# Heptagon/BZR manual

## July 5, 2012

## 1 Introduction and tutorial

### 1.1 Heptagon: short presentation

Heptagon is a synchronous dataflow language, with a syntax allowing the expression of control structures (e.g., switch or mode automata).

A typical Heptagon program will take as input a sequence of values, and will output a sequence of values. Then, variables (inputs, outputs or locals) as well as constants are actually variable or constant *streams*. The usual operators (e.g., arithmetic or Boolean operators) are applied pointwise on these sequences of values.

For example, the Heptagon program below is composed of one node **plus**, performing the pointwise sum of its two integer inputs:

```
node plus(x:int,y:int) returns (z:int)
let
    z = x + y;
tel
```

**x** and **y** are the inputs of the node **plus**; **z** is the output. **x**, **y** and **z** are of type **int**, denoting integer *streams*. **z** is defined by the equation z = x + y.

An execution of the node **plus** can then be:

| x          | 1 | 2 | 3 | 4 |  |
|------------|---|---|---|---|--|
| y          | 1 | 2 | 1 | 2 |  |
| plus(x, y) | 2 | 4 | 4 | 6 |  |

## 1.2 Compilation

The Heptagon compiler is named **heptc**. Its list of options is available by :

#### > heptc -help

Every options described below are cumulable.

Assuming that the program to compile is in a file named **example.ept**, then one can compile it by typing :

> heptc example.ept

However, such compilation will only perform standard analysis (such as typing, causality, scheduling) and output intermediate object code, but not any final or executable code.

The Heptagon compiler can thus generate code in some general languages, in order to obtain either a standalone executable, or a linkable library. The target language must then be given by the -target option:

```
> heptc -target <language> example.ept
```

Where <language> is the name of the target language. For now, available languages are C (c option) and Java (java option).

#### 1.3 Generated code

The generic generated code consists, for each node, of two imperative functions:

- one "reset" function, used to reset the internal memory of the node;
- one "step" function, taking as input the nodes inputs, and whose call performs one step of the node, updates the memory, and outputs the nodes outputs.

A standard way to execute Heptagon program is to compile the generated files together with a main program of the following scheme :

```
call the reset function
for each instant
  get the inputs values
  outputs ← step(inputs)
  do something with outputs values
```

Appendix A give specific technical details for each target language.

### 1.4 Simulation

A graphical simulator is available: hepts. It allows the user to simulate one node by providing a graphical window, where simulation steps can be performed by providing inputs of the simulated node.

This simulator tool interacts with an executable, typically issued of Heptagon programs compilation, and which await on the standard input the list of the simulated node's inputs, and prints its outputs on the standard output. Such executable, for the simulation of the node f, can be obtained by the -s <node> option:

```
> heptc -target c -s f example.ept
```

We can then directly compile the generated C program (whose main function stand in the \_main.c file):

```
> cd example_c
> gcc -Wall -c example.c
> gcc -Wall -c _main.c
> gcc -o f_sim _main.o example.o  # executable creation
```

This executable  $f_sim$  can then be used with the graphical simulator hepts, which takes as argument:

- The name of the module (capitalized name of the program without the .ept extension),
- the name of the simulated node,
- the path to the executable **f\_sim**.

```
> hepts -mod Example -node f -exec example_c/f_sim
```

## 2 Syntax and informal semantics

Heptagon programs are synchronous Moore machines, with parallel and hierarchical composition. The states of such machines define dataflow equations.

## 2.1 Nodes

Heptagon programs are structured in *nodes*: a program is a sequence of nodes. A node is a subprogram with a name f, inputs  $x_1, \ldots, x_n$ , outputs  $y_1, \ldots, y_p$ , local variables  $z_1, \ldots, z_q$  and declarations D.  $y_i$  and  $z_i$  variables are to be defined in D, using operations between values of  $x_j, y_j, z_j$ . The declaration of one variable comes with its type  $(t_i, t'_i \text{ and } t''_i)$  being the type of respectively  $x_i, y_i$  and  $z_i$ ).

```
node f(x_1:t_1;\ldots;x_n:t_n) returns (y_1:t'_1,\ldots,y_p:t'_p)
var z_1:t''_1,\ldots,z_q:t''_q;
let
D
tel
```

Heptagon allows to distinguish, by mean of clocks and control structures (switch, automata), for declarations and expressions, the discrete instants of activation, when the declarations and expressions are computed and progress toward further states, and other instants when neither computation nor progression are performed.

## 2.2 Expressions

### 2.2.1 Values and combinatorial operations

Heptagon is a dataflow language, i.e., every value, variable or constant, is actually a stream of value. The usual operators (e.g., arithmetic or Boolean operators) are applied pointwise on these sequences of values, as combinatorial operations (as opposed to *sequential* operations, taking into account the current *state* of the program: see delays in Section 2.2.2).

Thus, **x** denotes the stream  $x_1.x_2...$ , and **x** + **y** is the stream defined by  $(\mathbf{x} + \mathbf{y})_i = x_i + y_i$ .

| x     | $x_1$       | $x_2$       | $x_3$       | $x_4$       |  |
|-------|-------------|-------------|-------------|-------------|--|
| У     | $y_1$       | $y_2$       | $y_3$       | $y_4$       |  |
| x + y | $x_1 + y_1$ | $x_2 + y_2$ | $x_3 + y_3$ | $x_4 + y_4$ |  |

#### 2.2.2 Delays

Delays are the way to introduce some state in a Heptagon program.

• **pre**  $\mathbf{x}$  gives the value of  $\mathbf{x}$  at the preceding instant. The value at the first instant is undefined.

- x ->y takes the value of x at the first instant, and then the value of y;
- x fby y is equivalent to x ->pre y.

| x              | $x_1$   | $x_2$ | $x_3$ |
|----------------|---------|-------|-------|
| У              | $y_1$   | $y_2$ | $y_3$ |
| pre x          | $\perp$ | $x_1$ | $x_2$ |
| x ->y          | $x_1$   | $y_2$ | $y_3$ |
| x <b>fby</b> y | $x_1$   | $y_1$ | $y_2$ |

Here is a small example of a node that sums its inputs:

```
node sum(i:int) returns (o:int)
let
    o = 0 fby (o + i)
tel
```

#### 2.2.3 Clocks

It is possible to mix streams with different rates by having streams that are not present at each instant. This can be done using the **when** operator, that samples a stream according to a condition, either a boolean or an enumerated value. If x is a stream that is always present, then x when c (resp x when C(y)) is a stream equal to x but only present when c is true (resp. y = C). x whenot c is a shortcut for x when (not c). The split operator allows to sample a stream according to all possible values, i.e. split c (x) =x when c, x whenot c if x is a boolean and split y (x) =x when C1(y), ..., x when Cp(y) if y has an enumerated type defined by type t = $C1 | \ldots | Cp$ .

| x               | $x_1$ | $x_2$ | $x_3$ |
|-----------------|-------|-------|-------|
| С               | true  | false | true  |
| x <b>when</b> c | $x_1$ |       | $x_3$ |
| у               | C'    | C     | C'    |
| x when C(y)     | •     | $x_2$ |       |

The clock of an expression (resp. a stream) is a boolean stream that defines the instants when it is present. The clock of the streams that are always present is called **base** or .. If x has clock ck, denoted x :: ck, then x when c has clock ck on c.

The **merge** operator joins slow streams to create a faster stream.

| С           | true  | false | true  |
|-------------|-------|-------|-------|
| x           | $x_1$ | •     | $x_3$ |
| У           | •     | $y_2$ |       |
| merge c x y | $x_1$ | $y_2$ | $x_3$ |

**merge** expects complementary streams. If z = merge c x y and z :: ck, then we must have x :: ck on c and y :: ck onot c. It is thus different from **if** c **then** x **else** y that expects all its arguments to have the same clock. An analysis pass called *clock calculus* checks that these conditions are met.

Here is a first example of a bidirectional counter:

```
type modes = Up | Down
```

```
node two(m:modes;v:int) returns (o:int)
var x:int;
    x_up:int :: . on Up(m);
    x_down:int :: . on Down(m);
let
    o = 0 fby x;
    x_up = o when Up(m) + 1;
    x_down = o when Down(m) + 1;
    x = merge m (Up -> x_up) (Down -> x_down)
tel
```

Note that clocks are inferred by the compiler, so the clocking annotations are optional.

It is important to understand the interaction between sampling and delays. The value of a delay only changes when it is activated, that is, when its clock is true. As soon as a function f contains some delay operator, sampling its inputs is not equivalent to sampling its outputs, that is,  $f(x \text{ when } c) \neq (f x)$  when c.

| С                 | true | false | true | false |
|-------------------|------|-------|------|-------|
| counter(1)        | 0    | 1     | 2    | 3     |
| counter(1) when c | 0    | •     | 2    |       |
| counter(1 when c) | 0    |       | 1    |       |

## 2.3 Declarations

A declaration D can be either :

- an equation x = e, defining variable x by the expression e at each activation instants;
- a node application  $(y_1, \ldots, y_p) = f(e_1, \ldots, e_n)$ , defining variables  $y_1, \ldots, y_p$  by application of the node f with values  $e_1, \ldots, e_n$  at each activation instants;
- parallel declarations of  $D_1$  and  $D_2$ , denoted  $D_1$ ;  $D_2$ . Variables defined in  $D_1$  and  $D_2$  must be exclusive. The activation of this parallel declaration activate both  $D_1$  and  $D_2$ , which are both computed and both progress ;
- a switch control structure ;
- an automaton.

#### 2.3.1 Switch control structures

The switch control structure allows to controls which equations are evaluated:

```
type modes = Up | Down
node two(m:modes;v:int) returns (o:int)
var last x:int = 0;
let
    o = x;
    switch m
```

```
| Up do x = last x + v
| Down do x = last x - v
end
tel
```

The last keyword defines a memory which is shared by the different modes. Thus, last  $\mathbf{x}$  is the value of the variable  $\mathbf{x}$  in the previous instant, whichever was the activated mode.

#### 2.3.2 Automata

An automaton is a set of states (one of which being the initial one), and transitions between these states, triggered by Boolean expressions. A declaration is associated to each state. The set of variables defined by the automaton is the union, not necessarily disjoint (variables can have different definitions in different states, and can be partially defined : in this case, when the variable is not defined in an active state, the previous value of this variable is taken.

At each automaton activation instant, one and only one state of this automaton is active (the initial one at the first activation instant). The declaration associated to this active state is itself activated and progress in this activation instant.

**Example** The following example gives the node **updown**. This node is defined by an automaton composed of two states:

- the state  $\mathtt{Up}$  gives to  $\mathtt{x}$  its previous value augmented of 1
- the state Down gives to x its previous value diminued of 1

This automaton comprises two transitions:

- it goes from Up (the initial state) to Down when x becomes greater or equal than 10;
- it goes from Down to Up when x becomes less or equal 0.

```
node updown() returns (y:int)
var last x:int = 0;
let
y = x;
automaton
state Up
    do x = last x + 1
    until x >= 10 then Down
    state Down
    do x = last x - 1
    until x <= 0 then Up
    end
tel</pre>
```

| [ | current state | Up | Down | Down | Down |
|---|---------------|----|----|----|----|----|----|----|----|----|----|------|------|------|
|   | У             | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  | 9  | 10 | 9    | 8    | 7    |

Expressions on outgoing transitions of this active state are evaluated, so as to compute the next active state : these are weak transitions. Transitions are evaluated in declaration order, in the textual syntax. If no transition can be triggered, then the current state is the next active state.

## 2.4 Structured types

#### 2.4.1 Arrays

Arrays are denoted using the ^ symbol. For instance, int^10 is the type of arrays of size 10 containing integers. Arrays can be multidimensional, like int^3^2. One should note that indices appear in reverse order compared to C: int^3^2 should be understood as (int^3)^2, that is an array of size 2 containing arrays of size 3 (this would have been written int t[2][3] in C).

The following operators are defined for arrays:

- Accessing an element at a constant index: t[4]
- Accessing an element at a dynamic index with a default value: z = t.[x] default v is a stream defined by  $z_i = t_i[x_i]$  if  $0 \le x_i < n$  and  $z_i = v_i$  otherwise (where n is the size of t).
- Accessing an element at a truncated index: z = t[>x<] is defined by  $z_i = t_i[\min(\max(0, x_i), n-1)]$  (where n is the size of t).
- Modifying an element: t' = [t with [x] = v] is a new array equal to t, except for the element at index x which is equal to v, that is for all j in [0, n 1] such that  $j \neq x_i$ ,  $t'_i[j] = t[j]$  and  $t'_i[x_i] = v_i$ .
- Defining an array by copying one value:  $\mathbf{x}^n$  is the array of size  $\mathbf{n}$  whose elements are all equal to  $\mathbf{x}$ .
- Defining an array explicitely: [1, x, 3, y, 5].
- Extracting a slice: t[n..m] returns the sub array of t of size m-n+1 starting at index n and ending at index m.
- Concatening arrays: t1@t2.

It is also possible to use *iterators* to operate on arrays. For instance, one can add two arrays pointwise by doing t =map<<n>> (+)(t1, t2) (where n is the size of t1, t2 and t). The following iterators are defined:

- map applies a node pointwise to the input arrays and outputs one or several arrays. If f has type t1 \* t2 ->t'1 \* t'2 then map<<n>> f has type t1^n \* t2^n ->t'1^n \* t'2^n.
- mapi is the same as map but the iterated function should expect another integer argument, which will be equal to the index of the inputs in the input arrays. If f has type t1 \* t2 \* int ->t'1 \* t'2 then map<<n>> f has type t1^n \* t2^n ->t'1^n \* t'2^n.
- fold iterates a node on an array accumulating values. If f has type t1 \* t ->t then fold<<n>> f has type t1^n \* t ->t. For instance, fold<<2>> f(t, 0) is equivalent to f(t[1], f(t[0], 0)).
- foldi is to fold what mapi is to map.
- mapfold combines the result of map and fold, by accumulating a value and outputting a new array. If f has type t1 \* t ->t'1 \* t then mapfold<<n>> f has type t1^n \* t ->t'1^n \* t.

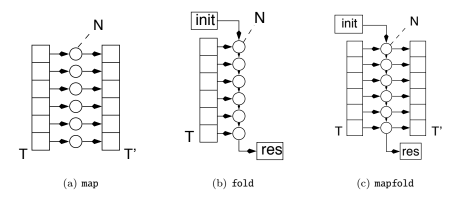


Figure 1: Iterators in HEPTAGON

Iterators can also be used for multidimensional arrays. In the case of **mapi** and **foldi**, the iterated function should then expect one argument for each dimension.

It is also possible to mimic partial application with iterators, by giving a list of arguments, not necessarily arrays, that will be given as first arguments to each application of the iterated function. For instance, o =map <<n>f <(a)>(t) is equivalent to o[i] =f(a, t[i]) for each element in the output array o.

Arrays are semantically functional: each modification on an array creates a new array. Implementing these arrays using separate arrays can lead to very unefficient code. That's why an optimization pass called *memory allocation* [1] tries to share arrays and avoid unnecessary copies within each node. It can be enabled using the **-memalloc** or **-0** option.

#### 2.4.2 Records

The syntax for declaring record types is the following:

type t = { f1: t1; f2: t2 }

Note that two different record types cannot have fields with the same name. It is possible to declare a new record ( $\{ f1 = x; f2 = 9 \}$ ), read a field of a record (x.f1) and modify one field  $(r' = \{ r with .f1 = v \}$  returns a new record where all fields are equal to the ones in r except for f1 which is equal to v).

#### 2.4.3 Types alias

It is possible to declare an alias for any type (like a typedef in C), for instance:

type meter = int
type matrix = int^10^10

This alias can then be used anywhere a type is expected.

## 2.5 Parametricity

The size of arrays can be parametrized by so-called *static expressions*, that is expressions that are reduced to constants at compile-time. They are either a literal, a global constant, a static

parameter of a node or a static operation on two static expressions (+, -, \*, /, +., <=, **or**, etc). Global constants are declared by:

const n : int = 100
const t0 : float^n = 1.0^n
const r0 = { f1 = 0; f2 = t0 }

The parameters of a node are given between << and >> at the declaration of the node and are instantiated with the same syntax:

```
node f<<m:int; t1: int^n>>(a:int^m) = (o:int^m)
let
    o = map<<m>> (+)(a, t1);
tel
node g(a:int^n) = (o:int^n)
let
    o = f<<n, t0>>(a);
tel
```

If the backend support parametricity (like in Java), static parameters are kept in the generated code. Otherwise, a pass of the compiler generates all the versions of the node that are needed. If a parametrized node defined in a file f1.ept is used in f2.ept, it is necessary to first compile f1.ept with the -c option (and without any -target), in order to generate a binary file f1.epo. The compilation of the second file, this time with the -target option, will generate code for all the necessary nodes, from f1.ept and f2.ept.

#### 2.6 Location annotations

Memory allocation [1] avoids unnecessary array copies within the nodes automatically (it can be enable with the -memalloc or -O options). In order to avoid copies when calling nodes, the user must add *location annotations*. We will give here only a short introduction, the interested reader can refer to [1] for more details. They express the fact that a function modifies in-place its argument. For instance, consider this node that swaps two elements in an array:

```
node swap(i, j:int; t_in:float^100 at r) returns (t_out:float^100 at r)
var t_tmp:float^100 at r;
let
   t_tmp = [ t_in with [i] = t_in[>j<] ];
   t_out = [ t_tmp with [j] = t_in[>i<] ];
tel</pre>
```

The location annotations are introduced by the keyword **at** followed by a location name. All the streams with the same location name are guaranteed to be stored together in the generated code, so the function generated for **swap** will directly modify its argument in-place.

Located variables are called *semilinear*: they can only be updated once, but they can be read many times. An update is a function that modify its argument in-place, for instance a node that inputs and outputs located variables, and a read is any other function. For instance, modifying one element in an array is an update, but accessing one element is a read.

Only located variables can be given to a function that expects located arguments. A located variable can only be obtained by updating another located variable or by explicitly initializing a new location with the **init** construction:

```
node shuffle(i_arr, j_arr : int^m; q : int) = (v : float)
var t, t_prev : float^n at r;
let
    init<<r>> t_prev = t_0 fby t;
    t = fold<<m>> swap(i_arr, j_arr, t_prev);
    v = t[>q<];
tel</pre>
```

## 2.7 Interfaces

Interface files, with the extension **.epi**, do not contain any definition but only the declaration (or signatures) of nodes, types and constants. In particular, they are useful to import external functions. For instance:

external fun sort(a:int^100) = (o:int^100)

The imported function should respect the calling convention given in appendix A. See the directory examples/extern\_C for a complete example.

## **3** BZR: Contracts for controller synthesis

Contracts are an extension of the Heptagon language, so as to allow to perform discrete controller synthesis on Heptagon programs. The extended language is named BZR.

We associate to each node a *contract*, which is a program associated with two outputs :

- an output  $e_A$  representing the environment model ;
- an invariance objective  $e_G$ ;
- a set  $\{c_1, \ldots, c_n\}$  of controllable variables used for ensuring this objective.

This contract means that the node will be controlled, i.e., that values will be given to  $c_1, \ldots, c_n$ such that, given any input trace yielding  $e_A$ , the output trace will yield the true value for  $e_G$ . Contracts are denoted :

```
node f(x_1:t_1;...;x_n:t_n) returns (y_1:t'_1;...;y_p:t'_p)
contract
var ...
let
...
tel
assume e_A
enforce e_G
with (c_1:t''_1;...;c_q:t''_n)
var ...
let
y_1 = f_1(x_1,...,x_n,c_1,...,c_q);
:
```

 $y_p = f_p(x_1, ..., x_n, c_1, ..., c_q);$ tel

## 4 BZR Running Example: Multi-task System

### 4.1 Delayable Tasks

We consider a multi-task system composed of n delayable tasks. Figure 2 shows a delayable task. A delayable task takes three inputs  $\mathbf{r}$ ,  $\mathbf{c}$  and  $\mathbf{e}$ :  $\mathbf{r}$  is the task launch request from the environment,  $\mathbf{e}$  is the end request, and  $\mathbf{c}$  is meant to be a controllable input controlling whether, on request, the task is actually launched (and therefore goes in the active state), or delayed (and then forced by the controller to go in the waiting state by stating the false value to  $\mathbf{c}$ ). This node outputs a unique boolean **act** which is true when the task is in the active state.

```
node delayable(r,c,e:bool) returns (act:bool)
let
    automaton
    state Idle
    do act = false
    until r & c then Active
        | a & not c then Wait
    state Wait
    do act = false
    until c then Active
    state Active
    do act = true
    until e then Idle
    end
    tel
```

Figure 2: Delayable task

The Figure 3 shows then a node **ntasks** where n delayable tasks have been put in parallel. The tasks are inlined so as to be able to perform DSC on this node, taking into account the tasks' states. Until now, the only interest of modularity is, from the programmer's point of view, to be able to give once the delayable task code.

This **ntasks** node is provided with a contract, stating that its composing tasks are exclusive, i.e., that there are no two tasks in the active state at the same instant. This contract is enforced with the help of the controllable inputs  $c_i$ .

#### 4.2 Contract composition

We want know to reuse the **ntasks** node, in order to build modularly a system composed of 2n tasks. The Figure 4 shows the parallel composition of two **ntasks** nodes. We associate to this composition a new contract, which role is to enforce the exclusivity of the 2n tasks.

It is easy to see that the contract of **ntasks** is not precise enough to be able to compose several of these nodes. Therefore, we need to refine this contract by adding some way to externally control the activity of the tasks.

```
node ntasks(r<sub>1</sub>,..., r<sub>n</sub>, e<sub>1</sub>,..., e<sub>n</sub>:bool)
returns (a<sub>1</sub>,..., a<sub>n</sub>:bool)
contract
let
ca<sub>1</sub> = a<sub>1</sub> & (a<sub>2</sub> or ... or a<sub>n</sub>);
:
ca<sub>n-1</sub> = a<sub>n-1</sub> & a<sub>n</sub>;
tel
enforce not (ca<sub>1</sub> or \ldots or ca<sub>n-1</sub>)
with (c<sub>1</sub>,..., c<sub>n</sub>:bool)
let
a<sub>1</sub> = inlined delayable(r<sub>1</sub>, c<sub>1</sub>, e<sub>1</sub>);
:
a<sub>n</sub> = inlined delayable(r<sub>n</sub>, c<sub>n</sub>, e<sub>n</sub>);
tel
```

Figure 3: ntasks node: n delayable tasks in parallel

## 4.3 Contract refinement

We first add an input c, meant to be controllable. The refined contract will enforce that:

- 1. the tasks are exclusive,
- 2. one task is active only at instants when the input c is true. This property, appearing in the contract, allow a node instantiating **ntasks** to forbid any activity of the n tasks instantiated.

The Figure 5 contains this new **ntasks** node.

However, the controllability introduced here is know too strong. The synthesis will succeed, but the computed controller, without knowing how c will be instantiated, will actually block every tasks in their idle state. Indeed, if the controller allows one task to go in its active state, the input c can become false at the next instant, violating the property to enforce.

Thus, we propose to add an assumption to this contract: the input c will not become false if a task was active an instant before. This new contract is visible in Figure 6.

We can then use this new **ntasks** version for the parallel composition, by instantiating the **c** input by a controllable variable and its negation. This composition can be found in Figure 7.

```
node main(r_1, \ldots, r_{2n}, e_1, \ldots, e_{2n}:bool)
returns (a_1, \ldots, a_{2n}:bool)
contract
let
ca<sub>1</sub> = a<sub>1</sub> & (a<sub>2</sub> or ... or a<sub>2n</sub>);
:
:
ca<sub>2n-1</sub> = a<sub>2n-1</sub> & a<sub>2n</sub>;
tel
enforce not (ca<sub>1</sub> or ... or ca<sub>2n-1</sub>)
let
(a_1, \ldots, a_n) = ntasks(r_1, \ldots, r_n, e_1, \ldots, e_n);
(a_{n+1}, \ldots, a_{2n}) = ntasks(r_{n+1}, \ldots, r_{2n}, e_{n+1}, \ldots, e_{2n});
tel
```

Figure 4: Composition of two ntasks nodes

```
node ntasks(c,r<sub>1</sub>,...,r<sub>n</sub>,e<sub>1</sub>,...,e<sub>n</sub>:bool) returns (a<sub>1</sub>,...,a<sub>n</sub>:bool)
contract
let
    ca<sub>1</sub> = a<sub>1</sub> & (a<sub>2</sub> or ... or a<sub>n</sub>);...
    ca<sub>n-1</sub> = a<sub>n-1</sub> & a<sub>n</sub>;
    one = a<sub>1</sub> or ... or a<sub>n</sub>;
tel
enforce not (ca<sub>1</sub> or ... or ca<sub>n-1</sub>) & (c or not one)
with (c<sub>1</sub>,...,c<sub>n</sub>:bool)
let
    a<sub>1</sub> = inlined delayable(r<sub>1</sub>,c<sub>1</sub>,e<sub>1</sub>);
    :
    a<sub>n</sub> = inlined delayable(r<sub>n</sub>,c<sub>n</sub>,e<sub>n</sub>);
tel
```

Figure 5: First contract refinement for the **ntasks** node

```
node ntasks(c,r<sub>1</sub>,...,r<sub>n</sub>,e<sub>1</sub>,...,e<sub>n</sub>:bool) returns (a<sub>1</sub>,...,a<sub>n</sub>:bool)
contract
let
    ca<sub>1</sub> = a<sub>1</sub> & (a<sub>2</sub> or ... or a<sub>n</sub>);...
    ca<sub>n-1</sub> = a<sub>n-1</sub> & a<sub>n</sub>;
    one = a<sub>1</sub> or ... or a<sub>n</sub>;
    pone = false fby one;
    tel
    assume (not pone or c)
    enforce not (ca<sub>1</sub> or ... or ca<sub>n-1</sub>) & (c or not one)
    with (c<sub>1</sub>,...,c<sub>n</sub>)
let
    a<sub>1</sub> = inlined delayable(r<sub>1</sub>,c<sub>1</sub>,e<sub>1</sub>);
    :
    a<sub>n</sub> = inlined delayable(r<sub>n</sub>,c<sub>n</sub>,e<sub>n</sub>);
tel
```

Figure 6: Second contract refinement for the ntasks node

```
node main(r_1, \ldots, r_{2n}, e_1, \ldots, e_{2n}:bool) returns (a_1, \ldots, a_{2n}:bool)
contract
let
ca_1 = a_1 & (a_2 or ... or a_{2n});
:
ca_{2n-1} = a_{2n-1} & a_{2n};
tel
enforce not (ca_1 or ... or ca_{2n-1})
with (c:bool)
let
(a_1, ..., a_n) = ntasks(c, r_1, ..., r_n, e_1, ..., e_n);
(a_{n+1}, ..., a_{2n}) = ntasks(\Not c, r_{n+1}, ..., r_{2n}, e_{n+1}, ..., e_{2n});
tel
```

```
Figure 7: Two ntasks parallel composition
```

## A Generated code

### A.1 C generated code

C generated files from an Heptagon program **example.ept** are placed in a directory named **example\_c**. This directory contains one file **example.c**. For each node **f** of the source program, assuming that **f** has inputs  $(x_1 : t_1, \ldots, x_n : t_n)$  and outputs  $(y_1 : t'_1, \ldots, y_p : t'_p)$ ,  $t_i$  and  $t'_i$  being the data types of these inputs and outputs, then the **example.c** file contains, for each node **f**:

• A **Example\_\_f\_reset** function, with an argument **self** being a memory structure instance:

void Example\_\_f\_reset(Example\_\_f\_mem\* self);

• A Example\_\_f\_step function, with as arguments the nodes inputs, a structure \_out where the output will be put, and a memory structure instance self:

```
void Example__f_step(t_1 x_1, ..., t_n x_n,
Example__f_out* \_out,
Example__f_mem* self);
```

After the call of this function, the structure **\_out** contains the outputs of the node:

typedef struct \{
 t'\_1 y\_1;
 ...
 t'\_p y\_p;
 Ly Example\_\_f\_ans;

#include "example.h"

An example of main C code for the execution of this node would be then:

```
int main(int argc, char * argv[]) \{
    Example__f_m mem;
    t<sub>1</sub> x<sub>1</sub>;
    ...
    t<sub>n</sub> x<sub>n</sub>;
    Example__f_out ans;
    /* initialize memory instance */
    f_reset(&mem);
    while(1) \{
        /* read inputs */
        scanf("...", &x<sub>1</sub>, ..., &x<sub>n</sub>);
        /* perform step */
        Example__f_step(x<sub>1</sub>, ..., x<sub>n</sub>, &ans, &mem);
        /* write outputs */
```

The above code is nearly what is produce for the simulator with the -s option (see Section 1.4).

## A.2 Java generated code

Java generated files from an Heptagon program **example.ept** are placed in a directory named **example\_java**. This directory contains one Java class  $\mathbf{f}$  (in the file  $\mathbf{f}$ .java) for each node  $\mathbf{f}$  of the source program. Assuming that  $\mathbf{f}$  has inputs  $(x_1 : t_1, \ldots, x_n : t_n)$  and outputs  $(y_1 : t'_1, \ldots, y_p : t'_p)$ ,  $t_i$  and  $t'_i$  being the data types of these inputs and outputs, then this  $\mathbf{f}$  class implements the following interface:

```
public interface f {
    public void reset();
    public fAnswer step(t<sub>1</sub> x<sub>1</sub>, ..., t<sub>n</sub> x<sub>n</sub>);
}
```

The **fAnswer** class being a structure containing the outputs:

```
public class fAnswer {
    t'_1 y_1;
    ...
    t'_p y_p;
}
```

## References

[1] Léonard Gérard, Adrien Guatto, Cédric Pasteur, and Marc Pouzet. A modular memory optimization for synchronous data-flow languages. In *LCTES'12*, 2012.