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WCET Nested-Loop Minimization in Terms of Instruction-Level-Parallelism

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Abstract—Several high-performance applications integrate loop bodies which represent the most critical sections. This aspect brings two challenges. First, the Worst Case Execution Time (WCET) must be determined in order to define the nested loop timing behaviour. The second challenge consists in raising the parallelism-level to enhance the performance. In particular, the Multidimensional Retiming (MR) is an important optimization approach that offers several instruction-level-parallelism solutions. Despite the fact that full parallelism allows achieving the optimal WCET, it leads to a high growth in processing cores, which is inadequate to embedded real-time implementations.

The main idea of this paper consists in driving the parallelism-level rise in terms of WCET development. First, the MR parameters that correspond to the nested loops are extracted. Thereafter, the WCET is formulated in term of the parallelism level rise. Then, an optimization heuristic is proposed which identifies the parallelism level that allows respecting the WCET constraint. Our experiments indicate the WCET prediction is accurate within an error rate of 8.54%. Secondly, the optimization heuristic implementations show an average improvement on number of cores of 27.18% compared to full parallel ones.

Keywords— HW/SW codesign, design space exploration, optimization, WCET, nested loops, parallelism.

I. INTRODUCTION

The loop bodies compose a large group of embedded real-time applications. They imply a higher rise in the execution time, which generally results in overtaking the real-time properties. Therefore, the performance is handled through all the design levels in terms of analysis and optimization, to ensure a safe final implementation.

Several WCET estimation approaches are proposed, which are generally classified in two kinds [5]. The first one consists in running test codes and estimating the WCET, based on the measured performances. However, those approaches are inefficient in the case of complex codes. The other approaches are based on a WCET static analysis. They model the implementation code in order to identify the data flow and the control flow parameters and use them on an equation system. The WCET static analyses are integrated on a general workflow that intercepts both code and micro-architecture parameters to provide a safe WCET [1,2]. Several workflows are implemented in efficient software tools and compilers for real-time applications [4, 5, 19].

Actually, the real-time application complexity is increased due to an intensive utilization of nested loops. Moreover, parallel architecture implies different parallel scheduling, hence different timing behaviours. In this context, many works have focused on analysing the nested loop structures in order to propose adequate equation systems for WCET estimation [6, 7, 8], which are in the scope of our work. The main idea is to explore the nested loop model to define the loop iteration numbers, the loop dependencies and the critical task inter-loops, etc. Some works have focused on specific loop bodies such as the one containing unfeasible paths like “if … else” instruction and “do … while” loop [2, 6], which are not considered in the paper.

Elsewhere, several parallelism approaches have led to enhance the nested loop performances. They generally represent the data flow and the control flow in a formal model such as the polyhedral model [9, 10] and the Multidimensional Data Flow Graph (MDFG) [11, 12]. Thereafter, they transform the model to parallelize the processing. The MDFG is an acyclic data flow graph that ensures an adequate representation of nested loops. It allows modelling the loop-carried dependencies across several loops. It is distinguished by a detailed instruction granularity compared to the polyhedral model. A lot of optimization approaches are proposed to parallelize the MDFG. The Multidimensional Retiming (MR) is a theoretical approach that guarantees parallelizing instructions inside nested loops. All MR techniques aim to iteratively increasing the instruction-level-parallelism until achieving a full parallel implementation, hence the minimal WCET [12, 13, 14]. However, the higher the parallelism level is, the more intensively the processing core number grows. Moreover, admitting that the application complexity keeps growing, the parallelism transformation implies a data overhead which requires a large cache memory size. As a result, even the optimal WCET is achieved, the MR provides implementations with important material resources.

In fact, the MR approach offers a significant solution space with a corresponding different parallelism level. Consequently, it is sufficient to apply a partial parallelism that enables attaining the WCET constraint, instead of applying a
full parallelism. However, it is inefficient to extract each solution and then define its WCET, due to the nested loop complexity. The works described in [17, 18] partially explore the solution space to provide a safe implementation, but they do not give optimal solutions. Few works have been interested in analyzing the WCET development in terms of parallelism techniques. As described in [15], the work combines the theory of both WCET model and loop unrolling to enhance the performance. Furthermore, an instruction-level-parallelism controlled by the WCET development is suggested for superscalar processors in [16]. However, both works were proposed for specific nested loops without modeling general loop-carried dependencies.

In this paper, we aim to drive the instruction-level-parallelism rising of nested loops by the WCET development. First of all, the WCET equation system is formulated in terms of MR parameters. Then, the parallelism level is increased iteratively until respecting the WCET constraint.

The rest of the paper is organized as follows. In section II, we present the basic concepts of modelling nested loops with MDFG, parallelizing the loop body with MR and predicting nested loop WCET. In section III, we present the theory of the WCET estimation and the optimization heuristic that drives MR in terms of WCET. The experimental results are given in section IV, followed by the concluding remarks in section V.

II. BASIC CONCEPTS

A. Multidimensional data flow graph

The MDFG is an extension of the classic data flow graph where each node presents an instruction and each edge presents a data dependency. The main characteristic is the ability to represent nested iterative and recursive structures. Indeed, one iteration is similar to executing all nodes once. The repetitive aspect is formulated by two concepts: the dimension $n$ of the MDFG, which is the nested loop number, and the delays, which are edge weights formulating the loop-carried dependencies. An MDFG is modelled as $G = (V, E, d, t)$, where $V$ is the node set, $E$ is the edge set, $d(e)$ is the multidimensional delay of edge $e$, and $t(u)$ is the computation time of the node $u$. For $e : u \rightarrow v$, $A d(e)$ edge delay is modelled by a vector with $n$ indexes such as $d(e) = (c_1, c_2, ..., c_n)$. Each index corresponds to a single loop in a way that $c_k$ presents the difference between the iteration executing $v$ and the other one executing $u$ of the loop $k$: If the node $u$ is executed in the iteration $x$ of the loop $k$, then the node $v$ is executed in the iteration $(x + c_k)$.

As an example, the Jacobi algorithm [25] shown in Fig. 1(a) is modelled by the MDFG in Fig. 1(b). It is composed by four nodes like the number of the innermost loop instructions. Each edge in the MDFG is labelled by a delay with two indexes, where $d(e) = (d.x, d.y)$. The 'd.x' and 'd.y' terms are in relation with the outermost loop and the innermost one, respectively. The A1 and A2 instructions are computed in the same iteration whether for the innermost or the outermost loops. Therefore, the A1 $\rightarrow$ A2 edge is labelled by the delay $d(e_1) = (0, 0)$ which is called “zero-delay edge”. For the data dependencies between D2 and A2, if D2 is executed in the iteration $i$ of the outermost loop, then A2 is executed in the iteration $i + 1$. Similarly to the innermost loop, D2 is executed in the previous iteration of the A2 execution. For this purpose, the delay value of the edge D2 $\rightarrow$ A2 is equal to $(1, -1)$.

The notation $p : v_i \rightarrow \ldots \rightarrow v_h$ is used to mean that $p$ is a path from $v_i$ to $v_h$. The delay and the computation time of a path $p$ are respectively equal to $d(p) = \sum_{k=e}^{n} d(e_k)$ and $t(p) = \sum_{k=e}^{n} t(e_k)$. A $p$ path, whose $d(p) = (0, ..., 0)$, is called “zero-delay path”. The critical paths $p_{cr}$ are the ones having the maximum computation time among all zero-delay paths in the MDFG ($t(p_{cr}) = \max t(p, d(p) = 0)$). The period during which all computation nodes in iteration are executed according to existing data dependencies and without resource constraints is called an Iteration Time $IT(G)$, where $IT(G) \geq t_{p_{cr}}$. For example, the critical paths is structured as $p : A1 \rightarrow A2 \rightarrow D1 \rightarrow D2$. Assuming that $t(A1) = t(A2) = 1$ time units (t. u.) and $t(D1) = t(D2) = 2$ t. u., the Jacobi algorithm $IT$ is equal to 6 t.u. The theoretical schedule is represented in Fig. 2 where the nodes belonging to the same iteration are modelled by the same pattern which can be executed using one core.

![Fig.1. The Jacobi algorithm: (a) code, (b) MDFG, (c) static schedule](image)

B. Multidimensional retiming

The MR is an instruction-level-parallelism approach of nested loops. It reduces the $IT$ by decreasing nodes in critical paths. With this objective, it shifts nodes from their original iteration in order to execute them in parallel with other nodes. For a retimed node, modifying its belonging iteration consists in modifying their edge delays with respect to a MR function. For the MR function $r(u) = (r_1, ..., r_n)$, the execution of the node $u$ in the iteration $i$ is moved to the iteration $i - r_u$ [12]. Thus, the MR is modelled as a graphical transformation that modifies the MDFG delays, while preserving the functional behaviour of the initial MDFG [12,13]. All MR techniques aim to achieve the $IT_{min}$. The incremental and chained MR techniques [12] apply the MR successively to each critical path nodes until having a full parallelism; i.e., all instructions in the same iterations are executed in parallel. However, the provided implementation requires processing cores as more as the nodes are.

Other MR techniques lead to schedule an MDFG with a $IT_{min}$, without getting a full parallelism [12, 14, 27]. Retiming the whole path is carried out, instead of a node only, in order
to minimize the processing cores when enhancing the performance. The delayed MR technique \cite{14, 27} proposes a theoretical approach to select and retime paths. It defines two terms to reflect the timing and data dependency characteristics of each path in the MDFG: the $D(u, v)$ which represents the minimum delay between the paths connecting $u$ and $v$, as described in Equation (1) (if just one path among those connecting $u$ and $v$ has a zero-delay then $D(u, v, v) = 0$); else, $D(u, v) \neq (0, \ldots, 0)$, and the $T(u, v)$ which defines the maximum execution time among the zero delay path connecting $u$ and $v$, as described in Equation (2).

$$D(u, v) = \min\{d(p), \text{where } p : u \rightarrow v, u \in V \text{ and } v \in V\}$$

$$T(u, v) = \max\{t(p), \text{where } p : u \rightarrow v, u \in V, v \in V \text{ and } d(p) = D(u, v)\}$$

Their values are ranged respectively in two matrices called $D$ and $T$ with $V \times V$ size, such as $V$ in the node set. Each $p : u \rightarrow v$ path is indexed by the cell with the line $u$ and the column $v$. Taking the Jacobi algorithm as an example, the MDFG in Fig. 1(b) is composed by four nodes, hence the $4 \times 4$ matrix dimension whose $D$ and $T$ matrices are respectively illustrated in Table I and Table II.

<table>
<thead>
<tr>
<th>$u, v$</th>
<th>$A1$</th>
<th>$A2$</th>
<th>$D1$</th>
<th>$D2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$A1$</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
</tr>
<tr>
<td>$A2$</td>
<td>(1, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
</tr>
<tr>
<td>$D1$</td>
<td>(1, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
</tr>
<tr>
<td>$D2$</td>
<td>(1, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
<td>(0, 0)</td>
</tr>
</tbody>
</table>

Thus, achieving the $IT_{min}$ is synonymous to providing an MDFG whose $D(u, v) = 0$ and $T(u, v) > IT_{min}$. Therefore, the delayed MR technique explores both matrices to define the sub-critical paths with respect to the previous condition. For the Jacobi algorithm, the delayed MR selects the MR function $r = (0, 1)$ to software pipeline the $P1: A1 \rightarrow A2$ path, which the retimed Jacobi algorithm is shown in Figure 2(a). Each $i^{th}$ occurrence of $P1$ path is shifted up and executed in the previous iteration of the innermost loop, where $1 \leq i \leq n$. The first $P1$ occurrence belonging to the first iteration of the innermost loop is shifted upstream the retimed loop as the first instruction in Fig. 2(a), which is called prologue. This implies that all $P1$ paths, belonging to $(i, 0)$ iteration, are executed outside the innermost loop whatever the $i$ index is. Correspondingly, the complementary instructions of the last iteration, which are called epilogue, are executed downstream the innermost loop. This transformation permits executing any $P1$ path in the innermost loop in parallel with other nodes, as shown in the static schedule of Fig. 2(c). Accordingly, the $IT$ is reduced from 6 to 4. In addition, the innermost loop is iterated $(N - 1)$ times, though requiring two cores to execute the implementation.

For $i$ from 1 to $N$ do
$A1: X := X + \frac{1}{2}(A1 + A2)$
$A2: Y := A2 + A1$
End for

For $i$ from 1 to $N$ do
$A1: X := X + \frac{1}{2}(A1 + A2)$
$A2: Y := A2 + A1$
End for

Thus, the second MR function is applied to the $D1$ node. The final MDFG is scheduled with the $IT_{min} = 2$ $t. u.$ using three processing cores as shown in the Fig.3.

$$WCRT = WCRT + WCST \quad (3)$$

The WCRT static analysis explores the software parameters of the loop nests. The equation system is always formulated using the computation time $t_i$ of all $i$ tasks and their instance numbers $c_i$ \cite{1, 4, 5, 21, 22, 23}. Each task is defined by exploring the control flow in order to define the critical path instructions. Therefore, its execution time $t_i$ is determined by associating a timing parameter for each computation instruction. The $c_i$ instance number is computed based on the iteration loop bounds that are identified from the code. In the case of parallel processing, the WCRT consists in selecting the task executed in parallel to identify the ones having the maximal computation time \cite{3, 20, 24}. Thus, the whole WCRT value is generally the maximal value of multiplying $t_i$ and $c_i$ parameters, as indicated in Equation 4. In the case of nested loops, Equation 4 is enabling its application by considering each loop as a task in its outermost one.

$$WCRT = \max \sum t_i \times c_i \quad (4)$$
The WCST represents the necessary time to schedule data to parallel cores, which is in terms of the thread number. Accordingly, the more the threads are, the higher the stall time is. Moreover, nested loops iterate the execution of the parallel processing which leads to a similar stall time rise, resulting in computing the WCST as indicated in Equation 5 [4, 20].

\[ WCST = N \times (T - 1) \times C (5) \]

Where \( N \) is the iteration number, \( T \) is the thread number and \( C \) is timing parameter that is experimentally defined as in [4, 20].

III. WCET DEVELOPMENT IN TERMS OF MULTIDIMENSIONAL RETIMING

A. Approach principle

The main idea of this paper consists in driving the instruction-parallelism-level rise in terms of WCET development. For this purpose, our approach requires an efficient WCET estimation.

As indicated in the last paragraph, the WCRT is expressed in terms of computation times and instance numbers of tasks. For the MDFG, those tasks corresponding to the instructions belong either to the innermost loop or the prologue/epilogue codes. Instance numbers are to be computed directly from their outermost loop bounds, irrespective of their number. As indicated in section II, to obtain iteration time, the identification of the critical paths is vital. For the WCST, it depends on the thread number that ensures implementing any code block, which is similar to the parallelism level. In this objective, the MDFG is to be swept in order to compute the maximal node count executed in parallel.

Elsewhere, the MR iteratively increases the parallelism level by software pipelining nodes. Accordingly, each MR function consists in decreasing the loop bounds, modifying the data paths either inside or both side loops, and increasing the parallelism level. Moreover, those transformations rise with respect to the applied MR function number. In this objective, the approach formulates the loop bound, the iteration time and the parallelism level developments into equations in terms of MR function and its application numbers. Thereafter, those equations are employed to estimate the WCET nested loops.

Based on the MR modification and equations 4 and 5, the MR leads to reduce the WCET despite increasing the parallelism level, hence the core rise. Thus, the approach proceeds iteratively to estimate the WCET and raises the parallelism level using the MR until achieving the WCET constraint or the \( IT_{\min} \).

Within this framework, figure 4 illustrates a step-by-step approach. It starts by providing an optimal MR function \( r \) and defining the \( IT_{\min} \) [12, 13, 14]. Then, it sweeps the MDFG to identify the maximal number of MR that can be applied and their corresponding data paths that will be shifted, as described in section B. In section C, those data paths are explored to compute the iteration times \( IT_{k} \) and the parallelism levels \( P_{LR} \), which correspond to each MR order. The development of the iteration loop bounds \( LB \) is formulated in terms of MR retiming function in section D. As a consequence, the WCET prediction of the whole nested loops is then detailed in section E. Finally, an optimization heuristic is proposed in section F, which drives the instruction-level-parallelism in terms of WCET development.

![Fig. 4. The approach flow graph](image)

B. Multidimensional retiming analysis

Each MR consists in reducing the critical path sizes by shifting nodes from the innermost loop and adding them in both sides of loop nests. Both actions affect the timing behaviour of the nested loops. To evaluate this modification, it is necessary to identify the possible MR modifications and the data-paths that are shifted. Those sub-paths can be identified by their latest nodes \( R \) with respect to the data dependency direction, where and \( R \subseteq V \). Accordingly, this section objective is defining a set \( V_{R} \) of vector \( (i, u) \) where \( u \) is the node to retime and \( i \) is its MR order. In fact, those nodes are defined based on the timing and data dependency characteristics of the MDFG which are modelled on the D and T matrices, as described in section II. Thus, both matrices are swept to define the maximal number of MR functions, the retimed nodes and their MR function order. The D and T matrices are incrementally swept in the direction of the data dependencies. The node set \( S \), having all incoming edges with non-zero-delays, are extracted in order to identify the last nodes \( R \) of the critical paths. So, each node is added to the set \( R \) if the computation time path is under \( IT_{\min} \) and \( D = (0, \ldots, 0) \) [13, 14]. Accordingly, the \( R \) list is filtered to remove redundancy and save the last ones. The first MR function adds delays to the \( R \) node outgoing edges. Hence, the \( R \) nodes are saved in the \( V_{R} \) list with their MR function order \( (k = 1) \). These steps are repeated by sweeping from the R successors and incrementing the \( k \) level, as described in algorithm 2.

For example, the initial MDFG of the Jacobi algorithm, whose the critical path is \( p_{ch}: A1 \rightarrow A2 \rightarrow D1 \rightarrow D2 \), is shown in Fig.1(b). As indicated in algorithm 1, sweeping leads to select the A2 node to be retimed first. The second iteration in algorithm 1 implies the selection of the D1 node to be the second one to retime. As a result, \( V_{R} = \{(A2, 1), (D1, 2)\} \).
C. Innermost IT and parallelism level in terms of MR

WCET prediction requires defining the innermost iteration time and the parallelism level for the initial MDFG and for each eventual MR function. Therefore, the parallelism levels are modeled as a set \( PL_{IT} \) of vector \( (PL_{loop},PL_{pro},PL_{epi}) \) where \( i \) is the MR order, and \( PL_{loop},PL_{pro} \) and \( PL_{epi} \) are the parallelism levels respectively of the innermost loop, the prologue and the epilogue. Similarly, the innermost ITs are modeled as a set \( IT_{IT} \) of vector \( (IT_{IT_1}) \). Based on the MR principles, the instructions added to the prologue are executed under \( IT_{IT_1} \). Therefore, the prologue and epilogue computation time are estimated in terms of \( IT_{IT_1} \) and the current MR order.

The IT is equal to the computation time of the critical path, as \( IT = \max \{ t(p), d(p) \} \). For the initial MDFG, the critical path is defined by sweeping it from the nodes \( V_D \), having all incoming edges with non-zero-delays, until the nodes \( V_A \), having all outgoing edges with non-zero-delays. Each MR modifies the critical-path in a way that it is defined from the retimed nodes \( V_R \), extracted by algorithm 1, to the \( V_A \) nodes. Similarly, the parallelism level of the initial MDFG corresponds to the maximal nodes that are executed in parallel, which are determined basing on their computation times. Taking as an example the Jacobi algorithm shown in Fig.1, the MDFG is swept to identify the path \( A_1 \rightarrow A_2 \rightarrow D \rightarrow D_2 \) critical path. Knowing that the first MR is to be applied to the \( A_2 \) node, sweeping is done from the \( A_2 \) successor nodes to the \( V_A \) nodes. Consequently, the \( D_1 \rightarrow D_2 \) path is swept twice. The longer the critical path is, the more repeatedly the nodes are swept. For this purpose, the main idea is to compute the execution time of critical path once, for all MR functions that can be applied. In this purpose, the MDFG should be swept in the opposition direction of data dependencies and incrementally defining the critical-path computation-time in each node, where the result is stored in the \( V_C_P \) list. To identify the execution time critical path after each MR function, the node having the same level is selected from the \( V_R \), to extract the one having the maximal-execution-time critical path. Elsewhere, for each MR function, the parallelism level \( p \) of retimed sub-paths is computed. Thereafter, \( p \) is added to the previous \( PL_{loop} \) and \( PL_{pro} \) and \( (P_{init\ loop} - p) \) is added to the previous \( PL_{epi} \).
In the case of the Jacobi algorithm MDFG shown in Fig.1(b), the D2 node is the last one for all zero-delay paths, which are saved in the L list. The MDFG is swept in the opposite direction to compute all the path execution times, which are finished by D2, whose values are stored in the VCP list, where $V_{CP} = \{(A1,4), (A2,5), (D1,4), (D2,2)\}$. The following loop in algorithm 2 ensures defining the innermost iteration times after applying respectively the first and second MR functions, where $IT_R = \{(0,0),(1,4),(2,2)\}$. The last loop ensures defining the set $PL_R = \{(0,1,0,0),(1,2,1,1),(2,3,2,2)\}$.

D. Loop bounds analysis

The MDFG allows modelling l nested loops, whatever the loop iteration bounds are. Therefore, all iteration loop bounds are modelled as $LB = (n_1,n_2,...,n_k)$, where $n_i$ is the loop bound of the $i$ loop and $1 \leq k \leq l$. As indicated in section II, the MR leads to reduce the loop bounds with respect to the MR function $r$ in a way that if $r = (r_1,r_2,...,r_k)$, the $n_i$ loop bound is reduced by $|r_i|$. This decline is ensured either the $r_i$ value, which is positive or negative. Thus, the nested loop bounds after the first $r$ MR function is as shown in Equation 6.

$$LB_{(i)} = (n_1 - |r_1|, n_2 - |r_2|,...,n_k - |r_k|)$$

Elsewhere, the $r$ function is enabled to be applied as well as the same parallelism is required. Therefore, the more the MR is applied, the loop bounds decrease with respect to the $r_i$ values. The LB loop bound is formulated in terms of the i MR function rank, as described in Equation 7.

$$LB_{(i)} = (n_1 - i \times |r_1|, n_2 - i \times |r_2|,...,n_k - i \times |r_k|)$$

After applying the MR function $r = (1,-1)$, the outermost and innermost loop bounds are respectively decreased by $|1|$ and $|-1|$. The same growth is implied after each MR function, as shown in Fig.3(b). Accordingly, the final $LB_{(r,i)}$ is formulated as described in Equation 8.

$$LB_{(i)} = (n_1 - i, n_2 - i)$$

E. WCET prediction

The WCET is predicted for all the nested loops in terms parallelism level that corresponds to the $i$ order of the MR function. Since the prologue and epilogue are added in both sides of each retimed loop, the nested loop WCET is incrementally computed from the innermost loop $WCET_{(1,l)}$ to the outermost one $WCET_{(l,1)}$. For the innermost loop, the $WCET_{(1,l)}$ represents the addition of the $WCR_{(1,l)}$ and the $WCST_{(1,l)}$ as described in Equation 3. The first term is equal to the iteration time $IT_i$ multiplied by its loop bound $LB_{(1,l)}$, which corresponds to the $i$th MR function. The second term is the multiplication of the loop bound $LB_{(1,l)}$ and the $C$ constant. If $r_1 \neq 0$ and $1 < l \leq k$, a prologue is added in both sides of the $l$ loop. As indicated in [13], the instructions shifted outside the loop are executed in $IT_{min}$. The greater $|r_1|$ is, the more the prologue execution time increases. Moreover, the prologue and epilogue WCSTs are computed in terms of $PL_{pro}, PL_{epi}$ and $LB_{(l,i)}$. Accordingly, the prologue and epilogue WCET is computed as described in Equation 9.

$$WCET_{pro-epi(l,i)} = (|r_1| \times IT_{min} + (LB_{(l,i)} \times C \times (PL_{pro(l,i)} + PL_{epi(l,i)}))$$

Therefore, each $l$ loop WCET $(l,i)$, where $1 < l \leq k$, is formulated in terms of the $WCET_{pro-epi(l,i)}$ and the $WCET_{(l-1,i)}$. Hence, the whole loop nested loop WCET is defined as shown in Equation 10.

$$WCET_{(l,i)} = \begin{cases} LB_{(l,i)} \times (IT_i + PL_{(l,i)} \times C), & \text{if } l = 1 \\ LB_{(l,i)} \times (WCET_{(l-1,i)} + WCET_{pro-epi(l,i)}), & \text{if } 1 < l \leq k \end{cases}$$

F. Optimization heuristic

Our approach aims to provide the MR function number that allows achieving the execution time constraint. In this context, it determines the MR parameters. Thereafter, the WCET is formulated in terms of MR parameters. Thus, the approach starts by computing the $IT_{min}$, the MR function $r$ and the D and T matrices, as indicated respectively in [12, 13]. Accordingly, the both suggested algorithms are called, where the first one generates the MR function number and the nodes to retime, while the second one provides the $IT_R$ and $PL_R$ sets. Hence, the approach iteratively computes the WCET in order to compare it to the constraint. The first iteration corresponds to the initial MDFG. The last one corresponds to the retimed MDFG with an increased MR function number. The whole steps of our proposed optimization heuristic are described in algorithm 3.

<table>
<thead>
<tr>
<th>ALGORITHM 3. MK for WCET constraint</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Inputs:</strong> a realizable MDFG $G = (V,G)$ and the execution time constraint $T_x$, the iteration bounds of the outermost loop and the innermost loop</td>
</tr>
<tr>
<td><strong>Outputs:</strong> the $i$ number of MR function</td>
</tr>
<tr>
<td>#Compute the execution time and the minimal cycle period #</td>
</tr>
<tr>
<td>Compute $D$ and $T$ matrices [13]</td>
</tr>
<tr>
<td>Define the $M_i$ function [7]</td>
</tr>
<tr>
<td>Identify the $V_R$ list and the MR number $K$ (as described in algorithm 1)</td>
</tr>
<tr>
<td>Identify the $IT_R$ and the $PL_R$ set (as described in algorithm 2)</td>
</tr>
<tr>
<td>$n_0 = 0$</td>
</tr>
<tr>
<td>Repeat</td>
</tr>
<tr>
<td>Compute $WCET_{(l,i)}$ (as indicated in Equation 10)</td>
</tr>
<tr>
<td>for $l$ from 1 to $n$ do</td>
</tr>
<tr>
<td>Compute $WCET_{(l,i)}$ (as indicated in Equation 10)</td>
</tr>
<tr>
<td>end for</td>
</tr>
<tr>
<td>$n_i = n_i + 1$</td>
</tr>
<tr>
<td>While $T_x - T_e &lt; 0$ and $n_i &lt; k$</td>
</tr>
</tbody>
</table>

Therefore, each $l$ loop WCET $(l,i)$, where $1 < l \leq k$, is formulated in terms of the $WCET_{pro-epi(l,i)}$ and the $WCET_{(l-1,i)}$. Hence, the whole loop nested loop WCET is defined as shown in Equation 10.
IV. EXPERIMENTAL RESULTS

In our experiments, the first step consists of verifying the WCET estimation, by comparing its values to the ones provided by the SWEET software tool [5, 19]. The second step, the optimization heuristic is compared to the delayed MR technique by evaluating the core number development of the provided implementations in terms of WCET constraints. Our benchmarks include several 2D nested loops which are the Infinite Impulse Response Filter (IIRF) [13], the Jacobi Algorithm (JA) [25], the Walsh-Fourier Transform (WFT) [12], and the Wave Digital Filter (WDF) [14]. The core numbers are respectively shown in Fig. 6.

All benchmarks are explored to define the MR function and the required function number to achieve the full parallelism, as indicated in algorithm 1. Then, the estimation model, described on section III, is applied to define the WCET in terms of MR function, whose values are cited in table III.

Table III. MR parameters and WCETs of applications

<table>
<thead>
<tr>
<th>Benchmarks</th>
<th>MR function</th>
<th>MR number</th>
<th>Execution time</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Without MR</td>
<td>MR function number</td>
<td></td>
</tr>
<tr>
<td>IIRF</td>
<td>(1, -1)</td>
<td>4</td>
<td>1000 890 720 560 316</td>
</tr>
<tr>
<td>JA</td>
<td>(0, 1)</td>
<td>2</td>
<td>540 400 240 -- --</td>
</tr>
<tr>
<td>WFT</td>
<td>(1, -1)</td>
<td>2</td>
<td>368 242 138 -- --</td>
</tr>
<tr>
<td>WDF</td>
<td>(0, 1)</td>
<td>2</td>
<td>1324 917 504 -- --</td>
</tr>
</tbody>
</table>

The SWEET software tool intercepts the nested loop code and the architectural timing parameter to provide the upper bound WCET. The verification main idea compares the SWEET provided values to those computed by the proposed equation system. This step is done for all benchmarks exposed in Table III, which values are shown in Fig. 5. As a consequence, the proposed WCET estimation offers an efficient prediction with an error rate equal to 8.54% compared to the SWEET tool.

Table IV. WCETs and core numbers in terms of WCET Constraints

<table>
<thead>
<tr>
<th>Benchmarks</th>
<th>WCET constraint</th>
<th>WCET</th>
<th>Core number</th>
</tr>
</thead>
<tbody>
<tr>
<td>IIRF</td>
<td>400</td>
<td>316</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>650</td>
<td>560</td>
<td>15</td>
</tr>
<tr>
<td></td>
<td>800</td>
<td>720</td>
<td>14</td>
</tr>
<tr>
<td></td>
<td>950</td>
<td>890</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>1100</td>
<td>1000</td>
<td>8</td>
</tr>
<tr>
<td>JA</td>
<td>300</td>
<td>240</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>400</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>500</td>
<td>400</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>600</td>
<td>540</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>700</td>
<td>540</td>
<td>1</td>
</tr>
<tr>
<td>WFT</td>
<td>200</td>
<td>138</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>250</td>
<td>242</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>300</td>
<td>242</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>350</td>
<td>242</td>
<td>2</td>
</tr>
<tr>
<td></td>
<td>400</td>
<td>368</td>
<td>2</td>
</tr>
<tr>
<td>WDF</td>
<td>600</td>
<td>504</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>800</td>
<td>504</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>1000</td>
<td>917</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>1200</td>
<td>917</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>1400</td>
<td>1324</td>
<td>2</td>
</tr>
</tbody>
</table>

After analyzing the optimization approach results, we compare its implementations to those provided by the delayed MR retiming in terms of core number, as shown in Fig. 6. As a consequence, the optimization approach presents a benefit in terms of core numbers regarding the delayed MR technique for an average improvement of 27.18%.

Fig. 5. Benchmark WCETs in terms of SWEET tool and WCET equation

Each application is submitted to five execution time constraints, as indicated in the first column of Table IV. Therefore, the optimization approach selects the MR parallelism and estimates its execution time that respects the target constraint, which are listed in the third column of Table IV. To get each constraint, the optimization approach provides implementation with a different parallelism level, which is in relation to the core number. The full parallel implementations are only achieved in the case of minor execution time constraints.

Fig. 6. The core number development in term of delayed MR technique and the optimization heuristic : (a) IIRF, (b) JA, (c) WFT and (d) WDF
V. CONCLUSION AND FUTURE WORKS

In this paper, we have proposed a nested loop WCET static analysis and an optimization heuristic that allows increasing the instruction-level-parallelism in order to achieve a WCET constraint. The main idea consists in providing an optimal parallelism level instead of a full parallel implementation. The experimental results show a noticeable efficiency of the WCET prediction and an important optimization of the heuristic in terms of processing cores.

In our future works, we aim to extend the approach to optimize the cache memory and overhead ratio, which allow enhancing the WCET estimation. Moreover, this work is enabled to be extended to other parallelism techniques such as the loop tiling and loop stripping, hence the whole parallelism solution space, which consequently leads to optimal implementations.

VI. REFERENCES


