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Computation Algebras[†]

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We introduce a framework that generalizes algebraic specifications by equipping algebras with descriptions of *evaluation strategies*. The resulting abstract mathematical description allows one to model the *implementation* of algebras on various platforms in a way that is independent of the function-oriented specifications.

We study algebras with associated data dependencies. The latter provide separate means for modelling computational aspects apart from the functional specifications captured by an algebra. The formalization of evaluation strategies (1) introduces increased portability among different hardware platforms, and (2) allows a potential increase in execution efficiency, since a chosen evaluation strategy may be tailored to a particular platform. We present the development process where algebraic specifications are equipped with data dependencies, the latter are refined, and, finally, mapped to actual hardware architectures.

1. Introduction

The creed of algebraic specification is *abstraction*: one models programs as algebras, thus abstracting from the operational details and focusing on the abstract functionality of the system. A single such specification may be implemented on different platforms. The implementation design, as well as the particular platform on which it is realized, need not be the primary issues. It is one of the central tenets of algebraic specification that the designer may initially model his system at a high level of abstraction without involving himself in low level implementation details, only introducing implementation concerns as appropriate, much later in the development process. At some point during the development process the choice of the set of constructors, the computation strategy, the target language, and, perhaps, the hardware platform begin to play a more important role. In particular,

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M. Walicki, M. Haveraaen and S. Meldal

because of its intentional abstraction from the operational aspects, the standard algebraic approach to specification has difficulties when faced with the possibilities of parallel implementations. It is at this point that our proposed method comes into play successfully.

We propose a framework of *computation algebras* addressing the issue of how to incorporate computation-related aspects within the framework of classical algebraic semantics. Computation algebras allow the developer to transform abstract requirements (expressed in an algebraic specification) into designs by adding implementation-specific information concerning the order and dependency between various subcomputations. Though occasionally tempting, we would like to avoid rebuilding the theory from scratch – we want to take advantage of the existing theoretical and tool-oriented results of algebraic specifications. Our aim is only to *extend* the classical framework with constructs that in an algebraic way address the choice of computational strategies.

This is achieved by associating additional structures, *data dependency* graphs, with standard algebras. The data depency graphs carry information about evaluation strategies for the operations of the algebras. We have to warn the reader that our 'data dependencies' are not exactly the same as those generally refered to in the literature, especially in the literature on imperative programs, such as, Miranker and Winkler (1983), Ferrante *et al.* (1987) and Kennedy and McKinley (1990). We proceed from the algebraic, that is, functional descriptions and dependencies relate functional terms and not program statements. In particular, a variable is only a (sub)term denoting a specific value – it is not an actual program variable of a program capable of storing various values at different points of computation.

We mainly address the semantic issues of this approach, without entering into a discussion of possible specification *formalisms*. Thus we will assume that data dependencies are given along with algebras, and study their interaction. Since data dependencies are standard algebraic objects, we expect that they can easily be described in a standard algebraic fashion. (We intentionally refrain from including an analysis of control dependencies – unlike data dependencies, they seem to be inherent features of actual programs rather than of abstract data types.) In some cases data dependencies may be recovered directly from an algebraic specification and Section 3 describes a way this can be done.

By allowing computational structures into the semantics of a specification, we provide the means not only for the development of programs, but for the development of *efficient* programs. We will show that data dependencies provide such a means, enabling one to assess the time/space resources required and to pursue optimization strategies such as syntactic and semantic memoisation. Above all, the computational structures carry the information necessary to achieve *parallel* implementations: the identification of dependencies among various parts of a computation enables us to determine its possible parallel distributions. In addition to algebras and data dependencies, we will introduce a representation of (parallel) machine architectures and show how mappings from dependencies to architectures may be utilized to yield parallel implementations. The framework enables us then to *port the implementations of computation algebras* along the morphisms between various hardware architectures. It should be emphasized, however, that we are *not* concerned with a general theory of concurrency. In this context, we view concurrency merely as a means of increasing the efficiency of the implementation.

Section 2 introduces the basic notions of computation algebras and their implementations, and illustrates them on a simple example. It also gives a general view of the envisaged development process using computation algebras. Section 3 addresses the issue of constructing dependencies for a function defined by a set of recurrence equations over a given algebra. It illustrates the ideas of the programming language SAPPHIRE developed at the University of Bergen for which computation algebras provide the semantic framework.

Section 4 refines the picture of development sketched in Section 2 – its results suggest a more specific metodology for development using computation algebras. Section 5 shows how non-determinism can be included into the proposed framework. Section 6 summarizes the results and indicates some directions for future work.

The paper focuses on the description of the *concepts*; its technical content is limited to definitions and a few results. It gives an improved version of the results from Walicki *et al.* (1996). The more technical aspects assume some elementary knowledge of the category theory (for example, Mac Lane (1971), Rydeheard and Burstall (1988) and Barr and Wells (1990)), with particular emphasis on fibrations.

2. The framework of computation algebras

This section introduces the framework of computation algebras. We define first some basic notions in Section 2.1, then data dependencies in Section 2.2, computation algebras in Section 2.3, and their implementation in Section 2.4. Section 2.5 summarizes these pieces presenting the view of the development process using computation algebras.

2.1. Signatures, algebras and graphs

We use the standard notion of a signature Σ as a pair $\langle \mathcal{S}, \mathcal{F} \rangle$ of sort and operation symbols. In Section 4.1 we will use the category Sig with signatures as objects and *injective* signature morphisms.

For a signature Σ , Alg(Σ) will denote a category of Σ -algebras. It is 'a category' because we do not make any assumptions as to what the algebras are, except that they are objects where one can interpret the Σ -terms. One may think of standard Σ -algebras with Σ -homomorphisms defined in the usual way. But one may as well think of the category of multiagebras with various multihomomorphisms, power algebras, unified algebras, *etc.* We will define various notions relatively to an arbitrary full subcategory $A \subseteq Alg(\Sigma)^{\dagger}$.

The primary objects of our interest will be simple directed acyclic graphs that

- have at most one edge between any pair of nodes, and
- have no isolated nodes (each node has at least one incident edge).

We call them *simple DAGs*. We will, however, need a more general notion of a mapping of such graphs than a simple graph homomorphism. Mapping an edge of the source graph to a *path* in the target graph corresponds to splitting the computation into several

[†] We hope that our use of a different font for categories means that our use of the symbol \subseteq for subcategory as well as subset will not create any confusion.

steps. On the other hand, mapping an edge to a *set* of edges (or paths) corresponds to distributing the computation.

To define this concept of a morphism, we introduce three categories: DAG, of DAGs; DAG°, of path closures; and DAG[⊕], of distributed paths (or just sets of paths). The latter two categories are used merely for the purpose of defining more general morphisms between DAG-objects. An object in DAG° also contains, in addition to the underlying graph G, all the paths over G – we will want to map an edge of one graph onto a path in a target graph[†]. An object in DAG[⊕] contains, in addition, 'parallel composition' of paths which, too, will be possible images of edges. The construction described in the rest of this subsection starts with the category DAG: the free functor F° adds freely to each object all the paths and then, the free functor F^+ adds all the sets of paths. The resulting composition $F^{\oplus} = F^\circ$; F^+ has left adjoint and our morphisms are Kleisli morphisms obtained from this adjunction.

Definition 2.1. DAG is the category of *directed acyclic graphs* where:

1 Objects are DAGs – algebras over signature $\Gamma = \langle I, E; s, t : E \rightarrow I \rangle$, where

- I is a set of nodes and E a set of edges;
- s/t return the source/target node of an edge;
- *E* viewed as a relation ($\subseteq I \times I$) is acyclic;
- there are no isolated nodes: for each node v, there is an edge e with s(e) = v or t(e) = v.

2 Morphisms are Γ -homomorphisms.

We will often write $i \rightarrow j$ to indicate the existence of an edge *e* from *i* to *j* (that is, with s(e) = i and t(e) = j). A Γ -algebra is a DAG (an object of DAG) if the transitive closure \rightarrow^+ of this relation is irreflexive.

A DAG may have several edges between a pair of nodes. Of particular importance to us will be *simple* DAGs – the ones with at most one edge between a pair of nodes.

Definition 2.2. A simple DAG is a DAG satisfying: $s(e_1) = s(e_2) \wedge t(e_1) = t(e_2) \Rightarrow e_1 = e_2$.

Definition 2.3. DAG° is the category of *path closures* of DAGs where:

- 1 Objects are algebras over signature $\Gamma^{\circ} = \Gamma \cup \{ \circ : E \times E \to E \}$ whose Γ -reducts belong to DAG for any $P \in Obj(DAG^{\circ}) : P|_{\Gamma} \in Obj(DAG)$. $_{-}\circ_{-}$ is a partial operation of path formation defined for any edges e_1, e_2 with $t(e_1) = s(e_2)$, and satisfying
 - $e_1 \circ (e_2 \circ e_3) = (e_1 \circ e_2) \circ e_3$
 - $s(e_1 \circ e_2) = s(e_1)$ and $t(e_1 \circ e_2) = t(e_2)$.

2 Morphisms are Γ° -homomorphisms.

Thus, a DAG^o-morphism $f : G \to H$ maps an edge $x \to y$ from G onto en edge from f(x) to f(y) in H – this latter edge, however, may correspond to a path obtained by composition of several edges in H. The following fact is easy to verify.

[†] The symbol 'o' is used for (sequential) composition in diagrammatic order – not for functional composition.

Proposition 2.4. Let

- F° : DAG \rightarrow DAG° be the free functor sending a $G \in Obj(DAG)$ onto its *path closure* G° by adding freely all possible compositions of edges from G; for a DAG-morphisms h, we let $F^{\circ}(h) = h^{\dagger}$.
- U° : DAG[°] → DAG be the forgetful functor sending a graph G° onto its Γ-reduct $G^\circ|_{\Gamma}$; for a DAG[°]-morphisms $h : U^\circ(h) = h$.

There is an adjunction $F^{\circ} \dashv U^{\circ}$ with the unit being the inclusion of the source graph into its path closure.

The objects in the next category will allow two nodes to be connected not only by paths but also by all finite, non-empty sets of paths.

Definition 2.5. DAG^{\oplus} is the category of *distributed* path closures of DAGs where:

1 Objects are algebras over signature $\Gamma^{\oplus} = \Gamma^{\circ} \cup \{ - \oplus - : E \times E \to E \}$ whose Γ° -reducts belong to DAG° – for any $P \in Obj(\mathsf{DAG}^{\oplus}) : P|_{\Gamma^{\circ}} \in Obj(\mathsf{DAG}^{\circ})$. Where $_{-} \oplus _{-}$ is a partial operation of parallel composition defined for any edges e_1, e_2 with $s(e_1) = s(e_2)$ and $t(e_1) = t(e_2)$, which is associative, commutative and idempotent:

$$- e_1 \oplus (e_2 \oplus e_3) = (e_1 \oplus e_2) \oplus e_3$$

$$- e_1 \oplus e_2 = e_2 \oplus e_1$$

$$- e \oplus e = e,$$

and composition distributes over \oplus :

 $- e_1 \circ (e_2 \oplus e_3) = (e_1 \circ e_2) \oplus (e_1 \circ e_3)$

$$- (e_1 \oplus e_2) \circ e_3 = (e_1 \circ e_3) \oplus (e_2 \circ e_3)$$

and, since \oplus is defined only for edges with the same source and target,

- $s(e_1 \oplus e_2) = s(e_1) = s(e_2)$ and $t(e_1 \oplus e_2) = t(e_1) = t(e_2)$.

2 Morphisms are Γ^{\oplus} -homomorphisms

The first three axioms allow formation of sets of edges (which here may be paths from the 'underlying' DAG); the next two ensure that composition of sets of paths corresponds to forming sets of composite paths. Canonically, an edge $x \rightarrow y$ is a non-empty set of edges (paths in the 'underlying' DAG) from x to y. Hence, a morphism $f : G \rightarrow H$ maps an edge $x \rightarrow y$ from G onto a non-empty set of paths between f(x) and f(y) in H. Again, we have an easy fact in the following proposition.

Proposition 2.6. Let:

- F^+ : DAG[°] → DAG[⊕] be the free functor sending a $G^\circ \in Obj(DAG^\circ)$ onto $G^\oplus \in Obj(DAG^\oplus)$ by adding freely all possible parallel (\oplus) compositions of edges from G° subject to the restrictions and axioms from the above Definition;
- U^+ : DAG^{\oplus} \rightarrow DAG^{\circ} be the forgetful functor sending a graph G^{\oplus} onto its Γ° -reduct $G^{\oplus}|_{\Gamma^{\circ}}$;

[†] Strictly speaking, $F^{\circ}(h)$ is h extended to paths: $F^{\circ}(h)(e_1 \circ e_2) = h(e_1) \circ h(e_2)$.



Fig. 1. A Gr-morphism from D to W.

 Both functors map morphisms onto identical morphisms in their respective target categories.

There is an adjunction $F^+ \dashv U^+$ with the unit being the inclusion of the source graph into its closure under parallel composition.

Now define two functors

$$F^{\oplus} = F^{\circ}; F^{+} : \mathsf{DAG} \to \mathsf{DAG}^{\oplus} \text{ and } U^{\oplus} = U^{+}; U^{\circ} : \mathsf{DAG}^{\oplus} \to \mathsf{DAG}.$$
 (1)

The two adjunctions from Propositions 2.4 and 2.6 give an adjunction $F^{\oplus} \dashv U^{\oplus}$. The morphisms we are looking for can be now defined as Kleisli morphisms between objects of DAG induced by this last adjunction. The resulting category is given by: the objects are simple DAGs, a morphism $f : G \to H$ is a DAG-morphism $f : G \to U^{\oplus}(F^{\oplus}(H))$. Morphisms are composed pointwise: for $f : A \to B$ and $g : B \to C$, f maps an edge $e = x \xrightarrow{A} y$ onto a set of paths $f(e) = \{p_1, p_2...p_n\}$ from f(x) to f(y) in B. Then, g maps each path p_i onto a set of paths $g(p_i) = \{r_{i1}, r_{i2}...r_{ik}\}$ from g(f(x)) to g(f(y)) in C, obtained by mapping each edge of p_i onto a set of paths and composing these paths along the p_i . The composition is then $g(f(e)) = \bigcup_{p_i \in f(e)} g(p_i)$. Formally, we have the following definition.

Definition 2.7. The category Gr is the full subcategory – with simple DAGs as objects – of the Kleisli category induced by the adjunction $F^{\oplus} \dashv U^{\oplus \dagger}$.

Since we will be working with the category Gr, in the following, we will use the notation G^{\oplus} – where $G \in Obj(Gr)$ – as an abbreviation for $U^{\oplus}(F^{\oplus}(G))$.

Example 2.8. Three examples of Gr-morphisms are given in Figures 1 and 2. The former maps edges onto paths while the latter maps an edge onto distributed paths.

A Gr-morphism $h : A \rightarrow B$ expresses the fact that B can simulate A – edges of A are not necessarily represented in B as edges, but possibly as paths, or even sets of paths. B may also contain additional edges not in the image of h.

Remark. DAG-embedding of A into B is a special case of a Gr-morphism $A \rightarrow B$. One might expect that a surjective (on the nodes) DAG-morphism $B \rightarrow A$ would yield a simpler

602

[†] This concise formulation expresses what we have described above. For the construction of Kleisli category, see, for example, Mac Lane (1971).



Fig. 2. Two Gr-morphisms from G to H.

notion. However, it would require our graphs to be reflexive (this is not a big problem) and, more importantly, would exclude many cases where B is a richer structure than A. For instance, there is no surjective DAG-morphism from

$$B = \bullet \swarrow$$

onto $A = \bullet \rightarrow \bullet \rightarrow \bullet$. But we would like to say that B does reflect the edges of A (and introduces some more) – and so there is a Gr-morphism $A \rightarrow B$.

From now on, we will treat 'graph' as a synonym for 'DAG' since DAGs are the only graphs we are going to deal with. Also, our graphs are primarily connected *simple* DAGs, in which case we will often identify a graph G and its edge relation E^G , written also $\stackrel{G}{\rightarrow}$. Then, we say that G is *well-founded* when the relation $\stackrel{G}{\rightarrow}$ is, and denote by G^+ the transitive closure $\stackrel{G}{\rightarrow}^+$. Notice that, in general, $G^\circ \neq G^+ \neq G^\oplus$. The following example (suggested by an anonymous referee) illustrates the difference between the three:



Occasionally, we may use the following terms:

- A graph is sequential if \rightarrow ⁺ is a total ordering of the nodes.
- A partitioning of a graph is a partitioning of the set of nodes $I = \bigcup_{k=1}^{k=n} I_k$ with $1 < n < \omega$ such that each I_k is sequential.
- An *st-graph* ('space-time') is a simple DAG with nodes indexed by pairs $(s, t) \in S \times \mathbb{N}$, such that:

M. Walicki, M. Haveraaen and S. Meldal

 $\begin{array}{l} 1 \ \forall s,t : \langle s,t \rangle \mapsto \langle s,t+1 \rangle \ ; \\ 2 \ \forall s,s',t,t' : \langle s,t \rangle \mapsto \langle s',t' \rangle \Rightarrow t' = t+1 \ ; \\ 3 \ \exists k \in \mathbb{N}_+ : \forall s,s',t : \langle s,t \rangle \mapsto \langle s',t+1 \rangle \Leftrightarrow \langle s,t+k \rangle \mapsto \langle s',t+k+1 \rangle. \end{array}$

The *st*-graphs are singled out because they will be used as the space-time representation of hardware architectures (Miranker and Winkler 1983). The *s* component identifies a processor and the *t* component a time-point during a computation. Edges represent possible communications. The first condition says that each processor performs a sequential computation with the possibility of maintaining its local data from one step of the computation to the next one. The second condition requires all elementary communications to happen in a single time-step. The last condition says that the communication topology is fixed throughout the whole computation, that is, the graph is actually a repetition of the pattern of the *k*-steps communications (for some fixed k > 0).

We note the following characterization.

Proposition 2.9. Let $k : A \to B \in Mor(Gr)$. k is mono $\Leftrightarrow k$ is injective on the edges.

Proof. Let T be an arbitrary object and $f, g: T \to A$ be two arbitrary morphisms.

- (⇒) If k is mono, then for any $f,g: T \to A$, $f; k = g; k \Rightarrow f = g$. In particular, for any edge $e \in E^T$: $k(f(e)) = k(g(e)) \Rightarrow f(e) = g(e)$. Taking T as a one-edge graph and considering all $f,g: T \to A$, implies that k must be injective on the edges.
- (\Leftarrow) Suppose k is injective on the edges and f; k = g; k. For each edge $e \in E^T$ we then have $k(f(e)) = k(g(e)) \Rightarrow f(e) = g(e)$. To verify that f = g, we have to check that also for each node i in T, f(i) = g(i). Assume that for some node $f(i) \neq g(i)$. Since T is a DAG (according to Definition 2.1 it has no isolated nodes), there is an edge e incident to i and then we would have $f(e) \neq g(e)$. Because of the injectivity on the edges, we then get $k(f(e)) \neq k(g(e))$, contrary to the assumption f; k = g; k.

2.2. Data dependencies

As remarked in the introduction, our data dependencies are intended to capture the information concerning what resources are going to be used in performing new steps of computation. Resources in this context refer exclusively to the values of some functions and the computations that compute such values. Thus our data dependencies relate functional terms and not program statements. For instance, computation of the value of the term f(s(x), 2) may depend on the value of s(x), which, in turn, may depend on x. However, x is just a term – a mathematical variable that, in any instance, has some specific value. It never depends on itself, as it might do if it was a program variable occurring in a statement x := x + 1.

For a signature Σ , a Σ data dependency is a graph (simple DAG) with nodes labelled by ground Σ -terms \mathscr{T}_{Σ} . The only requirement we put on the morphisms between such graphs is that they respect the sorting of labels.

Definition 2.10. (Data dependencies) $DD(\Sigma)$ is the category of Σ data dependencies where:

1 Objects are Σ data dependencies – pairs $\langle G, lab \rangle$ where $G \in Obj(Gr)$ is a simple DAG and $lab : I^G \to \mathscr{T}_{\Sigma}$ is a function labelling nodes of this graph with ground Σ -terms.

2 A DD(Σ)-morphism $m : \mathbf{C} = \langle C, lab_{\mathbf{C}} \rangle \rightarrow \mathbf{D} = \langle D, lab_{\mathbf{D}} \rangle$ is a Gr-morphism between the underlying graphs $m : \mathbf{C} \rightarrow D$, such that $\forall i \in I^{C} : Sort(lab_{\mathbf{D}}(m(i))) = Sort(lab_{\mathbf{C}}(i))^{\dagger}$.

In the following, D will denote an arbitrary subcategory of $DD(\Sigma)$. The functor

$$Gr: \mathsf{DD}(\Sigma) \to \mathsf{Gr}$$
 (2)

is the obvious forgetful functor, which, for a given Σ data dependency $\mathbf{D} = \langle D, lab \rangle$, returns its underlying graph D. We call such a graph the *shape of* **D**.

The relation $i \mapsto j$ (in the shape $Gr(\mathbf{D})$ of $\mathbf{D} = \langle D, lab \rangle$) indicates that the computation at node *j* depends on the computation at node *i*. If lab(i) = s and lab(j) = t, this means that the result of the evaluation of *t* at *j* depends on the result of the evaluation of *s* at *i*.

Example 2.11. To give a flavour of our aims, consider a specification of the Fibonacci function:

Fib is \mathscr{S} : Nat

 $\mathcal{F}: \dots \text{ the usual functions for } Nat$ $F: Nat \rightarrow Nat$ $\mathcal{A}: F(0) = 1$ F(1) = 1 F(n+2) = F(n) + F(n+1)

Ignoring the details for the moment (we will discuss them later), we may easily recognize the dependency given by the tree of recursive calls **R** in Figure 3. Certainly, Definition 2.10 allows us to map this tree, for instance, onto a linear order (respecting the ordering in the tree and avoiding loops). In the following subsection we will couple dependencies with algebras and thus specify the means of avoiding uninteresting morphisms that are admitted, in principle, by Definition 2.10. In Figure 3 we have indicated two dependency morphisms: d' corresponding to syntactic memoisation (identifying the nodes with identical labels), and d'' corresponding to semantic memoisation (identifying F(0) with F(1)) given the above specification[‡].

Here, we may consider the dependency **D** as a refinement (improvement) of **R**. Notice that **D** is sequential under the ordering F(n) < F(m) for n < m. Nevertheless, the partitioning of **D** indicated by the vertical arrows implies serious savings of storage space, even if the computations along both lines are heavily synchronized and far from independent.

Remark. As a matter of fact, there is also a dependency morphism $g : \mathbf{D} \to \mathbf{R}$ that embeds \mathbf{D} into the leftmost branch of \mathbf{R}^{\oplus} . This illustrates the fact that refinement of data dependencies is not necessarily an 'improvement' in the narrow sense of memoisation – it rather reflects the relation of one dependency being able to perform the computations of which another one is capable.

[†] For a signature $\Sigma = \langle \mathscr{S}, \mathscr{F} \rangle$, the function *Sort* : $\mathscr{T}_{\Sigma,X} \to \mathscr{S}$ returns the sort of the argument term.

^{\ddagger} In Section 4.2.1, we suggest a more specific notion of a dependency morphism *compatible* with a given class of algebras of which d'' is an example.



Fig. 3. The recursive tree dependency **R**, together with the 'actual' dependencies for computation of Fibonacci, **D**' and **D**, and the dependency morphism d = d'; d''.

A Σ -computation algebra is a Σ algebra with an associated Σ -dependency. The latter provides a clue for how various operations of the algebra are to be computed. In general, for $A \subseteq Alg(\Sigma)$, $D \subseteq DD(\Sigma)$, an (A, D)-computation algebra is an A-algebra with an associated D-dependency.

Definition 2.12 (Computation algebra). An (A, D)-computation algebra C is a triple $\langle A, B, ev \rangle$ where

1 $A \in Obj(A)$ 2 $\mathbf{B} \in Obj(D)$ 3 $ev : I^{Gr(\mathbf{B})} \to A$ is an evaluation function such that $lab_{\mathbf{B}}(i) = t \Rightarrow ev(i) = t^{A}$.

The *ev* function is uniquely determined by the algebra A and the labelling of **B**, so we will usually write a computation algebra as a pair $\langle A, \mathbf{B} \rangle$. When (A, D) does not matter or is clear from the context, we will simply write *computation algebra* (instead of (A, D)-computation algebra).

Dependencies may (and will) be interpreted as the requirements on the communications between different space-time points on an actual architecture. Consequently, we will refer to the dependency part **B** of a computation algebra $\langle A, \mathbf{B} \rangle$ as its *communication* part.

Definition 2.13 (Computation homomorphisms). An (A, D)-computation homomorphism m: $\langle A, A, ev_A \rangle \rightarrow \langle B, B, ev_B \rangle$ is a pair $\langle h, g \rangle$ such that (*cf.* Figure 4):

1 $h: A \to B \in Mor(A)$; 2 $g: \mathbf{B} \to \mathbf{A} \in Mor(D)$, and 3 $h(ev_A(g(i))) = ev_B(i)$, for all $i \in I^{Gr(\mathbf{B})^{\dagger}}$.

As with computation algebras, when (A, D) does not matter, or is clear from the context, we will speak about *computation homomorphisms*.

The primary intuition behind this definition of homomorphism is that it is to be used to model a refinement process – in the above definition, we would say that $\langle A, \mathbf{A} \rangle$

606

[†] The diagram in Figure 4 commutes, but not in a category (at least none we have defined explicitly), and so we write the composition of functions explicitly as a(b(i)).



Fig. 4. A homomorphism between two computation algebras.

refines $\langle B, \mathbf{B} \rangle$. Refinement concerns both the algebra and dependency part. The usual algebra homomorphism $h : A \to B$ corresponds to the classical notion of data refinement – A may have multiple representations for a single value from B. The (contravariant) homomorphism $g : \mathbf{B} \to \mathbf{A}$ represents the possible improvement of the dependency **B** by the dependency **A** (*cf.* Figure 3) – **A** may introduce paths, even sets of paths, in the place of single edges in **B**.

In Example 2.11, the standard algebra of natural numbers \mathcal{N} with function F, may give rise to three different computation algebras $\langle \mathcal{N}, \mathbf{R} \rangle$, $\langle \mathcal{N}, \mathbf{D}' \rangle$ and $\langle \mathcal{N}, \mathbf{D} \rangle$. The respective morphisms obtained from Figure 3, for example, $\langle id_{\mathcal{N}}, d \rangle : \langle \mathcal{N}, \mathbf{D} \rangle \rightarrow \langle \mathcal{N}, \mathbf{R} \rangle$, will represent gradual refinements of $\langle \mathcal{N}, \mathbf{R} \rangle$. On the other hand, a dependency morphism that sends, for instance, a node labelled with F(2) to a node labelled with F(4) will not yield a computation homomorphism unless algebra \mathcal{N} satisfies some additional (and here, unintended) equations.

Thus, refinement corresponds to a standard (data) refinement on the algebra part, while dependencies may be refined by 'stretching' (to paths), 'distributing' (to sets of paths) or augmenting with new dependencies. The following two examples illustrate these aspects of refinement.

Example 2.14. We let \vec{C} and \vec{D} be as in Figure 5, with $\vec{h}(a) = a$, $\vec{h}(b) = d$, and $\vec{h}(f) = f$. The commutativity requirement, $\bar{h}(ev_C(\vec{h}(i))) = ev_D(i)$, restricts the legal combinations of \vec{h} and \bar{h} . In particular, the labels in \vec{C} may differ from those in \vec{D} like with $\vec{h}(b) = d$ (taking, for simplicity, the names of the nodes in the figure to be their labels), only if d evaluates in \vec{C} to a value $ev_C(d) = x$ representing $ev_D(b)$, that is, such that $\bar{h}(x) = ev_D(b)$.

Also, refinement 'does not care' about what is computed in C at the nodes that are not in the image of $\vec{h}(\vec{Gr}(D))$. There may be new nodes and dependencies, also on nodes labelled by terms not occurring in the labelling of \vec{D} (like $c \rightarrow f$ in \vec{C}). In particular, if we admit a computation algebra with an empty dependency part (that is, empty carriers in the category D), a standard algebra \bar{A} may be considered a computation algebra with empty communication part, $A = \langle \bar{A}, \emptyset \rangle$. Then any algebra B with non-empty communication part will be a refinement of A provided there is a homomorphism $\bar{h}: \bar{B} \rightarrow \bar{A}$.

Example 2.15. As an example of a possible 'distribution', consider an algebra D with the sorts S, S_1, S_2 , and the dependency $t \rightarrow f(t)$ for two terms of sort S (Figure 6). Furthermore, let the sort S be the Cartesian product of the two other sorts $S = S_1 \times S_2$, with the obvious projections $\pi_i : S \rightarrow S_i$ and pairing function $\langle -, - \rangle : S_1 \times S_2 \rightarrow S$. Let \overline{D} satisfy the ordinary equalities for these functions and, in addition, let $\overline{D} \models t = \langle t_1, t_2 \rangle$, and $\overline{D} \models f(t) = \langle f_1(t_1), f_2(t_2) \rangle$. This allows us to design the refinement of the dependency



Fig. 5. Refinement of a computation algebra by adding new dependencies.



Fig. 6. Refinement of a computation algebra by distributing dependencies.

structure illustrated in Figure 6 (cf. Figure 2), which, together with the identity on \overline{D} , yields a computation homomorphism $\langle id_{\overline{D}}, \vec{h} \rangle$. The morphism \vec{h} refines the dependency $t \rightarrow f(t)$ in \vec{D} to both paths (through t_1 and t_2).

Computation algebras and their homomorphisms form a category.

Definition 2.16. (Category of computation algebras) CAlg(A, D) is the category with objects being (A, D)-algebras and morphisms being (A, D)-homomorphisms (and obvious pointwise composition). We will write CAlg(Σ) instead of When referring to the entire category CAlg(Alg(Σ), DD(Σ)), we will write CAlg(Σ) instead.

We define two obvious functors:

$$\overrightarrow{(-)} : \mathsf{CAlg}(\mathsf{A},\mathsf{D})^{op} \to \mathsf{D}$$
(3)

projects computation algebras onto their communication part, that is, $\langle \vec{A}, \vec{A} \rangle = \mathbf{A}$ and $\langle \vec{h}, g \rangle = g$ with the notation from Definition 2.13[†]. Similarly,

$$\overline{(_)} : \mathsf{CAlg}(\mathsf{A},\mathsf{D}) \to \mathsf{A} \tag{4}$$

projects computation algebras onto their algebra part ($\langle A, A \rangle = A$ and $\langle h, g \rangle = h$).

[†] Of course, each (A, D) has its own functor $(\overrightarrow{-})_{(A,D)}$, but we may as well view each such functor as a restriction of $(\overrightarrow{-})_{CAlg(\Sigma)}$ to the respective source subcategory.

We will now use these functors to simplify the notation for the components of such an algebra: a computation algebra A will denote the triple $\langle \bar{A}, \bar{A}, ev_A \rangle$, and a computation homomorphism h a pair $\langle \bar{h}, \bar{h} \rangle$.

2.4. Machines and distribution

We introduce the model of machine architectures and their mappings (simulations) in Section 2.4.1, and then define mappings of a computation algebra to a given architecture called 'distributions' in Section 2.4.2.

2.4.1. *Machine architectures* In order to relate the dependencies to hardware architectures, we will consider the latter as space-time unfoldings of the actual inter-processor communication structure (Miranker and Winkler 1983). Thus, an architecture is simply an *st*-graph, where each node represents a unique time-point at a given processor and edges represent the possible single-step communications. This perspective may be extended to cover dynamically changing architectures as well (it would merely require graphs rather than the *st*-graphs), but here we restrict our attention to the static case.

A morphism between two architectures expresses the possibility of using one architecture, Z, to *simulate* another, W, that is, that each communication of which W is capable can be simulated by a set of chains of communications in Z. This is captured by the existence of a mapping of W into the (possibly distributed) path closure of Z, that is, a Gr-morphism $W \rightarrow Z$. Consequently, we have the following definition.

Definition 2.17. (Machine architectures) The category of machine architectures is Gr.

Thus, in a way similar to the dependency morphisms, an architecture morphism $W \rightarrow Z$ tells us that, and how, Z can simulate communications of W.

We might have given a more restrictive definition of machine architectures. For instance, the unfoldings of actual processor-architectures will be st-graphs that are simply repeating patterns of one-step communications. Also, simulation morphisms might be restricted to Gr-monomorphisms. Nevertheless, we choose this more generous definition because it will result in a more uniform treatment – it will allow us to consider also shapes of data dependencies as objects, and distributions as morphisms of the same category Gr.

Example 2.18. An *n*-dimensional vector W has *n* processors connected so that each processor, except for W_1 and W_n , has two-way communication channels to its left and right neighbour: $W_1 \rightleftharpoons W_2 \rightleftharpoons \cdots \rightleftharpoons W_n$. A space-time unfolding for a 2-dimensional vector is illustrated in Figure 7 (a). The vertical arrows $(W_i, k) \mapsto (W_i, k+1)$ are added to represent the fact that each processor can carry its local data from one computation step to the next.

A ring Z with 3 processors where, in each step, processor Z_1 can communicate to Z_2 , Z_2 to Z_3 , and Z_3 to Z_1 is given in Figure 7 (b). This architecture may simulate the 2-dimensional vector W by the morphism m indicated in Figure 7 (c).



Fig. 7. Space-time representations of (a) 2-dimensional vector W, (b) 3-processor ring Z, and (c) a simulation m of W by Z.

2.4.2. Distribution Assume that we would like to implement a computation algebra on some specific machine architecture. Distribution is what, in our abstract setting, represents an implementation. It certainly does not capture all aspects of an actual implementation, and for this reason we have chosen to call it 'distribution'. This term reflects the intuition of distributing the dependencies of an algebra onto the communication structure of a given architecture.

The composition of functors (2) and (3) (\vec{L}) ; $Gr : CAlg(A, D)^{op} \to D \to Gr$ applied to an algebra returns the underlying graph of its communication part. We will write this composition as \vec{Gr} , that is, $\vec{Gr}(X) = Gr(\vec{X})$, for an object or morphism X in CAlg(Σ).

Distribution of an algebra $\langle \bar{C}, \bar{C} \rangle$ on a given architecture W amounts to a simulation of the (shape of the) dependency structure given by $Gr(\vec{C})$ on the communication structure W.

Definition 2.19. (Distribution) A distribution $m_{C \to W}$ of a computation algebra C on an architecture W is a Gr-morphism $m : \vec{Gr}(C) \to W$.

Example 2.20. The dependency **D** (with the shape $Gr(\mathbf{D})$ from Figure 3) is mapped onto a 2-processor vector W as shown in Example 2.8, Figure 1. Figure 8 is just a repetition of this earlier example: (a) is the data dependency **D** with the shape $Gr(\mathbf{D})$, (b) is a 2-processor vector W, and (c) is a distribution f of **D** on W.

Combining such distribution morphisms with the simulation morphisms between various architectures, leads to the possibility of porting the distributions.

Example 2.21. The mapping f from Figure 8 (c) is a distribution of a computation algebra $D = \langle N, \mathbf{D} \rangle$ (with N a Σ_{FIB} -algebra satisfying FIB) on the vector W. Then, the mapping m (Figure 7 (c)) gives a ported distribution of D on the ring Z. The results are shown in Figure 9.

A morphism between two distributions is defined as follows.

Definition 2.22. (Morphisms of distributions) A morphism between two (A, D) distributions, $m: f_{A \to W} \to g_{B \to Z}$, is a pair $\langle h, h' \rangle$, where:



Fig. 8. (a) Data dependency \mathbf{D} , (b) space-time representation of a 2-dimensional vector W, and (c) a distribution f of \mathbf{D} on W.



Fig. 9. (a) Data dependency **D**, (b) the distribution f of **D** on a 2-dimensional vector W, and (c) distribution f ported to the ring Z.

1 $h: A \rightarrow B \in Mor(CAlg(A, D))$; 2 $h': Z \rightarrow W \in Mor(Gr)$; and 3 $\tilde{Gr}(h); f = g; h'.$

The situation is depicted in Figure 10. The third condition is the requirement of the commutativity of the rightmost square in Gr^{\dagger} .

Definition 2.23. We let Dist(A, D) be the category

- 1 Objects distributions of (A, D) computation algebras (Definition 2.19),
- 2 Morphisms given in Definition 2.22 and pointwise composition[‡].

611

[†] We let the distribution morphisms go in the opposite direction to the respective Gr-morphisms between the distributions. This choice is to keep with the convention that morphisms between structures reflect some form of 'refinement' of the target by the source.

[‡] This works because the commutativity of the square in Figure 10 – that is, for each edge e in $\vec{Gr}(B)$, the (sets of) paths obtained in W by h'(g(e)) and by $f(\vec{Gr}(h)(e))$ are identical – implies commutativity for each (set of) paths in $\vec{Gr}(B)$. Hence composition with another such commuting square, given by $\vec{Gr}(n) : \vec{Gr}(C) \to \vec{Gr}(B)$, $n' : Y \to Z$, and $k : Gr(\vec{C}) \to Y$, will yield a commuting square $(\vec{Gr}(n); \vec{Gr}(h)); f = k; (n'; h')$.



Fig. 10. A morphism $\langle h, h' \rangle$ of distributions.

The forgetful functor

$$Arch: \text{Dist}(A, D) \to \text{Gr}^{op} \tag{5}$$

is the obvious projection of distributions onto their architecture part: in other words $Arch(f_{A \to W}) = W$ and $Arch(\langle h, h' \rangle) = h'$. (Again, we may view each $Arch_{\text{Dist}(A,D)}$ as a restriction of $Arch_{\text{Dist}(\text{CAlg}(\Sigma))}$ to Dist(CAlg(A,D)).)

2.5. The development process

The framework we have introduced is motivated by the view of the development of (possibly parallel) implementations by means of computation algebras. We envision the process starting with the usual refinement of algebraic specifications that proceeds through a series of subclasses of Alg(Σ) and results in one class of isomorphic algebras (an algebra \overline{A} in Figure 11 is a chosen representative from this resulting class). At this point, when the specification has become sufficiently concrete, one introduces the dependencies, that is, constructs a computation algebra $\langle \overline{B}, \overline{B} \rangle$. Dependencies can then be refined along with a further refinement of the algebra itself or, perhaps, with the algebra fixed.

Remark 2.24. Since there is no surjectivity requirement on ev, the communication part \vec{A} need not contain the full information about all operations of the algebra \bar{A} . (In the extreme case, the shape of \vec{A} might be the empty graph.) There are several reasons for the lack of such a requirement. First, it allows us to consider a standard algebra \bar{A} as a computation algebra with empty communication part. More importantly, it allows us to model the development process in which more and more operations of the algebra are gradually equipped with more detailed computational information. Furthermore, we may imagine a specification of computation algebra are of interest to the implementation, and the resulting computation algebra need only provide the dependency structure for these operations. Finally, we can use this setting in situations where (parts of) an algebra are implemented on some machine (for example, all built-in types constitute a given algebra), and one needs to compute some additional, newly defined functions – the communication part will then be needed only for these new functions.

When all the required operations of the algebra have been associated with the satisfactory dependencies, $\langle \bar{C}, \bar{C} \rangle$, one may choose an architecture on which to distribute the dependencies \bar{C} , $f_{C \to W}$. Once this is done, there remains only the question of portability, that is, of moving the distribution to other architectures.

612



Fig. 11. The development process.

One may imagine that a transition from one level (category) to the next means that the structure resulting from the development at the previous level is complete and remains fixed for the rest of the process. Thus, having reached the algebra \bar{B} , we begin to introduce and refine dependencies while \bar{B} remains unchanged (and so, \bar{C} is actually \bar{B}). Similarly, having designed $\langle \bar{C}, \bar{C} \rangle$ and distributing it on W, we may move to another architecture Z but $\langle \bar{C}, \bar{C} \rangle$ will be kept fixed (that is, $\langle \bar{D}, \bar{D} \rangle = \langle \bar{C}, \bar{C} \rangle$). Although this is probably the most natural scenario, Figure 11 illustrates the process in its full generality, where *all* the aspects may change at *all* levels.

Although the technical results below are rather straightforward and hardly surprising, they provide a sound justification for the above methodology as well as a clue as to how it may be applied. They yield a pleasing conceptual structure in which algebras can be thought of as indexed by various dependencies, and computation algebras by machine architectures. More importantly, they give us the possibility of modularizing the implementation process and reusing its results.

The simulation mappings between architectures (the Gr-morphisms) may be defined and stored independently of the actual programs implemented on them. Suppose that we have distributed a computation algebra D on an architecture W, $d_{D \to W}$ (Definition 2.19). In order to distribute D on another architecture Z, we need to define an appropriate distribution $c_{D \to Z}$. However, if we have a Gr-morphism $h: W \to Z$, it will tell us how Zcan simulate any computation on W. Thus it will allow us to port the distribution $d_{D \to W}$ by just using this general simulation h.

Such a reuse of (stored) simulation morphisms between architectures for porting distributions will be formalized by showing that the functor *Arch* from (5) yields a *fibration*. Intuitively, this means that we can view the category Dist(A, D) as classes of computation algebras indexed by machine architectures – the *fiber* over a given architecture W represents all computation algebras that can be distributed on W. Then, given an architecture morphism $h: W \to Z$ and an algebra D distributed on Z (in Z's fiber), there is a canonical way of defining a *cleavage*: an algebra C and its distribution $c_{C\to Z}$ on Z together with a distribution morphism $cl(h, D) : c_{C\to Z} \to d_{D\to W}$ that is 'compatible' with the simulation h. As a matter of fact, the triviality of the proof (that *Arch* is a fibration) coincides with the desired fact that D = C. Thus, cleavage gives explicitly the distribution resulting from porting a distribution of D from W to Z 'along' a given simulation h.

The concept of fibration seems a very natural model for porting distributions and we

will encounter it again later on. We now recall the notion of fibration, Definition 2.25, (for more on fibrations, see Bénabou (1985), Jacobs (1991) and Hermida (1993), and show that *Arch* is a fibration in Proposition 2.26.

Definition 2.25. (Fibration) Given two categories A and X and a functor $F : A \rightarrow X$:



- 1 A morphism $h : B \to C$ in A is *cartesian* if for any morphism $g : A \to C$ with F(g) = F(h), there exists a unique $f : A \to B$ such that f; h = g.
- 2 A functor F is a *fibration* if for every $C \in Obj(A)$ and $u : Y \to F(C) \in Mor(X)$, there exists a cartesian morphism $h : B \to C$, and the composite of two cartesian morphisms is cartesian.
- 3 *F* is a *cofibration* if $F^{op} : A^{op} \to X^{op}$ is a fibration; it is a *bifibration* if it is both fibration and a cofibration.
- 4 An $A \in Obj(A)$ is over Y if F(A) = Y; a morphism $g \in Mor(A)$ is over u if F(g) = u. For an object $Y \in Obj(X)$, the fiber over Y, $F^{-1}(Y)$, is the subcategory of A with objects all those over Y and morphisms all those over id_Y .
- 5 A particular choice of a cartesian lifting for every appropriate morphism $u \in Mor(X)$ is called a *cleavage* for *F*. Given a cleavage *cl*, a morphism $u : Y \to X \in Mor(X)$ and an object $C \in Obj(A)$ over *X*, we use cl(u, C) to denote the cartesian lifting of *u* whose codomain is *C*.

We will also define split fibrations:

- 6 Given a fibration F with a cleavage cl, any $u : Y \to X$ in X determines a reindexing functor $u^* : F^{-1}(X) \to F^{-1}(Y)$ as follows:
 - for an object C in $F^{-1}(X)$, $u^*(C)$ is the domain of cl(u, C)
 - for a morphism $f: D \to C$ in $F^{-1}(X)$, $u^*(f)$ is the unique morphism in $F^{-1}(Y)$ making $u^*(f)$; cl(u, C) = cl(u, D); f.
- 7 For every $Y \in Obj(X)$ there is a natural isomorphism $id_{F^{-1}(Y)} \rightrightarrows (id_Y)^*$, determined by the universal property of $cl(id_Y, A)$ for each $A \in Obj(A)$. For all $u : Y \to X$ and $w : X \to W$ there is a natural isomorphism w^* ; $u^* \rightrightarrows (u; w)^*$, determined by cl(u; w, A). If these isomorphisms are identities, the fibration is *split*.

Proposition 2.26. For any (A, D), *Arch* : Dist $(A, D) \rightarrow Gr^{op}$ is a (split) fibration.

Proof. The diagram to the left illustrates the fibration situation and the one to the right gives details for the present context (dashed arrows indicate the directions in the _^{op} categories):



For a Gr^{op}-morphism $h'^{op}: Z \to W$ and distribution $d = d_{p \to W}$, the cleavage is defined as

$$cl(h'^{op}, d_{D \to W}) \stackrel{\text{def}}{=} h : c_{D \to Z} \to d_{D \to W} \text{ where } \begin{cases} h = \langle id_D, h' \rangle \\ c_{D \to Z} = d; h'. \end{cases}$$
(6)

It is trivially a morphism in Mor(Dist(A, D)) since $id_{\tilde{Gr}(D)}$; c = d; h'.

Let $g : b_{B\to Z} \to d_{D\to W}$ be $\langle \langle \bar{g}, \bar{g} \rangle, h' \rangle$ – a distribution over h'^{op} . The unique $f : b \to c$ with $Arch(f) = id_Z$ making f; h = g is then $\langle \langle \bar{g}, \bar{g} \rangle, id_Z \rangle$. We have $f \in Mor(\text{Dist}(A, D))$ because $\bar{g}; b = d; h' = c; id_Z$, where the first equality holds since $g \in Mor(\text{Dist}(A, D))$.

Composition of cleavages is a cleavage: for $h': W \to Z, g': Z \to V$ and $d_{D \to W} \in Arch^{-1}(W)$ we have

$$cl(h'^{op}, d_{D \to W}) = \langle id_D, h' \rangle, cl(g'^{op}, (d; h')_{D \to Z}) = \langle id_D, g' \rangle,$$

so

$$cl(g'^{op},(d;h')_{D\to Z}); cl(h'^{op},d_{D\to W}) = \langle id_D,h';g' \rangle = cl(g'^{op};h'^{op},d_{D\to W}).$$

Hence Arch is a fibration.

It is split: we have $(id_W)^*(d_{D \to W}) = d_{D \to W}$, so the natural transformation

$$id_{Arch^{-1}(W)} \rightrightarrows (id_W)^*$$

is identity. Composition of cleavages gives a cleavage, so we get

$$(h'^{op})^*; (g'^{op})^* = (g'^{op}; h'^{op})^*.$$

Example 2.27. The collection of our examples so far illustrates this idea of development. In Example 2.11 we designed a dependency **R** that was then refined to **D** by the morphism $d : \mathbf{R} \to \mathbf{D}$ – assuming standard algebra for natural numbers \mathcal{N} . The dependency **D** was then distributed on a vector W by $f_{D \to W}$ (Figure 8) and ported to Z (Figure 9) using the simulation $m : W \to Z$ from Figure 7. Provided that the simulation m and distribution f were available from a library, all the work we had to do was to design the mapping d.



Fig. 12. The summary of the example.

3. Constructing dependencies for recursive functions

So far we have considered relationships between various aspects – functional specification (algebra), dependencies (communication part) and distribution – given independently from each other. In this section we show how – under certain conditions – dependencies can be derived from specifications using grammars. Although the described construction may indicate the possibilities of reconsidering implementation techniques for functional languages, it should be emphasized that we are here concerned only with *a particular example* of utilizing a computation algebra framework for designing efficient implementations of algebraic specifications.

We recall that a *context free grammar* G is given by a 4-tuple (T, N, \rightarrow, S) where T is a set of terminal symbols, N is a set of non-terminal symbols such that $N \cap T = \emptyset$, the set of production rules is $\rightarrow \subseteq N \times (T \cup N)^*$, and $S \in N$ is the start symbol. From the grammar G we get the *derives* relation $\Rightarrow \subseteq (N \cup T)^* \times (N \cup T)^*$ by $vXw \Rightarrow vuw$ for strings $v, w \in (N \cup T)^*$ and productions $X \rightarrow u$. The grammar defines the *language*, the set of strings, $\mathscr{L}(G) = \{w \in T^* | S \Rightarrow^* w\}$ where \Rightarrow^* is the reflexive, transitive closure of \Rightarrow .

In terms of Figure 11, we are at the transition point from A to CAlg(A, D): we have obtained a specific algebra A and introduce a new recursive function to be computed over A. For this new function, we construct the dependency relatively to the algebra A. The difference from the general setting described in Section 2.5 is that we want to compute a new function over a given algebra and not construct a dependency for the whole algebra (*cf.* Remark 2.24). This is quite a common scenario in practice and we will give a few examples.

Let Σ be a signature, $S_1...S_n$, S be some sorts from Σ , and $R : S_1 \times ... \times S_n \to S$ a new function symbol not in Σ . A generalized Σ -term recurrence for R is given by a set of equations of the form

$$R(\overline{\tau_0}(X)) = \Phi_{\overline{\tau_0}(X)}(X, R(\overline{\tau_1}(X)), R(\overline{\tau_2}(X)), \dots, R(\overline{\tau_z}(X)))$$
(7)

where X is a (sequence of) variable(s), for each $0 \le k \le z : \overline{\tau_k}(X)$ is a sequence of terms $t_{1,k}(X) \dots t_{n,k}(X)$ of appropriate sorts, that is, $t_{i,k}(X) \in \mathscr{T}_{\Sigma}(X)_{S_i}$, and $\Phi_{\overline{\tau_0}(X)}(X, y_1 \dots y_n) \in \mathscr{T}_{\Sigma}(X, y_1 \dots y_n)_S$ with all y_i of sort S. R is typically given by a set of such equations, that is, different equations for different (sequences of) term(s) $\overline{\tau_0}(X)$. (This form of generalized recurrence was studied in Haveraaen (1990), Haveraaen (1993), Čyras and Haveraaen (1995) and Haveraaen and Søreide (1998) as a definition schema for 'constructive recursive functions'.)

Given a recurrence (7) and a Σ -algebra A, we take as the set of nodes for the shape of the dependency, the domain of R in A, that is, the cartesian product $V = S_1^A \times ... \times S_n^A$.

We now construct a collection of context free grammars $G_{\Sigma,V}$, which will define the data dependency. As the non-terminal symbols of the grammar, we take the set $N = \{N_v : v \in V\}$ of distinct symbols indexed by the nodes $(N_u = N_v \Rightarrow u = v)$. Treating the non-terminals N_v as place-holders of type S, we may now define substitution rules from (7). For each variable assignment $\alpha : X \to A$, let $\alpha_i \in V$ denote the interpretation of $\overline{\tau_i}(X)$ under α , that is, $\alpha_i = \overline{\tau_i}^A(\alpha(X)) = \langle t_{1,i}^A(\alpha(X)), ..., t_{n,i}^A(\alpha(X)) \rangle$. For every expression form $\overline{\tau_0}$ (from the set of equations (7)) and every assignment $\alpha : X \to A$, we get the production

$$N_{\alpha_0} \longrightarrow_{G_{\Sigma,V}} \Phi^A_{\alpha_0}(\alpha(X), N_{\alpha_1}, N_{\alpha_2}, \dots, N_{\alpha_z})$$
(8)

Thus, for each node v, we obtain a node-specific grammar $G_{\Sigma,v} = \langle \mathscr{F}, N, \longrightarrow_{G_{\Sigma,v}}, N_v \rangle$, that is, where terminals are the function symbols \mathscr{F} from Σ , non-terminals are the N's, productions are given by (8), and the start symbol is N_v .

We define the shape of the dependency $\stackrel{G_{\Sigma,V}}{\longrightarrow}$ by introducing an edge for all pairs $u, v \in V$, according to the rule

$$u \xrightarrow{G_{\Sigma,V}} v \quad \Leftrightarrow \quad N_v \longrightarrow_{G_{\Sigma,V}} r N_u t, \text{ for some } r, t \text{ with } r N_u t \in \mathscr{F}_{\Sigma}(N).$$
(9)

This gives us a graph structure on V, and we can define a 'compatible' *labelling* function by requiring that

$$\forall v \in V : lab_{G_{\Sigma,V}}(v) \in \mathscr{L}(G_{\Sigma,v}) \tag{10}$$

where $\mathscr{L}(G_{\Sigma,v})$ is the language generated by the grammar $G_{\Sigma,v}$ of node v.

The result may not be a data dependency: the graph $\langle V, \stackrel{G_{\Sigma,V}}{\rightarrowtail} \rangle$ obtained from (9) may be cyclic, or the labelling (10) may not be unique problems often due to the recurrence (7) not being well-defined to begin with. The following conditions on $G_{\Sigma,V}$ ensure that the result will be a data dependency – a (simple) DAG with unique labelling.

Proposition 3.1. If a collection of node-specific grammars $G_{\Sigma,V}$ is such that:

1 for every non-terminal N_v there is exactly one rule $N_v \longrightarrow_{G_{\Sigma,V}} t$, and

2 the resulting graph $\Gamma_{G_{\Sigma,V}} = \langle V, \stackrel{G_{\Sigma,V}}{\rightarrowtail} \rangle$ is well founded,

then $G_{\Sigma,V}$ determines a unique Σ data dependency.

Proof. Point (2) implies that $\Gamma_{G_{\Sigma,V}}$ is acyclic. For every $v \in V$, the language $\mathscr{L}(G_{\Sigma,v})$ is a one-element set since by Point (1) we have exactly one $N_v \longrightarrow_{G_{\Sigma,V}} t$ for each v, and, since $\Gamma_{G_{\Sigma,V}}$ is well founded, each path from t will uniquely lead to a terminal node. Thus the unique data dependency is

$$D_{G_{\Sigma,V}} = \langle V, \stackrel{G_{\Sigma,V}}{\rightarrowtail}, lab_{G_{\Sigma,V}} \rangle$$

is the unique string in $\mathscr{L}(G_{\Sigma,V})$.

where $lab_{G_{\Sigma,V}}(v) = q$, where q is the unique string in $\mathscr{L}(G_{\Sigma,v})$.

In each particular case of a given recurrence (7) and an algebra A, the well-definedness of the resulting dependency has to be checked, for instance, by verifying the conditions of the above proposition.

Another obvious fact following from the assumptions of Proposition 3.1 is that if $u \xrightarrow{G_{\Sigma,V}} v$ then $lab_{G_{\Sigma,V}}(u)$ is a subterm of $lab_{G_{\Sigma,V}}(v)$.

Remark. The described construction obviously subsumes primitive recursion for which we have $\Sigma = \langle Nat, \mathscr{F} \rangle$, the only model \mathscr{N} being that of the naturals such that \mathscr{F} are the basic primitive recursive operations on naturals. Given recursive equation schema for an R with n + 1 variables

$$R(0, y_1, ..., y_n) = g(y_1, ..., y_n)$$

$$R(s(i), y_1, ..., y_n) = h(i, y_1, ..., y_n, R(i, y_1, ..., y_n)),$$

where g and h are primitive recursive operations, we get the set of nodes $V = \mathbb{N}^{n+1}$, nodes of the form $y = \langle y_0, y_1, \dots, y_n \rangle$, and the productions for any fixed values of $y_1 \dots y_n$:

$$\begin{array}{lcl} N_{\langle 0,y_1,\ldots,y_n \rangle} & \longrightarrow & g(y_1,\ldots,y_n) \\ \\ N_{\langle i+1,y_1,\ldots,y_n \rangle} & \longrightarrow & h(i,y_1,\ldots,y_n,N_{\langle i,y_1,\ldots,y_n \rangle}) \end{array}$$

This approach to the recursive programming of functions is a derivative of the semantic (that is, μ -recursive) approach to computability. Most functional languages, on the other hand, are compiled using (parallel) graph reduction techniques (Peyton and Simon 1987) or, more generally, a syntax-oriented computability model based on term substitution, for example, λ -calculus. Unlike these substitution based approaches, the construction described here leads often immediately to an efficient dependency structure that is amenable to a direct translation to efficient code on (parallel) machine hardware. The next two subsections illustrate this claim using two examples.

3.1. A simple example – the Fibonacci function

Take again the Fibonacci function from Example 2.11, which we want to compute in a standard algebra \mathcal{N} of natural numbers. That is, we have the signature $\Sigma = \langle \{Nat\}, \{s : Nat \rightarrow Nat, \oplus : Nat \times Nat \rightarrow Nat, z : Nat \} \rangle$, and \mathcal{N} is given by $Nat^{\mathcal{N}} = \mathbb{N}$, the set of natural numbers, $s^{\mathcal{N}} =$ the successor function, $\oplus^{\mathcal{N}} =$ addition, and $z^{\mathcal{N}} = 0$.

The recursive specification of the Fibonacci function $F : Nat \rightarrow Nat$ is

$$F(z) = s(z)$$

$$F(s(z)) = s(z)$$

$$F(i \oplus ss(z)) = F(i) \oplus F(i \oplus s(z)).$$
(11)

We choose \mathbb{N} as the set of nodes V (corresponding to the arity of F), that is, $V = \mathbb{N}$. With N_k , for $k \in \mathbb{N}$, as non-terminal symbols, we now obtain the production rules for the set of node-specific grammars (using $z^{\mathcal{N}} = 0$, $s(z)^{\mathcal{N}} = 1$ and $ss(z)^{\mathcal{N}} = 2$ for the indices):

$$N_0 \longrightarrow s(z)$$

$$N_1 \longrightarrow s(z)$$

$$N_{i+2} \longrightarrow (N_i \oplus N_{i+1}).$$
(12)



Fig. 13. The graph derived from the Fibonacci recurrence (11): on the left with the non-terminals, on the right with the language derived for each node (abbreviating s(z) by 1).

Each non-terminal has only one production and so each node will be labelled by a unique ground term generated by the corresponding grammar. The resulting graph and labelling (with parentheses added for readability) are shown in Figure 13.

Notice that although labelling uses fully unfolded terms, the graph itself is not the usual tree of recursive calls as, for example, **R** in Figure 3, but a much more efficient graph Γ_{Fib} .

3.2. An example of parallelization – the fast Fourier transform

To illustrate the application to actual parallelization of programs, we review the technique of the fast Fourier transform, FFT, due to Cooley and Tukey (1965). We first present the background problem and its solution given by a recurrence. The dependency for this recurrence obtained by our technique turns out to have the well-known shape of the *butterfly graph*. We then show a standard distribution of the butterfly on a hypercube architecture.

3.2.1. The numerical problem In many areas, computing of *n*-order polynomials $p(x) = \sum_{k=0}^{n} p_k x^k$, $0 \le n$, for certain argument values $x = x_1, \ldots, x_k$ is important. These areas include, for instance, signal processing, where evaluation of the polynomials for the complex roots of unity plays a central role.

An *n*'th complex root of unity is a complex number ω such that $\omega^n = 1$. For every $n \ge 1$ there are *n* distinct complex roots, spread at *n* regular intervals along the unit circle in complex space, numbered counterclockwise starting at the number 1, which is root number 0. The *j*'th such root is denoted ω_n^j , and satisfies

$$\omega_n^0 = 1$$
 $\omega_n^{j+1} = \omega_n^j \omega_n^1$ $\omega_n^j = \omega_n^{j \mod n}$

Useful facts about these roots are that $(\omega_{2n}^j)^2 = \omega_{2n}^{2j} = \omega_n^j$ and $\omega_{2n}^{j+n} = -\omega_{2n}^j$. The roots may be computed by $\omega_n^j = e^{(2\pi i/n)*j} = \cos(2\pi j/n) + \sin(2\pi j/n)i$, where $i = \omega_4^1$ is the imaginary unit, that is, $i^2 = -1$.

Cooley and Tukey (Cooley and Tukey 1965) discovered a very efficient algorithm, the FFT, for computing an *n*-order polynomial for all *n*'th complex roots of unity. Let us consider this method for the case where $n = 2^M$, that is, the one-dimensional binary version. Then the polynomial p(x) can be split into 'even' p_e and 'odd' p_o parts such that



Fig. 14. The butterfly pattern of the recurrence F.

 $p(x) = p_e(x^2) + x * p_o(x^2)$, where $p_e(x) = \sum_{k=0}^{2^{n-1}} p_{2k} * x^k$ and $p_o(x) = \sum_{k=0}^{2^{n-1}} p_{2k+1} * x^k$. Taking into account the properties of the roots of unity, we get

$$p(\omega_{2^{m}}^{j}) = p_{e}(\omega_{2^{m-1}}^{j}) + \omega_{2^{m}}^{j}p_{o}(\omega_{2^{m-1}}^{j})$$

$$p_{e}(\omega_{2^{m-1}}^{j}) = p_{e,e}(\omega_{2^{m-2}}^{j}) + \omega_{2^{m-1}}^{j}p_{e,o}(\omega_{2^{m-2}}^{j})$$

$$p_{o}(\omega_{2^{m-1}}^{j}) = p_{o,e}(\omega_{2^{m-2}}^{j}) + \omega_{2^{m-1}}^{j}p_{o,o}(\omega_{2^{m-2}}^{j})$$

This can be formulated as a recurrence equation

$$F(j,0) = p_j$$

$$F(j,m+1) = F(s_0(j,m+1),m) + \omega_{2m+1}^j F(s_1(j,m+1),m)$$
(13)

where $s_0(j,m) = j - (j \mod 2^m) + (j \mod 2^{m-1})$ sets the *m*'th bit of *j* to 0 and $s_1(j,m) = j - (j \mod 2^m) + 2^m + (j \mod 2^{m-1})$ sets the *m*'th bit of *j* to 1. Bits are the 0's and 1's in the binary representation of a number, and are counted with the least significant bit as number 1. The value of the polynomial at the *j*-th complex root of unity then equals $p(\omega_{2^M}^j) = F(j, M)$.

3.2.2. Constructing a dependency Applying our strategy, we start with the equations (13) defining the recurrence. We work with the algebra of natural and complex numbers, and as the set of nodes V we take the cartesian product $\{\langle j,m \rangle \in \mathbb{N} \times \mathbb{N} \mid 0 \leq j < 2^M, 0 \leq m \leq M\}$ – since F has two natural number arguments that are within these limits. The grammar resulting from the equations (13) is then (using \oplus as the complex addition operation and \otimes as the complex multiplication):

with the coefficients p_j of the polynomial treated as input since the roots of unity $\omega_{2^{m+1}}^j$ are fixed once M is known. The resulting shape of the data dependency for M = 3 is given in Figure 14. Again, observe how this method leads directly to an 'efficient' graph instead of generating the full tree of recursive calls.

Columns correspond to the j and rows to the m arguments with the leftmost column and bottom row numbered 0. The bottom row will be labelled with $p_0, p_1, p_2, ...$ according to the first kind of productions from (14). The labels in the upper rows will be generated



Fig. 15. Connections for hypercubes of dimension 3 (solid), and 4 (solid and dotted).

using the second kind of productions from (14). We do not write it explicitly (lengthy labelling can be easily recovered from the grammar) but only observe that the label at node $\langle j, m \rangle$ will represent the value of F(j,m). For instance, the node at row 2, column 1 (marked with the circle instead of bullet on the drawing) will represent the value of F(1,2).

3.2.3. Distribution on a hypercube The shape from Figure 14 is known as a butterfly graph. Increasing M by one, that is, taking $n = 2^{M+1}$, amounts to putting two butterflies of height M side-by-side, adding a new set of nodes on top (row M + 1), and connecting uppermost nodes (in the M'th row) of the one butterfly to its own and to the respective nodes of the other in row M + 1.

Now, Figure 15 shows the topology of hypercubes of dimension M = 3 and 4. Each of the 2^M processors has M bidirectional communication channels. A hypercube of dimension M + 1 is obtained from two hypercubes of dimension M by connecting each node of one hypercube to the corresponding node of the other.

The hypercube parallel program will then, on hypercube node j at time-step m + 1, compute F(j, m + 1) using input data supplied by nodes $s_0(j, m + 1)$ and $s_1(j, m + 1)$ at time-step m. One of the data supplying nodes will be the same as the node j, the other will be its neighbour along dimension m.

Figure 16 shows a distribution of the butterfly dependency of height 3 from Figure 14 on the *st*-graph of the hypercube of dimension 3 from Figure 15. (Solid arrows indicate



Fig. 16. Distribution of the butterfly on a hypercube.

the embedded edges of the butterfly; dotted lines the remaining edges of the hypercube.) Using the regularity in building a butterfly and a hypercube of higher dimensions, an analogous distribution is obtained for a butterfly of arbitrary height M into a hypercube of dimension M.

4. More about the development process

This section adds a few results concerning the development process using computation algebras. First, in Section 4.1, we show the counterpart of the standard result allowing one to view computation algebras over different signatures as objects of one category. Thus we obtain the means of refining the picture of the development process from Figure 11 by also admitting transitions between computation algebras over different signatures. Section 4.2 then identifies two subcategories of $CAlg(\Sigma)$ with more specific morphisms that make the coupling of the algebra- and the communication-part tighter than in the general case of computation homomorphisms from Definition 2.13. The latter of these subcategories was used in the design of the actual specification formalism SAPPHIRE. Section 4.3 states two further properties of the category of computation algebras that are of relevance for the methodology of development.

4.1. Computation algebras over different signatures

So far we have only considered computation algebras over one given signature. Thus, the development process illustrated in Figure 11 is not quite realistic since it is fairly certain that such a process will involve a change of signatures. We will now extend the standard way of flattening the categories of algebras indexed by signatures to the computation algebras and their distributions.

Let Sig be the category of signatures with injective signature morphisms[†]. We write a dependency **D** as a triple $\langle I^D, \xrightarrow{D}, lab_{\mathbf{D}} \rangle$.

Definition 4.1. (σ -reduct) Let $\sigma : \Sigma' \to \Sigma \in Mor(Sig)$ and $\mathbf{D} \in DD(\Sigma)$. The σ -reduct of $\mathbf{D}, \mathbf{D}|_{\sigma} \in DD(\Sigma')$, is defined in two steps (we assume the ordinary extension of σ to $\sigma : \mathcal{F}_{\Sigma} \to \mathcal{F}_{\Sigma'}$):

- 1 Let **X** be **D** with restricted re-labelling : $I^X = I^D$, $\stackrel{X}{\rightarrow} = \stackrel{D}{\rightarrow}$ and $lab_X(i) = \sigma^-(lab_D(i))$, where $\sigma^- : \mathscr{T}_{\Sigma} \to \mathscr{T}_{\Sigma'}$ is given by $\sigma^-(t) = \begin{cases} t' & \text{such that } \sigma(t') = t \text{ if it exists} \\ \bot & \text{otherwise.} \end{cases}$
- 2 $\mathbf{D}|_{\sigma}$ is then given by
 - $I^{D|_{\sigma}} = \{ i \in I^D : \sigma^{-}(lab_X(i)) \neq \bot \},\$
 - $lab_{D|_{\sigma}}(i) = lab_X(i)$, and

[†] We restrict ourselves to the injective signature morphisms for one main reason: defining a reduct functor over data dependencies for non-injective morphisms implies several choices that seem slightly arbitrary. We therefore prefer to postpone such a decision until we have more experience and reasons to choose one alternative over the others.

$$- i \xrightarrow{D|_{\sigma}} j \Leftrightarrow \begin{cases} i \xrightarrow{D} j & or \\ \exists \text{ path } i \xrightarrow{X} k_1 \cdots k_z \xrightarrow{X} j \text{ and } 1 \leqslant n \leqslant z \Rightarrow lab_X(k_n) = \bot \end{cases}$$

In the first step, we remove all the labels from D that are not in the image of σ , and re-label the remaining ones to their σ pre-image. In the second we convert paths with unlabelled (\perp -labelled) intermediary nodes into edges between the labelled end-points.

For $A \in Alg(\Sigma)$ and $\sigma : \Sigma' \to \Sigma \in Mor(Sig)$, we let $A|_{\sigma}$ denote the usual reduct. We then have the following extension of the standard result.

Proposition 4.2. $\sigma : \Sigma' \to \Sigma \in Mor(Sig)$ induces the functor $|_{\sigma} : CAlg(\Sigma) \to CAlg(\Sigma')$.

Proof. For an $A \in \mathsf{CAlg}(\Sigma)$, define $A|_{\sigma} = \langle \bar{A}|_{\sigma}, \bar{A}|_{\sigma} \rangle$. For a morphism $h : A \to B \in Mor(\mathsf{CAlg}(\Sigma))$, $h|_{\sigma} = \langle \bar{h}|_{\sigma}, \bar{h}|_{\sigma} \rangle$ where $\bar{h}|_{\sigma}$ is the image of \bar{h} under the usual reduct functor for algebras, while $\bar{h}|_{\sigma}$ is the restriction of \bar{h} to $\bar{B}|_{\sigma}$ under changed labelling: each node i of $\bar{B}|_{\sigma}$ is sent to the node $\bar{h}(i)|_{\sigma} = \bar{h}(i)$, and edge e of $\bar{B}|_{\sigma}$ onto the edge (path/set of paths) $\bar{h}(e)|_{\sigma}$ in $\bar{A}|_{\sigma}$. This is a well-defined $\mathsf{DD}(\Sigma')$ -morphism: commutativity \bar{h} ; ev_A ; $\bar{h} = ev_B$ implies the corresponding commutativity in the reduct, since the latter operations are restrictions of their pre-images.

Thus $_{-|_{-}}$: Sig^{op} \rightarrow CAT sending Σ to CAlg(Σ) and σ to $_{-|_{\sigma}}$ is an indexed category. By the standard application of the Grothendieck construction we may define the flattened category CAlg with (1) objects $\langle \Sigma, A \rangle$ where $A \in Obj(CAlg(\Sigma))$, (2) morphisms of the form $\langle \sigma, f \rangle : \langle \Sigma', A \rangle \rightarrow \langle \Sigma, B \rangle$, where $\sigma \in Mor(Sig)$ and $f : A \rightarrow B|_{\sigma} \in Mor(CAlg(\Sigma'))$, and (3) composition $\langle \sigma, f \rangle; \langle \rho, g \rangle = \langle \sigma; \rho, f; g|_{\rho} \rangle$.

Quite an analogous construction yields an indexed category $\|_{-}\|_{-}$: Sig^{op} \to CAT that sends Σ to Dist(Σ). For $\sigma : \Sigma' \to \Sigma$, the induced functor $\|_{\sigma} : \text{Dist}(\Sigma') \to \text{Dist}(\Sigma)$, sends a distribution $f_{A\to W} \in \text{Dist}(\Sigma)$ onto $(f\|_{\sigma})_{A|\sigma\to W} \in \text{Dist}(\Sigma')$, where $f\|_{\sigma}$ is the restriction of f to $\vec{A}|_{\sigma}$. The result of the flattening is the category Dist. We do not dwell on this general category, because we consider the categories $\text{Dist}(\Sigma)$ for particular Σ more central – porting *distributions* does not typically involve changes of signature.

4.2. More specific dependency morphisms

The class $CAlg(\Sigma)$ provides the general framework for possible specializations. Arbitrary dependency morphisms in $DD(\Sigma)$ make the connection between the algebra- and the communication-part of the objects in $CAlg(\Sigma)$ rather loose and indefinite. We mention two possible restrictions.

4.2.1. Computation algebras with compatible D-morphisms Let A be a fixed class of Σ algebras. It is natural to think of it as a model class of some (equational) specification. One may postulate that such a choice of A should restrict the relevant dependency morphisms. In particular, if A $\not\models s = t$, the D-morphisms should not be allowed to map nodes labelled by s to ones labelled by t (nor vice versa), since such mappings are not compatible with the algebras in A. Thus, compatibility of a DD(Σ) morphism $h : \mathbb{C} \to \mathbb{D}$ with respect to an algebra A means that h does not effect a re-labelling that is inconsistent with valuation in A. More precisely, we have the following definition. **Definition 4.3.** Let $h : \mathbb{C} \to \mathbb{D} \in Mor(DD(\Sigma))$ and $A \in A \subseteq Alg(\Sigma)$.

1 h is A-compatible iff $\forall i \in I^C : (lab_{\mathbf{D}}(h(i)))^A = (lab_{\mathbf{C}}(i))^A$;

- 2 h is A-compatible iff it is A-compatible for all $A \in Obj(A)$;
- 3 DD(A) is a wide subcategory of DD(Σ) with A-compatible morphisms.

Alternatively, $h : \mathbb{C} \to \mathbb{D}$ is A-compatible iff for each $A \in Obj(A)$, $\langle id_A, h \rangle$ is a computation homomorphism $\langle A, \mathbb{D} \rangle \to \langle A, \mathbb{C} \rangle$. We then have the following obvious proposition.

Proposition 4.4. Suppose that A has an initial object T_A , and let $h : \mathbb{C} \to \mathbb{D} \in Mor(DD(\Sigma))$. Then: h is A-compatible $\Leftrightarrow h$ is T_A -compatible.

Proof. (\Rightarrow) is obvious. For (\Leftarrow) assume *h* is T_A -compatible. This means that the lower triangle of the following diagram commutes, so *h*; $ev_{T_A}^{\mathbf{D}} = ev_{T_A}^{\mathbf{C}}$, where $ev_X^{\mathbf{Y}}$ assigns the values from an algebra *X* to the nodes of a dependency **Y** according to their labels (that is, $ev_X^{\mathbf{X}}(i) = (lab_{\mathbf{Y}}(i))^X$).



Since T_A is initial, for each $A \in A$ and dependency \mathbf{Y} , $ev_A^{\mathbf{Y}} = ev_{T_A}^{\mathbf{Y}}$; a where a is the unique homomorphism $T_A \to A$. But then h; $ev_{T_A}^{\mathbf{D}} = ev_{T_A}^{\mathbf{C}} \Rightarrow h$; $ev_{T_A}^{\mathbf{D}}$; $a = ev_{T_A}^{\mathbf{C}}$; $a \Rightarrow h$; $ev_A^{\mathbf{D}} = ev_A^{\mathbf{C}}$.

As a special case, the morphisms compatible with $A = Alg(\Sigma)$ are the ones that do not change the labelling at all, since they have to be compatible with the initial word algebra T_{Σ} .

For computation algebras with compatible dependency morphism, we also have the following fact about the restriction of the functor (\vec{L}) from (3):

Proposition 4.5. $(\ \)$: CAlg(A, DD(A))^{op} \rightarrow DD(A) is a split fibration.

Proof. We use the notation from the diagram to the right, which illustrates the fibration situation in the present context (the dashed arrows indicate the directions in _^{op} categories):



For a DD(A)-morphism $d : \mathbf{C} \to \mathbf{D}$ and $D = \langle \overline{D}, \mathbf{D}, ev_D \rangle$, the cleavage is defined as

$$cl(d, D) \stackrel{\text{\tiny def}}{=} \langle id_{\bar{D}}, d \rangle : C \dashrightarrow D$$
, where $C = \langle D, C, ev_C \rangle$ (15)

with ev_C given by the last point of Definition 2.12. Since *d* is A-compatible, *d*; ev_D ; $id_{\bar{D}} = ev_C$, and so the cleavage is a morphism in CAlg(A, DD(A))^{op}.

If $g : D \to B$ is over $d(g^{\bar{o}p} = d)$, the unique $f : C \to B$ over id_C making cl(d, D); f = g is $\langle \bar{g}, id_C \rangle$ (that is, $\langle id_{\bar{D}}, d \rangle$; $\langle \bar{g}, id_C \rangle = \langle \bar{g}, d \rangle$).

Obviously, the cleavages compose: for $c : \mathbf{B} \to \mathbf{C}$ and $d : \mathbf{C} \to \mathbf{D}$, we have $cl(d, D) = \langle id_{\bar{D}}, d \rangle$ and $cl(c, \langle \bar{D}, \mathbf{C} \rangle) = \langle id_{\bar{D}}, c \rangle$. So $cl(c, \langle \bar{D}, \mathbf{C} \rangle)$; $cl(d, D) = \langle id_{\bar{D}}, c \rangle$; $\langle id_{\bar{D}}, d \rangle = \langle id_{\bar{D}}, c; d \rangle = cl(c; d, D)$. Hence $(\)$ is a fibration.

It is split: $(id_{\mathbf{D}})^* \langle \langle \bar{D}, \mathbf{D} \rangle = \langle \bar{D}, \mathbf{D} \rangle$, so the natural transformation $id_{((\vec{D})^{-1})} \rightrightarrows (id_{\mathbf{D}})^*$ is identity $((\vec{D})^{-1}$ denotes the fiber over **D**). Also, since cleavages compose : $(d)^*$; $(c)^* = (c; d)^*$.

For the compatible dependency morphisms, we also have the following additional fact.

Proposition 4.6. $(\)$: CAlg(A, DD(A))^{op} \rightarrow DD(A) is a cofibration.

Proof. The proof and definition of cocleavage are essentially the same as in Proposition 4.5. We have the following picture:



For a DD(A)^{op}-morphism $d^{op} : \mathbf{C} \to \mathbf{D}$ and $D = \langle \overline{D}, \mathbf{D}, ev_D \rangle$, the cocleavage is defined as

$$ccl(d^{op}, D) \stackrel{\text{\tiny det}}{=} \langle id_{\bar{D}}, d \rangle : C \to D \text{ where } C = \langle \bar{D}, \mathbf{C}, ev_C \rangle,$$
 (16)

with ev_C induced by lab_C according to Definition 2.12.3. Trivial repetition of the arguments from the proof of Proposition 4.5 yields the conclusion.

This, together with Proposition 4.5, means that (-) is a bifibration and gives the following corollary.

Corollary 4.7. Reindexing functors induced by $((\ l, cl)$ have left adjoints.

Finally, we may refer back to Section 4.1 and observe that, for an injective signature morphism $\sigma : \Sigma' \to \Sigma$, if $d : \mathbb{C} \to \mathbb{D} \in Mor(DD(\Sigma))$ is A-compatible for an $A \in Obj(Alg(\Sigma))$, then $d|_{\sigma}$ is $A|_{\sigma}$ -compatible. Hence, the image under $_{-|_{\sigma}}$ of a CAlg(A, DD(A)) is a category $CAlg(A|_{\sigma}, D)$ where $Obj(A|_{\sigma}) = \{A|_{\sigma} : A \in Obj(A)\}$ and morphisms in D are $A|_{\sigma}$ -compatible. (In general, D may only be a subcategory of $DD(A|_{\sigma})$.)

4.2.2. *Graph-morphisms* We believe that refinement of data dependencies – at least as expressed by the compatible dependency morphisms in Definition 4.3 – may be a practical notion for the development of implementations and, perhaps, more abstract algorithms.

Nevertheless, one could argue that it is not needed in such generality since data dependencies are quite concrete objects introduced towards the very end of the development process. In particular, passing from one computation algebra to another should not imply the introduction of new dependencies, but only (and at most) purely syntactic memoisation. It is only the distribution morphisms that adjust the dependency graph to match it to the communication structure of a given architecture.

Accepting this view, one could work with a class of computation algebras (A, D) where $A \subseteq Alg(\Sigma)$ and the dependencies in D are restricted so that the target of the forgetful functor Gr projecting dependencies onto their graph-part (*cf.* (2) after Definition 2.10), is no longer the category Gr but its wide subcategory Gr^o: morphisms in Gr^o may map edges on paths but not on 'parallel compositions' of paths. (Gr^o can be defined as Gr in Definition 2.7 but with the adjunction $F^{\circ} \dashv U^{\circ}$ from Proposition 2.4 instead of $F^{\oplus} \dashv U^{\oplus}$.) Thus, objects in D are simple DAGs and morphisms are the A-compatible dependency morphisms h such that Gr(h) is a Gr^o-morphism.

In fact, restricting our attention to merely syntactic memoisation, we further restrict the morphisms in D to those that do not perform any relabelling but can, at most, identify some nodes from the source graph with identical labels. Thus, the morphisms are in fact Alg(Σ)-compatible. Let us denote this class of computation algebras by (A, D°(Σ)). Since D°(Σ)-morphisms do not perform any relabelling whatsoever, in this class, any pair $h \in Mor(D^{\circ}(\Sigma))$ and $g \in Mor(A)$ will give rise to an (A, D°(Σ))-computation morphism.

Since $Mor(D^{\circ}(\Sigma)) \subset Mor(DD(A))$, we can trivially repeat the constructions from Propositions 4.5 and 4.6 to see that $(\): (A, D^{\circ}(\Sigma))^{op} \to D^{\circ}(\Sigma)$ is a bifibration.

The class $(A, D^{\circ}(\Sigma))$, although theoretically perhaps not the most fascinating one, deserves to be mentioned because it has been used in practical applications and underlies the design of the specification formalism SAPPHIRE (Čyras and Haveraaen 1995; Haveraaen and Søreide 1998) (see Section 3).

4.3. *Two more facts about* CAlg(A, D)

The following two facts show further properties of CAlg(A, D) that can be relevant in the development process. The one in Section 4.3.1 allows us to treat refinement along the algebra- and communication-part relatively independently from each other. This indicates the possibility that once developed, chains of refinements of, say, dependency structures, could be reused in new contexts by coupling them with new actual algebras. The second, in Section 4.3.2, gives (some) conditions for the existence of initial computation algebras relative to the existence of initial objects in A and D.

4.3.1. CAlg(A, D) as a double category A morphism between computation algebras consists of a pair of one standard algebra-morphism and one dependency-morphism (with additional compatibility criterion, Definition 2.13). Yet, the coupling of the two is loose enough to allow us to separate them in a development process. More precisely, any morphism in CAlg(A, D) can be seen as applying an algebra morphism and then a dependency morphism (or *vice versa*). In a sense, CAlg(A, D) 'consists of' two categories: the 'horizontal' one being A and the 'vertical' one D.

Expressing this relative independence categorically, we can show that computation algebras form a double category with horizontal and vertical arrows, respectively, of the form $\langle \bar{h}, id \rangle$ and $\langle id, \bar{h} \rangle$ (and cells being the commuting squares $v \downarrow_{\substack{a \ b \ c \ g \ b}}^{A \longrightarrow B} h^{a}$, where h, g are horizontal and u, v vertical arrows).

We will not spell out the details of this construction, which is rather straightforward (for double categories, the reader may consult Ehresmann (1963) and Mac Lane (1971)). We only show the simple proposition illustrating that any morphism in CAIg(A, D) can be seen as consisting of two, relatively independent steps: one mapping the algebra-part and another mapping the communication-part. The nice implication of this is that we can work relatively independently with the algebra- and the communication-part of the objects in CAlg(A, D).

Proposition 4.8. Each morphism $h: C \to D$ in CAlg(A, D) can be factored so that

 $\begin{array}{l} 1 \hspace{0.2cm} h = \langle \bar{h}, id_{\vec{C}} \rangle; \hspace{0.2cm} \langle id_{\bar{D}}, \vec{h} \rangle, \\ 2 \hspace{0.2cm} h = \langle id_{\vec{C}}, \vec{h} \rangle; \hspace{0.2cm} \langle \bar{h}, id_{\vec{D}} \rangle. \end{array}$

Proof. The following diagram details the first case:



The intermediary computation algebra is as shown in the figure with $ev = ev_C$; \bar{h} . This makes $\langle \bar{h}, id_{\bar{C}} \rangle$ a computation homomorphism $\langle \bar{C}, \bar{C}, ev_C \rangle \rightarrow \langle \bar{D}, \bar{C}, ev \rangle$. Also, since h is a computation homomorphism, we have for all $i \in I^{\vec{D}}$: $\bar{h}(ev_C(\vec{h}(i))) = ev_D(i)$, that is, $ev(\vec{h}(i)) = ev_D(i)$, which means that $\langle id_{\bar{D}}, \vec{h} \rangle$ is a computation homomorphism $\langle \bar{D}, \bar{C}, ev \rangle \rightarrow \langle \bar{D}, \bar{D}, ev_D \rangle$. The second case is entirely analogous with the intermediary algebra being $\langle \bar{C}, \vec{D}, \vec{h}; ev_C \rangle$.

4.3.2. Initial objects Define functor (-): CAlg(A, D) \rightarrow D^{op} analogously to (-) in (3), that is, $\overleftarrow{A} = \overrightarrow{A}$ and $\overleftarrow{\langle f,g \rangle} = g$ (the communication part g of a morphism $\langle h,g \rangle$ in CAlg(A, D) is contravariant to the whole morphism). We then have the following general fact.

Proposition 4.9. If A has an initial object T_A , then the functor $F : D^{op} \to CAlg(A, D)$ defined by $F(\mathbf{C}) = \langle T_{\mathsf{A}}, \mathbf{C} \rangle$ and $F(g) = \langle id_{T_{\mathsf{A}}}, g \rangle$ is left adjoint to $(-), F \to (-)$, with identity as unit.



Fig. 17.

Thus, if D^{op} has an initial object, so does CAlg(A, D). However, in general, this will not be the case since interesting subcategories of $DD(\Sigma)$ do not have terminal objects (D^{op} do not have initial objects). In particular, \emptyset is initial in Gr and in $DD(\Sigma)$, but there are no terminal objects there. (If Z_A is terminal in A, then $\langle Z_A, \emptyset \rangle$ is terminal in $CAlg(A, DD(\Sigma))$.)

We do not focus on the notion of initiality and the relevance of the empty communication part is probably limited to the fact that it allows us to treat standard algebras as computation algebras with an empty dependency.

Nevertheless, the proposition gives us useful information in a slightly more restricted context. Typically, the initial object in a model class A of a specification is obtained as (or from) a term model. Then, given a particular (non-empty) dependency, the proposition provides an 'initial' evaluation strategy over such a term model.

5. Non-determinism

Since parallelism and distributed programs often are a source of non-determinism, we suggest here how this phenomenon can be incorporated into the present framework. There are at least two ways to do it. The first, given in Section 5.1, utilizes the notion of multialgebras, which are designed for incorporating non-determinism into algebraic specification as an abstraction mechanism (Hußmann 1993; Walicki and Meldal 1994; Walicki and Meldal 1997). The second, given in Section 5.2, is biased towards the operational view, according to which non-determinism is something that arises only during the actual computations.

5.1. The multialgebraic view

Revisiting the definition of computation algebras (Definition 2.12) and the subsequent constructions, one can observe that we have not made any significant assumptions about the underlying category of Σ -algebras. The only essential requirement was that *ev*-function assigns values from an algebra to each node of the dependency in a way respecting the *lab*elling of the node. In fact, instead of the usual Σ -algebras, we can work with a different category for Alg(Σ).

For modelling non-determinism, a natural choice is (some) category of multialgebras with multihomomorphisms (Walicki and Białasik 1997). A multialgebra is an algebra

628

where operations applied to single arguments may return sets of elements, which correspond to the possible results of a nondeterministic operation. Composition of operations is defined pointwise. A (ground) term is interpreted in a multialgebra as the set (of its possible results). The notion of a homomorphism $h : A \to B$ may be generalized in various ways (Walicki and Białasik (1997) surveys the possibilities), but the most common method replaces the usual homomorphism condition by the requirement $h(f(x)^A) \subseteq f^B(h(x))$, for each function symbol $f \in \Sigma$. There is also a choice of the primitive operations for writing specifications. The most common approaches use a primitive predicate, s < t (interpreted in an algebra A as set inclusion $s^A \subseteq t^A$, that is, as t being at least as nondeterministic as s), and/or the nondeterministic binary choice $\Box \Box : S \times S \to S$, for some/every sort S. We do not discuss the specification languages here, and in the examples to follow we will use the above, most common, definition of multihomomorphism.

The change of the category of algebras is the only modification needed. The computation homomorphism h (with \bar{h} being a multihomomorphism) allows us, for instance, to refine an algebra B to a more deterministic algebra A as illustrated in Figure 17. All operations except \sqcup are deterministic. \bar{h} may relabel $s \sqcup t$ to s because both terms have the same sort and, furthermore, because \bar{A} is more deterministic than \bar{B} : for $i \in I^B$ with $lab(i) = s \sqcup t$, we have that $\bar{h}(ev_A(\bar{h}(i))) = \bar{h}(s^A) = s^B \subseteq s^B \cup t^B = (s \sqcup t)^B = ev_B(i)$, that is, \bar{h} is a multihomomorphism.

5.2. The operational view

One source of non-determinism is purely operational – the differences in relative speed of various processors on various machines being unpredictable. A paradigmatic example of this is a processor P waiting for an input from one of several other processors. P processes the data that arrives first – without discriminating against any of the sources – and discards any later arrivals. The dependency

$$s \xrightarrow{s \sqcup t} t$$

from Figure 17 can be read as expressing just that: the value at the topmost node depends on the values at the other two nodes – it will simply be the one arriving first. Multialgebras provide an abstract model for this kind of phenomenon. We will now sketch a more detailed, low-level way of including them in the computation algebra framework.

The computation related information for a computation algebra $A = \langle \bar{A}, \dot{A} \rangle$ is represented by its communication part. Thus we may retain the standard notion of algebra \bar{A} and try to model non-determinism in \tilde{A} . We simply allow for labelling the nodes of \tilde{A} by sets of terms, rather than unique terms. The set labelling a node represents the possible values computed at the node. The source of the evaluation function ev_A is then a set of pairs $\langle i, t \rangle$ where *i* is some node and *t* one of the terms labelling $i : \bigcup_{i \in I^{\tilde{Gr}(A)}} \{\langle i, t \rangle : t \in lab^A(i)\}$. The nodes act as oracles – in an actual computation each of them will produce a unique value. But this value is not known in advance, so the labelling merely indicates the range of possibilities.





Definition 5.1. $nDD(\Sigma)$ is the category where (Γ is as in Definition 2.1):

- Objects are Σ n-data dependencies pairs G = ⟨G, lab⟩ where G ∈ Obj(Gr) is a simple DAG and lab : I^G → 𝒫⁺(𝒯_Σ) is a function labelling each node of this graph with a non-empty set of ground Σ-terms of the same sort, that is, ∀i ∀s, t ∈ lab(i) : Sort(s) = Sort(t).
 An nDD(Σ)-morphism h : B → A is a pair ⟨h, h_{lab}⟩ where
- \vec{h} is a (sort compatible) Gr-morphism $\vec{h} : Gr(\mathbf{B}) \to Gr(\mathbf{A})$,
- h_{lab} is a family $\{h_i : lab_{\mathbf{A}}(\vec{h}(i)) \rightarrow lab_{\mathbf{B}}(i)\}_{i \in I^B}$.

Sort compatibility means that the morphisms have to respect the sorting of labels: if labels of *i* are of sort *S*, then the labels of $\vec{h}(i)$ must be as well. The h_{lab} mapping (which we can treat as a partial function $I^A \times \mathscr{T}_{\Sigma} \to \mathscr{T}_{\Sigma}$ with the domain given by all the pairs $\langle \vec{h}(i), t \rangle$ for all $i \in I^B$ with $t \in lab_A(\vec{h}(i))$ picks the subset of labels of $lab_B(i)$ that is the pre-image of the label set $lab_A(\vec{h}(i))$. For instance, in Figure 18, **B** is a dependency for nondeterministic choice between *s* and *t*, which is then propagated as the argument to the function *f*. (The relabelling is merely a restriction and h_{lab} is the obvious inclusion.)

The node *i* in **B** with the label $\{s, t\}$ may compute *s* or *t*. Similarly, the topmost node *j* may compute f(s) or f(t). The dependency $i \rightarrow j$ says that when *i* computes one of *s* or *t*, *j* will compute one of f(s) or f(t). A possible deterministic refinement of **B** that always chooses *s* and computes only f(s) is given in **A** with the morphism *h*.

Observe, however, that the definitions of an n-dependency and an $nDD(\Sigma)$ -morphism do not take into account the possible semantic information about how the computation at *j* depends on the result delivered by *i*. In principle, it admits situations like **C**, where we obtain $s \rightarrow f(t)$. The morphism only captures the preservation of dependencies (in terms of the underlying graphs) and the non-increasing non-determinism. That the computation of f(t) at *j* may result only if the node below produces *t* is the kind of semantic information – how the computation at *j* depends on the input from *i* – that might (should) be incorporated into the specification of dependencies. In short, if the dependency $s \rightarrow f(t)$ in **C** is not of the intended kind, it should be prohibited by the specification (this topic is to be explored more fully in a future paper).

Definition 5.2. Given $A \subseteq Alg(\Sigma)$ and $D \subseteq nDD(\Sigma)$; an (A, D) *n-computation algebra* C is $\langle \overline{C}, \overline{C}, ev_C \rangle$ where

1 $\vec{C} \in Obj(D)$; 2 $\vec{C} \in Obj(A)$; 3 $ev_C : \vec{C} \times \mathscr{T}_{\Sigma} \to \vec{C}$ is a (partial) function such that $\forall t \in lab_{\vec{C}}(i) : ev_C(i, t) = t^{\vec{C}}$.

As for computation algebras, ev is uniquely determined by \overline{C} and the labelling of \vec{C} .



Fig. 19. An n-computation homomorphism.

Definition 5.3. An (A, D) *n*-computation homomorphism $h : C \to D$ is $\langle \bar{h}, \langle \bar{h}, h_{lab} \rangle \rangle$ where 1 $\bar{h} : \bar{C} \to \bar{D} \in Mor(A)$; 2 $\langle \bar{h}, h_{lab} \rangle : \bar{D} \to \bar{C} \in Mor(D)$, and 3 for each $i \in I^{\vec{Gr}(D)}$ and $t \in lab_{\mathbb{C}}(\bar{h}(i))$, we have $\bar{h}(ev_{\mathbb{C}}(\bar{h}(i), t)) = ev_{\mathbb{D}}(i, h_{lab}(\bar{h}(i), t))$.

The last condition (illustrated in Figure 19) is a generalization of the commutativity condition for the computation homomorphisms from the Definition 2.13. In the deterministic case, the two coincide, since then h_{lab} is uniquely determined by \vec{h} .

The category nCAlg(A, D) has (A, D) n-computation algebras as objects and (A, D) ncomputation homomorphisms as morphisms. Distributions of nCAlg(A, D) are defined as before (Definition 2.19), and so are the projection functors ($\vec{.}$) and Arch. Proposition 2.26 generalizes trivially to the nondeterministic context by an entirely analogous proof using the construction (6). Modifying the first point of Definition 4.3 to the requirement $\forall i \in I^{Gr(C)}, t \in lab_{\mathbf{D}}(\vec{h}(i)) : t^{A} = (h_{lab}(\vec{h}(i), t))^{A}$, we may define the class nCAlg(A, nDD(A)) of n-algebras with compatible dependency morphisms. Lemma 4.5 then generalizes equally easily to this class by a construction analogous to (16). We therefore do not give it explicitly here but only emphasize that the earlier results remain valid when we put n-'s at all relevant places.

6. Conclusions and further work

We have introduced a general notion of computation algebras that extends the standard algebraic semantics with additional structures, data dependencies, carrying information concerning the evaluation strategy. The dependency morphisms can be used to design more efficient implementations, in particular, by syntactic memoisation. The combination of dependencies with actual algebras and the notion of computation homomorphisms allows us to perform semantic memoisation as well.

The explicit information about the computation structure opens up the possibility of constructing parallel implementations from algebraic specifications. Dependency and distribution morphisms express, respectively, two aspects of parallelization: (a) the logical dependency decomposition, and (b) partitioning and routing on a physical medium. We have shown how indexing computation algebras with machine architectures yields a fibration. Its explicitly defined cleavage provides a way of porting distributions between various (parallel) architectures.

The general framework admits the definition of various subclasses of computation algebras. One such class – the algebras, CAlg(A, DD(A)), with compatible dependency morphisms – has been singled out and studied in more detail.

631

M. Walicki, M. Haveraaen and S. Meldal

Extending algebra(s) with the computation and control information resembles the otherwise widely studied attempts of adding such an information to term rewriting systems (Clavel *et al.* 1996; Borovansky *et al.* 1998a; Borovansky *et al.* 1998b; Clavel *et al.* 1999; Visser 1999). In this respect, the main difference is that we are not considering executable specifications (like term rewriting systems) and their efficiency or capabilities for modelling computations, but efficiency of an actual implementation of a (low level, detailed) specification. In particular, dependencies can be adjusted to actual (parallel) machine architectures, which are included as a part of the overall picture in our model of development process.

In Section 3 we suggested a method allowing one to construct data dependency from a given specification of a recursive function to be computed in a given algebra. A special case of this method, combined with a subcategory $(A, D^{\circ}(\Sigma))$ of computation algebras (*cf.* Section 4.2.2) and embedded in the programming language SAPPHIRE, has been worked out in more detail and used in practical applications for parallelization of programs (Haveraaen 1993; Čyras and Haveraaen 1995; Haveraaen and Søreide 1998; Haveraaen 2001). This construction of efficient data dependencies for recursive functions leads to the consideration of efficient implementations of functional programming languages, in particular, to basing such implementations on the semantic (μ -recursive), rather then syntactic (substitution oriented) computational model. This, however, is an open question for future research that falls outside the scope of the present paper.

Open issues

Computation algebras offer a mathematical framework for modelling efficient implementation of algebras. Application of this framework to actual architectures and (parallel) programming languages is the main topic deserving further study. We mention here three issues of more detailed character.

1 Space-time representation does not capture all aspects of parallel architectures For instance, the simulation m of a 2-dimensional vector on a 3-processor ring from Figure 7 (c) assumes that multiple data can be sent along the channels. (The third node in the rightmost column in Figure 7 (c) receives two pieces of data that both have to be passed to the node above it.) In general, this may be precluded on machines where sending multiple data requires several computation steps. Thus actual simulation morphisms have to respect such additional restrictions.

Similar issues concern the kind of allowed communications. Figure 20 (b) shows identical space-time representations of two different architectures: B – with broadcast, and S with stepwise communication (that is, a processor can communicate only along one channel at a time). The distribution (c) of **D** on B is not meaningful on S, which can be achieved in two steps as illustrated in (d).

Distribution morphisms can distinguish between these situations but since this information is not part of the st-representation, this is something that has to be done when designing actual morphisms. One should verify that space-time graphs, together with the simulation and distribution morphisms, offer sufficient representational power.

2 Portability vs. Efficiency Another issue concerns the efficiency of the ported imple-



Fig. 20. a) Data dependency \mathbf{D} , b) space-time representation of a broadcast, B, and a stepwise, S, architecture, c) distribution of \mathbf{D} on B, and d) on S.

mentations (distributions). For instance, the implementation achieved in Figure 9 (c) as the result of porting an implementation from a vector to a ring, is certainly not the most efficient one. There is, for instance, the unnecessary step from $(Z_3, 1)$ – holding F(1) – to $(Z_3, 2)$. This delay occured because we have used a generic simulation m of W by Z (from Figure 7 (c), that is, one that allows us to port any implementation from W to Z. Even if such a simulation is generically optimal (that is, uses minimum time on the target architecture for all possible simulations from the source architecture), for a particular case it may be possible to find a more efficient mapping. However, though efficiency of computation is always a concern, general portability and re-usability of software is at least equally important. What we have proposed is a sound methodology that guarantees correctness preservation of an implementation that is ported from one platform to another. We expect that the possible loss in computational efficiency will, hopefully, be richly compensated for by increased engineering efficiency. And, having successfully ported an implementation, there is nothing precluding some final optimization in order to achieve a more efficient product.

Code generation The most important issue concerns generation of actual code. It is 3 not at all obvious that the communication dependencies and the data dependencies should have the same or similar interpretation in terms of evaluation strategies. We may imagine that the hardware dependency $i \rightarrow j$ reflects the fact that j has to wait for a piece of data from i – once it is produced and received, j begins its computation. But we may also imagine a different scenario in which *j* ignores the input from *i* and simply evaluates whatever it is supposed to as soon as it has sufficient information to do so (that is, a non-strict interpretation of the dependency that is determined at run-time). A unifying view is to think of the nodes (in a dependency) as 'active agents' who make their local decisions as to how to treat the incoming information. Then the presence of an edge identifies a *potential* dependency. As a particularly interesting case, if we admit non-determinism in computations, it may turn out that the dependencies are introduced exactly in order to allow non-strict evaluations where the relative speed of processing and communication determines which dependencies are realized. The resulting implementation may turn out to be more time-efficient, allowing simultaneous computation of possible inputs.

Although not sufficiently detailed, our notion of distribution (Definition 2.19) does

M. Walicki, M. Haveraaen and S. Meldal

capture several aspects of parallelization. First, different 'columns' in the st-graph (the sets of nodes $\{\langle s,t \rangle : t \in \mathbb{N}\}$ for different s) of a given architecture will represent different processors and so a distribution mapping amounts to *partitioning* of the code onto these processors. Furthermore, the links between different components ('columns') of the partition correspond to possible communications between processors. Mapping a particular edge of a data dependency onto a path in a space-time graph then amounts to a particular choice of *routing*. (This is why we want to work with the path, and not transitive closures of graphs.) Finally, distribution morphisms represent (part of) the generation of the actual code, which consists of two parts: the communication part and the computation part. The shapes of dependencies serve as the source only of the first part, which is related to the communication structure, while their labelling indicates distribution of the actual code. $i \rightarrow j$ could correspond to the following commands being executed at i and j, respectively: 'i : compute X; send to j;', and 'j : wait for i; compute Y;', where X, Y are the pieces of code resulting from the labelling of iand j. (The directions of sending/receiving will often be identified not by the (name of) another processor but by the local (name of a) channel. This is another specific distinction, depending on the actual machine and programming language, which we have not addressed at all.)

Our model comprises only the abstract notion, not such a concrete view of implementation. Code generation for a special class of computation algebras specified equationally with the explicit specification of the dependency structures was studied in connection with the SAPPHIRE methodology for development of parallel programs, (Haveraaen 1993; Haveraaen and Søreide 1998).

Other questions for further research

- Although we have not discussed any specification formalism, it should be obvious that all the aspects of our setting are amenable to a description using algebraic techniques. In particular, data dependencies, architectures, and morphisms between them are standard algebraic objects. Future research should integrate the description of data dependencies into an actual specification framework.
- An interesting alternative would be to investigate the possibilities of extracting data dependencies from a given algebraic specification. Section 3 opens a way in this direction. Open questions concern, in particular, the syntactic conditions on the recursive definitions such as (7) that would ensure well-definedness of the resulting function in specific algebras. General construction of graphs and labellings from specifications is a broader issue worth closer investigation.
- A variety of dependency and, especially, simulation morphisms between existing architectures could be defined and stored for future use in concrete applications. It would be desirable to develop such a library and the tools for its use.
- We have indicated how the operational and multialgebraic notions of non-determinism may be included into the framework. However, the role of non-determinism will be explored further.
- We would like to make a closer study of the notion of dependency refinement and its

possible use for the development of algorithms. It is possible that the techniques of graph transformation might be relevant for this purpose.

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