thesis was the systematic synthesis of regular arrays. We have discussed some of the major benefits of synthesis methods such as the provision of a disciplined and rigorous approach to algorithm design (the various stages of the design process are clearly identified and supported by the methodology, and formal notation and transformations guarantee that the approach is well-founded), or the fact that they represent an effective way of engineering parallel algorithms (as computationally powerful methods can be developed based on the underlying mathematical model of affine Euclidean geometry with embedded lattice spaces). We have also exposed some of the limitations of the synthesis methods, primarily their restricted scope of application, and the fact that, even for problems in their scope, synthesis techniques have not (as yet) reached the maturity of a formal method which provides best practice without expertise [Hal95]. In particular, a fairly high level of expertise is required from the specifier throughout the design process to guide the transformations and provide efficient solutions under the given problem constraints.

The work of this thesis has aimed at overcoming some of the limitations of the existing techniques by widening their applicability to more general classes of problems. In doing so, the provision of practical solutions was one of our major concerns, and our techniques were developed within the traditional mathematical framework of synthesis methods. The two key issues in our developments have been: the identification of classes of non-affine problems; and their systematic transformations into regular problems to which classic mapping techniques can be applied. Related issues of scheduling and placement were also addressed.

A major achievement of this work is the characterisation and systematic treatment of classes of integral and dynamic problems. That these problems are of practical interest has been demonstrated by considering case studies from the literature to which our techniques apply.
In the thesis we have defined novel engineering solutions for regular arrays, rather than improving on existing methods, hence we cannot assess the solutions we have proposed by direct comparison with other methods. We can, however, make a number of observations on our development and stress points which deserve further investigation.

The emphasis of the work has been more on the feasibility of the approach than on its optimality. An obvious source of inefficiency both for integral and dynamic techniques is the computational overhead due to the way data routing is defined. Although we have partially addressed the problem through the provision of parametric uniformisation techniques, the reduction of routing overhead is an area of research for which further work would be beneficial. Note that, while regularising integral data dependencies mainly reduces to the definition of a suitable routing system, the treatment of dynamic data dependencies involves the definitions of two classes of computations: those computations necessary to establish the data dependence relation at run-time, and those computations necessary to route the data accordingly once such a relation has been established. Therefore, more efficient regularisation techniques for dynamic data dependencies should not just reduce the routing overhead, but also maximise the parallel execution of the two classes of computations. In our techniques, this is achieved by separating the computations for determining the coefficients of a dynamic index mapping from those for computing the new values, hereby increasing their potential for parallelism. In the case of Gaussian elimination with pivoting, a similar effect is obtained, at each iteration, by rearranging the entries of the current matrix while determining the pivot. Indeed, efficient regularisation techniques could be developed by considering special subclasses of problems. This type of approach is taken in [Fr-et-al93, Sw-et-al94] for so-called piece-wise linear data dependencies (a type of integral dependencies in which the index mappings define piece-wise linear transformations).

A major difference between transformations for integral (in general static) and dynamic problems is the necessity, in the latter case, to consider algorithm specifications more globally rather than applying the transformations to single data dependencies in isolation. The reason for such a difference is that the computations which establish the dynamic data dependencies at run-time have to be taken into account as part of the new algorithm specification. With our techniques this is achieved by combining the separation of some computations of the algorithm and their translation in the computation space. A limitation of our approach, formally captured by the requirement of separability of the computations (see Section 4.3.6), is that the computation domain of the non-uniform data dependence remains unchanged.
throughout the transformations. In other words, the techniques provide a systematic routing of the data between pairs of computation points which are fixed by the specification. This characteristic of our techniques is also shared by classic regularisation techniques for affine problems [QuVa89] from which our approach has developed. However, more powerful regularisation techniques could be developed which modify the original computation domains, so that separability is not necessary. For instance, such techniques may allow the domains to be expanded so to include the (possibly interleaved) computations of both the coefficients of the index mapping and the routing of the data. Such an approach would be particularly beneficial to problems such as Gaussian elimination with pivoting, in which the determination of the pivot and the corresponding updating of the matrix entries are interleaved throughout the algorithm. Indeed, such sophisticated techniques are likely to be complex and based on non-affine domain transformations, making more difficult the development of corresponding tool support. Also, a possible introduction of new non-uniform data dependencies in the specification due to the modification of the original computation domain needs to be addressed.

Our work on dynamic problems raises a number of questions. First of all, it could be argued whether our definition of dynamic data dependence provides an adequate abstraction for dynamic problems of practical interest, or whether other and more general forms of dynamic problems should be considered (a taxonomy of static and dynamic problems based on data dependencies and task generation can be found in [Me-et-al95]). We have shown that our approach is general enough to characterise interesting problems from the literature. However, a better insight is needed on its applicability in general. Also we could question whether the mathematical framework in which such an approach has been developed is adequate. Our treatment of dynamic problems, even in the restricted connotation which we have considered, shows that the existing framework lacks of some of the necessary basic notions, such as that of input. Moreover, we feel that the extension that we have provided, while realising our objective, does not constitute a particular elegant mathematical model for the treatment of dynamic problems. At this stage of the work, however, we have preferred to remain within the framework of the established synthesis methods (so that our techniques can be made immediately available to the engineering community) rather than attempting to redefine the whole theory.

Finally, there are a number of issues which have not been addressed in this work. Among them, whether the computability of integral and dynamic problems can be established. It
is possible that, because of the relationships among the various types of recurrences, some of the results known for other classes of recurrences (see the overview in Section 2.5) may be generalised to integral and dynamic problems. Also, although we have stressed the importance of providing synthesis techniques which can be mechanised, we have not considered the issue explicitly. Tool support for regular array synthesis exists (see the overview in Section 2.5) including algorithms for the manipulation of the recurrences, their domains and data dependencies. Those algorithms are based on known techniques from linear algebra, computational geometry and linear programming (such as determining standard basis of vector spaces, computing the convex hull of sets of points or solving linear optimisation problems - see Appendix F and work in [Ner63, PrSh85, Sch86]). The same type of algorithms could be applied for the manipulation of the specifications as defined by our regularisation techniques, which could then be easily integrated with existing libraries of transformations for regular array synthesis.
Appendix A

Notation

[set theory]

R real numbers
Q rational numbers
Z integer numbers
N natural numbers
N+ positive integer numbers
R^n n-dimensional Euclidean space
Z^n n-dimensional Euclidean lattice space
U generic set
\mathcal{P}(U) powerset of U
|U| cardinality of U
U^n n-fold Cartesian product of U, i.e., U \times \ldots \times U n times
pr_i : U^n \rightarrow U i^{th} projection mapping, for i = 1, \ldots, n
[U - U] set of all mappings from U to U
u n-tuple in U^n
\mathcal{S}(u) support set of u, defined as \mathcal{S}(u) = \cup_{i=1}^n \{pr_i(u)\}
f generic function
f(U) image of U under f, i.e., \{f(u) | u \in U\}
f^m composition of f with itself m times, i.e., f \circ \ldots \circ f. m times

[linear and affine algebra]

I_n n \times n identity matrix
M generic matrix
M_j j^{th} row of M
M^{-1} inverse of M
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$M^T$</td>
<td>transpose of $M$</td>
</tr>
<tr>
<td>$\text{null}(M)$</td>
<td>null space of $M$</td>
</tr>
<tr>
<td>$e_j$</td>
<td>$j^{th}$ vector in the standard basis of $\mathbb{R}^n$</td>
</tr>
<tr>
<td>$[\pi : \theta]$</td>
<td>hyperplane $\mathbb{R}^n$ with normal vector $\pi$ and coefficient $\theta$</td>
</tr>
<tr>
<td>$D$</td>
<td>generic set in $\mathbb{R}^n$</td>
</tr>
<tr>
<td>$\text{lin}(D)$</td>
<td>direction of $D$</td>
</tr>
<tr>
<td>$\text{aff}(D)$</td>
<td>affine hull of $D$</td>
</tr>
<tr>
<td>$\text{conv}(D)$</td>
<td>convex hull of $D$</td>
</tr>
<tr>
<td>$P$</td>
<td>generic convex polyhedral set in $\mathbb{R}^n$</td>
</tr>
<tr>
<td>$\text{vert}(P)$</td>
<td>set of points generating $P$</td>
</tr>
<tr>
<td>$\text{ray}(P)$</td>
<td>set of directions generating $P$</td>
</tr>
<tr>
<td>$C$</td>
<td>generic convex polyhedral cone in $\mathbb{R}^n$</td>
</tr>
<tr>
<td>$d_1, \ldots, d_m$</td>
<td>$m$ vectors in $\mathbb{R}^n$</td>
</tr>
<tr>
<td>$\text{cone}{d_1, \ldots, d_m}$</td>
<td>the smallest convex polyhedral cone generated by $d_1, \ldots, d_m$</td>
</tr>
<tr>
<td>$\langle d_1, \ldots, d_m \rangle$</td>
<td>linear space generated by $d_1, \ldots, d_m$</td>
</tr>
<tr>
<td>$S$</td>
<td>generic space (or set) in $\mathbb{R}^n$</td>
</tr>
<tr>
<td>$S^\perp$</td>
<td>orthogonal space of $S$</td>
</tr>
<tr>
<td>$T$</td>
<td>generic affine transformation in $\mathbb{R}^n$</td>
</tr>
<tr>
<td>$\mathcal{L}_T$</td>
<td>linear part of $T$</td>
</tr>
</tbody>
</table>

[basic concepts in regular array synthesis]

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\mathcal{CS}$</td>
<td>computation space</td>
</tr>
<tr>
<td>$\mathcal{PS}$</td>
<td>processors space</td>
</tr>
<tr>
<td>$\mathit{Var}$</td>
<td>universe of variables</td>
</tr>
<tr>
<td>$\mathcal{I}$</td>
<td>generic index mapping</td>
</tr>
<tr>
<td>$\Theta_{\mathcal{I}}$</td>
<td>dependence mapping defined by $\mathcal{I}$</td>
</tr>
<tr>
<td>$\Omega_{\mathcal{I}}$</td>
<td>dependence domain defined by $\mathcal{I}$</td>
</tr>
<tr>
<td>$\Theta^\perp_{\mathcal{I}}$</td>
<td>dependence cone defined by $\mathcal{I}$</td>
</tr>
<tr>
<td>$\perp$</td>
<td>undefined value (for variables)</td>
</tr>
</tbody>
</table>

[equations and systems]

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E$</td>
<td>generic recurrence or input equation</td>
</tr>
<tr>
<td>$D_E$</td>
<td>computation domain of $E$</td>
</tr>
<tr>
<td>$\cdot E$</td>
<td>result of $E$</td>
</tr>
<tr>
<td>$E^*$</td>
<td>arguments of $E$</td>
</tr>
<tr>
<td>$f_E$</td>
<td>applied function of $E$</td>
</tr>
</tbody>
</table>
APPENDIX A. NOTATION

$\mathcal{IM}_E$  
index mappings of $E$

$\text{Var}_E$  
set of variables of $E$

$S$  
generic system of equations

$D_S$  
domain of $S$

$\text{Var}_S$  
set of variables of $S$

$V$  
generic variable

$\text{Def}_E V$  
definition equations of $V$

$\text{Def}_D V$  
definition domain of $V$

$\text{Def}_S V$  
definition subsystem of $V$

[data dependencies and graphs]

$\mathcal{DD}$  
generic data dependence

$\mathcal{DD}_E$  
$i^{th}$ data dependence of $E$

$\mathcal{DD}_E$  
data dependencies of $E$

$\mathcal{DD}_S$  
data dependencies of $S$

$\mathcal{DD}_G$  
generic data dependence graph

$\mathcal{CDD}_G$  
generic complete data dependence graph

$\mathcal{R}_D G$  
generic reduced dependence graph

$\mathcal{ED}_G$  
generic extended dependence graph

[timing and allocation functions]

$t$  
generic timing function

$\lambda$  
vector defining of an affine timing function

$\mu$  
coefficient of an affine timing function

$a$  
generic allocation function

$\sigma$  
matrix defining of an affine allocation function

$[t, a]$  
space-time mapping determined by $t$ and $a$

$G^{t, a}$  
signal flow graph under $t$ and $a$

[miscellaneous]

$tr$  
translation in $\mathbb{R}^n$

$\text{ren}$  
variable renaming

$S^{\text{tr, ren}}$  
translated image of $S$

$\text{in}$  
generic input

$R, R_1, \ldots, S, S_1, \ldots$  
routing variables

$a, \beta, \gamma, \ldots$  
control variables

$G_1, \ldots, G_m$  
integer-valued variables
For simplicity, we usually represent an $n$-dimensional vector as an $n$-tuple, that is $x = (x_1, \ldots, x_n)$. The only exception to this convention is the case of matrix expressions, in which vectors are represented as column vectors.

As we restrict ourselves to lattice spaces, with some abuse of notation, we represent vector spaces, such as $\text{lin}(D)$ or $\text{null}(M)$, as lattice spaces.
Appendix B

Proofs of Basic Results

In this appendix we provide a formal proof of some of the basic results discussed in Chapter 2. For some of these results, a formal statement was given in Chapter 2. In these cases, we have maintained the same reference number, and indicate, in the right margin, the page at which the formal statement was given.

B.1 Timing and Allocation Functions

**Proposition B.1.1** [Compatibility] Let \( t \) be an affine timing function such that \( t(z) = \lambda \cdot z + \mu \), and \( a \) a linear allocation function with projection vector \( u \). Then \( t \) and \( a \) are compatible if and only if \( \lambda \cdot u \neq 0 \).

**Proof:** We first prove that if \( t \) and \( a \) are compatible then \( \lambda \cdot u \neq 0 \). Assume \( \lambda \cdot u = 0 \).

Then \( u \in \text{null}(\lambda) \). Therefore there exist \( z, z' \in [\lambda : c] \), for some \( c \in Z \), such that \( z' = z + mu \), with \( m \neq 0 \), and \( t(z) = t(z') \). Then:

\[
a(z') = \sigma \cdot z' = \sigma \cdot (z + mu) = \\
= \sigma \cdot z + m\sigma \cdot u = \sigma \cdot z = a(z).
\]

Hence, \( t \) and \( a \) are not compatible.

We now prove that if \( \lambda \cdot u \neq 0 \) then \( t \) and \( a \) are compatible. Assume \( t \) and \( a \) non compatible. Then there exist \( z, z' \) such that \( z \neq z' \), \( t(z) = t(z') \) and \( a(z) = a(z') \). However, if \( a(z) = a(z') \), then there exists \( m \neq 0 \) such that \( z' = z + mu \), as \( u \) is the projection vector. Therefore:

\[
t(z') = \lambda \cdot z' + \mu = \lambda \cdot (z + mu) + \mu = \\
= \lambda \cdot z + m\lambda \cdot u + \mu = t(z) + m\lambda \cdot u.
\]
and \( t(z') = t(z) \) only if \( \lambda \cdot u = 0 \).

\[ \square \text{B.1.1} \]

\section*{B.2 Condition for Uniformity}

**Proposition B.2.1** [Condition for Uniformity] Let \( \mathcal{D} = (\mathcal{D}, \mathcal{U}, \mathcal{V}, \mathcal{I}) \) be an affine data dependence, with \( \mathcal{I}(z) = A \cdot z + b \), for all \( z \in \mathcal{D} \). Then \( \mathcal{D} \) is uniform if and only if \( \text{lin}(\mathcal{D}) \subseteq \text{null}(I_n - A) \).

**Proof:** By definition, the affine closure of \( D \) is:

\[
\text{aff}(D) = \left\{ \sum \alpha_i z_i \mid z_i \in D, \sum \alpha_i = 1 \right\}
\]

and given \( d \in D \), the direction of \( D \) is:

\[
\text{lin}(D) = \text{aff}(D) - d = \{ z - d \mid z \in D \}.
\]

From the above definitions, it follows that \( \forall z \in D, z - d \in \text{lin}(D) \).

Let us prove that \( \text{lin}(D) \subseteq \text{null}(I_n - A) \) implies that there exists \( c \in \mathbb{Z}^n \) such that \( \forall z \in D, \Theta_I(z) = c \). For all \( z \in D \),

\[
\Theta_I(z) = (I_n - A) \cdot z - b = (I_n - A) \cdot (z - d + d) - b = (I_n - A) \cdot (z - d) + (I_n - A) \cdot d - b.
\]

As \( z - d \in \text{lin}(D) \) and, by assumption, \( \text{lin}(D) \subseteq \text{null}(I_n - A) \), then \( (I_n - A) \cdot (z - d) = 0 \) and the expression above reduces to

\[
\Theta_I(z) = (I_n - A) \cdot d - b,
\]

where \( (I_n - A) \cdot d - b \) is a constant vector in \( \mathbb{Z}^n \), independent from \( z \). In particular, when \( A = I_n \), then \( \Theta_I(z) = -b \).

We now prove that the existence of \( c \in \mathbb{Z}^n \) such that \( \forall z \in D, \Theta_I(z) = c \) implies \( \text{lin}(D) \subseteq \text{null}(I_n - A) \). As \( \forall z \in D, \Theta_I(z) = c \) then \( \forall z \in D, (I_n - A) \cdot z = b + c \), and, in particular, \( (I_n - A) \cdot d = b + c \). The (matrix) expression:

\[
(I_n - A) \cdot z = b + c
\]

defines a system of non-homogeneous linear equations whose solution space \( \text{null}(I_n - A) + (b + c) \) contains \( D \). Also \( \text{aff}(D) \) is contained in this space, as any affine combination of elements of \( D \) is a solution to the system. In fact, let \( z_i \in D \), for
i = 1 \ldots, m, and let $\sum a_i z_i$ be an affine combination of the points. with $\sum a_i = 1$.

Then:

$$(I_n - A) \cdot (\sum a_i z_i) = \sum a_i (I_n - A) \cdot z_i = \sum a_i (b + c) = b + c.$$ 

i.e., the affine combination is a solution to the system. By definition of $\text{lin}(D)$, for all $z \in \text{lin}(D)$, there exist $z' \in \text{aff}(D)$ and $d \in D$ such that $z = z' - d$. Then:

$$(I_n - A) \cdot z = (I_n - A) \cdot (z' + d) = 0$$

i.e., $z \in \text{null}(I_n - A)$. It follows that $\text{lin}(D) \subseteq \text{null}(I_n - A)$.

\[ \text{B.2.1} \]

**Corollary B.2.2** Let $\mathcal{D} = (D, U, V, I)$ be an affine data dependence. If $I$ is uniform then $\mathcal{D}$ is uniform. \[ \text{B.2.2} \]

**Corollary B.2.3** Let $\mathcal{D} = (D, U, V, I)$ be an affine data dependence with a non-uniform index mapping $I$. If $\mathcal{D}$ is uniform then $I$ can be replaced by a uniform index mapping $I'$.

**PROOF:** $I'$ is the index mapping $I'(z) = z - c$, where $c$ is the constant vector of Proposition B.2.1.

\[ \text{B.2.3} \]

**B.3 Affine Timing Function for Affine Data Dependencies**

Linear programming problems for optimal affine timing functions have been considered by several authors in the literature [RaFu87, RaFu89, QuVa89]. The main result of this section (Theorem 2.2.11) is based on the formulation in [QuVa89].

**Lemma B.3.1 [Valid Affine Timing Function]** Let $\mathcal{D} = (D, U, V, I)$ be an affine data dependence, $\Theta_I$ its dependence mapping and $\Omega_I = \text{range}_{\Theta_I}(D)$. Let $t$ be an affine timing function, such that for all $z \in \mathbb{Z}^n$, $t(z) = \lambda \cdot z + \mu$, with $\lambda \in \mathbb{Z}^n$ and with $\mu \in \mathbb{Z}$. $t$ is a valid timing function for $\mathcal{D}$ if and only if:

i) for all $v \in \text{vert}(\Omega_I)$, $\lambda \cdot v > 0$;

ii) for all $r \in \text{ray}(\Omega_I)$, $\lambda \cdot r \geq 0$. 

\[ \text{B.3.1} \]
**APPENDIX B. PROOFS OF BASIC RESULTS**

**Proof:** According to Definition 2.1.12, $t$ is a valid timing function for $\mathcal{D}_D$ if and only if for all $z \in D$, $t(z) > t(I(z))$. By expanding $t$ to its affine form we obtain:

$$\lambda \cdot z + \mu > \lambda \cdot I(z) + \mu$$

$$\lambda \cdot (z - I(z)) > 0$$

$$\lambda \cdot \Theta_T(z) > 0$$

i.e., $t$ is a valid timing function for $\mathcal{D}_D$ if and only if for all $z \in D$, $\lambda \cdot \Theta_T(z) > 0$.

First we prove that conditions i) and ii) implies $\lambda \cdot \Theta_T(z) > 0$, for all $z \in D$. As $\Omega_T$ is a convex polyhedron, it is finitely generated by a set of points and directions in $\mathbb{Z}^n$. Let $\text{vert}(\Omega_T) = \{v_1, \ldots, v_m\}$ and $\text{ray}(\Omega_T) = \{r_1, \ldots, r_p\}$. Then, for all $z \in D$, $\Theta_T(z)$ can be expressed as:

$$\Theta_T(z) = \sum_i a_i v_i + \sum_j b_j r_j,$$

with $a_i, b_j \geq 0$ and $\sum_i a_i = 1$. As $\Theta_T(z) \neq 0$ (otherwise the data dependence graph is cyclic), from i), ii) and the conditions on $a_i$ and $b_j$ it follows that:

$$\lambda \cdot \Theta_T(z) = \sum_i a_i \lambda \cdot v_i + \sum_j b_j \lambda \cdot r_j > 0.$$ 

We now prove that $\lambda \cdot \Theta_T(z) > 0$, for all $z \in D$, implies conditions i) and ii).

Condition i) is straightforward. As any vertex $v$ of $\Omega_T$ is in particular an element of the set, there exists $z \in D$ such that $v = \Theta_T(z)$. Therefore $\lambda \cdot v > 0$. Condition ii) can be proved by noticing that if $r$ is an infinite direction of $\Omega_T$ then it is the image of an infinite direction $r'$ of $D$ under the linear part $\mathcal{L}_{\Theta_T}$ of $\Theta_T$. Suppose that for all $z$, $\Theta_T(z) = \mathcal{L}_{\Theta_T}(z) + c$, for some $c \in \mathbb{Z}^n$. Assume that $\lambda \cdot r < 0$ and consider $z + ar' \in D$. Then:

$$\lambda \cdot \Theta_T(z + ar') =$$

$$= \lambda \cdot (\mathcal{L}_{\Theta_T}(z + ar') + c) =$$

$$= \lambda \cdot (\mathcal{L}_{\Theta_T}(z) + a\mathcal{L}_{\Theta_T}(r') + c) =$$

$$= \lambda \cdot (\Theta_T(z) + ar) =$$

$$= \lambda \cdot \Theta_T(z) + a\lambda \cdot r,$$

which is negative for $a$ sufficiently large, as $\lambda \cdot \Theta_T(z) > 0$. As this violates the validity of $t$, then the assumption $\lambda \cdot r < 0$ is false.  

\[ \text{B.3.1} \]
Corollary B.3.2 Let $\mathcal{DD} = (D, U, V, I)$ be a uniform data dependence, with index mapping $I(z) = z + b$, for all $z \in \mathbb{Z}^n$, and $b \in \mathbb{Z}^n$. Let $t$ be an affine timing function, such that for all $z \in \mathbb{Z}^n$, $t(z) = \lambda \cdot z + \mu$, with $\lambda \in \mathbb{Z}^n$ and $\mu \in \mathbb{Z}$. $t$ is a valid timing function for $\mathcal{DD}$ if and only if $\lambda \cdot b < 0$. \hfill\textit{B.3.2}

Lemma B.3.3 \textit{[Finite and Bounded Affine Timing Function]} Let $\mathcal{DD} = (D, U, V, I)$ be an affine data dependence and $t$ a valid affine timing function for $\mathcal{DD}$. Then:

i) $t$ is finite if and only if, for all $r \in \text{ray}(D)$, $\lambda \cdot r \neq 0$;

ii) $t$ is bounded (below) if and only if, for all $r \in \text{ray}(D)$, $\lambda \cdot r \geq 0$.

Proof: i) As $D$ is a convex polyhedral set, $D$ contains unbounded sets of the form $\{z + ar | a \geq 0\}$, for $z \in D$ and $r \in \text{ray}(D)$. Therefore, $t$ is not finite if and only if, for any such set, $t(z) = t(z + ar)$ for all $a > 0$. However, by expanding the definition of $t$,

$$t(z) = \lambda \cdot z + \mu$$

$$t(z + ar) = \lambda \cdot (z + ar) + \mu = \lambda \cdot z + \mu + a \lambda \cdot r$$

Therefore, $t(z) = t(z + ar)$ for all $a > 0$ if and only if $\lambda \cdot r = 0$.

ii) $t$ is not bounded below if and only if, given a set $\{z + ar | a \geq 0\}$, for $z \in D$ and $r \in \text{ray}(D)$, there exists an infinitely decreasing chain $t(z) > t(z + r) > \ldots > t(z + ar) > \ldots$. Such a chain exists if and only if $\lambda \cdot r < 0$. In fact, for all $a \geq 0$,

$$t(z + ar) > t(z + (a + 1)r)$$

if and only if

$$\lambda \cdot (z + ar) + \mu > \lambda \cdot (z + (a + 1)r) + \mu$$

$$\lambda \cdot z + a \lambda \cdot r + \mu > \lambda \cdot z + (a + 1)\lambda \cdot r + \mu$$

$$a \lambda \cdot r > (a + 1)\lambda \cdot r$$

i.e., if and only if $\lambda \cdot r < 0$. \hfill\textit{B.3.3}

Corollary B.3.4 Let $\mathcal{DD} = (D, U, V, I)$ be an affine data dependence and $t$ a valid affine timing function for $\mathcal{DD}$. Then $t$ is finite and bounded if and only if, for all $r \in \text{ray}(D)$, $\lambda \cdot r > 0$. \hfill\textit{B.3.4}
Corollary B.3.5  Let $\mathcal{DD} = (D, U, V, I)$ be an affine data dependence and $t$ a valid affine timing function for $\mathcal{DD}$. Then:

i) for all $r \in \text{ray}(D)$ and $z, z' \in D$ such that $z' = z + ar$, with $a \in \mathbb{Z}$, $a \neq 0$, if $t$ is finite then $t(z) \neq t(z')$;

ii) for all $r \in \text{ray}(D)$ and $z, z' \in D$ such that $z' = z + ar$, with $a \in \mathbb{Z}$, $a > 0$, if $t$ is bounded below then $t(z) < t(z')$.

Theorem 2.2.11  Let $\mathcal{DD} = (D, U, V, I)$ be an affine data dependence, $\Theta_I$ its dependence mapping and $\Omega_I = \text{range}_{\Theta_I}(D)$. Let $t$ be an affine timing function, such that for all $z \in \mathbb{Z}^n$, $t(z) = \lambda \cdot z + \mu$, with $\lambda \in \mathbb{Z}^n$ and with $\mu \in \mathbb{Z}$. Then $t$ is a non-negative, finite, bounded valid timing function for $\mathcal{DD}$ if and only if:

i) for all $v \in \text{vert}(\Omega_I)$, $\lambda \cdot v > 0$;

ii) for all $r \in \text{ray}(\Omega_I)$, $\lambda \cdot r \geq 0$;

iii) for all $v \in \text{vert}(D)$, $\lambda \cdot v + \mu \geq 0$;

iv) for all $r \in \text{ray}(D)$, $\lambda \cdot r > 0$.

Proof: First we prove that conditions i)-iv) implies that $t$ is a non-negative, finite, bounded, valid timing function for $\mathcal{DD}$. Condition i) and ii) implies that $t$ is valid (see Lemma B.3.1). Conditions iii) and iv) implies that $t$ is non-negative. In fact, let $\text{vert}(D) = \{v_1, \ldots, v_m\}$ and $\text{ray}(D) = \{r_1, \ldots, r_p\}$. Then, for all $z \in D$, $z$ can be expressed as:

$$z = \sum_i a_i v_i + \sum_j b_j r_j.$$  

with $a_i, b_j \geq 0$ and $\sum_i a_i = 1$. Because of conditions iii) and iv) $\sum_i a_i (\lambda \cdot v_i + \mu) + \sum_j b_j \lambda \cdot r_j \geq 0$. Therefore:

$$\sum_i a_i (\lambda \cdot v_i + \mu) + \sum_j b_j \lambda \cdot r_j =$$

$$= \sum_i a_i \lambda \cdot v_i + \sum_i a_i \mu + \sum_j b_j \lambda \cdot r_j =$$

$$= \lambda \cdot (\sum_i a_i v_i + \sum_j b_j r_j) + \mu =$$

$$= \lambda \cdot z + \mu = t(z) \geq 0.$$
Finally, condition iv) implies that $t$ is finite and bounded (see Lemma B.3.3).

Now we prove that if $t$ is a non-negative, finite, bounded, valid timing function for $\mathcal{DD}$ then conditions i)-iv) hold. $t$ valid implies conditions i) and ii) (see Lemma B.3.1). $t$ non-negative implies condition iii). In fact, as $v \in \text{vert}(D)$, in particular, $v$ is an element of $D$; then $t(v) \geq 0$, i.e., $\lambda \cdot v + \mu \geq 0$. Finally, $t$ finite and bounded implies condition iv) (see Lemma B.3.3).

**Corollary 2.2.13** Let $\mathcal{DD} = (D, U, V, \mathcal{I})$ be a uniform data dependence, with index mapping $\mathcal{I}(z) = z + b$, for all $z \in \mathbb{Z}^n$, and $b \in \mathbb{Z}$. Let $t$ be an affine timing function, such that for all $z \in \mathbb{Z}^n$, $t(z) = \lambda \cdot z + \mu$, with $\lambda \in \mathbb{Z}^n$ and $\mu \in \mathbb{Z}$. Then $t$ is a non-negative, finite, bounded valid timing function for $\mathcal{DD}$ if and only if:

i) $\lambda \cdot b < 0$;

ii) for all $v \in \text{vert}(D)$: $\lambda \cdot v + \mu \geq 0$;

iii) for all $r \in \text{ray}(D)$: $\lambda \cdot r > 0$.

**B.4 Dependence Cone and Affine Timing Function**

**Proposition 2.2.15** Let $\mathcal{DD} = (D, U, V, \mathcal{I})$ be a data dependence and $\Theta^*_T$ its dependence cone. If $\Theta^*_T$ is pointed then there exists a valid affine timing function for $\mathcal{DD}$.

**Proof:** The cone $\Theta^*_T$ is, by definition, a convex polyhedral set. If $\Theta^*_T$ is pointed then it contains no lines, and is finitely generated by its extremal rays. Let $\{r_1, \ldots, r_m\}$ be the set of the extremal rays of $\Theta^*_T$. A valid affine timing function for $\mathcal{DD} = (D, U, V, \mathcal{I})$ is determined by any non-null $\lambda \in \mathbb{Z}^n$ such that $\lambda \cdot r_i > 0$ for all extremal rays $r_i$. At least one such $\lambda$ exists because of the separation theorem (see Appendix E).

By definition of $\Theta^*_T$, for all $z \in D$, $\Theta_T(z) \in \Theta^*_T$. Hence, for all $z \in D$, $\Theta_T(z)$ can be written as a positive combination of the extremal rays of $\Theta^*_T$, i.e., $\Theta_T(z) = \sum_i a_i r_i$, with $a_i \geq 0$ and not all $a_i = 0$. Therefore, for all $z \in D$,

\[
\lambda \cdot \Theta_T(z) = \lambda \cdot \left( \sum_i a_i r_i \right) = \sum_i a_i \lambda \cdot r_i > 0.
\]
Therefore, any timing function defined as \( t(z) = \lambda \cdot z + \mu \) for some \( \mu \in \mathbb{Z} \) defines a valid timing function for \( \mathcal{D} \mathcal{D} \).

\[ \blacksquare \ 2.2.15 \]

## B.5 Pointed Cone with Unimodular Generators

The following result is due to Quinton and Van Dongen [QuVa89].

**Proposition 3.2.1 [QuVa89]** Let \( C \) be a pointed polyhedral convex cone of full dimension in \( \mathbb{Q}^n \). There exists a pointed polyhedral convex cone \( C' \) such that: \( C' \) contains \( C \) and its extremal rays constitute a unimodular basis of \( \mathbb{Z}^n \).

\[ \blacksquare \ 3.2.1 \]

**Proof:** The proof is based on the two following properties of convex polyhedral cones:

- if \( C \) and \( C' \) are convex polyhedral cones such that \( C \subseteq C' \), then their dual cones \( \hat{C} \) and \( \hat{C}' \) are such that \( \hat{C}' \subseteq \hat{C} \);

- if \( C \) is an \( n \)-dimensional cone in \( \mathbb{Q}^n \), with \( n \) extremal rays \( r_1, \ldots, r_n \) forming the columns of a matrix \( Q \), then if \( \hat{C} \) is an \( n \)-dimensional cone in \( \hat{Q}^n \), with \( n \) extremal rays forming the columns of the matrix \( \hat{Q} = -(Q^{-1})' \).

Suppose \( C \) has \( m \) extremal rays, with \( m \geq n \) (\( m \) cannot be less than \( n \) as \( C \) is assumed of full dimension in \( \mathbb{Q}^n \)). Let \( \hat{C} \) be its dual cone and let \( \hat{R} \) be the matrix having as columns the \( m \) extremal rays of \( \hat{C} \). Choose a sub-cone \( \hat{C}' \) of \( \hat{C} \) such that \( \hat{C}' \) has exactly \( n \) extremal rays which form a unimodular basis. Such rays are the columns of any \( n \times n \) unimodular matrix \( \hat{R}' \), which satisfies the matrix equation:

\[
\hat{R} \cdot P = \hat{R}'
\]

with \( P \) a non-negative \( m \times n \) rational matrix. Then the dual cone \( C' \) of \( \hat{C}' \) is the cone we are looking for, with extremal rays the columns of the matrix \( -(\hat{R}^{-1})' \).

\[ \blacksquare \ 3.2.1 \]
Appendix C

Case Studies

This appendix contains programs for the execution of the specifications of the case studies in Chapter 5. The programs are expressed in Mathematica. (Mathematica is a trade mark of Wolfram Research Inc. For a description of the syntax of Mathematica expressions and predefined functions, see [Wol88].) Input and output codes for the various case studies were produced with Mathematica and subsequently edited for inclusion in this thesis.

C.1 Notation and Conventions

In general, we have tried to be consistent with the notation used in Chapter 5. There are, however, a few exceptions. In particular, as C and D are protected symbols in Mathematica, variables with such names in Chapter 5 have been named CC and DD, respectively, in this appendix. Also, n has been used instead of the reserved symbol N.

In the following programs, domains are defined as systems of inequalities and represented in their matrix form, i.e., if a domain is defined by the system of inequalities $\pi \cdot z \geq \theta$, for a matrix $\pi$ and a vector $\theta$, the pair $(\pi, \theta)$ is provided to describe it.

Different from the specifications in Chapter 5, in the following programs we need to evaluate explicitly whether a point belongs to a certain domain before being able to evaluate any variable at that point. To this end we have defined a predicate in, which checks whether a point point belongs to a domain defined by a pair (pi, theta). The predicate computes the matrix expression (pi . point - theta) and checks whether all the entries of the resulting vector are non-negative. The predicate in is defined as follows:

\[
in[\text{pi_-}, \text{theta_-}, \text{point_-}] := \text{in[pi, theta, point]} = \\
\quad \text{Apply[And, Map[NonNegative, pi \cdot point - theta]]}
\]
where the predefined function `Map` applies the boolean predefined function `NonNegative` to each entry of `(pI.point-theta)`, and `Apply` realises the logical `And` of the results.

### C.2 Cyclic Reduction

The following Mathematica program refers to the case study presented in Section 5.1. The reduction phase of the algorithm is fully given, corresponding to the system of uniform equations obtained by applying regularisation techniques (as explained in Section 5.1.5) to all the variables of the specification in Section 5.1.3. In the code, we have included some comments which should help the reader to compare the program and the specification.

By recalling our discussion in Section 5.1.5, only two basic types of non-uniform data dependencies, which we called $D_1$ and $D_2$, were present in the original specification. Their decomposition introduced variables $P$ and $Q$ as well as the new non-uniform data dependencies $D_4$ and $D_5$. In turn, their uniformisation produced the routing variables $R^1$, $R^2$ and $S^1$, $S^2$, respectively, and the control variables $\alpha$, $\beta$, $\gamma$, and $\xi$, $\phi$, $\psi$, respectively. In the following program, all necessary decompositions and uniformisations relative to all the variables of the specification have been included. In order to maintain the relation with the discussion in Section 5.1.5 we have adopted the convention that the above decomposition and routing variables are prefixed with the name of the variable to which they refer. For instance, variables relative to $A$ have been named as $AP$, $AQ$, $AR_1$, etc. Similarly for $B$, $CC$ and $DD$. Note that only two sets of control variables ($\alpha$, $\beta$, $\gamma$, and $\xi$, $\phi$, $\psi$, respectively) are used for the routing of all the variables of the algorithm.

The correspondence between the identifiers in the following program and those in the specification should be straightforward, with the exception, perhaps, of `bargprime`, which corresponds to $g'$ and `bargdbprime`, which corresponds to $g''$.

#### C.2.1 Program

(* Cyclic Reduction - Reduction Phase *)

(* constants *)

\[
n = 2^r - 1
\]

\[
bargprime = \text{Floor}[(2^{(r-2)-1})/2]
\]

\[
bargdbprime = \text{Floor}[(2^{(r-1)-1})/2]
\]
(* domains *)

(* D0 *)

\[
\pi_0 = \begin{Bmatrix}
1, 0, 0, 0, -1, 0, 0, 0, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_0 = \begin{Bmatrix}
1, -2^{-r+1}, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]

(* D1 *)

\[
\pi_1 = \begin{Bmatrix}
1, 0, 0, 0, -1, 0, 0, 0, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_1 = \begin{Bmatrix}
3, -2^{-r+1}, 1, -r+1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]

(* D2 *)

\[
\pi_2 = \begin{Bmatrix}
1, 0, 0, 0, -1, 0, 0, 0, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_2 = \begin{Bmatrix}
2, -2^{-r+2}, 0, -r+2, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]

(* D21 *)

\[
\pi_{21} = \begin{Bmatrix}
1, 0, 1, -1, 0, -1, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_{21} = \begin{Bmatrix}
1, -2^{-r+2}, 0, -r+2, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]

(* D2(1,1) *)

\[
\pi_{211} = \begin{Bmatrix}
1, 0, 1, -1, 0, -1, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_{211} = \begin{Bmatrix}
1, -2^{-r+2}, 0, -r+2, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]

(* D2(1,2) *)

\[
\pi_{212} = \begin{Bmatrix}
1, 0, 1, -1, 0, -1, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_{212} = \begin{Bmatrix}
1, -(2^{-r})+2, 0, -r+2, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]

(* D22 *)

\[
\pi_{22} = \begin{Bmatrix}
1, 0, -1, -1, 0, -1, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_{22} = \begin{Bmatrix}
-2 \text{ bargprime}, -2^{-r}+2, 0, -r+2, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]

(* D2(2,1) *)

\[
\pi_{221} = \begin{Bmatrix}
1, 0, -1, -1, 0, -1, 0, 1, 0, 0, -1, 0, 0, 0, 1, 0, 0, -1
\end{Bmatrix}
\]

\[
\theta_{221} = \begin{Bmatrix}
-2 \text{ bargprime}, -2^{-r}+2, 0, -r+2, 1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0
\end{Bmatrix}
\]


\[ \pi_{222} = \begin{bmatrix} 1 & 0 & -1 \\ -1 & 0 & -1 \\ 0 & 1 & 0 \end{bmatrix} \]
\[ \theta_{222} = \{-2 \text{bargprime}, -2^{-r+2}, 0, -r+2, 0, 0\} \]

\[ \pi_{23} = \begin{bmatrix} 1 & 0 & 1 \\ -1 & 0 & -1 \\ 0 & 1 & 0 \end{bmatrix} \]
\[ \theta_{23} = \{1, -2^{-r+2}, 0, -r+2, 0, -\text{bargdbprime}\} \]

\[ \pi_{231} = \begin{bmatrix} 1 & 0 & 1 \\ -1 & 0 & -1 \\ 0 & 1 & 0 \end{bmatrix} \]
\[ \theta_{231} = \{1, -2^{-r+2}, 0, -r+2, 0, -\text{bargdbprime}+1\} \]

\[ \pi_{232} = \begin{bmatrix} 1 & 0 & 1 \\ -1 & 0 & -1 \\ 0 & 1 & 0 \end{bmatrix} \]
\[ \theta_{232} = \{1, -(2^{-r})+2, 0, -r+2, \text{bargprime}, -\text{bargprime}\} \]

\[ \pi_{24} = \begin{bmatrix} 1 & 0 & -1 \\ -1 & 0 & -1 \\ 0 & 1 & 0 \end{bmatrix} \]
\[ \theta_{24} = \{-2 \text{bargdbprime}, -2^{-r+2}, 0, -r+2, 0, -\text{bargdbprime}\} \]

\[ \pi_{241} = \begin{bmatrix} 1 & 0 & -1 \\ -1 & 0 & -1 \\ 0 & 1 & 0 \end{bmatrix} \]
\[ \theta_{241} = \{-2 \text{bargdbprime}, -2^{-r+2}, 0, -r+2, 1, -\text{bargdbprime}\} \]

\[ \pi_{242} = \begin{bmatrix} 1 & 0 & -1 \\ -1 & 0 & -1 \\ 0 & 1 & 0 \end{bmatrix} \]
\[ \theta_{242} = \{-2 \text{bargdbprime}, -2^{-r+2}, 0, -r+2, 0, 0\} \]

\[ \text{in}[\pi_-, \theta_-, \text{point}_-] := \text{in}[\pi, \theta, \text{point}] = \]
\[ \text{Apply}[\text{And}, \text{Map}[\text{NonNegative}, \pi . \text{point} - \theta ] ] \]

\[ \text{gprime}[i_-, j_, k_-] := 2^{-j} \]
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\[ \text{gdbprime}[i_-, j_-, k_] := 2^{(j+1)} \]

(* applied functions *)
\[ \text{inA}[i_-, j_-, k_] := a[i] \]
\[ \text{inB}[i_-, j_-, k_] := b[i] \]
\[ \text{inCC}[i_-, j_-, k_] := c[i] \]
\[ \text{inDD}[i_-, j_-, k_] := d[i] \]
\[ f[i_-, j_-, k_] := \text{If}[ \text{Mod}((i-2^j+1)/(2^j-1)), 2] = 0 \]
\[ \quad \& \& ((i-2^j+1)/(2^j-1) >= 2) \&\& ((i-2^j+1)/(2^j-1) <= 2^r-2), 1, 0] \]
\[ f1[t_-, a_-, b_-, c_] := \text{If}[ t == 1, -a*b/c, \text{undef}1] \]
\[ f2[t_-, a_-, b_-, c_-, d_-, e_-, f_-, g_] := \]
\[ \text{If}[ t == 1, a-(b*c/d)-(e*f/g), \text{undef}2] \]
\[ \text{inAlpha}[i_-, j_-, k_] := \text{Floor}[(\text{gprime}[i+bargprime, j, k-bargprime]-1)/2] \]
\[ \text{inBeta}[i_-, j_-, k_] := \text{Mod}((\text{gprime}[i+bargprime, j, k-bargprime]-1), 2] \]
\[ \text{inGamma}[i_-, j_-, k_] := \text{bargprime} \]
\[ \text{inXi}[i_-, j_-, k_] := \text{Floor}[(\text{gdbprime}[i+bargdbprime, j, k-bargdbprime]-1)/2] \]
\[ \text{inPhi}[i_-, j_-, k_] := \text{Mod}((\text{gdbprime}[i+bargdbprime, j, k-bargdbprime]-1), 2] \]
\[ \text{id}[a_] := a \]
\[ \text{dec}[a_] := a-1 \]
\[ f3[a_-, b_-, c_-, d1_-, d2_-, d3_] := \]
\[ \text{If}[a != c, d1, \text{If}[b == 0, d2, d3 ] ] \]

(* control variable T *)
\[ T[i_-, j_-, k_] := T[i, j, k] = \]
\[ \text{If}[ \text{in}[\pi 1, \theta 1, \{i, j, k\}], f[i, j, k], \text{undefT} ] \]

(* variable A *)
\[ A[i_-, j_-, k_] := A[i, j, k] = \]
\[ \text{If}[ \text{in}[\pi 0, \theta 0, \{i, j, k\}], \text{inA}[i, j, k], \]
\[ \quad \text{If}[ \text{in}[\pi 1, \theta 1, \{i, j, k\}], \]
\[ \quad \quad f1[T[i, j, k], \text{AP}[i-1, j-1, k], \text{AQ}[i-1, j-1, k], \]
\[ \quad \quad \text{BQ}[i-1, j-1, k]], \text{undefA} ] ] \]
(* routing of A - dependence type DD4 *)
(* variable AP *)
AP[i_, j_, k_] := AP[i, j, k] =
    If[ in[pi2, theta2, {i, j, k}], id[AR1[i, j, k]], undefAP ]

(* variable AR1 *)
AR1[i_, j_, k_] := AR1[i, j, k] =
    If[ in[pi21, theta21, {i, j, k}],
        f3[alpha[i, j, k], beta[i, j, k], gamma[i, j, k],
            AR1[i-1, j, k+1], AR2[i, j, k], AR2[i-1, j, k]],
        undefAR1 ]

(* variable AR2 *)
AR2[i_, j_, k_] := AR2[i, j, k] =
    If[ in[pi221, theta221, {i, j, k}], id[AR2[i-l, j, k]],
        If[ in[pi222, theta222, {i, j, k}], id[A[i, j, k]],
            undefAR2 ] ]

(* routing of A - dependence type DD5 *)
(* variable AQ *)
AQ[i_, j_, k_] := AQ[i, j, k] =
    If[ in[pi2, theta2, {i, j, k}], id[AS1[i, j, k]], undefAQ ]

(* variable AS1 *)
AS1[i_, j_, k_] := AS1[i, j, k] =
    If[ in[pi23, theta23, {i, j, k}],
        f3[xi[i, j, k], phi[i, j, k], psi[i, j, k],
            AS1[i-l, j, k+1], AS2[i, j, k], AS2[i-1, j, k]],
        undefAS1 ]

(* variable AS2 *)
AS2[i_, j_, k_] := AS2[i, j, k] =
    If[ in[pi241, theta241, {i, j, k}], id[AS2[i-l, j, k-1]],
        If[ in[pi242, theta242, {i, j, k}], id[A[i, j, k]],
            undefAS2 ] ]

(* variable B *)
B[i_, j_, k_] := B[i, j, k] =
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If[ in[pi0, theta0, {i, j, k}], inB[i, j, k],
    If[ in[pi1, theta1, {i, j, k}],
        f2[T[i, j, k], BP[i-1, j-1, k], AP[i-1, j-1, k],
        CCQ[i-1, j-1, k], BQ[i-1, j-1, k], CCP[i-1, j-1, k],
        AQ[i-1, j-1, k], B[i, j-1, k], undefB ] ]

(* routing of B - dependence type DD4 *)

(* variable BP *)

BP[i, j, k] := BP[i, j, k] =
    If[ in[pi2, theta2, {i, j, k}], id[BR1[i, j, k]], undefBP ]

(* variable BR1 *)

BR1[i, j, k] := BR1[i, j, k] =
    If[ in[pi21, theta21, {i, j, k}],
        f3[alpha[i, j, k], beta[i, j, k], gamma[i, j, k],
        BR1[i-1, j, k+1], BR2[i, j, k], BR2[i-1, j, k]], undefBR1 ]

(* variable BR2 *)

BR2[i, j, k] := BR2[i, j, k] =
    If[ in[pi221, theta221, {i, j, k}], id[BR2[i-1, j, k-1]],
        If[ in[pi222, theta222, {i, j, k}], id[B[i, j, k]], undefBR2 ] ]

(* routing of B - dependence type DD5 *)

(* variable BQ *)

BQ[i, j, k] := BQ[i, j, k] =
    If[ in[pi2, theta2, {i, j, k}], id[BS1[i, j, k]], undefBQ ]

(* variable BS1 *)

BS1[i, j, k] := BS1[i, j, k] =
    If[ in[pi23, theta23, {i, j, k}],
        f3[xi[i, j, k], phi[i, j, k], psi[i, j, k],
        BS1[i-1, j, k+1], BS2[i, j, k], BS2[i-1, j, k]], undefBS1 ]
(* variable BS2 *)
BS2[i_, j_, k_] := BS2[i, j, k] = 
    If[ in[pi241, theta241, {i, j, k}], id[BS2[i-1, j, k-1]],
        If[ in[pi242, theta242, {i, j, k}], id[B[i, j, k]], undefBS2 ] ]

(* variable CC *)
CC[i_, j_, k_] := CC[i, j, k] = 
    If[ in[pi0, theta0, {i, j, k}], inCC[i, j, k],
        If[ in[pi1, theta1, {i, j, k}],
            f1[T[i, j, k], CCP[i-1, j-1, k], CC[i, j-1, k],
                B[i, j-1, k], undefCC ] ]

(* routing of CC - dependence type DD4 *)
(* variable CCP *)
CCP[i_, j_, k_] := CCP[i, j, k] = 
    If[ in[pi2, theta2, {i, j, k}], id[CCR1[i, j, k]], undefCCP ]

(* variable CCR1 *)
CCR1[i_, j_, k_] := CCR1[i, j, k] = 
    If[ in[pi21, theta21, {i, j, k}],
        f3[alpha[i, j, k], beta[i, j, k], gamma[i, j, k],
            CCR1[i-1, j, k+1], CCR2[i, j, k],
                CCR2[i-1, j, k]], undefCCR1 ]

(* variable CCR2 *)
CCR2[i_, j_, k_] := CCR2[i, j, k] = 
    If[ in[pi221, theta221, {i, j, k}], id[CCR2[i-1, j, k-1]],
        If[ in[pi222, theta222, {i, j, k}],
            id[CC[i, j, k]], undefCCR2 ] ]

(* routing of CC - dependence type DDS *)
(* variable CCQ *)
CCQ[i_, j_, k_] := CCQ[i, j, k] = 
    If[ in[pi2, theta2, {i, j, k}],
        id[CCQ[i-1, j, k-1]], undefCCQ ]
id[CCS1[i, j, k]], undefCCQ ]

(* variable CCS1 *)
CCS1[i_, j_, k_] := CCS1[i, j, k] =  
  If[ in[pi23, theta23, {i, j, k}],
    f3[xi[i, j, k], phi[i, j, k], psi[i, j, k],
          CCS1[i-1, j, k+1], CCS2[i, j, k],
          CCS2[i-1, j, k]], undefCCS1 ]

(* variable CCS2 *)
CCS2[i_, j_, k_] := CCS2[i, j, k] = 
  If[ in[pi241, theta241, {i, j, k}], id[CCS2[i-1, j, k-1]],
    If[ in[pi242, theta242, {i, j, k}],
        id[CC[i, j, k]], undefCCS2 ]]

(* variable DD *)
DD[i_, j_, k_] := DD[i, j, k] = 
  If[ in[pi0, theta0, {i, j, k}], inDD[i, j, k],
    If[ in[pi1, theta1, {i, j, k}],
        f2[T[i, j, k], DDP[i-1, j-1, k], AP[i-1, j-1, k],
            DDQ[i-1, j-1, k], BQ[i-1, j-1, k], CCP[i-1, j-1, k],
                DD[i, j-1, k], B[i, j-1, k]], undefDD ]]

(* routing of DD - dependence type DD4 *)
(* variable DDP *)
DDP[i_, j_, k_] := DDP[i, j, k] = 
  If[ in[pi2, theta2, {i, j, k}], id[DDR1[i, j, k]], undefDDP ]

(* variable DDR1 *)
DDR1[i_, j_, k_] := DDR1[i, j, k] = 
  If[ in[pi21, theta21, {i, j, k}],
      f3[alpha[i, j, k], beta[i, j, k], gamma[i, j, k],
            DDR1[i-1, j, k+1], DDR2[i, j, k],
                DDR2[i-1, j, k]], undefDDR1 ]
(* variable DDR2 *)

\[
\text{DDR2}[i_, j_, k_] := \text{DDR2}[i, j, k] = \\
\begin{align*}
\text{If}\ [\ \text{in}[\pi221, \theta221, \{i, j, k\}], & \text{id}[\text{DDR2}[i-1, j, k-1]], \\
\text{If}\ [\ \text{in}[\pi222, \theta222, \{i, j, k\}], & \text{id}[\text{DD}[i, j, k]], \\
\text{id}[\text{DD}[i, j, k]], & \text{undefDDR2} ]
\end{align*}
\]

(* routing of DD - dependence type DDS5 *)

(* variable DDQ *)

\[
\text{DDQ}[i_, j_, k_] := \text{DDQ}[i, j, k] = \\
\begin{align*}
\text{If}\ [\ \text{in}[\pi2, \theta2, \{i, j, k\}], & \text{id}[\text{DDS1}[i, j, k]], \\
\text{ privileDDQ ]
\end{align*}
\]

(* variable DDS1 *)

\[
\text{DDS1}[i_, j_, k_] := \text{DDS1}[i, j, k] = \\
\begin{align*}
\text{If}\ [\ \text{in}[\pi23, \theta23, \{i, j, k\}], & f3[\xi[i, j, k], \phi[i, j, k], \psi[i, j, k], \\
\text{DDS1}[i-1, j, k+1], & \text{DDS2}[i, j, k], \\
\text{DDS2}[i-1, j, k]], & \text{undefDDSl ]
\end{align*}
\]

(* variable DDS2 *)

\[
\text{DDS2}[i_, j_, k_] := \text{DDS2}[i, j, k] = \\
\begin{align*}
\text{If}\ [\ \text{in}[\pi241, \theta241, \{i, j, k\}], & \text{id}[\text{DDS2}[i-1, j, k-1]], \\
\text{If}\ [\ \text{in}[\pi242, \theta242, \{i, j, k\}], & \text{id}[\text{DD}[i, j, k]], \\
\text{id}[\text{DD}[i, j, k]], & \text{undefDDS2 ]
\end{align*}
\]

(* control variables - dependence type DD4 *)

(* control variable alpha *)

\[
\text{alpha}[i_, j_, k_] := \text{alpha}[i, j, k] = \\
\begin{align*}
\text{If}\ [\ \text{in}[\pi211, \theta211, \{i, j, k\}], & \text{id}[\text{alpha}[i-1, j, k+1]], \\
\text{If}\ [\ \text{in}[\pi212, \theta212, \{i, j, k\}], & \text{id}[\text{DD}[i, j, k]], \\
in\text{alpha}[i, j, k], & \text{undefAlpha ]}
\end{align*}
\]

(* control variable beta *)

\[
\text{beta}[i_, j_, k_] := \text{beta}[i, j, k] = \\
\begin{align*}
\text{beta}[i, j, k]
\end{align*}
\]
If[ in[pi211, theta211, {i, j, k}], id[beta[i-1, j, k+1]],
   If[ in[pi212, theta212, {i, j, k}],
      inBeta[i, j, k], undefBeta ] ]

(* control variable gamma *)
gamma[i_, j_, k_]:= gamma[i, j, k] =
   If[ in[pi211, theta211, {i, j, k}], dec[gamma[i-l, j, k+l]],
      If[ in[pi212, theta212, {i, j, k}],
         inGamma[i, j, k], undefGamma ] ]

(* control variables - dependence type DD5 *)
(* control variable xi *)
xi[i_, j_, k_]:= xi[i, j, k] =
   If[ in[pi231, theta231, {i, j, k}], id[xi[i-1, j, k+1]],
      If[ in[pi232, theta232, {i, j, k}],
         inXi[i, j, k], undefXi ] ]

(* control variable phi *)
phi[i_, j_, k_]:= phi[i, j, k] =
   If[ in[pi231, theta231, {i, j, k}], id[phi[i-1, j, k+1]],
      If[ in[pi232, theta232, {i, j, k}],
         inPhi[i, j, k], undefPhi ] ]

(* control variable psi *)
psi[i_, j_, k_]:= psi[i, j, k] =
   If[ in[pi231, theta231, {i, j, k}], dec[psi[i-1, j, k+1]],
      If[ in[pi232, theta232, {i, j, k}],
         inPsi[i, j, k], undefPsi ] ]

C.2.2 Test

As the algorithm is static, we can obtain a symbolic computation (i.e., without providing actual inputs) of the program. We have tested the program for \(r = 3\). For brevity, we have restricted the output values below to the last computations on the domain \(D_1\) which occur
at the point \( (2^r - 1, r, 1) \) (point \( (7, 2, 0) \) in our example).

(* test *)
(* size parameter *)
r = 3

(* outputs *)
(* variable \( A \) *)
\[
A[7, 2, 0] = \\
-((a[1]*a[2]*a[3]*a[4])/ \\
\]

(* variable \( B \) *)
\[
B[7, 2, 0] = \\
(a[3]*a[4]*c[2]*c[3])/ \\
\]

(* variable \( CC \) *)
\[
CC[7, 2, 0] = \\
-((c[4]*c[5]*c[6]*c[7])/ \\
\]

(* variable \( DD \) *)
\[
DD[7, 2, 0] = \\
-((a[4]*d[3])/b[3]) + (a[3]*a[4]* \\
(\text{\(-((a[2]*d[1])/b[1]) + \text{\((d[2] - (c[2]*d[3])/b[3]))/ \\
(c[6]*d[7])/b[7])))\})
\]

C.3 M-to-1 Decimator

The program below corresponds to the specification of Section 5.2.3. The identifier \( Y_{\bar{}} \) is used to name \( \bar{Y} \).

C.3.1 Program

(* M-to-1 Decimator *)

(* domains *)

(* D1 *)
pi1 = {{1, 0}, {0, 1}, {0, -1}}
theta1 = {1, 0, 0}

(* D2 *)
pi2 = {{1, -1}, {0, 1}, {0, -1}}
theta2 = {0, 1, -n}

(* D3 *)
pi3 = {{1, 0}, {0, 1}, {0, -1}}
theta3 = {0, 0, 0}

(* D4 *)
pi4 = {{1, -1}, {-1, 1}, {0, 1}, {0, -1}}
theta4 = {-1, 1, 1, -n}

(* D5 *)
pi5 = {{1, -1}, {0, 1}, {0, -1}}
theta5 = {0, n+1, -n-1}

(* belongs to *)
in[pi_, theta_, point_] := in[pi, theta, point] =
    Apply[And, Map[NonNegative, pi . point - theta] ]
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(* applied functions *)
inX[i_, j_] := x[i-n]
inYbar[i_, j_] := 0
inH[i_, j_] := h[n-j]
inCC[i_, j_] := If[ Mod[i-n-1, M] == 0, 1, 0]
f1[a_, b_, c_] := a + b * c
f2[a_, b_] := If[b == 1, a, undef]
id[a_] := a

(* variable X *)
X[i_, j_] := X[i, j] =
   If[ in[pi1, theta1, {i, j}], inX[i, j],
      If[ in[pi2, theta2, {i, j}], X[i, j-1], undef ] ]

(* variable Ybar *)
Ybar[i_, j_] := Ybar[i, j] =
   If[ in[pi3, theta3, {i, j}], inYbar[i, j],
      If[ in[pi2, theta2, {i, j}],
         f1[Ybar[i-1, j-1], H[i-1,j], X[i, j-1]], undef ] ]

(* variable H *)
H[i_, j_] := H[i, j] =
   If[ in[pi4, theta4, {i, j}], inH[i, j],
      If[ in[pi2, theta2, {i, j}], id[H[i-1, j]], undef ] ]

(* control variable CC *)
CC[i_, j_] := Ctr[i, j] =
   If[ in[pi5, theta5, {i, j}], inCC[i, j], undef ]

(* variable Y *)
Y[i_, j_] := Y[i, j] =
   If[ in[pi5, theta5, {i, j}], f2[Ybar[i, j-1], CC[i, j]], undef ]
C.3.2 Test

We have executed the program for values of the size parameters $M = 3$ and $N = 5$. A few output values are provided for variables $Ybar$ and $Y$. Note how variable $Y$ decimates the values of $Ybar$ according to the parameter $M$.

(* test *)

(* size parameters *)

$M = 3$

$n = 5$

(* outputs *)

(* variable Ybar - for i=n,...,2n and j=n *)

$Ybar[5, 5] = h[4]*x[-4] + h[3]*x[-3] + h[2]*x[-2] + h[1]*x[-1] + h[0]*x[0]$

$Ybar[6, 5] = h[4]*x[-3] + h[3]*x[-2] + h[2]*x[-1] + h[1]*x[0] + h[0]*x[1]$


(* variable Ybar - for i=n,...,2n and j=n+1 *)

$Y[5, 6] = \text{undef}$

$Y[6, 6] = h[4]*x[-3] + h[3]*x[-2] + h[2]*x[-1] + h[1]*x[0] + h[0]*x[1]$

$Y[7, 6] = \text{undef}$

$Y[8, 6] = \text{undef}$


$Y[10, 6] = \text{undef}$

C.4 Knapsack Problem

This program corresponds to the specification of Section 5.3.3. The name barW is used to indicate $W$.

C.4.1 Program

(* Knapsack Problem *)
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(* constants *)
barW = Floor[c/(p+1)]
infty = 10000

(* domains *)
(* D1 *)
pi1 = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta1 = {0, 0, 1, -c, 0, 0}

(* D2 *)
pi2 = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta2 = {1, -n, 0, 0, 0, 0}

(* D2tr *)
pi2tr = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta2tr = {1, -n, -barW, barW, barW, -barW}

(* D3 *)
pi3 = {{1, 0, 0}, {-1, 0, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta3 = {1, -n, 1, 0, 0}

(* D4 *)
pi4 = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta4 = {1, -n, 1, -c, 0, 0}

(* D4tr *)
pi4tr = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta4tr = {1, -n, 1-barW, -c+barW, barW, -barW}

(* D4(1) *)
pi41 = {{1, 0, 0}, {-1, 0, 0}, {0, 0, 1}, {0, 0, -1}, {0, 1, 1}, {0, -1, -1}}
theta41 = {1, -n, 0, -barW, 1, -c}
(* D4(1,1) *)
\[
\pi_{411} = \{(1, 0, 0), (-1, 0, 0), (0, 0, 1), (0, 0, -1), (0, 1, 1), (0, -1, -1)\}
\]
\[
\theta_{411} = \{1, -n, 0, -\bar{w}+1, 1, -c\}
\]

(* D4(1,2) *)
\[
\pi_{412} = \{(1, 0, 0), (-1, 0, 0), (0, 0, 1), (0, 0, -1), (0, 1, 1), (0, -1, -1)\}
\]
\[
\theta_{412} = \{1, -n, \bar{w}, -\bar{w}, 1, -c\}
\]

(* D4(2) *)
\[
\pi_{42} = \{(1, 0, 0), (-1, 0, 0), (0, 0, 1), (0, 0, -1), (0, 1, -1), (0, -1, -1)\}
\]
\[
\theta_{42} = \{1, -n, 0, -\bar{w}, -2\bar{w}, -c\}
\]

(* D4(2,1) *)
\[
\pi_{421} = \{(1, 0, 0), (-1, 0, 0), (0, 0, 1), (0, 0, -1), (0, 1, -1), (0, -1, -1)\}
\]
\[
\theta_{421} = \{1, -n, 1, -\bar{w}, -2\bar{w}, -c\}
\]

(* D4(2,2) *)
\[
\pi_{422} = \{(1, 0, 0), (-1, 0, 0), (0, 0, 1), (0, 0, -1), (0, 1, -1), (0, -1, -1)\}
\]
\[
\theta_{422} = \{1, -n, 0, 0, -2\bar{w}, -c\}
\]

(* D4(2,3) *)
\[
\pi_{423} = \{(1, 0, 0), (-1, 0, 0), (0, 0, 1), (0, 0, -1), (0, 1, -1), (0, -1, -1)\}
\]
\[
\theta_{423} = \{1, -n, 0, -\bar{w}+1, -2\bar{w}, -c\}
\]

(* belongs to *)
\[
in[\pi_, \theta_, \text{point}_\_] := \text{in[pi, theta, point]} = \\
\quad \text{Apply[And, Map[NonNegative, pi . point - theta]]}
\]

(* applied functions *)
\[
inF1[i\_, j\_, k\_] := 0
\]
\[
inF2[i\_, j\_, k\_] := -\text{infty}
\]
\[
inV[i\_, j\_, k\_] := v[[i]]
\]
\[
inWtr[i\_, j\_, k\_] := w[[i]]
\]
\[
inGamma[i\_, j\_, k\_] := \bar{w}
\]
f[a_, b_, c_] := Max[a, b+c]
f1[alpha_, beta_, gamma_, a1_, a20_, a21_, a2p_] :=
    If[alpha != gamma, a1, If[beta == 0, a20, If[beta == 1, a21, a2p] ] ]

(* variable V *)
V[i_, j_, k_:] := V[i, j, k] =
    If[ in[pi2, theta2, {i, j, k}], inV[i, j, k],
        If[ in[pi4, theta4, {i, j, k}], V[i, j-1, k], undef ] ]

(* variable Wtr *)
Wtr[i_, j_, k_:] := Wtr[i, j, k] =
    If[ in[pi2tr, theta2tr, {i, j, k}], inWtr[i, j, k],
        If[ in[pi4tr, theta4tr, {i, j, k}], Wtr[i, j-1, k], undef ] ]

(* control variable alpha *)
alpha[i_, j_, k_:] := alpha[i, j, k] =
    If[ in[pi411, theta411, {i, j, k}], alpha[i-1, k],
        If[ in[pi412, theta412, {i, j, k}],
            Floor[Wtr[i, j, k]/(p+1)], undef ] ]

(* control variable beta *)
beta[i_, j_, k_:] := beta[i, j, k] =
    If[ in[pi411, theta411, {i, j, k}], beta[i-1, k],
        If[ in[pi412, theta412, {i, j, k}],
            Mod[Wtr[i, j, k], (p+1)], undef ] ]

(* control variable gamma *)
gamma[i_, j_, k_:] := gamma[i, j, k] =
    If[ in[pi411, theta411, {i, j, k}], gamma[i-1, k]+1,
        If[ in[pi412, theta412, {i, j, k}],
            inGamma[i, j, k], undef ] ]

(* variable R2 *)
R2[i_, j_, k_:] := R2[i, j, k] =
    If[ in[pi421, theta421, {i, j, k}], R3[i, j-1, k-1],
        "variable R2" ]
If[ in[pi422, theta422, {i, j, k}], F[i, j, k], undef ]

(* variable R3 *)
R3[i_, j_, k_] := R3[i, j, k] =
  If[ in[pi423, theta423, {i, j, k}], R2[i, j-1, k], undef ]

(* variable R1 *)
R1[i_, j_, k_] := R1[i, j, k] =
  If[ in[pi41, theta41, {i, j, k}],
      f1[alpha[i, j, k], beta[i, j, k], gamma[i, j, k], R1[i, j-1, k+1],
          R2[i, j, k], R2[i, j-1, k], R2[i, j-2, k], undef ]
  ]

(* variable F *)
F[i_, j_, k_] := F[i, j, k] =
  If[ in[pi1, theta1, {i, j, k}], inF1[i, j, k],
      If[ in[pi2, theta2, {i, j, k}], inF1[i, j, k],
          If[ in[pi3, theta3, {i, j, k}], inF2[i, j, k],
              If[ in[pi4, theta4, {i, j, k}],
                  f[F[i-1, j, k], R1[i, j, k], V[i, j-1, k]], undef ]
              ]
          ]
      ]
  ]

C.4.2 Test

The knapsack problem is a dynamic problem. Hence we cannot obtain an entirely symbolic computation of the algorithm. We have provided an actual weight distribution (variable w below) and obtained the corresponding optimal value \( F(4, 10, 0) \) (value of \( F[4, 10, 0] \) in our test case).

(* test *)
(* size parameters *)
c = 10
n = 4
p = 2

(* inputs *)
v = \{1, 3, 5, 9\}
\[ w = \{2, 3, 4, 7\} \]

(* outputs *)
\[ F[4, 10, 0] = 12 \]

**C.5 Gaussian Elimination with Partial Pivoting**

The program below corresponds to the first iteration of the algorithm as specified in Section 5.4.2.

**C.5.1 Program**

(* Pivoting *)

(* constants *)
\[ \text{infty} = 1000000 \]

(* domains *)
(* D1 *)
\[ \text{pi1} = \{\{1, 0, 0\}, \{-1, 0, 0\}, \{0, 1, 0\}, \{0, -1, 0\}, \{0, 0, 1\}, \{0, 0, -1\}\} \]
\[ \text{theta1} = \{0, 0, 1, -n, 1, -n\} \]

(* D2 *)
\[ \text{pi2} = \{\{1, 0, 0\}, \{-1, 0, 0\}, \{-1, 1, 0\}, \{1, -1, 0\}, \{0, 0, 1\}, \{0, 0, -1\}\} \]
\[ \text{theta2} = \{0, -n+2, n+1, -n-1, 1, -n\} \]

(* D3 *)
\[ \text{pi3} = \{\{1, 0, 0\}, \{-1, 0, 0\}, \{-1, 1, 0\}, \{1, -1, 0\}, \{0, 0, 1\}, \{0, 0, -1\}\} \]
\[ \text{theta3} = \{1, -n+1, -1, 1, 1, -n\} \]

(* D41 *)
\[ \text{pi41} = \{\{1, 0, 0\}, \{-1, 0, 0\}, \{-1, 1, 0\}, \{1, -1, 0\}, \{0, 0, 1\}, \{0, 0, -1\}\} \]
\[ \text{theta41} = \{1, -n+1, 0, -n, 1, -1\} \]

(* D42 *)
pi42 = {{1, 0, 0}, {-1, 0, 0}, {-1, 1, 0}, {1, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta42 = {1, -n+1, 0, -n, 2, -n}

(* D51 *)
pi51 = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta51 = {n, -n, n+1, -2n+1, 1, -1}

(* D52 *)
pi52 = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta52 = {n, -n, n+1, -2n+1, 2, -n}

(* D6 *)
pi6 = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta6 = {n, -n, n, -n, 1, -n}

(* D7 *)
pi7 = {{1, 0, 0}, {-1, 0, 0}, {0, 1, 0}, {0, -1, 0}, {0, 0, 1}, {0, 0, -1}}
theta7 = {n, -n, n+1, -2n+1, 1, -n}

(* belongs to *)
in[pi_, theta_, point_] := in[pi, theta, point] =
   Apply[And, Map[NonNegative, pi . point - theta]]

(* applied functions *)
inV1[i_, j_, k_] := a[[j, k]]
inV2[i_, j_, k_] := 0
inV3[i_, j_, k_] := 0
maxAbs[a_, b_] := If[Abs[a] >= Abs[b], a, b]
swV[a_, b_, c_] := If[c == 1, b, a]
inH[i_, j_, k_] := infty
minAbs[a_, b_] := If[Abs[a] >= Abs[b], Abs[b], Abs[a]]
swH[a_, b_, c_] := If[c == 1, a, b]
cmpAbs[a_, b_] := If[Abs[a] >= Abs[b], 0, 1]
id[a_] := a
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 upd[a_, b_, c_] := a - b * c
div[a_, b_] := a / b

(* variable V *)
V[i_, j_, k_] := V[i, j, k] =
  If[ in[pi1, theta1, {i, j, k}], inV1[i, j, k],
    If[ in[pi2, theta2, {i, j, k}], inV2[i, j, k],
      If[ in[pi41, theta41, {i, j, k}],
        maxAbs[V[i-1, j, k], H[i, j-1, k]],
        maxAbs[V[i, j, k], H[i, j, k]],
      maxAbs[V[i, j, k], H[i, j, k]]],
    maxAbs[V[i, j, k], H[i, j, k]]],
  maxAbs[V[i, j, k], H[i, j, k]]]

(* variable H *)
H[i_, j_, k_] := H[i, j, k] =
  If[ in[pi3, theta3, {i, j, k}], inH[i, j, k],
    If[ in[pi41, theta41, {i, j, k}],
      minAbs[V[i-1, j, k], H[i, j-1, k]],
    minAbs[V[i, j, k], H[i, j-1, k]]],
  minAbs[V[i, j, k], H[i, j, k]]]

(* control variable CC *)
CC[i_, j_, k_] := CC[i, j, k] =
  If[ in[pi41, theta41, {i, j, k}], cmpAbs[V[i-1, j, k], H[i, j-1, k]],
    cmpAbs[V[i, j, k], H[i, j-1, k]],
  cmpAbs[V[i, j, k], H[i, j-1, k]]]

(* row coefficient RC *)

\( RC[i_, j_, k_] := RC[i, j, k] = \)
\[
\text{If[ in[p151, theta51, \{i, j, k\}], div[V[i-1, j, k], H[i, j-1, k]],}
\]
\[
\text{If[ in[p152, theta52, \{i, j, k\}], id[RC[i, j, k-1]], undef ] ]}
\]

**C.5.2 Test**

Gaussian elimination with pivoting is a dynamic problem and a numerical execution of the program is required. In our test case we have considered a 5 x 5 input matrix (\( a \) below). The outputs show such the initial matrix (origmat), the matrix obtained by rearranging (rearmat) the entries once the pivot has been established, and the updated matrix (updmat) at the end of the iteration.

(* test case *)

(* size parameter *)

\( n = 5 \)

(* input matrix *)

\( a = \{\{1, 0, 1, 4, 2\}, \{2, 1, 0, 3, 0\}, \{1, 1, 2, 1\}, \{3, 1, 0, 2, 1\}, \{1, 5, 1, 2, 1\}\} \)

(* outputs - single iteration *)

(* original matrix *)

\( \text{origmat} = \text{Flatten[Table[V[i,j,k],\{i,0,0\}, \{j,i+1,i+n\}, \{k,1,n\}], 1]} \)
\( \text{MatrixForm[origmat]} \)
\[
\begin{array}{cccc}
1 & 0 & 1 & 4 & 2 \\
2 & 1 & 0 & 3 & 0 \\
1 & 1 & 1 & 2 & 1 \\
3 & 1 & 0 & 2 & 1 \\
1 & 5 & 1 & 2 & 1 \\
\end{array}
\]

(* rearranged matrix -- end of phase one *)

\( \text{rearmat} = \text{Flatten[Table[V[i,j,k],\{i,n-1,n-1\}, \{j,i+1,i+n\}, \{k,1,n\}], 1]} \)
\( \text{MatrixForm[rearmat]} \)
\[
\begin{array}{cccc}
3 & 1 & 0 & 2 & 1 \\
2 & 1 & 0 & 3 & 0 \\
\end{array}
\]
APPENDIX C. CASE STUDIES

\[
\begin{array}{cccc}
1 & 1 & 1 & 2 \\
1 & 5 & 1 & 2 \\
1 & 0 & 1 & 4 \\
\end{array}
\]

(* updated matrix *)

\[
\text{updmat = Flatten}\left[\text{Table}\left[V[i,j,k],\{i,n\},\{j,i,i+n-1\},\{k,1,n\}\right],1\right]
\]

\[
\text{MatrixForm}[\text{updmat}]
\]

\[
\begin{array}{cccc}
3 & 1 & 0 & 2 \\
0 & 1/3 & 0 & 5/3 & -(2/3) \\
0 & 2/3 & 1 & 4/3 & 2/3 \\
0 & 14/3 & 1 & 4/3 & 2/3 \\
0 & -(1/3) & 1 & 10/3 & 5/3 \\
\end{array}
\]
Appendix D

Graph Theory

This appendix is based on [Car79].

D.1 Graphs

A graph is a pair \( G = (N, A) \) such that \( N \) is a finite set and \( A \subseteq N \times N \). The set \( N \) is called the set of nodes of \( G \) and \( A \) its a set of arcs. The empty graph is the graph \( G = (\emptyset, \emptyset) \).

A graph \( G \) is said to be simple if and only if

- for all \( n_i \in N \), \( (n_i, n_i) \notin A \); and
- \( (n_i, n_j) \in A \) implies \( (n_j, n_i) \in A \).

Let \( G = (N, A) \) be a graph and \( \mathcal{P} = \{N_1, \ldots, N_n\} \) be a partition of \( N \). The condensation of \( G \) induced by \( \mathcal{P} \) is the graph \( G_{\mathcal{P}} = (\mathcal{P}, A_{\mathcal{P}}) \) such that \( A_{\mathcal{P}} = \{ (N_i, N_j) \in \mathcal{P} \times \mathcal{P} \mid N_i \neq N_j \} \) and \( \exists n_i \in N_i, n_j \in N_j \) such that \( (n_i, n_j) \in A \).

The simplification of a graph \( G \) is the graph \( G_S = (N, A_S) \) such that \( A_S = \{ (n_i, n_j) \in N \times N \mid n_i \neq n_j \text{ and either } (n_i, n_j) \in A \text{ or } (n_j, n_i) \in A \} \).

A path \( p \) of a graph \( G \) is a finite sequence \( (n_0, n_1), (n_1, n_2), \ldots, (n_{r-1}, n_r) \) such that \( \forall i, j \), \( (n_i, n_j) \in A \). The order of \( p \) is the number of arcs in the sequence. A path \( p \) is called a cycle if \( n_0 = n_r \). A loop is a cycle of order one.

Let \( G = (N, A) \) be a graph and \( n \in N \). Then:

- a descendant of \( n \) is a node \( n' \) such that there exists a path \( (n_0, n_1), (n_1, n_2), \ldots, (n_{r-1}, n_r) \) with \( n_0 = n \) and \( n_r = n' \). We denote by \( n_1 \) the set of all the descendants of \( n \); and
- a node accessible from $n$ is a node $n'$ such that $n' \in n$ or $n' = n$. We denote by $n^A$ the set of all nodes accessible from $n$.

### D.2 Connectivity Relations

Let $G = (N, A)$ be a graph. Then:

- the connectivity relation $\triangleright \subseteq N \times N$ of $G$ is the equivalence relation such that $n \triangleright m$ if and only if $m \in n^A$ on the simplification $G_S$ of $G$. When $n \triangleright m$, $n$ is said to be connected to $m$; and

- the connected components of $G$ are the subgraphs of $G$ generated by the equivalence classes of $\triangleright$. If $G$ has only one connected component, $G$ is said to be connected.

- the strong connectivity relation $\cong \subseteq N \times N$ of $G$ is the equivalence relation such that $n \cong m$ if and only if $m \in n^A$ and $n \in m^A$ on $G$. When $n \cong m$, $n$ is said to be strongly connected to $m$; and

- the strongly connected components of $G$ are the subgraphs of $G$ generated by the equivalence classes of $\cong$. If $G$ has only one strongly connected component, $G$ is said to be strongly connected.

Let $G = (N, A)$ be a graph and $\cong$ its strong connectivity relation. The reduced graph $G_R$ of $G$ is the condensation of $G$ induced by $N / \cong$.

### D.3 Graph Operations

Let $G = (N, A)$ be a graph and $N' \subseteq N$. The restriction of $G$ to $N'$ is the graph $G |_{N'} = (N', A')$ such that $A' = \{(n, n') \in A \mid n, n' \in N'\}$.

Let $G_j = (N_j, A_j)$ be graphs, for $j = 1, \ldots, r$. Their union is the graph $G = (N, A)$ such that $N = \bigcup_j N_j$ and $A = \bigcup_j A_j$. 
Appendix E

Convex Sets and Polyhedra

This appendix is based on [Roc70, Sch86]. In the appendix, for \( x = (x_1, \ldots, x_n), y = (y_1, \ldots, y_n) \in \mathbb{R}^n, x \cdot y \) denotes their scalar product \( x_1 y_1 + \ldots + x_n y_n \).

E.1 Combinations

Let \( x_1, \ldots, x_m \) be \( m \) points in \( \mathbb{R}^n \). A vector sum \( \lambda_1 x_1 + \ldots + \lambda_m x_m \), with \( \lambda_1, \ldots, \lambda_m \in \mathbb{R} \), is called:

- a linear combination of \( x_1, \ldots, x_m \):
- an affine combination of \( x_1, \ldots, x_m \), if \( \lambda_1 + \ldots + \lambda_m = 1 \);
- a convex combination of \( x_1, \ldots, x_m \), if the coefficients \( \lambda_i \) are all non-negative and \( \lambda_1 + \ldots + \lambda_m = 1 \);
- a positive (non-negative) linear combination of \( x_1, \ldots, x_m \), if the coefficients \( \lambda_i \) are all positive (non-negative);
- an integer combination of \( x_1, \ldots, x_m \), if the coefficients \( \lambda_i \) are all integers.

A vector sum \( \lambda_1 x_1 + \ldots + \lambda_k x_k + \lambda_{k+1} x_{k+1} + \ldots + \lambda_m x_m \) is called a convex combination of \( m \) points and directions if all the coefficients \( \lambda_i \) are non-negative and \( \lambda_1 + \ldots + \lambda_k = 1 \) for a fixed \( k \), with \( 0 \leq k \leq m \) (\( k = 0 \) means that there is no requirement about certain coefficients adding up to 1).
E.2 Affine Sets and Transformations

If \( x \) and \( y \) are distinct points in \( \mathbb{R}^n \), the set of points of the form \((1 - \lambda)x + \lambda y\), for \( \lambda \in \mathbb{R} \), is called the line through \( x \) and \( y \).

A subset \( M \) of \( \mathbb{R}^n \) is called an affine set if it contains the line through any pair of its points, i.e., if \((1 - \lambda)x + \lambda y\in M\) for every \( x \in M \), \( y \in M \) and \( \lambda \in \mathbb{R} \). The empty set \( \emptyset \), \( \mathbb{R}^n \) and all singleton sets are affine.

For \( M \subset \mathbb{R}^n \) and \( a \in \mathbb{R}^n \), the translate of \( M \) by \( a \) is \( M + a = \{ a + x \mid x \in M \} \). An affine set \( M \) is said to be parallel to an affine set \( L \) if \( M = L + a \) for some \( a \).

The subspaces of \( \mathbb{R}^n \) are the affine sets which contain the origin.

Each non-empty affine set \( M \) is parallel to a unique subspace \( L \), given by \( L = M - M = \{ x - y \mid x \in M, y \in M \} \). \( L \) is called the direction of \( M \) and is denoted by \( \text{lin}(M) \).

The dimension \( \dim(M) \) of an affine set \( M \) is defined as the dimension of the subspace \( L \) parallel to it, i.e., \( \dim(M) = \dim(\text{lin}(M)) \). By convention, \( \dim(\emptyset) = -1 \). Affine sets of dimension 0, 1 and 2 are called points, lines and planes, respectively. An \((n - 1)\)-dimensional affine set in \( \mathbb{R}^n \) is called a hyperplane.

Given \( \beta \in \mathbb{R} \) and a non-zero \( b \in \mathbb{R}^n \), the set \( H = \{ x \mid b^t \cdot x = \beta \} \) is a hyperplane in \( \mathbb{R}^n \). Every hyperplane may be represented in this way, with \( b \) and \( \beta \) unique up to a common non-zero multiple. \( b \) is called a normal to the hyperplane \( H \). We denote a hyperplane \( H \) by \([b : \beta] \).

Given \( b \in \mathbb{R}^m \) and an \( m \times n \) real matrix \( B \), the set \( H = \{ x \in \mathbb{R}^n \mid Bx = b \} \) is an affine set in \( \mathbb{R}^n \). Every affine set may be represented in this way. (Therefore, any affine subset of \( \mathbb{R}^n \) is an intersection of a finite collection of hyperplanes.)

The intersection of an arbitrary collection of affine sets is affine. Given \( S \in \mathbb{R}^n \), the intersection of all the affine sets containing \( S \) is called the affine hull of \( S \) and is denoted by \( \text{aff}(S) \). \( \text{aff}(S) \) is the unique smallest affine set containing \( S \) and consists of all the affine combinations of the elements of \( S \), i.e., \( \text{aff}(S) = \{ \sum \lambda_i x_i \mid x_i \in S, \sum \lambda_i = 1 \} \).

A set of \( m + 1 \) points \( b_0, b_1, \ldots, b_m \) is said to be affinely independent if \( \text{aff}(\{b_0, b_1, \ldots, b_m\}) \) is \( m \)-dimensional. Therefore \( b_0, b_1, \ldots, b_m \) are affinely independent if and only if \( b_1 - b_0, \ldots, b_m - b_0 \) are linearly independent.

The affine transformations from \( \mathbb{R}^n \) to \( \mathbb{R}^m \) are the mappings \( T \) of the form \( T(x) = L(x) + b \), where \( L \) is a linear transformation from \( \mathbb{R}^n \) to \( \mathbb{R}^m \) and \( b \in \mathbb{R}^m \).
E.3 Convex Sets

If \( x \) and \( y \) are distinct points in \( \mathbb{R}^n \), the set of points of the form \((1 - \lambda)x + \lambda y\), for \( 0 \leq \lambda \leq 1 \), is called the closed line segment between \( x \) and \( y \).

A subset \( C \) of \( \mathbb{R}^n \) is said to be convex if it contains the closed line segment between any two of its points, i.e., \((1 - \lambda)x + \lambda y \in C\) for every \( x \in M, y \in M \) and \( 0 < \lambda < 1 \). The empty set \( \emptyset \) and \( \mathbb{R}^n \) are convex.

Given \( \beta \in \mathbb{R} \) and a non-zero \( b \in \mathbb{R}^n \), the sets \( \{x \mid b^t \cdot x \leq \beta\} \) and \( \{x \mid b^t \cdot x \geq \beta\} \) are called closed half-spaces, and the sets \( \{x \mid b^t \cdot x < \beta\} \) and \( \{x \mid b^t \cdot x > \beta\} \) are called open half-spaces. Half-spaces are non-empty and convex.

A subset of \( \mathbb{R}^n \) is convex if and only if it contains all the convex combinations of its elements. The intersection of an arbitrary collection of convex sets is convex. Given \( S \in \mathbb{R}^n \), the intersection of all the convex sets containing \( S \) is called the convex hull of \( S \) and is denoted by \( \text{conv}(S) \). \( \text{conv}(S) \) is the unique smallest convex set containing \( S \) and consists of all the convex combinations of the elements of \( S \), i.e., \( \text{conv}(S) = \{\sum \lambda_i x_i \mid x_i \in S, \lambda_i \geq 0, \sum \lambda_i = 1\} \).

The dimension \( \text{dim}(C) \) of a convex set \( C \) is defined as the dimension of the affine hull \( \text{aff}(C) \) of \( C \).

E.4 Cones

A subset \( K \) of \( \mathbb{R}^n \) is called a cone if it is closed under non-negative scalar multiplication, i.e., \( \lambda x \in K \) for every \( x \in K \) and \( \lambda \geq 0 \). A convex cone is a cone which is convex.

Let \( K \) be a convex cone. Then there is a smallest subspace containing \( K \), namely \( K - K = \{x - y \mid x \in K, y \in K\} = \text{aff}(K) \), and there is a largest subspace contained within \( K \), namely \((-K) \cap K \). Cones are not necessarily pointed. For instance, subspaces are in particular convex cones. A cone \( K \) is pointed if and only if \((-K) \cap K = \{0\} \).

E.5 Recession Cone and Unboundedness

Unbounded closed convex sets have a simple behaviour at infinity. If \( C \) is an unbounded closed convex set and \( x \in C \), then \( C \) contains some entire half-lines starting at \( x \). The directions of such half-lines do not depend on \( x \): the half-lines of \( C \) starting at a different point \( y \) are just translates of those starting at \( x \).
The direction of the half-line \( \{ x + \lambda y \mid \lambda \geq 0 \} \), where \( y \neq 0 \), is defined as the set of all translates of the half-line, and is independent of \( x \). This is called the direction of \( y \). Two vectors have the same direction if and only if they are positive scalar multiples of each other. The zero vector has no direction.

Let \( C \) be a non-empty convex set in \( \mathbb{R}^n \). \( C \) recedes in the direction of \( y \) if \( C \) includes all the half-lines in the direction of \( y \) which start at points in \( C \), i.e., \( C \) recedes in the direction of \( y \), where \( y \neq 0 \), if and only if \( x + \lambda y \in C \) for every \( \lambda \geq 0 \) and \( x \in C \).

The recession cone of \( C \), denoted by \( 0^+C \), is the set of all vectors \( y \in \mathbb{R}^n \) satisfying the latter condition, including \( y = 0 \). The directions of the recession of \( C \) are directions in which \( C \) recedes.

Let \( C \) be a non-empty convex set. The recession cone \( 0^+C \) is a convex cone. It is the same as the set of vectors \( y \) such that \( C + y \subset C \).

A non-empty closed convex set \( C \) in \( \mathbb{R}^n \) is bounded if and only if its recession cone \( 0^+C \) consists of the zero vector alone.

If \( C \) is a non-empty convex set, the set \( (-0^+C) \cap 0^+C \) is called the linearity space of \( C \). It consists of the zero vector and all the non-zero vectors \( y \) such that, for every \( x \in C \), the line through \( x \) in the direction of \( y \) is contained in \( C \). The directions of the vectors \( y \) in the linearity space are called directions in which \( C \) is linear. If the linearity space has dimension 0, \( C \) is called pointed.

### E.6 Polyhedral Convex Sets

A polyhedral convex set in \( \mathbb{R}^n \) is a set which can be expressed as the intersection of some finite collection of closed half-spaces. Every affine set is polyhedral.

A polyhedral convex cone in \( \mathbb{R}^n \) is a set which can be expressed as the intersection of a finite collection of closed half-spaces whose boundary hyperplanes pass through the origin.

A finitely generated convex set is a set which is the convex hull of a finite set of points and directions. Thus \( C \) is a finitely generated convex set if and only if there exist vectors \( x_1, \ldots, x_m \) such that, for a fixed integer \( k \), with \( 0 \leq k \leq m \), \( C \) consists of all the vectors of the form

\[
x = \lambda_1 x_1 + \ldots + \lambda_k x_k + \lambda_{k+1} x_{k+1} + \ldots + \lambda_m x_m
\]

where \( \lambda_i \) are non-negative and \( \lambda_1 + \ldots + \lambda_k = 1 \).
The finitely generated convex sets which are cones are the sets which can be expressed this way with \( k = 0 \), i.e., with no requirement about certain coefficients adding up to 1. In such an expression, \( \{x_1, \ldots, x_m\} \) is called a set of generators of the cone. A finitely generated convex cone is the convex hull of the origin and finitely many directions.

The cone generated by the vectors \( x_1, \ldots, x_m \) is \( \text{cone}(x_1, \ldots, x_m) = \{\lambda_1 x_1 + \ldots + \lambda_m x_m \mid \lambda_1, \ldots, \lambda_m \geq 0\} \), i.e., it is the smallest convex cone containing \( x_1, \ldots, x_m \).

The finitely generated convex sets which are bounded are called polytopes.

The property of being polyhedral is a finiteness condition on the external representations of a convex set. The property of being finitely generated is a finiteness condition on the internal representations of a convex set. The two properties are equivalent, and polyhedral convex sets are the same as finitely generated convex sets.

The concepts of polyhedron and polytope are related under the so-called decomposition theorem. The theorem states that a set \( P \) of vectors in a Euclidean space is a convex polyhedron if and only if \( P = Q + C \) for some convex polytope \( Q \) and some polyhedral convex cone \( C \).

Also, if \( P = Q + C \) for some convex polytope \( Q \) and some polyhedral convex cone \( C \), then \( C \) is the recession cone of \( P \).

We say that \( P \) is generated by the points \( x_1, \ldots, x_m \) and the directions \( y_1, \ldots, y_l \) if

\[
P = \text{conv}(\{x_1, \ldots, x_m\}) + \text{cone}(\{y_1, \ldots, y_l\}).
\]

In this case we adopt the notation \( \text{vert}(P) \) to denote the set \( \{x_1, \ldots, x_m\} \) and \( \text{ray}(P) \) for the set \( \{y_1, \ldots, y_l\} \).

**E.7 Duality**

Let \( P \) be a set in \( \mathbb{R}^n \). The dual (or polar) cone of \( P \) is the cone \( \hat{P} = \{z \in \mathbb{R}^n \mid z \cdot x \geq 0, \forall x \in P\} \).

If \( P \) and \( P' \) are such that \( P \subseteq P' \), then \( \hat{P} \subseteq \hat{P'} \).

If \( C \) is a polyhedral convex cone, then \( \hat{C} \) is also a polyhedral convex cone. If \( C \) is finitely generated by the set \( \{r_1, \ldots, r_m\} \), then \( \hat{C} = \{z \in \mathbb{R}^n \mid z \cdot r_1 \geq 0, \ldots, z \cdot r_m \geq 0\} \).

There exists a one-to-one correspondence between the generators of \( C \) and those of \( \hat{C} \). In particular, let \( C \) be an \( n \) dimensional cone with \( n \) generators \( \{r_1, \ldots, r_n\} \), and let \( Q \) denote a
matrix having those generators as columns. Then the generators of $\tilde{C}$ are the column of the matrix $\tilde{Q} = -(Q^{-1})^t$, i.e., the opposite of the transpose of the inverse of $Q$.

**E.8 Separating Hyperplane Theorem**

If $C$ is a finitely generated cone in $\mathbb{R}^n$ and $d$ is a vector not in $C$, then there exists $\pi \in \mathbb{R}^n$ such that $\pi \cdot x \geq 0$ for all $x \in C$, and $\pi \cdot d < 0$. That is, the hyperplane $[\pi : 0]$ divides $\mathbb{R}^n$ into two parts, one containing $d$ and the other containing $C$. In addition, $C$ is pointed if and only if $\pi \cdot x > 0$ for all $x \in C$.

**E.9 Other Results**

Let $\mathcal{L}$ be a linear transformation from $\mathbb{R}^n$ to $\mathbb{R}^m$. Then $\mathcal{L}(C)$ is a polyhedral convex set in $\mathbb{R}^m$ for each polyhedral convex set $C$ in $\mathbb{R}^n$, and $\mathcal{L}^{-1}(D)$ is a polyhedral convex set in $\mathbb{R}^n$ for each polyhedral convex set $D$ in $\mathbb{R}^m$.

If $D$ is a convex polyhedron and $y_1, \ldots, y_m$ are directions in $\mathbb{R}^n$, the polyhedron generated by $D$ and $y_1, \ldots, y_m$ is defined as the set

$$P = \{ z + \lambda_1 d_1 + \ldots + \lambda_m d_m \mid z \in D, \lambda_i \geq 0 \}.$$
Appendix F

Aspects of Linear Algebra

This appendix is based on [Ner63, Ban93]. In this appendix, $V$ denoted a generic vector space over a generic field $F$. If $v_1, \ldots, v_p$ are vectors in $V$, then $(v_1, \ldots, v_p)$ denotes the subspace spanned by $v_1, \ldots, v_p$.

F.1 Elementary Row Operations and Elementary Matrices

There are three types of elementary row operations\footnote{Elementary column operations are defined in a similar way. In this work we consider elementary row operations only.} defined on (the rows of) a matrix $M$: i) multiply a row of $M$ by a non-zero scalar; ii) add a multiple of one row to another row; iii) interchange two rows.

An elementary matrix is any matrix obtained from an identity matrix by any elementary row operation. Therefore there is a one-to-one correspondence between elementary row operations and elementary matrices. An elementary matrix is a non-singular matrix and its inverse matrix is also an elementary matrix.

An elementary row operation on a matrix $M$ can be accomplished by premultiplying $M$ by the corresponding elementary matrix.

Any non-singular matrix $M$ can be written as a product of elementary matrices.
F.2 Hermite Normal Form

A matrix in Hermite normal form\(^2\) has the following form:

\[
\begin{bmatrix}
0 & \ldots & 0 & 1 & z & x & 0 & z & 0 & z & x \\
0 & \ldots & 0 & 0 & 0 & 1 & z & 0 & z & x \\
0 & \ldots & 0 & 0 & 0 & 0 & 1 & 0 & z & x \\
0 & \ldots & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
0 & \ldots & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\end{bmatrix}
\]

where an \(x\) denotes any number.

Any \(m \times n\) matrix can be reduced to its Hermite normal form, and such a form is unique. The reduction to Hermite normal form can be achieved by a series of elementary row operations. As the form is unique, it is independent from the the particular sequence of operations chosen.

If \(M\) is a non-singular square matrix, its Hermite normal form is the identity matrix.

The Hermite normal form has a number of important applications in linear algebra among which finding a standard basis for a subspace \(S\) and its orthogonal complement \(S^\perp\), determining the linearly independent vectors among the vectors of a set, solving systems of linear equations, inverting a matrix, etc. (see this appendix later on).

F.3 Integer Elementary Row Operations

As we mainly work in \(\mathbb{Z}^n\), then in general, we restrict ourselves to integer matrices and integer elementary row operations [Ban93]. There are three types of integer elementary row operations: i) multiply a row of \(M\) by \(-1\); ii) add an integer multiple of one row to another row; iii) interchange two rows.

F.4 Unimodularity

A square integer matrix \(M\) is unimodular if its determinant \(\det(M)\) is equal to ±1.

Each unimodular matrix is the result of a finite sequence of integer elementary row operations performed on the identity matrix (of the same size).

Unimodular matrices have the following properties: the transpose and the inverse of a unimodular matrix is unimodular; the product of two unimodular matrices is unimodular.

\(^2\)This form is often called row-echelon form.
F.5 Echelon Form

Let \( M \) be an \( m \times n \) integer matrix and \( l_i \) denote the column number of the leading element of row \( i \) (for a zero row, \( l_i \) is undefined). Then \( M \) is in Echelon form\(^3\) if for some integer \( p \), with \( 0 \leq p \leq m \), the following conditions hold:

- rows from 1 to \( p \) are non-zero rows;
- rows from \( p + 1 \) to \( m \) are zero rows;
- for \( 1 \leq i \leq p \), each element in column \( l_i \) below row \( i \) is zero;
- \( l_1 < l_2 < \ldots < l_p \).

The leading element of a row need not be equal to 1. A zero matrix is an echelon matrix for which \( p = 0 \). The rank of a matrix in Echelon form is equal to \( p \), i.e., \( \text{rank}(M) = p \).

A matrix in Echelon form has the following form:

\[
\begin{bmatrix}
0 & \cdots & 0 & x_1 & x & x & x & x & x & x \\
0 & \cdots & 0 & 0 & 0 & 0 & x_2 & x & x & x \\
0 & \cdots & 0 & 0 & 0 & 0 & 0 & 0 & x_p & x \\
0 & \cdots & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
\vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots & \vdots \\
0 & \cdots & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0
\end{bmatrix}
\]

where \( x, x_i \) denote integer numbers.

Any integer matrix can be reduced to Echelon form by a sequence of integer elementary row operations. Applying a finite sequence of elementary row operations to a matrix is equivalent to premultiplying the matrix by a suitable unimodular matrix.

F.6 Linear Functional and Duality

Let \( V \) be a vector space over a field of constants \( F \). A linear transformation \( L \) of \( V \) into \( F \) is called a linear form or linear functional on \( V \). The set of all linear functionals on \( V \) is a vector space, called the dual or conjugate space of \( V \), and denoted by \( \hat{V} \). Besides these spaces have the same number of dimensions, i.e., \( \dim(\hat{V}) = \dim(V) \).

\(^3\)In classical text of linear algebra, Hermite normal form and row-echelon form are exactly the same. The distinction we have made here is a little arbitrary. It would be more appropriate to call this form an integer Hermite normal form. However we adopt the terminology of [Ban93].
If $B = \{b_1, \ldots, b_n\}$ is a basis of $V$ then there exists a corresponding dual basis $\hat{B} = \{\hat{b}_1, \ldots, \hat{b}_n\}$ of $\hat{V}$. The relation between the two basis is characterised by the equations $\hat{b}_i(b_j) = \delta_{i,j}$, for all $i, j = 1, \ldots, n$, where $\delta_{i,j}$ is the Kronecker delta ($\delta_{i,j} = 1$ if $i = j$ and $\delta_{i,j} = 0$ if $i \neq j$).

Let $B' = \{b'_1, \ldots, b'_n\}$ be another basis of $V$. Let $P = [p_{i,j}]$ be the matrix of transition from $B$ to $B'$, i.e., $b'_i = \sum_{i=1}^n p_{i,j}b_i$. Then $P^t$ is the matrix of transition from basis $\hat{B}'$ to $\hat{B}$. Therefore $(P^t)^{-1} = (P^{-1})^t$ is the matrix of transition from $\hat{B}$ to $\hat{B}'$.

**F.7 Annihilators**

Let $V$ be an $n$-dimensional vector space and $\hat{V}^\ast$ its dual. If for $v \in V$ and $\hat{v} \in \hat{V}^\ast$, we have $v \cdot \hat{v} = 0$, we say that $v$ and $\hat{v}$ are orthogonal.

Let $W$ be a subset of $V$. The set of all the linear functional $\hat{v}$ such that $\hat{v} \cdot w = 0$ for all $w \in W$, is called the annihilator or orthogonal complement of $W$, denoted by $W^\perp$. Any element of $W^\perp$ is called an annihilator of $W$. $W^\perp$ is a sub-space of $\hat{V}$. Besides if $W$ is a sub-space of dimension $\rho$, then $\hat{V}$ is a sub-space of dimension $n - \rho$.

If $W_1$ and $W_2$ are two subspaces of $V$ and $W_1^\perp$ and $W_2^\perp$ are their annihilators in $\hat{V}$, then the annihilator of $W_1 + W_2$ is $W_1^\perp \cap W_2^\perp$, and the annihilator of $W_1 \cap W_2$ is $W_1^\perp + W_2^\perp$.

**F.8 Algorithmic Issues**

**F.8.1 Standard Basis of $V$**

Let $B = \{b_1, \ldots, b_n\}$ be a basis of a vector space $V$. A standard basis of $V$ can be found according to the following. Represent the basis as a matrix $B = [b_{i,j}]$ where each row is a vector of the basis. Reduce $B$ to its Hermite normal form $B'$. Then the rows of $B'$ represent a standard basis for $V$. This basis is standard because any basis of $V$ reduces to such a basis by Hermite normal form reduction.

**Set Spanning the Same Sub-space**

To decide whether two sets of vectors span the same subspace, we can simply reduce the corresponding matrices to their Hermite normal forms and compare their rows.
Linear Independence

Given a set of vectors \( \{x_1, \ldots, x_k\} \) in an \( n \)-dimensional space, their linear independence can be checked by defining a matrix \( M \) having such vectors as columns and reducing \( M \) to Hermite normal form. The standardised columns (those with just one non-null entry equal to 1) of the resulting matrix correspond to the linearly independent elements of the set.

F.8.2 Basis of \( W_1 + W_2 \)

Given two subspaces \( W_1, W_2 \) of a linear space \( V \), their sum is defined as \( W_1 + W_2 = \{ w_1 + w_2 \mid w_1 \in W_1, w_2 \in W_2 \} \).

If \( B_1 \) and \( B_2 \) are bases of \( W_1 \) and \( W_2 \), respectively, then \( B_1 \cup B_2 \) spans \( W_1 + W_2 \). Finding a basis for \( W_1 + W_2 \) amounts to discarding the dependent vectors of \( B_1 \cup B_2 \) until an independent spanning set remains. Therefore we can construct a matrix whose rows are the vectors in \( B_1 \cup B_2 \) and reduce it to Hermite normal form. The resulting rows constitute a basis of \( W_1 + W_2 \).

F.8.3 Basis of \( W_1 \cap W_2 \)

Instead of finding the intersection \( W_1 \cap W_2 \) directly, it is easier to find \( W_1^\perp \) and \( W_2^\perp \), then \( W_1^\perp + W_2^\perp \) and finally \( (W_1^\perp + W_2^\perp)^\perp \).

F.8.4 Basis of \( W^\perp \)

Given a basis \( \{b_1, \ldots, b_m\} \) of \( W \), a basis of \( W^\perp \) is a basis of the solution space of the homogeneous system of equations \( B \cdot y = 0 \), where \( B \) is the matrix having the vectors \( b_j \) as columns. This can be obtained by reducing \( B \) to Hermite normal form and finding the \( k \) independent columns of the resulting matrix, with \( k \leq n \). If \( k = n \) then \( W \) is full-dimensional and \( W^\perp = \{0\} \). Otherwise, \( W^\perp \) is a subspace of dimension \( n - k \). A basis for such subspace can be found by arbitrarily considering \( n - k \) variables as parameters and solving, with respect to them, the subsystem of equations resulting from the Hermite normal form of \( B \).

Basis of \( \text{null}(A) \)

Finding a basis for \( \text{null}(A) \), where \( A \) is an \( n \times n \) matrix, is equivalent to providing a basis for the solution space of the homogeneous system of linear equations \( A \cdot x = 0 \) for \( x \in \mathbb{R}^n \).
Therefore, finding a basis of $null(A)$ is an instance of the problem of finding a basis for the orthogonal complement of a given subspace.
Bibliography


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